Preliminary investigations of *Agrobacterium*-mediated transformation in indica rice MR219 embryogenic callus using *gusA* gene

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Preliminary steps in the genetic transformation of indica rice MR219 was investigated in the plant-*Agrobacterium tumefaciens* interaction. *Agrobacterium tumefaciens* strain LBA 4404 carrying a binary vector pCAMBIA 1305.2 harboring the modified GUS gene driven by the CaMV 35S promoter was used. Various transformation parameters influences were optimized using embryogenic calli via β-glucuronidase (GUS) as a reporter marker. Various transformation parameters were optimized including bacterial concentration, age of embryogenic callus, pre-culture period, wounding technique, co-cultivation period, immersion time and dry time before co-cultivation, acetosyringone (AS) concentration, pH of co-cultivation media and temperature of the co-cultivation period. The expression of the transient *gusA* gene in the plant genome was preliminary confirmed by histochemical GUS assay activity (as blue spots). The results from transient *gusA* gene expression of calli suggested that the *Agrobacterium*-mediated transfer system of T-DNA in indica rice MR219 was highly efficient. Therefore, the investigation of factors that influence T-DNA delivery is an important first step in the utilization of *Agrobacterium* in the transformation of indica rice MR219 calli.

Key words: Indica rice MR219, *Agrobacterium tumefaciens*, GUS expression.

INTRODUCTION

*Oryza sativa* L., indica-type rice, is an indubitably important staple food for many Asian regions. In Malaysia, indica rice variety MR219 was developed by Malaysian Agricultiral Research and Development Institute (MARDI) to boost rice yield up to 10 mt/ha; 10 to 15% higher than the previous MR84 variety. MR219 potential to produce high yields depends on the grain sizes which weigh 28 to 30 mg and the number of grain per panicle which can reach 200. In addition, MR219 is resistant to bacterial leaf blight and blast disease with short maturation period from 105 to 111 days (http://www.agnet.org/library/rh/2002010a/).

Nevertheless, further researches to improve the quality of MR219 are essential. Jiao et al. (2010) and Miura et al. (2010) recently showed that low number of tiller increased productivity with the increase of number of panicles and number of grains per panicles which is regulated by the over expression of OsSPL14 gene.

Genetic transformation has become an important means in the crop improvement strategies. The system is gene specific and less laborious with less time consumption as compared to the traditional rice breeding practice. *Agrobacterium tumefaciens* has been reported to be an excellent tool in various successful production of genetically modified (GM) crops and ornamentals including barley (Shrawat et al., 2007), banana (Sreeramanan et al., 2006), dragon fruit (Kavitha et al., 2010), cassava (Bull et al., 2009), japonica and indica rice cultivar Kasalath (Kumar et al., 2005; Nishimura et al., 2006; Hiei and Komari, 2008), and orchid

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Hui et al. (2004) reported that plant transformation requires continuous investigation. Hiei et al., 1997; Sreeramanan et al., 2006; Xing et al., 2007; Sreeramanan and Xavier, 2010). Hiei et al. (1997) reviewed various factors affecting the genetic transfer in rice transformation (Nishimura et al., 2006; Hiei and Komari, 2008). Early detection of plant transformation events is necessary for the optimization of the transformation system. Hiei et al. (1997) reviewed various factors affecting the genetic transfer in rice transformation (Nishimura et al., 2006; Hiei and Komari, 2008). Early detection of plant transformation events is necessary for the optimization of gene transfer into the rice genome. Reporter gene such as Escherichia coli gusA (uidA) gene encoding for enzyme β-glucuronidase (GUS) is used to provide a clear indication of the expression, transient or stable, of transferred genes in plant transgenic cells (Jefferson, 1987; Hiei et al., 1997; Sreeramanan et al., 2006; Xing et al., 2007; Sreeramanan and Xavier, 2010).

Various interactions and progress on different host plant for transformation require continuous investigation. Hui et al. (2004) reported that in vitro response is highly genotype dependent, especially in indica rice, which was difficult to have a uniform Agrobacterium-mediated transformation system. Hiei et al. (1997) reviewed various key factors required in Agrobacterium-mediated transfer of genes into rice such as vir gene induction, active cell division in the target tissue (s), medium composition, genotype and Agrobacterium strains and vectors. In the study, several parameters considered crucial for the transformation efficiency of MR219 rice variety using Agrobacterium-mediated system were assessed by using GUS as a reporter gene. There was no system report of Agrobacterium-mediated transformation carried out on this rice MR 219 variety until now.

**Figure 1.** Schematic diagram of the plasmid pCAMBIA 1305.2. The binary vector pCAMBIA1304 (CSIRO, Australia) harboring the reporter gusA and mgfps genes driven by the CaMV 35S promoter.

**MATERIALS AND METHODS**

Embryogenic calli used in this study were derived from mature rice seeds (Zuraida et al., 2007). The calli were maintained on agar solidified embryogenic media containing MS including B5 vitamin (Murashige and Skoog, 1982) and supplemented with 1 mg/L 2,4-D, 10 mg/L NAA, 30 g/L sucrose and 3.5 mg/L agar (Zuraida et al., 2007). The medium was adjusted to pH 5.7 with KOH prior to autoclaving. 3 to 6 weeks-old embryogenic callus was used for the transformation of indica rice MR219.

**Agrobacterium strain and plasmid**

*A. tumefaciens* strain LBA4404 harbouring the plasmid pCAMBIA1305.2 (http://www.cambia.org.au/) with the β-glucuronidase (GUS) gene (uidA version GUSPlus™ and the hpt gene) interrupted with a plant intron (GUS-INT) driven by the Cauliflower Mosaic Virus 35S (CaMV 35S) and nopaline synthase terminator as selectable marker gene was used for the transformation (Figure 1). This vector has hygromycin phosphotransferase (hptII) gene in T-DNA region driven by CaMV35S promoter and CaMV35S polyA terminator, which confers resistance to the antibiotic hygromycin as a plant selection marker. Neomycin phosphotransferase (nptII) gene is located outside the T-DNA region driven by the CaMV35S. Nopaline synthase (NOS) terminator confers resistance to the antibiotic kanamycin as a bacterial selection marker.

**Inoculation and co-cultivation with *A. tumefaciens***

Single colony of *Agrobacterium* from freshly subcultured plate was grown overnight in Luria-Bertani (LB) liquid medium containing 50 mg/L kanamycin and 100 mg/L streptomycin on a shaker at 180 rpm and at a temperature of 28°C. The following day, 250 μl of bacterial suspension was cultured into 10 ml liquid LB medium at 28°C and 280 rpm for 2 to 6 h until the desired density at OD_{600} nm was obtained.

**Optimization of transient GUS expression parameters**

Several factors affecting the *Agrobacterium*-mediated transformation frequency in indica rice MR219 callus were evaluated. The factors were: *Agrobacterium* concentration (0.1, 0.2, 0.4, 0.6, 0.8 and 1.0 at OD_{600} nm), age of embryogenic callus (3, 6, 9 and 12 weeks old), pre-culture period (1, 2, 3, 4, 5, 6 and 7 days), duration of dry time before co-cultivation (15, 30, 45, 60 and 90 min), acetylsyringone concentration in immersion medium (0.100, 200 and 300 μM), pH of co-cultivation media (5.0, 5.2, 5.4, 5.6, 5.8 and 6.0) and temperature of the co-cultivation period (22, 26 and 28°C). To determine the optimum conditions for transformation, factor of the standard conditions were changed each time and the effects on percentage of transient gusA gene expression were evaluated. At the end of the co-cultivation period, the embryogenic calli were evaluated and optimized on the basis of GUS expression (blue spots). Results obtained were based on the percentage of calluses expressing GUS spots over the total number of inoculated callus which was observed 3 days post-transformation. A sample was scored as GUS positive if the blue-spots on calluses mass were at least 25%. All experiments were carried out with 50 embryogenic callus samples and repeated three times.
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GUS histochemical assay

The putative transformants callus was histochemically detected for the presence of the GUS gene following the procedure of Jefferson et al. (1987). The samples from the putative transgenic callus were incubated in GUS buffer [containing the stocks of 2 mM X-glucuronide in DMSO, 100 mM Tris HCl (pH 7.0), 50 mM NaCl, 2 mM potassium ferricyanide and 0.1 % (v/v) triton X-100] overnight at 37°C and cleared in 75% ethanol for 4 h to clean the tissue prior observation. Transient GUS activity was recorded as blue spots (irrespective of size) using a stereomicroscope and photographed using a Nikon camera.

Statistical analysis

The analyses of variances were done and means were compared by the Duncan’s multiple range test (DMRT) using SPSS program 9.0 (SPSS Inc. USA).

RESULTS AND DISCUSSION

The efficiency of Agrobacterium-mediated system on rice calli was inspected through vigilant observations on the effect of several parameters considered to be critical steps. The success of transformation was assessed by the percentage of blue spots signifying transient expression of GUS gene.

The density of Agrobacterium directly affects transformation efficiency since gene transfer only occurs with proper Agrobacterium attachment to plant cells. Therefore high Agrobacterium concentration increased the number of plant cells being infected (Sahoo et al., 2003). Figure 2 shows significant increase in the transformation rate with the increases of Agrobacterium concentration from 0.1 to 0.6 of OD600 nm. The highest number of GUS staining was observed in Agrobacterium concentration OD600 nm at 0.6. More concentrated Agrobacterium suspension (when OD600 nm was 0.8-1.0) however, significantly reduced the number of transformed cells. This might be due to the fact that intense Agrobacterium infection caused severe damage to the plant cells, thus halting transformation process. Xing et al. (2007) reported that bacterium concentration at OD600 nm = 0.6 to 0.8 was the most efficient in sweet potato embryogenic callus Agrobacterium-mediated transformation. Karami (2008) reviewed that Agrobacterium concentration for transformation is dependent on multifactors including Agrobacterium strain and viability, plant species and cultivar and type of tissue used. Experiments with various rice genotypes and wheat showed that higher Agrobacterium density could increase transient GUS expression, but was not correlated with higher stable transformation frequency (Cheng et al., 1997; Hiei et al., 1997; Kumar et al., 2005; Nishimura et al., 2006).

The embryogenic callus of indica rice variety MR219 was selected as the target explants for transformations study. Hiei et al. (1997) demonstrated that embryogenic callus derived from mature seeds of japonica was the best target explants for Agrobacterium-mediated transformation due to its active cell division. Subsequently, freshly isolated immature embryos were later found to be the best explants type for indica rice (Aldemita and Hodges, 1996) and maize (Ishida et al., 2007). Bull et al. (2009) reviewed an overwhelming success of Agrobacterium-mediated transformation of friable embryogenic callus (FEC) in cassava; it is superior as a result of reduced risk of chimeric generation and reduced
The age of the callus is a crucial factor for transformation efficiency. Figure 3 illustrated the reduced transformation rate of embryogenic callus over time. Young calli of 3 weeks old highly enhanced transient GUS expressions as compared to older calli of 6 weeks old and above. Callus of more than 6 weeks old are more recalcitrant to *Agrobacterium* infection, due to increased immunity which is acquired by plant as defense to pathogen attack. Callus aged less than 3 weeks however, are too friable and fragile to endure bacterial infection (data not shown). Hiei and Komari (2008) reported that fresh and healthy immature embryos ensure successful japonica and indica rice transformation. Young embryogenic callus is also favourable due to its higher regeneration ability as compared to old calli (Raja et al., 2009).

Figure 4 shows the drastic increase (more than 70%) of gene transfer in wounded plant as compared to intact non-transformed plantlets escapes during selection as compared to organized tissues such as shoot organogenesis and cotyledon. The first efficient method for *Agrobacterium*-mediated transformation in japonica rice revealed the use of actively dividing, mature seeds derived embryogenic callus as suitable target explants (Nishimura et al., 2006).
Explant. In nature, *Agrobacterium* infected wounded plant result to tumourigenesis which is known as crown gall disease. There are many types of wounding applied in plant transformation such as simple small incision using blades, micro-wounding using particle gun or sonication and injection by syringe (Sreeramanan et al., 2006). Karami (2008) reviewed that the idea of wounding the plant is to create an entry pathway for the *Agrobacterium* infection, in order to compromise physical barrier that might block T-DNA transfer, such as the waxy cuticles on plant epidermis; to stimulate phenolic compound necessary for *vir* gene activation release; and that wounding makes plant cell to actively divide and makes cell walls to be less rigid for easy bacterium breaching, and active DNA replication at this point might facilitate T-DNA integration.

Shrawat et al. (2007) defined pre-culture period as the period that starts the moment immature embryo are first isolated and cultured until immediately before *Agrobacterium* infection. Figure 5 demonstrates that extending pre-culture period more than 3 days improved transformation frequency. Four (4) days of pre-culture proved to be the best increasing transformation efficiency by nearly 50% as compared to only 1 day pre-culture. Longer pre-culture time (5 and 6 days) although enhanced transformation, the efficiency however reduced after the 4th day of pre-culture. Extended pre-culture period allow embryogenic calli to maximize nutrient uptake from pre-culture media to stimulate cells to become competent for *Agrobacterium* attachment. In the transformation of *Vigna* sp., Sahoo et al. (2003) stated that during pre-culture, a physiological and developmental shift occurred in plant tissues to become regenerate-competent which consequently lead to the success of transformation.

Co-cultivation period is described as the time between the elimination of unattached *Agrobacterium* after inoculation, and when the calli are subjected to observation or selection of transgenic line. At this stage, embryogenic calli are cultured together with attached *Agrobacterium* for further infection process on medium lacking selection pressure and anti *Agrobacterium* component such as cefotaxime. Co-cultivation period of embryogenic callus also exhibit the same pattern to the pre-culture period (Figure 6). Significant increased of transient GUS expression were observed in 4 days of co-cultured calli. Lesser days (1 to 3 days) of co-cultivation period produced few transformed line due to insufficient time for complete or maximized transfer of *Agrobacterium* T-DNA into the plant genome. Extended time (5 and 6 days) of co-cultivation also significantly reduced transformation due to overgrown bacteria damage and suffocation of the explants. Huang and Wei (2005) reported that no transformation occurred in immature embryo of maize without co-cultivation period. They also stated that prolonged co-cultivation period reduced transformed immature embryo frequency due to abundant proliferation of bacterium.

The effect of infection period against drying period after infection was carried out and data are summarized in Table 1. In general, 30 min of drying period regardless of infection period was optimal for rice calli transformation in this study whereas, 90 min is the optimal infection period to ensure high transformation efficiency. The highest number of GUS activity was observed in calli infected for 90 min, followed by 30 min of drying on sterile filter paper.
prior to co-cultivation. The combination produce transformation rate recorded almost 85% higher than the combination of 30 min of infection and 60 min drying period which only recorded 26 ± 8 of the average blue dots.

The prolonged immersion period of 90 min allowed sufficient time for a large number of *Agrobacterium* to efficiently get attached to plant cells for infection process, and eventually highly influenced the transformation in calli. Drying period of 30 min is very suitable in MR219 embryogenic calli transformation of which the period is adequate to eliminate unattached bacteria from the explants and to preserve the cells freshness from losses of water content. Desiccation of precultured immature embryo in wheat after *Agrobacterium* infection was shown to lead to higher efficient transformation rate (Cheng et al., 2003). However, freshly isolated immature embryos and precultured immature embryos of sorghum showed higher competency in *Agrobacterium* infection as compared to those that had undergone desiccation process (Zhao et al., 2000).

Figure 7 illustrates the influence of various acetosyringone concentrations (0, 100, 200 and 300 µM) in the immersion medium on the transformation of embryogenic callus. Significant high GUS staining was achieved at the addition of 200 µM acetosyringone in the immersion medium. Similar result was obtained in the transformation of sweet potato when the effect of acetosyringone concentration in co-cultivation medium was investigated (Xing et al., 2007). Acetosyringone is a type of phenolic compound which is well known to increase *Agrobacterium* vir gene activity. However, like most monocot, rice is unable to synthesize such compound. Successful transformation record in barley proved that acetosyringone is not an essential factor; however inclusions of the compound in co-culture medium significantly enhanced *gus* or *gfp* expression in barley embryo (Shrawat et al., 2007) and *Phalaenopsis violacea* orchid (Sreeramanan and Xavier, 2010). They also stated that inclusion of acetosyringone is important when GUS activity is used to determine transformed embryo; whereas, successful gene delivery into embryo can still be observed based on *gfp* expression if acetosyringone is neglected.

Table 1. Effect of various drying periods after infection on indica rice MR219 embryogenic callus. All experiments were carried out with 50 embryogenic callus samples and repeated three times.

| Infection period (min) | Average of gusA blue spots for dry period after infection (min) |
|------------------------|---------------------------------------------------------------|
|                        | 15                | 30                | 45                | 60                | 90                |
| 30                     | 51±7              | 57±5              | 46±5              | 26±8              | 34±5              |
| 60                     | 63±5              | 74±12             | 64±6              | 49±4              | 57±12             |
| 90                     | 71±7              | 167±16            | 154±17            | 63±12             | 66±6              |
| 120                    | 101±17            | 123±10            | 46±8              | 56±6              | 41±7              |

Figure 6. Effect of co-culture periods on transient GUS expression in indica rice MR219 embryogenic callus. All experiments were carried out with 50 embryogenic callus samples and repeated three times.
Figure 7. Effect of acetosyringone concentrations on transient GUS expression in indica rice MR219 embryogenic callus. All experiments were carried out with 50 embryogenic callus samples and repeated three times.

Increasing GUS activity was observed in increasing pH of co-cultivation medium until pH 5.6 (Figure 8). The highest detection of GUS expression of the transformed explants was detected in pH 5.6 followed by pH 5.8. However, at pH 5.4, we could detect 98 blue spots as compared to at pH 5.2 where only 92 blue spots were produced, respectively (Figure 8). The pH of the co-cultivation period was found to be equally an important factor assisting the transformation efficiency. The highest *gfp* expressions of PLBs were observed in pH 5.5 which is a slightly acidic condition (Sreeramanan and Zuraida, 2010). Huang and Wei (2005) obtained the highest PPT-resistance maize calli at slight acidic pH of 5.4. It has been suggested that the acidic pH of 5.5 is suitable for the transformation process as it may induce the virulence genes required for gene transfers (Stachel et al., 1986;
The acidic pH of co-cultivation medium suitable for the Agrobacterium-mediated transformation process is due to its possible function in the induction of vir genes and initiation of T-DNA transfer (Huang and Wei, 2005).

Temperature has been considered a factor affecting the capacity of Agrobacterium to transfer the T-DNA to plant cells (Karami, 2008; Sreeramanan and Xavier, 2010). The investigation of the effect of temperature during cocultivation in MR219 embryogenic callus revealed that temperature plays an important role in transformation efficiency. Higher temperature of 26°C was found to be optimal to support the highest transient transformation frequency in this rice cultivar and there was no difference between 26 and 28°C (Figure 9). Dramatic transient expression reduction occurred when temperature decreased from 26 to 22°C. The results are contradictory to various studies in which optimal temperatures for transformation ranged from 19 to 25°C. However, due to high inconsistency across plant species and tissue used, it is possible to affirm that the optimal temperatures varied depending on these factors.

Therefore, an efficient Agrobacterium-mediated transformation protocol for indica rice MR219 was successfully established at this stage. The investigation of various factors that influence T-DNA delivery is an important first step in the utilization of Agrobacterium in the transformation of indica rice MR219 calli.

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