Inhibition of Cell Cycle and Induction of Apoptosis y Ethanol Leaves Extract of *Chrysanthemum cinerariifolium* (Trev.) In T47D Breast Cancer Cells

Roihatul Mutiah¹*, Alfiah Laily Inayatin¹, Rahmi Annisa¹, Yen Yen Ari Indrawijaya¹, Anik Listiyanapl

¹Department of Pharmacy, Faculty of Medical and Health Sciences, Maulana Malik Ibrahim State Islamic University of Malang Indonesia, East Java Indonesia 65144
²Department of Medical, Faculty of Medical and Health Sciences, Maulana Malik Ibrahim State Islamic University of Malang Indonesia, East Java Indonesia 65144

**ABSTRACT**

*Chrysanthemum cinerariifolium* (C.cinerariifolium) is a plant of the Asteraceae family, which has been applied by the community as an ornamental plant and traditional medicine. In this study, the effect of *C. cinerariifolium* leaves extract on inhibition of cell cycle and induction of apoptosis in T47D breast cancer cells was tested and compared to the standard chemotherapy agent. The citotoxic activity of *C. cinerariifolium* leaves extract against T47D cancer cells and Vero normal cells was tested by MTT method. Profile of apoptosis and cell cycle were observed by flow cytometry method. Based on chemical compounds profil which is tested used TLC showed that *C.cinerariifolium* leaves extracts contained flavonoid and terpenoid chemical compounds. The result of cytotoxic test showed that leaves extract of *C. cinerariifolium* was able to inhibit the growth of T47D cancer cell at IC₅₀ 418.8µg/mL. Doxorubicin, extracted from Streptomyces peucetius used as treatment in several cancers including breast cancer. Doxorubicin could inhibit the growth of T47D cancer cells in 115.1µg/mL. The results of cell cycle analysis showed that the *C. cinerariifolium* leaves extract inhibited cell cycle in G0-G1 and S phase, whereas doxorubicin was able to inhibit cell cycle in G0-G1 phase but experienced cell accumulation in G2-M phase. The percentage of apoptosis in cycle was showed in M1 (sub G1) and M5 (multinuclear) phase which treatment of *C. cinerariifolium* leaves extract was higher than doxorubicin. Therefore, *C. cinerariifolium* leaves extract has potential activity as anticancer agent causes inhibition of cell cycle and induction apoptosis.

**Keywords:** *Chrysanthemum cinerariifolium*, apoptosis, cell cycle, T47D cells

**INTRODUCTION**

Breast cancer is a malignancy that occurs in cells contained in breast tissue, and both derived from the components of the glands (ductal epithelial or globulus) and those derived from components other than glands such as tissue nerves in the breast, fat tissue, and blood vessels (Sari et al., 2017). American Cancer Society revealed in its latest data that by the year 2013, every year there are about 39,620 women died caused by breast cancer (DeSantis et al., 2014).

Breast cancer treatment efforts such as the use of chemotherapy agents, radiation, radiotherapy has widely used. Doxorubicin is a chemotherapy drug widely used in the treatment of breast cancer. However, the use of chemotherapy drugs is decreased because of the risk of side effects that often occur, such as resistance and cardiomyopathy (Octavia et al., 2012). Therefore, to minimize the occurrence of side effects used traditional medicine using herbs. One of them is chrysanthemum (*Chrysanthemum cinerariifolium*).

*Chrysanthemum cinerariifolium* (*C. cinerariifolium*) is a plant of the Asteraceae family that has been used by the community as an ornamental plant because of its beautiful flowers. Various plant organs can utilized as drugs such as antibacterial, anti-inflammatory, allergy, and also anticancer (Grdisa et al., 2009). Based on previous...
research indicates that the extract of Chrysanthemum zawadskii flower and leaves has pharmacological activity, as is an anti-inflammatory by inhibition of lipopolysaccharide on RAW264.7 cell induced by nitrite oxide (Kim et al., 2012).

The previous study showed that Chrysanthemum contain terpenoid compounds, flavonoids, and derivatives that are suspected of having anticancer activity (Ukiya et al., 2002). Previous research has also reported that C. cinerariifolium comprises a quercetin compound (Jeong et al., 2012; Alviana et al., 2016). The quercetin compound is thought to have an anticancer effect through the induction of the p21 gene which is a CDK inhibitor, along with a decline in Rb gene that inhibits cell cycle in G1 / S phase by inhibiting E2F (Yerlikaya et al., 2017).

Scientific evidence of anticancer activity of chrysanthemum plants with C. cinerariifolium species has not widely performed in both apoptotic and cell cycle testing. In this study reported the effects of C. cinerariifolium extract on apoptosis induction and regulation of breast cancer cell cycle T47D.

MATERIAL AND METHODS

The materials used in this study were C. cinerariifolium leaves, 96% ethanol, distilled water, n-Hexane p. a, ethyl acetate p.a, 10% H2SO4, T47D cells and Vero cells obtained from Parasitology Laboratory Faculty of Medicine Gajah Mada University, Yogyakarta, Complete Medium (CM) RPMI 1640 (Gibco, Invitrogen Canada), CM M199 (Gibco, Invitrogen Canada), PBS, Trypsin-EDTA, DMSO (EMSURE ACS, Japan), SDS (Merck, Berlin Germany), doxorubicin HCL 50 mg, MTT solution (Bio Basic Inc, Canada), trypsin-EDTA 0.25% (Gibco, Invitrogen Canada), RNAse (Gibco, Invitrogen Canada), propidium iodide (Sigma-Aldrich, USA), triton-X (pro GC Merck, Berlin, Germany) and Annexin V (Sigma-Aldrich, USA).

Plant determination

C. cinerariifolium (Trev.) plants were obtained from Nongkojajar, Pasuruan, East Java, Indonesia. Determination of C. cinerariifolium plant was conducted in Materia Medica Integrated Service Unit Batu City, East Java, Indonesia.

Ethical approval

This study has received ethical approval No. 002/EC/KEPK-FKIK/2018 from Medical Research Ethics Committee of Faculty of Medicine and Health Sciences Maulana Malik Ibrahim State Islamic University of Malang.

Sample preparation

Samples of C. cinerariifolium were harvested by cutting on the leaves using scissors. Then begins by sorting each section, washed, dried under the sun, and final sorting. Dry samples are mashed up with grinding machines and weighed C. cinerariifolium leaves powder.

Extraction of C. cinerariifolium

Leaves powder of C. cinerariifolium was put into the Erlenmeyer flask and 96% ethanol solvent was added with a ratio of 1: 20. Then extracted using UAE (Ultrasonication Assisted Extraction) for 2 min with three replications. The leaves filtrate C. cinerariifolium of the UAE evaporated the solvent using a rotary evaporator at 50°C temperature to produce a crude extract. The sticky extract was concentrated using an oven at 40°C temperature until the texture of the extract became concentrated. Then calculated extracted yield using the formula:

\[
\text{Extracted yield} = \frac{\text{Extract weight}}{\text{Raw material weight}} \times 100\%
\]

Identification of compounds used Thin Layer Chromatography (TLC)

In the identification of compounds, the silica gel 60 F254 used as a stationary phase with the n-Hexane p. a (Merck, Berlin, Germany) and ethyl acetate p.a (Merck, Berlin, Germany) (8: 2) periods of motion. The stain used was 10% H2SO4. Identify stain compounds using Thin Layer Chromatography (TLC) Visualizer.

Sample preparation for anticancer activity and toxicity test

Leaves extract was weighed as much as 10mg, dissolved with 100μg/mL DMSO and made seven serial concentrations in T47D cells were 1000; 800; 600; 400; 200; 100; 50μg/mL and in Vero cells were 1000; 500; 250; 125; 62.5; 31.25; 15.625µg/mL. While doxorubicin positive control was made seven serial concentrations, in T47D cells were 1087.04; 543.52; 271.76; 135.88; 67.94; 33.97; 16.985µg/mL and in Vero cells were 5435; 2717.60; 1358.80; 679.40; 339.70; 169.85; 84.925 µg/mL.

Anticancer activity and toxicity test

An anticancer activity test conducted in T47D cell culture with RPMI 1640 medium (Gibco,
Invitrogen Canada). Furthermore, toxicity test performed in Vero cell culture used the M199 medium (Gibco, Invitrogen Canada). T47D cell cultures and Vero cells were grown on 96 well plates and then incubated for 24h. After 24h the media was removed and washed used PBS, then each concentration of extracts was added into each well with three replications and incubated for 24h. After 24h the media was removed and washed used PBS, then added 100μL MTT reagents (Bio Basic Inc, Canada) to each well, including media control (without cells), then re-incubated for 4h in the CO\textsubscript{2} incubator.

After 4h the cell condition was observed under an inverted microscope, then a 100μL SDS 10% stopper was added and incubated at room temperature overnight. Furthermore the absorbance value is read using ELISA reader and calculated cell viability using the following formula:

\[ \text{Viability cell} = \frac{(T_a - M_c a)}{(C_c a - M_c a)} \times 100\% \]

\( T_a = \text{Treatment absorbance; } M_c = \text{Media control absorbance; } C_c a = \text{Control cell absorbance} \)

The result of viability cell obtained by IC\textsubscript{50} analysis for anticancer activity and CC\textsubscript{50} analysis for normal cell toxicity using Microsoft Excel (Mut’ah, 2017).

**Flowcytometry test**

The cultured T47D cells as much as 5x10\textsuperscript{5} cells/well were grown in RPMI medium at 6-well plate (for treatment and control cells) then incubated for 24h. The cell condition was observed in the microscope to see the cell distribution. Furthermore, the concentration of the sample and doxorubicin was made to the level of IC\textsubscript{50}. The 6-well plate that already contain cells taken from the incubator. Then, the media cell was removed by using a Pasteur pipette slowly and washed with PBS. Furthermore, the treatment conducted by inserting 2mL extract samples at the first well (for cell cycle), 2mL doxorubicin at the second well, and 2mL cell control at the third well and incubated for 24h. One conical was prepared for one type of treatment or one well. The medium is taken 1mL from the well with micropipette and transferred to the conical. To the each well, 1mL PBL was added 1mL PBS and moved into the conical. The conical was added 200μL of trypsin-EDTA 0.25% and incubated for 3min. Furthermore, the wells were added 1mL of control media into each well and resuspension until the cells disengaged one by one and the cells are transferred to the conical. The wells are added 2mL of PBS to retrieve the remaining cells, then moved into the conical. The conical is centrifuged at 2000 rpm for 5min and the supernatant was removed. The each well was washed with 1mL PBS again and moved to the conical. The conical is resuspended then moved to microtube. The microtubes were centrifuged at 2000rpm for 3min.

In cell cycle analysis, the supernatant is removed by pouring and added 500μL 70% alcohol into the conical while shaking slowly. The conical is kept at room temperature (37°C) for 35min and centrifuged at 600rpm for 5min to removed the added alcohol from the conical. The conical was added 500μL of PBS and centrifuged at 2000rpm for 3min. The washing was conducted twice used PBS and the conical is wrapped in aluminum foil and marked. For apoptotic analysis, the remained harvest cells was rinsed used PBS then it was centrifuged again and the PBS was removed from the conical. The conical was added of Propidium iodide reagent and allowed to stand for 30min. The sludge was added PI-Annexin V reagents carefully and immediately homogenized. The microtube containing the cell suspensions is wrapped in aluminum foil and incubated in a 37°C water bath for 5min. The cells suspension is homogenized and transferred into a flow cytometer tube using a nylon filter to test for cell cycle and cell apoptosis, then ready for analysis with a flow cytometer (CCRC, 2009).

**RESULTS AND DISCUSSION**

Identification of *C. cinerariifolium* leaves extract chemical compound

Chemical compounds identification was conducted to analyze qualitative content and expect phenolic and terpenoid compounds before the cytotoxic and apoptosis to be tested. Thin Layer Chromatography (TLC) is a physicochemical separation method based on two phases, which is a fluid phase as a mobile phase and a solid phase as a stationary phase (Mut’ah et al., 2013). Based on the optimization results showed that the best mobile phase were n-Hexane and ethyl acetate (6:2) as solvents. After elution and air drying of the plate, natural product reagent was sprayed using 10% H\textsubscript{2}SO\textsubscript{4} as universal staining. Furthermore, the plate was compared before and after to detection and observed the spots characteristic. The result of TLC visualizer identification used UV 366 rays shows the difference of compound separation between before and after spray (Figure 1).
It seen in the Rf results obtained, the Rf compounds on the TLC plate after spraying more than the Rf value on the TLC plate before spraying. The number of Rf TLC plate on the leaves extract after sprayed is 8.0. The yellow color with wavelength 341-389 nm after being sprayed showed the flavonol group compound, and the purple, red color showed terpenoid group compounds (Harborne, 1987). Meanwhile, purple
stains are suspected as sesquiterpenes compounds (Mutiah et al., 2013). Compounds of flavonoids and terpenoid groups play a major role in health, one of which is having anticancer activity in breast cancer (Bishayee et al., 2011; Weeb and Ebeler, 2004).

**Cytotoxic test of C. cinerariifolium leaves extract**

The anticancer activity test of *C. cinerariifolium* leaves extracts known by decreasing living cells percentage based on 50% Inhibitor Concentration (IC50) value. Cell morphological observations conducted under an inverted microscope after treatment on cells with each extract. Comparison of the inhibitory effect of T47D cell growth and Vero cell after treated with extract *C. cinerariifolium* (Figure 2). Morphological change between T47D control cells using the treatment of *C. cinerariifolium* leaves extract and doxorubicin positive control (Figure 2). The form of viable T47D cells was elongated, while the dead T47D cell was shaped rounded shrunk (Iin et al., 2014). In Vero cells, there was no apparent cell death due to the treatment of *C. cinerariifolium* leaves extract compared to control. Morphology of viable Vero cells was polygonal and flat (Goncalves, 2012). The treatment of *C. cinerariifolium* leaves extract possessed cytotoxic activity against human cancer cell lines T47D but does exert damage to normal vero cells. To determine the viability of cancer cells due to the treatment of *C. cinerariifolium* extract, then tested the cytotoxicity of T47D cells and Vero cells used the MTT method. The intensity of the purple color that formed is proportional to the number of living cells (Doyle and Griffiths, 2000). The higher the intensity of the purple color indicates a more significant amount of living cells (CCRC, 2009). Percentage of cell viability in T47D and Vero cells due to the treatment of doxorubicin and leaves extract (Figure 3).

**Cell cycle due to treatment of C. cinerariifolium leaves extract**

Inhibition in the phase of the cell cycle that observed was conducted using flow cytometry method. The flow cytometry method was a method that can detect every phase in the cell cycle based on the number of chromosomes on each phase (G1, S, and G2/M).
Through this flow cytometry (Figure 4) method, the distribution of cells at each phase in the cell cycle after treatment could be known. Furthermore, the pathway inhibition of *C. cinerariifolium* leaves extracts in blocking the cell cycle could be estimated.

**Apoptosis induction after treating of *C. cinerariifolium* leaves extract**

In flow cytometry test of *C. cinerariifolium* leaves extracted, the IC$_{50}$ was 418.8μg/mL and compared with doxorubicin at IC$_{50}$ 115.1μg/mL. The results of the apoptotic flowcytometry test against T47D cells (Figure 5). The colors formed in the cell dispersion data were analyzed using a Cell Quest program so that the colors formed can be separated according to the population. Living cells indicated by green, early apoptotic cells exhibited by yellow, late apoptosis indicated by pink, and necrosis indicated by red. The resulting colors obtained from cells that emit epi-fluorescence due to Annexin V or PI bonds which are then captured by UV rays (Indradmojo, 2015).

Anticancer activity on T47D and Vero cell and the effects of *C. cinerariifolium* leaves extract on apoptosis induction and regulation of breast cancer cycle T47D conducted as the purpose of this study. IC$_{50}$ result obtained from *C. cinerariifolium* leaves extract was 418.8μg/mL, while IC$_{50}$ doxorubicin result was 115.1μg/mL (Figure 3). The results of IC$_{50}$ obtained showed that 96% ethanol leaves extract of *C. cinerariifolium* to have anticancer activity against breast cancer (T47D). An extract has high anticancer activity if IC$_{50}$ <500μg/mL and has weak activity if IC$_{50}$> 500μg/mL (Costa et al., 2017). The IC$_{50}$ result in positive control equal to 115.1μg/mL and these result obtained was close to IC$_{50}$ researchers that IC$_{50}$ doxorubicin value against T47D cells is 250nM or 135.9μg/mL (Abdolmohammadi, 2008).
Doxorubicin has a low IC\textsubscript{50} result because it has high activity against breast cancer cells (Anjarsari, 2013). The results of cytotoxic activity test on Vero cell (Figure 3) that obtained from IC\textsubscript{50} showed that 96% ethanol leaves extract of \textit{C. cinerariifolium} has low toxicity to normal cells. The result of IC\textsubscript{50} leaves extract of \textit{C. cinerariifolium} in Vero cell was 676.182μg/mL, while IC\textsubscript{50} doxorubicin value was 1234.5μg/mL. The low toxicity of normal cells in vitro tests correlates with high levels of safety against normal cells (Mutiah \textit{et al.}, 2017).

Percentage analysis of the viability of living cells, apoptosis, and necrosis showed untreated cells (cell control) had a cell viability percentage of 94.47%, 3% apoptotic cells, and 1.82% necrosis cells (Figure 5). While doxorubicin-treated cells showed a decrease in cell viability to 77.78%, an increase in apoptosis by 17.13%, and an increase in the number of necrosis by 5.18% when compared with control cells. However, based on the percentage of cell viability when compared with the rate of apoptosis and necrosis cells, the percentage of living cells still more significant than dead cells. It might because doxorubicin was able to increase the activity of phosphorylation of P13K/ Akt then activate Bcl protein which is an antia apoptosis protein and could activate Bad protein which is a protein trigger apoptosis (Setiawati, 2011).

In the treatment of ethanol leaves extract of \textit{C. cinerariifolium} showed that T47D cell apoptosis induction was 49.88% and necrosis was 47.29%, and the living cell was 2.83% (Figure 4). The result of statistical analysis of three treatments showed a significant difference
between apoptosis percentage on cell control, after treatment of doxorubicin, and after treatment of *C. cinerariifolium* leaves extract with the considerable p-value (p<0.01). The results of the statistical analysis of the percentage of cell necrosis also showed similar results. The flow cytometry analysis conducted used cell quest program (Figure 5). Cell distribution in each phase of the cell cycle was colored used PI reagents as it was able to interact with DNA (Putri, 2014). Based on Figure 5 there was changed in the cell cycle between the *C. cinerariifolium* leaves extract and control cells that indicated by the reduction in cell numbers in the G0-G1 phase and S phase. These reductions meant the cessation of the cell. While in phase M1, the control cell was increased which indicates the presence of apoptotic cell death. Therefore, the cell may couldn’t proceed to the next phase (Mutiah et al., 2017). Doxorubicin led to cellular inhibition in the G0-G1 phase but pointed to the cell shift to the right at the G2-M phase indicating that the cell enters the phase (Mutiah, 2014). Accumulation cells in the G2-M phase causes the down regulation of Cdc24c, Cdk1, and Cyclin B (Su, et al., 2006). Inhibiting Cdc2/Cyclin B is one of the result in the effect of ATM-ATR signaling, causing G2-M arrest (Kolb, et al., 2012).

The increase of apoptosis by *C. cinerariifolium* leaves extract suspected due to the content of flavonoid compounds. Flavonoid compounds could induce apoptosis through p53 pathways. If flavonoids induce apoptosis through irreversible DNA damage, then apoptosis was possible through the p53 pathway (Meiyanto and Septisetyani, 2005). Anticancer activity of *C. cinerariifolium* leaves extract suspected by the mechanism of the flavonoid compounds contained in the leaves extract of *C. cinerariifolium*. Flavonoid compounds can inhibit the overall performance of CDK which is a cell cycle regulator. Inhibition that occurs through the working of enzymes CDK-activating kinase then could inhibit the formation of active CDK-cyclin complex (Supriatno and Rasmindar, 2014). Proteins that could inhibit the CDK-cyclin complex were INK4 and CIP (p21). INK4 plays a role in inhibiting progression in G1 phase. When a G1 phase occurs, the cell automatically cannot proceed to the next phase (Mutiah, 2014).

**CONCLUSION**

In conclusion, the results of the present study reveal that ethanol leaves extract of *C. cinerariifolium* could induce 49.88% T47D cell apoptosis at IC50 418.8μg/mL concentration, and could inhibit cell cycle in phase G0-G1 and S phase and increased cell number in phase M1. The flavonoid compounds that content in *C. cinerariifolium* supposed could induce apoptosis through p53 pathways. The ability of *C. cinerariifolium* ethanol leaves extracts to induce apoptosis to suggest that it could be a new candidate of anticancer therapy.

**ACKNOWLEDGEMENT**

This work supported by The Directorate General of Islamic Higher Education (DIKTIS) of Interdisciplinary Basic Research Grant numbers 3209/Un.3/HK.00.5/05/2018.

**REFERENCES**

Abdolmohammedi MH., Fouladdel SH., Shafee A., Amin GH., Ghaffari SM., Azizi E. 2008. Anticancer Effects and Cell Cycle Analysis on Human Breast Cancer T47D Cells Treated with Extracts of *Astrodaucus persicus* (Boiss.) Drude in Comparison to Doxorubicin. Journal Daru, 16:112-118.

Alviana N., Sidharta BR., Martini T. 2016. Test the effectiveness of antibacterial ethanol extract of *Chrysanthemum* of green (*Chrysanthemum morifolium Syn. Dendrathema grandiflora*) against *Staphylococcus aureus* and *Escherichia coli*. Yogyakarta: Universitas Atma Jaya Yogyakarta.

Anjarsari EY., Kristiani N., Larasati YA., Dewi D., Putri P, Meiyanto E. 2013. Synergistic Effect of Cinnamon Essential Oil (*Cinnamomum burmannii*) and Doxorubicin on T47D Cells Correlated with Apoptosis Induction. *IJCC*. 450-456.

Bishayee A., Ahmed S., Brandao GC., Dolabela MF. 2017. Anti-leishmanial Activity of *Handroanthus serratifolius* (Vahl) S. Grose (Bignoniaceae). Evidence-Based Complementary and Alternative Medicine Hindawi, 1:1-6.
DeSantis C., Ma J., Bryan L., Jemal A. 2013. Breast cancer statistics, 2013. A Cancer Journal for Clinicians, 64:52-62.

Doyle A., Griffiths JB. Cell and Tissue Culture for Medical Research. New York: John Willey and Sons Ltd 2000.

Goncalves D. 2012. The Micro-analysis of Human Burned Bones: Some Remarks. Cadernos do GEEvH, 1:32-40.

Grdisa M., Carovic-Stanko K., Kolak I., Satovic Z., 2009. Morphogical and Biochemical Diversity of Dalmatian Pyrethrum (Tanacetum cinerariifolium (Trevir.) Sch.Bip.). Agriculturae Conpectus Scientificus, 74(2):73-80.

Harborne JB. 1987. Phytochemical Methods Guide Modern ways to analyse plants. Bandung : ITB.

Jin T., Andrew J., Kisdjiamiatun. 2014. In Vitro toxicity of ethanolic acid fruit Parijoto (Medinilla speciosa, reinw.ex bl.) against breast cancer cells T47D. Jurnal Gizi Indonesia, 2:53-58.

Indradmojo C. 2015. Anti-cancer activities and pharmacological mechanisms of extracts and Benaul Nangka fraction (Macrosolen cochinchinensis) in breast cancer cells T47. Malang: UIN Maulana Malik Ibrahim Malang.

Jeong SW., Park S., Jin JS., Seo ON., Kim GS., Kim YH., et al., 2012. Influences of Four Different Light-Emitting Diode Lights on Flowering and Polyphenol Variations in the Leaves of Chrysanthemum (Chrysanthemum morifolium). J Agric Food Chem, 60:9793-9800.

Kim Y., Han J., Sung J., Lee J. 2012. Anti-inflammatory activity of Chrysanthemum zawadskii var. latilobum leaf extract through haem oxygenase-1 induction. Journal of Functional Foods, 4:474-479.

Kolb RH., Greer PM., Cao PT., Cowan KH., Yan Y. 2012. ERK1/2 signaling plays an important role in topoisomerase II poison-induced G2/M checkpoint activation. PLoS One, 7(11):e50281.

Meiyanto E., Septisetyani EP. 2005 Antiproliferative effects and Apoptosis of phenolic extract of etanolic leaves Gymnura procumbens (Lour.) Merr. Against HeLa cells. Artocarpus, 5:1-12.

Mutia'h R. 2017. Study of efficacy and safety of extracts of root and leaf Calotropis gigantea against colon cancer cells and breast cancer in Vitro. J Islamic Med, 2:67-75.

Mutiah R., Listiyana A., Indradmojo C., Griana TP., Dwi HH., Atmaja RR. 2017. Induction of Apoptosis and Phase-Cell Cycle Inhibition of G0-G1, S, G2-M of T47D Breast Cancer Cells on Treatment with Ethyl Acetate Fraction of Jackfruit Parasite Leaves (Macrosolen cochinesis). Journal of Applied Pharmaceutical Science, 7:138-143.

Mutiah R., Hayati EK, Triastutik Y. 2013. Separation and identification of sunflower leaf extract (Helianthus annus L.) used thin layer chromatography. Alchemy, 2:190-194.

Mutia'h, R., Widyawaruyanti A., Sukardiman. 2018. Calotropis gigantea A: Glycosides Terpenoids from Calotropis gigantea Induces Apoptosis of Colon Cancer WiDr Cells through Cell Cycle Arrest G2/M and Caspase 8 Expression. Asian Pacific Journal of Cancer Prevention, 19:1457-1464.

Mutia'h, R. 2014. Development of Phytopharmacy and anticancer (guidelines and techniques for development of Indonesian Herbal medicine into Phytofarmaka). Malang: UIN Maliki Press.

Octavia Y., Tocchetti CG., Gabrielson KL., Janssens S., Crijns HJ., Moens AL. 2012. Doxorubicin-induced cardiomyopathy: From molecular mechanisms to therapeutics strategies. J Mol Cell Cardiology, 52(6):1213-1225.

Putri H. 2014. Sample preparation for cell cycle by Flowcytometry method. Yogyakarta: Cancer Cemeprevention Research Center Pharmacy Faculty Gadjah Mada University.

Sari M., Dewi YL., Utami A. 2017. Family support relationship to the motivation of breast cancer patients in a chemotherapy in the room of Cendrawasih 1 RSUD Arifin Achmad Riau Province. J Ners Indonesia, 2:158-166.

Setiawati S., Meiyanto E. 2011. Increased cytotoxic effect of Doxorubicin by Hesperidin in cancer cells T47D. Bionatura-Jurnal Ilmu-Ilmu Hayati dan Fisik, 15:85-92.

SuCC., Lin JG., Li TM., Chung JC., Yang JS. et al., 2006 Curcumin-induced apoptosis of human colon cancer COLO-205 cells through the production of ROS, Ca2+ and the activation of caspase 3. Anticancer Res. 26, 4379-4389

Supriatno AH, Rasmindar M. 2014. Anticancer and antiproliferation activity of ant-nest ethanol fraction (Myrmecodia pendants) in human tongue cancer SP-CI. Dentofasial, 13:1-6.

Ukiya M., Akhisa T., Tokuda H., Suzuki H., Mukainaka T., et al., 2002. Constituents of Compositae plants III. Anti-tumor promoting
effects and cytotoxic activity against human cancer cell lines of triterpene diols and triols from edible chrysanthemum flowers. *Cancer Letters*, 177(1):7-12

Weeb MR, Ebeler SE. 2004. Comparative Analysis of Topoisomerase IIB Inhibition and DNA Intercalation by Flavonoids and Similar Compounds: Structural Determinates of Activity. *Biochem J.* 527-541.

Yerlikaya PO, Arisan ED, Gurkan AC, Unsu NP. 2017. Breast Cancer and Flavonoids as Treatment Strategy. *From Biology to Medicine*, 2:305-326.