Predicting the Stability of Homologous Gene Duplications in a Plant RNA Virus
Anouk Willemsen, Mark Zwart, Pablo Higueras, Josep Sardanyés, Santiago Elena

To cite this version:
Anouk Willemsen, Mark Zwart, Pablo Higueras, Josep Sardanyés, Santiago Elena. Predicting the Stability of Homologous Gene Duplications in a Plant RNA Virus. Genome Biology and Evolution, Society for Molecular Biology and Evolution, 2016, 8 (9), pp.3065-3082. 10.1093/gbe/evw219. hal-02004240

HAL Id: hal-02004240
https://hal.umontpellier.fr/hal-02004240
Submitted on 1 Feb 2019

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Predicting the Stability of Homologous Gene Duplications in a Plant RNA Virus

Anouk Willemsen*,1,2, Mark P. Zwart1,3, Pablo Higueras1, Josep Sardanyés4,5, and Santiago F. Elena*,1,6,7

1Instituto de Biología Molecular y Celular de Plantas (IBMCP), Consejo Superior de Investigaciones Científicas-Universidad Politécnica de Valencia, Campus UPV CPI 8E, Ingeniero Fausto Elio s/n, València, Spain
2Present address: MIVEGEC (UMR CNRS 5290, IRD 224, UM), National Center for Scientific Research (CNRS), 911 Avenue Agropolis, BP 64501, 34394 Montpellier, Cedex 5, France
3Present address: Institute of Theoretical Physics, University of Cologne, Zülpicher Straße 77, 50937 Cologne, Germany
4ICREA Complex Systems Laboratory, Universitat Pompeu Fabra, Barcelona, Spain
5Institut de Biologia Evolutiva (Consejo Superior de Investigaciones Científicas-Universitat Pompeu Fabra), Barcelona, Spain
6Instituto de Biología Integrativa y de Sistems (i2SysBio), Consejo Superior de Investigaciones Científicas-Universitat de València, Parc Científic de la Universitat de València, Paterna, València, Spain
7The Santa Fe Institute, Santa Fe, New Mexico

*Corresponding author: E-mail: anouk.willemsen@ird.fr; sfelena@ibmcp.upv.es.

Accepted: September 2, 2016

Data deposition: The sequences of the ancestral viral stocks were submitted to GenBank with accessions TEV-NIb1-NIb9: KT203712; TEV-NIb2-NIb9: KT203713; TEV-HCPro2-HCPro3: KX137150; TEV-NlaPro2-NlaPro8: KX137151; TEV: KX137149.

Abstract

One of the striking features of many eukaryotes is the apparent amount of redundancy in coding and non-coding elements of their genomes. Despite the possible evolutionary advantages, there are fewer examples of redundant sequences in viral genomes, particularly those with RNA genomes. The factors constraining the maintenance of redundant sequences in present-day RNA virus genomes are not well known. Here, we use Tobacco etch virus, a plant RNA virus, to investigate the stability of genetically redundant sequences by generating viruses with potentially beneficial gene duplications. Subsequently, we tested the viability of these viruses and performed experimental evolution. We found that all gene duplication events resulted in a loss of viability or in a significant reduction in viral fitness. Moreover, upon analyzing the genomes of the evolved viruses, we always observed the deletion of the duplicated gene copy and maintenance of the ancestral copy. Interestingly, there were clear differences in the deletion dynamics of the duplicated gene associated with the passage duration and the size and position of the duplicated copy. Based on the experimental data, we developed a mathematical model to characterize the stability of genetically redundant sequences, and showed that fitness effects are not enough to predict genomic stability. A context-dependent recombination rate is also required, with the context being the duplicated gene and its position. Our results therefore demonstrate experimentally the deleterious nature of gene duplications in RNA viruses. Besides previously described constraints on genome size, we identified additional factors that reduce the likelihood of the maintenance of duplicated genes.

Key words: gene duplication, genome stability, experimental evolution, virus evolution.

Introduction

Gene duplication results in genetic redundancy; in other words, the existence of genetic elements that encode for the same function. It is a powerful process that can regulate gene expression, increase the genetic and environmental robustness of organisms, and act as a stepping stone to the evolution of new biological functions. Therefore, it is not surprising that gene duplication is a frequent phenomenon in many organisms (Zhang 2003; Andersson and Hughes 2009). Despite these clear advantages, there are few examples of genetic redundancy in viral genomes. In general, viral genomes tend to be highly streamlined, with limited intergenic sequences and in many cases overlapping open reading frames. This tendency suggests that gene duplication is less common in viruses compared to eukaryotic cells. However, recent studies have shown that gene duplications can also occur in viruses, and that they can have significant evolutionary consequences. For example, virus-encoded gene duplications have been observed in many plant viruses, including Tobacco etch virus (TEV), a plant RNA virus that is well known for its role in the evolution of new biological functions. Therefore, understanding the factors that constrain the maintenance of gene duplications in viral genomes is important for elucidating the evolutionary potential of viruses and for understanding the mechanisms that govern the stability of genetic redundancy in RNA viruses.
frames (ORFs), suggesting that genome size is under strong selection (Lynch 2006). Therefore, it is not surprising that gene duplications appear to be relatively frequent only in the large DNA viruses (Shackelton and Holmes 2004; Suhre 2005). RNA viruses typically have smaller genomes (ranging from ~2 kb to ~33 kb) than DNA viruses (ranging from the ~1.7 kb of circoviruses to the ~2.5 Mb of the giant pandoraviruses) (Campillo-Balderas et al. 2015), and consequently there is an extremely low prevalence of gene duplication in RNA viruses (Belshaw et al. 2007, 2008; Simon-Loriere and Holmes 2013). For the reverse-transcribing viruses, three different gene duplication events have been reported within the Retroviridae family (Tristem et al. 1990; Lapierre et al. 1999; Kambol et al. 2003). This low prevalence of gene duplication in retroviruses is surprising, because repeated sequence elements of endogenous retroviruses are thought to mediate genomic rearrangements, including gene duplication (Hughes and Coffin 2001). For the ss-(+)RNA viruses, two different tandem gene duplications have been reported (Walker et al. 1992; Gubala et al. 2010; Blasdell et al. 2012; Simon-Loriere and Holmes 2013) within the Rhabdoviridae (infecting vertebrates, invertebrates, and plants). For the ss-(+)RNA viruses, single duplication events have been reported for four different domains: (i) a tandem duplication of the coat protein gene (CP) within the Closteroviridae (infecting plants) (Boyko et al. 1992; Fazeli and Rezaaian 2000; Kreuze et al. 2002; Tzanetakis et al. 2005; Tzanetakis and Martin 2007; Simon-Loriere and Holmes 2013); (ii) a tandem duplication of the genome-linked protein gene (VPg) in Foot-and-mouth disease virus from the Picornaviridae (infecting vertebrates) (Forss and Schaller 1982); (iii) a duplication of the third segment, generating an additional one, in Beet necrotic yellow vein virus from the Benyviridae (infecting plants) (Simon-Loriere and Holmes 2013); and (iv) a tandem duplication of the P1 gene in Cucumber vein yellowing virus from the Potyviridae (infecting plants) (Valli et al. 2007). In the latter case, the two P1 copies resemble two different genera types within the family, suggesting that both recombination and gene duplication have contributed to P1 evolution. To date, no cases of gene duplication in dsRNA viruses have been reported.

The variation in genome size and structure indicates that gene duplication must have played a role in the early diversification of viral genomes. However, the rapid evolution of RNA viruses and the potential fitness costs associated with harboring additional genetic material probably makes it unlikely to detect viruses with duplications, or even the signatures of recent duplication events. Strong selective constraints against increasing genome sizes are thought to play a role in the lack of gene duplications that we nowadays observe in RNA viruses (Holmes 2003; Belshaw et al. 2007, 2008). One of these constraints is the high mutation rates of RNA viruses, which is approximately one mutation per genome and per replication event (Sanjuán et al. 2010). Another constraint is the need for fast replication due to strong within-cell and within-host competition (Turner and Chao 1998). On the other hand, the small and streamlined RNA virus genomes also limit sequence space for the evolution of novel functions, and in turn adaptation to environmental changes.

Here, we therefore consider experimentally the evolutionary fate of gene duplications in viral genomes, in terms of their effects on fitness, the stability of the duplicated gene and the evolvability of these viruses. We choose to work with Tobacco etch virus (TEV; genus Potyvirus, family Potyviridae). This plant pathogen has a monopartite ss(+)-RNA genome, of about 9.5 kb in length (Revers and García 2015). It codes for a single ORF (i.e., polyprotein) that is further processed after translation into ten mature peptides, in addition the P3N-PIPO ORF is translated at a +2 frameshift within the P3 cistron (fig. 1A). We experimentally explore four cases of homologous duplication of genes within the TEV genome: (i) the multifunctional protein (HC-Pro) involved in aphid transmission, polyprotein cleavage, genome amplification, and suppression of RNA silencing; (ii) the main viral protease (Nla-Pro), (iii) the viral RNA-dependent RNA polymerase (Nlb), and (iv) the coat protein (CP). Potyviruses are a particularly interesting system for studying the evolution of gene duplications, as for each complete ss(+)-RNA there will be isostiochiometric expression of all genes. Assuming there are no unknown mechanisms that regulate gene expression, the scope for the regulation of gene expression in potyviruses could therefore be very limited. Gene duplication may represent a way to bypass these constraints and achieve higher expression of specific genes.

We speculated that the duplication of these four proteins might have widely different impacts on TEV fitness. As HC-Pro is a multifunctional protein, two copies of HC-Pro could lead to specific improvement of one or more of its functions. This potential improvement could possibly be caused by two mechanisms. First, by simply producing more protein there could be an immediate benefit for one of HC-Pro’s functions. Second, there could be improvement of protein function when the duplicated virus is evolved, because the two gene copies can diverge and specialize on different functions. Higher levels of Nla-Pro may result in a more efficient processing of the polyprotein, making more mature viral proteins available faster for the replication process. As potyviruses have only a limited number of post-translational mechanisms for regulating gene expression levels, we predicted that the overproduction of Nla-Pro will alter the equilibrium concentrations of all the different mature peptides and thus have a major impact in TEV fitness. Higher levels of Nlb may result in higher levels of transcription and faster replication of the virus and this could lead to higher levels of genome accumulation and potentially the within-host spread of infection by a greater number of virions. The cellular multiplicity of infection (MOI), which has been estimated to be as low as 1.14 virions per infected cell for TEV (Tromas, Zwart, Lafforgue, et al. 2014), might even increase. Higher levels of CP expression could allow for the encapsidation of more genomic RNA.
molecules without affecting the accumulation of all other mature peptides. However, in all these cases completion of the infectious cycle would still depend on the cytoplasmic amount of other limiting viral (e.g., P1, P3, CI, and VPg) or host proteins.

The duplication events that we explore here could therefore conceivably have beneficial effects on TEV replication, perhaps offsetting the costs inherent to a larger genome and thereby increasing overall fitness. Moreover, especially in the case of HC-Pro, they could perhaps lead to the evolution of greatly improved or novel functions. However, given the scarcity of gene duplications in RNA viruses, we expected that the fitness costs of duplication are likely to be high, and that further mutations could potentially help accommodate the duplicated gene, then this could lead to interesting evolutionary dynamics: will the duplicated gene be lost or will beneficial mutations that lead to stable maintenance of the gene occur first? (Zwart et al. 2014). Moreover, as they could potentially disrupt correct processing of the polyprotein, the possibility that some of the duplications would not be viable in the first place could also not be discounted (Majer et al. 2014). To address these issues we have constructed four viruses with gene duplications and tested their viability. We subsequently evolved these viruses and determined the stability of the duplicated gene, as well as looking for signals of accommodation of the duplicated gene. Finally, we built a mathematical model to estimate key parameters from the experimental data, such as the recombination rates responsible for the deletion of duplicated genes, and to explore the evolutionary dynamics and stability conditions of the system.

**Materials and Methods**

**Viral Constructs, Virus Stocks and Plant Infections**

The TEV genome used to generate the virus constructs, was originally isolated from *Nicotiana tabacum* plants (Carrington et al. 1993). In this study five different variants of TEV were used containing single gene duplications. Two of these virus variants, TEV-NIb1-NIb9 and TEV-NIb2-NIb9, were generated in a previous study (Willemsen et al. 2016). The other three variants were generated in this study: TEV-HCPro2-HCPro3, TEV-NIaPro2-NIaPro8, and TEV-CP10-CP11. The TEV-HCPro2-HCPro3, TEV-NIaPro2-NIaPro8, and TEV-CP10-CP11 viruses contain gene duplications in tandem.

**Serial Passages**

For the serial passage experiments, 500 mg homogenized stock tissue was ground into fine powder using liquid nitrogen and a mortar, and resuspended in 500 μl phosphate buffer (50 mM KH2PO4, pH 7.0, 3% polyethylene glycol 6000). From...
this mixture, 20 μl were then mechanically inoculated on the third true leaf of 4-week old *N. tabacum* plants. At least five independent replicates were performed for each virus variant. At the end of the designated passage duration (3 or 9 weeks) all leaves above the inoculated leaf were collected and stored at −80 °C. For subsequent passages, the frozen tissue was homogenized and a sample of the homogenized tissue was ground and resuspended with an equal amount of phosphate buffer (Zwart et al. 2014). Then, new *N. tabacum* plants were mechanically inoculated as described above. The plants were kept in a BSL-2 greenhouse at 24 °C with 16 h light.

**Reverse Transcription Polymerase Chain Reaction (RT-PCR)**

The wild-type TEV produces characteristic symptoms in the host plant. However, some of the altered genotypes show few or no symptoms and virus infection had to be confirmed by RT-PCR. To confirm infection and to determine the stability of the duplicated genes, RNA was extracted from 100 mg homogenized infected tissue using the InviTrap Spin Plant RNA Mini Kit (Stratagene). Reverse transcription (RT) was performed using MuLV reverse transcriptase (Thermo Scientific) and the reverse primer 5′-CGACACTACATGAGGAAATAG-3’ located in the 3′UTR of the TEV genome. PCR was then performed with Taq DNA polymerase (Roche) and primers flanking the region containing the duplicated gene copy (supplementary table S1, Supplementary Material online). To test whether the ancestral gene copy was intact this region was also amplified for TEV-NlaPro3-NlaPro8, TEV-Nlb2-Nlb9, and TEV-Nlb7-Nlb8 viruses, where the duplicated genes are not located in tandem (supplementary table S1, Supplementary Material online). PCR products were resolved by electrophoresis on 1% agarose gels. For those virus populations in which we detected deletions during the evolution experiment, we estimated the genome size based on the amplicon size and the genome size of the ancestral viruses.

**Fitness Assays**

The genome equivalents per 100 mg of tissue of the ancestral virus stocks and all evolved lineages were determined for subsequent fitness assays. The Invitrap Spin Plant RNA Mini Kit (Stratagene Molecular) was used to isolate total RNA from 100 mg homogenized infected tissue. Real-time quantitative RT-PCR (RT-qPCR) was performed using the One Step SYBR PrimeScript RT-PCR Kit II (Takara), in accordance with manufacturer instructions, in a StepOnePlus Real-Time PCR System (Applied Biosystems). Specific primers for the CP gene were used; forward 5′-TTGGTCTTGATGGCAACGTG-3′ and reverse 5′-CGACACTACATGAGGAAATAG-3′. The StepOne Software v.2.2.2 (Applied Biosystems) was used to analyze the data. The concentration of genome equivalents per 100 mg of tissue was then normalized to that of the sample with the lowest concentration, using phosphate buffer. For the accumulation assays, 4-week-old *N. tabacum* plants were inoculated with 50 μl of the normalized dilutions of ground tissue. For each ancestral and evolved lineage, at least three independent plant replicates were used. Leaf tissue was harvested 7 dpi. Total RNA was extracted from 100 mg of homogenized tissue. Virus accumulation was then determined by means of RT-qPCR for the CP gene of the ancestral and the evolved lineages. For each of the harvested plants, at least three technical replicates were used in the RT-qPCR.

To measure within-host competitive fitness, we used TEV carrying an enhanced green fluorescent protein (TEV-eGFP) (Bedoya and Daróš 2010) as a common competitor. TEV-eGFP has proven to be stable up to 6 weeks (using 1- and 3-week serial passages) in *N. tabacum* (Zwart et al. 2014), and is therefore not subjected to appreciable eGFP loss during our 1-week long competition experiments. All ancestral and evolved viral lineages were again normalized to the sample with the lowest concentration, and 1:1 mixtures of viral genome equivalents were made with TEV-eGFP. The mixture was mechanically inoculated on the same species of host plant on which it had been evolved, using three independent plant replicates per viral lineage. The plant leaves were collected at 7 dpi, and stored at −80 °C. Total RNA was extracted from 100 mg homogenized tissue. RT-qPCR for the CP gene was used to determine total viral accumulation, and independent RT-qPCR reactions were also performed for the eGFP sequence using primers forward 5′-CGACACTACATGAGGAAATAG-3′ and reverse 5′-GAATCTCAGGAGCAGGT-3′. The ratio (R) of the evolved and ancestral lineages to TEV-eGFP is then $R = \frac{(n_{CP} - n_{eGFP})}{n_{eGFP}}$, where $n_{CP}$ and $n_{eGFP}$ are the RT-qPCR measured copy numbers of CP and eGFP, respectively. Within-host competitive fitness can then be estimated as $W = \sqrt{R_t/R_0}$, where $R_0$ is the ratio at the start of the experiment and $R_t$ the ratio after $t$ days of competition (Carrasco et al. 2007). Note that the method for determining $R$ only works well when the frequency of the common is below ~0.75. This limitation was not problematic though, because in these experiments the fitness of the evolved virus populations remained the same or increased. The statistical analyses comparing the fitness between lineages were performed using R v.3.2.2 (R Core Team 2014) and IBM SPSS Statistics version 23.

**Sanger Sequencing**

For those evolved virus populations in which deletions were detected by RT-PCR, the exact positions of these deletions were determined. The genomes were partly sequenced by the Sanger method. RT was performed using AccuScript HiFi (Agilent Technologies) reverse transcriptase and a reverse primer outside the region to be PCR-amplified for sequencing (supplementary table S2, Supplementary Material online). PCR was then performed with Phusion DNA polymerase (Thermo-Fusion DNA polymerase). Phylogenetic trees were constructed using the Neighbor-Joining method and the Kimura 2-parameter model.
Read artifact filtering and quality trimming (30 ambiguous, 0 unknown) of the reads and 5’ end trimming tool kit/index.html, last accessed April 10, 2016) with an ABI3730XL DNA analyzer. For TEV- HCPro2- HCPro3, six sequencing reactions were done per lineage using the same outer reverse primer as used for PCR amplification plus five inner primers (supplementary table S2, Supplementary Material online). For TEV-NlaPro2-NlaPro8, three sequencing reactions were done per lineage using three inner primers (supplementary table S2, Supplementary Material online). For TEV-Nlb1-Nlb9 and TEV-Nlb2-Nlb9, six sequencing reactions were done per lineage using the same two outer primers as used for PCR amplification plus four inner primers (supplementary table S2, Supplementary Material online). For TEV-CP10-CP11, two sequencing reactions were done per lineage using the same two outer primers as used for PCR amplification (supplementary table S2, Supplementary Material online). Sequences were assembled using Geneious v.8.0.3 (www.geneious.com; last accessed April 10, 2016) and the start and end positions of the deletions were determined. Based on the ancestral reference sequences, new reference sequences were constructed containing the majority deletion variant for each of the evolved lineages.

Illumina Sequencing, Variants, and SNP Calling

For Illumina next-generation sequencing (NGS) of the evolved and ancestral lineages, the viral genomes were amplified by RT-PCR using AccuScript Hi-Fi (Agilent Technologies) reverse transcriptase and Phusion DNA polymerase (Thermo Scientific), with six independent replicates that were pooled. Each virus was amplified using three primer sets generating three amplifications of similar size (set 1: 5’-GCAATCAAGCATTCTACCTATTG CAGC-3’ and 5’-TATGGAGTTCTGTTGAGTTTCCAGATCC-3’; set 2: 5’-TGGACCGTACCAGGGATGTGG-3’ and 5’-AATGCTT CCAGAATATGCC-3’; set 3: 5’-TACACAAAAGCAGCT-TG-3’ and 5’-CGCCTTAATAGAGAATTTAG-3’). Equimolar mixtures of the three PCR products were made. Sequencing was performed at GenoScreen. Illumina HiSeq2500 2 x 100 bp paired-end libraries with dual-index adapters were prepared along with an internal PhiX control. Libraries were prepared using the Nextera XT DNA Library Preparation Kit (Illumina Inc.). Sequencing quality control was performed by GenoScreen, based on PhiX error rate and Q30 values.

Read artifact filtering and quality trimming (3’ minimum Q28 and minimum read length of 50 bp) was done using FASTX-Toolkit v.0.0.14 (http://hannonlab.cshl.edu/fastx_toolkit/index.html, last accessed April 10, 2016). De-replication of the reads and 5’ quality trimming requiring a minimum of Q28 was done using PRINSEQ-lite v.0.20.4 (Schmieder and Edwards 2011). Reads containing undefined nucleotides (N) were discarded. As an initial mapping step, the evolved sequences were mapped using Bowtie v.2.2.6 (Langmead and Salzberg 2012) against their corresponding ancestral sequence: TEV (GenBank accession number KX137149), TEV-HCPro2-HCPro3 ancestral (GenBank accession number KX137150), TEV-NlaPro2-NlaPro8 ancestral (GenBank accession number KX137151), TEV-Nlb1-Nlb9 ancestral (GenBank accession number KT203712), TEV-Nlb2-Nlb9 ancestral (GenBank accession number KT203713), and against the evolved lineages including the corresponding deletions in the lineages where they are present. Subsequently, mutations were detected using SAMTools’ mpileup (Li et al. 2009) in the evolved lineages as compared with their ancestral lineage. At this point, we were only interested in mutations at a frequency >10%. Therefore, we present frequencies as reported by SAMTools, which has a low sensitivity for detecting low-frequency variants (Spencer et al. 2014).

After the initial pre-mapping step, error correction was done using Polisher v2.0.8 (available for academic use from the Joint Genome Institute) and consensus sequences were defined for every lineage. Subsequently, the cleaned reads were remapped using Bowtie v.2.2.6 against the corresponding consensus sequence for every lineage. For each new consensus, Single nucleotide polymorphisms (SNPs) within each virus population were identified using SAMTools’ mpileup and VarScan v.2.3.9 (Koboldt et al. 2012). For SNP calling maximum coverage was set to 40000 and SNPs with a frequency <1% were discarded.

Modeling the Stability of Gene Insertions

We developed a mathematical model to fit with the experimental data for the 3-week and 9-week passages. We were particularly interested in better understanding the evolution of a viral population composed by two different viral types, the wild-type and one containing a duplication. The model consists of two coupled ordinary differential equations:

\[
\begin{align*}
\frac{dA}{dt} &= aA \left(1 - \frac{A + bB}{\kappa}\right) - \delta A \tag{1}
\end{align*}
\]

\[
\frac{dB}{dt} = bB \left(1 - \frac{aA + B}{\kappa}\right) + \delta A \tag{2}
\]

where A is the number of viruses containing a gene duplication, B is the number of viruses that had reverted to the wild-type with a single copy, a is the initial growth rate of type A virus, b is the initial growth rate of the type B, \(\mu = b/a\) is a constant for determining the effect of the presence of B on replication of A (Solé et al. 1998), \(\alpha = 1/\beta\) is a constant for determining the opposing effect of A on B, \(\kappa\) is the time-dependent carrying capacity of the host plant, and \(\delta\) is the recombination rate per genome and replication at which the extra copy of the gene is removed from A to produce B. We assume that \(\kappa\) increases linearly over time, being proportional to the estimated weight of collected plant tissue (2 g for the
whole plant at inoculation, 200 g for the collected leaves at 9 weeks). At the start of each round of infection, there is a fixed bottleneck size of \( \lambda \). The number of infecting virions of \( A \) is determined by a random draw from a Binomial distribution with a probability of success \( \lambda A/(A+B) \) and a size \( \lambda \), and the number of infecting virions of \( B \) is then \( \lambda \) minus this realization from the Binomial distribution.

Estimates of most model parameters could be obtained from previous studies (table 1). An estimate of \( b \) has been made (Zwart et al. 2012), whilst \( a \) can be determined knowing the competitive fitness of the virus with duplication relative to the wild-type virus. The value of \( b \) used is 1.344, and a values are 1.175, 1.234, 1.185, and 1.134 for TEV-HCPro2-HCPro3, TEV-NlaPro2-NlaPro8, TEV-Nlb1-Nlb9, and TEV-Nlb2-Nlb9, respectively. The only model parameter that needed to be estimated from the data is \( \delta \). To obtain an estimate of this parameter, we implemented the model as described in equations (1) and (2) in R 3.1.0. For each dataset to which we wanted to fit the model, we first simply ran the model for a particular realization (1) and (2) in R 3.1.0. For each dataset to which we wished to fit the model, we first simply ran the model for a large number of simulations were run for each parameter value. For each realization (1) and (2) used to characterize the dynamical properties, build the phase portraits and to obtain the transient times (supplementary text S2, Supplementary Material online) have been obtained using a fourth-order Runge–Kutta method with a time step size 0.1.

### Results

Genetic Redundant Constructs and the Viability of the Resulting Viruses

To simulate the occurrence of duplication events within the TEV genome (fig. 1A), different TEV genotypes were constructed using four genes of interest (fig. 1). Each of these genotypes therefore represents a single gene duplication event. Where necessary, the termini of the duplicated gene copies were adjusted, such that the proteins can be properly translated and processed. Cleavage sites are provided, similar to the original proteolytic cleavage sites at the corresponding positions. A description of every duplication event will be given in the same order as these genes occur within the TEV genome.

First, for duplication of the multifunctional HC-Pro gene, a second copy of HC-Pro was inserted in the second position within the TEV genome, between the P1 serine protease gene and the original HC-Pro copy, generating a tandem duplication (fig. 1B). This position is a common site for the cloning of heterologous genes (Zwart et al. 2011). Second, a copy of the Nla-Pro main viral protease gene was introduced between P1 and HC-Pro (fig. 1C). Third, two genotypes containing a duplication of the Nlb gene were generated (Willemsen et al. 2016), where a copy of the Nlb gene was inserted at the first position (before P1) and the second position in the TEV genome (fig. 1D). Fourth, for duplication of the CP we introduced a second copy at the tenth position between Nlb and CP, generating a tandem duplication (fig. 1E). Henceforth we refer to these five genetic redundant viruses as TEV-HCPro2-HCPro3, TEV-NlaPro2-NlaPro8, TEV-Nlb1-Nlb9, TEV-Nlb2-Nlb9, and TEV-CP10-CP11, respectively, with subscripts denoting the intergenic positions of the duplicated gene in question.

The viability of the genetic redundant viruses was tested in *N. tabacum* by inoculating plants with approximately 5 µg of
in vitro generated transcripts ("Materials and Methods" section). The TEV-HCPro2-HCPro3, TEV-NlaPro2-NlaPro8, TEV-Nlb1-Nlb9, and TEV-Nlb2-Nlb9 viruses were found to infect N. tabacum plants, as determined by RT-PCR of total RNA extracted from these plants. Multiple viability tests showed that the TEV-CP10-CP11 virus had a very low infectivity and that large amounts of RNA (>20 μg) were needed for a successful infection. When performing RT-PCR of the region containing the two CP copies, we detected either (i) a band corresponding to the wild-type virus, indicating that upon infection with RNA the second CP copy is deleted immediately, or (ii) a band that indicates the two CP copies are present. When taking the virus with the two CP copies as a starting population for experimental evolution, we found that this virus was highly unstable. Within the first 3-week passage, we detect a band corresponding to the wild-type virus, in six out of eight lineages, and we did not detect any infection in the remaining two lineages. When sequencing the region containing the deletions in the different lineages, using Sanger technology, exact deletions of the second CP copy were observed. We discontinued further experiments on this virus due to the extreme instability of the second CP copy.

Evolution of Genetically Redundant Viruses

After reconstitution of TEV-HCPro2-HCPro3, TEV-NlaPro2-NlaPro8, TEV-Nlb1-Nlb9, and TEV-Nlb2-Nlb9 from infectious clones, these viruses were evolved in N. tabacum plants, as determined by RT-PCR of total RNA extracted from these plants. Multiple viability tests showed that the TEV-CP10-CP11 virus had a very low infectivity and that large amounts of RNA (>20 μg) were needed for a successful infection. When performing RT-PCR of the region containing the two CP copies, we detected either (i) a band corresponding to the wild-type virus, indicating that upon infection with RNA the second CP copy is deleted immediately, or (ii) a band that indicates the two CP copies are present. When taking the virus with the two CP copies as a starting population for experimental evolution, we found that this virus was highly unstable. Within the first 3-week passage, we detect a band corresponding to the wild-type virus, in six out of eight lineages, and we did not detect any infection in the remaining two lineages. When sequencing the region containing the deletions in the different lineages, using Sanger technology, exact deletions of the second CP copy were observed. We discontinued further experiments on this virus due to the extreme instability of the second CP copy.

Viruses with a Gene Duplication Have Reduced Fitness Which Cannot Always Be Fully Restored after Deletion

For both the ancestral and evolved virus populations, we measured within-host competitive fitness (fig. 4) and viral accumulation (fig. 5). Comparing the ancestral viruses containing a gene duplication to the ancestral wild-type virus (solid circles in figs. 4 and 5), we observed statistically significant decreases in competitive fitness (fig. 4A: TEV-HCPro2-HCPro3: \( t_4 = 8.398, P = 0.001 \); fig. 4B: TEV-NlaPro2-NlaPro8: \( t_4 = 12.776, P < 0.001 \); fig. 4C: TEV-Nlb1-Nlb9: \( t_4 = 6.379, P = 0.003 \); TEV-Nlb2-Nlb9: \( t_4 = 8.348, P = 0.001 \)). Statistically significant decreases in accumulation levels were also observed for TEV-HCPro2-HCPro3, TEV-Nlb1-Nlb9, and TEV-Nlb2-Nlb9 (fig. 5A: TEV-HCPro2-HCPro3: \( t_4 = 3.491, P = 0.0251 \); fig. 5C: TEV-Nlb1-Nlb9: \( t_4 = 45.097, P < 0.001 \); TEV-Nlb2-Nlb9: \( t_4 = 8.348, P = 0.001 \)).
However, there was no difference in accumulation for the virus with a duplication of \( \text{NIa-Pro} \) (fig. 5B: \( t_4 = 2.099, P = 0.104 \)). All the four viruses with a gene duplication therefore have a reduced within-host competitive fitness, and three out of four viruses also have reduced accumulation levels. None of the possible benefits of these gene duplications therefore can compensate for their costs.

After evolving the viruses with gene duplications using three 9-week passages, within-host competitive fitness of all four viruses increased (open circles in fig. 4), compared with their respective ancestral viruses (fig. 4; asterisks indicate a significant increase, \( t \)-test with Holm–Bonferroni correction for multiple tests). When comparing the evolved lineages, within-host fitness for TEV-HCPro2-HCPro3 (fig. 4A) and TEV-NIaPro2-NIaPro8 (fig. 4B) was similar to that of the wild-type TEV. On the other hand, the evolved TEV-Nlb1-Nlb9 and TEV-Nlb2-Nlb9 lineages did not reach wild-type virus within-host fitness levels (fig. 4C). The within-host fitness of evolved lineages was compared by means of a nested ANOVA (table 2), allowing the independent lineages (at least 5) to be nested within the genotype and the independent plant replicates (3) to be nested within the independent lineages within the genotype. The nested ANOVA confirms that there is indeed an effect of the genotype for the TEV-Nlb1-Nlb9 and TEV-Nlb2-Nlb9 viruses (table 2; Willemsen et al. 2016), whereas for the TEV-HCPro2-HCPro3 and TEV-NIaPro2-NIaPro8 no effect was found. In summary, the fitness of TEV-HCPro2-HCPro3 and TEV-NIaPro2-NIaPro8 clearly increases to levels similar to the wild-type, whilst fitness did not increase for TEV-Nlb2-Nlb9 and TEV-Nlb1-Nlb9.

Together with within-host fitness, virus accumulation also increased significantly for the evolved TEV-Nlb1-Nlb9 and TEV-Nlb2-Nlb9 virus lineages (fig. 5C; asterisks), when compared with their respective ancestral viruses. However, accumulation levels did not increase significantly for most of the evolved lineages of the TEV-HCPro2-HCPro3 and TEV-NIaPro2-NIaPro8 genotypes. Nevertheless, these two genotypes have much higher initial accumulation levels than the genotypes with a duplication of the \( \text{NIb} \) gene. When comparing the accumulation levels of the evolved lineages to those of the wild-type, TEV-HCPro2-HCPro3 (fig. 5A), TEV-NIaPro2-NIaPro8 (fig. 5B), and TEV-Nlb2-Nlb9 (fig. 5C) do reach wild-type accumulation levels, whereas TEV-Nlb1-Nlb9 does not (fig. 5C). Comparing the accumulation levels of the evolved lineages by means of a nested ANOVA (table 2) confirms that there is an effect of the genotype for the TEV-Nlb1-Nlb9 virus (table 2; Willemsen et al. 2016), whereas no effect for the TEV-HCPro2-HCPro3, TEV-NIaPro2-NIaPro8, and TEV-Nlb2-Nlb9 viruses was found.

When comparing the within-host competitive fitness of the evolved TEV-NIaPro2-NIaPro8 9-week lineages to the 3-week lineages, we found that there is a linear relationship between genome size and within-host competitive fitness (fig. 6; Spearman’s rank correlation \( \rho = -0.795, 10 \text{ d.f.}, P = 0.006 \)). The evolved 9-week lineages, that contain genomic deletions, have a significant higher within-host competitive fitness (Mann–Whitney \( U = 4.5, P < 0.001 \)) than the evolved 3-week lineages without deletions.
Genome Sequences of the Evolved Lineages

All evolved and ancestral lineages have been fully sequenced using the Illumina technology. The sequences of the ancestral lineages were used as an initial reference for the evolved lineages. Furthermore, for the lineages where deletions were detected by RT-PCR (fig. 3), parts of the genome were sequenced by Sanger to determine the exact deletion sites. The majority deletions variants were used to construct new

Fig. 3.—The reduction in genome size over time. The different panels display how the genome size of the different viral genotypes with gene duplications (A–D) changes along the evolution experiments. The dotted grey lines indicate the genome sizes of the wild-type virus (below) and the ancestral viruses (above). The genome sizes of the 3-week lineages are drawn with dashed black lines and open symbols, and those of the 9-week lineages are drawn with continuous blue lines and filled symbols.
FIG. 4.—Within-host competitive fitness of the evolved and ancestral lineages. Fitness \( W \), as determined by competition experiments and RT-qPCR of the different viral genotypes with respect to a common competitor, TEV-eGFP. The ancestral lineages are indicated by filled circles and the evolved lineages by open circles. The different viral genotypes are color coded, where the wild-type virus is drawn in green. The asterisks indicate statistically significant differences of the evolved lineages as compared with their corresponding ancestral lineages (t-test with Holm–Bonferroni correction).
reference sequences for each of the evolved TEV-HCPro2-HCPro3, TEV-NlaPro2-NlaPro8, TEV-Nlb1-Nlb9, and TEV-Nlb2-Nlb2 lineages that contain deletions. After an initial mapping step, mutations were detected in the evolved lineages as compared with their corresponding ancestor (“Materials and Methods” section). Beside the large genomic deletions, different patterns of adaptive evolution were observed for each viral genotype (fig. 7 and table 3). For a more detailed description
of the mutations detected, see supplementary text S1 (Supplementary Material online). After remapping the cleaned reads against a new defined consensus sequence for each lineage, we looked at the variation within each lineage. Single nucleotide polymorphisms (SNPs) were detected at a frequency as low as 1%. In all four virus genotypes as well as in the wild-type virus, most of the SNPs were present at low frequency (frequency of SNPs at < 0.1: TEV-HCPro2-HCPro3 = 89.8%; TEV-NlaPro2-NlaPro8 = 84.8%; TEV-Nlb1-Nlb9 = 83.3%; TEV-Nlb2-Nlb9 = 81.2%; TEV = 85.2%), with a higher prevalence of synonymous (TEV-HCPro2-HCPro3: 54.7%, TEV-NlaPro2-NlaPro8: 57.7%, TEV-Nlb1-Nlb9: 66.4%, TEV-Nlb2-Nlb9: 64.5%, TEV: 59.7%) versus non-synonymous changes (supplementary fig. S1, Supplementary Material online). Moreover, a percentage of the non-synonymous changes for TEV-HCPro2-HCPro3 (16.7%) and TEV-NlaPro2-NlaPro8 (7.3%) as well as the wild-type TEV (14.8%), are actually leading to stop codons and therefore unviable virus variants. In the lineages of both TEV-HCPro2-HCPro3 and TEV-NlaPro2-NlaPro8, there are significant differences in the distribution of the frequency of synonymous and non-synonymous SNPs per nucleotide position (Kolmogorov–Smirnov test; TEV-HCPro2-HCPro3: D = 0.219, P < 0.001; TEV-NlaPro2-NlaPro8: D = 0.151, P = 0.009), whereas for TEV-Nlb1-Nlb9 and TEV-Nlb2-Nlb9 (Willemsen et al. 2016) and the wild-type virus this is not significant (Kolmogorov–Smirnov test; TEV: D = 0.084, P = 0.555). For more details on the frequency of the SNPs within every lineage, see supplementary tables S3–S5 (Supplementary Material online). 

### Genomic Stability of TEV with Duplications of Homologous Genes

To better understand the evolutionary dynamics of viruses with gene duplications, we developed a simple mathematical model of virus competition and evolution. Based on amplicon sizes, the genome size for all evolved lineages was estimated.
been drawn to emphasize the trend in the data. The evolved 3-week lineages without deletions. A linear regression has been fit to the data, and specifically how long the duplicated gene is maintained in the virus population. The model we developed describes how a population composed initially of only a virus variant with a gene duplication (variant A) through recombination, selection, and genetic drift acquires and eventually fixes a new variant that only retains the original copy of the duplicated gene (variant B). The model includes a genetic bottleneck at the start of each round of passaging (i.e., the initiation of infection in the inoculated leaf), with a fixed total number of founders and binomially distributed number of founders for variants containing the gene duplication. Following this genetic bottleneck, there is deterministic growth of both variants as well as deterministic recombination of A into B. The main question we addressed is whether knowing the fitness of duplicated viruses (i.e., a) is sufficient information to predict the stability of the inserted gene. Do the data support a context-dependent recombination rate, with the context being (i) identity and position of the duplication, (ii) passage length, or (iii) both? We considered four different situations that are represented in the following models:

Model 1: one recombination rate for all conditions (1 parameter);
Model 2: virus-genotype-dependent recombination rate (4 parameters);
Model 3: passage-duration-dependent recombination rates (2 parameters);
Model 4: virus-genotype- and passage-duration-dependent recombination rates (full model, 8 parameters).

The model estimates of $\delta$ are given in Table 4. Note that the parameter is often a minimum (when the virus is very unstable) or a maximum value (when the virus is very stable). If the optimum is represented by more than one parameter value (i.e., a range of parameter values that correspond to the lowest NLL), the mean of these values is given. Some of the models fit the data well (Table 5), and we therefore saw no reason to explore more complex models of virus evolution. When comparing these models, we found that Model 2 is the best-supported model (Table 5). Thus, only a genotype-dependent recombination rate is required to account for the data. The results strongly suggest that the fitness of a virus with a gene duplication does not provide sufficient information for predicting genomic stability. Rather, as the recombination rate is dependent on the genetic context, the supply of recombinants which have lost the duplicated gene will vary greatly from one genotype to another. High recombinogenic sites will remove the second copy fast, whereas low recombinogenic sites will preserve the copy for longer periods of time, after which it will be unavoidably removed, as confirmed by the numerical analysis of equations (1) and (2) (Supplementary text S2, Supplementary Material online).

On the other hand, passage duration has a strong effect on the observed stability of gene duplications. However, model selection shows that this phenomenon can be sufficiently explained by considering the combined effects of selection and genetic drift, and without invoking passage-duration-dependent recombination rates. Given that recombination and selection are deterministic in the model, deletion variants will always arise during infection. However, depending on the rates of recombination and selection, these deletion variants may not reach a high enough frequency to ensure they are sampled during the genetic bottleneck at the start of each round of infection. This effect will be much stronger in the 3-week passages—because there will be fewer recombinants and less time for selection to increase their frequency—explaining why some viruses there is such a marked difference in the observed genomic stability for different passage lengths.

The deterministic dynamics of the evolutionary model of stability of genomes containing gene duplications have also been investigated (Supplementary text S2, Supplementary Material online). These analyses were performed on the model as described in equations (1) and (2), albeit with a time-independent carrying capacity. The stability of three fixed points was analyzed: (i) the extinction of both A and B, (ii) the dominance of B over the population, and (iii) the coexistence of A and B in the population. The fixed points analysis indicates that when $a < b$ and $\delta > a - b$, the B virus subpopulation will outcompete the A subpopulation.
**Fig. 7.**—Genomes of the ancestral and evolved lineages. Mutations were detected using NGS data of the evolved virus lineages as compared with their ancestral lineages. The square symbols represent mutations that are fixed (>50%) and the circle symbols represent mutations that are not fixed (<50%). Filled symbols represent non-synonymous substitutions and open symbols represent synonymous substitutions. Black substitutions occur only in one lineage, whereas color-coded substitutions are repeated in two or more lineages, or in a lineage from another virus genotype. Note that the mutations are present at different frequencies as reported by SAMtools. Grey boxes indicate genomic deletions in the majority variant.
This parametric combination is the one obtained using the biologically meaningful parameter values shown in table 1. This means that, under the model assumptions, the population of virus variants containing a gene duplication is unstable, and the population will be asymptotically dominated by virus variants containing a single gene copy. Notice that, the rate of recombination (\( \delta \)) is also involved in this process of out-competition (see the bifurcation values calculated in supplementary text S2, Supplementary Material online). Specifically, if \( \delta > a - b \), coexistence of the two virus variants is not possible. Interestingly, the model also reveals the existence of a transcritical bifurcation separating the scenario between coexistence of the two variants and dominance of \( B \). Such a bifurcation can be achieved by tuning \( \delta \) as well as by unbalancing the fitness of \( A \) and \( B \) virus types. This bifurcation gives place to a smooth transition between these two possible evolutionary asymptotic states (i.e., unstable \( A \) population and coexistence of \( A \) and \( B \) populations). The bifurcation will take place when \( a = b + \delta \), or when \( \delta \) is above a critical threshold, \( \delta_c = a - b \), that can be calculated from the mathematical model (supplementary text S2, Supplementary Material online).

Finally, we characterized the time to extinction of the \( A \) subpopulation. We characterized these extinction times as a function of viral fitness and of \( \delta \). Such a time is found to increase super-exponentially as the fitness of the \( A \) subpopulation approaches the fitness of the \( B \) subpopulation. This effect is found for low values of recombination rates, similar to those shown in table 4. As expected, increases in \( \delta \) produce a drastic decrease in the time needed for the single-copy variants to outcompete the variants with a gene duplication within the population (supplementary text S2, Supplementary Material online).

**Discussion**

Present-day viruses display a great variation in genome size and structure, indicating that gene duplication must have played a role in the early diversification of virus genomes. However, nowadays gene duplications are rarely documented in viruses, especially in RNA viruses. Therefore, it is quite obvious to expect a priori that genomes with gene duplications will be unstable in present-day RNA viruses. However, precise gene duplications occur never or only rarely in nature, and as our model virus encodes for a polyprotein, partial gene duplications are most likely to be non-functional. The engineered viruses in this study, removed that hurdle and therefore we expected that there was a higher probability of the introduced gene duplications contributing to the enhancement of a virus function or even the exploration of new functions. Nevertheless, none of the duplication events explored in this study appeared to be beneficial for TEV, both in terms of their immediate effects and second-order effects on evolvability. Gene duplication resulted in either an unviable virus or a significant reduction in viral fitness. In all cases, the duplicated gene copy, rather than the ancestral gene copy, was deleted during long-duration passages. The earlier detection and more rapid fixation of deletion variants during longer-duration passages is congruent with results from a previous study (Zwart et al. 2014), where deletions of eGFP marker inserted in the TEV genome were usually observed after a single 9-week passage, but were rarely spotted even after nine 3-week passages. In this study, similar results were obtained for TEV-NlaPro\(_2\)-NlaPro\(_9\) where no deletions were spotted in the 3-week passages. On the other hand, the highly divergent results reported here for the stability of different duplicated genes suggest that passage duration is not the main factor determining whether gene duplication will be stable or unstable. Therefore, we postulated that the size of the duplicated gene, the nature of the gene, and/or the position for duplication could play a role in the stability of genomes containing a gene duplication.

The observation that gene duplication events result in decreases in fitness is not unexpected. The high mutation rate of RNA viruses is likely to constrain genome size (Holmes 2003), given that most mutations are deleterious (Sanjuán et al. 2004; Elena et al. 2006). This imposes the evolution of genome compression, where overlapping reading frames play a major role (Belshaw et al. 2007, 2008). In our model virus, we speculate that the fitness cost can be related to three processes: (i) the increase in genome size, (ii) the extra cost of more proteins being expressed in the context of using more cellular resources, and (iii) a disturbance in correct polyprotein processing. Although all the duplications considered here could conceivably have advantages for viral replication or encapsidation, our results suggest that the any such advantages are far outweighed by the costs associated with a larger genome, increased protein expression or the effects on polyprotein processing. However, the duplication of Nla-Pro does not affect the viral accumulation rate. This could be explained by the fact that the Nla-Pro gene is much smaller than the other duplicated genes. Consequently, in conditions where selection has the least time to act between bottleneck events associated with infection of a new host (3-week...
Table 4
Model Parameter Estimates for Deterministic Recombination Rate

<table>
<thead>
<tr>
<th>Model</th>
<th>Estimates of $\log_{10}[\theta]$ (Lower 95% fiducial limit, upper 95% fiducial limit)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$-6.2$ (*)</td>
</tr>
<tr>
<td>2</td>
<td>$2\text{HCPro}$</td>
</tr>
<tr>
<td></td>
<td>$2\text{NiaPro}$</td>
</tr>
<tr>
<td></td>
<td>$2\text{Nlb1}$</td>
</tr>
<tr>
<td></td>
<td>$2\text{Nlb2}$</td>
</tr>
<tr>
<td>3</td>
<td>$\geq -3.0$ (*)</td>
</tr>
<tr>
<td></td>
<td>$-9.65 (-10.1, -9.0)$</td>
</tr>
<tr>
<td></td>
<td>$\geq -2.9$ (*)</td>
</tr>
<tr>
<td></td>
<td>$-4.45 (-5.2, -3.7)$</td>
</tr>
<tr>
<td>4</td>
<td>$\geq -3.0$ (*)</td>
</tr>
<tr>
<td></td>
<td>$\leq -9.0$ (*)</td>
</tr>
<tr>
<td></td>
<td>$\geq -2.9$ (*)</td>
</tr>
<tr>
<td></td>
<td>$-4.45 (-5.5, -3.7)$</td>
</tr>
</tbody>
</table>

*The fiducial limit is identical to the parameter estimate, also when the parameter estimate is a range, $2\text{HCPro}$: TEV-HCPro2-HCPro3; $2\text{NiaPro}$: TEV-NiaPro2-NiaPro3; $2\text{Nlb1}$: TEV-Nlb1-Nlb2; $2\text{Nlb2}$: TEV-Nlb2-Nlb3; 3W: 3-week passages; 9W: 9-week passages.

Table 5
Model Selection for Models with Deterministic Recombination

<table>
<thead>
<tr>
<th>Model</th>
<th>Parameters</th>
<th>NLL</th>
<th>AIC</th>
<th>$\Delta$AIC</th>
<th>Akaike Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>254.284</td>
<td>510.569</td>
<td>443.470</td>
<td>0.000</td>
</tr>
<tr>
<td>2</td>
<td>4</td>
<td>29.549</td>
<td>67.099</td>
<td>-0.000</td>
<td>0.982</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>226.669</td>
<td>457.339</td>
<td>390.240</td>
<td>0.000</td>
</tr>
<tr>
<td>4</td>
<td>8</td>
<td>29.549</td>
<td>75.099</td>
<td>8.000</td>
<td>0.018</td>
</tr>
</tbody>
</table>

By fitting a mathematical model of virus evolution to the data, we found that knowing only viral fitness is not enough information to predict the stability of the duplicated genes: model selection suggests there is a context-dependent recombination rate, and specifically, the identity and position of the duplication also play a role. Given that the supply rate of variants with large deletions will be driven largely by homologous recombination, we had expected stability to depend on the genetic context. The estimates of the recombination rate per genome and generation in this study are far lower than previously reported for TEV, which was estimated to be $3.43 \times 10^{-5}$ per nucleotide and generation (Tromas, Zwart, Poulain, et al. 2014), which translates into $0.327$ per genome and generation. The estimates of this study (TEV-HCPro2-HCPro3; $1.00 \times 10^{-3}$; TEV-NiaPro2-NiaPro3; $2.24 \times 10^{-10}$; TEV-Nlb1-Nlb2; $1.26 \times 10^{-3}$; TEV-Nlb2-Nlb3; $3.55 \times 10^{-5}$) are much closer to the per nucleotide estimate. This large discrepancy could be related to two factors. First, Tromas, Zwart, Poulain, et al. (2014) considered recombination between two highly similar genotypes, which requires consideration of many details of the experimental system, including the rate at which cells will be coinfected by these genotypes. On the other hand, considering these details should lead to a general
estimate of the recombination rate (as opposed to the rate at which two different genotypes will recombine in a mixed infection) and hence this explanation is not very satisfactory. Second, only a small fraction of all recombination events will render viruses with a conserved reading frame, and a suitable deletion size: large enough to have appreciable fitness gains and be selected, but small enough not to disrupt the surrounding cistrons or polyprotein processing. Therefore, the parameter \(\delta\) described here is in fact the rate at which this particular subclass of recombinants occurs. This subclass is likely to be only a small fraction of all possible recombinants, and hence it is quite reasonable that these two estimates of the recombination rate vary by several orders of magnitude.

In addition to gene duplications, the model developed in this study can be applied to predict the stability of other types of sequence insertions, such as those brought about by horizontal gene transfer. Understanding the stability of gene insertions in genomes is highly relevant to the understanding genome-architecture evolution, but it also has important implications for biotechnological applications, such as heterologous genome engineering. This study can be applied to predict the stability of other types of insertions in genomes, which is highly relevant to the understanding of genome architecture evolution. In this study, we can be applied to predict the stability of other types of insertions in genomes, which is highly relevant to the understanding genome-architecture evolution.

Supplementary Material

Supplementary tables, figures, and texts are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org).

Acknowledgments

We thank Francisca de la Iglesia and Paula Agudo for excellent technical assistance. This work was supported by the John Templeton Foundation [grant number 22371 to S.F.E.]; the European Commission 7th Framework Program EvoEvo Project [grant number ICT-610427 to S.F.E.]; the Spanish Ministerio de Economía y Competitividad (MINECO) [grant numbers BFU2012-30805 and BFU2015-65037-P to S.F.E.]; the Botín Foundation from Banco Santander through its Santander Universities Global Division [J.S.]; the Secretaria d’Universitats i Recerca del Departament d’Economia i Coneixement de la Generalitat de Catalunya [J.S.]; and the European Molecular Biology Organization [grant number AST 625-2015 to A.W.]. The opinions expressed in this publication are those of the authors and do not necessarily reflect the views of the John Templeton Foundation.

Literature Cited


Thornbury DW, Patterson CA, Dessens JT, Pirone TP. 1990. Comparative sequence of the helper component (HC) region of Potato virus Y and a HC-defective strain, Potato virus C. Virology 178:573–578.


Associate editor: Eric Bapteste