Anticancer Property of Orthosiphon stamineus Benth. Extracts in Different Solvent Systems against T47D Human **Breast Cancer Cell Lines**

by Retno Widyowati

Submission date: 14-Apr-2023 03:45AM (UTC+0800)

Submission ID: 2063755009

File name: C-06_artikel.pdf (627.09K)

Word count: 5174

Character count: 28012

Anticancer Property of Orthosiphon stamineus Benth. Extracts in Different Solvent Systems against T47D **Human Breast Cancer Cell Lines**

Lusiana ARIFIANTI'D, Sukardiman SUKARDIMAN'D, Niken INDRIYANTI'D, Retno WIDYOWATI*****

Anticancer Property of Orthosiphon stamineus Benth. Extracts in Different Solvent Systems against T47D Human Breast Cancer Cell Lines

SUMMARY

Solvent system is an important factor in extraction process in order to obtain compounds that have pharmacological activity. The aim of this research is to develop a comprehensive extraction methods by modification of solvents used that might produce compounds possessing pharmacological activity for anticancer. In this study, Orthosiphon stamineus Benth. extract was used as sample in different solvent systems to observe their metabolite profiles. Extraction carried out using sonication techniques with ethanol solvents in three types of concentrations (96%, 70% and 50%). Then, the extracts introduced into anticancer activity profiles in order to find its active compounds. The anticancer activity had explored against breast cancer cells (T47D) using the MTT assay and doxorubicin as a positive control. The best IC50 value obtained from the 50% ethanol extract of Orthosiphon stamineus Benth. Based on the scanning chromatogram at 366 nm using Thin Layer Chromatography, each sample contains sinensetin and rosmarinic acid. The largest percentage of rosmarinic acid area was found on 70% ethanol extract of Orthosiphon stamineus Benth, while the highest percentage of sinensetin was found on 50% ethanol extract of Orthosiphon stamineus Benth. Thus, it can be concluded that sinensetin which has the most influence on anticancer activity.

Key Words: Orthosiphon stamineus, breast cancer, rosmarinic acid, sinensetin, T47D, anticancer

Orthosiphon stamineus Benth'in farklı Çözücü Ekstrelerinin T47D İnsan Meme Kanseri Hücre Hattına Karşı Antikanser Özelliği

ÖZ

Solvent sistemi, farmakolojik aktiviteye sahip bileşiklerin elde edilmesinde ekstraksiyon sürecinde önemli bir faktördür. Bu araştırmanın amacı, antikanser farmakolojik aktiviteye sabip bileşikler için çözücüleri modifiye ederek kapsamlı bir ekstraksiyon yöntemi geliştirmektir. Bu çalışmada Orthosiphon stamineus Benth. ekstraktı metabolit profillerini gözlemlemek için farklı solvent sistemlerinde numune olarak kullanılmıştır. Ekstraksiyon, üç farklı konsantrasyonda (% 96, % 70 ve % 50) etanol çözücülerle sonikasyon teknikleri kullanılarak gerçekleştirildi. Daha sonra ekstraktlar, aktif bileşiklerini bulmak için antikanser aktivite profillerine sokuldu. Antikanser aktivitesi, meme kanseri hücreleri (T47D) kullanılarak MTT analizi ile araştırılmıştır, pozitif kontrol olarak doksorubisin kullanılmıştır. Orthosiphon stamineus Benth'in elde edilen en iyi IC₅₀ değeri % 50 etanol ekstraktında bulunmuştur. İnce Tabaka Kromatografisi kullanılarak 366 nm'de elde edilen kromatogramına göre, her numune sinensetin ve rosmarinik asit içermektedir. Rosmarinik asit alanının en büyük yüzdesi Orthosiphon stamineus Benth'in % 70 etanol ekstresinde bulunurken, en yüksek sinensetin yüzdesi Orthosiphon stamineus Benth'in % 50 etanol ekstresinde bulunmuştur. Böylece antikanser aktivitesi üzerinde en fazla etkiye sahip olanın sinensetin olduğu sonucuna varılabilir.

Anahtar Kelimeler: Orthosiphon stamineus, meme kanseri, rosmarinik asit, sinensetin, T47D, antikanser

Received: 23.12.2019 Revised: 29.04.2020 Accepted: 02.05.2020

ORCID:0000-0003-1704-7908, Department of Pharmacognosy and Phytochemistry, Faculty of Pharmacy, Universitas Airlangga, Indonesia

[&]quot;ORCID:0000-0001-9689-5088, Department of Pharmacognosy and Phytochemistry, Faculty of Pharmacy, Universitas Airlangga, Indonesia "ORCID:0000-0003-3733-5018, Department of Pharmacology, Faculty of Pharmacy, Mulawarman University, Indonesia

[&]quot;" ORCID:0000-0003-0572-7551, Department of Pharmacognosy and Phytochemistry, Faculty of Pharmacy, Universitas Airlangga, Indonesia

INTRODUCTION

Recently, the efforts to treat cancer patients have focused on inhibiting the growth or killing cancer cells. The investigations to find an ideal drug that targets cancer cells with minimal side effects are ongoing. Some Indonesia medicinal plants may have a potential bioactive compound to be developed into an ideal drug for cancer. One of them is *Orthosiphon stamineus* Benth.

Orthosiphon stamineus Benth. (Lamiaceae) is a Indonesia plant native, traditionally used for diuretics, rheumatism, diabetes and hypertension (Fei et al., 2010). Previous study reported the activity of this plant extract inhibits human oral cancer cells (Younis et al., 2013). The methanol extract of this plant enhanced tamoxifen on breast cancer cell (MCF 7) proliferation (Sahib et al., 2009). In addition, 200 mg/kg ethanol extract of this plant showed no tumor cell growth compared to control group using xenograph method of tumor models that transplanted with breast cancer cells (MCF7) and colon cancer cells (HCT116) (Ahmad et al., 2010).

The bioactive compound will become lead compound to find an effective drug for cancer. One of the bioactive compounds found in this plant is sinensetin (5,6,6,7,3',4'-pentamethoxy flavone). It also contains orthosiphol D, orthosiphol E (Takeda et.al, 1993), orthosiphol A, orthosiphol B, 3'- hydroxy-5,6,7,4'-tetramethoxyflavone, neoorthosiphol A, neoorthosiphol B, α-amyrin, β-amyrin, maslinic acid, urosolic acid, orthosiphonone A, orthosiphonone B, myo-inositol, β-caryophyllene, caffeic acid, sinensetin, tetra-methyl scutellarein, eupatorin, cirsimaritin, acetovanillochromene, orthochromene A, methylripario chromene, agermacrene-D, β-selinen, α-cadinol, choline, betaine, O-cyamenea-terpineol, lyrol, valencene, nephthalin, camphor, α-elemene (Singh et al 2015), 5,6,7,8-tetra hydroxy-6-methoxy-flavones (Hossain et.al, 2008), potassium, flavonol glycosides, caffeic acid (rosmarinic acid) (Sumaryono, et al., 1991), essential oils, diterpenes, lipophilic flavones such as eupatorin, (6-hydroxy-5,7,4-trimethoxy flavone), and TMF (3'-hydroxy-5, 6, 7, 4 'tetrametoxy flavone) (Awale et.al, 2001), triterpenes such as betulinic, ursolic, oleanolic acids, β sitosterol (Tezuka et al., 2000) and flavonoids such as 5-hydroxy-6,7,30,40-tetrametoxiflavone, salvigenin, 6-hydroxy-5,7,40-trimethoxyflavone, 5,6,7,30-tetramethoxy-40-hydroxy-8-Cprenylflavone (Hossain and Rahman, 2015). The chloroform extract of this plant contained 1.48% of sinensetin, 2.26% of eupatorin, and 0.58% of 30-hydroxy- 5,6,7,40-tetrametoksiflavon (Mohamed et al., 2013, Yam et al., 2012).

Previous studies showed that the 50% methanol

extract of this plant using freezed and sprayed dried methods contained protein, polysaccharides and saponins (Siddiqui *et al.*, 2009). Research on metabolite profiles in this plants had been carried out using chromatographic and spectroscopic techniques combined with chemometrics (Akowuah *et al.*, 2004; Sumaryono, *et al.*, 1991; Saidan *et al.*, 2015a). Ethanol extract using maceration method contained high phenolics and flavonoids, (rosmarinic acid and eupatorin) as antioxidants, while 50% ethanol and methanol extracts using soxhlet contained high protein and glycosaponin. Water extracts using reflux and maceration showed high polysaccharides (Saidan *et al.*, 2015b).

Solvent system is an important factor in extraction process in order to obtain compounds that have pharmacological activity. In this study, the effect of different solvents on the metabolites profile in each extract had been determined. Extractions with 96, 70 and 50% ethanol were carried out according to the previous study with different method (Arifianti *et al.*, 2014). Arifianti had extracted this plant using maceration method while this study used ultrasonic method to accelerate the extraction process and % yield with optimum results at same concentrations. It correlated to the anticancer activity (breast cancer) and their secondary metabolites are responsible for their activity.

MATERIAL AND METHODS

General Experimental Procedures and Materials

The plant was extracted on CAMAG ultrasonic and then evaporated by BUCHI rotary evaporator. The metabolite profiles of Orthosiphon stamineus Benth leaves were measured on a CAMAG Scanner 3 Densitometer and Linomat 5. The solvent extracts were combination between ethanol p.a (Merck) and aquadest. Sinensetin and rosmarinic acid from Sigma used as standard. The material used for breast cancer activity of T47D cells were RPMI 1640 medium (Gibco, Invitrogen), Fetal Bovine Serum (FBS, Sigma), Penicillin-Streptomycin (Sigma), Amphotericine B (Sigma), Dimethyl sulfoxide (DMSO, Sigma), Phosphate Buffer Saline (PBS, Sigma), 3-(4,5-dimethylthiazol-2-yl)2-5- diphenyl tetrazolium bromide (MTT, ThermoFisher) and sodium dodecylsulfate (Sigma). The cancer cells inhibition was determined with Robonik Elisa Reader.

Plant Materials

Orthosiphon stamineus Benth leaves were obtained on late April 2018 from Balai Materia Medika, Malang and voucher specimens were deposited in Department of Pharmacognosy and Phytochemistry, Faculty of Pharmacy, Universitas Airlangga.

Extraction of plant materials

Extraction was carried out by ultrasonic method according to the previous published method by modification (Juliana et al., 2019). The dried leaves of Orthosiphon stamineus Benth (50 g) were extracted with 250 mL of each 96%, 70% or 50% ethanol (3 x 10 minutes) using CAMAG ultrasonic. Then the extracts were separated by filtration. The residue was re-extracted by using same procedure (3 times repeated). The filtrates were evaporated by BUCHI rotary evaporator to dryness under vacuo to get 96%, 70% or 50% ethanol extracts. The extracts were used to examine bioassay activity and phytochemical analysis. It performed by thin layer chromatography (TLC) to observe the sinensetin and rosmarinic acid profile.

Phytochemical analysis of plant extracts using TLC

The 10 mL of each 96%, 70%, 50% ethanol extracts of Orthosiphon stamineus Benth, sinensetin and rosmarinic acid were applied on a pre-coated TLC plate, silica gel 60F 254 (10 cm x 20 cm) as 7 mm bands using a Camag automatic TLC sampler (Linomat 5) spray-on band applicator equipped with 100 mL syringe, and the space between two spots was 2 mm of the plate. The extracts were each applied duplicates on the plate. The TLC plates were developed with chloroform-ethyl acetate in a ratio of 6:4 as mobile phase. Then, they were identified using a UV lamp at 365 nm. The phytochemical were analyzed based on chromatogram pattern using a CAMAG TLC Scanner 3 Densitometer and winCATS software, using a deuterium light source, the slit dimension was 6.00 x 0.45 mm. Peak areas were recorded and the spot of sinensetin and rosmarinic acid in the sample were confirmed by comparing the RF and spectra of the spot with that of sinensetin and rosmarinic acid standard (Arifianti et al., 2014, Hossain and Ismail, 2016).

Cell line

The T47D Human breast cancer cell lines were obtained from the CCRC (Cancer Chemoprevention Research Center), Gajah Mada University, Indonesia and a modification method described by Fresney Method (Freshney, 2005). The T47D Human breast cancer cells were maintained in RPMI 1640 that contained 10% of FBS, 2% of Penicillin-Streptomycin and 1% of Amphotericine B. It was stored at 37°C with humidified atmosphere of 5% CO₂ (Eppendorf). The cells were routinely observed to keep them from contamination.

Measurement of inhibition of cancer cell by MTT method

The MTT method used was a method that has been modified by Freshney (Freshney, 2005). The 5 x 104 cells/wells of T47D cells with or without samples (96%, 70%, 50% ethanol extracts of Orthosiphon stamineus Benth) were cultured in RPMI 1640 medium that contained 10% of FBS and 1 % (v/v) of penicillin-streptomycin into 96 well plate then incubated for 24 hours at 37°C and 5% CO, (70-80% confluent). The samples were dissolved in DMSO and further diluted with medium to make series of concentrations (15 - 1,000 mg/mL). The final concentration of DMSO in the test solution should not more than 1%. Control cell was treated with 1% DMSO. Cells were then treated with a serial dilution of tested samples. The doxorubicin concentrations of 2.5-100 mg/ml were used as positive control. After 24 h incubation, 0.5 mg/ml of MTT was added to each well and incubated for 4 hours. Then, the stopper solution (sodium dodecylsulfate 10% in 0.1 N HCl) was added to dissolve the formazan crystal and incubated overnight at room temperature and dark. Finally, the cells viability was measured using ELISA reader at 1 570 nm. The absorbance of each well then converted into percentage of viable cells using calculation below and the IC, values were determined by Probit analysis using SPSS software. Experiments were done in triplicates.

 $\% cellviability = \frac{samp\ absorbance - medium\ control\ absorbance}{cell\ control\ absorbance - medium\ control\ absorbance} \times 100\%$

RESULT AND DISCUSSION

Extraction

Several ways can be do to obtain phytochemicals from plants, one of which is extraction. Extraction efficiency is influenced by the chemical properties of the compound, the extraction method used, the particle size of the sample, the solvent used, and the presence of disturbing substances (Stalikas, 2007). Conventional extraction techniques are often associated with long heating times and a risk of bioactive

compounds degradation. This has led to sophisticated techniques such as ultrasonic extraction which are efficient in terms of extraction time and solvent consumption. In view of this method, ultrasonic cavitation produces shock waves that are able to disrupt the external structure of plant samples and release plant bioactives effectively (Budynas & Nisbett, 2008; Floros & Liang, 1994).

Ultrasonic extraction using ultrasonic frequencies at >20 kHz can accelerate the contact time between

samples and solvents at room temperature. It causes the mass transfer of bioactive compounds from plant cells to solvent to be faster. Sonication relies on sound energy that causes the cavitation. It forms small bubbles due to ultrasonic frequency transmission to help the diffusion of solvents into plant cell walls (Ashley *et al.*, 2001).

General extraction parameters such as concentration and ratio of solvents using ultrasonic method were first optimized. The solvent used was ethanol because it is non-toxic, good polarity for the sound energy and ultrasonic frequencies, so it is able to dissolve interesting bioactive compounds (Xiao et al.,2008). The extraction yield depends on the solvent with various polarity, pH, temperature, extraction time, and sample composition. At the effect of the same extraction time and temperature, the solvent and sample composition are the most important parameters. In this study, Orthosiphon stamineus Benth. extracts were obtained using ethanol and water at various concentrations (50%, 70%, and 96%). Their extraction yields were ranged between 15.64%, 12.39% and 3.44%, respectively (Table 1). The results of extraction with various solvents decreased in the following order: 50% ethanol> 70% ethanol> 96% ethanol. It showed that the extraction yield increases with increasing polarity of the solvent used in the extraction. Increasing the concentration of water in the solvent can increase the extraction yield. Compounds other than sinensetin and rosmarinic acid in the extract, it may have been extracted and contributed to higher yields. This might be caused by higher solubility of protein and carbohydrate in water-ethanol than pure ethanol (Zieliński and Kozłowska, 2000). The use of a combination of water in organic solvents can facilitate the extraction of water-soluble chemicals and/or organic solvents. This might be the reason why the ethanol extract yield is 50% higher than other extracts. The results of this study are in accordance with the results of extraction in Limnophila aromatica (Do et al., 2014) and several medicinal plants (Sultana et al., 2009).

Table 1. % yield of Orthosiphon stamineus Benth extracts

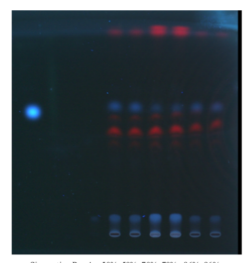
Sample	% yield
96% ethanol extract	3.444
70% ethanol extract	12.390
50% ethanol extract	15.636

The addition of water in the extraction solvent showed that the extraction yield is improved, because the presence of water increased heating efficiency due to its high dielectric constant (Sato & Buchner, 2004), and increased the permeability of plant matrices to encourage mass transfer and diffusion of bioactive compounds (Boeing *et al.*, 2014). The effects of aqueous ethanol have different effects, the optimum solvent concentration was found to be 50% (v/v) ethanol.

The phytochemical analysis

A number of sinensetin and rosmarinic acid found in the leaves of *Orthosiphon stamineus* Benth. TLC-densitometry is the current method for the quantization of some flavonoids and caffeic acid derivatives in pharmaceutical formulations. It is quickly gaining widespread acceptance in pharmaceutical analysis. This is due to simplicity, accuracy, cost effectiveness and possibility of simultaneous determination of a number of samples on a single TLC plate. HPTLC allows the identification and quantification of more than 20 samples in the same chromatographic process and requires more than 2 hours. Whereas TLC takes only 15-30 minutes because it does not require conditioning steps, such as in HPLC, and is cheaper.

A fingerprint chromatography was performed to describe components found in sinensetin & rosmarinic acid-rich extract using TLC densitometry according to the modified method of Hossain and Ismail (2016) as well as Hossain and Ismail (2009). Samples (96, 70 and 50% ethanol extracts of Orthosiphon stamineus Benth) and standard (sinensetin and rosmarine acid) were explored on a silica gel GF 254 and developed with chloroform-ethyl acetate in a ratio of 6:4. under UV observation at 366 nm by TLC visualizer (Fig. 1). The advantage of using the TLC-visualizer method is easy, fast, accurate, inexpensive and most suitable for natural material analysis. A sample chromatogram showed the presence of spots of the same color and at the same Rf value as the standard (Fig.1). A single peak at Rf 0.49 was observed in the chromatogram of sinensetin standard and Rf 0.06 as rosmarinic acid standard. At 70% ethanol extract had more intensity spot because it contained higest rosmarinic acid compare than other extracts.



Sinensetin Ros Ac. 50% 50% 70% 70% 96% 96%

Figure 1. The chromatograms of samples and standard without any spray reagent, in UV light at 366 nm

TLC analysis results showed bright blue fluorescent spots and the ultraviolet spectrophotometry showed the spectra images that were identical between samples with standards spectra. Two spectra are said to be identical if they have a MF (Match Factor) price > 95. In this study, the price of a match factor from the sample against sinensetin and rosmarinic acid standard obtained 0.99559 and 0.99985, respectively, so that they can be said to be identical (Fig. 2). The presence of sinensetin and rosmarinic acid in samples were proven by comparison of standards spectra with components that are separate from the samples UV-VIS spectra. Figure 2a showed the standard UV-VIS spectra of sinensetin (black) with samples, while in figure 2b was rosmarinic acid (pink) with samples. It can be observed the presence of sinensetin and rosmarinic acid peak in a sample at the same Rf value. It showed the similarity of spotting between sinensetin and rosmarinic acid standard in each sample. This data is supported by a standard spectrum profile of rosmarinic acid that has similarities with the spectrum of 96, 70 and 50% ethanol extracts of Orthosiphon stamineus Benth. Furthermore, a similar spectrum image was also obtained between sinensetin standard and them, but there was a slight shift in the sinensetin spectrum of 96% ethanol extract of Orthosiphon stamineus Benth (Fig.2).

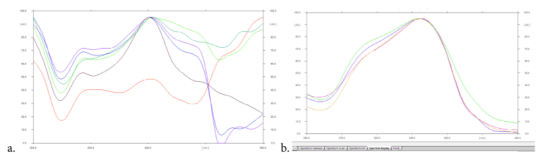


Figure 2. The UV-VIS Spectra overlay results of sinensetin (a, black) and rosmarinic acid (b, pink), together with samples of 50 % ethanol extract (blue), 70% ethanol extract (green) and 96% ethanol extract (yellow).

Quantitative determination was done by TLC-densitometry using the calibration curve method. The calibration curve was performed by the winCATS software program. In table 2, the percentage data of each chromatogram area refered to the sinensetin and rosmarinic acid standard. The largest percentage of ros-

marinic acid area was found on 70% ethanol extract (74.61±0.03), while the highest percentage of sinensetin was found on 50% ethanol extract (32.97±0.06) of *Orthosiphon stamineus* Benth. So 70% ethanol is the best solvent system for extracting rosmarinic acid while 50% ethanol for sinensetin.

Table 2. Peak	identification b	y TLC-densitometry
---------------	------------------	--------------------

Samples	Start position	Start Height	Max position	Max Height	Max %	End Position	End Height	Area	% Area	Mean % area
50% EtOH Ex_1	0.06 Rf	0.4 AU	0.11 Rf	102.3 AU	72.22%	0.16 Rf	0.0 AU	1330.3 AU	65.85%	67.04±0.02
	0.48 Rf	4.3 AU	0.55 Rf	39.4 AU	27.78%	0.60 Rf	2.9 AU	689.9 AU	34.15%	32.97±0.06
50% EtOH Ex_2	0.06 Rf	0.6 AU	0.11 Rf	103.4 AU	73.36%	0.14 Rf	1.2 AU	1338.7 AU	68.22%	
	0.49 Rf	3.3 AU	0.55 Rf	37.6 AU	26.64%	0.59 Rf	3.2 AU	623.7 AU	31.78%	
70% EtOH Ex_1	0.06 Rf	1.0 AU	0.12 Rf	192.9 AU	80.66%	0.18 Rf	0.0 AU	2774.9 AU	76.45%	74.61±0.03
	0.49Rf	7.0AU	0.56 Rf	46.3 AU	19.34%	0.60 Rf	2.8 AU	855.0 AU	23.55%	22.15±0.47
70% EtOH Ex_2	0.06 Rf	0.2 AU	0.12 Rf	191.2 AU	76.66%	0.15 Rf	1.2 AU	2821.9 AU	72.77%	
	0.50 Rf	5.9 AU	0.56 Rf	46.0 AU	18.45%	0.61 Rf	0.3 AU	804.1 AU	20.74%	
96% EtOH Ex_1	0.07Rf	0.0AU	0.11 Rf	73.6 AU	72.24%	0.15 Rf	0.1 AU	892.7AU	70.10%	71.37±0.02
	0.54Rf	6.3 AU	0.57 Rf	28.3 AU	27.76%	0.61 Rf	2.7 AU	380.8 AU	29.90%	28.63±0.02
96% EtOH Ex_2	0.07Rf	0.3AU	0.12 Rf	75.4 AU	74.20%	0.18 Rf	0.0 AU	942.1 AU	72.64%	
	0.56Rf	4.6AU	0.60Rf	26.2 AU	25.80%	0.64 Rf	0.4 AU	354.9 AU	27.36%	

Anticancer activity

The anticancer properties of the ethanol extracts of 96, 70 and 50% of *Orthosiphon stamineus* Benth. were determined by MTT test. This test was chosen because it is reliable, simple, applies to a variety of cells, and can be done in microtiter plates. The test was based on the reaction of colorimetry of 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide with the enzyme dehydrogenase in living cells to form a colored formazan corresponding to a viable cell numbers (McCauley et al., 2013).

Table 3. IC_{50} value of the samples on breast cancer activity

Sample	IC ₅₀ (ppm)
96% ethanol extract	259.016 ±18.3
70% ethanol extract	390.521 ± 14.5
50% ethanol extract	159.049 ± 12.9
Doxorubicin	63.916 ± 5.5

In this study, the anticancer activities of the extracts were examined against human breast carcinoma cells (T47D). Based on the bioactivity results, the best IC₅₀ value was obtained from 50% ethanol extract of *Orthosiphon stamineus* Benth. (table 3) and TLC-densitometry results also showed that this extract contains the higest sinensetin. Sinensetin was able to inhibit proliferation of gastric cancer cell, arterial blood gas (ABG) cancer cells by apoptosis mechanism through P53 and P21 regulation cell using Western Blot Technique (Dong *et al.*, 2011). While rosmarinic acid had known to prevent cell damage caused by free radicals, thereby reducing the risk of cancer and osteosclerosis (Fernando *et al.*, 2016) and is a major compound of polyphenol that can be used as a nutraceutical prod-

uct that helps improve body immunity in cancer patients (Moore *et al.*, 2016). Therefore, the 50% ethanol extract from this plant showed the highest cytotoxic activity against T47D breast cancer cells compared to other extracts and sinensetin has an important role for anticancer properties in the extracts.

CONCLUSION

The 50% ethanol extract from *Orthosiphon stamineus* Benth showed the highest cytotoxic activity against T47D breast cancer cells compared to other extracts. This extract contains highest of sinensetin (32.97 ± 0.06) compared to other extracts. This compound may be responsible for anticancer properties in the extracts.

Acknowledgements

The authors would like to acknowledge Faculty of Pharmacy, Universitas Airlangga for providing the facilities to do the research and funding Penelitian Dosen Pemula (PDP) Project.

CONFLICT OF INTEREST

The authors declare no conflict of interest, financial or otherwise.

REFERENCES

Ahmad, M. K., Ismail, Z., Aisheh, A. A., Majid, A. M. S. A., & Majid A. (2010). Orthosiphon stamine-us leaf extract can prevent the development of colorectal and breast cancer in vivo. Cancer Prevention Research, 3(12), A54.

Akowuah, A. G., Zhari, I., Norhayati, I., Sadikun, A., & Khamsah, S. (2004). Sinensetin, eupatorin, 3'-hydroxy-5, 6, 7, 4'-tetramethoxyflavone and rosmarinic acid contents and antioxidative effect of *Orthosiphon stamineus* from Malaysia. *Food Chemistry*, 87(4), 559-566.

- Arifianti, L., Oktarina, R. D., & Kusumawati, I. (2014) Pengaruh jenis pelarut pengektraksi terhadap kadar sinensetin dalam ekstrak daun Orthosiphon stamineus Benth. Planta Husada, 2(1), 1–4.
- Ashley, K., Andrews, R. N., Cavazosa, L., & Demange, M. (2001). Ultrasonic extraction as a sample preparation technique for elemental analysis by atomic spectrometry. *Journal of Analytical Atomic Spectrometry*, 16, 1147-1153.
- Awale, S., Tezuka, Y., Banskota, A. H., Kouda, K., Tun, K. M., & Kadota, S. (2001). Five novel highly oxygenated diterpenes of Orthosiphon stamineus from Myanmar. Journal Natural Product, 64(5), 592-596.
- Boeing, J. S., Barizão, É. O., Silva, B. C., Montanher, P. F., Almeida, V. D. C., & Visentainer, J. V. (2014). Evaluation of solvent effect on the extraction of phenolic compounds and antioxidant capacities from the berries: application of principal component analysis. Chemistry Central Journal, 8(1), 48.
- Budynas, R. G., & Nisbett, J. K. (2008). Shigley's mechanical engineering design (8th ed.). New York: McGraw-Hill.
- Do, Q. D., Angkawijaya, A. E., Tran-Nguyen, P. L., Huynh, L. H., Soetaredjo, F. E., Ismadji, S., & Ju, Y. (2014). Effect of extraction solvent on total phenol content, total flavonoid content, and antioxidant activity of *Limnophila aromatica*. *Journal of Food* and *Drug Analysis*, 22(3), 296-302.
- Dong, Y., Ji, G., Cao, A., Shi, J., Shi, H., Xie, J., & Wu, D. (2011). Effects of sinensetin on proliferation and apoptosis of human gastric cancer AGS cells. Zhongguo Zhong Yao, Za Zhi, 36(6), 790-794.
- Fei, Y. M., Vuanghao, L., Muhammad, S. I., Ziad, A. O., Fung, A. L., Noersal, R., Fawzy, A. M., Zuhair, A. G., Rusliza, B., Amirin, S., & Zaini, A. M. (2010). HPLC and anti- inflammatory studies of the flavonoid rich chloroform extract fraction of Orthosiphon Stamineus leaves. Molecules, 15, 4452-4466.
- Fernando, P. M. D. J., Piao, M. J., Kang, K. A., Ryu, Y. S., Hewage, S. R. K. M., Chae, S. W., Hyun, J. W. (2016). Rosmarinic acid attenuates cell damage against UVB radiation-induced oxidative stress via enhancing antioxidant effects in human HaCaT cells. Biomolecules & Therapeutics, 24(1), 75-84.
- Floros, J. D., & Liang, H. (1994). Acoustically, assisted diffusion through membranes and biomaterials. Food Technology and Biotechnology, 48(12), 79-84.
- Freshney, R.I. (2005). *Culture of animal cell: A manual of basic technique*. 5th Ed. John Wiley and Sons: New York. 120-135.

- Hossaina, M. A., Ismail, Z., Rahman, A., & Kang, S. C., (2008). Chemical composition and anti-fungal properties of the essential oils and crude extracts of Orthosiphon stamineus Benth. Industrial Crops and Products, 27(3), 328-334.
- Hossain, M. A., & Ismail, Z. (2009). High performance thin layer chromatographic determination of caffeic acid and rosmarinic acid from the leaves of Orthosiphon stamineus. Indonesian Journal of Chemistry, 9(1), 137-141.
- Hossain, M. A., & Ismail, Z. (2016). Quantification and enrichment of sinensetin in the leaves of Orthosiphon stamineus. Arabian Journal of Chemistry, 9(2), S1338-S1341.
- Hossain, M. A. & Rahman, S. M. M. (2015). Isolation and characterisation of flavonoids from the leaves of medicinal plant *Orthosiphon stamineus*. *Arabi*an *Journal of Chemistry* 8, 218-221.
- Juliana, S., Suciati, S. & Indrayanto, G. (2019). Sterol and triterpene profiles of the callus culture of Solanum mammosum. Makara J. Sci., 23(2), 72-78.
- McCauley, J., Zivanovic, A., Skropeta, D. (2013). Bioassays for anticancer activities. Methods in Molecular Biology, 1055, 191-205
- Mohamed, E. A. H., Yam, M. F., Ang, L. F., Mohamed, A. J. & Asmawi, M. A. (2013). Antidiabetic properties and mechanism of action of *Orthosiphon* stamineus Benth bioactive sub-fraction in streptozotocin-induced diabetic rats. Journal of Acupuncture and Meridian Studies, 6(1), 31-40.
- Moore, J., Yousef, M., & Tsiani, E. (2016). Anticancer effects of rosemary (*Rosmarius officinalis L.*) extract and rosemary extract polyphenols. Nutrients, 8(11), 731, 1-32.
- Sahib, I., Ismail, Z., Othman, N. H., & Majib, A. (2009). Orthosiphon stamineus methanolic extract enhances the anti proliferative effect of tamoxifen on human hormone dependent breast cancer. International Journal of Pharmacology, 5(4), 273-276.
- Saidan, N. H., Aisha, A. F., Hamil, M. S., Malik, A. M., & Ismail, Z. (2015a). A novel reverse phase high performance liquid chromatography method for standardization of *Orthosiphon stamineus* leaf extracts. *Pharmacognosy Research*, 7(1), 23-31.
- Saidan, N. H., Hamil, M. S. R., Memon, A. H., Abdelbari, M. M., Hamdan, M. R., Mohd, K. S., Majid, A. M. S. A., & Ismail, Z. (2015b), Selected metabolites profiling of *Orthosiphon stamineus* Benth leaves extracts combined with chemometrics analysis and correlation with biological activities, *BMC Complementary and Alternative Medicine*, 15(350), 1-12

- Sato, T., & Buchner, R. (2004). Dielectric relaxation processes in ethanol/water mixtures. *The Journal* of *Physical Chemistry A*, 108(23), 5007-5015.
- Siddiqui, M. J., Hafizoh, S. N., Ismail, Z., Sahib, H., Helal, M., & Majid, A. M. S. A. (2009) Analysis of total proteins, polysaccharides and glycosaponins contents of *Orthosiphon stamineus* Benth. in spray and freeze dried methanol:water (1:1) extract and its contribution to cytotoxic and antiangiogenic activities. *Pharmacognosy Research*, 1(5), 320-326.
- Singh, M. K., Gidwani, B., Gupta, A., Dhongade, H., Kaur, C. D., Kashyap, P. P. & Tripathi, D. K. (2015). A review of the medicinal plants of Genus Orthosiphon (Lamiaceae). International Journal of Biological Chemistry, 9(6), 318-331.
- Stalikas, C. D. (2007). Extraction, separation, and detection methods for phenolic acids and flavonoids. J. Sep Sci., 30, 3268-3295.
- Sultana, B., Anwar, F., & Ashraf, M. (2009). Effect of extraction solvent/technique on the antioxidant activity of selected medicinal plant extracts. *Molecules*, 14, 2167-2180.
- Sumaryono, W., Proksch, P., Wray, V., Witte, L., & Hartmann, T. (1991). Qualitative and quantitative analysis of the phenolic constituents from *Orthosi*phon aristatus. Planta Medica, 57, 176-80.
- Takeda, Y., Matsumoto, T., Terao, H., Shingu, T., Futatsuishi, Y., Nohara, T., & Kajimoto, T. (1993).
 Orthosiphol D and E, minor diterpenes from Orthosiphon stamineus. Phytochemistry, 33(2), 411-415.

- Tezuka, Y., Stampoulis, P., Banskota, A., Awale, S., Tran, K., & Saiki, I. (2000). Constituents of the Vietnamese medicinal plant Orthosiphon stamineus. Chemical and Pharmaceutical Bulletin, 48, 1711-1719.
- Yam, M. F., Mohamed, E. A. H., Ang, L. F., Pei, L., Darwis, Y., Mahmud, R., Asmawi, M. Z., Basir, R. & Ahmad, M. (2012). A simple isocratic HPLC method for the simultaneous determination of sinensetin, eupatorin, and 30-hydroxy-5,6,7,40-tetramethoxyflavone in *Orthosiphon stamineus* extracts. *Journal of Acupuncture and Meridian Studies*, 5(4), 176-181.
- Younis, L., Abu-Hassan, M. I., Zahar, F. F., & Husin, M. F., M. (2013). The use of Orthosiphon stamineus anticancer action against human oral squamous cell carcinoma. The Open Conference Proceedings Journal, 4, 217.
- Xiao, W., Han, L., & Shi, B. (2008). Microwave-assisted extraction of flavonoids from Radix astragali. Separation and Purification Technology, 62(3), 614-618.
- Zieliński, H., & Kozłowska, H. (2000). Antioxidant activity and total phenolics in selected cereal grains and their different morphological fractions. *Journal of Agricultural and Food Chemistry*, 48, 2008-2016.

Anticancer Property of Orthosiphon stamineus Benth. Extracts in Different Solvent Systems against T47D Human Breast Cancer Cell Lines

ORIGINALITY REPORT SIMILARITY INDEX **INTERNET SOURCES PUBLICATIONS** STUDENT PAPERS **PRIMARY SOURCES** humaniora.journal.ugm.ac.id 2% jurnal.untidar.ac.id Internet Source Chung-Hung Chan, Tiam-You See, Rozita Yusoff, Gek-Cheng Ngoh, Kien-Woh Kow. "Extraction of bioactives from Orthosiphon stamineus using microwave and ultrasoundassisted techniques: Process optimization and scale up", Food Chemistry, 2017 Publication F Novi, S Yati, E Siska, A G Agie. **1** % "Optimization Sonication Time and Dilution Factor in Determining the Concentration of **Endotoxin Challenge Vial with Kinetic** Turbidimetric Method", Journal of Physics: Conference Series, 2021 **Publication** eprints.ums.edu.my

Internet Source

6	Mircea Oroian, Isabel Escriche. "Antioxidants: Characterization, natural sources, extraction and analysis", Food Research International, 2015 Publication	1 %
7	www.ijcmas.com Internet Source	1 %
8	umpir.ump.edu.my Internet Source	1 %
9	journalarticle.ukm.my Internet Source	<1%
10	Ahmad Hazim Abdul Aziz, Nicky Rahmana Putra, Helen Kong, Mohd Azizi Che Yunus. "Supercritical Carbon Dioxide Extraction of Sinensetin, Isosinensetin, and Rosmarinic Acid from Orthosiphon stamineus Leaves: Optimization and Modeling", Arabian Journal for Science and Engineering, 2020 Publication	<1%
11	clinphytoscience.springeropen.com Internet Source	<1%
12	M. F. Yam, L. F. Ang, R. Basir, I. M. Salman, O. Z. Ameer, M. Z. Asmawi. "Evaluation of the anti-pyretic potential of Orthosiphon stamineus Benth standardized extract", Inflammopharmacology, 2008 Publication	<1%

13	H.B. Sahib, Z. Ismail, N.H. Othman, A.M.S. Abdul Maji. "Orthosiphon stamineus Benth. Methanolic Extract Enhances the Anti- Proliferative Effects of Tamoxifen on Human Hormone Dependent Breast Cancer", International Journal of Pharmacology, 2009 Publication	<1%
14	patents.justia.com Internet Source	<1%
15	topsecretapiaccess.dovepress.com Internet Source	<1%
16	www.degruyter.com Internet Source	<1%
17	Cheng-Hsun Wu, Shu-Chun Chen, Ting-Tsz Ou, Charng-Cherng Chyau, Yun-Ching Chang, Chau-Jong Wang. "Mulberry leaf polyphenol extracts reduced hepatic lipid accumulation involving regulation of adenosine monophosphate activated protein kinase and lipogenic enzymes", Journal of Functional Foods, 2013	<1%
18	biology.yonsei.ac.kr Internet Source	<1%
19	studentsrepo.um.edu.my Internet Source	<1%
20	www.thaiscience.info Internet Source	<1%

Dajing Shi, Hui Ding, Shimin Xu.
"Optimization of microwave-assisted extraction of wedelolactone from Eclipta alba using response surface methodology", Frontiers of Chemical Science and Engineering, 2014

<1%

Publication

Hossain, M. A., and S. M. Salehuddin.
"Simultaneous Quantification of Sinensetin and Tetramethoxyflavone in Misai Kucing Capsules using TLC-UV Densitometric Technique", Journal of Scientific Research, 2009.

<1%

Publication

Publication

K A Audah, J Amsyir, F Almasyhur, A M Hapsari, H Sutanto. "Development of extract library from indonesian biodiversity: exploration of antibacterial activity of mangrove leaf extracts ", IOP Conference Series: Earth and Environmental Science, 2018

<1%

Santosh Devkar, Yogesh Badhe, Suresh Jagtap, Mahabaleshwar Hegde. "
Quantification of major bioactive withanolides in (Ashwagandha) roots by HPTLC for rapid validation of Ayurvedic

<1%

products ", Journal of Planar Chromatography – Modern TLC, 2012

Publication

- repository.unair.ac.id <1% 26 Internet Source works.bepress.com Internet Source Bushra Saleem, Muhammad Islam, Hamid 28 Saeed, Fariha Imtiaz, Maryam Asghar, Zikria Saleem, Azra Mehmood, Surriya Naheed. "Investigations of Acacia modesta Wall. leaves for in vitro anti-diabetic, proliferative and cytotoxic effects", Brazilian Journal of Pharmaceutical Sciences, 2018 **Publication** Elsnoussi Ali Hussin Mohamed, Chung Pin <1% 29 Lim, Omar Saad Ebrika, Mohd. Zaini Asmawi, Amirin Sadikun, Mun Fei Yam. "Toxicity evaluation of a standardised 50% ethanol extract of Orthosiphon stamineus", Journal of Ethnopharmacology, 2011 Publication <1%
 - Jokić, Stela, Marina Cvjetko, Đurđica Božić, Sanja Fabek, Nina Toth, Jasna Vorkapić-Furač, and Ivana Radojčić Redovniković. "Optimisation of microwave-assisted extraction of phenolic compounds from broccoli and its antioxidant activity", International Journal of Food Science & Technology, 2012.

31	archive.org Internet Source	<1%
32	ejmcm.com Internet Source	<1%
33	journals.plos.org Internet Source	<1%
34	nmbu.brage.unit.no Internet Source	<1%
35	www.phcogres.com Internet Source	<1%
36	www.science.gov Internet Source	<1%
37	www.slideshare.net Internet Source	<1%
38	www.tandfonline.com Internet Source	<1%
39	Masmoudi, M "Optimization of pectin extraction from lemon by-product with acidified date juice using response surface methodology", Carbohydrate Polymers, 20081016 Publication	<1%
40	"Plant-derived Bioactives", Springer Science and Business Media LLC, 2020 Publication	<1%

Omar Z. Ameer, Ibrahim M. Salman, Mohammad Zaini Asmawi, Zaid O. Ibraheem, Mun Fei Yam. ": Traditional Uses, Phytochemistry, Pharmacology, and Toxicology ", Journal of Medicinal Food, 2012

Publication

42

Retno Widyowati, Neny Purwitasari, Rice Disi Oktarina, Wiwied Ekasari, Saarah Khairunnisa, Hsin-I. Chang. "Exploration of several plants from Baung Forest on bone formation cell models", Journal of Basic and Clinical Physiology and Pharmacology, 2021 Publication

<1%

<1%

Exclude quotes Off Exclude bibliography On

Exclude matches

Off

Anticancer Property of Orthosiphon stamineus Benth. Extracts in Different Solvent Systems against T47D Human Breast Cancer Cell Lines

GRADEMARK REPORT	
FINAL GRADE	GENERAL COMMENTS
/0	Instructor
PAGE 1	
PAGE 2	
PAGE 3	
PAGE 4	
PAGE 5	
PAGE 6	
PAGE 7	
PAGE 8	