Identification of a new plant extract for androgenic alopecia treatment using a non-radioactive human hair dermal papilla cell-based assay

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Abstract

Background: Androgenic alopecia (AGA) is a major type of human scalp hair loss, which is caused by two androgens: testosterone (T) and 5α-dihydrotestosterone (5α-DHT). Both androgens bind to the androgen receptor (AR) and induce androgen-sensitive genes within the human hair dermal papilla cells (HHDPCs), but 5α-DHT exhibits much higher binding affinity and potency than T does in inducing the involved androgen-sensitive genes. Changes in the induction of androgen-sensitive genes during AGA are caused by the over-production of 5α-DHT by the 5α-reductase (5α-R) enzyme; therefore, one possible method to treat AGA is to inhibit this enzymatic reaction.

Methods: RT-PCR was used to identify the presence of the 5α-R and AR within HHDPCs. A newly developed AGA-relevant HHDPC-based assay combined with non-radioactive thin layer chromatography (TLC) detection was used for screening crude plant extracts for the identification of new 5α-R inhibitors.

Results: HHDPCs expressed both 5α-R type 1 isoform of the enzyme (5α-R1) and AR in all of the passages used in this study. Among the thirty tested extracts, Avicennia marina (AM) displayed the highest inhibitory activity at the final concentration of 10 μg/ml, as the production of 5α-DHT decreased by 52 % (IC50 = 9.21 ± 0.38 μg/ml).

Conclusions: Avicennia marina (AM) was identified as a potential candidate for the treatment of AGA based on its 5α-R1-inhibitory activity.

Keywords: Androgenic alopecia, 5α-reductase inhibitory activity, Human hair dermal papilla cell-based assay, Anti-androgenic activity, Avicennia marina

Background

Androgenic alopecia (AGA) is a major type of scalp hair loss in both males and females that is becoming a worldwide issue. AGA is caused by androgens, namely, testosterone (T) and 5α-dihydrotestosterone (5α-DHT). T is the major circulating androgen and is converted to 5α-DHT by the 5α-reductase enzyme (5α-R) [EC 1.3.99.5], which is present in two isoforms i.e., 5α-reductase type 1 (5α-R1) and 5α-reductase type 2 (5α-R2) [1–6]. Both androgens bind to the androgen receptor (AR), forming a receptor-ligand complex that is translocated to the nucleus, where it acts as a transcription factor in regulating androgen-sensitive genes [7]. However, 5α-DHT exhibits five times higher binding affinity and 10-fold higher potency than T in inducing androgen-sensitive genes. The products of these androgen-sensitive genes act as growth factors, affecting hair growth [3, 5, 7–17]. During AGA, overproduction of 5α-DHT causes the down-regulation of growth factors, which results in the shortening of anagen/growth phase. This shortening causes the miniaturisation of large, thick-pigmented terminal hair to small, fine, unpigmented vellus hair with a diameter of less than 0.03 mm [1, 2].

Currently, minoxidil and finasteride are the two FDA-approved synthetic drugs used to treat AGA [18].
Minoxidil, a vasodilator and potassium channel opener, prolongs the anagen phase and converts vellus hair to terminal hair [19, 20]. However, minoxidil is only effective on 30-35 % of patients, and the treatment must be continued for life [18]. In addition, the side effects of 2 % and 5 % minoxidil solution include scalp irritation, pruritus, dryness, scaling, itchiness, redness, contact dermatitis and hypertrichosis [21, 22]. Finasteride, a synthetic azo-steroid, is a 5α-α-R2 inhibitor that binds irreversibly to the enzyme and inhibits the conversion of T to 5α-DHT, thereby reducing the serum 5α-DHT concentration by 68 % [3]. However, finasteride is effective in only 48 % of the patients [21] and has the observed side effects of impotence, abnormal ejaculation, abnormal sexual function, myalgia, tendinous pain and gynecomastia [23].

Because only two synthetic drugs are currently available, a search for new drugs to treat AGA is necessary. Among various sources, natural products provide an abundance of diverse chemical structures, representing a rich source for lead structures in drug development [18]. Recent studies have primarily focused on medicinal plants, and many natural product groups have been demonstrated to have 5α-R inhibitory activity. These include a sterol from Cuscuta reflexa [24]; fatty acids, including oleic, lauric, myristic and linoleic acid from Serenoa repens [25]; a triterpenoid, ganoderic acid from Ganoderma lucidum [26]; the saponins soyasaponin I and kaikasaponin III from Pueraria thomsonii [27]; and a catechin, epigallocatechin-3-gallate [EGCG] from Camellia sinensis [28]. In addition, various well-known flavonoids, such as myricetin, quercitin, alizarin, kaempferol, genistein and daidzein, have also been reported to exhibit inhibitory activity [28]. However, none of these reports have used dermal papilla cells for either their cell-based or enzyme-based (source of 5α-R) assay systems. Dermal papilla cells are a primary cell type that regulate hair growth, express 5α-R and are the only site of androgen action within the hair follicle [1, 18]. Therefore, the inhibition of 5α-R activity in this specific type of cells should have a direct effect on AGA treatment. Therefore, to obtain an inhibitory assay system for this type of hair loss, it is most relevant and reliable to use hair cells, specifically human hair dermal papilla cells (HHDPCs).

In this study, a non-radioactive HHDPC-based assay system, which is highly relevant to the AGA type of human scalp hair loss, was used to screen plant extracts for 5α-R inhibitory activity. The most potent crude extract was further tested for the anti-androgenic activity to evaluate its potential to overcome the effects of androgens on the expression of the aforementioned growth factors.

**Methods**

**Plant materials and extraction**

The list of the Thai plants and plant parts used in this study is shown in Table 1. All plant materials were purchased from of Chao Krom Poe Dispensary in Bangkok, Thailand. Voucher specimens of the plants were deposited in the herbarium collection of Thailand’s Forestry Department with the following specimen numbers (SNs): Avicennia marina, SN145917; Micromelum minutum, SN055802; Diospyros mollis, SN142793; Alternanthera sessilis, SN097887; Salacia verrucosa, SN099212; Crotalaria retusa, SN043656; Senna timoriensis, SN094415; Dalbergia parviflora, SN200944; Aegle sericea, SN124674; Bacopa monnieri, SN176524; Tarenna hoaensis, SN176524; Pterygota alata, SN059871; Scoparia dulcis, SN163093; and Senna garrettiana, SN008620. For preparation of crude extracts, the plant part used for each species was ground into powder and extracted through maceration using 100 % methanol at room temperature for two days. The methanolic extracts were then evaporated to dryness at 45 °C using a rotary evaporator (Buchi, Switzerland) and kept at -20 °C until used.
All of the organic solvents used were analytical grade and purchased from RCI Labscan (Bangkok, Thailand). Ultrapure grade dimethyl sulfoxide (DMSO) was purchased from Ameresco® (Framingham, USA). Testosterone (T) and 5α-dihydrotestosterone (5α-DHT) were purchased from Sigma-Aldrich (St. Louis, USA). Dutasteride was purchased from BDG Synthesis (Wellington, New Zealand). Agarose-LE was purchased from Affymetrix (Santa Clara, USA). Mesenchymal stem cell medium and its supplements were purchased from Sciencell Research Laboratories (Carlsbad, USA). Foetal bovine serum, 100X antibiotic-antimycotic solution, 10X PrestoBlue®, RPMI medium, 50X Tris-acetate-EDTA (TAE) buffer, 0.25 % trypsin-EDTA and Platinum® Taq polymerase were purchased from Invitrogen (Grand Island, USA). A GeneRuler 1-kb DNA ladder was purchased from Thermo Fisher Scientific (Pittsburgh, USA). RNeasy® mini kits were purchased from Qiagen (Valencia, USA). DNase I enzyme, first-strand cDNA synthesis kit, dATP, dTTP, dCTP and dGTP were purchased from Fermentas (Walthan, USA).

Culturing of human hair dermal papilla cells
HHDPCs, obtained from Sciencell Research Laboratories (Carlsbad, USA), were grown in mesenchymal stem cell medium containing 5 % foetal bovine serum (FBS), mesenchymal stem cell medium supplement and 1X antibiotic-antimycotic solution at 37 °C in 5 % CO₂. The cells between passages 2 to 6 were used in this study.

Evaluating the presence of 5α-R and AR in HHDPCs
Reverse-transcriptase polymerase chain reaction (RT-PCR) was used to identify the type of 5α-reductase enzymes (i.e., 5α-R1 and/or 5α-R2) and AR, expressed in passages 2, 4, 5 and 6 of HHDPCs. Total RNA was extracted from HHDPCs according to the manufacturer’s instructions for the RNeasy® mini kit. The RNA was subsequently treated with DNase I to remove any genomic DNA contamination.

### Table 1
List of Thai medicinal plants used in this study

| Medicinal plant            | Abbreviation | Family       | Plant part   |
|----------------------------|--------------|--------------|--------------|
| Afgekia sericea            | AS           | Fabaceae     | Aerial       |
| Alpinia galangal           | AG           | Zingiberaceae| Fresh Rhizome|
| Alternanthera sessils      | ASHE         | Amaranthaceae| Whole plant  |
| Avicennia marina           | AM           | Acanthaceae  | Heartwood    |
| Bacopa monnieri            | BM3          | Plantaginaceae| Aerial     |
| Balanophora abbreviatie    | BA           | Balanophoraceae| Aerial |
| Barleria cristata          | BC           | Acanthaceae  | Root         |
| Butea monosperma           | BM           | Fabaceae     | Stem         |
| Centella asiatica          | CA           | Mackinlayaceae| Aerial      |
| Citrus limonum             | CL           | Rutaceae     | Fresh Peel   |
| Crotonis retusa            | CR           | Fabaceae     | Root         |
| Dalbergia parviflora       | DP           | Leguminosae  | Heartwood    |
| Derris elliptica           | DE           | Leguminosae  | Stem         |
| Diospyros mollis           | DM           | Demberaceae  | Stem         |
| Kaempferia galangal        | KG           | Zingiberaceae| Dried Rhizome|
| Leersia hexandra           | LH           | Poaceae      | Stem         |
| Macleia cochinchen         | MC           | Moraceae     | Stem         |
| Micromelum minutum         | MM           | Rutaceae     | Stem         |
| Moriella mucofilla         | OM           | Oleandraceae | Stem         |
| Pterygota alata            | PA           | Malvaceae    | Stem         |
| Randia hondala             | RH           | Rubiaceae    | Root         |
| Salacia verrucosa          | SV           | Celastraceae | Stem         |
| Scoparia dulcis            | SD           | Plantaginaceae| Stem       |
| Senna garretiana           | SG           | Fabaceae     | Heartwood    |
| Senna timoriensis          | ST           | Fabaceae     | Stem         |
| Tarenna fragans            | TF           | Rubiaceae    | Leaves       |
| Tarenna hoaensis           | TH           | Rubiaceae    | Stem         |
| Telosma minor              | TM           | Asclepiadaceae| Stem      |
| Zanthoxylum limonella      | ZL           | Rutaceae     | Stem         |
| Zingiber officinale        | ZO           | Zingiberaceae| Fresh Rhizome|

### Table 2
Forward and reverse primers and expected sizes of AR, 5α-R enzymes and β-actin

| Name                      | Primer pair                                      | Expected size (bp) |
|---------------------------|--------------------------------------------------|--------------------|
| Androgen receptor (AR)    | F: 5' CGTGCGCGAAGTGATCCAGAA 3' \* R: 5' TGCCTGTGGTCTGACAGA            | 811                |
| GenBankNM_0000443.3       | R: 5' TGCCTGTGGTCTGACAGA \* R: 3'                | 380                |
| 5α-Reductase type 1 (5α-R1) | F: 5' ACTGATCTCTCTGCAGTCTG 3' \* R: 5' GGTGAGCTGGCCACCTGATGG 3' | 450                |
| GenBankNM_001047.2        | F: 5' ACTGATCTCTCTGCAGTCTG \* R: 3'              | 584                |
| 5α-Reductase type 2 (5α-R2) | F: 5' GGTGAGCTGGCCACCTGATGG \* R: 3'            |                    |
| GenBankNM_000348.3        | R: 5' GGTGAGCTGGCCACCTGATGG \* R: 3'            |                    |
| -actin (internal control) | F: 5' ATGATGATATCGCCGCGCTC 3' \* R: 3'          |                    |
| GenBankNM_001101.3        | R: 5' ATGATGATATCGCCGCGCTC \* R: 3'            |                    |
Complementary DNA (cDNA) was synthesised from the treated RNA using the first-strand cDNA synthesis kit. The cDNA obtained from this reaction was used as a DNA template for PCR reactions. The PCR reaction comprised of 1X PCR buffer, 2 mM MgCl₂, 0.4 mM dNTP mix, 0.4 μM each of forward and reverse primer and 2.5 units of Platinum® Taq polymerase. The forward and reverse primers for the two isoforms of 5α-R (5α-R1 and 5α-R2) and AR, shown in Table 2, were designed based on the protein region of the full-length sequence obtained from the NCBI GenBank using Clone manager (Scientific & Educational Software, USA) and made to order at 1st Base Laboratories (Selangor, Malaysia). The PCR-thermal profile started with an initial denaturation at 94 °C for 3 min, followed by 35 cycles of denaturation at 94 °C for 30 s, annealing for 30 s at 52 °C, and extension at 72 °C for 2 min, followed by a final extension at 72 °C for 10 min. The PCR products were analysed using 1 % agarose gel electrophoresis.

Cytotoxicity testing of methanolic plant extracts on HHDPCs
For the cytotoxicity test, HHDPCs were seeded at a cell density of 1x10⁵ cells/ml onto 96-well plates (100 l of 10,000 cells/well). After 24 h, the cells were separately treated with 100 l of 5, 10, 20, and 40 μg/ml of each extract to obtain the final concentrations of 2.5, 5, 10 and 20 g/ml, respectively, and 1 % DMSO (control). Cell viability was measured at 24 h after treatment using 1X PrestoBlue® reagent in RPMI medium. In the presence of viable cells, PrestoBlue® changes from a non-fluorescent blue colour to a fluorescent purple-pink colour, detected using Multimode Detector DTX 880 (Beckman Coulter®, Indianapolis, USA), a bottom-read fluorospectrophotometer with excitation/emission wavelengths of 535 and 615 nm, respectively. The highest final non-toxic concentration of each extract was then used for further studies.

Screening of plant extracts for 5α-R inhibitory activity using HHDPC-based assay combined with non-radioactive TLC detection
HHDPCs were seeded at a cell density of 1x10⁵ cells/ml onto 96-well plates (100 l of 10,000 cells/well). After 24 h, the cells were separately treated (in a total volume of 200 l) with the final concentrations of 0.1 mM T and 0.5 % DMSO (internal control), 0.1 mM T and 0.25-20 g/ml (non-toxic concentrations of the methanolic plant extracts), and of 0.5 % DMSO as negative control. After 48 h, the culture medium of each treatment was collected in Eppendorf tubes, and the attached treated cells were tested for cell viability using the 1X PrestoBlue® reagent in RPMI medium. The remained substrate T and the product 5α-DHT formed from the activity of cellular 5α-R were extracted from the culture medium using an equal volume of ethyl acetate. The ethyl acetate layer was dried and reconstituted using 20 l of methanol and spotted onto a TLC Silica gel 60 F₂₅₄ aluminium plate (Merck, Darmstadt, Germany). The TLC plate was developed using toluene:acetone (8:2) as the mobile phase [9]. The developed TLC plate was dipped briefly in a solution of 42.5 % phosphoric acid and heated at 120 °C for 20 min for the visual detection of 5α-DHT at 366 nm using a TLC Reprostar Imager (Camag, Switzerland), and the amount 5α-DHT formed was quantitated by scanning the product band of 5α-DHT at 366 nm using TLC Densitometer Scanner 3 (Camag, Switzerland).
Switzerland) to obtain its value of peak area which was then converted to the amount of 5α-DHT using a calibration curve. The calibration curve of 5α-DHT was constructed by plotting peak areas and various amounts of 5α-DHT which showed linearity from 50 to 500 ng with the coefficient of determination ($r^2$) value of 0.9874. The extract of AM displaying the highest potential in inhibiting the 5α-R enzyme was studied further to determine its effects on androgen-sensitive genes.

**Statistical analysis**
All of the experiments were performed in triplicate, and the data are shown as the means ± SD. One-way ANOVA statistical analysis was used, and a $P$-value <0.05 was considered to be statistically significant.

**Results**
**Expression of 5α-R1 and AR in HHDPCs**
The effect of androgens on hair growth is exerted through the interactions with 5α-R and AR in HHDPCs. Therefore, the presence of both the enzyme and the receptor is important for this study, and expression in HHDPCs was evaluated. The RT-PCR analysis revealed that the genes of the Type 1 enzyme, 5α-R1, and AR were both expressed in passages 2, 4, 5 and 6 of HHDPCs, whereas the Type 2 gene, 5α-R2, was not expressed in any of the passages (Fig. 1). β-actin, used as an internal control, was constantly expressed in all passages.

**Cytotoxicity of methanolic plant extracts on HHDPCs**
To obtain a suitable starting concentration for screening methanolic plant extracts for 5α-R inhibitory activity,
the cytotoxicity of each plant extract on HHDPCs at various concentrations was subsequently tested using Prestoblu{}, cell viability reagent. The results revealed that the extracts, at their final concentrations, exhibited different toxic levels on the cells (Fig. 2). Above the cell viability of 85 %, the extracts were considered to be non-toxic. The plant extracts of DM and SG were the most toxic, exhibiting toxicity above 2.5 g/ml, followed by the ST, ASC and DP extracts, which displayed toxicity above 5 g/ml (Fig. 2a). The other extracts with moderate toxicity above 10 g/ml were SD, OM, AM and BC (Fig. 2b), whereas the remaining extracts, including TF, CA, AS, LH, MF and BM, displayed no toxicity even at 20 g/ml (Fig. 2c). Based on these results, the highest final non-toxic concentration of each extract was used for the subsequent inhibitory activity analyses.

**Identification of plant extracts possessing 5α-R1 inhibitory activity using an HHDPC-based assay combined with non-radioactive TLC detection**

As previously mentioned, AGA is induced through the over-production of 5α-DHT, and the enzyme responsible for the conversion of T to 5α-DHT within HHDPCs is 5α-R. Thus, one potential method for reducing the effect of the androgens on hair growth is to inhibit this enzymatic reaction. HHDPCs expressed only 5α-R1, and therefore, the assay system was tested using a well-known 5α-R1 inhibitor, dutasteride, as a positive control. To obtain the starting concentration for testing the inhibitory activity, first, the cytotoxicity of dutasteride for HHDPCs was determined. It was observed that dutasteride exhibited toxicity (i.e., cell viability less than 85 %) above the final concentration of 10 g/ml (Fig. 3a). At this concentration, the cell-based assay displayed complete 5α-R1 inhibitory activity (Fig. 3b). Lowering the concentration of dutasteride to 0.01 and 0.001 g/ml, the assay displayed the 5α-R1 inhibition of 90.5 and 16.5 %, respectively, with an IC50 value of 0.005 g/ml or 9.7 nM (Fig. 3b). The assay system was then used to screen the methanolic plant extracts for 5α-R1 inhibitory activity. Using the highest final non-toxic concentration (Table 3) in the incubation mixture, each individual extract was evaluated for this activity. As shown in Figs. 4a-4c, each lane represents the inhibitory potential of each extract at its highest final non-toxic concentration. The lane “Cell + T” was the internal control for the conversion of T to 5α-DHT, and the lane “Cell- T” was the negative control. Among all extracts screened, the plant extracts of BM, DM, OM, MC, BC and BA displayed only 10 % inhibition, whereas the extracts of KG and MM displayed approximately 20 % inhibition. The highest inhibitory activity was observed from the crude extract of AM at a final concentration of 10 g/ml through the reduction in 5α-DHT formation by more than 50 % (Table 3). The IC50 value of AM was determined to be 9.21 ± 0.38 g/ml.

**Discussion**

This study focuses on identifying new 5α-R inhibitors specifically for treating AGA using a newly developed HHDPC-based assay system. A cell-based assay was chosen over a cell-free assay, where the 5α-R enzyme has been
reported to be isolated from the rat liver [9, 26, 29], prostate [23, 30], epididymis [27] or the human prostate [24], as the latter does not mimic the actual conditions of the enzymes and does not take into account the toxicity of the test compounds [28]. In addition, the reliability of the cell-free system is presumably low because the sequence similarities between the 5α-R1 and 5α-R2 from rats and humans are only 61 % and 75 % [6], respectively, which might affect the nature of active site formation and hence the activity. In addition, two isoforms of the enzymes, 5α-R1 and 5α-R2, are specifically distributed in different organs within the human body [6], and therefore, testing the enzyme isolated from the human prostate [24] might not be relevant to the type of enzyme present in human hair cells.

To obtain a cell-based assay system relevant for hair loss, it is more appropriate to use hair cells than transfected rat [28] or insect [25] cell lines. These cell lines of rats, insects and humans possess different levels of toxicity tolerance to the test compounds, such that a potent concentration of 5α-R inhibitor for the rat or insect cell line might be toxic to human hair cells. Hair comprises
at least fifteen distinct types of cells where the dermal papilla cell is a primary cell type that regulates hair growth, expresses 5α-R and is the only site of androgen action [1, 12, 18]. Therefore, human hair dermal papilla cells (HHDPCs) were used as a model for the identification of potent 5α-R inhibitors in this study. After 48 h of T treatment on HHDPCs in our optimised incubation mixture, the enzymatic product, 5α-DHT, was detectable and measurable on a TLC plate at 366 nm (Fig. 4) by simply dipping the developed TLC plate into 42.5 % phosphoric acid solution, heating it at 120 °C for 20 mins and quantitatively analysing it using Camag Scan-3 TLC densitometer. The sensitivity of this TLC detection is in the range of ng level (from 25 to 250 ng). Under the normal condition of our cell-based assay which is in the presence of 5.8 g of T (0.1 mM final concentration), it appears that HHDPCs (10,000 cells) present in the incubation mixture can convert T into 87 ng of 5α-DHT. In the presence of a positive crude extract such as AM, the amount of 5α-DHT formed was reduced to 42 ng, equivalent to 48 % of the conversion (Table 3), which is still in the detectable range.

Therefore, with the high sensitivity of detection, there is no need to use radioactively labelled T and complicated detectors as previously reported, such as a radioactive image analyser [26, 30], TLC radioactive scanners [28], HPLC radioactive detectors [25] or measuring the decrease in radiolabelled T concentration at 254 nm using HPLC [9, 23, 24, 27].

The RT-PCR results revealed the expression of only 5α-R1 (Fig. 1), which is consistent with the results of previous studies [6, 31–33]. Therefore, the non-radioactive HHDPC-based assay system was first evaluated with a positive control, dutasteride, which is a well-known specific 5α-R1 inhibitor [6]. The compound demonstrated complete inhibition of the enzyme activity at a final concentration of 10 g/ml with an IC50 value of 9.7 nM (Fig. 3). The assay system was then used for screening thirty methanolic plant extracts for 5α-R1 inhibitory activity. Among the tested extracts (Fig. 4a-c), it appeared that only the heartwood methanolic extract of Avicennia marina (AM) at the final concentration of 10 g/ml exhibited the highest potential for inhibiting the enzyme activity, as the 5α-DHT production was reduced to 48 % (Table 3). The rest of the crude extracts were found to have low or negative inhibitory effect (i.e., ranging from -10 to 20 %) compared with the internal control. To avoid false positive results, the attached treated cells in the 96-well plate were tested for their viability. The cells treated with 10 g/ml of AM and 10−4 M of T for 48 h displayed 100.5 ± 2.02 % (n = 3) viability relative to the internal control, confirming the positive effect of the AM extract.

For information about Avicennia marina (AM), this plant is commonly called grey or white mangrove. It is a species of mangrove trees belonging to the Acanthaceae family and has been traditionally used in Egypt to cure skin diseases [34]. Phytochemically, terpenoids and steroids, such as lupeol, botulin, -sitosterol and betulinic acid, have been identified from the bark of AM [34]. Whether any of these compounds are the active 5α-R1 inhibitors is still unknown. Therefore, activity-guided fractionation is currently being conducted to identify the active compounds in the AM extract.

Conclusions
The use of a non-radioactive HHDPC-based assay system in this study has led to the identification of Avicennia marina (AM) as a new potential candidate for the treatment of AGA.

Abbreviations
AGA: Androgenic alopecia; T: Testosterone; So-DHT: 5α-Dihydrotestosterone; AR: Androgen receptor; HHDPCs: Human hair dermal papilla cells; So-R: 5α-reductase; So-R1: 5α-reductase type 1; So-R2: 5α-reductase type 2; TLC: Thin layer chromatography; RT-PCR: Reverse-transcriptase polymerase chain reaction. The abbreviations of all plant names used in this study are shown in Table 1.

Competing interests
The authors declare that they have no competing interests.

Authors' contributions
RI: Experimental design and conduct, data analysis and manuscript preparation. OM: Preparation of plant crude extracts. PT: Co-supervisor of RJ, giving comments and suggestions. WD: Main supervisor of RJ, project leader, giving comments and suggestions and manuscript preparation. All authors read and approved the final manuscript.

Acknowledgments
RI would like to thank Chulalongkorn University for a Ph.D. scholarship under the Chula Dusadee Pipat Project. This study was financially supported through the PERDO’s Center of Excellence on Medical Biotechnology (CEMB) program.

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Received: 30 October 2013 Accepted: 19 January 2016

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