Alteration in the Doxorubicin-Induced DNA Damage in Cultured V79 Cells by Aegle marmelos (L.) Correa (Bael): A Comet Assay Study

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Abstract

The clinical applicability of wide spectrum chemotherapeutic drug, doxorubicin is limited due to induction of severe cardiomyopathy resulting from DNA damage. The present study was aimed to evaluate the protective effect of bael (Aegle marmelos) extract (AME) on the doxorubicin-induced molecular DNA damage in V79 cells. V79 (Chinese hamster lung fibroblasts) cells were treated with 0 or 25 µg/ml AME before exposure to 0, 1, 2.5, 5, 10, 25 or 50 µg/ml doxorubicin DOX. The DNA damage was studied at different post-doxorubicin treatment times using single cell gel electrophoresis. Doxorubicin caused a maximum DNA damage at 1 h post-DOX treatment indicated by a highest Olive Tail Movement (OTM) and tail DNA, whereas treatment of V79 cells with 25 µg/ml AME enhanced DNA repair at all assessment times with a maximum repair up to 8 h which did not alter thereafter. In another experiment DOX caused a concentration dependent increase in the DNA damage and treatment of V79 cells with 25 µg/ml AME significantly inhibited DOX-induced DNA damage at all post-DOX treatment times. The rate of DNA repair was higher in AME pre-treated cells than DOX-treatment alone. Assessment of cell survival showed a concentration dependent decline in the clonogenicity after DOX-treatment, whereas AME pre-treatment arrested the DOX-induced reduction in the cell survival. The DNA damage and clonogenicity of cells showed a close but inverse relationship, i.e., with increasing DNA damage there was a corresponding reduction in the cell survival. This relationship between cell survival and DNA damage was linear quadratic. Our study demonstrates that AME pretreatment reduced the DOX-induced DNA damage and hastened the DNA repair in V79 cells, thus demonstrating the chemoprotective potential of AME.

Keywords: V79 Cells, Aegle marmelos, Doxorubicin, DNA damage, Comet assay; Cell survival

Introduction

Doxorubicin (adriamycin), an antibiotic having a wide spectrum of anti-neoplastic activity was isolated from the cultures of mutant fungus, Streptomyces peucetius caesius [1]. It has been found to be active against several solid neoplasms [2], Hodgkin’s disease, leukemias, lymphomas [3] and rat tumors [4]. Despite the fact that doxorubicin is active against several tumors it causes severe cardiomyopathy limiting its clinical use [5]. Doxorubicin has been reported to induce micronuclei, chromatid and chromosome aberrations, DNA single and double strand breaks in vitro and in vivo [6-15]. In an attempt to control neoplastic disorders and distant metastases the chemotherapeutic agents are administered systemically, which adversely affects the DNA of other normal cells leading to genotoxicity and subsequently producing second malignancies [16,17]. Since doxorubicin is a wide spectrum chemotherapeutic drug, reduction in its toxicity will be beneficial in the better management of neoplasia and benefit the patients receiving doxorubicin therapy.

If the potential of doxorubicin has to be fully utilized in the treatment of cancer, its toxicity needs to be reduced. The use of antioxidants to reduce doxorubicin-induced toxicity has been advocated. The strategies have been devised to prevent doxorubicin-induced cardiotoxicity without affecting its antitumor activity by combining it with cardioprotective agents and antioxidants [18,19].

Dexazoxane, a bisdioxopiperazine compound is a clinically approved as a prophylaxis for the doxorubicin-induced cardiotoxicity in cancer patients [20]. Other agents like venoruton (a standardized mixture of flavonoids), propolis (bee glue) and many other natural antioxidants were clinically evaluated for their cardioprotective efficacy, but with limited success [21]. Recently, intensive researches on biological function of natural antioxidants to reduce doxorubicin-induced toxicity have been carried out with numerous plant materials worldwide, including those used as foods [15,22,23]. It is also well known that natural antioxidants, including phenolic or thiolic compounds could protect against damages caused by reactive oxidants by various biological mechanisms in living cells [22-25].

The antiquity of use of plants to treat various ailments in humans is as old as the human civilization. The Indian system of medicine, the Ayurveda gives a detailed account of the medicinal properties of numerous plants and their use to treat various disorders in humans. These plant-based systems continue to play an important role in healthcare and it has been estimated by the World Health Organization that approximately 80% of human population globally rely mainly on the traditional medicines for their primary healthcare [26]. Therefore, a need is felt to find alternative drugs to synthetic drugs, which could reduce and repair the deleterious effects of doxorubicin-induced DNA damage. This can be achieved by screening newer molecules or plant products, which may be effective at non-toxic dose levels.

Aegle marmelos, commonly known as bael, is a spinous tree belonging to family Rutaceae. Its edible leaf, root, bark, seeds and fruits are also valued in Ayurvedic medicine in India [27]. In fact as per
Preparation of extract single strand breaks, alkali-labile sites that are expressed as single damage by comet assay in cultured V79 cells exposed to Aegle marmelos was collected locally during the month of April-May insight into the protective nucleoids are incubated with bacterial repair endonucleases that protected the mice against doxorubicin-induced cardiotoxicity [19]. However, the mechanism of action of AME in protecting against the lymphocytes, V79 cells and mice bone marrow cells [14,15,33]. It also of these embedded cells at high pH results in structures resembling different kinds of damage in the DNA and convert lesions into several individual culture dishes [34]. The comet assay has found its utility in testing novel drugs for genotoxicity, human biomonitoring, cancer biology, molecular epidemiology, and fundamental research in DNA damage and repair. The sensitivity and specificity of the assay are greatly enhanced if the comet tail reflects the number of DNA breaks [34]. The comet assay has found its utility in testing novel chemicals for genotoxicity, monitoring environmental contamination with genotoxins, human biomonitoring, cancer biology, molecular epidemiology, and fundamental research in DNA damage and repair. The sensitivity and specificity of the assay are greatly enhanced if the nucleoids are incubated with bacterial repair endonucleases that recognize specific kinds of damage in the DNA and convert lesions into DNA breaks, thus increasing the amount of DNA in the comet tail. DNA repair can be monitored by incubating cells after treatment with damaging agent and measuring the DNA damage remaining at different intervals. The alkaline comet assay can detect double and single strand breaks, alkali-labile sites that are expressed as single strand breaks and single strand breaks associated with incomplete excision repair [35]. The use of comet assay may be helpful in assessing DNA damage at molecular level and ascertain the chemoprotective effect of bael. Therefore, the present study was undertaken to obtain an insight into the protective effect of bael on the cell survival and DNA damage by comet assay in cultured V79 cells exposed to different concentrations of DOX.

Materials and Method

Preparation of extract

The identification of Aegle marmelos (L.) Correa, family, Rutaceae was carried by Dr. Gopal Krishna Bhat (a well-known taxonomist), Department of Botany, Poorna Prajna College, Udupi, India and the herbarium specimen No. AG 032 is stored with us. The mature leaves of Aegle marmelos was collected locally during the month of April-May of the year, cleaned, shade dried and powdered. One hundred grams of the leaf powder was extracted with 50% ethanol in a Soxhlet apparatus. The extract was freeze-dried and stored at -80°C until further use. An approximate yield of 24% was obtained. Henceforth the bael extract will be called as AME.

Drug and chemicals

Doxorubicin hydrochloride (Adriamycin) was procured from Biochem Pharmaceutical Industries, Mumbai, India. Normal and low melting agarose (Cat No. A-4718), ethylenediamine tertra acetic acid (EDTA), Minimum essential medium (MEM), fetal calf serum, L-glutamine and gentamycin sulfate, trizama base, ethidium bromide and triton X-100 were procured from Sigma Chemical Co. St. Louis, USA. The other routine chemicals were procured from Ranbaxy fine Chemicals, Mumbai, India.

Preparation of drug

Doxorubicin hydrochloride (DOX) and AME were dissolved in MEM immediately before use.

Cell line and culture

V79, Chinese hamster lung fibroblasts, procured from the National Centre for Cell Sciences, Pune, India, have been used throughout the study. The cells were routinely grown in 75-cm² flasks (Falcon, Becton Dickinson, USA) with loosened caps, containing Eagle's minimum essential medium (MEM) supplemented with 10% fetal calf serum, 1% L-glutamine and 50 μg/ml gentamycin sulfate at 37°C in a CO₂ incubator (NuAire, Plymouth, MN, USA) in an atmosphere of humidified 5% CO₂ in 95% air.

Experimental design

A fixed number (5 x 10⁵) of exponentially growing cells inoculated into several individual culture flasks or embedded in agarose were divided into following groups:-

MEM+DOX

The cell cultures were incubated with DOX for 1 h, thereafter the drug containing media was replaced with a fresh drug-free MEM.

AME + DOX

The cultures of this group were treated with 25 μg/ml of AME for 1 h [15] and the AME containing media were replaced with MEM containing various concentrations of DOX for another 1 h.

Cell survival

Clonogenicity of cells was measured using colony-forming assay of Puck and Marcus [36]. The log phase cells were treated with 25 μg/ml AME for 1h before exposure to 0, 1, 2.5, 5, 10, 25 or 50 μg/ml DOX. One hour after the DOX treatment, the drug-containing medium was removed and the cells were washed twice with sterile PBS. The cells from each group of flasks were dislodged by trypsin EDTA treatment. Usually, 200 to 300 cells were inoculated into 25 cm² culture dishes (Nunc, Denmark) containing 5 ml medium in triplicate for each drug concentration for each group. The cells were allowed to grow for 9 days and the resultant colonies were stained with 1% crystal violet in methanol. The clusters containing 50 or more cells were scored as a colony. The plating efficiency of cells was determined and the surviving fraction was calculated and fitted on to non-linear polynomial functions.
DNA damage detection by comet assay

Since DOX acts on DNA, it was decided to investigate the effect of AME on the DOX-induced molecular damage to DNA by employing single cell gel electrophoresis (comet assay) in alkaline condition, where two individual experiments were carried out to study the effect of AME on the DOX-induced damage as described below.

DNA repair

The alteration in the DOX-induced DNA damage repair kinetics by AME was evaluated by single cell gel electrophoresis (comet assay). Exponentially growing V79 cells were incubated with MEM or 25 μg/ml of AME for 1h, washed, and then incubated in the medium with 10 μg/ml of DOX for 1 h at 37°C. To examine DNA repair kinetics at different times, the cells from both groups were washed twice and resuspended in fresh, drug-free medium. Aliquots of suspension from DOX treatment alone or AME+DOX groups were taken immediately at 0, ½, 1, 2, 4, 6, 12, 16 and 24 h post-DOX treatment and kept in an ice bath to stop the DNA repair.

DNA damage response

An another experiment was carried out to study the effect of 25 μg/ml AME on the DNA damage caused by different concentrations of DOX, where V79 cells were pre-treated with or without AME for 1h, washed, and then incubated in medium containing 0, 1, 2.5, 5, 10, 25 or 50 μg/ml DOX for 1 h at 37°C. Thereafter, the cells were washed twice and replaced with drug free medium. The cell cultures were terminated at 1, 4 or 6 h and the cells were collected and kept on ice bath for comet assay.

Alkaline comet assay

The single cell gel electrophoresis or comet assay is usually carried out by embedding cells in agarose, followed by lysis in an alkaline/neutral buffer. The application of electric current pulls the charged DNA out of the nucleus, where relaxed and damaged DNA fragments migrate away from the confines of the nucleus than intact DNA, which looks like the comet [Figure 1].

The images are captured and the extent of DNA damage can be precisely determined [34,37-39]. The comet assay was performed under alkaline conditions essentially according to the procedure of Singh et al. [37] with minor modifications [40]. The slides frosted at one side were covered with 100 μl of 0.6% low melting agarose (Sigma-Aldrich Co., St. Louis, USA; Cat No. A-4718) prepared in Ca- and Mg-free PBS at 37°C and the agarose was allowed to solidify under a cover slip on ice after which the cover slips were removed. One ml aliquots containing 1 × 10⁴ harvested V79 cells in culture medium were centrifuged at 1,500 rpm for 5 min. The pelleted cells were resuspended in 80 μl of 0.6% low melting agarose layered on to the first layer and allowed to solidify under a cover slip on ice. All the steps were conducted under a reduced light to prevent additional DNA damage.

The slides containing V79 cells and embedded in agarose were placed into cold lysis buffer (2.5 M NaCl, 100 mM Na2EDTA, 10 mM and Trizma base, pH 10 with freshly added 1% Triton X-100) for 2 h. This removed cellular proteins and left the DNA as nucleoids, thereafter the cell lysis buffer was drained from the slides. The slides containing cells were placed into a horizontal gel electrophoresis tank containing fresh alkaline (pH 13) electrophoresis buffer (300 mM NaOH, 1 mM Na2EDTA). The level of buffer was kept ~ 0.25 cm above the slides, which were kept in this condition for 20 min to allow unwinding of DNA. The electrophoresis was run for 20 min at 1.25 V cm⁻¹ and 300 mA in cold environment. The alkaline buffer was drained of from the slides, which were subsequently flooded slowly with neutralization buffer (0.4 M Trizma base, pH 7.5) for 5 min each three times. The slides were stained with 50 μl of ethidium bromide (2 mg/ml), covered with a coverslip and analyzed immediately.

The ethidium bromide stained DNA on each slide was visualized as "comets" with a fluorescent head and a tail at 40× magnification using an epifluorescence microscope (Olympus BX51, Olympus Microscopes, Tokyo, Japan) equipped with a 515-535 nm excitation filter, and a 590 nm barrier filter. The comet images were acquired using a CCD camera (CoolSNAP-Pro; Digital Color Camera Kit Ver 4.1, Media Cybergenetics, Silver Spring, Maryland, USA) fitted on the microscope. A total of 100 cells per sample were analysed to give a representative result for the population of cells [39]. The captured comet images were analyzed by Komet software (GraphPad Software, Version 5.5, Kinetic Imaging Ltd, Bromborough, UK). The mean olive tail moment (OTM) was selected as the parameter that best reflects DNA damage (defined as the distance between the profile centers of gravity for DNA in the head and tail). OTM was measured from three independent experiments, each containing quintuplicate measures and presented as Mean ± SEM.

Statistical analyses

The statistical analyses were performed using GraphPad Prism statistical software (GraphPad Software, San Diego, CA, USA). The significance among all groups was determined by one-way ANOVA and Bonferroni’s post-hoc test was applied for multiple comparisons. The experiments were repeated for confirmation of results. The results are the average of five individual experiments. The test of homogeneity was applied to find out variation among each experiment. The data of each experiment did not differ significantly from one another and hence, all the data have been combined and means calculated. A p value of <0.05 was considered statistically significant.

Results

The results of clonogenic survival are expressed as surviving fraction in Figure 2, whereas DNA damage are expressed in Tables 1 and 2 and Figures 3 and 4.
Effect of AME on DOX-induced decline in the cell survival

The alteration in the cytotoxicity of DOX by AME was studied by clonogenic assay. Treatment of V79 cells with different concentrations of DOX resulted in a concentration dependent decline in the cell survival as indicated by the reduction in the surviving fraction, that was lowest, i.e., 0.4 for 50 µg/ml DOX (Figure 2). Treatment of V79 cells with 25 µg/ml AME before exposure to different concentrations of DOX resulted in an elevation in the cell survival when compared with the DOX treatment alone (Figure 2).

One interesting fact was that the chemoprotective effect of AME increased with increasing concentration of DOX, where surviving fraction increased by 0.25 and 0.22 for 25 and 50 µg/ml DOX in AME +DOX group when compared with the concurrent DOX-treatment alone (Figure 4).

Alteration in the doxorubicin-induced DNA damage by AME

Cytotoxicity of doxorubicin is related to its capacity to induce DNA damage. In view of this we investigated whether AME will modulate the DNA strand breaks induced by doxorubicin using single cell gel electrophoresis (comet assay). The comet images are shown in Figure 1. The results are expressed as mean tail DNA and olive tail moment (OTM) in Tables 1 and 2.

Figure 2: Alteration in the survival of V79 cells treated with 25 µg/ml Aegle marmelos extract (AME) before exposure to different concentrations of doxorubicin. Squares: MEM+DOX and circles: AME+DOX.

Figure 3: Effect of 25 µg/ml AME on the olive tail moment in V79 cells exposed to various concentration of DOX at different post-DOX treatment times. Squares: MEM+DOX and circles: AME+DOX. a: 1 h, b: 4 h and c: 6 h.

Figure 4: Correlation between DNA damage and cell survival in V79 cells treated with 25 µg/ml AME before exposure to various concentrations of doxorubicin. Left panel DOX alone (upper 1 h, middle 4 h and lower 6 h). Right panel AME+DOX (upper 1 h, middle 4 h and lower 6 h).
Effect of AME on DNA repair kinetics

Baseline DNA damage did not change significantly with assay time (Table 1). In all cases, the DNA damage of control cells remained almost constant with time, indicating that preparation and subsequent processing of the V79 cells did not introduce significant damage to cellular DNA.

V79 cells exposed to 10 µg/ml DOX caused a significant increase in DNA damage as evident by the increased migration of DNA into the comet tails (Table 1).

The maximum DNA damage was observed at 1 h post-DOX treatment that showed a subsequent decline in the DOX-induced DNA damage with assay time, which was reflected in alleviation in tail DNA and OTM. The decline in OTM indicated repair of DNA damage that was maximum up to 8 h post-DOX treatment and remained almost unchanged thereafter up to 24 h post-DOX treatment. Treatment of V79 cells with 25 µg/ml AME before exposure to 10 µg/ml DOX caused a significant reduction in the DNA strand breaks triggered by doxorubicin at all post-DOX treatment times in AME+DOX group when compared to MEM+DOX treatment group (Table 1). The rate of DNA repair was higher in AME treated group for 1-6 h post-DOX treatment when compared with MEM+DOX treatment group (Table 1). Treatment of V79 cells with 25 µg/ml AME before exposure to different concentrations of DOX caused a dose-dependent reduction in the DNA damage by AME+DOX group when compared with MEM+DOX treatment (Figure 3). The reduction in the DNA damage by AME was statistically significant at all concentrations of DOX, when compared with DOX-treatment alone (Table 2). The amount of DNA damage in AME+DOX group at 1 h post-DOX treatment was almost equal to that of DOX-treatment alone at 6 h post-DOX treatment indicating that DNA repair rates were higher in AME treated group than the DOX treatment alone (Table 2).

Biological response

The biological response of treatments was determined by plotting OTM on the Y-axis, whereas the surviving fraction on the X-axis, respectively (Figure 4). A direct correlation between surviving fraction and DNA damage was apparent at all post-DOX treatment times. The increase in OTM resulted in a corresponding decline in the surviving fraction indicating an inverse correlation between DNA damage and cell survival (Figure 4). This correlation between surviving fraction and DNA damage was linear quadratic for all the post-DOX treatment times in both the DOX treatment alone and AME+DOX groups (Figure 4).

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**Table 1**: Alteration in the DNA damage in V79 cells exposed to 10 µg/ml DOX after treatment with 25 µg/ml of AME at various post-DOX treatment times, AME=Aegle marmelos extract, MEM=Minimum Essential Medium, DOX=Doxorubicin, SEM=Standard Error of the Mean. α=p<0.05; β=p<0.01; φ=p<0.001 and no symbol=non-significant. (When compared with 0 h), a=p<0.05, b=p<0.01, c=p<0.001, d=p<0.0001 and no symbol=non-significant (When compared with MEM+DOX).

| Post-treatment incubation time (h) | Tail DNA (mean ± SEM) | Olive Tail Moment (mean ± SEM) |
|-----------------------------------|------------------------|--------------------------------|
|                                  | MEM+DOX | AME+DOX | MEM+DOX | AME+DOX |
| 0                                 | 0.78 ± 0.055 | 25.32 ± 0.52 | 24.28 ± 0.52 | 0.78 ± 0.055 | 24.26 ± 0.52 | 23.32 ± 0.52 |
| 0.5                               | 0.91 ± 0.056 | 27.62 ± 0.52 | 25.32 ± 0.35a | 0.91 ± 0.056 | 24.32 ± 0.35 | 23.38 ± 0.52 |
| 1                                 | 0.90 ± 0.052 | 30.32 ± 0.43 | 26.24 ± 0.43b | 0.90 ± 0.052 | 25.85 ± 0.43 | 23.76 ± 0.43a |
| 2                                 | 0.90 ± 0.041 | 29.74 ± 0.28 | 24.25 ± 0.36b | 0.90 ± 0.041 | 24.25 ± 0.36 | 22.26 ± 0.28a |
| 4                                 | 0.92 ± 0.035 | 24.62 ± 0.64 | 22.48 ± 0.57a | 0.92 ± 0.035 | 21.74 ± 0.57 | 18.86 ± 0.64ab |
| 6                                 | 0.92 ± 0.035 | 22.36 ± 0.47a | 20.18 ± 0.47a | 0.92 ± 0.035 | 20.68 ± 0.47 | 18.12 ± 0.47ab |
| 8                                 | 0.90 ± 0.035 | 18.78 ± 0.53a | 19.72 ± 0.53β | 0.90 ± 0.035 | 19.58 ± 0.53a | 17.84 ± 0.53βa |
| 12                                | 0.90 ± 0.056 | 18.52 ± 0.55a | 19.36 ± 0.55β | 0.90 ± 0.056 | 18.82 ± 0.55 | 17.52 ± 0.55β |
| 16                                | 0.90 ± 0.043 | 18.04 ± 0.37β | 18.76 ± 0.32β | 0.90 ± 0.043 | 18.67 ± 0.32β | 17.36 ± 0.37β |
| 24                                | 0.84 ± 0.033 | 17.31 ± 0.49β | 16.74 ± 0.49φ | 0.84 ± 0.033 | 17.48 ± 0.49β | 17.31 ± 0.49β |
Discussion

Chemotherapy plays a major role in treating various cancers especially to control advanced stages of malignancies in clinical settings [41]. Most of these chemotherapeutic agents exhibit severe normal toxicity, and cause undesirable side effects. The clinical use of anthracyclines like DOX induces life threatening cardiomyopathy and congestive heart failure, which is a major hindrance for optimum use of DOX [42]. Clinical doses of doxorubicin and other anticancer drugs, which are sufficient to kill tumor cells are often toxic to normal cells/ tissues and lead to undesirable side effects like myelosuppression, neutropenia, cardiomyopathy, nausea, vomiting and induction of secondary tumors [17,18]. Pharmacological agents that protect myeloid and lymphoid systems from the suppressive effects of radiation or chemotherapy may be beneficial in attenuating the side effects induced by a standard therapy regimen/s. This might also allow drug’s dose escalation to effectively control the neoplastic diseases. Therefore, it is essential to screen pharmacological agents that can protect normal cells against DOX-induced cumulative toxicity. Plants might also allow drug’s dose escalation to effectively control the neoplastic diseases. Therefore, it is essential to screen pharmacological agents that can protect normal cells against DOX-induced cumulative toxicity. Plants have been reported to cause DNA strand breaks in a concentration-dependent manner in in vitro [50]. This increased DNA damage by doxorubicin may due to induction of 8-OhdG DNA adducts [23] and also due to induction of topoisomerase II. Doxorubicin differentiates across the cell membrane and intercalates between DNA base pairs and target topoisomerase II, resulting in DNA double strand breaks. Doxorubicin also generate free radicals that induce DNA single strand breaks, alkali labile sites and oxidized DNA bases [46,50-53].

Treatment of V79 cells with 25 µg/ml AME caused a time-dependent decline in the DOX-induced DNA damage as evidenced by decreased tail DNA and Olive tail moment, which is in agreement with earlier reports [7,14,15,47-49]. Similarly, idarubicin has been reported to cause DNA strand breaks in a concentration-dependent manner in in vitro [50]. This increased DNA damage by doxorubicin may due to induction of 8-OhdG DNA adducts [23] and also due to induction of topoisomerase II. Doxorubicin differentiates across the cell membrane and intercalates between DNA base pairs and target topoisomerase II, resulting in DNA double strand breaks. Doxorubicin also generate free radicals that induce DNA single strand breaks, alkali labile sites and oxidized DNA bases [46,50-53].

Table 2: Effect of 25 μg/ml AME on the DNA damage induced by different concentration of DOX at various times in V79 cells, AME=Aegle marmelos, MEM=Methyl Minimal Essential Medium, DOX=Doxorubicin, SEM=Standard Error of the Mean. a=p<0.05; b=p<0.01; c=p<0.001 and no symbol=non-significant. (When compared with 0 h), a=p<0.05, b=p<0.01, c=p<0.001, d=p<0.0001 and no symbol=non-significant (When compared with MEM+DOX)

| DOX (µg/ml) | Tail DNA (mean ± SEM) | Olive Tail Moment (mean ± SEM) |
|------------|-----------------------|-------------------------------|
|            | 1 h                   | 4 h                           | 6 h                           |
|            | MEM+DOX               | AME+DOX                       | MEM+DOX                       | AME+DOX                       |
| 0          | 0.82 ± 0.02           | 0.76 ± 0.04                   | 0.84 ± 0.04                   | 0.81 ± 0.04                   |
| 1          | 12.32 ± 0.36a         | 9.28 ± 0.18a                  | 9.52 ± 0.14b                  | 7.98 ± 0.526a                 |
| 2.5        | 17.62 ± 0.36b         | 14.48 ± 0.38b                 | 13.67 ± 0.32b                 | 11.84 ± 0.48bb                |
| 5          | 21.62 ± 0.42f         | 16.92 ± 0.28f                 | 19.21 ± 0.42f                 | 17.63 ± 0.29f                 |
| 10         | 29.34 ± 0.21f         | 27.59 ± 0.82f                 | 25.38 ± 0.84f                 | 23.58 ± 0.74f                 |
| 25         | 54.86 ± 0.58f         | 49.38 ± 0.63f                 | 49.53 ± 0.38f                 | 41.77 ± 0.46f                 |
| 50         | 86.32 ± 0.41f         | 79.96 ± 0.86f                 | 78.74 ± 0.726c                | 69.42 ± 0.28f                 |

**Table 2: Effect of 25 µg/ml AME on the DNA damage induced by different concentration of DOX at various times in V79 cells, AME=Aegle marmelos, MEM=Methyl Minimal Essential Medium, DOX=Doxorubicin, SEM=Standard Error of the Mean. a=p<0.05; b=p<0.01; c=p<0.001 and no symbol=non-significant. (When compared with 0 h), a=p<0.05, b=p<0.01, c=p<0.001, d=p<0.0001 and no symbol=non-significant (When compared with MEM+DOX)**

DNA double strand breaks in the cellular genome [46]. Therefore, we have investigated the ability of AME to protect against DOX-induced DNA strand breaks and repair in V79 cells exposed to different concentrations of DOX.

Exposure of V79 cells to DOX caused a significant DNA damage as evidenced by increased tail DNA and Olive tail moment, which is in agreement with earlier reports [7,14,15,47-49]. Similarly, idarubicin has been reported to cause DNA strand breaks in a concentration-dependent manner in in vitro [50]. This increased DNA damage by doxorubicin may due to induction of 8-OhdG DNA adducts [23] and also due to induction of topoisomerase II. Doxorubicin differentiates across the cell membrane and intercalates between DNA base pairs and target topoisomerase II, resulting in DNA double strand breaks. Doxorubicin also generate free radicals that induce DNA single strand breaks, alkali labile sites and oxidized DNA bases [46,50-53].

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| DOX (µg/ml) | Tail DNA (mean ± SEM) | Olive Tail Moment (mean ± SEM) |
|------------|-----------------------|-------------------------------|
|            | 1 h                   | 4 h                           | 6 h                           |
|            | MEM+DOX               | AME+DOX                       | MEM+DOX                       | AME+DOX                       |
| 0          | 0.82 ± 0.02           | 0.76 ± 0.04                   | 0.84 ± 0.04                   | 0.81 ± 0.04                   |
| 1          | 12.32 ± 0.36a         | 9.28 ± 0.18a                  | 9.52 ± 0.14b                  | 7.98 ± 0.526a                 |
| 2.5        | 17.62 ± 0.36b         | 14.48 ± 0.38b                 | 13.67 ± 0.32b                 | 11.84 ± 0.48bb                |
| 5          | 21.62 ± 0.42f         | 16.92 ± 0.28f                 | 19.21 ± 0.42f                 | 17.63 ± 0.29f                 |
| 10         | 29.34 ± 0.21f         | 27.59 ± 0.82f                 | 25.38 ± 0.84f                 | 23.58 ± 0.74f                 |
| 25         | 54.86 ± 0.58f         | 49.38 ± 0.63f                 | 49.53 ± 0.38f                 | 41.77 ± 0.46f                 |
| 50         | 86.32 ± 0.41f         | 79.96 ± 0.86f                 | 78.74 ± 0.726c                | 69.42 ± 0.28f                 |

**Table 2: Effect of 25 µg/ml AME on the DNA damage induced by different concentration of DOX at various times in V79 cells, AME=Aegle marmelos, MEM=Methyl Minimal Essential Medium, DOX=Doxorubicin, SEM=Standard Error of the Mean. a=p<0.05; b=p<0.01; c=p<0.001 and no symbol=non-significant. (When compared with 0 h), a=p<0.05, b=p<0.01, c=p<0.001, d=p<0.0001 and no symbol=non-significant (When compared with MEM+DOX)**
restored even after 20 h [47,55]. A dose-dependent increase in DNA damage with increasing concentration of DOX is indicated by a significant rise in the tail DNA and OTM. Berberine a topoisomerase II inhibitor has been found to increase DNA damage in HeLa cells [45]. AME has been reported to reduce DOX-induced DNA damage in V79 cells and mice bone marrow cells earlier [14,15]. Naringin has been reported to reduce the DOX-induced 8-OHdG DNA adducts [23]. Likewise, dexaxzone (ICRF-187), cycloheximide, 3,4-dihydroxybenzoic acid and lovastatin also inhibited DOX-induced DNA damage [55-58]. The reduction in the DOX-induced DNA damage by AME may be due to the presence of tannins including gallic acid and ellagic acid that have been reported to protect DNA breakages [59,60].

A concentration dependent decline in clonogenicity after DOX treatment is in agreement with earlier studies, where a similar effect has been reported after DOX treatment [15,61,62]. The protection against DOX-induced decline in cell survival by AME is in conformation with earlier studies on 3,4-dihydroxybenzoic acid and lovastatin [55,63]. An identical effect was also observed with 10 µM cycloheximide and 10 mM WR-2721 against DOX-induced cytotoxic effect [57,64]. Several other chemicals like thiol N-acetylcysteine (NAC) and hydrophilic flavonoids such as rutin and luteolin have been reported to reduce DOX-induced toxicity [65,66].

Biological response determination gives an indication of relationship between DNA damage and cell survival and we have observed an inverse correlation between the DNA damage and cell survival in both DOX and AME+DOX groups. The surviving fraction of cells declined with the increasing DNA damage indicating that the initial lesions in DNA got stabilized and became cell lethal. An identical effect has been observed earlier [15,55]. This relationship between DNA damage and surviving fraction fitted on a linear quadratic model at all assay times in both the groups. A linear quadratic relationship has been reported for micronuclei induction and cell survival earlier [11,67].

Several putative mechanisms may be involved in DOX induced DNA damage and cytotoxicity including intercalation into DNA [51,52], stabilization of topoisomerase II-DNA complex [46,53], free radical mediated toxicity caused by redox cycling of the semiquinone radical or formation of reactive oxygen species by the DOX iron complex [68-70]. DOX has been reported to be metabolically activated to a free radical state and interacts with molecular oxygen to generate superoxide radicals [70,71]. The superoxide radicals can react with hydrogen peroxide to form highly reactive hydroxyl radicals via the iron catalyzed Haber-Weiss reaction. Secondarily derived hydroxyl radicals can cause protein and DNA damage and initiate lipid peroxidation [73]. The inhibition of topoisomerase II by DOX may have caused stabilization of transient DNA strand breaks leading to increased DNA damage and subsequently the cell death in the present study. DOX has been reported to trigger the transcriptional activation of NF-κB as a response to DNA damage and elevate COX-II expression [74,75]. Our earlier study has reported increased PARP activity by DOX in vivo that was directly correlated with the increased formation of 8-OHdG DNA adducts that may have subsequently converted into strand breaks [23].

The exact mechanism of chemoprotective effect of AME is not known, several putative mechanisms may have contributed in various ways to reduce DOX-induced DNA damage and cytotoxicity in V79 cells. AME may have scavenged DOX-induced free radicals and/or inhibited iron-DOX complex formation. This contention is supported by our earlier reports, where AME has been reported to scavenge free radicals in vitro and in vivo [33,76]. Attrition in DOX-induced DNA damage may be attributed to the restoration of topoisomerase II activity by AME. The attenuation of DOX-induced lipid peroxidation by AME may have reduced the DNA damage and increased the survival in the AME+DOX group. AME has been reported to increase glutathione and reduce lipid peroxidation earlier [31,32]. The suppression of transcriptional activation of NF-κB, COX-II and PARP genes by AME may have also contributed in various ways to reduce DOX-induced DNA damage. AME may have also upregulated the transcription of Nr2 gene leading to reduced DNA damage and cell survival since DOX has been reported to alleviate its expression [77]. AME may have also suppressed the DOX-induced 8-OHdG DNA adducts thereby reducing the DNA damage and increasing the cell survival.

Conclusion

Our study indicates that AME inhibited DOX-induced DNA damage and subsequently the cytotoxicity. The reduction in DNA damage and cytotoxicity by AME may be due to free radical scavenging, iron chelation, restoration of topoisomerase II activity, inhibition of ROS generation and increased antioxidants status. AME may have suppressed the transcription of NF-κB, COX-II and PARP genes accompanied by the upregulation of Nr2 gene that may have increased the antioxidant status of DOX-treated cells and protected against cytotoxicity and DOX-induced DNA damage.

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