α-Mangostin from *Cratoxylum arborescens* demonstrates apoptogenesis in MCF-7 with regulation of NF-κB and Hsp70 protein modulation in vitro, and tumor reduction in vivo

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Abstract: *Cratoxylum arborescens* is an equatorial plant belonging to the family Guttiferae. In the current study, α-Mangostin (AM) was isolated and its cell death mechanism was studied. HCS was undertaken to detect the nuclear condensation, mitochondrial membrane potential, cell permeability, and the release of cytochrome c. An investigation for reactive oxygen species formation was conducted using fluorescent analysis. To determine the mechanism of cell death, human apoptosis proteome profiler assay was conducted. In addition, using immunofluorescence and immunoblotting, the levels of Bcl-2-associated X protein (Bax) and B-cell lymphoma (Bcl)-2 proteins were also tested. Caspaces such as 3/7, 8, and 9 were assessed during treatment. Using HCS and Western blot, the contribution of nuclear factor kappa-B (NF-κB) was investigated. AM had showed a selective cytotoxicity toward the cancer cells with no toxicity toward the normal cells even at 30 μg/mL, thereby indicating that AM has the attributes to induce cell death in tumor cells. The treatment of MCF-7 cells with AM prompted apoptosis with cell death-transducing signals. This regulated the mitochondrial membrane potential by down-regulation of Bcl-2 and up-regulation of Bax, thereby causing the release of cytochrome c from the mitochondria into the cytosol. The liberation of cytochrome c activated caspase-9, which, in turn, activated the downstream executioner caspase-3/7 with the cleaved poly (ADP-ribose) polymerase protein, thereby leading to apoptotic alterations. Increase of caspase 8 had showed the involvement of an extrinsic pathway. This type of apoptosis was suggested to occur through both extrinsic and intrinsic pathways and prevention of translocation of NF-κB from the cytoplasm to the nucleus. Our results revealed AM prompt apoptosis of MCF-7 cells through NF-κB, Bax/Bcl-2 and heat shock protein 70 modulation with the contribution of casapses. Moreover, ingestion of AM at (30 and 60 mg/kg) significantly reduced tumor size in an animal model of breast cancer. Our results suggest that AM is a potentially useful agent for the treatment of breast cancer.

Keywords: α-Mangostin, apoptosis, mitochondria, protein array, caspase 3/7, NF-κB

Introduction
Breast cancer is one of the most deadly cancers affecting women worldwide because of its high rate of incidence and mortality. According to the American Cancer Society, breast cancer is emphasized to account for 26% of all new cancer cases, which is the highest in percentage among all the cancers in American women.¹ In Malaysia, the National Cancer Registry reported that one in 20 Malaysian women are at a risk of acquiring breast cancer in their lifetime.² This incidence rate is still considered low
In this study, we evaluated the apoptotic cell death mechanism prompted by AM on breast cancer using MCF-7 cells as an in vitro model. In addition to that, we investigated the antitumor activity of AM in the animal model.

Materials and methods

Cell culture

Normal breast cells, MCF-10 A, and MCF-7 cells, were acquired from the American Type Culture Collection (Manassas, VA, USA) and then kept at 37°C in an incubator with 5% CO₂ saturation. They were grown in Roswell Park Memorial Institute medium (RPMI)-1640 (PAA Laboratories GmbH, Coelbe, Germany) together with 10% Fetal Bovine Serum (FBS).

Anti-proliferative effect of AM on MCF-7 cells

The inhibitory effect of AM was determined by MTT assay, in which 1×10⁵ of MCF-7 cells/µL were seeded in triplicate in 96-well plates and kept for 24 hours at 37°C with 5% CO₂ saturation. After 24 hours’ incubation, a serial dilution for different concentrations of AM was prepared and transferred to the MCF-7 cells and incubated for 24 hours in 37°C and 5% CO₂. 20 µL of 3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide (MTT solution, 5 mg/mL) was added to the treated cells in a dark place, covered with foil, and incubated for 4 hours. All media were discharged and a total of 100 µL volume of dimethyl sulfoxide was poured into each well until the purple formazan crystals dissolved. The plate was measured using a microplate reader (Tecan Group Ltd., Männedorf, Switzerland) at absorbance 570 nm. The experiment was conducted in triplicate to evaluate half-maximal inhibitory concentration for AM against the MCF-7 cell line.

Acridine orange propidium iodide double staining using fluorescent microscopy

MCF-7 cells were quantified using acridine orange (AO) and propidium iodide (PI) double-staining in conformity with the standard procedures, and inspected under a fluorescence microscope (Leica [Leica Microsystems, Wetzlar, Germany] with Q-Flo Software [Leica Microsystems]). Concisely, the treatment was conducted in a 25 mL culture flask, in which 1×10⁵ MCF-7 cells/mL were seeded and treated with different concentrations of AM (5, 10, and 20 mg/mL) for 24 hours. After 24 hours, the cells were centrifuged at 1,800 rpm for 5 minutes to discharge the old medium; then a cool PBS was used to wash...
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the cells two times before staining with AO/PI. Equal amounts, 10 µL (10 µg/mL), of fresh AO and PI were prepared and covered from light to stain the cell onto a glass slide, which was covered with a cover slip. The slides were viewed under a fluorescence microscope 30 minutes prior to the fluorescence fading.

**Assay of the apoptotic rate by Annexin V (AV)-fluorescein isothiocyanate (FITC) staining**

MCF-7 at concentration (1×10⁵ cells/mL) was exposed to 5, 10, and 20 µg/mL concentrations of AM, and the AV assay
was executed with the use of the BD Pharmingen™ Annexin V-FITC Apoptosis Detection Kit (APO Alert Annexin V, Clontech, Mountain View, CA, USA). The treated cells were centrifuged for 5 minutes at 1,800 rpm to eliminate the media. Subsequently, cells were washed with 1x binding buffer provided by the manufacturer. Then the cells were re-suspended in 200 µL of binding buffer; 5 µL of AV, and 10 µL of PI (Sigma-Aldrich Co, St Louis, MO, USA) were added before incubating at 37°C in the dark for 15 minutes. The binding buffer raised the reaction volume to 500 µL for the flow cytometric analysis. Flow cytometric analysis was conducted using FACS Canto II, (BD Biosciences, San Jose, CA, USA) cytometry b. The dimethyl sulfoxide-treated (0.1%, v/v) cells were employed as the control.

**Nuclear morphology, membrane permeability, mitochondrial membrane potential Δψm (MMP), and cytochrome c release analysis**

A Cellomics Multiparameter Cytotoxicity 3 Kit (Thermo Fisher Scientific, Waltham, MA, USA) was employed as previously described at length. The kit facilitates concurrent measurements of six independent parameters at the same time for the same cell population. Chromatin condensation, morphological change, detection of the MMP, release of cytochrome c, and cell permeability were used to test the loss of the MCF-7 cells. MCF-7 (1×10^6 cells/mL) were used to seed in 96 wells (Genetix Biotech Asia Pvt. Ltd., New Delhi, India) and treated with AM (5, 10, and 20 mg/mL). After 24 hours, the plates were processed according to the manufacturer’s instructions, and analyzed using the ArrayScan HCS system (Thermo Fisher Scientific). The images and the intensity data were reading the texture of the fluorescence inside each of the cells, together with the typical fluorescence of the cell population inside the well. The experiment was done in triplicate.

**Immunofluorescence analysis of Bax/Bcl-2**

A total of 5×10^3 MCF-7 cells were seeded in a 96 well plate (genetix), followed by 5, 10, and 20 µg/mL of AM treatment. Then, the cells were fixed for 15 minutes at 37°C by 4% paraformaldehyde after rinsing two times with PBS. PBS was used to rinse the cells three times, the cells were treated using a blocking buffer for 60 minutes incubation in 0.03% Triton X-100/PBS (Sigma-Aldrich, St Louis, MO, USA) and normal serum before the cells were rinsed once more with PBS. The diluted primary antibody solution containing 1× PBS/1% BSA/0.3% Triton X-100, was added after the aspirate blocking buffer. The cells underwent incubation overnight at 4°C. B-cell lymphoma (Bcl)-2 and Bcl-2-associated X protein (Bax) fluorochrome-conjugated secondary antibody diluted (Santa Cruz Biotechnology Inc., Dallas, TX, USA) in antibody dilution or in PBS only was added to the cells and incubated for 1 hour. After rinsing three times in PBS, the cells were treated with DAPI and examined using the CellReporter™ cytofluorimeter system (Molecular Devices LLC, Sunnyvale, CA, USA).

**Bioluminescent assay of caspaces 3/7, 8, and 9**

A concentration-dependent study of the activity of caspace-3/7, 8, and 9 was done using luminescence-based assay, Caspace-Glo™ 3/7, Caspace-Glo™ 8, and Caspace-Glo™ 9 Assay (Promega Corporation, Fitchburg, WI, USA). The cells were seeded in 96-well plates in 50 µL of RPMI 1640 together with 10% PBS and incubated for 24 hours. Then, the cells were treated with AM 5, 10, and 20 µg/mL and incubated for 24 hours at 37°C with 5% CO₂. After 24 hours, 100 µL of the assay reagent was added and the plate was incubated for 1 hour at 37°C. Luminescence was calculated using a Tecan Infinite® 200 Pro (Tecan, Männedorf, Switzerland) microplate reader.

**Western blot analysis**

MCF-7 cells were cultured in a 25 mL flask (TPP Techno Plastic Products AG, Trasadingen, Switzerland) and treated with AM 5, 10, and 20 µg/mL. The cells were collected, washed by cool PBS, and mixed with a lysis buffer (50 mM Tris-Hcl pH 8.0, 120 mM NaCl, 0.5% NP-40, 1 mM phenylmethylsulfonyl fluoride) to isolate the total proteins from the cells. A 40 µg measure of the extracted protein was separated using 12% sodium dodecyl sulphate-polyacrylamide gel electrophoresis, then moved to a polyvinylidene difluoride membrane (Bio-Rad Laboratories Inc., Hercules, CA, USA), and blocked with 5% non-fat milk in a TBS-Tween buffer 7 (0.12 M Tris-base, 1.5 M NaCl, 0.1% Tween 20) for 30 minutes at room temperature, and incubated with the appropriate primary antibody overnight at 4°C, then incubated with alkaline phosphatase-conjugated secondary antibody for 30 minutes at room temperature, and then washed in Tris-buffered saline and Tween 20 buffer. The primary antibodies, β-actin (sc-130300), Bax (sc-20067), heat shock protein (Hsp)70 (sc-69705), proliferating cell nuclear antigen (PCNA) (sc-25280), NF-κB/ P65 (sc-398442), cytochrome c (sc-13560), caspase-7 (sc-
81660), caspase-8 (sc-81657), and caspase-9 (sc-56073) were purchased from Santa Cruz Biotechnology Inc., but Bcl2 (ab38629) and poly (ADP-ribose) polymerase (PARP) (ab4830) were purchased from Abcam plc (Cambridge, UK). The membranes were then incubated for 1 hour at room temperature with alkaline phosphatase conjugated goat anti-mouse or goat anti-rabbit secondary antibodies in a ratio of 1:5,000 and then washed twice with TBST for 10 minutes three times on an orbital shaker. Then, the blots were developed using the BCIP/NBT (Santa Cruz Biotechnology Inc.) for sensitive colorimetric detection to quantify the target protein band.

**Determination of the intracellular reactive oxygen species (ROS) level**

ROS was determined with 2',7'-dichlorofluorescin diacetate (DCFH-DA). Concisely, 10 mM DCFH-DA stock solution (in methanol) was diluted 500-fold in Hank's Balanced Salt Solution with no serum or further additives to yield a 20 µM working solution. After exposure to AM for 4 hours the cells in the 96-well black plate were rinsed two times with Hank's Balanced Salt Solution before incubating in a 100 µL working solution of DCFH-DA at 37°C for 30 minutes. Using a fluorescence microplate reader, Tecan Infinite®200 Pro, the result was taken in emission at 485-nm excitation and 520-nm emission.

**DNA content and cell cycle phase distribution**

MCF-7 cells at a concentration of 1×10^5 cells/mL were seeded into a 25 mL flask containing RPMI 1640 augmented with 10% FBS and dosed with AM 5, 10, and 20 µg/mL. After incubation for 24 hours, the cells were collected, centrifuged to remove the old medium, and washed two times using warm PBS, then the pellet was fixed with 90% cool ethanol (kept at −20°C for 1 day before use) and kept at 4°C overnight. After incubation, the cells were spun down for 5 minutes at 1,800 rpm, the ethanol was removed and the pellets were washed with PBS to clear ethanol from the fixed cells. An amount of 600 µL warmed PBS was added and mixed gently. Then, 25 µL RNase A in a concentration of 10 µg/mL and 50 µL of PI at a concentration 1 µg/mL were decanted into the pellet and preserved in a dark place for 30 minutes. The DNA content of the cells was then examined using a FACSCanto II Becton-Dickinson Flow cytometer by studying at least 10,000 cells per sample. The percentage of cells in the G1, S, and G2 phases was investigated using Diva software (BD Biosciences, San Jose, CA, USA).

**Immunofluorescence of NF-κB activity**

HCS was utilized to assess the inhibitory effects of AM on tumor necrosis factor alpha (TNF-α)-induced NF-κB activation, ie, nuclear translocation of NF-κB. The experiments were conducted according to the company’s instructions for the NF-κB activation kit (Thermo Fisher Scientific). An ArrayScan reader was utilized to measure the variance between the strength of the nuclear and cytoplasmic NF-κB-associated fluorescence, and reported as a translocation parameter.

**Human apoptosis proteome profiler array**

To examine the pathways through which AM prompts apoptosis, we determined the apoptosis-related proteins utilizing the Proteome Profiler Array (RayBio® Human Apoptosis Antibody Array Kit, Raybiotech, Norcross, GA, USA), in accordance with the manufacturer’s instructions. In short, the cells were dosed with 20 µg/mL AM. Three hundred µg proteins from each sample were incubated overnight with the human apoptosis array. The data from the apoptosis array were measured by scanning the membrane using a Biospectrum AC ChemiHR 40 (UVP, LLC, Upland, CA, USA) and examination of the file of the array image was conducted using image analysis software in accordance with the instructions of the company.

**Mammary tumor implantation**

Rat LA7 mammary adenocarcinoma cells at around 75%–85% confluence were trypsinized, washed once in Dulbecco’s Modified Eagle’s Medium (supplemented with 5% FBS), spun down, and suspended in the same medium. A total of two million cells in a volume of 100 µL were injected into the mammary fat pads of 20 female Sprague-Dawley rats.

**Experimental design and treatment of implanted animals**

Ten days after implantation of LA7 mammary carcinoma, the animals were randomly separated into six groups (n=5 per group): normal control; mammary tumor control (LA7-induced non-treated); low dose (treated with 30 mg/kg/day of AM dissolved in Tween 20); high dose (treated with 60 mg/kg/day of AM dissolved in Tween 20); tamoxifen (treated subcutaneously with 10 mg/kg/day of tamoxifen dissolved in Tween 20); and AM (normal rats treated with 60 mg/kg/day of AM dissolved in Tween 20). All treatments by AM were selected based on the toxicological report by
Ibrahim et al, and given to the animals orally twice a week for 28 days using a gastric tube. At the end of treatment, the rats were euthanized and the entire tumor was removed, sectioned, and fixed in 10% paraformaldehyde. All experimental procedures performed on animals were done according to regulations set by the Institutional Animal Ethical Committee (FAR/20/04/2013/MYID), Faculty of Medicine, University of Malaya.

**Evaluation of effect of AM on tumor size**

The total tumor volume (mm$^3$) was quantified by multiplication of the slice sections and by summing the segmented areas.

**Statistical analysis**

The results were reported as the mean ± standard deviation for at least three analyses for each sample. The normality and homogeneity of variance assumptions were checked, and statistical analysis was performed according to the SPSS 16.0 package (SPSS Inc., Chicago, IL, USA) and GraphPad prism 5.0 (GraphPad Software, Inc., La Jolla, CA, USA). Analysis of variance was performed using the ANOVA procedure.

**Results**

**AM inhibits the growth of MCF-7 cells**

The cytotoxic effects of AM on the MCF-7 cells were ascertained by means of MTT assay. As shown in Table 1, AM inhibited the growth of MCF-7 cells and significant inhibition was exhibited at 10.26±0.25 μg/mL after 24 hours. Tamoxifen, which was used as standard control, showed an IC$_{50}$ of 2.35 μg/mL. Meanwhile, AM did not exhibit any cytotoxicity toward the normal breast cell line MCF-10A with IC$_{50} > 30$ μg/mL.

**AO-PI double staining cell morphological analysis**

To determine the viable cells, as well as those in early apoptosis, late apoptosis and secondary necrosis, the cells were counted using a fluorescence microscope. A total of 200 cells were counted arbitrarily and differentially, along with the negative control, which was untreated. Early apoptosis was detected using AO within the fragmented DNA with a bright green fluorescence. In addition, the green nuclear structure of the control cells was shown to be intact (Figure 1B). Following treatment with 5 μg/mL AM, moderate apoptosis was observed by blebbing and nuclear chromatin condensation (Figure 1C). In addition, in the later phases of apoptosis, alterations, including a reddish-orange color due to the binding of AO to the denatured DNA, were noted after AM 10 and 20 μg/mL of treatment (Figure 1D and E). The results indicated that the AM produced morphological features that are linked to apoptosis in a concentration-dependent manner. A statistically significant ($P<0.05$) difference was noted in the cell population through the differential recording of treated cells (200-cell population) (Figure 1F).

**Effect of AM on the ratio of apoptotic cells by AV-FITC staining**

Flow cytometric analysis, with AV/PI double staining, was used to confirm the induced apoptotic effect of AM by determining the percentage of apoptotic cells. The AV+/PI$^+$ staining signifies the early apoptotic cells because of the strength of the affinity between AV-FITC and phosphatidylserine that transported from the inner leaflet to the outer surface of the plasma membrane in early apoptosis. In contrast, AV+/PI$^+$ staining signifies the necrotic cells, as PI, which cannot penetrate an undamaged cell membrane, passes through the affected membrane of the dead cells or late apoptotic cells, and binds to the nucleic acid. In addition, viable cells are marked by AV$^-$/PI$^-$ and AV$^-$/PI$^+$ staining, which is symbolic of the late apoptotic cells. The characteristic dot plots of the flow cytometric assessment of apoptosis indicated that according to the evaluation of the untreated cells (control) and treated cells (with 5, 10, and 20 μg/mL AM), the percentages in early apoptosis and late apoptosis increased respectively (Figure 2). Moreover, the AM treatment clearly resulted in slightly fewer viable cells at 5 and 10 μg/mL. In addition, the results indicated that the anti-proliferative effect of AM in respect of MCF-7 cells was instigated by triggering cell apoptosis.

**AM-induced apoptosis in MCF-7 cells**

To verify the occurrence of apoptosis, we inspected the alterations in the nuclear morphology of the MCF-7 cells by determining the nuclear condensation and
α-Mangostin demonstrates apoptosis in MCF-7 cells

Fragmentation hallmark for apoptosis (Figure 3A). Hoechst 33342 staining indicated that some of the cells exhibited nuclear condensation with 20 µg/mL AM treatment. The intensity of the nucleus that directly relates to the apoptotic chromatin changes, such as blebbing, fragmentation, and condensation, are quantified in Figure 3B. A simultaneous increase in the cells’ permeability was also observed (Figure 3C).

**AM-induced MMP disruption and release of cytochrome c**

In the apoptosis, MMP is often disturbed by the forming of permeability transition pores or the insertion of pro-apoptotic proteins like Bax or BH3 interacting domain death agonist (Bid) in the membrane of mitochondria. Hence, we studied the effect of AM on the MMP of MCF-7 cells with the use of a mitochondria-specific voltage-dependent dye. As displayed in Figure 3D, the MMP in the cells treated with AM ($P<0.05$) showed a considerable reduction. In addition to that, a considerable decrease in the intensity of fluorescence (Figure 3D), which reflected the breakdown of MMP, AM also activated a major translocation of cytochrome c from the mitochondria into the cytosol. At 20 µg/mL, AM activated three times the amount of cytochrome c released in MCF-7 cells compared to the control ($P<0.05$) (Figure 3E). Moreover, since the release of cytochrome c from the mitochondria triggers the activation of pro-caspase 9, we measured the immunoblotting of cytosol cytochrome c. In a concentration-dependent manner, AM significantly increased the level of cytosol cytochrome c in MCF-7 cells compared to the control ($P<0.05$) (Figure 3D).
AM regulates the expression of Bcl-2 and Bax protein

The expression of pro-apoptotic proteins, such as Bax, is an early incident that sensitizes cells to undergo apoptosis. Compared with the controls, MCF-7 cells that were treated with AM displayed a considerable escalation in fluorescence intensity (Figure 4A) when stained using a specific Bax antibody. This outcome indicated that the expression of Bax is upregulated in MCF-7 cells after dosing with 20 µg/mL AM. Nevertheless, the anti-apoptotic, Bcl-2 levels were minimal during the period of treatment (Figure 3A). Figure 3B shows that both up and downregulation of Bax and Bcl-2 were statistically significant (P<0.05) in concentration dependent manner. To confirm the immunofluorescence result, the expression of Bax and Bcl-2 in the MCF-7 cells treated with AM were determined using Western blot analysis. AM treatment resulted in an upregulation of Bax and a downregulation of Bcl-2 in concentration dependent manner (Figure 4C), which resulted in an increase in the ratio of Bax to Bcl-2, therefore favoring apoptosis (Figure 4D).
**Effect of AM treatment on caspase-3/7, -8, and 9**

All the caspase enzymes being examined were shown to be induced by the treatment, and were concentration-dependent. A high level of caspase 3/7 was found in the greatest treatment concentration (20 µg/mL) with a significant variance from the control ($P<0.05$) for both types of cell. Although significant levels of caspase-8 and caspase-9 were found in both, the triggering of caspase was not found to be significantly prompted for caspase 3/7 at 5 and 10 µg/mL.

Our results provide further validation of the activation of all three caspaces by AM in MCF-7 cells (Figure 5A). To confirm the immunofluorescence result, the cell extracts were obtained after treatments and processed for Western blot analysis. The changes in protein levels in the MCF-7 cells treated with different concentrations of AM were determined using Western blot analysis. In a concentration dependent manner, AM treatment resulted in partial cleavage of pro-caspase-7, -9, and -8 (Figure 5B), which indicated that AM could increase the cleavage maturation of caspase-7, -9, and -8 (Figure 5C).

**Suppression of the PCNA and PARP protein expressions by AM**

The expression of PCNA and PARP cleavage on MCF-7 cells with or without AM treatment was tested by Western
ROS, up to twice as high as observed in the control, was observed by the oxidation-sensitive fluorescence of the dye DCFH-DA. The escalation of dose-dependence in DCF fluorescence was observed in the treated cells. We inspected the levels of ROS in MCF-7 cells treated with AM. ROS was observed by Western blot analysis. As shown in Figure 8D and E, after treatment with 5, 10, and 20 µg/mL of AM, the level of PCNA protein decreased significantly. Since PARP cleavage results in a cleaved product of 89 kDa, we have observed the reduction in the PARP (116 kDa) and concomitant presence of a cleaved product in a dose-dependent manner (Figure 8D and E).

**AM-induced cell death includes increased ROS formation**

The production of ROS was generally linked to the MMP disturbance and cell apoptosis. To determine the relation, we inspected the levels of ROS in MCF-7 cells treated with AM. ROS was observed by the oxidation-sensitive fluorescent dye DCFH-DA. The escalation of dose-dependence in DCF fluorescence was observed in the treated cells. In addition, a considerable and immediate formation of ROS, up to twice as high as observed in the control, was identified following the dosing of cells with AM at 10 and 20 µg/mL (Figure 5D). This result indicated that this compound triggered the intracellular ROS development of MCF-7 cells.

**Cell cycle analysis**

We conducted this trial to determine the influence of AM on the DNA content of MCF-7 cells by the cell cycle phase distribution (G$_0$/G$_1$, S, G$_2$, and M) after treatment (Figure 6). The results indicated that AM halted the cell cycle progression in the G$_0$/G$_1$ phase ($P<0.05$). The results displayed in Figure 6E show that there is a significant G$_0$/G$_1$ phase arrest in a concentration-dependent manner in the MCF-7 cells, which accounts for 52.44%, 55.28%, and 68.75% of cells following treatment with 5, 10, and 20 µg/mL, respectively, for 24 hours ($P<0.05$). Meanwhile, the cells in both the S and G$_2$/M phases diminished with an increase in the treatment concentration.
**α-Mangostin demonstrates apoptogenesis in MCF-7 cells**

The obstruction to apoptosis and cell proliferation was considered to be closely related to the activation of nuclear factor kappa-B (NF-κB). Hence, the role of AM in the suppression of activated NF-κB induced by the inflammatory cytokine, TNF-α, was examined using Alexa Fluor 488-conjugated anti-NF-κB antibody (Thermo Fisher Scientific, Waltham, MA, USA). Although a high NF-κB fluorescent intensity was observed in the cytoplasm in the control cells (medium alone) (Figure 7A), it was only faint in the nuclei, which suggests an absence of NF-κB activation in the control cells (medium alone). In addition, TNF-α alone considerably intensified the NF-κB fluorescent intensity in the nuclei. AM demonstrated a considerable inhibitory effect on the triggering of NF-κB (Figure 7B). In the cells treated with curcumin, a known inhibitor of NF-κB activation, a significant suppression of TNF-α-induced NF-κB nuclear translocation was observed.

**Figure 6** AM induced G₀/G₁ phase cell cycle arrest in MCF-7 cells.  
Notes: Histograms for cell cycle from analysis of MCF-7 cells treated with AM at 5 (B), 10 (C), and 20 μg/mL (D), where (A) is control. Results are representative of one of three independent experiments. (E) Induction of G₀/G₁ phase arrest in the cell cycle progression of MCF-7 cells by AM. *Indicates a significant difference (P<0.05).  
Abbreviation: AM, α-Mangostin.

**Inhibition of TNF-α-induced NF-κB nuclear translocation by AM**

The obstruction to apoptosis and cell proliferation was considered to be closely related to the activation of nuclear factor kappa-B (NF-κB). Hence, the role of AM in the suppression of activated NF-κB induced by the inflammatory cytokine, TNF-α, was examined using Alexa Fluor 488-conjugated anti-NF-κB antibody (Thermo Fisher Scientific, Waltham, MA, USA). Although a high NF-κB fluorescent intensity was observed in the cytoplasm in the control cells (medium alone) (Figure 7A), it was only faint in the nuclei, which suggests an absence of NF-κB activation in the control cells (medium alone). In addition, TNF-α alone considerably intensified the NF-κB fluorescent intensity in the nuclei. AM demonstrated a considerable inhibitory effect on the triggering of NF-κB (Figure 7B). In the cells treated with curcumin, a known inhibitor of NF-κB activation, a significant suppression of TNF-α-induced NF-κB nuclear translocation was observed.
as evidenced by the low nuclear NF-κB-related fluorescence intensity (Figure 7B). In parallel, the morphological alterations of NF-κB translocation, as shown by the immunofluorescence staining (Figure 7A), indicated the inhibitory effect of AM on TNF-α-induced NF-κB translocation in a concentration-dependent manner with significant inhibition for the 20 µg/mL concentration of AM. To affirm the immunofluorescence result, we measured the immunoblotting of the nuclear content of NF-κB p65. As shown in Figure 7C and D, treatment with AM significantly decreased the escalation of this nuclear protein induced by positive control (TNF-α) in a dependent manner, particularly at 20 µg/mL of AM.

**Human apoptosis protein array**

Following AM exposure for 24 hours, the MCF-7 cells were lysed, and the apoptotic markers were screened using a protein array (Figures 8 and S1). In Figure 8C, the images show representative changes detected. All the major markers responsible for the apoptosis signaling pathway, including Bax, Bcl-2, Bim, cytochrome c, caspace-3/7 and -8 were expressed in the two models. A significant chaperone Hsp70, which played a part in the apoptosis was also downregulated, while the cell proliferation repressor protein p27, as well as X-linked inhibitor of apoptosis protein (XIAP) were also induced in this in vitro model. To confirm the protein array result, the expression of Hsp70 in the MCF-7 cells treated with AM was determined using Western blot analysis. AM treatment resulted in a downregulation of this protein in a concentration dependent manner (Figure 8D and E).

**AM reduced tumor size in vivo**

Table 2 presents the volume and the reduction of tumor percentage (%) of control and experimental animals. The tumor growth was measured using a caliper. The tumors in the mammary tumor control group grew rapidly, reaching an average volume of 1,737±563 mm³ by day 28. Meanwhile, groups treated with AM 30 mg/kg and 60 mg/kg showed a significant (P<0.05) reduction in their tumor volume when compared with the mammary tumor.
control group. The AM 60 mg/kg treatment also resulted in greater reduction (79.2%) in tumor volume than the AM 30 mg/kg treatment (74.1%), whilst tamoxifen (10 mg/kg) treated groups reduced the mammary tumor by an average of 83.6% (Figure 9).

**Discussion**

The role of apoptosis, which is a highly regulated process of programmed cell destruction, is crucial in many functions, ranging from fetal development to adult tissue homeostasis. Tumors are attributed to uncontrolled...
proliferation as well as reduced apoptosis. One critical method by which cytotoxic drugs destroy cancer cells is the activation of apoptotic pathways.\textsuperscript{32} Herbal medicines have been a major source from which numerous apoptosis-inducing agents are derived,\textsuperscript{13} and, according to several reports, many of these naturally occurring compounds may contribute partially to human cancer prevention or therapy.\textsuperscript{34} These studies showed that bioactive compounds elicit apoptosis in cancer cells.\textsuperscript{35} Renewed interest in the application of oriental medicine for cancer treatment, along with auspicious clinical results, has led to much emphasis being placed on medicinal plants. Nevertheless, the chemical components as well as definite mechanisms of many herbal medicines remain obscure.

In this regard, \textit{C. arborescens} is one of the well-known plants used in Asian countries for preventing and treating different kinds of ailments.\textsuperscript{14} AM, as a natural compound, is a major prenylated xanthone isolated from this plant. Therefore, the present study elucidated the mechanism of apoptosis provoked by AM toward MCF-7 cells. According to Shier,\textsuperscript{26} compounds that demonstrate an IC\textsubscript{50} value of more than 30 \(\mu\)g/mL are considered as not potentially cytotoxic, while compounds with an IC\textsubscript{50} value of less than 5.0 \(\mu\)g/mL are considered very active. These findings show that AM acts differently on normal cells compared to cancer cells that are more cytotoxic toward mammary gland cancer cells than normal cells. Since we found that the cytotoxicity produced by AM is within a potential limit, we used AO and PI fluorescent dyes to observe the different stages of apoptosis, beginning from the condensation of the chromatin up to the formation of apoptotic bodies, with AM treatment. Although the morphological features were clearly noticed, the assay of AV was conducted in an attempt to quantify the cells of the apoptotic population. The present study established that AM treatment can induce cell death in MCF-7 cells through apoptosis. In addition, the results showed a significant dose-dependent increase occurring in the early stage of apoptosis.

Although both extrinsic and intrinsic pathways are involved in apoptosis, the sensitivity of the intrinsic pathways causes tumors to occur more frequently through this route.\textsuperscript{37} Mitochondria are the main cellular components for the intrinsic means of apoptosis due to their ability to directly initiate the apoptotic cellular program. The primary involvement of the mitochondria is in the cell’s redox status, although they can also execute multiple cellular functions, including energy production, as well as cell proliferation and death.\textsuperscript{38,39} The permeabilization of the outer membrane and alteration of the mitochondrial transmembrane potential (\(\Delta\psi\text{m}\)) by the mitochondria initiating the apoptotic cascade, releases soluble pro-apoptotic proteins including cytochrome c, which, ultimately, leads to the activation of caspaces 9 and 3.\textsuperscript{37} It is essential that the potential of the mitochondria membrane is decreased for this process to begin. The effect of AM on the mitochondria, which causes less MMP was revealed by the fluorescence-based high content screening analysis. Concurrently, the release of cytochrome c also increased. There is much evidence that suggests that there is a role of oxidative stress in the apoptosis and related mitochondrial changes.\textsuperscript{40} In order to check the involvement of ROS, we measured the ROS level upon AM treatment toward MCF-7. The results clearly underline this significant relation (\(P<0.05\)). There was elevated (3-fold) intracellular ROS with AM treatment (20 \(\mu\)g/mL) on MCF-7 cells and this could be due to the free radical generation during the cytotoxicity.

### Table 2 Effect of treatment with AM 30 mg/kg, AM 60 mg/kg and tamoxifen on tumor volume (mm\(^3\)) in experimental breast cancer in rats

<table>
<thead>
<tr>
<th>Treatment groups</th>
<th>Tumor volume (mm(^3))</th>
<th>Reduction of tumor percentage (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal control</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Mammary tumor control</td>
<td>1,773±563</td>
<td>–</td>
</tr>
<tr>
<td>AM 30 mg/kg LD</td>
<td>460±33.5(^*)</td>
<td>74.1(^%)*</td>
</tr>
<tr>
<td>AM 60 mg/kg HD</td>
<td>372±24.6(^*)</td>
<td>79.2(^%)*</td>
</tr>
<tr>
<td>Tamoxifen 10 mg/kg</td>
<td>290±21(^*)</td>
<td>83.6(^%)*</td>
</tr>
<tr>
<td>AM 60 mg/kg</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

Notes: Each value represents mean ± SD of given number of animals (n=5). The statistical significance is expressed as \(P<0.05\).

Abbreviations: AM, \(\alpha\)-Mangostin; LD, low dose; HD, high dose; SD, standard deviation.

Figure 9 Therapeutic effect of AM treatment on LA7-induced mammary tumors in rats (4 weeks).

Notes: Reduction of tumor size of rats in treated groups: low dose 30 mg/kg AM (B), high dose 60 mg/kg AM (C), and tamoxifen 10 mg/kg (D) compared to the mammary tumor control (A).

Abbreviation: AM, \(\alpha\)-Mangostin.
Concurrently, a number of Bcl-2 family members, including Bcl-2 and Bax, are known to have the ability to control apoptosis. In ascertaining the vulnerability of cells toward death, it is crucial for a balance to exist between anti-apoptotic and pro-apoptotic Bcl-2 family members. It has been shown in previous reports that susceptibility to mitochondria mediated apoptosis can be caused by the upregulation of Bax and downregulation of Bcl-2. Thus, we examined the effect of AM on the expression of proteins that belong to this family. Bcl-2 is a cytoplasmic protein that has a significant part in inhibiting apoptosis that is caused by factors, such as chemotherapeutic agents, irradiation, as well as the withdrawal of growth factors. The immunofluorescence results in this research demonstrated the Bcl-2 downregulation in MCF-7 cells after treatment with 20 µg/mL AM. This occurrence likely explains the apoptotic effects of AM on MCF-7 cells. The exertion of these effects on the expression of Bcl-2 may also be connected to the production of mitochondrial apoptotic factors, which, eventually, lead to apoptosis. Our findings regarding AM support previous findings that showed that the induction of apoptosis seems to be modulated by the levels of Bcl-2 and Bax in the head and neck squamous carcinoma cell line (HNSCC), human Caucasian colon adenocarcinoma cell line (COLO 205), undifferentiated human colon carcinoma cell line (MIP-101), and human colon epithelial adenocarcinoma cell line (SW 620).

Subsequently, with MMP disruption and the release of cytochrome c, AM treatment of MCF-7 cells triggered the activation of caspaces 3/7, 8, and 9. In fact, the stimulation of caspaces 9 and 8 happened with a minimum concentration of AM, while caspace 3/7 activation only took place at a maximum treatment concentration. Caspace 9 is located in the intermembrane space of the mitochondria. It is discharged in a Bcl-2-inhibitable manner following the induction of permeability transition, whereas, in cells, the release is upon the induction of apoptosis. The caspace 9 that has been released then activates the post-mitochondrial caspaces, which include caspaces 3 and 7; the disassembly of the cell takes place in what is termed the execution phase of apoptosis. Caspace 8 is closely implicated in apoptosis signaling via the extrinsic pathway, albeit it is found in the upstream and downstream of the mitochondria. Moreover, the engagement of caspace 8 in mitochondrial pathways through the cleavage of the Bcl-2 family member Bid to tBid has been shown on numerous occasions. Our results regarding AM are well correlated with a previous study on AM against BJMC3879 metastatic murine mammary adenocarcinoma cells that showed the activation of caspaces 3, 9, and 8.

PARP can detect DNA damage due to its sensitivity. This enzyme is strongly triggered by DNA strand breaks and has primary functions in processes, such as DNA repair and the maintenance of genome stability. In addition, it also performs a crucial role in the induction of cell death initiated by a variety of stimuli. The proteolytic degradation of PARP that takes place at the beginning of apoptosis is caused by the activated caspace 3. As demonstrated by the immunoblot analysis, AM treatment significantly reduced PARP. Based on the results, it was shown that the stimulation of caspaces 9 and -3 is induced by the released cytochrome c, which, in turn, cleaves 116 kDa PARP into 85 kDa fragments in MCF-7 treated cells. PARP plays a crucial role in DNA repair and interacts with many DNA replication/repair factors, including the PCNA, a protein involved in many DNA transactions. Our next aim was to detect the response of AM toward this interaction. The anti-proliferative efficacy of AM has been established as it was found to decrease the immunoblotting expression of this proliferative marker in MCF-7 cells.

The mode of apoptosis caused by numerous natural compounds is closely related to cell cycle arrest. It has been established that cell cycle control is a significant event in safe-guarding precise cellular division. Many carcinogenic processes are reported to cause the abnormalities of cell cycle regulators. For this reason, it is reasonable to target and alter the cell cycle regulators in cancer cells for the purpose of chemoprevention and treatment. Following dosage with various concentrations of AM, the examination of the cell cycle of MCF-7 cells exhibited a higher number of cells in the $G_0/G_1$ phase. On the other hand, the amount of cells in the S and $G_2/M$ phase was reduced in comparison with the untreated cells (Figure 6). These results indicate the ability of AM in inhibiting cellular proliferation via $G_0/G_1$ phase arrest. This corroborates a previous study concerning the effect of AM against BJMC3879 cells, which demonstrated the inhibition of cellular proliferation via $G_0/G_1$ phase arrest.

The nuclear factor kappa-light-chain-enhancer of activated B cells, also known as NF-kB, is a protein complex that plays a role in regulating DNA transcription. In addition, it has also been considered as an apoptosis inhibitor. Thus, repression of the NF-kB activity can induce apoptosis. In this study, we have demonstrated that AM can prevent the TNF-α-induced NF-kB translocation of the cytoplasm to the nucleus of the MCF-7 cells, suggesting the participation of an NF-kB inhibition mechanism in apoptosis.
Apoptosis involves a large number of proteins. To determine the role of the central apoptosis-related proteins, a protein array analysis was conducted. In this study, various proteins in the extrinsic and intrinsic pathways were examined. These proteins include those that are known to cause apoptosis, including Bax, caspase-3, cytochrome c, caspace-8, and second mitochondria-derived activator of caspase, as well as those that have been recognized as anti-apoptotic, including Bcl-2, XIAP, of which XIAP is a member of the inhibitors of apoptosis (IAP) family of proteins, and second mitochondria-derived activator of caspase is a pro-apoptosis protein that acts together with IAP to reduce their inhibitory effects. Together with the Bcl-2 family members, Hsps have also been deemed to be apoptosis inhibitors, due to their significant role in cell survival, through stopping the release of cytochrome c from mitochondria or by preventing the formation of apoptosome. The protein array analysis showed a significant reduction in Hsp70 following treatment with AM. This finding is consistent with the findings from an earlier study that indicated that apoptosis could be suppressed by an over-expression of Hsp70. The results from the protein array displayed a characteristic profile of protein levels related to the mitochondrial apoptosis in MCF-7 cells treated with AM.

In the second part of this study, we also evaluated the effect of AM on tumor growth in Sprague-Dawley rats induced with LA7 cells to develop breast cancer for a duration of 28 days. The AM-treated groups (30 mg/kg and 60 mg/kg) showed significant tumor size reduction compared with the mammary cancer untreated group, indicating the potential chemotherapeutic value of AM in vivo. Our results regarding AM are well correlated with a previous study on AM against breast cancer untreated group, indicating the potential chemotherapeutic value of AM in vivo. According to the observations mentioned in this report, it can be concluded that AM has the ability to induce apoptosis in MCF-7 cells with cell death-transducing signals that regulate the MMP through a downregulation of Bcl-2 and an upregulation of Bax, which, in turn, triggers the release of the cytochrome c from mitochondria to cytosol. The activation of caspace-9 is triggered once cytochrome c enters the cytosol, followed by the activation of the downstream executioner caspase-3/7. Subsequently, it cleaves specific substrates, making way for the occurrence of apoptotic changes. An increase of caspace-8 has shown the involvement of extrinsic pathways. This form of apoptosis was suggested to occur through both the extrinsic- and intrinsic-apoptosis pathways with regulation of NF-κB, Bax/Bcl-2, and Hsp70 protein modulation. In addition to that, ingestion of AM at (30 or 60 mg/kg) significantly reduced tumor size in an animal model of breast cancer.

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Disclosure
The authors declare that there are no conflicts of interest in this work.

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Supplementary materials

Figure S1 MCF-7 cells were lysed and protein arrays were performed.

Notes: Cells were treated with 20 μg/mL AM and the whole cell protein was extracted. Equal amounts of (300 μg) of protein from each sample were used for the assay (A and B).

Abbreviations: BAD, Bcl-2-associated death promoter; Bcl-w, Bcl-2-like protein 2; IGF-II, insulin-like growth factor 2; IGF-I, insulin-like growth factor 1; HTRA, high temperature requirement A; Hsp70, heat shock protein 70; Hsp27, heat shock protein 27; Fasl, Fas ligand protein; Fas, Fas receptor protein; DR6, death receptor 6; IGFBP, insulin-like growth factor binding protein; Livin, melanoma/kidney inhibitor of apoptosis protein; p21, cyclin-dependent kinase; p53, tumor suppressor p53; TNF-α, tumor necrosis factor alpha; AM, α-Mangostin.