pSITE Vectors for Stable Integration or Transient Expression of Autofluorescent Protein Fusions in Plants: Probing Nicotiana benthamiana–Virus Interactions

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Plant functional proteomics research is increasingly dependent upon vectors that facilitate high-throughput gene cloning and expression of fusions to autofluorescent proteins. Here, we describe the pSITE family of plasmids, a new set of Agrobacterium binary vectors, suitable for the stable integration or transient expression of various autofluorescent protein fusions in plant cells. The pSITE vectors permit single-step Gateway-mediated recombination cloning for construction of binary vectors that can be used directly in transient expression studies or for the selection of transgenic plants on media containing kanamycin. These vectors can be used to express native proteins or fusions to monomeric red fluorescent protein or the enhanced green fluorescent protein and its cyan and yellow-shifted spectral variants. We have validated the vectors for use in transient expression assays and for the generation of transgenic plants. Additionally, we have generated markers for fluorescent highlighting of actin filaments, chromatin, endoplasmic reticulum, and nucleoli. Finally, we show that pSITE vectors can be used for targeted gene expression in virus-infected cells, which should facilitate high-throughput characterization of protein dynamics in host–virus interactions.

Additional keywords: agroinfiltration, confocal microscopy, nucleolus, SYNV, transformation.

Nicotiana benthamiana Domin (Goodspeed 1954) has emerged as a critically important model host plant for a broad diversity of plant pathogens (Badel et al. 2006; Barajas et al. 2006; Chague et al. 2006; Goodin et al. 2005; Jimenez et al. 2006; Kanneganti et al. 2006; Shung et al. 2006; Sreedharan et al. 2006; Vijaya Palani et al. 2006; Xiao et al. 2003). Additionally, due to its tractability for cell biology and biochemical research, N. benthamiana often is used to complement studies initiated with Arabidopsis thaliana, the long-standing paradigm for plant molecular genetics (Deng et al. 2006; Latz et al. 2006; Moeder et al. 2007; Rahier et al. 2006; Zhao et al. 2006), or other plant systems (Wang and Kumar 2006). Moreover, N. benthamiana holds a unique position in plant–microbe interactions research because it has become the predominant experimental host for plant viruses, many of which do not infect A. thaliana (Christie and Crawford 1978; Coutts and Buck 1985; Quacquarelli 1975). Therefore, research tools and resources to support functional genomics research in N. benthamiana are essential to advance our understanding of plant–virus interactions. To this end, microarray-based transcriptional profiling to determine changes in N. benthamiana gene expression in response to virus infection have been conducted (Senthil et al. 2005; Whitham et al. 2006). However, integrated plant functional genomics investigations to establish the relationship between changes in the level of host gene expression and the subcellular localization of the encoded proteins in response to pathogen infection requires facile vector systems that permit i) high-throughput construction of recombinant expression vectors, ii) protein expression in either transient assays or transgenic plants without the need for subcloning into different vectors, iii) the ability to efficiently deliver proteins and their interacting targets or substrates to the same cell, iv) expression of proteins in pathogen-infected cells, and v) the ability to monitor membrane or protein dynamics in a large number of cells so as to permit rigorous statistical analyses.

Here, we describe the construction of a novel set of binary vectors that are highly amenable to a wide variety of functional genomics projects in N. benthamiana. These “pSITE” vectors, which can be used for stable integration or transient expression of autofluorescent protein (AFP) fusions in plant cells, are derivatives of the exceptional pSAT series described recently (Tzﬁra et al. 2005), for which modiﬁcations are being constructed to suit particular applications in plant biology (Chung et al. 2005; Citovsky et al. 2006). Our rationale for undertaking this project was motivated by the need to eliminate the two-step cloning procedure required to mobilize genes-of-interest first into pSAT vectors and then into RCS1 or RCS2 binary vectors (Goderis et al. 2002; Tzﬁra et al. 2005). Therefore, to simplify the pSAT system, we replaced the multiple cloning site in the pSAT-6 AFP cassettes with the destination “DEST” fragment required for Gateway LR-Clonase-mediated recombination cloning (Hartley et al. 2000; Walhout et al. 2000). Briefly, this synthetic construct contains the F plasmid-derived ccdB gene, the expression of which is toxic in most lab strains of Escherichia coli. In vitro sequence-specific recombination reactions, directed by flanking “att” sites, result in the
replaced the *ccdB* gene with any gene-of-interest (Hartley et al. 2000; Walhout et al. 2000). Subsequent transformation of *E. coli* results in a strong negative selection against *ccdB*-containing plasmids, resulting in highly efficient recovery of recombinant clones. This conversion of the pSAT vectors to be compatible with recombination-based expression systems should promote high-throughput protein localization studies. Ultimately, the pSITE system provides a highly desirable system that can support both transient and stable expression studies without the need for subcloning into different vectors.

Using genes encoded by *Sonchus yellow net virus* (SYNV), genus *Nucleorhabdovirus*, and a variety of markers for cellular loci, we validated the pSITE vectors in terms of the five criteria for vectors described above. First, the pSITE vectors have been constructed to permit Gateway-compatible recombination-mediated cloning to produce constructs for the expression of amino-(N-series vectors) or carboxy-terminal (C-series vectors) fusions to four different AFPs. Gateway technology, which is amenable to automation and high-throughput expression studies (Gong et al. 2004), has been incorporated into several plant expression systems (Chen et al. 2006; Chung et al. 2005; Curtis and Grossniklaus 2003; Goderis et al. 2002). Moreover, there exists a plethora of Gateway-compatible vectors for protein expression in bacteria, yeast, or insect cells that can be employed to support plant biology research. Second, the pSITE vectors carry a *nopaline phosphotransferase* (*nptII*) gene for the selection of transgenic plant tissue on media containing kanamycin. Third, using a synthetic construct composed of a green fluorescent protein (GFP) and red fluorescent protein (RFP) fusion containing a *Tobacco vein mottling virus* (*TVMV*) protease cleavage site between the two AFPs, we evaluated the efficiency of targeting a protease and its substrate to the same cell. Fourth, we provide high-resolution micrographs of localization patterns for a variety of proteins expressed transiently in mock-inoculated or SYNV-infected plants. Additionally, we have developed transgenic plants expressing RFP fused to the N. *benthamiana* histone 2B (RFP:NbH2B) or Importin-α1 proteins (NbImp-α1:RFP). Fifth, we demonstrate that infection by SYNV has a statistically significant effect on the localization of nuclear marker proteins.

Taken together, our data suggest that the pSITE vectors are a significant and much needed addition to the pSAT family of vectors (Chung et al. 2005; Citovsky et al. 2006; Tzfira et al. 2005) and are particularly well suited for dissecting the cell biology of plant–pathogen interactions.

**RESULTS**

**Twelve pSITE vectors for plant biology research.**

We have constructed 12 pSITE vectors for use in a wide variety of research applications (Table 1; Fig. 1). In addition to AFP-containing plasmids, two derivatives of pSAT-MCS were converted to Gateway compatibility to permit expression of native proteins. These Gateway-adapted cassettes were then mobilized into the pRCS2-ocs-nptII (Chung et al. 2005) (Gen-Bank accession number DQ005456) binary vector containing an *nptII* gene under the control of *octopine synthase* promoter and terminator for the selection of transgenic plant material on media containing kanamycin.

Available AFPs in the pSITE vectors include the cyan (CFP), green, and yellow-green (YFP) spectral variants of the GFP, as well as the monomeric RFP. These CFP, GFP, YFP, and RFP genes are found in pSITE vectors 1, 2, 3, and 4, respectively (Table 1; Fig. 1). Due to potential problems that can arise when using tetrameric AFPs, (Goodin et al. 2002), we omitted DsRed2 in the pSITE vectors, though it was used in development of the pSAT series (Tzfira et al. 2005). The AFP cassettes in the pSITE vectors are positioned relative to the duplicated 35S promoter so as to produce amino-terminal or carboxy-terminal fusions to AFPs. Finally, to ensure translation initiation at the start codon of the gene-of-interest, we removed the *Ncol* sites in the pSITE-0A and pSITE-NA series vectors because they contained ATG codons in a good translational context.

**Validation of the pSITE vectors using proteins encoded by SYNV.**

The pSITE vectors should perform in a fashion identical to that of pSAT derivatives reported earlier (Tzfira et al. 2005). However, we were interested in comparing results obtained with these new vectors with data reported using the pGD vectors, from which expression is controlled by a single 35S promoter (Goodin et al. 2002). We were particularly interested in knowing whether the localization patterns of SYNV-encoded proteins were significantly different than previously reported (Goodin et al. 2001, 2002). Therefore, we expressed the SYNV nucleocapsid (N) (Zuidema et al. 1987), matrix (M) (Hillman et al. 1990), and phosphoprotein (P) (Heaton et al. 1987) proteins as fusions to RFP using pSITE-4CA in leaves of mGFP5-endoplasmic reticulum (ER) plants (Fig. 2, A through X3). Prior to imaging, the leaves were infiltrated with 4’,6-diamidino-2-phenylindole (DAPI)-containing buffer to permit staining of nuclei. In contrast to the results previously reported using DsRed2 (Goodin et al. 2002), RFP-P fusions did not aggregate in the nucleus (Fig. 2, I through P). More importantly, RFP-M fusions localized exclusively to nuclei, whereas the DsRed2 fusion to this protein was highly prone to aggregation (Fig. 2, Q through X) (Goodin et al. 2002). The RFP-N protein, like M, was also found only in the nucleus, where it excluded DAPI at its major site of accumulation (Fig. 2, Y through X3). Further, we note that, during the course of

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<th>Fluorescent taga</th>
<th>DEST-terminal</th>
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<tr>
<td>pSITE-0A</td>
<td>Nil</td>
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<td>Yes</td>
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<tr>
<td>pSITE-0B</td>
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a DEST = destination fragment; ECFP, EGFP, EYFP, and RFP = enhanced cyan, green, yellow, and red fluorescent proteins, respectively.

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validating the pSITE vectors, 39 different constructs were generated. Of these, 35 (90%) were detectable by fluorescence microscopy. Of the 10 proteins expressed as amino-terminal fusions (N series vectors), 7 (70%) were detectable by fluorescence microscopy. In contrast, 28 of 29 (96.5%) proteins expressed as carboxy-terminal fusions were detectable.

**Fluorescent markers for cellular loci.**

Dyes such as DAPI can be infiltrated into leaf tissue to mark particular cellular loci such as the nucleus. In addition, fluorescent proteins with known sites of localization are useful aids in cell biology research (Cutler et al. 2000; Goodin et al. 2002; Panavas et al. 2003; Saint-Jore et al. 2002). Therefore, we are developing RFP- and GFP-tagged markers for all major plant cellular loci in pSITE vectors, and the cognate transgenic *N. benthamiana* plants, in order to provide a standard set of reagents in a common vector background. The first five of these markers and two derivative RFP-tagged lines are shown in Figures 3 and 4.

Using overlap-polymerase chain reaction (PCR), we replaced the GFP in mGFP5-ER (endoplasmic reticulum) with RFP so as to produce a red fluorescent marker for the ER, hereafter referred to as RFP-ER. The *A. thaliana* basic chitinase signal peptide and Histidine-Aspartate-Glutamate-Leucine (HDEL)-ER retention signal present in mGFP5-ER (Haseloff et al. 1997) are conserved in RFP-ER. Transient expression of this marker in mGFP5-ER plants showed that both proteins accumulated in a similar manner in SYNV-induced intranuclear membranes (Fig. 3B through D) and in the ER (Fig. 3E through G). Fluorescent recovery after photobleaching (FRAP) experiments established that the RFP-ER marker was more sensitive to photobleaching than GFP but otherwise showed statistically identical FRAP kinetics in short-time-course experiments (Fig. 3A).

Fluorescence in transgenic plants expressing RFP fused to the *N. benthamiana* histone 2B (RFP:NbH2B) was detected only in the nucleus (Fig. 3K through N). Consistent with chromatin targeting, RFP:NbH2B could be used to track all stages of cell division, including anaphase (Fig. 3K through N). RFP:NbH2B transgenic plants were phenotypically indistinguishable from wild-type *N. benthamiana* plants, suggesting no adverse effect on these plants due to histone overexpression (data not shown). Likewise, expression of GFP:Talin in *N. benthamiana* resulted in labeling of actin filaments in a manner consistent with published reports (Kost et al. 1998; McCann and Craig 1997). Similar results were obtained with RFP:Talin (data not shown).

In contrast to the predicted results for RFP:ER, GFP:Talin, and RFP:NbH2B, we recovered transgenic plants in which RFP fused to *N. benthamiana* Importin-α1 (NbImp-α1;RFP) accumulated preferentially in guard cells (Fig. 3H) or in equal levels in all nuclei (Fig. 3I). Mature leaves of the latter plants were curled relative to wild-type plants (Fig. 4A through D), whereas plants with expression only in guard cells were extremely distorted (data not shown).

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**Fig. 1.** Schematic representations of pSITE vectors. A, All modified pSAT6 cassettes were cloned into the pRCS2-ocs-nptII binary vector at the PI-Psp site. The ability to select transgenic plant cells is conferred by the nptII gene, the expression of which is controlled by the ocs promoter (P-OCS) and terminator (T-OCS). B, “C-series” pSITE vectors for Gateway recombination-mediated construction of binary vectors for expression of proteins-of-interest fused to the carboxy termini of autofluorescent proteins (AFPs). C, “N-series” pSITE vectors for Gateway recombination-mediated construction of binary vectors for expression of proteins-of-interest fused to the amino termini of AFPs. D, “0-series” pSITE vectors for Gateway recombination-mediated construction of binary vectors for expression of native proteins. Protein expression is controlled by a duplicated *Cauliflower mosaic virus* (CaMV) 35S promoter (2X35S) and a *Tobacco etch virus* translational leader (TL). All vectors employ the CaMV35S transcriptional terminator (TER). NcoI*: this restriction site was deleted to create the pSITE-NB and pSITE-0B vectors. The “DEST” fragment, flanked by LR clonase-mediated recombination sites (attR1 and attR2), contains the *ccdB* gene derived from the F plasmid. The roles of the *ccdB* gene and *att* sites in generating recombinant vectors are defined in the text.
Immunodetection of AFP fusions expressed from pSITE vectors.

Most protein fusions were easily detectable by Western immunoblotting when transiently expressed in leaf tissues following agroinfiltration (Fig. 5A and B). We could routinely detect fusions up to 97 kDa (RFP:SYNV-G) by simple colorimetric detection without the need for more sensitive detection methods. In order to assess the efficiency for co-delivery of two fusions into the same cell, we constructed a GFP:RFP fusion that contained a TVMV protease cleavage site between the two AFPs (Figs. 5D, lane 1, and 4E, lane 1). Coexpression with the TVMV protease, either as a native protein (pSITE-0B-TVMVprotease),...
amino-terminal fusion to CFP (pSITE-1NB-TVMVprotease), or carboxy-terminal fusion to CFP (pSITE-1CA-TVMVprotease), resulted in complete digestion of the GFP:RFP reporter. Processing of the reporter occurred even though the TVMV-protease:CFP fusion was not as readily detectable as the amino fusion. Both CFP protein fusions were, however, easily detectable by fluorescence microscopy (data not shown). In general, expression levels were lower when the same fusion was expressed in the context of transgenic plants compared with levels produced via transient expression (Fig. 5F and G). This determination was based on data from 10 and 2 independent transgenic lines of RFP:NbH2B and NbImp-α1:RFP, respectively. Interestingly, we were able to recover transgenic plants expressing NbImp-α1:RFP (expressed from pSITE-4NA) but not RFP:NbImp-α2 (expressed from pSITE-4CA), despite the fact that both fusions expressed to high levels in transient assays (data not shown).

**Nucleolar dynamics in SYNV-infected cells.**

With the advent of the pSITE and related vectors, high-throughput characterization of the changes in the subcellular localization of host protein in response to virus infection is now possible. Prior to engaging in elucidation of the differential localization of the *N. benthamiana* proteome in mock-inoculated versus virus-infected cells, we conducted pilot studies to map viral proteins in these two cell types (Goodin et al. 2007), as well to determine the effect of SYNV infection on the localization of host proteins (this study). In addition to results to be presented elsewhere, these experiments revealed changes in the localization pattern of RFP:AtFib1 in virus-infected nuclei (Fig. 6). Rather than aberrant localization, we discovered that there was an increase in the percentage of nuclei with multiple nucleoli in SYNV-infected cells (Fig. 6). Briefly, three patterns of nucleolar expression were established. Category 1 included those nuclei in which only one nucleolus could be observed. Category 2 included those nuclei in which accumulation of RFP:AtFib1 could be observed at two loci, which occurred mostly in an “earth and moon” relative size relationship (Fig. 6A through C). This was the predominant category in mock-inoculated cells. Category 3 included those nuclei that contained three or more nucleoli. Patterns of nucleolar distribution were determined in 63 and 69 independent sets of serial optical sections (“z-stacks”) of nuclei in

![Fig. 4.](image)

A, Wild-type *Nicotiana benthamiana* plant. B, Transgenic *N. benthamiana* plant expressing NbImp-α1:RFP. C, Fully expanded leaf of *N. benthamiana* plant shown in A. D, Fully expanded leaf of *N. benthamiana* plant shown in B.
mock-inoculated and virus-infected nuclei (Goodin et al. 2005), respectively. Quantitative analyses of nucleoli in mock-inoculated and virus-infected nuclei revealed that there was an increase in the percentage of nucleoli with multiple (more than three) nucleoli per nucleus in virus-infected cells. There was no significant change in the percentage of nucleoli in category 1 (6 and 7% in mock-inoculated and SYNV-infected nuclei, respectively). Category 2 showed a reduction from 76 to 49% in mock-inoculated and SYNV-infected nuclei, respectively. Finally, category 3 increased from 18 to 49% in mock-inoculated and SYNV-infected nuclei, respectively. (Fig. 5D through G). A χ² analysis showed that the differences in localization patterns were statistically significant (P value = 0.01). Interestingly, we detected an increase in only the number of large nucleolar bodies, whereas the number of small nucleolar bodies remained at one per nucleolus.

DISCUSSION

Plant functional genomics research is increasingly dependant upon vector systems that can be used in facile high-throughput protein expression studies. Additionally, *N. benthamiana* increasingly is being used to dissect the cell biology of host–pathogen interactions and to validate results obtained in *A. thaliana*. Therefore, we have modified the pSAT system to create the pSITE vectors which employ single-step Gateway cloning to produce recombinant binary vectors for the expression of native proteins, or their autofluorescent derivatives, in plant cells (Tzfira et al. 2005). These vectors can be used in transient assays or for the generation of transgenic plants. Unlike other reports of novel binary vectors for plant biology research (Chen et al. 2006; Curtis and Grossniklaus 2003; Earley et al. 2006), we have validated the pSITE vectors in the context of virus-infected cells, thereby demonstrating their suitability for use in probing plant–pathogen interactions. Although this study addressed the changes in the localization of plant proteins in response to virus infection, a parallel study demonstrating the differential localization of SYNV-encoded proteins in virus-infected versus mock-inoculated cells has been reported elsewhere (Goodin et al. 2007).

The pSITE vectors excelled in their use for standard protein localization studies, as demonstrated with RFP fusions of SYNV-encoded proteins. Several differences of note were observed when comparing these results with similar data obtained with pGD vectors (Goodin et al. 2002). First, using a monomeric form of DsRed proved superior to the use of DsRed2 because previously encountered problems due to tetramerization or aggregation were eliminated. Additionally, we found that a greater amount of SYNV-M and -N accumulated in the nucleus when these proteins were expressed from pSITE vectors. Perhaps this is due to the use of a duplicated 35S promoter and translational enhancer in the pSITE vectors compared with a single 35S promoter in the pGD vectors. As for the three aforementioned SYNV proteins, RFP-ER, RFP-NbH2B, NbImp-a1:RFP, and GFP-Talin, all were targeted to their predicted cellular loci. In no case did we observe obvious mislo-
calization of protein fusions. Therefore, we conclude that the pSITE vectors should be extremely useful for general localization studies. Moreover, the compatibility of pSITE vectors with Gateway cloning technology suggests that these vectors should be highly amenable to high-throughput projects (Li et al. 2006; Simpson and Pepperkock 2003; Tian et al. 2004; Wiemann et al. 2004). As noted above, 90% of the constructs made in AFP-containing pSITE vectors yielded protein fusions that were detectable by fluorescence microscopy. Approximately 70% of amino-terminal fusions (N-series vectors) were detectable by fluorescence microscopy, whereas 96.5% of carboxy-terminal fusions (C-series vectors) were detected. Despite this apparent greater utility of the C-series vectors, we advise users of these vectors to test expression of genes-of-interest from both the N- and C-series of vectors. Such tests are simple to conduct, given the presence of stop codons in all three reading frames after the 3′ lambda phage “att” recombination site in the C-series vectors. Therefore, the same pDONR plasmids, in which genes-of-interest lacking stop codons are cloned, can be used with the N- and C-series pSITE vectors.

In general, detection of AFP fusions, either in transgenic plants or in transient expression assays, was straightforward. However, stability of the fusions appeared to be highly variable, particularly in the case of the SYNV-N protein; however, this most likely is a consequence of the fusion partner and not the AFPs in the pSITE vectors.

Importantly, our experiments, using the TVMV protease and a fluorescent reporter substrate, strongly indicate that two proteins can be delivered to the same cell with high efficiency. The GFP-RFP reporter was digested to completion, such that the fusion protein was undetectable in co-infiltrated tissue, demonstrating that the TVMV protease was functional whether it was expressed as a native protein or with CFP fused to its carboxy- or amino-terminus. In a more extreme test case, the expression of NbH2B (approximately 18 kDa) as a 35-kDa RFP fusion provided a marker for chromatin dynamics entirely consistent with that expected for a histone marker (Boisnard-Lorig et al. 2001). Although all proteins or enzymes may not tolerate such additions, these data, along with a wealth of published reports, support the contention that many proteins are functional as AFP fusions.

We note that, for confocal microscopy studies requiring a nuclear marker, the transgenic RFP:NbH2B lines can be imaged using the 543-nm laser line of the common He-Ne lasers. Therefore, this should provide an exceptional alternative to the use of propidium iodide which, being highly cell impermeant, requires incubation of plant tissue in harsh buffers when it is used as a nuclear counterstain (Kumar et al. 2006). Additionally, the use of the cell-permeant DNA-selective dye DAPI requires a UV or near-UV laser which, if unavailable, forces the use of propidium iodide. Thus, the RFP:NbH2B lines may circumvent two major technical limitations that often reduce the quality of micrographs of nuclear localized proteins in plant cells.

Following rigorous technical validation of the pSITE vectors, we employed them to examine the effect of virus infection on the localization of host proteins. These initial experiments revealed a statistically significant increase in the number of nucleoli in SYNV-infected nuclei. Several examples of changes in nucleolar morphology and localization of nucleolar proteins in response to viral infection or interaction with viral proteins have been reported (Hiscox 2007). However, presently it is unclear whether the SYNV-induced increase in nucleolar copy number is a general response to stress or is required for particular aspects of the SYNV life cycle, such as recruitment of ribosomal subunits (Walsh and Mohr 2006), inhibition of specific functions associated with viral proteins (Canto et al. 2006; Uhrig et al. 2004), or other as yet undefined associations. Further experiments utilizing transgenic plants expressing RFP:AtFib1 and a large set of AFP-tagged nucleolar proteins (Pendle et al. 2005) are in progress to provide further insight into the effect of SYNV on nucleolar protein localization and function (data not shown).

In summary, the pSITE vectors described here meet all necessary criteria for support of plant functional genomics projects, namely i) the ability to generate recombinant expression vectors in high-throughput, which we have addressed by constructing Gateway-compatible vectors; ii) the ability to express proteins in either transient assays or in transgenic plants without the need for subcloning into different vectors, which we have demonstrated for a number of proteins; iii) the ability to efficiently deliver proteins and their interacting targets or substrates to the same cell, which we have addressed using TVMV
protease and a fluorescent substrate; iv) facile expression of protein fusions in mock-inoculated and virus-infected cells, which we have demonstrated using SYNV-encoded proteins as well a number of markers for cellular loci; and v) the ability to monitor membrane or protein dynamics in a large number of cells so as to produce statistically significant results, which we have addressed primarily with FRAP analyses of RFP-ER and the localization of RFP:AtFib1 in mock-inoculated and SYNV-infected leaves.

**MATERIALS AND METHODS**

Plant material and growth conditions.

*N. benthamiana* “16c” plants (referred herein as mGFP5-ER plants) (Brigneti et al. 1998; Haseloff et al. 1997; Ruiz et al. 1998) constitutively express GFP targeted to the ER under the control of the *Cauliflower mosaic virus* 35S promoter. These plants, and the nontransgenic parental line, were grown in a greenhouse under ambient conditions. SYNV inoculations were conducted as described by Goodin and associates (2005).

**Construction of the pSITE vectors.**

Gateway-compatible binary vectors for expression of proteins of interest fused to the carboxy- (C-series vectors) or amino-termini (N-series vectors) of AFPs were derived from the pSAT6 vectors (Tzfira et al. 2005). Construction of the C-series vectors first required digestion of pSAT6 vectors containing the enhanced green fluorescent protein (EGFP) gene or its cyan (ECFP) or yellow-shifted (EYFP) spectral variants in pSAT6-DEST-EGFP-C1 (AY818372) and pSAT6-EGFP-C1 (AY818380), and pSAT6-ECFP-C1 (AY818374) and pSAT6-RFP-C1 (DQ005474) with BglII and NdeI. The 1,865-bp fragment released from each vector was ligated to a 4,387-bp BglII and NdeI fragment from pSAT6-DEST-ECFP-C1 to create the intermediate vectors pSAT6-DEST-EGFP-C1 (EF212303), pSAT6-DEST-ECFP-C1 (EF212304), and pSAT6-DEST-RFP-C1 (EF212305), respectively. Next, binary vector RCS2 with nptII plant selection marker (DQ005456) and the intermediate vectors pSAT6-DEST-EGFP-C1 (EF212303), pSAT6-DEST-ECFP-C1 (EF212304), pSAT6-DEST-RFP-C1 (EF212305), and pSAT6-EGFP-C1 (AY818372) were digested with PI-PspI. The 3.5-kb fragments from the intermediate vectors were ligated to the binary vector backbone RCS2 (DQ005456) and pSITE-3CA (EF212293), pSITE-1CA (EF212302), pSITE-4CA (EF212292), and pSITE-2CA (EF212294).

Similarly, construction of the N-series pSITE vectors required digestion of pSAT6-DEST-EGFP-N1 (AY818370), pSAT6-MCS (AY818383), pSAT6A-MCS (EF212312), pSAT6-RFP-N1 (DQ005475), pSAT6A-EGFP-N1 (DQ005470), pSAT6A-ECFP-N1 (DQ005469), and pSAT6A-RFP-N1 (DQ005472) with BglII and Apol. The 1,723-bp fragment released from pSAT6-DEST-EGFP-N1 (AY818370) was ligated to each aforementioned vector, thereby replacing the MCS with the Gateway destination fragment sequence to make pSAT6-DEST (EF212311), pSAT6A-DEST (EF212310), pSAT6-DEST-RFP-N1 (EF212309), pSAT6A-DEST-EGFP-N1 (EF212308), pSAT6A-DEST-ECFP-N1 (EF212307), and pSAT6A-DEST-RFP-N1 (EF212306) intermediate vectors, respectively. Finally, intermediate vectors were digested with PI-PspI and a 3.5-kb fragment release from each of them; 2.8 kb from pSAT6-DEST (EF212311) and pSAT6A-DEST (EF212310) was ligated to binary vector backbone RCS2 (DQ005456) to make pSITE-0A (EF212300), pSITE-0B (EF212301), pSITE-4NA (EF212293), pSITE-2NA (EF212299), pSITE-2NB (EF212296), pSITE-1NB (EF212297), and pSITE-4NB (EF212298).

The presence of the NcoI site in pSITE-NA vectors results in a 17-amino-acid extension at the amino termini of fusion proteins. In order to permit translational initiation from the start codon of proteins-of-interest, the NcoI site in the “A” versions of the pSITE-N and pSITE-0 vectors was removed by sequential digestion with NcoI and mung bean nuclease followed by ligation with T4DNA ligase (Table 1). Vectors lacking an NcoI site carry the “B” designation (Table 1). Thus, using the same gene-of-interest, protein fusions expressed from a pSITE-NA vector are approximately 2 kDa larger than those expressed from the cognate pSITE-NB vector. Similarly, using the same gene-of-interest, protein fusions expressed from a pSITE-NB vector are approximately 2 kDa smaller than those expressed from the cognate pSITE-CA vector, because the NcoI site is retained in these latter vectors (Fig. 1).

**Gateway cloning of genes into pSITE vectors.**

A two-step procedure was used to incorporate recombinant *att* sites into PCR products in order to make them suitable for use as substrates for Gateway BP clonase-mediated (Invitrogen Corp, Carlsbad, CA, U.S.A.) recombination into pDONR 221 (Invitrogen). Briefly, genes-of-interest were amplified using gene-specific forward and reverse primers that each contained 5' extensions corresponding to half of the *att* site. The PCR products were then reamplified using “universal” primers that incorporated the remainder of the *att* sites. PCR amplification was conducted with Phusion High Fidelity DNA Polymerase (Finnzymes, Espoo, Finland) according to manufacturer’s recommendations in a Bio-Rad (Hercules, CA, U.S.A.) thermal cycler.

The PCR products containing complete *att* sites were purified and then cloned into pDONR221 vector by overnight incubation in the presence of Gateway BP clonase, per the manufacturer’s recommendation, at 25°C. In this manner, “entry clones” for SYNV-N (L32603), SYNV-P (AY971951), SYNV-sc4 (L32604), SYNV-M (M35689), SYNV-G (L32603) *N. benthamiana* histone 2B (EF189156), *N. benthamiana* Importin-α homologues 1 and 2 (EF137253 and EF137254) (Kanegami et al. 2007), A. *thaliana* Fibriilin 1 (AtFib1 and NM_124626.3) (Barneche et al. 2000), TMV protease (NC_001768), and RFP with N-terminal fusion of TMV protease cleavage site were created. Next, the plasmid DNA from the positive entry clones was introduced into a pSITE vector of choice by overnight incubation of Gateway LR clonase recombinant reaction (as per the manufacturer’s recommendation) at 25°C. Finally, each recombinant pSITE vector was transformed into *Agrobacterium tumefaciens* LBA4404.

**Transient expression of proteins in leaf epidermal cells.**

Suspensions of *A. tumefaciens* were infiltrated into leaves of *N. benthamiana* as previously described (Goodin et al. 2002; Tsai et al. 2005). In order to express proteins in SYNV-infected cells, symptomatic leaves of plants were infiltrated at the peak of symptom expression, typically 10 to 14 days post inoculation. Following a 48-h incubation of infiltrated plants under constant illumination at 25°C, water-mounted sections of leaf tissue were examined by confocal microscopy.

**Laser scanning confocal microscopy.**

All confocal microscopy was performed on an Olympus FV1000 (Olympus America Inc., Melville, NY, U.S.A.). DAPI, CFP, GFP, YFP, and RFP were excited using 405-, 440-, 488-, 515-, and 543-nm laser lines, respectively. When using multiple fluor simultaneously, images were acquired sequentially line-by-line in order to eliminate excitation and emission cross talk. The primary objective used to produce the micrographs in this study was an Olympus water immersion PLAPO60XWLSM (NA 1.0). Image acquisition was conducted at a resolution of 512-by-512 pixels and a scan rate of 10
μs/pixel. Control of the microscope, as well as image acquisition and exportation as TIFF files, was controlled by Olympus Fluoview software version 1.5. Image analysis was conducted using Photoshop 7.0 (Adobe Systems Incorporated, San Jose, CA, U.S.A.) and Canvas 8.0 (Deneba Software, Miami).

FRAP.
FRAP experiments were conducted using leaf tissue harvested 10 to 14 days post inoculation from SYNV-infected mGFP5-ER plants. Mock-inoculated plants of similar age were used as controls. FRAP experiments were performed using the Olympus FV1000 described above. Briefly, 25-mm² sections of leaf tissue were mounted on glass slides in water and covered with a glass coverslip. Imaging for FRAP experiments was conducted using a x60 objective and 488-nm laser line from a multiline argon laser set at 0.3% power. Regions of interest (ROIs) were photobleached for 50 ms using a 405-nm diode laser, set at full power, which was delivered via the FV1000 Simultaneous scanner. Images for FRAP analyses were acquired at a resolution of 512-by-512 pixels and a scan-rate of 2 μs/pixel, which was necessary to monitor fast protein dynamics. Two images were acquired prior to photobleaching followed by an additional seven images to monitor fluorescence recovery. Quantitative fluorescence data, in Excel format, and confocal images, in TIFF format, were exported using Olympus Fluoview software. FRAP experiments were repeated three times for each ROI, with 2 min between bleaching events in order to allow full recovery of fluorescence. Replicated fluorescence intensity data were averaged and these data were normalized across experiments. Mean and standard deviations for fluorescence intensity at each time point were calculated and plotted using Excel software (Microsoft Corporation, Redmond, WA, U.S.A.).

A. tumefaciens-mediated plant transformation.
Due to its widespread familiarity in the plant biology community, all transgenic plants developed in this study were derived from the parental line used to generate the “16c” line of N. benthamiana mGFP5-ER plants (Brigneti et al. 1998, Haseloff et al. 1997; Ruiz et al. 1998). The genetic variation among N. benthamiana accessions in use is poorly defined at present. Therefore, we have standardized all of our N. benthamiana resources by working exclusively with the 16c parent. The transformation procedure employed was an adaptation of the methods described by Horsch and associates (1985) and Kalanidis and associates (2002). Briefly, A. tumefaciens LBA4404 carrying the pSITE-4CA-N. benthamiana histone 2B (RFP:NbH2B) or pSITE-4NB-N. benthamiana Importin-α (NbImp-α1:RFP) binary vectors were grown overnight at 28°C. Surface-sterilized leaves from greenhouse-grown N. benthamiana plants were infected with the Agrobacterium culture. The explants were co-cultivated for 2 days on Murashige and Skoog (1962) media (MS) supplemented with benzylaminopurine and indole-3-acetic acid at 2 and 0.5 mg/liter, respectively. Putative transgenic shoots from the leaf explants were induced on the same medium supplemented with cefotaxime (500 mg/liter) and kanamycin (150 mg/liter). Regenerated shoots were transferred to rooting media that included MS with cefotaxime (250 mg/liter) and kanamycin (50 mg/liter). After rooting, the plants were transferred to soil in pots and were kept in culture room at 25°C with a 16-h photoperiod. Later, seed were collected from selfed T0 plants.

Western immunoblotting.
Leaf disc samples were collected in triplicate from infiltrated or transgenic N. benthamiana leaves using a 1-cm cork borer. The tissue samples were ground in 200 μl of 2× sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) loading buffer (0.5 M Tris-HCl, 20% glycerol [vol/vol], 2% SDS [wt/vol], 1.4 M β-mercaptoethanol, 0.001% bromophenol blue [wt/vol], pH 6.8). Following electrophoretic separation on 12% SDS-PAGE gels, proteins were transferred onto nitrocellulose membranes, which were incubated in buffers containing polycyalonal antibodies to DsRed (1:2000 dilution; Clontech, Mountain View, CA, U.S.A.) or GFP (1:2000 dilution; Aves Labs, Tigard, OR, U.S.A.), as appropriate. Detection of N. benthamiana importin-α was performed using polyclonal antibodies raised against the recombinant A. thaliana importin-α (1:3000 dilution; Rose Biotechnology, Palo Alto, CA, U.S.A.). After incubation in secondary antibodies (1:20,000 dilution; Sigma, St. Louis) conjugated to alkaline phosphatase, the immunoblots were developed in sodium bicarbonate buffer containing the chromogenic substrate 5-bromo-4-chloro-3-indolyl phosphate and oxidant nitro blue tetrazolium chloride. We note that detection of AFP fusions was highly dependent upon the cloned “gene-of-interest.” For example, RFP:SYNV-G expression levels always were lower than RFP alone or RFP:SYNV-P. Therefore, the loading of protein samples corresponding to each construct was titrated to determine the optimal volume to produce immunoblots with clearly identifiable bands. Composite immunoblots then were produced to show that fusions of the predicted size were produced in planta. Due to an emphasis on characterizing protein localization, expression levels for each construct relative to the total amount of protein in extracts from infiltrated leaves was not determined.

Statistical analyses.
A quantitative analysis of the localization of RFP:AtFib1, a nucleolar marker protein, in virus-infected and mock-inoculated cells was conducted using χ² analysis (P = 0.01; 2 degrees of freedom) (Steel and Torrie 1980).

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LITERATURE CITED


