Unveiling Crucivirus Diversity by Mining Metagenomic Data

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33 evolution

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34 ABBREVIATIONS

35 CruV: Crucivirus; CRESS: Circular Rep-Encoding Single Stranded; CP: Capsid; RdRP:

36 RNA-dependent RNA Polymerase; ORF: Open Reading Frame; SJR: Single Jelly Roll;

37 PI: Pairwise Identity; MDA: Multiple Displacement Amplification; S3H: Superfamily 3

38 helicase; RCR: Rolling Circle Replication; CDS: Coding Sequence; CruCGE: Cruci-

39 like Circular Genetic Element

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40 ABSTRACT

41 The discovery of cruciviruses revealed the most explicit example of a common protein 42 homologue between DNA and RNA viruses to date. Cruciviruses are a novel group of 43 circular Rep-encoding ssDNA (CRESS-DNA) viruses that encode capsid proteins (CPs) 44 that are most closely related to those encoded by RNA viruses in the family 45 Tombusviridae. The apparent chimeric nature of the two core proteins encoded by 46 crucivirus genomes suggests horizontal gene transfer of CP genes between DNA and 47 RNA viruses. Here, we identified and characterized 451 new crucivirus genomes and 48 ten CP-encoding circular genetic elements through de novo assembly and mining of 49 metagenomic data. These genomes are highly diverse, as demonstrated by sequence 50 comparisons and phylogenetic analysis of subsets of the protein sequences they encode. 51 Most of the variation is reflected in the replication associated protein (Rep) sequences, 52 and much of the sequence diversity appears to be due to recombination. Our results 53 suggest that recombination tends to occur more frequently among groups of cruciviruses 54 with relatively similar capsid proteins, and that the exchange of Rep protein domains 55 between cruciviruses is rarer than gene exchange. Altogether, we provide a 56 comprehensive and descriptive characterization of cruciviruses.

57 **IMPORTANCE**

58 Viruses are the most abundant biological entities on Earth. In addition to their impact on 59 animal and plant health, viruses have important roles in ecosystem dynamics as well as 60 in the evolution of the biosphere. Circular Rep-encoding single-stranded (CRESS) DNA 61 viruses are ubiquitous in nature, many are agriculturally important, and are viruses that 62 appear to have multiple origins from prokaryotic plasmids. CRESS-DNA viruses such 63 as the cruciviruses, have homologues of capsid proteins (CPs) encoded by RNA viruses. 64 The genetic structure of cruciviruses attests to the transfer of capsid genes between disparate groups of viruses. However, the evolutionary history of cruciviruses is still 65 66 unclear. By collecting and analyzing cruciviral sequence data, we provide a deeper 67 insight into the evolutionary intricacies of cruciviruses. Our results reveal an unexpected 68 diversity of this virus group, with frequent recombination as an important determinant 69 of variability.

70 INTRODUCTION

71 In the last decade, metagenomics has allowed for the study of viruses from a 72 new angle; viruses are not merely agents of disease, but abundant and diverse members 73 of ecosystems (1, 2). Viruses have been shaping the biosphere probably since the origin 74 of life, as they are important drivers of the evolution of the organisms they infect (3–5). 75 However, the origin of viruses is not entirely clear. Viruses, as replicons and mobile 76 elements, are also subject to evolution. Virus variability is driven by various mutation 77 rates, recombination and reassortment of genetic components (6). These attributes, 78 coupled with types of genomes (RNA or DNA, single or double stranded and circular or 79 linear), lead to a large genetic diversity in the 'viral world'.

80 Viruses are generally classified based on the nature of their transmitted genetic 81 material (7). Viral genetic information is coded in either RNA or DNA. Moreover, these 82 genomes can be single (positive or negative sense) or double stranded, linear or circular, 83 and can be comprised of a single or multiple molecules of nucleic acid (monopartite or 84 multipartite, respectively). These different groups of viruses have different replication 85 strategies, and they harbor distinct taxa based on their genome arrangement and 86 composition (1). The striking differences between viral groups with disparate genome 87 types suggest polyphyletic virus origins (8).

88 For example, the highly abundant circular Rep-encoding single-stranded DNA 89 (CRESS-DNA) viruses may have been derived from plasmids on multiple occasions by 90 acquiring capsid genes from RNA viruses (9–11). Eukaryotic CRESS-DNA viruses 91 constitute a diverse and widespread group of viruses with circular genomes -some of 92 them multipartite- that contains the families Geminiviridae, Circoviridae, Nanoviridae, 93 Alphasatellitidae, Genomoviridae, Bacilladnaviridae. Smacoviridae and 94 Redondoviridae (ICTV classification for some groups is pending at this time), in 95 addition to vast numbers of unclassified viruses (12, 13). Universal to all CRESS-DNA 96 viruses is the Rep, which is involved in the initiation of the virus' rolling-circle 97 replication. Rep homologues are also encoded in plasmids (13, 14). Some pathogenic 98 CRESS-DNA viruses are agriculturally important, such as porcine circoviruses, and 99 nanoviruses and geminiviruses that infect a wide range of plant hosts (12). However, 100 many CRESS-DNA viruses have been identified in apparently healthy organisms, and 101 metagenomics have revealed their presence in most environments (12).

102 In 2012, a metagenomic survey of a hot and acidic lake in the volcanic Cascade 103 Range of the western USA uncovered a new type of circular DNA virus (15). The genome of this virus is a CRESS-DNA virus based on the circularity of its sequence, the 104 105 presence of a *rep* gene, and a predicted stem-loop structure with a conserved nucleotide 106 sequence (ori) that serves as an origin for CRESS-DNA virus rolling-circle replication 107 (RCR; reviewed in (16, 17)). Interestingly, the sequence of CP encoded by this genome 108 resembles those encoded by the RNA viruses in the family Tombusviridae (15). It was 109 hypothesized that this virus originated by the acquisition of a capsid gene from an RNA 110 virus through a yet to be demonstrated RNA-DNA recombination event (15, 18). Since 111 the discovery of this putatively "chimeric virus", 80 circular sequences encoding a Rep 112 and a CP that share homology to tombusvirus CPs have been found in different 113 environments around the globe (19, 20, 29–31, 21–28). This growing group of viruses 114 have been branded "cruciviruses", as they imply the *crossing* between CRESS-DNA 115 viruses and RNA tombusviruses (27). Cruciviruses have been found associated with 116 forams (20), alveolates hosted by isopods (26), arthropods (19, 22), and in peatland 117 ecosystems (27), but no host for cruciviruses have been elucidated to date.

- 118 The circular genome of known cruciviruses is variable in size, ranging from 2.7 119 to 5.7 kb and often contains ORFs in addition to the Rep and CP, which have been 120 found in either a unisense or an ambisense orientation (20, 27). The function of 121 additional crucivirus ORFs is unclear due to the lack of sequence similarity with any 122 characterized protein. The genome replication of CRESS-DNA viruses is initiated by 123 the Rep protein, that binds to direct repeats present just downstream of the stem of the 124 ori-containing stem-loop structure and nicks the ssDNA (32, 33). The exposed 3'OH 125 serves as a primer for cellular enzymes to replicate the viral genome via RCR (33–35). 126 The exact terminating events of CRESS-DNA virus replication are poorly understood 127 for most CRESS-DNA viruses, but Rep is known to be involved in the sealing of newly 128 replicated genomes (33, 35-37).
- Rep has a domain in the N-terminus that belongs to the HUH endonuclease superfamily (38). This family of proteins is characterized by a HUH motif (Motif II), in which two histidine residues are separated by a bulky hydrophobic amino acid, and a Tyr-containing motif (Motif III) that catalyzes the nicking of the ssDNA (38–41). CRESS-DNA virus Reps also contain a third conserved motif in the N-terminal portion of the protein (Motif I), likely responsible for dsDNA binding specificity (42). In many

135 CRESS-DNA viruses, the HUH motif has been substituted for a similar motif that lacks 136 the second histidine residue (e.g. circoviruses have replaced HUH with HLQ) (10, 38). 137 The C-terminal portion of eukaryotic CRESS-DNA virus Reps contain a Superfamily 3 138 helicase domain (S3H) that may be responsible for unwinding dsDNA replicative 139 intermediates (43, 44). This helicase domain is characterized by Walker A and B motifs, 140 Motif C and an Arg finger. Previous studies have identified evidence of recombination 141 in the endonuclease and helicase domains of Rep, which contributes to the potential 142 ambiguity of Rep phylogenies (45). Interestingly, the Rep proteins of different 143 cruciviruses have been shown to be similar to CRESS-DNA viruses in different 144 families, including circoviruses, nanoviruses, and geminiviruses (20, 27). In some cruciviruses, these differences in phylogeny have been observed between the individual 145 146 domains of a single Rep protein (21, 27). The apparent polyphyly of crucivirus Reps suggests recombination events involving cruciviruses and other CRESS-DNA viruses, 147 148 even within Reps (20, 21).

149 All characterized CRESS-DNA viruses package their DNA into small capsids 150 with icosahedral symmetry or their geminate variants, built from multiple copies of the 151 CP encoded in their genome (12). The CP of these CRESS-DNA viruses appears to fold 152 into an eight-strand ß-barrel that conforms to the single jelly-roll (SJR) architecture, 153 which is also commonly found in eukaryotic RNA viruses (46). The CP of cruciviruses 154 has no detectable sequence similarity with the capsid of other CRESS-DNA viruses, and 155 is predicted to adopt the SJR conformation found in the CP of tombusviruses (15, 20, 156 21). Three domains can be distinguished in tombusviral CPs (47, 48). From the N- to 157 the C-terminus: i) The RNA-interacting or R-domain, a disordered region that faces the interior of the viral particle to interact with the nucleic acid through abundant basic 158 159 residues (49, 50), ii) the shell or S-domain containing the single jelly-roll fold and the 160 architectural base of the capsid (48) and iii) the protruding or P-domain, that decorates 161 the surface of the virion and is involved in host transmission (51). In tombusviruses, the S-domain of 180 CP subunits interact with each other to assemble around the viral RNA 162 163 in a T=3 fashion, forming a \emptyset ~35 nm virion (48, 52).

The study of cruciviruses suggests evidence for the transfer of capsid genes between disparate viral groups, which can shed light on virus origins and the phenotypic plasticity of virus capsids. Here, we document the discovery of 461 new crucivirus genomes (CruV) and cruci-like circular genetic elements (CruCGE) identified in

168 metagenomic data obtained from different environments and organisms. This study 169 provides a comprehensive analysis of this greatly expanded dataset and explores the

- extent of cruciviral diversity –mostly due to Rep heterogeneity– impacted by rampant
- 171 recombination.

172 MATERIALS AND METHODS

173 **Recovery of viral genomes from assembled viromes.**

174 A total of 461 crucivirus-related sequences were identified from 1168 metagenomic surveys (Supp. Tables 1 and 2). 1167 viromes from 57 published datasets 175 176 and one unpublished virome were obtained from different types of environments: i) 177 aquatic systems (freshwater, seawater, hypersaline ponds, thermal springs and 178 hydrothermal vents), ii) engineered systems (bioreactor, food production), and iii) 179 eukaryote-associated flora (human, insect and other animal feces, human saliva and 180 fluids, cnidarians and plants). New cruciviral sequences were identified in these viromes 181 by screening circular contigs for the presence of CPs from previously known 182 cruciviruses (20) and tombusviruses, using a BLASTx bit-score threshold of 50.

Additionally, sequences CruV-240, CruV-300, CruV-331, CruV-338 and CruV-367 were retrieved from Joint Genome Institute (JGI)'s IMG/VR repository (53), by searching scaffolds with a function set including the protein family pfam00729, corresponding to the S-domain of tombusvirus capsids. The sequences with an RdRP coding region were excluded, and the circularity of the sequences, as well as the presence of an ORF encoding a tombusvirus-like capsid, were confirmed with Geneious 11.0.4 (Biomatters, Ltd).

190 Annotation of crucivirus putative genes.

191 The 461 cruciviral sequences were annotated and analyzed in Geneious 11.0.4. 192 Coding sequences (CDSs) were semi-automatically annotated from a custom database 193 (Supp. Table 3) of protein sequences of published cruciviruses and close homologs 194 obtained from GenBank, using Geneious 11.0.4's annotation function with a 25% 195 nucleotide similarity threshold. Annotated CDS were re-checked with GenBank 196 database using BLASTx to identify sequences similar to previously described 197 cruciviruses and putative relatives. Sequences containing in-frame stop codons were 198 checked for putative splicing sites (54), or translated using a ciliate genetic code only 199 when usage rendered a complete ORF with similarity to other putative crucivirus CDSs. 200 Predicted ORFs longer than 300 bases with no obvious homologs and no overlap with 201 CP or Rep-like ORFs were annotated as "putative ORFs".

202 **Putative Stem-loop annotation.**

203 Stem-loop structures that could serve as an origin of replication for circular 204 ssDNA viruses were identified and annotated using StemLoop-Finder (Pratt et al., 205 unpublished, (55, 56)). The 461 cruciviral sequences were scanned for the presence of 206 conserved nonanucleotide motifs described for other CRESS-DNA viruses 207 (NANTANTAN, NAKWRTTAC, TAWWDHWAN, & TRAKATTRC) (12). The 208 integrated ViennaRNA 2.0 library was used to predict secondary structures of DNA 209 around the detected motif, including the surrounding 15-20 nucleotides on either side 210 (57, 58). Predicted structures with a stem longer than four base pairs and a loop 211 including seven or more bases were subjected to the default scoring system, which 212 increases the score by one point for each deviation from ideal stem lengths of 11 base 213 pairs and loop lengths of 11 nucleotides. A set of annotations for stem-loops and 214 nonanucleotides was created with StemLoop-Finder for those with a score of 15 or 215 below. Putative stem-loops were excluded from annotation when a separate stem-loop was found with the same first base, but attained a greater score, as well as those that 216 217 appeared to have a nonanucleotide within four bases of its stem-loop structure's first or last nucleotide. 218

219 Conservation analysis and visualization

Pairwise identity matrices. The pairwise identity (PI) between the protein sequence
 from translated cruciviral genes was calculated with SDTv1.2 (59), with MAFFT
 alignment option for CPs and S-domains, and MUSCLE alignment options for Reps.

- Sequence conservation annotation. CP sequence conservation represented in Fig. 2A
 was generated with Jalview v2.11.0 (60), and reflects the conservation of the
 physicochemical properties for each column of the alignment (61).
- Sequence logos. Sequence logos showing frequency of bases in nonanucleotides at the
 origin of replication or residue in conserved Rep motifs were made using the weblogo
 server (http://weblogo.threeplusone.com/; (62)).
- Structural representation of capsid conservation. The 3D structure of CP was modeled
 with Phyre² (63). The generated graphic was colored by sequence conservation with
 Chimera v.1.13 (64), from the alignment of the 47 capsid sequences from each of the
 CP-based clusters (Fig. 3B).

233 Phylogenetic analyses.

234 Multiple sequence alignments. CP sequences were aligned using MAFFT (65) in 235 Geneious 11.0.4, with a G-INS-i algorithm and BLOSUM 45 as exchange matrix, with 236 an open gap penalty of 1.53 and an offset value of 0.123, and manually curated. Rep 237 protein sequences were aligned using PSI-Coffee [http://tcoffee.crg.cat/; (66)]. Rep 238 alignments were manually inspected and corrected in Geneious 11.0.4, and trimmed 239 using TrimAI v1.3 with a strict plus setting (67). To produce individual alignments of 240 the endonuclease and helicase domains the full length trimmed alignments were split at 241 the Walker A motif (45).

Phylogenetic trees. Phylogenetic trees containing the entire dataset of cruciviral
sequences were built on Geneious using the FastTree plugin (68). For the analysis of
sequence subsets, trees were inferred with PhyML 3.0 web server [http://www.atgcmontpellier.fr/phyml/; (69)], using an aLRT SH-like support (70). The substitution
model for each analysis was automatically selected by the program.

Intergene and interdomain comparison. Tanglegrams were made using Dendroscope
v3.5.10 (71) to compare the phylogenies between different genes or domains within the
same set of crucivirus genomes.

Sequence similarity networks. A total of 540 CP amino acid sequences, and 600 Rep
amino acid sequences were uploaded to EFI–EST web server for the calculation of PIs
[https://efi.igb.illinois.edu/efi-est/; (72)]. A specific alignment score cutoff was
established for each dataset analyzed, and xgmml files generated by EFI-EST were
visualized and edited in Cytoscape v3.7.2 (73).

- 255 Accession numbers
- 256 *Pending*

257 **RESULTS & DISCUSSION**

258 Expansion of the crucivirus group.

259 To broaden our understanding of the diversity and relationships of cruciviruses, 260 461 uncharacterized circular DNA sequences containing predicted CDSs with sequence 261 similarity to the CP of tombusviruses were compiled from metagenomic sequencing 262 data. The data came from published and unpublished metagenomic studies, carried out 263 in a wide variety of environments, from permafrost to temperate lakes, and on various 264 organisms from red algae to invertebrates (Metagenomes and their metadata are 265 provided in Supp. Table 2). The selected genomes are assumed to be complete and 266 circular based on the terminal redundancy identified in *de novo* assembled genomes.

The cruciviral sequences were named sequentially, beginning with the smallest genome, which was named CruV-81 to account for the 80 crucivirus genomes reported in prior literature (15, 19, 28–31, 20–27). The average GC content of the newly described cruciviral sequences is 42.9 ± 4.9 % (Fig. 1B) with genome lengths spanning from 2,474 to 7,947 bases (Fig. 1A), some exceeding the size of described bacilladnaviruses ($\leq 6,000$ nt (74)), the largest CRESS-DNA viruses known (12).

Of the 461 sequences that contain a CP ORF, 451 have putative coding regions with sequence similarity to Rep of CRESS-DNA viruses (10). The CP and Rep ORFs are encoded in a unisense orientation in 40% of the genomes and an ambisense orientation in 58% of the genomes. The remaining ~2% correspond to ten CruCGEs with no clear Rep CDS. Five of these CruCGEs contain a predicted origin of rollingcircle replication (RCR) (Supp. Table 1), indicating that they are circular genomes that undergo RCR characteristic of other CRESS-DNA virus genomes (16, 17).

280 One possible reason for the lack of a Rep ORF in certain sequences is that some 281 of these may be sub-genomic molecules or possible components of multipartite viruses 282 (75). Some CRESS-DNA viruses, such as geminiviruses and nanoviruses, have 283 multipartite genomes (76). Moreover, some ssRNA tombunodaviruses; including 284 *Plasmopara halstedii* virus A and *Schlerophthora macrospora* virus A – viruses that 285 contain the most similar capsid sequences to cruciviral capsids (15, 27)- also have 286 multipartite genomes (77). Unfortunately, no reliable method yet exists to match 287 different sequences belonging to the same multisegmented virus in metagenomes,

288 making identification of multipartite or segmented viruses from metagenomic data289 challenging (76).

290 Stem-loop structures with conserved nonanucleotide motifs as putative origins 291 of replication were predicted and annotated in 277 cruciviral sequences with StemLoop-292 Finder (Pratt et al., unpublished). In some cases, more than one nonanucleotide motif 293 with similar scores were found for a single genome, resulting in more than one stem-294 loop annotation. Of the annotated genomes, 223 contain a stem-loop with a 295 nonanucleotide with a NANTANTAN pattern, with the most common sequence being 296 the canonical circovirus motif TAGTATTAC, found in 64 of the genomes (Supp. Table 297 1; (78)). The majority of the 54 sequences that do not correspond to NANTANTAN 298 contain a TAWWDHWAN nonanucleotide motif, typical of genomoviruses (79). The 299 frequency of bases at each position in the nonanucleotide sequence is given in Fig. 1C, 300 and reflects similarity to motifs found in other CRESS-DNA viruses (10).

301 Crucivirus capsid protein (CP)

302 The CP of cruciviruses is predicted to have a single jelly-roll (SJR) architecture, 303 based on its homology to tombusvirus CPs for which 3D structures have been 304 determined [Fig. 2A; (80-82)]. The SJR conformation is found in CPs of both RNA and 305 DNA viruses (46). The SJR CP of tombusviruses and cruciviruses contains three 306 distinct domains: the RNA-binding or R-domain, the shell or S-domain, and the 307 protruding or P-domain (Fig. 2A). All 461 crucivirus CPs analyzed in this study contain 308 a complete S-domain. This domain contains a distinct jelly-roll fold and interacts with 309 the S-domain of other capsid subunits in the virion of related tombusviruses (48). The 310 S-domain has greater sequence conservation than the remaining regions of the CP (Fig. 311 2A), likely due to its functional importance in capsid structure. In tombusviruses, the S-312 domain contains a calcium binding motif (DxDxxD), which was not identified in 313 previously described cruciviruses (83). However, we detected this Ca-binding motif in 314 68 CPs of the newly identified cruciviral sequences. These crucivirus sequences form a 315 distinct cluster, shown in red in Fig. 3B. The S-domain is flanked on the N-terminus by 316 the R-domain, which in cruciviruses appears variable in size (up to 320 amino acids 317 long), and appears to be truncated in some of the CP sequences (e.g. CruV-386 and 318 CruV-493). The R-domain is characterized by an abundance of basic residues at the N-

319 terminus, followed by a Gly-rich tract (Fig. 2A). The P-domain, on the C-terminal end 320 of the CP sequence, is generally the largest domain, with the exception of CruV-385, 321 where it appears to be truncated. The conservation of CP suggests a similar structure for 322 all cruciviruses. However, those cruciviruses with larger genomes may assemble their 323 capsids in a different arrangement to accommodate their genome. While the capsids of 324 tombusviruses have been shown to adopt a T=1 icosahedral conformation, rather than 325 the usual T=3, when the R-domain is partially or totally removed (82), we have not seen 326 a correlation between the length of CP domains and genome size in our dataset that 327 could be indicative of alternative capsid arrangements. Furthermore, no packaging 328 dynamics relating genome size and virion T-number arrangement have been determined 329 in CRESS-DNA viruses, although sub-genomic elements of geminiviruses can be 330 packaged in non-geminate capsids (84, 85).

331 Interestingly, CruV-420 contains not one tombusvirus-related CP, but two. A 332 recent compilation of CRESS-DNA viruses from animal metagenomes also contains 333 four genomes with two different CPs in their capsid (31). Whether these viruses use two 334 different CPs in their capsid (as some RNA viruses do), or whether these are intermediates in the exchange of CP genes, as predicted from the gene capture 335 336 mechanism proposed by Stedman (2013) (18), is unclear. If the latter is true, CP gene 337 acquisition by CRESS-DNA viruses may be much more common than previously 338 thought.

339 Crucivirus Rep

340 The Reps of CRESS-DNA viruses typically contain an endonuclease domain 341 characterized by conserved motifs I, II and III, and a helicase domain with Walker A 342 and B motifs, motif C, and an Arg-finger [Fig. 2B; (12)]. The majority (85.9%) of the 343 crucivirus genomes described in this dataset contain all of the expected Rep motifs 344 (Supp. Table 4). However, five genomes (CruCGE-110, CruCGE-296, CruCGE-436, 345 CruCGE-471 and CruCGE-533) with overall sequence homology to other Reps (35.8, 32.7, 49.7, 60.2 and 57.2 % PI with other putative Reps in the databases, respectively), 346 347 lack any detectable conserved motifs within their sequence. Thus, these sequences are 348 considered CP-encoding cruci-like circular genetic elements (CruCGEs).

The endonuclease catalytic domain of Rep (motif II), including HUH, was identified in 441 of the genomes of which 95.2% had an alternative HUH, with the most common arrangement being HUQ (70.0%), also found in circoviruses and nanoviruses [(10, 25, 39); Fig. 2B]. 26.2% of the crucivirus motif II deviate from the HUH motif by additionally replacing the second hydrophobic residue (U) with a polar amino acid (Fig. 2B; supp. Table 4), with 53 of Reps with the sequence HYQ (12.0%), also found in smacoviruses (10, 23, 45).

356 We identified thirteen putative Reps in these crucivirus genomes that lack all four motifs typically found in S3H helicases (e.g. CruV-166, CruV-202, CruV-499; 357 358 Supp. Table 4). Recent work has shown that the deletion of individual conserved motifs 359 in the helicase domain of the Rep protein of beak and feather disease virus does not 360 abolish ATPase and GTPase activity (86). The absence of all four motifs may prevent 361 these putative Reps from performing helicase and ATPase activity using previously 362 characterized mechanisms. However, it is possible that crucivirus Reps that lack these motifs are still capable of ATP hydrolysis and associated helicase activity. 363 Alternatively, these activities may be provided by host factors (87), or by a viral 364 replication-enhancer protein – as is the case with the AC3 protein of begomoviruses 365 (88). 366

367 We identified 36 crucivirus genomes whose putative rep genes contain in-frame stop codons or the HUH and SF3 helicase are in different frames, suggesting that their 368 369 transcripts may require intron splicing prior to translation. Acceptor and donor splicing 370 sites identical to those found in maize streak virus (54) were found in all these 371 sequences, and the putatively spliced Reps annotated accordingly. In five of the 36 372 spliced Reps, we were unable to detect any of the four conserved motifs associated with 373 helicase/ATPase activity, which are encoded in the predicted second exon in most cases. 374 CruV-513 and CruV-518 also contain predicted splicing sites in their CP gene.

No GRS motifs –which have been identified as necessary for geminivirus replication (89), and have also been found in genomoviruses (90)– were detected in Reps in our dataset. We were unable to detect any conserved Rep motifs present in cruciviruses that are absent in other CRESS-DNA viruses. Given the conservation of Rep motifs in these newly-described cruciviruses, we expect most to be active in RCR.

380 Crucivirus CPs share higher genetic identity than their Rep proteins

To assess the diversity in the proteins of cruciviruses, the percentage pairwise identity (% PI) between the protein sequences was calculated for CP and Rep using SDTv1.2 (Fig. 3). The average % PI for CP was found to be 33.1 ± 4.9 % PI (Figs. 3A and 3D), likely due to the high levels of conservation found in the S-domain (40.5 ± 8.4 % PI; Figs. 3B and 3D), while the average % PI for Rep is quite low at 24.7% (± 5.6 SD; Figs. 3C and 3D). The high variation of the Rep protein sequence relative to CP in cruciviruses correlates with a previous observation on a smaller dataset (20).

388 To compare cruciviruses to other viral groups with homologous proteins, 389 sequence similarity networks were built for CP and Rep (Fig. 4). For the CP, related 390 protein sequences from tombusviruses and unclassified RNA viruses were included. 391 The virus sequences were connected when the similarity between their protein sequence 392 had an e-value < 1e-20, sufficient to connect all cruciviruses and tombusviruses, with 393 the exception of CruV-523 (Fig. 4A). However, using BLASTp, CruV-523 showed 394 similarity to other RNA viruses with an e-value < 1e-9, which were not included in the 395 analysis. The CP sequence similarity network analysis demonstrates the apparent 396 homology of the CPs in our dataset with the CP of RNA viruses; specifically to 397 unclassified RNA viruses that have RdRPs similar to either tombusviruses -also 398 described as tombus-like viruses (77, 91, 92)- or to nodaviruses. The latter RNA viruses 399 are proposed to belong to a chimeric group of viruses named tombunodaviruses (93).

400 For sequence similarity network analysis of Rep, sequences from CRESS-DNA 401 viruses belonging to the families Circoviridae, Nanoviridae, Alphasatellitidae, 402 Geminiviridae, Genomoviridae, Smacoviridae and Bacilladnaviridae were used (Fig. 403 4B). Due to the heterogeneity of Rep (Fig. 3C), the score cutoff for the network was 404 relaxed to an e-value < 1e-10; nonetheless, ten divergent sequences lacked sufficient 405 similarity to form connections within the network. While the Rep of the different viral 406 families clustered in specific regions of the network, the similarity of cruciviral Reps 407 spans the diversity of all CRESS-DNA viruses, and blurs the borders between them. 408 Though there are cruciviruses that appear to be closely related to geminiviruses and 409 genomoviruses, these connections are less common than with other classified CRESS-410 DNA families (Fig. 4B). While still highly divergent from each other, the conserved 411 motifs in the Rep still share the most sequence similarity with CRESS-DNA viruses 412 (Fig. 2B).

413 The broad sequence space distribution of cruciviral Rep sequences has been 414 proposed to reflect multiple Rep acquisition events through recombination with viruses 415 from different CRESS-DNA viral families (20). However, the apparent larger diversity 416 of cruciviral Reps relative to classified CRESS-DNA viruses can be due to the method 417 of study, as most classified CRESS-DNA viruses have been discovered from infected 418 organisms and are grouped mainly based on Rep similarity (1). By contrast, here 419 crucivirus sequences are selected according to the presence of a tombusvirus-like CP. 420 Moreover, the Rep of cruciviruses could be subject to higher substitution rates than CP 421 (26). It is possible that sequence divergence in CP is more limited than in the Rep due to 422 structural constraints.

423 Horizontal gene transfer among cruciviruses.

To gain insight into the evolutionary history of cruciviruses, we carried out phylogenetic analyses of their CPs and Reps. Due to the high sequence diversity in the dataset, two smaller subsets of sequences were analyzed:

427 i) CP-based clusters. Clusters with more than six non-identical CP sequences whose S-428 domains share a % PI greater than 70% were identified from Fig. 3B. This resulted in 429 the identification of seven clusters, and a more divergent, yet clearly distinct, cluster 430 was included (pink in Fig. 3B). A total of 47 genomes from the eight different clusters 431 were selected for sequence comparison. The protein sequences of CP and Rep were 432 extracted, aligned, and their phylogenies inferred and analyzed using tanglegrams (Fig. 433 5A). The CP phylogeny shows that the eight CP-based clusters form separate clades 434 (Fig. 5A). On the other hand, the phylogeny of Rep shows a different pattern of 435 relatedness between those genomes (Fig. 5A). This suggests different evolutionary 436 histories for the CP and Rep proteins, which could be due to recombination events 437 between cruciviruses, as previously proposed with smaller datasets (20, 21).

ii) *Rep-based clusters*. To account for the possible bias introduced by selecting genomes
from CP cluster groups and to increase the resolution in the phylogeny of the Rep
sequences, clusters with more than six Rep sequences sharing PI > 45 and < 98% were
identified. A total of 53 genomes from six clusters (Fig. 3C) were selected, and their CP
and Rep protein sequences analyzed. The phylogeny of Reps shows distinct clades
between the sequences from different clusters (Fig. 5B). When the phylogeny of Rep

444 was compared to that of their corresponding CPs, we observed the presence of groups of 445 cruciviruses that clade with each other in both the Rep and the CP. Discrepancies in 446 topology between Rep and CP were observed as well, particularly in the CP clade 447 marked with an asterisk in Fig. 5B. This clade corresponds to the highly-homogeneous 448 red CP-based cluster shown in Fig. 3B, and suggests that gene transfer is more common 449 in cruciviruses with a more similar CP, likely infecting the same type of organism. On 450 the other hand, the presence of cruciviral groups with no trace of genetic exchange may 451 indicate that lineages within the cruciviral group may have undergone speciation in the 452 course of evolution.

453 To investigate possible exchanges of individual Rep domains among 454 cruciviruses, the Rep alignments of the analyses of the CP-based and Rep-based clusters 455 were split at the beginning of the Walker A motif to separate endonuclease and helicase 456 domains. From the analysis of the CP-based clusters, we observed incongruence in the 457 phylogenies between endonuclease and helicase domains (Fig. 6A), suggesting 458 recombination within crucivirus Reps, as has been previously hypothesized with a much 459 smaller dataset (21). This incongruency is not observed in the analyzed Rep-based 460 clusters (Fig. 6B). This is likely due to the higher similarity between Reps in this subset 461 of sequences, biased by the clustering based on Rep. We do observe different topologies 462 between the trees, which may be a consequence of different evolutionary constraints to 463 which the endonuclease and helicase domains are subject. The detection of CP/Rep 464 exchange and not of individual Rep domains in Rep-based clusters suggests that the rate 465 of intergenic recombination is higher than intragenic recombination in cruciviruses.

466 Members of the SAR supergroup are potential crucivirus hosts

467 While no crucivirus host has been identified to date, the architecture of the Rep 468 protein found in most cruciviruses suggests a eukaryotic host. The fusion of an 469 endonuclease domain to a S3H helicase domain is observed in other CRESS-DNA 470 viruses which are known to infect eukaryotes (38). This is distinct from Reps found in prokaryote-infecting CRESS-DNA viruses -which lack a fused S3H helicase domains 471 472 (94)- and other related HUH endonucleases involved in plasmid RCR and HUH 473 transposases (38). Additionally, the CP of cruciviruses, a suggested determinant of 474 tropism (95, 96), is homologous to the capsid of RNA viruses known to infect

475 eukaryotes. The RNA viruses with a known host with capsids most similar to cruciviral
476 capsids (tombunodaviruses) infect oomycetes, a group of filamentous eukaryotic
477 stramenopiles (77).

478 Cruciviruses have been found as contaminants of spin columns made of 479 diatomaceous silica (21), in aquatic metagenomes enriched with unicellular algae (20), 480 in the metagenome of Astrammina rara -a foraminiferan protist part of the rhizaria-(20), and associated with epibionts of isopods, mainly comprised of apicomplexans and 481 482 ciliates, both belonging to the alveolates (26). These pieces of evidence point toward the 483 stramenopiles/alveolates/rhizaria (SAR) supergroup as a candidate taxon to contain 484 potential crucivirus hosts (97). No host prediction can be articulated from our sequence 485 data. However, at least five of the crucivirus genomes only render complete translated 486 CP and Rep sequences when using a relaxed genetic code. Such alternative genetic 487 codes have been detected in ciliates, in which the hypothetical termination codons UAA and UAG encode for a glutamine (98). The usage of an alternative genetic code seems 488 evident in CruV-502 -found in the metagenome from seawater collected above diseased 489 490 coral colonies (99)- that uses a UAA codon for a glutamine of the S-domain conserved 491 in 33.5% of the sequences. While the data accumulated suggest unicellular eukaryotes 492 and SAR members as crucivirus-associated organisms, the host of cruciviruses remains 493 elusive, and further investigations are necessary.

494 Classification of cruciviruses

495 Cruciviruses have circular genomes that encode a Rep protein probably involved 496 in RCR. The single-stranded nature of packaged crucivirus genomes has not been 497 demonstrated experimentally; however, the overall genomic structure and sequence 498 similarity underpins the placement of cruciviruses within the CRESS-DNA viruses.

The classification of the CRESS-DNA viruses is primarily based upon the phylogeny of the Rep proteins, although commonalities in CP and genome organization are also considered (13). This taxonomic criteria is challenging in cruciviruses, whose Rep proteins are highly diverse and apparently paralogous. Whether the use of proteins involved in replication for virus classification should be preferred over structural proteins has been previously questioned (100).

505 The capsid of cruciviruses, as well as the capsid of other CRESS-DNA virus 506 families like circoviruses, geminiviruses and bacilladnaviruses, possess the single-jelly 507 roll architecture (46). However, there is no obvious sequence similarity between the CP 508 of cruciviruses and that of classified CRESS-DNA viruses. The crucivirus CP -509 homologous to the capsid of tombusviruses- is an orthologous trait within the CRESS-510 DNA viruses. Hence, CP constitutes a synapomorphic character that demarcate this 511 group of viruses from the rest of the CRESS-DNA viral families. Thus, the 512 classification of cruciviruses is challenging.

513 CONCLUDING REMARKS

514 Cruciviruses are a growing group of CRESS-DNA viruses that encode CPs that 515 are homologous to those encoded by tombusviruses. Over 500 crucivirus genomes have 516 been recovered from various environments across the globe. These genomes vary in 517 size, sequence and genome organization. While crucivirus CPs are relatively 518 homogeneous, the Reps are relatively diverse amongst the cruciviruses, spanning the 519 diversity of all classified CRESS-DNA viruses. It has been hypothesized that 520 cruciviruses emerged from the recombination between a CRESS-DNA virus and a 521 tombus-like RNA virus (15, 18). Furthermore, cruciviruses seem to have recombined 522 with each other to exchange functional modules between them, and probably with other 523 viral groups, which blurs their evolutionary history. Cruciviruses show evidence of 524 genetic transfer, not just between viruses with similar genomic properties, but also 525 between disparate groups of viruses such as CRESS-DNA and RNA viruses.

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540 **BIBLIOGRAPHY**

- Simmonds P, Adams MJ, Benk M, Breitbart M, Brister JR, Carstens EB, Davison AJ, Delwart E, Gorbalenya AE, Harrach B, Hull R, King AMQ, Koonin E V., Krupovic M, Kuhn JH, Lefkowitz EJ, Nibert ML, Orton R, Roossinck MJ, Sabanadzovic S, Sullivan MB, Suttle CA, Tesh RB, Van Der Vlugt RA, Varsani A, Zerbini FM. 2017. Consensus statement: Virus taxonomy in the age of metagenomics. Nat Rev Microbiol 15:161–168.
- 546 2. Chow C-ET, Suttle CA. 2015. Biogeography of Viruses in the Sea. Annu Rev Virol 2:41–66.
- 5473.Koonin E V., Dolja V V. 2013. A virocentric perspective on the evolution of life. Curr Opin Virol5483:546–557.
- 4. Koonin E V., Krupovic M. 2018. The depths of virus exaptation. Curr Opin Virol 31:1–8.
- 550 5. Berliner AJ, Mochizuki T, Stedman KM. 2018. Astrovirology: Viruses at Large in the Universe.
 551 Astrobiology 18:207–223.
- 5526.Domingo E, Sheldon J, Perales C. 2012. Viral Quasispecies Evolution. Microbiol Mol Biol Rev55376:159–216.
- 554 7. Baltimore D. 1971. Expression of animal virus genomes. Bacteriol Rev 35:235–241.
- 555 8. Koonin E V., Senkevich TG, Dolja V V. 2006. The ancient virus world and evolution of cells.
 556 Biol Direct 1.
- 557 9. Krupovic M, Ravantti JJ, Bamford DH. 2009. Geminiviruses: A tale of a plasmid becoming a virus. BMC Evol Biol 9:1–11.
- 55910.Kazlauskas D, Varsani A, Koonin E V., Krupovic M. 2019. Multiple origins of prokaryotic and
eukaryotic single-stranded DNA viruses from bacterial and archaeal plasmids. Nat Commun
10:1–12.56110:1–12.
- 56211.Krupovic M. 2012. Recombination between RNA viruses and plasmids might have played a
central role in the origin and evolution of small DNA viruses. BioEssays 34:867–870.
- 56412.Zhao L, Rosario K, Breitbart M, Duffy S. 2019. Eukaryotic Circular Rep-Encoding Single-565Stranded DNA (CRESS DNA) Viruses: Ubiquitous Viruses With Small Genomes and a Diverse566Host Range. Adv Virus Res, 1st ed. 103:71–133.
- 567 13. Rosario K, Duffy S, Breitbart M. 2012. A field guide to eukaryotic circular single-stranded DNA viruses: Insights gained from metagenomics. Arch Virol 157:1851–1871.
- 56914.Cheung AK. 2015. Specific functions of the Rep and Rep' proteins of porcine circovirus during570copy-release and rolling-circle DNA replication. Virology 481:43–50.
- 57115.Diemer GS, Stedman KM. 2012. A novel virus genome discovered in an extreme environment572suggests recombination between unrelated groups of RNA and DNA viruses. Biol Direct 7:13.
- 573 16. Cheung AK. 2012. Porcine circovirus: Transcription and DNA replication. Virus Res 164:46–53.
- 574 17. LAUFS J. 1995. Geminivirus replication: Genetic and biochemical characterization of rep protein
 575 function, a review. Biochimie 77:765–773.
- 57618.Stedman K. 2013. Mechanisms for RNA capture by ssDNA viruses: Grand theft RNA. J Mol577Evol 76:359–364.

- 19. Rosario K, Dayaram A, Marinov M, Ware J, Kraberger S, Stainton D, Breitbart M, Varsani A.
 2012. Diverse circular ssDNA viruses discovered in dragonflies (Odonata: Epiprocta). J Gen Virol 93:2668–2681.
- 58120.Roux S, Enault F, Bronner G, Vaulot D, Forterre P, Krupovic M. 2013. Chimeric viruses blur the582borders between the major groups of eukaryotic single-stranded DNA viruses. Nat Commun5834:2700.
- 584 21. Krupovic M, Zhi N, Li J, Hu G, Koonin E V., Wong S, Shevchenko S, Zhao K, Young NS. 2015.
 585 Multiple layers of chimerism in a single-stranded DNA virus discovered by deep sequencing.
 586 Genome Biol Evol 7:993–1001.
- 587 22. Hewson I, Ng G, Li WF, LaBarre BA, Aguirre I, Barbosa JG, Breitbart M, Greco AW, Kearns
 588 CM, Looi A, Schaffner LR, Thompson PD, Hairston NG. 2013. Metagenomic identification,
 589 seasonal dynamics, and potential transmission mechanisms of a Daphnia-associated single590 stranded DNA virus in two temperate lakes. Limnol Oceanogr 58:1605–1620.
- 591 23. Steel O, Kraberger S, Sikorski A, Young LM, Catchpole RJ, Stevens AJ, Ladley JJ, Coray DS,
 592 Stainton D, Dayaram A, Julian L, van Bysterveldt K, Varsani A. 2016. Circular replication593 associated protein encoding DNA viruses identified in the faecal matter of various animals in
 594 New Zealand. Infect Genet Evol 43:151–164.
- 59524.McDaniel LD, Rosario K, Breitbart M, Paul JH. 2014. Comparative metagenomics: natural596populations of induced prophages demonstrate highly unique, lower diversity viral sequences.597Environ Microbiol 16:570–585.
- 59825.Dayaram A, Galatowitsch ML, Argüello-Astorga GR, van Bysterveldt K, Kraberger S, Stainton599D, Harding JS, Roumagnac P, Martin DP, Lefeuvre P, Varsani A. 2016. Diverse circular600replication-associated protein encoding viruses circulating in invertebrates within a lake601ecosystem. Infect Genet Evol 39:304–316.
- 602 26. Bistolas K, Besemer R, Rudstam L, Hewson I. 2017. Distribution and Inferred Evolutionary
 603 Characteristics of a Chimeric ssDNA Virus Associated with Intertidal Marine Isopods. Viruses
 604 9:361.
- 60527.Quaiser A, Krupovic M, Dufresne A, Francez A-J, Roux S. 2016. Diversity and comparative606genomics of chimeric viruses in Sphagnum- dominated peatlands. Virus Evol 2:vew025.
- 607 28. Salmier A, Tirera S, de Thoisy B, Franc A, Darcissac E, Donato D, Bouchier C, Lacoste V,
 608 Lavergne A. 2017. Virome analysis of two sympatric bat species (Desmodus rotundus and
 609 Molossus molossus) in French Guiana. PLoS One 12:e0186943.
- 610 29. de la Higuera I, Torrance EL, Pratt AA, Kasun GW, Maluenda A, Stedman KM. 2019. Genome
 611 Sequences of Three Cruciviruses Found in the Willamette Valley (Oregon). Microbiol Resour
 612 Announc 8.
- 613 30. Kraberger S, Argüello-Astorga GR, Greenfield LG, Galilee C, Law D, Martin DP, Varsani A.
 614 2015. Characterisation of a diverse range of circular replication-associated protein encoding DNA
 615 viruses recovered from a sewage treatment oxidation pond. Infect Genet Evol 31:73–86.
- Tisza MJ, Pastrana D V, Welch NL, Stewart B, Peretti A, Starrett GJ, Pang Y-YS, Krishnamurthy
 SR, Pesavento PA, McDermott DH, Murphy PM, Whited JL, Miller B, Brenchley J, Rosshart SP,
 Rehermann B, Doorbar J, Ta'ala BA, Pletnikova O, Troncoso JC, Resnick SM, Bolduc B,
 Sullivan MB, Varsani A, Segall AM, Buck CB. 2020. Discovery of several thousand highly
 diverse circular DNA viruses. Elife 9:555375.

- 621 32. Brown DR, Schmidt-Glenewinkelg T, Reinberg D, Hurwitz J. 1983. DNA sequences which support activities of the bacteriophage phiX174 gene A protein.
- 33. Steinfeldt T, Finsterbusch T, Mankertz A. 2006. Demonstration of Nicking/Joining Activity at the
 Origin of DNA Replication Associated with the Rep and Rep' Proteins of Porcine Circovirus
 Type 1. J Virol 80:6225–6234.
- 626 34. Gassmann M, Focher F, Buhk HJ, Ferrari E, Spadari S, Hübscher U. 1988. Replication of single627 stranded porcine circovirus DNA by DNA polymerases α and δ. BBA Gene Struct Expr.
- 628 35. Roth MJ, Brown DR, Hurwitz J. 1984. Analysis of Bacteriophage phi174 Gene A Protein629 mediated Termination and Reinitiation of 6X DNA Synthesis 11. Structural characterization of
 630 the covalent 4X A protein-DNA complex.J. Biol. Chern.
- 631 36. Cheung AK. 2007. A stem-loop structure, sequence non-specific, at the origin of DNA replication
 632 of porcine circovirus is essential for termination but not for initiation of rolling-circle DNA
 633 replication. Virology 363:229–235.
- 63437.Stenger DC, Revington GN, Stevenson MC, Bisaro DM. 1991. Replicational release of635geminivirus genomes from tandemly repeated copies: Evidence for rolling-circle replication of a636plant viral DNA. Proc Natl Acad Sci U S A.
- 637 38. Chandler M, de la Cruz F, Dyda F, Hickman AB, Moncalian G, Ton-Hoang B. 2013. Breaking
 638 and joining single-stranded DNA: the HUH endonuclease superfamily. Nat Rev Microbiol 11:525–538.
- 64039.Brown DR, Schmidt-Glenewinkelg T, Reinberg D, Hurwitz J. 1983. DNA sequences which641support activities of the bacteriophage phiX174 gene A protein 258:8402–8412.
- 642 40. Ilyina T V., Koonin E V. 1992. Conserved sequence motifs in the initiator proteins for rolling
 643 circle DNA replication encoded by diverse replicons from eubacteria, eucaryotes and
 644 archaebacteria. Nucleic Acids Res 20:3279–3285.
- 645 41. Koonin E V, Ilyina T V. 1993. Computer-assisted dissection of rolling circle DNA replication.
 646 BioSystems 30:241–268.
- 647 42. Londoño A, Riego-Ruiz L, Argüello-Astorga GR. 2010. DNA-binding specificity determinants of
 648 replication proteins encoded by eukaryotic ssDNA viruses are adjacent to widely separated RCR
 649 conserved motifs. Arch Virol 155:1033–1046.
- 65043.Gorbalenya AE, Koonin E V. 1993. Helicases: amino acid sequence comparisons and structure-651function relationships. Curr Opin Struct Biol 3:419–429.
- 652 44. Clerot D, Bernardi F. 2006. DNA Helicase Activity Is Associated with the Replication Initiator
 653 Protein Rep of Tomato Yellow Leaf Curl Geminivirus. J Virol 80:11322–11330.
- 65445.Kazlauskas D, Varsani A, Krupovic M. 2018. Pervasive chimerism in the replication-associated655proteins of uncultured single-stranded DNA viruses. Viruses 10:1–11.
- 46. Krupovic M, Koonin E V. 2017. Multiple origins of viral capsid proteins from cellular ancestors.
 Proc Natl Acad Sci U S A 114:E2401–E2410.
- 47. Hopper P, Harrison SC, Sauer RT. 1984. Structure of tomato bushy stunt virus. V. Coat protein sequence determination and its structural implications. J Mol Biol 177:701–713.
- 48. Sherman MB, Guenther R, Reade R, Rochon D, Sit T, Smith TJ. 2019. Near-Atomic-Resolution
 661 Cryo-Electron Microscopy Structures of Cucumber Leaf Spot Virus and Red Clover Necrotic
 662 Mosaic Virus: Evolutionary Divergence at the Icosahedral Three-Fold Axes. J Virol 94.

- 49. Alam SB, Reade R, Theilmann J, Rochon DA. 2017. Evidence for the role of basic amino acids in
 the coat protein arm region of Cucumber necrosis virus in particle assembly and selective
 encapsidation of viral RNA. Virology 512:83–94.
- 666 50. Park SH, Sit TL, Kim KH, Lommel SA. 2013. The red clover necrotic mosaic virus capsid
 667 protein N-terminal amino acids possess specific RNA binding activity and are required for stable
 668 virion assembly. Virus Res 176:107–118.
- 669 51. Ohki T, Akita F, Mochizuki T, Kanda A, Sasaya T, Tsuda S. 2010. The protruding domain of the coat protein of Melon necrotic spot virus is involved in compatibility with and transmission by the fungal vector Olpidium bornovanus. Virology 402:129–134.
- 52. Llauró A, Coppari E, Imperatori F, Bizzarri AR, Castón JR, Santi L, Cannistraro S, De Pablo PJ.
 2015. Calcium Ions Modulate the Mechanics of Tomato Bushy Stunt Virus. Biophys J 109:390–
 397.
- 53. Paez-Espino D, Chen I-MA, Palaniappan K, Ratner A, Chu K, Szeto E, Pillay M, Huang J,
 Markowitz VM, Nielsen T, Huntemann M, K. Reddy TB, Pavlopoulos GA, Sullivan MB,
 Campbell BJ, Chen F, McMahon K, Hallam SJ, Denef V, Cavicchioli R, Caffrey SM, Streit WR,
 Webster J, Handley KM, Salekdeh GH, Tsesmetzis N, Setubal JC, Pope PB, Liu W-T, Rivers AR,
 Ivanova NN, Kyrpides NC. 2017. IMG/VR: a database of cultured and uncultured DNA Viruses
 and retroviruses. Nucleic Acids Res 45:D457–D465.
- 68154.Wright EA, Heckel T, Groenendijk J, Davies JW, Boulton MI. 1997. Splicing features in maize682streak virus virion- and complementary-sense gene expression. Plant J 12:1285–1297.
- 55. Steinfeldt T, Finsterbusch T, Mankertz A. 2006. Demonstration of Nicking/Joining Activity at the
 Origin of DNA Replication Associated with the Rep and Rep' Proteins of Porcine Circovirus
 Type 1. J Virol 80:6225–6234.
- 686 56. Hafner GJ, Dale JL, Harding RM, Wolter LC, Stafford MR. 1997. Nicking and joining activity of
 687 banana bunchy top virus replication protein in vitro. J Gen Virol 78:1795–1799.
- 688 57. Lorenz R, Bernhart SH, Höner zu Siederdissen C, Tafer H, Flamm C, Stadler PF, Hofacker IL.
 689 2011. ViennaRNA Package 2.0. Algorithms Mol Biol 6:26.
- 690 58. Mathews DH, Sabina J, Zuker M, Turner DH. 1999. Expanded sequence dependence of
 691 thermodynamic parameters improves prediction of RNA secondary structure. J Mol Biol
 692 288:911–940.
- 693 59. Muhire BM, Varsani A, Martin DP. 2014. SDT: A Virus Classification Tool Based on Pairwise
 694 Sequence Alignment and Identity Calculation. PLoS One 9:e108277.
- 69560.Waterhouse AM, Procter JB, Martin DMA, Clamp M, Barton GJ. 2009. Jalview Version 2-A696multiple sequence alignment editor and analysis workbench. Bioinformatics 25:1189–1191.
- 697 61. Livingstone CD, Barton GJ. 1993. Protein sequence alignments: A strategy for the hierarchical analysis of residue conservation. Bioinformatics.
- 699 62. Crooks GE, Hon G, Chandonia JM, Brenner SE. 2004. WebLogo: A sequence logo generator.
 700 Genome Res.
- Kelley LA, Mezulis S, Yates CM, Wass MN, Sternberg MJE. 2015. The Phyre2 web portal for
 protein modeling, prediction and analysis. Nat Protoc 10:845–858.
- 703 64. Pettersen EF, Goddard TD, Huang CC, Couch GS, Greenblatt DM, Meng EC, Ferrin TE. 2004.
 704 UCSF Chimera A visualization system for exploratory research and analysis. J Comput Chem 25:1605–1612.

- 706 65. Katoh K, Standley DM. 2013. MAFFT Multiple Sequence Alignment Software Version 7:
 707 Improvements in Performance and Usability. Mol Biol Evol 30:772–780.
- Floden EW, Tommaso PD, Chatzou M, Magis C, Notredame C, Chang J-M. 2016. PSI/TM-Coffee: a web server for fast and accurate multiple sequence alignments of regular and transmembrane proteins using homology extension on reduced databases. Nucleic Acids Res 44:W339–W343.
- 712 67. Capella-Gutiérrez S, Silla-Martínez JM, Gabaldón T. 2009. trimAl: A tool for automated
 713 alignment trimming in large-scale phylogenetic analyses. Bioinformatics.
- Price MN, Dehal PS, Arkin AP. 2009. Fasttree: Computing large minimum evolution trees with
 profiles instead of a distance matrix. Mol Biol Evol 26:1641–1650.
- 69. Guindon S, Dufayard JF, Lefort V, Anisimova M, Hordijk W, Gascuel O. 2010. New algorithms
 717 and methods to estimate maximum-likelihood phylogenies: Assessing the performance of PhyML
 718 3.0. Syst Biol 59:307–321.
- 719 70. Anisimova M, Gascuel O. 2006. Approximate Likelihood-Ratio Test for Branches: A Fast,
 720 Accurate, and Powerful Alternative. Syst Biol 55:539–552.
- 721 71. Huson DH, Scornavacca C. 2012. Dendroscope 3: An interactive tool for rooted phylogenetic
 722 trees and networks. Syst Biol.
- 723 72. Zallot R, Oberg N, Gerlt JA. 2019. The EFI Web Resource for Genomic Enzymology Web Tools:
 724 Leveraging Protein, Genome, and Metagenome Databases to Discover Novel Enzymes and
 725 Metabolic Pathways. Biochemistry acs.biochem.9b00735.
- 726 73. Shannon P, Markiel A, Ozier O, Baliga NS, Wang JT, Ramage D, Amin N, Schwikowski B,
 727 Ideker T. 2003. Cytoscape: A software Environment for integrated models of biomolecular
 728 interaction networks. Genome Res 13:2498–2504.
- 729 74. Kazlauskas D, Dayaram A, Kraberger S, Goldstien S, Varsani A, Krupovic M. 2017.
 730 Evolutionary history of ssDNA bacilladnaviruses features horizontal acquisition of the capsid gene from ssRNA nodaviruses. Virology 504:114–121.
- 732 75. Sicard A, Michalakis Y, Gutiérrez S, Blanc S. 2016. The Strange Lifestyle of Multipartite
 733 Viruses. PLoS Pathog 12:1–19.
- 734 76. Varsani A, Lefeuvre P, Roumagnac P, Martin D. 2018. Notes on recombination and reassortment
 735 in multipartite/segmented viruses. Curr Opin Virol 33:156–166.
- 736 77. Grasse W, Spring O. 2017. ssRNA viruses from biotrophic Oomycetes form a new phylogenetic
 737 group between Nodaviridae and Tombusviridae. Arch Virol 162:1319–1324.
- 738 78. Rosario K, Mettel KA, Benner BE, Johnson R, Scott C, Yusseff-Vanegas SZ, Baker CCM,
 739 Cassill DL, Storer C, Varsani A, Breitbart M. 2018. Virus discovery in all three major lineages of
 740 terrestrial arthropods highlights the diversity of single-stranded DNA viruses associated with
 741 invertebrates. PeerJ 6:e5761.
- 742 79. Varsani A, Krupovic M. 2017. Sequence-based taxonomic framework for the classification of uncultured single-stranded DNA viruses of the family Genomoviridae. Virus Evol 3.
- 744 80. Harrison SC, Olson AJ, Schutt CE, Winkler FK, Bricogne G. 1978. Tomato bushy stunt virus at
 745 2.9 Å resolution. Nature 276:368–373.
- 746 81. Chelvanayagam G, Heringa J, Argos P. 1992. Anatomy and Evolution of Proteins Displaying the
 747 Viral Capsid Jellyroll Topology. J Mol Biol 228:220–242.

- Katpally U, Kakani K, Reade R, Dryden K, Rochon D, Smith TJ. 2007. Structures of T=1 and T=3 Particles of Cucumber Necrosis Virus: Evidence of Internal Scaffolding. J Mol Biol 365:502–512.
- 751 83. Campbell JW, Clifton IJ, Greenhough TJ, Hajdu J, Harrison SC, Liddington RC, Shrive AK.
 752 1990. Calcium binding sites in tomato bushy stunt virus visualized by laue crystallography. J Mol Biol 214:627–632.
- 754 84. Casado CG, Javier Ortiz G, Padron E, Bean SJ, McKenna R, Agbandje-McKenna M, Boulton MI.
 755 2004. Isolation and characterization of subgenomic DNAs encapsidated in "single" T = 1 isometric particles of Maize streak virus. Virology 323:164–171.
- 85. Bennett A, Rodriguez D, Lister S, Boulton M, McKenna R, Agbandje-McKenna M. 2018.
 Assembly and disassembly intermediates of maize streak geminivirus. Virology 525:224–236.
- 759 86. Chen JK, Hsiao C, Wu JS, Lin SY, Wang CY. 2019. Characterization of the endonuclease
 760 activity of the replication-associated protein of beak and feather disease virus. Arch Virol 164:2091–2106.
- Rizvi I, Choudhury NR, Tuteja N. 2015. Insights into the functional characteristics of geminivirus rolling-circle replication initiator protein and its interaction with host factors affecting viral DNA replication. Arch Virol 160:375–387.
- 765 88. Pasumarthy KK, Choudhury NR, Mukherjee SK. 2010. Tomato leaf curl Kerala virus
 766 (ToLCKeV) AC3 protein forms a higher order oligomer and enhances ATPase activity of
 767 replication initiator protein (Rep/AC1). Virol J 7:128.
- Nash TE, Dallas MB, Reyes MI, Buhrman GK, Ascencio-Ibanez JT, Hanley-Bowdoin L. 2011.
 Functional Analysis of a Novel Motif Conserved across Geminivirus Rep Proteins. J Virol 85:1182–1192.
- 771 90. Varsani A, Krupovic M. 2017. Sequence-based taxonomic framework for the classification of
 772 uncultured single-stranded DNA viruses of the family Genomoviridae. Virus Evol 3.
- Shi M, Lin XD, Tian JH, Chen LJ, Chen X, Li CX, Qin XC, Li J, Cao JP, Eden JS, Buchmann J,
 Wang W, Xu J, Holmes EC, Zhang YZ. 2016. Redefining the invertebrate RNA virosphere.
 Nature 540:539–543.
- Dolja V V., Koonin E V. 2018. Metagenomics reshapes the concepts of RNA virus evolution by revealing extensive horizontal virus transfer. Virus Res 244:36–52.
- 778 93. Greninger AL, DeRisi JL. 2015. Draft Genome Sequence of Tombunodavirus UC1. Genome
 779 Announc 3.
- 780 94. Krupovic M. 2013. Networks of evolutionary interactions underlying the polyphyletic origin of ssDNA viruses. Curr Opin Virol 3:578–586.
- 782 95. Allison AB, Organtini LJ, Zhang S, Hafenstein SL, Holmes EC, Parrish CR. 2016. Single
 783 Mutations in the VP2 300 Loop Region of the Three-Fold Spike of the Carnivore Parvovirus
 784 Capsid Can Determine Host Range. J Virol.
- 785 96. Carbonell A, Maliogka VI, Pérez J de J, Salvador B, León DS, García JA, Simón-Mateo C. 2013.
 786 Diverse Amino Acid Changes at Specific Positions in the N-Terminal Region of the Coat Protein
 787 Allow Plum pox virus to Adapt to New Hosts. Mol Plant-Microbe Interact 26:1211–1224.
- 788 97. Beakes GW, Glockling SL, Sekimoto S. 2012. The evolutionary phylogeny of the oomycete
 789 "fungi." Protoplasma 249:3–19.

790	98.	Hanyu N, Kuchino Y, Nishimura S, Beier H. 1986. Dramatic events in ciliate evolution:
791		alteration of UAA and UAG termination codons to glutamine codons due to anticodon mutations
792		in two Tetrahymena tRNAs Gln . EMBO J 5:1307–1311.

- 99. Soffer N, Brandt ME, Correa AMS, Smith TB, Thurber RV. 2014. Potential role of viruses in white plague coral disease. ISME J 8:271–283.
- 100. Krupovic M, Bamford DH. 2010. Order to the Viral Universe. J Virol 84:12476–12479.

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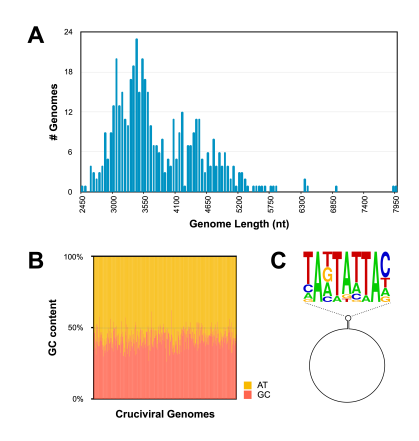


Figure 1. Genome properties of 461 new cruciviral circular sequences. (A) Histogram of cruciviral genome lengths categorized in 50 nt bins. (B) Percentage of G + C content versus A + T in each of the sequences described in this study (C) Relative abundance of nucleotides in the conserved nonanucleotide sequence of the 211 stem-loops and putative origins of replication represented predicted with StemLoop-Finder (Pratt *et* al., unpublished) in Sequence Logo format.

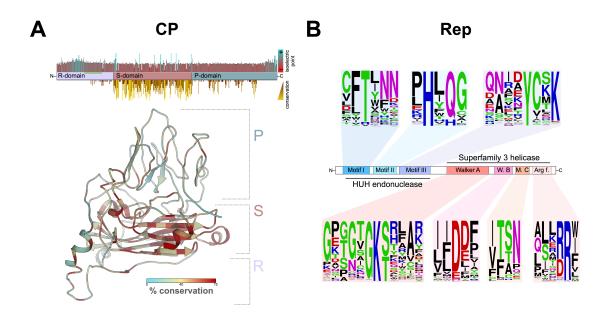


Figure 2. Protein conservation in cruciviruses. (A) Top: distribution of domains, isoelectric point and conservation in a consensus capsid protein (CP). 461 CP protein sequences were aligned in Geneious 11.0.4 with MAFFT (G-INS-i, BLOSUM 45, open gap penalty 1.53, offset 0.123) and trimmed manually. The conservation of the physico-chemical properties at each position was obtained with Jalview v2.11.0, and the isoelectric point was estimated in Geneious 11.0.4. The region of CP rich in glycine is highlighted with a green bar. **Bottom:** Structure of a cruciviral CP (CruV-359) as predicted by Phyre² showing sequence conservation based on an alignment of the 47 CP protein sequences from the CP-based clusters. (B) Conserved motifs found in cruciviral Reps after aligning all the extracted Rep protein sequences using PSI-Coffee. Sequence logos were generated at http://weblogo.threeplusone.com to indicate the frequency of residues at each position.

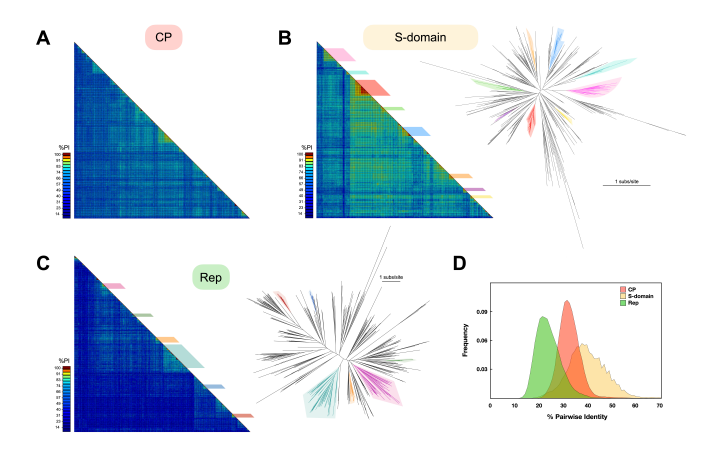


Figure 3. Diversity of cruciviral proteins. (A) CP diversity. Pairwise amino acid identity (PI) between the CPs predicted for 461 cruciviral sequences. The alignment and analysis were carried out with SDT, using the integrated MAFFT algorithm. (B) S-domain diversity. Left: PI matrix between the capsid protein (CP) predicted S-domain of the 461 sequences described in this study. The alignment and analysis were carried out with SDT, using the integrated MAFFT algorithm. The colored boxes indicate the different clusters of sequences used to create the CP-based clusters sequence subset. **Right:** Unrooted phylogenetic tree obtained with FastTree from a manually curated MAFFT alignment of the translated sequences of the S-domain (G-INS-i, BLOSUM 45, open gap penalty 1.53, offset 0.123). The colored branches represent the different clusters observed in the matrix. Scale bar indicates substitutions per site. (C) Rep diversity. Left: Pairwise identity (PI) matrix between all Reps found in cruciviral genomes in this study. The alignment and analysis were carried out with SDT, using the integrated MUSCLE algorithm. Right: Unrooted phylogenetic tree obtained with FastTree from an PSI-Coffee alignment of the translated sequences of Rep trimmed with TrimAl v1.3. The colored branches represent the different clusters that contain Rep-based clusters sequence subset. Scale bar indicates substitutions per site. (D) PI frequency distribution. The frequency of PI values for each of the putative proteins or domains analyzed is shown.

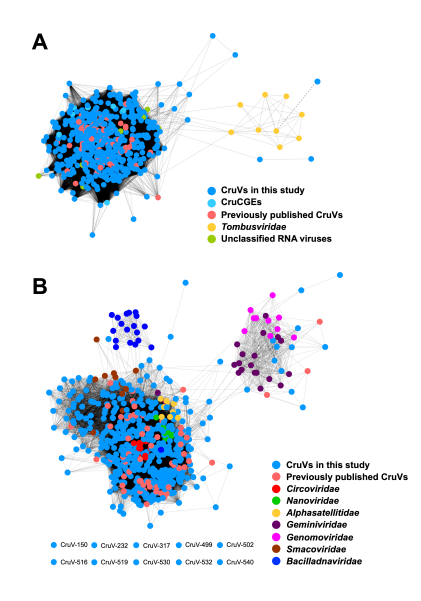


Figure 4. Similarity networks of cruciviral proteins with related viruses. (A) Capsid proteins (CP) represented by colored dots are connected with a solid line when the similarity between them is greater than e-value= $1e^{-20}$. The dashed line represents an e-value = $6e^{-7}$ between the nodes corresponding to the CP of CruV-523 and turnip crinkle virus, as given by BLASTp. (B) Replication-associated protein (Rep) translations, represented by colored dots, are connected with a solid line when the similarity between them greater than e-value= $1e^{-10}$. The eight nodes at the bottom left did not connect to any other node. All networks were carried out with pairwise identities calculated in the EFI–EST web server and visualized in Cytoscape v3.7.2.

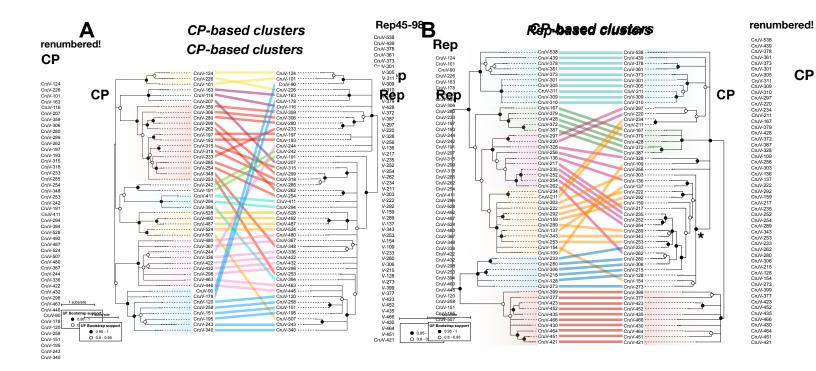


Figure 5. Comparison of phylogenies of CP and Rep proteins of representative cruciviruses. (A) Tanglegram calculated with Dendroscope v3.5.10 from phylogenetic trees generated with PhyML from Cp (PhyML automatic model selection LG+G+I+F) and Rep (PhyML automatic model selection RtREV+G+I) alignments. The tips corresponding to the same viral genome are linked by lines that are color-coded according to the clusters obtained from Fig. 3A (CP-based clusters). (B) Tanglegram calculated with Dendroscope v3.5.10 from phylogenetic trees generated with PhyML from Cp (PhyML automatic model selection LG+G+I+F) and Rep (PhyML automatic model selection RtREV+G+I) alignments. The tips corresponding to the same viral sequence are linked by lines that are color-coded according to the clusters obtained from Fig. 3B (Rep-based clusters). The clade marked with an asterisk is formed by members of the red cluster of subset A. Branch support is given according to aLRT SH-like (Anisimova & Gascuel, 2006). All nodes with an aLRT SH-like branch support inferior to 0.8 were collapsed with Dendroscope prior to constructing the tanglegram.

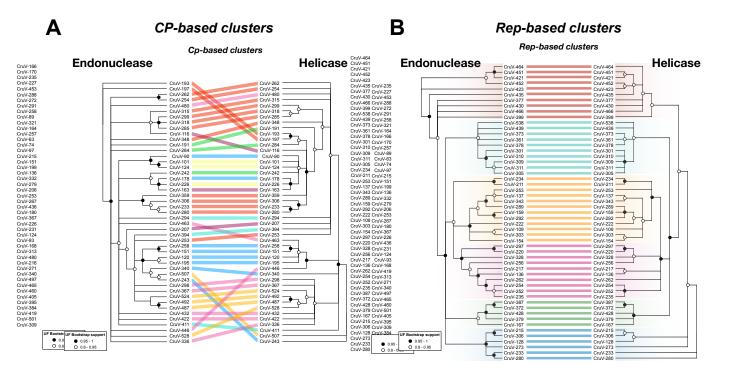


Figure 6. Comparison of phylogenies between the endonuclease and helicase domains of Reps from representative cruciviruses. (A) Tanglegram calculated with Dendroscope v3.5.10 from phylogenetic trees generated with PhyML from separate alignments of Rep endonuclease and helicase domains. The tips corresponding to the same viral genome are linked by lines that are color-coded according to the clusters obtained from Fig. 3A (CP-based clusters). (B) Same as A but with sequences from the clusters obtained from Fig. 3B (Rep-based clusters). All nodes with an aLRT SH-like branch support inferior to 0.8 were collapsed with Dendroscope v3.5.10 prior to constructing the tanglegram.