Cucumber RDR1s and cucumber mosaic virus suppressor protein 2b association directs host defence in cucumber plants

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Abstract
RNA-dependent RNA polymerases (RDRs) regulate important aspects of plant development and resistance to pathogens. The role of RDRs in virus resistance has been demonstrated using siRNA signal amplification and through the methylation of viral genomes. Cucumber (\textit{Cucumis sativus}) has four RDR1 genes that are differentially induced during virus infection: \textit{CsRDR1a}, \textit{CsRDR1b}, and duplicated \textit{CsRDR1c1/c2}. The mode of action of \textit{CsRDR1s} during viral infection is unknown. Transient expression of the cucumber mosaic virus (CMV)-2b protein (the viral suppressor of RNA silencing) in cucumber protoplasts induced the expression of \textit{CsRDR1c}, but not of \textit{CsRDR1a/b}.

Results from the yeast two-hybrid system showed that \textit{CsRDR1} proteins interacted with CMV-2b and this was confirmed by bimolecular fluorescence complementation assays. In protoplasts, \textit{CsRDR1s} localized in the cytoplasm as punctate spots. Colocalization experiments revealed that \textit{CsRDR1s} and CMV-2b were uniformly dispersed throughout the cytoplasm, suggesting that \textit{CsRDR1s} are redistributed as a result of interactions. Transient overexpression of individual \textit{CsRDR1a/b} genes in protoplasts reduced CMV accumulation, indicating their antiviral role. However, overexpression of \textit{CsRDR1c} in protoplasts resulted in relatively higher accumulation of CMV and CMV\textDelta 2b. In single cells, \textit{CsRDR1c} enhances viral replication, leading to CMV accumulation and blocking secondary siRNA amplification of \textit{CsRDR1c} by CMV-2b protein. This suggests that CMV-2b acts as both a transcription factor that induces \textit{CsRDR1c} (controlling virus accumulation) and a suppressor of \textit{CsRDR1c} activity.

KEYWORDS
cucumber mosaic virus suppressor 2b, cucumber RDR1, host defence, protein–protein interaction, protoplast

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1 | INTRODUCTION

Viruses pose serious threats to economically important crops by reducing yield and quality. Plants employ a variety of defence strategies to counter viral infection processes. RNA silencing, also known as RNA interference (RNAi), is a host defence mechanism that provides antiviral resistance/immunity by targeting the viral genome. RNAi degrades the viral genome into small RNAs (20–24 nucleotides [nt]), known as small interfering RNAs (siRNA), using Dicer-like (DCLs) endonucleases in a sequence-complementarity manner (Baulcombe, 2004; Ding & Voinnet, 2007; Pumpkin & Voinnet, 2013; Wang et al., 2010).

To achieve potent silencing, host RNA-dependent RNA polymerases (RDRs) are crucial for manifestations of the RNAi process as they amplify viral double-stranded (ds) RNA and transmit silencing signals (Ahlquist, 2002; Qu, 2010; Vance & Vaucheret, 2001). RDRs are characterized by the presence of a catalytic DLDGD domain and can synthesize dsRNA from single-stranded (ss) RNA templates by primer-independent and primer-dependent mechanisms (Willmann et al., 2011). The resulting dsRNA is then reincorporated into a cyclical RNA degradation pathway, in which RNA silencing leads to the amplification of silencing signals for the degradation of the target RNA (Lipardi et al., 2001; Sijen et al., 2001; Wang et al., 2010). These RDRs are classified into three major clades: RDRa, RDRb, and RDRγ; plants contain only RDRa and RDRγ (Zong et al., 2009). The Arabidopsis genome encodes six different RDRs, of which RDR1, RDR2, and RDR6 play important roles in the biogenesis of small RNA (Willmann et al., 2011). RDR1 and RDR6 are known to play important roles in antiviral immunity by producing viral dsRNA (Guo et al., 2019) and preventing meristem invasion (Schwach et al., 2005). Recent studies have identified two host factors, antiviral RNAi-defective 1 (AVI1) and AVI2, that enhance the activity of RDR1 and RDR6 for the generation of viral siRNAs (Guo et al., 2017, 2018). Moreover, RDR1/RDR6 loss-of-function mutants show increased susceptibility to many RNA viruses (Guo et al., 2018; Qi et al., 2009; Wang et al., 2010). After infection by cucumber mosaic virus (CMV) or turnip mosaic virus, RDR1 was found to be responsible for the biogenesis of a distinct class of endogenous virus-activated siRNAs that confer broad-spectrum antiviral responses by silencing host genes (Cao et al., 2014). RDR2 plays an important role in the biogenesis of host-derived 24-nt repeat-associated siRNAs (rasiRNAs) (García-Ruiz et al., 2010).

CMV is a widespread pathogenic member of the Bromoviridae family with a broad host range that includes many economically important crops (Palukaitis & García-Arenal, 2003). Its genome consists of three positive-sense, ssRNA segments (RNAs 1 to 3), that together encode five proteins. The 1a and 2a proteins, which are encoded by RNA-1 and RNA-2, form part of the replication complex that is responsible for the replication of the viral genome. The 2b protein is encoded by a subgenomic RNA (RNA-4A) derived from RNA-2 (Ding et al., 1994) and is a multifunctional protein that acts as a viral suppressor of RNA silencing (VSR; Díaz-Pendón et al., 2007) and aids the long-distance movement of the virus and symptom induction (Ding et al., 1995). RNA-3 encodes two proteins, movement protein (MP) that translates from the 5′ open reading frame and is responsible for cell-to-cell virus movement, and the 3′ open reading frame that translates the viral capsid or coat protein (CP) through subgenomic RNA-4 (Palukaitis & García-Arenal, 2003).

To evade the host defence machinery, viruses encode VSR proteins, which can interfere with the RNA silencing pathway at several points, to suppress this process (Csorba et al., 2015). The CMV-2b protein (referred to hereafter as “2b”) is a well-known VSR that obstructs posttranscriptional gene silencing and transcriptional gene silencing (Guo & Ding, 2002; Hamera et al., 2012). The 2b protein plays a role in binding both 21-24 nt siRNA duplexes and long (55 nt) dsRNAs (Duan et al., 2012). Argonaute (AGO) proteins are important for the gene-silencing process. The 2b protein has been shown to interact with the AGO PAZ and PIWI domains, interfering with AGO functions (Duan et al., 2012; Hamera et al., 2012; Zhang et al., 2006). An earlier report by Hamera et al. (2012) showed that, during virus infection, 2b interferes with genome methylation or transcriptional gene silencing and changes the expression profile of many Arabidopsis genes. In addition, Duan et al. (2012) demonstrated the remodelling of the Arabidopsis thaliana genome methylation pattern due to the binding of the 2b protein with 21-, 22-, or 24 nt siRNA. Transgenic A. thaliana plants expressing the 2b protein show a genome-wide reduction of CHH and CHG methylation in their transposable elements and coding genes (Hamera et al., 2012; Zhao et al., 2016). However, this suppression of DNA methylation is independent of the AGO-binding activity of the 2b protein (Duan et al., 2012). The 2b protein also plays an important role in symptom development and the systemic movement of CMV (Lewsey et al., 2010; Wang et al., 2004).

Interaction of the 2b protein with Arabidopsis catalase 3 decreases catalase activity (specifically the scavenging of cellular hydrogen peroxide), causing necrotic symptom development during CMV infection and thus favouring virus infection (Inaba et al., 2011). A CMVΔ2b mutant lacking the 2b gene was reported to accumulate at high levels along with severe symptoms in Arabidopsis RDR1 or RDR6 loss-of-function mutants (rdr1, rdr1/rdr6, and rdr1/rdr2/rdr6), indicating the importance of the RDR1/RDR6 genes for immunity to CMV (Wang et al., 2010). Recently, it was found that autophagy reduces CMV accumulation via 2b degradation and contributes to resistance that is mediated by AGO1-dependent and DCL1/4-dependent RNA-silencing mechanisms (Shukla et al., 2020).

Most plant families encode single or duplicated RDR1 genes (Chen et al., 2010; Qin et al., 2017; Willmann et al., 2011); exceptionally, the cultivated Cucurbitaceae encode a number of putative RDR1 genes (Leibman et al., 2018). The cucumber (Cucumis sativus) genome encodes four RDR1 genes, namely CsRDR1α, CsRDR1β, and duplicated CsRDR1c1/c2, that have been shown to be differentially induced during infection by various viruses (Leibman et al., 2018).

These studies demonstrate the role of the 2b protein in pathogenicity, based on its suppression of a variety of RNA-silencing activities. In this study, the role of RDR1s in defence activities against CMV infection was evaluated in cucumber and we explored the...
exact roles of individual RDR1 genes during virus infection. In the present study, the hypothesis of association between cucumber CsRDR1s and the 2b protein, and the impact of that interaction on the progress of CMV infection, were evaluated using various assays. This study demonstrates the interaction and localization of the 2b with CsRDR1s, despite suggesting regulatory differential regulation of CsRDR1s. Higher levels of expression of CsRDR1s can effectively suppress virus infection and associated symptoms to aid the development of tolerance/resistance.

2 | RESULTS

2.1 | CMVΔ2b failed to induce CsRDR1 expression

Cucurbits possess a unique RDR1 family comprising four genes (RDR1a, RDR1b, RDR1c1, and RDR1c2), which are differentially expressed during virus infection (Leibman et al., 2018). Here, we investigated the role of cucumber RDR1s (CsRDR1s) during infection by CMV and a 2b-deficient CMV mutant (CMVΔ2b) in cucumber cv. Bet Alfa, a very virus-susceptible cultivar (Wang et al., 2003). To produce the CMVΔ2b clone, the RNA-2 biologically active cDNA clone was modified. Specifically, two nucleotide changes were made in each of the first three ATG codons (ATG→ACC) of the 2b gene, as a CMVΔ2b clone with only one nucleotide change (ATG→ACG) reverted to wild type in cucumber. Thus, 2b gene expression was disrupted without affecting 2a protein translation. During CMV infection, expression of CsRDR1a remained higher (c.4-fold at 7 days postinoculation [dpi]) in infected cv. Bet Alfa plants than in healthy plants; that difference decreased to c.2-fold at 14 dpi (Figure 1a). CsRDR1b expression levels were increased up to 2-fold at 7 dpi and 3-fold at 14 dpi during CMV infection, relative to healthy control plants (Figure 1a). In contrast, CsRDR1a and CsRDR1b expression levels remained unchanged (or slightly decreased) in CMVΔ2b-infected cv. Bet Alfa plants (Figure 1a). Duplicate versions of CsRDR1a (CsRDR1c1 and CsRDR1c2) with 98% identity were observed in infected plants but were not expressed in healthy plants. Previously, CsRDR1c1 and CsRDR1c2 were found to be similarly expressed during virus infection (Leibman et al., 2018), so expression of CsRDR1c1 alone was examined and hereafter CsRDR1c1+2 is referred to as CsRDR1c. In contrast to the results obtained for CsRDR1a/1b, very high levels of CsRDR1c expression were observed on CMV infection, with an 8,000-fold increase at 7 dpi and a 3,000-fold increase at 14 dpi in cv. Bet Alfa plants (Figure 1a).

During CMVΔ2b infection, CsRDR1a and CsRDR1b expressions were not induced. The level of CsRDR1a expression also remained unchanged at 7 dpi, but it showed a 10-fold increase at 14 dpi relative to healthy cv. Bet Alfa plants. These results demonstrate that CsRDR1a, CsRDR1b, and CsRDR1c expression change is associated with either the CMV level or presence of the 2b protein, as no or less induction was observed during CMVΔ2b infection as compared to CMV (at 7 and 14 dpi; Figure 1a). Similar results were obtained in western blotting, where higher CMV coat protein (CP) accumulation was observed at 7 and 14 dpi in CMV-infected cv. Bet Alfa as compared to CMVΔ2b mutant-infected samples (Figure 1b). A similar trend was observed in a tolerant cucumber cv. Shimshon during CMV and CMVΔ2b infection (Figure 1b). These results clearly suggest that the expression of the CMV-2b gene is important for the induced expression of CsRDR1a, CsRDR1b, and CsRDR1c during infection.

2.2 | CMV-2b enhances CsRDR1c expression in single cells

The role of the 2b protein in the induction of CsRDR1s was assessed by transient expression of the 2b protein in cucumber protoplasts. Cucumber protoplasts isolated from cv. Bet Alfa plants were transfected with the expression vectors pSAT6-CMV2b-GFP (CP), pSAT6-CMV2b-GFP (2b), and the pSAT6EV-GFP empty vector control (EV; Figure 2).

After 72 hr of incubation, the expression levels of endogenous CsRDR1a, CsRDR1b, and CsRDR1c were determined using reverse transcription (RT) PCR. It was observed that 2b protein itself could induce CsRDR1c, whereas CP overexpression had no such effect (Figure 2). However, the CsRDR1b and CsRDR1a genes were expressed in cucumber leaf protoplasts and the levels of their expression were not affected by the expression of 2b or CP, as compared to polyethylene glycol (PEG)-treated controls (Cont) and pSAT-EV (empty vector, EV) (Figure 2). These results indicate that the 2b protein that was localized to the nucleus acted directly or indirectly as an inducer (“transcription factor”) for CsRDR1c regulation.

2.3 | Effect of CsRDR1 overexpression against CMV at the single-cell level

To investigate the activity of CsRDR1s against CMV, CsRDR1a/1b/1c were expressed transiently in cucumber protoplasts, along with CMV in vitro transcripts. Cucumber protoplasts transfected with CMV, CMV+pSAT6-EV-GFP, and nontransfected protoplasts were used as controls. Relative CMV accumulation was determined 68 hr posttransfection by quantitative RT-PCR (RT-qPCR; Figure 3). Transient expression of CsRDR1a effectively suppressed the accumulation of CMV and CMVΔ2b compared to control protoplasts transfected with either CMV alone or CMV+pSAT6-EV-GFP (Figure 3a). Similarly, transiently expressed CsRDR1b effectively suppressed the accumulation of CMV about 6-fold. However, a smaller increase in CMVΔ2b accumulation (c.2-fold) was observed (Figure 3b). In contrast to CsRDR1a and CsRDR1b, transient expression of CsRDR1c was associated with increased accumulation of CMV (close to 14-fold), whereas a c.7-fold increase was observed for the CMVΔ2b mutant (Figure 3c). The ectopic expression of the CsRDR1s was confirmed by RT-qPCR with reference to the CMV- and CMVΔ2b-transfected protoplasts (Figure 3d-f). Endogenous CsRDR1s expression remained lower during this period in cucumber protoplasts, which suggests that ectopically expressed CsRDR1a and CsRDR1b triggered an antiviral response against CMV by 68 hr posttransfection. Interestingly, transient expression of CsRDR1c during CMV infection was 54 and 74
times greater than the expression of CsRDR1a and CsRDR1b, respectively, and 39 and 97 times greater during CMVΔ2b infection than for CsRDR1a and CsRDR1b, respectively (Figure 3d–f). This highly increased level of CsRDR1c expression in protoplasts is probably due to autopositive regulation of the endogenous CsRDR1c expression in the presence of CMV, as was observed in cucumber leaves (Figure 1). The unexpected increase in the levels of CMV and CMVΔ2b due to transient expression of CsRDR1c (Figure 3c), in contrast to transient expression of CsRDR1a or CsRDR1b, suggests that CsRDR1c abundance alone in CMV-infected cells plays a role in enhancing viral replication at the cell level for a short time posttransfection.

2.4 | 2b protein, a viral suppressor of RNA silencing, directly interacts with CsRDR1 proteins in yeast

Because the 2b gene was found to induce CsRDR1c expression (Figure 2), the possibility of a direct interaction of the 2b protein with CsRDR1 proteins was evaluated using the yeast two-hybrid (Y2H) GAL4-based Gold system. The 2b protein was used as bait to screen against three cucumber prey proteins: CsRDR1a, CsRDR1b, and CsRDR1c. In general, RDRs are characterized by an RNA recognition motif (RRM) at the N-terminus and the RNA-dependent RNA polymerase catalytic domain (RdRp) in the centre of the protein (Figure 4). Therefore, RRM and RdRp regions were selected from CsRDR1a and CsRDR1b to generate prey constructs. The RRM region is not well defined in CsRDR1c but a region (N′) from the CsRDR1c N-terminus was selected (Figure 4a). The CMV 2b interacted strongly with CsRDR1a, CsRDR1b, and CsRDR1c (up to 10⁻³/10⁻⁴ OD dilutions in quadruple-dropout selection medium), whereas negative controls (empty bait vector, pGBKT7, and pGBKT7-Lam) showed no growth on triple/quadruple-dropout selection plates with any CsRDR1 proteins (Figure 4b). The binding between 2b and the RRm of CsRDR1a/b and N′ of CsRDR1c was stronger than with the RdRp motif or the intact CsRDR1 proteins. Therefore, these three CsRDR1 proteins are able to interact with 2b in yeast cells. The specificity of these interactions was confirmed by testing other CMV proteins.
CP [Figure 4b] and movement protein [MP; Figure S1]) as negative control bait proteins. None of the additional viral proteins had any interaction with CsRDR1s, as no growth was observed on selection medium plates (Figures 4b and S1). Thus, it was observed that CMV 2b protein specifically bound the CsRDR1a/1b/1c proteins in yeast cells.

We also identified different regions of the 2b protein that are important for interactions. The 2b protein has distinct domains for different functions: amino acids 1–61 have nuclear localization signals (NLS), a dsRNA/siRNA-binding region and phosphorylation sites, the amino acid region 38–110 binds AGO proteins, and the region containing amino acids 62–110 lacks AGO-binding affinity (Figure 5a). Therefore, various 2b protein deletion AD-fusions (pGADT7), named 2b1–61, 2b1–95, 2b38–95, 2b38–110, 2b62–95, and 2b62–110, were generated to check binding with CsRDR1s (Figure 5b). In this study, it was found that CsRDR1s were able to interact with only a full-length 2b protein (Figure 4b) and that deletion of the 2b C-terminal region abolished the interaction with CsRDR1s, as no interaction was found with the 2b1–61, 2b1–95, 2b38–95, or 2b62–95 baits (Figure 5b). Thus, the 2b C-terminus region was shown to be essential for the interaction in yeast (2b-CsRDR1s). We were unable to use 2b38–110 and 2b62–110 as bait due to their autoactivation in yeast cells (Figure S2). In contrast, full-length 2b, 2b1–61, 2b1–95, 2b38–95, and 2b62–95 did not show any autoactivation in the GAL4-based Matchmaker Gold Y2H system (Figure S2). CsRDR1's RRM and RdRp domains showed stronger affinity for the

**FIGURE 2** (a) CsRDR1c expression in cucumber protoplasts induced in the presence of CMV 2b. Cucumber protoplasts were transfected with pSAT6-2bGFP (2b) and pSAT6-CPGFP (CP) under the 35S promoter and levels of expression of CsRDR1a, CsRDR1b, and CsRDR1c were analysed at 72 hr posttransfection by reverse transcription PCR. The sizes of amplified PCR products are written next to arrows. Cucumber protoplasts treated with empty pSAT6GFP vector (EV) or polyethylene glycol (PEG; Cont) served as controls. EF-1α expression (CsEF1α) served as a housekeeping-gene reference. Expression of the 2b protein fused to green fluorescent protein (GFP) in protoplasts was observed under (b) a bright field and (c) UV light (three protoplasts are marked with white arrows in both fields)

**FIGURE 3** Effects of CsRDR1s on the accumulation of CMV and CMVΔ2b in cucumber protoplasts. (a–c) CMV and CMVΔ2b accumulation in cucumber protoplasts transfected with pSAT6-eCFP constructs of CsRDR1aeCFP, CsRDR1beCFP, and CsRDR1ceCFP or empty pSAT6-eCFP vector (EV) along with in vitro transcripts of CMV or CMVΔ2b mutant. CMV and CMVΔ2b levels were determined by reverse transcription (RT) PCR of cucumber protoplasts at 68 hr after transfection. Virus accumulation was calculated with reference to CMV-transfected control samples. (d–f) Accumulation of CsRDR1a, CsRDR1b, and CsRDR1c was verified by quantitative RT-PCR and normalized to that of EF-1α. The experiment was repeated twice with similar results and data from one of the replicates is shown
2b protein than did full-length CsRDR1s (Figure 4b). This may be due to failure of exposure of important motifs due to improper folding of full-length proteins in yeast. However, no interaction was observed with the 2b C-terminus deletion baits (2b<sub>1-61</sub>, 2b<sub>1-95</sub>, 2b<sub>28-95</sub>, 2b<sub>62-95</sub>; Figure 5b). These results demonstrate the importance of the 2b C-terminus in this interaction, as summarized in Table 1.
2.5 | CsRDR1s interact with CMV 2b in planta and are localized in the cytoplasm

To verify the 2b–CsRDR1s interaction in planta, we performed bimolecular fluorescence complementation (BiFC) assays in Nicotiana benthamiana leaves. Growth media carrying Agrobacterium tumefaciens BiFC-bearing constructs of the N- and C-terminus halves of a yellow fluorescent protein (YFP), fused to the 2b or CsRDR1 coding sequences, were combined and agroinfiltrated into N. benthamiana leaves. Strong YFP reconstitution fluorescence was observed during CsRDR1a/1b/1c’s interaction with 2b in the cytoplasm 48 hr postagroinfiltration (Figure 6). However, no YFP reconstitution was observed with an empty vector (not shown) or CMV CP, which were used as negative controls. These results, obtained using intact leaves, further confirm that 2b specifically associates with CsRDR1s.

2.6 | CMV-2b protein colocalizes with CsRDR1s in the cytoplasm of protoplasts

The subcellular localization of CMV 2b-enhanced green fluorescent protein (2b-eGFP) and CsRDR1s-enhanced cyan fluorescent protein (CsRDR1a-eCFP, CsRDR1b-eCFP, and CsRDR1c-eCFP) was examined in cucumber protoplasts after separate transfection. At 48 hr posttransfection, expression of 2b was observed in the nucleus and cytoplasm (Figure 7a), whereas CsRDR1a, CsRDR1b, and CsRDR1c were localized in the cytoplasm as granular structures (Figure 7a). The coexpression of these CsRDR1s with 2b was associated with colocalization of signals throughout the cytoplasm (Figure 7b), which was quite distinct from the localization of the individual CsRDR1s (Figure 7a), that is, coexpression led to cytoplasmic redistribution of these proteins.

3 | DISCUSSION

Plant RDR1s are critical enzymes for RNA silencing that play a crucial role in the amplification of virus-silencing signals. Natural and artificial RDR1 mutants display induced susceptibility to various viruses (Rakhshandehroo et al., 2009; Wang et al., 2010; Yang et al., 2004). To overcome RNA silencing-mediated host resistance, viruses encode VSR proteins, which interfere with different steps of the RNA-silencing mechanism (Burgyán & Havelda, 2011).

We have previously shown that the members of the cucumber CsRDR1 gene family share 55%–60% homology. CsRDR1a and CsRDR1b, which are positioned in a head-to-head orientation, were expressed constitutively in healthy plants and a high level of CsRDR1b expression was associated with virus resistance in cucumber (Leibman et al., 2018). Here, we investigated the direct influence of the CMV silencing suppressor protein 2b on the expression of and interaction with members of the CsRDR1 gene family in cucumber. First, we analysed the expression of the four CsRDR1s (a, b, c1, and c2) during infection by CMV and CMVΔ2b in cucumber. We observed that CMV infection increased CsRDR1a and CsRDR1b levels 2- to 4-fold and increased CsRDR1c levels several thousand-fold. We have shown previously that infection of cucumber with CMV and viruses belonging to different families also activates CsRDR1c transcription (Leibman et al., 2018). In the presence of the CMVΔ2b mutant, CsRDR1a and CsRDR1b levels did not change, but the CsRDR1c level increased c.10-fold. In addition, in contrast to CsRDR1a and CsRDR1b, CsRDR1cΔ1/2 were not expressed in healthy cucumber or melon plants (data not shown). Therefore, based on the current study, we propose that higher levels of expression of CsRDR1c are dependent on either the virus titre or CMV 2b as reflected by CMV accumulation levels in plants (cv. Bet Alfa) that were infected with CMV or CMVΔ2b mutant. However, relative higher expression of CsRDR1c at 14 dpi despite lower virus titre compared to 7 dpi in CMVΔ2b mutant infected plants could be due to its accumulation with time after virus trigger or due to its autopositive regulation as observed in protoplasts on ectopic expression of CsRDR1c.

These results suggest that CsRDR1c has a different antiviral regulation and functional mechanism than either CsRDR1a or CsRDR1b and that the induced expression of CsRDR1c probably relies on the presence of the 2b protein. This mechanism must be absent from CsRDR1a and CsRDR1b. In Arabidopsis, AtRDR1 is regulated by a broad range of phytohormones (Hunter et al., 2013). Moreover, CMV infection induces and enriches plant immunity pathway genes, but not the expression of AtRDR1 (Zhao et al., 2018), suggesting differences between the regulatory systems and that CsRDR1c plays a different role in cucumber than it does in Arabidopsis. Therefore, we examined the effect of 2b on the expression of genes from the CsRDR1 family at the cellular level in cucumber protoplasts.

We showed, for the first time, that a suppressor like 2b can directly or indirectly induce CsRDR1c gene expression at the cellular level in protoplasts (after 68 hr) and in plants. Because 2b encodes two NLS motifs leading to nuclear localization (Wang et al., 2004).
**FIGURE 5**  (a) Schematic representation of important domains in the 2b protein (CMV-Fny strain; marked with black arrows) that were used for yeast two-hybrid (Y2H) analysis. 2b protein sequences from Subgroup I were aligned to analyse conserved domains. 2b1-61, 2b38-110, 2b62-110, 2b1-95, 2b38-95, and 2b62-95 represent the selected target regions for bait constructs. The 2b1-61 domain includes nuclear localization signals NLS1 and NLS2 (marked with blue lines), and phosphorylation sites (shown by blue triangles). (b) Interaction of CMV-Fny 2b protein deletion mutants with CsRDR1s, both full-length and specific domains, in yeast cells. CMV 2b protein fragments were fused to the GAL4 DNA-binding domain (DNA-BD). Full-length cucumber RDR1 proteins (RDR1a, RDR1b, and RDR1c) and specific domain regions (RNA recognition motif, RRM; RNA polymerase domain, RdRp) were expressed as fusions with the GAL4 activation domain (AD). After yeast mating, diploid transformants were selected on double-dropout selection (DDO) plates at 28 °C for 3 days. Then a dilution assay was performed on double-dropout (DDO), triple-dropout (TDO), and quadruple-dropout (QDO) selection plates (supplemented with the antibiotic aureobasidin A) involving up to $10^{-4}$ dilutions. Plates were kept for 3–4 days before imaging. The 2b protein, along with empty vector control, and CsRDR1s were used as negative controls.
and has DNA-binding abilities (Sueda et al., 2010), we propose that 2b acts like a transcription factor in the regulation of CsRDR1c gene expression.

CMV-2b is a well-known virulence factor that compromises host defences either by suppressing salicylic acid (SA)-mediated resistance or by binding long/small dsRNAs, DNA, AGO proteins, and CMV 1a protein (Diaz-Pendon et al., 2007; Ji & Ding, 2001; Lewsey et al., 2010; Watt et al., 2020; Zhao et al., 2018). This is why a CMVΔ2b mutant with a nonfunctional 2b protein is weaker in suppression of host defence and accumulates to only very low levels due to the slow systemic movement of the virus (Lewsey et al., 2009). Although 2b deletion was found to affect the initial rate of replication in protoplasts, the lack of 2b did not prevent the replication of a mutant virus, which later accumulated to similar levels (Soards et al., 2002).

In light of this, it is possible that 2b has dual activity: inducing CsRDR1c transcription to control virus accumulation and acting as a suppressor to block CsRDR1c activity. Previous studies have shown that the 2b protein inhibits the generation of secondary viral siRNA produced by RDR1 by binding CMV siRNAs and dsRNA precursors (Diaz-Pendon et al., 2007), inhibiting the slicing activity of AGO1/4 by direct interaction with these proteins (Diaz-Pendon et al., 2007; Fang et al., 2016; Hamera et al., 2012) and suppressing aphid resistance (Tungadi et al., 2020). Recently, CMV CP was also found to inhibit 2b and antagonize its viral suppressor function while maintaining a balance between virus accumulation and plant development (Zhang et al., 2017). Thus, the present study demonstrates

### Table 1

Summary of protein–protein interactions between 2b and CsRDR1s in yeast cells

|       | 2bI1<sup>-110</sup> | 2bI1<sup>-95</sup> | 2bI2<sup>-95</sup> | 2bI2<sup>-95</sup> | 2bI6<sup>-61</sup> |
|-------|---------------------|-------------------|-------------------|-------------------|-------------------|
| CsRDR1a | +                   | -                 | -                 | -                 | -                 |
| CsRDR1a<sub>RRM</sub> | +                   | -                 | -                 | -                 | -                 |
| CsRDR1b<sub>RdRp</sub> | +                   | -                 | -                 | -                 | -                 |
| CsRDR1b | +                   | -                 | -                 | -                 | -                 |
| CsRDR1b<sub>RRM</sub> | +                   | -                 | -                 | -                 | -                 |
| CsRDR1b<sub>RdRp</sub> | +                   | -                 | -                 | -                 | -                 |
| CsRDR1c | +                   | -                 | -                 | -                 | -                 |
| CsRDR1c<sub>N</sub> | +                   | -                 | -                 | -                 | -                 |
| CsRDR1c<sub>RdRp</sub> | +                   | -                 | -                 | -                 | -                 |

**Figure 6** Bimolecular fluorescence complementation assay showing interaction between the CMV-2b protein and cucumber RDR1 proteins (CsRDR1a, CsRDR1b, and CsRDR1c) in Nicotiana benthamiana leaf epidermal cells. 2b and CsRDR1 proteins were fused to the YFP<sup>C</sup> terminus in the pSPYCE(M) vector and at the YFP<sup>N</sup> terminus of the pSPYNE173R vector, respectively, and infiltrated in the following combinations: 2b-YFP<sup>C</sup>/CsRDR1a-YFP<sup>N</sup>, 2b-YFP<sup>C</sup>/CsRDR1b-YFP<sup>N</sup>, and 2b-YFP<sup>C</sup>/CsRDR1c-YFP<sup>N</sup>. CMV coat protein (CP) was also used as a negative control to analyse the interaction with CsRDR1 proteins in the following combinations: CP-YFP<sup>C</sup>/CsRDR1a-YFP<sup>N</sup>, CP-YFP<sup>C</sup>/CsRDR1b-YFP<sup>N</sup>, and CP-YFP<sup>C</sup>/CsRDR1c-YFP<sup>N</sup>.
a new role for 2b in virus self-attenuation based on its control of CsRDR1c expression in cucumber.

The transient expression of CsRDR1a and CsRDR1b proteins in cucumber protoplasts resulted in the suppression of both CMV and CMVΔ2b titres, suggesting roles for those proteins in plant defence. This observation is also supported by earlier studies of the role of RDR1 in plant defence against CMV infection, in which RDR1 was shown to be important for the amplification of siRNAs from the 5′ region of CMV genomic RNA to enhance antiviral immunity (Wang et al., 2010). Later, RDR1 was also reported to play a role in biogenesis of virus-activated siRNAs in Arabidopsis, which target endogenous genes during CMVΔ2b infection (Cao et al., 2014). The observation that CMV and CMVΔ2b RNA levels increase in cucumber protoplasts on CsRDR1c overexpression suggests that CsRDR1c encourages viral replication at the cell level and that the presence of the 2b protein is important for blocking CsRDR1c activity at the plant level. These data suggest that CsRDR1c might play a dual role: contributing to SA-mediated antiviral defences and suppressing gene silencing. Similarly, overexpression of Nt-RDR1 in N. benthamiana enhanced virus hypersusceptibility, suggesting RDR6-mediated antiviral RNA silencing (Ying et al., 2010). Earlier studies also showed that CMVΔ2b is unable to induce disease symptoms and demonstrated reduced virus accumulation (Lewsey et al., 2009). Thus, the present study presumes that, in addition to the CMV CP regulation of 2b (Zhang et al., 2017) and CMV 1a-mediated regulation of the 2b–AGO1 association (Watt et al., 2020), the equilibrium between CMV-2b and CsRDR1c protein levels controls virus accumulation and the infection progress.

Direct interaction was observed between CsRDR1s and the 2b protein. Previously, the 2b protein was shown to reduce the catalytic activity of AGO1 and AGO4 via direct interaction with their PIWI domains to counter host defence activity (Hamara et al., 2012; Zhang et al., 2006). In the present study, the 2b protein directly associated with the RRM and RdRp domains (Shao & Lu, 2014; Willmann et al., 2011) of the CsRDR1a, CsRDR1b, and CsRDR1c proteins in yeast cells and in N. benthamiana leaves, as observed via BiFC assays. This interaction occurred only with the intact 2b protein, in contrast with the interaction with AGO proteins, and may indicate that the protein structure is essential for such contact. The 2b tetramer form is the biologically active moiety associated with siRNA duplexes (Gellért et al., 2012) and such a structure may be necessary for the interaction with RDR1 proteins.

In cucumber protoplasts, 2b was localized to the nucleus and also found throughout the cytoplasm; the CsRDR1a/b/c proteins were
found throughout the cytoplasm in punctate spots. This finding is congruent with findings from previous studies of the 2b protein (Duan et al., 2012; Wang et al., 2004). Information about the subcellular distribution of RDR1 was not available previously and only RDR6 was known to be localized in cytoplasmic bodies (Kumakura et al., 2009). The colocalization of 2b and CsRDR1s in plant cells strengthens the assumption of a direct interaction between 2b and CsRDR1s, as demonstrated in the yeast system.

Coexpression of CMV-2b and CsRDR1s in cucumber protoplasts resulted in the redistribution of these proteins throughout the cytoplasm into a very distinct pattern that differed from the pattern of the localization of the individual proteins. Moreover, redistribution of proteins in the presence of an interacting partner has also been observed in previous studies, specifically 2b coexpression with AGO1/AGO4 (Duan et al., 2012) and AtRDR6 with rice yellow stunt virus P6 protein (Guo et al., 2013). Nucleus/nucleolar localization of the 2b is essential for CMV virulence, while its cytoplasm-localized form is more efficient in silencing suppression. The 2b protein partitions between the nucleus and cytoplasm, and thereby assists in maintaining the balance between CMV accumulation and injury to the host plant. Thus, 2b-mediated redistribution of AGO1/4 to the nucleus benefits the virus by interfering with RDR-dependent degradation of viral genomic RNA in the cytoplasm (Du et al., 2014). The present study suggests that CMV has evolved another approach to counter host antiviral defence: it directly interferes with host RDR1 activity to prevent secondary amplification of viral siRNAs. From a plant-defence perspective, virus-mediated induction of CsRDR1s in cucumber seems to play an important role in host defence against CMV by suppressing 2b virulence.

Taken together, these results present a dual role for the CMV-2b protein (Figure 8). In one role, it physically interacts with host-protein RNA-silencing components (i.e., CsRDR1s) to interfere with the amplification of viral siRNAs. In the other role, 2b acts like a transcription factor with its DNA-binding properties to affect the expression of CsRDR1c, and to balance plant growth and virus accumulation. These results also lead to the concept that higher induction of CsRDR1a/1b/1c genes during infection counters viral infection, but that CMV has evolved a strategy to counter this function that involves a 2b-mediated interaction. Furthermore, it may be suggested that a shift in equilibrium between 2b and CsRDR1c protein levels directly affects virus accumulation and the infection progress.

4 | EXPERIMENTAL PROCEDURES

4.1 | Plant growth conditions and virus inoculations

N. benthamiana plants were grown in a growth chamber at 25 °C with a 14-hr photoperiod. These plants were used for agroinfiltration at the three- to four-leaf stage (Shiboleth et al., 2007). Cucumber cultivars Bet Alfa and Shimshon were maintained under growth conditions similar to those described above and plants at the cotyledon stage with an emerging first true leaf were inoculated mechanically with CMV-Fny (Wang et al., 2003). For inoculum, CMV-infected leaves were homogenized in 50 mM sodium phosphate buffer, pH 7.0, and mock controls were inoculated with buffer only. The samples were collected and frozen in liquid nitrogen until further analysis of virus titre. For protoplast isolation, the fully expanded first cucumber leaves (c.15 days after germination) from plants grown as above were used.

![FIGURE 8](image-url) A model describing the regulation of the cucumber defence system as an equilibrium between CMV-2b virulence and the RNA-silencing activity of cucumber RDR1s. CMV-2b interferes with cucumber RDR1s’ (1a, 1b, 1c) induction of antiviral silencing by directly binding these proteins and thereby interfering with the RDR1-mediated viral siRNA amplification process. The presence of CMV-2b also acts as a trigger for the induction of CsRDR1c expression. Hence, oscillations in the equilibrium between cucumber RDR1c and CMV-2b protein levels direct virus accumulation. Only steps important for CMV pathogenicity are shown in this figure. dsRNA, double-stranded RNA; AGO1, argonaute protein 1; RDR, RNA-dependent RNA polymerase protein; SA, salicylic acid; 2b, CMV-2b protein; CP, CMV coat protein.

Enhanced siRNA amplification activity

More viral siRNA = Less virus accumulation

Suppressed siRNA amplification activity

Less viral siRNA = More virus accumulation
4.2 Construct preparation

To prepare Y2H bait constructs, PCR products of CMV-Fny 2b (full-length and various domains), CP, and MP (accession D10538) were cloned into the pGBK7 vector (Clontech) at the appropriate restriction sites (Table S1). Similarly, full-length and domain regions of CsRDR1a (accession KT316427), CsRDR1b (accession KT316425), and CsRDR1c1 (accession KT316426) were cloned into the pGADT7 vector (Clontech).

For the BiFC assay (Waadt et al., 2008), 2b and CP were cloned at BamHI/Xhol and BamHI/Smal sites, respectively, into the pSPYCE(M) vector to generate 2b-pSPYCE and CP-pSPYCE. CsRDR1a/1b/1c1 were cloned at BamHI/Smal, BamHI/Xhol, or BamHI/SacI sites in pSPYNE(R) 173 to produce CsRDR1a-pSPYNE, CsRDR1b-pSPYNE, and CsRDR1c-pSPYNE, respectively. For subcellular localization in protoplasts, the 2b and CP genes were cloned at Ncol/BamHI sites in the pSAT6-EGFP-N1 vector (Chung et al., 2005). CsRDR1a/1b/1c1 genes were cloned in the Sall/BamHI, Xhol/BamHI, or EcoRI/BamHI sites, respectively, in pSAT6-ECFP-C1.

4.3 Generation of CMV-2b mutants

Infectious clones of CMV-Fny (pFny109, pFny209, and pFny309; Rizzo & Palukaitis, 1990) and pFny209Δ2b were used to produce in vitro transcripts. pFny209Δ2b contains one nucleotide change in the first ATG of the coding frame, that is, ATG was changed to ACG (methionine to threonine), so that methionine (met1) would not be available for starting translation of the 2b gene. However, we found that this CMVΔ2b reverts to wild type 7 dpi in cucumber, whereas the mutation was found to be stable in N. benthamiana and N. tabacum. Therefore, a new CMVΔ2b infectious clone (with the first three codons for methionine mutated ATG→ACC) was generated as described by Du et al. (2007); that mutation was stable in cucumber.

Briefly, overlap extension PCR was used to generate fragments that had terminal HindIII and PstI restriction sites for subsequent cloning into the pFny209 plasmid. The first PCR product was amplified using primers CMVRNA2-1893F (with HindIII) and the reverse primer specific for CMVΔ2b (3CC2b2419R); the second PCR product was amplified using RNA2-2471F and CMVRNA2R (with a PstI site) on the PCR template pFny209. The third PCR procedure was performed by mixing the PCR fragments of the first and second PCR procedures and involved the use of CMVRNA2-1893F and CMVRNA2R primer pairs.

4.4 Yeast two-hybrid assay

The protein–protein interaction between the 2b VSR and the cucumber RDR1 defence proteins was carried out using the GAL4-based Matchmaker Gold Yeast Two-Hybrid System (Clontech; Y2H). Y2H Gold and Y187 yeast strains were used for the transformation of bait and prey vectors, respectively. Small-scale transformation was performed according to the manufacturer’s protocols and plates were incubated at 28 °C for 3 days. Yeast mating was carried out with positive colonies of 2b and CsRDR1s as per the manufacturer’s instructions. Transformants were selected on selection medium plates: SD/-Leu/-Trp/+AbA, followed by SD/-Leu/-Trp/-His/-Ade/+AbA. For the dilution assay, double-transformants were grown in SD/-Leu/-Trp broth overnight and different dilutions at equal OD600 were spotted on selection plates SD/-Leu/-Trp/-His/-Ade/+AbA, followed by incubation at 28 °C for 3–4 days.

4.5 BiFC assay

In planta confirmation of interactions between 2b and CsRDR1a/1b/1c1 (with CP used as a control) was performed in N. benthamiana by BiFC assay. pSPYNE(R) 173 and pSPYCE (M) vectors were used for cloning (Waadt et al., 2008) and these BiFC constructs were transformed into the A. tumefaciens GV3101. Further agroinfiltration was carried out by resuspending A. tumefaciens cells in MES-MgCl2 induction buffer (10 mM MES, 10 mM MgCl2, pH 5.6, 200 µM acetosyringone) at a final OD600nm of 0.5/0.5 for target proteins and 0.3 for the p19 suppressor, using a needleless syringe. YFP fluorescence was observed in infiltrated leaves at 48 hr after treatment under a confocal microscope.

4.6 Protoplast isolation and transfection

Protoplasts were isolated from cucumber leaves as described previously (Owen et al., 2016), with minor modifications to the enzymolysis and transfection buffers. In the enzyme buffer, 0.5% Onozuka cellulase R10, 0.25% Onozuka macerase, bovine serum albumen (0.1%), and CaCl2 (10 mM) were added to MMC buffer (600 mM mannitol, 5 mM MES, 10 mM CaCl2, pH 5.8) and the solution was heated to 55 °C for 10 min. For the transfection buffer, 35% polyethylene glycol (PEG; 4,000) and 5 mM CaCl2 solution were used. For a single transfection, 15 µg of the respective plasmid or transcripts were used. For colocalization, 9 µg of each were used. Briefly, for transfection, 200 µl of protoplasts containing 2 × 10⁶ cells was mixed gently with plasmids (30 µl) and 500 µl of PEG (35%) was added to this mixture. The mixture was then incubated at room temperature for 5 min. Transfected protoplasts were incubated in the dark at 26 ± 2 °C for 68 hr.

4.7 Confocal microscopy

For confocal microscopy, agroinfiltrated leaf sections were cut and mounted in water under a cover glass. The protoplasts were observed on a slide containing a depression in the centre under a cover glass. Simultaneous detection of two fluorophores and
chloroplast autofluorescence was performed by sequential scanning at their respective excitation and emission spectra. Excitation/emission wavelengths for GFP, CFP, YFP, and the chloroplasts were 488 nm/497–565 nm, 442 nm/450–500 nm, 514 nm/520–565 nm, and 514 nm/680–740 nm, respectively. Each fluorescent protein was scanned individually to avoid crosstalk.

4.8 | RT-qPCR

Total RNA was extracted from mock control and CMV-inoculated cucumber leaf samples using TRizol. One microgram of total RNA was treated with DNase (Thermo Fisher Scientific) according to the manufacturer’s instructions. First-strand cDNA synthesis was performed using the qPCRBIOS cDNA synthesis kit (PCR Biosystems). Quantitative PCR was performed using 2.5 µl of diluted cDNA, 7.5 µl of SYBR green 2x master mix (Applied Biosystems) and 1 µl of forward and reverse primer (3 µM each), in a final volume of 15 µl of reaction mixture in a StepOne Real-Time system (Applied Biosystems). The reaction conditions included denaturation at 94 °C for 20 s, followed by 40 cycles of 94 °C for 3 s and 60 °C for 30 s. The host EF-1α gene was used as an internal control for cDNA normalization. For each replicate, total RNA was extracted from pooled samples of three test plants. Relative expression was quantified using the 2^−ΔΔCt method. Three biological replicate leaf discs (50 mg fresh weight) were taken for analysis and each sample was a pool of three plants.

4.9 | Protein extraction and western blotting

For detection of CMV CP in inoculated plants, total protein was extracted from plant samples using urea-SDS extraction buffer (7.5 mM Tris-HCl pH 6.8, 9 M urea, 4.5% sodium dodecyl sulphate [SDS], 7.5% β-mercaptoethanol). The homogenized samples were incubated at 100 °C for 10 min and then centrifuged in a microfuge at maximum speed for 10 min. The supernatant was analysed on 12% polyacrylamide gel. Three biological replicate leaf discs (50 mg fresh weight) were taken for analysis and each sample was a pool of three plants.

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CONFLICT OF INTEREST
Authors declare no competing financial interest.

DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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SUPPORTING INFORMATION
Additional Supporting Information may be found online in the Supporting Information section.

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