INTRODUCTION

*Tinospora crispa* is a climber plant widely distributed from the Southwestern part of China to Southeast Asia including Indonesia, Malaysia, Vietnam, Thailand, and India. This bitter tasted plant locally known as brotowali [1,2]. It has been used as traditional medicine in Indonesia, Malaysia, and Thailand to treat fever, diabetes, cholera, rheumatism, reducing thirst, hypertension, increasing appetite, and protection from mosquito bites. The crushed leaves are applied on wounds and made into a poultice for itch [1,3]. It was reported that the aqueous extract of *T. crispa* has been showed anti-inflammatory activity using the carrageenan-induced paw edema test [4] and also reported to suppress the synthesis and release of nitric oxide (NO) in lipopolysaccharides (LPS)-stimulated murine macrophages [5]. The methanolic and aqueous extract of *T. crispa* displayed a dose-dependent cytotoxic effect on MCF-7, HeLa, and Caov-3 cell lines with an half-maximal inhibition concentration (IC_{50}) value of 33.6, 165, and 100 µg/ml, respectively [6,7].

From methanol extract of dried pulvered *T. crispa* stems, we have isolated tinocrisposide (C_{36}H_{36}O_{11}), a furano diterpene glycoside with very bitter taste. This work has been reported in Arch Pharm [8]. An anti-inflammatory effect of tinocrisposide has been investigated in vivo in our previous study and reported in 43rd GA-Congress in Halle, Germany [9].

Inflammation is a physiological response of a body to infection or tissue injury [10] which can be caused by different stimuli such as bacterial infection, physical and chemical factors, immunological reactions, and tissue damage [11,12]. Inflammations may be acute or chronic. Acute inflammation is believed to be a defensive mechanism to aid in the killing of bacteria, virus, and/or parasites, eliminate irritants, and maintain normal physiological functions [13,14]. The cardinal signs are characterized by redness (*rubor*), heat (*calor*), pain (*dolor*), swelling (*tumor*), and loss of function [15]. Macrophage plays an important role in the inflammatory response. LPS from the outer membrane of Gram-negative bacteria cell walls activates multiple signal pathways in macrophages such as the nuclear factor-κB pathway that causes inflammation through enhanced cytokine, NO, and prostaglandin production [16-19]. These inflammatory cytokines and mediators play an important role in preventing the spread of infection and also required for repair damaged tissue [20]. However, excessive pro-inflammatory mediators can also promote tissue damage resulting in chronic inflammation [21]. The persistent and prolonged chronic inflammation increases the risk associated with certain generative diseases such as rheumatoid arthritis, atherosclerosis, heart disease, asthma, diabetes, and even cancer [22]. Therefore, finding new anti-inflammatory agents can be a concrete strategy in fighting not only different inflammatory diseases but also cancer.

The anti-inflammatory effect of a new compound or extracts can be conducted by various methods. Uma *et al.* assessed the *in vivo* anti-inflammatory activity of crude extract of *Corbicichona decumbens* by carrageenan- and histamine-induced paw edema method [23]. Soundarajan *et al.* used human keratinocyte cells (HaCaT cells) to identify the inflammatory effect of monocrotophos on cell viability, NO secretion, lactate dehydrogenase release, malondialdehyde release, nuclear changes, reactive oxygen species generation, and cytokine expression [24]. Kusmadi *et al.* investigated the anti-inflammatory effects of pomegranate peel ethanol extract on mice colon. The study
was done through inflammation pathway and it showed reduction in inflammation score on mice model with chronic inflammation induced by dextran sodium sulfate [25]; meanwhile, Kurian et al. explored an anti-inflammatory property of melanin in vitro in RAW 264.7 cell line using cyclooxygenase, lipoxigenase, and cellular nitrite inhibitory assays [26]. The aim of our study was to investigate in vitro the anti-inflammatory activity of tinocrisposide using the LPS-stimulated RAW 264.7 macrophage cells.

METHODS

Chemicals and reagent
Dulbecco’s modified eagle medium (DMEM), fetal bovine serum (FBS), penicillin-streptomycin (PenStrep), trypsin, and 0.4% Trypan blue stain were purchased from Gibco (Invitrogen Corporation, USA). Phosphate-buffered saline (PBS) was purchased from Sigma-Aldrich Chemical Co. (St. Louis, USA). 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) was purchased from Calbiochem, USA. Dimethylsulfoxide (DMSO) was purchased from Fisher Chemical (Thermo Fisher Scientific, USA). LPS and Griess reagent were purchased from Sigma-Aldrich Chemical Co. (St. Louis, USA). Dichloromethane (DCM), n-hexane, acetic acid, ammonia, ethyl acetate, chloroform, and methanol were analytical grade and purchased from Merck, Germany. Silica gel 60 was purchased from Merck Darmstadt, Germany.

Instruments
Inverted Microscope (TC 5400, Meiji Techno, Japan), Microplate reader (Dynatech MR5000 TECAN, Teca Group Ltd., Switzerland), Hemocytometer (Hirschmann, EM Techcolor, Germany), Class II biological safety cabinet (Jouan MSC 12, Thermo Fisher Scientific, USA), Light microscope (Motic B2 series, Motic Asia, Hong Kong), Autoclave (HV-85, Hirayama, Japan), CO2 incubator with a density of 100,000 U/ml, and 1% streptomycin (10,000 µg/ml) in 37°C incubator with 5% of CO2 atmosphere.

Cell viability
The viability of RAW 264.7 cells was assessed by MTT according to Mosmann method [27]. This experiment was performed to elaborate interval concentration of tinocrisposide that exerts no toxic effect to the cells. RAW 264.7 cells were seeded in 96-well flat bottom tissue culture plate with a density of 2×104 cell/well and grown until confluence at 37°C, 5% CO2. Supernatants were discarded, and a complete medium contained various concentration of tinocrisposide solutions in DMSO was then added (3.125–100 µg/ml) into each well. The plate was incubated for 24 h at 37°C, 5% CO2. Supernatants were discarded, and the cell was then washed 2 times with PBS. 20 µl MTT stock solution (5 mg/ml PBS) was added to each well, and the plates were further incubated for 4 h at 37°C, 5% CO2. 100 µl of DMSO was added to each well to dissolve the formed water-insoluble purple formazan crystal. After allowing 1 h, the absorbency of each well was measured with a microplate reader (TECAN) at a wavelength of 570 nm.

Solution of sodium nitrite (NaNO2) reference
6.9 mg of NaNO2 was weighed correctly and transferred into 100 ml volumetric flask, dissolved in 25 ml of distilled water. The volume was made up with distilled water until 100 ml to get a stock solution of NaNO2 with concentrations of 100 µM. A certain volume of stock solution was further diluted with distilled water to get the standard solutions of NaNO2 with concentrations of 100, 50, 25, 12.5, 6.25, and 3.125 µM, respectively. 4 µl of each concentration of standard solutions was mixed with 25 µl of each concentration of standard solutions of NaNO2 with concentrations of 100, 50, 25, 12.5, 6.25, and 3.125 µM, respectively. 25 µl of each concentration of standard solutions was mixed with 25 µl of Griess reagent (0.2% naphthylethenediamine dihydrochloride and 2% sulphanilamide in 5% phosphoric acid), allowed to rest at room temperature for 10 min, and the absorbency of each solution was measured with microplate reader at wavelength, λ 560 nm.

NO determination (Griess assay)
Nitrite accumulation was used as an indicator of NO production in the medium. Released NO (nitrous oxide) to a medium was assayed according to Lee et al. [28] with slight modification. RAW 264.7 cells were plated in 96-well plate with a density of 1×104 cell/well until confluence. Supernatants were discarded and 100 µl medium containing various concentrations of tinocrisposide (1, 5, 25, 50, and 100 µM) was added to the cells. The cells were allowed to rest for 1 h and then incubated with LPS (1 g/ml) for 15 h at 37°C in a humidified atmosphere containing 5% CO2. 25 µl supernatant of each well was transferred into new 96-well plate and was mixed with 25 µl Griess reagent (0.2% naphthylethenediamine dihydrochloride and 2% sulphanilamide in 5% phosphoric acid), allowed to rest at room temperature for 10 min, and then, the absorbency of each solution was measured with microplate plate reader at wavelength of 560 nm.

Experimental procedures

Extraction and isolation
The dried and powdered stem (3.2 kg) of T. crispa was extracted using n-hexane and methanol, successively through maceration process for 3 times for 3 days each at room temperature. The crude methanol extracts (273 g) were dissolved in acetic acid 5% for overnight, decanted, and then partitioned, sequentially, with n-hexane and DCM. Each fraction was dried to yield n-hexane and DCM fractions. The water fraction was basified with ammonia until pH 9 and then partitioned with DCM to get DCM base fraction. The neutral (24.56 g) and base DCM dried fractions (2.64 g) were subjected to column chromatography over silica gel 60 with mobile phase the mixture of DCM and MeOH using step gradient elution system (1–15% of MeOH in DCM). The fractions were combined on the basis of thin-layer chromatography (TLC) monitoring. TLC was carried out using Silica gel F254 plate (Merck) as the stationary phase and the mixture of MeOH:DCM (1:9) as the mobile phase. The fractions, which spots have the same Rv values, were combined and further purified by second-column chromatography on silica gel 60 to obtain pure tinocrisposide as white amorphous powder.

Plant materials
Fresh T. crispa stem samples were collected from Lubuk Minturun, Padang, Indonesia. This plant was identified and authenticated by Dr. Nurainas at the Department of Biology, Faculty of Science, Andalas University. A voucher specimen was deposited in the Herbarium ANDA (No. 178/K-ID/ANDA/V/2015, University of Andalas).

Cell lines
RAW 264.7 mouse macrophage cells were purchased from American Type Culture Collection (ATCC), Virginia, USA.

Basic procedures in cell culture
Cells were cultured in complete DMEM medium containing 10% FBS, 1% penicillin (10,000 U/ml), and 1% streptomycin (10,000 µg/ml) in 37°C incubator with 5% of CO2 atmosphere.
Statistical analysis
All experiments were repeated 3 times (N=3). The results were represented as mean±standard deviation. All experimental data were treated by SPSS program (SPSS version 21, SPSS Institute, Chicago, IL, USA). One-way ANOVA and followed by Dunnet post hoc test were used to examine the difference between groups. A p<0.05 was considered as a statistically significant.

RESULTS AND DISCUSSION
In this study, we used tinocrisposide as a test compound which was isolated from methylene chloride fractions of T. crispa stems. The isolation process was conducted according to our previous method by Pachaly and Adnan [8]. The structure of the compound was identified spectrosopically and confirmed with previous data in our publication.

The compound was namely identified by Fourier-transform infrared (FTIR) and ultraviolet (UV) spectrophotometry and analyzed by gas chromatography–mass spectrometry (GC-MS). Spectral data of the compound were as follows, FTIR (KBR pellet, cm⁻¹): 3411 was for O-H stretch, 1654 was for C=O stretch, 1708 for γ-lactone, 2924, was for Ar-H, 1516 and 1437 were for C=C stretch, 874, 816, and 760 were for C-H bending. UV in MeOH nm: 210 (furan ring), GC-MS fragments (retention time 25000): 83, 98, 238, 310, 327.

RAW 264.7 cell viability
Various concentrations of tinocrisposide were treated with RAW 264.7 cells for 24 h with MTT method to find the appropriate concentrations that exert no toxic and a false-positive effect. This experiments showed that tinocrisposide concentrations of 3.125, 6.25, 12.5, 25, 50, and 100 µM gave % viability cells of 107.4±4.10, 99.09±3.76, 91.32±1.57, 84.40±0.64, 80.00±1.49, and 73.74±1.95%, respectively (Table 1). All tinocrisposide-treated concentrations showed % viability of 264.7 cells higher than 50%. It can be assumed that tinocrisposide was a non-toxic compound against 264.7 RAW cells. Therefore, the tinocrisposide concentration in the inhibition of NO production experiment was adjusted in the interval of 1–100 µM.

Nitrite reference curve
25 µl of NaNO₂ solution from each concentration of 100, 50, 25, 12.5, 6.25, and 3.125 µM were mixed with 25 µl Griess reagent (40 mg/ml) and allowed at room temperature for 10 min. Absorbancy of solutions was measured using microplate plate reader at wavelength, λ 560 nm. Concentrations and absorbency data of experiments were presented in Table 2.

Concentrations and absorbance data of nitrite standard solution were processed with Microsoft Excel Software to get a calibration curve of nitrite solutions as presented in Fig. 2 and a regression equation, y=0.002x+0.055. It showed a linear correlation between nitrite solution in interval concentration of 3.25–100 µM with their absorbance, with coefficient correlation, R²=0.998.

Effect of tinocrisposide on NO production in LPS-stimulated RAW 264.7 cells
Tinocrisposide effect on inhibition of NO production in LPS-stimulated RAW 264.7 cells was assayed by quantification of nitrite accumulation in medium culture using Griess reagent. This method based on diazotization reaction that was firstly described by Griess in 1879. Griess reagent contains sulfanilamide and N/1-naphthyl) ethylenediamine dihydrochloride (NED) in acid condition (H₃PO₄) to quantitate the major metabolites of NO (nitrous oxide) such as nitrite and nitrate, in a variety of biological fluids, such as plasma, serum, urine, and culture medium.

Formed nitrite (NO₂⁻) in medium reacts under acidic condition with sulfanilamide to form a diazonium cation which subsequently couples to the aromatic amine NED to produce red-violet, water-soluble azo dye that can be quantified spectrophotometrically at a wavelength, λ 560 nm. In this experiment, nitrite was released into the supernatant of the medium as a response to inflammation of 264.7 RAW cells that have been stimulated by LPS. RAW cells were treated with various concentrations of tinocrisposide 1, 5, 25, 50, and 100 µM, allowed for 1 h, stimulated with LPS, and incubated for 15 h at 37°C in a humidified atmosphere containing 5% CO₂.

The intensity of azo dye that formed from the reaction of dissolved nitrite in cell supernatant with Griess reagent was quantified with a microplate reader at a wavelength, λ 560 nm. Reading absorbance of samples was calculated using the regression equation of nitrite solution standard that gave a linear correlation between concentrations of nitrite solution and the absorbance of produced red-violet azo dye, to get nitrite concentration in the medium sample and the medium control.

LPS-stimulated 264.7 RAW cells were used as positive control, and 264.7 RAW cell that was not stimulated with LPS and not treated with tinocrisposide was used as negative control. Dexmethasone was used as nitrite inhibitor standard. Our experiment found that treating of tinocrisposide concentration of 1, 5, 25, 50, and 100 µM against LPS-stimulated RAW cells gave nitrite production concentrations of 39.23, 34.0, 28.9, 20.25, and 16.3 µM, respectively, and with % inhibition of 22.67, 33.00, 43.03, 60.1, and 68.00, respectively (Fig. 3).

Anti-inflammatory effect of tinocrisposide was studied by measuring NO production in LPS-stimulated RAW 264.7 cells. Cellular model
Table 1: Effect of tinocrisposide on 264.7 cell viability

<table>
<thead>
<tr>
<th>Concentration (µM)</th>
<th>Average of % cell viability</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.125</td>
<td>107.43±1.10</td>
</tr>
<tr>
<td>6.25</td>
<td>99.09±3.76</td>
</tr>
<tr>
<td>12.5</td>
<td>91.3±1.57</td>
</tr>
<tr>
<td>25</td>
<td>84.6±0.64</td>
</tr>
<tr>
<td>50</td>
<td>80.0±1.49</td>
</tr>
<tr>
<td>100</td>
<td>73.7±1.95</td>
</tr>
</tbody>
</table>

The results were represented as mean±SD, n=3. SD: Standard deviation

Table 2: Absorbance data of nitrite standard

<table>
<thead>
<tr>
<th>Concentration (µM)</th>
<th>Absorbance</th>
<th>Average</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>3.125</td>
<td>0.0591</td>
<td>0.0588</td>
</tr>
<tr>
<td>6.25</td>
<td>0.0676</td>
<td>0.0665</td>
</tr>
<tr>
<td>12.5</td>
<td>0.0853</td>
<td>0.0820</td>
</tr>
<tr>
<td>25</td>
<td>0.1152</td>
<td>0.114</td>
</tr>
<tr>
<td>50</td>
<td>0.1718</td>
<td>0.171</td>
</tr>
<tr>
<td>100</td>
<td>0.2747</td>
<td>0.2756</td>
</tr>
</tbody>
</table>

of LPS-stimulated macrophage is extensively used to measure anti-inflammatory activity. NO production was determined by nitrite (NO₂⁻), which is one of two primary, stable, and non-volatile breakdown products of NO. This assay depends on a diazotization reaction that was originally described by Griess (1879). This reagent uses sulfanilamide and NED under acidic (phosphoric acid) conditions to detect NO₂⁻ in many of biological and experimental liquid matrices such as plasma, serum, urine, and tissue culture medium. NO₂⁻ produced in biological and experimental fluid turns Griess reagent into pink-purple color when mixed together. The intensity of the color is measured and higher intensity of pink-purple color indicates higher NO₂⁻ production. In this study, nitrite was released into cell culture supernatants in response to inflammation.

This study compares NO production of LPS-stimulated cell treated with tinocrisposide with a positive control, a negative control, and a standard drug. Positive control used was cell exposed to LPS only. Since there is no treatment toward the cell, it was expected that the positive control produces the highest NO production. Meanwhile, a negative control of the study is untreated cell without LPS induction. The untreated cell was expected to produce lowest NO since there was no exposure to the inflammatory stimulus. Dexamethasone, a well known for its anti-inflammatory activity was used as a standard drug with exposed LPS. Results of this study showed that tinocrisposide able to reduce NO production in LPS-induced inflammatory in RAW 264.7 cells in a dose-dependent manner. During inflammatory process, upregulation of inducible NO synthase (iNOS) genenates NO which is one of the pro-inflammatory mediators. Upregulation of iNOS can be contributed by inflammatory stimuli such as cytokines, IL, and bacterial endotoxin. This study stimulates the RAW 264.7 cells using LPS. LPS is bacterial endotoxin that found in the outer membrane of Gram-negative bacteria cell walls. Thus, in response to inflammatory stimuli, high level of NO is generated [28]. Exposure of RAW 264.7 cells to LPS upregulates iNOS and pre-treatment of the cell with varying compound concentration (1, 5, 25, 50, and 100 µM) before LPS induction was able to reduce the NO production.

Tinocrisposide used was able to decrease NO production even at low concentration when NO production is compared against the positive control. At high concentration used (100 µM), NO produced is 16.3 µM and dexamethasone is 13.68 µM, this means that there was no significant difference between treated and reference. This suggests that compound possesses good potential as an anti-inflammatory agent. This was based on trend seen in NO generated. Meanwhile, the effect to dose used in this study did not affect cell viability since the test dose chosen (1, 5, 25, 50, and 100 µM) was based on cell viability study conducted before NO determination study. Therefore, the NO reduction effect observed is based on tinocrisposide action and not due to a reduction in the cell population.

MTT assay was conducted to determine the cytotoxic effects of tinocrisposide on RAW 264.7 cells. The cell lines were incubated at various concentrations for 24 h, and cytotoxicity was determined using a microplate reader. Meanwhile, the IC₅₀ value was determined from the linear regression of the experimental data. MTT assay widely employed in cell biology for the study of growth factor, cytokines, and cytotoxicity of chemotherapeutic agents since it provides a simple and quantitative method for cell population’s assessment in response to external factors exposure [29].

Based on the results of one-way ANOVA, the overall p<0.05 (p=0.00) this conclude that there were statistically significant differences between groups. The Dunnett’s test is used to compare one control group against several treatment groups. In the Griess Assay, tinocrisposide at a concentrations of 50 and 100 µM showed p<0.05 (p=0.249 and 0.898, respectively); this means that there were no statistically significant differences between treatment group and dexamethasone. In the cytotoxic assay against RAW 264.7 cells, tinocrisposide at concentration 3.125 µg/ml also showed p>0.05 (p=0.134 and 0.241), this means that there were no statistically significant differences between treatment group and control.

CONCLUSION

In summary, the present study showed that tinocrisposide, an isolated compound of T. crispa stem, decreased NO production in a dose-dependent manner, where the significant inhibition was shown at concentration of 50 and 100 µM which NO production was 20.25 and 16.3 µM, respectively, and with inhibition of 60.1% and 68%, respectively. This result indicated that there was no significant difference between inhibition data of sample with the data of dexamethasone standard.

ACKNOWLEDGMENTS

This work was supported by Rector of Andalas University to enhance research activity at Faculty of Pharmacy.

AUTHOR CONTRIBUTIONS


CONFICT OF INTEREST

None.

REFERENCES

6. Froemming GA. Antiproliferative and antioxidant effects of Tinospora


