Evaluation of cytotoxic profile of hydroalcoholic extract of fruit rinds of *Garcinia pedunculata* on human embryonic kidney and breast carcinoma cells

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INTRODUCTION

The dried fruit rinds of *Garcinia pedunculata* (GP) have many important medicinal properties and have been used in different disease conditions. *G. pedunculata* Roxb. Fam. Clusiaceae trees are commonly seen in north eastern states of India, Bangladesh and Andaman Nicobar islands. The tribal communities of these regions were regularly used the dried form of fruit rinds in their diet and found medicinal values. The folklore use claims that it has good therapeutic effect in fever, cough, bronchial asthma, rheumatoid arthritis, obesity, atherosclerosis, cardiovascular disease, infectious and inflammatory diseases.¹ It has been reported to possess the following vital phytochemical compositions such as garcinol, pedunculol, cambogacin and (−)-hydroxy citric acid.²,³ The hydro alcoholic extract of *G. pedunculata* (HAGP) has been reported to show cardio protective activities against isoprenaline induced myocardial infarction in rats.⁴ The HAGP has been found that the hepatoprotective activity...
against paracetamol induced hepatotoxicity. Previous pharmacological studies have stated that it has potent antimicrobial and anti-inflammatory properties with potent antioxidant and free radical scavenging activities. It has shown to have a protective activity against cisplatin induced nephrotoxicity in Wistar albino rats and protected kidney tissues from cisplatin induced severe oxidative stress. The hexane and chloroform fraction of GP has shown antimutagenicity properties.

The use of synthetic cancer chemotherapy is associated with high degree of adverse effects and overweighs its potential therapeutic benefits. Most of the chemotherapeutic agents kill both normal and tumorous cells and hence posed with numerous toxicity profiles. In this regard many phytomedicines derived from herbal sources were screened for their significant cytotoxic effects. It has been reported that the plant derived alkaloids such as vincristine, vinblastine and epipodophyllotoxines possess good cytotoxic effects in various solid tumors. Meanwhile, phytochemistry possess potent anti-oxidant and cytoprotective activity. However, there are no studies reported on cytotoxic effect of G. pedunculata. Thus, the aim of the present study was to screen the effect of hydro alcoholic extract of fruit rinds of G. pedunculata on normal (human embryonic kidney cells) and metastatic breast cancer cell lines.

METHODS

3-(4, 5-dimethylthiazol-2-yl)-2, 5-diphenyltetrazolium bromide (MTT) -Sigma Aldrich, dimethyl sulfoxide (DMSO), phosphate buffer saline (PBS), 96 well plate, human embryonic kidney (HEK-293), minimum essential medium eagle, fetal bovine serum, M.D. Anderson metastatic breast cancer cell lines (MDA-MB), and Leibovitz’s medium. All chemicals and reagents used were of analytical grade.

Plant material

The fruit rinds of G. pedunculata were collected from Assam during the month of April 2016. It was authenticated in Pharmacognosy laboratory at SDM Centre for Research in Ayurveda and Allied Sciences, Udupi and the voucher specimen (No. 13100501) has been deposited for future reference.

Extract preparation

The fruit rinds were shade dried and powdered at SDM Pharmacy Udupi. The powder obtained from a single batch was used throughout the study. Fruit powder of G. pedunculata weighing 100 g was soaked in 0.4 litre of cold distilled water for 24 hour was filtered and concentrated by evaporation. The concentrated extract was used for cell viability studies.

Cytotoxicity study

In the present study two cell lines, HEK-293 and MDA-MB-231 were opted. The confluent cell line was taken in flask and trypsinized the cells. The cells were washed twice with PBS and centrifuged it. The pellet was suspended in MEM (E) and Leibovitz medium with 10% foetal bovine serum respectively. The cells were counted using haemocytometer. Cells around 10,000 cells/well were plated to 96 well plate and incubated at 37°C in CO2 incubator for 24 hrs. After 24 hours of incubation old medium was discarded from 96 well plates. Dissolved the different concentrations of drug in suitable serum free medium and add to the different test groups. The cells were incubated for 24 hours at 37°C in CO2 incubator. After completion of incubation time added 20 µL of MTT dye (5 mg/ml in PBS) to all wells. The plates were covered with aluminum foil and incubate in CO2 incubator for four hours. After four hours added 100 µl of DMSO to all the wells and mixed it by careful shaking. The absorbance was recorded using multiwell plate reader record at 540 nm (or 540 nm with reference to 630 nm) and the percentage viable cells were calculated using following formula% of viable cells = [(Test sample-blank) / (Control-blank)] ×100.

RESULTS

Effect of HAGP on HEK-293

HAGP at graded dose has shown increase in the percentage viability of MDA-MB 231 cells. HAGP at 10, 20, 50, 100, 200, 500 and 1000 µg/ml concentrations has shown the% live cells of 80.768, 80.770, 83.222, 83.082, 95.203, 108.498 and 112.503 respectively. Control group has shown 100% live cells whereas; the Cisplatin at 500 µg/ml has shown 32.99% live cells (Figure 1).

Effect of HAGP on MDA-MB 231

HAGP at graded dose has shown decrease in the percentage viability of MDA-MB 231 cells. HAGP at 10, 20, 50, 100, 200, 500 and 1000 µg/ml concentrations has shown the% live cells of 91.067, 85.071, 82.256, 82.875, 88.102, 95.076 and 98.503 respectively. Control group has shown 100% live cells whereas; the Cisplatin at 500 µg/ml has shown 32.99% live cells (Figure 1).

Figure 1: Effect of HAGP on HEK-293.
81.309, 79.223 and 77.490 respectively. Control group has shown 100% live cells whereas; the Cisplatin at 500 μg/ml has shown 32.066% live cells (Figure 2).

![% Viability of MDA-MB 231 cells](image)

**Figure 2: Effect of HAGP on MDA-MB 231.**

**DISCUSSION**

The present study was conducted to evaluate the cytotoxic potentials of hydro alcoholic extract of *G. pedunculata* (HAGP) at various concentrations on both normal cell line such as human embryonic kidney cells (HEK-293) and metastatic breast carcinoma (MDA-MB 231) cells using direct counting of percentage viability of cells.

In the present study, we observe that there is a considerable increase in the percentage viable of HEK-293 cells when incubated at higher concentration of HAGP as compared to the lower doses. The minimum dose of the test drug selected for the present study was 10μg/mL and we observed 80.768 percentage of viable cells. However, at higher dose such as 500 μg/ml and 1000 μg/ml, we found 108.498 and 112.503 percentage viability respectively. The increase in percentage of viable HEK-293 cells was considerably higher when compared to the normal control group. These preliminary observations indicate that the test drug HAGP has a potential to stimulate normal cell proliferation. The exact mechanism of action and the important phyto-constituents responsible for increase in percentage of viable HEK-293 cells in the present finding could confirm only by further detailed investigations.

The effect of HAGP on Metastasis breast cancer cell line 231 has shown a considerable decrease in the percentage viability at various dose levels of HAGP. We observed 91.067 percentage viability at lowest concentration (10 μg/ml) and a graded decrease in the percentage viability was observed in the following order 85.071, 82.256, 82.875, 81.309, 79.223 and 77.490 at 20, 50, 100, 200, 500 and 100μg/ml. The result indicates that there is a significant decrease in the percentage viability of metastatic breast cancer cell line, when these were incubated with various dose levels of HAGP. The earlier study reported that *G. Pedunculata* has shown potent antioxidant and free radical scavenging activities. In addition, it has potent role in the protection of liver, kidney and cardiac tissues against various toxicants. The hexane fraction of *G. pedunculata* fruit rinds possess anti mutagenic properties. The phytochemicals such as garcinol, (-)- hydroxy citric acid and phenolic compounds such as pedunulol were responsible for its various pharmacological activities. The present preliminary study result indicates that HAGP has potential to kill or inhibit the rate of proliferation of cancer cells at various levels. However, there is a need of detailed exploration of present findings like the important phyto-constituent responsible for cytotoxic activity and dose selection.

**CONCLUSION**

The present study concludes that HAGP has a potent cytotoxic activity in the cancerous cell lines and has promising proliferator effect on normal cell line. Thus it can be a better alternative chemotherapeutic agent if it was refined and explored the specific phytomedicine responsible for cytotoxic effects on abnormal cell lines.

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