

Resistance against melon chlorotic mosaic virus and tomato leaf curl New Delhi virus in melon

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► **To cite this version:**

Gustavo Romay, Michel Pitrat, Hervé Lecoq, Catherine Wipf-Scheibel, Pauline Millot, et al.. Resistance against melon chlorotic mosaic virus and tomato leaf curl New Delhi virus in melon. Plant Disease, American Phytopathological Society, 2019, 103 (11), pp.2913-2919. 10.1094/PDIS-02-19-0298-RE . hal-02619468

HAL Id: hal-02619468

<https://hal.inrae.fr/hal-02619468>

Submitted on 25 May 2020

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1 **Resistance against melon chlorotic mosaic virus and tomato leaf curl New Delhi virus in**
2 **melon**

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4 G. Romay^{1,3}, M. Pitrat², H. Lecoq¹, C. Wipf-Scheibel¹, P. Millot¹, G. Girardot¹ and C. Desbiez^{1*}

5
6 ¹ INRA, Unité de Pathologie Végétale, Domaine St Maurice, Allée des Chênes, CS60094,
7 84143 Montfavet cedex, France

8 ² INRA, Unité de Génétique et Amélioration des Fruits et Légumes, Domaine St Maurice,
9 Allée des Chênes, CS60094, 84143 Montfavet cedex, France

10 ³ Université catholique de Louvain, Earth and Life Institute, Phytopathology, Croix du Sud 2
11 Bte L07.05.03, 1348 Louvain-la-Neuve Belgium.

12

13 * Corresponding author: cecile.desbiez@inra.fr

14

15 **Key words:** *Cucumis melo*, begomovirus, MeCMV, ToLCNDV, genetic control, inheritance

16

17 **Abstract**

18 Thirty-one melon accessions were screened for resistance to the begomoviruses *Melon*
19 *chlorotic mosaic virus* (MeCMV) and *Tomato leaf curl New Delhi virus* (ToLCNDV). Five
20 accessions presented nearly complete resistance to both viruses. Accession IC-274014,
21 showing the highest level of resistance to both viruses, was crossed with the susceptible
22 cultivar Védrantais. The F₁, F₂, F₃/F₄ and both back-cross progenies were mechanically
23 inoculated with MeCMV. Plants without symptoms nor virus detection by ELISA and/or PCR
24 were considered as resistant. The segregations were compatible with two recessive and one

25 dominant independent genes simultaneously required for resistance. Inheritance of
 26 resistance to ToLCNDV in the F₂ was best explained by one recessive gene and two
 27 independent dominant genes simultaneously required. Some F₃ and F₄ families selected for
 28 resistance to MeCMV were also resistant to ToLCNDV, suggesting that common or tightly
 29 linked genes were involved in resistance to both viruses. We propose the names
 30 *begomovirus resistance-1* and *Begomovirus resistance-2* for these genes (symbols *bgm-1* and
 31 *Bgm-2*). Resistance to MeCMV in IC-274014 was controlled by *bgm-1*, *Bgm-2* and the
 32 recessive gene *melon chlorotic mosaic virus resistance (mecmv)*; resistance to ToLCNDV was
 33 controlled by *bgm-1*, *Bgm-2* and the dominant gene *Tomato leaf curl New Delhi virus*
 34 *resistance (Tolcndv)*.

36 Introduction

37 Melon (*Cucumis melo* L.) is one of the major crop species in the family *Cucurbitaceae*.
 38 Global production of melon is circa 30,000,000 tons per year (www.fao.org/faostat).
 39 Cucurbit diseases caused by begomoviruses (genus *Begomovirus*, family *Geminiviridae*) are a
 40 new threat to cucurbit production worldwide. At least six and eight begomovirus species
 41 from the New World (NW) and Old World (OW), respectively, have been described infecting
 42 cucurbits in the past 30 years (Brown *et al.* 2015). Some of them have a narrow geographic
 43 distribution despite a high local prevalence and agronomic impact, like the NW begomovirus
 44 *Melon chlorotic mosaic virus* (MeCMV) observed only in Venezuela so far (Romay *et al.*
 45 2010). Others have a larger geographic range: the NW begomovirus *Squash leaf curl virus*
 46 (SLCV) and the OW begomovirus *Watermelon chlorotic stunt virus* (WmCSV) were introduced
 47 into the Eastern Mediterranean Basin in the early 2000s (Lapidot *et al.* 2014). The OW
 48 *Tomato leaf curl New Delhi virus* (ToLCNDV), first described in India, was more recently

49 reported in Europe and in the north of Africa (Fortes et al. 2016) where it has rapidly
50 become a major agronomic problem (Moriones et al. 2017). Control of begomoviruses relies
51 on the limitation of their vector *Bemisia tabaci* –mostly by intensive insecticide treatments-,
52 and on genetic resistance when available. Considering the frequent emergence of highly
53 damaging begomoviruses, breeding for resistance is a key factor for a durable control,
54 particularly if broad-spectrum resistances effective against several viruses at once can be
55 found.

56 In melon, resistances to different viruses and pathogens are often found in the same
57 genotype even when the resistance factors are distinct (Lecoq et al. 1998, Table 1).
58 Resistances to WmCSV (Yousif et al. 2007), cucurbit leaf crumple virus (CuLCrV) (Hagen et al.
59 2008b; McCreight et al. 2008), and more recently ToLCNDV (Lopez et al. 2015), have been
60 described. However there is no evidence for general resistance factors against several of
61 these viruses

62 In this work, we looked for resistance against the genetically distant MeCMV (NW) and
63 ToLCNDV (OW) and checked whether common genetic factors could target these two
64 viruses. In a first step we screened 31 melon accessions, previously described as resistant to
65 at least one virus or representative of the phenotypic melon diversity. We also tested the
66 inheritance of resistance to MeCMV in the most resistant accession IC-274014 and its
67 relationship with resistance to ToLCNDV, in order to find efficient and generic resistance
68 factors that could be used in breeding programs.

69

70 **Material and methods**

71 Virus strains and infectious clones

72 MeCMV was collected from watermelon (*Citrullus lanatus* (Thumb.) Matsum. and Nakai)
73 in Zulia, Venezuela. A multimeric infectious clone of this virus was obtained in a previous
74 work (Romay et al. 2015). A ToLCNDV (ES13-35) isolate was obtained from a leaf sample of
75 zucchini squash (*Cucurbita pepo* L) from Murcia, Spain. Total DNA was extracted from the
76 infected plant as described by Gilbertson et al. 1991, and ToLCNDV detection was confirmed
77 by sequencing of a PCR fragment amplified using universal primers for begomoviruses
78 (Wyatt and Brown, 1996). The full-length genome sequence of this isolate was obtained for
79 further constructions of a multimeric infectious clone.

80 The complete genome of ES13-35 isolate was amplified by rolling circle amplification
81 (RCA) according to the manufacturer's recommendations (Romay et al. 2015). The DNA-A
82 fragment generated by RCA was digested with *Xba*I and *Xho*I (~1 kb) and inserted into
83 pBluescript II SK (+) (Stratagene). Then, a complete DNA-A was linearized with *Bam*HI and
84 inserted into the plasmid containing the *Xba*I-*Xho*I fragment. To generate a multimeric clone
85 of DNA-B, a 1.6kb fragment was obtained by digestion of RCA product with *Pst*I and insertion
86 into pBlueScript. Then, a complete DNA-B fragment after *Mlu*I digestion of the RCA product
87 was ligated to the previously clone containing the 1.6kb fragment and linearized with *Mlu*I.
88 For cloning of both DNA components, *Escherichia coli* strain DH5 alpha was used as host of
89 plasmid constructs.

90 The infectivity of the clones was confirmed by bombardment of melon seedlings
91 according to Romay et al. 2015.

92

93 Viral inoculation assays

94 Melon seedlings of the susceptible cultivar Védrantais, inoculated with infectious clones
95 of ToLCNDV and MeCMV, were used as virus sources for mechanical inoculations. One g of

96 young infected leaf tissue was ground in 5 ml of an ice-cold solution of 0.03M Na₂HPO₄ +
97 0.2% DIECA (diethyldithiocarbamate). Activated charcoal and carborundum were added to
98 the sap extract (Romay et al. 2015). Two successive inoculations were performed for each
99 melon seedling: a first one on the cotyledons at 6 days post sowing, followed by a second
100 inoculation onto the first true leaf six days later, as described by Romay et al. 2015.
101 Symptoms were checked visually at weekly intervals. At 30 days post-inoculation (dpi),
102 young leaves were collected from all plants for accessions that did not show any symptoms,
103 and one plant per accession in the accessions that displayed symptoms, for serological or
104 molecular virus detection. All inoculated plants were kept in biosafety level 3 (S3)
105 greenhouse.

106

107 Virus detection (ELISA, PCR)

108 Begomovirus detection was performed by PCR as described in (Romay et al. 2015), with
109 primers 64ACR5 and 64debAC3 (Romay et al. 2014) for MeCMV and primers TLCNDV-CP-5'
110 (5'-ATGKYGAAGCGACCAGCAGA-3') and TLCNDV-CP-3' (5'-CCGAATCATARAARTAGATCCG-3')
111 for ToLCNDV.

112 For the studies on F₂ and backcrosses (BC) with MeCMV, all plants were tested in DAS-ELISA
113 with an antiserum against SLCV (Cohen et al. 1983; courtesy of J.E. Duffus) that presents a
114 strong cross-reactivity with MeCMV.

115 For the studies on F₃ and F₄ progenies, asymptomatic plants were tested by DAS-ELISA
116 with a ToLCNDV antiserum (Agdia EMEA, Grigny, France) also presenting a strong cross-
117 reactivity with MeCMV (data not shown).

118

119 Plant material

120 *Screening for resistance*

121 Thirty one melon accessions were selected from a germplasm collection at Institut
122 National de la Recherche Agronomique (INRA, Avignon, France) for the resistance screening
123 experiments. Most of the selected accessions had been previously reported to be resistant
124 or tolerant to at least one plant pathogen; others were representative of the phenotypic
125 melon diversity (Pitrat, 2016a,b, Table 1). Ten plants of each melon accession were used for
126 each virus in two independent tests.

127

128 *Inheritance studies*

129 The melon accession IC-274014 belongs to the group momordica (Pitrat, 2016b) and was
130 supplied by N.P.S. Dhillon (Dhillon et al. 2007). It was crossed with the cultivar Védraçais
131 belonging to the group cantalupensis sub-group charentais (obt. Vilmorin). The F₁ hybrid was
132 selfed to obtain the F₂ progeny and backcrossed to IC-274014 or Védraçais to obtain
133 respectively the BC_R and BC_S progenies. The parents, F₁, F₂ and both BC were mechanically
134 inoculated with MeCMV in two independent tests. A third test was conducted for MeCMV
135 with only the susceptible parent Védraçais and the F₂ progeny. Ten resistant F₂ plants (no
136 symptom, no virus detected by ELISA and PCR after inoculation with MeCMV) were selfed to
137 produce ten F₃ progenies. As only few seeds with low germination ability have been
138 obtained in the F₃ progeny, two to four plants, without selection for virus resistance or
139 susceptibility, from each F₃ progeny were selfed to obtain F₄ families.

140 The F₁ and F₂ progenies (IC-274014 × Védraçais) were mechanically inoculated with
141 ToLCNDV. Besides, plantlets of the F₃ progenies deriving from the F₂ selected for resistance
142 to MeCMV, and of the corresponding F₄, were mechanically inoculated with ToLCNDV to
143 determine if the same genes were involved in resistance to both viruses.

144

145 Correlation with resistance to other cucurbit-infecting viruses

146 *Resistance of the F₄ (IC-274014 × Védraçais) progenies to zucchini yellow mosaic virus*
147 *(ZYMV)*

148 Ten plantlets of Védraçais, IC 274014 and of the F₄ progenies at the cotyledonary stage
149 were mechanically inoculated with strain E15 of ZYMV (Lecoq and Pitrat, 1984). Symptoms
150 were recorded weekly and an ELISA test was performed 4 weeks after inoculation on the
151 asymptomatic plants.

152

153 *Behaviour of melon accession and of 6 of the F₄ (IC-274014 × Védraçais) progenies towards*
154 *cucurbit aphid-borne yellows virus (CABYV)*

155 A field assay was performed to test for the resistance to CABYV. An experimental plot was
156 planted in July 2018 with 20 plants of Védraçais and 10 plants each of 11 accessions (HSD
157 2445-005, IC-274014, PI 164323, PI 164723, PI 179901, PI 234607, PI 236355, PI 414723, PI
158 482420, WM7 and WM9), the F₁ hybrid (IC-274014 × Védraçais) and 6 of the F₄ (IC-
159 274014 × Védraçais) progenies. The plantlets were planted in 4 rows of 50 plants each,
160 with 1.5 m between ranks and 0.8 m between plants in a rank. Two blocks of 5 plants each
161 were planted for each accession including the resistant control PI 414723, and 4 blocks for
162 the susceptible control Védraçais. Two months after planting, an ELISA test was performed
163 with antisera against CABYV, ZYMV, watermelon mosaic virus (WMV) and cucumber mosaic
164 virus (CMV).

165

166

167 **Results**

168 Infectivity of MeCMV and ToLCNDV clones

169 The full-length genome of the ToLCNDV isolate ES13-35 (GenBank accession numbers
 170 MK279352 and MK279353 for DNA-A and DNA-B, respectively) showed at least 99%
 171 nucleotide identity with isolates previously reported in southern Spain (Juarez et al. 2014;
 172 Moriones et al. 2017; Ruiz et al. 2015). After biolistic inoculation of seedlings of the
 173 susceptible cultivar Védraçais with multimeric clones of ToLCNDV and MeCMV, viral
 174 symptoms could be observed between the first and second week post-inoculation: severe
 175 mosaics, leaf crumpling and stunting (more severe for MeCMV than for ToLCNDV) with
 176 occasional enations on the lower leaf surface for MeCMV only. The presence of each virus
 177 was confirmed by PCR at 30 dpi. Mechanical inoculation of both begomoviruses from these
 178 sources yielded 100% infection in Védraçais in all experiments.

179

180 Resistance to MeCMV and ToLCNDV in melon accessions

181 Among the 31 melon lines tested, 17 were susceptible to both MeCMV and ToLCNDV
 182 (Table 1). In the susceptible accessions as well as in Védraçais, symptoms induced by
 183 MeCMV were usually more severe than those of ToLCNDV with severe stunting, leaf
 184 crispation and occasionally leaf enations (data not shown).

185 Among the 14 remaining accessions, heterogeneous response
 186 (susceptibility/intermediate resistance/resistance) or intermediate resistance was observed
 187 in 8 melon lines for MeCMV and in 7 melon lines for ToLCNDV. The symptoms that were
 188 predominantly observed in intermediate resistant melon were systemic chlorotic spots.
 189 Interestingly, several accessions exhibited a “recovery” phenotype, with more symptoms in
 190 the first leaves at 7 dpi than in young leaves at 30 dpi. This was the case for the melon lines
 191 PI 164723, PI 414723 and 90625 for ToLCNDV, AM 87, PI 414723 and PI 179901 when

192 inoculated with MeCMV. PCR analyses confirmed the presence of the viruses in all
193 symptomatic plants –including the “recovered” ones that showed almost no symptoms at
194 their apex, but also in a few asymptomatic ones (data not shown).

195 Accessions IC-274014, WM 7, WM 9, PI 124112 and PI 282448 did not show any
196 symptoms after inoculation by ToLCNDV or MeCMV. PCR results indicated the occasional
197 presence of viruses in asymptomatic plants (data not shown). Accessions AM 87 and PI
198 179901 presented a complete resistance to ToLCNDV but a heterogeneous behaviour
199 towards MeCMV, with some plants displaying yellow spots or mosaics (Table 1).

200

201 Determinism of IC-274014 resistance to MeCMV

202 Védrantais exhibited severe mosaic symptoms, stunting and enations when inoculated
203 with MeCMV and the virus was clearly detected by ELISA or PCR. IC-274014 exhibited no
204 symptom and the virus was not detected. The F_1 (IC-274014 \times Védrantais) exhibited
205 systemic chlorotic spots and the virus was detected in apical leaves (Figure 1), indicating a
206 recessive genetic control for virus infection. As expected for a recessive genetic control, the
207 BC_5 was fully susceptible (Table 2). All the BC_5 plants exhibited a range of symptoms: severe
208 mosaic and stunting, mild mosaic or mottle or systemic chlorotic spots and the virus was
209 detected by ELISA in all the plants. In the BC_R progeny, some plants had no symptoms but
210 the majority exhibited symptoms varying from systemic chlorotic spots to mottle; the virus
211 was not detected by ELISA and/or PCR in 29/95 plants (Table 2). This observed segregation
212 fitted a 3 susceptible : 1 resistant ratio expected for two independent recessive genes, both
213 required to confer resistance.

214 In the F_2 progeny, all types of symptoms were observed (mosaic, mottle, a majority with
215 systemic chlorotic spots) and some plants had no visible symptoms. The virus was detected

216 by ELISA and/or PCR in all the plants with symptoms and in a few plants without clear
217 symptoms. The observed segregation (409 susceptible vs. 20 resistant plants) was
218 compatible with a 15 susceptible (positive in ELISA or PCR) vs.1 resistant (negative in ELISA or
219 PCR) ratio ($\chi^2 = 1.846$; $df = 1$; Prob = 17%) corresponding to a genetic control by two
220 independent recessive genes required simultaneously for resistance.

221 After selfing F_2 resistant plants, two F_3 families (1A6 and 1A8) appeared resistant to
222 MeCMV (Table 3). The 8 other F_3 families obtained by selfing of resistant F_2 plants were
223 segregating, with some plants showing symptoms of mottling or systemic chlorotic spots and
224 virus systemic spread as detected by ELISA. If only two recessive genes were required and
225 sufficient to confer resistance to MeCMV, all the F_3 and F_4 plants issued from resistant F_2
226 plants should be resistant. It is the case only for 1A8 and the associated F_4 progenies. The F_3
227 1A6 appeared fully resistant but only six plants were tested and the corresponding F_4 were
228 either segregating (1A6A and 1A6B) or susceptible (1A6C). The F_3 1A6 can therefore be
229 considered as also segregating. The high frequency of segregating F_3 could be explained by
230 the need of a third dominant gene. This hypothesis of a third dominant gene does not
231 change the expected segregation in the F_1 , BC_S and BC_R progenies. Indeed, the F_1 and BC_S are
232 all susceptible because the recessive resistance alleles are not present in a homozygous state
233 whatever the state of the dominant allele. For the BC_R , the dominant resistance allele is
234 present in all plants either in a homozygous or a heterozygous state, so the resistance or
235 susceptibility depends only on the state of the two recessive alleles. Under the hypothesis of
236 two recessive and one dominant genes simultaneously required for resistance, the expected
237 segregation in the F_2 would be 61 susceptible vs. 3 resistant. The observed segregation (409
238 susceptible vs. 20 resistant plants) fits this hypothesis (Table 2).

239 In the case of two recessive and one dominant genes simultaneously required for
240 resistance, the F₂ resistant plants must be homozygous for the two recessive genes but can
241 be homozygous or heterozygous for the third dominant gene. If the 3 genes are
242 independent, 1/3 of the F₂ resistant plants should be homozygous for the dominant gene
243 producing homogeneous resistant F₃ progenies and 2/3 of the F₂ should be heterozygous,
244 producing segregating F₃ progenies. The observed segregation (1 homozygous F₃ progeny vs.
245 9 segregating ones) fits this hypothesis: $\chi^2 = 2.450$ (df = 1; Prob = 12%) even if the power of
246 the statistical analysis is low due to the small number of F₃ families available. Excluding the
247 resistant 1A8 family, the observed segregation for the number of susceptible or resistant
248 plants (43 susceptible vs 93 resistant, Table 3) among the 9 segregating F₃ families fits the
249 hypothesis of 1 susceptible vs 3 resistant corresponding to the expected segregation for a
250 single dominant gene ($\chi^2 = 3.177$; df = 1; Prob = 7%).

251 Similarly, in the F₄ generation obtained by selfing F₃ plants without selection for
252 resistance or susceptibility, it was expected a 3 susceptible vs. 5 resistant ratio in the number
253 of F₄ plants. As the number of plants in the F₄ families varies from 20 (corresponding to the
254 F₃ 1A4 or 1A11) to 80 (corresponding to the F₃ 1A5), it had to be adjusted to the same
255 number of F₄ plants for each F₃ progeny in order to test the 3:5 segregation (Supplementary
256 Table S1). The observed segregation fits this hypothesis ($\chi^2 = 0.085$, df = 1; Prob = 77.1%).

257 In summary, the segregations observed for resistance to MeCMV in all the progenies (F₁,
258 BC, F₂, F₃ and F₄) fitted the hypothesis of two recessive and one dominant independent
259 genes which can be provisionally symbolized *a*, *b* and *C*.

260

261 Resistance to ToLCNDV

262 After mechanical inoculation with ToLCNDV, Védrantais exhibited severe mosaic
 263 symptoms and the virus was detected in ELISA, while IC-274014 exhibited no symptom and
 264 the virus was not detected in ELISA. As for MeCMV, the F₁ was considered as susceptible
 265 since the virus was detected in all inoculated plants, indicating a recessive genetic control
 266 but the symptoms were only mild chlorotic spots.

267 In the F₂ progeny, 25 plants among 200 inoculated plants exhibited no symptom and were
 268 negative in ELISA, what did not correspond to a mono-or digenic inheritance (Supplementary
 269 Table S2). However this observed segregation fitted a 55 susceptible vs. 9 resistant expected
 270 segregation corresponding to one recessive and two dominant genes simultaneously
 271 required for resistance ($\chi^2 = 0.404$; df = 1; Prob = 52.5%). These genes can be provisionally
 272 symbolized *x*, *Y* and *Z*.

273

274 Relationship between resistance to MeCMV and ToLCNDV

275 The F₃ and F₄ progenies derived from F₂ plants resistant to MeCMV were inoculated with
 276 ToLCNDV (Table 3). Among 157 F₃ and 250 F₄ plants, 95 and 70 respectively were resistant to
 277 ToLCNDV. The F₃ 1A8 and the two corresponding F₄ 1A8A and 1A8B which were
 278 homogeneously resistant to MeCMV were also homogeneously resistant to ToLCNDV. This
 279 suggested a linkage or common genes for resistance to both viruses. A first hypothesis was a
 280 linkage between one of the genes for MeCMV resistance and one of the genes for ToLCNDV
 281 resistance with four subcases: (i) *a* (or *b*) linked with *x*, (ii) *a* (or *b*) linked with *Y* (or *Z*), (iii) *C*
 282 linked with *x*, and (iv) *C* linked with *Y*(or *Z*). A second hypothesis was two linkages with five
 283 subcases: (i) *a* linked with *x* and *b* linked with *Y* (or *Z*), (ii) *a* linked with *Y* and *b* linked with *Z*,
 284 (iii) *a* linked with *Y* (or *Z*) and *C* linked with *x*, (iv) *a* linked with *Y* and *C* linked with *Z*, and (v) *a*
 285 linked with *x* and *C* linked with *Y* (or *Z*). The expected frequencies of resistant and

286 susceptible plants to ToLCNDV were calculated under the conditions that *a* and *b* were
 287 homozygous and that *C* was heterozygous CC^+ or homozygous CC for respectively 2/3 or 1/3
 288 of the F_3 progenies. Among these nine possibilities, two fit the observed segregation in the
 289 F_3 progenies: linkages between *a* and *x* and between *b* and *Y* (Prob $\chi^2 = 54.7\%$) and linkages
 290 between *a* and *x* and between *C* and *Y* (Prob $\chi^2 = 20.5\%$) (Supplementary table S3). For the F_4
 291 progenies, the probabilities of the χ^2 were respectively $2.9 \cdot 10^{-7}$ and 4.3% with in both
 292 cases an excess of observed susceptible plants (Supplementary Table S3). The best
 293 hypothesis is thus: *a* linked with *x*, and *C* linked with *Y*, whereas gene *b* involved in MeCMV
 294 resistance and gene *Z* involved in ToLCNDV resistance are independent. F_3 plants
 295 homozygous resistant to both viruses such as the F_3 1A8 could be expected with a frequency
 296 of 1/12 (1/3 for *C-Y* and 1/4 for *Z*, *a-x* and *b* being homozygous for resistance to MeCMV)
 297 which corresponds to the observed frequency of one homozygous resistant F_3 out of 10. The
 298 expected segregation for the number of susceptible : segregating : resistant F_4 families
 299 towards ToLCNDV was 23 : 16 : 9. The observed values (respectively 8:13:2 in Table 3) fit this
 300 hypothesis ($\chi^2 = 5.778$; $df = 2$; Prob = 5.6%).

301

302 Correlation with resistance to other viruses

303 IC-274014 was found resistant to ZYMV after mechanical inoculation (Supplementary
 304 Table S4). Among the 23 F_4 (IC-274014 \times Védraçais) progenies, 14 were symptomless, 5
 305 displayed more or less severe symptoms of mosaics, yellowing and stunting, and 4 had an
 306 heterogeneous behaviour (no symptoms, chlorotic spots or mosaics). Among the 14
 307 symptomless F_4 progenies, 5 were positive in ELISA for at least one plant, being thus
 308 considered as heterogeneous (F_4 progenies 1A1A, 1A3B, 1A5D and 1A7B) or susceptible (F_4
 309 progeny 1A1B) (Supplementary Table S4 and data not shown).

310 In conditions of natural infection, IC-274014 was resistant to CABYV whereas Védraçais,
 311 the F₁ (IC-274014 × Védraçais) and the six tested F₄ progenies were susceptible to the virus
 312 based on ELISA (Table 4). Among the tested accessions, HSD 2445-005, PI 414723 and WM9
 313 were resistant to CABYV, whereas PI 164323, PI 164723, PI 179901, PI 234607, PI 236355, PI
 314 482420, Védraçais and WM7 were susceptible based on ELISA (Table 4). PI 414723 and
 315 Védraçais were confirmed as resistant and susceptible respectively. All accessions were
 316 susceptible to CMV and WMV, whereas ZYMV was not present in the experimental plot
 317 (data not shown)

318

319 Discussion and Conclusions

320 Among the 31 melon accessions tested against MeCMV and ToLCNDV, most had similar
 321 behaviours towards both viruses. MeCMV and ToLCNDV are quite distant molecularly (less
 322 than 60% and 50% identity in DNA-A and DNA-B, respectively), and belong to different
 323 clades (NW vs. OW). However, MeCMV and ToLCNDV share a high infectivity when
 324 mechanically inoculated, a rare trait among begomoviruses even though it has been
 325 described in a few ones particularly in the NW clade (Morales and Niessen, 1988; Wege and
 326 Pohl, 2007). This suggests that, like a few other begomoviruses, they are not exclusively
 327 phloem-limited, at least at some stages of their host development (Sudarshana et al. 1998).
 328 A common resistance mechanism may thus target the same key parameter in the cycle of
 329 these two viruses. Both viruses were also found associated with atypical alphasatellites in
 330 natural conditions (Anwar, 2017; Romay et al. 2010). ToLCNDV has also been found
 331 associated with a betasatellite (Anwar, 2017), contrary to MeCMV. The two viruses differ
 332 strikingly by their host range. MeCMV appears restricted to a few cucurbit hosts (Romay et
 333 al. 2015), whereas ToLCNDV has a broad host range including more than 40 species from a

334 range of families including *Cucurbitaceae*, *Solanaceae*, *Malvaceae*, *Amaranthaceae*,
335 *Euphorbiaceae* and *Fabaceae* (Zaidi et al. 2017). In several accessions, intermediate
336 resistance mechanisms appeared related to a “recovery” phenotype, as observed after
337 melon or watermelon infection with CuLCrV (Hagen et al. 2008a). This suggests that the
338 resistance could be related in these cases to RNA silencing (Ghoshal and Sanfaçon, 2015;
339 Hagen et al. 2008a) although the actual mechanism remains unknown.

340 Accession IC-274014 was resistant (no symptoms and no virus detection by ELISA or PCR)
341 to MeCMV. Different types of symptoms were observed in the F₁, F₂ and BC progenies
342 between IC-274014 and the susceptible cultivar Védrantais: severe mosaic, mottle, systemic
343 chlorotic spots. Inheritance of these different types of symptoms was difficult to analyse in
344 terms of genetic control. The F₁ exhibited systemic chlorotic spots which were not observed
345 in the parents. In the F₂ or BC progenies, “recovery” phenotypes were often observed, with
346 much higher symptom severity in the old leaves than in young ones. In plants with the
347 different types of symptoms, the virus was always detected by ELISA or PCR. When using this
348 criterion (presence vs. absence of virus), the observed segregations in the analysed
349 progenies fitted the hypothesis of one dominant and two recessive independent genes
350 simultaneously required for resistance.

351 Resistance of IC-274014 to ToLCNDV could be controlled by one recessive and two
352 dominant genes. As some progenies selected at the F₂ stage only for MeCMV resistance
353 were also resistant to ToLCNDV, we concluded that common genes were involved in
354 resistance to both viruses, even if the hypothesis of genes belonging to the same clusters
355 cannot be completely ruled out, clusters of resistance genes being present in melon (Garcia-
356 Mas et al. 2012; Gonzalez et al. 2014). We propose the names *begomovirus resistance-1* and
357 *Begomovirus resistance-2* for these genes (symbols *bgm-1* and *Bgm-2*) corresponding

358 respectively to *a-x* and *C-Y* in the results section. For resistance to MeCMV, one more
 359 recessive gene was required corresponding to the locus *b* in the results section; the name
 360 *mecmv resistance* (symbol *mecmv*) is proposed. Full resistance to MeCMV was controlled by
 361 *bgm-1*, *Bgm-2* and *mecmv*. Resistance to ToLCNDV was controlled by *bgm-1*, *Bgm-2* and an
 362 additional dominant gene (proposed name *ToLCNDV resistance*, symbol *Tolcndv*)
 363 corresponding to the gene *Z* in the results section.

364 The susceptible F_4 plants after inoculation with MeCMV or ToLCNDV exhibited mild
 365 symptoms (mottling or systemic chlorotic lesions) and not the severe symptoms (mosaic and
 366 stunting) observed in the susceptible control Védraçais. Consequently, *bgm-1* and/or *Bgm-2*
 367 could be involved in the control of the severe symptoms.

368 Accession IC-274014 is also resistant to *Zucchini yellow mosaic virus* (ZYMV) and *Cucurbit*
 369 *aphid-borne yellow virus* (CABYV), but no correlation was observed between resistance to
 370 ZYMV or CABYV on the one hand, and to MeCMV or ToLCNDV on the other hand
 371 (Supplementary Table S4).

372 Genes conferring resistance to several viruses from the same family, or even to
 373 completely different pathogens, have already been described. This can be dominant genes
 374 (e.g. the *L* gene conferring resistance to 7 tobamoviruses in pepper, see Moury and Verdin,
 375 2012) or recessive ones. Recessive resistance is often related to the loss or modification of a
 376 susceptibility factor required for the viral cycle, notably eukaryotic initiation factor 4E
 377 (Kuwata, 2016; Robaglia and Caranta, 2006). A recessive resistance gene to the begomovirus
 378 tomato yellow leaf curl virus, *ty-5*, corresponds to the tomato homolog of the messenger
 379 RNA surveillance factor Pelota (Lapidot et al. 2015) and conferred resistance in field
 380 conditions to at least five tomato-infecting begomovirus species (Al-Shihi et al. 2018). The
 381 resistance abolished symptoms and reduced viral accumulation (Anbinder et al. 2009).

382 Besides *ty-5*, additional minor quantitative trait loci contributed to the resistance (Anbinder
383 et al. 2009). A similar situation may happen for MeCMV and ToLCNDV resistance in melon,
384 although the nature of *bgm-1* and the mechanisms of resistance remain unknown so far.

385
386 A pool of melon accessions with resistance to one or several begomoviruses is emerging
387 from the literature (Table 4). Most of them are from India and belong to the acidulus,
388 momordica or kachri groups (Pitrat, 2016b). The most recent publication on genetic control
389 was on the resistance to ToLCNDV of the accession WM 7 from India belonging to the kachri
390 group: a major dominant gene (on linkage group LG XI) and two minor quantitative trait loci
391 (QTLs) (on LG II and LG XII) (Saez et al. 2017). From our results, resistance in IC-274014 is also
392 controlled by three genes, one recessive (*bgm-1*) and two dominant ones (*Bgm-2* and
393 *Tolcndv*) suggesting that common genes could be involved in WM 7 and IC-274014. It would
394 be interesting to search for the candidate regions for the different resistance genes.

395 Several accessions belonging to different horticultural types (cantaloupes, honeydew,
396 cassaba...) were resistant to CuLCrV (Hagen et al. 2008b) after agroinoculation. Resistance in
397 the accession PI 313970 after inoculation by *B. tabaci* was controlled by one recessive gene,
398 *culcrv* (McCreight et al. 2008). Allelism tests indicate that the same gene was also probably
399 present in other resistant accessions such as MR-1, PI 124111, PI 124112, PI 179901, PI
400 234607, and PI 236355 (McCreight et al. 2008); PI 236355 which exhibited the best level of
401 resistance to CuLCrV was susceptible to MeCMV and ToLCNDV (Table 4). So the gene *culcrv*
402 is probably not involved in resistance to these two last viruses, and different from *bgm-1* or
403 *mecmv*.

404 Resistance to WmCSV after *Agrobacterium*-mediated inoculation was observed in several
405 accessions (Yousif et al. 2007). Accessions resistant to WmCSV were also partially or

406 completely resistant to MeCMV and ToLCNDV (Table 4). The inheritance of resistance to
 407 WmCSV has not been published. It would be interesting to test IC-274014 and the F₄ (IC-
 408 274014 × Védraçais) resistant or susceptible to MeCMV and ToLCNDV for their behaviour
 409 when inoculated with WmCSV.

410 Our results suggest that although some resistance factors to begomoviruses are specific,
 411 common broad-spectrum resistance genes may be present in melon germplasm and could
 412 constitute interesting tools for breeders.

413

414 **Acknowledgements**

415 We thank the Experimental Infrastructure team of INRA-Montfavet for their help in the
 416 greenhouse experiments. We thank D. Besombes for obtaining part of the melon accessions
 417 and crosses and Dr J.E. Duffus (USDA, Salinas) for providing the SLCV antiserum.

418 This work was funded in part through the EMERAMB project in the ARIMNet2 2015 Call.
 419 ARIMNet2 (2014-2017) is an ERA-NET coordinated by INRA (France). It has received funding
 420 from the European Union's Seventh Framework Programme for research, technological
 421 development and demonstration under grant agreement no. 618127.

422

423 The authors declare that they have no conflict of interest

424

425

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- 540

Melon accession	Known resistance against cucurbit viruses ^b	MeCMV ^a	ToLCNDV ^a
Védrantais		S	S
Ouzbèque 1		S	S
Ouzbèque 2		S	S
Isabelle		S	S
Anso 77		S	S
AR Hale's Best Jumbo		S	S
Canton		S	S
HSD 93-20-A	CVYV (systemic necrosis) ^c	S	S
PMR-5	MNSV ^d	S	S
WMR29	PRSV ^e , MWMV ^e , MRMV ^e	S	S
PI 161375	CMV ^e , MNSV ^e	S	S
PI 164323	CVYV ^c	S	S
PI 179905	ZYMV ^c	S	S
PI 234607	<u>CuLCrV</u> ^f	S	S
PI 255478	CABYV ^g	S	S
PI 482420	CYSDV ^e , WMV ^c	S	S
PI 236355	<u>CuLCrV</u> ^f	S	S
PI 164723	<u>WmCSV</u> ^h	S	IR
AM 87		H (S/IR/R)	R
HSD 2445-005	<u>WmCSV</u> ^h	H (S/R)	H (R/IR)
HSD 2458 B	CVYV ^c	H (S/IR/R)	H (IR/R)
Tibish Kordofan		H (S/R)	H (S/IR/R)
MR-1	<u>CuLCrV</u> ^f , LIYV ^e	H (S/IR/R)	IR
PI 179901	<u>CuLCrV</u> ^f	H (S/IR/R)	R
PI 313970 (90625)	CABYV ^g , <u>CuLCrV</u> ^f , CYSDV ^c , LIYV ^c , <u>WmCSV</u> ^h	H (S/R)	H (S/IR/R)
PI 414723	CABYV ^g , <u>CuLCrV</u> ^f , PRSV ^e , <u>ToLCNDV</u> ⁱ , <u>WmCSV</u> ^h , WMV ^e , ZYMV ^e	H (S/IR/R)	H (IR/R)
IC-274014	CABYV ^c , ZYMV ^c	R	R
PI 124112	CABYV ^g , <u>CuLCrV</u> ^f , PRSV ^e , <u>ToLCNDV</u> ⁱ , <u>WmCSV</u> ^h	R	R
PI 282448	CABYV ^g , <u>WmCSV</u> ^h	R	R
WM7	<u>ToLCNDV</u> ⁱ	R	R
WM9	<u>ToLCNDV</u> ⁱ	R	R

541
542 Table 1. Behaviour of 31 melon accessions after artificial inoculation with two
543 begomoviruses: melon chlorotic mottle virus and tomato leaf curl New Delhi virus.

544 ^a R: resistant; S: susceptible; IR: intermediate resistance; H: heterogeneous.

545 ^b CABYV: cucurbit aphid-borne yellow virus (polerovirus), CMV: cucumber mosaic virus
546 (cucumovirus), CuLCrV: cucurbit leaf crumple virus (begomovirus), CYSDV: cucurbit yellow
547 stunting disorder virus (crinivirus), CVYV: cucumber vein yellowing virus (ipomovirus), LIYV:

548 lettuce infectious yellows virus (crinivirus), MNSV: melon necrotic spot virus (carmovirus),
549 MRMV: melon rugose mosaic virus (tymovirus), MWMV: Moroccan watermelon mosaic virus
550 (potyvirus), PRSV: papaya ringspot virus (potyvirus), WMV: watermelon mosaic virus
551 (potyvirus), WmCSV: watermelon chlorotic stunt virus (begomovirus), ZYMV: zucchini yellow
552 mosaic virus (potyvirus). Begomoviruses (CuLCrV, MeCMV, ToLCNDV, WmCSV) are
553 underlined.

554 ^cPitrat, 2016a

555 ^dMallor et al. 2003

556 ^eLecoq et al. 1998

557 ^fMcCreight et al. 2008

558 ^gDogimont et al. 1996

559 ^hYousif et al. 2007

560 ⁱLopez et al. 2015

561

	Test	Total	Susc. ^a	Res. ^a	Segreg. ^b	Chi-square	
						Value	Probability
Védrantais	1	30	30	0			
	2	30	30	0			
	3	10	10	0			
	Total	70	70	0			
IC-274014	1	25	0	25			
	2	25	0	25			
	Total	50	0	50			
F ₁ = IC- 274014 × Védrantais	1	25	25	0	1:0		
	2	22	22	0	1:0		
	Total	47	47	0	1:0		
F ₂ = F ₁ × F ₁	1	99	97	2	61:3	1.577	21%
	2	80	78	2	61:3	0.857	35%
	3	250	234	16	61:3	1.641	20%
	Total	429	409	20	61:3	0.006	98%
BC _S = F ₁ × Védrantais	1	50	50	0	1:0		
	2	47	47	0	1:0		
	Total	97	97	0	1:0		
BC _R = F ₁ × IC- 274014	1	50	40	10	3:1	0.667	41%
	2	45	26	19	3:1	7.119	0.8%
	Total	95	66	29	3:1	1.547	21%

562
563 Table 2. Number of plants observed after inoculation with MeCMV in the parents
564 (Védrantais and IC-274014), F₁, F₂ and back-cross (BC) progenies
565 ^a Susceptible = with clear virus symptoms and/or positive detection by ELISA or PCR;
566 Resistant = no symptom and no virus detection by ELISA or PCR.
567 ^b Expected segregation (Susceptible : Resistant) under the hypothesis of two recessive and
568 one dominant genes.

569

Melon genotypes	MeCMV		ToLCNDV		Melon genotypes	MeCMV		ToLCNDV	
	Susc.	Res.	Susc.	Res.		Susc.	Res.	Susc.	Res.
Védrantais	16	0	16	0	Védrantais	20	0	10	0
IC-274014	0	15	0	20	IC-274014	0	20	0	10
F ₃ progenies					F ₄ progenies				
1A1	3	7	4	6	1A1A	4	6	10	0
					1A1B	14	5	14	6
1A2	4	14	13	5	1A2A	17	3	20	0
					1A2B	1	9	9	1
1A3	5	15	7	13	1A3A	5	5	6	4
					1A3B	0	25	3	7
1A4	14	3	6	11	1A4A	3	7	10	0
					1A4B	7	3	8	2
1A5	5	4	5	4	1A5A	1	19	9	1
					1A5B	0	20	10	0
					1A5C	1	19	10	0
					1A5D	0	20	10	0
1A6	0	6	3	4	1A6A	1	7	10	0
					1A6B	1	4	9	1
					1A6C	16	4	10	0
1A7	6	14	6	14	1A7A	4	4	5	5
					1A7B	1	19	3	7
1A8	0	17	0	18	1A8A	0	20	0	10
					1A8B	0	20	0	10
1A10	2	17	5	15	1A10A	3	7	2	8
					1A10B	8	2	6	4
1A11	4	13	13	5	1A11A	1	19	7	3
					1A11B	3	13	9	1

570

571 Table 3: Number of plants susceptible or resistant to MeCMV and ToLCNDV in the parents

572 (Védrantais and IC-274014) and in the F₃ and F₄ progenies from F₂ plants resistant to

573 MeCMV.

574

27

Accession	MeCMV ^a	ToLCNDV ^a	CuLCrV ^d	WmCSV ^e	CABYV ^{a,b}
Védrantais	S	S	nt	S	S
HSD 2445-005	Segreg (R/S)	Segreg (R/S)	nt	R	R
IC-274014	R	R	nt	nt	R
Kharbuja	nt	IR ^f	nt	nt	nt
PI 124111 (MR-1)	IR	IR	R	nt	nt
PI 124112	R	IR ^f /R	IR	R	R ^g
PI 164323	S	S	nt	nt	S
PI 164723	S	IR	R/IR	R	S
PI 179901	IR	R	IR	nt	S
PI 234607	S	S	R	nt	S
PI 236355	S	S	R	nt	S
PI 282448	R	R	nt	R	R ^g
PI 313970 (90625)	R	IR	R	R	R ^g
PI 414723	IR	IR	IR	R	R
WM 7	R	R ^f /R	nt	nt	S ^c
WM 9	R	IR ^f /R	nt	nt	R

575 Table 4. Behaviour of melon accessions (S = susceptibility, R = resistance, IR = partial
576 resistance, nt = not tested) towards the four Begomoviruses melon chlorotic mosaic virus
577 (MeCMV), tomato leaf curl New Delhi virus (ToLCNDV), cucurbit leaf crumple virus (CuLCrV)
578 and watermelon chlorotic spot virus (WmCSV) and the polerovirus cucurbit aphid-borne
579 yellows virus (CABYV).

580 ^aThis work, unless otherwise specified

581 ^b results from field trial based on ELISA test

582 ^c low ELISA values

583 ^dMcCreight et al. 2008

584 ^eYousif et al. 2007

585 ^fLopez et al. 2015

586 ^gDogimont et al. 1996

587

588 Figure 1: Symptoms of MeCMV: curling, yellowing and stunting in the susceptible cv.
589 Védrantais (a); no symptom and no virus multiplication in the resistant accession IC-274014
590 (b); systemic chlorotic spots in the F₁ (IC-274014 × Védrantais)
591
592

593 Supplementary Table S1: Observed number of plants susceptible and resistant to MeCMV in
594 the F₄ (IC-274014 × Védrrantais) progenies after selection of the F₂ for resistance to this
595 virus, and adjusted number for the same size of F₄ plants for the different F₃ progenies.

596

597 Supplementary Table S2: Expected segregations for resistance to ToLCNDV in the F₂ (IC-
598 274014 × Védrrantais) under different hypotheses

599

600 Supplementary Table S3: Expected vs. observed frequency for susceptible vs. resistant plants
601 to ToLCNDV in the F₃ and F₄ progenies derived from F₂ (IC-274014 × Védrrantais) resistant to
602 MeCMV, and probability of the chi-square under different hypotheses of linkage (or identity)
603 of genes involved in resistance to MeCMV (*a*, *b* and *C*) and to ToLCNDV (*x*, *Y* and *Z*).

604

605 Supplementary table S4: behaviour of Védrrantais, IC-274014, F₁ and F₄ (IC-274014
606 × Védrrantais) progenies towards CABYV and ZYMV, in relation to their susceptibility to the
607 begomoviruses MeCMV and ToLCNDV. The F₄ progenies are from F₂ plants resistant to
608 MeCMV.

609

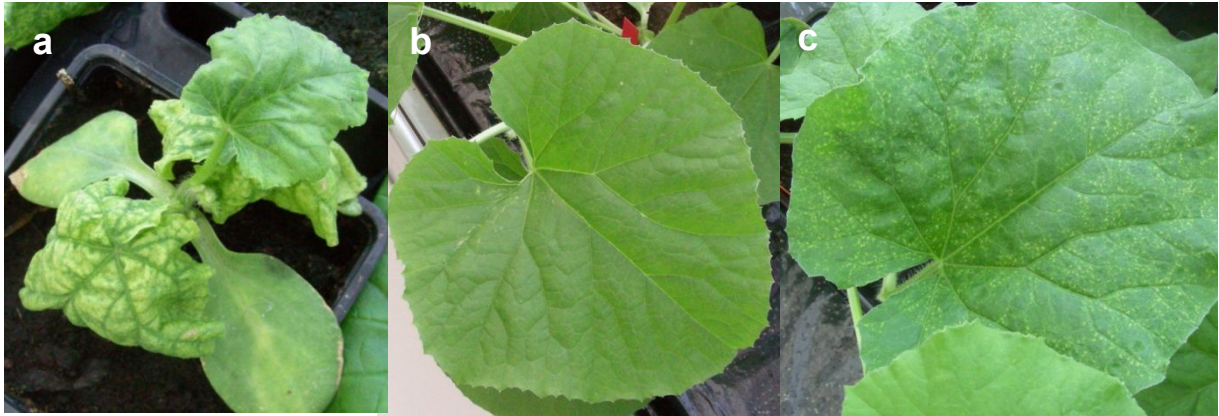


Figure 1

F ₄ progeny	Observed			Adjusted		
	Susceptible	Resistant	Total	Susceptible	Resistant	Total
1A1A	4	6	10	21.8	13.3	35.1
1A1B	14	5	19			
1A2A	17	3	20	21.1	14.0	35.1
1A2B	1	9	10			
1A3A	5	5	10	5.0	30.1	35.1
1A3B	0	25	25			
1A4A	3	7	10	17.6	17.6	35.1
1A4B	7	3	10			
1A5A	1	19	20	0.9	34.2	35.1
1A5B	0	20	20			
1A5C	1	19	20			
1A5D	0	20	20			
1A6A	1	7	8	19.1	16.0	35.1
1A6B	1	4	5			
1A6C	16	4	20			
1A7A	4	4	8	6.3	28.8	35.1
1A7B	1	19	20			
1A10A	3	7	10	19.3	15.8	35.1
1A10B	8	2	10			
1A11A	1	19	20	3.9	31.2	35.1
1A11B	3	13	16			
Total	91	220	311	113.12	197.88	311

Supplementary Table S1: Observed number of plants susceptible and resistant to MeCMV in the F₄ (IC-274014 × Védraçais) progenies after selection of the F₂ for resistance to this virus, and adjusted number for the same size of F₄ plants for the different F₃ progenies.

	Number of plants		Segregation	Chi-square	
	Susceptible	Resistant		Value	Prob
Observed	175	25			
Hypothesis					
1 recessive	150	50	3 Sus:1 Res	16.667	< 0.0001
2 recessive	187.5	12.5	15 Sus:1 Res	13.333	0.0003
1 recessive+1 dominant	162.5	37.5	13 Sus:3 Res	5.128	0.0235
1 recessive + 2 dominant	171.875	28.125	55 Sus:9 Res	0.404	0.525

Supplementary Table S2: Expected segregations for resistance to ToLCNDV in the F₂ (IC-274014 × Védraçais) under different hypotheses

	F ₃		F ₄	
	Susceptible:Resistant	Proba χ^2	Susceptible:Resistant	Proba χ^2
Observed frequency				
-uncorrected	62:95		180:70	
-adjusted*	64.08:92.92		166.46:83.54	
One linkage				
C with Y (or Z)	103:25	6.4 10 ⁻¹³	835:189	1.36 10 ⁻⁴
a (or b) with Y (or Z)	49:15	1.3 10 ⁻¹⁰	193:63	0.030
a (or b) with x	39:25	3.6 10 ⁻⁴	175:81	0.67
C with x	103:25	6.4 10 ⁻¹³	835:189	1.36 10 ⁻⁴
Two linkages				
a with x	3:5	0.547	7:9	2.9 10 ⁻⁷
and b with Y (or Z)				
a with Y	5:3	1.2 10 ⁻⁴	9:7	0.018
and b with Z				
a (or b) with Y (or Z)	11:5	6.6 10 ⁻⁷	43:21	0.886
and C with x				
a (or b) with Y	11:5	6.6 10 ⁻⁷	43:21	0.886
and C with Z				
a (or b) with x	23:25	0.2054	37:27	0.0432
and C with Y (or Z)				

Supplementary Table S3: Expected vs. observed frequency for susceptible vs. resistant plants to ToLCNDV in the F₃ and F₄ progenies derived from F₂ (IC-274014 × Védraçais) resistant to MeCMV, and probability of the chi-square under different hypotheses of linkage (or identity) of genes involved in resistance to MeCMV (*a*, *b* and *C*) and to ToLCNDV (*x*, *Y* and *Z*).

*The observed frequencies were adjusted to take into account the difference in the number of plants tested for the different F₃ and F₄ progenies.

The F₂ plants used to obtain the F₃ and F₄ progenies were selected for MeCMV resistance, so they were homozygous for genes *a* and *b* (and for the linked or identical genes for ToLCNDV resistance) and homozygous *CC* (for 1/3 of the plants) or heterozygous *CC*⁺ (for 2/3 of the plants) for the third dominant gene.

If there are some linkages between the genes involved in resistance to MeCMV and ToLCNDV, genes linked with *a* or *b* must be homozygous in the F₂ selected for resistance to MeCMV, and no more segregating in the F₃ and F₄ progenies. Genes linked with *C* are homozygous and heterozygous in 1/3 and 2/3 respectively of the MeCMV-resistant F₂. The other genes segregate independently, with frequencies of ¼ homozygous dominant, ½ heterozygous and ¼ homozygous recessive in the F₂. The expected frequencies in the F₃ and

F₄ progenies obtained without further selection for virus resistance were calculated based on these frequencies in the F₂ plants.

	MeCMV	ToLCNDV	ZYMV	CABYV
Védrantais	S ^a	S	S	S
IC-274014	R	R	R	R
F1 (IC × Ved)	IR	IR	R	S
1A1A	H	S	H	nt
1A1B	S	H	S	S
1A2A	H	S	S	nt
1A2B	H	H	S	nt
1A3A	H	H	R	nt
1A3B	R	H	H	S
1A4A	H	S	H	nt
1A4B	H	H	H	nt
1A5A	R/H	H	R	nt
1A5B	R	S	R	S
1A5C	R/H	S	R	nt
1A5D	R	S	H	S
1A6A	H	S	S	nt
1A6B	H	H	H	nt
1A6C	S/H	S	H	nt
1A7A	H	H	R	nt
1A7B	R/H	H	H	nt
1A8A	R	R	R	S
1A8B	R	R	R	S
1A10A	H	H	R	nt
1A10B	H	H	R	nt
1A11A	R/H	H	S	nt
1A11B	R/H	H	S	nt

Supplementary table S4: behaviour of Védrantais, IC-274014, F₁ and F₄ (IC-274014 × Védrantais) progenies towards CABYV and ZYMV, in relation to their susceptibility to the begomoviruses MeCMV and ToLCNDV. The F₄ progenies are from F₂ plants resistant to MeCMV.

^aS = with clear virus symptoms and/or positive detection by ELISA; R = no symptoms, no virus detection by ELISA; H= heterogeneous based on symptoms and/or ELISA.

nt = not tested