Development of RNAi-Based Vector Aims at Creating Antiviral Soybean Plants in Vietnam

Lo Thi Mai Thu, Le Van Son, Chu Hoang Ha, and Chu Hoang Mau

Abstract—Soybeans (Glycine max (L.) Merrill) are easily infected by soybean mosaic virus (SMV) and bean yellow mosaic virus (BYMV), which may reduce their yields from 66%-80%. At present, preventive measures are still the main methods to reduce losses caused by these two viruses. RNAi is thus considered as a modern and promising technique to generate plants resistant to viruses. In this study, we cloned and characterized the Coat protein (CP) coding gene of SMV cultivar SL1 and SL2, which is 720 bp in length and encodes 240 amino acids. The two conserved regions of CP gene from these two viruses (SMV and BYMV) were integrated into a 573 bp fragment, called CPi(SMV-BYMV). Vector pK7GW-CPi (SMV-BYMV), carrying the CPi (SMV-BYMV) fragment in sense and antisense orientation separated by an intron, was successfully designed by Gateway technology (Invitrogen). Once being transformed into plant cells, this vector is expected to be transcribed into a hairpin RNA structure that could induce RNAi mechanism to degrade the CP genes of SMV and BYMV, thus limit their infection. Therefore, this vector could be a potential tool to generate transgenic soybean plants that are resistant to both two virus species SMV and BYMV in Vietnam.

Index Terms—BYMV, CPi, CP gene, Glycine max, SMV, RNAi, transgenic vector.

I. INTRODUCTION

Soybeans (Glycine max (L.) Merrill) are industrial crops which have high economic and nutritional value, as well as as well as play an important role in improving soil fertility. Soybeans, however, are easy to be infected by numerous virus species (more than 100 species according to statistics) [1], in which some could cause severe damage such as soybean mosaic virus (SMV) and bean yellow mosaic virus (BYMV) [1], [2]. SMV and BYMV can simultaneously infect into a soybean plant and their symptoms are difficult to distinguish. Their simultaneous infection could reduce soybean yields from 66% - 80% [3], [4]. SMV and BYMV can be spread by aphids, in addition to longer diseases transmitted through seeds.

At present, main methods to reduce losses caused by these two viruses are still traditional preventive measures, such as selection of the disease-resistant plants, field sanitation and crop rotation. These traditional methods are often complicated, time consuming, ineffective and unsustainable. Therefore, scientists have to seek new efficient methods to control these viruses. One of the most potential approaches is RNA interference (RNAi). RNAi is a process that inhibits gene expression by the double stranded RNA (dsRNA) that can cause the degradation of target mRNA or blocking translation [5]-[7]. Based on this principle, some author has successfully applied RNAi mechanism in the strategy to create transgenic plants resistant to viruses [8]-[10]. In the past few year, RNAi has been broadly used as a new biotechnological tool for the control of virus diseases in plants.

Genomes of both SMV and BYMV consists of one single-stranded, linear, positive-sense RNA molecule. As in all members of the genus Potyvirus, their single open reading frame is expressed as a large polyprotein which is autoproteolytically cleaved to yield ten functional proteins: P1 proteinase (P1), helper component proteinase (HC-Pro), Protein P3 (P3), 6 kDa protein 1 (6K1), Cytoplasmic inclusion protein (CI), 6 kDa protein 2 (6K2), the viral genome linked protein (VPg), Nuclear inclusion protein A (NIA), Nuclear inclusion protein B (NIB), and coat protein (CP) [11], [12]. The Coat Protein plays vital roles in protection of the RNA inside, infection, movement of virus from cell to cell, transmission by aphids, and symptom formation [13]. Therefore, silencing the CP gene can suppress the infection of SMV and BYMV.

The RNAi-based vector containing CPi fragment inverted repeats could be used to transfer into plants and would express into hairpin RNA structure (hairpin RNA, hpRNA), thereby stimulating RNAi mechanism when there is penetration of SMV or BYMV into plants. Scientists noticed that the buffer zone of hpRNA be repeated with a intron sequences (ihpRNA) will inducing gene silencing with the highest level [14], [15]. Kenny et al. (2007) have created lines of transgenic pea plants antiviral bean golden mosaic virus with resistance ability up to 93% [16]. Lim et al. (2007) studied on HC-Pro gene and transferred into soybean plants has noticed that, when transgenic plants were infected SMV after 2 weeks, the leaf mosaic symptoms disappear and SMV content has decreased significantly, however HC-Pro gene of SMV has altered leaf morphology and reduce granulating in transgenic soybean plants [17]. Generally, most of the transgenic plants slow down the accumulation of virus and slow or reduce disease symptoms. However, the disease resistance of crops still heavily depends on the function and location of the transgene. In Vietnam, the first successful application of RNAi in generation of plants resistant to viruses was the generation of transgenic tobacco plants resistant to Cucumber mosaic virus (CMV) and Tobacco mosaic virus (TMV). The transgenic tobacco lines containing

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simultaneously two CP gene fragments of TMV and CMV showed resistance ability to two virus strains up to 60% [18]. Next was the success in creation of transgenic papaya plants resistant to papaya ringspot virus (PRSV) [19], [20]. However, the authors also found that the rates of disease resistance in transgenic plant lines depend on gene fragments selected for designing RNAi-based vector. Mosaic disease in soybean plant caused by SMV and BYMV have symptoms that are difficult to distinguish, and soybean plants can also be coinfected by these two viruses at the same time. Thus, in this study, we present some results on amplification of CP gene from mRNA isolated from SMV lines SL1, SL2 in Son La province, Vietnam. After that, we designed an RNAi-based vector carrying the conserved region of CP genes of both SMV and BYMV, with the desire to create transgenic soybean plants that are resistant to both these two virus species.

II. MATERIALS AND METHODS

A. Materials

Plant materials: Leaves of soybean plants which are suspected to be infected by SMV collected in several localities of Son La province, Vietnam were used for total RNA extraction of SMV. Tobacco cultivars N. tabacum K326 was supplied by Department of Plant Cell Technology, Institute of Biotechnology, Vietnam.

Bacteria strains: E. coli DH5α, E. coli One Shot® TOP 10 (Invitrogen) and strains A. tumefaciens CV58C1 carrying plasmid pGV2660 cytotoxic.

Plasmid: Vector cloning PBT, transgenic vector pK7GW1WG2(II), vector cloning pENTRY™ and pENTRYTM Directional TOPO® Cloning KIT (Invitrogen), KIT Gateway Gateway® LR reaction Clonase™ II.

B. Methods for Cloning CP Gene and Synthesis of CPI Gene Region of SMV and BYMV

Total RNA was isolated using Trizol Reagent KIT, cDNA was synthesized using the Maxima ® First Strand cDNA Synthesis KIT. The cDNA was used as template for PCR to amplify coding region of CP gene of SMV. From the information of SMV CP gene sequence on GenBank, code number X63771, we designed the primer pair SMVcp-F/SMVcp-R to amplify the coding region of SMV CP gene:

<table>
<thead>
<tr>
<th>Prime</th>
<th>5’-</th>
<th>3’</th>
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<tbody>
<tr>
<td>SMVcp-F</td>
<td>TTACCATCGAAGGAGCAAGGAAG-3’</td>
<td>5’-TGACATTGAGGAAACATCTCAACT-3’</td>
</tr>
</tbody>
</table>

PCR products were then examined by electrophoresis on 1.0% agarose gel and purified by JET™ Gene Gel Extraction KIT. Recombinant vector was transformed into E. coli DH5α cells with variable heat shock. Bacteria carrying recombinant vector were selected on antibiotic selective environment (carbenicillin, kanamycin). The colonies carrying the recombinant vector were selected by colony-PCR with specific primers, then cultured in LB supplemented with antibiotic to collect biomass. Recombinant plasmids were obtained by extracting using the molecular cloning methods [21]. The two CP gene sequences of SMV lines SL1, SL2 were determined on the automated DNA ABI PRISM 3100 Avant Genetic Analyzer machine. It utilized the software biological Clustal to test the sequence of CP gene.

From the sequences of CP genes of SMV and BYMV, we identified their conserved regions using BLAST in NCBI. Primer pair SMV-CPI-F/SMV-CPI-R was used to integrate these two regions from these two viruses:

<table>
<thead>
<tr>
<th>Prime</th>
<th>5’-</th>
<th>3’</th>
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</thead>
<tbody>
<tr>
<td>MV-CPI-F</td>
<td>CACCGGCGAAGGACGTTACA-3’</td>
<td>5’-ATGTTCCGAACCCCAAGCAAT-3’</td>
</tr>
</tbody>
</table>

C. Methods of Designing Transgenic Vector Carrying RNAi Structure Containing CPI Gene Region by Gateway Technique

Designing of transgenic vector in plants carrying RNAi structures is performed as described by Karimi et al. (2002) [10].

Fragment CPI (SMV-BYMV) is amplified by PCR with primer pair MV-CPI-F/BYMV-CPI-R. PCR products were examined by electrophoresis on 1.0% agarose gel and purified by GeneJET PCR Purification Kit (Thermo Scientific). The CPI fragment was then cloned into vector pENTRY/D to create recombinant vector pEN-CPI containing the recombinant position attL (Recombinant position with attrR on vector pK7WG2 (II)). Recombinant vector pEN-CPI transformed into E. coli One Shot® TOP 10 cells with variable heat shock. The colonies carrying the recombinant vector were selected by colony-PCR with specific primers SMV-CPI-F/BYMV-CPI-R and primer pair M13-F/M13-R.

Vector carrying RNAi structure containing CPI gene region (pK7GW-CPI [SMV-BYMV]) was created by Gateway technique (Invitrogen). Transgenic vector carrying RNAi structure containing CPI gene region pK7GW-CPI(SMV-BYMV) was transferred into Agrobacterium tumefaciens cells by electrical pulses (2.5 kV, 25 μF, 200 Ω). The colonies A. tumefaciens carrying vector pK7GW-CPI(SMV-BYMV) were selected by colony-PCR technique with specific primers SMV-CPI-F/BYMV-CPI-R.

III. RESULTS AND DISCUSSION

A. Cloning and Sequence Determining of CP Gene of SMV, and Generating CPI (SMV-BYMV) Fragment

From the leaves of soybean plants which were suspected to be infected by SMV, we extracted total RNA and synthesized cDNA with random primers (random hexamer primers). CP gene was amplified by PCR with specific primer pair SMVcp-F/SMVcp-R and RT-PCR products obtained a very special band with the length of approximately 0.7 kb as expected (Fig. 2 a). PCR products were purified and inserted into cloning vector pBT_2705bp and transformed into E. coli DH5α. We selected clones carrying recombinant plasmid, extracted recombinant plasmids and checked the size of the insert by colony-PCR. The results confirmed that plasmid carrying an insert with expected size (Fig. 2 b).

The results of gene sequencing showed that CP gene fragment is 720bp in length, encoding 240 amino acids.
Results comparing with gene sequences CP published on GenBank by using BLAST in NCBI show that sequences of CP gene fragment isolated from SMV lines SL1 and SL2 are similar to that in CP gene sequence (code number: X63771) on Genbank with 100% (for SMV line SL1) and 99.3% (for SMV line SL2) of similar coefficients. Results comparing two sequences of CP gene fragment of lines SL1 and SL2 show there is 99.3% of similarity, and different in only 5 nucleotides (Table I).

<table>
<thead>
<tr>
<th>Nucleotide positions</th>
<th>X63771</th>
<th>Line SL1</th>
<th>Line SL2</th>
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<tbody>
<tr>
<td>31</td>
<td>A</td>
<td>A</td>
<td>T</td>
</tr>
<tr>
<td>38</td>
<td>C</td>
<td>C</td>
<td>G</td>
</tr>
<tr>
<td>686</td>
<td>A</td>
<td>A</td>
<td>G</td>
</tr>
<tr>
<td>690</td>
<td>G</td>
<td>G</td>
<td>T</td>
</tr>
<tr>
<td>694</td>
<td>T</td>
<td>T</td>
<td>A</td>
</tr>
</tbody>
</table>

In addition, the differences between amino acid sequences were found in 5 amino acid positions (11, 13, 220, 230, 232) and three positions of amino acid in poty-coat region. This result confirmed that we have successfully cloned CP gene from the SMV lines SL1 and SL2.

Genomes of SMV and BYMV include genes on single-stranded RNA and the inhibition of mRNA by RNAi mechanism occurs only when there is a high homology between the siRNA and the target mRNA. On the other hand, the antiviral effect of transgenic plants depends on the function of the target viral genes, so selected the genes to design the structure for RNAi technique is an important step leading to the success of this technique. We collected information about the CP genes of SMV and BYMV in GenBank, used BLAST in NCBI to identify conserved regions to design CPi gene region containing conserved regions of the two CP genes, denoted as CPi(SMV-BYMV), with the size of 573 nucleotides.

**B. Designing an RNAi-Based Vector Containing CPi(SMV-BYMV) Fragment**

CPi region CPi(SMV-BYMV) was amplified by PCR with primer pair SMV-CPi-F/BYMV-CPi-R and PCR products obtained a very special band with the length of 573bp as expected (Fig. 3).
PCR products were purified and inserted into cloning vector pENTR™ Directional TOPO® Cloning KIT to create recombinant plasmid carrying the CPi (pEN-CPi (SMV-BYMV)) and insert into E. coli TOP 10 One Shop®. Positive colonies were checked by colony-PCR with specific primers SMV-CPi-F/BYMY-CPi-R and primer pair M13-F/M13-R (Fig. 4).

RNAi-based vector containing CPi gene region, denoted as pK7GW-CPi (SMV-BYMV) was generated by adding 3μl LR reaction (plasmid pEN-CPi (50–150ng), vector pK7GWIWG2(II) (150 ng/μl), H2O deionised) into 50μl E. coli cell One Shot ® TOP 10, with variable heat shock at 42°C in 45 seconds. Recombinant plasmid pK7GW-CPi (SMV-BYMV) was checked by restriction enzymes Xho I and colonly-PCR. Recombinant vector pK7GW-CPi (SMV-BYMV) was cut by Xba I to obtain DNA fragment with the approximate size of 1.1kb, in which the 1.1kb DNA fragment contains our target gene CPi (SMV-BYMV). Structure pK7GW-CPi (SMV-BYMV) was transformed into A. tumefaciens and the transformed A. tumefaciens cells were selected and checked by colony-PCR reaction with primers SMV-CPi-F/BYMY-CPi-R (Fig. 5). Positive A. tumefaciens lines were stored in glycerol at -80°C in order to generate transgenic soybean plants that are resistant to SMV and BYMV.

IV. CONCLUSION

SMV CP gene has been isolated and cloned from SMV cultivar SL1, SL2 (Vietnam). CP gene fragment has the size of 720bp, encoding 240 amino acids. The two conserved regions of CP gene from these two viruses (SMV and BYMV) were integrated into a 573 bp fragment, called CPi(SMV-BYMV). Vector pK7GW-CPi (SMV-BYMV), carrying the CPi (SMV-BYMV) fragment in sense and antisense orientation separated by an intron, was successfully designed by Gateway technology (Invitrogen). This vector will be used to create transgenic soybean plants that are resistant to both SMV and BYMV.

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