Garcinone-E exhibits anticancer effects in HeLa human cervical carcinoma cells mediated via programmed cell death, cell cycle arrest and suppression of cell migration and invasion

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Abstract
Xanthones are an important class of natural compounds bearing huge bioactivity profiles. Garcinone-E is one among most active xanthones showing potential anticancer activity against various human cancer cell lines. Therefore, the current study was performed to explore the anticancer potency of naturally occurring garcinone-E xanthone against human HeLa cervical cancer cells. The underlying mechanism of action was also tried to be explored via testifying its induction of programmed cell death, arrest of cell cycle, suppression of cell migration, cell invasion and cell adhesion. MTT assay was implemented to estimate the viability of HeLa cells after garcinone-E exposure and clonogenic assay was used to analyze the effect on clonogenic potential. Acridine orange/ethidium bromine (AO/EB) staining assay was performed for monitoring of programmed cell death along with western blotting. Flow cytometric studies were carried out to analyze cell cycle check points. Transwell chambers assays were carried out for studying the impact of garcinone-E on migration and invasion potency of HeLa cells. Western blotting was used to study the expressions of apoptosis linked proteins in HeLa cells. Results indicated that garcinone-E remarkably decreased the viability to minimum in HeLa cells in both dose and time-reliant manner. The clonogenic capacity of HeLa cells was efficiently reduced by garcinone exposure. AO/EB staining showed that the anti-viability action of garcinone-E was apoptosis allied which was supported by western blotting as well. The cell cycle check points study indicated cell cycle arrest at G2/M-phase. HeLa cell migration and invasion were reduced efficiently after being subjected to garcinone-E treatment in a dose reliant fashion. In conclusion, garcinone-E has a remarkable potential to act as anti-cervical cancer chemopreventive provided further in vivo studies are required.

Keywords: Xanthones, Garcinone-E, Anticancer, Cervical cancer, Cell adhesion

Introduction
Natural products have long served human kind in copious aspects including food and medicine (Chen et al. 2018). Mangosteen, a tropically grown fruit in countries like Thailand, Malaysia and Indonesia, is a reservoir of a number of naturally occurring xanthones. Different parts of the plant like roots, barks and pericarps have long been used as herbal medicine in most of the Southeast Asian countries (Obolskiy et al. 2009). In traditional Thai medicine (TTM) mangosteen pericarps were used to cure inflammation, gonorrhea, leucorrhoea and abdominal pain (Pedraza-Chaverri et al. 2008). Recent studies have reported that isolated compounds out of the fruit
bear tremendous pharmacological applications like anti-microbial, antitumor, anti-inflamatory and antioxidant (Fang 2016; Xie et al. 2015; Sani 2015; Kritsanawong et al. 2016; Sidahmed 2016). Besides mangosteen contains several secondary metabolites, xanthones in particular have revealed efficient anticancer activity. Xanthones are a major class of natural polyphenols which contain xanthene-9-one structural skeleton. α-mangostin is a primary xanthone that has been extensively investigated but garcinone-E have also revealed potential pharmacological and biological properties (Ho et al. 2002). Garcinone-E has been reported to exert anticancer effects against different human cancer cell lines including colorectal, breast and hepatocellular carcinoma (Mohamed et al. 2017). Although, the anticancer nature of garcinone-E have been reported, its underlying mechanism of action is yet to be understood completely. Cervical cancer (CC), is a dangerous life threatening and a frequent gynecological disorder prevalent in women, globally. It has been ranked as 4th lethal malignancy in women with approximately 0.27 million deaths in the year of 2015 (Bray et al. 2018). 90% of deaths were recorded in LMIC (low and middle-income countries) and this mortality is 18 times the mortality in developed nations (WHO 2018). Among all subtypes of cervical cancer, almost all include a major-risk factor of long HPV infection (human papillomavirus infection). Vaccination programmes and screening HPV are the major and efficient strategies towards disease prevention (Crosbie et al. 2013). CC bears two histological subtypes: adenocarcina and squamous cell carcinoma, accounting for 25% and 75% of overall cervical cancer cases, respectively (Ries et al. 2007). Despite advances in CC treatment, diagnosis, screening and prevention, in recent past, substantial global and regional disparities in CC outcomes has led global gynecological cancer societies to issue evidence-based guidelines to improve quality of life in CC patients (Cibula et al. 2018). Keeping in view the need for novel chemotherapeutics for CC and anticancer potential of garcinone-E, present study was designed to investigate anticancer activity of garcinone-E against drug-resistant HeLa human CC cells. Its effects of selective and potent anticancer activity mediated via inducing programmed cell death, G2/M phase cell cycle arrest and suppressing cellular migration, cell invasion and cell adhesion, was also investigated.

Materials and method

Cell culture conditions

Human HeLa cervical carcinoma cells and normal HCeRePiC human cervical epithelial cells were procured from American Type Culture collection (ATCC, MD, United States). Both of the cell line were placed in Dulbecco’s Modified Eagle’s medium (DMEM; GIBCO, CA, United States) medium and cultured with 100 μg/mL of streptomycin (GibCO). Cell lines were maintained in a CO2 humidified incubator containing 95% air and 5% CO2. Exponentially growing cells were harvested for further performed experiments. Garcinone-E (98%) was procured from Wuhan Chem Faces Biochemical Co Ltd. Hubei, China.

Examination of cell proliferation

To carry out the proliferation assessment in garcinone-E (Wuhan Chem Faces Biochemical Co Ltd. Hubei, China) treated cervical HeLa cancer cells, MTT assay was used. HeLa cancer cells and normal HCeRePiC cells were harvested and placed with a density of 3 × 10⁵ cells/well onto 96-well plates and precultured overnight. Afterwards, each well was supplemented with variant garcinone-E drug doses viz 0, 8, 16, 32, 64, and 128 μM and cultured for different time intervals of 24 h, 48 h and 72 h. Further, each well was placed with a stock solution 100 μL of MTT (1 mg/mL) (Sigma, MO, United States) and incubated for 4 h. Cell suspension was completely cleared off and each well was added with DMSO (100 μL). Finally, SpectraMAX M5 microplate reader (Molecular devices, CA, United States) was used to record absorbance for assessment of optical density.

Examination of clonogenic potential

To estimate the effects of garcinone-EW on the cell clonogenic potency of HeLa cervical cancer cells, clonogenic assay was used. Exponentially growing cells were placed onto 6 mm cultural dishes with 450 colonies each dish. Each dish was added with different garcinone-E doses viz 0, 16, 64, and 128 μM and left on incubation for 48 h. Afterwards, cultural medium was substituted with fresh medium and HeLa colonies were left on incubation for further 12 days. Then HeLa colonies were fixed using paraformaldehyde and staining was performed with crystal violet (Sigma-Aldrich) for visualization of colonies. Finally, HeLa colonies were numbered under a light microscope (OLYMPUS, Japan).

Apoptosis assay

To analyze apoptosis in garcinone-E treated HeLa cancer cells, Ariffin et al. method of AO/EB staining was employed. HeLa cells were harvested at 80% confluence and subjected to garcinone-E exposure at varying doses viz 0, 16, 64 and 128 μM. Thereafter, AO/EB staining was accomplished with 1 μL of AO/EB stock solution containing 100 μg/mL each of AO and EB (Sigma, St. Louis, MO). Visualization of fluorescent HeLa cells was performed with a confocal Laser-Scanning microscope.
Examination of cell cycle
Flow cytometric studies were carried out to analyze various cell cycle check points in HeLa cells after garcinone-E exposure. HeLa cells were plated onto p60 tissue culture dishes with a concentration of $1 \times 10^4$ cells/mL and cultured with variant garcinone-E doses (0, 16, 64 and 128 μM) for 24 h. After garcinone-E treatment, cells were harvested and washed with PBS followed by centrifugation of 5 min. Ethanol (70%) was then used to fix the centrifuged HeLa cells followed by double washing with PBS. After washing, suspensions were treated with RNase and propidium iodide (Sigma-Aldrich) concentration of 50 μL and 25 μL respectively. Finally, Muse flow cytometry (Millipore, MA, United States) was used for DNA content quantification.

Transwell chambers assay
Transfection of cervical cancer HeLa cells was performed in transwell chambers by garcinone-E drug with variant doses viz 0, 16, 64 and 128 μM. The lower transwell chambers were placed with FBS (10%) and RMPI-1640 cultural medium (Corning Incorporated, Corning, NY, United States). In addition to this, upper chambers were placed with test drug and a concentration of $1 \times 10^5$ cells/well of target cells. Drug treatment was given for 12 h followed by 10 min of incubation at 4 °C. Un-migrated cells on the membranes were cleaned off by using a cotton swab and migrated once were stained. For staining crystal violet dye was used for 5 min. The migrated cells were pictured by using a light microscope (TS100; Nikon Corporation, Tokyo, Japan) under 200× of magnification power. Similar method was followed for invasion assay except chambers were coated with Matrigel.

Western blotting analysis
The expressions of different genes in garcinone-E treated cervical cancer cells (HeLa cells) were observed through western blotting assay. The cells were exposed to variant garcinone-E drug doses viz 0, 16, 64 and 128 μM for 24 h. Afterwards, cells were lysed used lysis buffer (Sigma Aldrich) and quantification of protein content within each lysate was performed with BCA assay. About 40 μg of proteins were separated through SDS-PAGE and transferred to PVDF membranes (Millipore, MA, United States) electrophoretically. These membranes were subjected to primary antibodies treatment and antibodies were used against Bax, Bl-2, and Caspases (3, 8 and 9). This treatment lasted for 12 h at 4 °C followed by secondary antibodies treatment (anti-rabbit IgG conjugated to HRP) with incubation at 4 °C overnight. Finally, the bands of proteins were observed through enhanced chemiluminescence (Pierce, Rockford, IL, United States).

Statistical analysis
The experimental data were represented as standard deviation and mean values after executing individual procedures in triplicates. Tukey’s multiple comparison test and ANOVA was used to analyze significance through GraphPad Prism (Demo, Version 5). p value of less than 0.05 was taken as statistically significant.

Results
Garcinone-E inhibited proliferation rate
Cancer cells proliferate at higher frequency in an uncontrolled manner. Proliferation suppression remains a huge breakthrough in cancer research and targeting it remains first priority of chemopreventives. Proliferation rate in cervical cancer HeLa cells was analyzed through MTT assay after treatment with variant garcinone-E (Fig. 1) doses viz 0, 16, 64 and 128 μM for 24 h, 48 h and 72 h. In case of control HeLa cells the proliferation remained almost unaffected. The proliferation in case of garcinone-E treated HeLa cancer cells (0–128 μM) showed significant decrease in viability from 100% to almost 10% (Fig. 2). The proliferation rate in normal cervical HCerEpiC cells also decreased but only to a lesser extent. The viability percentage decreased with increasing the time of garcinone-E drug exposure. The viability of garcinone-E treated HeLa cells were limited to almost 35%, 20% and 5% after 24 h, 48 h and 72 h respectively (Fig. 3). Therefore, it was evident from MTT assay that garcinone-E reduced proliferation rate selectively in cancerous HeLa cells rather than normal HCerEpiC cells in a concentration as well as time dependent manner.

Fig. 1 Chemical structure of garcinone-E molecule
Garcinone-E inhibited the HeLa cell colony formation
The ability of cancerous cells to develop as colonies was tested against HeLa cancer cells after being treated with garcinone-E. Variant garcinone-E drug doses were supplemented to HeLa cells for 12 days and then number of colonies against controls were calculated. Results indicated tremendous potential of garcinone-E as colony suppressant against HeLa cancer cells. The number of colonies was observed to reduce from 450 to almost 50 at higher drug concentrations (0–128 μM) (Fig. 4).

Garcinone-E induced apoptotic cell death
There are several pathways that serve as potential targets for chemopreventives to exert their anticancer effects. Apoptosis is a leading target and is known as type-I programmed cell death. The underlying mechanism of action of garcinone-E inducing antiproliferative effects was testified for inducing apoptosis. For that acridine orange/ethidium bromide (AO/EB) staining assay was performed. Results reported increased number of apoptotic (early and late) cells and necrotic cells with increased test drug doses (Fig. 5). Western blotting indicated that the expression of proapoptotic Bax increased in comparison to Bcl-2 which decreased effectively with drug doses. The expression of Caspases was also reported to be increasing with increasing garcinone-E doses (Fig. 6). Therefore, AO/EB staining and western blotting indicated that garcinone-E induced caspase-dependent apoptotic cell death in HeLa cancer cells.

Garcinone-E induced cell cycle arrest
The frequent proliferation rate indicates that cancer cells goes through uncontrolled mitosis. Mitosis goes through a series of check points and it is a lead target of chemopreventives to inhibit frequent proliferation. Flow cytometric analysis was carried out to monitor different cell cycle check points. Results indicated that the different phases of cell cycle were going normally in comparison to G2/M-phase. G2/M-phase cells accumulated on increasing the drug concentration from 0 to 128 μM (Fig. 7). Which indicated that garcinone-E induced G2/M-phase cell cycle arrest in HeLa cervical cancer cells.

Garcinone-E inhibits cell migratory and cell invasive potential in HeLa cells
The ability of HeLa cells to migrate and invade were examined via transwell chambers assay. Cell migration and invasion are the two vital features of metastatic
Fig. 3  MTT assay results indicating the rate of proliferation of cervical cancer HeLa cells and normal HCerEpiC human cervical epithelial cells. Both the cell lines were exposed to different time intervals of 24 h, 48 h and 72 h and variant concentrations (0–128 μM) of garcinone-E. Individual separate experiments were implemented three times considering p < 0.05 as statistically significant and data was signified as ±SD.

Fig. 4  Clonogenic studies of HeLa cells after being subjected to garcinone-E exposure for 12 days. Individual separate experiments were implemented three times considering p < 0.05 as statistically significant and data was signified as ±SD.
Fig. 5 Apoptosis assessment via AO/EB staining assay representing the early, late apoptotic and necrotic cell percentage after HeLa cells were subjected to garcinone-E exposure. Individual separate experiments were implemented three times.

Fig. 6 Western blotting assay indicating the expressions of apoptosis allied proteins and caspase (-3, -8 and -9) proteins. Individual separate experiments were implemented three times.

Fig. 7 Flow cytometric analysis of HeLa cells after being subjected to garcinone-E at indicated doses. Individual separate experiments were implemented three times considering $p < 0.05$ as statistically significant and data was signified as $\pm$SD.
malignant cancer cells. Therefore, targeting cell migration and invasion is among dynamic approaches to combat cancer metastasis. After treatment with varying garcinone-E doses (0–128 μM) HeLa cells were exposed to transwell chambers assay. Migrated cells were observed to decline with increasing garcinone-E doses in contrast to controls (Fig. 8). The effect of garcinone-E on invasion potency of HeLa cervical cancer cells displayed tremendous suppression of invaded cells doses-reliantly (Fig. 9). Therefore, garcinone-E reduced both cell invasion as well as cell migration potency of HeLa cervical cancer cells dose-reliantly.

Discussion
Cervical cancer is a harmful diseases, accounting for huge mortality and morbidity among women, globally. The lethality gets increased with asymptotic behavior of the disease at earlier stages. Lack of availability of effective chemopreventives and management strategies creates an emergency for promising anticancer agents for cervical cancer management (Campos et al. 2017). Naturally occurring xanthones have been reported with potential bio-activity profiles (Khursheed et al. 2016; Vieira and Kijjoa 2005). Herein, garcinone-E a naturally occurring xanthone in mangosteen fruit was investigated for its anticancer activity against cervical cancer HeLa cells. An attempt was also made to investigate the underlying mechanism of action. Apoptosis is one of the fundamental pathways in a cell that activates death to eliminate damaged cells and maintain normal tissue hemostasis. The stimulation of apoptosis takes place through two variant pathways that is intrinsic and extrinsic (Ng and Bonavida 2002; Gorczyca 1993). Extrinsic apoptosis gets activated by the death receptors and intrinsic apoptosis is mediated through mitochondria. The reduction in mitochondrial membrane potential results in leakage of cytochrome c into cytoplasm and initiates a number of reactions that eventually result in apoptosis (Wang et al. 1999). Herein, the current research was designed to explore the anticancer nature of garcinone-E against cervical cancer. The effects of inducing programmed cell death (PCD), G2/M-phase cell cycle arrest, suppression of cell migration, suppression of cell invasion and cell adhesion. The MTT assay was executed to gage the

Fig. 8 Transwell chambers assay results indicating the number of migrated HeLa cells after garcinone-E treatment. Individual separate experiments were implemented three times
antiproliferative nature of garcinone-E and revealed that garcinone-E subdue the proliferation rate in HeLa cells with a time as well as dose clinging fashion. Afterwards, studies were carried out to unleash the basic underlying mechanism of action behind the antiproliferative effects of garcinone-E. It was observed that garcinone-E induced PCD via induction of dose reliant apoptosis in HeLa cells. The levels of proteins (proapoptotic and antiapoptotic) were examined through western blotting and activity of proapoptotic proteins got enhanced after being subjected to garcinone-E drug. Cancer cells frequently undergo mitosis in an uncontrolled manner, which results in further spread of the disease (Matsukawa 1993). Targeting cell cycle in a cancerous cell is among major therapeutic targets in cancer treatment. Garcinone-E was observed to induce G2/M-phase cell cycle arrest in HeLa cells through flowcytometric analysis. Migration and invasion of cancer cells from source to distant places results in cancer metastasis which enhances the lethality of the disease. Thus, cell migration and invasion was checked in HeLa cancer cells after garcinone-E exposure through transwell chambers assay indicated subjugating of both. Results also indicated that cell adhesion was also limited to by garcinone-E drug in HeLa cells in a concentration reliant manner. Taking together, the current study evidenced that naturally occurring garcinone-E exhibits selective and potent anticancer properties in drug-resistant HeLa human cervical cancer cells. Garcinone-E induced programmed cell death, G2/M phase cell cycle arrest and suppressed cellular migration, cell invasion and cell adhesion.

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