Transformation of Amorphadiene Synthase and Antisilencing P19 Genes into Artemisia annua L. and its Effect on Antimalarial Artemisinin Production

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Abstract

Purpose: The low content of artemisinin related to the biosynthetic pathway is influenced by the role of certain enzymes in the formation of artemisinin. The regulation of genes involved in artemisinin biosynthesis through genetic engineering is a choice to enhance the content. This research aims to transform ads and p19 gene as an antisilencing into Artemisia annua and to see their effects on artemisinin production.

Methods: The presence of p19 and ads genes was confirmed through polymerase chain reaction (PCR) products and sequencing analysis. The plasmids, which contain ads and/or p19 genes, were transformed into Agrobacterium tumefaciens, and then inserted into leaves and hairy roots of A. annua by vacuum and syringe infiltration methods. The successful transformation was checked through the GUS histochemical test and the PCR analysis. Artemisinin levels were measured using HPLC.

Results: The percentages of the blue area on leaves by using vacuum and syringe infiltration method and on hairy roots were up to 98, 92.55%, and 99.00% respectively. The ads-p19 sample contained a higher level of artemisinin (0.18%) compared to other samples. Transformed hairy root with co-transformation of ads-p19 contained 0.095% artemisinin, where no artemisinin was found in the control hairy root. The transformation of ads and p19 genes into A. annua plant has been successfully done and could enhance the artemisinin content on the transformed leaves with ads-p19 up to 2.57 folds compared to the untransformed leaves, while for p19, co-transformed and ads were up to 2.25, 1.29, and 1.14 folds respectively.

Conclusion: Antisilencing p19 gene could enhance the transformation efficiency of ads and artemisinin level in A. annua.

Introduction

Malaria has been a serious problem in the world. In 2016, there have been more than 133 million people infected by Plasmodium spp, a malarial caused parasite, about 445000 of them died. The use of antimalarial drugs, such as chloroquine, tends to be reduced because of drug resistance so that more effective drugs for malaria disease are needed. WHO has recommended the ACTs (artemisinin-based combined therapies) as a choice for treatment of malaria. Artemisinin, a sesquiterpene produced by Artemisia annua L. has an excellent effect on malaria in multi-drug resistant Plasmodium strains. Artemisinin together with its derivatives, especially dihydroartemisinin and artesunate, was reported to have good activity against P. falciparum.

Recently, artemisinin has been reported to have potential effects for systemic lupus erythematosus. It could ameliorate renal damage, reduce the symptoms, and increase antibodies as well as proteinuria. To date, A. annua is the only source for artemisinin with a low yield. Because of its unique complex structure, the chemical synthesis is difficult, and it becomes less prospective. Other approaches to enhance the production of artemisinin are through cell culture and genetic engineering for the key enzymes of artemisinin biosynthesis in plant cell and yeast. Cell culture technique has advantages as an alternative system for recombinant pharmaceuticals. Farnesyl pyrophosphate is a precursor of artemisinin derivative biosynthesis. It is synthesized from one isoprenoid unit derived from the non-mevalonate pathway and two C-5 isoprenoid units derived from the mevalonate pathway in the cytosol. Farnesyl pyrophosphate is used by amorpha-4,11-diene synthase (ads) as a precursor to produce cyclic amorpha-
Enzymes coded genes which have the key roles in the artemisinin biosynthesis have been cloned. Therefore, the enhancement of artemisinin production can be performed, using genetic engineering of these genes, and transform them into plants or microbes. Transient expression system of a gene in plants using agrofiltration has been developed as an alternative to optimize protein expression. Agro-infiltration has a flexible nature in the production of recombinant proteins in plant tissue and only need few days to get the results.

Transient expression system with plant virus vector via Agrobacterium-mediated transformation has been performed for the production of recombinant protein with a high level and short time. The bacterium infects the plant cells and integrates a region of a large tumor-inducing (Ti) plasmid resident in Agrobacterium into the plant's nuclear genome. An Ads gene-encoded amorpha-4,11-diene synthase, which is a key enzyme in artemisinin biosynthesis, has been transformed using vector pCAMBIA1303 resulting in plasmid pCAMBIA 1303-ads. The plasmid has been transformed into A. annua. A. tumefaciens strain AGL1, which is the most efficient transformation among others with up to 70.91% from the total explants of A. annua leaves. Although genetic transformation has been successfully done in plants, DNA of A. tumefaciens may activate the protection response in the plants, also called RNA silencing. Post-transcriptional gene silencing (PTGS) or RNA silencing is a natural protective response of plants from foreign nucleic acids, such as viral infection and transgene expression in plant cells, which can invade plants. In this process, the double-stranded, short-interfering RNA is cleaved from single-stranded RNA (ssRNA) and double-stranded RNA (dsRNA) or viral sequences by plant RNAse III-type.

The existence of PTGS will destroy the RNA of A. tumefaciens infected the plants so that the DNA transfer process in A. tumefaciens to the plants is not maximal. However, several plant viruses have the silencing suppressors which can inhibit the protection mechanism of plants. One of silencing suppressors is p19 gene from tomato bushy stunt virus. The purpose of this research is to evaluate the effect of a P19 gene in recombinant A. annua containing amorpha-4,11-diene synthase.

Materials and Methods

There are a Luria-Bertani (LB) medium containing NaCl 1%, tripton 1%, yeast 0.5%, bacto agar 1.5%, and a liquid LB medium without bacto agar as a growing medium. TAE 1X (dilution from TAE 50X (Tris base 24.2% (Promega), acetic acid glacial 5.71 %, EDTA 0.5 M pH 8.0 10%)); DNA 1 kb ladder (Fermentas); agarose (Top vision); DreamTaq Green MM (Fermentas); forward primers (5’-AAA CTC GAG ATG GAA CGA GCT ATA CAA G-3’), reverse primer (5’-AAA CTC GAG TTA CTC GCC TTT TTT TTT G-3’), Reagent for plasmid isolation containing solution 1 (glucose 50 mM, TrisCl 25 mM, EDTA pH 8 10mM, deionized up to 100 %), solution 2 (NaOH 0.2 N, SDS1 % w/v), and solution 3 (sodium acetate 5 M, acetic acid glacial), isopropanol, ethanol, and TE-RNase.

The preparation of p19 gene, plasmid pCAMBIA 1303 and pCAMBIA 1303-ads

Ads and synthetic p19 genes were confirmed using polymerase chain reaction (PCR) with the composition of PCR reaction consisting of 12.5 μL of Dream Tag DNA polymerase, 1.25 μL of each forward and reverse primer as well as DNA template with the total volume of 25 μL after adding free-water nuclease. PCR products were detected using agarose gel 1.5% and then purified using a gel purification kit.

Plasmid pCAMBIA 1303 and pCAMBIA 1303-ads were cut by Xho1 and incubated at 37°C for 9 minutes. Reactions were done at 37°C for 16 hours. Restriction results were checked by electrophoresis with agarose gel 1% for 30 min with 100 voltage. Bands with the size above 10000 bp were purified. The p19 gene whose sequence was confirmed was cut from pGEM-T Easy using Xho1. Bands with a size of 527 bp were purified. Pure p19 genes were measured of their concentrations and ready to be continued for transformation.

The ligation of p19 gene into plasmid pCAMBIA 1303 and pCAMBIA 1303-ads

P19 gene was ligated into plasmid pCAMBIA 1303 and pCAMBIA 1303-ads, having been cut by Xho1. The composition of ligation reaction consists of 1 μL of T4 DNA Ligase (0.03 unit), 1.6 μL of 10X Rapid ligation buffer, 6 μL of Plasmid CAMBIA 1303/pCAMBIA 1303-ads (24 ng), 4 μL of p19 gene (16 ng) and nuclease-free water until the final concentration of 16 μL. Ligation was done at 4°C for 16 hours. Ligation reactions were transformed into E. coli DH5α in a LB medium containing kanamycin 50 ppm and incubated at 37°C overnight.

Plasmid isolation

A single colony of transformants was selected and suspended in a liquid LB medium containing kanamycin 50 ppm and incubated in the shaker 200 rpm, 37°C overnight. Plasmids were collected and centrifuged to separate cell pellets. The pellets were re-suspended with buffer 300 μL then homogenized. The solution was centrifuged with 14000 rpm for 5 minutes. 700 μL of supernatant was transferred to 1.5 mL microtube and added cold isopropanol (700 μL), then incubated with 14000 rpm for 5 minutes. The supernatant was discarded. 40 μL of alcohol was added to the pellet and centrifuged for 5 minutes. Pellets were dried for 30-40 minutes, then re-dissolved with 30 μL of TE buffer containing RNase and incubated for 1 hour at 37°C. The solution was stored at -20°C for further analysis. Plasmid pCAMBIA 1303-p19 and pCAMBIA 1303-ads-p19 were confirmed by
electrophoresis, PCR product analysis, restriction analysis, and sequencing.

The transformation of pCAMBIA 1303-p19 and pCAMBIA 1303-ads-p19 into Agrobacterium tumefaciens and its confirmation

Recombinant plasmids were transformed into A. tumefaciens AG1. One hundred microliters of A. tumefaciens AG1 was thawed and added 1 µg of plasmid, and then incubated on ice and liquid nitrogen for 5 minutes. Cells were incubated at 37°C for 25 minutes and added a liquid yeast-extract-peptone (YEP) medium 1 mL and incubated again at 37°C for 3 hours. Cells were centrifuged in 14 000 rpm for 1 minute. Pellets were re-suspended in 100 µL medium and inoculated in solid YEP containing ampicillin 100 ppm and kanamycin 50 ppm. A single colony was taken and suspended in liquid YEP medium containing ampicillin 100 ppm and kanamycin 50 ppm, and then incubated in a shaker at 250 rpm in the room temperature for 2x16 hours in a dark condition. The plasmid was isolated and characterized using electrophoresis 0.8% agarose and PCR product analysis with a specific primer of p19 gene.

The transformation of a plasmid containing A. tumefaciens into Artemisia annua L.

Vacuum infiltration method

Fourteen-day-old of A. annua was used for the explants. Leaves and hairy roots were incubated in 150 mL of MS basal medium for transformation mediated by A. tumefaciens AG1L. 10 mL of A. tumefaciens culture cells in YEP medium was grown until reached OD_600 = 1. Pellets were re-suspended in 50 mL MS normal medium supplemented with acetosyringone and 0.002% surfactant Silwet S-408. The selected leaves were incubated in a MS medium and continued with the transformation of a plasmid containing A. tumefaciens into A. annua L. by vacuum infiltration method for 20 minutes at 8°C in a dark condition. Infected leaves and hairy roots were blotted using sterile filter paper and co-cultivated for 3 days in room temperature and a dark condition.

Syringe infiltration method

The selected leaves were incubated in a MS medium and continued with the transformation of plasmid-containing A. tumefaciens into A. annua L. with a syringe infiltration method in a dark condition. Syringe infiltration known as agroinfiltration, that is, when a gene is inserted into leaves transiently using a syringe without the needle. Infected leaves were blotted using with sterile filter paper and co-cultivated for 3 days in a room temperature and a dark condition.

The analysis of GUS transient expression

GUS transient expressions in A. tumefaciens infected explants were checked using histochemical method with 5-bromo-4-chloro-3-indolyl glucuronide (X-Gluc) staining.28 Each infected explant was rinsed in phosphate buffer pH 6.8 for 5 minutes and then incubated at 37°C for 24 hours in a dark condition. To strengthen the blue color, explants were washed with ethanol 70% until chlorophyll disappeared.23

The analysis of artemisinin content

Leaves of wild type, hairy roots, and transgenic of A. annua were dried and powdered, and then extracted with ethyl acetate 3x10 mL. Ethyl acetate was evaporated. The residue was dissolved with methanol 1.5 mL for derivatization. One hundred of methanol extract was added 400 µL of NaOH 0.05 N and then incubated for 30 minutes, continued with the addition of acetic acid 0.2 N, and then incubated once more on ice for 10 minutes. The methanol was added up to 1 mL, and then filtered with membrane filter 0.45 µm in size. The samples were injected to high-performance liquid chromatography (HPLC) system with Hewlett Packard Hwallet RP-18 (100 mm x 4.6 and particle size 5 µm) column. The mobile phase was the mixture of phosphate buffer (5 mM, pH 7): methanol: acetonitrile (60:30:10). The elution was gradient with flow of 0.6 mL/min, column temperature of 30°C. The detector was Diode Array Detector with 260 nm wavelength.29

Results and Discussion

Synthetic p19 gene from tomato bushy stunt virus which has a function in disturbing plant protection as PTGS has been amplified with PCR using DreamTaq Master Mix DNA Polymerase and a specific primer of p19 gene. On the other hand, the p19 gene is a virus gene which is easily mutated when it is amplified. Used primers were designed with extension for Xho restriction enzyme sites to be suitable for restriction analysis for both genes and pCAMBIA 1303. PCR results for confirmation of p19 from synthetic gene and ads gene from pCAMBIA-ads plasmid showed that both genes are detected in the right sizes (Figure 1). Electropherogram showed that band p19 has both size and intensity the same as the mass ruler DNA ladder 10 ng/µL. It is assumed that the concentration of the p19 gene is around 10 ng/µL. This p19 gene was further cloned into a cloning vector and transformed into E. coli DH5α. Furthermore, each p19 gene and pCAMBIA-ads plasmid was cut to prepare the cloning.

Ligation into cloning vector was done to prepare DNA insert in a large amount and the same condition. This was to minimalize the mutation event in the next ligation and to check whether the DNA insert in the right condition to be ready for cloning in an expression vector. Cloning is a process to duplicate the parent material, resulting in a large copy of genetic material which is completely the same as its parent material. To confirm the size of the DNA p19 gene, the plasmid from a cloning result was cut by EcoR1 and Xho1 enzyme. The result showed that the p19 gene was successfully cloned into PGEM-T Easy.
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This has also been confirmed using sequencing analysis as shown in Figure 2.

Both pCAMBIA 1303 and pCAMBIA 1303-ads plasmids for ligation were prepared by discarding hyg gene (1000 bp) using XhoI enzyme. Hyg gene minus-plasmid was further purified and measured the concentration. The concentration of pCAMBIA 1303 and pCAMBIA 1303-ads plasmids after purification was 4 ng/μL each. p19 gene was ligated into pCAMBIA 1303 and pCAMBIA 1303-ads plasmids. The availability of kanamycin resistance gene in a pCAMBIA 1303 vector could be used for the selection of transformants in colonies. To check whether the ligation reaction was successfully done or not, the results were analyzed by migration, PCR product, and restriction analysis. A single colony in solid LB medium was taken and re-suspended for isolation. From migration, restriction, and sequencing analysis, all of them showed that the p19 gene was successfully transformed to pCAMBIA 1303 vector, and pCAMBIA-ads plasmid resulted in pCAMBIA 1303-p19 and pCAMBIA-ads-p19 plasmids. Furthermore, both pCAMBIA-p19, pCAMBIA-ads-p19 and pCAMBIA 1303 were transformed into A. tumefaciens with a heat shock method. A successful transformation was checked by PCR using DNA templates from A. tumefaciens harboring each plasmid. PCR analysis results as in Figure 3 showed that the p19 gene could be detected in size 537 pb for pCAMBIA-p19 plasmid (lane 1, 2 and 10) and pCAMBIA-ads-p19 (lane 5 and 6). This band was also found in the positive controls (lane 7-8), while there were no bands in the same size found in A. tumefaciens wild type (lane 3), pCAMBIA-ads plasmid (lane 4) and negative control (lane 9). This confirmed that the p-19 gene in pCAMBIA 1303-p19 and pCAMBIA-ads-p19 were successfully transformed into A. rhizogenes.

No band in the A. rhizogenes wild type and harboring pCAMBIA-ads confirmed that p-19 gene did not come from the contaminated A. tumefaciens and pCAMBIA ads plasmid. To confirm the existence of ads gene in genomic DNA of A. rhizogenes harboring pCAMBIA-P19 (1-2), wild type (3), pCAMBIA-ADS (4), pCAMBIA-ADS-P19 (5-6), positive controls (7-8), negative control (9), pCAMBIA-P19 (10) and DNA Ladder 1 kb (11).

Figure 2. Sequence of p19 gene resulted from cloning with pGEM-T easy.

No band in the A. rhizogenes wild type and harboring ads-p19 plasmid (lane 1,2), pCAMBIA-ads and positive control, while there were no bands in the same size found in negative controls (lane 3-5 and 10) and A. tumefaciens wild type (lane 8), indicating that the ads came only from the transformed plasmids.

Naturally, A. tumefaciens could transfer DNA fragment of its tumor-induced (Ti) plasmid into a plant genome.
to be expressed in plants. A. tumefaciens AGL1 as a carrier for recombinant plasmid that contained inserted gene was grown until reaching OD value 1. OD value 1 means the optimum condition where A. tumefaciens AGL1 can effectively be transformed. Two-week old A. annua plant cell cultures were prepared to accept DNA insert of the p19 gene. The addition of surfactant will improve penetration into cuticle, so it will stimulate material transfer to plant cells. Surfactant Silwet S-408 has a significant effect in improving transformation efficiency in hairy root culture of A. annua up to 27.84% out of the total infected explants.

The transformation of A. tumefaciens AGL1 recombinants into A. annua was done using vacuum infiltration method for 20 minutes. Several research projects have proven this method could improve the expression frequency of A. tumefaciens into plants. Vacuum infiltration can exhaust the air in the intra cell so that the medium can enter the intra cell's spaces. This method was also selected since it is easy to transfer genetic material to A. annua leaves with a small size where the syringe method is not suitable. The infected plants were furthermore incubated for 3 days to maximize the transient transformation into plants. The transformation was also done using syringe infiltration method. Syringe infiltration or agroinfiltration is a method used in plant biology to stimulate the expression of genes in a plant transiently to express the desired protein. This method is widely used technique to transform foreign genes into plant cells since it is simple, rapid, and versatile. The most popular method for agroinfiltration is syringe infiltration. This method is a simple procedure with no need for specialized equipment. A needleless syringe is used to apply Agrobacterium into plant leaves or other plant organs. In this method, the suspension of A. tumefaciens harboring the gene containing plasmids is infected into the plant leaves by direct injection. Furthermore, the bacteria transfer the inserted gene into the plant cells via T-DNA transfer. The benefit of the syringe infiltration method is not time-consuming and convenience. The yields of the recombinant protein are generally more consistent and much higher when compared to other traditional plant transformations. For the agroinfiltration method, Tween-20 could significantly improve the transformation efficiency with the optimal concentration of 0.03% (v/v). To check the transformation results which contain the p19 gene with the plasmid pCAMBIA 1303 and plasmid pCAMBIA 1303-ads, histochemical method using GUS transient expression analysis was done. GUS gene is a gene attached to plasmid pCAMBIA and has an important role as reporting genes in genetic analysis. The expression of reporting gene could have the roles in several aspects, such as protein localization reporting, an indicator for translation activity, or a transduction signal, and successfully gene insert. Existence detection of β-glucuronidase (GUS) could be done qualitatively with histochemical GUS and quantitatively with spectrophotometric GUS. Based on histochemical test, the blue color appeared after adding substrate X-gluc (5-bromo-4-chloro-3-indoyl glucuronide). The percentage of blue area out of total area of leaves and hairy root showed that co-transformation pCAMBIA 1303-ads with pCAMBIA 1303-p19 (Table 1, Figure 5) gave transformation efficiency value higher than direct transformation pCAMBIA 1303-p19-ads and the control pCAMBIA 1303-ads, while the transformation of p19 gave the value higher than control pCAMBIA 1303. These results confirmed that the ads transformation could not optimally be expressed in A. annua due to the RNA silencing process by plant cells. Ads expression level was enhanced by co-transformation together with p19 meaning that the p19 gene suppressed the RNA silencing mechanism by plant cells. Other reports showed that p19 gene of Cymbidium ring spot virus could inhibit RNA silencing through small RNA-binding activity. In the in vitro RNA-silencing system, small RNAs, bound by p19 in plants, are double-stranded siRNAs and they are competent in silencing. During virus infection, p19 could reduce the amount of free siRNA in cells through forming p19–siRNA complexes; therefore, siRNAs are inaccessible for effector complexes of RNA-silencing machinery. The p19-mediated sequestration of siRNAs in virus-infected cells inhibits the spread of the mobile, systemic signal.
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The enhancement of the expression level was also shown by transformed hairy root with the antisilencing p19 gene.

**The analysis of artemisinin content using HPLC**

The assay of artemisinin in samples was calculated by the equation lines derived from artemisinin standard calibration curve. The optimum conditions were obtained by using HP Hwallet column RP-18 (100 mm x 4.6 mm id, particle size 5 μm), the mobile phase a mixture of phosphate buffer (pH 7.5 mm - methanol-acetonitrile (60:30:10, v/v) and a flow rate of 0.6 mL/min. Based on research conducted previously, it is stated that the column temperature is set at 30°C, which aims to improve measurement precision, improving separation, maintaining the retention time repeatability, sharpen chromatogram peak, increasing the efficiency of the column, as well as lowering the pump pressure.

Based on the chromatogram of standard solution, artemisinin appeared at the fifth minute so that for the analysis of samples, it was focused on the fifth minute as well. The next step, the area is calculated in the linear regression equation. Chromatograms of the samples were compared with the control chromatogram. The controls were leaves without transformation and leaves that are transformed by AGL I-non containing genes (AGL I wildtype).

According to the obtained graphs, the hypothesis on the addition of the p19 gene which can increase gene expression was proven, in this case, the gene ads. This is demonstrated by the samples that were transformed with ads gene only were not higher than the genes which were inserted simultaneously with p19. This means that the mechanism of PTGS actually occurs in *A. annua* plant and silencing suppressor p19 suppresses the protection mechanism.

The transformation of plasmids containing genes ads, ads-p19, p19 and co-transformation have been successfully performed on wildtype leaves and hairy root of *A. annua*, based on histochemical GUS and artemisinin content analysis using HPLC with chromatogram, as shown in Figure 6. The levels of artemisinin derived from analysis by HPLC for samples without transformation, agl, ads, ads-p19, p19 and co-transformation using vacuum infiltration method were 0.07, 0.074, 0.08, 0.18, 0.16, and 0.083% respectively, while using syringe infiltration method were 0.07, 0.07, 0.08, 0.17, 0.09, and 0.07% (Figure 7). For the hairy root culture, the co-transformation of ads and p19 in each plasmid could produce artemisinin 0.095%, while no artemisinin was found in the untransformed hairy root. It can be concluded that the ads-p19 gene using vacuum infiltration method.
A method could increase the artemisinin compound in *A. annua* wildtype compared with single *ads* gene plasmid or single *p19* gene plasmid.

**Conclusion**

Amorpha-4,11-diene synthase (*ads*), a key enzyme of antimalarial artemisinin, has been transformed in *A. annua* leaves, mediated by *A. tumefaciens*. In addition, the antisilencing of the *p19* gene has also been transformed into this plant to increase the expression level of *ads* gene. The transformation of *ads* and *p19* genes into leaves of *A. annua* has enhanced the artemisinin content on transformed leaves with *ads-p19* up to 2.57 folds compared to untransformed leaves, while for *p19*, co-transformed and *ads* were up to 2.25, 1.29, and 1.14 folds respectively.

**Ethical Issues**

Not applicable.

**Conflict of Interest**

The authors confirm that this article content has no conflicts of interest.

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