

# Shrinkage of Genome Size in a Plant RNA Virus upon Transfer of an Essential Viral Gene into the Host Genome

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## Abstract

Nonretroviral integrated RNA viruses (NIRVs) are genes of nonretroviral RNA viruses found in the genomes of many eukaryotic organisms. NIRVs are thought to sometimes confer virus resistance, meaning that they could impact spread of the virus in the host population. However, a NIRV that is expressed may also impact the evolution of virus populations within host organisms. Here, we experimentally addressed the evolution of a virus in a host expressing a NIRV using *Tobacco etch virus* (TEV), a plant RNA virus, and transgenic tobacco plants expressing its replicase, *NIb*. We found that a virus missing the *NIb* gene, TEV- $\Delta NIb$ , which is incapable of autonomous replication in wild-type plants, had a higher fitness than the full-length TEV in the transgenic plants. Moreover, when the full-length TEV was evolved by serial passages in transgenic plants, we observed genomic deletions within *NIb*—and in some cases the adjacent cistrons—starting from the first passage. When we passaged TEV and TEV- $\Delta NIb$  in transgenic plants, we found mutations in proteolytic sites, but these only occurred in TEV- $\Delta NIb$  lineages, suggesting the adaptation of polyprotein processing to altered *NIb* expression. These results raise the possibility that NIRV expression can indeed induce the deletion of the corresponding genes in the viral genome, resulting in the formation of viruses that are replication defective in hosts that do not express the same NIRV. Moreover, virus genome evolution was contingent upon the deletion of the viral replicase, suggesting NIRV expression could also alter patterns of virus evolution.

**Key words:** experimental evolution, genome evolution, nonretroviral integrated RNA viruses, plant virus, RNA virus, virus evolution.

## Introduction

The genome size of RNA viruses is highly variable, ranging from less than 4 kb for members of the *Narnaviridae* to more than 30 kb for some *Coronaviridae*. However, most RNA viruses have genome sizes in the 5–15-kb range (Holmes 2009). What could explain this wide range of genome sizes in RNA viruses? Small genomes probably replicate faster than large ones, while genome size limits enzymatic complexity. This limited complexity results in lower-fidelity replication enzymes, which in turn leads to higher variability (Smith and Szathmáry 1995). On the other hand, larger genomes will replicate more slowly, while being less constrained in the complexity of the proteins they code for. Therefore, larger genomes allow for evolutionary innovations such as polymerases with greater replication fidelity (Denison et al. 2011) or the modulation of host immune responses (Shackelton and Holmes 2004). However, the evolution of larger genomes and greater complexity may be

impeded by Eigen's paradox: to obtain proofreading activity, an RNA virus must already have a large genome (Eigen 1971; Smith and Szathmáry 1995). It has therefore been suggested that RNA virus genome size evolves to the upper limit imposed by mutation rate (Swetina and Schuster 1982).

If genome size of RNA viruses is limited by a tradeoff between RNA-dependent RNA polymerases (RdRp) fidelity and decreased replication rates, then examples of enlargement in genome size should be scarce for RNA viruses. There are, however, some examples of the integration of host genetic material into viral RNA genomes, leading to the largest known RNA viruses. For instance, the genome of coronaviruses contains homologous sequences of at least five cellular enzymes that are associated with RNA processing and that may be involved in modulating mutation rate (Snijder et al. 2003). Acquisition of a host's ubiquitin gene by certain strains of bovine viral diarrhoea pestivirus resulted in increased pathogenicity (Meyers et al. 1989). Plant closterovirus genomes

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encode for a homologue of cellular molecular chaperone HSP70 (Dolja et al. 2006) that is required for virion assembly (Satyanarayana et al. 2000). Finally, a particularly interesting case is the ubiquitous presence of DNA/RNA repair AlkB-like proteins in several groups of plant viruses, including closterovirus, flexivirus, tymovirus, and even one potyvirus (*Blackberry virus Y*; Susaimuthu et al. 2008), whose function has been postulated to be both repair of nucleic acid alkylation and suppression of posttranscriptional gene silencing (Aravind and Koonin 2001; Bratlie and Drabløs 2005). Similarly, the insertion of sequences into a viral genome generally leads to their rapid removal, as has been seen in the case of heterologous genes (Chapman et al. 1992; Guo et al. 1998; Chung et al. 2007; Paar et al. 2007; Zwart et al. 2014) and repetitions of endogenous noncoding sequences (Nagai et al. 2003; Gritsun and Gould 2006). Finally, most viruses have a tendency to evolve defective interfering viruses under conditions promoting cellular coinfection (Huang 1973; Roux et al. 1991), again suggesting a link between genome size and replication. Although demographic conditions may allow for the maintenance of extraneous sequences (Dolja et al. 1993; Zwart et al. 2014), overall it appears as if genome size is limited and under purifying selection, and that the incorporation or maintenance of foreign sequences comes at a cost to viruses (Sakai et al. 1999; Marks et al. 2005; Holmes 2009; Zwart et al. 2014).

On the other hand, genome size does not appear to be under the same selective pressure in eukaryotes, probably due to the smaller effective population sizes in higher organisms (Lynch et al. 2006). Although the existence of many retroviral sequences in eukaryotic genomes was already well documented (Gifford and Tristem 2003), it has been recently found that there are also many nonretroviral RNA virus genes in eukaryotic genomes. These nonretroviral integrated RNA viruses (NIRVs) have been found in mammals (Geuking et al. 2009; Belyi et al. 2010; Horie et al. 2010; Katzourakis and Gifford 2010), plants (Hong et al. 1998; Chiba et al. 2011), and fungi (Taylor and Bruenn 2009). In fungi, NIRVs are very common and have in some cases become functional genes (Koonin 2010). In one case, the entire viral RdRp was integrated and conserved, probably conferring viral resistance to the host (Taylor and Bruenn 2009). It has also been suggested that NIRVs can confer resistance (Bertsch et al. 2009) or tolerance (Flegel 2009) against the original virus to plant or animal hosts. Indeed, it has been a common practice to generate commercial transgenic plants resistant to viruses by inserting full or partial viral genes into the genome of the plants of agronomic interest (reviewed in Kundu and Mandal [2001]); these transgenic plants can be considered as artificial cases of NIRVs.

Because the reading frame is conserved in many NIRVs (Taylor and Bruenn 2009) they can in principle be expressed. In practice, this will depend on their location in the genome and the state of the host cell. Some NIRVs are, however,

clearly expressed by the host (Taylor and Bruenn 2009). Host NIRV expression has been proposed to mediate host immunity against the cognate virus (Bertsch et al. 2009; Flegel 2009; Taylor and Bruenn 2009). However, host NIRV expression may also have important implications for the evolution of the cognate virus. The absence of an effective NIRV-induced host immune response and sufficiently high NIRV expression levels could lead to complementation of host proteins, making the NIRV-expressed genes in the cognate virus functionally redundant. What would be the impact of NIRV expression on viral replication? Is there any selective advantage to using in trans the protein produced from the NIRV as a functional viral protein? How will a virus population evolve in such a situation of functional redundancy? Finally, how do such large-scale genomic perturbations affect the regulation of expression and accumulation of all other viral components? To our knowledge these questions have not been addressed in any study, while given the prevalence of NIRVs in natural host populations and biosafety issues linked to the use of transgenic plants expressing viral genes for resistance, it is a highly relevant topic.

Here, we evaluated the patterns of genomic evolution of *Tobacco etch virus* (TEV, genus *Potyvirus*, family Potyviridae) in transgenic *Nicotiana tabacum* plants expressing Nilb, the TEV RdRp (Li and Carrington 1995). This system is an experimental model for exploring what the effects of high levels of NIRV expression are on virus infection and evolution. During infection of these transgenic plants, plant-derived Nilb expression makes virus-derived Nilb-expression redundant (Li and Carrington 1995). We therefore expect that there may be relaxed selection for maintenance of the *Nilb* cistron in the viral genome, which might lead to selection of variants with genomic deletions at the *Nilb* locus, due to the replicative advantage associated with a smaller genome size. Ultimately, the expression of *Nilb* might therefore induce genome shrinkage and result in a virus population that can no longer autonomously infect wild-type plants. We first determined whether the deletion of viral *Nilb* led to an increase in viral fitness in Nilb-expressing plants. Then, we passaged a full-length wild-type virus in Nilb-expressing plants and used next-generation sequencing (NGS) to check for deletions in TEV *Nilb* locus. Finally, we preempted the complete deletion of TEV *Nilb* in Nilb-expressing plants and studied the effects of Nilb complementation in trans on genomic changes.

## Materials and Methods

### Virus Genotypes

The pTEV7DA infectious clone (Dolja et al. 1992) was used as a source for TEV. A TEV genotype was produced that lacked the full replicase gene ( $\Delta Nilb$ ) by inverse polymerase chain reaction (PCR) using *Pfu* turbo DNA polymerase (Stratagene) and primers conserving the proteolytic Nla/Nilb

( $\Delta Nlb$ -F) and  $Nlb/CP$  ( $\Delta Nlb$ -R) sites (supplementary table S1, Supplementary Material online). The resulting clone was named pTEV7DA- $\Delta Nlb$  (Tromas and Elena 2010).

### Direct Competition between TEV and TEV- $\Delta Nlb$ Genotypes

Concentrated saps of TEV and TEV- $\Delta Nlb$  were obtained by grinding 500 mg of 7 days postinoculation (dpi) infected tissue in a mortar with 500  $\mu$ l of grinding buffer. *Nicotiana tabacum* 35S::*Nlb* plants were inoculated with a 1:1 mixture of homogenates of both viruses. Twenty plants were inoculated by abrasion of the third true leaf with 30  $\mu$ l mixed homogenate, and 20 plants were taken as a mock-inoculated control. After 7, 21, and 63 dpi, total RNA was extracted from the youngest leaves of five plants. The region flanking the *Nlb* cistron was reverse transcribed using *Moloney Murine leukemia virus* (M-MLV) reverse transcriptase (RT) (Fermentas) and primer 97-101-R (supplementary table S1, Supplementary Material online). PCR amplification was performed using the high-fidelity Phusion DNA polymerase (Finnzymes) and the 73-80-F/92-96-R primers (supplementary table S1, Supplementary Material online). PCR products were resolved on a 1% agarose gel to verify the presence of both genotypes. RT-qPCR was performed to compare viral accumulation in case of direct competition between the two genotypes. RT-qPCR for the *CP* cistron was used to determine the total level of viral accumulation (i.e., accumulation of both genotypes), using the specific primers TEV-*CP*-qPCR-F and TEV-*CP*-qPCR-R (supplementary table S1, Supplementary Material online). RT-qPCRs were performed using One-Step SYBR PrimeScript RT-PCR kit II (Takara) following the instructions provided by the manufacturer. *CP* was chosen because it locates in the 3' end of TEV genome and hence would only quantify complete genomes but not partial incomplete amplicons. Each RNA sample was quantified three times in independent experiments. Amplifications were done using the ABI PRISM Sequence Analyzer 7000 (Applied Biosystems). The thermal profile was as follows: RT phase consisted of 5 min at 42 °C followed by 10 s at 95 °C; and PCR phase of 40 cycles of 5 s at 95 °C and 31 s at 60 °C. Quantification results were examined using SDS7000 software v. 1.2.3 (Applied Biosystems). RT-qPCR for *NlaPro/Nlb* cistrons was then performed to determine viral accumulation of the TEV genotype only, using specific primers TEV-*NlaPro/Nlb*-F and TEV-*NlaPro/Nlb*-R (supplementary table S1, Supplementary Material online) and otherwise identical conditions when compared with the quantification of the *CP*. The ratio of TEV- $\Delta Nlb$  to TEV ( $R$ ) is then  $R = (n_{CP} - n_{NlaPro/Nlb}) / n_{NlaPro/Nlb}$ , where  $n_{CP}$  and  $n_{NlaPro/Nlb}$  are the copy numbers of the *CP* and *NlaPro/Nlb*, respectively, as measured by RT-qPCR. To estimate the replicative advantage  $W$  (Carrasco et al. 2007) of TEV- $\Delta Nlb$ , we performed linear regression on the infection time in days and the log-transformed ratio data. The antilog-transformed slope is  $W$ .

### Experimental Evolution Protocol

Infectious plasmids pTEV7DA and pTEV7DA- $\Delta Nlb$  were linearized with *Bgl*II (Takara) and transcribed into 5'-capped RNAs using the SP6 mMESSAGE mMACHINE kit (Ambion Inc). Transcripts were prepared as described previously (Carrasco et al. 2007). Two batches of five 4-week-old *N. tabacum* 35S::*Nlb* plants were inoculated mechanically on the third true leaf either with TEV transcripts (4–7  $\mu$ g) in inoculation buffer (3% polyethyleneglycol 8000, pH = 7.0) or TEV- $\Delta Nlb$  transcripts (20–25  $\mu$ g) in the same inoculation buffer. In all cases, TEV symptoms appeared 6–7 dpi.

Two evolution experiments were performed with these viruses. In the first experiment, conceived to study the appearance of genomic deletions de novo, four 3-week passages of TEV and TEV- $\Delta Nlb$  were performed in *N. tabacum* 35S::*Nlb* plants. At 21 dpi, all symptomatic leaves of each of the five lineages per virus were collected. Five hundred milligrams of symptomatic tissue was ground in 500  $\mu$ l of grinding buffer (50 mM potassium phosphate pH = 7.0, 3% polyethyleneglycol 6000). Ten microliters of the resulting sap was then mixed with inoculation buffer and used to infect a batch of five healthy transgenic plants by abrasion for each TEV genotype. In the second experiment, conceived to evaluate the subsequent evolution and fitness optimization of genotypes missing the *Nlb* cistron, 30 one-week passages of TEV and TEV- $\Delta Nlb$  were performed in *N. tabacum* 35S::*Nlb* plants. At 7 dpi, symptomatic leaves were collected for each of the five lineages per virus. Infectious sap was prepared as described earlier.

### TEV Genomic RNA Extraction and Purification for Sequence Analysis

To analyze the lineages of TEV- $\Delta Nlb$  and TEV evolved in *N. tabacum* 35S::*Nlb* (second experiment above), total RNA was extracted using RNAeasy Plant Mini Kit (Qiagen) from symptomatic leaves of five *N. tabacum* 35S::*Nlb* infected by the evolved lineages of TEV and five *N. tabacum* 35S::*Nlb* infected by the evolved lineages of TEV- $\Delta Nlb$ . To verify the presence of deletions, the full genome of each genotype was reverse transcribed using M-MLV RT (Fermentas) and primer 97-101-R (supplementary table S1, Supplementary Material online), then PCR amplified using the Ultra High-Fidelity DNA polymerase Phusion (Finnzymes) and primers 73-80-F and 97-101-R (supplementary table S1, Supplementary Material online). Deletions were confirmed by resolution of PCR products on 1% agarose gels.

Subsequently, the full genome was reverse transcribed using M-MLV RT (Fermentas) and primers 97-101-R, 67-77-R, and 40-45-R and PCR amplified using the high-fidelity DNA polymerase Phusion (Finnzymes) and three pairs of primers 2-10-F/40-45-R, 45-48-F/67-77-R, and 73-80-F/97-101-R (supplementary table S1, Supplementary Material online). By using these pairs, we ensured that the mRNA from the

transgene was not amplified and that we obtained only TEV sequences from viral genomes. PCR products were resolved on 1% agarose gels to verify the presence of deletions. For both genotypes, TEV and TEV- $\Delta Nlfb$ , the three amplicons were purified and sequenced by GenoScreen (Lille, France) using BIGDYE 3.1 and a 96-capillars ABI3730XL sequencing system (Applied Biosystems) with overlapping readouts using the same set of primers as in Agudelo-Romero et al. (2008). Contigs were assembled using GENEIOUS version 4.8.

### High-Throughput Sequencing and Data Analysis

To analyze the diversity of deletions that could occur in *Nlfb* cistron during the evolution of TEV in *N. tabacum* 35S::*Nlfb* (first experiment described above), the full genome of each genotype was reverse transcribed using M-MLV RT (Fermentas) and primer R97-101, then PCR amplified using the Ultra High-Fidelity DNA polymerase Phusion (Finnzymes) and primers F73-80 and R97-101 (supplementary table S1, Supplementary Material online). Seven independent RT-PCR replicates were performed for each reaction and then pooled, to limit possible PCR artifacts. These PCR amplicons were sequenced by GenoScreen (Lille, France) with an Illumina HiSeq2000 equipment (Illumina Inc.) using a paired-end ( $2 \times 100$  b) protocol. The reads were already demultiplexed and adapters removed. Data were then analyzed with GSNAP (Wu and Nacu 2010). GSNAP aligned both single-end and paired-end reads. GSNAP is designed to detect short and long-distance splicing, but in our case, we used it to detect deletions instead of introns. GSNAP was used with its default set of parameters, and results were parsed using custom Perl scripts: 1) we excluded reads that are mapped on 100 consecutive bases on TEV genome; 2) for the remaining sequences, we identified sequences that mapped in two different virus genome regions with a minimum of eight bases in the 5' and 3' ends; 3) we excluded reads poorly represented ( $< 10$ ) and showing a low CIGAR (concise idiosyncratic gapped alignment report) string diversity ( $< 10$ ) in order to eliminate possible artifacts; 4) we calculated the distance between the two different regions to define deletion size and position; 5) we calculated the frequency of each deletion detected; and 6) finally, we obtained the number of reads for each deletion variant in R1 and R2 and we compared it to the total number of reads for each lineage.

### Analysis of TEV- $\Delta Nlfb$ Mutants

Each mutation was introduced in the pMTEV infectious clone (Bedoya and Daròs 2010) by site-directed mutagenesis using the Quickchange II XL kit (Stratagene) and following the indications given by the manufacturer. To minimize unwanted errors during the mutagenesis process, the kit incorporates *Pfu* Ultra High-Fidelity Turbo DNA polymerase (Stratagene). The amplification conditions were 1 min at 95 °C (initial denaturation), followed by 18 cycles consisting of 30 s at 95 °C,

45 s at 65 °C and 18 min at 68 °C, and a final extension step of 28 min at 68 °C.

Infectious plasmids of each mutant were linearized with *Bgl*II (Takara) and transcribed into 5'-capped RNAs. Transcripts were prepared as described previously. Two batches of five 4-week-old *N. tabacum* 35S::*Nlfb* plants were, respectively, inoculated mechanically on the third true leaf with TEV transcripts (20–30  $\mu$ g) in inoculation buffer. In all cases, the first symptoms appeared 6–7 dpi. Seven dpi, infectious saps were obtained by grinding 500 mg of TEV mutant-infected tissue in a mortar with 500  $\mu$ l of grinding buffer; then, three 4-week-old *N. tabacum* 35S::*Nlfb* plants for every TEV mutants were, respectively, inoculated mechanically on the third true leaf with 15  $\mu$ l of the corresponding sap.

Total RNA was extracted from the youngest leaves of each of the three plants per TEV mutant 2.67 dpi and then RNAs were purified. RT-qPCR with the *CP* primers was performed to compare viral accumulation of each mutant to the TEV- $\Delta Nlfb$  ancestor as described previously. For each genotype, the Malthusian growth rate per day was computed as  $m = (1/t) \log Q_t$ , where  $Q_t$  is amount of TEV RNA (in pg) per 100 ng of total plant RNA quantified at  $t = 2.67$  dpi. Absolute fitness was then defined as  $W = e^m$  (Crow and Kimura 1970).

## Results and Discussion

### TEV- $\Delta Nlfb$ Has Greater Within-Host Fitness than TEV in *N. tabacum* 35S::*Nlfb*

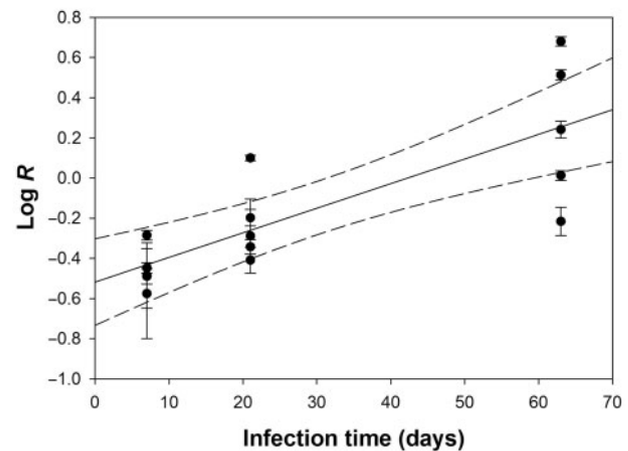
Li and Carrington (1995) generated transgenic *N. tabacum* cv. Xanthi plants expressing TEV *Nlfb* under control of the *Cauliflower mosaic virus* (CaMV) 35S promoter, referred to here as *N. tabacum* 35S::*Nlfb*. TEV variants with disruption of the polymerase GDD amino acid motif, the nuclear localization signal, or the *Nlfb* N-terminal cleavage site could replicate in these transgenic plants, whereas replication was not supported in wild-type plants. Moreover, a TEV variant missing *Nlfb* altogether could also infect these transgenic plants and induce symptoms after 7 dpi, similar to the wild-type virus. Tromas and Elena (2010) also generated a TEV variant missing the *Nlfb* cistron, referred to here as TEV- $\Delta Nlfb$ . TEV- $\Delta Nlfb$  can infect *N. tabacum* 35S::*Nlfb*, but not the wild-type *N. tabacum*, in agreement with previous results (Li and Carrington 1995). TEV is the full-length wild-type virus, capable of infecting both wild-type *N. tabacum* and 35S::*Nlfb* transgenic plants. When wild-type *N. tabacum* plants were coinoculated with different mixtures of TEV:TEV- $\Delta Nlfb$  (1:100 to 100:1), we detected only infection by TEV. Coinfected wild-type plants were probably not observed because the low cellular multiplicity of infection of TEV in *N. tabacum* (Tromas et al. 2014) precludes cellular coinfection, especially early in infection.

TEV- $\Delta Nlfb$  has a 16% smaller genome than TEV. If 1) a relationship between genome size and virus replication exists (Sakai et al. 1999; Zwart et al. 2014) and 2) dependence

on in trans *Nlb* expression for replication does not impose a large fitness burden to the virus, then *Nlb* deletion in the virus genome will cause a faster replication and a selective advantage over a full-length wild-type virus. We therefore inoculated 15 *N. tabacum* 35S::*Nlb* plants with a 1:1 mixture of ground infectious tissue with TEV and TEV- $\Delta$ *Nlb* and measured the ratio of the two variants at 7, 21, and 63 dpi using quantitative reverse transcription PCR (RT-qPCR). Five plants were analyzed for each time point, meaning that each data point represents an independent measurement. On the basis of these data, we estimated the selective advantage for TEV as the per day selection rate constant  $W$  (Materials and Methods). A significant, upward-sloping linear regression was observed between time and the ratio of the TEV- $\Delta$ *Nlb*:TEV (fig. 1;  $r^2 = 0.632$ ,  $F_{1,14} = 22.330$ ,  $P < 0.001$ ), meaning that the ratio of the viruses shifts significantly toward TEV- $\Delta$ *Nlb*, and this virus therefore has a significantly higher competitive fitness. We estimated that  $W = 1.028 \pm 0.006 \text{ d}^{-1}$  ( $\pm 1$  SD), suggesting that TEV- $\Delta$ *Nlb* will slowly displace the wild-type virus in transgenic plants. If genome size was inversely proportional to fitness and the 16% reduction in genome size were the only determinant of fitness for TEV- $\Delta$ *Nlb*, we would expect  $W = 1/(1-0.16) = 1.19 \text{ d}^{-1}$ . The 95% confidence interval for our estimate of  $W$  is 1.016–1.040  $\text{d}^{-1}$ , suggesting that 1) there is a cost to depend on in trans *Nlb* expression, but 2) this cost is partially compensated for by faster replication of TEV- $\Delta$ *Nlb*, resulting in a higher  $W$  than the wild-type virus. Our data therefore support the idea of a relationship between genome size and fitness, and suggest that viruses harboring deletions in *Nlb* will outcompete the wild-type virus in *Nlb*-expressing plants. Therefore, NIRV expression might also lead to conditions favoring the replication of virus variants, with deletions in the gene(s) encoded for by the NIRV. However, the accumulation of viruses with large genomic deletions due to in trans expression of virus' genes will also depend on mutational supply, specifically how often recombination generates viruses with deletions in *Nlb* that conserve the reading frame in downstream genes.

#### Full-Length Genomes Were Not Recovered by Recombination between TEV- $\Delta$ *Nlb* and the Transcripts from the 35S::*Nlb* Transgene

Borja et al. (1999) reported that inoculation of transgenic *N. benthamiana* plants expressing the CP protein of tomato bushy stunt tobamovirus (TBSV) with TBSV mutants carrying defective CP resulted in the restoration of wild-type virus throughout a double-recombination event between the viral genome and the mRNA transcribed from the transgene. In sharp contrast, several authors have failed to detect recombination events between transgenes and potyviruses (Chung et al. 2007; Dietrich et al. 2007). To test for possible recombination events in our system, we performed an evolution

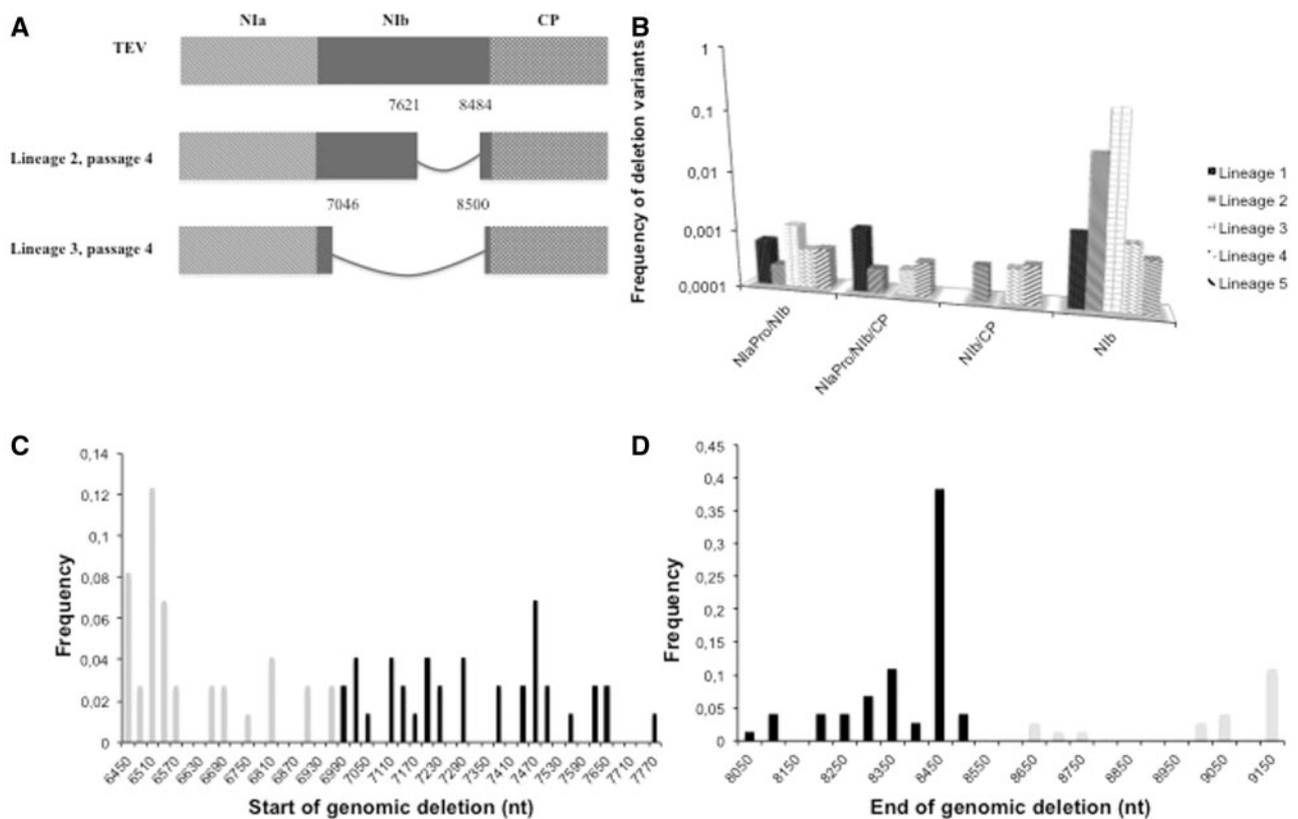


**Fig. 1.**—Plot of the log-transformed ratio of TEV:TEV- $\Delta$ *Nlb* (ordinate) over time (abscissa), given in weeks postinoculation. Error bars represent  $\pm 1$  standard deviation, the solid line represents the linear regression and the dashed lines the 95% CI of regression.

experiment consisting of five independent lineages of TEV- $\Delta$ *Nlb* serially passaged four times in *N. tabacum* 35S::*Nlb* transgenic plants, with each passage being 3 weeks long. At the end of the evolution experiment, we analyzed the evolved lineages by RT-PCR using a combination of primers that will specifically amplify full-genome viruses (supplementary table S1, Supplementary Material online). We found no evidence for recombination between TEV- $\Delta$ *Nlb* and host-derived *Nlb* transcripts. Therefore, we conclude that, in our experimental set up, regeneration of the wild-type genome by recombination is an unlikely phenomenon. This outcome is congruent with the estimates of within-host fitness for TEV and TEV- $\Delta$ *Nlb*; recombinants analogous to the wild-type virus would be displaced by TEV- $\Delta$ *Nlb*, and the process could occur rapidly as de novo variation will initially be present at low frequencies in the virus population. On the other hand, this result may reflect the possibility that recombination between TEV- $\Delta$ *Nlb* and the transgene transcripts is very rare, as shown for other potyviruses (Chung et al. 2007; Dietrich et al. 2007).

#### Rapid Occurrence of Large Genomic Mutations in TEV upon Passaging in *N. tabacum* 35S::*Nlb* Plants

To test whether there will indeed be rapid accumulation of virus variants with large deletions as suggested by estimates of  $W$ , we performed serial passage experiments. Here, we chose to perform four serial passages, with each passage having 3 weeks of duration, because genome size appears to be under stronger selection in long passages (Zwart et al. 2014). Five independent lineages of TEV were evolved. After four passages, deletions were detected by RT-PCR in two lineages: Lineage 2 and Lineage 3 (fig. 2A). These two deletions (nucleotide positions 7621–8484 and 7046–8500) conserved the



**FIG. 2.**—(A) TEV and deletion variants detected by RT-PCR in different lineages at passages one and four. After four 3-week passages, *Nlb* deletion variants are observed in different lineages as, for example, Lineage 4 and Lineage 5. All deletions observed after RT-PCR are situated within *Nlb* and retain the reading frame. (B) Histogram of the frequency of different deletion variants observed in the five evolved lines. (C) Frequency distribution of the start nucleotide position of genomic deletions in the evolved lineages. (D) Frequency distribution of the end nucleotide position of genomic deletions in the evolved lineages. In (C) and (D), black bars indicate positions within the *Nlb* cistron, gray bars indicate regions outside *Nlb*, that is, in (C) within the *NlaPro* cistron and in (D) within the *CP* cistron.

reading frame and were situated within the *Nlb* cistron (positions 6982–8517 of the wild-type genome).

To detect virus deletion variants with much greater sensitivity, Illumina NGS was performed on these five TEV lineages evolved for four 3-week passages. A TEV genome region encompassing the *NlaPro*, *Nlb*, and *CP* cistrons (positions 6377–9473 of the wild-type genome) was amplified and sequenced (Materials and Methods). We developed a new approach for identifying deletions, adapting GSNAP (Wu and Nacu 2010) to detect deletions instead of introns (Materials and Methods). As a negative control, the same region was analyzed using NGS data for five lineages of TEV-eGFP evolved for three 9-week passages in wild-type *N. tabacum* plants (Zwart et al. 2014). No deletions were detected in the entire region, suggesting that our NGS analysis does not generate artifacts. The frequencies of deletion variants (see supplementary table S1, Supplementary Material online) in systemically infected leaves of transgenic plants were significantly higher than the frequencies found in our negative controls (Fisher’s exact test,  $P < 0.001$ ). The NGS results were consistent with RT-PCR data; the deletion variant detected by RT-PCR in Lineage 2

(positions 7621–8484) and Lineage 3 (positions 7046–8500) was the variant found at the highest frequency by NGS, 0.249 and 0.246, respectively. Sixty-four different deletion variants were observed in total, ranging in frequency from  $8.44 \times 10^{-5}$  (positions 6538–8478; Lineage 4) to 0.249 (positions 7621–8484; Lineage 2). Interestingly, 6.25% of deletion variants disrupted the reading frame of the coding sequence. Moreover, 43.75% of deletion variants extended into genes flanking *Nlb*, meaning that probably only the remaining 56.25% variants could be viable when transcomplemented by the *Nlb* expressed by the plant.

In 17 deletion variants, the 5’ end of the deletion extended into *NlaPro*; the frequencies of these variants in the virus populations range from  $8.44 \times 10^{-5}$  (positions 6538–8478; Lineage 4) to  $1.73 \times 10^{-3}$  (positions 6460–8478; Lineage 4). Furthermore, in eight other variants the deletion started within *NlaPro* (upmost 5’ position being 6516) and ended within *CP* (downmost 3’ position being 9163), varying in frequency from  $1.02 \times 10^{-4}$  (positions 6923–9013; Lineage 4) to  $1.88 \times 10^{-3}$  (positions 6569–8746; Lineage 1). The 43.75% of genotypes carrying deletions larger than the *Nlb* cistron will produce

viruses with truncated cistrons *NlaPro*, *CP*, or both. Moreover, deletions beyond the *Nib* cistron will lead to the loss of proteolytic cleavage sites and result in fusion of the truncated proteins. These genomes will probably be defective particles and some may additionally interfere with replication of the full-length virus. These defective particles can only replicate if they are complemented by a full-length TEV or possibly other defective genomes, and their spread may therefore be limited. The highest frequency of putatively defective deletion variants does not exceed 1%, in line with our expectations: two *NlaPro*-to-*CP* deletion variants (positions 6569–8746 and 6774–8795; Lineage 1) show frequencies of  $1.88 \times 10^{-3}$  and  $1.58 \times 10^{-3}$ , respectively. All lineages are dominated by variants with *Nib* deletion only, which is consistent with the fact that these variants are viable. Although defective viruses are often maintained in virus populations by complementation (Roux et al. 1991; Moreno et al. 1997; Froissart et al. 2004; García-Arriaza et al. 2004), the low level of coinfection (Dietrich and Maiss 2003; Tromas et al. 2014) may rapidly eliminate defective viruses from potyvirus populations.

The ends of the genomic deletions found by NGS (fig. 2C and D) were not uniformly distributed (Kolmogorov–Smirnov tests: 5' end  $n = 42$ ,  $P < 0.001$ ; 3' end  $n = 23$ ,  $P < 0.001$ ). Surprisingly, the 5' start of the genomic deletion was clustered in *NlaPro*, suggesting a possible recombination hotspot outside of the *Nib* cistron. For the 3' end of the deletion, ending sites clustered toward the 3' end of *Nib* cistron. This clustering is consistent with the idea that we are more likely to detect deletion variants that only affect part of *Nib* and are, therefore, likely viable by complementation in trans.

Our results demonstrate that rapid deletions of *Nib* occurred upon passaging in *N. tabacum* 35S::*Nib*. In two lineages, these variants reached high frequencies, comprising approximately a fourth of the population. These results are therefore a proof of principle that NIRV expression could have a similar effect on a virus population: allowing deletion mutants to be maintained in the virus populations, while also providing conditions in which these mutants could reach high frequencies. The immediate effect of these deletions will be that a part of the virus population is replication defective in host individuals that do not express the NIRV. This could have important ramifications for viral spread, and virus–host coevolution. On the other hand, the further evolution of a virus population in a host expressing virus genes may also be contingent upon the presence of genomic deletions in the viral genome.

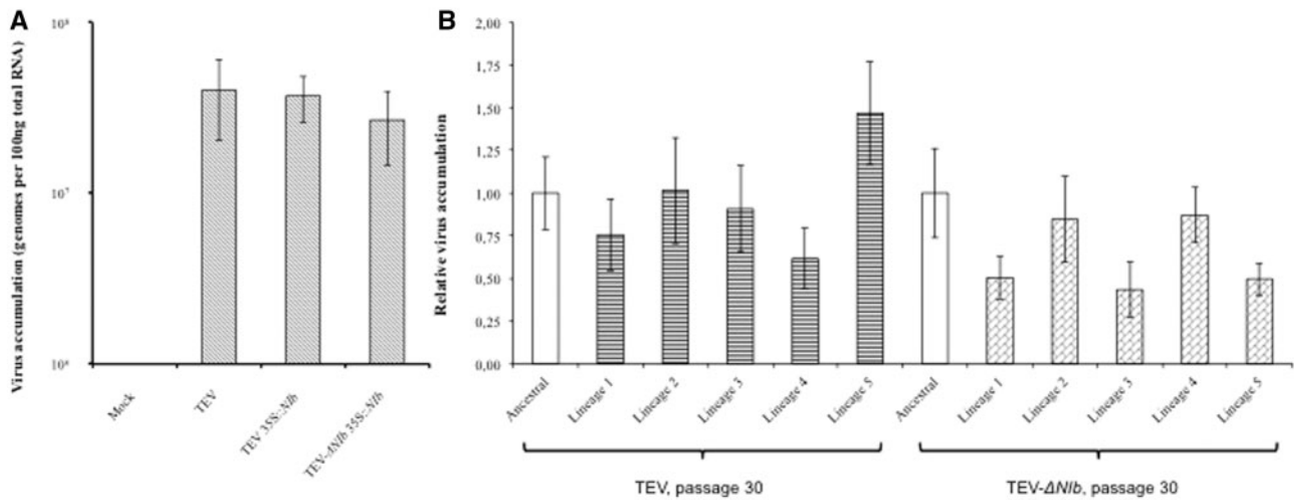
#### TEV Evolution in *N. tabacum* 35S::*Nib* Plants Is Contingent upon the Presence of *Nib* in the Viral Genome

As we just described, serial passages of TEV in *N. tabacum* 35S::*Nib* led to the rapid occurrence of a whole spectrum of deletion variants. How would these deletion mutants further

evolve in transgenic plants, and would their evolution be different from that of a full-length TEV? To address this question and gain insight into the long-term evolution of viruses harboring NIRV-induced deletions, we performed a second evolution experiment for both TEV and TEV- $\Delta$ *Nib* passaged in *N. tabacum* 35S::*Nib*, with five independent lineages for each virus. Li and Carrington (1995) showed that beta-glucuronidase (GUS) activity in leaves systemically infected by the TEV- $\Delta$ *Nib*-GUS mutant was comparable to that of the parental TEV-GUS. We found a similar result for the accumulation of TEV and TEV- $\Delta$ *Nib* in *N. tabacum* 35S::*Nib* using RT-qPCR (fig. 3A): there was not a significant difference in virus accumulation at 7 dpi ( $t_8 = 1.394$ ,  $P = 0.201$ ). The infection dynamics of the two viruses should therefore be similar, allowing us to compare the evolutionary outcomes. However, to compare the evolution of the full-length TEV and TEV- $\Delta$ *Nib*, we had to devise a setup in which the integrity of the *Nib* gene is ensured for TEV. Two previous studies have found that marker genes were highly stable in the viral genome when many 1-week passages were performed, although the markers were quickly lost in longer passages (Dolja et al. 1993; Zwart et al. 2014). By performing 30 one-week passages with TEV and TEV- $\Delta$ *Nib* in *N. tabacum* 35S::*Nib*, we therefore hoped to avoid the occurrence of large genomic deletions in TEV. On the other hand, we realize that selection may not operate as effectively as during longer passages, due to short passage duration and frequent bottleneck associated to the mechanical transmissions among plants.

After 30 weeks of evolution, we measured virus accumulation of the evolved lineages (fig. 3B). The data were analyzed by ANOVA (table 1), and we found no effect of the virus type (TEV or TEV- $\Delta$ *Nib*) on accumulation, although there was a strong effect of lineage. The significant lineage-dependent effects suggest that random factors have played an important role during evolution, implicating genetic drift effects due to short passage duration. When pairwise comparisons were made between the evolved lineages and their respective ancestral viruses, Lineage 1 of TEV- $\Delta$ *Nib* had a significantly lower accumulation ( $t$ -test with Holm–Bonferroni correction:  $t_7 = 4.001$ ,  $P = 0.005$ ). Accumulation of all other lineages was not significantly different from the ancestral virus, although the data suggest Lineage 3 and Lineage 5 of TEV- $\Delta$ *Nib* also have low accumulation (fig. 3B).

All evolved lineages of TEV and TEV- $\Delta$ *Nib* were fully sequenced to determine whether there were differences in their patterns of molecular evolution. We were particularly interested in understanding whether the use of *Nib* in trans may have affected the regulation of expression and accumulation of all other viral components. Mutational spectra for both viral genotypes were dominated by point mutations as not a single insertion or deletion was observed (table 2). As anticipated, the passaging setup used leads to the retention of *Nib* cistron by all TEV lineages, and by contrasting the evolution of TEV and TEV- $\Delta$ *Nib*, we are therefore considering the evolutionary



**Fig. 3.**—(A) TEV and TEV-ΔNlb accumulation evaluated by absolute RT-qPCR after one passage in *N. tabacum* 35S::Nlb transgenic plants. No significant difference in the accumulation of TEV and TEV-ΔNlb was observed. (B) Relative TEV and TEV-ΔNlb virus accumulation in *N. tabacum* 35S::Nlb after 30 passages.

**Table 1**  
Nested ANOVA for Virus Accumulation of Evolved TEV and TEV-ΔNlb

Source of Variation	SS	d.f.	MS	F	P
Virus	0.768	1	0.768	1.663	0.233
Lineage within virus	3.721	8	0.465	3.779	0.003
Error	4.431	36	0.123		

NOTE.—Viral accumulation data were normalized by the accumulation of the ancestral virus. SS is the sum of squares, d.f. is the degrees of freedom and MS is the mean square.

effects of having a full-length or truncated genome. There were no differences between ratios of transitions to transversions; the ratio was 8.5 for TEV and 6.0 for TEV-ΔNlb (Fisher’s exact test:  $P = 1.000$ ). Given that it is biochemically easier to produce transitions than transversions, it is not surprising to observe these biased ratios. Likewise, there was not a significant difference in the ratio of synonymous to nonsynonymous substitutions between the two viruses (Fisher’s exact test:  $P = 0.330$ ); for TEV, this ratio was 1.375 whereas for TEV-ΔNlb it was 0.636. For both genotypes, mutations were evenly distributed along the genome (fig. 4A; Kolmogorov–Smirnov test: TEV  $n = 19$ ,  $P = 0.330$ ; TEV-ΔNlb  $n = 21$ ,  $P = 0.688$ ). An interesting example of convergent evolution was mutation A6805G (K2269E) that appeared in 80% of TEV lineages and in 40% of TEV-ΔNlb lineages. This nonsynonymous mutation is localized in a superficial domain of NlaPro (Nunn et al. 2005), and it could affect the proteinase function or modulate the interactions of NlaPro with other proteins. This same mutation has also been previously reported in serial-passage evolution experiments of TEV in *N. tabacum*: in 10% of lineages after 15 one-week passages (Bedhomme et al. 2012), and in 30% of lineages after three 9-week passages (Zwart et al. 2014).

**Table 2**  
Mutation Spectrum and Substitution Matrix for TEV and TEV-ΔNlb Evolved Genome after 30 one-week Passages in *N. tabacum* 35S::Nlb Transgenic Plants

	Type of Mutations	
	TEV	TEV-ΔNlb
Total	19	21
Base substitution	19	21
Transitions	17	18
Transversions	2	3
Synonymous	11	7
Nonsynonymous	8	11
Untranslated regions	0	3

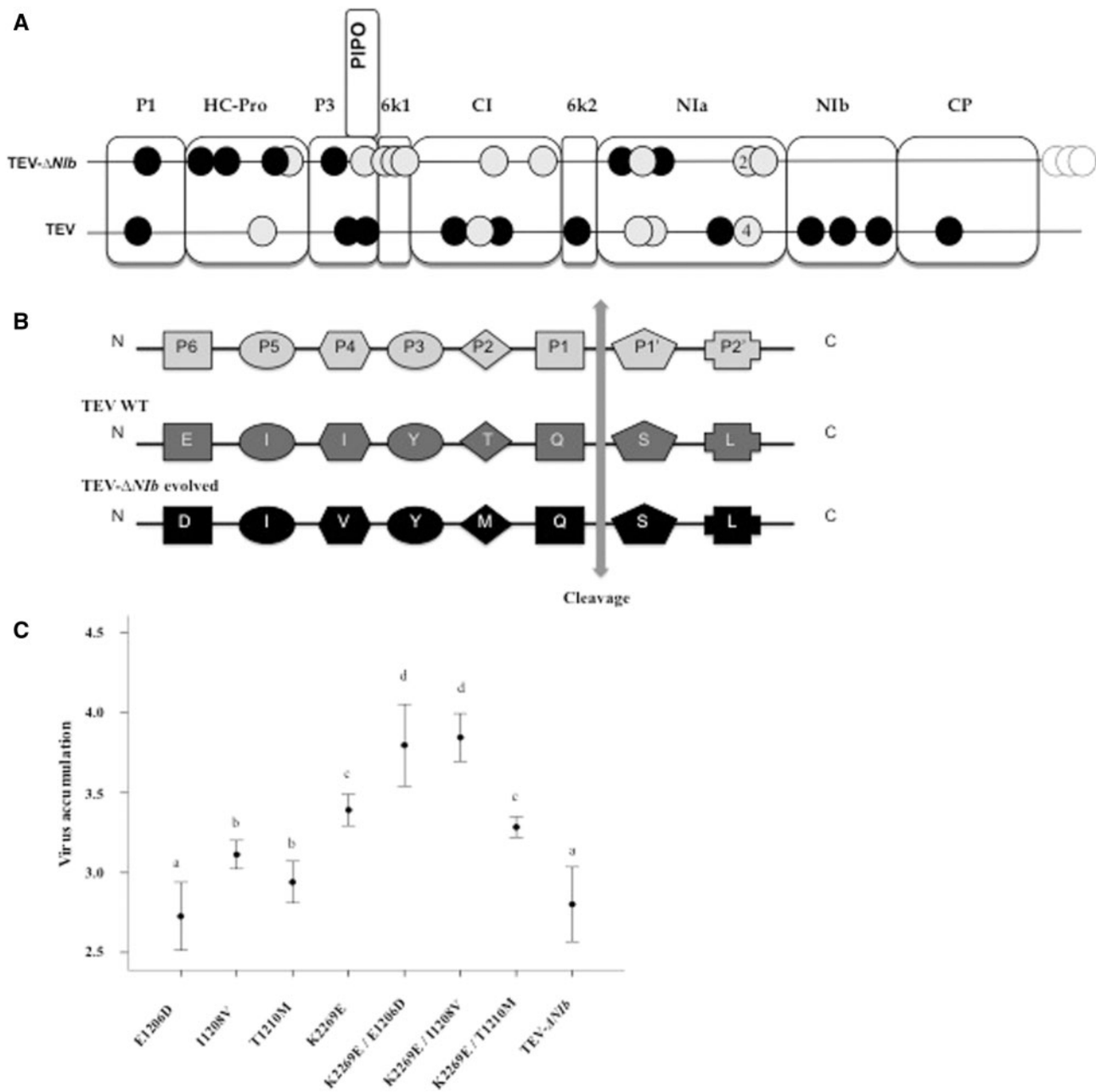
  

	Substitution matrices			
	A	U	G	C
A	—	0	5	0
U	2	—	0	2
G	6	0	—	0
C	0	4	0	—

	Substitution matrices			
	A	U	G	C
A	—	0	9	0
U	0	—	0	3
G	4	1	—	0
C	1	3	0	—

In three lineages of TEV-ΔNlb, we observed three nonsynonymous mutations localized in the 6K1/CI junction (fig. 4B): G3618U (E1206D) (Lineage 1), A3622G (I1208V) (Lineage 5), and C3629U (T1210M) (Lineage 3), affecting residues P6, P4, and P2 of the proteolytic site, respectively (Adams et al. 2005). A conserved heptapeptide sequence has been identified at this site [E-X-X-Y-X-Q/S (or G)] at the 6K1/CI junction (Carrington and Dougherty 1987, 1988; Carrington et al. 1988; Adams et al. 2005). Each mutation was observed in a single lineage, and combinations thereof were not observed.



**FIG. 4.**—(A) Mutations found in TEV- $\Delta$ Nlb and TEV consensus genome after 30 weekly passages in *N. tabacum* 35S::Nlb transgenic plants. Black circles represent synonymous mutation, gray circles nonsynonymous mutation, and white circle mutations localized in UTRs. (B) TEV- $\Delta$ Nlb mutations on the NIaPro proteolytic 6K1/CI site. P6 to P2' (E-X-X-Y-X-Q-G/S) are key amino acids constituting the NIaPro binding site. Cleavage site is between P1 and P1' (vertical arrow). (E) (26.5%) and (D) (12.8%) are the most frequent amino acids at P6 position in the family Potyviridae; V (80%) is the most frequent in position P4; H (40%) and F (15%) are the most common in position P2 among the Potyviridae, M is represented with a frequency of only 2%. Generally, mutations at positions P2/P4 may involve regulation of the cleavage rate (Dougherty et al. 1989). (C) Accumulation of mutants E1206D, I1208V, T1210M, K2269E, and the combinations K2269E/E1206D, K2269E/I1208V, and K2269E/T1210M. Mutant viruses belong to four groups (A–D) with homogeneous accumulation (Holm-Bonferro post hoc test;  $P < 0.05$ ).

The E1206D mutation at the P6 site may only weakly affect the proteolytic processing as amino acids E and D are widely represented in Potyviridae family at this site (Adams et al. 2005). On the other hand, I1208V at position P4 should

have a stronger impact on processing activity; in only 7% of all Potyviridae is I present at this site, whereas V is present in 80% (Adams et al. 2005). Similarly, T1210M in P2 may also have a strong impact on processivity: T is found at the P2 site

in only 2% of the Potyviridae (2%). Generally, mutation at sites P2/P4 may involve regulation of the cleavage rate (Dougherty et al. 1989).

We conclude that the evolution of TEV and TEV- $\Delta Nlb$  in *N. tabacum* 35S::Nlb was similar when transmission bottlenecks are close in time. Although there was a significant lineage-dependent effect on accumulation, only one evolved lineage of TEV- $\Delta Nlb$  had a significantly lower accumulation than its ancestral virus. Moreover, patterns of molecular evolution were similar. There were no significant differences in the ratios of transitions to transversions, synonymous to nonsynonymous mutations, and the overall distribution of mutations. The one mutation that occurred in multiple lineages, A6805G, occurred in both lineages of TEV and TEV- $\Delta Nlb$ . Therefore, the only conspicuous difference between the two viruses is mutations localized in the 6K1/CI junction.

#### Analysis of Mutations in the 6K1/CI Junction of TEV- $\Delta Nlb$

The K2269E, E1206D, I1208V, and T1210M mutations were introduced in the pMTEV infectious clone by site-directed mutagenesis. As K2269E is common among the different lineages, we also generated clones containing this mutation in combination with the other three. As a proxy for within-host fitness, we measured viral accumulation of each simple and double mutant at 2.67 dpi by RT-qPCR in three biological replicates. Data were fitted to a GLM with a log-link function and gamma distributed error structure, using replicate as factor nested within virus genotype. A significant effect of virus genotype on the observed viral accumulation was found ( $\chi^2 = 134.648$ , 7 d.f.,  $P < 0.001$ ), and a Holm–Bonferroni post hoc test identified four different homogenous groups (fig. 4C), with *a* being the group with the lowest viral accumulation value and *d* the highest. The results are generally in agreement with our expectations based on the prevalence of residues in the Potyviridae. E1206D has the same accumulation as the ancestral virus. The other single mutations all increased accumulation, being represented in groups *b* and *c*. The double mutations resulted in the highest accumulation levels, with two being the sole members of group *d* and one being represented in group *c*.

Thus, we conclude that all mutations except E1206D increase accumulation of TEV- $\Delta Nlb$  early in infection and that this effect was even stronger for double mutations. High levels of early accumulation are likely to occur for a virus variant that expands rapidly, and therefore suggest an increase in the within-host fitness of the evolved lineages. The mutations in the 6K1/CI proteolytic site were not observed in the wild-type virus, suggesting that this is an instance of adaptive evolution contingent upon deletion of *Nlb*.

TEV is translated into a large polyprotein, thus the cleavage process may be a mechanism to regulate the effective

concentration of viral proteins in infected cells (Merits et al. 2002). Most of these mutations analyzed in this section showed a significant higher viral accumulation than the ancestral genotype. The mutations we found in evolved lineages may positively regulate the cleavage 6K1/CI in order to produce more mature 6K1 and CI proteins. Nlb and CI are constituents of the viral replication complex (Li et al. 1997), and thus increasing the effective CI concentration may counterbalance the lower production of Nlb from the transgene.

#### Concluding Remarks

NIRVs are thought to impact viruses by—in some cases—conferring virus resistance to host organisms (Koonin, 2010). However, these virus-derived sequences could also impact the evolution of virus populations. Here, we used an experimental model system to study the effects of NIRV expression on virus populations, using TEV and transgenic tobacco plants expressing its replicase, Nlb. We found that 1) the loss of *Nlb* cistron leads to higher within-host fitness in transgenic plants, 2) that genomic deletions quickly accumulate in a full-length virus passaged in transgenic plants, and 3) that the evolution of a virus population can be contingent upon the deletion of virus genes, although many aspects of phenotypic and molecular evolution were unchanged. Overall, our results therefore suggest that the expression of NIRVs could have far-reaching consequences for virus evolution. Moreover, these effects can occur on relatively short timescales (i.e., deletion variants were generated de novo and reached frequencies of ~0.25 within four 3-week passages). These changes on short timescales support the notion that plausible epidemiological models of virus spread in NIRV-expressing host populations need to take virus evolution into consideration. For example, if large genomic deletions occur rapidly in NIRV-expressing hosts, then eventually many virus-infected hosts will only be infectious to those hosts expressing the NIRV. The generation of various classes of infectious hosts could in this case decrease viral transmission between hosts, an effect that could be further augmented by the clustering of NIRV-expressing hosts.

Our results provide further support for the notion of a relationship between genome size and replication/within-host competitive fitness (Sakai et al. 1999; Marks et al. 2005; Zwart et al. 2010, 2014), although there are clearly limitations to this idea (Bull et al. 2004). Not only could TEV- $\Delta Nlb$  outcompete TEV in transgenic plants, also all independent TEV lineages accumulate various *Nlb* deletions upon passaging. We identified 64 different deletion variants, most of which conserved the reading frame. In 17 deletion variants the deletion extended far into the upstream cistron, *NlaPro*, while for eight deletion variants, the deletion also extended into the downstream cistron, *CP*. Because both *NlaPro* and *CP* proteins are essential for virus replication and these deletions removed one or more proteolytic

cleavage sites, the virus variants probably cannot replicate autonomously, even in the transgenic plants. Therefore, the virus population in the simulated NIRV-expressing host appears to have generated two types of defective viruses: 1) those with deletions in *Nlb* only and can replicate autonomously in the transgenic plants but not in the wild-type plants and 2) those with deletions extending beyond the *Nlb* cistron, which cannot replicate autonomously in either host. On the basis of these observations, we speculate that the NIRV-expressing hosts may be a source of defective genotypes in natural virus populations. If a deletion genotype arises de novo in a NIRV-expressing host, it will not require coinfection with a full-length virus to be maintained in the population. Consequently, it may more easily reach intermediate to high frequencies in a NIRV-expressing plant, allowing the defective virus to be sustained even in non-NIRV hosts by coinfection with full-length viruses (Montville et al. 2005). Complementation can slow down the rate at which deleterious alleles are eliminated from the virus population (Froissart et al. 2004; Sardanyés and Elena 2010) and promote selection for defective-interfering particles as cheater genotypes that reduce mean population fitness (Turner and Chao 1999).

Our study has important implications for the development and agronomic use of transgenic plants resistant to viruses generated by inserting complete viral genes into the plant genome, in particular by inserting viral replicase genes (reviewed in Kundu and Mandal [2001]). From a biosafety standpoint, it has been known for several years that recombination between viral sequences transgenically expressed by plants and virus genomes may occur (Turturo et al. 2008; Morroni et al. 2009), raising concerns about whether this transgenic strategy would favor the emergence of new viruses (Tepfer 2002; Thompson and Tepfer 2010). Here, we have illustrated a different evolutionary scenario: the expression by the plant of functional viral proteins creates the conditions in which viruses with long or even full deletions of the complemented protein may dominate the population, thus precluding their spread in nontransgenic plants.

Finally, it is fair to mention that our experimental system remains artificial to some extent. Although many examples of DNA viruses using host enzymes for replication exist (e.g., geminiviruses and polydnviruses), no RNA virus has been described so far that does not encode for its own functional RdRp, even though certain plants encode functional RdRps of viral origin in their genomes, for example, the narnavirus-like RdRp found in the mitochondria of *Arabidopsis thaliana* (Hong et al. 1998). In this respect, a critical reader may argue that RdRp was not the best choice for studying the pseudogenization of viral genes in the presence of a homologous NIRV. However, our results should be strong enough to convince such a reader that even a highly constrained gene, such as RdRp can be lost from viral genomes if provided in trans by the host.

## Supplementary Material

Supplementary table S1 is available at *Genome Biology and Evolution* online (<http://www.gbe.oxfordjournals.org/>).

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