Pseudorecombination between Two Distinct Strains of *Cucumber mosaic virus* Results in Enhancement of Symptom Severity

Mi Sa Vo Phan¹, Jang-Kyun Seo⁲, Hong-Soo Choi⁳, Su-Heon Lee⁴ and Kook-Hyung Kim¹,4*

¹Department of Agricultural Biotechnology and Plant Genomics and Breeding Institute, Seoul National University, Seoul 151-921, Korea
²Crop Protection Division, National Academy of Agricultural Science, Rural Development Administration, Suwon 441-707, Korea
³Department of Applied Biology, Kyungpook National University, Daegu 702-701, Korea
⁴Research Institute of Agriculture and Life Sciences, Seoul National University, Seoul 151-921, Korea

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Recently, a *Cucumber mosaic virus* (CMV) strain, named as CMV-209, was isolated from *Glycine soja*. In this study, symptom expression of CMV-209 was analyzed in detail in *Nicotiana benthamiana* by comparing with that of CMV-Fny, which is a representative strain of CMV. Using infectious cDNA clones of CMV strains 209 and Fny, symptom expression of various pseudorecombinants between these two strains were examined in the early and late infection stages. In the early infection stage, the pseudorecombinants containing Fny-RNA2 induced stunting and leaf distortion on the newly emerged leaves whereas the pseudorecombinants containing 209-RNA2 caused no obvious symptoms. In the late infection stage, the pseudorecombinants containing 209-RNA1 and Fny-RNA2 induced severe leaf distortion and stunting, while CMV-209 induced mild symptom and CMV-Fny caused typical mosaic, general stunting, and leaf distortion symptoms, indicating that RNA 2 encodes a symptom determinant(s) of CMV, which is capable of enhancing symptoms. Furthermore, our results support the possibility that natural recombination between compatible viruses can result in emergence of novel viruses causing severe damages in crop fields.

**Keywords**: *Cucumber mosaic virus*, symptom expression, pseudorecombination
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al., 2007). In a few cases, the determinants of expression symptom has been mapped to RNA1 (Divéki et al., 2004) or both RNA1 and RNA2 (Zhang et al., 1994).

CMV-209 is a CMV strain, which was recently isolated from Glycine soja (Vo Phan et al., 2014). In this study, symptom expression of CMV-209 was analyzed in Nicotiana benthamiana based on pseudorecombination of genomic RNAs of CMV-209 with those of CMV-Fny, which is a typical strain of CMV. Our results demonstrated that, in the early infection stage, the pseudorecombinants harboring Fny-RNA2 caused stunting and leaf distortion on the newly emerged leaves although the pseudorecombinants holding 209-RNA2 exhibited no obvious symptoms. In the late infection stage, the pseudorecombinants consisting of 209-RNA1 and Fny-RNA2 induced severe leaf distortion and stunting, while CMV-209 induced mild symptom and CMV-Fny caused typical mosaic, general stunting, and leaf distortion symptoms, indicating that the symptom determinant(s) of CMV is included in RNA2 and is capable of enhancing symptom severity. Additionally, our results support the possibility that novel viruses causing severe damages in crop fields can be emerged by natural recombination between compatible viruses.

The full-length cDNAs of genomic RNA of CMV-209 and -Fny were amplified using the cDNA clones of CMV-209 (i.e. pCMV-209R1, pCMV-209R2 and pCMV-209R3) (Vo Phan et al., 2014) and Fny [i.e. pCR1 (+), pCR2 (+) and pCR3 (+)] (Seo et al., 2009) as templates. The primers, CMV-R1R2-5-end-T7, CMV-R3-5-end-T7, CMV-3-end-Smal (Table 1), were used for PCR, thereby the resulting products contained a T7 promoter and Smal site at the 5′ and 3′ ends, respectively. The resulting PCR products were cloned directly into pGEM®-T Easy vector (Promega, USA). The final cDNA clones of CMV-209 and -Fny constructed for in vitro transcription were named pT7-CMV-209R1, pT7-CMV-209R2, pT7-CMV-209R3, pT7-CMV-FnyR1, pT7-CMV-FnyR2, and pT7-CMV-FnyR3, respectively (Fig. 1).

The CMV-209 and -Fny clones were linearized by Smal. The cDNA-derived infectious RNAs of CMV-209 and -Fny were generated by in vitro transcription with T7 RNA polymerase (Takara, Japan) in the presence of 7-methylguanosine “cap” structure (Promega, USA) according to the manufacturer’s protocol. The in vitro transcripts from the full-length cDNA clones of CMV-209 and -Fny RNAs 1, 2, and 3 were named W1, W2, W3, F1, F2, and F3, respectively (Fig. 1) and mixed to mimic the original CMV RNAs (i.e. CMV-209, W1W2W3; CMV-Fny, F1F2F3). Six different pseudorecombinants, W1W2F3, W1F2W3, F1W2F3, F1F2W3, F1W2F3, and W1F2F3, were also generated by mixing the in vitro transcripts. All of the parental and pseudorecombinant RNAs were diluted in the inoculation buffer (0.05 M potassium phosphate, pH 7.5) and total 1.5 µg of RNA transcripts (500 ng of each RNA transcript 1, 2 and 3) were inoculated per plant. All the N. benthamiana plants inoculated with transcripts were lightly dusted with carborundum before inoculation. Mock-treated plants were inoculated with the inoculation buffer. The inoculated plants were maintained in an insect-free greenhouse at 25°C under a 14 h photoperiod. The disease

### Table 1. Primers used for CMV-detection and construction of the CMV cDNA clones

| Name                  | Sequence (5′ → 3′) |
|-----------------------|--------------------|
| CMV-3-end-Smal        | GCCCCCCGTTGCTCCTTTTTRGAGRCC |
| CMV-R1R2-5-end-T7    | taatacgactcaataTGGTTATTTCAAGAGGCAGCG |
| CMV-R3-5-end-T7      | taatacgactcaataGTAATCTCTACCAGTGTG |
| CMV-detR1-F          | TTTTTGAAACGCCTCCACTGACATGA |
| CMV-detR1-R          | AGTGAAGCCTTTACGGCTTGGG |

*The Smal sequence is shown in bold face and the sequence of T7 promoter is written in italic.*
symptoms were observed at two stages, the early infection stage [7 days post inoculation (dpi)] and the late infection stage (12 dpi). CMV infection of the inoculated plants was confirmed by RT-PCR at 7 dpi using primers CMV-detR1-F and CMV-detR1-R (Table 1).

The *N. benthamiana* plants infected with CMV-Fny (F1F2F3) showed general stunting, vein clearing, and leaf distortion on the infected systemic leaves in the early infection stage (Fig. 2C). In contrast, the plant inoculated with CMV-209 (W1W2W3) developed no obvious symptoms and indistinguishable from healthy plants although they were systemically infected with CMV-209 as confirmed by a specific RT-PCR (Fig. 2D). The pseudorecombinants containing F2 (i.e. F1F2W3, W1F2F3, and W1F2W3) produced leaf distortion and stunting symptoms (Fig. 2E, G and I). However, two pseudorecombinants of W1W2F3 and F1W2F3 induced no obvious symptoms (Fig. 2F and H). In summary, the pseudorecombinants containing F2 induced stunting and leaf distortion on the newly emerged leaves, while those containing W2 (except for F1W2W3, which induced vein clearing) caused no obvious symptoms in the early infection stage. Therefore our results suggested that CMV RNA2 contains a symptom determinant(s) causing stunting and leaf distortion in the early infection stage.

We further observed the infected plants to investigate whether the symptoms induced by the pseudorecombinants could be intensified in the late infection stage. To this end, the *N. benthamiana* plants infected with each pseudorecombinant were observed at 12 dpi. Similar to the observation in the early infection stage, the plant inoculated with CMV-209 (W1W2W3) exhibited no obvious symptoms and indistinguishable from healthy plants (Fig. 4B), whereas CMV-Fny (F1F2F3) induced vein clearing, general leaf distortion, and severe mosaic symptoms on the infected systemic leaves at 12 dpi (Fig. 4C). The pseudorecombinants, W1F2F3 and F1W2W3, also did not cause any obvious symptoms as similar to CMV-209 (Fig. 4F and H). However, the pseudorecombinants, W1F2W3 and W1F2F3 containing CMV-Fny RNA2 induced severe leaf distortion and stunting symptoms (Fig. 4E and G).
ceptionally, the plant infected with F1F2W3, showed vein clearing and mild mosaic symptoms (Fig. 4I), while those infected with F1W2W3 developed vein clearing and mild leaf distortion in the late infection stage (Fig. 4D). Our results implied that the compatibility among CMV genomic RNA segments plays a role in symptom expression and, as observed in the cases of W1F2W3 and W1F2F3, pseudorecombination between distinct strains of a virus species can result in emergence of virus progenies carrying higher infectivity or causing severe damages.

In general, disease symptom development of *N. benthamiana* plant induced by CMV-Fny could be divided into two visually distinct stages. Under our plant growth condition, the newly emerged leaves after inoculation of CMV-Fny initially showed vein clearing and general stunting and leaf distortion at 7 dpi (Early infection stage, Fig. 2C) and then the next leaves emerged at 12 dpi showed green mosaic (Late infection stage, Fig. 4C). CMV-Fny RNA1 might play a role in causing vein clearing in the early infection stage, because it was evident that the pseudorecombinants, F1W2W3 and F1F2W3 induced vein clearing while the other recombinants containing W1 did not (Fig. 2). Zhang et al. (1994) also suggested that both RNAs 1 and 2 of CMV-Fny were involved in determining the severity of systemic symptom on tobacco working with CMV-Fny and CMV-LS (Zhang et al., 1994). However, Rao and Franki (1982) handled with eighteen pseudorecombinants constructed *in vitro* from three strains of CMV (CMV-U, CMV-M, and CMV-K) and showed that RNA1 had little effect on symptom induction (Rao and Franki, 1982). The fact that involvement of CMV RNA1 in symptom determination was reported rather rare, except for mapping of the genetic determinant for necrosis of CMV-Ns to the protein 1a (Divéki et al., 2004).

Our results clearly show that CMV-Fny RNA2 played a crucial role in symptom induction in both the early and late infection stage. F1W2F3 caused no obvious symptom (Fig. 2H, 4H) while W1F2W3 induced severe stunting and leaf distortion (E, G, respectively). W1F2W3 and W1F2F3 induced severe stunting and leaf distortion (E, G, respectively). Mock represents healthy plant. Photographs were taken at 12 days after inoculation.

### Table 2. Characteristics of symptoms induced by CMV-209, CMV-Fny and their pseudorecombinants on *Nicotiana benthamiana* plants

| Inoculum | Disease symptom / severity of systemic symptom |
|----------|-----------------------------------------------|
|          | 7 dpi                                         | 12 dpi |
| W1W2W3   | ns/−                                          | ns/−   |
| W1W2F3   | ns/−                                          | ns/−   |
| W1F2W3   | ld, gst/+++                                   | sdl, sst/++++ |
| F1W2F3   | vc/−                                          | vc, mld/+ |
| F1F2F3   | vc, ld, gst/++                                | vc, mld, smo /+++ |
| F1F2W3   | vc, ld, gst/++                                | vc, mmo/++ |
| F1W2F3   | ns/−                                          | ns/−   |
| W1F2F3   | ld, gst/++                                    | sdl, sst/++++ |

*Inoculums of CMV-209, CMV-Fny and their pseudorecombinants were generated by mixing of *in vitro* RNA transcripts.

*Symbols: ns, no obvious symptom; ld, leaf distortion; sdl, severe leaf distortion; mld, mild leaf distortion; gst: general stunting; sst, severe stunting; vc, vein clearing; smo, severe mosaic; mmo; mild mosaic.

−, no obvious symptom; +, mild symptom; ++, intermediate symptom; ++++, severe symptom; +++++, very severe symptom.

### Table 2. Characteristics of symptoms induced by CMV-209, CMV-Fny and their pseudorecombinants on *Nicotiana benthamiana* plants

| Inoculum | Disease symptom / severity of systemic symptom |
|----------|-----------------------------------------------|
|          | 7 dpi                                         | 12 dpi |
| W1W2W3   | ns/−                                          | ns/−   |
| W1W2F3   | ns/−                                          | ns/−   |
| W1F2W3   | ld, gst/+++                                   | sdl, sst/++++ |
| F1W2F3   | vc/−                                          | vc, mld/+ |
| F1F2F3   | vc, ld, gst/++                                | vc, mld, smo /+++ |
| F1F2W3   | vc, ld, gst/++                                | vc, mmo/++ |
| F1W2F3   | ns/−                                          | ns/−   |
| W1F2F3   | ld, gst/++                                    | sdl, sst/++++ |

*Inoculums of CMV-209, CMV-Fny and their pseudorecombinants were generated by mixing of *in vitro* RNA transcripts.

*Symbols: ns, no obvious symptom; ld, leaf distortion; sdl, severe leaf distortion; mld, mild leaf distortion; gst: general stunting; sst, severe stunting; vc, vein clearing; smo, severe mosaic; mmo; mild mosaic.

−, no obvious symptom; +, mild symptom; ++, intermediate symptom; ++++, severe symptom; +++++, very severe symptom.

Fig. 4. Symptoms induced in *N. benthamiana* by CMV-209, CMV-Fny, and their pseudorecombinants constructed *in vitro* from the three genomic segments of the two virus strains. W1W2W3 (CMV-209), F1W2F3, W1W2F3 and F1W2F3 caused mind symptom (B, D, F, H, respectively). F1F2F3 (CMV-Fny) and F1F2W3 caused general stunting, mosaic, and leaf distortion (C, I, respectively). W1F2W3 and W1F2F3 induced severe stunting and leaf distortion (E, G, respectively). Mock represents healthy plant. Photographs were taken at 12 days after inoculation.
al., 1996; Du et al., 2007; Shi et al., 2002; Soards et al., 2002). Recently, the importance of specific domains within the 2b protein for symptom induction has been investigated carefully (Ding et al., 1996; Goto et al., 2007; Lewsey et al., 2009). Lewsey et al. (2009) demonstrated that two nuclear localization signals (NLS1 and NLS2) and the N-terminal domain (5T) of CMV-Fny 2b protein were required and essential for symptom induction, respectively. The C-terminal domain (3T) and two serine residues within a putative phosphorylation domain (KSPSE) of CMV-Fny 2b protein were modulated symptom severity (Fig. 5). The comparison of amino acids sequence of 2b protein of CMV-Fny and CMV-209 revealed that most of various amino acids were located at the N-terminal domain, two of them at NLS1 domain and one at C-terminal domain (Fig. 5). It is worth to determine which amino acid residues are responsible for symptom induction. Interestingly, although CMV-209 caused latent symptom on *N. benthamiana* (Fig. 2B, 4B), the amino acid residues of the 2b protein at position 55 being Leu was conserved in CMV-209 (Fig. 5), which has been reported to be crucial for the RNA silencing suppression and the induction of viral symptoms (Xu et al., 2013). It remains to be examined whether this conserved amino acid residue is actually responsible for suppressing silencing.

Reassortment of genomic segments is one of the mechanisms for genetic variation and new strain generation of multipartite RNA viruses (Morse, 1992). There is evidence that reassortment occurs in natural populations of plant virus (Chen et al., 2007; Fraile et al., 1997; Lin et al., 2004; Maoka et al., 2010). In spite of some studies demonstrated that the reassortment between CMV subgroups was a rare event (Bonnet et al., 2005; Fraile et al., 1997; Jacquemond, 2012), it did not mean that reassortment was not important in CMV evolution (Lin et al., 1996; Roossinck, 2002; White et al., 1995). For example, CMV-Tsh was a natural reassortant between CMV subgroup IA and II strains, in which RNAs 1 and 3 of CMV-Tsh was derived from one or two subgroup II strain(s), while RNA2 was derived from a subgroup I strain, revealed an evolution than its parents. It was probably that RNA2 of subgroup II was replaced by RNA2 of subgroup IA due to its low efficiency in suppression of host defense mechanism during the mixed infection of CMV-Tsh parental viruses (Chen et al., 2007). In the present work, we found that the reassortment between CMV-209 and CMV-Fny caused more severe symptom than wild-type strains (e.g. W1F2W3 and W1F2F3 inducing severe leaf distortion and stunting (Fig. 4E, G). In contrast, W1W2W3 showed latent symptom (Fig. 4B) and F1F2F3 exhibited mosaic, leaf distortion, and general stunting (Fig. 4C)). In addition, we also observed that a pseudorecombinant comprising of RNA1 of CMV-209 and RNAs 2 and 3 of CMV-12, an isolate form Azuki bean, showed more severe symptom than its parents (data not shown). These results indicated that natural reassortment between compatible viruses can result in emergence of novel viruses causing severe damages in crop fields.

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