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Cymbidium mosaic potexvirus isolate-dependent host movement systems reveal two movement control determinants and the coat protein is the dominant

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ABSTRACT

Little is known about how plant viruses of a single species exhibit different movement behavior in different host species. Two Cymbidium mosaic potexvirus (CymMV) isolates, M1 and M2, were studied. Both can infect Phalaenopsis orchids, but only M1 can systemically infect Nicotiana benthamiana plants. Protoplast inoculation and whole-mount in situ hybridization revealed that both isolates can replicate in N. benthamiana; however, M2 was restricted to the initially infected cells. Genome shuffling between M1 and M2 revealed that two control modes are involved in CymMV host dependent movement. The M1 coat protein (CP) plays a dominant role in controlling CymMV movement between cells, because all chimeric CymMV viruses containing the M1 CP systemically infected N. benthamiana plants. Without the M1 CP, one chimeric virus containing the combination of the M1 triple gene block proteins (TGBps), the M2 5′ RNA (1-4333), and the M2 CP effectively moved in N. benthamiana plants. Further complementation analysis revealed that M1 TGBp1 and TGBp3 are co-required to complement the movement of the chimeric viruses in N. benthamiana. The amino acids within the CP, TGBp1 and TGBp3 which are required or important for CymMV M2 movement in N. benthamiana plants were mapped. The required amino acids within the CP map to the predicted RNA binding domain. RNA-protein binding assays revealed that M1 CP has higher RNA binding affinity than does M2 CP. Yeast two-hybrid assays to detect all possible interactions of M1 TGBp1 and CP, and only TGBp1 and CP self-interactions were observed.

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Introduction

Viruses have a defined, restricted host range, which varies among different viruses, or sometimes even different isolates of the same virus species. For example, Cucumber mosaic virus (CMV) has a host range of more than 1000 plant species in 85 families, including monocotyledonous and dicotyledonous plants, and by contrast, Barley stripe mosaic virus (BSMV) has a confined natural host to barley (ICTVDB: The Universal Virus Database of the International Committee on Taxonomy of Viruses [http://www.ictvdb.iacr.ac.uk/ictv/fr-index.htm]; [Timian, 1974]). Disease may occur when a particular virus adapts and/or shifts to a new host, for which there are many examples in plants and even humans (e.g., severe acute respiratory syndrome).

Successful infection of a plant host requires that plant viruses achieve (i) replication in the initially infected cells, (ii) movement through plasmodesmata to adjacent cells from the initial infected site and (iii) long-distance spread in a host through the vascular tissue. Plant viruses possess only limited genetic information; hence, to complete their infection cycles, interactions between virus proteins and necessary host factors is critical. For a virus to successfully infect different plant hosts, the limited number of virus-encoded proteins must have the ability to recognize and interact with factors in different hosts for a successful infection. For some plant viruses, host plants restrict viral cell-to-cell movement rather than viral replication, which results in a subliminal infection (Cheo, 1970; Sulzinski and Zaitlin, 1982). In other cases, viruses may move from cell-to-cell, but the plant prevents viral long-distance spread (Fujita et al., 1996; Saenz et al., 2002; Wang et al., 1998). Failure to accomplish any of the stages can be thought of as resistance.

In our previous studies, we collected several isolates of Cymbidium mosaic virus (CymMV) for construction of virus-induced gene silencing vectors for orchid functional genomic studies (Lu et al., 2007). CymMV is a prevalent orchid virus and belongs to the genus Potexvirus in the family Flexiviridae (Adams et al., 2005b). The recorded natural host range is largely limited to species belonging to Orchidaceae (ICTVDB: The Universal Virus Database of the International Committee on Taxonomy of Viruses [http://www.ictvdb.iacr.ac.uk/ictv/fr-index.htm];[Adams et al., 2005a; Zettler et al., 1990]). Interestingly, we found two distinct CymMV isolates, M1 and M2, both of which can infect Phalaenopsis orchids, but only M1 can systemically infect Nicotiana benthamiana plants. Because both isolates can systemically infect orchids, this suggests that viral-encoded
replication and movement proteins are functional in orchids. However, because only M1 can systemically infect *N. benthamiana* plants this suggests that M1-encoded proteins could interact successfully with factors in both plant species to allow a systemic infection, but M2-encoded proteins do not. M1 and M2 share very high nucleotide sequence identity (97%), which suggests that subtle differences contribute to the differences in host reaction.

Viruses in the genus *Potexvirus* have monopartite, positive-sense single-strand RNA genomes (Adams et al., 2005a). Studies of potexviruses have contributed importantly to our understanding of the complexity of the mechanisms of plant virus movement within plants (Lucas, 2006; Verchot-Lubicz, 2005; Verchot-Lubicz et al., 2007). Comprehensive studies of potexviruses has revealed that triple gene block proteins (TGBp) and coat proteins (CP) work cooperatively to promote viral movement between cells and long-distance in their plant hosts (Bayne et al., 2005; Beck et al., 1991; Chapman et al., 1992; Fedorkin et al., 2001; Forster et al., 1992; Ju et al., 2007; Krishnamurthy et al., 2003; Lin et al., 2006; Lough et al., 1998, 2000; Mitra et al., 2003; Tamai and Meshi, 2001; Verchot-Lubicz, 2005). The current model suggests that TGBp1 increases the plasmodesmal size-exclusion limit and moves into neighboring cells to suppress RNA silencing (Bayne et al., 2005; Howard et al., 2004); both TGBp1 and CP bind viral RNA (vRNA) forming viral ribonucleoprotein complexes (vRNP; Karpova et al., 2006; Lough et al., 2000, 2001); the complex is then delivered by TGBp2 induced vesicles (containing TGBp3) to plasmodes mata, and through plasmodes mata to neighboring cells; later, after the free form of TGBp1 binds membrane-associated TGBp2 and/or TGBp3, then TGBp1 (on the vRNP complex) binds with the membrane-associated TGBp1 complex (TGBp1 with TGBp2 or TGBp3) to form a membrane-bound complex and begin new rounds of replication (Verchot-Lubicz, 2005).

Interactions between TGBp1 and CP (Karpova et al., 2006; Lough et al. 2000), and TGBp1 self-interactions (Samuels et al., 2007) have been reported, and those between TGBp1 and TGBp2, and between TGBp1 and TGBp3 have been suggested (Karpova et al., 2006; Krishnamurthy et al., 2002; Lough et al., 2000; Verchot-Lubicz, 2005; Yang et al., 2000). Interestingly, trans-complementation of TGBp between potexvirus species was not successful, and it was suggested that species-specific interactions among potexvirus movement proteins are obligatory for cell-to-cell movement (Lin et al., 2006). However, trans-complementation of movement defective potexviruses by movement proteins derived from *Tobamovirus*. *Crucifer tobamovirus*, *Nicotiana benthamiana* and *Red clover necrotic mosaic dianthovirus* have been reported (Morozov et al., 1997).

Although the movement of plant viruses, including potexviruses, has been extensively studied, little is known about how plant viruses of a single species exhibit different movement behavior in different host species. In this study, we aimed to use our CymMV-M1, -M2 and *N. benthamiana* system to study the molecular basis of CymMV isolate-dependent host movement determinants, and to resolve the mechanism involved in CymMV translocation. Our studies not only help to identify the molecular determinants of CymMV isolate-dependent host movement, but other phenomena that are not easily resolved by studies using a single isolate. Finally, our efforts revealed two control modes.

**Results**

**Construction of CymMV infectious clones and derived expression vectors**

The construction of the infectious clone of the isolates M1 (pCymMV-M1, Fig. 1) and its derived expression vector (pCymMV-M1-GFP, Fig. 1) engineered with the green fluorescent protein (GFP) was previously described (Lu et al., 2007). We followed the same strategy to construct the M2 infectious clone (pCymMV-M2, Fig. 1) and its derived expression vector (pCymMV-M2-GFP, Fig. 1). *Phalaenopsis amabilis* var. *formosa* and *Nicotiana benthamiana* plants were inoculated with transcripts from pCymMV-M1 and pCymMV-M2, and nucleic acids were extracted from the inoculated and upper, non-inoculated leaves of pCymMV-M1 and pCymMV-M2-inoculated plants 14 days post-inoculation. Similar to our previous findings, both pCymMV-M1 and pCymMV-M2 systemically infected *Phalaenopsis* orchids (Fig. 1B), but only pCymMV-M1 systemically infected *N. benthamiana* (Fig. 1C).

**Replication of CymMV isolates in *N. benthamiana* protoplasts**

To test whether CymMV-M2 can replicate in *N. benthamiana*, protoplasts were inoculated with transcripts from pCymMV-M1, pCymMV-M2, pCymMV-M1-GFP and pCymMV-M2-GFP. Illumination of GFP was observed from pCymMV-M1-GFP and pCymMV-M2-GFP.
infected protoplasts at 14 h post-inoculation (Fig. 2). Northern blot hybridization also indicated that pCymMV-M1 and pCymMV-M2 replicated in N. benthamiana protoplasts (Fig. 3I).

Subliminal infection of CymMV-M2 in N. benthamiana

To next analyze why M2 failed to accomplish systemic infection in N. benthamiana plants, more plants were inoculated with transcripts from pCymMV-M1-GFP and pCymMV-M2-GFP. Because of limited expression of GFP of both pCymMV-M1-GFP and pCymMV-M2-GFP in plants of N. benthamiana, we therefore inoculated N. benthamiana with transcripts of pCymMV-M1 and pCymMV-M2 and assayed by whole-mount in situ hybridization 14 days post-inoculation. pCymMV-M1 moved from the inoculation foci and infected a broader area of leaves (Fig. 2B), whereas pCymMV-M2 was limited to the initial inoculated cells and did not spread from cell to cell, thus resulting only in subliminal infections (Fig. 2B).

Genome shuffling between CymMV-M1 and -M2 for complementation analysis

PVX TGBps 1–3 and CP proteins are known to be required for virus cell-to-cell movement (Bayne et al., 2005; Chapman et al., 1992; Krishnamurthy et al., 2002, 2003; Lough et al., 2000; Morozov and Solovyev, 2003; Tamai and Mashi, 2001; Verchot-Lubicz, 2005). To determine if specific M1 gene(s) can complement M2 for virus movement, we constructed a series of M1 and M2 chimeric viruses (Fig. 3). N. benthamiana protoplasts and plants were inoculated with transcripts derived from each construct. Northern blot hybridization revealed that each virus replicated in N. benthamiana protoplasts (Fig. 3I), and statistics analysis (ANOVA) of real-time RT-PCR quantification of average relative percentage (from 3 independent experiments) of CymMV RNA from CymMV clone-infected protoplasts at 24 h post-inoculation revealed no significant difference in percentage between these clones ($P = 0.88$).

Systemic infections were monitored by performing RT-PCR on leaves distal from the inoculation site at 2 weeks post-inoculation (Figs. 3A–H). These analyses showed that if the CymMV construct contained the M1 CP, the virus systemically infected N. benthamiana plants (Figs. 3A, C, E, H). One notable exception was the pCymMV-M2/TGB-M1. This construct contained the M2 CP and 5′ RNA sequence (1–4333; include the whole RNA-dependent RNA polymerase encoding region), but also the M1 triple gene block ORFs (Fig. 3F).

To identify whether TGBps 1–3 were all necessary to complement the systemic infection of pCymMV-M2/TGB-M1, we performed genome shuffling between M1 and M2 TGBps 1–3, and constructed chimeric viruses (Figs. 4A–H). N. benthamiana protoplasts and plants were inoculated with transcripts derived from each construct. All constructs were replication competent in protoplasts as assayed by northern blot hybridization (Fig. 4I), and statistics analysis (ANOVA) of real-time RT-PCR quantification of average relative percentage (from 3 independent experiments) of CymMV RNA from CymMV clone-infected protoplasts at 24 h post-inoculation revealed no significant difference in percentage between these clones ($P = 0.91$).

Systemic infection, as detected by RT-PCR at 2 weeks post-inoculation (Figs. 4A–H), indicated that only chimeric viruses containing the M1 TGBp1 and TGBp3 moved systemically (Fig. 4G). Because these results were complicated, we used the Karnaugh Maps (Harrison, 1969) tabular logical calculation to explore possible explanations (data not shown). The results of these analyses are summarized in Fig. 5. The analysis revealed that two control modes, which we termed dominant and matching control modes, are involved in CymMV movement. The M1 CP plays a dominant role in controlling the trafficking of the infection agents within plants. Without the M1 CP, another control mode, which required correct
matching of the particular movement accessory components including the M2 5′ RNA (1–4333), M1 TGB p1, M1TGBp3, and M2 CP was also identified.

**Mapping the amino acids on CymMV CP that are important for systemic infection of N. benthamiana plants**

Our studies indicated that the M1 CP plays a major role in determining CymMV movement in *N. benthamiana*. Because M1 and M2 share a very high nucleotide sequence identity (97%), we attempted to identify the individual, important amino acids of the CP that allowed for the CymMV systemic infection of *N. benthamiana* plants. Sequence alignment of the M1 and M2 CPs revealed only four amino acid differences (Figs. 6A, B). Site-directed mutagenesis was then performed to construct a series of CymMV-M2 CP mutants containing specific M1 CP amino acids (Figs. 6C–I). All constructs were replication competent in protoplasts as assayed by northern blot hybridization (Fig. 6J), and statistics analysis (ANOVA) of real-time RT-PCR quantification of average relative percentage (from 3 independent experiments) of CymMV RNA from CymMV clone-infected protoplasts at 24 h post-inoculation revealed no significant difference in percentage between these clones (*P* = 0.83). Transcripts derived from each construct were used to inoculate *N. benthamiana* plants. These analyses showed that the CP amino acid G82A and L89P changes were both required for M2 to systemically infect *N. benthamiana* plants (Figs. 6C, H). Interestingly, these amino acids are located within the previously predicted RNA binding domain of the CP (Fig. 7A).

Fig. 3. Schematic representation of genome organization and infectivity assay of the parental CymMV-M1 and CymMV-M2 and the derived chimeric constructs. (A–H). Rectangles represent open reading frames encoded by CymMV genomic RNA, RNA-dependent RNA polymerase (RdRp), triple gene block (TGB) ORFs 1, 2, and 3 and capsid protein (CP). Sequences corresponding to pCymMV-M1 and pCymMV-M2 are indicated by gray and white rectangles, respectively. The restriction enzyme sites for constructing chimeric viruses are indicated. (Although HpaI sites are located 129 nt downstream of CP translation start sites, the amino acid sequences in the regions between M1 and M2 are identical.) Clones competent in protoplast accumulation and systemic infection in *N. benthamiana* are indicated by +, and the ratio of systemic infected to total inoculated plants is indicated. Systemic infection was detected 2 weeks post-inoculation by RT-PCR. (I) Protoplast infectivity was detected 24 h post-inoculation by northern blot hybridization, and the ribosomal RNA used for a loading control are indicated. Genomic RNA (*G*), TGBp, and CP subgenomic RNA are indicated. The pCymMV-R used as a negative control is illustrated in Fig. 1. The average percentage of relative real-time RT-PCR quantification (from 3 independent experiments) of CymMV RNA from CymMV clone-infected protoplasts at 24 h post-inoculation is indicated. The accumulation of pCymMV-M1 was set at 100% for relative quantification. Numbers at the left correspond to positions of marker RNAs (sizes in 1000 nucleotides) analyzed in the same gel.
It has been previously reported that the CP is involved in forming ribonucleoprotein complexes and is an important step for potexvirus movement (Karpova et al., 2006; Lough et al., 2000, 2001). Our results indicated that the important amino acids of the CP that allowed for the CymMV systemic infection are located within the previously predicted RNA binding domain (Fig. 7A; 1). Thus, it might be that the binding affinity between CPs and RNAs of CymMV M1 and M2 may play roles in CymMV movement, and therefore we attempted to analyze and compare their binding affinities. Because the whole genomic RNA of CymMV is 6227 nucleotides in length, which made conventional gel-shift assays difficult, we performed modified RNA–protein pull-down assays for our analyses. Our data indicated that CymMV M1 CP has a higher binding affinity as compared to CymMV M2 CP (Fig. 7D, lanes 1–4). We then also performed CP and CymMV RNA binding assays on CymMV M2 CP with the amino acid changes G82A and L89P (derived from clone pCymMV-M2-CP-GL/AP; Fig. 6H). The results indicated that with these amino acid changes (G82A and L89P), the binding affinity was higher as compared to wild-type CymMV-M2 CP (Fig 7D, lanes 3, 4, 5, and 6).

Interaction analysis of TGBps by use of yeast two-hybrid assay

Our analysis also indicated that both M1 TGBp1 and TGBp3 were co-required for CymMV M2 chimeric viruses to move systemically in *N. benthamiana* plants (Fig. 4G), and was consistent with previous suggestions indicating interactions between potexvirus TGBp1 and TGBp3 (Lucas, 2006; Verchot-Lubicz, 2005; Verchot-Lubicz et al., 2007). Therefore, we attempted to determine if specific interactions occurred within and between CymMV TGBps and CP by use of the
and H at TGBp1 positions 44 and 94 were required for M2 (pCymMV-M2) to systemically infect N. benthamiana, either individually or simultaneously (Figs. 8I, J). Similarly, site-directed mutagenesis was conducted on pCymMV-M2/TGB1-M1 (Fig. 4C) and pCymMV-M2/TGB3-M1 (Fig. 4D). It has been previously reported that in systemic hosts of PVX, when the binding between CP-vRNA is weaker, the need for TGBps in systemic infection was analyzed by RT-PCR (Fig. 8). The chimeric viruses with Q8L and/or S71A in M2 TGBp3 did move systematically, the ratio of systemically infected plants was much reduced (Figs. 8G, H).

Mapping the amino acids on TGBp1 and TGBp3 of CymMV that are important for systemic infection of N. benthamiana

Sequence alignment analysis revealed 4 and 2 amino acid differences within the TGBp1 and TGBp3 between CymMV-M1 and CymMV-M2, respectively (Figs. 8A, B). Loss-of-function assays were performed by using site-directed mutagenesis to substitute TGBp1 and TGBp3 with the 4 and 2 amino acids derived from M2 TGBp1 and TGBp3, respectively (Figs. 8C–H). N. benthamiana plants were inoculated with transcripts derived from each construct, and systemic infection was analyzed by RT-PCR (Fig. 8). The chimeric viruses with either amino acids Y44H and H94R changes in M1 TGBp1 failed to move systematically (Figs. 8C, D). Although the chimeric viruses with either amino acids L8Q or A71S changes in the M1 TGBp3 did move systematically, the ratio of systemically infected plants was much reduced (Figs. 8G, H).

To further characterize whether Y44 and/or H94 in TGBp1 are required for M2 to systemically infect N. benthamiana plants, site-directed mutagenesis was conducted on pCymMV-M2/TGB3-M1 (Fig. 4E) to construct M2 clones substituted with H44Y and R94H at TGBp1, either individually or simultaneously (Figs. 8I–K). Both Y and H at TGBp1 positions 44 and 94 were required for M2 (pCymMV-M2/TGB3-M1) to systemically infect N. benthamiana plants (Fig. 8K). Mutants having only one of these changes failed to move systematically (Figs. 8I, J). Similarly, site-directed mutagenesis was conducted on pCymMV-M2/TGB1-M1 (Fig. 4C) and pCymMV-M2/TGB3-M1 (Fig. 4D) to construct clones substituted with Q8L and/or S71A at TGBp3 (Fig. 8L–O). Both Q8L and/or S71A in M2 TGBp3 were important but not required for M2 to systemically infect N. benthamiana. Mutants having only one of these changes moved systematically, but the ratio of systemically infected plants was much reduced (Figs. 8G–O). The Y44 (position 44) and H94 in TGBp1 are located within the previous predicted NTP helicase domain but not in the identified conserved motifs (Kadare and Haenni, 1997; Kalinina et al., 2002; Leshchiner et al., 2006), and L8 and A71 in TGBp3 are also not located in the previously identified transmembrane domain (Fig. 9B, 21).

All constructs used in Fig. 8 were replication competent in protoplasts as assayed by northern blot hybridization (Fig. 8P), and statistics analysis (ANOVA) of real-time RT-PCR quantification of the average relative percentage (from 3 independent experiments) of CymMV RNA from CymMV clone-infected protoplasts at 24 h post-inoculation revealed no significant difference in percentage between these clones (P = 0.92).

Discussion

Our studies comparing CymMV isolate-dependent host movement provides new and important information on the molecular basis of how viruses of the same species can move in different host plants. We also demonstrate complex aspects of potexvirus movement not easily identified by studies using a single isolate. The more detailed resolution of these phenomena, as we demonstrate in this paper, will further help to dissect the mechanisms of potexvirus movement in plants.

Our data indicated that M1-encoded CP plays a major role for CymMV in systemic infection of N. benthamiana plants (Fig. 5). In all our CymMV clones, if the constructs contained the M1 CP, the virus systemically infected N. benthamiana plants (Figs. 3A, C, E, H). Interestingly, our data also showed that if CP was not properly functional (e.g., M2 CP), alterations in TGBps (e.g., M1 TGBp1 and 3) may compensate for the function of CP and allow CymMV to systemically infect N. benthamiana plants (Fig. 3F). In addition, when comparing pCymMV-M2/TGB-M1 and pCymMV-M1/CP-M2 (Figs. 3F and G), only pCymMV-M2/TGB-M1 was movement competent. The TGBps and CP are the same in both chimeric viruses. These data indicate that another control mode which required correct matching of the particular movement accessory components (M2 5′ RNA, TGBp1 and TGBp3 and M2 CP) also allowed the systemic infection of CymMV in N. benthamiana (Fig. 3F and Fig. 5). Thus, our data suggested two control modes are involved in CymMV movement in N. benthamiana.

The nature of the CymMV vRNP that moves between cells remain a subject for study (Verchot-Lubicz et al., 2007), and two forms, one is the linear vRNP formed by TGBp1, vRNA and CP (Lough et al., 2000) and another is virion or altered virion with a single-tailed particle comprising RNA, CP and TGBp1 (Cruz et al., 1998; Karpova et al., 2006). However, no matter what the proposed form of vRNP that moves between cells, all data suggest that potexvirus CP and vRNA are involved in forming the vRNP which traffics to the plasmodesmata. Our analysis indicated that the CymMV CP-vRNA binding affinity likely plays an important role in CymMV movement in N. benthamiana plants. Therefore, it is possible that the binding between CP-vRNA may affect the nature of vRNP; thus affecting the movement of CymMV in different hosts.

Our data also suggested that different combinations of CP-vRNA required different degrees of help from TGBps for movement (Figs. 3, 4). When the binding between CP-vRNA is higher (Fig. 7D), the need for TGBs in CymMV movement is more flexible (Figs. 3A, C, E, H); however, when the binding between CP-vRNA is weaker, the need for TGBs in movement is critical and only certain combinations of CymMV movement accessory components allowed for CymMV movement (Figs. 3B, F, 4G). It has been previously reported that in systemic hosts of PVX, the movement of TGBps are host dependent (Krishnamurthy et al., 2002; Yang et al., 2000). Because in different hosts the cellular conditions and morphology are different, it is possible that potexviruses may use different strategies to adapt to different hosts. Adjusting movement accessory components could be an efficient way to generate the diversity of functions in response to changes of cellular conditions and provide flexibility for virus movement in different hosts, and thus could be an advantage for viruses with modular designed movement proteins (Morozov and Solovyev, 2003).
Although interactions between potexvirus TGBp1 and TGBp2 and between TGBp1 and TGBp3 have been suggested (Karpova et al., 2006; Krishnamurthy et al., 2002; Lough et al., 2000; Verchot-Lubicz, 2005; Yang et al., 2000), the interactions have yet to be proven. Interestingly, our data showed that CymMV M1-encoded TGBp1 and TGBp3 are co-required for M2 systemic infection of *N. benthamiana* (Fig. 4G); which support the hypothesis for interactions between TGBp1 and TGBp3. However, the results of our yeast two-hybrid analysis were similar to those that have been recently reported showing that TGBp1 was self-associated, but showed no interactions between TGBp1 and TGBp2 or TGBp3 (Samuels et al., 2007). Thus, the interactions may not occur physically, or the interactions could be transient and not easily detected.

**Fig. 6.** Schematic representation of capsid protein (CP) mutants and infectivity assays. (A–I). Rectangles represent open reading frames encoded by CymMV genomic RNA, RNA-dependent RNA polymerase (RdRp), triple gene block ORFs 1, 2, and 3 and capsid protein (CP). Black and gray lines in the CP regions indicate the different amino acids found between pCymMV-M1 and pCymMV-M2, respectively. Position of amino acids is indicated. Chimeric viruses competent in protoplast accumulation and systemic infection in *N. benthamiana* plants are indicated by +, and the ratio of systemic infected/total inoculated plants is indicated. Systemic infection was detected 2 weeks post-inoculation by RT-PCR. (J) Protoplast infectivity was detected 24 h post-inoculation by northern blot hybridization, and the ribosomal RNA used for a loading control are indicated. Genomic RNA (G), TGBp, and CP subgenomic RNA are indicated. The pCymMV-R- used for a negative control is illustrated in Fig. 1. The pCymMV-V- used as a negative control is illustrated in Fig. 1. The average percentage of relative real-time RT-PCR quantification (from 3 independent experiments) of CymMV RNA from CymMV clone-infected protoplasts at 24 h post-inoculation is indicated below the gels. The accumulation of pCymMV-M1 was set at 100% for relative quantification. Numbers at the left correspond to positions of marker RNAs (sizes in 1000 nucleotides) analyzed in the same gel.

Within the potexvirus TGBp1, the NTPase/helicase domains have been predicated, and 7 conserved regions including two canonical motifs of NTPase, DEY and GKS, were identified (Fig. 9; Kadare, and Haenni, 1997; Kalinina et al., 2002; Leshchiner et al., 2006). Although, currently the roles of NTPase/helicase domains in potexvirus movement have not been well established (Verchot-Lubicz et al., 2007), mutations eliminating the DEY motif affects the protein subcellular targeting and mutations eliminating the DEY and GKS motifs inhibit the ability of TGBp1 to increase plasmodesmata size-exclusion limits as well as virus movement (Angell et al., 1996; Lough et al., 1998; Morozov et al., 1999). The amino acids identified in CymMV TGBp1 that are required for CymMV-M2 movement in *N. benthamiana* are within the NTPase/helicase domain, but not in the conserved motifs (Fig. 9). The amino acids are also not in the positions within the NTPase/helicase domain previously reported as essential for Bamboo mosaic potexvirus movement (Lin et al., 2004). Within the potexvirus TGBp3, a transmembrane domain has been previously reported. PVX
with substitution mutations within the transmembrane domain is restricted to a single cell, and PVX with mutations outside the transmembrane domain show reduced movement between cells and may not move systemically (Krishnamurthy et al., 2003). The amino acids identified in TGBp3 which are important for CymMV-M2 movement in N. benthamiana are not located in the transmembrane domain (Fig. 9; Krishnamurthy et al., 2003). Because all amino acids identified in CymMV TGBp1 and TGBp3 are not located in the conserved motifs previously identified among potexviruses, nor within the amino acids reported as essential for potexvirus movement, we speculate that changes in these regions may only slightly affect but not abolish the function (as we see in the case of CymMV-M2 CP) of TGBp1 and TGBp3, and the differential function required for virus movement is host dependent. This may explain why M1 and M2 can both systemically infect Phalaenopsis orchid, but only M1 can systemically infect N. benthamiana.

Fig. 7. The amino acid sequence alignment of predicted RNA binding domain of potexvirus CPs and slot-blot detection of biotin-labeled CymMV RNAs. (A) The amino acid sequence alignment of predicted RNA binding domain were conducted by use of clustal X 1.83 (Thompson et al., 1997). The previously identified conserved positive charged amino acids are indicated by stars (Abouhaidar and Lai, 1989). The arrows indicate the amino acids important for pCymMV-M2 to systemically infect N. benthamiana. The viruses, abbreviation and accession number used in alignment are described below: Cymbidium mosaic virus (CymMV, accession number AY571289); Potato aucuba mosaic virus (PAMV, accession number NC_003632); Narcissus mosaic virus (NMV, accession number NC_001441); Scallion virus X (ScaVX, accession number NC_003400); Papaya mosaic virus (PapMV, accession number NC_001748). (B). The concentration of recombinant CPs used in this experiment derived from clones pCymMV-M1 (M1-CP; Fig. 1), -M2 (M2-CP; Fig. 1) and -M2-CP-GL/AP (M2-CP-GL/AP; Fig. 6) and Glutathione-S-transferase (GST) derived from clones (pGEX 2T-1, Pharmacia Biosciences, Inc., New Jersey, USA) are shown. (C) Control experiments using M1 and M2 biotin-labeled RNA only (lane 1 and 2, respectively), or M1 and M2 RNA incubated with GST (lane 3 and 4, respectively) are shown. Because no RNA–protein binding occurred in these control experiments, RNA was detected only in S. (D) The results of experiments using RNA–protein combinations, M1 RNA/M1-CP (lane 1), M2 RNA/M1-CP (lane 2), M1 RNA/M2-CP (lane 3), M2 RNA/M2-CP (lane 4), M1 RNA/M2-CP-GL/AP (lane 5) and M2 RNA/M2-CP-GL/AP (lane 6), are shown. Different concentrations of sodium chloride (in millimolars; mM) added in incubation buffer are indicated. For convenience, the RNA–protein combinations used in the experiments are also indicated in the closed boxes. We repeated this experiment three times and one result is shown. The average S/P ratios were derived from the average of three independent experiments, and data were analyzed by Dunnett’s T test. *indicates significant difference (P<0.01).
It is worth noting that correct matching of the particular movement accessory components also includes the CymMV 5' RNA sequence and/or the RNA-dependent RNA polymerase (RdRp). The potexvirus RdRp has not been reported to be required for virus movement, however, it has been suggested that the 5' untranslated region of the potexvirus genomic RNA plays a role in viral cell-to-cell movement (Lough et al., 2006).

Both CymMV-M1 and -M2 replicated well in N. benthamiana protoplasts or in Phalaenopsis orchids. It seems more likely that RNA rather than the RdRp plays a role in CymMV movement. However, recently it was reported that the TMV replicase and movement protein function together in altering plasmodesmat (Guenoune-Gelbart et al., 2008). Therefore, we cannot rule out the possibility that the RdRp plays a role in controlling CymMV movement in N. benthamiana plants.

### Materials and Methods

#### RNA extraction and northern blot hybridization

RNA was extracted from plants as described for northern blot analysis and RT-PCR (Tian et al., 1996). T7 RNA polymerase and HpaI-digested pCymMV-M1 plasmids (CymMV probe; corresponding to the 590 nt of CymMV at its 3’ end) were used to generate the negative-sense DIG-labeled probes (Roche Applied Science; Mannheim, Germany). Northern blot hybridization was performed as described (Klaassen et al., 1996), and hybridization signals were detected by use of the chemiluminescent substrate CDP STAR (Roche Applied Science) and exposing blots to Fuji medical X-ray film (Fuji, Tokyo, Japan).

#### Construction of CymMV infectious clones and derived expression vectors

The construction of pCymMV-M1 and pCymMV-M1-green fluorescent protein (Fig. 1A) was as described (Lu et al., 2007). The same approaches were used to construct pCymMV-M2 and pCymMV-M2-GFP. Both pCymMV-M1 and -M2 had been completely sequenced. pCymMV-M1-R is a spontaneous mutation clone obtained during the cloning of pCymMV-M1. pCymMV-M1-R contains a mutation in the RNA-dependent RNA polymerase (RdRp) region, which causes pre-termination of RdRp (Fig. 1A). The 5’ mutated RdRp region of the pCymMV-M1-R has been sequenced.

### Table 1

Determination of interactions among CymMV encoded triple gene block proteins by yeast two-hybrid assay.

| AD fusiona | BD fusiona | Trp− and Leu− mediumb | Trp−, Leu−, His− and Ade− mediumc | Histochemical assayd | Histochemical assayd (X-gal) |
|------------|------------|------------------------|----------------------------------|----------------------|-----------------------------|
| T-antigen  | p533      | + + +                  | + + +                            | + + +                |
| TGBp1      | *          | + − +                  | + − +                            | − − −                |
| TGBp2      | *          | + − +                  | + − +                            | − − −                |
| TGBp3      | *          | + − +                  | + − +                            | − − −                |
| CP         | *          | + − +                  | + − +                            | − − −                |
| *          | TGBp1 1    | + − +                  | + − +                            | − − −                |
| *          | TGBp2 1    | + − +                  | + − +                            | − − −                |
| *          | TGBp3 1    | + − +                  | + − +                            | − − −                |
| *          | CP         | + − +                  | + − +                            | − − −                |
| TGBp1 1    | TGBp2 1    | + + +                  | + + +                            | + + +                |
| TGBp1 1    | TGBp3 1    | + + +                  | + + +                            | + + +                |
| TGBp1 1    | CP         | + + +                  | + + +                            | + + +                |

a “−”, plasmid expression AD domain (pGADT7) and BD domain (pGBK7) are added.
b “+”, yeast colonies that grew on SD medium (Lacking Trp and Leu), “−”, yeast colonies that did not grow on SD medium (lacking Trp and Leu).
c “+”, yeast colonies that grew on SD medium (Lacking Trp, Leu, His and Ade), “−”, yeast colonies that did not grow on SD medium (Lacking Trp, Leu, His and Ade).
d “+”, yeast colonies that were blue, “−”, yeast colonies that did not grow and no blue histochemical stain was observed.
e Plasmids containing T-antigen-AD and murine p53-BD were provided by the manufacturer and used as positive controls.

#### Construction of CymMV recombinant clones

The infectious clones pCymMV-M1 and pCymMV-M2 (Fig. 1A) were digested with Nael and NheI individually. Two digested fragments derived from each reaction were gel purified by use of a gel extraction kit (Qiagen; Hilden, Germany); the small fragments derived from pCymMV-M1 and -M2 were exchanged and ligated by use of T4 DNA ligase (Promega; Madison, WI, USA) to construct pCymMV-M1/RdRp-M2 and pCymMV-M2/RdRp-M1 (Figs. 3C and D). The gel purification and ligation conditions followed the manufacturer’s manual. The construction of pCymMV-M1/TGB-M2, pCymMV-M1/TGB-M2, pCymMV-M1/CP-M2 and pCymMV-M1/CP-M1 clones (Figs. 3E-H) was similar to that of pCymMV-M1/RdRp-M2 and pCymMV-M2/RdRp-M1, except the restriction enzymes Nael and HpaI were used to construct pCymMV-M1/TGB-M2 and pCymMV-M2/TGB-M1, and the restriction enzymes HpaI and SalI were used to construct pCymMV-M1/CP-M2 and pCymMV-M2/CP-M1. All recombinant clones were sequenced in the modified region to ensure that the clones were correct.

#### Construction of TGBps recombinant clones

pCymMV-M1 and pCymMV-M2 were used as the initial template, and the primer pairs CymMV F3783/CymMV-TGB1 R and CymMV-TGB1 F/CymMV CPR (Supplementary Table S1) were used, respectively in the PCR reactions to amplify the two overlapping fragments. The amplified fragments were gel purified and mixed together at a ratio of 1:1, then used in another PCR reaction. The PCR reaction cycles were 94 °C for 5 min for 1 cycle, then 94 °C for 30 s, 55 °C for 30 s and 72 °C for 2 min for a total of 5 cycles. Then the primer pair CymMV F3783/CymMV CPR was added for another 30 cycles. The PCR cycles were 94 °C for 5 min for 1 cycle, then 94 °C for 30 s, 55 °C for 30 s and 72 °C for 2 min for a total of 30 cycles. The amplified products were digested with Nael and HpaI (Fig. 3), and then separated on a 1% agarose gel to purify the 1.3-kb fragment. The pCymMV-M2 was digested with Nael and HpaI, then separated on a 1% agarose gel to purify the 8-kb fragment. Both digested fragments were gel purified and mixed together at a ratio of 1:1, then used in another PCR reaction. The PCR reaction cycles were as described above. The amplified fragments were gel purified and mixed together at a ratio of 1:1, then used in another PCR reaction. The PCR reaction cycles were as described above. The amplified products were digested with Nael and HpaI (Fig. 3), and then separated on a 1% agarose gel to purify the 1.3-kb fragment. The pCymMV-M2 was digested with Nael and HpaI, then separated on a 1% agarose gel to purify the 8-kb fragment. Both digested fragments were ligated to construct

#### Site-directed mutagenesis

Each of the mutated clones derived from pCymMV-M1/M2 with amino acid substitution(s) in the CP, TGBp1 and TGBp3 (Figs. 6, 8) was constructed by site-directed mutagenesis as described (Lu et al., 2007). pCymMV-M1 was used as the initial template, and the primer pairs CymMV F3783/TGB1 Y444H R and TGB1 Y444H F/CymMV CPR were used in the first PCR reactions. The amplified fragments were gel purified and mixed together at a ratio of 1:1, then used in another PCR reaction. The PCR reaction cycles were as described above. The amplified products were digested with Nael and HpaI (Fig. 3), and then separated on a 1% agarose gel to purify the 1.3-kb fragment. The pCymMV-M2 was digested with Nael and HpaI, then separated on a 1% agarose gel to purify the 8-kb fragment. Both digested fragments were ligated to construct

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Fig. 8. Schematic representation of TGBp1 mutants and infectivity assay between pCymMV-M1 and pCymMV-M2. (A–O) Rectangles represent ORFs encoded by CymMV genomic RNA, RNA-dependent RNA polymerase (RdRp), triple gene block ORFs 1, 2, and 3 and capsid protein (CP). Black and gray lines indicate different amino acids between pCymMV-M1 and pCymMV-M2, respectively. Positions of amino acids are indicated. Chimeric viruses competent in protoplast accumulation and systemic infection in N. benthamiana plants are indicated by +, and the ratio of systemic infected to total inoculated plants is indicated. Systemic infection was detected 2 weeks post-inoculation by RT-PCR. (P) Protoplast infectivity was detected 24 h post-inoculation by northern blot hybridization, and the ribosomal RNA used for a loading control are indicated. Genomic RNA (G), TGBp, and CP subgenomic RNA are indicated. The pCymMV-R used as a negative control is illustrated in Fig. 1. The average percentage of relative real-time RT-PCR quantification (from 3 independent experiments) of CymMV RNA from CymMV clone-infected protoplasts at 24 h post-inoculation is indicated below the gels. The accumulation of pCymMV-M1 was set at 100% for relative quantification. Numbers at the left correspond to positions of marker RNAs (sizes in 1000 nucleotides) analyzed in the same gel.
pCymMV-M1-TGB1 Y/H containing a single substitution (Y/H) at amino acid position 44 of TGBp1 of pCymMV-M1 (Fig. 8C). The construction of other TGB site-directed mutagenesis clones (Fig. 8) was essentially the same as that for pCymMV-M1-TGB1 Y/H, except that different primer pairs and templates were used. The primer CymMV F3783/TGB1 and CymMV CPR were used in construction of all mutant clones. The remaining primers are in Supplementary Table S1. The mutated clones with names beginning with pCymMV-M1- or -M2- indicate that the initial template was pCymMV-M1 or pCymMV-M2, respectively.

Modifications of nucleotide sequence in an overlapping region between TGBp1 and TGBp2 (one nucleotide substitution) and between TGBp2 and TGBp3 (two nucleotides substitution) do not cause amino acid sequence changes in TGBp2. All mutant clones were sequenced in the modified region to ensure that the mutations were correct.

Preparation and transfection of N. benthamiana protoplasts

Capped transcripts corresponding to the wild-type virus and the constructed vectors of CymMV were synthesized by use of the mMESSAGE mMACHINE T3 high yield capped RNA transcription kit (Ambion, Inc., Austin, TX). In total, 5 μg of pCymMV-M1 and its derivative plasmids were digested with SpeI and in vitro transcription was conducted according to manufacturer’s instruction. Protoplasts prepared from Nicotiana benthamiana plants and RNA transfection were as previously described (Satyanarayana et al., 2002), except that 10 μg of each transcript and 2×10^5 cells were used for each inoculation, and after inoculation, protoplasts were incubated at 26 °C. The inoculated protoplasts were collected as previously described (Klaassen et al., 1996). Aliquots containing approximately 1×10^5 cells were collected by centrifugation (1300 × g) at different times post-inoculation, and RNA was isolated by use of TRIzol Reagent (Invitrogen, San Diego, CA) according to the manufacturer’s recommendations. The RNA was dissolved in 30 μl DEPC-treated water, and 5 μg RNA (equivalent to 5×10^4 cells) was used for northern hybridization analysis.

Plants and virus inoculation

An amount of 5 μg transcripts was dissolved in inoculation buffer (0.05 M NaH₂PO₄/Na₂HPO₄ pH 7.0) and rubbed to carborundum-dusted Phalaenopsis var. formosa (6 leaf stage) or N. benthamiana (5 leaf stage) by hands wearing latex gloves. The inoculated leaves were washed with excess distilled water, and the viruses were detected 2 weeks post-inoculation. All plants were kept in an insect-proof and...
thermal-controlled (25 °C–28 °C) greenhouse and with 12 h of light (5.22 μmol/s·m−2) for two weeks. P. amabilis var. formosa (10 cm in height) was purchased from the Taiwan Sugar Research Institute (Tainan, Taiwan). Each set of inoculations (Figs. 3, 4, 6 and 8) was repeated at least three times. RT-PCR was used to amplify the modified region (primer pairs used for RT-PCR were described in Supplementary Table 1) of each mutated CymMV from two randomly selected plants of each inoculation set followed by sequencing, all detected progeny viruses still maintained the original modification.

RT-PCR

RNA extracted from CymMV-infected plants was used as a template for synthesis of cDNAs by Moloney murine leukaemia virus (MMLV) reverse transcriptase following the manufacturer’s instructions (Promega, Inc., Madison, WI, USA). The PCR amplification conditions were as described (Rubio et al., 2000). The cDNAs were PCR amplified in a mixture containing 1.5 mM MgCl2, 1 mM of each of the 4 dNTPs, 2.5 U of Taq DNA polymerase (Promega, Inc., Madison, USA), and 50 ng of each oligonucleotide. The PCR cycles were 94 °C for 4 min for 1 cycle, then 94 °C for 30 s, 55 °C for 30 s and 72 °C for 1 min for 30 cycles, then an extension at 72 °C for 10 min.

Construction of CymMV CP expression clones

pCymMV-M1 and pCymMV-M2 were used as the initial template, and the primer pairs CymMV CP-BamHI F and CymMV CP-BamHI R (Supplementary Table S1) were used in the PCR reactions to amplify the two CymMV CP fragments. The fragments were gel purified by use of a gel extraction kit (Qiagen) and digested with BamHI. The fragment derived from pCymMV-M1 and pCymMV-M2 was incubated with the BamHI-digested expression vector pGEX-2T and ligated by use of T4 DNA ligase (Promega) to construct pGEX-2T-2-M1CP and pGEX-2T-2-M2CP. Both clones had been sequenced completely.

Expression and purification of CymMV CP

A total of 5 ml of overnight cultures of Escherichia coli XL1-blue transformed with pGEX-2T-2-M1CP, pGEX-2T-2-M2CP and pGEX-2T, were grown until they reached the OD600 = 0.5 at 37 °C. 200 ml LB medium containing ampicillin (50 μg/ml) was added, and bacteria were grown until they reached the OD600 = 1.0 at 37 °C. 200 ml LB medium containing ampicillin (50 μg/ml) was added, and bacteria were grown until they reached the OD600 = 0.5 at 37 °C. Then the fusion protein expression was induced with 1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) for 2 h at 37 °C. After centrifugation at 5000 × g for 5 min at 4 °C, the supernatant was removed and the pellet was re-suspended in incubation buffer of 1 M of sodium chloride and by use of a LightShift Chemiluminescent EMSA kit (Promega, Inc., Rockford, IL) followed the manufacturer’s protocol for chemical illumination. Fluorescence signals were captured by use of a Biospectrum AC Imaging system (UVP, LLC, Upland, CA, USA).

Whole-mount RNA analysis

Whole-mount RNA analysis was as described with modification (Zachgo et al., 2000). Sample fixing was extended to 2 h RT in a glass vial in PBS containing 0.1% Tween 20, 0.08 M EGTA, 10% DMSO and 5% paraformaldehyde. CymMV CP probe (see above) was used for hybridization.

Construction of CymMV TGBp and CP yeast two-hybrid clones

pCymMV-M1 was used as the initial template, and the primer pairs TGB1F/TGB1R, TGB2F/TGB2R, TGB3F/TGB3R, CPF/CRP (Supplementary Table S1) were used in PCR reactions to amplify the TGBp1, TGBp2, TGBp3 and CP fragments. The fragments were gel purified, incubated with the Smal digested pgDAT7 and pGBK7T (Clontech, Mountain View, CA), and ligated by use of T4 DNA ligase (Promega) to construct pTGB1-AD, pTGB2-AD, pTGB3-AD, pCP-AD, pTGB1-BD, pTGB2-BD, pTGB3-BD and pCP-BD. All clones had been sequenced completely.

Yeast two-hybrid system

Yeast two-hybrid assays were done by use of the Matchmaker Yeast Two-Hybrid System 3 (Clontech) following the manufacturer’s instructions. Yeast was transformed with bait vectors and prey vectors and selected on SD plates lacking Leu and Trp. After 2 days of growth at 30 °C, the yeast colonies were transferred to two different selection plates containing SD medium lacking Leu, Trp, and SD medium (containing X-α-GAL) lacking Leu, Trp and His.

Statistical analysis

Statistical analysis involved ANOVA or Dunnett’s T test with use of MINITAB14 (Minitab Inc., State College, PA, USA).

Appendix A. Supplementary data

Supplementary data associated with this article can be found in the online version, at doi:10.1016/j.jvirol.2009.02.049.

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