Primordial capsid and spooled ssDNA genome structures penetrate 1 ancestral events of eukaryotic viruses 2

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24 Abstract

25 Marine algae viruses are important for controlling microorganism communities in the marine 26 ecosystem, and played a fundamental role during the early events of viral evolution. Here, we have 27 focused on one major group of marine algae viruses, the ssDNA viruses from the Bacilladnaviridae 28 family. We present the capsid structure of the bacilladnavirus, Chaetoceros tenuissimus DNA virus 29 type II (CtenDNAV-II), determined at 2.3 Å resolution. Structural comparison to other viruses 30 supports the previous theory where bacilladnaviruses were proposed to have acquired their capsid 31 protein via horizontal gene transfer from a ssRNA virus. The capsid protein contains the widespread 32 virus jelly-roll fold, but has additional unique features; a third β-sheet and a long C-terminal tail. 33 which are located on the capsid surface and could be important for virus transmission. Further, low-34 resolution reconstructions of the CtenDNAV-II genome reveal a partially spooled structure, an

arrangement previously only described for dsRNA and dsDNA viruses.

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37 Key words: algae virus, asymmetric reconstruction, CRESS-DNA, Cressdnaviricota, cryo-EM,

38 Bacilladnaviridae, bacilladnavirus, diatom virus, evolution, genome structure, spooled genome

39 Introduction

40 Marine algae viruses prevail massively in the oceans and greatly affect the global ecosystem by 41 causing mortality and lysis of microbial communities, releasing organic carbon and other nutrients 42 back into the environment (the 'viral shunt'), thereby affecting the global oxygen level, and the marine nutrient and energy cycling (Fuhrman, 1999; Suttle, 2007; Wilhelm and Suttle, 1999). Algae blooms 43 44 are global phenomena of dynamic appearance and disappearance of certain algae species. The 45 recurring patterns of algae blooms are associated with the presence of marine diatom/algae viruses 46 (Bratbak et al., 1993; Johannessen et al., 2017; Tarutani et al., 2000). The viruses typically have a very 47 narrow host range, thus causing host-specific mortality and control of algae host populations 48 (Brussaard, 2004), and are therefore attractive for being used as a microbial agent against harmful 49 algae blooms (Nagasaki et al., 1999).

50 Another incentive for studying marine algal viruses is for penetrating early evolutionary scenarios of primordial eukaryotic viruses. Since unicellular marine organisms were the earliest eukaryotes on 51 52 earth, they were presumably host of the most ancient viruses (Dolja and Koonin, 2018; Koonin et al., 53 2015). The extant unicellular organisms, such as unicellular algae, therefore likely retain genetic and 54 structural characteristics from their ascendant (Koonin and Dolja, 2014; Munke et al., 2020). 55 Structural comparison between so-called primordial viruses, such as diatom viruses, and the 56 phylogenetically closely related viruses in the same evolutionary lineages can reveal acquired 57 functional structures in viruses that infect higher eukaryotes such as human, animal and crop 58 pathogens (Munke et al., 2020; Okamoto et al., 2020, 2016).

59 The Bacilladnaviridae viruses in this study are greatly involved in the discussions of both the 60 global ecosystem model and early virus evolution. The Bacilladnaviridae viruses infect a diverse 61 range of marine bloom-forming algae species (Kimura and Tomaru, 2015; Tomaru et al., 2013, 2011). 62 These Bacilladnaviridae viruses are only infectious to their specific algae host. They lyse the host 63 algae cells and reduce the number significantly during their propagation (Kimura and Tomaru, 2015; 64 Tomaru et al., 2011, 2013). Understanding their host-specific transmission mechanism is the key to 65 clarify how these ssDNA viruses affect the dynamics of bloom-forming algae species. 66 Bacilladnaviruses carry a circular ssDNA genome of ~ 6 kb, which is partially double-stranded (~ 700 -67 800 bp) and encodes three proteins; one coat protein, one replication protein (Rep), and a third protein 68 with unknown function (Kimura and Tomaru, 2015). Until recently, bacilladnaviruses have been 69 included in the informal group CRESS DNA viruses (for circular Rep-encoding ssDNA viruses), but 70 this group has now formed the phylum *Cressdnaviricota* (Krupovic et al., 2020).

71 Viruses with different nucleic acid types have traditionally been organised into different classes, 72 however an increasing number of evidence has emerged during the last decade suggesting that RNA-73 DNA recombination is much more prevalent than previously recognized (Diemer and Stedman, 2012; 74 Kazlauskas et al., 2017; Krupovic, 2013; Tisza et al., 2020). In terms of bacilladnaviruses, Kazlauskas 75 et al suggested based on sequence homology that their capsid proteins likely have been acquired 76 through horizontal gene transfer (HGT) from an ancestral noda-like virus (Kazlauskas et al., 2017). 77 This is not an unreasonable scenario considering the prevalence of noda-like viruses in the aquatic 78 environment (Wolf et al., 2020). The icosahedral capsid of the Bacilladnaviridae viruses has adapted 79 to pack a ssDNA genome instead of a ssRNA genome, however the lack of available capsid and 80 genome structures of icosahedral ssDNA viruses have prevented addressing this theory.

Here, we present 3D reconstructions that reveal both the capsid and genome organization of the bacilladnavirus *Chaetoceros tenuissimus* DNA virus type II (CtenDNAV-II). An atomic model of the capsid protein could be constructed from the 2.3 Å resolution capsid structure. The capsid protein contains a jelly-roll fold with two additional extensions; a third seven-stranded β -sheet that is intertwined within the jelly-roll and C-terminal tails in two of the three proteins that form the 86 icosahedral asymmetric unit. Both extensions, which are unique to the bacilladnaviruses, are located

87 on the capsid surface and display a higher intraspecies sequence variability compared to other parts of

the capsid protein, suggesting a functional role during virus transmission. The virus transmission is

89 further discussed with support from a hemagglutination inhibition assay, adsorption test and sequence

90 data. The DALI program (Holm, 2020a) was used to demonstrate that the capsid of the

bacilladnaviruses indeed are structurally more similar to capside of RNA viruses than to those of other

ssDNA viruses, corroborating the HGT theory in early virus evolution. In addition, the first genome

structure of a ssDNA virus presented here suggests a spooled genome packaging mechanism, which

94 has previously only been described for dsRNA and dsDNA viruses.

95 **Results**

96 Summary of structure determination

97 The structure of the CtenDNAV-II viron was determined using cryo-electron microscopy (cryo-EM).

98 A Titan Krios microscope (Thermo Fisher Scientific) equipped with a K2 Summit direct electron 99 detector (Gatan) (see Materials and Methods) was used to record micrographs (Supplementary Fig. 100 1A) and confirmed mature virons containing a DNA genome. The particles had a diameter of 101 approximately 35 nm and appeared to be morphologically reproducible. The three-dimensional (3D) reconstruction of CtenDNAV-II was performed with RELION (Scheres, 2012). The capsid was 102 103 reconstructed by imposing icosahedral symmetry (I4) and using 33,507 particles, whereas the genome 104 structure was reconstructed without imposing any symmetry (C1) and using 21,559 particles. The 105 overall resolution of the final maps were estimated using the "gold standard" Fourier shell correlation 106 (FSC) 0.143 criterion (Henderson et al., 2012; Scheres and Chen, 2012) to 2.3 Å for the capsid 107 (Supplementary Fig. 1C) and 13 and 23 Å for the two genome reconstructions (Supplementary Fig. 2). 108 The local resolution of the capsid reconstruction was distributed between 2.3 and 9.1 Å and estimated 109 using ResMap (Kucukelbir et al., 2014) (Supplementary Fig. 1D). An atomic model was built, refined,

- 110 and validated according to our cryo-EM map. Data acquisition and processing, refinement, and
- 111 validation statistics are summarized in Table 1.
- 112

113 Table 1. Data collection, processing and refinement statistics

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300 -1.00 to 2.00				

114 The capsid protein displays different conformations within the asymmetric unit

115 The CtenDNAV-II capsid displays T = 3 symmetry, i.e. 180 capsid protein protomers assemble such 116 that the asymmetric unit comprises 3 capsid subunits in 3 quasi-equivalent positions termed A, B, and 117 C (Fig. 1A). For subunit A residues 64-72 and 77-371 were modelled, for subunit B residues 64-371 118 were modelled, and for subunit C residues 64-357 were modelled. The C-terminus of the A and B 119 subunits form long tails that end on the capsid surface around the 3- and 5-fold axes respectively. 120 Here, the last 19 residues could not be modelled (Fig. 2C). Additional density was however visible in 121 the map when the contour level was decreased (Supplementary Fig. 2), indicating that the C-terminus 122 forms flexible protrusions on the capsid surface. Another 14 residues were unmodelled in the C-123 terminus of the C subunit (red rectangle in Fig. 2C), nevertheless it is clear that, in contrast to the A-124 and B-subunits, the C-terminus is directed towards the capsid interior (Fig. 1C). Unmodelled density is 125 observed on the inside of the capsid below the C subunits (Supplementary Fig. 3), which presumably 126 originates from the C-terminus of the C subunit, since corresponding unmodelled density was not 127 visible below the A and B subunits (Supplementary Fig. 3A). Different conformations and/or 128 flexibility were also observed between the three subunits in the N-termini (Fig. 1C), which are all 129 located on the inside of the capsid. The first 63 residues of the N-terminus could not be modelled in 130 any of the three subunits (Fig. 2C). The modelled termini and corresponding cryo-EM map are 131 displayed in Fig. 1D. Apart from the termini, the three subunits have very similar structures (Fig. 1C). In addition, unmodelled density, presumably originating from an ion or small molecule, was visible in 132 133 the interface of the three subunits that constitute one protomer (Supplementary Fig. 4). The density is 134 surrounded by six arginine residues (R86 and R272), which is an unusual finding among viruses. 135 Similar arginine rings have however been described for the HIV virus, in which the arginines interact 136 with polyanions such as nucleoside triphosphates (dNTPs) and myo-inositol hexaphosphate (IP6) 137 (Mallery et al., 2018).





141 Figure 1. Atomic model of the CtenDNAV-II capsid. The three subunits, A, B and C, are coloured 142 purple, green and yellow, respectively, in panel A-C. (A) The entire capsid rendered with a surface 143 representation viewed down an icosahedral two-fold axis. (B) The secondary structure of one single 144 asymmetric unit viewed from the outside. (C) Superimposition of the three subunits. (D) Refined side 145 chains of representatives of secondary structural elements and areas that differ between the three 146 subunits.

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148 The capsid protein jelly-roll fold has unique features

149 The canonical viral jelly-roll consists of eight anti-parallel β-strands that are named from B to I and 150 arranged in two four-stranded sheets (BIDG and CHEF). The loops connecting each strand are named 151 BC, CD, etc. (Harrison et al., 1978; Rossmann et al., 1985). For CtenDNAV-II, the two sheets are 152 formed by strands BIDD'G and C''C'CHEF, respectively (Fig. 2A-C), thus containing three 153 additional strands (D', C' and C'') compared to the standard viral jelly-roll fold. In addition, a third 154 antiparallel β -sheet with seven strands is intertwined within the jelly-roll, i.e. strands from the third 155 sheet are formed by extensions of loops CD, EF and GH of the jelly-roll. The third sheet, which is 156 located on the capsid surface, is thus composed of two C-strands (C" and C""), two E-strands (E' 157 and E"), and three G-strands (G', G" and G"") (Fig. 2B). In conclusion, the jelly-roll fold of 158 CtenDNAV-II has three unique features: three additional strands, an extra surface-exposed β -sheet, 159 and C-terminal tails in subunit A and B.



- 160 TSDLSTY<mark>HIA NKRARMVEAK L</mark>DKKNNTDAA GEPFVPGSSR
- 161 Figure 2. Capsid protein topology and structural organisation. The β -strands are coloured 162 according which β -sheet they belong to. The β -strands within the green and blue β -sheets are named 163 alphabetically according to the conventional jelly-roll fold nomenclature (B to I) (A) The secondary 164 structure of subunit A. (B) Schematic diagram of the secondary structure. (C) The amino acid 165 sequence of the CtenDNAV-II capsid protein, starting from residue 1. Each line has 70 residues and is 166 further subdivided into blocks of 10 residues by spaces within the sequence. The residue numbering is the same as in PDB entry 7NS0. Modelled and unmodelled residues are coloured black and grey, 167 168 respectively. Residues highlighted with red rectangles were unmodelled in one of the subunits: F73-P76 in subunit A and H358-L371 in subunit C. The assigned secondary structure is shown 169 170 schematically above the sequence. The underlined residues in the unmodelled N-terminus indicate 171 positively charged residues.
- 172

173 Capsid proteins from CtenDNAV-II and ssRNA viruses are similar

Previous structures of so-called CRESS DNA viruses (phylum *Cressdnaviricota*) (Krupovic et al., 2020) include viruses from families *Ciroviridae* (eg. 3R0R and 5ZJU) and *Geminiviridae* (eg.6F2S

- and 6EK5). The capsid proteins in viruses from these two families also contain a jelly-roll domain, but
- 177 lack the third surface-exposed β -sheet and N-terminal tail found in CtenDNAV-II. A search with the

178 DALI program (Holm, 2020a) revealed that the CtenDNAV-II capsid protein is more similar to capsid 179 proteins of ssRNA viruses than to ssDNA viruses (Fig. 3A), corroborating the previous HGT theory 180 by Kazlauskas et al (Kazlauskas et al., 2017). The closest ssDNA virus is that of Beak and feather 181 disease virus (Circoviridae), which ends up on 11th place (z-score 8.4) behind ten RNA viruses. 182 Highest similarity is found between ssRNA viruses from families Carmotetraviridae, 183 Alphatetraviridae and Nodaviridae that resulted in z-scores of 14.4-15.1 (Fig. 3A). For interpretation 184 of the results from DALI see Holm (2020) (Holm, 2020b). Superimpositions between capsid proteins 185 from CtenDNAV-II and representatives from each family show that those RNA viruses with highest z-186 score all have surface projections in addition to the jelly-roll, albeit with different folds, that occupy 187 similar positions as the third β -sheet of CtenDNAV-II (Fig. 3B). However, in none of the other viruses 188 the projection is as intertwined with the jelly-roll as for CtenDNAV-II, and instead the protrusion 189 often forms a separate domain with a linker region, such as the Immunoglobulin (Ig)-like domain of 190 Providence virus (Carmotetraviridae) (left most panel in Figure 3B) (Speir et al., 2010). The C-191 terminal surface protruding tail in subunit A and B of CtenDNAV-II is a unique feature that is not 192 observed in any of the other virus families (Fig. 3B).

193 In addition to the C-terminal tail and β -sheet surface projection, superimpositions performed by the 194 DALI server also revealed non-conserved loops (HI and C'C'') from the jelly-roll (Supplementary

Fig. 5A, B). These areas are all located on the capsid surface clustered together around the five- and

- 196 three-fold axes (Supplementary Fig. 5C).
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Α						
PDB	Name	Family	Genome	z-score	rmsd	Residues
2qqp	Providence virus*	Carmotetraviridae	ssRNA	15.1	2.9	209
3s6p	Helicoverpa Armigera Stunt virus*	Alphatetraviridae	ssRNA	14.9	2.9	208
4ftb	Flock House virus*	Nodaviridae	ssRNA	14.9	3.1	201
2bbv	Black Beetle virus	Nodaviridae	ssRNA	14.7	2.9	199
1ohf	Nudaurelia capensis omega virus	Alphatetraviridae	ssRNA	14.5	3.0	207
1f8v	Pariacoto virus	Nodaviridae	ssRNA	14.5	3.2	204
1nov	Nodamura virus	Nodaviridae	ssRNA	14.4	3.1	202
3ide	Infectious pancreatic necrosis virus*	Birnaviridae	dsRNA	12.1	3.0	192
2df7	Infectous brusal disease virus	Birnaviridae	dsRNA	12.1	3.3	198
2izw	Ryegrass mottle virus*	Solemoviridae	ssRNA	8.8	3.3	162
5zju	Beak and feather disease virus*	Circoviridae	ssDNA	8.4	4.1	155
В	·	· _	-	_	_	-



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Figure 3. Structural comparison of the CtenDNAV-II capsid protein with corresponding proteins from other viruses. (A) DALI-based structural comparison where the top 11 unique viruses (i.e., not unique PDB entries) with highest z-score are listed. Viruses indicated with * were used for the superimpositions in (B). For each comparison the z-score, rmsd and the number of residues that were used for the alignment are listed. (B) Superimpositions of the CtenDNAV-II capsid protein (gray) with representatives from each family listed in (A).

206 Intraspecies sequence variability in surface-exposed regions

207 Since the presented structure is the first of a virus from the Bacilladnaviridae family and its most 208 structurally close-related viruses are ssRNA viruses (Fig. 3), we sequenced (see Supplementary 209 methods) 15 CtenDNAV-II strains with the anticipation of improving our understanding of the 210 structure. The alignment (Supplementary Fig. 6) revealed 11 areas with mutations, of which 9 are 211 surface-exposed primarily on the C-terminal tail (R365, V367, A380 and R390) or beta-sheet 212 projection (A211-T213, T285 and S291) (Fig. 4). Thus, the mutated areas correlate with the unique 213 structural features for the CtenDNAV-II capsid protein that was identified from the superimpositions 214 with the structurally closely related viruses (Fig. 3B and Supplementary Fig. 5A). Interestingly, two 215 strains, 2-6 11V10 and 2-6 11V04, shared the same C-terminus sequence, which were completely 216 different from the other 13 strains (Fig. 4 and Supplementary Fig. 6).





Figure 4. Location of mutations identified among CtenDNAV-II strains. Subunit A (left), and a pentamer viewed from the outside looking down the five-fold axis (right) with mutations having the same colour code as in Supplementary Figure 6. Two mutations (A380 and R390), in dashed circle, are located in the unmodelled region of the C-terminus.

224 Chaetoceros tenuissimus viruses employ different transmission strategies

225 Sialic acids, acidic sugars that terminate glycan chains on the cell surface, are one of the common viral 226 receptors, and well-described for vertebrate viruses such as influenza, corona and polyoma. The 227 existence of sialic acids in algae and the discussions on their importance for algae virus interactions is 228 however more recent (Fulton et al., 2014; Munke et al., 2020; Wagstaff et al., 2018). Wagstaff and co-229 workers could demonstrate that Prymnesium paryum is capable of de novo synthesis of the deaminated 230 sialic acid Kdn, and through bioinformatics analysis, that sialic acid biosynthesis is widespread among 231 microalgae. However, we failed to detect homologous sequences in C. tenuissimus (Accession 232 numbers BLLK01000001-BLLK01000085) for any of the synthesases identified in Wagstaff et al. To 233 further corroborate our findings, we performed a classical hemagglutination inhibition (HI) assay (see 234 Supplementary methods) to investigate the sialic acid-binding capability of CtenDNAV-II. An RNA 235 virus (CtenRNAV-II) (family Marnaviridae) that infects the same diatom host was also tested as a representative of another major group of diatom viruses. The HI test demonstrated that CtenRNAV-II
shows partial HI activity, especially at high concentrations (no and 2 times dilution), while
CtenDNAV-II did not show any HI activity (Supplementary Fig. 6). In conclusion, both the synthesase
sequence analysis and HI assay indicate that the transmission of CtenDNAV-II is not sialic acidmediated.

241 Chaetoceros tenuissimus viruses display different host cell affinities

242 The adsorption kinetics of CtenDNAV-II and CtenRNAV-II were examined with two diatom species 243 including three culture strains (see Supplementary methods). The adsorption of CtenDNAV-II was not 244 detected for Chaetoceros socialis f. radians NIES-3713 (Supplementary Fig. 8). This result is 245 reasonable, since the DNA virus does not lyse C. socialis (Kimura and Tomaru 2015). Although 246 CtenDNAV-II lyses both C. tenuissimus strains NIES-3714 and NIES -3715 (Kimura and Tomaru 247 2015), significant adsorption of the virus was only observed with the former strain (Supplementary 248 Fig. 8). As for CtenRNAV-II, significant adsorption was detected for C. tenuissimus strain NIES-249 3715, but not to other diatom species and strains (Supplementary Fig. 8). A previous study reported 250 that CtenRNAV-II lyses C. tenuissimus strain NIES-3715, but not NIES-3714, and has a low lytic 251 activity on C. socialis f. radians (Kimura and Tomaru 2015). Therefore, the present result would be 252 consistent with the phenotypic relationship and suggest that the specific adsorption of CtenRNAV-II is 253 determined at host strain level. In conclusion, the degree of the virus and diatom host cell affinities 254 would be determined by combinations of the virus species and host strain level.

255 CtenDNAV-II genome structure is partially spooled

256 The genome organization of CtenDNAV-II is shown in Figure 5. The CtenDNAV-II genome consists 257 of an ordered outer layer (Fig. 5B and E) that is partially spooled around a disordered core (Fig. 5C). 258 The outer layer (EMDB-12555) was reconstructed at 13 Å by masking out the capsid, while the core 259 (EMDB-12556) was reconstructed at 23 Å by using a spherical mask (see Materials and Methods for 260 details). The reconstruction of the outer genome layer displays a coil of three turns that are positioned 261 between the icosahedral 5-fold axes (Fig. 5E). On each side of the three turns are additional DNA 262 fragments that do not follow the same spooling arrangement (Fig. 5B). The spooled genome 263 packaging has previously only been described in viruses with double stranded genomes (Ilca et al., 264 2019; Liu et al., 2019), whereas ssRNA viruses instead have a branched network formed by the 265 genome secondary structure (Dai et al., 2017; Gorzelnik et al., 2016; Koning et al., 2016). The distance between the parallel turns are about 28 Å (Fig. 5E), which is about the same distance as what 266 267 has been described for dsRNA viruses (Ilca et al., 2019). Below each icosahedral 5-fold axis of the 268 capsid the reconstruction of the outer genome layer displays protrusions towards the capsid (Fig. 5B 269 and E). However, these protrusions are not necessarily true at every position and could partially be 270 artefacts from the reconstruction, which is supported by the fact that some of the protrusions loose 271 connections to the remaining outer layer when the contour level is decreased while other protrusions 272 remain intact. Connections between the outer genome layer and the core seem to be confined to two 273 specific areas on opposite sides of the outer layer (Supplementary Fig. 9). Supplementary Movie 1 274 shows the asymmetric reconstruction in relation to the capsid.



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276 Figure 5. Asymmetric reconstructions of CtenDNAV-II radially coloured from red to blue. (A) 277 The capsid (blue) reconstructed with icosahedral symmetry and the asymmetric reconstructions 278 sighted below viewed down the icosahedral five-fold axis. (B) Same view as (A) but with the 279 icosahedral reconstruction removed. A protrusion (blue) from the outer genome layer (cyan) is 280 visualised at each five-fold axis (labelled as 5). (C) Same view as in (A) and (B) but with the 281 icosahedral and outer layer reconstructions removed. (D) A thin slice of the reconstructions shown in 282 (A). (E) The picture in (B) has been rotated 90°. The resolution (FSC=0.143) of the outer layer and 283 core are 13 Å and 23 Å, respectively. The FSC curves are shown in Supplementary Figure 2.

284 **Discussion**

285 Transmission mechanisms of Chaetoceros tenuissimus viruses

286 This paper describes the first structure (Figs. 1-2) of a Bacilladnaviridae virus, which is one of the 287 major groups of marine bloom-forming algae viruses. The CtenDNAV-II capsid protein contains the 288 widespread virus fold, the jelly-roll, but has in addition a third β -sheet and a C-terminal extension, 289 both of which form surface domains in close proximity to each other (Supplementary Fig.5) on the 290 capsid surface and that to our knowledge are unprecedented in other jelly-roll fold viruses determined 291 to date (Fig. 3, Supplementary Figs. S2 and S5). We have previously determined the structure of a 292 C. tenuissimus infecting RNA virus (family Marnaviridae), which has a primordial surface loop and 293 lacks the receptor-binding motif found in other structurally and phylogenetically related animal 294 viruses (Munke et al., 2020). This implies that these host-specific algae viruses have unique receptor-295 binding mechanisms. The molecular details on the infection mechanisms of diatom viruses are 296 unknown. However, a putative transmission route could exist through interactions with host sialic 297 acids, since sialic acid biosynthesis has been shown to be widespread among microalgae (Wagstaff et 298 al., 2018). In addition, many structurally related jelly-fold viruses interact with sialic acids for cell 299 entry, such as polyomaviruses via their conserved surface loops (Ströh et al., 2020). Likewise, the 300 C-terminal P-domain of the betanodavirus capsid protein interacts with sialic acids, and its deletion 301 affects virulence and host specificity (Liu et al., 2005; Moreno et al., 2019). The combined results of no homologous sequences in C. tenuissimus for any of the synthesases identified in Wagstaff et al 302 303 (Wagstaff et al., 2018), with the lack of HI activity (Supplementary Fig. 6), strongly refute a sialic 304 acid interaction between CtenDNAV-II and its host. This implies that the CtenDNAV-II virus 305 employs an alternative receptor and transmission strategy. The RNA virus infecting the same diatom 306 host (CtenRNAV-II) did however interact weakly with the mammalian sialic acid, which could 307 indicate that microalgae might have sialic acids or structurally related acidic sugars that have not yet 308 been described in the literature. The CtenRNAV-II virus is phylogenetically closely related to 309 Picornaviridae viruses, some of which use sialic acids as their entry receptor (Zocher et al., 2014). 310 The different receptor-binding strategies employed by the DNA and RNA virus could explain their 311 different host cell affinities (Supplementary Fig. 8).

312 Nevertheless, the CtenDNAV-II capsid protein has two unique surface features, the C-terminal tail 313 and β -sheet projection, compared to the most structurally close-related viruses (Fig. 3). These 314 extensions from the jelly-roll fold were in addition found to have an increasing number of mutations 315 between different virus species compared to other parts of the structure (Fig. 4 and Supplementary Fig. 316 6). Host specificity of algae viruses have previously been correlated with surface-exposed amino acid 317 substitutions (Nagasaki et al., 2005). Future research on CtenDNAV-II strains and their infection 318 patterns might reveal similar relationships. The locations of the mutations do however make the β -319 sheet and C-terminal extensions putative interaction candidates.

320 Structural insights on eukaryotic jelly-roll fold viruses in evolution

321 About 30% of all viruses adopt the single jelly-roll fold (Krupovic and Koonin, 2017), and the results 322 presented herein provide structural insights on the evolution of this fold. Our findings corroborate the 323 theory that the jelly-roll capsid proteins of ssDNA, ssRNA and dsRNA viruses originate from the 324 same virus with a primordial jelly-roll fold (Fig. 3) (Kazlauskas et al., 2017; Krupovic and Koonin, 325 2017). According to the HGT theory, the capsid protein of CRESS-DNA viruses could have been 326 acquired from the ssRNA protist viruses in early evolutionary events (Kazlauskas et al., 2019, 2017). 327 Indeed, the capsid protein of CtenDNAV-II is structurally more similar to ssRNA and dsRNA viruses 328 than to other CRESS-DNA viruses (Fig. 3). Because the rolling-circle replication proteins of CRESS-329 DNA viruses are polyphyletic, the capsid protein genes have probably been acquired from ssRNA viruses on several independent occasions and at different time-points in evolution (Krupovic, 2013).
Similarly, icosahedral dsRNA viruses are thought to have originated from ancestral ssRNA
nodaviruses and tetraviruses (Coulibaly et al., 2005). Yet another example includes the large CRESSDNA virus FLiP (Flavobacterium-infecting, lipid-containing phage) that shows a dsDNA adenoviruslike capsid structure (Laanto et al., 2017), which potentially links CRESS-DNA and dsDNA viruses.
To put our new structural insights into a broader context, we have updated the putative emergence of

- the ssDNA, ssRNA and dsRNA viruses (Fig. 6).
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Figure 6. A scheme depicting the putative emergence of virus capsids carrying different nucleic acid genomes. The figure illustrates based on current knowledge the most relevant virus groups and the evolutionary relationship between their virus capsids. Events that likely took place at an early stage of evolution in ancient algal pools are circled in blue. The dashed blue line represents that ssRNA and ssDNA virus recombination has occurred at several independent occasions, at different

time points in evolution. The figure is based on results described in this paper as well as previous results and discussions (Chiba et al., 2020; Coulibaly et al., 2005; Diemer and Stedman, 2012; Kazlauskas et al., 2019; Krupovic, 2013; Munke et al., 2020; Okamoto et al., 2020, 2016). The viron pictures were derived from ViralZone, SIB Swiss Institute of Bioinformatics (Hulo et al., 2011) (https://viralzone.expasy.org/) licensed under a Creative Commons Attribution 4.0 International Licens.

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351

Animal and plant ssRNA, dsRNA and CRESS-DNA viruses with icosahedral capsids that contain the jelly-roll fold should have acquired additional structural motifs for infecting their hosts. The primordial jelly-roll capsid protein structure of the CtenDNAV-II highlights the acquired features of these viruses that infect multicellular eukaryotes. Examples of such acquisitions are the Ig-like domain of tetraviruses and the P-domain of nodaviruses and birnaviruses (Fig. 3). These structures have been described as putative receptor-binding domains of those viruses; however, an exhaustive characterization still remains. The external Ig-like domain in tetraviruses (family *Carmotetraviridae*

359 and *Alphatetraviridae*) is believed to function in receptor binding to host cells (Helgstrand et al., 2004; 360 Penkler et al., 2016). Some Nodaviridae viruses, so-called gammanodaviruses, have an external P-361 domain that is important for virus attachment and entry (Ho et al., 2018), whereas the capsid proteins 362 of other nodaviruses retain primordial features that are structurally much more similar to the 363 CtenDNAV-II capsid (Johnson et al., 2001) (Fig. 3). The dsRNA Birnaviridae viruses display an 364 external highly antigenic P-domain that is likely involved in the initial attachment to the host cells 365 (Coulibaly et al., 2005; Garriga et al., 2006). The CRESS-DNA polyomaviruses possess a conserved 366 jelly-roll fold VP1 protein, and has acquired extra minor proteins VP2 and VP3 that has an ER-to-367 cytosol penetration function during cell entry (Inoue and Tsai, 2011). Several studies report on the 368 presence of a large number of diverse unclassified animal CRESS-DNA viruses, including putative 369 human pathogenic viruses (Ng et al., 2015; Phan et al., 2015; Tisza et al., 2020). Structural 370 comparison of their capsid proteins with that of CtenDNAV-II could reveal the acquired features of 371 these animal CRESS-DNA viruses.

A spooled genome requires structural adaptations of the capsid

The technological and computational improvements in cryo-EM structure determination have led to an increasing number of virus genome structures during the last decade. However, determining the genome structure within the symmetric icosahedral capsid still remains challenging, especially for single-stranded genomes (Koning et al., 2016). By employing a previously described method of subtracting the contribution of the capsid (Ilca et al., 2019), we were here able to demonstrate the first entire genome structure of a ssDNA virus (Fig. 5 and Supplementary Movie 1).

379 Previous cryo-EM studies have revealed three types of genome organization for non-enveloped 380 icosahedral viruses. The dsDNA and some dsRNA viruses such as Bacteriophage $\Phi 6$ forms spooled 381 genome structures with certain interfilament spacing (Ilca et al., 2019; Liu et al., 2019; Wang et al., 382 2019). Segmented dsRNA reoviruses instead form non-spooled or partially-spooled genomes with 383 pseudo-D3 symmetry that interact with the RNA-dependent RNA polymerase (RdRp) (Ding et al., 384 2019; Liu and Cheng, 2015; Wang et al., 2018; Zhang et al., 2015). The third variant is found in 385 Leviviridae viruses where the secondary structure of their ssRNA genomes form a branched network 386 of stem-loops (Dai et al., 2017; Gorzelnik et al., 2016; Koning et al., 2016), however it remains 387 unknown if a similar organisation is found in other ssRNA viruses.

388 The structure presented here is the first entire genome structure for a ssDNA virus. The genomes of 389 ssDNA viruses have the capability to form biologically functional secondary structures similar to 390 ssRNA viruses (Muhire et al., 2014) and since the capsid gene of bacilladnaviruses has been 391 horizontally transferred from ssRNA viruses, a similar branched structural network is imaginable also 392 for ssDNA viruses. However, the structure of the CtenDNAV-II genome is much more similar to those 393 of the spooled dsDNA or the partially spooled dsRNA genomes (Fig. 5 and Supplementary Movie 1). 394 Two major questions arise for the genome packaging of bacilladnaviruses. The first question is how to 395 stabilize the spooled ssDNA genome without a secondary structure organization? Potentially, a partial secondary structure might exist, and could together with the dsDNA segment that exist in 396 397 bacilladnaviruses (Kimura and Tomaru, 2015; Tomaru et al., 2011, 2013), be sufficient for stabilizing 398 the ssDNA genome inside the capsid. Another possibility is that the dsDNA gene of bacilladnaviruses 399 help stabilizing the ssDNA without having any secondary structure. The second question is how the ssRNA virus-like capsid protein was adapted to packing the dsDNA/dsRNA-like spooled genome of 400 401 the CtenDNAV-II? The spooled genome structures of the dsDNA and dsRNA viruses are packaged by 402 flexible interactions between the capsid protein and the genome. The interactions are mediated by 403 small contacts with hydrophobic and/or positively-charged amino acid residues of their capsid proteins 404 (Ding et al., 2019; Ilca et al., 2019; Liu and Cheng, 2015, 2015; Wang et al., 2019, 2018; Zhang et al., 405 2015). In contrast, the secondary structure of the (+)ssRNA viral genome binds specifically to the

406 capsid (Dai et al., 2017; Gorzelnik et al., 2016; Hesketh et al., 2018; Koning et al., 2016). Our 407 hypothesis is that the unmodelled N-terminus, and possibly the C-terminus of the C-subunit, which both possess numerous positively charged amino acids, interact with the ssDNA genome, in a flexible 408 409 and non-specific manner similar to dsDNA and dsRNA viruses. In contrary, the structurally related 410 tetra- and nodaviruses seem to interact more specifically with their ssRNA genome, since small RNA 411 segments (Speir et al., 2010) or parts of the genome (Johnson et al., 2001) have been revealed even 412 when the icosahedral symmetry has not been broken during the structure determination. This could 413 explain why the comparison between CtenDNAV-II and the RNA viruses revealed different structural 414 features of their capsids' interior, where the CtenDNAV-II was unmodelled and the ssRNA viruses 415 had additional α -helices (Fig. 3). A 7 nucleotide-stem loop gene fragment of the ssDNA geminivirus 416 has been reported to interact with its capsid (Hesketh et al., 2018), which could indicate that not all 417 ssDNA genomes are spooled.

The spooling genome structure is efficient for packaging long genomes of dsRNA and dsDNA viruses (Ilca et al., 2019; Liu and Cheng, 2015). The acquisition of the T=3 capsid gene from a ssRNA virus could facilitate the packaging of a larger CRESS-DNA genome that cannot be accommodated by capsids of small T=1 CRESS-DNA viruses (Kazlauskas et al., 2017). Many other ssDNA and dsDNA viruses such as animal tumor polyomaviruses and papillomaviruses, and plant geminiviruses are thought to be originated from primordial CRESS-DNA viruses (Kazlauskas et al., 2017; Koonin et al.,

424 2015; Koonin and Dolja, 2014). Considering the polyphyletic origin of ssDNA viruses and their

425 acquisitions of different RNA capsid proteins during the course of evolution, it will be interesting

426 when future ssDNA virus structures are unravelled.

427 Materials and methods

428 Virus production and purification

429 CtenDNAV-II was produced as previously described (Kimura and Tomaru, 2015). The crude virus 430 suspension was loaded onto 15 to 50% (w/v) sucrose density gradients and centrifuged at 24,000 431 $\times rpm$ (102,170 $\times g$) for 18 h at 4°C (Sw40Ti rotor, Beckman Coulter). The fractions of the sucrose 432 gradient were applied to SDS-PAGE. The VP2 capsid protein fractions were pooled and subjected to 433 centrifugation at 28,000 rpm (139,065 $\times g$) for 3 h at 4°C (Sw40Ti rotor, Beckman Coulter). The pellet

434 was resuspended in 50 mM Tris (pH 7.4), 100 mM NaCl, and 0.1 mM EDTA.

435 Cryo-EM and data collection

436 An aliquot (3μ) of purified CtenDNAV-II virons (10 mg ml^{-1}) was deposited onto freshly glow-437 discharged holey carbon-coated copper grids (Quantifoil R 2/2, 300 mesh, copper) followed by 2 s of 438 blotting in 100% relative humidity for plunge-freezing (Vitrobot Mark IV) in liquid ethane. Images 439 were acquired using a Titan Krios microscope (Thermo Fisher Scientific) operated at 300 kV and 440 equipped with a K2 Summit direct electron detector (Gatan) and an energy filter.

441 Image processing and 3D reconstruction

442 The micrographs were corrected for beam-induced drift using MotionCor2 1.2.6 (Zheng et al., 2017), 443 and contrast transfer function (CTF) parameters were estimated using Gctf 1.06 (Zhang, 2016). The 444 RELION 3.1 package (Zivanov et al., 2018) was used for particle picking, 2D and 3D classifications, 445 de novo 3D model generation and refinement. A reconstruction of the capsid was generated in I4 446 symmetry using 33,507 particles, which were obtained by performing 9 consecutive 2D classification 447 steps. The two genome reconstructions were generated in C1 symmetry using 21,559 particles, which 448 were generated by performing 6 consecutive 3D classifications of the 33,507 particles that were 449 obtained from the 2D classification step. Resolutions were estimated using the gold standard Fourier 450 shell correlation (threshold, 0.143 criterion) (Henderson et al., 2012; Scheres and Chen, 2012). The 451 data set and image processing are summarized in Table 1.

452 To reconstruct the genome without icosahedral symmetry a similar procedure to what has been 453 described by Ilca et al (Ilca et al., 2019) was carried out. The contribution of the capsid was first 454 subtracted from the map created during the final iteration of the I4 refinement job using the Particle 455 subtraction function in Relion. To create a mask for the particle subtraction, the capsid model was first 456 transformed to a density map using the molmap command in Chimera and then an inverted soft edged 457 mask was created from the density map using relion mask create with the --invert option. The 458 subtraction was followed by 3D classification (C1 symmetry), which generated the subset of 21,559 459 particles that was used for the final C1 refinement. The 3D classifications revealed clear density of an 460 outer layer, and an additional subtraction was therefore performed using a spherical mask of 100 Å 461 before the final refinement. The spherical mask was created using relion mask create with the --462 denovo and --outer radius options. The two maps (the capsid density created by molmap and the 463 circular map) were combined using Chimeras vop command before creating a new inverted mask 464 using relion mask create, which was used for subtraction before the final 3D refinement. To 465 reconstruct the core alone the spherical mask was used for the subtraction procedure.

466 Model building and refinement

467 The atomic model of CtenDNAV-II capsid protein was manually built into the density map using Coot 468 (Emsley and Cowtan, 2004). The model was further improved through cycles of real-space refinement 469 in PHENIX (Adams et al., 2010) with geometric and secondary structure restraints, and subsequent 470 manual corrections by Coot were carried out iteratively. Refinement statistics are summarized in 471 Table 1. Figures were generated using the programs UCSF Chimera (Pettersen et al., 2004) and UCSF 472 Chimera X (Goddard et al., 2018). Structural comparison of the CtenDNAV-II capsid protein was
473 carried out by the DALI server as a heuristic search against all structures (as of 2020-06-03) in the
474 PDB (Holm, 2020a).

475 **Data availability**

- 476 The atomic coordinates of CtenDNAV-II have been submitted to the Protein Data Bank under
- 477 accession no. 7NS0. The cryo-EM density maps of the CtenDNAV-II capsid and genome has been
- 478 deposited at the Electron Microscopy Data Bank under no. EMD-12554, 12555 and 12556,
- 479 respectively.

480 Acknowledgements

481 The data were collected at the Cryo-EM Swedish National Facility funded by the Knut and Alice482 Wallenberg, Erling Persson Family, and Kempe Foundations, SciLifeLab, Stockholm University and

483 Umeå University. We thank Julian Conrad and Dustin Morado for help with data collection. We also

- 484 want to thank Afonso Vieira for valuable discussions.
- 485

Funding was provided by the following agencies: Vetenskapsrådet (VR)/The Swedish Research Council (to K.O., grant no. 2018-03387), the Swedish Foundation for International Cooperation in Research and Higher Education (STINT) (to Janos Hajdu and K.O., grant no. JA2014-5721), FORMAS research grant from the Swedish Research Council for Environment, Agricultural Sciences, and Spatial Planning (to K.O., grant no. 2018-00421), the Royal Swedish Academy of Sciences (to K.O., grant no. BS2018-0053), and the Japan Society for the Promotion of Science KAKENHI (to Keizo Nagasaki, K.K., and Y.T., grant no. 16H06429, 16K21723, 16H06437, and 19H00956).

- 493
- 494 A.M., K.K., Y.T., and K.O. prepared the cryo-EM samples. A.M. and K.O. designed the experiments.
- 495 A.M. and K.O. collected the cryo-EM data and analysed the data. H.W performed the HI assay. Y.T.
- 496 performed the adsorption test. Y.H., K.Y., S.M. and K.K. performed the sequence analyses. A.M. and
- 497 K.O. wrote the manuscript. All of the authors discussed the results and proofread the manuscript.
- 498
- 499 We declare no competing financial interests.

References 500

- 501 Adams PD, Afonine PV, Bunkóczi G, Chen VB, Davis IW, Echols N, Headd JJ, Hung L-W, Kapral
- 502 GJ, Grosse-Kunstleve RW, McCoy AJ, Moriarty NW, Oeffner R, Read RJ, Richardson DC,
- 503 Richardson JS, Terwilliger TC, Zwart PH. 2010. PHENIX: a comprehensive Python-based system for
- 504 macromolecular structure solution. Acta Crystallogr D Biol Crystallogr 66:213–221.
- 505 doi:10.1107/S0907444909052925
- 506 Bratbak G, Egge JK, Heldal M. 1993. Viral mortality of the marine alga Emiliania huxlevi
- 507 (Haptophyceae) and termination of algal blooms. Mar Ecol Prog Ser 93:39-48.
- 508 Brussaard CPD. 2004. Viral Control of Phytoplankton Populations-a Review1. J Eukaryot Microbiol
- 509 51:125–138. doi:10.1111/j.1550-7408.2004.tb00537.x
- 510 Chiba Y, Tomaru Y, Shimabukuro H, Kimura K, Hirai M, Takaki Y, Hagiwara D, Nunoura T,
- 511 Urayama S. 2020. Viral RNA Genomes Identified from Marine Macroalgae and a Diatom. Microbes 512 Environ 35:n/a. doi:10.1264/jsme2.ME20016
- Coulibaly F, Chevalier C, Gutsche I, Pous J, Navaza J, Bressanelli S, Delmas B, Rey FA. 2005. The 513
- 514 Birnavirus Crystal Structure Reveals Structural Relationships among Icosahedral Viruses. Cell 515 120:761-772. doi:10.1016/j.cell.2005.01.009
- Dai X, Li Z, Lai M, Shu S, Du Y, Zhou ZH, Sun R. 2017. In situ structures of the genome and 516
- 517 genome-delivery apparatus in a single-stranded RNA virus. Nature 541:112-116.
- 518 doi:10.1038/nature20589
- 519 Diemer GS, Stedman KM. 2012. A novel virus genome discovered in an extreme environment
- 520 suggests recombination between unrelated groups of RNA and DNA viruses. Biol Direct 7:13. 521 doi:10.1186/1745-6150-7-13
- 522 Ding K, Celma CC, Zhang X, Chang T, Shen W, Atanasov I, Roy P, Zhou ZH. 2019. In situ structures

523 of rotavirus polymerase in action and mechanism of mRNA transcription and release. Nat Commun 524 **10**:2216. doi:10.1038/s41467-019-10236-7

- 525 Dolja VV, Koonin EV. 2018. Metagenomics reshapes the concepts of RNA virus evolution by
- 526 revealing extensive horizontal virus transfer. Virus Res 244:36-52. doi:10.1016/j.virusres.2017.10.020
- 527 Emsley P, Cowtan K. 2004. Coot: model-building tools for molecular graphics. Acta Crystallogr D
- 528 Biol Crystallogr 60:2126–2132. doi:10.1107/S0907444904019158
- 529 Fuhrman JA. 1999. Marine viruses and their biogeochemical and ecological effects. Nature 399:541-
- 530 548. doi:10.1038/21119
- 531 Fulton JM, Fredricks HF, Bidle KD, Vardi A, Kendrick BJ, DiTullio GR, Van Mooy BAS. 2014.
- 532 Novel molecular determinants of viral susceptibility and resistance in the lipidome of *E miliania*
- 533 huxleyi: Novel lipids in Emiliania huxleyi. Environ Microbiol 16:1137–1149. doi:10.1111/1462-534 2920.12358
- Garriga D, Querol-Audí J, Abaitua F, Saugar I, Pous J, Verdaguer N, Castón JR, Rodriguez JF. 2006. 535
- 536 The 2.6-Angstrom Structure of Infectious Bursal Disease Virus-Derived T=1 Particles Reveals New
- 537 Stabilizing Elements of the Virus Capsid. J Virol 80:6895-6905. doi:10.1128/JVI.00368-06
- 538 Goddard TD, Huang CC, Meng EC, Pettersen EF, Couch GS, Morris JH, Ferrin TE. 2018. UCSF
- 539 ChimeraX: Meeting modern challenges in visualization and analysis: UCSF ChimeraX Visualization 540 System. Protein Sci 27:14-25. doi:10.1002/pro.3235
- 541 Gorzelnik KV, Cui Z, Reed CA, Jakana J, Young R, Zhang J. 2016. Asymmetric cryo-EM structure of
- 542 the canonical *Allolevivirus* OB reveals a single maturation protein and the genomic ssRNA in situ. Proc Natl Acad Sci 113:11519-11524. doi:10.1073/pnas.1609482113 543
- 544 Harrison SC, Olson AJ, Schutt CE, Winkler FK, Bricogne G. 1978. Tomato bushy stunt virus at 2.9 Å
- 545 resolution. Nature 276:368-373. doi:10.1038/276368a0
- 546 Helgstrand C, Munshi S, Johnson JE, Liljas L. 2004. The refined structure of Nudaurelia capensis ω
- 547 Virus reveals control elements for a T = 4 capsid maturation. *Virology* **318**:192–203.
- 548 doi:10.1016/j.virol.2003.08.045
- 549 Henderson R, Sali A, Baker ML, Carragher B, Devkota B, Downing KH, Egelman EH, Feng Z, Frank
- 550 J, Grigorieff N, Jiang W, Ludtke SJ, Medalia O, Penczek PA, Rosenthal PB, Rossmann MG, Schmid
- 551 MF, Schröder GF, Steven AC, Stokes DL, Westbrook JD, Wriggers W, Yang H, Young J, Berman
- 552 HM, Chiu W, Kleywegt GJ, Lawson CL. 2012. Outcome of the First Electron Microscopy Validation
- 553 Task Force Meeting. Structure 20:205-214. doi:10.1016/j.str.2011.12.014

- Hesketh EL, Saunders K, Fisher C, Potze J, Stanley J, Lomonossoff GP, Ranson NA. 2018. The 3.3 Å
- structure of a plant geminivirus using cryo-EM. *Nat Commun* 9:2369. doi:10.1038/s41467-018-04793 6
- 557 Ho KL, Gabrielsen M, Beh PL, Kueh CL, Thong QX, Streetley J, Tan WS, Bhella D. 2018. Structure
- 558 of the Macrobrachium rosenbergii nodavirus: A new genus within the Nodaviridae? *PLOS Biol* 559 16:2000028 doi:10.1271/journal.nbia.2000028
- 559 **16**:e3000038. doi:10.1371/journal.pbio.3000038
- Holm L. 2020a. DALI and the persistence of protein shape. *Protein Sci* **29**:128–140.
- 561 doi:10.1002/pro.3749
- Holm L. 2020b. Using Dali for Protein Structure Comparison In: Gáspári Z, editor. Structural
- 563 Bioinformatics, Methods in Molecular Biology. New York, NY: Springer US. pp. 29–42.
- 564 doi:10.1007/978-1-0716-0270-6_3
- Hulo C, de Castro E, Masson P, Bougueleret L, Bairoch A, Xenarios I, Le Mercier P. 2011.
- ViralZone: a knowledge resource to understand virus diversity. *Nucleic Acids Res* 39:D576–D582.
 doi:10.1093/nar/gkq901
- 568 Ilca SL, Sun X, El Omari K, Kotecha A, de Haas F, DiMaio F, Grimes JM, Stuart DI, Poranen MM,
- Huiskonen JT. 2019. Multiple liquid crystalline geometries of highly compacted nucleic acid in a
 dsRNA virus. *Nature* 570:252–256. doi:10.1038/s41586-019-1229-9
- 571 Inoue T, Tsai B. 2011. A Large and Intact Viral Particle Penetrates the Endoplasmic Reticulum
- 572 Membrane to Reach the Cytosol. *PLoS Pathog* 7:e1002037. doi:10.1371/journal.ppat.1002037
- Johannessen T, Larsen A, Bratbak G, Pagarete A, Edvardsen B, Egge E, Sandaa R-A. 2017. Seasonal
- 574 Dynamics of Haptophytes and dsDNA Algal Viruses Suggest Complex Virus-Host Relationship.
 575 *Viruses* 9:84. doi:10.3390/v9040084
- Johnson JE, Tang L, Johnson KN, Ball LA, Lin T, Yeager M. 2001. The structure of Pariacoto virus
 reveals a dodecahedral cage of duplex RNA. *Nat Struct Biol* 8:77–83. doi:10.1038/83089
- 578 Kazlauskas D, Dayaram A, Kraberger S, Goldstien S, Varsani A, Krupovic M. 2017. Evolutionary
- history of ssDNA bacilladnaviruses features horizontal acquisition of the capsid gene from ssRNA
 nodaviruses. *Virology* 504:114–121. doi:10.1016/j.virol.2017.02.001
- 581 Kazlauskas D, Varsani A, Koonin EV, Krupovic M. 2019. Multiple origins of prokaryotic and
- 582 eukaryotic single-stranded DNA viruses from bacterial and archaeal plasmids. *Nat Commun* 10:3425.
 582 dai:10.1028/c41467.010.11422.0
- 583 doi:10.1038/s41467-019-11433-0
- 584 Kimura K, Tomaru Y. 2015. Discovery of Two Novel Viruses Expands the Diversity of Single-
- Stranded DNA and Single-Stranded RNA Viruses Infecting a Cosmopolitan Marine Diatom. *Appl Environ Microbiol* 81:1120–1131. doi:10.1128/AEM.02380-14
- 587 Koning RI, Gomez-Blanco J, Akopjana I, Vargas J, Kazaks A, Tars K, Carazo JM, Koster AJ. 2016.
- Asymmetric cryo-EM reconstruction of phage MS2 reveals genome structure in situ. *Nat Commun*7:12524. doi:10.1038/ncomms12524
- 590 Koonin EV, Dolja VV. 2014. Virus World as an Evolutionary Network of Viruses and Capsidless
- 591 Selfish Elements. *Microbiol Mol Biol Rev* 78:278–303. doi:10.1128/MMBR.00049-13
- 592 Koonin EV, Dolja VV, Krupovic M. 2015. Origins and evolution of viruses of eukaryotes: The
- 593 ultimate modularity. *Virology* **479–480**:2–25. doi:10.1016/j.virol.2015.02.039
- 594 Krupovic M. 2013. Networks of evolutionary interactions underlying the polyphyletic origin of
- 595 ssDNA viruses. *Curr Opin Virol* **3**:578–586. doi:10.1016/j.coviro.2013.06.010
- Krupovic M, Koonin EV. 2017. Multiple origins of viral capsid proteins from cellular ancestors. *Proc Natl Acad Sci U S A* 114:E2401–E2410. doi:10.1073/pnas.1621061114
- 597 Nati Acaa Sci U S A 114:E2401–E2410. doi:10.10/3/pnas.1621061114 E09 Krumenie M. Versenie A. Kerslevelees D. Breideert M. Delwert E. Beserie K. V
- 598 Krupovic M, Varsani A, Kazlauskas D, Breitbart M, Delwart E, Rosario K, Yutin N, Wolf YI,
- 599 Harrach B, Zerbini FM, Dolja VV, Kuhn JH, Koonin EV. 2020. Cressdnaviricota : a Virus Phylum
- 600 Unifying Seven Families of Rep-Encoding Viruses with Single-Stranded, Circular DNA Genomes. J
- 601 *Virol* 94:e00582-20, /jvi/94/12/JVI.00582-20.atom. doi:10.1128/JVI.00582-20
- 602 Kucukelbir A, Sigworth FJ, Tagare HD. 2014. Quantifying the local resolution of cryo-EM density
- 603 maps. *Nat Methods* 11:63–65. doi:10.1038/nmeth.2727
- Laanto E, Mäntynen S, De Colibus L, Marjakangas J, Gillum A, Stuart DI, Ravantti JJ, Huiskonen JT,
- Sundberg L-R. 2017. Virus found in a boreal lake links ssDNA and dsDNA viruses. *Proc Natl Acad*
- 606 *Sci* **114**:8378–8383. doi:10.1073/pnas.1703834114
- Liu H, Cheng L. 2015. Cryo-EM shows the polymerase structures and a nonspooled genome within a
- 608 dsRNA virus. *Science* **349**:1347–1350. doi:10.1126/science.aaa4938

- 609 Liu W, Hsu C-H, Hong Y-R, Wu S-C, Wang C-H, Wu Y-M, Chao C-B, Lin C-S. 2005. Early
- 610 endocytosis pathways in SSN-1 cells infected by dragon grouper nervous necrosis virus. J Gen Virol 86:2553-2561. doi:10.1099/vir.0.81021-0 611
- 612 Liu Y-T, Jih J, Dai X, Bi G-Q, Zhou ZH. 2019. Cryo-EM structures of herpes simplex virus type 1
- 613 portal vertex and packaged genome. Nature 570:257-261. doi:10.1038/s41586-019-1248-6
- Mallery DL, Márquez CL, McEwan WA, Dickson CF, Jacques DA, Anandapadamanaban M, Bichel 614
- 615 K, Towers GJ, Saiardi A, Böcking T, James LC. 2018. IP6 is an HIV pocket factor that prevents
- 616 capsid collapse and promotes DNA synthesis. eLife 7. doi:10.7554/eLife.35335
- 617 Moreno P, Souto S, Leiva-Rebollo R, Borrego JJ, Bandín I, Alonso MC. 2019. Capsid amino acids at
- 618 positions 247 and 270 are involved in the virulence of betanodaviruses to European sea bass. Sci Rep 619 **9**:14068. doi:10.1038/s41598-019-50622-1
- 620 Muhire BM, Golden M, Murrell B, Lefeuvre P, Lett J-M, Gray A, Poon AYF, Ngandu NK, Semegni
- 621 Y, Tanov EP, Monjane AL, Harkins GW, Varsani A, Shepherd DN, Martin DP. 2014. Evidence of
- 622 Pervasive Biologically Functional Secondary Structures within the Genomes of Eukaryotic Single-
- 623 Stranded DNA Viruses. J Virol 88:1972-1989. doi:10.1128/JVI.03031-13
- 624 Munke A, Kimura K, Tomaru Y, Okamoto K. 2020. Capsid Structure of a Marine Algal Virus of the
- 625 Order Picornavirales. J Virol 94:e01855-19, /jvi/94/9/JVI.01855-19.atom. doi:10.1128/JVI.01855-19
- 626 Nagasaki K, Shirai Y, Takao Y, Mizumoto H, Nishida K, Tomaru Y. 2005. Comparison of genome
- 627 sequences of single-stranded RNA viruses infecting the bivalve-killing dinoflagellate Heterocapsa
- 628 circularisquama. Appl Environ Microbiol 71:8888-8894. doi:10.1128/AEM.71.12.8888-8894.2005
- 629 Nagasaki K, Tarutani K, Yamaguchi M. 1999. Growth Characteristics of Heterosigma
- 630 akashiwo Virus and Its Possible Use as a Microbiological Agent for Red Tide Control. Appl 631 Environ Microbiol 65:898.
- 632 Ng TFF, Zhang W, Sachsenröder J, Kondov NO, da Costa AC, Vega E, Holtz LR, Wu G, Wang D,
- 633 Stine CO, Antonio M, Mulvaney US, Muench MO, Deng X, Ambert-Balay K, Pothier P, Vinjé J,
- 634 Delwart E. 2015. A diverse group of small circular ssDNA viral genomes in human and non-human 635
- primate stools. Virus Evol 1:vev017. doi:10.1093/ve/vev017
- 636 Nishihara T, Kurano N, Shinoda S. 1986. Calculation of Most Probable Number for Enumeration of
- 637 Bacteria on a Micro-Computer. Eisei Kagaku 32:226-228. doi:10.1248/jhs1956.32.226
- 638 Okamoto K, Ferreira RJ, Larsson DSD, Maia FRNC, Isawa H, Sawabe K, Murata K, Hajdu J, Iwasaki
- 639 K, Kasson PM, Miyazaki N. 2020. Acquired Functional Capsid Structures in Metazoan Totivirus-like
- 640 dsRNA Virus. Structure 28:888-896.e3. doi:10.1016/j.str.2020.04.016
- 641 Okamoto K, Miyazaki N, Larsson DSD, Kobayashi D, Svenda M, Mühlig K, Maia FRNC, Gunn LH,
- 642 Isawa H, Kobayashi M, Sawabe K, Murata K, Hajdu J. 2016. The infectious particle of insect-borne
- 643 totivirus-like Omono River virus has raised ridges and lacks fibre complexes. Sci Rep 6:33170. 644 doi:10.1038/srep33170
- 645 Penkler DL, Jiwaji M, Domitrovic T, Short JR, Johnson JE, Dorrington RA. 2016. Binding and entry
- 646 of a non-enveloped T =4 insect RNA virus is triggered by alkaline pH. Virology 498:277–287.
- 647 doi:10.1016/j.virol.2016.08.028
- 648 Pettersen EF, Goddard TD, Huang CC, Couch GS, Greenblatt DM, Meng EC, Ferrin TE. 2004. UCSF
- 649 Chimera? A visualization system for exploratory research and analysis. J Comput Chem 25:1605– 650 1612. doi:10.1002/jcc.20084
- 651 Phan TG, Mori D, Deng X, Rajindrajith S, Ranawaka U, Fan Ng TF, Bucardo-Rivera F, Orlandi P,
- 652 Ahmed K, Delwart E. 2015. Small circular single stranded DNA viral genomes in unexplained cases
- 653 of human encephalitis, diarrhea, and in untreated sewage. Virology 482:98-104.
- 654 doi:10.1016/j.virol.2015.03.011
- 655 Rossmann MG, Arnold E, Erickson JW, Frankenberger EA, Griffith JP, Hecht H-J, Johnson JE,
- 656 Kamer G, Luo M, Mosser AG, Rueckert RR, Sherry B, Vriend G. 1985. Structure of a human
- 657 common cold virus and functional relationship to other picornaviruses. *Nature* **317**:145–153.
- 658 doi:10.1038/317145a0
- 659 Scheres SHW. 2012. RELION: Implementation of a Bayesian approach to cryo-EM structure
- 660 determination. J Struct Biol 180:519-530. doi:10.1016/j.jsb.2012.09.006
- 661 Scheres SHW, Chen S. 2012. Prevention of overfitting in cryo-EM structure determination. Nat
- 662 Methods 9:853-854. doi:10.1038/nmeth.2115

- 663 Speir JA, Taylor DJ, Natarajan P, Pringle FM, Ball LA, Johnson JE. 2010. Evolution in Action: N and
- 664 C Termini of Subunits in Related T = 4 Viruses Exchange Roles as Molecular Switches. *Structure* 665 19:700, 700, doi:10.1016/j.ctr.2010.03.010
- 665 **18**:700–709. doi:10.1016/j.str.2010.03.010
- 666 Ströh LJ, Rustmeier NH, Blaum BS, Botsch J, Rößler P, Wedekink F, Lipkin WI, Mishra N, Stehle T.
- 667 2020. Structural Basis and Evolution of Glycan Receptor Specificities within the Polyomavirus
- 668 Family. *mBio* 11:e00745-20, /mbio/11/4/mBio.00745-20.atom. doi:10.1128/mBio.00745-20
- 669 Suttle CA. 2007. Marine viruses major players in the global ecosystem. *Nat Rev Microbiol* 5:801–
- 670 812. doi:10.1038/nrmicro1750
- 671 Suttle CA. 1993. Enumeration and isolation of virusesHandbook of Methods in Aquatic Microbial672 Ecology. Boca Raton: Lewis Publishers.
- 673 Tarutani K, Nagasaki K, Yamaguchi M. 2006. Virus adsorption process determines virus
- 674 susceptibility in Heterosigma akashiwo (Raphidophyceae). Aquat Microb Ecol 42:209–213.
- 675 doi:10.3354/ame042209
- 676 Tarutani K, Nagasaki K, Yamaguchi M. 2000. Viral Impacts on Total Abundance and Clonal
- 677 Composition of the Harmful Bloom-Forming PhytoplanktonHeterosigma akashiwo. *Appl Environ* 678 *Microbiol* 66:4916–4920. doi:10.1128/AEM.66.11.4916-4920.2000
- Tisza MJ, Pastrana DV, Welch NL, Stewart B, Peretti A, Starrett GJ, Pang Y-YS, Krishnamurthy SR,
- 680 Pesavento PA, McDermott DH, Murphy PM, Whited JL, Miller B, Brenchley J, Rosshart SP,
- 681 Rehermann B, Doorbar J, Ta'ala BA, Pletnikova O, Troncoso JC, Resnick SM, Bolduc B, Sullivan
- 682 MB, Varsani A, Segall AM, Buck CB. 2020. Discovery of several thousand highly diverse circular
- 683 DNA viruses. *eLife* **9**:e51971. doi:10.7554/eLife.51971
- Tomaru Y, Kimura K. 2016. Rapid quantification of viable cells of the planktonic diatom *Chaetoceros*
- tenuissimus and associated RNA viruses in culture. *Plankton Benthos Res* **11**:9–16.
- 686 doi:10.3800/pbr.11.9
- 687 Tomaru Y, Shirai Y, Toyoda K, Nagasaki K. 2011. Isolation and characterisation of a single-stranded
- 688 DNA virus infecting the marine planktonic diatom Chaetoceros tenuissimus. *Aquat Microb Ecol* 689 64:175–184. doi:10.3354/ame01517
- 690 Tomaru Y, Toyoda K, Suzuki H, Nagumo T, Kimura K, Takao Y. 2013. New single-stranded DNA
- virus with a unique genomic structure that infects marine diatom Chaetoceros setoensis. *Sci Rep* 3:3337. doi:10.1038/srep03337
- 693 Wagstaff BA, Rejzek M, Field RA. 2018. Identification of a Kdn biosynthesis pathway in the
- 694 haptophyte *Prymnesium parvum* suggests widespread sialic acid biosynthesis among microalgae. J
- 695 Biol Chem 293:16277–16290. doi:10.1074/jbc.RA118.004921
- 696 Wang F, Liu Y, Su Z, Osinski T, de Oliveira GAP, Conway JF, Schouten S, Krupovic M, Prangishvili
- 697 D, Egelman EH. 2019. A packing for A-form DNA in an icosahedral virus. Proc Natl Acad Sci
- 698 **116**:22591–22597. doi:10.1073/pnas.1908242116
- Wang X, Zhang F, Su R, Li X, Chen W, Chen Q, Yang T, Wang J, Liu H, Fang Q, Cheng L. 2018.
- 700 Structure of RNA polymerase complex and genome within a dsRNA virus provides insights into the
- 701 mechanisms of transcription and assembly. *Proc Natl Acad Sci* **115**:7344–7349.
- 702 doi:10.1073/pnas.1803885115
- Wilhelm SW, Suttle CA. 1999. Viruses and Nutrient Cycles in the Sea. *BioScience* 49:781–788.
 doi:10.2307/1313569
- 705 Wolf YI, Silas S, Wang Y, Wu S, Bocek M, Kazlauskas D, Krupovic M, Fire A, Dolja VV, Koonin
- EV. 2020. Doubling of the known set of RNA viruses by metagenomic analysis of an aquatic virome.
 Nat Microbiol 5:1262–1270. doi:10.1038/s41564-020-0755-4
- 708 Zhang K. 2016. Gctf: Real-time CTF determination and correction. *J Struct Biol* **193**:1–12.
- 709 doi:10.1016/j.jsb.2015.11.003
- 710 Zhang X, Ding K, Yu X, Chang W, Sun J, Hong Zhou Z. 2015. In situ structures of the segmented
- 711 genome and RNA polymerase complex inside a dsRNA virus. *Nature* **527**:531–534.
- 712 doi:10.1038/nature15767
- 713 Zheng SQ, Palovcak E, Armache J-P, Verba KA, Cheng Y, Agard DA. 2017. MotionCor2: anisotropic
- correction of beam-induced motion for improved cryo-electron microscopy. *Nat Methods* 14:331–332.
- 715 doi:10.1038/nmeth.4193

- 716 Zivanov J, Nakane T, Forsberg BO, Kimanius D, Hagen WJ, Lindahl E, Scheres SH. 2018. New tools
- for automated high-resolution cryo-EM structure determination in RELION-3. *eLife* 7:e42166.
- 718 doi:10.7554/eLife.42166
- 719 Zocher G, Mistry N, Frank M, Hähnlein-Schick I, Ekström J-O, Arnberg N, Stehle T. 2014. A Sialic
- 720 Acid Binding Site in a Human Picornavirus. *PLoS Pathog* **10**:e1004401.
- 721 doi:10.1371/journal.ppat.1004401



724

725 Supplementary Figure 1. Data collection and reconstruction of the capsid. (A) Cryo-EM raw 726 image of CtenDNAV-II. (B) Cryo-EM 3D reconstruction of the CtenDNAV-II capsid. The capsid is 727 viewed down the five-fold axis and radially coloured from blue to red. The icosahedral five-fold, 728 three-fold, and two-fold axes are labelled as 5, 3, and 2, respectively. (C) The gold standard FSC 729 resolution curves of masked (blue) and unmasked (green) reconstructions of the CtenDNAV-II capsid. 730 Possible effects of the masking were compensated for by noise randomization (red), to create the final 731 FSC curve (black). The resolution at which the correlation drops below the FSC = 0.143 (gold 732 standard threshold (Henderson et al., 2012; Scheres and Chen, 2012) is 2.3 Å. (D) Local resolution of 733 the final reconstruction determined by Relion. The left panel shows a histogram of the local resolution, 734 and the middle and right panel shows the reconstruction coloured from red to white to blue that 735 corresponds to local resolutions 2.6, 2.4 and 2.2, respectively. The icosahedral five-fold, three-fold, 736 and two-fold axes are labelled as 5, 3, and 2, respectively in the middle panel. The front half of the 737 reconstruction is removed in the right most panel to visualize the inside of the capsid. 738









contour level. Unmodelled densities around the three- and five-fold axes are indicated with blue

arrows and circles, which correspond to the C-terminal ends of subunit A and B. The capsid is viewed

down the five-fold axis and radially coloured from blue to red. The positions of the five-, three-, and

two-fold axes are shown by a white pentamer, triangle, and ellipse, respectively.

749

750 Supplementary Figure 3. Unmodelled density is visualized on the inside of the capsid below the

- C subunit. (A) Inside view of the cryo-EM reconstruction coloured based on the atomic model of the
 three subunits. By decreasing the contour level, unmodelled densities are visualized in grey. (B)
- 753 Close-up view of the C subunit, where the reconstruction is transparent to reveal the model in ribbon
- representation. The last modelled residues in the C-termini of the two 2-fold symmetry related C
- subunits are indicated with black arrows.





Supplementary Figure 4. Putative ion located in the subunit interface. The model of a single 758 icosahedral protomer viewed from the inside of the capsid and corresponding cryo-EM map is 759 visualized to the left. To the right, is a close-up view of the subunit interface, displaying density in the 760 centre possibly originating from an ion and the surrounding arginine residues.



Supplementary Figure 5. Superimposition performed by the DALI server of the CtenDNAV-II 763 764 capsid protein against corresponding capsid proteins from viruses in Fig. 3B. The figures show 765 CtenDNAV-II subunit A coloured according to (A) structural conservation and (B) sequence 766 conservation where conservation mapping goes from blue for the highest values through green to red 767 for the lowest values. (A) The three areas where structural conservation has the lowest values; the C-768 terminal tail, β-sheet projection, and jelly-roll loops C'-C" and H-I are highlighted in green, magenta 769 and cyan dashed lines. (C) The entire capsid rendered with a surface representation viewed down an 770 icosahedral two-fold axis. The amino acids with lowest structural conservation (red in panel A) are 771 coloured green, magenta and cyan that corresponds to the areas N-terminal tail, β-sheet projection and 772 jelly-roll loops labelled in (A). The insert is zooming in on one of the five-folds and highlights one A-773 subunit and the C-terminus (green) coming from the B-subunit in an adjacent five-fold related 774 protomer.

Strain		PDGRMLESLP	RRCQLVTEIR	NNVTVGSNPT	YILVAPSLGL	AFQAYQDTNV	PGGLDSSVYG
SS10V-8V							
SS10V-35V							
2-6_11V10							
2-6_11V12							
2-6_10V07							
2-6_10V05							
2-6_10V17							
2-6_10V20							
SS10V-24V							
2-6_10V27							
2-6_10V33							
2-6_11V04							
2-6_11V08							
2-6_11V06							
2-6_11V17							
	1						
Strain	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
SS10V-8V	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
Strain SS10V-8V SS10V-35V	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
Strain SS10V-8V SS10V-35V 2-6_11V10	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV 	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07 2-6_10V05	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
StrainSS10V-8VSS10V-35V2-6_11V102-6_11V122-6_10V072-6_10V052-6_10V17	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV 	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07 2-6_10V05 2-6_10V17 2-6_10V20	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV 	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
StrainSS10V-8VSS10V-35V2-6_11V102-6_10V072-6_10V072-6_10V052-6_10V172-6_10V20SS10V-24V	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV 	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
StrainSS10V-8VSS10V-35V2-6_11V102-6_10V072-6_10V072-6_10V172-6_10V172-6_10V20SS10V-24V2-6_10V27	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
StrainSS10V-8VSS10V-35V2-6_11V102-6_10V072-6_10V052-6_10V172-6_10V20SS10V-24V2-6_10V272-6_10V33	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
StrainSS10V-8VSS10V-35V2-6_11V102-6_10V072-6_10V052-6_10V172-6_10V20SS10V-24V2-6_10V272-6_10V332-6_11V04	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV 	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
StrainSS10V-8VSS10V-35V2-6_11V102-6_10V072-6_10V052-6_10V172-6_10V20SS10V-24V2-6_10V272-6_10V332-6_11V042-6_11V08	LQNRGCTVRA	NLSATSIENY I	NDIAKWRIVS	QGINLKLNNV	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_10V07 2-6_10V05 2-6_10V17 2-6_10V20 SS10V-24V 2-6_10V33 2-6_11V04 2-6_11V08	LQNRGCTVRA	NLSATSIENY	NDIAKWRIVS	QGINLKLNNV	EDENDGWYEA	CRFQHDWTPD	ELCLRSTEND

Strain	ASTISQDEDL	VMGVISSSFL	NGALNTIGNN	MVEQRGYESG	LLKNIHKRMF	QLHNNTSAIR E	PKTLQGQFNY
SS10V-8V		M	1111111111	1111111111			
SS10V-35V		<mark> </mark>				1111111111	
2-6_11V10						1111111111	
2-6_11V12						1111111111	
2-6_10V07						1111111111	
2-6_10V05						1111111111	
2-6_10V17	TNN					1111111111	
2-6_10V20						1111111111	
SS10V-24V	T					1111111111	
2-6_10V27						1111111111	
2-6_10V33						1111111111	
2-6_11V04						1111111111	
2-6_11V08	NS						
2-6_11V06	NS					1111111111	
2-6_11V17						1111111111	
	1						
Strain	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I	IQNLELQYSP
<u>Strain</u> SS10V-8V	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I	IQNLELQYSP
SS10V-8V SS10V-35V	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I	IQNLELQYSP
Strain SS10V-8V SS10V-35V 2-6_11V10	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I	IQNLELQYSP
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I IIIIIIIIIIII IIIIIIIIIIII R AKRPHCKR Y R IIIIIIIIII	QNLELQYSP
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I I R AKRPHCKR Y R I	IQNLELQYSP
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07 2-6_10V05	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	IQNLELQYSP SKFGAAIFS
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07 2-6_10V05 2-6_10V17	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I 	IQNLELQYSP SKFGAAIFS
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07 2-6_10V05 2-6_10V17 2-6_10V20	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I R AKRPHCKR Y R 	QNLELQYSP (SKFGAAIFS
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07 2-6_10V05 2-6_10V17 2-6_10V20 SS10V-24V	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	IQNLELQYSP SKFGAAIFS
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07 2-6_10V05 2-6_10V17 2-6_10V20 SS10V-24V 2-6_10V27	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I R AKRPHCKR Y R 	IQNLELQYSP SKFGAAIFS
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07 2-6_10V05 2-6_10V17 2-6_10V20 SS10V-24V 2-6_10V27 2-6_10V33	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I R AKRPHCKR Y R 	IQNLELQYSP SKFGAAIFS
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07 2-6_10V05 2-6_10V17 2-6_10V20 SS10V-24V 2-6_10V27 2-6_10V33 2-6_11V04	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I 	QNLELQYSP
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07 2-6_10V05 2-6_10V17 2-6_10V20 SS10V-24V 2-6_10V27 2-6_10V33 2-6_11V04 2-6_11V08	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I 	QNLELQYSP
Strain SS10V-8V SS10V-35V 2-6_11V10 2-6_11V12 2-6_10V07 2-6_10V05 2-6_10V17 2-6_10V20 SS10V-24V 2-6_10V27 2-6_10V33 2-6_11V04 2-6_11V08 2-6_11V06	GSEITFSGTE	SEARFTDVPS	NRQLVDSLWH	NDYDCILIKL	YPRENTGAAG	QTGSALIVNA I IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	QNLELQYSP (SKFGAAIFS

Strain	TSDLSTYHIA NKRARMVEAK LDKKNNTDAA GEPFVPGSSR*
SS10V-8V	
SS10V-35V	
2-6_11V10	N*
2-6_11V12	
2-6_10V07	
2-6_10V05	
2-6_10V17	P T*
2-6_10V20	
SS10V-24V	
2-6_10V27	
2-6_10V33	
2-6_11V04	N*
2-6_11V08	
2-6_11V06	
2-6_11V17	

782 Supplementary Figure 6. Amino acid alignment of capsid proteins from 15 CtenDNAV-II strains. The consensus sequence is displayed in grey and the 783 mutations in colour. The amino acids are organised in the same scheme as in Figure 2 with divisions into 70 residues per line, which are further subdivided 784 into blocks of 10 residues by spaces within the sequence. The N-termini were not successfully sequenced and the first residue is therefore P81. This paper 785 describes the structure of strain SS10V-8V.



787

Supplementary Figure 7. Hemagglutination inhibition assay. Twofold serially diluted 1 mg/mL
BSA, influenza virus (flu), CtenDNAV-II and CtenRNV-II are tested for HI activity. The BSA and flu

were used for negative and positive controls for the assay.

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Supplementary Fig. 8. Adsorption tests between viruses and diatoms. The results are shown in 795 slope of free virus titer in a culture for 2h (see materials and methods). Significant differences were 796 detected in CtenDNAV-II vs. NIES-3714, and CtenRNAV-II vs. NIES-3715 (t-test, p<0.05, n=5). Their adsorption coefficients were 1.4±0.9×10⁻⁹ mL min⁻¹ and 2.7±1.1×10⁻⁹ mL min⁻¹, respectively. 797 798 NIES-3713: Chaetoceros socialis f. radians; NIES-3714: Chaetoceros tenuissimus; NIES-3715: 799 Chaetoceros tenuissimus.

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801 **Supplementary Figure 9. Connections between the outer genome layer and the core.** The outer 802 genome layer where the sites that connect to the core are highlighted in magenta. The connections 803 between the outer genome layer and the core seem to be confined to two specific areas on 804 approximately opposite sides. One area is shown in the left panel and the second is shown to the right, 805 which is rotated 180°.

806

807 Supplementary methods

808 Sequencing of viral capsid protein gene

809 Two hundred µL of C. tenuissimus NIES-3715 strain culture was added to each well of a 96-well 810 culture plate, and 10 uL of virus strains (SS10V-8V, SS10V-24V, SS10V-35V, 2-6 10V-05, 2-811 6 10V-07, 2-6 10V-17, 2-6 10V-20, 2-6 10V-27, 2-6 10V-33, 2-6 11V-04, 2-6 11V-06, 2-6 11V-812 08, 2-6 11V-10, 2-6 11V-12, 2-6 11V-17) isolated from C. tenuissimus were added to each separate well. Diatom cultures with viral solution were grown at 15 °C in modified SWM3 medium (Kimura 813 814 and Tomaru, 2015) under a 12/12-h light-dark cycle of ca. 110-150 mmol of photons / m2 / s1 by 815 using white fluorescent illumination. After 24 hours of viral inoculation, the cultures were transferred 816 to microtubes and the cells were pelleted by centrifugation. One hundred uL of TE were added to each 817 cell pellet, and they were heated at 100 °C for 10 min. These solutions were used as DNA samples. 818 The PCR reaction was conducted as follows: 1 cycle at 94 °C for 2 min, 25 cycles each at 94 °C for 30 819 s, 60 °C for 30 s, 72 °C for 1 min, and 1 cycle at 72 °C for 10 min. The primer set of middle part of the 820 (P31 CtDV-Ari-F2: 5'-AGCAACCACAAATCCCCG-3'; P32 CtDV-Ari-R1: 5'gene 821 ATTTTGAATAGCGTTTACAATGAG-3') and the primer set of last part of the gene 822 (CtD2V VP2inF1: 5'-AATACTAGTGCCATTCGTCC-3', CtD2V VP2outR2: 5'-823 ACTACCATCATGATTGAGACTG-3'), they detect the gene for the viral capsid protein were used. 824 Each PCR product was purified with the ISOSPIN PCR Product (Nippon gene, Tokyo) and sequenced 825 on an ABI 3130 DNA Analyzer using the Sanger dideoxy method. Each DNA sequence was 826 combined and converted to amino acid sequences. The partial capsid protein amino acid sequences of 827 each virus were automatically aligned using Clustal W.

828 Hemagglutination inhibition (HI) assay

829 The HI test was performed according to classical procedures. The purified BSA, influenza virus (flu), 830 CtenDNAV-II and CtenRNAV-II were adjusted to 1 mg/mL in PBS. Twofold serial dilutions of these 831 samples were prepared for the assay, of which 25 μ L of each dilution was incubated with 50 μ L of 125 832 times PBS-diluted red blood cells from sheep blood (Håtunalab). The assay was conducted in a V-833 bottom, 96-well plate. The plate was incubated for 1 hr at RT, and used for observing the 834 hemagglutination of the red blood cells.

835 Adsorption test

836 Host diatom

- 837 The virus adsorption kinetics of CtenDNAV-II and CtenRNAV-II was examined using the following
- 838 diatom strains: Chaetoceros socialis f. radians NIES3713, C. tenuissimus NIES-3714 and C.
- 839 *tenuissimus* NIES-3715. Previously reported compatibilities between host diatoms and viruses based 840 on (Kimura and Tomaru, 2015) are as following:

	CtenDNAV-II	CtenRNAV-II
C. socialis f. radians NIES3713 (L-4)	Not lysed	Lysed
C. tenuissimus NIES-3714 (2-6)	Lysed*	Not lysed
C. tenuissimus NIES-3715 (2-10)	Lysed	Lysed*

841 *original host-virus combination

842 Viral inocula

- 843 Exponentially growing cultures of C. tenuissimus NIES-3715 were inoculated with CtenDNAV-II and
- 844 CtenRNAV-II (0.1% v v⁻¹), which were stored at 4°C in the dark, and incubated for 7 d under the
- growth conditions described above. Lysates were passed through a 0.2-µm polycarbonate membrane

filter (Whatman[®] Nuclepore Track-Etched Membranes, Merck KGaA, Darmstadt, Germany) to
remove cellular debris and stored at 4°C. Filtered lysates were used as experimental inocula.

848 Virus enumeration

849 The number of infectious viral units was determined using the extinction dilution method (Suttle, 1993). Briefly, stored filtrates were rapidly thawed at 25°C, and then diluted with SWM3 medium in a 850 series of 10-fold dilution steps. Aliguots (100 µL) of each dilution were added to 8 wells in cell-851 852 culture plates with 96 flat-bottom wells and mixed with 150 µL of an exponentially growing host 853 algae culture, C. tenuissimus NIES-3715. The cell culture plates were incubated at 20°C under a 12:12 854 h light:dark cycle of 130–150 μ mol photons \cdot m⁻²s⁻¹ (cool white fluorescent illumination) and cultures 855 were monitored using an optical microscope (Nikon Ti, Tokyo, Japan) for up to 14 d for culture lysis. 856 Culture lysis due to viral infection was usually observed as a complete degradation of the host cells in 857 a well. Virus abundance was calculated from the number of wells in which algal lysis occurred using a 858 BASIC programme as previously described (Nishihara et al., 1986).

859 Adsorption test

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860 The host cell cultures were grown in SWM3 medium under a 12:12 h light:dark cycle of approximately 400 μ mol of photons m⁻²s⁻¹ using white LED illumination at 25°C for *C. tenuissimus* 861 862 (NIES-3714 and -3715) and 20°C for C. socialis f. radians (NIES-3713). Cell counts were performed immediately with no fixation using Fuchs-Rosenthal hemocytometer or an image-based cytometer 863 864 (Tali[®] Image Cytometer, Thermo Fisher Scientific Ltd., Waltham, MA, USA) (Tomaru and Kimura, 2016). When the cultures reached stationary phase, cell concentration of $>10^6$ cells mL⁻¹, 20 mL of the 865 866 culture was poured into a sterilized polystyrene flask (n=5). An aliquot of the culture was passed 867 through a 0.2µm pore sized filter to remove the diatom cells. The resultant filtrate was poured into the 868 polystyrene flask as a control (n=5). The virus suspensions were inoculated into these cell-suspended 869 and filtrate cultures at 0.1%v/v (20µL). The virus inoculations were conducted during 1h and 2h after the light cycle started. The final virus titers in the cultures were set at $10^5 \sim 10^6$ infectious units mL⁻¹ 870 871 (multiplicity of infections = $10^{-1} \sim 10^{0}$).

Aliquots of the cultures were sampled at just after virus inoculations (0h), 1h and 2h of post
inoculation. The samples were used for cell counts and unadsorbed virus enumeration. The samples
used for viral titer estimations (infectious units mL⁻¹) were passed through 0.2-μm filters (DISMIC[®]25CS020AS, Advantec, Tokyo, Japan) to remove cellular debris. Filtrates were stored at -80°C until
analysis.

The adsorption coefficient (Cd; mL min⁻¹) was determined as follows (Tarutani et al., 2006):

$$C_d = \frac{\alpha - \alpha_C}{N}$$

879 where α (min⁻¹) and α_c (min⁻¹) is the slope determined by linear regression for the natural logarithm of 880 the percentage of free viruses in diatom cell culture and control culture against sampling time, 881 respectively. N (cells mL⁻¹) is the host cell number.