The geminivirus nuclear shuttle protein is a virulence factor that suppresses transmembrane receptor kinase activity

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Despite the large number of leucine-rich-repeat (LRR) receptor-like-kinases (RLKs) in plants and their conceptual relevance in signaling events, functional information is restricted to a few family members. Here we describe the characterization of new LRR-RLK family members as virulence targets of the geminivirus nuclear shuttle protein (NSP). NSP interacts specifically with three LRR-RLKs, NIK1, NIK2, and NIK3, through an 80-amino acid region that encompasses the kinase active site and A-loop. We demonstrate that these NSP-interacting kinases (NIKs) are membrane-localized proteins with biochemical properties of signaling receptors. They behave as authentic kinase proteins that undergo autophosphorylation and can also phosphorylate exogenous substrates. Autophosphorylation occurs via an intermolecular event and oligomerization precedes the activation of the kinase. Binding of NSP to NIK inhibits its kinase activity in vitro, suggesting that NIK is involved in antiviral defense response. In support of this, infectivity assays showed a positive correlation between infection rate and loss of NIK1 and NIK3 function. Our data are consistent with a model in which NSP acts as a virulence factor to suppress NIK-mediated antiviral responses.

[Keywords: Receptor-like kinases; nuclear shuttle protein; geminivirus; defense signaling]

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Receptor-like protein kinases [RLKs] are a diverse group of membrane-spanning proteins with a predicted signal sequence and a cytoplasmic kinase domain that have been implicated in a wide range of signal transduction pathways. The complexity of this family in the plant kingdom may reflect an intensified communication of the plant cell with its environment by perception and transduction of external signals throughout development. In Arabidopsis, whose complete genomic sequence has been determined [The Arabidopsis Genome Initiative 2000], the RLKs make a large gene family comprising 417 family members with receptor configurations [Shiu and Bleecker 2001]. These putative transmembrane receptors are structurally organized into a divergent extracellular domain at the N-terminal portion that is thought to confer ligand-binding specificity followed by an internal transmembrane segment and a predicted cytoplasmic signal-transducing domain that includes a juxtamembrane domain, a conserved serine/threonine kinase domain, and a C-terminal region. On the basis of their extracellular domain, these receptors have been organized into different structural classes.

The RLK subfamily with extracytoplasmic leucine-rich repeats [LRRs] represents the largest class of putative receptor-encoding genes in the Arabidopsis genome [Shiu and Bleecker 2001; Dievart and Clark 2004]. They are classified into 13 subfamilies [LRR I–XIII] on the basis of the LRR organization, ranging from three to 26 LRRs. Despite their conceptual relevance in cell signaling events, biological function has been assigned to only a few plant LRR-RLKs, in development and defense responses. Examples of the developmental LRR-RLK genes include the Arabidopsis ERECTA and CLAVATA1 genes that determine floral organ shape and size [Torii et al. 1996; Clark et al. 1997], HAESA that regulates floral abscission [Jinn et al. 2000], SERK1 that is involved in ovule development and early embryogenesis [Hecht et al. 2001], and BR11 and BAK1 that are involved in brassinosteroid perception and signaling [Li et al. 2002; Nam and Li 2002]. The defense genes are represented by Xa-21 from Oryza sativa, which confers resistance to the bacterial pathogen Xanthomonas oryzae pv. Oryzae [Song et al. 1995] and FLS2 involved in the perception of the bacterial elicitor flagellin [Gomez-Gomez and Boller 2000]. Recently, we demonstrated that a nuclear shuttle protein [NSP] from the tomato-infecting geminiviruses TG MV [Tomato golden mosaic virus] and TCrLYV [Tomato crinkle leaf yellows virus] inter-
acts stably with tomato and soybean proteins belonging to the LRR-RLK family and designated as LeNIK (Lycoopersicium esculentum NSP-interacting kinase) and GmNIK (glycine max NIK) [Mariano et al. 2004]. However, the biological significance of this interaction remains to be elucidated.

Geminiviruses constitute a large group of plant viruses whose genomes are packed as single-stranded DNA circles in small, twinned isometric particles and are converted to double-stranded forms in nuclei of differentiated plant cells [Hanley-Bowdoin et al. 1999]. Members of the genus Begomovirus, such as TGMV and TCrLYV, possess two genomic components, DNA-A and DNA-B. The DNA-A has the potential to code for five gene products and is involved in replication, transcriptional activation of viral genes, and encapsidation of the viral genome [Elmer et al. 1988; Kallender et al. 1988; Sunter et al. 1990; Fontes et al. 1992, 1994a; Sunter and Bisaro 1992]. The DNA-B component encodes two movement proteins, the nuclear shuttle protein NSP (BV1) and the movement protein MP (BC1), both required for systemic infection [Lazarowitz and Beachy 1999; Gafni and Epel 2002]. NSP facilitates the intracellular trafficking of viral DNA between the nucleus and the cytoplasm, whereas MP potentiates its cell-to-cell movement.

Current models of viral cell-to-cell movement hold that the translocation of plant virus into adjacent uninfected cells occurs via plasmodesmata or viral-induced, endoplasmic reticulum (ER)-derived tubules [Lazarowitz and Beachy 1999; Gafni and Epel 2002]. From the biochemical activities of MP and NSP from Squash leaf curl virus (SLCV) and Bean dwarf mosaic virus (BDMV), it has been proposed that geminiviruses use both mechanisms for cell-to-cell trafficking of viral DNA [Noueiry et al. 1994; Sanderfoot and Lazarowitz 1996]. In the first model, NSP facilitates the intracellular movement of the viral genome from the nucleus to the cytoplasm, where it is replaced by MP that transports the viral DNA to adjacent cells via plasmodesmata [Noueiry et al. 1994]. Accordingly, MP from BDMV has been shown to act as a classical movement protein with the capacity to bind viral DNA in vitro, to increase plasmodesmal size exclusion limit, and to promote translocation of the viral genome to adjacent cells in microinjection studies [Noueiry et al. 1994; Rojas et al. 1998]. In the second model, MP facilitates NSP-mediated intracellular transport of viral DNA from the nucleus to the cytoplasm and then mediates the transport of the NSP–DNA complex to adjacent cells via ER-derived tubules induced by the viral infection [Lazarowitz and Beachy 1999]. Consistent with this model, MP from the phloem-limited SLCV has been immunolocalized to unique tubules that extend across the walls of developing phloem cells and has been shown to relocate NSP from the nucleus to the cell periphery [Sanderfoot and Lazarowitz 1995; Sanderfoot et al. 1996; Ward et al. 1997]. In addition, SLCV MP does not bind DNA in vitro, whereas NSP does [Pascal et al. 1994]. The apparent discrepancies between these two models have been explained by the phloem limitation of SLCV, whereas BDMV can also infect nonvascular tissues. However, both models rely on the observations that NSP shuttles viral DNA between the nucleus and the cytoplasm and acts in concert with MP to promote the cell-to-cell spread of viral DNA [Lazarowitz and Beachy 1999; Gafni and Epel 2002].

The localization of NSP and its proposed role in cell-to-cell movement of the viral DNA predict that interactions with host factors may occur in both the cytoplasm and the nucleus. Thus, the interaction of NSP with transmembrane factors could involve an active recognition in the nuclear pore, plasma membrane, or plasmodesmata [Mariano et al. 2004]. The recent demonstration that NSP from tomato-infecting geminiviruses associates physically with LRR-RLKs may provide insight into the functions of this complex class of putative transmembrane receptors. In this investigation, we took advantage of the capacity of the geminivirus CaLCuV (Cabbage leaf curl virus) to infect Arabidopsis [Hill et al. 1998]. In Arabidopsis, reverse genetic studies are possible [Alonso et al. 2003] to decipher the function of these NSP-interacting putative transmembrane receptors and to determine the biological relevance of the NIK–NSP interaction. We demonstrate that the Arabidopsis NSP-interacting LRR-RLKs are authentic protein kinases with biochemical properties of signaling receptors. Binding of NSP to NIK inhibits its kinase activity, and loss of NIK gene function enhances susceptibility to geminivirus infection. Our data suggest that NIKs are involved in antiviral defenses and that NSP potentially counters the proactivation mechanism of this pathway by inhibiting NIK kinase activity.

Results

NSP interacts with LeNIK- and GmNIK-related proteins from Arabidopsis that belong to the LRRII subfamily of the RLK family

We demonstrated previously that NSP interacts with the kinase domain of a putative transmembrane receptor from tomato (LeNIK) and soybean (GmNIK) [Mariano et al. 2004]. The full-length host proteins exhibit a modular organization that resembles members of the Arabidopsis LRRII subfamily of the RLK family [Shiu and Bleecker 2001]. Among them, GmNIK is most closely related to At5g16000-encoded product (U19571, 76% sequence identity), followed by At3g25560 (70%) and At1g60800 (61%), whose functions are unknown. The sequence conservation increases to 85% identity if just the kinase domain is used as the basis for comparison. It also shares significant conservation of primary structure with the other members of this family (40%–55% identity). We selected the three GmNIK-most-related LRRII-RLKs to evaluate their capacities to interact with NSP from tomato-infecting geminiviruses (TGMV and TCrLYV) and from an Arabidopsis-infecting geminivirus (CaLCuV). Interactions were assessed by the two-hybrid system monitored for histidine and uracil prototrophy. NSPs interacted with both intact (KD) and nucleotide-binding site (NBS)-deleted kinase (ΔKD) domains of all tested
members of the LRRII-RLK subfamily (Table 1). Interactions of two partners were confirmed by monitoring β-galactosidase activity in yeast protein extracts. The NSP interactions were specific to members of the LRRII-RLK subfamily, because NSP did not interact with either an active or an inactive kinase domain of BRI1, a 25 LRR-containing receptor kinase that belongs to the LRR IX subfamily of the RLK family (Shiu and Bleecker 2001; Dievart and Clark 2004) (Fig. 1B). Based on primary sequence conservation as compared with GmNIK and LeNIK, as well as from their capacity to interact with NSP, At5g16000, At3g25560, and At1g60800 are designated here as NIK1, NIK2, and NIK3, respectively. These results validated our previous interpretation that NSP–NIK complex formation was neither virus specific nor host specific (Mariano et al. 2004).

Specificity of NSP interaction among members of the LRR-RLK family

To further confirm the interaction of NSP with NIKs, we performed an in vitro protein binding assay (Fig. 1A). Purified GST-NSP or GST was incubated with the in vitro translated [35S]Met-labeled intact NIK1 or NIK3, as well as the kinase domain (KD) or the NBS-deleted kinase domain (KD) of NIK1, NIK2, NIK3, and AtSERK1, as indicated in Figure 1A. The resulting complexes were isolated on glutathione-Sepharose beads. Except for KD-SERK1, all the other in vitro labeled proteins, either as intact or truncated kinases, bound to GST-NSP but not to GST alone, indicating that NSP specifically and stably interacts with NIK1, NIK2, and NIK3, but not with AtSERK1 (Fig. 1A). Although the truncated kinase domain of AtSERK1 (KD-SERK1) interacted with NSP in yeast, as monitored by His and Ura prototrophy of double-transformed yeast cells, quantification of β-galactosidase activity resulted in a weak signal [0.18 and 0.21 units], whereas NSP interaction with KDNIK1, KD-NIK2, and KDNIK3 gave higher values [Table 1]. Collectively, these results suggested that NSP discriminates among the members of the LRRII-RLK subfamily. Because NSP interaction with AtSERK1 was weak in yeast and barely detected in vitro, AtSERK1 was not considered further.

To determine the region of NIK1 responsible for specific interaction with NSP, we fused various truncations of its kinase domain to GAL4 activation domain and assayed them for their interaction with NSP in yeast. These results are shown in Figure 1B, and the kinase deletions that scored positively for histidine and uracil prototrophy are schematized in Figure 1C. The deletion analysis mapped the NSP-interacting domain in NIK1 to an 80-amino acid region of the kinase domain [amino acids 422–502] that encompasses the putative active site for Ser/Thr kinases (subdomain VIb–HrDvKssNxLLD) and the activation loop [subdomain VII–DFGAk/rx, plus subdomain VIII–GtxGyiaPEY] (Fig. 1C).

The NIKs are plasma membrane-localized proteins that exhibit biochemical properties consistent with a receptor signaling function

The NSP-interacting LRRII-RLKs possess an internal transmembrane helix and a peptide signal that may target the protein to the secretory apparatus. Consistent with this observation, NIK1 fractionated with leaf microsomal preparations composed primarily of endomembrane vesicles and plasma membranes [data not shown].

| Table 1. Interaction of NSP from Arabidopsis and tomato-infecting geminiviruses with members of the LRRII-RLK family in yeast |
|----------------|-----------------|-----------------|-----------------|-----------------|
| Bait           | pDBLeu          | pBD-NSPCLCV     | pBD-NSPTGMV     |
| Prey           | +H +U           | −H +3AT         | −U β-Galα       | +H +U           | −H +3AT         | −U β-Galα       |
| pAD-KDNIK1     | +               | −               | −               | 0.0             | +               | +               | +               | +               | +               | +               | 0.642 ±         |
| pAD-ΔKDNIK1    | +               | −               | −               | 0.0             | +               | +               | +               | 0.512 ±         | +               | +               | 0.642 ±         |
| pAD-KDNIK2     | +               | −               | −               | 0.0             | +               | +               | +               | 0.528 ±         | +               | +               | 0.505 ±         |
| pAD-KDNIK3     | +               | −               | −               | 0.0             | +               | +               | +               | 0.425 ±         | +               | +               | 0.328 ±         |
| pAD-ΔKDNIK3    | +               | −               | −               | 0.0             | +               | +               | +               | 0.812 ±         | +               | +               | 0.724 ±         |
| pAD-ΔKDSERK1   | +               | −               | −               | 0.0             | +               | +               | +               | 0.812 ±         | +               | +               | 0.799 ±         |
| pAD-SBP        | +               | −               | −               | 0.0             | +               | +               | +               | 0.215 ±         | +               | +               | 0.187 ±         |

The bait proteins were expressed as GAL4 DNA-binding domain fusions, and the prey proteins were expressed as GAL4 activation domain fusions in yeast. KD corresponds to the C-terminal kinase domain of the protein and ΔKD is NBS-deleted kinase domain NIK1, NIK2, NIK3, and SERK1 are members of the Arabidopsis LRRII-RLK family. SBP is an unrelated sucrose-binding protein from soybean used as a control in the two-hybrid assays. (U) Uraclie; (H) histidine; (3AT) 25 mM 3-aminotriazole; (β-Gal) β-galactosidase activity. *Values for activity are the mean ± standard deviation from four replicas.
Furthermore, NIK1, NIK2, and NIK3 fused to green fluorescent protein (GFP) were targeted to the cell surface in transient assays of bombarded young leaves from Arabidopsis (Fig. 2A). Localization of NIK–GFP fusions was also analyzed in cauline cells of stably transformed plants thorough confocal laser scanning microscopy (Fig. 2B). Intense GFP fluorescence was observed in the cell surface delimitating the cell boundaries. Because of the chlorophyll autofluorescent background, chloroplasts were also labeled, although to a lesser extent.

The C-terminal kinase domain of NIKs contains all of the 11 conserved subdomains of protein kinases, in addition to specific signatures of serine/threonine kinases in subdomains V1b and VIII [Hanks et al. 1988; Hanks and Quinn 1991; Taylor et al. 1995]. To investigate whether NIKs possess kinase activity, we expressed their C-terminal regions containing an intact kinase domain (KD) or a truncated kinase domain (ΔKD), in which the critical subdomains I and II were deleted, as GST fusions in Escherichia coli and affinity-purified them on glutathione-Sepharose. Kinase activity was assayed in vitro by incubation of purified recombinant proteins with [γ-32P]ATP and analysis of 32P incorporation. As shown in Figure 3A, GST-KDNIK1, GST-KDNIK2, and GST-KDNIK3 undergo autophosphorylation in a Mg2+- or Mn2+-dependent manner and no activity was observed with CaCl2. In fact, inclusion of CaCl2 in Mg2+-based reactions reduced the kinase activity. In contrast, neither GST alone (data not shown) nor the truncated kinase domain fusions (GST-ΔKDNIK1 and GST-ΔKDNIK3) exhibited kinase activity (Fig. 3B). GST-ΔKDNIK1 did not show detectable autophosphorylation, consistent with its lack of the critical Ser/Thr kinase subdomain I and II. These results provide direct biochemical evidence that NIK1, NIK2, and NIK3 cytoplasmic domains possess a functional kinase activity that requires the presence of typical conserved domains for Ser/Thr kinases.

In plants, autophosphorylation of Ser/Thr kinases occurs via either an intra- or intermolecular mechanism [Schulze-Muth et al. 1996; Roe et al. 1997; Sessa et al. 1998; Shah et al. 2001b]. To examine the mechanism of
NIKs autophosphorylation, we tested the ability of the GST-fused kinase domains (KD) to phosphorylate inactive, truncated versions of transmembrane kinases (∆KD-NIK1, ∆KD-NIK3, and ∆KD-SERK1). As shown in Figure 3B, GST-KDNIK1 was able to phosphorylate the inactive ∆KD-NIK1 in trans. Transphosphorylation was not restricted to the cognate receptor, as ∆KD-NIK3 and ∆KD-SERK1 were also phosphorylated by NIK1. Likewise, NIK2 [Fig. 3B] and NIK3 (data not shown) could trans-phosphorylate the inactive, truncated kinase domains of the transmembrane receptors. These results indicate that NIK autophosphorylation occurs through an intermolecular mechanism. Furthermore, they suggest that NIKs phosphorylate exogenous substrates, because AtSERK1, which has been shown to be phosphorylated on serine and threonine residues [Shah et al. 2001b], was efficiently phosphorylated by the kinases.

In the mechanism of transphosphorylation-mediated receptor kinase activation, oligomerization precedes the activation of kinase domain [Lemmon and Schlessinger 1994; Roe et al. 1997]. To determine whether oligomerization is involved in regulation of NIK1 kinase activity, we took advantage of the capacity of GST to dimerize and the existence of a properly located thrombin-cleavage site on the recombinant protein that allows separation of GST from its fusion partner, KDNIK1. The kinase activity of the thrombin-cleaved GST-KDNIK1 was lower than that of intact GST-KDNIK1 [Fig. 3C, left panel]. After normalizing to the amount of protein, the difference was fourfold [Fig. 3C, bottom panel]. Progressive dilution of the recombinant protein into the kinase assay had a much greater impact on the specific kinase activity of the thrombin-cleaved GST-KDNIK1 than on that of intact GST-KDNIK1. Thus, in the absence of the GST dimer, the specific activity of the cleaved kinase portion was concentration dependent. These results support a mechanism for activation of the kinase domain involving trans-autophosphorylation mediated by oligomerization of the protein.

**NSP inhibits NIK kinase activity**

Given that NIKs are authentic serine/threonine kinase proteins and that Squash NSP is phosphorylated in infected plants [Pascal et al. 1994], we next examined whether the viral NSP is an NIK substrate. Neither NIK1 nor NIK2 phosphorylated GST-NSP from TGMV and CaLCuV [Fig. 4A]. In contrast, inclusion of the recombinant viral proteins in the kinase assays inhibited both NIK1 and NIK2 activity. Incorporation of 32P into GST-NIK1 and GST-NIK2 was reduced six- and twofold, respectively, by either NSP from CaLCuV or TGMV [Fig. 4A, bottom panel]. NSP from TCrLYV inhibited the kinase activity of the NIK1, NIK2, and NIK3 to the same extent as did TGMV-NSP or CaLCuV-NSP [data not shown]. Inhibition of NIK activity by GST-fused viral proteins was not due to the presence of the GST domain because His-tagged CaLCuV-NSP, which is not able to dimerize with GST-KDNIK1 via the GST tag, also inhibited the kinase activity of NIK1 [Fig. 4B]. A threefold molar excess of His-NSP was sufficient to abolish autophosphorylation of NIK1. The inhibitory activity of His-NSP was higher than that of GST-NSP [Fig. 4C] and this difference may be due to conformational constraints imposed by the GST tag.

Despite the strong conservation of the NSP-binding domain among the members of the LRR-RLK subfamily [Fig. 1C], NSP associated weakly with AtSERK1 [Fig. 1A] and did not inhibit its kinase [Fig. 4A]. Furthermore, NSP can be phosphorylated in vitro by reticulocyte kinases [data not shown] and has also been shown to be phosphorylated in vivo [Pascal et al. 1994]. Together, these observations suggest that the NSP inhibition of NIK is not due to casual interactions of NSP with kinases, and instead reflects an inherent, specific activity of the viral protein.

**NIK1, NIK2, and NIK3 exhibit partially overlapping but distinct expression patterns**

In Arabidopsis, CaLCuV infects leaf tissues with high efficiency, inducing symptoms of severe stunting and leaf epinasty [Hill et al. 1998]. To determine whether the onset of geminivirus infection correlates with NIK expression, we analyzed the expression pattern of NIK1, NIK2, and NIK3 by semiquantitative RT–PCR with gene-specific primers [Fig. 5A]. All RT–PCRs were repeated with different numbers of cycles to ensure a quan-
titative linear amplification. In addition, the integrity and amount of cDNA from different organs were routinely assessed with actin-specific primers. Both NIK1 and NIK3 transcripts were highly expressed in seedlings and leaves and accumulated to a lower level in roots. In contrast, NIK2 transcripts predominated in roots and were barely detected in seedlings and leaves. The specificity of the primers was confirmed in expression analysis of T-DNA insertion nik1, nik2, and nik3 alleles (see below) (Fig. 5B).

NIK knockout lines exhibit enhanced susceptibility to geminivirus infection

To assess directly the biological significance of NIK–NSP interactions, we identified T-DNA insertion mutants in each gene (Fig. 5C). RT–PCR was performed on leaf (L) and root (R) RNA samples from wild-type (NIK1*, NIK2*, and NIK3*), nik1, nik2, and nik3 knockout (KO) lines. With the gene-specific primers, we detected no accumulation of the corresponding transcript in the respective homozygous T-DNA insertion mutant, confirming that they are null alleles (Fig. 5B). nik1, nik2, and nik3 mutant plants and wild-type Col-0 lines were inoculated with CaLCuV DNA-A and DNA-B. Both Col-0 and nik1-KO lines developed typical CaLCuV symptoms of similar intensity. However, removal of coat protein sequences from CaLCuV DNA-A attenuated the virus in Col-0 plants but not in the nik1 null background (Fig. 6A, bottom panels). In this case, disease symptoms varied in severity from extreme stunting with severe epinasty and chlorosis in nik1-KO to mild stunting with epinasty and moderate chlorosis in Col-0 lines. This attenuated form of the virus was used to analyze the course of infection in the mutant lines. As judged by symptom appearance and viral DNA accumulation, inactivation of NIK1 and NIK3 alleles accelerated the onset of virus infection in comparison to Col-0- and NIK1*-infected plants, in which the T-DNA insertion segregated out recovering homozygous NIK1 wild-type alleles.
The infection rate for \textit{nik1} mutant and \textit{NIK1*} plants correlated with accumulation of viral DNA (Fig. 6B). In addition to being detected earlier in the \textit{nik1} mutant, viral DNA-A and DNA-B accumulated to higher levels in these lines compared with \textit{NIK1*}. The enhanced susceptibility phenotype was not observed to the same extent in \textit{nik2} null alleles. The infection rate for \textit{nik2-KO} was slightly higher than that for Col-0 but still considerably less than those for \textit{nik1-KO} and \textit{nik3-KO} (Fig. 6B). These results are consistent with the pattern of \textit{NIK1}, \textit{NIK2}, and \textit{NIK3} expression. In other experiments, the infectivity data, expressed as days postinoculation to get 50% of infected plants (DPI50%), further confirmed the results (Fig. 6D).

Discussion

A complete knowledge of the size and complexity of the LRR-RLK family has emerged with the sequence of the \textit{Arabidopsis} genome but functional information is limited to a few family members that are amenable to genetic approaches. Here we describe the biochemical characterization of new members of this family, identified as virulence targets of the geminivirus NSP. NSP interacted with three members of this family, \textit{NIK1}, \textit{NIK2}, and \textit{NIK3}, which were shown to be authentic Ser/Thr kinases with biochemical properties consistent with a receptor signaling function. NSP inhibited the kinase activity of \textit{NIK1}, 2, and 3, but not other kinases, suggesting their involvement in antiviral defense response. This interpretation was corroborated in vivo by infectivity assays showing a positive correlation between infection rate and inactivation of \textit{NIK1} and \textit{NIK3} gene expression. Furthermore, the failure of \textit{NIK2} null alleles to exhibit an enhanced susceptibility phenotype strengthened the argument that suppression of NIK-mediated defenses is NSP dependent, because the expression pattern of \textit{NIK2} is not spatially coordinated with the onset of CaLCuV infection and, hence, with NSP accumulation.

From primary structural analysis, \textit{NIK1}, \textit{NIK2}, and \textit{NIK3} are classified as LRR-RLKs and, thus, predicted to be transmembrane serine/threonine receptor kinases. We demonstrated that the NIK proteins exhibit a series of biochemical properties characteristic of authentic Ser/Thr receptor kinases that mediate signal transduction through membranes. First, the NIK proteins were local-
IZED TO THE PLASMA MEMBRANE. SECOND, WE HAVE SHOWN THAT THEIR CYTOPLASMIC DOMAIN IS A FUNCTIONAL KINASE WITH AUTOPHOSPHORYLATION PROPERTIES. DELETION OF THE CONSERVED SER/THR KINASE SUBDOMAINS I AND II ABOLISHED AUTOPHOSPHORYLATION OF GST-FUSION NIK PROTEIN, CONFIRMING THAT INCORPORATION OF $^{32}$P INTO THE FUSION PROTEIN IS DUE TO AN INTRINSIC KINASE ACTIVITY AND NOT TO A CONTAMINATING PROTEIN. THIRD, THE NIK KINASE DOMAIN WAS ABLE TO PHOSPHORYLATE EXOGENOUS SUBSTRATES. IN FACT, WE SHOWED THAT ATSERK1, WHICH HAS BEEN DEMONSTRATED TO BE PHOSPHORYLATED ON SERINE/THREONINE RESIDUES (SHAH ET AL. 2001B), IS A SUBSTRATE FOR NIK. FINALLY, THE MOLECULAR FEATURES OF NIK AUTOPHOSPHORYLATION DESCRIBED HERE PROVIDED INSIGHT INTO THE LIKELY MECHANISM OF NIK RECEPTOR ACTION. THE PHOSPHORYLATION OF AN INACTIVE, DELETED-KINASE DOMAIN BY THE Cognate RECEPTOR DEMONSTRATED THAT AUTOPHOSPHORYLATION OCCURS VIA AN INTERMOLECULAR REACTION. THIS TRANSPHOSPHORYLATION MECHANISM WAS FURTHER SUPPORTED BY THE FINDING THAT THE AUTOPHOSPHORYLATION REACTION EXHIBITED SECOND-ORDER KINETICS (FIG. 4C). INTERMOLECULAR AUTOPHOSPHORYLATION IS CONSISTENT WITH A RECEPTOR SIGNALING MECHANISM IN WHICH LIGAND-MEDIATED OLIGOMERIZATION IS REQUIRED FOR KINASE ACTIVATION. CONSISTENT WITH THIS MECHANISM, REMOVAL OF THE GST TAG, WHICH FUNCTIONS AS AN EXTRA DIMERIZING COMPONENT, FROM THE RECOMBINANT NIK PROTEIN REDUCED ITS KINASE ACTIVITY BY FOURFOLD. IT IS LIKELY THAT OLIGOMERIZATION OF NIK PRECEDES ACTIVATION OF THE CYTOPLASMIC KINASE DOMAIN THAT IS MEDIATED BY A TRANSPHOSPHORYLATION EVENT. DIMERIZATION AND INTERMOLECULAR AUTOPHOSPHORYLATION IS A WELL-DOCUMETED MECHANISM FOR MAMMALIAN RECEPTORS WITH TYROSIINE KINASE ACTIVITY (LEMMON AND SCHLESSINGER 1994) AND, MORE RECENTLY, HAS BEEN SHOWN TO DICTATE LRRRII-RLK ACTION IN PLANTS (SHAH ET AL. 2001A,B).

WE DEMONSTRATED THAT NSP EFFECTIVELY INHIBITS AUTOPHOSPHORYLATION OF NIK RECEPTORS, SUGGESTING THAT NSP COUNTERS THE PROACTIVATION MECHANISM OF A NIK-MEDIATED SIGNALING PATHWAY. ACCORDINGLY, THE NSP-INTERACTION DOMAIN WAS MAPPED ON NIK1 TO AN 80-AMINO ACID STRETCH CONTAINING THE PUTATIVE ACTIVE SITE AND THE ACTIVATION LOOP FOR SER/THR KINASES (FIG. 1C). THE A-LOOP PLAYS A CRITICAL ROLE IN CONTROLLING THE ACTIVITY OF PROTEIN KINASES, AS WELL AS IN SUBSTRATE RECOGNITION (ELLIS ET AL. 1986; JOHNSON ET AL. 1996), AND ITS PHOSPHORYLATION CONSTITUTES A KEY REGULATORY MECHANISM THAT INCREASES THE KINASE CATALYTIC ACTIVITY (FOR REVIEWS, SEE JOHNSON ET AL. 1998; HUBBARD AND TILL 2000). ALTHOUGH THE MOLECULAR MECHANISM FOR NSP INHIBITION OF NIKs REMAINS UNRESOLVED, THE OVERLAP OF THE NSP-INTERACTING DOMAIN WITH REGULATORY DOMAINS FOR KINASE ACTIVITY SUGGESTS PRIORITIES FOR STRUCTURE-BASED MUTAGENESIS STUDIES ON NIK. FOR EXAMPLE, INTERMOLECULAR AUTOPHOSPHORYLATION OF A CONSERVED THR 468 RESIDUE IN THE A-LOOP OF ATSERK1 IS REQUIRED FOR KINASE ACTIVATION (SHAH ET AL. 2001B). GIVEN THE REMARKABLE CONSERVATION OF THIS ESSENTIAL THREONINE IN THE A-LOOP OF ALL LRRRII-RLKS (FIG. 1C), IT IS LIKELY THAT NIKs USE A SIMILAR REGULATORY MECHANISM FOR KINASE ACTIVATION. IT WOULD BE INTERESTING TO VALIDATE EXPERIMENTALLY THIS PREDICTION AND TO KNOW WHETHER NSP BINDING TO NIK WOULD IMPAIR INTERMOLECULAR PHOSPHORYLATION OF THE NIK A-LOOP AND HENCE KINASE ACTIVATION.

ALTHOUGH WE HAVE BEEN UNABLE TO DETECT DIRECT INTER-
action of NIK and NSP in infected plants, as both proteins are expressed at extremely low levels, an in vivo functional link between these proteins has been provided by infectivity assays in loss-of-function mutants. 

nik1 and nik3 mutant plants displayed enhanced susceptibility phenotypes on geminivirus infection, whereas nik2 mutants were similar to Col-0, with comparable infection rates. These results implicated NIK1 and NIK3 as components of antiviral defenses, whereas NIK2 might not respond as effectively to geminivirus infection. Although we have shown that in vitro NSP-binding activity is a shared property of all three transmembrane receptors, NIK2 expression is barely detectable in leaves where the process of infection takes place with high efficiency [Hill et al. 1998]. In contrast, NIK1 and NIK3 are highly expressed in leaves, indicating that opportunistically they are capable of eliciting effectively defense strategies against geminivirus infection. Except for NSP that has been demonstrated to target and inhibit these transmembrane kinases specifically, none of the other viral-encoded proteins were found to interact with these transmembrane kinases in yeast [Mariano et al. 2004]. Collectively, these results support the argument that the geminivirus NSP acts as virulence factor to suppress NIK-mediated defenses.

Mutant alleles of nik1 and nik3 displayed an enhanced susceptibility phenotype on geminivirus infection, implicating these transmembrane receptors in host defense responses against geminivirus infection. A question that remains unanswered is how NIK-mediated responses act to impair viral infectivity and how NSP interaction would fit in this mechanism. Although NSP–NIK interaction exhibited some properties consistent with the guard hypothesis of plant resistance, our infection results are inconsistent with this possibility [Van der Biezen and Jones 1998]. According to the elicitor-receptor model of resistance, NSP would function as a virulence factor by binding to the transmembrane receptor NIK and impairing its interaction with a putative R protein, thereby preventing a resistance response. This model also predicts that in the absence of its putative R protein partner, suppression of NIK expression in these susceptible lines would not affect the efficiency of geminivirus infection. However, both nik1 and nik3 null alleles were isolated in the susceptible Col-0 genetic background and, in contrast to the guard hypothesis for NIK function, they exhibited enhanced infection rates. More likely, the transmembrane receptor NIK functions as an upstream signaling component of an alternative innate antiviral host defense to impact virus replication or movement negatively. Enhanced efficiency of viral DNA replication or cell-to-cell movement in nik1 and nik3 mutants would account for the increased infection rates and viral DNA accumulation in these lines. The action of NSP would potentially counter these NIK-mediated defense responses to geminivirus infection by inhibiting autophosphorylation of NIK and receptor kinase activation. According to this model, NSP binding to the kinase domain of NIK would prevent activation of a NIK-mediated signaling pathway, creating an intracellular environment that is more favorable to virus proliferation and spread.

Recently, the geminivirus TraP (transactivator protein) has been shown to interact with and inactivate adenosine kinase (ADK) and SNF1 kinases [Hao et al. 2003; Wang et al. 2003]. Both SNF1 and ADK have been
proposed to be components of innate antiviral defenses, and inactivation of these kinases by TraP appears to be a countereffensive measure. The finding here that NSP also targets and inhibits transmembrane kinases indicates that geminiviruses may effectively adopt a broad-spectrum cellular kinase inhibition-based strategy for countering antiviral defenses. This raises the question as to whether these viral proteins act synergistically to suppress integrated defense pathways or adaptively to inactivate multiple antiviral defenses. The discovery that NIKs mediate a defense response to geminivirus infection associated with the consequent loss-of-function phenotypes makes the NIK-integrated signaling pathway amenable to genetic characterization. This will complement biochemical approaches to the isolation of interactive proteins and downstream components. These studies also suggest that the NSP–NIK interaction may be a molecular target for the development of a broad-spectrum resistance strategy against geminivirus infection.

Materials and methods

Plasmid construction

The GmNIK and LeNIK cDNA homologs from Arabidopsis, NIK1 (U19571), NIK2 (U21612), NIK3 (U10870), and AtSERK1 (U13033) were obtained from Arabidopsis Biological Resource Center. To create plasmids for the two-hybrid analysis, we amplified the NIK1 C-terminal kinase domain (KD), encoding amino acids 298–638 and its truncated versions [ΔKD298–502, ΔKD422–502, ΔKD502–638] by PCR from U19571 cDNA and cloned them into EcoRI and SstI sites of pEXAD502 (Invitrogen Life Technologies, Inc.), as GAL4 activation domain fusions. Likewise, intact [KD] and truncated versions (ΔKD) of NIK2 ([KDNIK2345–538, ΔKDNIK2415–635]), NIK3 ([KDNIK3257–632, ΔKDNIK3325–502, ΔKDNIK3408–625]), and AtSERK1 ([KD1SERK1408–625]) kinase domains as well as an active (KDBRI1-M) kinase domain of BRI1 were fused in-frame to the GAL4 activation domain in pEXAD502. The NSP coding regions were amplified by PCR from TGMV DNA-B (Fontes et al. 1994b), TCrLYV DNA-B (Galvão et al. 2003), or CaLCuV-B (Hill et al. 1998) with Pfu DNA polymerase (Stratagene) and cloned into SacI and SstI sites of pBDLeu (Invitrogen Life Technologies, Inc.). Purification of His-tagged and GST-fusion proteins

All recombinant GST-fused proteins and His-tagged proteins were prepared using the expression vectors pDEST15 (GST fusions) and pDEST17 (6× His fusions) (Invitrogen, Life technologies, Inc.). Plasmids containing different fusions were transferred into Escherichia coli strain BL21 and the synthesis of the recombinant proteins was induced by 0.5 mM isopropyl-β-D-thiogalactopyranoside for 12 h at 20°C. The GST fusions and His-tagged proteins were affinity-purified using GST-Sepharose beads (Pharmacia) and Ni²⁺-agarose (Qiagen), respectively, according to the manufacturer’s instructions.

In vitro protein binding assay

To express intact and truncated versions of NIK in vitro, we used 1 μg of recombinant plasmid containing the appropriate insert in pDEST14 (as described earlier) in an in vitro transcription and translation system supplemented with [35S]-methionine and T7 RNA polymerase (Promega). The 35S-labeled proteins were incubated for 1 h at 4°C with 50 μL of glutathione-Sepharose beads to which purified GST or GST-NSP had been adsorbed. The beads were pelleted by centrifugation and washed five times with 1 mL of 50 mM Tris·HCl (pH 7.5), 120 mM NaCl, and 0.1% Nonidet P-40. Bound proteins were resolved by SDS/PAGE on an 8%–20% polyacrylamide gel and visualized by fluorography.

Fluorescence microscopy

Fluorescence deconvolution microscopy was conducted on a DeltaVision deconvolution microscope system (Applied Precision) consisting of an Olympus IX70 inverted microscope. Specific filter was used for GFP [490 ± 10 excitation, 510 ± 5 emission] and NIK–GFP patterns were observed in epidermal cells of Arabidopsis leaves bombarded with pK7F–NIK1, pK7F–NIK2, and pK7F–NIK3. For confocal laser scanning microscopy, stably transformed cauli samples were directly examined with a plan Neofluar 40× objective (NA 0.75, Zeiss Corp.) and a LSM 510 microscope system (Carl Zeiss) equipped with an argon-krypton laser as excitation source (excitation of GFP molecules at 488 nm). GFP emission was detected by using a 505–530-nm filter.
filter. Images were assembled with Adobe Photoshop (Adobe Systems).

**Protein kinase assay**

Purified GST-KDNIK1, GST-KDNIK2, or GST-KDNIK3 fusion proteins were incubated alone or with GST-ΔKDNIK1, GST-ΔKDNIK3, or GST-ΔKDNIK2 for 30 min at 25°C in 30 µl of kinase buffer containing 18 mM HEPES [pH 7.4], 10 mM MgCl₂, 10 mM MnSO₄, 1 mM DTT, 10 µM ATP, and 5 µCi [γ-³²P]ATP. Phosphoproteins were resolved by SDS-PAGE. The gel was stained with Coomassie brilliant blue to verify protein loading, dried, and subjected to autoradiography. Incorporation radioactivity in protein bands was quantified by phosphoimaging.

**RT-PCR**

Total RNA from seedlings, leaves, flowers, and roots was extracted using an RNACeasy kit (Qiagen). First-strand cDNA was synthesized from 2–5 µg of total RNA using the SuperScript III Kit (Invitrogen Life Technologies, Inc.) according to the manufacturer’s instructions. PCR assays were performed with NIK1, NIK2, and NIK3-specific primers as described (Cascardo et al. 2000). PCR was also carried out with actin gene-specific primers to assess the quantity and quality of the cDNA.

**CalLCuV inoculation and analysis of infected plants**

*Arabidopsis thaliana* plants at the seven-leaf stage were inoculated with plasmids containing partial tandem repeats of CalLCuV DNA-A and DNA-B by biolistic delivery as described (Schaffer et al. 1995). Total nucleic acid was extracted from systemically infected leaves and viral DNA was detected by PCR with DNA-A or DNA-B begomovirus-specific primers (Rojas et al. 1993) and/or by DNA gel blot analysis (Fontes et al. 1994b).

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The geminivirus nuclear shuttle protein is a virulence factor that suppresses transmembrane receptor kinase activity

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