

A new flavi-like virus identified in populations of wild carrots

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Diversity of polerovirus-associated RNAs in the virome of wild and cultivated carrots

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1 Abstract

The self-replicating polerovirus-associated RNAs have recently been classified as tombusvirus-2 like associated RNAs (tlaRNAs). In a metagenomic comparison of the virome of wild (Daucus 3 carota ssp. carota) and cultivated (D. carota ssp. sativus) carrots, four tlaRNAs were identified: 4 carrot red leaf virus-associated RNA (CtRLVaRNA), beet western vellows virus-associated 5 RNA (BWYVaRNA), and two other tlaRNAs detected for the first time in carrots, arracacha 6 latent virus E-associated RNA (ALVEaRNA) and a new tlaRNA tentatively named carrot red 7 leaf virus-associated RNA 2 (CtRLVaRNA-2). Their genomic sequences show a typical 8 9 tlaRNA genome organization with a very short 5' UTR, a long 3' UTR and consecutive ORFs, ORF1a and ORF1b separated by an amber stop codon, leading to the production of 10 ORF1a/ORF1b RdRp fusion protein. The genetic diversity and prevalence of these various 11 12 agents were analysed in carrot populations in France and Spain. CtRLVaRNA was the most frequently detected in both wild and cultivated carrot populations and was genetically more 13 14 heterogeneous than ALVEaRNA and CtRLVaRNA-2. ALVEaRNA was detected both in France and Spain and showed preferential association with cultivated carrots. In contrast, 15 CtRLVaRNA-2 was only detected from one region of France and was preferentially associated 16 with wild carrot populations. Analysis of the virome of individual plants showed the presence 17 of CtRLV but not that of the expected helper virus for ALVEaRNA and BWYVaRNA 18 suggesting a degree of flexibility in their associations with helper viruses, which in turn may 19 have biological and epidemiological consequences. 20

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Key words: associated RNA, satellite, high-throughput sequencing, *Polerovirus, Daucus carota*

24 INTRODUCTION

An important group of plant pathogens are satellites, which are subviral agents known to 25 interact with co-infecting viruses and the host cellular machinery (Gnanasekaran & 26 27 Chakraborty, 2018). A common feature of satellites is their dependence on a helper-virus to complete a full transmission cycle (Gnanasekaran and Chakraborty, 2018; Badar et al., 2021). 28 Satellites are currently divided into satellite viruses, encoding their own capsid protein but 29 relying on the helper virus for replication, and satellite nucleic acids, which vary in their 30 dependence on the helper virus for either encapsidation, cell-to-cell movement and/or 31 replication (Badar et al., 2021). 32

A class of single stranded satellite nucleic acids is often found in association with poleroviruses 33 and displays an autonomous replication capability (Campbell et al., 2020). However, these 34 35 satellites depend on co-infection partners of the family Solemoviridae (genus Polerovirus or Enamovirus) for encapsidation and systemic movement (Badar et al., 2021; Campbell et al., 36 37 2020). Recently, these Polerovirus-associated RNAs have been classified as tombusvirus-like 38 associated RNAs (tlaRNA) due to their phylogenetic affinities to viruses of the Tombusviridae family (Campbell et al., 2020). Satellite tlaRNAs are single-stranded, positive-sense RNAs of 39 about 3 kb harboring two open reading frames (ORFs), ORF1a and ORF1b. ORF1a encodes a 40 protein of about 30 kDa and ORF1b is expressed by the readthrough of ORF1a termination 41 codon, resulting in a fusion protein encoding the satellite RNA-dependent RNA polymerase 42 (RdRp) (Campbell et al., 2020; Peng et al., 2021). There are few studies addressing the 43 specificity of the association between satellites and their helper virus (Oncino, et al., 1995; 44 Kurath et al. 1993). This may be due to the fact that such disease complexes have been largely 45 46 analysed under controlled conditions, focusing on a specific virus-satellite association in a single host species (Peng et al., 2021; Mo et al., 2011), leading to a consequential knowledge 47 gap on such complexes in more natural infection conditions. The fact that isolates of beet 48

western yellows virus-associated RNA (BWYVaRNA) and Turnip yellows virus-associated 49 RNA (TuYVaRNA) show over 93% identity over their entire genome and the recent report of 50 the partial sequence of a potato leafroll virus-associated RNA (MF133518, Campbell et al., 51 2020) showing 99.5% nucleotide identity with carrot red leaf virus-associated RNA 52 (CtRLVaRNA) suggests however some level of promiscuous associations involving tlaRNAs. 53 An agriculturally important member of tlaRNAs is CtRLVaRNA, which is involved in the 54 55 carrot motley dwarf disease (CMD) complex, causing severe damage to commercial carrot crops (Adams, et al., 2014). CtRLVaRNA has a worldwide distribution, being reported from 56 the US (Watson., 1998; Campbell et al., 2020), Japan (Yoshida, 2020), New Zealand (Tang, et 57 58 al.,2009) and the UK (Adams, et al. 2014). Other viruses involved in the CMD complex are carrot red leaf virus (CtRLV, Polerovirus) and carrot mottle virus (CMoV) and/or carrot mottle 59 mimic virus (CMoMV), both of which belong to the genus Umbravirus. Umbraviruses (family 60 Tombusviridae) are well studied for their dependence for encapsidation/aphid transmission on 61 helper viruses of the family Solemoviridae (previously Luteoviridae). In addition to their RdRp, 62 63 umbraviruses encode two additional ORFs, ORF3 and ORF4, involved in cell-to cell and longdistance movement, respectively. In the CMD complex, the capsid protein of CtRLV (helper 64 virus) is used by the coinfecting umbraviruses and satellites (dependent viruses) for 65 66 transcapsidation of their genome, thus allowing their transmission by the aphid vectors of CtRLV. Satellites generally do not share any sequence similarity with their helper virus but 67 often modulate (exacerbate or attenuate) helper-virus induced symptoms, as well as helper virus 68 accumulation in infected hosts (Gnanasekaran, P. & Chakraborty, S., 2018; Badar, et al., 2021). 69 The accumulation of CtRLV has, for example, been shown to be reduced in plants coinfected 70 71 by CtRLVaRNA and CMoV (Yoshida, 2020). In contrast, BWYVaRNA, first described as an autonomously replicating subviral RNA (Chin et al., 1993), has been shown to stimulate the 72 73 accumulation of its helper virus, beet western yellows virus (BWYV, genus Polerovirus).

Similarly, the recently described pod pepper vein yellows virus-associated RNA has been shown to stimulate the accumulation of its helper virus, leading to increased virulence (Peng et al., 2021). Other examples of such tlaRNAs are tobacco bushy top disease-associated RNA (TBTDaRNA), which is involved in the tobacco bushy top disease complex (Mo et al., 2011) or arracacha latent virus E-associated RNA (ALVEaRNA) which was recently identified in the virome of *Arracacia xanthorrhiza*, characterized by high throughput sequencing (partial genomic sequences MF136436 and MF073199).

In a comparative metagenomic study aiming to understand specific virome fingerprints in cultivated and wild relative species, the viromes of different cultivated (*Daucus carota* ssp. *sativus*) and wild carrot (*D. carota* ssp. *carota*) populations were investigated. Four tlaRNAs were thus identified, two of which were detected for the first time in carrots, including a novel tlaRNA distantly related to CtRLVaRNA. The various tlaRNAs showed differences in their distribution and prevalence and, for some of them, preferential association with wild or cultivated carrot populations.

88 MATERIALS AND METHODS

89 **Plant samples collection and preparation**

90 In summer 2019, 16 different populations of cultivated carrots (Daucus carota ssp. sativus) and wild carrots (Daucus carota ssp. carota) were sampled locally in the Nouvelle-Aquitaine region 91 92 of France, including four cultivated populations, four off-type populations growing within the fields (their early bolting phenology and root morphology suggest that they originate from 93 pollen or seed contaminations of the planted commercial varieties and they are therefore 94 95 referred to here as off-type) and eight wild carrot populations. In summer 2020, a similar sampling was conducted in multiple areas of France, covering a north-south country gradient. 96 In total, 29 carrot populations were collected, comprising 11 cultivated carrots populations, 97

three off-type populations and 15 wild populations. A total of 45 carrot populations were 98 therefore sampled in France over two years. A similar sampling was performed in early summer 99 2021 near Segovia (Central Spain), including five cultivated and one wild carrot populations. 100 101 The characteristics of all sampled carrot populations are provided in Supplementary Table S1. Plants were sampled independently of the presence of viral symptoms but plants with obvious 102 fungal infestation or necrosis were excluded. Leaf samples were stored desiccated of anhydrous 103 104 CaCl₂ until used. For each population, a pool corresponding to 50 plants was assembled (about 0.1g of leaf/plant). 105

106 Double-stranded RNAs purification, Illumina sequencing and virome assembly and 107 annotation

108 Double-stranded (ds) RNAs were purified from each pool of carrot leaves. In the case of two French carrot populations (one wild and one cultivated), dsRNAs were also extracted from 109 leaves of the 50 individual plants (75 mg per plant). Double-stranded RNAs were purified by 110 two rounds of CF41 cellulose chromatography and nuclease treatment as described (Marais et 111 112 al., 2018). A negative extraction control was included in each extraction by using only buffer. Purified dsRNAs were converted to cDNA and random amplified while simultaneously adding 113 MID tags (François et al., 2018; Marais et al., 2018). PCR products were purified using the 114 115 MinElute PCR Purification Kit (Qiagen SAS France, Courtaboeuf, France) and their concentration determined spectrophotometrically (Marais et al., 2018). Amplification products 116 were sequenced in multiplexed format (2×150 bp) on an Illumina NovaSeq 6000 system at the 117 GetPlaGe platform (GenoToul INRAE Toulouse, France). 118

119 Sequencing reads were imported into CLC Genomics Workbench, version 21.0.3, 120 demultiplexed and trimmed on quality and length using default setting and a minimum read 121 length of 60 nucleotides (nt) and subsequently assembled *de novo* with the following 122 parameters: word size: 50, bubble size: 300, minimal contig length: 250 nt. Viral contigs were

identified by a comparison against the Virus REFSEQ GenBank database using BlastX 123 (Altschul et al., 1990). Contigs or scaffold thus identified for tlaRNAs were extended by rounds 124 of mapping of remaining reads. To confirm the HTS sequencing results, specific primers were 125 designed for each of the identified tlaRNAs (Supplementary Table S2) and pools or individual 126 plant samples were tested by two-step RT-PCR (Marais et al., 2011). If necessary, the 5' and 127 3' terminal sequences of individual tlaRNAs were confirmed using specific primers 128 (Supplementary Table S2) and the SMARTer RACE Kit (Takara Bio Europe SAS, Saint-129 Germain-en-Laye, France). 130

131 Analysis of tlaRNAs genomic sequences and Phylogenetic analyses

Multiple alignments of the fully reconstructed genomes of the known and new tlaRNAs 132 133 identified in the carrot virome and of tlaRNAs sequences retrieved from GenBank were prepared using ClustalW as implemented in Mega 7 (Kumar et al., 2018). ORFs were identified 134 in tlaRNAs sequences with CLC GW and compared with sequences in GenBank using the 135 BlastP search tool to verify ORF predictions. Recombination events in tlaRNAs were analysed 136 based on a multiple alignment of complete genomic sequences and using the RDP4 package 137 (Martin et al., 2015). Only recombination events detected by at least four out of seven 138 implemented algorithms were considered. Neighbour-joining trees were inferred from 139 alignments of whole genomes or of the RdRp central conserved domain and strict nucleotide 140 (nt) or amino acid (aa) identities calculated using MEGA 7. The significance of branches was 141 estimated with 1000 bootstrap replicates. 142

143 **RESULTS**

144 Identification of known and novel tlaRNAs in the virome of wild and cultivated carrot
145 populations

Following de novo assembly of cleaned reads from plant pools or from individually sequenced 146 147 plants of wild and cultivated carrots, a large number of viral contigs were identified via BlastXbased contigs annotation. Among them, multiple contigs showed significant homology to 148 149 known tlaRNAs. Preliminary analyses revealed that the identified tlaRNAs formed four distinct groups corresponding to isolates closely related to CtRLVaRNA, BWYVaRNA and the partial 150 reported sequence for ALVEaRNA. A fourth group of sequences was more distantly related to 151 152 CtRLVaRNA. Besides CtRLVaRNA and BWYVaRNA, no other associated RNA had been reported in carrots before. Full genomes of CRLVaRNA and BWYVaRNA were obtained by 153 extension of contigs or scaffolds using CLC Genomics Workbench v21 as explained in 154 Materials and Methods. In the case of some isolates, full genome sequences were directly 155 obtained as judged from comparison with the conserved 3' and 5' untranslated regions (UTR) 156 (Campbell et al., 2020) of GenBank reference sequences, obviating the need to perform RACE 157 experiment. For some others, a few 5' or 3' nucleotides were missing but no effort was made to 158 determine the 5' end 3' genome ends. The sequences of CRLVaRNA and BWYVaRNA isolates 159 thus determined have been deposited in GenBank under accession numbers ON603907 -160 161 ON603916.

162 The sequence of an isolate representative of the fourth group only distantly related to

163 CtRLVaRNA was assembled from the sequencing library from an individually extracted wild

164 carrot (FR19-9, Table S1). The assembly covered the entire genome, which was confirmed by

sequence comparisons with the conserved 3' and 5' (UTR) of CtRLVaRNAs so that RACE

166 experiments were not needed. The finalized genome (2,840 nt long, GenBank ON603908)

included 355,574 reads (~4 % of total reads for that sample) with a high average coverage of

168 14,177x. The name carrot red leaf virus-associated RNA-2 (CtRLVaRNA-2) has been

169 proposed for this novel molecule (see below).

8

The full genome of an isolate representative of the group showing close homology (93% nt 170 171 identity) with ALVEaRNA was obtained from the assembly derived from the sequencing reads from a single cultivated carrot plant (FR19-7, Table S1). The initial large contig identified was 172 extended through repeated round of residual reads mapping and the 5' and 3' genome ends 173 determined by RACE experiments. The full genome (GenBank ON603911) is 2,864 nt long 174 and integrates 82,202 reads corresponding to 4.51% of total reads for this sample with an 175 average coverage of 3,242x. Detailed information on all full genomes of tlaRNA isolates 176 identified in the present study are provided in Supplementary Table S3. 177

178 Genome organization of the identified tlaRNAs

The genome organization of all identified tlaRNAs isolates shows the typical genome 179 180 organization for these molecules (Sanger, 1994; Campbell et al., 2020 & Yoshida, 2020) with a very short 5' UTR (Campbell, 2020), two conserved ORFs, ORF1a and ORF1b in the same 181 reading frame and a long 3' UTR (Figure 1). The 5' UTR of the novel CtRLVaRNA-2 (GenBank 182 ON603908) is 14 nt long, while the situation is more complex with ALVEaRNA (GenBank 183 ON603911) since it shows three AUG initiation codons in close succession at its 5' genome end 184 at nt positions 5-7, 8-10 and 29-31. None of these is in a good Kozak initiation context but the 185 most likely candidate appears to be that at positions 29-31, since it leads to a protein starting 186 with the MCAALS sequence which is similar to the MQAALS sequence of CtRLVaRNA-2 187 (Figure 1). The ORF1a of CtRLVaRNA-2 and ALVEaRNA (696 and 757 nt, respectively) 188 encode proteins of about 25 kDa. For all analyzed isolates, this ORF is terminated by an amber 189 190 stop codon immediately followed in CtRLVaRNA and in other tlaRNAs by a GGL triplet. The sole exception is ALVEaRNA for which the GGL triplet is located upstream of the amber stop 191 codon (Figure S1). Readthrough of the amber stop codon generates the ORF1a-ORF1b RdRp 192 fusion protein. The ORFs of CtRLVaRNA-2 and ALVEaRNA are roughly colinear with 193 respective length and genome positions of 2289 nt (position 15-2303) and 2343 nt (position 29-194

197 *Recombination analysis and phylogenetic affinities of the various tlaRNAs identified in the*198 *carrot virome*

199 To reveal their phylogenetic affinities and taxonomic position, the complete genomes of the various tlaRNAs identified in the carrot virome were aligned with complete tlaRNA genomes 200 retrived from the GenBank database. A recombination analysis using the RDP4 package and 201 202 this whole genome alignment revealed a single significant recombination event, which was detected by all seven algorithms (corrected probability for the null hypothesis of 5e⁻³³) and 203 involving a ca. 100 nt segment of CtRLVaRNA integrated near the 5' end of CABYVaRNA 204 205 (genome positions 22-120). This recombination event had been previously identified (Campbell et al., 2020). 206

The same multiple alignment was then used to reconstruct the neighbour joining phylogenetic 207 tree of Figure 2. As previously shown by Campbell et al. (2020), tlaRNAs cluster into three 208 different clades supported by high bootstrap values and that show a high between-group average 209 divergence of 51.3-54.4% +/- 1%. The most diverse 210 nt clade assembles TuYVaRNA/BWYVaRNA plus CABYVaRNA, PeVYVaRNA, PoPeYVaRNA, TBTDaRNA 211 and TVDVaRNA (Figure 2). The second clade includes CtRLVaRNA and CtRLVaRNA-2, 212 213 with an average nt divergence between them of 31.1% + 0.7% (27.0% + 1.3% aa divergence for the viral fusion protein) and the last clade includes only ALVEaRNA, with an average 214 divergence between the partial sequences from Arracacha and those from carrot of only 6.8% 215 216 +/- 0.5% (4% +/- 0.7% aa average divergence). The CtRLVaRNA variants reported here show an average divergence of 9.2% +/- 0.3% (5.9% +/- 0.5% at protein level) with those already 217 present in GenBank, while the corresponding value for the carrot BWYVaRNA and the 218

BWYVaRNA/TuYVaRNA isolates already in GenBank is 8.7% +/- 0.3% (6.2% +/-0.5% at
protein level).

Distribution, prevalence and variability of tlaRNAs in wild and cultivated carrot populations 221 In France, CtRLVaRNA was present in all 16 populations sampled in the Nouvelle-Aquitaine 222 region in 2019 (four cultivated, four off-type and eight wild carrot populations). A similarly 223 high prevalence was observed in the 2020 sampling, with CtRLVaRNA present in 10 cultivated 224 populations, 14 wild populations and three off-type populations, while only one cultivated and 225 one wild population (FR20-27 and FR20-22, Table S1) from northern France (Normandy and 226 Hauts-de-France) had no CtRLVaRNA infection. ALVEaRNA was present in 11/16 of the 2019 227 sampled populations including four cultivated, four off-type and three wild populations. Among 228 229 the populations sampled in 2020, eight cultivated, three off-type populations (all located in the 230 Nouvelle-Aquitaine region) and eight wild populations (two of which situated in northern France, in Normandy and Île-de-France, respectively) were infected with ALVEaRNA. Over 231 the two years sampling in France, the new CtRLVaRNA-2 was present in 12 wild populations 232 and only one cultivated population that bordered a wild population in which CtRLVaRNA-2 233 was present (FR20-25 and FR20-26, respectively, Table S1). All CtRLVaRNA-2 infected 234 populations were from the Nouvelle-Aquitaine region. In contrast, the prevalence of 235 BWYVaRNA was low since it was detected only in two off-type populations and one cultivated 236 population from the Nouvelle-Aquitaine region, as well as from two wild populations from 237 Normandy and Hauts-de-France. Overall rates of detection of the various tlaRNAs in French 238 239 carrot populations are given in Table 1.

Populations sampled in Spain showed a similarly high prevalence of CtRLVaRNA, which was
present in all five cultivated populations but not in the wild one. ALVEaRNA was similarly
frequent as it was detected in all six sampled populations. BWYVaRNA and the new
CtRLVaRNA-2 tlaRNA were not detected in any of these six Spanish populations.

11

Plants from two populations [one wild (FR19-9) and one cultivated (FR19-7)] of the 2019 244 sampling in France were analyzed by HTS individually, which revealed a difference in the 245 prevalence of the analysed tlaRNAs. CtRLVaRNA showed a high prevalence in cultivated 246 247 carrots (86% infection, 46/50) but only infected 20% (10/50) of wild ones. ALVEaRNA had a lower prevalence but still showed preferential infection of cultivated plants with infection 248 detected in 28% (14/50) of cultivated plants but only 2% (1/50) in the wild ones. This trend was 249 recently confirmed by PCR screening of single plants in additional populations (D.S. 250 unpublished results). CtRLVaRNA-2 was only identified in a single plant of the wild population 251 while BWYVaRNA had not been identified from the pools of the two individually analysed 252 253 populations and was, as expected, not identified in the corresponding single plants.

In an effort to get a broader insight into the variability of tlaRNA isolates in carrot populations 254 all tlaRNA contigs spanning the region corresponding to positions 1135-1845 of the central 255 256 domain of ORF1b of CtRLVaRNA (AF020617) were retrieved from populations and individual plant assemblies. This region was selected striking a balance between the length of the region 257 (and therefore the amount of phylogenetic information) and the number of isolates for which 258 data could be obtained from HTS contigs. The obtained sequences were used to create a 259 multiple alignment with the corresponding region of GenBank isolates. Figure 3 presents the 260 Neighbor-Joining phylogenetic tree reconstructed from this multiple alignment. The isolates of 261 CtRLVaRNA-2 identified in this study from a highly homogenous ensemble with a low 262 diversity in the analyzed region of 1.6% + 0.7% (0.8% + 0.7% a divergence). Similarly, the 263 identified ALVEaRNA isolates found in both French and Spanish carrot populations are very 264 similar with a diversity of 1.5% +/- 0.7% (0.9% +/- 0.3% aa divergence). The sole exception 265 concerns two French isolates (one from cultivated and one from off-type carrots) that form a 266 third cluster within ALVEaRNA, distinct from both the arracacha South-American isolates and 267 the European carrot isolates. In contrast, the variability between of CtRLVaRNA isolates is 268

significant, with an average nt diversity of 6.3% +/- 0.5% (1.6% +/- 0.7% aa diversity). Indeed, 269 270 the CtRLVaRNA isolates cluster in two ensembles with strong bootstrap support and a highly divergent sigma isolate (KM486093). In turn the two clusters each separate into subclusters, 271 some of which had not been identified from the full-length genomes available in GenBank or 272 reported here (Figure 3). No obvious correlation between the type of the carrot hosts and the 273 phylogeny of CtRLVaRNA or ALVEaRNA isolates could be identified (Supplementary 274 Figures S2 and S3) suggesting that the populations of these tlaRNAs in wild and cultivated 275 276 carrots largely function as single metapopulations.

277 DISCUSSION

CtRLVaRNA and BWYVaRNA were the only tlaRNA detected in carrots so far. In the present 278 study, two additional tlaRNAs were identified in the virome of cultivated and wild carrots. All 279 280 identified tlaRNAs show a typical genome organization for this group of molecules, with a very short 5' UTR, a long 3' UTR and ORF1a and ORF1b separated by an amber stop codon, leading 281 to the production of a ORF1a/ORF1b RdRp fusion protein of ~90kDa. Similar to umbraviruses, 282 tlaRNAs are dependent on helper viruses of the genus Polerovirus for encapsidation and 283 transmission. However, unlike the RdRp of tlaRNAs, which is expressed through a readthrough 284 event, umbraviruses express their RdRp by a -1 frameshift. They also possess two additional 285 ORFs for cell-to-cell and long-distance movement (Liu et al., 2021). The role of umbraviruses 286 in some specific disease complexes is well documented (Taliansky & Robinson, 2003; Murant 287 288 et al., 1969), while less is known about the contribution of tlaRNAs to such disease complexes. 289 The individually analysed plants in this study were frequently coinfected with CtRLVaRNA, CMoV, and/or CMoMV but interestingly, of the 10 individually analysed wild plants showing 290 291 tlaRNAs-umbravirus coinfection, about a third (4/10) did not show evidence of the presence a a helper virus of the Polerovirus or Enamovirus genera. Eventhough, it is not possible to draw 292

a firm conclusion from this observation, tlaRNAs are known to be self-replicating (Campbell et al., 2020), so it could be hypothesized that on occasion they could be transmitted by vectors without a concomitant transmission of the helper virus, in particular if presence of the tlaRNA or of umbravirus(es) had a negative impact on helper virus accumulation (Yoshida, 2020). In such a scenario, there exists the possibility of an assistance for movement of tlaRNAs by coinfecting umbraviruses but, in the absence of encapsidation and of the aphid transmission it permits, this situation could represent an epidemiological dead end.

Currently, there are no molecular criteria established by the International Committee on 300 301 Taxonomy of Viruses (ICTV) for satellites species demarcation. CtRLVaRNA-2 forms a well 302 separated cluster from CtRLVaRNA isolates with on average less than 70% nucleotide identity (average nt divergence $31.1\% \pm 0.7\%$). Its RdRp is similarly divergent, with on average 27.0%303 +/- 1.3% as compared to that of CtRLVaRNA isolates (Figure 3). At both nucleotide and protein 304 levels, these 305 values are close to those separating CABYVaRNA from TuYVaRNA/BWYVaRNA (nt: 33.6% +/- 1.8%; aa: 29.7% +/- 0.8%). Based on these elements, 306 the CtRLVaRNA-2 isolates reported here are proposed to belong to a new, distinct species. 307

Until the present work ALVEaRNA was only known through two partial sequences submitted 308 in GenBank and identified during virome studies of Arracacia xanthorrhiza (Arracacha, family 309 310 Apiaceae) in Peru. ALVEaRNA isolates were predominantly obtained from field-grown carrots accounting for 89.3% (25/28) of the identified isolates (Supplementary Table S4), while the 311 remaining three isolates come from wild carrots (10.7%; 3/28). These isolates form a separate 312 cluster from the partial sequences from Peru (Figure 3) but show a high degree of identity with 313 them (6.8% \pm - 0.5% average nt divergence) and there is little doubt that they should be 314 315 considered as belonging to the same satellite species. Arracacha latent virus E (ALVE, GenBank MF136435, MF073198), the helper virus of ALVEaRNA in Arracacha, is reported 316 in GenBank as an unclassified Luteoviridae, but its RdRp phylogenetic affinities and genomic 317

organization place it in the Enamovirus genus. No evidence for the presence of ALVE was 318 found in any of the individual carrot plants or carrot populations analyzed in the present study, 319 indicating that (an) other virus(es) fullfil the role of helper virus for ALVEaRNA in Europe. 320 321 CtRLV is a strong candidate because the ALVEaRNA-infected plants (15 of the total 100 individual plants examined) were always coinfected with CtRLV and only one of these plants 322 was infected with a possible other helper virus in the form of a novel *Enamovirus*. These 323 observations reinforce the notion that at least some tlaRNAs present a degree of promiscuity 324 325 and can be assisted by different helper viruses. The near identity of tlaRNAs associated with BWYV and TuYV (Figure 2) point in the same direction and so does the partial sequence of a 326 327 tlaRNA associated with potato leafroll virus (MF133518; Campbell et al., 2020) which is 99.5% identical to CtRLVaRNA isolates. Similarly, another tlaRNA identified during the present 328 study is very closely related to BWYVaRNA isolates or TuYVaRNA isolates, with an average 329 330 8.7% nt divergence (6.2% aa divergence). Remarkably, similar to the present finding, TuYVaRNA was detected in British carrots in the absence of BWYV or TuYV but in the 331 presence of CtRLV (Adams et al. 2014), suggesting once again promiscuity of tlaRNAs and an 332 333 ability of TuYVaRNA to be assisted by CtRLV.

CtRLVaRNA was detected in 43 out of 45 populations from four different French regions (Île-334 335 de-France, Normandy, Occitanie and Nouvelle-Aquitaine) and in five out of six carrot populations sampled in central Spain, confirming its wide distribution. At plant population level 336 it did not show preferential detection in one or another carrot type. However, when analyzing 337 single plants, it was much higher in cultivated than in wild carrots (86% vs 20%) a situation 338 that has been confirmed by the PCR testing of individual plants from additional populations 339 340 (not shown). For their part ALVEaRNA and CtRLVaRNA-2 showed preferential association to one type of carrot population. ALVEaRNA was detected more frequently in cultivated carrot 341 populations than in wild ones (79.9% vs 47.8%) and had a higher prevalence in cultivated single 342

plants than in wild one (28% vs 2%). The situation is reversed for CtRLVaRNA-2 which was 343 detected much more frequently in wild (52.2%) than in cultivated (6.7%) populations (Table 344 1). Interestingly, the single cultivated population in which it was detected was bordered by a 345 CtRLVaRNA-2 positive wild population, and in a 500 nt shared region the contigs from the 346 two populations showed only 3 single nucleotide polymorphisms. Taken together these results 347 suggest a flow from the wild carrots to the cultivated ones. CtRLVaRNA-2 also showed a 348 349 limited geographical distribution being detected only in the Nouvelle-Aquitaine region of France. 350

The off-type populations differed phenologically and morphologically from cultivated carrots, 351 352 showing early bolting and not or very poorly tuberized roots, both traits linking them to wild carrots. However, in a number of cases their roots had the orange color typical of cultivated 353 carrots, suggesting they might represent pollen or seed contamination during commercial seed 354 production resulting in hybrid genotypes. Even though their phenotype was evidently closer to 355 that of wild carrots, off-type populations showed tlaRNAs detection frequencies close to those 356 357 of cultivated carrots growing side by side rather than to those of wild carrots (Table 1), suggesting that whatever the underlying mechanism(s), differences in tlaRNAs prevalence are 358 likely more influenced by growth conditions than by plant genotype. 359

The results reported here extended our knowledge of the host range and diversity of tlaRNAs and provide further insights in the variability of associations they are able to establish with their helper viruses, demonstrating the flexibility of such systems of assistance and complementation. With four different tlaRNAs identified, the carrot virome has proven particularly rich in these unusual agents, some of which like CtRLVaRNA and ALVEaRNA appear to show a wide distribution and a high prevalence. Given these elements, it will be of much interest to analyse the specificity of associations and the complex interplay between co367 dependent viruses (poleroviruses, umbraviruses and satellites of various types) in the carrot368 pathosystem and their epidemiological consequences.

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378 Data availability: The genome sequences for the tlaRNAs reported here have been deposited379 in GenBank. The raw sequence datasets are available on request from the authors

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- 450

452 **Table 1:** Rate of detection of the different tlaRNAs in carrot populations sampled in France

Figure 1: Genome organization of CtRLVaRNA-2 and ALVEaRNA. ORF1a of CtRLVaRNA2 and ALVEaRNA codes for a putative protein of 25.8kDa and 27.8 kDa, respectively, and
ends in an amber stop codon (UAG). ORF1b is expressed by readthrough of ORF1a and
encodes the viral RNA-dependent RNA polymerase.

Figure 2: Neighbour-joining phylogenetic tree reconstructed from the alignment of the full 457 genome sequence of tlaRNAs using a strict nucleotide identity distance. Bootstrap values 458 greater than 70% (1,000 repeats) are shown. Novel tlaRNA sequences reported here are 459 indicated by black shapes of different shapes for each tlaRNA. The accession numbers of 460 reference sequences retrieved from GenBank are indicated. The three clusters of tlaRNAs are 461 indicated with different shadings. The scale bar corresponds to 10% nucleotide divergence. 462 463 ALVEaRNA: arrachacha latent virus E-associated RNA; TuYVaRNA: turnip yellows virusassociated RNA; BWYVaRNA: beet western yellows virus-associated RNA; PeVYVaRNA: 464 pepper vein yellows-associated RNA; PoPeYVaRNA: pod pepper yellows virus-associated 465 RNA; TBTDaRNA: tobacco bunchy top disease-associated RNA; TVDVaRNA: tobacco vein 466 distortion virus-associated RNA; CABYVaRNA: cucurbit aphid borne virus-associated RNA; 467 ctRLVaRNA: carrot red leaf virus-associated RNA; ctRLVaRNA-2: carrot red leaf virus-468 associated RNA 2 469

Figure 3: Neighbour-joining phylogenetic tree of tlaRNAs reconstructed from the alignment
of nucleotide sequence of the central part of ORF1b corresponding to positions 1135-1845 of
CtRLVaRNA AF020617. Bootstrap values greater than 70% (1,000 repeats) are shown.
Sequences derived from the novel complete genomes shown in Figure 2 are indicated by black

diamonds, those from reference sequences obtained from GenBank are indicated white
diamonds, together with the relevant accession number. The four tlaRNAs are indicated by
different shadings, and that corresponding to the novel CtRLVaRNA-2 indicated on the right
side. The scale bar corresponds to 10% nucleotide divergence.

478 Supplementary information

Figure S1: multiple alignments of of the readthrough region of tlaRNAs-encoded P1-P2
fusion protein with the GLL motif highlighted

481 Figure S2: Neighbour-joining phylogenetic tree of CtRLVaRNA color coded for the type of
482 carrot population (cultivated, off-type, wild) from which the associated RNA was identified.

483 The tree was reconstructed from the alignment of nucleotide sequence of the central part of

484 ORF1b corresponding to positions 1135-1845 of CtRLVaRNA AF020617. Bootstrap values

485 greater than 70% (1,000 repeats) are shown.

Figure S3: Neighbour-joining phylogenetic tree of ALVEaRNA color coded for the type of
carrot population (cultivated, off-type, wild) from which the associated RNA was identified.
The tree was reconstructed from the alignment of nucleotide sequence of the central part of
ORF1b corresponding to positions 1135-1845 of CtRLVaRNA AF020617. Bootstrap values
greater than 70% (1,000 repeats) are shown.

491 **Table S1**: characteristics of all carrot populations sampled in the present study

492 **Table S2**: specific detection and RACE primers used in the present study

493 **Table S3**: origin, sequence assembly parameters and accession numbers for the full length

494 genomes of tlaRNA isolates reconstructed during the present study

Table S4: origin and accession numbers for the partial ORF1b sequences of tlaRNA isolatesanalyzed in the present study

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498 Figure 1







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tlaRNA	Cultivated carrot	Wild carrots	Off-type ^a carrot
	populations	populations	populations
CtRLVaRNA	14/15 (93.3%)	22/23 (93.3%)	7/7 (100%)
ALVEaRNA	12/15 (79,9%)	11/23 (47,8%)	7/7 (100%)
CtRLVaRNA-2	1/15 (6.7%)	12/23 (52.2%)	0/7 (0%)
BWYVaRNA	1/15 (6.7%)	2/23 (8.7%)	2/7 (28,6 %)

506 Table 1

^acarrot plants that were phenologically and morphologically more similar to wild carrots but

508 grew within cultivated carrot fields