Review

Mimiviridae: a rising family of highly diverse large aquatic dsDNA viruses infecting a wide variety of eukaryotes.

Jean-Michel Claverie1, * and Chantal Abergel2

1 Aix-Marseille Université & CNRS, Structural and Genomic Information Laboratory (UMR7256), Mediterranean Institute of Microbiology (FR3479), 163 Avenue de Luminy, Marseille F-13288, France; jean-michel.claverie@univ-amu.fr
2 Aix-Marseille Université & CNRS, Structural and Genomic Information Laboratory (UMR7256), Mediterranean Institute of Microbiology (FR3479), 163 Avenue de Luminy, Marseille F-13288, France; chantal.abergel@igs.cnrs-mrs.fr
* Correspondence: jean-michel.claverie@univ-amu.fr; Tel.: +33-49-182-5420

Abstract: Since 1998, when Jim van Etten’s team initiated its characterization, Paramecium bursaria Chlorella virus 1 (PBCV-1) had been the largest known DNA virus, both in terms particle size and genome complexity. In 2003, The Acanthamoeba-infecting Mimivirus unexpectedly superseded PBCV-1, opening the era of giant viruses, i.e. with virions large enough to be visible by light microscopy and genomes encoding more proteins than many bacteria. During the 15 following years, the isolation of many Mimivirus-relatives, have made the Mimiviridae one of the largest and most diverse family of eukaryotic viruses isolated from aquatic environments. Metagenomic studies keep suggesting that many more remain to be isolated. As Mimiviridae members are found to infect an increasing range of phytoplankton species, their taxonomic position compared to the traditional Phycodnaviridae (i.e. etymologically “algal viruses”) became a source of confusion in the literature. Following a rapid history of the key discoveries that established the Mimiviridae family, we describe its current taxonomic structure and propose a set of operational criteria to help in the classification of future isolates.

Keywords: Mimiviridae; Algal virus; Giant virus; Phycodnaviridae; Aquatic virus

1. Introduction

The viral nature of Mimivirus was finally recognized in 2003 [1, 2], more than 10 years after Timothy Rowbotham, an investigator for Public Health England, first isolated it from a cooling tower in the city of Bradford in search for a pneumonia-causing pathogen [3]. For ten years, the newly isolated Acanthamoeba-infecting microbe, dubbed Bradfordcoccus, was unsuccessfully investigated as an intracellular parasitic bacterium, resisting all standard characterization approaches. The difficulties however, were not technical, but epistemological [4, 5]. After more than a century of virology dedicated to the study of “filtering” infectious agents originally defined as those not retained by sterilizing filters and invisible by light microscopy, the notion that a virus could be propagated by particles as big as bacteria was a conceptual leap that no bona fide virologist was ready to make. It was thus for a team of bacteriologists to identify Mimivirus as the first “giant” virus, then allowing the subsequent discovery of many more of them, now encompassing at least 4 different families and opening a new and booming era in virology [4].

The initial awe caused by the mere size of the giant virus particles was further amplified by the discovery that they packed giant genomes encoding more genes than many bacteria and the smallest parasitic eukaryotic microorganism [4, 6, 7]. This revived numerous reflections and debates on the concept of virus [4, 6, 8-15], on their position relative to the Tree of Life [16-25], and on their role in
the evolution of the cellular life forms [4, 10, 14, 26-28]. The giant virus revolution is clearly not near its end, and more surprises are to come. One of them is the continuing and even accelerating discovery of Mimivirus relatives in fresh, brackish, and sea water environments, already making the Mimiviridae a prominent family among aquatic viruses (Table 1). Members of this rapidly increasing family are both remarkable by the phylogenetic diversity of their hosts and the unmatched variability of their particle and genome sizes. Following a short history of the early developments, we then review the most recent discoveries of aquatic Mimiviridae, before ending by a set of guidelines to help their correct classification amidst other established families of aquatic viruses.

2. Results

The saga of the giant viruses truly began with the pioneering work of Jim van Etten and his collaborators who discovered, isolated and extensively characterized the first large DNA infecting a protist. This protist was the unicellular fresh-water green algae Chlorella (Trebusiophyceae, Viridaeplantae), small round cells 3 µm in diameter, nowadays quite popular as a detoxifying nutritional supplement in health food stores. Ironically, chlorella are linked with the history of virology from the very beginning since they were discovered by the same famous Dutch microbiologist, Martinus W. Beijerinck who forged the term “virus” (even though its concept of “liquid” infectious agent was quite wrong)[29]. Thus, with a little twist of fate, Beijerinck could have discovered Chlorella-infecting viruses, instead of simply reproducing Ivanovsky’s work on the filterable tobacco mosaic virus [30]. No doubt that the history of virology would then have been entirely different [5] and the discovery of giant viruses not delayed until the 21st century.

2.1. Pioneering work on large aquatic viruses

The first descriptions of very large icosahedral viruses infecting chlorella were published in 1981 [31, 32], thus very soon after Torrella and Morita first reported on the large incidence of virus-like particle in coastal see water in their landmark 1979 article [33]. An interesting twist (most often forgotten) of their landmark work is that they observed populations of unusually large viruses, collected on a 0.2 µm pore size filter, meant to retain bacteria. Unfortunately, the whole field of marine virology was subsequently based on the analysis of “viral fractions” defined as the filtrates in such filtration protocols [34], thus coming back to the historical definition of viruses as “filterable” microbes. This again, probably delayed the discovery of giant marine viruses by decades.

### Table 1. Physically isolated Mimiviridae with fully sequenced genomes

<table>
<thead>
<tr>
<th>Name1</th>
<th>accession</th>
<th>Genome size (kb)</th>
<th>Virion type</th>
<th>Virion size (nm)</th>
<th>Host phylum2</th>
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<tr>
<td>Mimivirus2</td>
<td>NC_014649</td>
<td>1,181 icosahedron</td>
<td>750</td>
<td>Amoebozoa</td>
<td></td>
</tr>
<tr>
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<td>680</td>
<td>Amoebozoa</td>
<td></td>
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<tr>
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<td>NC_020104</td>
<td>1,021 icosahedron</td>
<td>620</td>
<td>Amoebozoa</td>
<td></td>
</tr>
<tr>
<td>CroV</td>
<td>NC_014637</td>
<td>693 icosahedron</td>
<td>300</td>
<td>Heterokonta</td>
<td></td>
</tr>
<tr>
<td>PgV</td>
<td>NC_021312</td>
<td>460 icosahedron</td>
<td>150</td>
<td>Haptophyceae</td>
<td></td>
</tr>
<tr>
<td>CeV</td>
<td>NC_028094</td>
<td>474 icosahedron</td>
<td>160</td>
<td>Haptophyceae</td>
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</tr>
<tr>
<td>TetV</td>
<td>KY322437</td>
<td>668 icosahedron</td>
<td>240</td>
<td>Chlorophyceae</td>
<td></td>
</tr>
<tr>
<td>BsV</td>
<td>MF782455</td>
<td>1,386 icosahedron</td>
<td>300</td>
<td>Excavata</td>
<td></td>
</tr>
<tr>
<td>AaV</td>
<td>NC_024697</td>
<td>371 icosahedron</td>
<td>140</td>
<td>Heterokonta</td>
<td></td>
</tr>
<tr>
<td>TupanSL</td>
<td>KY523104</td>
<td>1,439 icos. + tail</td>
<td>450+550</td>
<td>Amoebozoa</td>
<td></td>
</tr>
<tr>
<td>TupanDO</td>
<td>MF405918</td>
<td>1,516 icos. + tail</td>
<td>450+550</td>
<td>Amoebozoa</td>
<td></td>
</tr>
</tbody>
</table>

1: Abbreviations: CroV: Cafeteria roentgenbursis virus; PgV: Phaeocystis globosa virus; CeV: Chrysochromulina ericina virus; TetV: Tetraselmis virus; BsV: Bodo saltans virus; AaV: Aureococcus anophagefferens virus; TupanSL: Tupanvirus soda Lake; TupanDO: Tupanvirus deep Ocean. 2: several strains very similar to these prototypes have been isolated and fully sequenced.
Once Van Etten and collaborators realized that the easiest virus to work with was one infecting a symbiotic chlorella of paramecium (the now famous *Paramecium bursaria* chlorella virus, PBCV-1), its characterization proceeded quickly. Much before its complete genome sequencing in 1997 [35], they correctly estimated as soon as 1982 that itsicosahedral particle 190 nm in diameter contained a dsDNA genome more than 300 kb-long [36]. The stage was then already set for the new notion of “giant” viruses defined in reference to both their large particle sizes and large genomes. Variable thresholds are used in today’s literature, but are around 350 nm for the particle size (i.e. making them visible by light microscopy) and 300 kb for the genome length [4]. Historically, the terms “giant” viruses was first used in 1995 by Jim Van Etten in the title of an article published in a Korean journal [37], then again in 1999 [38]. Following the subsequent threshold values introduced by the revolution of the truly giant viruses initiated by Mimivirus [1-4], Jim Van Etten nowadays qualifies the chlorella viruses as “very large” instead of “giant”. As we progress in our knowledge about their structure and metabolic capacity, it becomes clear that no fundamental biological difference distinguishes “very large” from “giant” dsDNA viruses that even co-exist in the same Mimiviridae family.

### 2.2. From *Acanthamoeba polyphaga* Mimivirus (APMV) to the first marine Mimiviridae

Our determination of the complete genome sequence of Mimivirus in 2004 [2] happened to coincide with a number of unrelated developments in the field of aquatic/marine microbiology that led us to change the course of our own research. One was the seminal publication of the first environmental “shotgun” sequencing of the Sargasso Sea by Venter’s team [39]. The second one was the fourth Algal Virus Workshop (4th AVW) [40] to which we participated at the invitation of Jim Van Etten, whom we had contacted as the leading expert in “very large” viruses. In collaboration with Dr. E. Ghedin (then from the TIGR Institute), we screened Venter’s first marine metagenomics sequence database with the freshly determined Mimivirus sequence. To our surprise, 15% of predicted Mimivirus ORFs had their closest homologs in these marine environmental sequences [41]. A few months later, our attendance to 4th AVW (and exchanges with its amazingly congenial research community) made us aware that unicellular algae were indeed host of many large viruses. Following these two hints, we first confirmed *in silico* the presence of Mimivirus relatives in the sea, using metagenomic datasets [42], gene-targeted amplicons sequences [43] and the partial genome sequences of previously isolated viruses infecting phylogenetically distant unicellular marine algae [44]. We finally succeeded in isolating the closest marine Mimivirus relative, *Megavirus chilensis*, from water sampled off the coast of central Chile [45]. In contrast with the other known marine Mimiviridae members, *M. chilensis* is infecting *Acanthamoeba*, i.e. the original host of Mimivirus. With its 1.26 Mb genome predicted to encode 1120 proteins, *M. chilensis* remained the largest Acanthamoeba-infecting Mimiviridae until the recent isolation and characterization of two new aquatic isolates called Tupanviruses [46](see below)(Table 1).

### 2.3. *Cafeteria roenbergensis* virus: the prototype of a new subfamily within the Mimiviridae

A large virus was isolated from the coastal waters of Texas in the early 1990s. Its host, originally misidentified as *Bodo* sp., was *Cafeteria roenbergensis*, a 2 to 6µm-long bicosoecid heterokont phagotrophic flagellate (Stramenopiles) that is widespread in marine environments [47]. In 2010, the complete genome sequence of *C. roenbergensis* virus (CroV) suggested the existence of a distinct clade within the Mimiviridae [48] (Figure 1 and 2). It was also the first demonstration that Mimivirus-relatives could infect very distant hosts (*Acanthamoeba* and *C. roenbergensis*), belonging to the earliest diverging branches within the Eukaryota, the unikonts and the bikonts [50]. As it is devoid of the thick fiber layer surrounding the icosahedral capsids characteristic of the *Acanthamoeba*-infecting Mimiviridae, CroV’s particle is only 300 nm in diameter. Yet, its 693 kb genome (predicted to encode 544 proteins) remained the largest of all marine plankton-infecting viruses until recently (see section 2.5). As a new, distinct member of the family, CroV played an important role in delineating both the conserved features and the expected diversity of the Mimiviridae in terms of gene content, particle size and morphology, and replication process. These conserved features can be used as criteria to
recognize and classify future family members, despite the further diversity exhibited by emerging clades. These features are presented in section 3.2.

2.4. Smaller Mimiviridae infecting bona fide microalgae: yet another subfamily

Prior to the complete genomic characterization of Mimivirus, a number of viruses infecting various species of unicellular algae had been isolated. These viruses exhibiting large dsDNA genomes packed in largeicosahedral particles were propagated on cultures of Pyramimonas orientalis (Chlorophyta, Prasinophyceae), Phaeocystis pouchetii and P. globosa (Haptophyta, Prymnesiophyceae), Chrysochromulina ericina (Haptophyta, Prymnesiophyceae) [51-53]. Targeted amplicons of genes encoding DNA polymerase B [43] or capsid proteins [54] produced partial sequences, some of which suggested a phylogenetic affinity between the largest algal viruses and Mimivirus. However, given the possibility of lateral gene transfers, complete genomes sequences were required to confirm the existence of a distinct clade of algae-infecting Mimiviridae. Such a clade was clearly confirmed following the analyses of the complete genome sequences of P. globosa virus (PgV) in 2013 [55], of Aureococcus anophagefferens virus in 2014 [56], and C. ericina (now Haptolina ericina) virus (CeV), in 2015 [57] (Table 1, Figure 1 and 2). The host spectrum of phytoplankton-infecting Mimiviridae was recently extended to bona fide green algae (e.g. members of the Chlorophyta lineage) with the detailed characterization of a virus (TetV) infecting Tetraselmis [58]. TetV exhibits the largest genome (668 kb) sequenced to date for a virus that infects a photosynthetic organism (Table 1). Yet the diameter of its icosahedral virion (≈240 nm) remains smaller than virus-like particles (up to 500 nm) observed long ago in other green algae [59, 60]. This suggests that more findings are expected as new algal hosts will be investigated. As of today, the known host spectrum of algae-infecting Mimiviridae (green branches in Figure 1 and 2) thus encompasses several classes of the heterokonta superphylum: Haptophyceae (PgV, CeV), Pelagophyceae (AaV) and Chlorodendrophyceae (TetV).

2.5. Recent additions to truly giant new members of the Mimiviridae

As we started to think that the upper limits of Mimiviridae genome size and gene content were reached with the 1,259,197-bp and 1,120 predicted proteins of M. chilensis, the year 2018 brought in three record-breaking viruses. Two of them, called Tupanvirus soda lake (TupanSL) (isolated from an alkaline lake in Brazil) and Tupanvirus deep ocean (TupanDO) (isolated from 3,000 meters deep marine sediments) can be propagated in the laboratory on A. castellanii and Vermamoeba vermiformis [46]. Their linear genomes consist of 1,439,508 bp and 1,516,267 bp, predicted to encode 1276 and 1425 proteins, respectively. Not far behind, but infecting a very different host, Deeg et al. [61] isolated the kinetoplastid-infecting Bodo saltans virus (BsV), exhibiting a 1.39 Mb genome predicted to encode 1227 proteins. Interestingly, both types of viruses are hinting to the existence of further subdivisions in the growing Mimiviridae family (Figure 1 and 2).

2.6. Metagenomic contributions to the expansion of the Mimiviridae family

We strongly believe that the description and taxonomic analysis of a brand new virus family, such as the Mimiviridae, should imperatively rely on laboratory cultivable isolates the availability of which is required for a detailed description of the infectious cycle that should be similar throughout the family. The exchange of virus isolates between laboratories should remain a key requirement for independent scientific validation, and is obviously needed for collaboration purposes. It should also be emphasized that most viral sequences produced from environmental DNA are so alien that they
can only be detected and interpreted by comparison with sequences determined from previously isolated viruses. This being said, the complete history of the Mimiviridae could not be told without mentioning some significant contributions from metagenomic studies.

We previously cited the contribution of the early Sargasso Sea data to establish the presence of Mimivirus relatives in the sea [41]. This was further confirmed in the larger Global Ocean Sampling data set [62] and associated to algal viruses through their partially sequenced DNA polymerases [43].

Once a special version of the DNA mismatch repair protein (MutS7) -curiously shared with the octocoral mitochondria [63] - was recognized as a distinctive feature of the Mimiviridae, it also served to estimate their abundance in the marine environment [44].

The metagenomics analysis of a hypersaline lake in Antarctic by Yau et al. [64] inaugurated a new era by claiming the discovery of two new Mimiviridae members based on two large contigs assembled from environmental sequences. Their claim was further supported by the full genome assembly of a virophage, then known as a unique parasite of Mimivirus [65, 66], from the same data. This article was quite influential in prompting for the search for additional virophages in a large variety of environments [67-70]. However, it was also a source of taxonomic confusion, as the authors named their two new large viruses “Organic Lake Phycodnavirus 1 and 2” (abbreviated as OLPV-1, OLPV-2), based on their phylogenetic affinity with viruses (CeV and PgV) infecting the unicellular algae Chrysochromulina and Phaeocystis. Like other algae-infecting Mimivirus relatives these viruses are now believed to define a Mimiviridae subfamily (see section 2.4) (Figure 1 and 2) although they often remain clumsily referred to as “OLPV-like” in the literature.

The more recent planet-wide TARA OCEANS expeditions [71, 72] confirmed the abundance of marine Mimiviridae (also called Megaviridae) in the ocean, although most of the sequences classified in this family showed large divergence from known viral genomes. This suggests that we need more isolates to adequately span the diversity of the family (as confirmed by the most recent findings, see section 2.5). The specific search for viral contigs exhibiting genes coding for the glutamine-dependent asparagine synthase uniquely found in Mimiviridae also hinted for the existence of additional clades for which isolated representatives are lacking [73].

The need for such missing isolates is well illustrated by a recent study performed by Schulz et al. [74]. Following a metagenomics analysis of a wastewater treatment plant in Klosterneuburg (Austria), these authors identified a divergent Mimivirus-relative (called “Klosneuvirus”) with an estimated genome size of 1.57 Mb dwarfing the size of all previously isolated and fully sequenced Mimiviridae. In absence of a physical isolate, such a genome size remains to be confirmed as it is obtained by summing 22 separate contigs (none of them larger than 452 kb) predicted to be part of the same genome by a heuristic bioinformatics procedure called “binning”. A similar uncertainty is shared by three other predicted Klosneuvirus relatives (i.e. Catovirus, Hokovirus, Indivirus). Despite the lack of physical isolates, the authors have proposed to classify them in a new subfamily (the “Klosneuvirinae”) within the family Mimiviridae. Fortunately, the isolation of BsV (section 2.5) one year later appears to substantiate this new clade (Figure 1 and 2). However, despite its phylogenetic similarity, BsV is conspicuously lacking the key feature of the postulated Klosneuvirinae: their large sets of amino-acyl tRNA synthetases (up to 19 for Klosneuvirus, only 2 for BSV).

The latest unexpected addition to the family Mimiviridae might be a virus that can cause a lethal disease in lake sturgeon Acipenser fulvescens [75], called Namao virus. The genomic data presumably associated to this virus consists of two non-overlapping contigs totaling 306,448 bp, sequenced from
DNA extracted from a homogenate of 3 tissues of 16 infected fishes [76]. There is yet no proof that these two contigs belong to the same virus. The remote possibility also remains that these sequences might correspond to ancestral viral sequences integrated in the fish genome, as previously seen in other organisms [77, 78]. Once definitely established by the full characterization of a cloned Namao virus isolate, the confirmation that some Mimiviridae members could include vertebrate species would make the host range of this family (from algae to fish!) truly unmatched among known (aquatic) viruses.

3. Discussion

3.1. Algae-infecting Mimiviridae versus Phycodnaviridae

The discovery of an increasing number of Mimivirus relatives infecting algae has generated some confusion in the literature about their taxonomic assignment. This confusion is propagated through the non-official classification assigned to their associated genomic entries by leading reference databases such as NCBI's [79]. Completely sequenced algae-infecting Mimiviridae can be found erratically classified as members of the Phycodnaviridae family in the genus Prymnesiovirus (PgV), as “unclassified Phycodnaviridae” (AaV, CeV), or as “unclassified virus” (TetV-1). Among partially sequenced Mimiviridae candidates (inferred from metagenomics assemblies), the naming and classification of two “Organic Lake Phycodnaviruses” (OLPV-1 and OLPV-2) has been particularly damaging. Others include Phaeocystis pouchetii virus (PpV-01) and Pyramimonas orientalis virus (PoV-01B). These erroneous classifications are not detected by non-specialists and do affect the outcome of automated large-scale phylogenetic analyses and gene contents comparisons, such as the delineation of core gene sets.

The difficulties in classifying new algae-infecting large DNA viruses are also inherent to the historical definition of the Phycodnaviridae family. According to its official ICTV definition, the family is meant to include algae-infecting viruses with large icosahedral virion (Ø > 100 nm) and large genome (> 100 kb). The family was then further divided into 5 genera based on the algal host (Table 2). We believe there are some fundamental flaws in this historical design. First, as algae are among the most phylogenetically diverse organisms (both by the origin of their chloroplast and that of the eukaryotic recipient), we could not expect that a single viral family (nowadays suggesting a monophyletic group for most authors) would adequately represent the diversity of their viruses. Indeed, the phycodnaviruses from the various genera appear increasingly different as we learn more about their infection process, their replication cycle, their genome structure, and their gene content. No other established viral family would encompass viruses with enveloped versus non-enveloped virions, with different genome structures (circular, linear), or encoding or not encoding a transcriptional apparatus. To our opinion, today’s genera composing the Phycodnaviridae should become separate families.

The current genus partition, designed according to the taxonomy of the algal host, is similarly unsatisfactory as it suggests a one to one correspondence between virus and host type, which we know is obviously wrong (e.g. viruses from many different families infect human). The diversity of the viruses infecting the Prymnesiophyceae (Haptophyceae) is a good example of this problem. Two types of very different dsDNA Prymnesioviruses have been shown to infect the unicellular algae Phaeocystis [53]. This is also the case of Tetraselmis which can be infected by a smaller virus (60 nm diameter, 31 kb genome) [80] (in addition to RNA viruses, not discussed here). It is now clear that the
largest virion/genome type (highlighted in orange in Table 2) belongs to the Mimiviridae [55], as well as other Haptophyceae-infecting viruses and the so-called Organic Lake “phycodnaviruses” [57].

Moreover, the Phycodnaviridae family includes a “Coccolithovirus” genus corresponding to a third type of virus infecting *Emiliania huxleyi* that is also a Prymnesiophyceae. Thus, even within a single genus, the current partition does not correctly reflect the diversity of the “Phycodnaviridae”. More problems can be anticipated with the green dinoflagellate *Heterocapsa circularisquama* (Dinophyceae), infected by a large DNA virus which might be close to the Asfarviridae [81].

Table 2. Structure of the Phycodnaviridae family as presently recognized by ICTV.

<table>
<thead>
<tr>
<th>Genus</th>
<th>Prototype</th>
<th>Encoded RNA pol</th>
<th>Genome Size (kb)</th>
<th>(G+C) %</th>
<th>Accession</th>
<th>Virion Ø (nm)</th>
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<tr>
<td>Chlorovirus</td>
<td>EhV-86</td>
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<td>407</td>
<td>40.2</td>
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<td>Prasinovirus</td>
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<td>336</td>
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<td>NC_002687</td>
<td>200</td>
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<td>OtV-5</td>
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<td>187</td>
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<td>460</td>
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<td>?</td>
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<td>HaV-1</td>
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<td>275</td>
<td>30.4</td>
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<td>202</td>
</tr>
</tbody>
</table>

1. Enveloped capsid [82]; 2. Two distinct virus groups [53]. The highlighted line corresponds to the proposed Mesomimivirinae subfamily within the Mimiviridae [57].

3.2. How to recognize and classify future members of the Mimiviridae

Our opinion is that a virus family, should only include viruses sharing a number of key properties likely to have been inherited from a common “founding” ancestor. These common properties will in turn imply the presence and conservation of orthologous proteins each of which should have its most similar homolog within the family (with rare exceptions caused by horizontal transfers). These key properties/proteins will be linked to a similar structure of the virion (e.g. capsid proteins), the genome replication machinery (DNA replication and repair enzymes), the virus cycle dependency of nuclear functions (e.g. the transcription machinery), or the intracellular location (e.g. nuclear/cytoplasmic) of particle production. Proteins/enzymes uniquely present in a group of viruses will be particularly useful as family “signature” even though they will be useless in the context a broader phylogenetic analyses. Finally, we do expect that the physiological traits shared by these viruses will make their global gene contents qualitatively similar. Members of the same virus family should then appear well clustered in a cladistic analysis/tree [61, 83].

The number of available Mimiviridae complete genome sequences is now large enough to make each new member easily recognizable by global protein level similarity searches. Even though the four most recent Mimiviridae isolates (TetV, BsV, Tupanviruses) exhibit a variable fraction of predicted protein best matching viral database entries, more than 60% are from previously characterized members of the family.

Confirmatory evidence are readily provided by the positioning of Mimiviridae candidates in the phylogenetic trees built from the sequence alignments or key enzymes such as B-type DNA polymerase, virion packaging ATPase, major capsid protein [AaV, TetV, CeV], three types of proteins (respectively NCVOG0038; NCVOG0022; NCVOG0249) with homologs in other families of large dsDNA viruses [84].
The previous genomic characterizations of Mimiviridae members led to the discovery of various key features apparently unique to the family. The association of any of them with a candidate virus (even partially sequenced) will considerably strengthen its tentative classification as a new member of the Mimiviridae. These key findings includes:

1. A high prevalence of the strictly conserved AAAATTGA motif in the promoter regions of early-transcribed genes [85].
2. A high prevalence of hairpin-forming transcription termination motifs in the 3' end of genes [86].
3. The co-isolation (or sequencing) of a virophage. Virophages are small dsDNA viruses with 17-18 kb genomes that can only replicate in host cells undergoing an infection by a Mimiviridae [66, 87]. Not all family members have been associated with a virophage [Table 3].
4. The detection of a transpoviron. Transpovirons are 7 kb-long plasmid-like linear dsDNA molecules found in association with Mimivirus-relatives [88].
5. The presence of a Mimiviridae-specific version of glutamine-dependent Asparagine synthetase (AsnS) gene. A different, easily distinguishable, homolog of this enzyme is also found in some Prasinoviruses (e.g. OtV5) [73].
6. The presence of aminoacyl tRNA synthetases (aaRS). The finding of such central components of the translational apparatus in the Mimivirus genome [2] was considered revolutionary, as the historical definition of viruses denied them the capacity to synthetize proteins [4]. Conflicting hypotheses on the evolutionary origin of these viral enzymes generated a still ongoing controversial debate. The presence of these aaRS remains a remarkable feature of the Mimiviridae, although not unique to them anymore [7]. Unfortunately, the number of virus-encoded aaRS varies greatly among the Mimiviridae members (from 0 up to 20 different ones in Prasinoviruses [46]) (Table 3), which reduces the utility of their detection for classification purpose. However, the presence of aaRS in a virus genome remains a strong complementary argument for their classification within the Mimiviridae.

The statistical nature of features 1 and 2 make them less decisive than features 3 to 6, although none of the later are present in all members (Table 3).

As of today, the sole protein that is unique to the Mimiviridae and found in all fully sequenced genome is a special version of a homolog to a DNA mismatch repair enzyme (MutS7). MutS proteins are ubiquitous in cellular organisms. However, the Mimiviridae MutS7 version is curiously related to the one found in e-proteobacteria and in the mitochondrial genomes of octocorals [12, 44] (Figure 2). Detecting MutS7 homologs in DNA sequences of newly isolated viruses is thus a strong argument in favor of their classification within the Mimiviridae. Thanks to their strong phylogenetic signals, both AsnS and MutS7 can be used as “baits” to detect presumed Mimiviridae members in large metagenomic sequence data sets [44, 73].

4. Conclusion

15 years after the publication of its prototype Mimivirus, the Mimiviridae now appears as a major family of aquatic viruses [89]. Retrospectively, it is quite fortunate that the name “Mimivirus” (for Microbe Mimicking virus) was forged without reference to the original Acanthamoeba host. Viruses from the Mimiviridae family are now known to infect the most phylogenetically diverse spectrum of eukaryotic unicellular organisms, ranging across 6 major phyla: Amoebozoa (Mimivirus), Haptophyta (PgV), Chlorophyta (TetV), Excavata (BsV), Heterokonta (CroV), and perhaps Opisthokonta (Sturgeon virus). In addition to its host diversity, the Mimiviridae family is also the one exhibiting the broadest distribution of genome sizes (from 370 kb for AaV to 1.51Mb for TupanDO), as well as of particle sizes (from 750 nm to 140 nm for icosahedral virions, up to 2.3 µm...
for the tailed Tupanviruses). Despite these huge differences, all currently known members of the family appear to propagate in particles with similar architecture (including an internal lipid membrane), and replicate in the host cytoplasm using their own well-conserved transcription machinery. Yet, their gene contents are also extremely variable, the intersection of which only amounts to 30 strictly conserved core genes [57]. Future work may thus lead to a partition of the current Mimiviridae family into smaller more homogenous distinct families, as we suggested for the Phycodnaviridae.

Table 3. Characteristic features of various Mimiviridae

<table>
<thead>
<tr>
<th>Name</th>
<th>DNA Pol B</th>
<th>RNA Pol II</th>
<th>aaRS²</th>
<th>MCP¹</th>
<th>MutS7</th>
<th>AsnS</th>
<th>Trans-poviron</th>
<th>Viro-phage</th>
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</thead>
<tbody>
<tr>
<td>Mimivirus</td>
<td>Yes</td>
<td>8</td>
<td>4</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td>Megavirus</td>
<td>Yes</td>
<td>8</td>
<td>7</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Moumouivirus</td>
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<td>8</td>
<td>5</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>CroV</td>
<td>Yes</td>
<td>8</td>
<td>1</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>PgV</td>
<td>Yes</td>
<td>8</td>
<td>0</td>
<td>2</td>
<td>Yes</td>
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<td>No</td>
<td>Yes</td>
</tr>
<tr>
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<td>Yes</td>
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<td>0</td>
<td>3</td>
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<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
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<td>8</td>
<td>0</td>
<td>1</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>BsV</td>
<td>Yes</td>
<td>7</td>
<td>2</td>
<td>4</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>AaV</td>
<td>Yes</td>
<td>8</td>
<td>0</td>
<td>2</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>TupanSL</td>
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<td>8</td>
<td>20</td>
<td>3</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>TupanDO</td>
<td>Yes</td>
<td>7</td>
<td>20</td>
<td>3</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
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</tr>
<tr>
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<td>D</td>
<td>0</td>
<td>1</td>
<td>No</td>
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<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>

Abbreviations are the same as in Table 1.

¹: YlmV: Yellowstone lake mimivirus (NC_028104, 73689 bp). This metagenomic assembly is erroneously listed as “complete”, while it lacks most of the Mimiviridae core genes, including the essential B-type DNA polymerase and most of the transcriptional apparatus. ²: Number of distinct encoded RNA polymerase subunits. YlmV only exhibits the minor subunit D. ³: Number of encoded amino-acyl tRNA synthetases. ⁴: Number of encoded major capsid protein paralogs.
373

374 3.2. Figures

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Phylogeny of the Mimiviridae based on the DNA polymerase B. Diverse algae-infecting “Phycodnaviruses” (prasinovirus OtV5, OlV1; raphidovirus HaV1; chlorovirus PbCV1, AtCV1) clearly cluster as an outgroup (with high confidence). The Mimiviridae members cluster in different proposed subfamilies (from the bottom up): The “Megavirinae” (in red, closest to Mimivirus), a new clade including BsV (in blue, mostly defined by Klosneuvirus-like metagenomics assemblies), the Mesomimivirinae (in green, with PgV and CeV). These proposed subfamilies are highly supported (100% bootstrap values). In contrast, CroV (in indigo), AaV and TetV (in green), remain isolated. This tree was produced from an alignment of 23 sequences (531 sites), using neighbor joining and the JTT substitution model. Bootstrap values are indicated on each branch. The computations were performed on the MAFFT online service [49]. NCBI accession numbers are given for each sequence. The prefix “Meta” indicates sequences from non-isolated viruses.
Figure 2. Phylogeny of the Mimiviridae based on MutS7. MutS sequences from Epsilon proteobacteria (from the genus Arcobacter, and Sulfurospirillum) are used as root, as they are most likely the source of the Mimiviridae MutS7 gene (MutS homologs are present throughout the epsilon division). The tree suggests with great confidence that the mitochondrial MutS present in octocorals (in grey) and that present in all Mimiviridae have a common origin. The clustering of the Mimiviridae into several subfamily is mostly consistent with that suggested by the DNA polymerase phylogeny (Figure 1). The algae-infecting Mimiviridae (in green) now appears monophyletic. This tree was produced from an alignment of 23 sequences (479 sites), using neighbor joining and the JTT substitution model. Bootstrap values are indicated on each branch. The computations were performed on the MAFFT online service [49]. NCBI accession numbers are given for each sequence. The prefix “Meta” indicates sequences from non-isolated viruses. The prefix “Partial” sequences from amplicons.

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Conflicts of Interest: The authors declare no conflict of interest. The founding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the decision to publish the results”.

References


