Ethanolic Extract of *Citrus reticulata* Peel Inhibits the Migration of WiDr Colon Cancer Cells

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Abstract

Colorectal cancer is third rank on the cancer cases in Indonesia. To cure the cancer needs big cost and lot of effort. On the other side, the side effect of medicine or chemotherapy on patient need to reduce. Cancer cell spread to other tissue based on its migration and invasion ability. *Citrus reticulata* peel contains flavonoid such as Tangeretin and Nobiletin, both of this compounds have anticancer activity. The aims of this study is to reveals the potency of ethanolic extract of *Citrus reticulata* peel on the inhibition of migration on WiDr colon cancer cells. The toxicity of ethanol extract of *Citrus reticulata* peel on WiDr colon cancer line was measured using 3-(4,5-dimethyltiazol-2-il)-2,5-diphenyltrazolium bromide (MTT) assay and investigate the cell migration was using scratch wound healing assay. The ethanol extract of *Citrus reticulata* peel showed the value of inhibitory concentration 50 (IC₅₀) was 184.5 μg/mL, this result categorize as moderate cytotoxic. Meanwhile the migration assay showed that the deceleration of migration occurred on 0.5 IC₅₀, 0.33 IC₅₀ and 0.25 IC₅₀ during 24 h and 36 h incubation, event thought there were not significant different (p>0.05). The ethanol extract of *Citrus reticulata* peel has a potential migration inhibition on WiDr cell line.

Keywords: *Citrus reticulata*, WiDr cell line, migration

INTRODUCTION

Cancer is one of the causes of death in the whole world. The evidence of cancer around 56.8% and the death cause of cancer (64.9%) grow on the develop country in 2012 and was predicted increasing 19.3 new cases per year in 2025 (WHO, 2013). Colorectum cancer is ranked third after breast cancer and lung cancer (WHO, 2013). The highest mortality of colorectum cancer at Central and East Europe (20.3 per 100,000 for male and 11.7 per 100,000 for female). Colorectum cancer in Indonesia has a third position as 12.8 per 100,000 for adult community and has 9.5% mortality of total cancer evidence (IARC, 2012).

Most of colorectal patient come to the hospital with worse condition due to unknown early signs of cancer or ignored this sign event. (Ministry of Health of RI, 2017). Cancer started from uncontrolled growth cell. Repair DNA failed to work and then induced mutation on genome of soma cell. Colorectal cancer was malignancies on colon tissue, especially on part of colon and rectum. The malignancy is begun from proliferation and migration of cell, invasion cell to other tissue such as blood and to other organ and growing fast after...
constructs new vessel to fulfill nutrient demand for their metabolism.

The inhibition of cancer cell migration is very important effort to prevent invasion to other tissue. There were many ways to cure colon cancer such as chemotherapy, take medicine, and surgery. In addition, the side effect during cancer therapy still a major problem. The exploration of adjuvant medicine to reduce the side effect is necessary to do the most important effect to prevent the malignancies of cell and invasion cancer cell to other tissue. Many plants used to as basic material to prevent from many diseases, including cancer. For example, *Citrus reticulata* peel has been reported contain of tangeritin and nobelitin, member of flavonoids (Shu-Fen, et al., 2004 and Morley, 2007).

Tangeritin was able to inhibit HL-60 Leukemia cancer cell (Hirano, et al., 1995) and to induce cell-cycle G1 arrest. Tangeritin was reported has the lowest inhibitory concentration 50 (IC_{50}) than other flavonoids on COLO 205 colon cancer cell (Pan, et al., 2002). Nobelitin has activity to suppress the production of MMP1.9 on HT1080 human fibrosarcoma cancel cell line (Sato, et al., 2002), inhibit activity of MCF breast cancer cell line and NCI-H460 lung cancer cell line, basal skin carcinoma (Wu, et al., 2015).

**MATERIALS AND METHODS**

**Extraction of *Citrus reticulata* Using Maceration Methods**

Sixty grams (60 g) of *Citrus reticulata* peel powder was put in three of 1000 mL Erlenmeyer (20 g each) extracted using 200 mL 95% ethanol solvent. Samples were extracted for 24 h and shaken for 3 h at room temperature. The dissolved fraction in ethanol is separated, put into a flask and the pulp obtained was macerated using same solvent. This extraction was carried out three times. Second repetition and third, the volume of immersion solvent is 150 mL. The filtrate obtained was concentrated with a vacuum rotary evaporator at a 60°C and stopped when the extract was thick enough, marked by stopping the solvent dropping on the round bottom flask. The thick extract was stored at a temperature of less than 20°C to prevent damaged.

**WiDr Cancer Colon Maintenance**

WiDr cancer colon from Cancer Chemoprevention Research Center, Faculty of Pharmacy, Universitas Gadjah Mada were maintenance in RMPI, enriched with FBS, fungizone, Streptomycin/penicillin. The cell growth until 80% confluence for treatment. Trypsin and EDTA was used to detach the cell.

**Cytotoxic Test using 3-(4,5-dimethyltiazol-2-il)-2,5-diphenyltrazolium bromide (MTT) assay**

The principle of the 3-(4,5-dimethyltiazol-2-il)-2,5-diphenyltrazolium bromide (MTT) method is the reduction of yellow tetrazolium MTT salt to be purple formazan crystals by the succinate dehydrogenase enzyme due to cellular metabolic activity by NAD(P)H-dependent cellular enzyme oxidoreductase in the cell respiration pathway in the mitochondria (Berridge, et al., 2005). Addition of stopper reagent (which is detergent) will dissolve this colored crystal then absorbed using an Enzyme-
linked immunosorbent assay (ELISA) reader. The intensity of the purple color is proportional to the number of living cells. If the intensity of the purple color is greater, it means that the number of living cells is increasing.

The sample to be used in the cytotoxic test consisted of nine treatment groups. First group was WiDr cells as negative controls that were not given any treatment and only allowed to grow in the growth media. Second group was WiDr cells as positive controls given doxorubicin as a standard cancer drug with doses of 7.5 μg/mL, 15 μg/mL, and 20 μg/mL. The 3rd, 4th, 5th, 6th and 7th group were WiDr cells that were given ethanol extracts of *Citrus reticulata* peel with dosages 10 μg/mL, 40 μg/mL, 80 μg/mL, 100 μg/mL. The positive and negative control groups were repeated three times in each treatment group. Meanwhile, the cell groups that were given ethanol extracts of *Citrus reticulata* peel were repeated five times for each treatment group.

**Migration Assay using Scratch Wound Healing Assay**

Aproximately 5x10^4 of WiDr cell line were seeded on the six well plate. This cell were devide into six treatment groups with negative control (cell with media only) and positive control (using doxorubicin). Treatment doses were 1/2 IC_{50}, 1/3 IC_{50}, 1/4 IC_{50}, 1/8 IC_{50} of ethanol extract of *Citrus reticulata* peel. After cell attached and spread then growing around 80% confluent, make a scratch using sterile yellow tip. A scratch will removed the cell, and became a discrete area so each edge cell can migrate to the empty area. Then add 5 mL treatment solution as well. On 0, 12 and 24 h took a picture to every treatment and analyze using image J software. The migration test was repeated three times for each treatment.

**Data Analysis**

Data was represented as mean±SD, the significant value continued with LSD. Data analyzed using SPSS program. Ethical clearance for this research was issued with number 643/EP-FKIK-UMY/XII/2017.

**RESULTS**

The average of WiDr colon cancer cell viability showed depending of dose on ethanol extract *Citrus reticulata* as shown in Table 1. The increasing of dose was followed by the decreasing of WiDr colon cancer cell viability.

**Table 1. The average of viability WiDr cell line on depending of dose.**

<table>
<thead>
<tr>
<th>Dose of Ethanol Extract of <em>Citrus reticulata</em> (µg/mL)</th>
<th>WiDr Colon Cancer Cell Viability (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>90.7±0.034</td>
</tr>
<tr>
<td>40</td>
<td>72.7±0.121</td>
</tr>
<tr>
<td>80</td>
<td>69.2±0.07</td>
</tr>
<tr>
<td>100</td>
<td>63.6±0.05</td>
</tr>
<tr>
<td>240</td>
<td>9.2±0.023</td>
</tr>
</tbody>
</table>

The adding 30, 60 and 140 µg/mL of dose were followed by decreasing as 22%; 3%; 8.6% and 53.8% of WiDr colon cancer cell line viability. Figure 1 showed the IC_{50} of ethanol extract of *Citrus reticulata* peel was 184.5 µg/mL.

**Figure 1. Percentage of WiDr cell line viability depending dose of *Citrus reticulata* peel ethanololic extract.**
Meanwhile as a positive control was doxorubicin on some doses experiment as shown in Table 2. The IC\textsubscript{50} of doxorubicin was 7.5 μg/mL.

**Table 2 The result of doxorubicin treatment on WiDr colon cancer cell viability.**

<table>
<thead>
<tr>
<th>Doxorubicin (μg/mL)</th>
<th>WiDr Colon Cancer Cell Viability (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>7.5</td>
<td>52.74±0.06</td>
</tr>
<tr>
<td>15</td>
<td>47.95±0.04</td>
</tr>
<tr>
<td>20</td>
<td>16.09±0.02</td>
</tr>
</tbody>
</table>

The migration WiDr colon Cancer cell per 12 h shown on Figure 2. Figure 2 showed that all groups have decreasing on their migration area depending of time. The anova analyzed showed that there were difference between all groups. Event it’s not significance (p>0.05).

**Figure 2. The inhibition of migration from *Citrus reticulata* peel extract on WiDr Colon Cancer Cell using scratch wound healing assay.** WiDr cell were seeded in six well plate after confluence then scratch with blue tip and wash with PBS. The cell then treated with doxorubicin (positive control), 1/8 IC\textsubscript{50}, 1/4 IC\textsubscript{50}, 1/3 IC\textsubscript{50} and 1/2 IC\textsubscript{50} of ethanol extract of *Citrus reticulata*. This assay was measured for 12, 24 and 36 h.

The percentage WiDr colon cancer cell area migration as shown on Table 3. On the 12 h the smallest area migration on the 1/2 IC\textsubscript{50} and the largest area migration on 1/4 IC\textsubscript{50}. On the 24 h the smallest area migration on doxorubicin and the largest on the 1/4 IC\textsubscript{50}. On the 36 h, the smallest area migration on the 1/2 IC\textsubscript{50} and the largest area migration on the 1/4 IC\textsubscript{50}.

**Table 3. Percentge of WiDr colon cancer cell area migration.**

<table>
<thead>
<tr>
<th>Migration Time (hour)</th>
<th>Negative Control (%)</th>
<th>Doxorubicin (%)</th>
<th>Citrus reticulata Peel Extract (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12</td>
<td>11.6</td>
<td>8.4</td>
<td>13.9</td>
</tr>
<tr>
<td>24</td>
<td>19.9</td>
<td>12.7</td>
<td>22.4</td>
</tr>
<tr>
<td>36</td>
<td>28.8</td>
<td>21.0</td>
<td>36.1</td>
</tr>
</tbody>
</table>

**Table 4. Interval percentage of WiDr colon cancer cell area migration.**

<table>
<thead>
<tr>
<th>Interval on the Migration Time (hour)</th>
<th>Control (%)</th>
<th>Doxorubicin (%)</th>
<th>Citrus reticulata Peel Extract (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12-24</td>
<td>8.3</td>
<td>4.3</td>
<td>8.5</td>
</tr>
<tr>
<td>24-36</td>
<td>8.9</td>
<td>8.3</td>
<td>13.7</td>
</tr>
</tbody>
</table>

**DISCUSSION**

This study reveal that extract ethanol *Citrus reticulata* peel have antimigration effect, as compared with doxorubicin. The 1/2 IC\textsubscript{50} of extract ethanol *Citrus reticulata* have the largest effect on antimigration of WiDr colon cancer cell than others dose, as shown had the lowest area tomi grate. This result in line with Celano, et al. (2015) that reported the extract caused a reduction of cell migration, associated with decreased activity of the metalloproteinase MMP-2. These findings show that the flavonoid fraction of *Citrus reticulata* juice exerts in vitro antiproliferative effects on ATC cells (Human Tyroid cancer cell line), associated with a reduction of migration. This result also in line with Ajikumaran S Nair, et al. (2018) reported that *Citrus reticulata* peel water extract at 25, 50 μg/mL showed 100% cell death in DLA cells (Dalton’s Lymphoma Acites) also had antitumor on mice.

Table 3. showed the interval per 12 h that control group have stable acceleration of migration, almost similar with the 1/3 IC\textsubscript{50} group but doxorubicin have twice acceleration of migration (from 4.3 to 8.3). The decreasing migration on 1/2 IC\textsubscript{50} (from 6.9 to 4).
Meanwhile Tahsin, et al. (2017) showed that the extract of n-hexane (Hex) of Citrus reticulata Blanco bark, contain of scoparone, xanthyletin, lupeol, β-amyrin, stigmasterol, β-sitosterol and palmitic acid, had IC$_{50}$ values as 53.0; 52.4 and 49.1 μg/mL against the cancer cell lines A54 (human lung adenocarcinoma cell line), MCF7 (human breast adenocarcinoma cell line) and PC3 (human Caucasian prostate adenocarcinoma). In line with Kim, et al. (2005) reported that Citrus reticulata had an apoptosis effect on Human Gastric Cancer Cells SNU-668 by the high expression on Bax, caspase 3 and low expression of BCl2 on dose and time dependent manner. The inhibition of migration, invasion and proliferation liver cancer cell via down regulation of MMP-2/9, N-cadherin, and vimentin induced by combination between Citrus reticulate peel and black tea (Wen, et al., 2019). Similar result showed by Chang, et al. (2015), that Citrus reticulate has inhibitory effect on ovarian cancer metastasis, also inhibit TGF-β1-induced EMT through the canonical TGF-β1-SMAD-Snail/Slug axis. As known that Citrus reticulata contains of some flavonoids such as tangeritin, nobilin (Morley, 2007) also narirutin, hesperidin (Tumbas, et al., 2010). Citrus reticulata have many bioactivity i.e., as anticancer (Garcia and Castillo, 2008; Wu, et al., 2015), antibacteria activity, (Hamdan et al., 2016) and antiinflammatory (Boughendjoua and Boughendjoua, 2017). Tangeritin and nobelitin in Citrus reticulata can up-regulate p53 expression. Flavonoids can act as all three types of agent, as to inhibit enzymes involved in cell activation. Attempts to control cancer involve a variety of means, including the use of suppressing, blocking, and transforming agents. Suppressing agents prevent the formation of new cancers from procarcinogens, and blocking agents prevent carcinogenic compounds from reaching critical initiation sites, while transformation agents act to facilitate the metabolism of carcinogenic components into less toxic materials or prevent their biological actions (Gracia and Castillo, 2008). Event Tangeritin and Nobelitin already proved have a role on suppress the apoptosis cancer cell also as antiangiogenetic on CAM (Chrisnanto, et al., 2008).

This research is pre elementary research with the small repetition (five repetition) on each group. The following research should conduct to prove the sufficient of dose that controlled the migration of cell cancer consider preventing the cancer expression, especially for the people who has high risk to express cancer. Either to use the primer human colon cancer cell to explore the real result for human that has high risk of colon cancer.

**CONCLUSION**

The present research is pre elemiary for antimigration inhibition of ethanolic extract Citrus reticulata. The concentration of ethanolic extract Citrus reticulata on 1/2 IC$_{50}$ able to inhibits the migration of WiDr Colon Cancer Cell. Hopefully, the future research will reveals about the mechanism and pathway of this activity.

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