Dicer-Like 4 Is Involved in Restricting the Systemic Movement of Zucchini yellow mosaic virus in Nicotiana benthamiana

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Zucchini yellow mosaic virus (ZYMV) induces serious diseases in cucurbits. To create a tool to screen for resistance genes, we cloned a wild ZYMV isolate and inserted the visual marker Rosea1 to obtain recombinant clone ZYMV-Ros1. While in some plant-virus combinations Rosea1 induces accumulation of anthocyanins in infected tissues, ZYMV-Ros1 infection of cucurbits did not lead to detectable anthocyanin accumulation. However, the recombinant virus did induce dark red pigmentation in infected tissues of the model plant Nicotiana benthamiana. In this species, ZYMV-Ros1 multiplied efficiently in local inoculated tissue but only a few progeny particles established infection foci in upper leaves. We used this system to analyze the roles of Dicer-like (DCL) genes, core components of plant antiviral RNA silencing pathways, in ZYMV infection. ZYMV-Ros1 local replication was not significantly affected in single DCL knockdown lines nor in double DCL2/4 and triple DCL2/3/4 knockdown lines. ZYMV-Ros1 systemic accumulation was not affected in knockdown lines DCL1, DCL2, and DCL3. However in DCL4 and also in DCL2/4 and DCL2/3/4 knockdown lines, ZYMV-Ros1 systemic accumulation dramatically increased, which highlights the key role of DCL4 in restricting virus systemic movement. The effect of DCL4 on ZYMV systemic movement was confirmed with a wild-type version of the virus.

When plant viruses manage to enter an initial cell in their hosts, frequently with the help of a vector organism, they express and replicate their genomes to produce progeny that first move cell to cell to reach the host vascular tissue and then move long distance to distal areas. During this process, viruses hijack multiple elements, complexes, and structures from host plants. They must also surpass all barriers and neutralize plant defensive responses. Viral and host components establish a complex interaction network that frequently leads to infection and disease response but to resistance at other times. One of the main goals of plant virology is to understand this network in order to be able to modify the equilibrium to favor resistance.

Plants use RNA-mediated gene silencing pathways to fight invading viruses (Hamilton and Baulcombe 1999). Plant Dicer-like (DCL) RNases recognize virus-specific double-stranded or highly-structured RNAs to produce 21- to 24-nucleotide (nt) small interfering RNA (siRNA) duplexes (Aliyari and Ding 2009; Zhang et al. 2015). One of the strands is selectively loaded by an argonaute protein to guide the RNA-induced silencing complex to target and repress sequence complementary viral RNAs (Carbonell and Carrington 2015). The amount of viral double-stranded RNA that triggers the antiviral RNA silencing pathways in the host plant is amplified by RNA-dependent RNA (RDR) polymerase activities by using viral siRNAs as primers (Wang et al. 2010). In order to counteract this defensive response, evolution has shaped the proteins dedicated to suppress RNA silencing in virtually all plant viruses (Csorba et al. 2015).

Zucchini yellow mosaic virus (ZYMV) is a prominent pathogen of many plant species of the family Cucurbitaceae, which includes different important crops (Lecoq and Desbiez 2012). It was first isolated in Italy in 1973 (Lisa et al. 1981), although it is currently present worldwide (Desbiez and Lecoq 1997). ZYMV belongs to the genus Potyvirus in the family Potyviridae and its genome consists of approximately 10,000-nt-long single-stranded RNA molecules of plus polarity that encodes two versions of a large polypeptide (Wu et al. 2010). The genomic RNA of potyviruses (genus Potyvirus) is covalently attached, at its 5’ end, to a viral protein genome-linked (VPg), contains a polyadenylated tail at its 3’ end, and is encapsidated by approximately 2,000 units of the viral coat protein (CP) in an elongated and flexuous virion (Revers and García 2015). Potyviral proteins, which include the P1 protease, the helper component protease (HC-Pro), the P3 protein and P3N-PIPO, small hydrophobic polypeptide 6K1, the cylindrical inclusion (CI) protein, a second small hydrophobic polypeptide 6K2, the nuclear inclusion a (NiA) protein, which is further split into its two VPg and protease (NIAPro) domains, viral RDR polymerase or nuclear inclusion b (NiB) protein, and the CP (Fig. 1A), are produced from a regulated cascade of proteolytic processing through the activity of three viral proteases, namely, P1, HC-Pro, and NIAPro.

For the purpose of obtaining a viral clone to facilitate screening for resistance in cucurbitaceous plants, we cloned a wild isolate of ZYMV and constructed a recombinant clone that...

Nucleotide sequence data for the Vera isolate of Zucchini yellow mosaic virus has been deposited in the GenBank database under accession number KX499498.

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expresses the snapdragon (*Antirrhinum majus* L.) *Rosea1* marker gene, which activates anthocyanin (a class of flavonoid pigments) biosynthesis and allows the visual tracking of viral infection in different plant species (Bedoya et al. 2012). Anthocyanin accumulation is cell-autonomous and only occurs in those cells infected by the marked virus in which *Rosea1* is expressed. The amount of anthocyanins correlates with viral load (Bedoya et al. 2012). Unlike fluorescent protein markers, pigment accumulation in this system is clearly visible to the naked eye (Majer et al. 2013). We observed that this tool proved quite useless in zucchini (*Cucurbita pepo* L.) and other cucurbits, *ZYMV* natural hosts, given a negligible accumulation of colored anthocyanins in infected tissues. However, in experimental host *Nicotiana benthamiana* Domin, efficient virus-local multiplication in inoculated tissues and an inefficient systemic movement to distal tissues were observed for the recombinant clone. We reasoned that the *Rosea1*-marked *ZYMV* clone and *N. benthamiana* combination could represent an excellent experimental system to study the virus sequence determinants and the host factors involved in the long-distance movement of this virus. By means of this novel tool, we particularly aimed to analyze the differential contributions of the four *N. benthamiana* DCL genes, core components of the host RNA silencing pathways, in virus systemic movement in *N. benthamiana*. Our results show that, while

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**Fig. 1.** Symptoms induced by *Zucchini yellow mosaic virus* (*ZYMV*) and *ZYMV-Ros1* in zucchini plants. A, Schematic representation of the *ZYMV* genome. Lines represent the 5′ and 3′ untranslated regions (UTR) and boxes represent the different viral cistrons as indicated. In *ZYMV-Ros1*, a cDNA that codes for *Antirrhinum majus* *Rosea1* was inserted between the Nib (nuclear inclusion b) and CP (coat protein) cistrons. The cDNA included sequences that corresponded to extra amino and carboxy terminal peptides, as indicated, to mediate proteolytic release from the polyprotein. B, Representative zucchini plants agroinoculated with the empty binary plasmid, *ZYMV*, or *ZYMV-Ros1*. To better appreciate symptoms, a picture of selected leaves is also shown. All the pictures were taken at 21 days postinoculation (dpi). C, Western blot analysis of *ZYMV* CP accumulation in upper noninoculated tissues of three independent plants agroinoculated with the empty plasmid (lanes 1 to 3), wild-type *ZYMV* (lanes 4 to 6), and *ZYMV-Ros1* (lanes 7 to 9) at 15 dpi. The position and size (expressed in kiloDaltons) of marker proteins are indicated at the left of the panel. D, Bar graph representing the average *ZYMV* CP accumulation, quantified as luminescence arbitrary units (AU) by Western blot, in the upper noninoculated tissues of the previously described (C) zucchini plants. Error bars represent the standard error median.
individual DCL genes barely contributed to the inhibition of virus multiplication in inoculated tissues. DCL4 plays a major role in restricting ZYMV systemic movement in *N. benthamiana*.

**RESULTS**

A ZYMV infectious clone that includes the visual Rosea1 marker suboptimally moves long distance in *N. benthamiana*.

The Rosea1 marker system has been successfully applied to several combinations of viruses and host plants (Bedoya et al. 2012). However, it cannot be considered as a universal system, as it is based on the activity of a heterologous transcription factor on a host endogenous metabolic pathway. We wondered whether this system could be applied to ZYMV to track infection in cucurbitaceous plants. If so, the system would be a most valuable tool to facilitate high-throughput screening for resistance in cucurbit breeding programs. To this end, we cloned a wild ZYMV isolate from a zucchini plant (cv. Scallop), grown in 2013 in Horta de Vera (Valencia, Spain), which presented severe infection symptoms. Two internal cDNA fragments from the viral genome were amplified by reverse transcription-polymerase chain reaction (RT-PCR), whereas the 5' and 3' terminal cDNAs were amplified by a rapid amplification of cDNA ends (RACE) strategy. The sequence information from all these cDNAs was used to design a set of primers (Supplementary Table S1) to amplify the whole genome of the Vera isolate of ZYMV in three cDNA fragments flanked by the recognition site of a type-IIS restriction enzyme. Finally, these fragments were assembled into a binary plasmid in which the full-length ZYMV cDNA was flanked by Cauliflower mosaic virus (CaMV) 35S promoter and a 50-nt-long poly(A) stretch followed by the CaMV 35S terminator. This ZYMV cDNA was sequenced and the resulting full-length sequence was deposited in GenBank as the Vera isolate of ZYMV (GenBank accession number KX499498). A standard nt BLAST search displayed the highest identity with a Taiwanese isolate of ZYMV (GenBank accession number AF127929.2) (Lin et al. 2001). An alignment analysis using ClustalW exhibited 94.6% nt identity between both sequences with 517 nt differences.

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To confirm that recombinant clone ZYMV-Ros1 expressed a functional copy of Rosea1, we agroinoculated *N. benthamiana* plants using the same *Agrobacterium tumefaciens* cultures. ZYMV replicates with no symptoms in the inoculated leaves of *N. benthamiana* and, depending on the strain, moves systemically and induces latent infection (Desbiez and Lecoq 1997). We previously showed that this species produces intense reddish pigmentation when infected with several viruses that express Rosea1, including other potyviruses like Tobacco etch virus (TEV) (Bedoya et al. 2012). Unlike the tissues agroinoculated with the empty vector or with ZYMV, tissues agroinoculated with ZYMV-Ros1 displayed intense dark red pigmentation at 7 dpi (Fig. 2A; Supplementary Fig. S4). This result indicates that ZYMV-Ros1 expresses a functional Rosea1 transcription factor that efficiently induces accumulation of reddish anthocyanins in *N. benthamiana*. As expected, none of the *N. benthamiana* plants showed infection symptoms. However, a prolonged observation of these plants revealed that, in those agroinoculated with ZYMV-Ros1, some pigmented foci constantly appeared to be scattered on the upper noninoculated leaves (Fig. 2B). This result suggests that the Vera isolate of ZYMV is able to move a long distance in *N. benthamiana* but inefficiently. It is noteworthy that we detected infection foci on the upper noninoculated leaves, due to the vivid pigmentation induced by the Rosea1 transcription factor. We realized that the combination of the ZYMV-Ros1 clone and *N. benthamiana* could represent a convenient experimental system to study elements involved in ZYMV systemic movement.

**Analysis of the contribution of the four host DCL genes to ZYMV systemic movement in *N. benthamiana***.

With this experimental system at hand, we next aimed to study the effect of the four *N. benthamiana* DCL genes, which are core components of host RNA-mediated silencing pathways, on the systemic movement of ZYMV in this species. To this end,

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Fig. 2. ZYMV-Ros1 suboptimally moves long distance in *Nicotiana benthamiana*. **A**, Leaves agroinoculated with the empty plasmid, Zucchini yellow mosaic virus (ZYMV), or ZYMV-Ros1. Pictures were taken at 7 days postinoculation (dpi). **B**, Comparison of systemic leaves from three *N. benthamiana* plants agroinoculated with the empty binary plasmid or ZYMV-Ros1. Arrows indicate the pigmented infectious foci induced by ZYMV-Ros1 in systemic leaves. The picture was taken at 21 dpi.
we used a set of *N. benthamiana* transgenic plants in which the different *DCL* genes were down-regulated by expressing specific hairpin constructs (Dadami et al. 2013; Katsarou et al. 2016). RT-quantitative (q)PCR and Northern blot hybridization analyses of transgenic lines DCL1.13i, DCL2.11i, DCL3.10i, and DCL4.9i showed a specific reduction in the *DCL1*, *DCL2*, *DCL3*, and *DCL4* mRNAs levels, respectively. Apart from the plant knockdown in the single *DCL* genes, we also used line DCL2/4.5i, which expresses a hairpin to simultaneously downregulate *DCL2* and *DCL4*, and line DCL3.10 × 2/4.5i, the heterozygous progeny that results from crossing DCL3.10i as a female and DCL2/4.5i as a male (Dadami et al. 2013; Katsarou et al. 2016).

We first questioned whether these genes had an effect on virus accumulation in inoculated tissue. For this purpose, we agroinoculated two leaves of three *N. benthamiana* plants that corresponded to the wild type and the *DCL* knockdown lines DCL1.13i, DCL2.11i, DCL3.10i, DCL4.9i, DCL2/4.5i, and DCL3.10 × 2/4.5i. The agroinoculated tissues were harvested at 6 dpi and proteins were extracted. ZYMV CP was analyzed by electrophoretic separation, followed by Western blot using a specific anti-CP antibody (Fig. 3). Wild-type noninoculated controls were added to the analysis. The quantification of the Western blot signals is summarized in Supplementary Table S2. Figure 3 shows the three Western blots, as well as the graph of ZYMV CP accumulation in the agroinoculated tissue of the different *N. benthamiana* lines. According to the amount of CP, ZYMV-Ros1 accumulation in the inoculated tissues of the DCL1.13i, DCL2.11i, DCL4.9i, and DCL2/4.5i lines was similar to that of the wild-type plants. ZYMV-Ros1 accumulation was lower in the DCL3.10i line (0.6-fold on average) and higher in the DCL3.10 × 2/4.5i line (1.75-fold on average) compared with the wild-type plants (Fig. 3B). However, none of the differences in ZYMV-Ros1 accumulation between wild type and each of the *DCL* knockdown lines was statistically significant (P < 0.05 for all pair-wise Student’s t test comparisons).

Next, we analyzed the effect of the *DCL* downregulation on ZYMV systemic movement in *N. benthamiana*. Sets of three plants that corresponded to the wild type and the different knockdown lines were agroinoculated with ZYMV-Ros1 in three leaves. In the *N. benthamiana* lines DCL1.13i, DCL2.11i, and DCL3.10i, we obtained the same outcome previously obtained in the wild-type plants. Very few infection foci were detected in the upper noninoculated tissues. However in *N. benthamiana* line DCL4.9i and in lines DCL2/4.5i and DCL3.10 × 2/4.5i, ZYMV-Ros1 was able to efficiently move a long distance into the upper noninoculated tissue (Fig. 4A and B). Pictures of the three independent inoculated plants as well as the selected leaves that corresponded to each line are shown in Supplementary Figure S5. We previously showed that anthocyanin accumulation very precisely correlates with viral load in *Rosea1*-marked viruses (Bedoya et al. 2012). Therefore, in order to make a more quantitative estimate of ZYMV-Ros1 systemic movement in all these plants, we harvested all the aerial tissues above the agroinoculated leaves at 27 dpi and quantified the accumulation of reddish anthocyanins by a spectrophotometric analysis (Fig. 4C; Supplementary Table S3)

While anthocyanin accumulation in the DCL1.13i, DCL2.11i, and DCL3.10i lines was negligible and indistinguishable from the wild-type plants, the aerial tissues of lines DCL4.9i, DCL2/4.5i, and DCL3.10 × 2/4.5i accumulated substantial amounts of these pigments (Fig. 4C). It was noteworthy that the anthocyanin accumulations in the double DCL2/4.5i and in the triple DCL3.10 × 2/4.5i knockdown lines were 1.8- and 2.9-fold higher, respectively, on average, than in the single knockdown DCL4.9i line. Taken together, these results support a crucial role of *DCL4* in restricting the systemic movement of ZYMV in *N. benthamiana* that may be functionally complemented by *DCL2* and *DCL3*. To confirm this result, a similar experiment was conducted with wild-type ZYMV under same experimental conditions. Western blot and RT-qPCR analyses of virus accumulation at 28 dpi in the whole upper noninoculated tissues confirmed the crucial role of *DCL4* in restricting virus systemic movement (Fig. 4D and E).

**DISCUSSION**

The goal of this research was to create a tool to facilitate screening ZYMV resistance in the breeding programs of cucurbit plants. Although the initial aim failed, we were able to generate a convenient experimental system to analyze the contribution of ZYMV genetic determinants and host factors to viral systemic movement, which should ultimately help to understand and develop resistance to infection by this virus.

A new Mediterranean isolate of ZYMV that mostly resembles a sequence variant from Taiwan.

We constructed an infectious clone from a Spanish isolate of ZYMV that infected a zucchini plant. Our clone mostly resembles sequence variant AF127929.2, which has been reported in Taiwan and was isolated in 1993 from sponge gourd (*Luffa cylindrica* Roem.) (Lin et al. 2001). The two variants differ in 517 nt positions (94.6% identity), including the insertion of a U at position 9,465, which corresponds to the 3′ untranslated region. Our finding of a 2013 Mediterranean isolate that mostly resembled a 1993 Taiwanese sequence variant, which belongs to phylogenetic group A-IV, mainly composed of East Asian isolates (Couotts et al. 2011), supports the easy worldwide dispersion of this virus.

The ZYMV-mediated expression of *Rosea1* does not produce visible anthocyanin accumulation in cucurbit plants.

We constructed a recombinant ZYMV clone that expresses *A. majus* R2R3 MYB transcription factor Rosea1 (ZYMV-Ros1) (Fig. 1A). This recombinant clone induced the accumulation of reddish anthocyanins in the infected tissues of *N. benthamiana* (Fig. 2) but not in cucurbitaceous species like zucchini (Fig. 1B) or melon. Tomato plants engineered to over-express the two *A. majus* transcription factors Rosea1 and Delila under the control of a fruit-specific promoter produced purple tomatoes with high anthocyanin content (Butelli et al. 2008; Su et al. 2016). In plants, anthocyanin biosynthesis is controlled at the transcriptional level by members of three protein families: R2R3 MYB transcription factors, bHLH transcription factors, and WD repeat proteins. They interact to form a ternary complex that activates a series of genes that lead to anthocyanin biosynthesis and accumulation in vacuoles (Zhang et al. 2014). We previously showed that the virus-mediated expression of Rosea1 and Delila in tobacco tissues also induces the accumulation of large amounts of anthocyanins in infected tissues (Bedoya et al. 2010). Next, we reported that the sole virus-mediated expression of Rosea1 suffices to induce pigment accumulation that is readily detectable to the naked eye in infected tissues in several host plant-virus combinations. This finding suggests that this transcription factor is a convenient marker to visually track plant virus infection and movement (Bedoya et al. 2012). In terms of the size, Rosea1 is only slightly larger than the most conventional reporter gene used in plant virology, green fluorescent protein (GFP) (Tilsner and Oparka 2010). Although the impact of Rosea1 in recombinant virus fitness is stronger than that of GFP, the stabilities of both markers in the viral genome are similar (Majer et al. 2013). We succeeded in producing a visible reddish pigmentation of the infected tissues in solanaceous plants (*N. benthamiana* or *N. tabacum* L.) and also in the nonsolanaceous *Arabidopsis thaliana* L., using recombinant potyviruses...
such as TEV or *Turnip mosaic virus* (*TuMV*) and also with viruses like *Tobacco mosaic virus* or *Potato virus X* that belong to different families (Bedoya et al. 2012). *Narcissus mosaic virus* (genus *Potexvirus*) has also been shown to induce visible pigment production in *N. benthamiana* plants when expressing *Arabidopsis thaliana* R2R3 MYB transcription factor AtMYB75 (PAP1) (Zhang et al. 2013). Lack of pigment accumulation in the tissues of cucurbitaceous plants infected with ZYMV-Ros1 may result from an incompatibility between *A. majus* Rosea1 and the endogenous companion transcription factors of the bHLH and WD repeat types. Not much is known about the flavonoid pathway in cucurbits. Other flavonoids, such as flavone derivatives, have been detected in cucumber (*Cucumis sativus* L.) and melon leaves (Krauze-Baranowska and Cisowski 2001). Flavonol derivatives have also been reported in the reproductive organs of some cucurbits (Imperato 1980). Naringenin chalcone is the main flavonoid that accumulates in the fruit rind of some yellow melon varieties (Feder et al. 2015; Tadmor et al. 2010).

Zucchini plants infected with ZYMV-Ros1 showed milder symptoms than those infected with wild-type ZYMV (Fig. 1B). Some leaves in these plants presented a distinctive beautiful pattern that consisted in dark green perinerval stripes on a light-green background (Fig. 1B). The possibility that these distinctive symptoms might still arise from some unknown activity of the Rosea1 transcription factor cannot be ruled out.

**ZYMV-Ros1 inefficiently moves long distance in *N. benthamiana***.

According to the anthocyanin production induced by the Rosea1 marker, we observed that ZYMV-Ros1 efficiently accumulated in the agroinoculated tissues of *N. benthamiana* plants, but very few particles from the progeny were able to establish systemic infection foci (Fig. 2). As an alternative to agroinoculation, we obtained the same result from mechanical inoculation of *N. benthamiana* plants with an extract of ZYMV-Ros1–infected zucchini. It is worth noting that the visual marker was crucial for this observation, since very few systemic infection foci kept appearing in the first weeks after inoculation. The Rosea1-induced pigmented foci, which were directly observable without using specialized instrumentation such as a UV lamp (Bedoya et al. 2012), easily attracted our attention. It has been previously described that some ZYMV strains induce latent infection in *N. benthamiana* either systemically or in a limited manner to inoculated tissue (Desbiez and Lecoq 1997; Lesemann et al. 1983; Wang et al. 1992).

We reasoned that the combination of our recombinant ZYMV-Ros1 virus and the *N. benthamiana* host could represent an excellent experimental system to study the virus genetic determinants and host factors involved in ZYMV systemic movement. *N. benthamiana*, particularly the lineage used in most research laboratories, is susceptible to a large number of plant virus species from very different taxonomic groups. This is most probably because this lineage, which was originally harvested in an extreme habitat of central Australia, is a natural *rdr1* mutant (Bally et al. 2015; Carbonell 2015). Consequently, this species is frequently adopted as a model plant in many research works into plant viruses. In our system, the amount of viral particles capable of reaching upper noninoculated tissues was easily determined by monitoring the dark red pigmentation of these tissues. The efficiency of viral systemic movement was also quantified in systemic tissues by counting infection foci or by measuring anthocyanin accumulation by a simple colorimetric analysis of methanol extracts. We previously showed that, for Rosea1–marked viruses, anthocyanin accumulation correlates with viral load in infected tissues (Bedoya et al. 2012). This experimental system should help to analyze genetic determinants in the virus genome that affect systemic movement. More importantly, this system should allow analyzing the host factors involved in ZYMV systemic movement. In this way, by inoculating knockout or knockdown *N. benthamiana* mutants, host factors involved in favoring or restricting ZYMV systemic movement could be identified and analyzed. Similarly, by inoculating *N. benthamiana* plants in which the candidate factors from cucurbit species are expressed by stable genetic transformation or by transient expression through *Agrobacterium tumefaciens* or viral vectors,
the host factors that are recruited by the virus to mediate its systemic movement in the natural hosts could be screened. The identification and analysis of all these factors will provide an understanding of the mechanisms that underlie ZYMV systemic movement. These factors may also be used as targets to breed or engineer resistance in cucurbitaceous plants by blocking virus systemic movement.

*DCL4 is involved in restricting ZYMV systemic movement in N. benthamiana.*

Since DCL proteins initiate the antiviral RNA silencing response in plants (Aliyari and Ding 2009; Zhang et al. 2015), we used our new experimental system, based on ZYMV-Ros1 and *N. benthamiana*, to analyze the effects of *DCL* genes on ZYMV systemic accumulation. *N. benthamiana*, like *Arabidopsis thaliana*,...
leaves (Mermigka et al. 2016). In any case, viral systemic movement in plants is a rather complex and prolonged process and the specific mechanisms by which antiviral silencing blocks viral systemic spread need to be further clarified.

**MATERIALS AND METHODS**

**Amplification of ZYMV cDNAs.**

Total RNA was purified by silica gel chromatography (Zymo Research) from a piece of symptomatic leaf from a zucchini plant (cv. Scallop) growing in 2013 in Horta de Vera (Valencia, Spain) that showed typical symptoms of viral infection. From this RNA preparation, cDNAs were initially synthesized using RevertAid RT (Thermo Fisher Scientific) and oligodeoxynucleotide primers P1 and P4, designed on the basis of ZYMV GenBank reference sequence variant NC_003224.1. The two cDNAs were amplified with Phusion high-fidelity DNA polymerase (Thermo Fisher Scientific) and primers P2 and P3 and P5 and P6. New cDNAs corresponding to the 5' and 3' viral ends were amplified by RACE. To amplify the 3' end, we took advantage of the native polyadenylate tail of ZYMV genomic RNA. Using primer P7, we synthesized a cDNA that was next amplified by two subsequent PCRs using primers P8 and P9 and P10 and P11. To amplify the 5' end, we first synthesized a cDNA using primer P12. A polytidine tail was next added to the 3' end of this cDNA, using calf thymus terminal transferase (Thermo Scientific). Finally, the 5' end was amplified in two consecutive PCRs using primers P13 and P14 and P15 and P16. All these cDNAs were inserted into EcoRV-digested pBluescript II KS(+) (GeneBank accession number X52327.1) and were sequenced. Experimental sequences served to design primers (P17 to P29) to amplify the whole ZYMV genome in three fragments (5', central, and 3') by RT-PCR. We applied a nested PCR strategy in which 1 µl of the first reaction was used as a template for the second reaction. These three ZYMV cDNAs were ligated to EcoRV-digested pBluescript II KS(+), using T4 DNA ligase (Thermo Fisher Scientific), and *Escherichia coli* DH5α electroporated with the products of ligation.

**Construction of ZYMV infectious clones.**

The cloned cDNAs corresponding to the 5', central, and 3' fragments of the Vera isolate of ZYMV were recovered by digestion with the type-I13 restriction enzyme Eco311I (Thermo Fisher Scientific) from the pBluescript II KS(+) derivatives (described above) and were assembled (Engler et al. 2009) into the binary vector pG35Z, also digested with Eco311I. pG35Z is a binary vector derived from pCLEAN-G181 (GenBank accession number EU186083) (Thole et al. 2007) that we constructed as a previous step to assemble the ZYMV full-length clone. The map and sequence of pG35Z is in Supplementary Fig. S6. The resulting plasmid harboring the full cDNA of the Vera isolate of ZYMV (GenBank accession number KX499498) was named pGZYMV. Using PCR with the Phusion high-fidelity DNA polymerase, digestion with Eco311I, and ligation with T4 DNA ligase, a cDNA corresponding to the coding region of *A. majus Roseal* transcription factor (GenBank accession number DQ275529.1) was inserted between the Nlb and CP cistrons (positions 8,541 and 8,542 of KX499498). This cDNA was flanked with sequences coding for amino acids to complement the native Nlb/CP proteolytic site that was split in two. The resulting plasmid was named pGZYMV-Ros1.

**Plant agroinoculation.**

*Agrobacterium tumefaciens* C58C1 harboring the helper plasmid pCLEAN-S48 (Thole et al. 2007) was electroporated with pG35Z (empty plasmid), pGZYMV, or pGZYMV-Ros1. Liquid cultures of transformed *Agrobacterium tumefaciens* were grown to optical density at 600 nm (OD600) of approximately
1.0. Cells were recovered by centrifugation and were resuspended at an OD_{600} of 0.5 in 10 mM MES-NaOH, pH 5.6, 10 mM MgCl{2}, and 150 µM acetylsyringone. Cultures were induced for 3 h at 28°C and were used to agroinoculate zucchini (cv. MU-CU-16), melon (cv. Piel de Sapo), or N. benthamiana plants (Bedoya and Darós 2010). The inbreeding line MU-CU-16 belongs to the zucchini morphotype of the subspecies pepo of Cucurbita pepo and was provided by the Cucurbits Breeding Group of the Institute for the Conservation and Breeding of Agricultural Biodiversity, Universitat Politècnica de València (Blanca et al. 2011; Esteras et al. 2012). We agroinoculated 3-day-old zucchini, approximately one-month-old melon plants, and 4.5- or 5.5-week-old N. benthamiana plants.

**Western blot analysis.**

Infiltrated tissues of *N. benthamiana* plants (5.5 weeks old) were harvested 6 dpi (between 0.45 and 0.89 g, depending on the sample) and were ground with a mortar and pestle in the presence of liquid N{2}. Three volumes of buffer TET (60 mM Tris-HCl, pH 6.8, 2% sodium dodecyl sulfate [SDS], 100 mM dithiothreitol, 10% wt/vol glycerol, and 0.01% bromophenol blue) were added and the extracts were incubated at 95°C for 5 min. Extracts were clarified by centrifugation for 15 min and 40 µl of the supernatants (equivalent to 13 mg of fresh tissue), separated by discontinuous polyacrylamide gel electrophoresis (PAGE), in 12.5% polyacrylamide gels (5% polyacrylamide for the stacking gel) containing 0.05% SDS. Proteins were electroblotted to polyvinylidene fluoride membranes (GE Healthcare), which were blocked for 1 h in 5% nonfat milk in buffer TBS-T (0.05% SDS). Membranes were incubated overnight at 4°C with an antibody against ZYMV CP in upper noninoculated leaves of *N. benthamiana*. Membranes were washed three times with WB and alkaline phosphatase detected with CSPD (Roche Life Science). Luminescence was recorded and quantified with a LAS-3000 image analyzer (Fujiﬁlm). This protocol was also used to analyze the accumulation of ZYMV RNA in upper noninoculated leaves of *N. benthamiana*.

**RT-qPCR analysis of ZYMV RNA.**

RNA preparations were puriﬁed from *N. benthamiana* tissue samples, using the RNeasy plant mini kit (Qiagen) and were quantiﬁed using a NanoDrop ND-1000 spectrophotometer. cDNAs were synthesized in 20-µl reactions, including 100 ng of total RNA, 50 U RevertAid RT, and 5 pmol of primer P30. Two microliters of the products of these reactions were subjected to 20-µl qPCR amplification reactions by triplicate, using the Maxima SYBR Green/ROX qPCR master mix (Thermo Scientiﬁc) and 6 pmol primers P31 and P32 in a StepOnePlus real-time PCR system (Applied Biosystems). The amount of ZYMV RNA molecules present in 100 ng of RNA preparation was calculated from a calibration line obtained in the same condition with a RNA standard corresponding to the ZYMV 3’ genome fragment (from position 8,542 to 9,592 of KX499498), obtained by in vitro run off transcription and quantiﬁed by spectrophotometric analysis. StepOne Software v.2.2.2 (Applied Biosystems) was used to analyze the data.

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**LITERATURE CITED**


