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# Past and present giant viruses diversity explored through permafrost metagenomics

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### Past and present giant viruses diversity

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### 9 Abstract

10 Giant viruses are abundant in aquatic environments and ecologically important 11 through the metabolic reprogramming of their hosts. Lesser is known about giant 12 viruses from soil although two of them, belonging to different families, were reactivated 13 from 30,000-y-old Siberian permafrost samples, suggesting an untapped diversity of Nucleocytoviricota in this environment. Through permafrost metagenomics, we reveal 14 15 a high heterogeneity in the abundance of giant viruses representing up to 12% of the 16 total community in one sample. *Pithoviridae* and *Orpheoviridae*-like viruses were the 17 most important contributors, followed by Mimiviridae. A comparison to other terrestrial 18 metagenomes confirmed that the diversity pattern in these samples is quite unique. In 19 contrast, *Pandoraviridae* sequences remained scarce. Using a stringent methodology, we were able to assemble large genomes, including a complete circular 1.6 Mb 20

*Pithoviridae*-like from a 42,000-y-old sample. The uncovered *Pithoviridae* diversity also provided insights into the family evolution. The phylogenetic reconstruction of specific functions not only revealed gene transfers between cells and viruses, but also between viruses from different families. Finally, the extensive annotation of the permafrost viral sequences revealed a patchwork of predicted functions amidst an even larger reservoir of anonymous genes of unknown functions.

### 28 Introduction

29 The permafrost, soil remaining continuously frozen for at least 2 years, covers 15% of 30 the Northern hemisphere (1) and gathers complex communities of living organisms and variable soil types. The microbial community of the surface cryosol is in some 31 cases subject to freezing and thawing every year (2) whereas communities from 32 33 deeper layers are trapped in the syngenetic (as old as the sediment) or epigenetic 34 (more recent than the sediment) permafrost. Pleistocene permafrost has been showed to harbor up to  $5x10^7$  cells per wet gram of soil of which about a fifth is alive (3). The 35 36 permafrost has thus the ability to preserve organisms for tens if not hundreds 37 thousands of years and acts as a huge reservoir of ancient microorganisms. It has 38 been shown for instance that numerous bacteria isolated from permafrost samples 39 remained viable (4, 5), even potentially up to 1.1 million years (6). Even in low 40 biomass-containing frozen environments such as glacier ice, metagenomics approaches have recently revealed hundreds of distinct bacterial genera (7). 41 42 Unicellular (8–10) and even multicellular (11, 12) eukaryotes can also be preserved 43 for thousands of years and be revived from such frozen environments.

44 Besides cellular organisms, metagenomics studies have revealed bacteriophages 45 communities archived in surface (13) or deeper (7) glacier ice, the majority of which 46 being taxonomically unassigned. Following the high bacterial abundance (14), bacteriophages are expected to be the most abundant viruses in the permafrost. 47 However, in the unfiltered size fraction, the eukaryotic viruses Nucleocytoviricota 48 49 (formerly known as Nucleocytoplasmic large DNA viruses or NCLDVs) are also highly 50 represented (14). This phylum gathers large double stranded DNA viruses such as 51 Pokkesviricetes (*Poxviridae* and *Asfarviridae*) as well as all the known giant viruses

52 (i.e. viruses visible by light microscopy): the Megaviricetes (Phycodnaviridae, Mimiviridae and Pimascovirales). Likewise, a handful of scaffolds of potential 53 54 *Phycodnaviridae*, also belonging to this phylum, were identified in a metagenomic study of glacial environments (13). More importantly, among Nucleocytoviricota, two 55 56 giant viruses, namely Pithovirus sibericum and Mollivirus sibericum, were reactivated 57 from a 30,000-y-old permafrost sample on Acanthamoeba castellanii (15, 16). Together with the presence of numerous protists and in particular amoeba in 58 permafrost (9), this hints at the existence of many more giant viruses in such 59 60 environments.

61 Recently, several studies specifically targeting the viral dark matter from 62 environmental metagenomics data have started to grasp the diversity and gene-63 content of the Nucleocytoviricota (17–19). It became clear that the genomes of these viruses code for various auxiliary metabolic genes, making them capable of 64 65 reprogramming their host's metabolism and hence are potentially important drivers of 66 global biogeochemical cycles (17, 18, 20). They also seem to be widespread in aquatic 67 environments. More specifically, Mimiviridae (in particular the proposed 68 Mesomimivirinae sub-family (21)) and Phycodnaviridae are major contributors of the 69 marine viromes all over the world, as revealed by thousands of metagenome-70 assembled viral genome (MAG) sequences (17-19). They also have been found active at the surface layer of the ocean by metatranscriptomics (22). The 71 72 Nucleocytoviricota ecological functions and diversity in terrestrial samples on the other hand is far less known, with the exception of *Klosneuvirinae* sequences recovered 73 74 from forest soil samples (23) and of *Pithoviridae* sequences assembled from the Loki's castle deep sea sediments sequences (24). The overwhelming proportion of marine-75

related as compared to terrestrial Nucleocytoviricota sequences from metagenomic studies is most likely due to the difficulty at revealing their hidden diversity in these environments (23). Soils host highly complex microbial communities making metagenomic studies notoriously challenging as population heterogeneity with closely related strains can hamper sequence assembly (25, 26).

81 Current giant viruses' metagenomic studies rely on the detection of Nucleocytoviricota 82 core genes (17, 18, 23, 24). However, among the very few shared genes some are 83 highly divergent or even completely absent from certain viral families. For instance, a 84 packaging ATPase, presumably encoded by a "core" gene in large DNA viruses, is absent in *Pithoviridae* (27). Likewise, the Major Capsid Protein (MCP) often used as a 85 86 marker gene to detect Nucleocytoviricota within metagenomic assemblies (18) is only 87 present in a divergent form in *Pithoviridae* (15) and completely absent from 88 *Pandoraviridae* (27, 28). Thus, the probability to detect these types of non-icosahedral 89 giant viruses is drastically lowered.

90 Although two distinct non-icosahedral giant viruses were initially isolated from 91 permafrost samples (15, 16) little is known on the Nucleocytoviricota diversity in this 92 type of environment. Here we propose an analysis of these viruses from eleven 93 permafrost samples ranging from the active layer up to 49,000-y-old (14). We show that the permafrost is a great source of viral diversity. Although the samples are very 94 95 heterogeneous in Nucleocytoviricota content, they reach up to 5% of the assembled 96 sequences and 12% of the total coverage in one deep permafrost sample. We found 97 here that *Pithoviridae* and *Orpheoviridae*-like families as well as *Mimiviridae* are the 98 main contributors of the giant virus diversity of the deep permafrost.

### 99 **Results**

### 100 Cryosol metagenomes assemblies

We gathered permafrost and surface cryosol raw metagenomic data produced by (14) on the three surface samples from Kamchatka (C-D-E, Table S1) that are also the samples from which Cedratvirus kamchatka (29) and Mollivirus kamchatka (30) were isolated, and on eight deep samples from the Yukechi Alas area radio-carbon-dated from 53 to over 49,000-y-old, seven of which are syngenetic (Table S1).

106 We first performed an assembly of the reads (Table S2) followed by binning and 107 assessed the quality of the bins, mostly composed of prokaryotic sequences (90%), 108 using Checkm (32). This revealed potential chimeras (Fig. S1A). We thus chose not 109 to consider them as unique organisms but instead used binning as a procedure to 110 decrease complexity in our datasets. The reads were first separated according to the 111 bin they belonged to and a second *de novo* assembly was made within each bin. This resulted in significantly longer scaffolds and a larger total assembly (Table S2). 112 113 Applying Checkm to this final dataset identified nearly no chimera (Fig. S1B). Thus, our method significantly gained in reliability by lowering the proportion of chimeras in 114 115 comparison to conventional binning.

### **Discriminating Nucleocytoviricota in metagenomic samples**

From this dataset we then sought to extract Nucleocytoviricota sequences. Our method is based on the detection of both Nucleocytoviricota genes (including the ones specific to the non-icosahedral *Pithoviridae* and *Pandoraviridae*) and cellular ones. Clearly the combination of the two showed a very distinct pattern for Nucleocytoviricota compared to cellular genomic sequences (Fig. 1A), as revealed by a control metagenomic mimicking database containing reference Nucleocytoviricota genomes 123 from (31), cellular genomes randomly sampled from Genbank in addition to amoeba and algae genomes (known to be the hosts of Nucleocytoviricota) as well as amoeba-124 125 hosted intracellular bacteria (Babela massiliensis Parachlamydia and acanthamoebae). The control database was also used to find the optimal parameters 126 discriminating Nucleocytoviricota sequences (slope = 0.1, intercept = 1; Fig. 1), 127 yielding high classification performance (sensitivity = 98.16% and specificity  $\geq 99.53\%$ ; 128 129 Fig. S2). For comparison we also tested the Viralrecall tool (35) that confirmed 1848 out of the 1973 (94%) scaffolds we detected. Finally, further controls for contamination 130 131 in the Nucleocytoviricota dataset involved a search for ribosomal sequences, none of which were found. Manual functional annotation also allowed the identification of 7 132 scaffolds potentially belonging to intracellular bacteria, a phage and a nudivirus that 133 134 were removed. At the end, our Nucleocytoviricota identification method on the 135 permafrost dataset resulted in 1966 scaffolds ranging from 10 kb up to 1.6 Mb, corresponding to 1% of all scaffolds over 10 kb in size (Fig. 1B). 136



#### 139 Figure 1: Extraction method of viral scaffolds

138

140 Each point corresponds to one scaffold. Viral matches (y-axis) were counted as the number of ORFs 141 matching a Nucleocytoviricota-specific HMM. These HMMs come from a previous study (18) to which 142 were added specific HMMs from the VOG database and HMMs constructed on Pandoraviridae and 143 Pithoviridae genomes. Cellular matches (x-axis) are the number of Diamond blastP matches against 144 the cellular Refseq database with a threshold of 35% of sequence identity. The dashed lines represent 145 the chosen threshold excluding all point under or on the line. (A) Control dataset. The inset is a zoom 146 of the bottom-left corner of the plot. For clarity, 1 bacterial point with over 1000 cellular matches and 1 147 viral match are not shown. (B) Permafrost data. For clarity, 5 points with over 1000 cellular matches are 148 not shown.

As said, Nucleocytoviricota metagenomic studies often rely on the MCP as a bait, making it hard, if not impossible, to catch some of the non-icosahedral viruses. By adding *Pithoviridae* and *Pandoraviridae* HMMs to the original profiles (18) and VOG's HMMs, we gained 5% (n = 110) more scaffolds that were mainly unclassified or from *Pithoviridae* and divergent *Pithoviridae* families (see further for phylogenies).

### 154 Large viral genomes from deep permafrost

155 Although our strategy to exclude conventional binning was primarily designed to 156 capture high confidence MAGs at the price of completeness, we were still able to 157 recover large Nucleocytoviricota genomes in single scaffolds with no apparent 158 chimera. Eight of them, assembled from 16m to 19m deep permafrost samples (R, N 159 and M, Table S1) dating from 42,000 to 49,000 years, reached over 500kb (Fig. 2). 160 The largest one of 1.6 Mb, referred to as Hydrivirus alas, is most likely complete as it was successfully circularized. Although these large scaffolds are deeply sequenced 161 162 (with an average coverage in between 14 and 72), they are far from belonging to the most abundant viruses in their samples (the highest coverages are of 53, 181 and 163 164 1572 in samples M, N, R respectively).



# Figure 2: Gene content of the large genomes recovered from ancientpermafrost samples

For each genome, the position of ORFans (ORFs with no match in the NR database), cellular and viral matches are recorded along the genome. The positions of tRNAs are also showed as red arrows. The pie charts present the proportion and taxonomy of viral matches with slices ≥ 5% labeled. The environmental *Pithoviridae/Orpheoviridae*-like category contains metagenomic sequences from (23, 24). The Hydrivirus alas genome was circularized.

174 These MAGs vary in divergence from known genomes, having from 22% up to 72% of ORFans for Staryvirus yedoma (Fig. 2). As always for newly discovered giant 175 176 viruses, their genomes also match cellular genes from all domains of life (with very few Archaea). When looking at the viral matches, two scaffolds seem close to 177 Megamimivirinae (Pleistovirus alas and Yakustkivirus alas), one to Klosneuvirinae 178 (Marosvirus alas) and four to Pithoviridae/Orpheoviridae (Hydrivirus alas, Stamnovirus 179 180 alas, Krossosvirus alas and Pelikevirus alas). The most divergent, Staryvirus yedoma, shows an even distribution of viral best BlastP matches with no specific family standing 181 182 out (Fig 2). Together with its high ORFan content, this suggests that it belongs to a Nucleocytoviricota viral family with no previous isolate so far. 183

184 The complete 1.6 Mb Hydrivirus alas genome reaches a size similar to the isolated Orpheovirus (32). The other 715 to 855 kb scaffolds (Stamnovirus alas, Krossosvirus 185 186 alas and Pelikevirus alas) are slightly larger than isolated *Pithoviridae* (ca. 600 kb) (15. 187 33, 34). However they were not circularized as expected for a *Pithoviridae* genome structure (15) and are potentially even larger. Still, in the four of them, most of the core 188 189 genes are present (Table S3). Furthermore, except for Pelikevirus alas, all the 190 Pithoviridae-like large genomes and Marosvirus alas have a near complete base 191 excision repair system.

# Heterogeneous Nucleocytoviricota abundance of the Russiancryosol

The permafrost samples were very heterogeneous in Nucleocytoviricota relative abundance (Fig. 3) and number of scaffolds, ranging from 2 found in the sample O (core permafrost under a lake in Yedoma, frozen for 40,000 years) to 721 scaffolds found in sample R (core permafrost under a drained thermokast lake, frozen for over 42,000 years). This sample was also the richest in eukaryotes with mostly 199 Streptophyta (35%), Dikarya (14%), Platyhelminthes (9%), Eumycetozoa (8%) and 200 Longamoebia (7%). In other permafrost core samples, the most abundant eukaryote 201 clades were Streptophyta (34%), Dikarya (18%), Chordata (7%), Arthropoda (5%) and 202 Eumycetozoa (3%) (Fig. S3). Interestingly, amoebas (Longamoebia) are a lot more 203 abundant in sample R than in other samples (Fig. S3).

204



#### 205

# Figure 3: Relative abundance of Nucleocytoviricota and Eukaryota across samples

The relative abundance is calculated as the sum of coverages belonging to the given group divided by the total sample coverage. Sample names in red are surface samples from Kamchatka while samples in blue, green and purple indicate that they come from three different forages in the Yukechi Alas area. The pie charts indicate the taxonomy of the Nucleocytoviricota in different samples (see further for phylogeny). Only classified scaffolds were considered. 213 The relative proportion of giant viruses abundance (Fig. 3) and the number of scaffolds 214 were correlated to the ones of Eukaryota, with Spearman correlation coefficients of 215  $\rho$ =0.72 (p-value=0.017) and  $\rho$ =0.83 (p-value=0.003), respectively. Such correlation could simply be explained by host-parasites dynamics. Alternatively, one could 216 hypothesize that Nucleocytoviricota scaffolds correspond to endogenized viruses in 217 218 eukaryotes (GEVE), as previously shown in green algae (35). The confusion is 219 possible as 57% (193 out of 338) of the GEVE pseudo-contigs (see Methods) were captured by our Nucleocytoviricota detection method. To explore this possibility, we 220 221 thus checked for endogenization signs in the viral scaffolds using Viralrecall (36) (example in Fig. S4) but none was found. In addition, Nucleocytoviricota largely 222 outnumber eukaryotes with a 4:1 Nucleocytoviricota/Eukaryota ratio in the sum of 223 224 coverages (mean=4.06, sd=4.22) and number of scaffolds (mean=4.40, sd=3.34). 225 Altogether, this suggests that most of the discovered permafrost Nucleocytoviricota scaffolds correspond to bona fide free viruses. 226

### 227 Exploration of the sequence diversity

To further investigate which viral families were present in the samples, we conducted a phylogenetic analysis based on the 7 marker genes (Table S3) from (31) We excluded the transcription elongation factor TFIIS from our analysis as its evolution was unclear and not coherent with the known Nucleocytoviricota clades (Fig. S5). It should also be noted that the primase D5 revealed an unexpected grouping of the Cedratviruses with *Phycodnaviridae* instead of *Pithoviridae*, suggesting that this gene was acquired from an unknown source in Cedratviruses (Fig. S5).

235 With this method, 369 Nucleocytoviricota scaffolds (19%) were taxonomically 236 classified (Fig. 4) corresponding to 40% of the 72 Mb of total Nucleocytoviricota

identified sequences. *Pithoviridae* and *Orpheoviridae*-like viral families were clearly 237 238 the most diverse followed by Megamimivirinae. In contrast, Marseilleviridae, Alphairidoviridae, Betairidoviridae and Ascoviridae were completely absent from our 239 240 samples. *Poxviridae* were not included in the phylogeny as they were absent from our 241 samples and adding their marker genes lowered the tree bootstraps values. In addition 242 to our strategy to combine different marker genes, we also computed a phylogenetic 243 tree from a single conserved one, the DNA polymerase, confirming that *Pithoviridae* 244 and Orpheoviridae-like sequences were the most diverse families in our samples (Fig. 245 S6).



### 248 Figure 4: Nucleocytoviricota phylogeny in all samples

Consensus of 1000 bootstrapped trees calculated by a partitioned analysis on 7 marker genes. Black dots represent branch bootstrap support  $\ge$  90. The colored labels indicate the reference genomes and the large viruses identified in this study (marked with a star). The colored clades were manually created to be monophyletic based on reference genomes. The marker genes used for this phylogeny are indicated as colored squares. Empty squares correspond to marker genes absent from the reference genomes. Black bars show the normalized mean coverage of the scaffold. Pimascovirales are defined as the clade composed of all the *Ascoviridae*, *Iridoviridae*, *Marseilleviridae*, but also *Pithoviridae*, *Orpheoviridae* and the metagenomic intermediate clades. The Extended\_phycodnaviridae group includes *Pandoraviridae* and Mollivirus. The Extended\_klosneuvirinae group includes the Cafeteria roenbergensis virus.

The permafrost data appears to reveal a whole new clade branching before the *Phycodnaviridae* and with no previously isolated representatives (Fig. 4). This is probably an artefact due to the divergent Cedratviruses primase D5 gene closer to the *Phycodnaviridae* primase. On the primase D5 tree (Fig. S7), the clade is split between Cedratviruses and the other half (probably *Phycodnaviridae*) that remains in the same position in the tree. A second unknown clade branching right before *Phycodnaviridae* (Fig. 4) had four members and was mildly supported by the bootstrap analysis (71%).

266 In order to explore the genome content diversity, we next analyzed the best BlastP matches against the NR database. Sequences unclassified by our phylogenetic 267 268 approach were dominated (50.6%) by ORFans, in the same range than 269 phylogenetically classified permafrost scaffolds (from 25.2% to 73.9% with on average 270 54%, Fig. S8A). This suggests that these sequences are not more divergent to known 271 relatives than any other Nucleocytoviricota sequence. They remained unclassified simply because they lack the marker genes. Secondly, even though viral Blast results 272 273 alone are only crude taxonomy indicators, they were nevertheless consistent with our 274 phylogenetic analyses. with unclassified scaffolds mainly composed of Megamimivirinae (43.4%), Klosneuvirinae (26.2%) and Pithoviridae (22.6%) related 275 276 sequence (Fig. S8B).

277 Not only *Pithoviridae* were unexpectedly diverse (Fig. 4), they were also the most 278 abundant Nucleocytoviricota according to their normalized coverage (Fig. 3).

279 Pithoviridae/Orpheoviridae-like families appear in all samples and particularly in R and N where they are very abundant (Fig. 3). The most covered sequences in five samples 280 281 (C, N, R, K and Q) come from these, and from extended Phycodnaviridae, Megamimivirinae and Klosneuvirinae in other samples. Along with the relative 282 283 abundance, the diversity of Nucleocytoviricota is guite heterogeneous with the 284 exception of samples N and R from the same borehole (16 and 19m respectively) 285 having a similar distribution (Fig. 3). Most viruses are specific to the sample they were recovered from, in particular the ones from surface samples (Fig. S9). Surprisingly, we 286 287 also found viruses that were common to samples from close locations in Central Yakutia but from different ages (samples K, L, M, N, P, Q and R; Table S1). This 288 289 indicates that part of the viral community was maintained over time.

### 290 Worldwide Nucleocytoviricota distribution

291 The *Pithoviridae* diversity and abundance observed in two samples from the Russian permafrost highlight the richness of this viral family in this environment, or alternatively, 292 293 a Nucleocytoviricota detection method more adapted to non-icosahedral viruses. To 294 investigate the presence of Nucleocytoviricota in other environments we applied the 295 same methodology to the Mgnify database (37), resulting in 3564 classified contigs. Since biomes are unevenly present in this database, with marine samples being 296 297 largely predominant, we found more Nucleocytoviricota in such samples (Fig. 5A). The 298 phylogenetic distribution of the scaffolds confirmed previous results highlighting the high diversity of *Mesomimivirinae* and *Phycodnaviridae* in oceanic samples (17–19). 299 300 On the other hand, *Pithoviridae* and *Orpheoviridae*'s diversity was much lower, with 301 corresponding sequences mostly found in engineered samples (bioreactors and 302 wastewater).



#### 305 Figure 5: Worldwide Nucleocytoviricota phylogenetic distribution

306 (A) 3664 contigs assembled from 427 datasets of the EBI Mgnify database and (B) 804 contigs 307 assembled from 147 terrestrial datasets of the JGI IMG/M database. Viral contigs were detected using 308 the previously described method and placed on tree using at least one of the seven marker genes. The 309 tree was made using Cyprinid herpesvirus 2 as outgroup. Clades containing the reference sequences 310 were manually drawn. Colored circles at tips represent reference genomes and the outer circle shows 311 the corresponding biome.

Terrestrial biomes being completely absent from the Mgnify database, we completed 312 this analysis by using 1835 terrestrial datasets collected from the JGI IMG/M database 313 314 (38). The vast majority of the samples exhibited no Nucleocytoviricota at all and few contigs over 10 kb in general (Fig. S10), probably due to the difficulty at assembling 315 316 sequence data from these complex environments. Our Russian samples, along with 317 few outliers from this database, stood out for having a high number of viral and total 318 contigs. Mesomimivirinae was the most represented sub-family in this terrestrial dataset (Fig. 5B), mainly due to its presence in two deep shales samples also rich in 319 320 Phycodnaviridae. Noteworthy, Pandoravirus-like sequences were found in sand and a 900 kb contig grouping next to *Pandoraviridae* and *Molliviridae* in peat permafrost
 samples. Pimascovirales were found in a variety of soil samples.

Overall, *Pithoviridae* and *Orpheoviridae* were more abundant in terrestrial samples than in aquatic samples (Fig. 5A and Fig. 5B). Russian permafrost samples were particularly and highly significantly enriched in these viruses, followed by forest soil, bioreactors and wastewater samples (Fig. S11).

# Functions encoded in the permafrost Nucleocytoviricotasequences

329 A total of 64,648 viral ORFs over 50 amino acids were manually annotated and 330 assigned to functional categories. Most of the permafrost metagenomes predicted 331 proteins are of unknown function (81%), as expected from the high proportion of ORFs 332 of that category in reference genomes (64%, Fig. S12). With a stringent minimal ORF size of 150 amino acids, the proportions are still of 76% and 56%, respectively. 333 334 Unspecific annotations such as Ankyrin repeat proteins, F-box proteins and FNIP repeat proteins, represent 1.4, 0.2 and 0.5% of the permafrost viral proteins while they 335 336 represent 4.6, 1.8 and 0.6% in the reference ones. Most genes with a known function 337 are involved in DNA replication, recombination and/or repair. There are also auxiliary 338 metabolic genes that are scattered within the different viral families (Fig. S13). The 339 distribution of functional categories found in the permafrost is the same as in the 340 Nucleocytoviricota references (Fig. S12). Overall, our analysis highlights a patchwork of functions encoded by these viruses (Fig. S13). 341

Looking at the most shared functions (i.e. present in most families) among the reference genomes and permafrost MAGs, we identified the known core genes (Fig. 6). Interestingly, the highly conserved mRNA capping enzyme is absent from the

Iridoviridae/Ascoviridae clade. The patatin phospholipase, suspected to be conserved 345 346 among Nucleocytoviricota (39), is confirmed as a core gene, only absent from Alphairidoviridae (Fig. 6). Its role in viral infection is still unclear but such proteins 347 348 participate to cell invasion in parasitic bacteria and eukaryotes (40, 41). Also, according to our data, the A32-like packaging ATPase is no longer a universal 349 350 Nucleocytoviricota marker gene, as it is not only lacking from the reference 351 *Pithoviridae* genomes but also absent from all clades ranging from Pitho-orpheo div8 352 to Pithoviridae (Fig. 6). Surprisingly, the Glutamine and Glutamine-dependent asparagine synthases known to characterize Mimiviridae (42) were also found in a 353 354 permafrost Pithoviridae.



356

### 357 Figure 6: Most shared functions among Nucleocytoviricota families

Functions were selected among the annotations found in at least 10 clades. Metagenomic sequences are marked as black rectangles at the bottom of the plot while blank spaces correspond to reference genomes. Groups with less than 300 ORFs were marked as "Other". The size of the dots represents the normalized ORFs counts (i.e ORF counts/total number of ORFs in the group). The right-most 362 column indicates the number of distinct clades having the function. The lines are sorted according to363 this value.

### 364 **DNA structure-related genes**

Giant DNA viruses from different clades exhibit either circular (15, 43) or linear (16, 365 366 28) genome structures. Marseilleviridae have a chromatin-like genome organized around virally encoded histones (44). Here we expanded the range of viral histones, 367 368 identifying them in *Pithoviridae*, *Megamimivirinae*, Pithoviridae div1, Pithoorpheo div7 and extended Phycodnaviridae sequences. We reconstructed their 369 phylogenetic histories which turned out to involve many independent HGTs of different 370 ages (Fig. 7A). In some cases, such as for Pleistovirus alas and a Pitho-orpheo div7 371 scaffold, the viral H3 histone is of ancient origin with a deep branching before the 372 373 eukaryotic ones. The latter also forms a histone doublet with fused H3-H4 domains as 374 already observed for the Marseilleviridae H2A-H2B and H3-H4 histones (44, 45). 375 Other viral histones appear more recently acquired from eukaryotes, like for the 400 kb M bin2028 k2 scaffold belonging to the extended Phycodnaviridae (close to 376 377 Pandoraviridae and Molliviridae) that encodes a H2A histone from Viridiplantae and a H3 one from an unknown eukaryote (Fig. 7A). Even more recently, the reference 378 379 Pandoraviruses (P. salinus and P. dulcis) and Medusavirus acquired H2B histones of 380 amoebic origin.



382 Thermoplasmata

## Figure 7: Phylogeny of three functions found in the permafrost Nucleocytoviricota.

385 All trees were computed by lqtree. Only ORFs of permafrost classified scaffolds were used in the trees. 386 (A) Sequences of core histones were retrieved from the HistoneDB database. The types of the viral 387 histones were confirmed through an Hmmsearch alignment of reference histone on the viral ORFs. (B) 388 The ATP synthase subunit F tree was build using sequences matching the PF1990 Pfam domain as 389 well as Pithoviridae div1 proteins with this annotated function and the best BlastP matching proteins 390 against NR. (C) The truncated hemoglobin tree was computed using the proteins from this study 391 combined with the IPR00146 Interpro domain sequences and BlastP matches of Nucleocytoviricota 392 sequences.

### 393 Auxiliary metabolic genes

Unexpectedly, an ATP synthase subunit F was found in a Pithoviridae div1 sequence 394 395 of nearly 200kb (Fig. 7B). The viral ORF matches the PF01990 Pfam domain that gathers prokaryotic ATP synthases as well as subunits of the eukaryotic vacuolar 396 397 ATPase. In eukaryotes, these proteins can serve many roles depending on the 398 organism and cell type but a common function is to acidify cellular compartments such as lysosomes (46). The ATP synthase subunit found in Pithoviridae div1 appears to 399 be of ancient origin (Fig. 7B). Two other subunits of the ATP synthase (one 400 401 Delta/Epsilon and one Beta) were also found in unclassified Nucleocytoviricota from 402 this study.

403 Other auxiliary metabolic genes found in this study include viral truncated hemoglobins 404 that are absent from reference Nucleocytoviricota. They likely come from three 405 different HGT events (Fig. 7C). A first one occurred between a prokaryote and 406 Pithoviridae\_div1 viruses. A second bacteria-to-virus HGT involved a *Pithoviridae* or 407 an Extended\_Phycodnaviridae that subsequently exchanged the truncated

408 hemoglobin gene. These proteins are able to bind oxygen and protect cells against409 oxidative stress from NO or other oxygen reactive molecules (47).

### 410 **Translation-related genes**

411 We found 20 different types of virally-encoded aminoacyl-tRNA synthetases (aaRSs) 412 in the permafrost metagenomic scaffolds. *Klosneuvirinae* is the clade with the most translation-related gene content followed by Megamimivirinae. For instance the 413 414 Klosneuvirinae Marosvirus alas found in this study (Fig 2 and Fig. 4) contains an expanded translation-related gene repertoire (10 translation initiation factors, 4 415 416 translation elongation factors, a translation termination factor and as much as 11 different aaRSs) as well as 5 tRNAs clustered together (Fig. 2). Besides Mimiviridae, 417 418 ten different types of aaRSs were found in the Pithoviridae div1 clade, including 7 419 different ones in Hydrivirus alas (Fig 2 and Fig. 4) that also encodes 9 tRNA, 3 420 translation initiation and elongation factors, and a translation termination factor.

421 We investigated the phylogeny of the different types of aaRSs found in our datasets that revealed entangled evolutionary pathways between viruses and cellular 422 423 organisms (Fig. S14, S15 and S16). In most cases, the viral aaRSs came from a 424 probable HGT from Eukaryotes (tryptophan, leucine, glutamine, threonine, 425 methionine, isoleucine, arginine, aspartate, serine and phenylalanine) (Fig. S14 and 426 S16). One clear example is the exchange of a threonine-tRNA synthetase from 427 Dictyostelia (Amoebozoa) to Hydrivirus alas (Fig. S14). The exchanges concerned both the mitochondrial (for instance arginine, phenylalanine) or the cytoplasmic copies 428 429 (Fig. S16). There were also some more rare cases of HGT from a Prokaryote to a virus as for the glycine- and tyrosine-tRNA synthetases that were transferred from an 430 431 Archaea (Fig. S15). Genes have also passed from Bacteria to Nucleocytoviricota as

432 for the glycine-tRNA synthetase of Hydrivirus alas and the valine-tRNA synthetase of 433 a permafrost *Megamimivirinae*. For the latter, the bacterial sources were Rickettsiales 434 that are endosymbionts of amoeba (48), thus probably sharing the same host. The 435 source of the tryptophan-tRNA synthetase in Hydrivirus alas is less clear but one can 436 see that a duplication event occurred at the same locus right after the gene was 437 acquired (Fig. S14).

While the vast majority of Nucleocytoviricota genes have no identifiable homologs, the ones with cellular homologs usually deeply branch in the phylogenetic trees (17, 49), in accordance with their suspected ancient origin (31, 50). We found here several viral aaRSs that belong to divergent families tightly clustered together within the cellular homologs (Fig. S16). So not only viral aaRSs are of cellular origin, spanning all domains of life, they were also probably exchanged between viruses of different families.

### 445 **The Major Capsid Protein**

446 Little is known about the structural proteins that constitute the particle of non-447 icosahedral giant viruses and what function their encoded MCP might have. In 448 Ascoviridae, the MCP is still a major protein in the virion (51), while Pandoraviridae 449 simply lack the gene. The MCP present in the related Mollivirus sibericum is only the 450 seventh most abundant virion protein and is thought to be involved in scaffolding 451 during virion assembly (52), as observed in *Poxviridae* (53). The annotated MCP-like 452 genes in Pithoviridae and Orpheoviridae genomes are so divergent that a Blast 453 homology search against the NR database fails to identify other Nucleocytoviricota homologs, even at a low confidence E-value threshold of 10<sup>-2</sup>. Furthermore, the protein 454 455 is not detected in Pithovirus sibericum virion proteome (15). Thus, the homology

between Nucleocytoviricota MCPs and the MCP-like of Pithoviridae/Orpheoviridae is 456 worth being explored. We therefore constructed a BlastP network of all annotated 457 458 large eukaryotic DNA viruses MCPs (Fig. 8A). As expected, the Pithoviridae MCPslike are the most divergent and disconnected from the rest of the network, where 459 460 icosahedral viruses and Ascoviridae form a strong cluster. But when adding the MCPs encoded in the permafrost metagenomics scaffolds, the Pithoviridae share 461 connections to the other Nucleocytoviricota (Fig. 8B). More specifically, the MCP 462 encoded in the Pitho-orpheo div clades (Fig. 4) fill the gap between the Pithoviridae 463 and the other Pimascovirales genes (Fig. 8B-C). Increasing the BlastP E-value 464 stringency places the *Pithoviridae*-like MCPs apart from other Megaviricetes except 465 for Pitho-orpheo div8, indicating that its MCP is closer to Marseilleviridae than to 466 *Pithoviridae* (Fig. 8D). From this we can conclude on the homology between the 467 icosahedral Nucleocytoviricota and the non-icosahedral Pithoviridae MCPs. 468

469



471 Figure 8: Major capsid protein network

The network was made with a BlastP of all vs all annotated large eukaryotic DNA virus major capsid proteins with varying E-value cutoffs and visualized in cytoscape. The edges were calculated from the bitscore. Circles correspond to reference genomes and rectangles to MAGs from this study. The red arrows depict the MCP identified in a Pitho-orpheo\_div8 scaffold (see Fig. 4 for phylogeny) and the dashed ellipses highlight the *Pithoviridae/Orpheoviridae*-like MCPs. The NR E-value was calculated based on the difference of database size.

### 478 **Discussion**

479 Recent large scale metagenomic data analyses strikingly revealed that Nucleocytoviricota are widespread in various environments (17, 18, 23, 24). Our 480 481 analysis of the cryosol and permafrost samples, as well as other datasets (JGI IMG/M 482 and EBI Mgnify databases), confirms this ubiquity. Nevertheless we pointed out an 483 important heterogeneity in Nucleocytoviricota proportion across environments. Some of the permafrost datasets appeared to be among the most enriched in 484 485 Nucleocytoviricota, reaching up to 12% of the sequenced organisms. The relative DNA 486 sequence coverage (Fig. 3) even suggests that they outnumber their hosts, in the 487 same way bacteriophages often outnumber bacteria in the ocean (54, 55). This high abundance is also the result of the high Nucleocytoviricota diversity in the samples, as 488 489 it does not come from a single virus. Furthermore, by taking advantage of the 490 permafrost ability to preserve ancient organisms, we showed that some 491 Nucleocytoviricota strains were not only abundant but had been present in the active community for a long time (Fig. S9). Considering only syngenetic permafrost samples, 492 493 we found Nucleocytoviricota shared in samples of up to 14,000 years difference. This 494 indicates that they are important players of this particular area of central Yakutia.

495 The Nucleocytoviricota diversity explored in this study strikingly revealed many Pithoviridae-like sequences that are very divergent from the reference genomes and 496 497 constitute new clades within the Pimascovirales. This includes large genomes, in particular the complete 1.6 Mb Hydrivirus alas genome. So, next to Pandoraviruses 498 499 (28), Orpheoviruses (32), Klosneuviruses (56) and Mimiviruses (57), this provides with 500 yet another example of a viral genome largely over 1 Mb. The nature of the 501 evolutionary forces pushing some viruses to retain or acquire so many genes remains a matter of debate (58-61). Horizontal gene transfers from cellular hosts is 502 503 hypothesized by some authors to account for their large gene content (56, 62). We indeed found examples of cellular genes gained by HGT in this study (Fig. 7 and Fig. 504 505 S14-S16) but this only accounts for a small proportion of their gene content, the vast 506 majority having no identifiable cellular homologs. Gene duplication, on the other hand, 507 a well-known source of functional innovation since the pioneering work of Susumu Ohno (63), may contribute to the genome inflation of giant viruses (49, 64). Another 508 509 possible source of genetic innovation is the *de novo* gene creation from intergenic 510 regions (49, 65). The present work expanded the Nucleocytoviricota families' 511 pangenomes, in particular the *Pithoviridae*-like and *Mimiviridae*, with an overwhelming proportion of ORFans. This militates for the *de novo* gene creation hypothesis that 512 remains to be further tested. 513

514 Despite the isolation of Mollivirus kamchatka (30) and Cedratvirus kamchatka (29) in 515 the studied samples (samples C-D-E), their genomes were not identified in our 516 assembled metagenomic data. Such a discrepancy had already been observed for 517 Pithovirus sibericum and Mollivirus sibericum, where metagenomic sequence reads 518 confirmed their presence in the samples but at a coverage too low to obtain assembled

519 contigs (16). Concerning the Pandoraviridae, although different strains of these viruses were isolated from various geographical locations, including soil (28, 49, 65-520 521 68), very few Pandoraviridae-like sequences were identified in our cryosol data and in 522 a vast array of environmental samples (Fig. 5), as already noticed from previous 523 metagenomic studies (18). This underlines the importance of exploring complex 524 environment communities with complementary approaches to unravel the true diversity of less studied giant virus families. This includes adapted metagenomic 525 526 pipelines such as our attempt to reveal non-icosahedral viruses combined to direct 527 isolation.

528 The functional annotation performed in this work highlights the paucity of functions 529 strictly shared between Nucleocytoviricota. Even a central protein like the A32 530 Packaging ATPase is absent from the entire *Pithoviridae*-like clade (Fig. 8). Likewise 531 the MCP is not encoded in the *Pandoraviridae* genomes. Regarding the highly divergent *Pithoviridae/Orpheoviridae* MCP-like genes, our analysis helped to reveal 532 533 their homology with the other Pimascovirales (Fig. 8). These genes could then either 534 come from a shared ancestor (69), as suggested by the core genes phylogenies (Fig. S5), or was acquired very early in the *Pithoviridae*/Orpheoviridae evolution. We can 535 speculate that the MCP quickly started to lose or change its function, before the 536 537 divergence of Pitho-orpheo div7 and the other family members (Fig. 4), perhaps with a progressive change in virion morphology. 538

539 Besides the few functions shared by the Nucleocytoviricota, our work also highlights 540 a patchwork of functions encoded by these genomes. When looking at specific 541 functions, we detected independent cases of HGT from Eukaryotes to viruses but also 542 between viruses belonging to different families (Fig. 7 and Fig. S14-S16). This is

543 probably the testimony of coinfections, as members of the *Marseilleviridae*, 544 *Mimiviridae*, *Pithoviridae*, *Pandoraviridae* and *Molliviridae* families can infect the same 545 host. In line with this hypothesis, we recently showed that DNA methylation, 546 widespread in giant viruses, is mediated by methyltransferases and Restriction-547 Modification systems that are frequently horizontally exchanged between viruses from 548 different families (29).

549 The functional patchwork, the gene exchanges between viruses of different families, 550 together with the very few shared genes, may challenge the monophyly of the recently established Nucleocytoviricota phylum by the International Committee on Taxonomy 551 of Viruses (ICTV) (70). Except for the DNA primase of Cedratviruses, our trees of 552 553 seven marker genes would indeed indicate a shared ancestry of the different 554 Nucleocytoviricota families analyzed in this work (Fig. S5). However, when cellular 555 genes are integrated to the phylogenetic trees, only three of the five most shared genes strictly support the monophyly of the Nucleocytoviricota (71): the viral late 556 557 transcription factor 3, the Holliday junction resolvase and the A32 packaging ATPase. 558 The latter has also been shown to be exchanged between *Mimiviridae* and Yaravirus, 559 an Acanthamoeba infecting virus that does not belong to the phylum (71, 72). The other core genes such as the DNA polymerase is separated by several cellular clades 560 561 between Pokkesviricetes and Megaviricetes (73). Likewise the two largest subunits of the RNA polymerase of Asfarviridae and Mimiviridae have a different history than the 562 other Nucleocytoviricota (31). These examples question the consistency of the 563 564 phylum.

565 The primary objective of this study was to assess the diversity of large DNA viruses in 566 permafrost. Our analyses revealed an unexpected number of new viral sub-groups

and clades among some of the previously established families of the Nucleocytoviricota phylum, mixing an intricate patchwork of functions amidst a majority of anonymous genes of unknown functions. The in-depth study of these genes will allow to better understand their physiology but also to rule on the existence or not of a common ancestor for its deepest branches.

### 572 Materials and Methods

#### 573 Data preparation

Illumina sequencing reads from all samples (Table S1) were assembled into contigs 574 using Spades (v3.14) (74) and then binned using Metabat2 (v2.15) (75) with a minimal 575 contig length  $\geq$  1500 and bin length  $\geq$  10,000. Reads corresponding to each contig 576 577 were retrieved and gathered from their respective bins using an in house script. The read subsets were then reassembled using Spades (v3.14) in default mode or with the 578 "--meta" option. Reads were mapped on the resulting scaffolds  $\geq$  10kb using Bowtie2 579 580 (v2.3.4.1) (76) with the "--very-sensitive" option. Scaffold relative coverage was computed as the mean scaffold coverage divided by the total sample coverage. Bins, 581 582 contigs and scaffolds were verified with Checkm (v1.1.2) (77) using the lineage workflow. 583

### 584 **Control database preparation**

585 Reference Nucleocytoviricota were chosen following a former phylogenetic study (31).
586 The corresponding genomes were gathered from the NCBI repository. Lausannevirus,
587 Melbournevirus, Ambystoma tigrinum virus, Infectious spleen and kidney necrosis
588 virus, Invertebrate iridovirus 22, Invertebrate iridovirus 25 and Singapore grouper
589 iridovirus were removed to avoid an overrepresentation of their families. We added the

590 genomes of Acanthamoeba castellanii medusavirus (AP018495.1), Bodo saltans virus 591 (MF782455.1), Cedratvirus kamchatka (MN873693.1) and Tetraselmis virus 1 592 (KY322437.1). Genomes from Archaea, Eukaryota and Bacteria (Table S4) were 593 retrieved from Genbank. For each genome, non-overlapping sequences were cut with an in house script following a distribution similar to our dataset to simulate 594 595 metagenomic contigs. Genes were then predicted by Genemark (v3.36) (74) using the 596 metagenomic model. For the Nucleocytoviricota phylogeny, core genes previously 597 identified (31) were used in addition to the ones found by Psiblast (from BLAST+ 598 v2.8.1) (75). We also added Amsacta moorei entomopoxvirus (AF250284.1). Variola 599 virus (NC 001611.1) and Cyprinid herpesvirus 2 (MN201961.1) as outgroup.

### 600 Nucleocytoviricota specific profiles databases

601 The database constructed by (18) was completed with specific signatures of 602 Pithoviridae using the genomes of Cedratvirus A11 (34), Cedratvirus kamchatka (29), 603 Cedratvirus lausannensis (76), Cedratvirus zaza (77), Brazilian cedratvirus (77), 604 Pithovirus massiliensis (33), Pithovirus sibericum (15), Orpheovirus (32), all the metagenomic *Pithoviridae* released from one study of Loki's Castle hydrothermal 605 vents (24), the divergent Orpheoviridae/Pithoviridae SRX247688.42 (17), the 606 GVMAG-S-1056828-40 (18) other Cedratvirus/Pithovirus 607 and sequences 608 (supplementary data files). For Pandoraviridae we gathered sequences from 609 Pandoravirus braziliensis (78), P. celtis (65), P. dulcis (28), P. inopinatum (67), P. macleodensis (49), P. neocaledonia (49), P. pampulha (78), P. guercus (65), P. 610 611 salinus (28), Mollivirus kamchatka (30) and M. sibericum (16). The ORFs were then predicted using Genemark (v4.32) with the "--virus" option and ORFs  $\geq$  50 amino-acids 612 were kept. Orthogroups were calculated with Orthofinder (79) and HMM profiles were 613 614 built using the Hmmer suite (v3.2.1) (80) for each one. HMMs were further aligned to 33

the Refseq protein database (from March 2020) using the same suite. Only HMMs specific to *Pithoviridae*, *Orpheoviridae*, *Pandoraviridae* or Molliviruses with E-value  $\leq$ 10<sup>-10</sup> were kept to complete the database. To these were added Nucleocytoviricotaspecific VOG orthogroups (https://vogdb.org/).

### 619 Retrieving viral sequences

The Nucleocytoviricota-specific profile database was searched against the control and 620 621 permafrost ORFs using Hmmsearch. To check for cellular signatures, all the ORFs were aligned to the Refseq protein database using Diamond blastp (v0.9.31.132) with 622 623 the "--taxonlist 2,2759,2157" option and hits  $\geq$  35% sequence identity were checked. On the control metagenomic simulated dataset, the amount of false positives and false 624 625 negatives were assessed according to the cellular and viral matches for each group 626 (Nucleocytoviricota, Archaea, Bacteria, Eukaryota). We set the threshold at less than 627 1% of false eukaryotic positives. The same threshold was applied to the permafrost data to retrieve viral contigs. 628

### 629 Functional annotation

All the ORFs  $\geq$  50 amino-acids were queried against the NR database (from June 630 631 2020) using Blastp, the VOG database using Hmmsearch, the Pfam database using Interproscan (v.5.39-77) and against EggNOG (81) using the online version of 632 633 Emapper-1.03. For all, the E-value threshold was set to 10<sup>-5</sup>. Functional annotations 634 of each predicted protein were defined manually, first based on the matching domains 635 annotations, then by considering the full sequence alignments (Blast, EggNOG and 636 VOG). EggNOG categories were also set manually for each gene. When existing, the 637 functional annotations of reference viral genomes (see control database preparation) were retrieved from Genbank. Grouper iridovirus, Heliothis virescens ascovirus 3e and 638

639 Invertebrate iridescent virus 6 were manually reannotated using the same protocol as640 for the permafrost ORFs.

### 641 **Contamination control**

The functional annotation step helped to remove non-Nucleocytoviricota scaffolds 642 based on the presence of typical viral/phage genes or with ORFs consistently 643 matching cellular organisms. The scaffolds were checked for the presence of 644 645 ribosomes using Barrnap (v0.9) (82). Finally, we checked for possible GEVEs (Giant Endogenous viral elements) in our curated scaffolds. We made pseudo-contigs from 646 647 the GEVEs identified by (35) and applied our method on them. As 57% (193 out of 338) of the GEVEs peudo-contigs were caught, we proceeded to check for 648 649 endogenization signs in our permafrost scaffolds. This was done by plotting the 650 domain of the Blastp hits as well as the VOG matches for each scaffold with the results 651 of the Viralrecall (v2.0) rolling score (36). Scaffolds with at least one region with a negative Viralrecall score were visually inspected. For comparison, we also tested 652 653 Viralrecall with the "--contiglevel" option.

### 654 Large genomes assembly verification and circularization

The eight largest MAGs ( $\geq$  500kb) were scrutinized for possible chimeric assemblies. We used the Integrative Genome Viewer (83) to assess potential coverage drops (mainly due to ambiguous bases added during scaffolding), but in each case read pairs overlapped the low coverage intervals. For circularization, we created a model contig concatenating both ends of the MAG, mapped the reads using Bowtie2 and checked the uniformity of the coverage at the junctions.

### 661 Abundance estimation and mapping

Metagenomics reads were mapped to the viral scaffolds using Bowtie2 with the –verysensitive option and filtered with Samtools (-q 3 option). Reads  $\leq$  30 nucleotides were discarded. The relative mean coverage of the scaffolds were then used as estimators of the scaffold abundance in the sample. For in-between sample comparisons, reads were size-filtered and then mapped to the viral scaffold with a minimum quality filter of 30. Then, only scaffold  $\geq$  10kb in size were considered.

#### 668 **Phylogenetic analysis**

For the selected marker genes, individual gene trees were built from reference genomes only. Multiple alignments were performed using MAFFT (v7.407) (84), removal of divergent regions with ClipKIT (85) and models estimations (86) and tree inference using lqtree (v1.6.12) (87) (options "-bb 1000" (88), "-bi 100" and "-m MFP"). The best model was VT+F+R4 for the TFIIS tree, LG+F+G4 for the MCP and LG+F+R5 for all the other marker genes. A global tree was calculated by a partitioned analysis (89) to include genomes with missing data.

To identify the marker genes in the permafrost data, Psiblast was used to align reference marker genes to the viral ORFs (initial E-value  $\leq 10^{-5}$ ). Next, in order to avoid using a paralog of the marker genes, we defined a second stringent E-value threshold the following way: E-values of all second matches for scaffolds with multiple copies were sorted in ascending order, then the stringent threshold was defined based on the first quartile (Table S5). Finally, only the best match per scaffold was kept for phylogenetic reconstruction if it was better than the stringent threshold for this gene.

683 The 7 marker genes were aligned using PASTA (90), clipped with ClipKIT and 684 concatenated by Catsequences (91). The global tree with ultrafast bootstraps was then inferred by lqtree with options "-spp, -bb 1000" and "-bi 200 -m MFP" that
calculates the best model per marker gene. Tree visualization was handled using
Figtree (http://tree.bio.ed.ac.uk/software/figtree/) and the Itol web server (92).

### 688 Worldwide Nucleocytoviricota distribution

The EBI Mgnify (37) protein database from March 11<sup>th</sup> 2021 was downloaded and 689 proteins from environmental or engineered biomes were extracted for further analysis. 690 691 We only selected proteins from contigs  $\geq$  10 kb in size. We also downloaded 1835 terrestrial assemblies from the JGI IMG/M (38) database (Table S6), of which 1502 692 693 exhibited at least one contig  $\geq$  10 kb. The ORFs were predicted using Metagenemark 694 as previously. Nucleocytoviricota sequences were extracted from both databases as 695 described above (see Retrieving viral sequences). The same method than previously 696 described (see phylogenetic analysis) was applied to search for marker genes for 697 phylogeny. Reference and metagenomic marker genes were aligned using MAFFT with the "---auto" option. Amsacta moorei entomopoxvirus, Variola virus and Cyprinid 698 699 herpesvirus 2 were included in the analysis. The alignments were clipped with ClipKIT 700 and concatenated for a partitioned analysis. Empirical models for each partition were 701 inferred by Modelestimator (93). Finally, the trees were computed using lqtree (with bb 1000 -bi 200). 702

### 703 **Phylogenetic analyses of selected functions**

For each function, a dataset of proteins was built using a combination of Nucleocytoviricota ORFs, corresponding Blast matched proteins from the NR database and reference proteins from specific databases. The latter includes Uniprot reviewed proteins of domains PF01990 (ATP synthase subunit F), IPR001412 (class I aminoacyl-tRNA synthetases), IPR006195 (class II aminoacyl-tRNA synthetases)

and IPR001486 (truncated hemoglobin). The reference core histone proteins were
also retrieved from the HistoneDB 2.0 database (94) in addition to reviewed archaeal
core histones from Uniprot (clustered using CDhit (95)). For all the functions, the
multiple alignments were performed using PASTA (90) or MAFFT (84) and trimmed
with ClipKit (85). The tree was then computed by lqtree (87) with options -bb 5000 -bi
200 -m TEST.

### 715 Major Capsid Protein network

All proteins annotated as "Large eukaryotic DNA virus major capsid protein" or "Divergent major capsid protein" were gathered with the reference MCPs and aligned against each other with BlastP (E-value  $\leq 10^{-5}$ ). The network was created using Cytoscape (v3.8.2) (96). The edge-weighted Spring Embedded layout was used and the bitscores were chosen as weights in the heuristic mode. The E-value threshold was progressively decreased to  $10^{-30}$  and changes in the network were observed along the way.

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