Aqueous Extracts of Dried Fruits of *Zanthoxylum armatum* DC., (Rutaceae) Induce Cellular and Nuclear Damage Coupled with Inhibition of Mitotic Activity *in Vivo*

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**ABSTRACT**

Bioactive compounds derived from plant natural compounds have proven to be valuable sources of metabolites which can seldom be obtained from other sources. Plants belonging to the genus *Zanthoxylum* have been valued across various cultures for their curative properties. *Zanthoxylum armatum* DC., belonging to the family Rutaceae is extensively used in traditional practices in North-Eastern India and neighbouring regions including South-East Asia. However, the potential cytogenetic effects of *Zanthoxylum armatum* under *in vivo* conditions, and their causative mechanisms have not yet been studied in detail. The current study was undertaken to evaluate the cytotoxic and genotoxic potential of aqueous extracts of fruits of *Z. armatum* under *in vivo* conditions using the *Allium* test. Physiological and cellular data indicate that the extracts induce clumped chromosomes at metaphase stage of cell division coupled with mitotic arrest. Electron microscopy data reveal membrane damage of cellular organelles, chromatin condensation and chromatin marginalisation in cell of roots incubated in the extracts. The extracts also induce concentration dependent protein precipitation and genomic DNA degradation.

**Keywords:** Clumped Chromosomes; Chromatin Condensation and Marginalisation; TEM; Toxicity; *Zanthoxylum armatum*

1. **Introduction**

Plant derived natural compounds such as quinine, morphine, digitoxin, artemisinin and various others have made important contributions to modern civilisation. Advances in natural products research have helped in developing leads towards the identification and isolation of compounds having significant biological activity [1]. Bioactive secondary metabolites which are produced by plants for their own defense have proven to be valuable sources of new drugs some of which cannot be obtained from other sources [2-4]. Research in plants therefore represents an invaluable source for discovering new substances, considering that most plants can contain a large number of secondary metabolites. From approximately more than 300,000 plant species reported, only a small percentage has been the subject of phytochemical and biological activity studies [5].

Plants belonging to the genus *Zanthoxylum* have been reported to harbour medicinal properties which justifies their use in traditional medicines worldwide [6,7]. Many species have been used in different parts of the world especially in Asia, Africa and America to treat a number of diseases in humans and animals [8-11]. Extracts of *Z. nitidum* exhibit antibacterial activity *in vitro*, as well as antioxidative properties [12,13]. Aqueous extracts of *Z. macrophylla* roots have been reported to show anti-sickling effects in human erythrocytes [14]. Extracts of *Z. piperitum* show scavenging activities towards active oxygen species [15]. Crude extracts of *Z. acanthopodium* and *Z. alainthoides* are reported to possess antioxidative properties [16,17]. *Z. armatum* contains numerous chemical constituents reported to exhibit several biological activities [18,19]. The genus *Zanthoxylum* is also reported to harbour several metabolites that exhibit anti-proliferative activity [20,21]. While the putative curative properties of medicinal plants are widely reported, there are also reports of the *in vitro* toxicity effects of plant extracts utilised in traditional medicines [22-24]. *In vitro* assays indicate that extracts of such plants induce significant damage on cellular and nuclear material. Spiridonov *et al.*, [25] reported acute *in vitro* cytotoxicity of *Potentilla erecta* extracts, which is used for alleviating symptoms of disease in cancer patients. The extracts of aerial parts of *Rumex acetosa* used in traditional Korean and Japanese medicine is also reported to show toxicity...
in vitro [26]. Similar reports are also available for Plantago sp used in traditional Spanish medicine [27]. Plants extracted in herbal remedies are also reported to show cytotoxic potential when used either singly or as concoctions with extracts of other plants [22].

Zanthoxylum armatum DC. (syn. Z. alatum Roxb.) belonging to the family Rutaceae is extensively used in traditional practices of the Khasi tribe in North-Eastern India and in neighbouring regions including South-East Asia. According to reports of ethnobotanical properties of the Zanthoxylum genus, the most commonly used extraction methods use mainly water as a solvent. Herbal formulations prepared from various parts of the plant are reported to be anti-helminthic and hypoglycaemic, and are used as treatment of cholera, tonic for fever, remedy for skin diseases, diseases of the mouth and teeth and others [28,29]. However based on currently available data, the potential cytogenetic effects of Z. armatum under in vivo conditions, and their causative mechanisms have not yet been studied in detail. The current study was undertaken to evaluate the cytotoxic and genotoxic potential of aqueous extracts of Z. armatum in an in vivo experimental system and gain some insight into the mechanism(s) by which these extracts exert their toxic potential.

2. Materials and Methods

2.1. Plant Material and Growth Conditions

The plant specimen used in this study was collected from Shillong, Meghalaya and identified with the help of Dr. P. B. Gurung, Herbaria curator, Department of Botany, North Eastern Hill University, Meghalaya. A voucher specimen was deposited in the herbaria of the Department of Botany, St. Edmund’s College and assigned a voucher number SEC/BOT/EK/101 dated 01/07/2010.

For the Allium test, onion bulbs of approximately 45 cm in diameter were incubated in double distilled water in cylindrical opaque plastic vials (25 mm × 50 mm) for 48 hr with water being changed every 24 hr. After 48 hr the water was substituted with the appropriate solutions and incubated for another 24 hr after which roots of equal length were harvested for analyses. The light conditions for growth were maintained at 14 hr light and 10 hrs dark cycles. Light (200 µmol·m⁻²·s⁻¹) was supplied via cool white fluorescent tubes. Temperature was maintained at 25°C ± 1°C respectively throughout the period of incubation. For each experiment a minimum of 5 bulbs were used.

2.2. Preparation of Aqueous Extract

Fruits of locally growing Zanthoxylum armatum DC., were collected, washed and air dried in the shade until constant weight was obtained. The dried fruits were then ground to a coarse powder in a grinder at short bursts of 30 sec each. 5 gm of powdered material was soaked in 100 mL double distilled water and incubated in the dark for 48 hr at 25°C ± 1°C with frequent agitation. After 48 hr the extract was filtered through Whatmann filter paper No 1 and immediately used. Drying of the aqueous extract in a rotary vacuum evaporator at 40°C resulted in sticky brownish slurry, which could not be used further. Therefore, the fresh aqueous extracts, hereon referred to as Aqueous Extract of Dried Fruits (AEDF), were quantified as total phenolic content [30] and used directly for further experiments.

2.3. Determination of Total Phenolic Content

Total phenolic content of AEDF was estimated according to Makkar et al., [30] with slight modifications. Fifty micro litre (50 µl) of AEDF extract for each sample was taken in a test tube and the volume was made to 1.0 ml with double distilled water. Then, 0.5 ml of 1 N Folin Ciocalteu reagent was added and mixed thoroughly after which 2.5 ml of 0.7 M Na₂CO₃·H₂O solution was added. The resulting solution was mixed thoroughly and incubated for 40 minutes at 25°C ± 1°C. Absorbance was measured spectrophotometrically at 725 nm and total phenolic content was estimated as tannic acid equivalent and expressed as mg/ml. Tannic acid was used for generating the standard curve.

2.4. Root Growth Measurement and Mitotic Squash Preparations

Root lengths were measured at 0 hr, 24 hr, 48 hr and 72 hr after incubation. The initial incubation in double distilled water was considered as 0 (zero) hr for all experiments. Five of the longest roots per bulb were considered. All measurements are represented as the mean ± S.D. of 5 roots per bulb with a minimum of 5 bulbs per experiment.

Root squash preparations were prepared for analyses of mitotic cells. At 72 hr of incubation, 5 - 10 of the longest roots from each bulb were harvested at a distance of 5 mm from the root tip. The roots were immediately boiled in 2% (w/v) aceto-orcein. From each root a segment of 1 mm from the root tip was dissected out. This was used for squash preparation for analyses of mitotic cells. For each analysis a minimum of 1000 cells were examined per root tip with a minimum of 5 roots per bulb. Images of cells were taken with a microscope (Weswox TRHL-66A) fitted with a digital camera (Sony, Japan).

2.5. Transmission Electron Microscope Studies

After incubation for 24 hr in AEDF, the roots were har-
vested for TEM studies. Root tips of approximately 3 mm in length, measured from the tip of roots incubated in AEDF and the corresponding regions of the control roots were fixed in Karnovsky’s fixative for 1.5 hr. The fixed roots were then washed thrice by immersion in 0.1 M cacodylate buffer for 15 min. The fixed roots were further processed and examined using 100 CXII JEOL transmission electron microscope at the Sophisticated Analytical Instrumentation Facility, North-Eastern Hill University, Shillong as described earlier by Choudhury and Sharan [31]. A minimum of 3 roots each from 5 different bulbs were used. Bulbs grown only in double distilled water were taken as negative controls.

2.6. Protein Precipitation Studies

Bovine serum albumin (Fraction V) diluted in double distilled water to a final concentration of 1 mg/mL was taken in different tubes. Increasing amounts of AEDF quantified as total phenolic content were added into each tube, mixed thoroughly and centrifuged at 2000 g for 10 minutes at room temperature. From each tube equal volumes of solution were taken for total protein quantification and the remaining solutions were incubated for 24 hr at 25°C ± 1°C in the dark. After incubation, the tubes were centrifuged at 2000 g for 10 minutes at room temperature and from each tube equal volumes of solution were taken for total protein quantification. Total protein concentration was determined by the Bradford assay [32].

2.7. Genotoxicity Studies

For in vivo studies, total genomic DNA was extracted from roots of Allium bulbs incubated in increasing concentrations of AEDF for 24 hrs as described earlier. Genomic DNA extraction was modified from Križman et al., [33]. Briefly, fresh root tissue was homogenised with CTAB buffer (20 mM EDTA, 100 mM Tris-HCl [pH 8.0], 1.4 M NaCl, 2% [w/v] N-Cetyl-N,N,N, Trimethylammonium bromide (CTAB) and 1% [w/v] Polyvinylpyrrolidone) in a ratio of 0.2 gm:1.5 ml (tissue: buffer) alongwith β-Mercaptoethanol (50 μL/ml buffer) and 0.5% (w/v) activated charcoal and incubated at 55°C for 30 min with constant agitation followed by centrifugation at 12,500 g for 10 min at room temperature. The resulting supernatant was mixed with equal volumes of chloroform:isoamyl alcohol (24:1) solution and centrifuged at 12,500 g for 10 min at room temperature. The supernatant was treated with 5 μl RNaseA (1 mg/ml) for 30 min and the DNA was precipitated with cold isopropanol, washed twice with 70% cold ethanol and dissolved in 100 μl of TE buffer. Total genomic DNA was quantified by measuring the absorbance at A260 and electrrophoresed in 1% (w/v) agarose.

For studying the effects of AEDF on genomic DNA, total genomic DNA from roots of Allium bulbs incubated only in distilled water was extracted as described above. 5 μg of DNA was incubated with increasing concentrations of AEDF for 12 hrs and electrophoresed in 1% (w/v) agarose.

3. Results

3.1. Analysis of Mitotic Activity and Changes in Phase Index of Allium Root Cells under Incubation with AEDF

Incubation of 2-day-old onion roots in AEDF for 24 hr inhibited root growth by approximately 83% in comparison with the negative controls (Figure 1(b)). The inhibition in root growth was comparable to that induced by the Fenton’s reagent [34] which is a commonly used inducer of oxidative stress. A dose response analysis of the effect of AEDF on root growth indicated that half maximal effective concentration (EC50) for inhibition of root growth was observed at total phenolics concentration of 0.4 mg/ml (Figure 1(c)). Analysis of mitotic activity indicated that incubation of roots in AEDF caused changes in the percentage of cells in either metaphase phases’ in comparison to the roots in the control. While both the Fenton’s reagent and AEDF inhibited root growth along with alterations in the mitotic indices, the characteristic effect caused by AEDF, was an extensive increase in the metaphase index during the period of incubation (Figure 1(d)). Furthermore, incubation in AEDF induced clumping of chromosomes in all metaphase cells examined while the cells at interphase showed extensive vacuolisation of the cell nuclei (Figures 1(e)-(j)). The meristematic cells containing clumped chromosomes, accounted for 5% of the total cells examined which corresponded to the percentage of cells in either metaphase or anaphase stage of mitotic division in the negative controls.

3.2. Transmission Electron Microscope Studies

Roots of Allium bulbs incubated in AEDF for 24 hr showed inhibition of growth which was dependent on the concentration of AEDF used. In most of the roots, the inhibition in root growth was accompanied by swelling of the region near the root tips (Figures 2(a)-(g)). Transmission electron microscope studies of the roots incubated in AEDF for 24 hr showed extensive alterations in the ultra-structure of the cellular organelles investigated compared to the negative controls. The cell wall and cell membrane of the controls was intact and continuous. In contrast, in the experimental roots, while the integrity of
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Figure 1. (a) Voucher specimen of *Zanthoxylum armatum* DC. Inset shows dried fruit; (b) Root growth in *Allium* bulbs after 24 hr, 48 hr and 72 hr of incubation; (c) Dose response analysis of root growth in varying concentrations of AEDF; (d) Mitotic phase indices of *Allium* roots after 24 hr incubation in the respective solutions; (e)-(j): Root tip cells of *Allium* after incubation in dd H2O (e), (f), Fenton’s reagent (g), (h) and AEDF (1 mg/ml) (i), (j). *Allium* bulbs were incubated in dd H2O for 48 hr and then transferred into the respective solutions for another 24 hr. Images are representative of cells observed after 24 hr incubation in the respective solutions. Arrows indicate normal mitotic stages with visibly distinct chromosomes (e), (f), cells with cellular and nuclear damage (g), (h) cells with clumped chromosomes and nuclear vacuolisation (i), (j). In (b) and (c), *denotes point of incubation in AEDF as described in text. Bars = 50 µm.

Figure 2. (a) Morphology of root tips after incubation in different concentrations AEDF (as indicated) for 24 hr. Concentrations were determined as total phenolic content (mg/ml); (b)-(g) Representative images of roots showing distinct swelling under increasing concentrations of AEDF. Concentrations used were same as that of panel (a); (h)-(p) TEM micrographs of roots incubated in AEDF (1mg/ml) for 24 hr; (h)-(j) cell wall and cell membrane showing disrupted continuity in the treated roots (i), (j); (k)-(m) mitochondria showing cristolysis and with ruptured or destroyed outer membrane in the treated roots (i), (m); (n)-(p) nuclei with condensed or marginalized chromatin in the treated roots (o), (p); (b), (k), (n) = control roots incubated in ddH2O. CW-cell wall, CM-cell membrane, IS-intercellular space, MT-mitochondria, MC-mitochondrial membrane; RER-rough endoplasmic reticulum, NM-nuclear membrane, NC-nucleoli, CR-chromatin. Bars = 500 µm (b)-(g), 1 µm (h)-(j), 0.5 µm (k)-(m), 2 µm (n)-(p).

the cell wall appeared to be intact, the cell membrane was ruptured and discontinuous at several places (Figures 2(h)-(j)). The cell membrane also appeared to be detached from the cell wall. The root cells of the controls had a large number of mitochondria of varying shapes and sizes with an outer membrane and very well defined cristae (Figure 2(k)). The mitochondria of the cells in roots incubated in AEDF, however, exhibited extensive cristolysis and disarrangement of the cristae (Figures 2(l) and (m)). Furthermore, the integrity of outer membrane of the mitochondria in the root cells exposed to AEDF was also disrupted, while in some cases the entire outer membrane was destroyed (Figure 2(m)) whilst maintaining the structural identity of the cristae. In the control roots, the nuclei of the cells at interphase were spherical...
in shape with an intact double-layered nuclear envelope and well-defined nucleolus with distinctly visible chromatin (Figure 2(n)). As observed with the other cellular organelles, the integrity of the nuclear envelope in the cells of the roots incubated in AEDF was also disrupted. The disruption of the nuclear envelope was accompanied with marked chromatin condensation. In most of the cells, the electron-dense nuclear material was aggregated peripherally under the nuclear membrane (chromatin marginalisation) while in others a uniformly dense nuclei was prevalent (Figures 2(o) and (p)).

3.3. Effect of AEDF on Protein and DNA

24 hr of incubation of protein solution (BSA) with increasing amounts of AEDF resulted in a gradual decrease in the absorbance of the incubated solutions at 595 nm (Figure 3(a)). Normalisation of the absorbance obtained for the different solutions with the absorbance at the same wavelength prior to incubation, showed protein precipitation to be proportional to the concentration of AEDF used (Figure 3(b)).

Incubation of roots of Allium bulbs in increasing concentrations of AEDF for 24 hr did not show any visible damage to the integrity of genomic DNA (Figure 3(c)). Interestingly incubation of fresh genomic DNA with AEDF for 12 hr, showed increased degradation of DNA with increasing AEDF concentration (Figure 3(d)).

4. Discussion and Conclusions

The current study demonstrates that the aqueous extracts of dried fruits (AEDF) of Z. armatum DC., can induce nuclear and cellular damage coupled with mitotic arrest in Allium roots. AEDF induced clumping of chromosomes in the meristematic cells thereby possibly inhibiting further mitotic activity. Such a condition would inhibit the progression of cell division beyond metaphase, thereby leading to subsequent mitotic arrest [35]. In murine cell lines, drugs that induce cytotoxic effects concurrently induce clumped metaphase chromosomes, mitotic arrest at metaphase and subsequent cell death [35]. In cell lines, it has been suggested that clumped chromosomes during cell division may arise due to inhibition of telomerase resulting in end-to-end fusion of the chromosomes [36]. Various plant extracts used in traditional medicinal practices have also been reported to inhibit telomerase activity [37-39]. It is interesting to note that in plants, germinating seedlings and root tips are sites of maximum telomerase activity [40]. It could therefore be possible that AEDF alters or inhibits telomerase activity in the dividing cells resulting in clumped chromosomes at metaphase. While it is still premature at this stage to conclusively ascertain how AEDF induces the observed clumped chromosomes at metaphase, the results clearly suggest that chromosome clumping is one of the mechanisms by which AEDF induces genotoxicity in vivo.

Assessing cell membrane integrity is one of the most common ways to measure cell viability and cytotoxic effects. Compounds that exhibit cytotoxic effects often compromise cell membrane integrity. The results from the current study conclusively show that AEDF can cause loss of structural membrane integrity in cellular organelles such as mitochondria, nuclei and cell membranes, all of which are critical for proper functioning of the cell. Lipid peroxidation is known to trigger the loss of membrane integrity, structural damage to DNA, and cell death. Since AEDF induced disruption of cell and organelle membrane coupled with chromatin condensation and marginalisation, it was of interest to investigate whether the observed cellular damages could be attributed to lipid peroxidation effected by AEDF. The formation of substances, which react with thiobarbituric acid (TBARS), is characteristic of the terminal stage of lipid peroxidation and indicates the breakdown of peroxidised lipids. Analysis of roots of Allium bulbs incubated in varying concentrations of AEDF exhibited a concentration dependent inhibition of a yellow coloured complex, previously reported to be an aldehyde-TBA complex [41,42] (result not shown). This is in agreement with previously published reports of the inhibition of lipid peroxidation by Zanthoxylum extracts [43,44] suggesting that the cel-
lular damages induced by AEDF possibly does not involve induction of lipid peroxidation and might occur via an alternative mechanism(s).

Chromatin condensation and marginalisation are nuclear events that have been associated with the disruption of cell division eventually resulting in cell death [45,46]. Chromatin condensation and nuclear DNA fragmentation occur in plant cells during senescence [47] and/or in the presence of cytotoxic agents [48] and prolonged exposure to higher concentrations of cytotoxic agents can induce loss of membrane integrity. Electron microscopy analyses indicate that AEDF drastically alters the morphology of the cellular organelles investigated. TEM results reveal ultrastructural morphological characteristics such as 1) electron-dense nuclei (marginalisation in the early phase); 2) disorganized cytoplasmic organelles; and 3) disrupted organelle membrane which indicate that the cytotoxic potential of AEDF could be attributed to its ability to possibly induce necrosis since the ultrastructural alterations induced by AEDF bear striking resemblances to typical necrotic hallmarks.

The results presented in this study indicates that AEDF does not seem to induce any visible damage to the integrity of nuclear DNA under in vivo conditions which is in contrast to that reported for apoptotic cells or cells undergoing programmed cell death, but can induce degradation of isolated genomic DNA. The reason for such variation could be that under in vivo conditions, the proportion of DNA degraded by AEDF is much lesser compared to the intact genomic DNA, thereby masking the genotoxic potential of AEDF. This could be partially due to the reason that not all the cells are as easily accessible to AEDF as the meristematic cells present at the root tips. Since the meristematic cells constitute only a small percentage of the total tissue from which the genomic DNA was isolated, therefore, the integrity of the nuclear DNA seemed unaffected by AEDF. However, under in vitro conditions when all the nuclear DNA present was accessible to AEDF, it induced extensive DNA degradation. This implies that AEDF does have the potential to induce acute genotoxic effects. In conclusion, the current study clearly demonstrates that the aqueous extracts of dried fruits (AEDF) of Zanthoxylum armatum DC., which is used extensively in traditional herbal remedies exert its toxic potential in vivo by inducing membrane damage of cellular organelles, chromatin condensation, chromatin marginalisation, chromosome clumping and nuclear DNA damage resulting in subsequent mitotic arrest.

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REFERENCES


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