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A complex *eIF4E* locus impacts the durability of *va* resistance to *Potato virus Y* in tobacco

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**SUMMARY**

Many recessive resistances against potyviruses are mediated by eukaryotic translation initiation factor 4E (*eIF4E*). In tobacco, the *va* resistance gene commonly used to control *Potato virus Y* (*PVY*) corresponds to a large deletion affecting the *eIF4E-1* gene on chromosome 21. Here, we compared the resistance durability conferred by various types of mutations affecting *eIF4E-1* (deletions of various sizes, frameshift or nonsense mutations). The 'large deletion' genotypes displayed the broadest and most durable resistance, whereas frameshift and nonsense mutants displayed a less durable resistance, with rapid and frequent apparition of resistance-breaking variants. In addition, genetic and transcriptomic analyses revealed that resistance durability is strongly impacted by a complex genetic locus on chromosome 14, which contains three other *eIF4E* genes. One of these, *eIF4E-3*, is rearranged as a hybrid gene between *eIF4E-2* and *eIF4E-3* (*eIF4E-2* and *eIF4E-3* in the genotypes showing the most durable resistance, while *eIF4E-2* is differentially expressed between the tested varieties. RNA-seq and quantitative reverse transcriptase-polymerase chain reaction experiments demonstrated that *eIF4E-2* expression level is positively correlated with resistance durability. These results suggest that besides the nature of the mutation affecting *eIF4E-1*, three factors linked with a complex locus may potentially impact *va* durability: loss of an integral *eIF4E-1*, presence of *eIF4E-2* and overexpression of *eIF4E-2*. This latter gene might act as a decoy in a non-productive virus–plant interaction, limiting the ability of *PVY* to evolve towards resistance breaking. Taken together, these results show that *va* resistance durability can in large part be explained by complex redundancy effects in the *eIF4E* gene family.

**Keywords:** *eIF4E*, tobacco, potyvirus, *va* resistance, durability, Ethyl methanesulfonate mutant.

**INTRODUCTION**

*Potato virus Y* (*PVY*) has been classified in the top 10 of the most economically and scientifically important plant viruses (Scholthof et al., 2011). It is the type member of the genus *Potyvirus* in the *Potyviridae* family, one of the largest genera of plant viruses (Wylie et al., 2017). *PVY* is transmitted by aphids and infects a large range of host plants worldwide, mostly within the *Solanaceae* family. In particular, it causes one of the most damaging diseases in cultivated tobacco (Quenouille et al., 2013). *PVY* isolates are distributed into three biotypes, with the *PVYN* and *PVYP* biotypes separated by their ability or inability, respectively, to cause systemic vein necrosis on tobacco (Moury, 2010; Singh et al., 2008). Although mosaic symptoms impact less dramatically the yield and quality of tobacco crops, severe leaf necrosis induced by *PVYN* isolates are a major concern for tobacco producers (Lacroix et al., 2010; Rolland et al., 2009; Tian et al., 2011).

As for other potyviruses, the *PVY* genome consists of a single RNA molecule, polyadenylated at its 3' end and covalently linked to a 25 kDa virus-encoded VPg protein at its 5' end (Leonard et al., 2000; Revers and García, 2015). The completion of the viral cycle involves a complex interplay between virus- and
host-encoded factors, also called susceptibility factors. Absence or non-adequacy of a single susceptibility factor can result in plant resistance, a phenomenon described as ‘loss-of-susceptibility’ (van Schie and Takken, 2014). Such recessive resistance genes against potyviruses have mainly been identified among translation initiation factors, including eukaryotic initiation factors 4E (eIF4E) and 4G (eIF4G) or their isoforms (Robaglia and Caranta, 2006; Sanfaçon, 2015; Truniger and Aranda, 2009; Wang and Krishnaswamy, 2012). Translation initiation factors 4E are essential components encoded by a small multigene family that bind to the mRNA cap structure at the 5’ end of most mRNAs (Browning and Bailey-Serres, 2015). Natural resistance to potyviruses in crops such as lettuce (Nicaise et al., 2003), pepper (Ruffel et al., 2002), pea (Gao et al., 2004) and tomato (Ruffel et al., 2005) has been shown to result from non-synonymous substitutions in eIF4E genes. Even if the exact mechanism(s) involved in eIF4E-mediated resistance are still unclear (Wang and Krishnaswamy, 2012), it has been shown that the viral genome linked protein (VPg) can interact with eIF4E, mimicking the 5’ cap structure of messenger RNAs (mRNAs). In addition, potyviruses can evolve towards resistance breaking through the acquisition of mutations that either restore compatibility of their VPg with the host mutant eIF4E or allow an interaction with another member of the eIF4E family (Charron et al., 2008; Gallois et al., 2010, 2018; Lebaron et al., 2016; Takakura et al., 2018).

In tobacco, the main source of resistance against PVY is the va gene, which originates from the Virgin A Mutant (VAM) tobacco line obtained after X-ray irradiation-induced mutagenesis (Koelle, 1961; Singh et al., 2008). A large deletion of almost 1 Mbp has been characterized in VAM using random amplified polymorphic DNA (RAPD) markers (Noguchi et al., 1999) and a physical map of the corresponding region of chromosome 21 was recently developed (Dluge et al., 2018). Julio et al. (2015) showed that a deletion of a particular copy of elf4E (hereafter named elf4E-1) encoded by the S10760 gene is responsible for the va resistance. In particular, mutants obtained by ethyl methane sulfonate (EMS) mutagenesis with nonsense mutations in the S10760 gene showed PVY resistance, confirming its direct involvement in va resistance (Julio et al., 2015).

However, the resistance of elf4E-1 EMS-KO mutants is unstable, with the rapid and frequent emergence of PVY resistance-breaking (RB) variants (Julio et al., 2015). These observations suggest a possible role of the particular type of mutation affecting elf4E-1 or of additional genetic factor(s) in the durability of the va resistance. Julio et al. (2015) identified 35 tobacco accessions carrying a va resistance allele in a germplasm collection. Depending on the type of mutation at the va locus, four groups of resistant genotypes were distinguished (Fig. 1). The first group, named LD for ‘large deletion’, corresponds to tobacco varieties carrying the 1 Mbp deletion on chromosome 21. The second group, named SD for ‘small deletion’, gathers tobacco varieties carrying a smaller deletion on chromosome 21 (as judged by the presence of markers absent in the LD group) that also results in a complete deletion of elf4E-1. The third category, named FS for ‘frameshift’ does not display any chromosomal deletion but a natural 2 bp deletion in the elf4E-1 gene, leading to a C-terminal truncated elf4E-1 protein of only 163 amino acids. The fourth group includes two EMS stop codons mutants, EMS1 and EMS2, that encode C-terminal truncated elf4E-1 proteins of only 50 and 53 amino acids, respectively [corresponding to mutants W50* and W53* in Julio et al. (2015)].

The objectives of the present study were to better characterize the differences in resistance durability observed between tobacco genotypes differing in the va allele they

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**Fig. 1** Type of mutation at the elf4E-1 locus on chromosome 21 in the different N. tabacum va accessions. The va accessions listed on the right are classified into four categories (LD, SD, FS and EMS) according to the type of mutation at the elf4E-1 locus. The EMS1 and EMS2 mutants correspond to the W50* and W53* mutants described in Julio et al. (2015). ‘Unknown’ indicates that the length of the deletion in the SD genotypes is not known.
possess. Through genetic and transcriptomic analyses we show here that resistance durability is strongly influenced by a complex genetic locus on chromosome 14, which contains other eIF4E copies. One of these copies is absent and replaced by a hybrid copy in the genotypes with the most stable resistance. In addition, the mRNA expression level of another copy is positively correlated with resistance durability. Taken together these results show that va-mediated resistance durability can be explained by redundancy/competition effects between members of the elf4E gene family, parallel to what was recently shown in tomato (Gauffier et al., 2016).

RESULTS

Phenotypic evaluation of the resistance conferred by different types of mutations at the va locus

Our previous results showed that the resistance of EMS-KO eIF4E-1 mutants is unstable, with the rapid and frequent emergence of PVY RB variants (Julio et al., 2015). These observations suggest a possible role of the particular type of mutation affecting eIF4E-1 or of additional genetic factor(s) in the durability of the va-mediated resistance. In order to investigate the potential role of the mutation type affecting eIF4E-1, we challenged 13 accessions representing different classes of va mutations (Fig. 1) with nine PVY isolates avirulent on VAM and representative of the virus genetic diversity (five belonging to the N clade and four to the O clade, see the experimental procedures section). The susceptible BB16 genotype was used as a positive control for infection. For each of the 117 tobacco genotype–PVY isolate combinations, the infection rates (number of infected plants over number of inoculated plants) at 15 and 30 days post inoculation (dpi) are detailed in Table S1. Based on these results, Fig. 2 presents the distribution of the numbers of infected plants per genotype at 30 dpi globally for the nine PVY isolates used. On the BB16 susceptible control, systemic viral accumulation was detected in 94% of inoculated plants at 15 dpi and in 100% of plants at 30 dpi, regardless of the PVY isolate used (Table S1). On the opposite, the resistant LD VAM and TN86 accessions displayed the lowest proportion of plants showing PVY accumulation. Indeed, at 15 dpi PVY accumulation could not be detected in a single inoculated plant, while at 30 dpi only 2% and 5% of the plants showed detectable PVY accumulation for VAM and TN86, respectively (Table S1). The two other varieties with the large deletion, Wisslica and PBD6, displayed a slightly higher proportion of plants with PVY accumulation. Accessions of the SD group displayed even higher frequencies of infected plants (Fig. 2), with a median between six and seven plants, except for the variety Little C (median of 11 plants). Finally, the groups showing the
most variable results between PVY isolates and the highest frequency of infected plants correspond to the FS group (Burley DC, Semoy and Skro.L56 accessions) and the two EMS mutants (Table S1). Taken together, these results show that plants carrying a mutation affecting elf4e-1 vary greatly in their behaviour towards PVY, ranging from very efficient resistance for LD genotypes to very ineffective resistance for FS and EMS genotypes (Fig. 2).

In order to confirm that within the LD group, VAM and TN86 display a higher resistance efficiency than Wislica and PBD6, a larger number of plants of the four LD genotypes were challenged in three independent experiments by two PVY isolates, SN3 and O139 (selected because they showed the highest infection rate in all challenged tobacco accessions). The EMS1 mutant and the susceptible BB16 genotype were used as controls. Table 1 shows that at 15 dpi, the four LD genotypes displayed significantly higher resistance efficiency than EMS1 and BB16, with 0% to 32% of infection for the LD genotypes, compared to 57% to 93% of infection for EMS1 and BB16, respectively (Table 1). At 30 dpi, there was no significant difference in the infection rates on PBD6 and EMS1 genotypes for the two PVY isolates, varying from 46% to 78%. The VAM genotype displayed significantly lower percentages of infection (1–2%) than the other LD genotypes (Table 1). Depending on the isolate, TN86 and Wislica displayed a significantly higher level of resistance than PBD6, confirming our previous observations showing that the efficiency of the resistance differs between LD genotype, with the VAM genotype displaying a significantly lower percentage of infection (Fig. 2).

We challenged one tobacco genotype representative of each group (VAM for LD, Elka 245 for SD, Skro.L56 for FS and EMS1) with a larger collection of PVY isolates representative of the O and C clades. Table S2 shows that overall the resistance effectiveness is still statistically higher in VAM than in the other tested genotypes, with only nine infected plants out of 192 inoculated plants (5%) when taking into consideration all PVY isolates. The three other tested tobacco accessions, representative of the LD, FS and EMS mutant groups provided very similar results, with overall infection rates of 83–88%, which are not statistically different (Table S2). As before, 100% infection was recorded in BB16 used as a susceptible control.

### Infection of va tobacco genotypes is associated with the emergence of RB variants

Previously, Masuta et al. (1999) and Lacroix et al. (2011) identified amino acid changes in the central part of the VPg of PVY that were associated with the ability to overcome the va resistance. In the present study, we suspected that the varying number of infected plants, increasing with time of infection, corresponds to the emergence of RB variants. To confirm this hypothesis, the nucleotide sequence of the region encoding the central portion of the VPg was determined for a number of infected plants representing the vast majority of the viral isolate/tobacco genotype combinations tested. In total, the sequence of 91 independent viral progenies was determined for PVYN isolates propagated in va tobacco accessions (Tables 2 and S3) and for 90 PVYβ progenies (Tables 2 and S4). The sequences obtained were then compared with the sequence of the corresponding inoculum. In the susceptible BB16 control accession, no mutation was observed in the 40 progenies analysed, irrespective of the PVY isolate (Tables S3 and S4). On the va accessions, a very different situation was

#### Table 1  Response of the four LD va-genotypes, EMS1 mutant and susceptible BB16 tobacco to PVY isolates O139 and SN3.

| PVY isolate | Experiment | VAM 15 dpi | VAM 30 dpi | TN6 15 dpi | TN6 30 dpi | Wislica 15 dpi | Wislica 30 dpi | PBD6 15 dpi | PBD6 30 dpi | EMS1 15 dpi | EMS1 30 dpi | BB16 15 dpi | BB16 30 dpi |
|-------------|------------|------------|------------|------------|------------|----------------|----------------|------------|------------|------------|------------|------------|------------|
| PVYO-O139   | 1          | 0/30       | 1/30       | 2/30       | 3/30       | 6/30          | 8/30          | 2/30       | 17/30      | 9/30       | 10/30      | 10/13      | 13/13      |
|             | 2          | 0/30       | 0/30       | 3/30       | 16/30      | 15/30         | 19/30         | 16/30      | 29/30      | 30/30      | 30/30      | 15/15      | 15/15      |
|             | 3          | 1/30       | 1/30       | 0/30       | 2/30       | 2/30          | 10/30         | 2/30       | 16/30      | 27/30      | 30/30      | 15/15      | 15/15      |
|              | Infection rate (%) | 1<sup>a</sup> | 2<sup>a</sup> | 6<sup>a</sup> | 23<sup>b</sup> | 26<sup>b</sup> | 41<sup>b</sup> | 22<sup>b</sup> | 69<sup>c</sup> | 73<sup>c</sup> | 78<sup>c</sup> | 93<sup>c</sup> | 100<sup>1</sup> |
| PVYO-SN3    | 1          | 0/30       | 0/30       | 0/30       | 0/30       | 1/30          | 1/30          | 1/30       | 2/30       | 0/30       | 0/30       | 10/13      | 10/13      |
|             | 2          | 0/30       | 0/30       | 7/30       | 17/30      | 22/30         | 24/30         | 22/30      | 25/30      | 30/30      | 30/30      | 15/15      | 15/15      |
|             | 3          | 0/30       | 1/30       | 0/30       | 3/30       | 3/30          | 8/30          | 6/30       | 14/30      | 21/30      | 30/30      | 15/15      | 15/15      |
|              | Infection rate (%) | 0<sup>c</sup> | 1<sup>c</sup> | 8<sup>c</sup> | 23<sup>b</sup> | 29<sup>c</sup> | 37<sup>c</sup> | 32<sup>c</sup> | 46<sup>c</sup> | 57<sup>c</sup> | 67<sup>c</sup> | 93<sup>c</sup> | 93<sup>c</sup> |

<sup>1</sup>Number of infected plants/number of inoculated plants for each independent experiment (1, 2 or 3). Infection was analysed by ELISA at 15 and 30 dpi. Multiple chi-squared tests for pairwise comparisons were performed using the R software v. 3.2.5. For each PVY isolate, the infection rates labelled with the same letter (lower case or upper case for infection rates observed at 15 or 30 dpi, respectively) are statistically identical.
observed. For the five PVY isolates belonging to the N clade, mutations in the central portion of the VPg were observed in all progenies, with amino acid substitutions affecting the lysine (K) at position 105 observed in 66 of the 91 progenies (72%, Tables 2 and S3). This position has previously been identified as a major RB determinant (Janzac et al., 2014; Masuta et al., 1999). In addition to VPg residue 105, point mutations at positions 101, 108, 109 and 119 were also observed, alone or in combination and always at much lower rates (Table 2 and Table S3). This position has previously been identified as a major RB determinant (Janzac et al., 2014; Masuta et al., 1999). In addition to VPg residue 105, point mutations at positions 101, 108, 109 and 119 were also observed, alone or in combination and always at much lower rates (Table 2 and Table S3). Positions 101, 108 and 109 have also been previously reported as being associated with va resistance breaking (Janzac et al., 2014; Lacroix et al., 2011). For the four PVYO isolates, a total of 65 out of 90 analysed progenies (72%, Tables 2 and S4) displayed mutations in the central portion of the VPg, with again mutations at residue 105 being by far the most frequent (64%, Table 2). A few viral variants were also observed with mutations affecting VPg residues 101, 119, 120 and 121 (Tables 2 and S4). However, despite the presence of symptoms and detectable viral accumulation, no VPg mutation was observed in 28% of the PVY0 progenies coming from different va genotypes, and in particular in four EMS2 mutant progenies (Tables 2 and S4). On further propagation on EMS2 plants of two of these first passage progenies (derived, respectively, from the SN3 and LA4 isolates) a high infection rate was noted (92%) and a mutation at position 105 of the VPg was detected in these second passage progenies, suggesting that RB variants were probably already present, although in low proportion, in the first passage progenies. Altogether, these results indicate that the PVY infections observed on the different va tobacco accessions correspond to the emergence of RB variants having mutations at already identified key positions in the central region of the VPg. As we observed a high differential in RB frequency between tobacco accessions with the PVYO isolate, it was chosen to evaluate the durability of the va resistance in later experiments.

### Table 2 Amino acid substitutions identified in the VPg central region (amino acids 101–123) of the progenies of PVYN and PYYO isolates after passaging in the 13 different va tobacco accessions.

| Amino acid change       | Frequency |
|-------------------------|-----------|
|                         | PVYN variants | PVYO variants |
| K105E, T, Q, M          | 73% (66/91) | 64% (58/90) |
| K105T + G119H           | –          | 1% (1/90)   |
| S101G                   | 24% (22/91) | 4% (4/90)   |
| S101G + V108I           | 1% (1/91)  | –           |
| V108I + G119C           | 1% (1/91)  | –           |
| E109D                   | 1% (1/91)  | –           |
| S120R                   | –          | 1% (1/90)   |
| N121Y                   | –          | 1% (1/90)   |
| No mutation identified  | –          | 28% (25/90) |
| TOTAL                   | 91         | 90          |

The amino acid positions are numbered according to the VPg sequence of PVY-N605 (GenBank X97895) and PVY-O (GenBank U09509) for PVYN and PVYO isolates, respectively. Single and double mutations are listed. The nomenclature ‘K105E’ indicates the substitution of a lysine (K) by a glutamic acid (E) at position 105. For single mutations at position 105, the four substitutions observed (K105E, K105T, K105Q or K105M) are regrouped. C, cysteine; D, aspartic acid; E, glutamic acid; G, glycine; H, histidine; I, isoleucine; K, lysine; M, methionine; N, asparagine; Q, glutamine; R, arginine; S, serine; T, threonine; V, valine; Y, tyrosine. –, no mutation detected in the progenies at this position.

A complex eIF4E locus impacts va durability

We believe that the differences in va durability could be associated with genetic background effects and redundancy between eIF4E family members. We therefore decided to analyse the eIF4E family members to see if they might correlate with resistance durability. Tobacco is an allotetraploid derived from the diploids *N. sylvestris* and *N. tomentosiformis*, which are, respectively, susceptible and resistant to PVY. Sequences annotated as eIF4E or its isoforms elfiso4E and novel cap-binding protein (nCBP) have been identified in the transcriptomes of the two parental genomes and of tobacco (Julio et al., 2015). Among the six eIF4E copies, four are derived from the *N. tomentosiformis* (T) genome, T015277, T021658, T021287, and T025160, and two from the *N. sylvestris* (S) genome, S10760 and S05588. In the present study, eIF4E-1 stands for the S10760 gene (va locus) and eIF4E-2, -3, -4, -5 and -6 for the T021658, T025160, T021287, S05588 and T015277 genes, respectively (Fig. 3). The Blast function available on the Sol Genomics Network server (https://solgenomics.net/) (Edwards et al., 2017) showed a localization of the eIF4E-2, 3 and 4 genes on chromosome 14 of the tobacco genotype K326 (PI552505), starting, respectively, at positions 109418388, 109399608 and 109292645, indicating these genes are located within a ca. 132 kbp region (Fig. S1). The short distance between eIF4E-2 and eIF4E-3 was confirmed using simple sequence repeat (SSR) markers (Bindler et al., 2011) (see experimental procedures). Therefore, all three genes can be considered as a single large locus on chromosome 14. The two other eIF4E-5 and eIF4E-6 copies map on chromosome 17 (positions 79984931 and 70652013, respectively). Chromosome 17 may have undergone rearrangements between the *N. tomentosiformis* and *N. sylvestris* genomes (Edwards et al., 2017), which may explain the simultaneous presence of eIF4E-5 gene (S05588, 5-genome origin) and eIF4E-6 (T015277, T-genome origin).

The analysis of RNA-seq data (Julio et al., 2015) failed to identify amino acid polymorphisms in any of the six eIF4E copies between accessions with durable or unstable va resistance.
vips. Indeed, only a partial coverage of the eIF4E-3 gene was observed when mapping reads from the VAM, Start and Sk70 LD accessions, while complete coverage was observed for the other LD accessions (TN86, Wislica and PBD6) in the EMS1 mutant and in BB16 (Fig. S3).

A specific amplification of eIF4E-3 with the forward primer T025160F6 (Table S5) designed in the seemingly 5’ missing region in VAM and the reverse primer T025160R6-2 (expressed region) was performed on the genomic DNA of a panel of tobacco accessions. Compared to other accessions (i.e. TN86, Wislica, PBD6, TN90, EMS1 mutant and BB16) from which the expected polymerase chain reaction (PCR) product of 324 bp was readily amplified, no amplification product was observed in VAM, Start and Sk70, confirming the existence of a genomic deletion affecting the 5’ region of eIF4E-3 in these accessions. This was further confirmed by high-throughput sequencing of the VAM genomic DNA, which revealed instead the presence of a hybrid eIF4E copy, with the 5’ part of eIF4E-2 fused to the 3’ part of eIF4E-3, hereafter named eIF4E-2-3 (Fig. S4). This peculiar gene structure was confirmed by PCR in the VAM, Sk70 and Start genotypes (Fig. 4) using the various primer pairs shown in Fig. 5. Surprisingly, two other LD accessions, TN86 and TN90, were shown to contain both eIF4E-2-3 and an entire eIF4E-3 gene (Fig. 4).

Altogether, those results confirm the presence of a complex eIF4E locus on chromosome 14 (hereafter named Chr14-eIF4E) differing between durable LD and non-durable LD accessions.

**Genetic analysis indicates that va durability is influenced by the type of mutation at the eIF4E-1 locus and by the complex Chr14-eIF4E locus**

We first checked whether the eIF4E-2-3 allele at the complex Chr14-eIF4E locus co-segregates with resistance durability. The F1 progeny of a cross between VAM and EMS1 showed a 12% infection rate at 30 dpi, comparable to that of VAM (Table 3), showing that the resistance durability trait is dominant. The pheno-typing of 256 F2 progeny plants confirmed that the eIF4E-2-3 allele co-segregates with resistance durability. Using the same F2 progeny, we also checked whether the type of mutation at the eIF4E-1 locus [large deletion (eIF4E-1LD) or point mutation (eIF4E-1KO)] influences resistance durability. Indeed, homozygous eIF4E-1LD/eIF4E-1LD and heterozygous eIF4E-1LD/eIF4E-1KO F2 plants display infection rates comparable those of VAM and F1 hybrids, respectively (Table S6). Interestingly, the homozygous eIF4E-1KO/eIF4E-1KO F2 progenies display a significantly higher level of resistance durability than the EMS1 parent, with infection rates of 47% and 98%, respectively. This difference may reflect the influence of the locus Chr14-eIF4E.

**Overexpression of the eIF4E-2 copy in the LD genotypes correlates with a higher durability of va-mediated resistance**

An initial analysis of RNA-seq data suggested that eIF4E-2 (T021658) was more highly expressed in the resistant VAM accession (Julio et al., 2015), suggesting the possible existence of an expression compensation mechanism between eIF4E orthologs derived from the two parental genomes in the
amphidiploid tobacco. The results of a more extensive RNA-seq analysis on a range of tobacco accessions carrying different types of mutations affecting the eIF4E-1 gene are shown in Table 4. They show that for all seven accessions with the large genomic deletion a higher expression of eIF4E-2 is observed as compared to the other va genotypes. It is also worth noting that among the LD accessions, three (VAM, Sk70 and Start) show a particularly striking overexpression (Table 4). Real-time PCR confirmed these RNA-seq results, using two different housekeeping genes for normalization (Fig. 6). The expression level of the eIF4E-2 gene was systematically increased by a factor 4 to 7 as compared to BB16 in TN86 and VAM, respectively, and by a factor of 1.5 in Wislica and PBD6. In contrast, there was no significant overexpression in the EMS1 mutant (Fig. 6A). The same trend was observed using another housekeeping gene (EF1-α) as reference (Fig. 6B). Overall, both RNA-seq data and quantitative reverse transcriptase-polymerase chain reaction (Q-RT-PCR) indicate that eIF4E-2 overexpression is significantly stronger in VAM and TN86 as compared to Wislica and PBD6. The high overexpression level detected by RNA-seq for Start and Sk70 was also confirmed by real-time PCR, while TN90 showed an intermediate behaviour (Fig. 7A,B). The RNA-seq data show the FS mutants and the small deletion accessions to behave similarly to EMS1 and BB16 in this respect. Taken together, these results demonstrate the existence of a very good correlation between resistance durability and the expression level of eIF4E-2. This correlation is further reinforced by the observation that, similar to VAM, the Sk70 and Start accessions show the highest level of eIF4E-2 overexpression as well as a very high level of resistance durability when confronted with the PVY-O139 isolate (Fig. 7C). Conversely, with an intermediate eIF4E-2 overexpression level, TN90 displayed an intermediate resistance durability (Fig. 7C).

DISCUSSION

The durability of the resistance differs largely among va-tobacco accessions

Here, we compared the durability of the resistance to PVY in 13 va tobacco accessions differing by their genetic background and the type of mutations affecting the eIF4E-1 gene (va locus),
## Table 3
The 5′ deletion of eIF4E-3 gene at the locus eIF4E-2-3 on chromosome 14 co-segregates with the resistance durability character.

| Genotype | Phenotype | eIF4E-1 locus | eIF4E-3 locus | P value (χ²) |
|----------|-----------|---------------|---------------|-------------|
| VAM      | eIF4E-1LD/eIF4E-1LD | eIF4E-2-3/eIF4E-2-3 | 6% (2/33) | |
| EMS1     | eIF4E-1KO/eIF4E-1KO | eIF4E-3/eIF4E-3 | 98% (32/33) | |
| F1       | eIF4E-1LD/eIF4E-1KO | eIF4E-3/eIF4E-2-3 | 12% (4/33) | 0.16 $ |
| F2       | eIF4E-1LD/eIF4E-1LD | eIF4E-3/eIF4E-2-3 | 5% (3/59) | 1 † |

The allele eIF4E-2-3 corresponds to the hybrid form eIF4E-2-3 on chromosome 14 as present in the VAM accession. The allele eIF4E-3 corresponds to the complete eIF4E-3 gene seen in most tobacco accessions. The allele eIF4E-1LD corresponds to the large deletion on chromosome 21, which results in the complete deletion of the eIF4E-1 gene as seen in the VAM accession. The allele eIF4E-1KO corresponds to the point mutation seen in the EMS1 mutant and leads to a C-terminally truncated eIF4E-1 proteins of 50 aa.

## Table 4
RNASeq expression analysis of all eIF4E orthologs in the four groups of resistant accessions.

| Genotype | Accession | eIF4E-1 | eIF4E-2 | eIF4E-3 | eIF4E-4 | eIF4E-5 | eIF4E-6 |
|----------|-----------|---------|---------|---------|---------|---------|---------|
| Large deletion | VAM      | 0.4     | 173.1   | 4.0     | 4.9     | 16.9    | 20.0    |
|          | TN86     | 1.0     | 50.9    | 7.4     | 2.5     | 8.5     | 13.2    |
|          | Wislica  | 0.0     | 34.2    | 34.5    | 2.0     | 18.2    | 21.0    |
|          | PB06     | 0.0     | 38.3    | 32.1    | 0.0     | 22.0    | 32.6    |
|          | Sk70     | 2.3     | 147.5   | 8.5     | 0.0     | 17.5    | 6.5     |
|          | Start    | 0.4     | 107.8   | 2.4     | 2.7     | 19.8    | 15.5    |
|          | TN90     | 0.0     | 78.5    | 18.5    | 1.9     | 15.4    | 12.7    |
| Small deletion | Elka     | 0.4     | 11.3    | 11.8    | 2.1     | 14.2    | 15.6    |
|          | Little C. | 0.0     | 19.2    | 13.4    | 1.2     | 16.8    | 14.9    |
|          | Philippin| 0.0     | 17.6    | 13.0    | 1.2     | 22.6    | 14.6    |
|          | Wika     | 0.0     | 13.9    | 14.4    | 2.4     | 16.6    | 9.1     |
| Frameshift | Burley DC | 4.7     | 18.2    | 12.4    | 0.8     | 14.3    | 14.8    |
|          | Semoy    | 5.5     | 19.6    | 11.7    | 1.5     | 16.6    | 18.1    |
|          | Skro. L56| 4.6     | 17.8    | 9.8     | 1.0     | 17.7    | 18.1    |
| EMS mutant | EMS1     | 0.4     | 21.4    | 15.5    | 3.2     | 16.3    | 18.5    |

Normalized expression levels were obtained from RNA-seq transcriptome analysis for uninfected tobacco accessions representative of the four categories of resistant plants (Julio et al., 2015). Data were analysed with the 'Set Up Experiment' function and 'two groups comparison' by comparing characterized resistance to PVY (resistant versus susceptible). Data were normalized with the 'Normalize' function using default parameters (Method: Scaling; Values: Original Expression Value; Normalization value: Mean, CLC Genomics Workbench v5.5 software). Normalized data for each contig are shown.
such as genomic deletions smaller than the large 1 Mbp found in VAM, natural frameshift mutations and EMS nonsense mutations. The 13 tobacco genotypes were challenged by a panel of PVY isolates representative of the virus diversity, and infection rates were monitored up to 30 days after inoculation to better analyse resistance breakdown. The PVY infections observed on the va genotypes correspond to the emergence of RB variants having, as previously shown, mutations in the VPg central region (Lacroix et al., 2011; Masuta et al., 1999; Moury et al., 2004; Xu et al., 2017). This suggests that in the absence of a functional eIF4E-1 copy, RB PVY variants may use another eIF4E isoform, similar to the situation when TuMV overcomes an eIFiso4E loss-of-function in Arabidopsis through VPg mutations allowing recruitment of eIF4E1 (Bastet et al., 2018; Gallois et al., 2018). However, our data suggest that the situation is even more complex in tobacco, and that besides the potential use of an eIF(iso)4E copy by RB PVY, as suggested by Takakura et al. (2018), at least two other eIF4E copies may play a role in va resistance durability. Owing to their close proximity in the tobacco genome, however, it is not possible to genetically discriminate the potential contributions to durability of the over-expression of eIF4E-2 and the presence of the hybrid eIF4E-2-3 gene (and/or lack of an integral eIF4E-3 copy).

Redundancy among eIF4E isoforms might mediate va resistance durability

A compatible eIF4E isoform is required by potyviruses for their genome translation, replication, stabilization, and intercellular and systemic transport (Contreras-Paredes et al., 2013; Robaglia and Caranta, 2006; Sanfaçon, 2015; Wang and Krishnaswamy, 2012). Some potyviruses can also use several eIF4E isoforms in a given host (Gaufﬁer et al., 2016; Jenner et al., 2010; Mazier et al., 2011; Xu et al., 2017). This suggests that in the absence of a functional eIF4E-1 in va-resistant genotypes, RB PVY variants may use another eIF4E isoform, similar to the situation when TuMV overcomes an eIFiso4E loss-of-function in Arabidopsis through VPg mutations allowing recruitment of eIF4E1 (Bastet et al., 2018; Gallois et al., 2018). However, our data suggest that the situation is even more complex in tobacco, and that besides the potential use of an eIF(iso)4E copy by RB PVY, as suggested by Takakura et al. (2018), at least two other eIF4E copies may play a role in va resistance durability. Owing to their close proximity in the tobacco genome, however, it is not possible to genetically discriminate the potential contributions to durability of the over-expression of eIF4E-2 and the presence of the hybrid eIF4E-2-3 gene (and/or lack of an integral eIF4E-3 copy).

Two genomic regions are involved in the durability of va-mediated resistance in tobacco

Phenotypic, genetic and transcriptomic analyses showed that the type of mutation at the eIF4E-1 locus, together with the functionality and expression levels of other eIF4E copies at a complex locus on chromosome 14, impact va-mediated resistance durability. The eIF4E-2, eIF4E-3 and eIF4E-4 genes are all derived from the T genome and are the closest orthologs of the S genome eIF4E-1 (Fig. 3). The presence of these three closely related genes on chromosome 14 likely results from a triplication event in N. tomentosiformis, with some further chromosomal rearrangement(s) leading in LD accessions to the disruption of eIF4E-3, in some cases to the appearance of eIF4E-2-3 and to an increase in eIF4E-2 copy number.

Our observation that the resistance in VAM is significantly more durable than in most other va accessions confirms the results of Acosta-Leal and Xiong (2008), who observed a rapid selection of RB variants in NC745, but not in VAM. They further suggested that functional components of the resistance were independently controlled by two segregating recessive genes in VAM: va itself (renamed va1) responsible for limiting potyvirus cell-to-cell movement and another locus, va2, limiting virus accumulation. They also suggested that va1 and va2 conditioned the stronger resistance of VAM (i.e. the resistance durability phenotype) and that the functional gene product of va1, va2 or both loci, like most host factors interacting with VPg, might be an eIF4E factor. The fact that va encodes the eIF4E-1 copy was demonstrated by Julio et al. (2015) and the present results strongly suggest that the chr14-eIF4E complex locus could correspond to the va2 locus, encoding gene products that may indeed interact with PVY VPg, as suggested by Acosta-Leal and Xiong (2008).

Redundancy among eIF4E isoforms might mediate va resistance durability

A compatible eIF4E isoform is required by potyviruses for their genome translation, replication, stabilization, and intercellular and systemic transport (Contreras-Paredes et al., 2013; Robaglia and Caranta, 2006; Sanfaçon, 2015; Wang and Krishnaswamy, 2012). Some potyviruses can also use several eIF4E isoforms in a given host (Gaufﬁer et al., 2016; Jenner et al., 2010; Mazier et al., 2011; Xu et al., 2017). This suggests that in the absence of a functional eIF4E-1 in va-resistant genotypes, RB PVY variants may use another eIF4E isoform, similar to the situation when TuMV overcomes an eIFiso4E loss-of-function in Arabidopsis through VPg mutations allowing recruitment of eIF4E1 (Bastet et al., 2018; Gallois et al., 2018). However, our data suggest that the situation is even more complex in tobacco, and that besides the potential use of an eIF(iso)4E copy by RB PVY, as suggested by Takakura et al. (2018), at least two other eIF4E copies may play a role in va resistance durability. Owing to their close proximity in the tobacco genome, however, it is not possible to genetically discriminate the potential contributions to durability of the over-expression of eIF4E-2 and the presence of the hybrid eIF4E-2-3 gene (and/or lack of an integral eIF4E-3 copy).
Our results apparently diverge from the recent study of Takakura et al. (2018) showing that mutating the T-genome eIF(iso)4E reduces the susceptibility to an RB PVY variant. However, these authors use an RB variant and cannot therefore evaluate how the tobacco genotype may affect the frequency with which such variants may appear, as was analysed in our study. The two series of results and conclusions are therefore in fact compatible and even complementary. It remains to be evaluated whether a mutation in eIF(iso)4E would impact the frequency of emergence of RB mutants.

Overall, a tight correlation network seems to link the superior durability seen in LD accessions such as VAM, Sk70 and START, the overexpression of eIF4E-2, the presence of multi-copies of eIF4E-2, the presence of the eIF4E-2-3 recombined gene and the deletion of eIF4E-1. It is not currently possible to order in a cause and effect model these elements but, with its multiple levels, the durability/eIF4E-2 correlation appears to be the most promising. It should be noted that homeostatic control feedback mechanisms between eIF4E family members to compensate loss of function has been demonstrated in plants (Gallois et al., 2018). This was previously reported in Arabidopsis, where eIF4E-1 protein concentration increases to compensate for loss of eIF(iso)4E in the KO-eIF(iso)4E (Duprat et al., 2002). In tobacco cv Samsun, the depletion of eIF(iso)4E by an antisense down-regulation strategy resulted in a compensatory increase in eIF4E protein levels (Combe et al., 2005), while in Brassica rapa, overexpression of an eIF(iso)4E transgene induces the expression of endogenous eIF4E (Kim et al., 2014), revealing another regulation mechanism.

**Fig. 7** Relative quantification of eIF4E-2 gene expression by Q-RT-PCR and resistance level in various LD tobacco accessions. Eif4E-2 transcript levels were normalized to the expression of (A) RL2 (60S ribosomal protein L2) and (B) EF1-α (elongation factor 1 alpha) housekeeping genes, respectively. Gene expression was quantified from two pools of six independent plants. The relative expression level (fold induction) derived from the mean of the two pools by comparing the two housekeeping genes to obtain ΔCt (Ct(eIF4E-2) – Ct(RL2) or EF1-α) and BB16 (susceptible reference) value was used as the calibrator to determine the ΔΔCt and then calculated using the formula 2^-ΔΔCt (as described by Schmittgen and Livak, 2008). The standard deviation between the two pools is indicated by vertical lines. Statistical analysis was performed using the Kruskal–Wallis rank sum test in R software v. 3.2.5. Significantly different values are noted with capital letters (P value ≤ 0.05) when comparing the expression level of eIF4E-2 gene between each genotype. GenBank accession numbers of the genes used: eIF4E-2 (KM202068), RL2 (X62500), EF1-α (XM_009784954). (C) The different tobacco genotypes were challenged with PVY-O139 isolate. †Number of infected plants/number of inoculated plants at 30 dpi. Infection rates correspond to the total number of infected plants/number of inoculated plants for two experiments (statistical analysis was performed using the multi-comparison chi-squared test in R software v. 3.2.5. Infection rates labelled with the same letter are statistically identical (P value < 0.05).
tomato, it has also been suggested that protein–protein interactions between elf4E-1 and elf4E-2 lead to a yet unexplained elf4E-2 degradation (Gauffier et al., 2016). In such a scenario, the deletion of elf4E-1, the loss of elf4E-3 and the presence of elf4E-2 can all contribute, alone or in combination, to the overexpression of elf4E-2.

The elf4E-2 copy could act as a decoy increasing va resistance durability

In the scenario depicted in Fig. 8, PVY preferentially uses elf4E-1 to complete its infectious cycle in susceptible tobacco. In the resistant va accessions, either elf4E-1 is absent (LD and SD genotypes) or a non-functional C-terminally truncated elf4E-1 is expressed (FS and EMS genotypes). The absence of elf4E-1 or the failure of its truncated form to bind the viral VPg, a step that appears to be necessary although not sufficient to establish viral infection, could represent the key element in the va-mediated resistance. However, the resistance is more durable in the LD genotypes because of the less frequent emergence of RB mutants. Assuming these RB mutants do not pre-exist in the inoculum, their emergence requires a minimal level of replication, possibly involving imperfect interaction with (an)other isoform(s) such as elf(iso)4E-T (Takakura et al., 2018). One hypothesis to explain va durability is that PVY VPg could also interact in a non-productive way with other isoforms, in particular with elf4E-2. The overexpression of elf4E-2 observed in LD accessions and exacerbated in accessions lacking elf4E-3 and carrying elf4E-2 could monopolize the VPg in a non-functional interaction, limiting the possibility for the virus to evolve towards resistance breaking.

Consistent with this model, our results show that the va resistance durability trait is dominant and that elf4E-2 strongly differs from elf4E-1 in the loop 1 region close to the cap binding pocket (Fig. S4A), which contains amino acid sites known to be crucial for susceptibility to potyviruses (German-Retana et al., 2008; Robaglia and Caranta, 2006). All these elements support the model that overexpression of elf4E-2 would act in a dominant negative manner (Chandler, 2003) similar to the resistance afforded through the transgenic overexpression of recessive resistance alleles in a susceptible background (Cavatorta et al., 2011; Kang et al., 2007). Other hypotheses are also possible, for example a direct involvement of the elf4E-2 copy, also acting as a decoy.

Despite its wide use, several adverse effects of the va gene have been observed. The resistant lines tend to have short and narrow leaves, and lower productivity (Noguchi et al., 1999). This may reflect the presence of important genes in the large deletion containing the va gene. At the same time, this deletion may have negative effects on recombination during backcrosses and could impose a linkage drag, preventing restoration of a normal phenotype during breeding efforts (Yamamoto, 1992). Efforts have

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**Fig. 8** Model of elf4E-mediated va resistance durability in tobacco. In tobacco, the main elf4E copy used by PVY is elf4E-1. In the resistant va accessions, either elf4E-1 is absent (crossed-out) or a mutant C-terminally truncated elf4E-1 protein is expressed (star). The appearance of RB variants, carrying mutations in the central region of the VPg (star), suggests that those RB variants can recruit another elf4E factor for their infection cycle in the absence of the main susceptibility factor elf4E-1 or in the presence of a C-terminal truncated protein. This factor can be elf(iso)4E-T or elf4E-2 or elf4E-3. The solid arrows represent productive recruitment for the viral infection whereas the dotted arrows represent unproductive recruitment. The resistance is more durable when elf4E-1 is absent (LD accessions). This could be explained by the overexpression of elf4E-2 copy in the absence of elf4E-1, which is even exacerbated in the absence of elf4E-3 and presence of the hybrid elf4E-2. The PVY or RB-PVY variants could not use the elf4E-2 copy, which, when more abundant than other endogenous elf4E copies, would monopolize translation initiation machinery. A surplus of elf4E-2 protein presumably makes ribosomal entry unavailable for the other endogenous elf4E copy that may be used by the RB virus or could also monopolize the VPg in a nonfunctional interaction, limiting the possibility for the virus to evolve towards resistance breaking, leading to higher resistance durability.
been developed to find new markers for background selection in backcross breeding programmes (Tajima et al., 2002) but these have been incompletely successful in allowing the reduction of the negative effects of va. The second locus contributing to resistance durability, chr14-elf4E, could be transferred by marker-assisted selection (MAS) in elite lines. A strategy to achieve a fully durable resistance could be to combine in the same plant a complete deletion of elf4E-1 and elf(iso)4E-T with the chr14-elf4E locus. As new breeding techniques become also available for genome editing (Abdallah et al., 2015; Andersen et al., 2015), such approaches could be envisioned for introducing sequence-specific deletions in the elf4E-1 and elf(iso)4E-T genes in a chr14-elf4E background, opening new perspectives to improve our ability to manipulate and deploy durable tobacco resistance against one of its major viral pathogens, Potato virus Y.

EXPERIMENTAL PROCEDURES

Plant material

The PVY-susceptible N. tabacum cv. Xanthi was used to produce viral inoculum for biological characterization experiments. The 163 accessions used for RNA-seq analysis belong to the Imperial Tobacco LTD collection and were described in Julio et al. (2015) as well as the elf4E-1 EMS stop codons mutants EMS1 and EMS2 corresponding to W50* and W53* mutations (with tryptophan amino acid at position 50 or 53 in the sequence replaced by a stop codon). Another elf4E-2 mutant (named E3-762) was obtained from the EMS collection described in Julio et al. (2015). Among the tobacco accessions of known PVY resistance status previously characterized using S10760-linked markers as well as other markers associated with the deletion affecting the va genomic region, the VAM, TN86, Wiliska, PBD6, START, SK70 and TN90 accessions display a va-resistance allele associated with a large deletion on chromosome 21 (Julio et al., 2015). In this paper, those accessions belong to the LD group (Fig. 1). In two dark air-cured (Little C. and Philippin) and two flue-cured (Wika and Elka 245) tobaccos, the S10760 (elf4E-1) gene could not be amplified, which suggests that it is either deleted or mutated in a way that prevents its amplification (Julio et al., 2015). However, the other markers linked to va are not deleted in these plants, which suggests that they carry a smaller form of the deletion affecting va. Those accessions are therefore classified in this study in the SD group. The last group of accessions concerns resistant plants in which the S10760 elf4E-1 gene is present and which do not appear to have a deletion on chromosome 21. However, a 2-bp deletion in the S10760 gene has been identified in two dark air-cured varietal types (Semoy and Skro. L56) and in one burley type (Burley DC), which would truncate the encoded protein and explain the observed resistance, as for the two EMS mutants analysed in Julio et al. (2015). In this study, those three accessions are classified into the FS group (Fig. 1). Healthy and infected plants were maintained in a separate insect-proof greenhouse compartment (18/25 °C night/day) or in a climate chamber (18/20 °C, 8 hours night/16 hours day).

Viral material

The PVY isolates used to challenge the different tobacco accessions belong to the clade N (CSA1, LA7, CSA6, MaSan4, 1108) and the clade O (O139, LA4, SN3 and SAV8), depending on their ability to induce (N) or not (O) systemic vein necrosis symptoms in tobacco (Jakov et al., 1997; Moury, 2010; Singh and Singh, 1996) (Table S1). Those isolates were selected according to their pathogenic and biological properties in tobacco after a field survey carried out in tobacco plots in France in 2007 (Janzac et al., 2014; Lacroix et al., 2010). The other PVY isolates listed in Table S2 belong to the C clade, more specifically to the C1 (SON41, Alger1, Marti13, LYE84.2, CA1A57, CA1A41, CA1A6, LYE72-Puc2P13) or C2 (Cadgen, LYE90v) groups. Some isolates are PVY recombinants between O and C clades (CA1A56, LYE245), others belong to the groups PVY \textsuperscript{NTH} (NTN-H) and PVY \textsuperscript{WI} (WilgaP) (Glais et al., 2002). PVY isolates belonging to the N clade are Pologne 6 puc3 Pl2 and N605, and one isolate comes from Brazil, Brasil 1054. Some of the isolates are referenced in (Ben Khalifa et al., 2012; Moury, 2010; Moury et al., 2004, 2011; Woloshuk et al., 1993).

Virus inoculation and detection

Each PVY isolate was first propagated into N. tabacum cv Xanthi plants in order to prepare viral inoculum. Fifteen days after inoculation, the systemically infected apical leaves (~12 g) were ground in liquid nitrogen and the powder stored at -80 °C. Total RNAs were extracted from 1 mg of powder, using the SV Total RNA Isolation System (Promega, Madison, USA) extraction kit. The viral RNAs were then quantified by real-time PCR on a light cycler 480 (Roche, Bâle, Switzerland) using the AgPath-ID kit. The viral inoculum was then calibrated at a concentration of 10^8 copies of viral RNA/100 µL in the inoculation buffer (50 mM Na_2HPO_4, 50 mM KH_2PO_4, supplemented with 40 mM sodium diethylthiocarbamate, pH = 7.2).

Tobacco plants with two fully expanded true leaves were inoculated manually approximately 2 weeks after sowing. The two last developed leaves were inoculated with 100 µL of the calibrated viral inoculum supplemented with carborundum. Evaluation of PVY infection in non-inoculated leaves was performed by DAS enzyme-linked immunosorbent assay (ELISA) (Clark and Adams, 1977; Lacroix et al., 2010) at 15 and 30 dpi. Absorbance values at 405 nm (A_{405}) with the background subtraction of buffer samples were considered for analysis. Samples were considered positive when their A_{405} was higher than three times the mean A_{405} of non-inoculated samples. Polyclonal antibodies detecting all PVY isolates were used.
isolates, monoclonal antibodies specific for PVY\textsuperscript{N} isolates (kindly provided by Maryse Guillet, INRA-FN3PT, Le Rheu, France) and monoclonal antibodies specific for PVY\textsuperscript{O/C} isolates (Neogen Ltd, Lansing, Michigan, USA) were used.

Analysis of VPg sequence in the viral progenies

The viral progenies present in infected plants were amplified by immuno-capture RT-PCR (IC-RT-PCR) as described previously (Glais et al., 1998) using the 3′ NTR-reverse primer for the production of cDNA and primer pair RJ2-F and Nla-R (Table S5) to amplify the VPg region. Thermal cycling conditions were as follows: 2 min at 95 °C followed by 40 cycles of 1 min at 94 °C, 1 min at 57 °C and 1 min at 72 °C, followed by an elongation step of 10 min at 72 °C. Each PCR product was sent to Genoscreen (Lille, France) for sequencing.

Analysis of eIF4E-2 gene expression by quantitative RT-PCR

Plant samples were ground with a mixer mill (Retsch, Haan, Germany) at −80 °C and total RNAs were isolated using the NucleoSpin® RNA plant Kit (Macherey Nagel, Düren, Germany). The RNAs obtained were treated with TURBO DNA-free™ Kit (Invitrogen, Thermo Fischer Scientific, Waltham, Massachusetts, USA) and the concentration and purity were determined by measuring absorbance at 230, 260 and 280 nm in a microplate UV-Vis spectrophotometer (EPOCH™ BioTek instrument, Winooski, Vermont, USA). The total RNA was adjusted to 50 mg/mL and was reverse transcribed according to the manufacturer’s instructions using the RevertAid H Minus enzyme (Thermo Scientific) and oligo(dT) primers. The cDNA was used to perform the real-time Q-RT-PCR on the Light Cycler® 480 Instrument II (Roche, Bâle, Switzerland), using the Light Cycler®480 SYBR Green I MASTER Kit (Roche, Bâle, Switzerland), using 2 min at 95 °C followed by 40 cycles of 1 min at 94 °C, 1 min at 57 °C and 1 min at 72 °C, followed by an elongation step of 10 min at 72 °C. Each PCR product was sent to Genoscreen (Lille, France) for sequencing.

Copy number analysis for eIF4E-2 and eIF4E-3 and eIF4E-2-3

In order to evaluate the copy number of genomic DNA targets for eIF4E2, eIF4E3 and eIF4E-2-3, we used the TaqMan® Copy Number Assay based on gold-standard TaqMan MGB probe chemistry. TaqMan® Copy Number Assays are run together with a TaqMan Copy Number Reference Assay in a duplex qPCR reaction; the copy number assay detects the target sequence and the reference assay detects a sequence that is known to be present in two copies in the diploid genome. Here, the reference sequence was the nitrate reductase gene (GenBank XM_016606716) for all 3′ regions common to eIF4E-2 and eIF4E-2-3 amplified in a multiplex reaction with the primers E2-E2E3-F and E2-E2E3-R and the probe E2-E2E3 (Fig. 6, Table S5). The region specific to eIF4E-2 was amplified in multiplex with the primer pair 4E2-F and 4E2-R and the probe 4E2 (Table S5). The 3′ region specific to eIF4E-3 was amplified in a multiplex reaction with the primer pair E3-E2E3-F and E3-E2E3-R and the probe E3-E2E3
(Fig. 6, Table S5). All PCR reactions were carried out in a 10 μL volume containing 5 ng of DNA, 1x TaqPath™ ProAmp™ Master Mix (Applied Biosystems™, Foster City, California, USA), 900 nM of each primer and 250 nM of each probe. The amplification was conducted using a QuantStudio-3 Real-Time PCR (Applied Biosystems, Foster City, California, USA) as follows: 95 °C for 10 min, 40 cycles of 95 °C for 15 s and 60 °C for 1 min. Results were analysed by the relative quantitation method using CopyCaller® Software v2.0 (Applied Biosystems, Foster City, California, USA).

Sequencing of tobacco VAM accession genomic DNA

VAM DNA was extracted independently from eight plantlets using a Qiacube Dnaseasy kit, checked for concentration and mixed together. Libraries were prepared with an Illumina TruSeq PCR-free kit and sequenced on one lane of HiSeq 3000 (paired-end together. Libraries were prepared with an Illumina TruSeq PCR-free kit and sequenced on one lane of HiSeq 3000 (paired-end to-}

Screening for mutation in the eIF4E-2 gene

A new EMS mutant collection was developed from seeds of the TN90 resistant variety as described in Julio et al. (2008) and used to detect eIF4E-2 mutants. Specific primers for eIF4E-2 gene were designed to amplify an exon region in each gene. Primers T021658E6TF and T021658E3T2R (Table S5) were used to amplify an amplicon of 329 bp of eIF4E-2. PCR amplification was carried out with FAM and VIC dye-labeled primer pairs in a 10 μL volume containing 2 μL of 10 × AmpliTaq® Polymerase (Applied Biosystems, Foster City, California, USA), 1 μL of AmpliTaq buffer, 0.5 μL of dNTPs (Applied Biosystems, 2.5 mM each) and 7 ng of each primer. PCR was conducted using a thermal cycler (GeneAmp® PCR System 9600, Applied Biosystems) as follows: 28 to 32 cycles of 94 °C for 30 s, 62 °C for 45 s and 72 °C for 1 min, followed by 7 min at 72 °C for final extension. eIF4E-2 PCR products were analysed by capillary electrophoresis-single strand conformation polymorphism and sequenced as described in Julio et al. (2008).

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**SUPPORTING INFORMATION**

Additional supporting information may be found in the online version of this article at the publisher’s web site:

Fig. S1 Position of the eIF4E-2, eIF4E-3 and eIF4E-4 genes on the *N. tabacum* (var. K326; PI552505) chromosome 14.

Fig. S2 Amino acid sequence alignment of the various eIF4E proteins of *Nicotiana tabacum*.

Fig. S3 Mapping of the RNASeq reads against the eIF4E-3 consensus sequence for the LD accessions VAM, TN86, Wislica, PBD6, Sk 70 and Start, the EMS1 mutant and the susceptible BB16 accession.

Fig. S4 Amino acid sequence polymorphism between eIF4E 1, eIF4E 2, eIF4E 3 and the hybrid eIF4E 2-3 copy present in some LD genotypes of *Nicotiana tabacum*.

Table S1 Response of *va* tobacco genotypes to PVY isolates.

Table S2 Response of representative *va* tobacco genotypes to various PVY isolates.

Table S3 Amino acid changes in the VPg central region (amino acids 101–123) of the progenies of five PVY° isolates, following infection of 13 different *va* tobacco accessions, in comparison with the sequence of the parental isolates.

Table S4 Amino acid changes in the VPg central region (amino acids 101–123) of the progenies of four PVY° isolates, following infection of 13 different *va* tobacco accessions, in comparison with sequence of the parental isolates.

Table S5 List of primers and probes used in this study.

Table S6 The deletion at the locus eIF4E-1LD on chromosome 21 co-segregates with the resistance durability character.