Christia vespertilionis extract induced antiproliferation and apoptosis in breast cancer (MCF7) cells

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Abstract

Background C. vespertilionis extracts were evaluated for antiproliferative and apoptosis effect on breast cancer (MCF7) cells.

Methods and results The leaves extracts were analysed for its antiproliferative effect on breast cancer (MCF7) cells and normal epithelial breast (MCF 10A) cells using Sulforhodamine B (SRB) assay. The selective extract was evaluated for its ability to induce apoptosis using Annexin V-FITC apoptosis staining and the expression of molecular genes using qualitative reverse transcription-polymerase chain reaction (RT-PCR) against MCF7 cells. Gas chromatography–mass spectrometry (GC–MS) was used to identify the compounds from the selective extract. The findings showed that dichloromethane fraction (CV-Dcm) extract had high antiproliferative effect against MCF7 cells (IC50 = 24 µg/mL, selective index (SI) = 8.17). The percentages of apoptosis cells in CV-Dcm-treated MCF7 cells was 58.8%. The CV-Dcm extract induced downregulation of PCNA level. The apoptotic genes were also triggered in both extrinsic and intrinsic signaling pathways, affecting a 1.5-fold increase in BAX, 1.4-fold increase in cytochrome c, 1.3-fold increase in caspase-8, 1.7-fold increase in caspase-3 and 0.5-fold-decrease in BCL-2. Treated MCF7 cells also activated P53-dependent apoptotic death pathway.

Conclusions The present work strongly suggests that high efficacy of CV-Dcm extract was attributed to its antiproliferative and apoptosis-inducing activation in MCF7 cells, most likely due to its favourable compounds.

Keywords Christia vespertilionis · Antiproliferative · Apoptosis · Antioxidant assays · Gene expression · Gas chromatography–mass spectrometry

Introduction

Cancer is caused by rapid proliferation and development of abnormal cells [1]. Global Cancer Incidence, Mortality and Prevalence (GLOBOCAN) (2020) reported that women breast cancer (11.7%) surpassed lung cancer (11.4%) as the most commonly diagnosed cancer [2]. Breast cancer is the most vulnerable among women in Malaysia, with 1 in 19 women are at risk [3]. Krauss and Stickeler [4] estimated that 75.0% of breast cancer cases have expressed oestrogen receptor as this hormone contributes to uncontrolled growth of breast cells. According to Holliday and Speir [5], breast cancer cell line known as Michigan Cancer Foundation-7 (MCF7) is an ideal model to evaluate hormone response studies.

Christia vespertilionis (L.f.) Bakh. F, which is a member of the Fabaceae family. This plant is used to enhance blood circulation and treat snake bites, bone fractures, tuberculosis, colds and bronchitis. This plant is widely distributed in Southeast Asia and is native to China, Vietnam, Indonesia, Malaysia, Cambodia and Thailand [6]. It is also locally referred as “Butterfly wing” or “Daun Rerama” because of its uniquely shaped trifoliate leaf [7]. Lee et al. [7] showed that C. vespertilionis extracts were effective against MCF7 cells. However, there is a knowledge gap to elucidate the...
antiproliferative and apoptosis molecular mechanism against MCF7 cells. Thus, the objective of this study was to investigate the antiproliferative and apoptosis effect against treated MCF7 cells. Additionally, the gene expression of treated MCF7 cells was also validated through qualitative reverse transcription-polymerase chain reaction (RT-PCR). In addition, the compounds from the selective extract were identified using gas chromatography–mass spectrometry (GC–MS).

**Methodology**

**Plant collection and identification**

*C. vespertilionis* was obtained from Guar Perahu Herbal Valley, Penang, Malaysia (5.426404°N, 100.478224°E). This plant was identified using universal DNA barcode primers (*rbcL, matK, and trnH-psbA*) as described by Ismail et al. [8]. The identified DNA sequences have been submitted to National Center for Biotechnology Information (NCBI)’s GenBank. The accession numbers of *C. vespertilionis* have been retrieved from NCBI’s GenBank (MK138830 to MK138849). This plant was authenticated by School of Biological Sciences, Universiti Sains Malaysia and the voucher collection was 11,777 (Fig. S1).

**Preparation of extracts**

The leaves extract was prepared according to the method by Mohd Fisall et al. [9]. The following analytical grade solvents for plant extraction have been obtained from Qrec (Asia) Sdn. Bhd, Malaysia: Dichloromethane, methanol, n-hexane, chloroform, and n-butanol. The fresh leaves (50 g) were soaked and rinsed with distilled water. The leaves were air dried for five days at room temperature (25–30 °C). The crude extract was extracted using 80% methanol by maceration method using orbital shaker (Buch & Holm, Denmark) at 170 rpm. After 24 h, the crude methanolic extract was filtered using Whatman filter paper No. 1 (Sigma-Aldrich, United States) and the filtrate was extracted again with the same solvent. The maceration process was repeated for three days. The rotary evaporator (Buchi, United States) was used to evaporate the methanol from the crude extract. Then, the crude extract was freeze-dried using freeze dryer (Martin Christ, Germany).

For fraction extract, a portion of the crude extract (10 g) was dissolved in 100 mL of distilled water. The dissolved crude extract was transferred to a separatory funnel and subsequently fractionated using n-hexane, dichloromethane, chloroform, and n-butanol solvents based on the solvent’s polarity. The fractionation for each solvent was repeated three times. The solvents from different polarity fraction extracts were evaporated using a rotary evaporator. The crude and fraction extracts were preserved at -20 °C for further used.

**Cell culture**

The breast cancer, MCF7 cells and normal epithelial breast (MCF 10A) cells were purchased from American Type Culture Collection (ATCC), United States. The cell culture complete media for MCF7 cells and MCF 10A cells were purchased from Nacalai Tesque, Japan. The MCF7 cells were cultivated in the Roswell Park Memorial Institute culture medium (RPMI-1640) growth medium supplemented with 10% fetal bovine serum (FBS) and 1% of penicillin–streptomycin (PenStrep). The MCF 10A cells were cultivated in Dulbecco’s modified Eagle medium (DMEM) growth medium supplemented with 20 ng/mL epidermal growth factor (EGF), 0.5 mg/mL hydrocortisone, 5% horse serum, 10 µg/mL insulin and 1% Penstrep.

The cells were incubated in the humidified 5% carbon dioxide (CO2) incubator (Nuaire, United States) at 37 °C. When the cells were ready to be subcultured, the old media was discarded, and the attached cells were rinsed with 4 mL of phosphate buffered saline (PBS) (Amresco, United States). The 200 µL of 0.25% trypsin–EDTA (GIBCO, United States) was added to the cells. After 5 min, the flask was gently tapped a few times to enhance the trypsinisation process. In order to deactivate the trypsin, 3 mL of complete media were added to the cells. The cell suspension was transferred to a 15 mL tube and the tube was centrifuged at 300 × g for 5 min. The supernatant was decanted, and the cells pellet were resuspended with new complete media.

**Antiproliferative assay**

The effect of *C. vespertilionis* leaves extracts on antiproliferative activity of MCF7 and MCF 10A were assessed using Sulforhodamine B (SRB) assay, as described by Skehan et al. [10] with minor modifications. In each 96-well plates, the MCF7 and MCF 10A cells (1 x 104 cells/mL) were seeded and incubated in a humidified 5% CO2 incubator for 24 h. Dimethyl sulfoxide (DMSO) was used to dissolve *C. vespertilionis* extracts. Then, *C. vespertilionis* extracts were serially diluted into six concentration (0 to 200 µg/mL). The 0.1% DMSO was used as working solutions of *C. vespertilionis* extracts to ensure that the concentration of DMSO in the cell culture was within the acceptable limit (0.1–0.5%).
The 0.1% DMSO and Tamoxifen (Nacalai Tesque, Japan) were used as a negative and positive control, respectively. The treated cells were incubated in 96-well plates in the humidified 5% CO₂ incubator at 37 °C for 72 h.

After 72 h, both cells were fixed with 50 µL of 50% cold trichloroacetic acid (fixed solution) for 30 min at room temperature. Then, the fixed solution was decanted from each well and the plate was washed with distilled water for four times. The cells in each well were stained with 100 μL of 0.4% SRB in 1% acetic acid for 30 min. The plate was washed again with 1% acetic acid for four times. The 100 µL of 10 mM Tris buffer was transferred to each well and the plate was shaken for 5 min. The cell viability was measured using PowerWave microplate spectrophotometer (Biotek, United States) at 570 nm. The half-maximal inhibitory concentration (IC₅₀) was determined from previous study by Bendale et al. [12]:

\[
\text{Cell viability (\%)} = \frac{\text{Absorbance (sample – blank)}}{\text{Absorbance (control – blank)}} \times 100\%
\]

**Selective index (SI)**

The SI values were calculated using Abd Razak et al. [13]'s equation: 

\[
\text{SI} = \frac{\text{IC}_{50} \text{ of normal cells (MCF 10A)}}{\text{IC}_{50} \text{ of cancer cells (MCF7)}}
\]

The highest SI value indicates that the extract was less toxic to normal cells. The selective extract was further analysed for the antiproliferative activity of MCF7 cells at different incubation times (24, 48, and 72 h).

**Cell morphology**

The MCF7 cells (7.5 × 10⁴ cells/mL) were allowed to grow in 6-well plate. After 24 h, the media was discarded and replaced with complete media mixed with 0.1% DMSO (negative control) and IC₅₀ of selected extract. The cells were incubated in humidified 5% CO₂ incubator. After 72 h, the media was decanted and the cells were stained with 2 mL acridine orange/propidium iodide (AO/PI) dye (10 µg/mL), which was mixed with the complete media, and incubated for 15 min. The treated MCF7 cells were observed using fluorescence microscope (Olympus, United States) at 40× magnification.

**Apoptosis analysis**

The MCF7 cells (2 × 10⁵ cells/mL) were allowed to grow in the 25 cm² cell culture flasks for 24 h. The untreated control cells were incubated with 0.1% DMSO and treated cells were incubated with the IC₅₀ of selected extract. The cells were stained using Annexin V-FITC Kit (Miltenyi Biotec, Germany) and the cells were quantified using MoFlo XDP Cell Sorter (Beckman Coulter, United States). The results were analysed in graph plots and percentages of different cell stages.

**Gene expression assessment by real-time PCR**

The MCF7 cells were treated as mentioned in apoptosis analysis method. The ribonucleic acid (RNA) of MCF7 cells was extracted using RNeasy Mini Kit (Qiagen, Malaysia). The purity of the extracted RNA was measured using NanoDrop 2000 UV–Vis Spectrophotometer (Thermo Fisher Scientific, United States). The Tetro™ cDNA Synthesis Kit (Bio line, United Kingdom) has been used to synthesise the complementary DNA (cDNA) according to the manufacturer’s instruction. The SensiFAST™ SYBR® Hi-ROX Kit (Bio line, United Kingdom) was used to determine the expression of proliferating cell nuclear antigen (PCNA), P53, BAX, BCL-2, caspase-8 and caspase-3 genes in MCF7 cells. The β-actin gene was used as an internal control. The gene sequences retrieved from De et al. [14], Devarajan et al. [15], Ferreira and Cronjé [16], Ngoc et al. [17], Priyadarsini et al. [18] were presented in Table S1.

The final volume of each reaction mixture was 10 µL, comprised of 5µL of 2× SensiFAST SYBR® Hi-ROX Mix, 1 µL of synthesized cDNA (10 ng/µL), 1 µL of forward and reverse primers of each gene (10 µM) and 3.2 µL of sterile distilled water. The reaction for each gene was biologically repeated three times. The StepOne Plus™ Real-Time PCR System instrument (Applied BioSystems, United States) was set up to run under following program: an initial denaturation step at 95 °C for 2 min followed by 40 cycles at 95 °C for 5 s, 60 °C for 5 s, and 72 °C for 20 s. The gene expression was measured using 2⁻ΔΔCT formula to obtain fold change of targeted genes between the untreated and treated MCF7 cells.

**Statistical analysis**

The analyses were performed triplicate and Statistical Package for the Social Sciences (SPSS) version 24 was used to carry out the analyses of variance (ANOVA) Tukey’s multiple comparison test and student’s paired t-test. The p < 0.05 were considered as significant difference.
The 1 mg/mL of selected extract was dissolved with high-performance liquid chromatography (HPLC)-grade methanol (Merck, Malaysia). The compound extract was identified using an Agilent gas chromatograph model 6890 equipped with an Agilent 19090S-433 capillary column, HP-5MS (0.25 mm × 30 m × 0.25 µm) (California, United States). The carrier gas, helium was used at 1.0 mL per minute. The oven was set to the following temperature: 70 °C was retained for 2 min, then it was elevated to 280 °C at the rate of 20 °C per min for 20 min. The GC–MS analysis was run for 32.5 min. The compounds of selected extract were identified by comparing the similarity index of at least 80% with the compounds from National Institute of Standards and Technology (NIST) library.

Results

Extraction yield

Based on Fig. 1, the percentage yield of crude methanolic (CV-Crd) extract was 11.5%. For fraction extract, the aqueous residue fraction (CV-Aqu) extract was the best at achieving higher extraction yields of 43.8%. The n-butanol fraction (CV-But) extract had the second highest percentage yields of 35.1%. Besides that, the dichloromethane fraction (CV-Dcm) extract also had relatively high extraction yield of 32.5%. The extraction yield value for the n-hexane fraction (CV-Hex) extract was much lower at 13.4%. However, the lowest extraction yield was chloroform fraction (CV-Chl) extract with a value of 4.7%. The order of the extraction yield of *C. vespertilionis* extracts were as follow; CV-Aqu > CV-But > CV-Dcm > CV-Hex > CV-Crd > CV-Chl. The results showed that the extraction yield of different solvents differed significantly.

Antiproliferative activity

The extracts were analysed on MCF7 and MCF 10A cells for their antiproliferative effect. Figure 2 (a) and Fig. 2 (b) reveal the effects of crude and fraction extracts at various concentration (6.25 to 200 µg/mL) on cell viability of MCF7 and MCF 10A cells. The findings showed that CV-Dcm had the highest inhibitory effects against MCF7, where the extract inhibited MCF7 cells at the lowest IC\textsubscript{50} value of 24 µg/mL followed by CV-Chl extract with IC\textsubscript{50} of 74 µg/mL, CV-Aqu extract (IC\textsubscript{50} = 114 µg/mL) and CV-But extract (IC\textsubscript{50} = 190 µg/mL).

*C. vespertilionis* extract were evaluated for the cell viability of MCF 10A cells (Fig. 2 (b)). The CV-Dcm extract inhibited the proliferation of MCF 10A cells with IC\textsubscript{50} value of 196 µg/mL and had the highest SI value of 8.17 (Fig. 2 (c)). The order of SI values were as follow: CV-Dcm > CV-Hex > CV-Chl > CV-Aqu > CV-Crd > CV-But. In Fig. 2 (c), Tamoxifen inhibited the proliferation of MCF7 cells with IC\textsubscript{50} of 6.6 µg/mL. According to the results obtained, CV-Dcm selectively inhibited MCF7 cells proliferation while being less toxic to MCF 10A cells. Figure 2 (e) illustrated the antiproliferative activity of MCF7 treated with CV-Dcm extract at 24, 48 and 72 h. The antiproliferative effect of CV-Dcm-treated MCF7 cells occurred in a dose and time-dependent manner, where CV-Dcm extract inhibited proliferation of MCF7 cells at IC\textsubscript{50} value of 80 µg/mL, 68 µg/mL and 24 µg/mL at 24 h, 48 h and 72 h, respectively.
Apoptosis effect

The morphological changes of MCF7 cells treated with CV-Dcm extract was displayed in Fig. 3. The AO/PI staining showed typical apoptosis morphology features such as cell membrane blebbing and nuclear chromatin fragmentation. Annexin V-FITC and propidium iodide (PI) were used to analyse the cells condition in the treated MCF7 cells using flow cytometry. Figure 4 displays the apoptotic level of treated MCF7 cells after 72 h treatment. The rate of total apoptosis (early and late apoptosis) in treated MCF7 cells was 58.8 ± 2.3% at 72 h.

The RT-PCR of the selected gene expression further supports the antiproliferative and apoptosis effect of treated MCF7 cells. The fold change of the targeted genes was determined using the $2^{-\Delta\Delta CT}$ formula and normalised to housekeeping gene (β-actin). As shown in Fig. 4, the CV-Dcm-treated MCF7 cells had caused significant upregulation of $P53$ (1.4 ± 0.13), $BAX$ (1.5 ± 0.12), cytochrome c (1.4 ± 0.12), caspase-8 (1.3 ± 0.12) and caspase-3 (1.7 ± 0.12) and downregulation of $BCL-2$ (0.5 ± 0.13) compared to untreated control cells. Thus, this study exhibited that CV-Dcm extract significantly activated the intrinsic and extrinsic of apoptosis pathway. Interestingly, $PCNA$ level in treated MCF7 cells decreased by 0.63 ± 0.13 compared with untreated control MCF7 cells.

Phytochemistry screening

Five compounds were obtained from GC–MS analysis where the similarity indices of the compounds were more than 80% with known NIST library (Fig. S2). The GC–MS showed the presence of phytochemical compounds known as pentadecyl acrylate (28.9%), methyl palmitate (15.9%), cyclododecane (20.9%), methyl linolenate (23.7%) and phytol (10.7%) (Table 1).

Discussion

Non-polar to polar solvents were used as fraction solvents to remove and purify various components found in the crude methanolic extract. In this study, fraction solvents were consecutively fractionated from the crude methanolic extract due to the fact that fraction extracts may have high inhibition effects against cancer cells [19]. The extraction yields of C. vespertilionis extracts were in line with the study by Ismail et al. [20, 21], which showed the similar results with the percentage yield of Clinacanthus nutans extracts. CV-Aqu extract was best at achieving higher extraction than other solvents. The highest yields of CV-Aqu could be attributed to high impurities like organic acids, sugars and soluble proteins with polar compounds. These findings were in line with Anwar et al. [22], who reported that the highest extraction yield came from the aqueous extract from Brassica oleracea.

The CV-Dcm extract exhibited strongest IC50 value when treated with MCF7 cells. The finding was concurrent with a report from Sidek et al. [23] who revealed that the dichloromethane fraction extract of C. vespertilionis exhibited the strongest extract when treated with cervical cancer (HeLa) cells at 72 h. The IC50 of CV-Dcm extract also met the requirement of the National Cancer Institute, which requires the extracts to have an IC50 less than 30 µg/mL for preliminary assay [24]. Furthermore, CV-Dcm extract had the highest SI value in which the extract with SI value more than 2 was considered to have high selectivity [25].

It was found that the treated MCF7 cells induced apoptosis when phosphatidylserine (PS) of the cellular membrane is exposed to the Annexin V [26]. Hence, it is important to validate the expression of molecular mechanism of the CV-Dcm-treated MCF7 cells. The CV-Dcm extract upregulated the gene expressions of $P53$, $BAX$, cytochrome C, caspase-8 and caspase-3 and downregulate the gene expressions of $BCL-2$ and $PCNA$ in the MCF7 cells (Fig. 4 (d)). Their expressions give compelling evidence that these genes and their interaction play important mediating roles in the extract’s ability to induce apoptosis. The result showed that the increased level of $P53$ suggested that the extract could lead to apoptosis through extrinsic pathway [27].

Haupt et al. [28] stated that the extrinsic apoptotic pathway was found to be triggered by $P53$ expression by inducing transmembrane death receptor proteins such as Fas, DR5, PERP, and TRAIL, which then transmitted the apoptotic death signal from the cell surface to intracellular signalling paths. Additionally, $P53$ is a well-known gene target for anti-cancer development, in which it can exerts antiproliferative effects [29], DNA damage, mitosis failure [30], senescence and apoptosis [31, 32].

$P53$ tumour-suppressor interacts with $BAX$ or $BAK$ to trigger the intrinsic pathway [33]. The present study showed increased level of $BAX$ and decreased level of $BCL-2$ in treated MCF7 cells. This study validates that $BCL-2$ in treated MCF7 cells could not protect the integrity of mitochondrial membrane [34] and enhanced the level of $BAX$ which is necessary for the pores creation in the mitochondrial membrane, resulting in the cytochrome c released, followed by the caspase-3 activation. Importantly, caspase-8...
a

![Graph showing percentage cell viability against concentration for different extracts.]

b

![Graph showing percentage cell viability against concentration for different extracts.]

c

![Graph showing percentage cell viability against concentration for MCF7 and MCF 10A.]

d

| Type of extracts | IC$_{50}$ (µg/mL) MCF7 | IC$_{50}$ (µg/mL) MCF10A | Selective index |
|------------------|-------------------------|--------------------------|-----------------|
| CV-Crd           | 94                      | 134                      | 1.4             |
| CV-Hex           | 100                     | 190                      | 1.9             |
| CV-Dcm           | 24                      | 196                      | 8.2             |
| CV-Chl           | 74                      | 130                      | 1.8             |
| CV-But           | 190                     | 190                      | 1.0             |
| CV-Aqu           | 114                     | 198                      | 1.7             |
| Tamoxifen        | 6.6                     | > 10                     | -               |

e

![Graph showing percentage cell viability against concentration for different time points.]

[Springer]
can also regulates the expression of the extrinsic apoptotic pathway [35]. Notably, the significant PCNA downregulation explained the proliferative inhibition of MCF7 cells induced by CV-Dcm extract which has not been noted in any *C. vespertilionis* studies. PCNA level in treated MCF7 cells was approximately lower compared to untreated control MCF7 cells. The findings suggested that the decreased in PCNA expression could be responsible for the suppression in MCF7 cells growth. This study showed the remarkable effectiveness of the CV-Dcm extract on MCF7 cells due to activation of the apoptosis and antiproliferative effects.

The GC–MS analysis revealed five phytochemical compounds found in CV-Dcm extract. Previous studies revealed that pentadecyl acrylate has antioxidant activity [36]. Additionally, the had cytotoxicity activities on T-cell leukemia cell line [37]. Methyl palmitate may be the basis of the antiproliferative and apoptosis effect on treated MCF7 cells.

Cyclododecane is said to be antimicrobial and antitumour [38] and methyl linolenate is reported as anti-inflammatory, cancer preventive, hypocholesterolaemia and hepatoprotective [39]. Phytol has been shown to contain a variety of medicinal properties such as antitumoral and anticarcinogenic effects [40]. However, the chemical compounds may differ because most of the plants are largely influenced by climate variables such as humidity, water content, temperature, soil pH changes and nutrient content [41] that will also interfere with plant genetics, leading to genetic variations that can influence the content of plant materials [42].

**Conclusion**

In conclusion, the results showed that CV-Dcm suppressed MCF7 cells proliferation due to the downregulation of PCNA level. This extract also induced apoptosis against MCF7 cells through both intrinsic and extrinsic pathways and caused *P53*-dependent apoptotic deaths. Hence, the *C. vespertilionis* extract seems to be a potential source for anticancer agents. research.
Figure a: Scatter plots showing the distribution of Propidium Iodide and Annexin V-FITC levels in untreated and treated samples. The plots indicate the presence of apoptotic cells.

Figure b: Similar to Figure a, but with a different set of data points indicating a higher concentration of apoptotic cells in the treated group.

Figure c: Bar graph comparing the percentage of apoptotic cells between untreated and treated samples. The graph shows a significant increase in the percentage of apoptotic cells in the treated group.

Figure d: Bar graph comparing the relative fold change of different genes (P53, BAX, BCL-2, Cytochrome C, Caspase-8, Caspase-3, PCNA) in untreated and treated samples. The graph indicates a significant decrease in the expression of pro-apoptotic genes and an increase in anti-apoptotic genes in the treated group.
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Author contributions NZI and HA designed the experiment. NZI performed the experiments. NZI, IAA, WASM, NNMZ and HA analysed the data. NZI, WASM and IAA wrote the manuscript. NZI, IAA, WASM, NNMZ and HA have given the final shape of manuscript and all authors approved the manuscript.

Declarations

Conflict of interest The authors have declared that there is no conflict of interest.

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