An Efficient Hairy Root System for Validation of Plant Transformation Vector and CRISPR/Cas Construct Activities in Cucumber (Cucumis sativus L.)

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Hairy root induction system has been applied in various plant species as an effective method to study gene expression and function due to its fast-growing and high genetic stability. Recently, these systems have shown to be an effective tool to evaluate activities of CRISPR/Cas9 systems for genome editing. In this study, Rhizobium rhizogenes mediated hairy root induction was optimized to provide an effective tool for validation of plant transformation vector, CRISPR/Cas9 construct activities as well as selection of targeted gRNAs for gene editing in cucumber (Cucumis sativus L.). Under the optimized conditions including OD650 at 0.4 for infection and 5 days of co-cultivation, the highest hairy root induction frequency reached 100% for the cucumber variety Choka F1. This procedure was successfully utilized to overexpress a reporter gene (gus) and induce mutations in two Lotus japonicus ROOTHAIRLESS1 homolog genes CsbHLH66 and CsbHLH82 using CRISPR/Cas9 system. For induced mutation, about 78% of transgenic hairy roots exhibited mutant phenotypes including sparse root hair and root hair-less. The targeted mutations were obtained in individual CsbHLH66, CsbHLH82, or both CsbHLH66 and CsbHLH82 genes by heteroduplex analysis and sequencing. The hairy root transformation system established in this study is sufficient and potential for further research in genome editing of cucumber as well as other cucumis plants.

Keywords: cucumber, hairy root, Rhizobium rhizogenes, CRISPR/Cas9, root hair-less, CsbHLH66, CsbHLH82, ROOTHAIRLESS1

INTRODUCTION

Originating in the Indian subcontinent, cucumber (Cucumis sativus L.) has become one of the top consumed vegetables widely grown in the world (Paris et al., 2012; Taha et al., 2020). However, cucumber cultivation currently has suffered from biotic and abiotic stresses (Liu et al., 2013; Chojak-Koźniewska et al., 2018; Fan et al., 2020; Taha et al., 2020). There
are various approaches used to improve cucumber production such as optimizing cultivation conditions (Taha et al., 2020), plant grafting (Elsheery et al., 2020), and breeding. Of which, modem breeding with the help of genetic engineering has been known as the most effective method because genetic variability of cucumber varieties is low 3–8% (Chee, 2001; Plader et al., 2007). The genome editing boosted by CRISPR/Cas systems has been widely applied on various crops as a powerful tool to improve plant traits and gene function studies (Zhang et al., 2021). In cucumber, there have been several reports of using CRISPR/Cas9 for targeted gene knockout (Chandrasekaran et al., 2016; Hu et al., 2017). However, applications of genome editing in cucumber have been challenged by less effective transformation procedures that are time consuming and labor extensive processes (Hu et al., 2017). Therefore, the improvement of procedures and the development of supporting tools are necessary to accelerate genetic engineering in cucumber.

*Rhizobium rhizogenes*-mediated hairy root cultures have been established and developed in different plant species for biomass, recombinant protein production, gene function studies, and secondary metabolite researches (Gutierrez-Valdes et al., 2020). In addition, hairy roots also could be used as materials for generating whole plants in certain reports (Trulson et al., 1986; Bueno dos Reis et al., 2007; Butler et al., 2020; Zhang et al., 2020). Recently, the hairy root transformation has been widely utilized to validate and optimize induced mutagenesis by the CRISPR/Cas9 system (Michno et al., 2015; Du et al., 2016; Jacobs and Martin, 2016; Li et al., 2019; Le et al., 2020). Both *in vitro* and *in vivo* cucumber hairy root cultures have been established for different research purposes (Shi and Lindemann, 2006; Anuar et al., 2011; Fan et al., 2020). However, the application of cucumber hairy root transformation for genome editing using CRISPR/Cas systems has not been reported yet. Moreover, it has been reported that hairy root induction frequency is affected by cucumber varieties, bacterial strains, and transformation methods (JingLong et al., 2017; Fan et al., 2020).

In this study, an efficient procedure for *in vitro* cucumber hairy root induction and transformation was established for a local cucumber cultivar. This system was also applied for the expression of a reporter gene (*gus*) as well as validation of CRISPR/Cas9 construct activity. The targeted mutations including heterozygous, homozygous, and biallelic indels in two targeted genes *CsbHLH66* and *CsbHLH82* was obtained at primary and propagated transgenic hairy roots. These results provide an effective tool for gene expression, gene function, and genome editing studies in cucumber.

**MATERIALS AND METHODS**

**Plant Material**

Seeds of a local cucumber (*Cucumis sativus* L.) cultivar Choka F1 were surface sterilized in 15% commercial bleach (bleaching solution containing 5% sodium hypochlorite, Duc Giang, VN) for 15 min, then washed 4–5 times in sterile water (Hoang et al., 2021). The sterilized seeds were soaked in autoclaved water for 3 h at 26°C. After removing seed coats, unwounded seeds were placed on MS30 medium (Table 1) for 2 days in the dark.

**CRISPR/Cas9 Vector Construction for Hairy Root Assay**

The *CsbHLH66* and *CsbHLH82* and other related protein sequences were discovered by the BlastP (NCBI) using the *Lotus japonicus* *ROOTHAIRLESS1* (*LjRHL1*) protein sequence (Karas et al., 2009) as a reference. To induce targeted mutations in *CsbHLH66* and *CsbHLH82* genes, gRNAs were generated by the CCTop program (Stemmer et al., 2015). The CRISPR/Cas9 plant transformation vectors were constructed based on the pKSE401 vector (addgene plasmid # 62202), a gift from Qi-Jun Chen laboratory (Xing et al., 2014), in which, the *neomycin phosphotransferase II* (*nptII*) gene was replaced by the *basta resistance* (*bar*) gene. The coding sequence of *bar* gene was amplified from the binary vector pZY102 (Zeng et al., 2004) using primer pairs: *NcoI*-bar-F and *RsrII*-bar-R (*Supplementary Table S1*). The PCR amplifications were conducted using the Phusion High-Fidelity DNA Polymerase kit (Cat. No. F553L – Thermo Scientific) following the manufacturer’s instructions. After purification, the PCR products were inserted into pKSE401 at *NcoI* and *RsrII* sites. For gRNA insertion, *BsaI*-*CsbHLH66* gRNA-gRNA scaffold-U6 terminator-*BsaI* and *BsaI*-U6 promoter-*CsbHLH82*-gRNA-*BsaI* fragments were generated from the pKSE401 template by PCR using the listed primers (*Supplementary Table S1*). These two fragments were assembled into the destination vector pKSE401 using *BsaI* enzyme sites. The final constructed vector was confirmed by sequencing and named pKSE401-CsbHLH82-66. The pKSE401-CsbHLH82-66 and pZY102 binary vectors were mobilized into *R. rhizogenes* strain K599 and used for cucumber hairy root transformation.

**Rhizobium rhizogenes-Mediated Transformation and Hairy Root Induction**

*Rhizobium rhizogenes* strains including ATCC 15834 (pRi15834) and K599 (pRi2659) wild-type or carrying binary vectors were used for bacterial mediated transformation. Briefly, single colonies

| Medium | Composition or references |
|---|---|
| MS | Murashige and Skoog, 1962 |
| MS30 | MS, 30 g/L sucrose and 7.5 g/L agar; pH 5.8 |
| Re-suspension | 0.5X MS, 30 g/L sucrose and 200 μM/L AS; pH 5.8 |
| Co-cultivation | MS30, 200 μM AS, pH 5.8 |
| Hairy root culture | MS30, 300 mg/L cefotaxime; pH 5.8 |
| Hairy root selection | Hairy root culture medium added selective agents |

AS, antibiotics and selective agents were added to medium after autoclaving.
were cultured in 5 ml liquid yeast extract peptone (YPE) medium supplemented with appropriate antibiotics (25 mg/L rifampicin for the R. rhizogenes strain K599; no antibiotic for the R. rhizogenes strain ATCC 15834; 25 mg/L rifampicin plus 50 mg/L kanamycin for the R. rhizogenes strain K599 harboring pKSE401-CsbHLH82-66 vector; 25 mg/L rifampicin plus 100 mg/L spectinomycin for the R. rhizogenes strain K599 harboring pZY102 vector). The culture was shaken at 28°C and 200 rpm for 4–6 h, transferred to 100 ml of fresh medium and cultured overnight until OD$_{650}$ reached 0.8–1.0. On the next day, the cells were harvested by centrifugation at 3,000 rpm for 15 min and re-suspended in the resuspension medium (Table 1) to get the OD$_{650}$ of 0.4.

The cotyledons from 2-day-old seedlings were collected and used as explants for infection. The distal parts and proximal petioles were excised and removed. Then, the proximal parts of the cotyledons were cut in half along the midrib to get small pieces (Figures 1A, B). Cotyledon segments were kept on wet filter papers during the cutting time to avoid water loss. About 60–70 explants were soaked in 30 ml bacterial suspension for 30 min at room temperature (Figure 1C). Then, explants were removed from the extra liquid and placed on the co-cultivation medium (Table 1) covered by a filter paper (Whatman # 1001-090) on top. The culture plates were kept at 25 ± 2°C in the dark for 5 days. Next, explants were washed by autoclaved water, blotted by sterile filter papers, and placed on the hairy root culture medium (Table 1) for 15–20 days. Root tip fragments (2 cm) were cut and transferred to fresh medium for elongation and development. Transgenic hairy roots were obtained on the hairy root selection medium (Table 1) for 10–15 days (Figure 1E). Cucumber hairy root induction and selection were conducted at 25 ± 2°C under a light density of 2,000 lux with a photoperiod of 16/8 h.

GUS Histochemical Analysis

The pZY102 transformed hairy roots survived on the selection medium were collected for GUS histochemical analysis using 5-bromo-4-chloro-3-indolyl glucuronide buffer at 37°C for 16 h (Jefferson et al., 1987). GUS expression in cucumber hairy roots was observed and captured by a regular camera.

Identification and Characterization of Induced Mutant Hairy Roots

Root hair phenotypes were analyzed using the method described by Karas et al. (2005, 2009), in which root hairs were observed under ZEISS Stemi 305 stereo microscope and captured with AxioCam ERc 5 s digital camera. Genomic DNA was extracted from hairy roots using the CTAB method (Doyle and Doyle, 1987). Hairy roots with induced mutations in CsbHLH genes were detected by heteroduplex analysis on polyacrylamide gel electrophoresis (PAGE; Chen et al., 2012) with some modifications. Briefly, the spanning target regions on the CsbHLH genes were amplified using flanking specific primers (Supplementary Table S1). The PCR amplifications were conducted using DreamTaq Green PCR Master Mix (Cat. No. K1081 – Thermo Scientific) with the following conditions: initial denaturation at 94°C for 4 min; 30 cycles at 94°C for 20 s, 55°C for 20 s and 72°C for 30 s, and a final extension.

**FIGURE 1** | Main steps of cucumber hairy root induction. (A) The 2-day-old seedlings; (B) cotyledon explants preparation. The red arrows indicate the cotyledon fragments used for bacterial infection; (C) cotyledon fragments were soaked in bacterial suspension; (D) the infected explants on co-cultivation medium; (E) hairy root formation at 20 days after co-cultivation; (F) hairy root fragments were transferred to a new medium; (G) hairy roots developed on hairy root culture medium.
at 72°C for 5 min. The mixture containing equal amplicons from wild type (WT) and transgenic hairy roots was denatured at 95°C for 10 min and slowly cooled down to room temperature to form heteroduplexes. The heteroduplexes were detected by PAGE using 12% polyacrylamide gel. The PCR amplicons were further purified and ligated to the pJET1.2 cloning vector (Thermo Fisher Scientific, United States) for sequencing. The sequencing was performed on the ABI3500XL system (Applied Biosystems). CsbHLH mutant sequences were compared to wild-type sequences using BioEdit 7.2.

Statistical Analysis
All statistical analyses were performed using SPSS software version 24.1. The significant analysis (value of \( p < 0.05 \)) between experiments was performed by Student’s t-test or one-way ANOVA followed by post-hoc Duncan’s new multiple range tests.

RESULTS

Optimization of Cucumber Hairy Root Induction Procedure
In this study, cotyledon pieces from 2-day-old seedlings of ChokaF1 cultivar were first inoculated with \( R. \) rhizogenes K599 strain to evaluate the effects of co-cultivation duration on induced hairy root efficacy. After 3, 5, and 7 days of co-culture, explants were transferred onto hairy root induction medium. There were 21.1 ± 1.6% explants of the 5-day co-cultivation treatment that showed hairy root formation at 5 days on induction medium, and it dramatically reached 100% at 20 days after co-cultivation (Figure 2A). While the highest rate of explants in 3- and 7-day co-cultivation producing hairy roots after 20 days were 84.4 ± 1.6% and 65.6 ± 3.14%, respectively, and no more explants produced hairy roots if having the culture period prolonged. Bacterial overgrowth was also observed in the 7-day co-cultivation treatment (data not shown). Therefore, 5-day co-cultivation was sufficient for inducing hairy roots from the local cucumber ChokaF1 and further used for all cucumber hairy transformation experiments.

The effects of \( R. \) rhizogenes strain (ATCC 15834), a widely used \( R. \) rhizogenes strain to induce hairy roots in various plant species including cucumber (Shi et al., 2006; Shi and Lindemann, 2006), were also evaluated on the local cucumber cultivar in comparison to the K599 strain (Figure 2B). In this experiment, explants inoculated with ATCC 15834 strain produced hairy roots earlier than those infected with the K599 strain. About 33.3 ± 11.9% explants induced hairy roots (2–3 cm) after 1 week on hairy root induction medium, whereas no hairy root was observed from explants infected with K599. After 10 days, the rate of explants producing hairy roots was 34.4 ± 4.2% for K599, while it was double (61.1 ± 18.1%) for the ATCC 15834 strain. The number of ATCC 15834 inoculated explants generating hairy root was continuously higher than K599 treated ones after 15 days. However, after 20 days the highest hairy root induction frequency from ATCC 15834 infected explants was lower (83.3 ± 16.6%) than the ones from K599 (100%). In addition, the formation of calli were observed from wounding sites as well as whole explants which were inoculated with the ATCC 15834 strain. Whereas very few calli formed from K599 inoculation (Supplementary Figure S1).

From all the above results, an optimized procedure for cucumber in vitro hairy root induction was established, in which cotyledons of 2-day seedlings were infected with \( R. \) rhizogenes K599 and co-cultivated for 5 days. The total time to conduct the procedure is less than 1 month and the hairy root induction rate could reach up to 100%. Details of the procedure are presented in Figures 1, 3.

Utilizing Cucumber Hairy Root System for Validating Plant Transformation Vector Activities and Transgene Expression
Utilizing the developed hairy root system, the expression of transgenes was conveniently assessed. K599 strain harboring binary vector pZY102 carrying a reporter gene (gus) and
selective gene (*bar*) was transferred into cucumber cotyledon pieces. The induced hairy roots (2 cm) were removed from explants and cultured on the selection medium containing herbicide phosphinothricin (PPT) concentrations ranging from 0.5 to 5 mg/L. The herbicide-resistant hairy roots showed normal morphology and development, while non-resistant hairy roots were completely inhibited and turned yellow without elongation (Figure 4A). The analysis of *gus* gene expression in putative transgenic hairy roots (herbicide resistant ones) after 10 days on selection medium indicated that not all of them were positive for GUS staining (Figures 4B,C; Supplementary Figure S2). The number of selection-escape hairy roots was varied and depended on PPT concentration. On the medium with 3 or 5 mg/L PPT, 6.7 ± 1.6% of survived hairy roots escaped. This number raised up to 46 ± 7.2% and 48.7 ± 7.6% in the medium containing 1.0 and 0.5 mg/L PPT, respectively. On the 15th day in selection medium of 3 mg/L PPT, the number of GUS positive/negative hairy roots was not significantly different from hairy roots on the 10th day (Figure 4C). This result indicates that transgenic hairy roots could be effectively obtained in 3 mg/L PPT medium in 10 days with the lowest rate of escaping from the selection. Therefore, the activity of the binary vector and the expression of transgenes could be effectively analyzed using the cucumber *in vitro* hairy root system for 10–15 days.
Identification of Targeted Genes for CRISPR/Cas9 Induced Mutagenesis of Cucumber Hairy Root

In addition to testing the transgene expression, the developed hairy root induction system in this study was applied for genome editing studies utilizing the CRISPR/Cas9 system. Four candidate sequences, namely, CsbHLH82, CsBHLH66, CsUNE12, CsBHLH84, were identified when the LjRHL1 protein sequence (from *Lotus japonicus*, Karas et al., 2009) was used as a reference for searching on cucumber database (NCBI: ASM407v2) by protein BLAST tool. The phylogenetic tree was constructed using the four candidate sequences, the LjRHL1 sequence, and 3 ortholog sequences from *Arabidopsis thaliana* by the maximum likelihood method and JTT matrix-based model in MEGA X (Kumar et al., 2018). The result indicated that CsbHLH82 and CsBHLH66 were highly close to LjRHL1 (Figure 5A). Moreover, the bHLH domain sequences of the CsbHLH82 and CsBHLH66 were identical to the one of the LjRHL1 (Figure 5B). Therefore, both CsbHLH82 and CsBHLH66 were selected as the targets for mutagenesis in the cucumber hairy roots.

Inducing Targeted Mutations in Cucumber Hairy Roots by CRISPR/Cas9 System

The DNA sequences of both CsbHLH82 and CsbHLH66 genes were reconfirmed in the local cucumber variety for developing a CRISPR/Cas9 vector. Two gRNAs targeting the first exon of CsbHLH66 and CsbHLH82 were identified using the CCTop tool (Figure 6A) and inserted in the CRISPR/Cas9 expression binary vector (Figure 6B) with the purpose to induce individual and simultaneous targeted mutations of the two tested genes.

Targeted mutations of CsbHLH82 and CsbHLH66 were induced in hairy roots using the K599 strain that harbored the CRISPR/Cas9 vector. Each induced hairy root considered as an independent line was transferred to the selection medium. Phenotyping based on more than 80 hairy root lines indicated that there were three morphological groups (Figure 6C, Supplementary Figure S3) including (1) abundant root hairs like wild-type hairy roots (22.04%), (2) sparse and short root hairs (52.54%), and (3) root hair-less (25.42%).
Heteroduplex analysis of 13 randomly selected hairy root lines from group 2 and 3 (Figures 7A, B) showed that all tested lines contained mutations on CsbHLH82 upon the shifted DNA band observation on the PAGE gel as compared to the wild type hairy roots (Figure 7A). In the case of CsbHLH66, only 4 hairy root lines (3, 6, 10, 30) exhibited induced mutations (Figure 7B). Importantly, these 4 lines all belonged to group 3 having the root hair-less phenotype.

The induced mutations in CsbHLH genes of 3 lines (4, 15, 29) from the spare root hair group and 4 lines (2, 6, 9, 10) from the root hair-less group were further characterized via sequencing targeted amplicons (Figures 7C, D). For the CsbHLH82 gene, induced mutations occurred in all tested hairy root lines (Figure 7C). Various types of indels were observed at the target locations, in which the deletion sizes ranged from −1bp up to −33bp and the insertions were from +1 to +10bp. Different from CsbHLH82, targeted mutations in CsbHLH66 were confirmed in only 5 of 7 tested root hair lines, including 2 mutant alleles in lines 2 and 15, which were undetectable in heteroduplex analysis because of only 1bp indels (1bp insertion or deletion). The 1bp insertion and big deletions (−16 to −49bp) in the CsbHLH66 targeted gene were found in 3 tested hair-less root lines, while 1bp deletion was found in one sparse root hair line 15 only (Figure 7D). Moreover, two mutant alleles (biallelic mutation) of targeted genes were also observed in many hairy root lines such as lines 6 and 10 for CsbHLH66 gene and lines 2, 6, 15, and 29 of CsbHLH82 gene (Figures 7C, D). The homozygous mutant allele was found in line 9 (for CsbHLH82 gene). Three different alleles (chimeric alleles) of the CsbHLH82 gene were also observed in hairy root line 4. Importantly, lines 6 and 10 contained null mutations of both the CsbHLH82 and CsbHLH66 genes. As a consequence, it is obvious that our developed hairy root system has a great potential for genome editing studies in cucumber.

The stability of induced mutations of targeted genes was further confirmed in in vitro propagated hairy roots. The root fragments from line 6 and 10 were cut and propagated in a new medium for the next generations. Heteroduplex, Sanger, and amplicon sequencing analysis were conducted using DNA
extracted from the third propagation root lines. DNA band patterns in PAGE and sequencing data indicated that the mutations observed in the propagated lines were the same as in the primary roots (first propagation) for both \textit{CsbHLH82} and \textit{CsbHLH66} (Supplementary Figure S4). This result demonstrated that stably induced mutations were conducted in cucumber hairy roots.

\section*{DISCUSSION}

\subsection*{Establishment of an Efficient Procedure for Cucumber \textit{in vitro} Hairy Root Induction}

In this study, an efficient procedure for cucumber \textit{in vitro} hairy root induction was established. The hairy root induction frequency could reach 100\% after 20 days of co-cultivation.
using the *R. rhizogenes* K599 strain. Different factors including explants, co-inoculation, and co-cultivation conditions have been reported to influence plant transformation efficacy as well as hairy root induction. In our results, the 5 days co-cultivation was selected because of the highest hairy root induction efficiency. Co-cultivation duration extended to 7 days negatively influenced cucumber hairy root formation due to the overgrowth of bacteria after inoculation. This result is in agreement with Shi et al. (2006), in which 5 days of co-cultivation was considered to be optimal for cucumber in vitro hairy root induction. One other optimization in this study was the use of cotyledon pieces from 2-day-old seedlings as initial explants instead of 10-day-old seedlings as in previous studies (Shi et al., 2006; Shi and Lindemann, 2006) for hairy root generation. Such kind of explants was reported to be optimal for cucumber transformation and regeneration only (Nanasato et al., 2013). Apart from increasing the number of hair roots produced, the use of younger cotyledons also helps to shorten the time needed for hairy root induction procedure (<1 month).

**Rhizobium rhizogenes K599 Strain Is Effective for Cucumber Hairy Root Induction**

The hairy roots were effectively induced from the local cultivar Choka F1 using *R. rhizogenes* K599 strain. Bacterial strains have been known as an important factor for hairy root induction and transformation. In previous studies, different *R. rhizogenes* strains were used for in vitro hairy root induction in cucumber (Trulson et al., 1986; Shi et al., 2006; Shi and Lindemann, 2006; Anuar et al., 2011; JingLong et al., 2017) and the hairy root induction rates were variable depending on cucumber cultivars and transformation procedures. The hairy root induction rate of over 90% was reported by Shi et al. (2006) and Shi and Lindemann (2006) using the ATCC 15834 strain. Recently,
R. rhizogenes K599 strain had been utilized for inducing hairy roots in cucumber with the frequency of 95% under in vivo conditions (Fan et al., 2020). In our study, the K599 strain could facilitate 100% infected cucumber explants to produce strong hairy roots, much higher than that of ATCC 15834 strain (83.3 ± 16.6%). The formation of callus instated of hairy roots by ATCC 15834 strain (Supplementary Figure S1) might be a reason to lower the number of hairy roots produced. The fewer callus formed from K599 inoculation were coupled with the better hairy root morphology. Therefore, the K599 strain is most suitable for hairy root induction and transformation of the local cultivar.

A Convenient Tool to Assess the Activities of CRISPR/Cas9 Constructs in Cucumber

Hairy root systems have been used for genome editing of various plant species. For cucumber, the hairy root system has been applied for studying transgenic plant generation (Trulson et al., 1986), gene expression (Fan et al., 2020), promoter evaluation (Anuar et al., 2011), recombinant protein production (Shi and Lindemann, 2006), but not for genome editing yet. In this study, an efficient procedure for in vitro hairy root induction of a local cucumber cultivar (ChokaF1), which was successfully utilized for stable transformation (Hoang et al., 2021). This procedure was well performed for the expression of a reporter gene (gus) in a short time. The developed hairy root system was also effective for gene editing of cucumber using CRISPR/Cas9 construct. Targeted mutations of two endogenous genes (CsbHLH66 and CsbHLH82) were induced accurately. Different indels of targeted genes were observed in tested hair roots including insertions, and larger and small deletions (Figure 7). All homozygous, heterozygous, and biallelic, as well as chimeric alleles could be obtained in the primary hairy roots (Figure 7). In addition, the targeting capability of different sgRNAs (single guide RNA) could be evaluated within 1 month, instated of several months or longer if using the transgenic plant approach. The targeted mutations induced in transgenic hairy roots were also observed in stable transgenic cucumbers (Chandrasekaran et al., 2016; Hu et al., 2017).

CsbHLH66 and CsbHLH82 Genes Are Involved in Root Hair Performance of Cucumber

The root hair-less phenotype was observed in Lotus japonicus and characterized as a consequence of natural mutations in the SjRHL1 gene (Karas et al., 2009). In A. thaliana, this phenotype was also found in the triple mutant of AtLRL1, AtLRL2, and AtLRL3, while sparse root hair phenotypes were caused by single or double mutations of these three genes (Karas et al., 2009). In our study, induced mutations in CsbHLH82 resulted in the root hair spare phenotype (lines 15, 29), while simultaneous mutations of both CsbHLH82 and CsbHLH66 genes caused the root hair-less phenotype (lines 6, 10; Figures 7C, D). This result suggests that two bHLH transcription factors encoded by CsbHLH82 and CsbHLH66 are redundantly involved in regulating the root hair formation in cucumber. However, further research needs to be performed to characterize and confirm the contributions of CsbHLH66, CsbHLH82, and other CsbHLH genes in cucumber root hair formation at whole plant levels.

CONCLUSION

In this study, an effective protocol for cucumber hairy root induction and transformation was developed for a local cucumber cultivar. This hairy root system has been successfully used to evaluate the expression of transgenes and the activities of CRISPR/Cas9 systems within 1 month. The obtained results obviously provides a potential tool for transgene expression and genome editing studies in cucumber as well as other Cucumis plants.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding authors.

AUTHOR CONTRIBUTIONS

PD and HC conceived and supervised the study. PD, DN, and TH designed the experiments, analyzed the data and approved the submitted version. CN conducted the phylogenetic tree and designed the gRNAs. NL and HT performed CRISPR/Cas9 vector constructions. TH conducted cucumber hairy root transformation. DN performed genotyping and phenotyping of transgenic hairy roots. HC, Y-HM, and CN revised and proofread the manuscript. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2021.770062/full#supplementary-material
Supplementary Table S1 | Primers used in this study.

Supplementary Figure S1 | Cucumber hairy root induction by Rhizobium rhizogenes strain K599 and ATCC 15834. The photos were captured at 10 days after co-cultivation.

Supplementary Figure S2 | GUS histochemical analysis of the herbicide resistant hairy roots. WT: non-transgenic hairy roots grown on the PPT free medium; 1–14: Hairy root lines survived and developed on the selection medium with PPT.

Supplementary Figure S3 | Distribution of root hair phenotypes of tested cucumber hairy roots.

Supplementary Figure S4 | The stability of induced mutations in propagated cucumber hairy roots. (A) Heteroduplex analysis of Cas9HLH66 gene. (B) Heteroduplex analysis of Cas9HLH62 gene. M: 100bp DNA Ladder; WT: wild type; P6 and P10: The original hairy roots of lines 6 and 10; 6–1, 6–10: 1–10, 6–1–10: Propagated roots from lines 6 and 10 at the second and third generations. (C, D) Expanding targeted sequences of Cas9HLH62 and Cas9HLH66 genes from propagated roots of the hairy root lines 6 and 10. Amplicon sequencings were conducted by the Sanger method. The expected sequences of mutant alleles were denoted upper chromatograms.

(B) Clone sequencing of Cas9HLH62 and Cas9HLH66 targeted regions of propagated cucumber hairy roots at the third generation. Red and blue letters indicate target and PMI sequences, respectively. Green letters indicate inserted nucleotides.
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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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