Effects of Chinaberry Tree Extract on Proliferation, Apoptosis, Migration and Cell Cycle Distribution of Human Breast Cancer Cells
To study the effects of chinaberry tree extract on proliferation, apoptosis, migration and cell cycle distribution of human breast cancer cells, the MCF-7 breast cancer cells obtained were divided into blank group, low concentration group, medium concentration group and high concentration group. Phosphate buffered saline solution and sterile physiological saline were added to the blank group for cell culture. 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide and flow cytometry were used to detect cell proliferation and apoptosis at different time periods. Transwell assay was used to detect cell migration; flow cytometry was used to detect cell cycle distribution, and intercellular cell adhesion molecule-1, matrix metalloproteinase-2, matrix metalloproteinase-9 were detected by Western blotting. The proliferation rate of MCF-7 breast cancer cells in the low, medium and high concentration group was significantly lower than that in the blank group at different time periods, while the apoptosis rate was significantly higher than that in the blank group, and the effect was more obvious with the increase of concentration (p<0.05). The number of cell migration of MCF-7 breast cancer cells in the low, medium and high concentration group was significantly reduced compared with the blank group, and the cell migration was significantly reduced with the increase of the extract concentration of chinaberry tree (p<0.05). The higher the extract concentration of chinaberry tree, the lower the proportion of MCF-7 cells in G0/G1 and S cycle and the higher the proportion of MCF-7 cells in G2/M cycle (p<0.05). The expressions of intercellular cell adhesion molecule-1, matrix metalloproteinase-2 and matrix metalloproteinase-9 in the low, medium and high concentration groups were significantly lower than those in the blank group, and the expressions of intercellular cell adhesion molecule-1, matrix metalloproteinase-2 and matrix metalloproteinase-9 in the high concentration group were significantly lower than those in the low and medium concentration groups (p<0.05). The extract of chinaberry tree can effectively inhibit the proliferation and migration promote the apoptosis and reduce the growth of breast cancer cells by changing the periodic distribution of cancer cells, and its ability is concentration dependent.

Key words: Extract of chinaberry tree, human breast cancer cells, cell cycle, matrix metalloproteinase
these were fetched out. The temperature 40° was maintained with the help of a digital water bath (BH series precision). Then 2 ml of medium (10 % PBS, 1 % of 500 ml double-antibody mycillin and RPMI-1640 medium) was added followed by centrifugation at 1000 rpm for 5 min. The supernatant was discarded. Cells were resuspended in complete medium before they were transferred to culture flasks for sub culturing with 5 ml of the medium. These cells were placed in a CO₂ incubator (37°C, saturated humidity 95:5 % CO₂) and on the next day, as the cellular fusion rate was up to 80-90 % the cells were treated for continuous passage. MCF-7 human breast cancer cells used were divided into the control group, low concentration group, moderate concentration group and high concentration group. Twenty four hours after the inoculation, the medium was replaced by serum-free medium accompanied with the addition of Szechwan Chinaberry extracts for pretreatment for 24 h. Later, the cells were being starved for another 24 h before the medium was replaced by serum-free media. On daily basis, Szechwan Chinaberry extracts were applied followed by extraction of cellular proteins 48 h after. Within a total of 4 wells, 1 µl of PBS solution, and sterile saline solution, 25, 50 and 75 nM Szechwan Chinaberry extracts were placed respectively for cultivation.

The prepared cell suspension using the cultured cells above was added into wells of a 96-well plate, 90 µl each. The observation points for cells in the control, low, moderate and high concentration groups were at 12, 24, 36, 48, 60 and 72 h after the injection. Five replicate wells were set aside for each group to be added with different concentrations of Szechwan Chinaberry extracts. Cells of the control group were treated in each well with 10 µl of the culture medium and then cultivated for the same temporal periods as that in groups subjected to 3 different concentrations of extracts. Once the time was up, 20 µl of MTT was added into each well for 4-h incubation. Then, as the supernatant in wells was absorbed, 150 µl of DMSO was added followed by shaking for 10 s. The OD values of groups at 570 nm were measured in a microplate reader. The proliferation rates of MCF-7 human breast cancer cells were calculated using the following formula, % proliferation rate = (OD value/reference value-1)×100.

Cells of the 4 groups were placed into a 5-well plate for cultivation at 37°C, CO₂ concentration of 5 % with saturated humidity, within 12, 24, 36, 48, 60 and 72 h respectively, before 0.25 % of trypsin solution was added for digestion. Later, the cells were collected by centrifugation at 2000 rpm for 5 min and washed with PBS buffer twice for 3 min each time. Once again, centrifugation at 2000 rpm was applied to separate cells which were collected and dissolved in 1 ml of PI staining solution, placed aside for 1 h in the dark at room temperature. Once being labeled with specific fluorescence, the cells surrounded by rapidly flowing sheath fluids emitted photons which may be detected by a flow cytometer in terms of the photon numbers. The apoptosis was in this way measured and the technique used was the TUNEL technique (Volker, Cancer letters 1949:1;125-131,2003).

Breast cancer cells in the logarithmic phase were treated into a cell density of 3×10⁵ cells/ml before they were resuspended in blood serum medium of 1 %. In each chamber 100 µl of the cell suspension prepared was injected, and the Transwell chambers was placed on the culture plate followed by the addition of serum-free medium, set aside for 30 min until the completion of matrigel rehydration. After 24 h of cultivation with the cell suspension prepared and blood serum medium in saturated humid environment, the cells in chambers were washed with PBS buffer and fixed by paraformaldehyde and then subjected to crystal violet staining for 15 min in each group. Once again, PBS buffer was used to rinse the cells 3 times. Finally, 5 fields randomly selected were photographed under the microscope to count the number of cells, which shall be repeated 3 times in each group.

Seventy two hours after the cultivation in various concentrations of Szechwan Chinaberry extracts, MCF-7 cells were collected and 1 ml of 1 % ethanol solution was added for immobilization at 4° overnight. PI (50 µg/ml) was used to stain, before the incubation in dark for 0.5 h at 4°. The distribution of cell cycles was detected using the flow cytometer.

The collected samples were rinsed 3 times with PBS buffer, which was then separated out, and applied with IP cell lyse for 35 min before the total proteins were extracted and determined by biocinonic acid assay (BCA method). At a density of 20 µg/well, the collected proteins that have been treated with concentrated SDS-PAGE sample loading buffer for 15 min were subjected to SDS-PAGE gel electrophoresis at 100 V, 10 min. Later, the membrane was soaked in 10 % milk and sealed on the shaking table at 37°, 1.5 h. TBST was used to dilute the primary antibody β-actin (internal...
control) at a ratio of 1:1000. In combination with the primary antibody, the membrane was incubated at 4°C overnight and stored. On the next day, it was rinsed with a TBST buffer and treated with the second antibody for incubation within 1 h, followed by repeated rinsing in TBST buffer. At the end, following the instructions of developing and fixing kit, the developing agent was applied for substrate coloring.

SPSS21.0 statistical software was used for data analysis. Measurement data were expressed in mean±standard deviation (SD). In terms of comparison among multiple groups, F tests were adopted. As to pairwise comparison, independent t-test shall be used. P<0.05 was considered as statistical significance.

The proliferation rates of breast cancer cells in the low, moderate and high concentration group as detected at different temporal points were significantly lower than those in the control group, and the rates in the high concentration group inferior to those in the low concentration group, with the differences statistically significant (p<0.05), as shown in fig. 1.

The apoptosis rate of cells in the low, moderate and high concentration group as detected at different temporal points were obviously greater than those in the control group, in addition to the rates in the high concentration group superior to those in the low concentration group. The differences were statistically significant (p<0.05), as shown in fig. 2.

As it can be seen from fig. 3, the number of migrated cells in the low, moderate and high concentration group notably decreased as compared to that in the control group. Actually, the number of migrated cells decreased with the increase of concentration of Szechwan Chinaberry extracts.

Fig. 4 showed that the proportion of cells in G_0/G_1 and S period decreased significantly, while those in G_2/M period increased notably, with the increase of concentration of Szechwan Chinaberry extracts. The differences revealed statistical significance as well (p<0.05).

With the intervention of Szechwan Chinaberry extracts at different concentrations, the expressions of ICAM-1, MMP-2 and MMP-9 in cells of the 3 groups obviously decreased as compared to those in the control one (p<0.05), and the levels of the high concentration group were lower than that of the low or moderate concentration group (p<0.05), as shown in fig. 5.

At present, the therapy of breast cancer included operation, radiotherapy, chemotherapy, endocrinotherapy and targeted therapy[5,6], among which the operative treatments, although they are usually the preferred choice, may be denied by most female patients because this affects body aesthetics[7-9]. Treatment for cancers with plant extracts has been a hotspot in the medical field. Given the single monomeric composition of plant extracts, those anticancer components can be easily separated and purified, with respect to breast cancer, would inhibit cancer growth in a more scientific and accurate way[10,11]. In this investigation, MCF-7 human breast cancer cells were treated with Szechwan Chinaberry extracts, followed by detection of cell proliferation, apoptosis, migration and cyclic distribution using MTT assay, immunofluorescence, trans well assay and flow cytometry respectively, so as to determine the intervening effects of the extracts on these cells and their clinical applications.

Szechwan Chinaberry extracts contain triterpenoids with certain toxicity that at the earliest used in agricultural desensitization. They have been applied as an anthelmintic for more than 50 y actually[12-14]. Studies showed that Szechwan Chinaberry extracts exhibited biological activities in blocking neuromuscular junctions, antibotulism, and affecting efferent nerves, central nervous and the respiratory system[15-17]. Recently, several experts and scholars pointed out that Szechwan Chinaberry extracts have antineoplastic effects, including inhibition to hepatocellular carcinoma cells, human leukemia cells and human breast cancer cells, by primarily inducing the apoptosis of relevant cells[18-20]. In this paper, the Szechwan Chinaberry extracts were added at 25, 50 and 75 nM concentrations to MCF-7 cells. The cells in the control group were exposed to saline solution. After being
Fig. 2: Apoptosis of breast cancer cells
Apoptosis of breast cancer cells at different time points under different concentrations of Szechwan Chinaberry extracts. (▬●▬) Control group, (▬●▬) low dose group, (▬●▬) middle dose group, (▬●▬) high dose group.

Fig. 3: Migration of breast cancer cells
The migration of breast cancer cells under different concentrations of Szechwan Chinaberry extracts. Methylrosanilinium chloride solution×100, (A) control group, (B) low dose group, (C) middle dose group, (D) high dose group.
cultivated for a certain period at 6 temporal points, these cells were detected in terms of proliferation and apoptosis using MTT and immunofluorescence assays. As the results showed, the proliferation rate of breast cancer cells decreased with time. The falling range in the control group was not as apparent as good, but the proliferation rates in the other groups exposed to Szechwan Chinaberry extracts have significantly decreased, indicating the Szechwan Chinaberry extracts exerted inhibitory effect on breast cancer cells. Besides, this was supported by the results of apoptosis experiment where the apoptotic rates of cells treated with the Szechwan Chinaberry extracts at 3 different concentrations were obviously increased. It also found that the inhibiting forces imposed on cells applied with the high concentration of Szechwan Chinaberry extracts were much more powerful than that in the low or moderate concentration group, in addition to the notably increased apoptotic rates. These suggested the Szechwan Chinaberry extracts inhibit breast cancer cells in a concentration-dependent way. The higher the concentration, the more obvious is the inhibition.

Transwell assays were applied to determine the migration of breast cancer cells. As the results showed, different concentrations of extracts produced varied migration effects. The maximum of migrated cells was observed in the control group, while the minimum found in the high concentration group, indicating some blocking effects on the migration of breast cancer cells by the Szechwan Chinaberry extracts. In terms of cyclic distribution, with the intervention of Szechwan Chinaberry extracts, the proportion of cells in $G_0/G_1$ and $S$ period markedly decreased and the proportion of cells in $G_2/M$ period increased. These results revealed that the inhibition of Szechwan Chinaberry extracts on breast cancer cells may also be realized by altering the distribution of cell cycles.

Studies have pointed out that tumor metastasis, by which the tumor cells break away from their own cell population to a new organ where they form new tumors, has made the planned therapeutic regime more difficult to implement. Control of metastasis therefore has been an effective method for cancer treatment\cite{21}. ICAM-1, the member of immunoglobulin superfamily, is one kind of important surface adhesion molecules. They were rarely expressed or even never expressed in normal tissues, unlike the over-expression of them in malignant tumors, which may indicate that ICAM-1 plays some important roles in the metastasis of breast cancer cells\cite{22-24}. MMPs are membrane-binding proteins that degrade extracellular matrix and show vital functions in promoting cancer metastasis\cite{25-27}. As some experts revealed, MMP-2 and MMP-9 function well in invasion and metastasis of breast cancer cells. During any metastasis, the breast cancer cells must go through the basal membrane in the vascular walls, which means that an elevated expression level of MMP-2 and MMP-9 indicates certain diffusion and metastasis of tumor cells\cite{28-30}. In this paper, it was found the level of...
expressions of ICAM-1, MMP-2 and MMP-9 in breast cancer cells treated with Szechwan Chinaberry extracts significantly decreased. This told that the inhibition of Szechwan Chinaberry extracts to metastasis of breast cancer cells may be realized by regulation of ICAM-1, MMP-2 and MMP-9 expressions.

In conclusion, the Szechwan Chinaberry extracts effectively inhibited the proliferation and migration of breast cancer cells in a concentration-dependent manner by changing the distribution of cell cycles, in addition promoted the apoptosis of tumor cells. The research in this paper shall provide vital theoretical basis for future treatment of breast cancers.

**Conflict of interest:**

No conflict of interest between any of the authors.

**REFERENCES**


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