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Assessing the impact on virus transmission and insect vector
behavior of a viral mixed infection in melon

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1 **Summary**

2 Mixed viral infections in plants are common, and can cause synergistic or
3 antagonistic interactions. Except in complex diseases with severe symptoms, mixed
4 infections frequently remain unnoticed, and their impact on insect vector transmission is
5 largely unknown. In this work, we considered mixed infections of two unrelated viruses
6 commonly found in melon plants, the crinivirus *Cucurbit yellow stunting disorder virus*
7 (CYSDV) and the potyvirus *Watermelon mosaic virus* (WMV), and evaluated their
8 vector-transmission by whiteflies and aphids, respectively. Their dynamics of
9 accumulation were analyzed until 60 days post-inoculation (dpi) in mix-infected plants,
10 finding reduced titers of WMV and much higher titers of CYSDV compared to single
11 infections. At 24 dpi, corresponding to the peak of CYSDV accumulation, similar
12 whitefly transmission rates were obtained when comparing either individual or mix-
13 infected plants as CYSDV sources, although its secondary dissemination was slightly
14 biased towards plants previously infected with WMV, regardless of the source plant.
15 However, at a later time point mix-infected plants partially recovered from the initially
16 severe symptoms, and CYSDV transmission became significantly higher. Interestingly,
17 aphid transmission rates both at early and late time points were unaltered when WMV
18 was acquired from mix-infected plants despite its reduced accumulation. This lack of
19 correlation between WMV accumulation and transmission could result from
20 compensatory effects observed in the analysis of the aphid feeding behavior by Electrical
21 Penetration Graphs. Thus, our results showed that mix-infected plants could provide
22 advantages for both viruses, directly favoring CYSDV dissemination while maintaining
23 WMV transmission.

24

1 **Introduction**

2 Transmission to susceptible hosts is a key step in the cycle of pathogens that relies
3 mainly on the activity of insect vectors, like many plant viruses (Whitfield, Falk, and
4 Rotenberg 2015). As in other parasitic relationships, plant pathogenic viruses could affect
5 the host-vector interaction in order to improve their transmissibility, inducing changes in
6 the host morphology or physiology (Culver and Padmanabhan 2007), or manipulating the
7 vector behavior (Lefèvre et al. 2009a; Mauck 2016a; Mauck, Chesnais, and Shapiro
8 2018). Here, we aimed to evaluate how mixed viral infections in plants could affect vector
9 transmission traits.

10 Plant viruses are distributed worldwide having a significant impact on plant survival and
11 production of crops (Alexander et al. 2014). Considering the important ecological roles
12 played by viruses, their relationships with hosts and organisms acting as vectors are being
13 explored (Eigenbrode et al. 2002; Mauck, De Moraes, and Mescher 2010; Mauck et al.
14 2015). Indeed, recent reports documented cases in which plant viruses can manipulate
15 their hosts, and consequently modify the attractiveness for vectors and modulate
16 transmission (Peñaflor et al. 2016; Mauck et al. 2012). Observed effects include
17 modification of host defense responses (Petek et al. 2014; Zhang et al. 2012), changes in
18 the nutritional quality of the plant (Alexander et al. 2014; Su et al. 2015), or alterations
19 of visual traits and production of volatiles conditioning attraction (Ingwell, Eigenbrode,
20 and Bosque-Pérez 2012; Liu et al. 2013). Since transmission to new hosts determines in
21 part the fitness of the pathogen, vector performance may have a direct impact on the
22 ecology and evolution of the viruses they transmit (Gutiérrez et al. 2013; Stafford,
23 Walker, and Ullman 2011).

24 Nevertheless, our understanding of how such interactions function in complex ecological
25 environments like natural and agricultural ecosystems is still limited. In these situations,
26 plants frequently interact with several pathogens simultaneously, and the composition
27 and dynamics of ecological communities can be significantly affected (Mauck, De
28 Moraes, and Mescher 2015; Mauck et al. 2015; Stout, Thaler, and Thomma 2006). In
29 particular, mixed infections of plant viruses are common, and current evidence suggests
30 that mixed infections are the rule and not the exception in nature (Mascia and Gallitelli
31 2016; Roossinck, Martin, and Roumagnac 2015; DaPalma et al. 2010). In addition,
32 several important viral diseases of plants are the outcome of interactions between several
33 viral agents (Syller 2012).

1 In nature, viral mixed infections can be generated by the transmission of different viruses
2 by different vectors, or by the same vector carrying the different viruses (Syller 2014).
3 The result of a mixed infection varies according to the type of interaction between the
4 viruses involved, from synergism to antagonism (Mascia and Gallitelli 2016; Syller
5 2012). In short, in a synergistic interaction, at least one of the viruses is benefited by the
6 presence of the other(s). The benefit is often quantifiable as an increase in the viral titer,
7 and consequently sometimes the resulting symptoms are more severe compared to a
8 single infection. On the contrary, during an antagonistic interaction, the presence of more
9 than one virus is detrimental for at least one of them, and this is often reflected in a
10 decrease of the viral titer. The order of infection might influence the outcome in virulence
11 and titers (Saldaña, Elena, and Solé 2003; Chávez-Calvillo et al. 2016).

12 As mentioned, few researchers have explored mixed viral infections with respect to their
13 effect on host-pathogen-vector relationships in agricultural ecosystems (Salvaudon, De
14 Moraes, and Mescher 2013; Syller 2012; Syller and Grupa 2016). In the present work we
15 focused our research in a mixed viral infection to shed some light in the possible
16 ecological outcomes regarding vector transmission. The work was carried out in melon
17 (*Cucumis melo*, L.), an economically relevant crop with global production close to 30
18 million tons in 2016 (www.fao.org). As in other crops, a significant part of the potential
19 production of melon could be lost due to diseases caused by different pathogens,
20 including viruses (Oerke 2006). Moreover, field surveys have reported high occurrence
21 of multiple viral infections in melon and other cucurbits (Juarez et al. 2013). To bring our
22 research closer to current problems, we worked with two viruses that frequently infect
23 melon: the crinivirus *Cucurbit yellow stunting disorder virus* (CYSDV), and the
24 potyvirus *Watermelon mosaic virus* (WMV), transmitted respectively by whiteflies and
25 aphids (Desbiez et al. 2011; Navas-Castillo, López-Moya, and Aranda 2014). Nowadays,
26 CYSDV can be found throughout many tropical and subtropical areas where cucurbits
27 are cultivated (Wintermantel et al. 2017). It has a bipartite single-strand plus-sense RNA
28 genome. As other criniviruses, CYSDV is restricted to the phloem of the susceptible
29 plants, being transmitted by whiteflies, such as *Bemisia tabaci* Gennadius, in a
30 semipersistent manner (Célix et al. 1996; Navas-Castillo, López-Moya, and Aranda
31 2014). On the other hand, WMV (formerly also known as WMV2) presents a worldwide
32 distribution, predominantly in temperate regions, and has a large host range, being able
33 to infect more than 170 plant species (Desbiez and Lecoq 2004). As other potyviruses,
34 WMV presents a positive-sense, single-stranded RNA genome that is translated into a

1 polyprotein subsequently processed by viral-encoded proteinases into functional products
2 (Valli, García, and López-Moya 2015). The infection occurs in most plant tissues, and its
3 transmission is mediated by different aphids, including *Myzus persicae* Sulzer, in a
4 nonpersistent manner (Díaz-Pendón et al. 2005).

5 The objective of our study was to evaluate mixed infections of CYSDV and WMV in
6 melon, in particular to know if the presence of both viruses could influence their vector
7 transmission. To answer this, we first compared the viral load of each virus either in single
8 or in mix-infected plants, following their accumulation along a period of 60 days, and
9 complemented the quantification measurements with observations of symptoms. Then,
10 transmission experiments were performed under laboratory conditions to evaluate the
11 influence of different plants (single- or mix-infected) as virus sources. Furthermore, the
12 crinivirus transmission was also analyzed under field conditions. Regarding the
13 transmission of WMV, the acquisition from single or mix-infected plants was compared
14 using Electrical Penetration Graph (EPG) to measure the number and duration of the
15 different phases and sub-phases.

16 Our results reveal that mixed infections could affect the transmissibility of these two
17 viruses. This knowledge could provide scientific-based advice to minimize virus damage,
18 thus improving melon production and quality. More generally, our findings may help to
19 recognize the importance and ecological impact of mixed viral infections on vector
20 transmission and virus dissemination.

21 22 **Results**

23 **Mix-infected plants accumulated more CYSDV and less WMV compared to single** 24 **infections**

25 Levels of CYSDV and WMV were compared between single and mix-infected
26 plants during 60 days after virus inoculation. To mimic natural conditions, infected plants
27 were obtained using viruliferous whiteflies for CYSDV and aphids for WMV. Samples
28 were collected every 12 days to quantify viral loads. In all time-points the titer of CYSDV
29 was higher in mix-infected compared to single infected plants, with the major peak of
30 accumulation occurring at 24 dpi during mixed infections (Fig. 1). On the other hand,
31 WMV titers were lower in mix-infected plants in comparison to single infections, for
32 most of the analyzed time points. Statistical comparisons were performed first for paired
33 values, using t-test for those cases where normality and homoscedasticity was confirmed,

1 and Mann-Whitney for the rest, finding significance at all time points except at 12 dpi
2 (not shown). The dynamics of the curves of virus accumulations were also analyzed by
3 ANOVA, finding dependency of viral accumulation both for single vs. mixed infections
4 in the two cases (CYSDV and WMV), and of time for CYSDV.

5
6 **Mix-infected plants at late time points recovered from the severe symptoms caused**
7 **by WMV alone**

8 Observation of symptoms were performed during 60 days to compare single and
9 mix-infected plants. In our growing conditions plants with individual infections showed
10 the expected symptoms described for each virus: severe mosaic and puckering with
11 distortion of all leaves in WMV infected plants; and mild leaf mottling, interveinal
12 chlorosis and yellowing, mainly visible in older leaves in CYSDV infected plants (Fig
13 2a). Interestingly, symptoms in mix-infected plants were very similar to those of a single
14 infection by WMV during approximately the first half of the experiment (Fig 2b), but
15 during the following thirty days, plants recovered and moderately regained growth and
16 vigor, with a final aspect that was more similar to CYSDV single infected plants (Fig 2b
17 and c). The same pattern of symptoms development was observed and reproduced in all
18 repetitions of the experiments.

19
20 **CYSDV transmission rates were similar at 24 dpi when comparing single or mixed**
21 **infected source plants, and both under controlled or field conditions**

22 To determine if the mixed infections had some impact on the dispersal of the
23 viruses by their natural vectors we performed transmission experiments.

24 First, we analyzed the transmission of CYSDV under controlled laboratory conditions,
25 comparing whiteflies that acquired the virus either in single or in mix-infected plants.
26 Plants at 24 dpi were chosen as virus sources due to the highest CYSDV accumulation in
27 mix-infected plants (Fig 1). Insects were allowed to acquire the virus during 48 h on
28 infected plants, and then the viruliferous whiteflies were transferred to uninfected
29 receptor plants for virus inoculation during 48 h. The transmission rates were determined
30 at 18 dpi, testing CYSDV presence by RT-PCR. No significant differences were found
31 (Chi-square) between transmission rates for each individual experiment when the
32 whiteflies acquired the virus in mixed infected compared to single infected plants (Fig 3),
33 although the pooled values reached a significative ($P < 0.05$) higher transmission rate for

1 CYSDV, a result totally unexpected if we take into account the much lower viral load of
2 CYSDV alone compared to CYSDV+WMV at the chosen time point for the acquisition.
3 Transmission experiments were also performed in field conditions during two
4 consecutive years (see Experimental Procedures) in order to determine if the transmission
5 rates of CYSDV were influenced either by the source plants (single or mixed infected at
6 24 dpi) and/or by the receptor plants (uninfected or WMV-infected). In these field
7 conditions, the results (Figs 4 and 5, supplemental tables S2 and S3) showed larger
8 differences in transmission rates depending on the year, probably reflecting higher
9 mobility and activity of the vectors caused by diverse environmental clues (most likely
10 temperature), thus the comparisons were done separately for each year. Images of the
11 experimental setup, and the complete set of values are provided as supporting information
12 (Fig S1, Tables S2 and S3).

13 In the first experiment, transmission rates were almost identical when the viral source
14 plants were either single or mix-infected plants, suggesting that the acquisition of
15 CYSDV was not altered by the presence of WMV in mixed infections (Fig 4). Although
16 transmission was slightly higher to WMV-infected than to healthy receptor plants from
17 the two kinds of inoculum sources, the differences were not statistically significant
18 ($t=1.06$ and 1.62 for single or mixed infected source plants).

19 The experimental design in the second year was modified to allow comparison of
20 inoculation choices to either uninfected or WMV-infected plants alternated in every net-
21 structure. The results (Fig 5, sections a and b) confirmed absence of significant
22 differences ($t=1.13$ and 1.06 for single or mixed infected sources) between the
23 transmission rates to WMV-infected plants compared to uninfected plants, but the slight
24 preference of viruliferous whiteflies to transmit the crinivirus to plants previously
25 infected with the potyvirus was observed in two of the three repetitions performed,
26 regardless of the source of virus acquisition. In order to quantify this putative preference
27 to pre-infected plants, a model was tested for the dependance of percentage of infection
28 as linear function of the pre-infected state of the receptor plant, which resulted in 14% of
29 the infection rate variability explained by the model, although without statistical
30 significance ($p=0.23$). Therefore, further data will be required in order to confirm the
31 potential pre-infection effect.

32

33 **Reduced WMV levels in mix-infected plants does not result in less transmission of**
34 **WMV by aphids at 24 dpi**

1 To test if the mixed infection affected transmission of WMV, a different
2 methodology was adopted, taking into consideration the differences of the non-persistent
3 and semipersistent transmission processes. Apterous aphids were fed on a leaf of a single
4 or mixed infected melon plant to acquire the virus, and subsequently manually transferred
5 to test plants for virus inoculation. After insecticide treatment, the plants were observed
6 until appearance of WMV symptoms. Despite the lower viral titers in the mix-infected
7 plants at 24 dpi (Fig 1, and upper panel of Fig 6), the transmission rates when the insects
8 acquired the virus in a mixed infected plant compared to a single infected one were not
9 significantly different in two out of the three individual experiments, neither in the pooled
10 values (Fig 6).

11 **Aphid feeding behavior was altered in mix-infected plants**

12 The feeding behavior of aphids was monitored using EPG to determine if it was
13 influenced by single or mixed infections in the source plants. We fed wired aphids on i)
14 uninfected control plants; ii) single-infected plants with CYSDV; iii) single-infected
15 plants with WMV; and iv) mixed-infected plants with both CYSDV+WMV. The results
16 indicated that the presence of CYSDV alone did not have any major effect on the behavior
17 of aphid vectors compared to control uninfected plants, with the most relevant parameters
18 showing similar values or minor changes not associated with transmission-relevant
19 activities (Table S4). However, we observed that aphids spent more time without probing
20 on plants infected with WMV compared to double-infected plants. Also, longer duration
21 of the intracellular punctures (detectable as potential drops or Pds) was observed in plants
22 infected with CYSDV+WMV, with differences after grouping the data both by insect and
23 by event (Table 5). The statistical analysis of the duration of each of the sub-phases of
24 the Pds confirmed the existence of significant differences in the case of the sub-phase II-
25 3 (Fig 7 a), which was previously associated with acquisition of non-persistently
26 transmitted viruses (Collar, Avilla, and Fereres 1997). The duration of subphase II-3 of
27 the Pds (Fig 7 b, and complete data set in Table S5), and also the number of pulses (Fig
28 7 c) were significantly higher in plants infected with CYSDV+WMV.

29 **At a late time point, mix-infected plants used as source produced similar rates of 30 aphid transmission of WMV, but higher rates of whitefly transmission of CYSDV.**

31 The observed recovery of plants with mixed infections of WMV + CYSDV at late
32 time points (Fig 2) prompted us to perform additional measurements of vector
33
34

1 transmission rates. For these experiments, five groups of 10 test plants were considered
2 as biological repetitions for each one of the four cases: whitefly-transmission of CYSDV
3 and aphid-transmission of WMV, using as virus sources for acquisition either single
4 (CYSDV or WMV, respectively) or mixed (CYSDV+WMV) infected plants at 52 dpi.
5 Results are shown in Fig 8.

6 The transmission of CYSDV was much higher (98%) from mix-infected plants compared
7 to single infected source (62%), with significant differences (Chi-square, $df=1$, $P <$
8 0.0001), strongly contrasting with the results obtained at 24 dpi (Fig 3). Comparing the
9 source plants at these two time points, the recovered double-infected plants correlated
10 with a better performance as virus source for whitefly transmission. On the other hand,
11 WMV transmission rates were 74 and 67 % respectively from single and mixed infected
12 sources, not showing significant differences (Chi-square test, $df=1$, P value 0.4429),
13 reaching values in the same range as those found at 24 dpi (Fig 6), despite the clear
14 differences in WMV accumulation at late time points of infection between single and
15 mix-infected plants.

17 **Discussion**

18 Plants naturally infected by more than one virus has been reported often (Juarez
19 et al. 2013; Peñaflor et al. 2016; Roossinck et al. 2010; Tugume, Mukasa, and Valkonen
20 2016; Zinga et al. 2013). However, not many studies addressed the ecological impact of
21 such complex pathosystems (Elena, Fraile, and García-Arenal 2014). The interactions
22 between hosts, pathogens and vectors could result in an equilibrium with significant
23 effects on each of the three players (Mauck 2016b), as expected in any parasitic
24 interaction (Lefèvre et al. 2009b). Antecedents pointing to the importance of these
25 relationships during mixed infections have been described. For instance, potentially
26 relevant ecological repercussions of complex host-pathogens interactions have been
27 found between unrelated pathogens, like bacteria and viruses (Shapiro et al. 2012, 2013).
28 Concerning viruses, the recently reviewed ecological implications of host manipulation
29 (Mauck, Chesnais, and Shapiro 2018) derive mainly from reports related to single
30 infections (Eigenbrode et al. 2002; Ingwell, Eigenbrode, and Bosque-Pérez 2012; Mauck,
31 De Moraes, and Mescher 2010; Stafford, Walker, and Ullman 2011), although a recent
32 work considered double infections caused by two persistently transmitted viruses (Chen
33 et al. 2018). Repercussions on other combinations of viral infections, such as non-

1 persistently and semipersistently transmitted viruses (Bragard et al. 2013), are mostly
2 unknown.

3 In the case of cucurbits, available surveys revealed that mixed viral infections are quite
4 common (Gil-Salas et al. 2012; Sufrin-Ringwald and Lapidot 2011; Juarez et al. 2013).
5 We focused our work on melon plants infected by two virus that generate important
6 economic losses: WMV and CYSDV (Abrahamian and Abou-Jawdah 2014; Juarez et al.
7 2013). Mixed viral infections were established experimentally using aphids and
8 whiteflies, and the viral load for each individual virus was determined during 60 dpi. Our
9 results showed that CYSDV accumulated to higher titers in mixed compared to single
10 infected plants at all time-points with a peak at 24 dpi, while the viral load of WMV was
11 consistently lower in mix-infected plants than in single infections. These results
12 confirmed similar kinetics reported during mixed infections involving a potyvirus and
13 other plant viruses, including criniviruses: the most common pattern revealed that in
14 mixed infections the accumulation of the potyvirus tend to remain similar or lower
15 compared to single infections, while accumulation increases for the non-potyvirus partner
16 (Murphy and Bowen 2006; Wang et al. 2009). This usual outcome might be due, at least
17 partially, by attenuation of the host RNA silencing-related antiviral defense by the strong
18 activity of the RNA silencing suppressor helper-component protease HC-Pro present in
19 most potyviruses (Valli et al. 2018). This hypothesis is also indirectly supported by the
20 atypical case of the mixed infection between the potyvirus *Sweet potato feathery mottle*
21 *virus* (SPFMV) and the crinivirus *Sweet potato chlorotic stunt virus* (SPCSV) in sweet
22 potato plants, in which the potyvirus does not generate a synergism in the usual direction,
23 a fact that could be related with the almost complete lack of RSS activity in the SPFMV
24 HCPro (Mingot et al. 2016). In the case of a mixed infection of two potyviruses like
25 WMV and *Zucchini yellow mosaic virus* (ZYMV) in squash (*Cucurbita pepo*), authors
26 found that ZYMV replicated at similar rates in single and mixed infections, whereas
27 WMV accumulated to significantly lower levels in the presence of ZYMV, although it
28 was still readily transmitted from mix-infected plants (Salvaudon, De Moraes, and
29 Mescher 2013).

30 The virus-virus interaction during mixed infection is considered essential for the
31 combined disease. Outcomes can be classified as antagonistic or synergistic, with
32 synergisms being caused in many cases by combinations of unrelated viruses (Syller
33 2012). At the same time, the possible effects in the plant caused from mixed infections
34 could range from beneficial, to neutral or detrimental. CYSDV and WMV belong to very

1 distinct viral groups, crinivirus and potyvirus, being transmitted respectively by non-
2 related insects, semipersistently by whiteflies, and non-persistently by aphids. A
3 superficial analysis of these differences can lead to the assumption that they might not
4 share the same niches (such as host tissues being infected), and thus might not compete
5 for vector transmission or host resources, which could lead to synergism rather than
6 antagonism (DaPalma et al. 2010). In fact, this is roughly what we observed until around
7 the midpoint of the kinetics of virus accumulation. Unexpectedly, the situation appeared
8 to be somehow reversed after a prolonged double-infection period, revealing a more
9 complex interaction between the two viruses. The severity of symptoms caused by
10 CYSDV infection have been directly correlated to the viral load in the plant (Eid et al.
11 2006; Marco et al. 2003), and thus the higher titers of CYSDV in our mix-infected plants
12 could be one of the factors leading to stronger disease symptoms during the initial stages
13 of the double infection. However, we observed that mix-infected plants began to recover
14 gradually, starting around 36-48 dpi, and by 60 dpi the symptoms were considerably less
15 severe in the mix-infected plants compared to the ones infected only with WMV.
16 Symptom recovery has been described in some viral infections as related to RNA
17 silencing mechanisms (Lim et al. 2011), but further studies will be needed to determine
18 if this is also the case for CYSDV+WMV.

19 When we analyzed the CYSDV transmission rates to healthy plants in laboratory
20 conditions at 24 dpi, we found no significant differences when comparing single or mix-
21 infected plants as virus sources, despite the higher titers in the second case. In fact, a
22 report showed that accumulation of CYSDV associated with an increase of symptoms
23 might negatively influence transmission, at least in non-cucurbit hosts (Wintermantel et
24 al. 2016). However, the situation was totally reversed at 52 dpi, with mix-infected plants
25 becoming a better source for virus transmission. It is not easy to explain these differences
26 in transmission rates at 24 dpi and 52 dpi, especially considering that the viral load was
27 lower in the late time point (Fig 1). We can speculate that the severely deformed leaves
28 of the mix-infected plants at the early time point could cause steric difficulties for the
29 whiteflies to reach the phloem and acquire the virus, while the larger and less deformed
30 leaves of CYSDV single infected plants were more accessible for vectors. This
31 circumstance might disappear later after the recovery of the plants with WMV+CYSDV,
32 leading to a more comparable topology of leaves where the higher virus load would
33 become determinant for transmission success. Such interactions have been described in
34 certain conditions, for instance when the presence of a satellite modifies transmissibility

1 of the helper virus (Escriu, Perry, and García-Arenal 2000). However, we cannot rule out
2 other explanations. For instance, in other combinations involving phloem-restricted
3 viruses, co-infection with viruses not restricted to the vasculature can remove phloem
4 limitations and make the virus more accessible for vectors, as in the case of *Potato leafroll*
5 *virus* (PLRV) and the potyvirus *Potato virus A* (PVA) (Savenkov and Valkonen 2001).
6 Additional work will be needed to verify these or other hypothesis.

7 When CYSDV transmission was tested in field conditions, similar rates were obtained
8 when comparing single or mixed-infected plants, suggesting that the acquisition process
9 was not affected, regardless of the presence or absence of WMV in the source plants.
10 However, in these experiments we observed a slightly higher transmission towards WMV
11 pre-infected plants compared to uninfected test plants (Fig 4). This observation suggested
12 that the dissemination of CYSDV might be biased towards plants previously infected by
13 potyviruses. Indeed, a repetition of the field experiments allowing free-choice to the
14 viruliferous whiteflies during the inoculation period confirmed that the virus was again
15 preferentially inoculated to WMV pre-infected plants than to uninfected controls (Fig 5
16 a, b). In other words, the influence of the potyvirus presence in mixed infection could be
17 operating not only in the acquisition and transmission of the unrelated crinivirus at late
18 time points, but also in the inoculation process to favor mixed infections. Such effects of
19 vector attraction are more likely to be expected in semipersistently transmitted viruses,
20 while for non-persistently transmitted viruses such as WMV, effects on acquisition will
21 be more relevant for the transmission outcome. The explanation of this possible
22 preference for WMV pre-infected plants during inoculation of CYSDV by viruliferous
23 whiteflies will require further experimentation, but we can hypothesize that mechanisms
24 like a reduced expression of JA-responsive genes, shown in potyvirus-infected plants
25 (Westwood et al. 2014), could increase attraction to whiteflies.

26 The results of WMV transmission experiments showed mostly unaltered rates both at
27 early and late time points, regardless of the observed changes in viral load with reduced
28 virus accumulation in mix-infected plants. The transmission process was further analyzed
29 by EPG to determine the aphid feeding behavior. This technique has been used in
30 different studies (Garzo et al. 2016; Moreno et al. 2012; Munster et al. 2017), but to our
31 knowledge this is the first case in which plants single or mix-infected with viruses were
32 compared. The behavior of aphids was recorded when feeding on uninfected, CYSDV-
33 infected, WMV-infected, or mixed (CYSDV+WMV) infected plants, and significant
34 differences were found for transmission relevant parameters between WMV-single and

1 double-infected plants. Our results clearly showed that aphids, when feeding on mix-
2 infected plants, spent more time in the subphase correlated with acquisition of non-
3 persistent viruses (Collar, Avilla, and Fereres 1997), and also produced more pulses
4 during intracellular stylet punctures (Pds). Interestingly, no differences in number and
5 duration of Pd waveforms per insect were observed between uninfected plants and single
6 infected plants with either WMV or CYSDV alone, suggesting that the observed changes
7 in aphid behavior on mix-infected plants might be indeed attributed to the simultaneous
8 presence of both viruses. For the future, it remains to be determined if these changes
9 respond to differential attractiveness and/or nutritional conditions of plants, or alterations
10 in the defense mechanisms of plants during mixed infections, and how these specific
11 changes are caused by the simultaneous presence of the two viruses and not by each one
12 individually. It is relevant to recall that the transmission mechanisms are different for
13 criniviruses and potyviruses, which can have implications for the manipulative strategies
14 of vector-borne pathogens, as suggested recently (Mauck, Chesnais, and Shapiro 2018).
15 In fact, manipulative strategies by vectors have been reported only for a few cases of
16 mixed infections (Rochow 1972; Froissart, Michalakos, and Blanc 2002; Bourdin and
17 Lecoq 1991; Perry and Francki 1992), but the present work, addressing the
18 transmissibility from co-infections of a potyvirus and a crinivirus, suggests that the
19 possible effects of mixed infections on vector transmission might be more general than
20 previously considered.

21 To summarize, we observed that titers of CYSDV remained higher in doubly infected
22 plants, therefore increasing the chances of being transmitted, while the plants were able
23 also to support more whitefly vectors after recovering from the initial stronger symptoms.
24 Indeed, our experiment at 52 dpi showed a clear increase of the transmission rate for
25 CYSDV when acquired from mix-infected plants. Regarding the other virus partner,
26 transmission experiments indicate that WMV could also benefit from its presence in
27 mixed infections, even if the viral titers were lower compared to single infected plants in
28 most time points, because in this case the manipulation of aphids could somehow
29 compensate the lower titers by extending the feeding subphase related to virus
30 acquisition. In other words, during mixed infections CYSDV might enhance its
31 accumulation in plants less severely affected (and thus, prone to survive longer and in
32 better shape) and consequently its chances of dissemination, and at the same time the less
33 abundant WMV might be maintaining the same transmission rates thanks to alterations
34 in the feeding behavior of its vector. Thus, and from an ecological point of view, mix-

1 infected plants would confer adaptive advantages to both viruses for their respective
2 dissemination. The implications of multiple infections for the evolution of pathogens
3 (Tollenaere, Susi, and Laine 2016) are being explored in some virus pathosystems (Ali
4 and Roossinck 2017), and our results suggest that vectors might be key elements in the
5 outcome, along with some effects derived from the frequent presence of more than one
6 pathogen in mixed infections. Although additional work will be needed to elucidate all
7 the mechanisms operating during mixed infections, we believe that our results might be
8 useful to explore how the interactions and their dynamics can conditions the available
9 strategies to control viral diseases in crops.

11 **Experimental Procedures**

12 **Plants, insects and viruses**

13 Seeds of the virus-susceptible melon cultivar "Piel de sapo" PS T111, provided by
14 Semillas Fitó (Spain) were germinated after 5 min treatment with the fungicide Captan in
15 aqueous solution (3g/l). Plants were grown in chambers at 22-25°C and 16/8 h light/dark
16 cycles.

17 A clonal population of the aphid *Myzus persicae*, named MP89 and originally from the
18 ICA-CSIC (Madrid, Spain), was maintained on tobacco plants (*Nicotiana tabacum*, cv
19 Xanthi). A colony of *Bemisia tabaci* biotype MEAM1 (formerly biotype B) was provided
20 by Dr. R. Gabarra (IRTA-Cabrils, Barcelona, Spain), and was raised on tomato plants
21 (*Solanum lycopersicum*, cv Micro-Tom). Plants supporting insects for laboratory
22 experiments were kept in separated growth chambers at 23-25°C and 16/8 h light/dark
23 cycles. For field experiments with whiteflies, individuals from a colony of *B. tabaci*
24 biotype MED (former Q) were used (Rodríguez-López et al. 2012).

25 The Spanish isolate of the crinivirus CYSDV, named CYSDV-AILM, was kindly
26 provided by Dr. M.L. Gómez-Guillamón (IHSM "La Mayora", Málaga, Spain), and was
27 maintained through periodical whitefly transmission on susceptible host plants.

28 The initial infection of WMV was achieved using an infectious full-length clone kindly
29 provided by Dr. C. Desbiez (INRA-PACA, Avignon, France), following described
30 procedures (Desbiez et al. 2011), and the virus was later propagated through aphid
31 transmission.

33 **CYSDV and WMV transmission experiments**

1 Both CYSDV and WMV isolates were transmitted to melon plants using their
2 natural vectors. Experimental conditions for CYSDV transmission were set after testing
3 temperature and the number of viruliferous whiteflies, with acquisition and inoculation
4 access periods of 48h each, choosing a temperature of 25 °C and a ratio of 20 whiteflies
5 per plant. A controlled-vacuum hand-trap and clamp-cages were used for handling insects
6 during transmission to individual plants. In experiments with a large number of test
7 plants, enough viruliferous whiteflies to reach the chosen rate were released inside insect-
8 proof cages containing the plants to be inoculated.

9 The transmission of WMV by aphids was performed essentially as described before for
10 other potyviruses (Ruiz-Ferrer et al. 2005). Briefly, aphids were fasted for 3h hours and
11 then allowed to acquire and inoculate the virus during periods of 10-30 min in each case,
12 transferring them manually from the source to the receptor plant with the help of a
13 paintbrush, until 10 viruliferous aphids were placed on each individual test plant. Vectors
14 were then killed with Confidor insecticide, and the plants were maintained to determine
15 transmission rates by presence of visual symptoms for WMV, or through molecular
16 diagnostic for CYSDV (see below).

17 The CYSDV field transmission experiments were performed inside net-structures (insect
18 proof walk-in 5 × 5 × 2 m) within a tunnel net house and under natural temperature and
19 light conditions (Fig S1) (Rodríguez-López et al. 2012). Tree plants infected with
20 CYSDV or co-infected with CYSDV+WMV were used as inoculum source, surrounded
21 by receptor plants (n=20), either uninfected or previously infected with WMV, distributed
22 in a circle of about 2 m diameter. About 800 non-viruliferous whiteflies were released in
23 the center of the source plants, leading to a vector density of 40 whiteflies/plant. After 4
24 days the receptor plants were treated with insecticide and transferred to a greenhouse for
25 up to 30 days, using tissue print to determine the presence of CYSDV. Field experiments
26 were performed in two consecutive years, using three independent net structures for each
27 combination of source (CYSDV or CYSDV+WMV) and receptor (uninfected or WMV)
28 plants (no choice, total 12 net structures). In free-choice experiments, 10 uninfected plants
29 and 10 WMV-infected plants were distributed in alternated positions (3 net structures for
30 each type of source plants).

31 32 **Sampling, RNA extraction and detection of viruses**

33 Sampling of representative tissues was standardized to account for growth
34 differences between uninfected, single, and double virus-infected plants. Two leaf discs

1 (1 cm diameter) were taken from the second to the latest fully-expanded leaf at each
2 sampling time point, regardless of size and position within the plant. Samples were
3 immediately frozen in liquid nitrogen and ground in Tissue lyser (QIAGEN). Total RNA
4 was extracted using TRIzol reagent (Invitrogen) according to provider instructions,
5 including an additional ethanol precipitation to improve purity. Quality and concentration
6 were estimated in NanoDrop® spectrophotometer (ND-8000).

7 The High-Capacity cDNA Reverse Transcription kit (Applied Biosystems™) was used
8 to generate cDNA from 1µg of total RNA, following the manufacturer protocol. The
9 cDNAs served as templates for amplification of viral sequences by polymerase chain
10 reaction (PCR) using TaKaRa ExTaq® commercial kit. The number of cycles and
11 conditions for amplification (temperature and extension time) were optimized for each
12 combination of primers. Virus specific oligos (Table S1) were used to amplify an 813 nt
13 fragment in the CP region of WMV, and a 563 nt fragment near the 3' end of the genomic
14 RNA2 of CYSDV (Marco et al. 2003). DNA fragments were analyzed by electrophoresis
15 on agarose gels.

17 **Quantification of viral load by RT-qPCR**

18 Virus quantification was performed with cDNAs generated from 1 µg of total
19 RNAs, that were previously treated with DNase (ThermoScientific), by RT-qPCR and
20 comparison to standard curves representing known dilutions of *in vitro* transcripts.

21 To generate virus transcripts, specific oligos were designed to amplify and clone
22 fragments of approximately 1.5 kb located near the 5' ends of viral genomic RNAs in
23 each case (Table S1). The plasmid pGEMT-Easy (Promega) was used to clone fragments
24 deriving from the 5' ends of RNA 1 and RNA 2 of CYSDV, generating constructs
25 RNA1ORF1CYSDV and RNA2Hsp70hCYSDV with the viral sequences under the
26 control of T7 promoter to allow *in vitro* transcription with genome orientation (+ sense).
27 For WMV, the complete P1 coding region was cloned into pENTR/D-TOPO, and
28 transferred by Gateway recombination to the vector pDEST 14 (Invitrogen) under the
29 control of T7 promoter in genome orientation (+ sense). Constructs were verified by
30 restriction analysis and sequencing. Transcripts were obtained with linearized plasmids
31 as templates using the MEGAscript T7 kit (Ambion), quantified and used as standards in
32 RT-qPCR.

33 Two pairs of specific oligos were designed ([https://www.genscript.com/ssl-](https://www.genscript.com/ssl-bin/app/primer)
34 [bin/app/primer](https://www.genscript.com/ssl-bin/app/primer)) to amplify fragments ranging from 80 to 150 base pairs (bp) for each

1 viral RNA (S1 Table), and those leading to better linear regression with the reference
2 were selected in each case. SYBRGreen (Roche) was used to detect PCR products in a
3 Ligth Cycler 480 (Roche) equipment, using 50 ng of cDNAs as template in the PCR
4 reactions

6 **Screening of CYSDV infected plants through tissue print and molecular** 7 **hybridization**

8 The presence of CYSDV before the onset of symptoms was determined by
9 tissue printing and hybridization with a virus-specific probe (Más and Pallás 1995).
10 Briefly, the plasmid pLM15/24 (Marco et al. 2003) kindly provided by Dr. M. Aranda
11 (CEBAS-CSIC, Murcia), containing a fragment of 5437 bp from the virus RNA1 under
12 the control of the SP6 promoter, was linearized with the restriction enzyme XbaI, and *in*
13 *vitro* transcription was carried out using the DIG NORTHERN kit (Roche), following
14 manufacturer's instructions, to obtain negative sense transcripts labeled with digoxigenin.
15 The transcripts were hydrolyzed to shorter size using a carbonate buffer treatment,
16 ethanol precipitated, and resuspended until use (Marco et al. 2003).

17 Cross sections of petioles were printed by pressing on positively charged Nylon
18 membranes (Roche), and the nucleic acids fixed by crosslinking with 120 mJ/cm²
19 (UVC500 crosslinker, Amersham Biosciences). The membrane was incubated with
20 prehybridization solution (20x SSC, 10% N-Lauryl sarcosine, 50% Formamide, 10%
21 SDS and 2% commercial blocking agent from Roche) at 65°C for 3h, before adding the
22 denatured probe (heated for 10 minutes at 65 °C) and incubated for 14-16 h at 65°C.
23 Processing included: two 10 min rinses (2x SSC, 0.1% SDS); two 15 min rinses (0.1x
24 SSC, 0.1 % SDS); 5 min in washing solution (0.1 Maleic acid, 0.15M NaCl pH7.6, 0.3%
25 Tween 20); 1h blocking (0.1 Maleic acid, 0.15 M NaCl pH 7.6, 1% commercial blocking
26 agent of Roche); 1 h incubation with anti-DIG antibody (Roche) at the recommended
27 concentration; two 15 min rinses in washing solution. The signal was finally visualized
28 in a ChemiDoc chemiluminescence equipment (Bio-Rad), using CDP-Star substrate
29 (Roche) in detection buffer (0.1M TrisHCl pH 9.5, 0.1M NaCl).

31 **Monitoring of the feeding behavior of aphids by EPG**

32 Feeding behavior of *M. persicae* on melon plants was analyzed by EPG
33 (Tjallingii 1978). Activities up to 1-hour were recorded from individual insects feeding
34 on plants: i) not inoculated, ii) infected with CYSDV, iii) infected with WMV, or iv)

1 infected with CYSDV+WMV. Adult apterous aphids were connected to a gold wire (3
2 cm long, 20 μm of diameter) with a drop of silver conductive paint (Pelco® Colloidal
3 Silver no. 16034, Ted Pella Inc., Redding, CA, USA) on the dorsum. The wire was glued
4 to a copper nail and connected to an EPG probe following standard procedures (Carmo-
5 Sousa et al. 2014). Each wired insect was placed on the leaf of a plant, with the circuit
6 closed through the plant substrate. Four plants were tested in parallel using a 4-channel
7 device (Giga-4; EPG Systems, Wageningen, The Netherlands), and 20 to 30 repetitions
8 were recorded. EPG data acquisition was carried out using the Stylet+ computer program
9 (EPG-Systems, Wageningen, The Netherlands), to identify EPG waveforms described for
10 aphids (Tjallingii 1988) corresponding to non-probing (np), intercellular apoplastic stylet
11 pathway (C), and intracellular stylet puncture (Pd). No other EPG waveforms were
12 observed during the 1h recording time. Behavioural variables were processed using a
13 previously described MS Excel Workbook (Sarria et al. 2009).
14 EPG variables (mean \pm SE) were calculated (Backus et al. 2007) to determine: proportion
15 of individuals that produced a specific waveform type (PPW); number of wave-form
16 events per insect (NWEI); total waveform duration (min) per insect (WDI); and total
17 waveform duration (min) per event (WDE). Data were analyzed after transformation
18 using $\ln(x+1)$. Because of the non-Gaussian distribution of the data, a Mann-Whitney U
19 test was conducted in GraphPad prism 6.0.

20

21

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33
34

1 **Table Caption**

2

3 **Table 1.** Selected EPG variables describing the feeding behavior of *M. persicae* on single
4 and mixed virus infected melon plant

5

6

7 **Figure Legends**

8

9 **Figure 1. Kinetics of viral loads in single and mix-infected plants.** Quantification of
10 CYSDV (left) and WMV (right), showing viral loads in single (blue) and mixed (red)
11 infected plants. Each point represents 2 independent biological replicates of individually
12 infected plants (dataset n=15), collected at 12, 24, 36, 48 and 60 d.p.i. The mean values
13 are indicated by the horizontal lines, connected through an adjusted curve, and surrounded
14 by gray areas representing 95% confidence intervals.

15

16 **Figure 2. Symptomatology of infected plants.** A) Details of leaves with symptoms
17 caused by single infection with WMV (upper panel) or CYSDV (lower panel). B)
18 Representative single and mix-infected plants photographed at the indicated time-points.
19 C) Side-by-side representative plants of the four categories (from left to right: non-
20 infected, CYSDV, WMV, CYSDV+WMV) photographed at 24 and 48 dpi.
21 corresponding with the development used as virus sources during vector-transmission
22 experiments

23

24 **Figure 3. CYSDV transmission in controlled conditions at 24 dpi.** Pooled values of
25 virus accumulation are plotted on the upper panel (compared by U Mann-Whitney test)
26 above the tabulated results of the three repetitions of transmission, with number of
27 infected plants/number of plants tested for each independent experiment, followed by the
28 values of two-tailed Chi-square tests (df=1). Transmission rates are showed graphically
29 using pink for CYSDV infected vs. green for uninfected test plants.

30

31 **Figure 4. Transmission of CYSDV under field conditions.** Average of transmission
32 rates in three individual net-structures, with 20 test plants each, either uninfected or
33 previously infected by WMV, and surrounding single infected CYSDV plants (left panel),

1 or mixed infected CYSDV+WMV plants (right panel) as sources of inoculum. No
2 significant differences were found (t-test).

3
4 **Figure 5. Free-choice transmission of CYSDV under field conditions.** A) Graphical
5 representation of comparative transmission of CYSDV to alternated plants, 10 uninfected
6 and 10 previously infected with WMV, using single infected CYSDV plants as source of
7 inoculum; results for each individual net-structure are represented individually. B)
8 Similar experiment as in a) with mixed infected CYSDV+WMV plants as inoculum
9 source. Analysis of data was performed with an association test.

10
11 **Figure 6. WMV transmission in controlled conditions at 24 dpi.** Pooled values of virus
12 accumulation are plotted on the upper panel (compared using t-test analysis) above the
13 tabulated results for the three repetitions of transmission, with number of infected
14 plants/number of plants tested for each independent experiment, followed by the values
15 of two-tailed Chi-square tests (df=1). Transmission rates are showed graphically using
16 blue for WMV infected vs. green for uninfected test plants.

17
18 **Figure 7. EPG monitoring of aphid feeding behavior.** A) Example of a representative
19 EPG record with a potential drop (Pd) in which the different sub-phases (II-1, II-2 and II-
20 3) can be identified. B) Comparison of the duration of the sub-phases corresponding to
21 Pds between aphids fed on single or mix-infected plants c) Comparison of number of
22 pulses of subphase II-3 between aphids fed on single or mix-infected plants. The error
23 bars represent ± 1 SEM. ** P<0.005.

24
25 **Figure 8. CYSDV and WMV transmission in controlled conditions at 52 dpi.** The
26 transmission rate of CYSDV by whiteflies (upper panel) and WMV by aphids (lower
27 panel) is shown for five independent experiments using 10 plants each, with the number
28 of infected plants/number of plants tested for each independent experiment, and the
29 pooled total followed by the value of a two-tailed Chi-square test (df=1). Statistical
30 significant differences are indicated with P-value in bold. Transmission rates are depicted
31 graphically using the same color code as in previous figures: pink for CYSDV-infected,
32 blue for WMV-infected, and green for uninfected test plants.

1 **Supporting Information Legends**

2

3 **Table S1.** List of primer sequences and their uses

4

5 **Table S2.** CYSDV transmission in field conditions from single and mixed-infected plants
6 to uninfected or WMV-infected in different net-houses.

7

8 **Table S3.** CYSDV transmission in field conditions from single and mixed-infected plants
9 to uninfected or WMV-infected plants in the same net-structures, in alternate positions.

10

11 **Table S4.** Comparative analysis of EPG parameters in aphids feeding on uninfected,
12 CYSDV, WMV, and WMV+CYSDV infected plants

13

14 **Table S5.** Monitoring of intracellular subphases during potential drops (Pd) detected in
15 aphids feeding behavior by EPG

16

17 **Figure S1.** Secondary CYSDV transmission in field conditions.

18

19

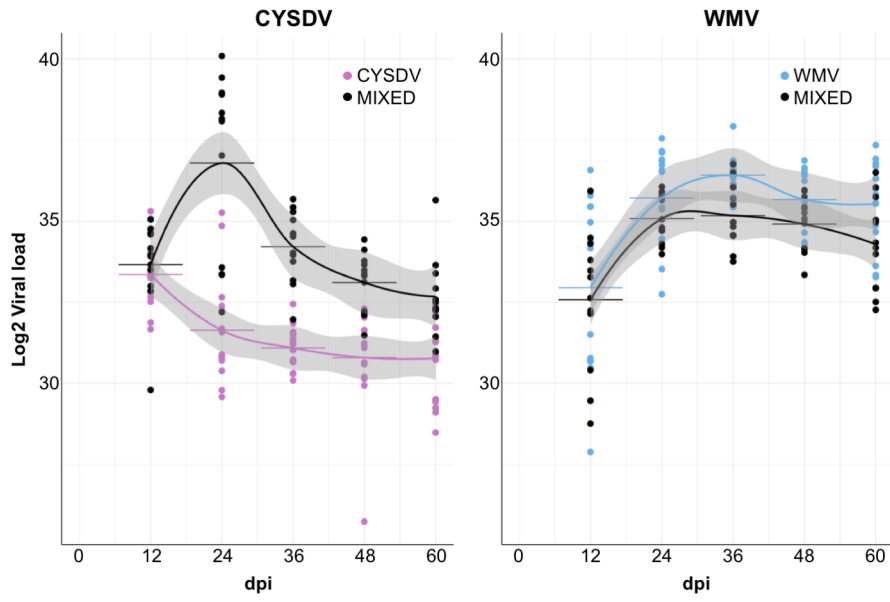
Table 1. Selected EPG variables describing the feeding behavior of *M. persicae* on single and mixed virus infected melon plant

Variables ¹	Treatment	PPW	NWEI	P ²	WDI	P ²	WDE	P ²
Np	WMV	30/30	0.80±0.17 (0.48-1.04)	0.73	3.07±0.27 (2.28-3.32)	0.04	2.10±0.53 (0.65-3.18)	0.03
	CYSDV + WMV	28/28	0.82±0.23 (0.30-1.18)		2.96±0.28 (2.19-3.39)		2.00±0.46 (0.84-2.98)	
C	WMV	30/30	0.77±0.18 (0.30-1.00)	0.48	2.75±0.31 (2.08-3.25)	0.06	1.69±0.55 (0.94-3.05)	0.74
	CYSDV + WMV	28/28	0.80±0.22 (0.30-1.15)		2.91±0.27 (2.36-3.26)		1.68±0.59 (0.92-3.26)	
Pd	WMV	30/30	1.14±0.26 (0.60-1.59)	0.12	1.75±0.25 (1.11-2.13)	0.02	0.70±0.08 (0.52-1.12)	<0.0001
	CYSDV + WMV	28/28	1.24±0.23 (0.70-1.62)		1.90±0.22 (1.29-2.20)		0.74±0.11 (0.54-1.18)	

¹ For each treatment, the following parameters were measured: Np, non-probe activity; C, intercellular stylet pathway; Pd, short intracellular punctures. PPW means proportion of individuals that produced the waveform type, with NWEI being the number of waveform events per insect, WDI the waveform duration per insect, and WDE the waveform duration per event. Durations are expressed in minutes for Np and C, and in seconds for Pd. Values are indicated as mean ± 1 SEM (range given in parentheses).

² Statistical comparisons between the two treatments were performed for each parameter with non-parametric U-Mann Whitney test, considering non Gaussian distribution variables. Bold-type indicates significant differences (p<0.05)

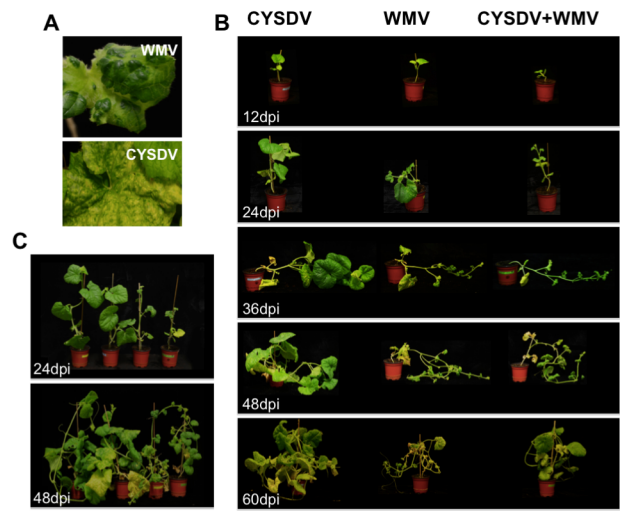
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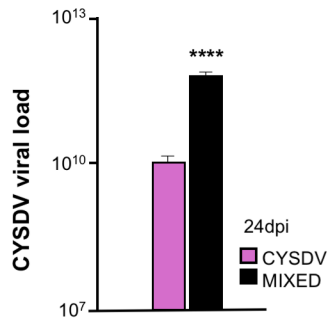
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3 Fig.1

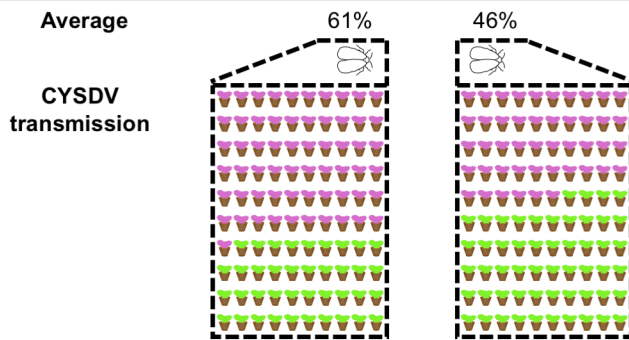
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1
2 Fig.2
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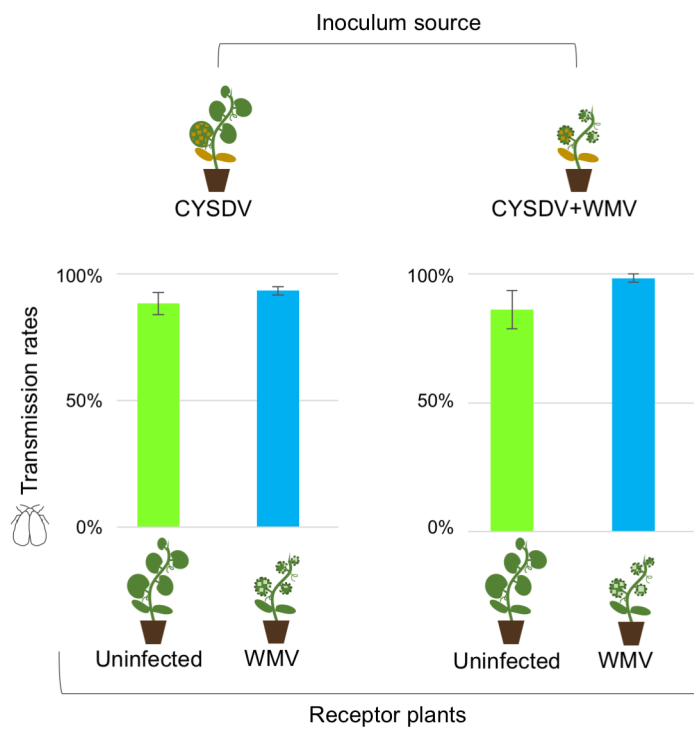
Inoculum source		CYSDV	CYSDV+WMV	Chi-square , df	P value
Transmission rates	Exp.1	27/40	19/40	3.27 , 1	0.07
	Exp. 2	15/39	14/40	0.10 , 1	0.75
	Exp. 3	30/40	22/40	3.52 , 1	0.06
Total		72/119	55/120	5.16 , 1	0.02



1

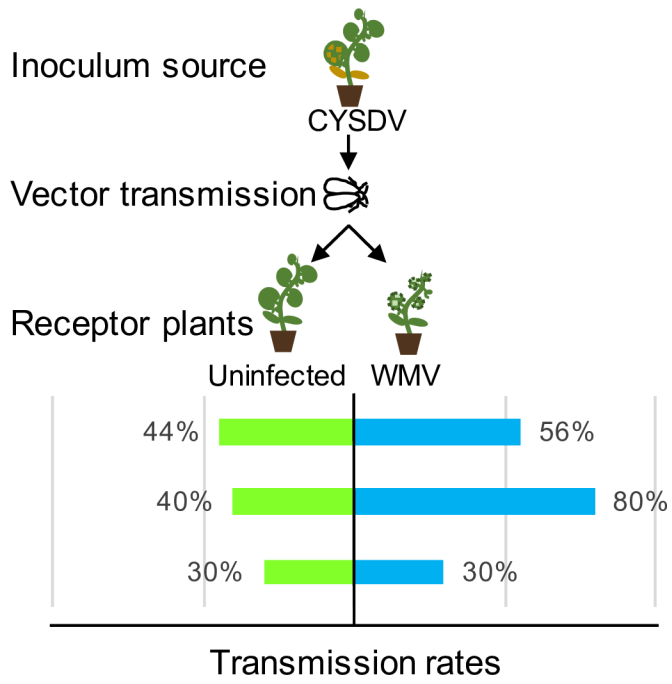
2 Fig.3

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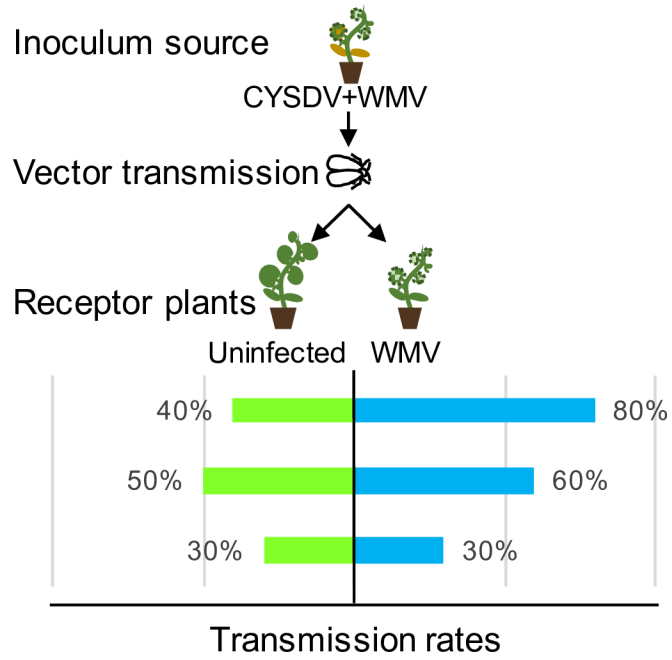


1
2 Fig.4
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A



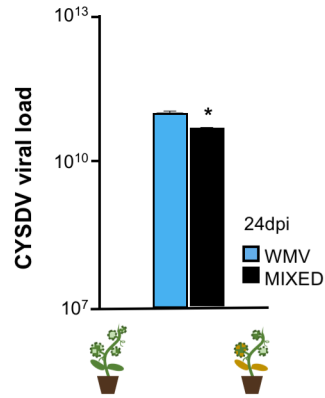
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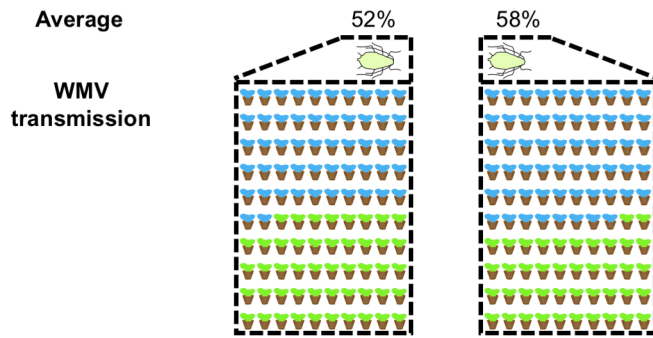
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2 Fig. 5

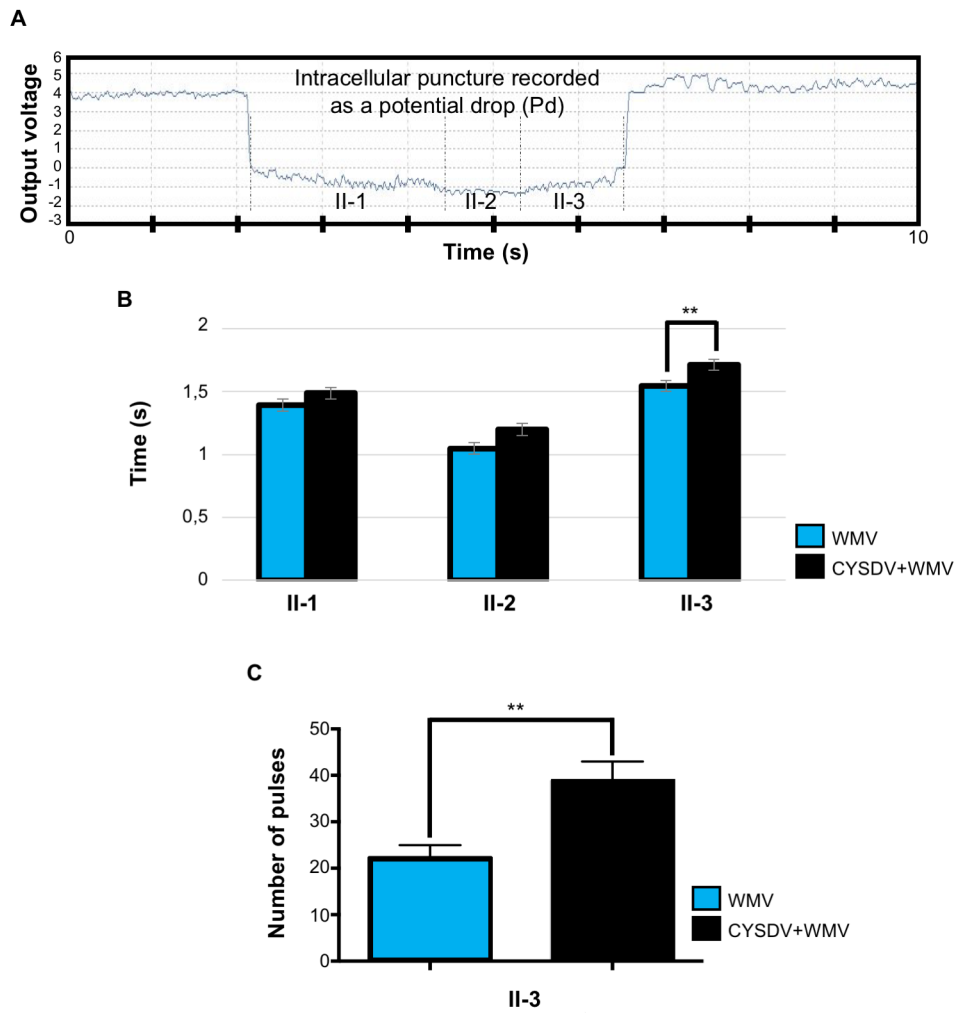
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Inoculum source		WMV	CYSDV+WMV	Chi-square , df	P value
Transmission rates	Exp.1	23/40	28/40	1.35 , 1	0.25
	Exp. 2	15/40	23/38	4.14 , 1	0.04
	Exp. 3	21/34	13/32	2.95 , 1	0.09
Total		59/114	64/110	0.93 , 1	0.33





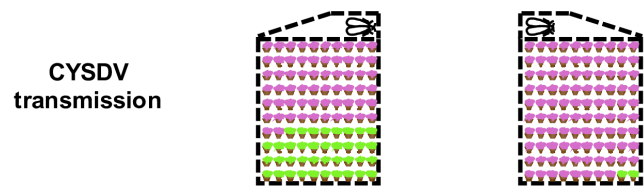
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2 Fig. 6
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

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2 Fig.7
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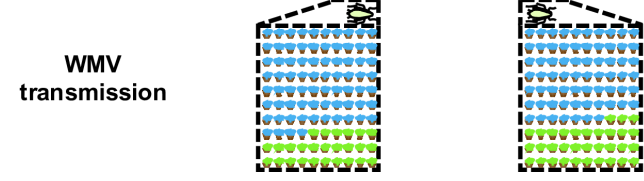
A

Inoculum source	 CYS DV	 CYS DV+WMV		
	5/10	10/10		
	8/10	9/10		
Transmission rates	8/10	10/10		
	4/10	10/10		
	6/10	10/10		
Total	31/50	49/50	Chi-square, df	P value
			20.25, 1	< 0.0001
Average %	62%	98%		



B

Inoculum source	 WMV	 CYS DV+WMV		
	8/10	5/9		
	7/8	8/10		
Transmission rates	6/9	7/10		
	5/10	7/10		
	9/10	6/10		
Total	35/47	33/49	Chi-square, df	P value
			0.59, 1	0.44
Average %	74%	67%		



1

2 Fig. 7

3

1 **Supplemental Table 1.** List of primer sequences and their uses

Name	Sequence	Use
RNA1 CYSDV Fw	5'ATGTCGTCGTCAGTTGC 3'	Template for <i>in vitro</i> transcription ¹
RNA1 CYSDV Rv	5'GTCGAGACCACTCCAGTACA 3'	
RNA2 CYSDV Fw	5'AGGTGGGTAGGTGTTGACAG 3'	Template for <i>in vitro</i> transcription ¹
RNA2 CYSDV Rv	5'TGGGAAGTACCGTCTCCTTACC 3'	
P1 WMV Fw	5' CACCATGGCAACAATCATGTTTGGAG 3'	Template for <i>in vitro</i> transcription ¹
P1 WMV Rv	5'TCAATAATGTTGAATATCTTCTATCTCC 3'	
qPCR RNA1 CYSDV Fw	5'GACTCAAACAAGCGCGATGA 3'	Analysis of viral load (CYSDV RNA1)
qPCR RNA1 CYSDV Rv	5'GCGTTGCCGTTACTGAGGAT 3'	
qPCR RNA2 CYSDV Fw	5'TTTGTCCGCTGACGACTGG 3'	Analysis of viral load (CYSDV RNA2)
qPCR RNA2 CYSDV Rv	5'GGTTCAATGCAACGACTCAGA 3'	
qPCR P1 WMV Fw	5'TGCAACTATTGTGGCCTTCG 3'	Analysis of viral load (WMV)
qPCR P1 WMV Rv	5'CATGGCACCTCCACCTCATA 3'	
WMV Fw	5'CTTATGGTTGTCATTGCTATG 3'	Detection of WMV
WMV Rv	5' CCCACCAACTGTTGGAAG 3'	
MA156 Fw	5' GAAGAATTCCAGGCAAGG 3'	Detection of CYSDV ²
MA129 Rv	5' TCACATCATCAATCCAAAAG 3'	

2 ¹ Primers were used for cloning partial virus sequences to generate *in vitro* transcripts as templates
3 for quantitative comparisons in RT-qPCR. Non-viral sequences used for cloning purposes are
4 highlighted in bold.

5 ² Primers described by Marco et al. 2003

6

7

1 **Supplemental Table 2.** CYSDV transmission in field conditions from single and mixed-
 2 infected plants to uninfected or WMV-infected in different net-houses.

Source	Receptors	Repetition¹	Infected plants²	Total	Average³
CYSDV	Uninfected	1	19/20	53/60	88.33%
		2	18/20		
		3	16/20		
	WMV	1	19/20	56/60	93.33%
		2	19/20		
		3	18/20		
CYSDV+WMV	Uninfected	1	15/20	50/58	86.2%
		2	20/20		
		3	15/18		
	WMV	1	20/20	59/60	98.33%
		2	19/20		
		3	20/20		

3 ¹ Three independent net-structures, each with 20 receptor plants, all of the same type, are shown
 4 ² Results obtained in each independent net-house are indicated for each treatment, and pooled
 5 values and percentages are shown in the last columns.
 6 ³ Statistical analysis were performed by two-tail t-Test comparisons of every pair of treatments
 7 (after verifying inequality of variances by F-Test). None of the paired comparisons indicated
 8 significant differences between the samples means.
 9
 10

1 **Supplemental Table 3.** CYSDV transmission in field conditions from single and mixed-
 2 infected plants to uninfected or WMV-infected plants in the same net-structures, in
 3 alternate positions.

Source	Repetition¹	Receptors	Infected plants	Average
CYSDV	1	Uninfected	3/10	30%
		WMV	3/10	30%
	2	Uninfected	4/10	40%
		WMV	8/10	80%
	3	Uninfected	4/9	44,44%
		WMV	5/9	55,55%
Total	Uninfected	11/29	37,93%	
	WMV	16/29	55,17%	
CYSDV+WMV	1	Uninfected	3/10	30%
		WMV	3/10	30%
	2	Uninfected	5/10	50%
		WMV	6/10	60%
	3	Uninfected	4/10	40%
		WMV	8/10	80%
Total	Uninfected	12/30	40%	
	WMV	17/30	56,66%	

4 ¹ The results of three independent net-structures (with 10 uninfected and 10 previously
 5 WMV-infected receptor plants in alternated positions) are shown for each treatment, with
 6 the pooled values for each category in the "Total" rows.

7 ² Statistical analysis were performed by two-tail t-Test comparisons of every pair of
 8 treatments (after verifying inequality of variances by F-Test). None of the paired
 9 comparisons indicated significant differences between the samples means.

10
 11

1 **Supplemental Table 4.** Comparative analysis of EPG parameters in aphids feeding on
 2 uninfected, CYSDV, WMV, and WMV+CYSDV infected plants

Parameter¹	Treatment	PPW	NWEI	WDI	WDE
NP	Uninfected	20/20	0.73±0.05 (0.30-1.04) a	2.88±0.06 (2.24-3.16) a	2.04±0.04 (0.84-2.87) a
	CYSDV	23/23	0.88±0.03 (0.60-1.20) a	2.91±0.06 (2.34-3.33) a	1.90±0.04 (0.86-2.95) b
	WMV	30/30	0.80±0.03 (0.48-1.04) a	3.07±0.05 (2.28-3.32) b	2.10±0.04 (0.65-3.18) ac
	CYSDV +WMV	28/28	0.82±0.04 (0.30-1.18) a	2.96±0.05 (2.19-3.39) a	2.00±0.03 (0.84-2.98) abd
	C	Uninfected	20/20	0.71±0.05 (0.30-1.04) a	2.96±0.04 (2.49-3.26) a
	CYSDV	23/23	0.87±0.03 (0.60-1.18) b	2.90±0.07 (1.96-3.21) ab	1.75±0.05 (0.84-3.14) b
	WMV	30/30	0.77±0.03 (0.30-1.00) ab	2.75±0.06 (2.08-3.25) b	1.69±0.04 (0.94-3.05) b
	CYSDV +WMV	28/28	0.80±0.04 (0.30-1.15) ab	2.91±0.05 (2.36-3.26) ab	1.68±0.05 (0.92-3.26) b
Pd	Uninfected	20/20	1.28±0.04 (0.85-1.53) a	1.86±0.04 (1.38-2.09) ab	0.69±0.004 (0.51-1.14) ab
	CYSDV	23/23	1.26±0.04 (0.85-1.59) a	1.85±0.04 (1.45-2.21) ab	0.69±0.004 (0.50-1.04) a
	WMV	30/30	1.14±0.05 (0.60-1.59) a	1.75±0.05 (1.11-2.13) a	0.70±0.005 (0.52-1.12) b
	CYSDV +WMV	28/28	1.24±0.04 (0.70-1.62) a	1.90±0.04 (1.29-2.20) b	0.74±0.005 (0.54-1.18) c

3 ¹ Parameters measured include: NP, non-probe activity; C, intercellular stylet pathway; Pd, short
 4 intracellular punctures. PPW means proportion of individuals that produced the waveform type,
 5 with NWEI being the number of waveform events per insect, WDI the waveform duration per
 6 insect, and WDE the waveform duration per event. Durations are expressed in minutes for NP
 7 and C, and in seconds for Pd. Values are indicated as mean ± SE with extremes given in
 8 parentheses. The statistical comparisons were performed in pairs for each parameter with non-
 9 parametric U-Mann Whitney tests, considering non Gaussian distribution variables. Only values
 10 followed by different letters were found significantly different (p<0.05).
 11

1 **Supplemental Table 5.** Monitoring of intracellular subphases during potential drops
 2 (Pd) detected in aphids feeding behavior by EPG

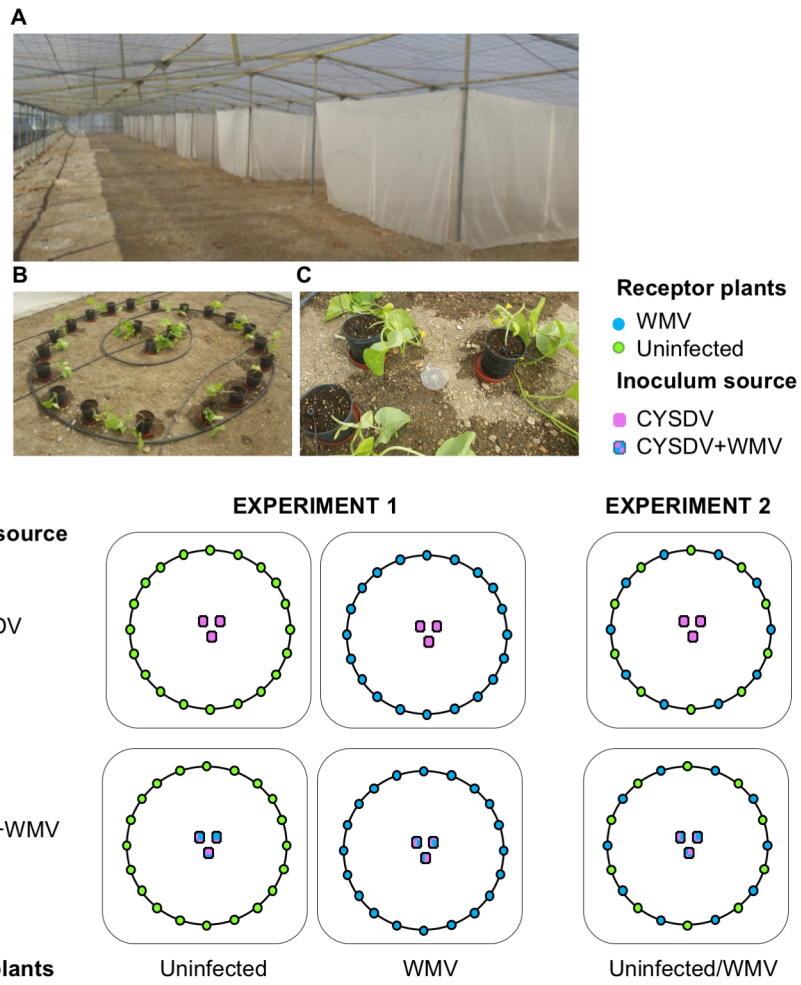
Sub-phase¹	Treatment	PPW	WDI	P²
II-1	WMV	30/30	1,36±0,05(0,75-1,82)	0,09
	CYSDV+WMV	28/28	1,49±0,05(0,81-1,90)	
II-2	WMV	30/30	1,03±0,05(0,52-1,46)	0,08
	CYSDV+WMV	28/28	1,15±0,05(0,53-1,53)	
II-3	WMV	30/30	1,34±0,04(0,77-1,69)	0,002
	CYSDV+WMV	28/28	1,53±0,04(0,91-1,84)	

3 ¹ The sub-phases of pd, associated with uptake (acquisition, II-3) and release (inoculation, II-1)
 4 of non-persistently transmitted viruses are identified following Martin et al., 1997. PPW means
 5 proportion of individuals that produced the waveform type, and WDI is the waveform duration
 6 per insect in seconds. Values are indicated as mean ± SE with extremes given in parentheses.

7 ² Statistical comparisons between the two treatments was performed for each parameter with non-
 8 parametric U-Mann Whitney test, considering non Gaussian distribution variables. Bold-type
 9 indicates significant differences (p<0.05).

10
 11

1



2

3

S1 Figure