Induction of Apoptosis and Antiproliferative Activity of Naringenin in Human Epidermoid Carcinoma Cell through ROS Generation and Cell Cycle Arrest

Md Sultan Ahamad1,*, Sahabjada Siddiqui2,*, Asif Jafri3, Sheeba Ahmad3, Mohammad Afzal4, Md Arshad2*

1 Department of Zoology, Shibli National (PG) College, Azamgarh, Uttar Pradesh, India, 2 Molecular Endocrinology Lab, Department of Zoology, University of Lucknow, Lucknow, Uttar Pradesh, India, 3 Department of Zoology, D S College, Aligarh, Uttar Pradesh, India, 4 Human Genetics and Toxicology Laboratory, Department of Zoology, Aligarh Muslim University, Aligarh, Uttar Pradesh, India

Abstract

A natural predominant flavanone naringenin, especially abundant in citrus fruits, has a wide range of pharmacological activities. The search for antiproliferative agents that reduce skin carcinoma is of great importance. The objective of this study was to analyze the anti-proliferative and apoptotic mechanism of naringenin using MTT assay, DNA fragmentation, nuclear condensation, change in mitochondrial membrane potential, cell cycle kinetics and caspase-3 as biomarkers and to investigate the ability to induce reactive oxygen species (ROS) initiating apoptotic cascade in human epidermoid carcinoma A431 cells. Results showed that naringenin exposure significantly reduced the cell viability of A431 cells (p<0.01) with a concomitant increase in nuclear condensation and DNA fragmentation in a dose dependent manner. The intracellular ROS generation assay showed statistically significant (p<0.001) dose-related increment in ROS production for naringenin. It also caused naringenin-mediated epidermoid carcinoma apoptosis by inducing mitochondrial depolarization. Cell cycle study showed that naringenin induced cell cycle arrest in G0/G1 phase of cell cycle and caspase-3 analysis revealed a dose dependent increment in caspase-3 activity which led to cell apoptosis. This study confirms the efficacy of naringenin that lead to cell death in epidermoid carcinoma cells via inducing ROS generation, mitochondrial depolarization, nuclear condensation, DNA fragmentation, cell cycle arrest in G0/G1 phase and caspase-3 activation.

Introduction

Apoptosis plays a crucial role in the normal development and pathology of a wide variety of tissues [1]. However, most cancer cells do not undergo apoptosis due to impairment of apoptotic signal transmission [2]. Triggering apoptosis in tumor cells can therefore be an effective strategy in anticancer therapy. Apoptosis can be initiated through two different mechanisms, the extrinsic death-receptor dependent pathway and a mitochondria-dependent or intrinsic pathway [3]. The extrinsic pathway is activated by cell-surface ligand-gated death receptors such as the tumor necrosis factor receptor super-family. Intrinsic apoptosis is initiated by disparate extracellular and intracellular stimuli. Mitochondrial release of cytochrome c directly triggers caspase-3 activation. Caspase-3 then cleaves cellular targets to promote apoptosis [4,5].

Over 4000 different flavonoids have been identified in the human diet and the daily intake ranges from 23 mg to 1 g [6]. It shows an inverse association between consumption of flavonoids and the risk of human cancers at many sites [7]. The flavanone naringenin, especially abundant in citrus fruits such as grapefruit (Citrus paradisi), oranges (Citrus sinensis) and in tomato (Solanum lycopersicum) [8], is reported to have anti-proliferative effects in different cancer cell lines [9]. Generally cytotoxic and antitumor agents are non-selective and also kill normal proliferating cells. Identification of active cancer specific compounds remains a thrust area in drug screening and drug discovery mechanisms. Naringenin has been reported to induce apoptosis in various human cancer cells [10,11] and its higher concentrations (>300 μM) cause normal cell toxicity in human lung embryonic fibroblasts [12].

Oxidative stress induced DNA damage or deficient DNA-repair is reported to have an etiological or prognostic role in cancer [13]. Research also suggests that electrophiles could play a key role in chemical carcinogenesis. The oxidative inactivation of enzymes by free radicals and the accumulation of oxidized proteins may play a critical role in the alteration of cellular function and cell death.

Citation: Ahamad MS, Siddiqui S, Jafri A, Ahmad S, Afzal M (2014) Induction of Apoptosis and Antiproliferative Activity of Naringenin in Human Epidermoid Carcinoma Cell through ROS Generation and Cell Cycle Arrest. PLoS ONE 9(10): e110003. doi:10.1371/journal.pone.0110003

Editor: Rizwan H. Khan, Aligarh Muslim University, India

Received March 24, 2014; Accepted September 16, 2014; Published October 16, 2014

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Data Availability: The authors confirm that all data underlying the findings are fully available without restriction. All relevant data are within the paper.

Funding: This study is supported by the University Grant Commission (UGC), New Delhi, India as major research projects (No. 42–500/2013SR and 37-436/2009SR). Author Sahabjada Siddiqui is thankful to ICMR, New Delhi, India for Senior Research Fellowship (No. 45/26/2013/BMS/TRM). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* Email: molendolab@gmail.com

These authors contributed equally to this work.
Oxidative damage can generate large amounts of carbonyl products and hence measurement of these components could reflect oxidative protein damage [14]. However, the mechanism of action of naringenin to suppress cell growth is still ambiguous, since this compound appears to have multiple cellular targets including cytochrome P450 enzymes [15].

Several mechanisms have been proposed for naringenin induced anti-proliferative effects i.e. antioxidant activities, kinase and glucose uptake inhibition, and to hamper cell proliferation via estrogen receptor [16,17]. However, the composite (combined) involvement of reactive oxygen species (ROS), cell apoptosis, DNA fragmentation, mitochondrial membrane potential (MMP), cell cycle kinetics and caspase-3 activities in the molecular mechanisms of naringenin-induced anti-proliferative effects in human epidermoid carcinoma A431 cell line remains to be investigated. The aim of the present study was two-fold: (a) to analyze the anti-proliferative and apoptotic effects of naringenin through ROS mediated 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT), nuclear condensation, mitochondrial membrane potential and DNA fragmentation, and (b) cell cycle kinetics and caspase-3 induction as biomarkers in cancer cell viz. human epidermoid carcinoma A431 cell.

Materials and Methods

Cell line culture

Normal skin cell (HaCaT) and Human epidermoid carcinoma (A431) cell line were obtained from Cell Repositories, National Centre for Cell Sciences (NCCS), Pune, India. A431 was cultured in Dulbecco’s modified eagle medium (DMEM) and HaCaT cell was cultured in DMEM:F12 (1:1) medium supplemented with 10% (v/v) fetal calf serum (Himedia), 2.0 mM L-glutamine, 1.5 g/l NaHCO₃, 0.1 mM non-essential amino acids, 1.0 mM sodium pyruvate and 1% antibiotic solution. Cells were maintained at 37°C, 5% CO₂ in a humidified air.

MTT assay for cell viability in HaCaT and A431 cells

This assay is based on the enzymatic reduction phenomenon of 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) dye and provides a direct relationship between the viable cells and absorbance. The effect of naringenin on cell viability was evaluated by MTT reduction assay as performed earlier [18]. Selective doses viz. 50 μM, 100 μM, 200 μM, 300 μM, 400 μM, 500 μM and 750 μM of naringenin were prepared in dimethyl sulfoxide (DMSO) in final volume of 100 μl media. After 24 h exposure, 10 μl of MTT solution (5 mg/ml stock solution) was added in each well re-incubated for 3 h at 37°C until formazan blue crystal developed. Media was discarded from each well and 100 μl of DMSO was added to dissolve formazan crystals for 10 min at 37°C. The absorbance was recorded at 540 nm by microplate reader (BIORAD-680) and relative percentage cell viability was evaluated.

Cell morphology analysis

The effect of naringenin was analyzed for morphological changes in the cultured cells [19]. The cells were seeded at a density of 1×10⁴ cells/well in 96-well culture plate. After overnight incubation, the cells were treated with different concentrations of naringenin for 24 h. The cellular morphology was observed under inverted phase contrast microscope (Nikon ECLIPSE Ti-S, Japan).

Reactive oxygen species (ROS) activity assay

Microscopic fluorescence imaging was used to study ROS generation in A431 cells after exposure to different concentrations of naringenin [20]. Cells (1×10⁴ per well) were seeded as described above for the MTT assay. Cells were then exposed to 50 μM, 100 μM, 200 μM, 300 μM, 400 μM, 500 μM and 750 μM concentrations of naringenin for 12 h. Cells were incubated with 2,7-Dichlorodihydrofluorescein diacetate (DCFH-DA) (10 μM) for 30 min at 37°C. The reaction mixture was aspirated and replaced by 200 μl of phosphate-buffered saline (PBS) in each well. The plate was kept on a shaker for 10 min at room temperature in the dark. An inverted fluorescence microscope (Nikon ECLIPSE Ti-S, Japan) was used to visualize intracellular fluorescence of cells and to capture images. For quantitative ROS analysis, cells (1×10⁴ per well) were re-seeded in 96-well black bottom culture plate and allowed to adhere for 24 h in a CO₂ incubator at 37°C. A431 cells were treated with different concentrations of naringenin for 12 h. After exposure, cells were incubated with DCFH-DA (10 μM) for 30 min at 37°C. Fluorescence intensity was measured by multwell microplate reader (Synergy H1 Hybrid Multi-Mode Microplate Reader, BioTek) at excitation wavelength of 485 nm and emission wavelength of 528 nm. Values were expressed as the percentage of fluorescence intensity relative to the control wells.

DAPI staining for apoptosis analysis

The apoptotic effect of compounds was analyzed by using fluorescent nuclear dye 4’,6-diamidino-2-phenylindole dihydrochloride (DAPI) [21]. The cells were seeded and treated as mentioned previous. Cells were then washed with PBS and fixed in 4% paraformaldehyde for 10 min. Subsequently the cells were permealized with permealizing buffer (3% paraformaldehyde and 0.5% Triton X-100) and stained with DAPI dye. After staining, the images were captured and numbers of cells were quantified using a fluorescent microscope (Nikon ECLIPSE Ti-S, Japan).

Assessment of mitochondrial membrane potential

Flouroprobe 5,5’,6,6’-tetrachloro-1’,3’,3’-tetraethylbenzimidazol-carbocyanine iodide (JC-1) is a cationic, lipophilic dye and therefore has been extensively used to study the loss of the mitochondrial membrane potential which occurs during apoptosis [22]. In normal cells, due to high membrane potential (polarized mitochondria), the dye concentrates in the mitochondrial matrix, and it forms red fluorescent aggregates (J-aggregates). Any event that dissipates the mitochondrial membrane potential (depolarized mitochondria) prevents the accumulation of the JC-1 dye in the mitochondria and thus, the dye is dispersed throughout the entire cell leading to a shift from red (J-aggregates) to green fluorescence (JC-1 monomers). A decrease in red/green ratio is indicative of apoptosis. The cells were grown in 24-well plate and treated with different concentrations of naringenin. After 24 h exposure, the treated cells were washed with PBS and stained with 2 μg/ml of JC-1 dye in DMEM media without phenol red at 37°C in dark for 30 min. The photographs were then taken by inverted fluorescent phase contrast microscope and the mitochondrial depolarization patterns of cells for cells quantification were examined by using imaging software NIS-Elements F 4.00.00.

DNA extraction and fragmentation assay

Assay was done by the method of [23] with minor modifications. At the end of treatment period, cells were harvested and washed twice with cold PBS. The cell pellets were lysed in 500 μl DNA lysis buffer (20 mM EDTA, 10 mM Tris-HCl pH 8.0, 0.2% Triton X-100 and 100 μg/ml Proteinase K) for 1.5 h at 37°C.
The samples were centrifuged at 6000×g for 5 min to collect the supernatant containing DNA. The supernatant was added with equal volume of isopropanol and 25 μl 4M NaCl to make 1.0 M final concentration and incubated overnight at −20°C. Sample was then centrifuged at 6000×g for 25 min at room temperature. The pellet was dissolved in 50 μl diH2O and 2 μl RNase A (10 mg/ml) and reincubated at 37°C for 1 h. The extracted DNA was subjected to electrophoresis on 1.5% agarose gel containing ethidium bromide. The DNA bands were read under ultraviolet illumination gel-doc system (QUANTUM-ST-IV-1326.WL/20MX XPRESS; France).

Analysis of cellular DNA contents by flow cytometry
Cell cycle phase distribution with cellular DNA contents were carried out using flow cytometry. A431 cells were seeded into 6-well plate at density 1×10⁴ cells/ml and treated with 100 μM, 300 μM and 500 μM of naringenin for 24 h in 5% CO₂ incubator and at 37°C temperature [24]. After 24 h incubation, the cultured cells were harvested, washed with cold PBS, fixed in 70% ethanol and treated with RNase A (10 mg/ml). Fixed cells were stained with propidium iodide (PI) dye followed by incubation for 30 min at room temperature in dark. The PI fluorescence of individual nuclei was measured using flow cytometer (BD FACS Calibur, Becton Dickinson, USA). Data were analyzed with the Cell Quest Pro V 3.2.1 software (Becton Dickinson, USA).

Analysis of Caspase-3 activity
The caspase-3 activity was assayed using Caspase-3 Colorimetric Assay Kit (Catalog No. K106, BioVision, USA). Approximately 3×10⁶ (treated and untreated) of cells were resuspended with chilled lysis buffer. The cell lysate was incubated on ice for 10 min prior to centrifugation (10,000×g for 1 min). The reaction buffer, with 10 mM DTT was added to the supernatant of cell lysate and incubated further for 30 min on ice. The cell lysate (50 μl) was aliquoted into wells of 96-well microplate and 50 μl of reaction buffer containing 10 mM DTT was then added to the lysate. About 5 μl of 4 mM DEVD-pNA substrate was added in each wells and incubated at 37°C for 2 h. Absorbance at 405 nm was then read in a microplate reader. The absorbance of treated cells was compared with untreated control to find the change in caspase-3 activity.

Statistical analysis
Data were expressed as the mean ± SD from three independent experiments. One-way ANOVA and Dunnett’s multiple comparison test were performed using Graph Pad prism (Version 5.01) software for significance test, using a p value≤0.05.

Results
Effect of naringenin in normal HaCaT cells
To find out the experimental doses of naringenin, cytotoxic tests were performed on normal skin cells HaCaT using MTT assay and cell morphology assessment. Morphological study suggests that higher concentration 750 μM of naringenin changes cell morphology and induces cell death. However, doses 30 μM, 100 μM, 200 μM, 300 μM, 400 μM and 500 μM of naringenin did not affect significantly on cell morphology as revealed in Fig. 1C. Cell viability data showed that percent cell viability in normal cells was 91.97% and 85.40% at 400 μM and 500 μM concentrations of naringenin respectively i.e. cell death was statistically insignificant, however, significant cell death was observed at 750 μM of naringenin (Fig. 1D).

Naringenin induces morphological changes and inhibits cell viability of A431 cells
The cell morphology was assessed using inverted phase contrast microscopy under 20× objective (Fig. 1A). Photograph clearly revealed that morphological shapes of the cells were drastically changed being round in shape in the treatment groups which is characterized by cellular shrinkage, detachment from surface and deformation of cell bodies at higher concentrations. A dose dependent cell morphological changes were observed in A431 cells at 24 h treatment of naringenin. The effect of different concentrations of naringenin on cell viability was shown in Fig. 1B. The percent cell viability data indicates that at 50 μM, 100 μM and 200 μM of naringenin concentrations, cell viability reduced to 93.01%, 77.92%, and 63.69% respectively as compared to control, which further dropped to 46.35%, 30.82%, 24.47% and 15.44% (p<0.001) at 300 μM, 400 μM, 500 μM and 750 μM, respectively. Results revealed that last three doses are more cytotoxic. We have selected three optimum doses (100 μM, 300 μM and 500 μM) for further studies.

Naringenin causes nuclear changes with apoptosis within A431 cells
Nuclear apoptosis were observed by using fluorescent DAPI dye. Condensed and fragmented nuclei were regarded as apoptotic cells. As observed from photomicrograph, naringenin induces the nuclear condensation in a dose dependent manner as indicated by arrow (Fig. 2A). Maximum chromatin condensation was observed in the cells treated at 500 μM of naringenin. Furthermore, approximately 14.00% and 27.30% and 58.00% apoptotic cells were observed at 100 μM, 300 μM and 500 μM of naringenin, respectively (Fig. 2B).

Naringenin induces intracellular ROS generation in A431 cells
ROS generation was assessed at 12 h after naringenin treatment. Florescent micrograph of stained A431 cells (Fig. 3) depicts the effect of naringenin-induced intracellular ROS generation. The photomicrograph suggests that A431 cells treated with naringenin elevates ROS intensity in a dose dependent manner as compared to control (Fig 3A). Maximum ROS generation was observed in the cells treated at 500 μM of naringenin. The quantitative measurement of ROS intensity showed that 100 μM of naringenin elevates significant ROS production, approximately 7.43% (**p<0.01) as compared to control. However, ROS production was sustained increasingly to approximately 17.46% and 51.52% (**p<0.001) at the concentrations 300 μM and 500 μM of naringenin, respectively (Fig 3B). Increased ROS generation is involved in the induction of apoptosis through various pathways.

Naringenin modulates mitochondrial membrane potential (ΔΨm) of A431 cells
The increased and decreased florescent intensity of red and green florescent caused by JC-1 indicates the change in mitochondrial membrane potential (ΔΨm). Result indicated that cells treated with 100 μM, 300 μM and 500 μM of naringenin induced a strong green fluorescence in a dose dependent manner (Fig 4A). The merge images of JC-1 red and JC-1 green cell viability data indicates that at 50 μM, 100 μM and 200 μM of naringenin concentrations, cell viability reduced to 93.01%, 77.92%, and 63.69% respectively as compared to control, which further dropped to 46.35%, 30.82%, 24.47% and 15.44% (p<0.001) at 300 μM, 400 μM, 500 μM and 750 μM, respectively. Results revealed that last three doses are more cytotoxic. We have selected three optimum doses (100 μM, 300 μM and 500 μM) for further studies.

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66.25% and 113.80% at 100 μM, 300 μM and 500 μM of naringenin treatment, respectively (Fig. 4B).

**Naringenin induces DNA fragmentation of A431 cells**

Naringenin was tested to ascertain the DNA damage following exposure of various concentrations of compound to A431 cells. The results obtained from the DNA fragmentation assay (Fig. 5) showed the undamaged DNA in control cells whereas naringenin treated cells represent DNA fragmentation which was increased in a dose dependent manner. Small shearing was observed in the cells treated with 100 μM and 300 μM of naringenin, though maximum fragmentation was observed at 500 μM of naringenin exposure.

**Naringenin modulate cellular DNA content and induces G0/G1 phase arrest**

Cell cycle analysis with cellular DNA content was performed by flow cytometry. Apoptotic cells were estimated by calculating the number of sub-diploid cells in the cell cycle histogram. When cells were exposed to 100 μM of naringenin, apoptotic cells reported were 6.24% as compared to control. However, at 300 μM and 500 μM of naringenin treatment, apoptotic cells were markedly increased to approximately 14.39% and 26.32% respectively. Analysis of cell cycle revealed that the cells in G0/G1 phase were observed to be 35.86%. However, 100 μM of naringenin arrest the hypodiploid cells approximately 53.75% in G0/G1 phase while...
300 μM and 500 μM of naringenin arrest the cells 54.03% and 48.20% as compared to control (Fig. 6A and 6B).

Naringenin induces caspase-3 activity in A431 treated cells

The activity of caspase (cysteine-dependent aspartate-specific proteinase), an important biochemical feature in apoptotic signaling, was further investigated to determine whether the apoptosis was induced by naringenin. Caspase-3 is the main downstream effector caspase in apoptotic pathway and hence we compared its activity in treated and untreated control cells. The result showed that caspase-3 was significantly increased to 29.57% and 85.10% (***p<0.001) at concentrations 100 and 300 μM of naringenin. However, caspase-3 activity was dramatically increased to 114.02% (***p<0.001) at 500 μM naringenin concentration (Fig. 7).

Discussion

Phyto-chemicals have been demonstrated significant potential as anticancer therapeutic agents due to their ability to inhibit tumor growth, angiogenesis, and metastasis without many side effects [25]. Much of these activities come from flavonoids, which
are principal components of many such phytochemicals and demonstrate the capacity to inactivate carcinogens, inhibit angiogenesis, and halt cell proliferation and promote apoptosis [26]. Naringenin is a flavonoid abundant in citrus fruits, especially their peels and rinds, and has been suggested to exhibit antiproliferative effects against human tumor cells. As naringenin is generally present in foods bound to sugars as β-glycosides (i.e., naringenin), it was originally thought that absorption from the diet would be negligible. However, a number of studies have detected naringenin in human urine and plasma following oral doses of pure naringenin or the grapefruit juice [27,28,29,30].

Here we have endeavored to work out the effects of the naringenin on epidermoid carcinoma A431 cell line. Cell viability analysis suggests that naringenin halts proliferation of cells (Fig. 1B). Further morphological analysis of photomicrograph by phase contrast microscopy (Fig. 1A) showed that the morphological shapes of the cells were drastically changed being round in shape in the treatment groups which is characterized by cellular shrinkage and detached from surface, forming clusters. These results suggested that the treatment of naringenin significantly reduces the cell viability of cancer cell line in a dose dependent manner.

Figure 5. DNA fragmentation analysis. DNA fragmentation of A431 cell treated with 100 μM, 200 μM and 300 μM of naringenin. 1st well showing the 3 kb DNA marker. DNA laddering pattern showed that naringenin induces DNA fragmentation in a dose dependent manner.

doi:10.1371/journal.pone.0110003.g005

Figure 6. Effect of naringenin on different phases of cell cycle. A431 cells were treated with different concentrations of naringenin (100 μM – 300 μM) for 24 h, stained with propidium iodide and measured by flow cytometry (A) Representative photomicrograph showing the apoptosis and phase distribution of cell population (B) Quantitative distribution of percentage of A431 cells in different phases of cell cycle treated with different concentrations of naringenin. Data is representative of three independent experiments.

doi:10.1371/journal.pone.0110003.g006
The accumulation of free radical mediated damage may be a possible mechanism of cancer development [31]. Excessive ROS production can lead to oxidation of macromolecules and play an important role in mtDNA mutations, ageing, and cell death [32]. Therefore, increased intracellular ROS production might be implicated in cellular damage of skin carcinoma. Our experiments for the qualitative and quantitative analysis of ROS generation suggest that naringenin induced ROS mediated apoptosis induction in A431 cells. As compared the cell viability assay to ROS generation assay inhibiting cell viability and induction of apoptosis, both mechanisms showed similar trends. Mechanistically, it is well understood that the ROS damage almost all classes of sub-cellular components, are produced in numerous pathophysiological states and have been recognized as participating in the development of multistage carcinogenesis [33]. Oxidative stress is associated with damage to a wide range of micro and macromolecular species including lipids, proteins, and nucleic acids, thereby, producing major interrelated derangement of cellular metabolism including peroxidation of lipids, formation of protein carbonyls, and single strand breaks in DNA [34]. In addition, the intrinsic increase of basal ROS in cancer cell are highly dependent on antioxidant enzymes systems viz. glutathione (GSH) and glutathione peroxidase (GPX) under the influence of anti-oxidant agents which perturb the anti-oxidant system and lead apoptosis [35]. Previous report has demonstrated the dietary agent could effectively inhibited B(a)P induced lung carcinogenesis by offering protection from protein damage and also by suppressing cell proliferation [36]. Fragmented and condensed nucleus in the cells is a hallmark of apoptotic induction [37]. Our result also supporting such statement where fragmented and condensed nucleus stained with DAPI dye in A431 cells suggested that naringenin caused cell death by an apoptotic process. The loss of mitochondrial membrane potential ($\Delta \Psi_{m}$) is as an early event in apoptotic process as shown by earlier workers [38,39]. Our data also support this finding as shown in Fig. 4A and 4B that naringenin depolarizes the membrane potential significantly in a dose dependent manner. The results indicate that the sub fraction of naringenin possesses promising apoptotic potential.

DNA fragmentation is generally considered to be the hallmark of apoptosis. Our DNA fragmentation data support earlier observation that naringenin induced the DNA fragmentation and subsequent cellular damage (Fig. 5). Indeed, the nuclear DNA of apoptotic cells shows a characteristic laddering pattern of oligonucleosomal fragments. Several studies have demonstrated a positive correlation between DNA fragmentation and apoptosis [22,24].

The results support the findings that key changes in cell death can occur without DNA fragmentation and it alone should not be considered as a criterion for assessing apoptotic cell death [40]. To test the other parameters for apoptosis, we have performed cell cycle kinetic assay and observed that naringenin have antitumor actions, causing the cell cycle arrest, induction of apoptosis or a combination of these mechanisms, as earlier workers have also reported similar finding through different methods [41,42]. In order to elucidate the type of death induced by naringenin, apoptotic characteristics including hypodiploidy (sub-G1 peak) of the cells were analyzed by flow cytometry using PI staining. G1 and G2 phases of the cell cycle are the major check points and have an important role in cell cycle progression [43]. Result of cell cycle studies indicate that naringenin arrests the cells in G0/G1 phase of cell cycle and consequently lead to apoptosis (Fig. 6A and 6B). Caspase-3 is the primary effector caspase of the cell and plays an important role in the execution of apoptotic cell death by cleaving the cellular substrates [44], we then compared its activity in treated and untreated cells which favors that naringenin induced the caspase-3 activities and induction of caspase-3 activity leads to cell apoptosis (Fig. 7).

In conclusion, the present study provides a novel insight into the mechanism of action of naringenin-induced apoptosis in human skin carcinoma cells without/little affecting normal skin cells and shows a link between antiproliferative and apoptotic induction, and the cell death was due to the induction of ROS mediated mitochondrial membrane depolarization, nuclear condensation and DNA fragmentation. We also observed that naringenin induced cell cycle arrest at G0/G1 phase and caspase-3 activity. Our data confirm the potential of naringenin as an agent of chemotherapeutic and cytostatic activity in human skin carcinoma cancer and therefore, may be potentially valuable for application in drug developments. For further confirmation, molecular mechanism will be carried out to elucidate the molecular pathways.

Acknowledgments
Authors are thankful to research scholars of Molecular Endocrinology Lab, Department of Zoology, University of Lucknow, Lucknow and Human Genetics and Toxicology Laboratory, Department of Zoology, AMU, Aligarh (especially, Mohd Farred) for valuable discussion. Authors also acknowledge to AL Vishwakarma, SAIF-CDRI, Lucknow, for technical assistance in FACS analysis.

Author Contributions
Conceived and designed the experiments: M. Ahamad SS M. Arshad. Performed the experiments: SS M. Ahamad AJ M. Arshad. Analyzed the data: SS M. Ahamad SA M. Arshad. Contributed reagents/materials/analysis tools: M. Ahamad M. Arshad. Contributed to the writing of the manuscript: M. Ahamad SS SA M. Arshad. Language check, valuable suggestion and moral support: M. Afzal.

References
1. Hengartner MO (2000) The biochemistry of apoptosis. Nature 407: 770–776.
2. Evan GI, Vousden KH (2001) Proliferation, cell cycle and apoptosis in cancer. Nature 411: 342–349.
3. Ashkenazi A, Dixit VM (1998) Death receptors: signaling and modulation. Science 281: 1305–1308.
4. Fulda S, Debatin KM (2006) Extrinsic versus intrinsic apoptosis pathways in anticancer chemotherapy. Oncogene 25: 4798–4811.
5. Green DR, Kroemer G (2004) The pathophysiology of mitochondrial cell death. Science 305: 626–629.
6. Hertog MGL, Hollman PCH, Katan MB, Kromhout D (1993) Intake of anticancer chemotherapy. Oncogene 25: 4798–4811.
7. Frydoonfar HR, McGrath DR, Spiegelman AD (2002) The variable effect on proliferation of a colon cancer cell line by citrus fruit flavonoid naringenin. Colorectal Disease 5: 149–152.
8. Hertog MGL, Hollman PCH, Katan MB, Kromhout D, Science 305: 626–629.
9. Ashkenazi A, Dixit VM (1998) Death receptors: signaling and modulation. Science 281: 1305–1308.
10. Park JH, Jin CY, Lee BK, Kim GY, Choi YH, et al. (2008) Naringenin induces apoptosis through downregulation of Akt and caspase-3 activation in human leukemia THP-1 cells. Food Chem Toxicol 46: 3684–3690.
11. Matsuo M, Sasaki N, Saga K, Kaneko T (2005) Cytotoxicity of flavonoids toward cultured normal human cell. Biol Pharm Bull 28 (2): 253–259.
20. Alarifi S, Ali D, Ya AO, Ahamed M, Siddiqui MA, et al. (2013) Oxidative stress. Biochim Biophys Acta 1832: 169–176.

21. Kapoor R, Kakkar P (2012) Protective role of morin, a flavonoid, against high glucose induced oxidative stress and cell death. Apoptosis 12: 913–922.

22. Ashfaq M, Singh S, Sharma A, Verma N (2013) Cytotoxic evaluation of the seed kernel of Oenothera paradoxa against mongstad goldfish (Carassius auratus) [bambangan]. Food Chem Toxicol 48: 579–591.

23. Saraste A, Pulkk K (2006) Morphologic and biochemical hallmarks of apoptosis. Carcinogenesis 27: 328–337.

24. Ly JD, Grubb DR, Lawen A (2003) The mitochondrial membrane potential (deltapsi(m)) in apoptosis; an update. Apoptosis 8: 115–128.

25. Thirunavukkarasu C, Sakthivelan D (2001) Apigenin inhibits oxidative stress-induced macromolecular damage in Nitrosodimethylamine (NDMA)-induced hepatocellular carcinogenesis in Wistar albino rats. Mol Carcino 41: 11–20.

26. Ross JA, Kasum CM (2002) Dietary flavonoids: bioavailability, metabolic effects, and safety. Annu Rev Nutr 22: 19–34.

27. Ameer B, Weintraub RA, Johnson JV, Yost RA, Rouseff RL (1996) Flavanone absorption and plasma glucolipid effects in man. Nutrition 12: 275–280.

28. Fuhr U, Kummert AL (1995) The fate of naringin in human. A key to grapefruit juice-drug interactions? Clin Pharmacol Ther 58: 365–373.

29. Ishii K, Faruta T, Kasuya Y (1997) Determination of naringin and naringenin in human urine by high performance liquid chromatography utilizing solid-phase extraction. J Chromatogr B Biomed Appl 704: 299–305.

30. Lee YS, Reidenberg MM (1998) A method for measuring naringenin in biological fluids and its disposition from grapefruit juice by man. Pharmacology 56: 314–317.

31. Thirunavukkarasu C, Sakthivelan D (2001) Effect of selenium on N-nitrosodimethylamine induced multistage hepatocarcinogenesis with reference to lipid peroxidation and enzymic antioxidants. Cell Biochem Funct 19: 27–35.

32. Ott M, Gogvadze V, Orrenius S, Zhivotovsky B (2007) Mitochondria, oxidative stress and cell death. Apoptosis 12: 913–922.

33. Jeyabal PV, Syed MB, Venkataraman M, Sambhandham JK, Sakthivelan D (2005) Apigenin inhibits oxidative stress-induced macromolecular damage in Nitrosodimethylamine (NDMA)-induced hepatocellular carcinogenesis in Wistar albino rats. Mol Carcino 41: 11–20.

34. Young BS, Woodside JV (2001) Antioxidants in health and disease. J Clin Pathol 54: 176–186.

35. Trachootham D, Alexandre J, Huang P (2009) Targeting cancer cells by ROS-mediated mechanisms: a radical therapeutic approach? Nat Rev Drug Discov 8: 579–591.

36. Selvendiran K, Muntaz Banu S, Sakthivelan D (2004) Protective effect of piperine on benzo(a)pyrene-induced lung carcinogenesis in Swiss albino mice. Clinica Chimica Acta 350: 73–78.

37. Saraste A, Pulkk K (2006) Morphologic and biochemical hallmarks of apoptosis. Carcinogenesis 27: 328–337.

38. Ly JD, Grubb DR, Lawen A (2003) The mitochondrial membrane potential (deltapsi(m)) in apoptosis; an update. Apoptosis 8: 115–128.

39. Zhao M, Mydlarz WK, Zhou S, Califano J (2008) Head and neck cancer cell lines are resistant to mitochondrial-depolarization-induced apoptosis. Otolaryngol Head Neck Surg 139: 257–263.

40. Cohn SM, Sun XM, Snowden RT, Dinsdale D, Skillern DN (1992) Morphological features of apoptosis may occur in the absence of micro-nucleosomal DNA fragmentation. Biochim Biophys Acta 1169: 331–334.

41. Shen SC, Chen YC, Hsu FL, Lee WR (2003) Differential apoptosis-inducing effect of quercetin and its glycosides in human promyelocytic leukemia HL-60 cells by alternative activation of the caspase 3 cascade. J Cell Biochem 89: 1044–1055.

42. Abu Bakar MF, Mohamad M, Rahmat A, Farr SA, Fry JR (2010) Cytotoxicity, cell cycle arrest, and apoptosis in breast cancer cell lines exposed to an extract of the seed kernel of Mangifera pajang (bambangan). Food Chem Toxicol 48: 1688–1697.