Effects of *Pueraria tuberosa* Linn Hydroalcoholic Tuber Extract on Expression of Apoptosis Associated Proteins in HT – 29 Human Colon Carcinoma Cell Line

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**Abstract**

Medicinal plants have been used as an excellent source for years together as an alternative medicine for treating various disorders including anticancer, cardiovascular disorders and immunological disorders. *Pueraria tuberosa* linn (PT) commonly known as Indian Kudzu is a folklore medicine used in Ayurvedic formulations against various disorders. The present study reports the anticancer potential of the hydroalcoholic tuber extract of PT which is rich in isoflavones. The cytotoxicity of PT tuber extract was evaluated by MTT assay. Further the apoptotic potential of the extract was confirmed by Ethidium bromide/acridine orange (EtBr/AO) staining assay. DNA fragmentation which is considered as the hallmark of apoptosis was also evaluated for the hydroalcoholic tuber extract of PT. The cellular DNA was stained and the distribution of cells in the sub-G1, G0/G1, S and G2/M phases were assessed by flow cytometry. The extract showed a 50% inhibition of cell viability against HT-29 cells at a concentration of 63.91µg/mL. HT-29 cells exhibited apoptotic cell death, increased the expression of caspases, p53 and Bax. Altogether, the results suggest that the hydroalcoholic PT tuber extract has a better anticancer activity against the tested human colon carcinoma cell line.

**Keywords**

Indian Kudzu, *Pueraria tuberosa*, HT – 29, Anticancer, Apoptosis, Cell cycle

**Introduction**

Over the past few decades, there has been tremendous progress in our understanding of the molecular biology of cancer. Nonetheless, we have not conquered this dread disease yet. Like the majority of other human disorders, cancer is basically preventable. One of the most promising approaches to reduce the risk of cancer is chemoprevention (Greenwald, 2001; Kelloff *et al.*, 2000). Chemo prevention is the attempt to use natural and synthetic compounds to intervene in the early precancerous stages of carcinogenesis, before malignancy manifests. Recently, there have been considerable efforts to search for naturally occurring substances for the intervention of carcinogenesis (Surh, 2003).
To date, some phytomedicines have been postulated to be potential cancer chemo preventive agents through their ability to prevent, inhibit or reverse carcinogenesis. This ability is attributed to the presence of phytomedicines of specific families including polyphenols, polysaccharides, terpenes and sulfur compounds, which may be the possible reason for the anticancer mechanism.

_Pueraria tuberosa_ commonly known as Indian kudzu is climber with woody tuberculated stem (Chopra, 1956), growing throughout tropical parts of India, mostly in moist regions. The plant constitutes major chemical constituents including isoflavone like daidzein, genistein, puerarin, puerarone, coumestan, tuberosin, pterocarpin tuberosin, puertuberosanol and hydroxyl tuberosone (Pandey et al, 1998). The tuberous root of PT is brown in color and slightly curved is used for rejuvenation treatment. The tubers of PT are described as sweet, refrigerant, emollient, laxative, aphrodisiac, galactogogue, diuretic, emetic, cardiotonic, expectorant, febrifuge and used for the treatment of various ailments (Devaiah, 2008). Moreover, it is also useful in emaciation of children, debility and poor digestion (Chopra, 1956; Vaidyaratnam, 1997). Its crude powder, ethanolic and butanolic extracts possess significant estrogenic activity as well as progestational and mild antiprogestational activity (Prakash et al., 1985). The antimicrobial activity effects of PT extracts from the tuber, leaf and root showed special attention towards hospital infection (Sadguna et al., 2015).

Antioxidant and anti – inflammatory activity of the tubers of PT and its active component puerarin was also reported recently (Pandey et al., 2007, Nidhi., 2010). Protection against stress induced myocardial ischemia by PT was also evaluated and it was the first clinical case to report cardio protective potential of PT tuber extract by its clinical improvement in angina and cardiac performance as well as favourable alterations in biochemical parameters without any untoward side effects (Verma., 2009).

Although the crude tuber extract of PT has been reported for various disorders such as hypoglycemic, antioxidant, cardioprotective, neuroprotective, anti – inflammatory and antiprostational no study have reported concerning the anticancer activity of tuber extract of PT against human colon carcinoma. Hence the present study is an attempt to explore the anticancer potential of tuber extract of PT since it has already been proven a good antioxidant.

**Materials and Methods**

Rosewell Park Memorial Institute (RPMI) 1640 medium Dimethyl sulfoxide (DMSO), penicillin, streptomycin, tryspin-EDTA, MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide). Fetal bovine serum (FBS) was obtained from GIBCO BRL (Gaithersburg, MD). Bcl-2, Bax, procaspase-3, cleaved caspase-3 and caspase - 9 were purchased from Cell Signaling Technology (Beverly, MA). β-actin and HRP-conjugated secondary antibodies were purchased from Santa Cruz Biotechnology (Santa Cruz, CA). All other chemicals were of analytical grade.

**Cell culture and drug preparations**

The Human Colon Adenocarcinoma cell lines (HT-29) were purchased from National centre for cell sciences (NCCS) Pune. Cells were maintained in RPMI 1640 medium (GIBCO BRL) supplemented with heat inactivated 10% fetal bovine serum (GIBCO BRL) and 2mM L-glutamine (Sigma Chemical Co), 100units/ml penicillin and 100μg/ml streptomycin and maintained at 37°C in a atmosphere of 5% CO₂ incubator at 95% air humidified.
Cell proliferation assay or MTT assay

Proliferation of HT - 29 cells was assessed by MTT assay (Safadi et al., 2003). The proliferation test is based on the colour reaction of mitochondrial dehydrogenase in living cells by MTT. Cells were plated in 96-well plate at a concentration of 5 × 10⁴ cells/well 24 h after plating. After 24h of cells incubation, the medium was replaced with 100μl hydroalcoholic extract containing medium at different concentrations (2 – 1024μg/ well) and incubated for 24h. Untreated cells served as control and received only 0.1% DMSO in which the fraction was prepared. At the end of treatment period, media from control and extract -treated cells was discarded and 20μl of MTT (5mg/ml PBS) was added to each well. Cells were then incubated for 4h at 37°C in CO₂ incubator. MTT was then discarded and the coloured crystals of produced formazan were dissolved in 200μl of DMSO and mixed effectively by pipetting up and down. Spectrophotometrical absorbance of the purple blue formazan dye was measured using an ELISA reader (BIORAD) at 570nm. Optical density of each sample was compared with control optical density and graphs were plotted.

Ethidium bromide/acridine orange (Dual staining)

Ethidium bromide/acridine orange staining was carried out by the method of Gohelet al., 1999. HT - 29 cells were plated at a density of 1×10⁶ in 48-well plates. They were allowed to grow at 37°C in a humidified CO₂ incubator until they were 70–80% confluent. Then cells were treated with 64μg/ml and128μg/ml (selected based on the IC₅₀ concentration) of hydroalcoholic PT extract for 24h. The culture medium was aspirated from each well and cells were gently rinsed twice with PBS at room temperature. Then equal volumes of cells from control and drug treated were mixed with 100μl of dye mixture (1:1) of ethidium bromide and acridine orange) and viewed immediately under Nikon inverted fluorescence microscope (Ti series) at 10x magnification. A minimum of 300 cells was counted in each sample at two different fields. The percentage of apoptotic cells was determined by [% of apoptotic cells = (total number of apoptotic cells/total number of cells counted) ×100].

DNA fragmentation assay

DNA extraction and agarose gel electrophoresis were performed using the following method. Briefly, 1×10⁶ cells were plated in 100-mm Petri dishes with DMEM containing 10% FBS. Cells were incubated for 24 h in5% CO₂ and 95% air at 37°C. Control cells received0.1% DMSO containing DMEM and PT extract –treated cells received 50 and 75μg/ml of PT extract –containing DMEM. After 24 h, the cells were trypsinized and combined with the cells in the medium by centrifugation at1500 r.p.m. for 5 min, and then they were washed twice with PBS. The resulting pellet was resuspended in 0.25mlof lysis buffer, transferred to a microfuge tube, and incubated for 1h at 37°C. To this 4μl of proteinase K was added and tubes were then incubated at 50°C for3h. To each tube, 0.5ml of phenol: chloroform: isoamylalcohol (25:24:1) was added, mixed and centrifuged at 13000 r.p.m. at 4°C to separate the DNA containing upper aqueous phase. To the resultant aqueous phase, two volumes of ice-cold absolute ethanol and 1/10the volume of 3M sodium acetate were added and kept at –20°C overnight to precipitate DNA. The DNA was pelleted by centrifuging at 13000r.p.m. for 30 min at 4°C to separate the DNA containing upper aqueous phase. To the resultant aqueous phase, two volumes of ice-cold absolute ethanol and 1/10the volume of 3Msodium acetate were added and kept at –20°C overnight to precipitate DNA. The DNA was pelleted by centrifuging at 13000r.p.m. for 10 min at 4°C and the supernatant was aspirated and the pellet washed in 1ml of 70% ethanol. After repeating the above centrifugation step and removing last traces of the supernatant fraction, the pellet was allowed to dry at room temperature for approximately 30 min before being resuspended in 50μl of Tris-EDTA buffer.
DNA was quantified by ultraviolet-visible spectroscopy and 10μg of DNA was electrophoresed in 1.5% agarose gel containing ethidium bromide in an i mini gel tank containing Tris-borate-EDTA buffer for 2h at 90 V. The gel was then examined under ultraviolet light and photographed.

Flow cytometry

Flow cytometric analysis was carried out as described by Rasola and Geuna (2001). Briefly, 1×10^6 cells were plated in 100-mm Petri dishes with DMEM containing 10% FBS. Cells were incubated for 24 h in 5% CO₂ and 95% air at 37°C. Control cells received 0.1% dimethyl sulphoxide (DMSO) containing DMEM, and PT extract-treated cells received 50 and 75μg/ml of PT extract containing DMEM. After 24 h, the cells were trypsinized and combined with floating cells in the medium they were used for flowcytometry assay. The treatment protocol is as follows: 1×10^6 cells were taken from control and from PT extract-treated plates and were centrifuged at 1000g for 5 min. Supernatant was removed and cells were washed twice with PBS. The pellet was resuspended in approximately 500μl of ice-cold PBS and cells were mixed by aspiration 20 times using a pipette. Cells were fixed by adding 5 ml of cold ethanol drop by drop and were kept at −20°C over night. After overnight fixation, ethanol was removed by centrifuging at 1000g for 10 min. The pellet was washed twice with PBS + 1% BSA (ethanol-fixed cells were difficult to pellet; adding BSA or serum to the wash medium overcame this). The pellet was resuspended in 800μl of PBS containing 1% BSA. One hundred micro litre of 10×propidium iodide solution was added (500μg/ml propidium iodide in PBS, pH 7.4) and one hundred micro litre of RNase A was added (10 mg/ml prepared in 10 mm Tris-HCl, pH 7.5) and incubated at 37°C for 30 min.

Western blotting

Western blotting was carried out as described previously (Tu et al., 2004). Briefly, HT-29 cells (1×10^6) were seeded onto 100-mm culture dishes in the presence or absence of extract, and were treated for 24h. The medium was removed and the cells were washed with PBS (0.01M, pH7.2) for several times and lysed on ice in lysis buffer containing 100μg/ml phenyl methyl sulfonyl fluoride (PMSF), 50 mM Tris–base at pH 8.0, 150mM NaCl, 0.02% NaN₃, 1% NP-40, and 1μg/ml aprotinin. The supernatants were collected by centrifugation at 10,000xg for 5 min at 4°C, and were used as the cell protein extracts. The harvested protein concentration was measured using a protein assay kit (Bio-Rad). Equal amounts of total protein (30μg/well) were subjected to electrophoresed on 10% SDS–polyacrylamide gel and electro transferred onto PVDF membrane. Proteins were blocked overnight with 5% non-fat dried milk in PBS-T at 2-8°C. After washing in PBS containing 0.1% Tween 20 for 3 times, the membrane was incubated with the specific primary antibodies [Bcl-2 (rabbit polyclonal antibody at a dilution of1:500), Bax (rabbit polyclonal antibody at a dilution of1:500), procaspase-3 and cleaved caspase 3 (goat polyclonalantibody at a dilution of 1:250), and Caspase 9 (1:1000)] in 5% (w/v) skim milk in PBST. After overnight incubation at 4°C, the membrane was then washed three times with TBST, incubated further with alkaline phosphatase conjugated goat anti-mouse antibody, or anti-rabbit antibody at room temperature for 2 hours, and then washed three times with TBST. After reaction with horseradish peroxidase-conjugated goat anti-mouse antibody, the immune complexes were visualized by using the chemiluminescence ECL PLUS detection reagents following the manufacturer’s procedure (Amersham Bioscience).
Statistical analysis

Data were expressed as mean±S.E.M and analyzed by Tukey’s test to determine the significance of differences between groups. A \( p \) value lower than 0.05, 0.01 or/and 0.001 was considered to be significant.

Results and Discussion

Cytostatic effects of Pt hydroalcoholic extract against HT – 29 colon carcinoma cells

The in vitro cytotoxic effect of hydroalcoholic PT extract was screened against HT-29 cell line and viability of tumour cells was confirmed using MTT assay. The PT hydroalcoholic extract was able to reduce viability of the HT - 29 cells in a dose-dependent manner as shown in (Figure 1). The IC\textsubscript{50} value was found to be 63.91µg.

PT tuber extract induced apoptosis

Acridine Orange/Ethidium Bromide (AO/EB) was done to evaluate the type of cell death induced by the PT extract in HT - 29 cells; the morphological variations after double staining were investigated. Live cells stained with AO emitted green fluorescence. Early apoptotic cells had fragmented DNA which exhibited intense green coloured nuclei. Late apoptotic and necrotic cell’s DNA were fragmented and stained orange and red. From the data it was clear that with increasing concentration of drug, the number of viable cells decreased tremendously. The percentage of apoptotic cells after treatment with 64µg/ml and 128µg/ml of drug was drastically increased (\( p<0.001 \)) to 37% and 66% respectively (Figure 2).

DNA fragmentation

Evaluation of apoptosis was further carried out by determining the DNA laddering as a result of DNA fragmentation, indicative of the late stage of apoptosis. DNA isolated from HT – 29 cells treated with PT extract showed evident fragmented DNA in a dose – dependent manner, when compared with control cells (Figure 3).

Extract of hydroalcoholic pt tuber inhibited cell cycle progression

The effect of the hydroalcoholic extract was studied on the cell cycle phases of the studied HT - 29 cells. After 24h of incubation, stability in all the cell cycle populations is generally noticed and compared with the control cell line without treatment. Consistent with its effect on cell growth inhibition and induction of apoptosis, PT extract induced cell cycle arrest significantly arrested the cell growth at G2/M phase. PT extract treatment increased the cells in the sub G0/G1 phase from 0.18% to 22.59% in 64µg treated cells and 33.16% in 128µg treated cells respectively (Figure 4). An increased cell population in the sub G0/G1 phase with a concomitant decrease in the G1 and S phase compared to the untreated cells suggest that the PT tuber extract inhibited the cell cycle progression in G2/M phase and subjected the cells to apoptosis which is evident from cell accumulation in sub G0/G1 phase.

Alteration in Bax/Bcl 2 ratio, BID and activation of Caspase 9 and effector Caspases 3 and 8

Further the mechanism by which PT hydroalcoholic extract induced apoptosis was examined. Figure 5 demonstrates the protein analysis of Bax, Bid, Bcl2, Bid, Caspase 3, Caspase 8 and Caspase 9, as assessed by western blotting and its corresponding densitometric analysis. It is evident from the protein expression studies that PT extract increased the expression of proapoptotic...
factors Bax, Bid, Caspase 3, Caspase 8 and Caspase 9 and a subsequent decrease in anti-apoptotic protein Bcl2 was observed dose-dependently (Figure 5).

Uncontrolled cellular growth, as a consequence of defects in the cell cycle and apoptotic machinery, is responsible for the development of most cancers. Therefore, agents that can regulate the cell cycle and apoptosis may be useful in the management and therapy of tumors (McDonald et al., 2000; Owa et al., 2001, Kotha Anilkumar et al., 2017). Natural products exerting diverse bioactivities and possessing unique structural properties are important sources for the development of novel anti-tumor drugs (Newman DJ et al., 2000). The present study aimed to determine the ability of *P. tuberosa* hydroalcoholic extract to induce apoptosis and to identify the related biochemical mechanisms in a human colon cancer HT-29 cell line.

**Figure 1** Anticancer activity of hydroalcoholic PT extract - MTT assay

Values are expressed as Mean ± SEM (n = 3)

**Figure 2** Acridine Orange / Ethidium Bromide staining of HT-29 cells

Morphological changes in the nuclei following hydroalcoholic PT extract induced apoptosis detected by EtBr/AO staining. A: Negative control (without treatment). B & C: Treatment with hydroalcoholic PT extract 64µg and 128µg respectively for 24h. D depicts representative bar graph of percentage number of apoptotic cells counted in three different fields.
**Figure 3** DNA fragmentation assay of HT-29 cells treated with hydroalcoholic PT extract

The isolated DNA was loaded into a well on a 2% agarose gel and electrophoresed. M refers to the 100 bp DNA marker.

**Figure 4** HT-29 cells DNA fragmentation during apoptosis by flow cytometry treated with hydroalcoholic PT extract

Control Cell, B & C. PT extract treatment increased the cells in the sub G0/G1 phase from 0.18% to 22.59% in 64µg treated cells and 33.16% in 128µg treated cells respectively.
Figure 5 Effect of hydroalcoholic PT extract induced apoptosis in Caspase dependent pathway markers by immunoblotting analysis

PT extract increased the expression of proapoptotic factors Bax, Bid, Caspase 3, Caspase 8 and Caspase 9 and a subsequent decrease in anti-apoptotic protein Bcl2

The present study showed that PT hydroalcoholic extract exhibited strong anticancer activity on HT–29 human colon cancer cells via inhibition of proliferation and induction of apoptosis, which was dose-dependent with an IC50 value of 63.91µg/ml. The induction of apoptosis was confirmed by Etbr/AO staining procedure which showed that apoptosis percentage for 64µg/ml and 128µg/ml of drug was drastically increased (p< 0.001) to 37% and 66% respectively. Also PT hydroalcoholic treatment resulted in apoptosis, as evidenced by specific shearing of DNA, which is considered to be the hallmark of apoptosis. Cell cycle arrest at G2/M phase with subsequent accumulation of cells at subG0 phase further confirms the apoptotic potential of PT hydroalcoholic tuber extract. Puerarin was also found to inhibit proliferation and induces apoptosis in human glioblastoma cell lines (Yang et al., 2015).

Apoptosis is initiated by a cascade of proteases known as caspases and these have become potentially attractive targets for the development of new cancer therapies. Two pathways of caspase activation have been described. The first one is mediated by death receptors, controlled by caspases 8/10 and the second one is the mitochondrial mediated pathway and one of the primary regulators of this pathway to apoptosis is the family of Bcl-2 proteins (Green and Reed 1998). In addition, caspase-8 activated by apoptotic stimuli, converts Bid to truncated Bid (tBid), leading to conformational changes in Bax, mitochondrial depolarization, and cytochrome c release from mitochondria. This leads finally to the activation of caspase-3 and induction of apoptosis via a complex of apoptotic protease activating factor-1 (Apaf-1), pro-caspase-9, and cytochrome c after translocation of tBid to the mitochondria (S.
Fulda et al., 2006). Thus, translocation of pro-apoptotic Bax proteins from cytosol to mitochondria represents a key event for the activation of apoptosis. To clarify the mechanisms of induction of apoptosis by PT hydroalcoholic extract, we investigated the expression of apoptosis-inhibiting and apoptosis-promoting proteins in HT-29 cells. In the present study, quercetin induced a strong expression of Bax and tBid proteins, on the other hand, the level of expressions of Bcl-2 decreased in a dose dependent manner. The findings of the present investigation were found to be similar to the study conducted by Gan and Yin, (2015).

Death receptors (DRs) of the tumor necrosis factor receptor super family contain a conserved “death domain” in their intracellular region (Locksley et al., 2001). Fas associates with adaptor molecules, Fas Associated protein with Death Domain (FADD), through homotypic death domain interaction (Chinnaiyan et al., 1995). FADD then recruits caspase-8 through homotypic interactions of death effector domains (DEDs), leading to caspase-8 activation and induce apoptosis (Boldin et al., 1996). In the present study caspase-8 expression was significantly increased in both 64µg/ml and 128µg/ml treatment respectively. Therefore, PT hydroalcoholic extract induces the activation of extrinsic pathway which is confirmed by the expression of caspase-8. The PT extract induced apoptosis is a kind of caspase-3 DNA fragmentation. Caspase-3 has been shown to play important role in chemotherapy (Ibrado et al., 1996). Activation of caspase-3 appears to be a critical event in the execution of neuronal apoptosis (Honarpour et al., 2000). In the intrinsic pathway an apoptosis some complex is formed by Apaf-1, caspase-9, and cytochrome C as a result caspase-9 is activated which in turn activates caspase-3 dependent and caspase-3 activation is an upstream of The data further confirms that extrinsic pathways have contributed to PT hydroalcoholic extract induced apoptosis of HT-29 cells, as evident by increased expression of fragmented caspase-3, and -9 which was caused by caspase-dependent cell death and antitumor inhibition activity for various types of human cancer cells was considered (Yukiko et al., 2018).

Therefore the results in the present study concludes that PT hydroalcoholic extract possess potent anticancer property that induces apoptosis by activating caspase cascade pathway on human colon cancer HT-29 cell line. Thus it may be concluded that PT hydroalcoholic extract acts as a potential drug to induce apoptosis through the activation of caspases and other apoptotic related proteins.

References


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