1	High virulence does not necessarily impede viral adaptation to a
2	new host: A case study using a plant RNA virus
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#### 25 Abstract

26 **Background:** When between-host selection pressures predominate, theory suggests that high 27 virulence could hinder between-host transmission of microparasites, and that virulence 28 therefore will evolve to lower levels that optimize between-host transmission. Highly virulent 29 microparasites could also curtail host development, thereby limiting both the host resources 30 available to them and their own within-host effective population size. High virulence might 31 therefore curtail the mutation supply rate and increase the strength with which genetic drift 32 acts on microparasite populations, thereby limiting the potential to adapt to the host and 33 ultimately perhaps the ability to evolve lower virulence. As a first exploration of this 34 hypothesis, we evolved Tobacco etch virus carrying an eGFP fluorescent marker in two semi-35 permissive host species, Nicotiana benthamiana and Datura stramonium, for which it has a 36 large difference in virulence. We compared the results to those previously obtained in the 37 typical host, *Nicotiana tabacum*, where we have shown that carriage of *eGFP* has a high 38 fitness cost and its loss serves as a real-time indicator of adaptation.

39 **Results:** After over half a year of evolution, we sequenced the genomes of the evolved lineages and measured their fitness. During the evolution experiment, marker loss leading to 40 41 viable virus variants was only observed in one lineage of the host for which the virus has low 42 virulence, D. stramonium. This result was consistent with the observation that there was a 43 fitness cost of *eGFP* in this host, while surprisingly no fitness cost was observed in the host for which the virus has high virulence, N. benthamiana. Furthermore, in both hosts we 44 45 observed few lineages with increases in viral fitness, and host-specific convergent evolution 46 at the genomic level was only found in *N. benthamiana*.

47 Conclusions: The results of this study do not lend support to the hypothesis that high
48 virulence impedes microparasites' evolution. Rather, they exemplify that jumps between host
49 species can be game changers for evolutionary dynamics. When considering the evolution of

50	genome architecture, host species jumps might play a very important role, by allowing
51	evolutionary intermediates to be competitive.
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53	Keywords: Adaptation, Experimental evolution, Host-pathogen interactions, Virulence, Virus
54	evolution, Genome architecture evolution
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## 72 Background

From both applied and fundamental perspectives, virulence is a key phenotypic trait of 73 74 microparasites. In medicine and agriculture, it is crucial to understand mechanistically how 75 microparasites harm the host, in order to devise effective interventions. From a more 76 fundamental perspective, evolutionary biologists have long been interested in understanding 77 why many microparasites are highly virulent. It has been suggested that virulence reduces 78 between-host transmission, and that selection would therefore act to maximize between-host 79 transmission by reducing virulence [1, 2]. High virulence would signal maladaptation, for example following a host-species jump, and eventually be selected against. The ubiquity of 80 81 microparasitic virulence and the fact that many apparently well-adapted microparasites have 82 high virulence led to a more sophisticated framework: the hypothesis that there are tradeoffs 83 between virulence and transmission [2-5]. This framework posits that high levels of 84 replication could increase the probability of a microparasite being transferred to a new host, 85 whilst also increasing the probability that the host would die quickly and the temporal 86 window for transmission would be very brief. Under this more plausible framework, 87 virulence evolves to the level that optimizes between-host transmission [4, 6, 7].

88 The tradeoff hypothesis forms the cornerstone for theoretical frameworks considering the 89 evolution of virulence in many different pathosystems. Many important additions to the 90 framework have been made, for example recognizing that within-host competition and 91 opportunism can lead to increases in virulence [8–10]. Moreover, the importance of other 92 factors at the between-host level have been given consideration, such as self-shading [11]. 93 Self-shading occurs when the host population is structured, and a highly virulent 94 microparasite kills all host organisms in a subpopulation before transmission to another 95 subpopulation can be achieved. The effects of evolution on microparasitic virulence have

96 therefore been given considerable attention, although the number of experimental studies that97 address this issue is still rather limited, especially for viruses [12].

98 The effects of evolution on microparasite virulence have been widely considered. However, 99 virulence itself could also have profound effects on evolution, including its own evolutionary 100 dynamics [13]. This reversed causality is already apparent from the tradeoff model, under 101 which microparasites with suboptimal virulence will undergo reduced between-hosts 102 transmission. All other things equal, if a smaller number of hosts are infected effective 103 population size will be decreased, increasing the strength of genetic drift and decreasing the 104 mutation supply rate. In addition, the evolution to optimum virulence may be slow as this 105 optimum is not static and can shift towards lower virulence as the density of susceptible hosts 106 decreases [14]. Moreover, a wide range of virulence can be associated with each step of 107 evolution towards the optimum, where selection favors genotypes with higher fitness that may 108 improve transmission but not necessarily improve virulence [13]. Besides these effects of 109 virulence on evolution, it is conceivable that a similar within-host effect could also occur, 110 when virulence curtails host development and thereby limits the host resources available to 111 the microparasite. Virulence would then limit the microparasite effective population size 112 within hosts, again reducing the mutation supply and thereby slowing the rate of adaptation. 113 Interestingly, all of these mechanisms could limit the rate at which lower virulence evolves, 114 meaning that high virulence might persist longer than suggested by the simple tradeoff model 115 [13].

There are many reasons why high virulence in host-pathogen interactions could emerge, but the most likely avenue is probably a change of host species. For example, infection of Ebola virus in bats is asymptomatic, while in humans and other primates the death rate is high [15]. Changes in virulence have been explained by the host phylogeny, where similar levels of virulence are displayed by closely related hosts and host jumps across large genetic distances

may result in high virulence [16]. However, if a microparasite is confronted with a new host
environment in which its level of virulence is altered, how does virulence affect its ability to
adapt to the new host?

124 Here we address this question using Tobacco etch virus (genus: Potyvirus, family: 125 *Potyviridae*), a (+)ssRNA virus that infects a wide-range of host plants, and an experimental 126 evolution approach. To consider the effect of virulence on virus adaptation, we looked for 127 two natural host species in which (i) there was some evidence that TEV potential for 128 adaptation would be roughly similar, and (*ii*) there was a large difference in virulence. The 129 distribution of mutational fitness effects (DMFE) of TEV has been compared in eight host 130 species, and this study concluded that there were strong virus genotype-by-host species 131 interactions [17]. For many host species distantly related to the typical host of TEV, 132 Nicotiana tabacum, the DMFE changed drastically; many mutations that were neutral or 133 deleterious in N. tabacum, became beneficial. However, for two closely related host species, 134 Nicotiana benthamiana and Datura stramonium, most mutations tested remained neutral or 135 deleterious [17], implying that the fraction of beneficial mutations in both hosts is small. 136 Moreover, virus accumulation after one week of infection is also similar for both hosts [18]. 137 On the other hand, TEV infection of *N. benthamiana* will typically result in heavy stunting 138 and the death of the plant within a matter of weeks, whereas TEV infection of D. stramonium 139 is virtually asymptomatic. Whilst there are many similarities between TEV infection in these 140 two hosts, one key difference is host-pathogen interactions and therewith levels of viral 141 virulence brought about.

As a first exploration of the effects of virulence on microparasite evolution, we therefore decided to evolve TEV in *N. benthamiana* and *D. stramonium*. By serially passaging each independent lineage in a single plant, our study maximizes within-host selection. This setup allows us to exclusively focus on effects of within-host selection, although for our model

146 system we expect to see large differences in the resulting population size and the scope of 147 virus movement within the host. Moreover, to immediately gauge whether adaptive evolution 148 might be occurring, we passaged a TEV variant expressing a marker protein (Fig. 1), the 149 enhanced GFP (eGFP). Upon long-duration passages in N. tabacum, this exogenous sequence 150 is quickly lost due to its strong fitness cost, and its loss is reliably indicated by a loss of eGFP 151 fluorescence [19]. According to the above hypothesis that high virulence may impair the rate 152 of microparasite evolution, we expect that adaptive evolution would occur more quickly in the 153 host species for which TEV has lower virulence, D. stramonium, than in the host species for 154 which it has high virulence, N. benthamiana. Hence, we expected that in D. stramonium (i) the eGFP marker would be lost more rapidly, (ii) there would be more sequence-level 155 156 convergent evolution, and (iii) there would be larger increases in within-host competitive 157 fitness. However, the results clashed with these simple hypotheses, exemplifying the extent 158 to which a host species jump can be a game changer for RNA virus evolutionary dynamics.

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#### 160 Methods

#### 161 Virus stocks, plants and serial passages

162 The TEV genome used to generate TEV-eGFP virus, was originally isolated from N. tabacum 163 plants [20]. To generate a virus stock of the ancestral TEV-eGFP, the pMTEV-eGFP plasmid 164 [21] was linearized by digestion with BglII prior to in vitro RNA synthesis using the mMESSAGE mMACHINE<sup>®</sup> SP6 Transciption Kit (Ambion), as described in [22]. The third 165 166 true leaf of 4-week-old N. tabacum L var Xanthi NN plants was mechanically inoculated with 167 5 µg of transcribed RNA. All symptomatic tissue was collected 7 days post-inoculation (dpi). For the serial passage experiments, 500 mg homogenized stock tissue was ground into fine 168 169 powder and diluted in 500 µl phosphate buffer (50 mM KH<sub>2</sub>PO<sub>4</sub>, pH 7.0, 3% polyethylene 170 glycol 6000). From this mixture, 20 µl were then mechanically inoculated on the sixth true

171 leaf of 4-week old *N. benthamiana* Domin plants and on the third true leaf of 4-week old *D*. 172 stramonium L plants. Ten independent replicates were used for each host plant. Based on a 173 previous study done in *N. tabacum* [19], passages of TEV-eGFP in *D. stramonium* were done every 9 weeks. In N. benthamiana the virus induces host mortality, and therefore the 174 175 passages had to be restricted to 6 weeks for this host. At the end of the designated passage 176 duration all leaves above the inoculated one were collected and stored at -80 °C. For 177 subsequent passages the frozen tissue was homogenized and a sample was ground and 178 resuspended with an equal amount of phosphate buffer [19]. Then, new plants were 179 mechanically inoculated as described above. Three passages were performed for lineages 180 evolved in *D. stramonium* and five passages for lineages evolving in *N. benthamiana*, making 181 the number of generations of evolution similar in both hosts. All plants were kept in a 182 biosafety level 2 greenhouse at 24° C with 16 h light:8 h dark photoperiod.

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#### 184 Reverse transcription polymerase chain reaction (RT-PCR)

185 To determine whether deletions occurred at the *eGFP* locus, RNA was extracted from 100 mg 186 homogenized infected tissue using the InviTrap Spin Plant RNA Mini Kit (Stratec Molecular). 187 Reverse transcription (RT) was performed using M-MuLV reverse transcriptase (Thermo 188 Scientific) and reverse primer 5'-CGCACTACATAGGAGAATTAG-3' located in the 3'UTR of the TEV genome. PCR was then performed with Taq DNA polymerase (Roche) and 189 190 primers flanking the *eGFP* gene: forward 5'-GCAATCAAGCATTCTACTTC-3', and reverse 191 5'-CCTGATATGTTTCCTGATAAC-3'. PCR products were resolved by electrophoresis on 192 1% agarose gels.

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#### 194 Virus accumulation and within-host competitive fitness assays

195 Prior to performing assays, the genome equivalents per 100 mg of tissue of the ancestral virus 196 stocks and all evolved lineages were determined for subsequent fitness assays. The InviTrap 197 Spin Plant RNA Mini Kit (Stratec Molecular) was used to isolate total RNA of 100 mg 198 homogenized infected tissue. Real-time quantitative RT-PCR (RT-qPCR) was performed 199 using the One Step SYBR PrimeScript RT-PCR Kit II (Takara), in accordance with 200 manufacturer instructions, in a StepOnePlus Real-Time PCR System (Applied Biosystems). 201 Specific primers for the *CP* gene were used: forward 5'-TTGGTCTTGATGGCAACGTG-3' 202 and reverse 5'-TGTGCCGTTCAGTGTCTTCCT-3'. The StepOne Software v.2.2.2 (Applied 203 Biosystems) was used to analyze the data. The concentration of genome equivalents per 100 204 mg of tissue was then normalized to that of the sample with the lowest concentration, using 205 phosphate buffer.

206 For the accumulation assays, 4-week-old N. benthamiana and D. stramonium plants were 207 mechanically inoculated with 50 µl of the normalized dilutions of ground tissue. Inoculation 208 of each viral lineage was done on the same host plant on which it had been evolved, plus TEV 209 and the ancestral TEV-eGFP virus on each of the hosts, using three independent plant 210 replicates per lineage. Leaf tissue was harvested 10 dpi. Total RNA was extracted from 100 211 mg of homogenized tissue. Virus accumulation was then determined by means of RT-qPCR 212 for the CP gene of the ancestral and the evolved lineages. For each of the harvested plants, at 213 least three technical replicates were used for RT-gPCR.

To measure within-host competitive fitness, we used TEV carrying a red fluorescent protein: TEV-mCherry as a common competitor. This virus has a similar insert size and within-host fitness compared with TEV-eGFP [19]. 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-mCherry [21]. The mixture was mechanically inoculated on the same host plant on which it had been evolved, plus TEV and the ancestral TEV-eGFP 220 virus on each of the hosts, using three independent plant replicates per viral lineage. The 221 plant leaves were collected at 10 dpi, and stored at -80 °C. Total RNA was extracted from 222 100 mg homogenized tissue. RT-qPCR for the CP gene was used to determine total viral 223 accumulation, and independent RT-qPCR reactions were also performed for the mCherry 224 sequence using specific primers: forward 5'-CGGCGAGTTCATCTACAAGG-3' and reverse 225 5'-TGGTCTTCTTCTGCATTACGG-3'. The ratio of the evolved and ancestral lineages to 226 TEV-mCherry (R) is then  $R = (n_{CP} - n_{mCherry})/n_{mCherry}$ , where  $n_{CP}$  and  $n_{mCherry}$  are the 227 RT-qPCR measured copy numbers of CP and mCherry, respectively. Then we can estimate the within-host competitive fitness as  $W = \sqrt[t]{R_t/R_0}$ , where  $R_0$  is the ratio at the start of the 228 229 experiment and  $R_t$  the ratio after t days of competition [22]. The statistical analyses 230 comparing the fitness between lineages were performed using R v.3.2.2 [23] and IBM SPSS 231 Statistics version 23.

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#### 233 Illumina sequencing, variants, and SNP calling

234 For Illumina next-generation sequencing (NGS) of the evolved and ancestral lineages, the 235 viral genomes were amplified by RT-PCR using AccuScript Hi-Fi (Agilent Technologies) 236 reverse transcriptase and Phusion DNA polymerase (Thermo Scientific), with six independent 237 replicates that were pooled. Each virus was amplified using three primer sets, generating 238 three amplicons of similar size (set 1: 5'-GCAATCAAGCATTCTACTTCTATTGCAGC-3' 239 and 5'-CCTGATATGTTTCCTGATAAC-3'; set 2: 5'-ACACGTACTGGCTGTCAGCG-3' 240 and 5'-CATCAATGTCAATGGTTACAC-3'; set 3: 5'-CCCGTGAAACTCAAGATAG-3' and 5'-CGCACTACATAGGAGAATTAG-3'). Equimolar mixtures of the three PCR 241 242 Sequencing was performed at GenoScreen (Lille, France: products were made. 243 www.genoscreen.com). Illumina HiSeq2500 2×100bp paired-end libraries with dual-index 244 adaptors 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.

247 Read artifact filtering and quality trimming (3' minimum Q28 and minimum read length of 50 248 bp) was done using FASTX-Toolkit v.0.0.14 [24]. De-replication of the reads and 5' quality 249 trimming requiring a minimum of Q28 was done using PRINSEQ-lite v.0.20.4 [25]. Reads 250 containing undefined nucleotides (N) were discarded. Initially, the ancestral TEV-eGFP 251 sequence was mapped using Bowtie v.2.2.6 [26] against the reference TEV-eGFP sequence 252 (GenBank accession: KC918545). Error correction was done using Polisher v2.0.8 (available 253 for academic use from the Joint Genome Institute) and a consensus sequences was defined for 254 the ancestral TEV-eGFP lineage. Subsequently, the cleaned reads of the evolved sequences 255 were mapped using Bowtie v.2.2.6 against the new defined consensus sequence. Single 256 nucleotide mutations for each viral lineage were identified using SAMtools' mpileup [27] and 257 VarScan v.2.3.9 [28], where the maximum coverage was set to 40000 and mutations with a 258 frequency < 1% were discarded. Note that the single nucleotide mutations detected here can 259 be fixed (frequency > 50%) in the evolved lineages, as the detection was done over the 260 ancestral population. Hence, it allows us to compare the mutations that arose by evolving 261 TEV-eGFP in the different hosts.

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#### 263 **Results**

#### 264 Experimental setup and fluorescent marker stability upon passaging of TEV-eGFP

TEV-eGFP was mechanically passaged in *N. benthamiana* and *D. stramonium*. In a previous
study we noted that 9-week long passages led to rapid deletion of *eGFP* as well as rapid
convergent evolution in *N. tabacum* [19]. Although 9-week passages could be performed in *D. stramonium*, for *N. benthamiana* this was not possible due to virus-induced host mortality.
These plants died after 6 weeks of infection, and therefore we were forced to collect tissue at

270 this time point. As D. stramonium grows to similar heights as N. tabacum when infected with 271 TEV, and N. benthamiana does not grow much after infection, we chose to maximize 272 infection duration to make the results comparable to those obtained in *N. tabacum* [19]. We 273 performed three 9-week passages in D. stramonium and - to keep the total evolutionary time 274 comparable – five 6-week passages in N. benthamiana. In D. stramonium all ten lineages 275 initiated were completed, whereas in N. benthamiana only 6/10 lineages were completed. 276 The remaining four *N. benthamiana* lineages failed to cause infection in subsequent rounds of 277 passaging, and were therefore halted. Initial symptomatology of TEV-eGFP in N. 278 benthamiana was very mild, while this symptomatology was more severe in the second and 279 subsequent passages, possibly indicating adaptation of the virus to this alternative host. In D. 280 stramonium the symptomatology was constant along the evolution experiment.

281 Based on previous results, we expected that the exogenous *eGFP* gene sequence would be 282 rapidly purged [19, 29, 30], and as such would serve as a first indicator of the occurrence of 283 TEV adaptation. However, the usefulness of fluorescence for determining the integrity of the eGFP marker was limited in both hosts, by (i) the high levels of autofluorescence in the 284 285 highly symptomatic N. benthamiana leaves, and (ii) the patchy fluorescence in the D. 286 stramonium tissue. Therefore, unlike for TEV-eGFP in N. tabacum, the fluorescent marker 287 was of limited use here. Nevertheless, all N. benthamiana lineages appeared to have some 288 fluorescence until the end of the evolution experiment, and we observed a loss of fluorescence 289 in only 1/10 D. stramonium lineages in the third 9-week passage.

After each passage, RNA was extracted from the collected leaf tissue, and RT-PCR with primers flanking the *eGFP* insert was performed. This RT-PCR assay can therefore detect deletions in the *eGFP* gene, even when deletions extend well into the downstream HC-Pro cistron [19]. In general, the RT-PCR results confirmed the fluorescence microscopy results: A large deletion was detected only in the one *D. stramonium* lineage with a loss of 295 fluorescence (Fig. 2A; 9-weeks passage 2 L8). This deletion variant went to a high frequency 296 in the subsequent passage (Fig. 2A; 9-weeks passage 3 L8). For *N. benthamiana* lineages, we 297 did detect a low-frequency deletion in the eGFP cistron in one lineage (Fig. 2B; 6-weeks 298 passage 4 and 5 L4), but this deletion is so large that this variant is most likely no longer 299 capable of autonomous replication. The deletion size is around 1500 nt, which means that 300 after deleting the entire eGFP, around 800 nt are deleted from HC-Pro, which has a size of 301 1377 nt in total. This deletion extends well into the central region of HC-Pro, beyond the 302 well-conserved FRNK box, which is essential for virus movement and RNA-silencing 303 suppressor activity [31, 32]. We performed an extra round of passaging with all N. 304 benthamiana lineages to check whether this variant would remain at a low frequency, and 305 found exactly this result (Fig. 2B; 6-week passage 6 L4). Furthermore, we detected a small 306 deletion in one lineage (Fig. 2B; 6-week passage 5 and 6 L1) that was maintained at a low 307 frequency in subsequent passages of the virus population.

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#### 309 Whole-genome sequencing of the evolved lineages

All evolved and the ancestral TEV-eGFP lineages were fully sequenced by Illumina 310 311 technology (SRA accession: SRP075180). The consensus sequence of the ancestral TEV-312 eGFP population was used as a reference for mapping the evolved lineages. The deletion 313 observed by RT-PCR (Fig. 2A) in one of the D. stramonium lineages was confirmed by a low 314 number of reads mapping inside the *eGFP* region (median coverage *eGFP*: 111.5), compared 315 to a higher average coverage outside this region (median coverage P1 gene: 19190, median 316 overall genome coverage: 18460). The large deletion included the N-terminal region of HC-317 *Pro*, as observed for other deletions that occur after gene insertions before this gene [19, 34]. 318 For all other lineages in *D. stramonium* and *N. benthamiana*, coverage over the genome was

319 largely uniform and similar to the ancestral virus population, indicating that there were indeed320 no genomic deletions present at appreciable frequencies.

321 Single nucleotide mutations were detected from a frequency as low as 1%, comparing the 322 evolved TEV-eGFP lineages in N. benthamiana and D. stramonium to the ancestral 323 population (Fig. 3). This detection was also performed for evolved TEV-eGFP lineages in N. 324 tabacum, that were sequenced in a previous study [19] (SRA accession: SRP075228). In the 325 evolved N. benthamiana lineages 165 unique mutations were found, with a median of 34.5 326 (range: 27 - 47) mutations per lineage. In the evolved D. stramonium lineages 239 unique 327 mutations were found, with a median of 31.5 (range: 16 - 35) mutations per lineage. In the evolved N. tabacum lineages, 183 unique mutations were found, with a median of 21.5 328 329 (range: 17 - 36) mutations per lineage. Note that the single nucleotide mutations detected 330 here can be fixed (frequency > 50%) in the evolved lineages, as the detection was done over 331 the ancestral population. Hence, it allows us to compare the mutations that arose by evolving 332 TEV-eGFP in the different hosts.

We detected only one mutation (U6286C; CI/Y2096H) that is shared between all three hosts. However, this mutation was present at a low frequency and not detected in all *D. stramonium* and *N. tabacum* lineages (Fig. 3 and Table 1). The *N. benthamiana* and *D. stramonium* lineages share more mutations (15) than either *N. benthamiana* or *D. stramonium* share with *N. tabacum* (4 and 9, respectively). However, most of these mutations are present in only a few lineages and at low frequency (Fig. 3 and Table 1).

The synonymous mutations U7092C, A7479C and A8253C, that are shared between *D. stramonium* and *N. tabacum*, are present in the highest number of lineages and reach higher frequencies comparing all shared mutations detected in the three hosts. Despite of these mutations already being present in the ancestral population, the frequencies at which these mutations are present display interesting patterns. In both *D. stramonium* and *N. tabacum* the mutations A7479C and A8253C are always detected at the same frequency within each lineage, suggesting a strong linkage between these mutations (Additional file 1: Fig. S1). Furthermore, the U7092C mutation never appears together with the former two mutations (Additional file 1: Fig. S1), suggesting that this mutation occurs in another haplotype and that there may be sign epistasis between these two combinations of synonymous mutations. Interestingly, the ancestral U7092C, A7479C and A8253C mutations were not detected in the *N. benthamiana* lineages, demonstrating the differences in host-pathogen interactions.

Host-specific mutations were mostly found in the evolved TEV-eGFP lineages of *N*. *benthamiana* (Fig. 3 and Table 2). In this host, a total number of 7 specific mutations were detected, all of them being nonsynonymous. In *D. stramonium* no host-specific mutations were detected. And in *N. tabacum* only one host-specific mutation was detected in the 3'UTR (Table 2). Note that host specific mutations were defined as mutations detected in at least half of the evolved lineages. For more information on the mutations found in the three hosts please see Additional file 2: Tables S1-S3.

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# 359 Viral accumulation and within-host competitive fitness

We measured virus accumulation 10 dpi, by RT-qPCR for a region within the coat protein gene (*CP*). In both host species, we found no statistically significant differences (*t*-test with Holm-Bonferroni correction) between TEV, TEV-eGFP and the lineages of TEV-eGFP evolved in that host (Fig. 4).

We then measured within-host competitive fitness by means of head-to-head competition experiments with TEV-mCherry, a virus with a different marker but similar fitness to TEVeGFP [21]. Here we observed interesting differences between TEV and TEV-eGFP in the two different hosts. Whereas the TEV-eGFP had lower fitness than the wild-type virus in *D. stramonium* (Fig. 5, compare TEV and ancestral TEV-eGFP; *t*-test:  $t_4 = 13.438$ , *P* < 0.001), there was no difference in *N. benthamiana* (Fig. 5, compare TEV and ancestral TEV-eGFP; *t*test:  $t_4 = -1.389$ , P = 0.237). Our results therefore suggest that although there is a fitness cost associated with the *eGFP* cistron in *N. tabacum* [19] and *D. stramonium*, there is none in *N. benthamiana*. Interestingly, *N. tabacum* and *N. benthamiana* are more closely related to each other than either species is to *D. stramonium*, and yet the host species has a strong effect on the costs of a heterologous gene.

375 For the lineages evolved in D. stramonium, only for 1/10 lineages there was a significant 376 increase in competitive fitness compared to the ancestral TEV-eGFP observed (Fig. 5, L8; t-377 test with Holm-Bonferroni correction:  $t_4 = -6.890$ , P = 0.002). This lineage is the only one to 378 have a deletion in the eGFP insert. In N. benthamiana, 1/6 lineages had a significant increase 379 in within-host fitness (Fig. 5, L4; *t*-test with Holm-Bonferroni correction:  $t_4 = -5.349$ , P =380 0.006). However, this increase in fitness probably is not associated with the large genomic 381 deletion for three reasons: (i) the wild-type TEV without the eGFP cistron has a similar 382 fitness compared to the ancestral TEV-eGFP, suggesting no deletions in eGFP would be 383 beneficial, (ii) the RT-PCR results show that this variant occurs at a low frequency in the 384 population, and therefore is unlikely to effect strongly the results of the competition assay, 385 and (*iii*) this deletion variant remains at low frequency during the next round of passaging 386 (Fig. 2B), suggesting that while frequency-dependent selection might occur, its fitness is not 387 higher than the coevolving full-length TEV-eGFP. Moreover, another lineage of N. 388 benthamiana where we did not detect any deletions, also appeared to have increased in fitness (Fig. 5, L6; *t*-test:  $t_4 = -4.0792$ , P = 0.015), however, after the Holm-Bonferroni correction 389 390 not significantly. Interestingly, the lineage that did increase its fitness significantly (L4) is the 391 only lineage that contains mutations in the 6K2 protein in this host (Additional file 2: Table 392 S1). Therefore we speculate that single-nucleotide variation is one of the main driving forces 393 for an increase in TEV-eGFP fitness in N. benthamiana.

These fitness measurements show that most lineages failed to adapt to the new host species. However, in the two cases that there were significant fitness increases, the underlying genetic changes were consistent with the expected route of adaptation. In *D. stramonium*, where *eGFP* has a high fitness cost, this sequence was deleted. In *N. benthamiana*, where *eGFP* apparently has not fitness cost, host-specific single-nucleotide variation was observed.

399

#### 400 **Discussion**

401 We set out to explore the hypothesis that differences in virulence for different hosts could 402 have an effect on the rate of virus adaptation in each host [13]. Although we find this 403 hypothesis simple and provocative, the observed patterns in our experiments suggest that even 404 in a controlled laboratory environment, reality will often be complex and hard to predict. We 405 used a virus expressing an eGFP fluorescent marker in the hope that the loss of this marker 406 could serve as a real-time indicator of adaptation. However, there were complications with 407 this method, and a loss of fluorescence was only observed in a single D. stramonium lineage. 408 RT-PCR and Illumina sequencing confirmed the loss of the eGFP marker in this case, and its 409 integrity in all other lineages. The data of our competitive fitness assay demonstrate why the 410 maker sequence was probably rather stable in *N. benthamiana*; *eGFP* does not appear to have 411 a cost in this host species, in sharp contrast to the strong fitness cost observed in D. 412 stramonium as well as previously observed in the more closely related host N. tabacum [19]. 413 We expect that the marker will eventually be lost, but only due to genetic drift and therefore 414 at a slow rate.

What mechanisms might underlie the difference in the fitness costs of *eGFP* marker in these two host plants? In a previous study, we showed that the loss of the *eGFP* marker occurred more rapidly as the duration of each passage was increased [19]. During long passages transmission bottlenecks are more spaced on time, and much larger census population sizes 419 are reached. However, there is also much greater scope for virus movement into the newly 420 developing host tissues. As for N. tabacum [19], here we again observed that the eGFP 421 marker does not affect virus accumulation, whereas it does lower competitive fitness in D. 422 These observations suggest that the effects of the *eGFP* marker on virus stramonium. 423 movement are the main reason for selection against the marker. However, marker loss in D. 424 stramonium appears to occur much slower compared to N. tabacum [19], indicating poor 425 virus adaptation to this alternative host. Given the high virulence of TEV for *N. benthamiana*, 426 including strong stunting, there will be limited virus movement during infection and thus 427 significantly lower population census sizes. Hence, we speculate that in N. benthamiana the 428 limited scope for virus movement and accumulation -due to the virus' virulence itself- might 429 mitigate the cost of the *eGFP* marker. Alternatively, cell-to-cell and systemic virus 430 movement in *N. benthamiana* might be so slow that the addition of the *eGFP* marker matters 431 little. A slow systemic virus movement may also explain why in the second round of 432 infection four lineages failed to re-infect N. benthamiana, as initial virus accumulation 433 appeared to be very low until possible virus adaptation by means of point mutations occurred. 434 These results are at odds with our expectations, but they nevertheless have some interesting 435 implications. First, host species changes can apparently ameliorate the costs of exogenous 436 genes. Although strong virus genotype-by-host species interactions have been previously 437 shown for TEV [17], we did not anticipate that a such a simple difference (the presence of 438 eGFP) could also be subjected to such an interaction. These results suggest that when 439 considering the evolution of genome architecture, host species might play a very important 440 role, by allowing evolutionary intermediates to be competitive. For example, for TEV we 441 have shown that the evolution of an alternative gene order through duplication of the NIb 442 replicase gene is highly unlikely, as this intermediate step leads to significant decreases in fitness, making the trajectory to alternative gene orders inaccessible [34]. If NIb duplication 443

444 has a similar interaction with host species as the eGFP insert has, then an alternative host 445 species could act as a stepping-stone and hereby increase the accessibility of the evolutionary 446 trajectory to alternative gene orders. Similar effects of environmental change have been noted 447 in other studies [35]. The generality of these results has not been addressed vet using other 448 viruses with altered genome architecture, but the possibilities are tantalizing. Second, our 449 results could also have implications for assessing the biosafety risks of the genetically 450 modified organisms. Our results suggest that extrapolating fitness results from a permissive 451 host to alternative hosts can be problematic, even when the scope for unexpected interactions 452 appears to be limited, as would be the case for the addition of eGFP expression. In other 453 model systems, unexpected interactions between heterologous genes and host species have 454 also been reported [36].

455 Our results were not consistent with the hypothesis that high virulence could slow down the 456 rate of adaptation, as in each host only a single lineage had evolved higher fitness. The low 457 rate of adaptation observed was consistent with a previous report [18], although we used 458 passages of a longer duration here and had therefore expected more rapid adaptation [19]. 459 Given the low rate at which lineages adapted in this experiment, however, we do not consider 460 that our results provide strong evidence against the hypothesis. Nevertheless, our results do 461 stress that differences in host biology can have a much stronger effect on evolutionary 462 dynamics than differences in virus-induced virulence between host species. An alternative 463 way to tackling the question of the effects of virulence on adaptation might be to use a 464 biotechnological approach; hosts which have different levels of virulence can be engineered, 465 to ensure the main difference between host treatment is microparasite-induced virulence. For 466 example, plant hosts could be engineered to express antiviral siRNAs at low levels. Such an 467 approach would allow for a more controlled test of the hypothesis suggested here, whilst 468 probably not being representative for natural host populations. On the other hand, such 469 experiments could perhaps help shed light on the effects of virulence on adaptation in470 agroecosystems or vaccinated populations.

471

### 472 **Conclusions**

473 A host species jump can be a game changer for evolutionary dynamics. A non-functional 474 exogenous sequence -eGFP- which is unstable in its typical host, has shown to be more 475 stable in two alternative host species for which TEV has both lower and higher virulence than 476 in the typical host. In addition, eGFP does not appear to have any fitness effects in the host 477 for which TEV has high virulence. These observations clashed with the hypothesis that high 478 virulence slows down the rate of adaptation. Moreover, when considering the evolution of 479 genome architecture, host species jumps might play a very important role, by allowing 480 evolutionary intermediates to be competitive.

481

#### 482 **Declarations**

- 483 Ethics approval and consent to participate
- 484 Not applicable
- 485
- 486 **Consent for publication**
- 487 Not applicable
- 488

### 489 Availability of data and material

- 490 The raw read data from Illumina sequencing is available at SRA with accession: SRP075180.
- 491 The fitness data has been deposited on LabArchives with doi: 10.6070/H4N877TD.
- 492
- 493 Competing interests

494 The authors declare that they have no competing interests

495

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504

### 505 Author's contributions

AW, MPZ and SFE designed the study. AW and MPZ performed the experiments. AW, MPZ
and SFE analyzed the data and wrote the manuscript. All authors read and approved the final

508 manuscript.

509

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512

### 513 Additional files

#### 514 Additional file 1:

515 Figure S1. Frequency of mutations found in both *D. stramonium* and *N. tabacum*. Mutations 516 detected in both *D. stramonium* and *N. tabacum* that were present in all the lineages of either 517 one of these hosts. The frequency of these mutations in either the ancestral population (anc)

- 518 or the different lineages (L1-L10) is given by the color-coded points. The points are 519 connected by the broken lines to emphasize the trend in the data.
- 520
- 521 Additional file 2:
- 522 Table S1. Mutations detected in the *Nicotiana bentiamiana* lineages as compared to the523 ancestral lineage
- 524 **Table S2.** Mutations detected in the *Datura stramonium* lineages as compared to the 525 ancestral lineage
- 526 Table S3. Mutations detected in the *Nicotiana tabacum* lineages as compared to the ancestral527 lineage
- 528

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620

# 621 Tables

# 622 **Table 1.** TEV-eGFP mutations shared in the different hosts

			N. benthamiana		D. stramonium		N. tabacum	
Nucleotide	Amino acid		Number of		Number of		Number of	
change	change	Gene	lineages	Frequency range	lineages	Frequency range	lineages	Frequency range
U6286C	Y2096H	CI	6/6	0.013 - 0.131	2/10	0.012 - 0.031	4/10	0.019 - 0.140
A208G	M70V	P1	1/6	0.012	1/10	0.010	-	-
C1039U	H347Y	P1	1/6	0.013	1/10	0.035	-	-
G1332A	M444I	eGFP	1/6	0.176	1/10	0.139	-	-
U1556G*	V519G	eGFP	1/6	0.011	6/10	0.010 - 0.016	-	-
U1836G	synonymous	HC-Pro	5/6	0.093 - 0.108	1/10	0.089	-	-
A1917G	synonymous	HC-Pro	1/6	0.015	1/10	0.017	-	-
A6278G	E2093G	CI	1/6	0.012	2/10	0.014 - 0.031	-	-
C6547U	H2183Y	VPg	1/6	0.013	1/10	0.014	-	-
U6747C	synonymous	VPg	1/6	0.012	1/10	0.110	-	-

A6776G	D2259G	VPg	2/6	0.533 - 0.746	1/10	0.023	-	-
G6803A	S2268N	VPg	1/6	0.014	1/10	0.024	-	-
A6438G	synonymous	6K2	1/6	0.012	1/10	0.024	-	-
C8405G*	T2802R	NIb	5/6	0.010 - 0.018	5/10	0.010 - 0.016	-	-
U9474C	synonymous	СР	1/6	0.013	1/10	0.061	-	-
C9837U	synonymous	СР	1/6	0.070	1/10	0.010	-	-
U3803C	I1268T	P3	2/6	0.656 - 0.881	-	-	2/10	0.010 - 0.019
U3872C	V2191A	P3	1/6	0.064	-	-	1/10	0.016
G4411A	V1471I	CI	1/6	0.030	-	-	1/10	0.016
C4989U	synonymous	CI	1/6	0.018	-	-	1/10	0.066
C548U	T183I	P1	-	-	1/10	0.011	1/10	0.024
G2928A*	synonymous	HC-Pro	-	-	1/10	0.017	1/10	0.999
U7092C*	synonymous	NIa-Pro	-	-	10/10	0.091 - 0.755	2/10	0.176 - 0.999
A7479C*	synonymous	NIa-Pro	-	-	10/10	0.132 - 0.790	7/10	0.960 - 0.999
A7567G	K2523E	NIa-Pro	-	-	4/10	0.012 - 0.053	10/10	0.015 - 0.871

G7710A	synonymous	NIa-Pro	-	-	1/10	0.014	1/10	0.024
A8253C*	synonymous	NIb	-	-	10/10	0.136 - 0.801	7/10	0.805 - 0.998
G9117A	synonymous	NIb	-	-	1/10	0.321	2/10	0.022 - 0.025
U9249C	synonymous	NIb	-	-	2/10	0.011 - 0.231	1/10	0.040
* 1	:							

\*Also detected in the ancestral population

	Nucleotide	Amino acid		Number of	Frequency
	change	change	Gene	lineages	range
N. benthamiana	G3797A	G1266E	P3	3/6	0.291 - 0.664
	G4380U	E1460D	6K1	3/6	0.012 - 0.093
	U4387C	Y1463H	6K1	4/6	0.011 - 0.016
	C4391U	T1464M	6K1	6/6	0.041 - 0.138
	G4397A	S1466N	CI	6/6	0.012 - 0.019
	A6771U	L2257F	VPg	4/6	0.027 - 0.201
	G8909U*	W2970L	NIb	5/6	0.026 - 0.042
D. stramonium	-	-	-	-	-
N. tabacum	G10253A		3'UTR	10/10	0.025 - 0.040

# 623 **Table 2.** Host specific mutations in the evolved TEV-eGFP lineages.

<sup>\*</sup>Also detected in the ancestral population.

624

# 625 Figure Legends

Figure 1. Schematic representation of TEV-eGFP. The *eGFP* gene is located between *P1*and *HC-Pro* cistrons. Proteolytic cleavage sites were provided at both ends of *eGFP*.

628

629 Figure 2. Deletion detection in the *eGFP* gene. Agarose gels with RT-PCR products of the region encompassing the eGFP gene. The TEV and TEV-eGFP are shown for comparative 630 631 purposes. The negative controls are healthy plants and PCR controls (C-). (A) TEV-eGFP in 632 D. stramonium has 10 independent lineages (L1-L10). A deletion encompassing the eGFP 633 gene was detected in one lineage (L8) in the second 9-week passage. This deletion went to a 634 high frequency in the subsequent passage. (B) TEV-eGFP in N. benthamiana has six independent lineages (L1-L6). A deletion bigger than the size of eGFP was detected in one 635 636 lineage (L4) in the fourth 6-week passage. This deletion was not fixed in the two subsequent 637 passages. A small deletion was detected in the fifth and sixth 6-week passage in L1.

638

639 Figure 3. Genomes of the TEV-eGFP lineages evolved the three different hosts as compared 640 to the ancestral lineage. Mutations were detected using NGS data of the evolved lineages 641 (L1-L10), as compared to their ancestral population. The square symbols represent mutations 642 that are fixed (> 50%) and the circle symbols represent mutations that are not fixed (< 50%). 643 Filled symbols represent nonsynonymous substitutions and open symbols represent 644 synonymous substitutions. The triangle symbols represent mutations that are present in either 645 the 3'UTR or 5'UTR. Black substitutions occur only in one lineage, whereas color-coded 646 substitutions are repeated in two or more lineages. Note that the mutations are present at 647 different frequencies as reported by VarScan 2. Grey boxes with continuous black lines 648 indicate genomic deletions in the majority variant of the virus population. The grev 649 transparent box with dotted black lines in L4 of N. benthamiana indicates a genomic deletion

in a minority variant. The latter box was drawn to indicate the size of the deletion, assuming
that the deletion starts at the first position of *eGFP*. For more information on the frequency
of the mutations please see Additional file 2: Tables S1-S3.

653

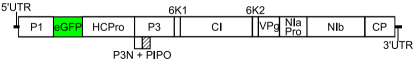
**Figure 4.** Virus accumulation of the evolved and ancestral lineages. Virus accumulation, as determined by accumulation experiments and RT-qPCR at 10 dpi, of TEV, the ancestral TEV-eGFP, and the evolved TEV-eGFP lineages in the corresponding hosts. TEV and the evolved lineage with a deletion in the *eGFP* gene are indicated with the orange bars. The ancestral TEV-eGFP and the evolved lineages with an intact *eGFP* gene are indicated with the green bars. Error bars represent SD of the plant replicates.

660

661 **Figure 5.** 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 662 663 respect to a common competitor; TEV-mCherry. W was determined at 10 dpi, of TEV, the 664 ancestral TEV-eGFP, and the evolved TEV-eGFP lineages in the corresponding hosts. TEV 665 and the evolved lineage with a deletion in the *eGFP* gene are indicated with the orange bars. 666 The ancestral TEV-eGFP and the evolved lineages with an intact eGFP gene are indicated 667 with the green bars. The orange asterisks indicate statistical significant differences of the 668 evolved lineages as compared to TEV (*t*-test with Holm-Bonferroni correction). The green 669 asterisks indicate statistical significant differences of the evolved lineages as compared to the 670 ancestral TEV-eGFP (*t*-test with Holm-Bonferroni correction). Error bars represent SD of the 671 plant replicates.



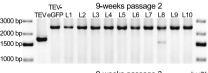
TEV-eGFP

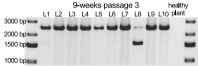


#### Figure 2

Α

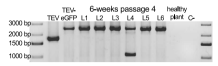
#### Datura stramonium





#### В

#### Nicotiana benthamiana



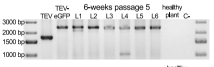
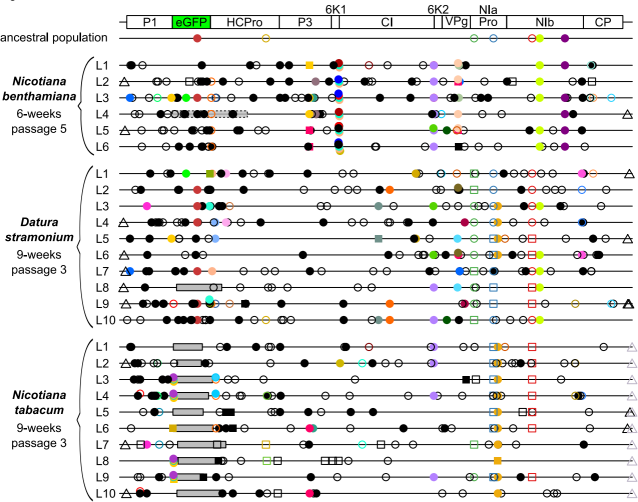
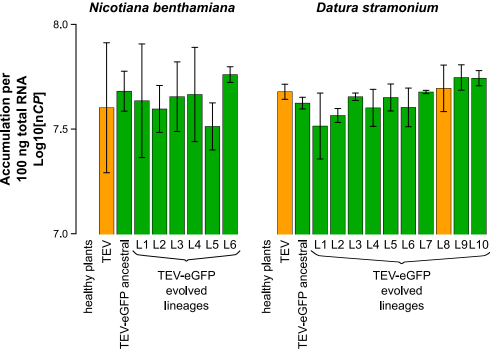




Figure 3







#### Datura stramonium



#### Nicotiana benthamiana

Datura stramonium

