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Origins and Evolution of the Global RNA Virome

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ABSTRACT Viruses with RNA genomes dominate the eukaryotic virome, reaching enormous diversity in animals and plants. The recent advances of metaviromics prompted us to perform a detailed phylogenomic reconstruction of the evolution of the dramatically expanded global RNA virome. The only universal gene among RNA viruses is the gene encoding the RNA-dependent RNA polymerase (RdRp). We developed an iterative computational procedure that alternates the RdRp phylogenetic tree construction with refinement of the underlying multiple-sequence alignments. The resulting tree encompasses 4,617 RNA virus RdRps and consists of 5 major branches; 2 of the branches include positive-sense RNA viruses, 1 is a mix of positive-sense (+) RNA and double-stranded RNA (dsRNA) viruses, and 2 consist of dsRNA and negative-sense (−) RNA viruses, respectively. This tree topology implies that dsRNA viruses evolved from +RNA viruses on at least two independent occasions, whereas −RNA viruses evolved from dsRNA viruses. Reconstruction of RNA virus evolution using the RdRp tree as the scaffold suggests that the last common ancestors of the major branches of +RNA viruses encoded only the RdRp and a single jelly-roll capsid protein. Subsequent evolution involved independent capture of additional genes, in particular, those encoding distinct RNA helicases, enabling replication of larger RNA genomes and facilitating virus genome expression and virus-host interactions. Phylogenomic analysis reveals extensive gene module exchange among diverse viruses and horizontal virus transfer between distantly related hosts. Although the network of evolutionary relationships within the RNA virome is bound to further expand, the present results call for a thorough reevaluation of the RNA virus taxonomy.

IMPORTANCE The majority of the diverse viruses infecting eukaryotes have RNA genomes, including numerous human, animal, and plant pathogens. Recent advances of metagenomics have led to the discovery of many new groups of RNA viruses in a wide range of hosts. These findings enable a far more complete reconstruction of the evolution of RNA viruses than was attainable previously. This reconstruction reveals the relationships between different Baltimore classes of viruses and indicates extensive transfer of viruses between distantly related hosts, such as plants and animals. These results call for a major revision of the existing taxonomy of RNA viruses.

KEYWORDS RNA virus, evolution, virome, RNA-dependent RNA polymerase, capsid protein

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Early evolution of life is widely believed to have first involved RNA molecules that functioned both as information storage devices and as catalysts (ribozymes) (1, 2). Subsequent evolution involved the emergence of DNA, the dedicated genomic material, and proteins, the ultimate operational molecules. RNA molecules remained central for translating information from genes to proteins (mRNA), for the functioning of the translation machinery itself (rRNA and tRNA), and for a variety of regulatory functions (various classes of noncoding RNA that are increasingly discovered in all life forms) (3). Viruses with RNA genomes (referred to as “RNA viruses” here) that do not involve DNA in their genome replication and expression cycles (4, 5) can be considered to represent the closest extant recapitulation and, possibly, a relic of the primordial RNA world.

RNA viruses comprise 3 of the 7 so-called Baltimore classes of viruses that differ with respect to the nature of the genome (i.e., the nucleic acid form that is packaged into virions) and correspond to distinct strategies of genome replication and expression: positive-sense (+) RNA viruses, double-stranded (ds) RNA viruses, and negative-sense (–) RNA viruses (6). The +RNA viruses use the simplest possible strategy of replication and expression as the same molecule functions as both genome and mRNA (7). Most likely, the first replicators to emerge in the RNA world, after the evolution of translation, resembled +RNA viruses (8). Because the +RNA released from a virion can be directly used for translation to produce viral proteins, virions of +RNA viruses contain only structural proteins, in addition to the genome. In contrast, –RNA and dsRNA viruses package their transcription and replication machineries into their virions because these are necessary to initiate the virus reproduction cycles (9, 10).

RNA viruses comprise a major part of the global virome. In prokaryotes, the known representation of RNA viruses is narrow. Only one family of +RNA viruses (*Leviviridae*) and one family of dsRNA viruses (*Cystoviridae*) are formally recognized, and furthermore, their members have limited host ranges. No –RNA viruses have been isolated from prokaryotes (4, 11, 12). Although recent metagenomic studies suggest that genetic diversity and host range of prokaryotic RNA viruses could be substantially underestimated (13, 14), it appears that the scope of the prokaryotic RNA virome is incomparably less than that of the DNA virome. As discussed previously, potential causes of the vast expansion of the RNA virome in eukaryotes might include the emergence of the compartmentalized cytosol that provided a hospitable, protective environment for RNA replication that is known to be associated with the endoplasmic reticulum and other membrane compartments (11). Conversely, the nuclear envelope could be a barrier that prevents the access of DNA viruses to the host replication and transcription machineries and thus partially relieves the stiff competition that DNA viruses of prokaryotes represent for RNA viruses.

In sharp contrast, the 3 Baltimore classes of RNA viruses dominate the eukaryotic virosphere (11, 15). Eukaryotes from all major taxa are hosts to RNA viruses, and, particularly in plants and invertebrates, these viruses are enormously abundant and diverse (15–17). Until recently, the study of RNA viromes had been heavily skewed toward viruses infecting humans, livestock, and agricultural plants. Because of these limitations and biases, the results of attempts to reconstruct the evolutionary history of RNA viruses were bound to be incomplete. Nonetheless, these studies have yielded important generalizations. A single gene encoding an RNA-dependent RNA polymerase (RdRp) is universal among RNA viruses, including capsidless RNA replicons but excluding some satellite viruses (18). Even within each of the 3 Baltimore classes, virus genomes do not include fully conserved genes other than those encoding RdRps (15). However, several hallmark genes are shared by broad ranges of RNA viruses, including, most notably, those encoding capsid proteins of icosahedral and helical virions of +RNA viruses (19, 20) and those encoding key enzymes involved in virus replication such as distinct helicases and capping enzymes (15).

The RdRp gene and encoded protein are natural targets of evolutionary analysis because they are the only universal gene and encoded protein associated with RNA viruses. However, obtaining strongly supported phylogenies for RdRps is a difficult task due to the extensive sequence divergence, apart from several conserved motifs that are

required for polymerase activity (21–23). RdRps belong to the expansive class of polymerases containing so-called Palm catalytic domains along with the accessory Fingers and Thumb domains (24, 25). In addition to viral RdRps, Palm domain polymerases include reverse transcriptases (RTs) of retroelements and reverse-transcribing viruses and the DNA polymerases that are responsible for genome replication in cellular organisms and diverse DNA viruses. Within the Palm domain class of proteins, RT and the +RNA virus RdRps are significantly similar in sequence and structure and appear to comprise a monophyletic group (22, 24–26). More specifically, the highest similarity is observed between the +RNA virus RdRps and the RTs of group II introns. These introns are widespread retrotransposons in prokaryotes that are thought to be ancestral to the RTs of all other retrotransposons as well as retroviruses and pararetroviruses (recently jointly classified as the order *Ortervirales*) of eukaryotes (27–30).

A phylogenetic analysis of the +RNA virus RdRps revealed only a distant relationship between the leviviruses and the bulk of the eukaryotic +RNA viruses, leaving the ancestral relationships uncertain (31). The origin of eukaryotic +RNA viruses from their prokaryotic counterparts is an obvious possibility. However, given the dramatically greater prevalence of +RNA viruses among eukaryotes compared to the narrow spread of leviviruses and “levi-like viruses” in bacteria, an alternative scenario has been proposed. In this scenario, RdRps of the prokaryotic and eukaryotic +RNA viruses independently descended from distinct RTs (11, 31). Among eukaryotic +RNA viruses, phylogenetic analysis of the RdRps strongly supports the existence of picornavirus and alphavirus “supergroups,” which are further validated by additional signature genes (7, 15). However, both the exact compositions of these supergroups and the evolutionary relationships among many additional groups of viruses remain uncertain. Some RdRp phylogenies suggest a third supergroup combining animal “flavi-like viruses” and plant tombusviruses, but this unification is not supported by additional shared genes and thus remains tenuous (7, 15, 21).

The similarity of RdRps among dsRNA viruses is limited, but these RdRps are similar to various degrees to +RNA virus RdRps. Therefore, different groups of dsRNA viruses might have evolved from different +RNA viruses independently, on multiple occasions (15, 32). Although it is not entirely clear how prokaryotic dsRNA viruses fit into this concept, evidence of an evolutionary affinity between cystoviruses and reoviruses has been presented (33, 34).

For a long time, the evolutionary provenance of –RNA virus RdRps remained uncertain due to their low sequence similarity to other RdRps and RTs (23). However, recent protein structure comparisons point to a striking similarity between the RdRps of –RNA orthomyxoviruses and those of +RNA flaviviruses and dsRNA cystoviruses (35). All these findings notwithstanding, the overall evolutionary relationships among the RdRps of +RNA, –RNA, and dsRNA viruses and RTs remain unresolved. In particular, whether the RdRps of +RNA and –RNA viruses are mono- or polyphyletic is unclear.

Many deep evolutionary connections between RNA virus groups that were originally thought to be unrelated have been delineated using the results of pre-metagenomic-era evolutionary studies. These discoveries culminated in the establishment of RNA virus supergroups (7, 9, 36). However, the evolutionary provenance of many other RNA virus groups remained unclear, as did the relationships between the RNA viruses of the 3 Baltimore classes and retroelements and their ultimate origins. The prospects of substantial progress appeared dim because of the extreme sequence divergence among RNA viruses, which could amount to irrevocable loss of evolutionary information.

Recent revolutionary developments in virus metagenomics (metaviromics) dramatically expanded knowledge of the diversity of RNA viruses and provided an unprecedented amount of sequence data for informed investigation into RNA virus evolution (11, 17, 37). The foremost development was the massive expansion of the known invertebrate virome, which was achieved primarily through meta-transcriptome sequencing of various holobionts. The subsequent phylogenetic analysis revealed previously unknown lineages of +RNA and –RNA viruses and prompted reconsideration of

high-rank virus unifications, such as +RNA virus supergroups (14, 38–41). The RNA viromes of fungi and prokaryotes also underwent notable expansion, albeit it was not as extensive as that of invertebrates (13, 42, 43).

Here we reexamine the evolutionary relationships among and within the 3 Baltimore classes of RNA viruses through a comprehensive analysis of the available genomic and metagenomic sequences. In particular, to build a phylogenetic tree of thousands of viral RdRps, we designed an iterative computational procedure that alternates phylogeny construction with refinement of the underlying multiple alignments. Although RNA viruses have relatively short genomes (~3 to 41 kb), the combined gene repertoire (pangenome) of these viruses includes numerous genes that are shared, to various degrees, by related subsets of RNA viruses. To obtain further insight into virus evolution, we therefore attempted to reconstruct the history of gain and loss of conserved proteins and domains in different virus lineages. We also investigated evolution of the single jelly-roll capsid protein (SJR-CP), the dominant type of capsid protein among +RNA viruses. Our analysis revealed patterns that are generally congruent with the RdRp phylogeny and provide further insights into the evolution of different branches of RNA viruses. Finally, we analyzed a bipartite network in which RNA virus genomes are connected via nodes representing virus genes (44, 45) to identify distinct modules in the RNA virosphere. The results shed light on the evolution of RNA viruses, revealing, in particular, the monophyly of –RNA viruses and their apparent origin from dsRNA viruses, which seem to have evolved from distinct branches of +RNA viruses on at least two independent occasions.

RESULTS

Comprehensive phylogeny of RNA virus RNA-dependent RNA polymerases: overall structure of the tree and the 5 major branches. Amino acid sequences of RdRps and RTs were collected from the nonredundant National Center for Biotechnology Information (NCBI) database and analyzed using an iterative clustering-alignment-phylogeny procedure (see Fig. S1 in the supplemental material) (see Materials and Methods for details). This procedure ultimately yielded a single multiple alignment of the complete set of 4,617 virus RdRp sequences (see Data Set S1 in the supplemental material) and 1,028 RT sequences organized in 50 + 2 clusters (50 clusters of RdRps and 2 clusters of RTs; see Materials and Methods for details). This sequence set did not include RdRps of members of the families *Birnaviridae* and *Permutatetraviridae*, distinct groups of RNA viruses that encompass a circular permutation within the RdRp Palm domain (46) and therefore could not be confidently included in the alignment over their entire lengths.

The phylogenetic tree of RdRps and RTs (Fig. 1; see also Data Set S2) was assembled from a set of trees that represent three hierarchical levels of relationships. At the lowest level, full complements of sequences from each cluster were used to construct cluster-specific trees. At the intermediate level, up to 15 representatives from each cluster were selected to elucidate supergroup-level phylogeny. At the highest level, up to 5 representatives from each cluster were taken to resolve global relationships (Data Set S3A and S4). The final tree (Data Set S2) was assembled by replacing the cluster representatives with the trees from the previous steps.

The large number and immense diversity of the viruses included in our analysis create serious challenges for a systematic, phylogeny-based nomenclature of the identified evolutionary lineages of RNA viruses. Many such lineages consist of viruses newly discovered by metaviromics and are not yet formally classified by the International Committee on Taxonomy of Viruses (ICTV) and therefore cannot be assigned formal names. For the purpose of the present work, we adopted a semiarbitrary naming scheme using the following approach. (i) We use taxon names that had been fully accepted by the ICTV as of March 2018 (47) whenever possible. These names are recognizable through their capitalization and italicization and rank-specific suffixes (e.g., *-virales* for orders and *-viridae* for families). As is the common practice in virus taxonomy, the officially classified members of each ICTV-approved taxon are referred to

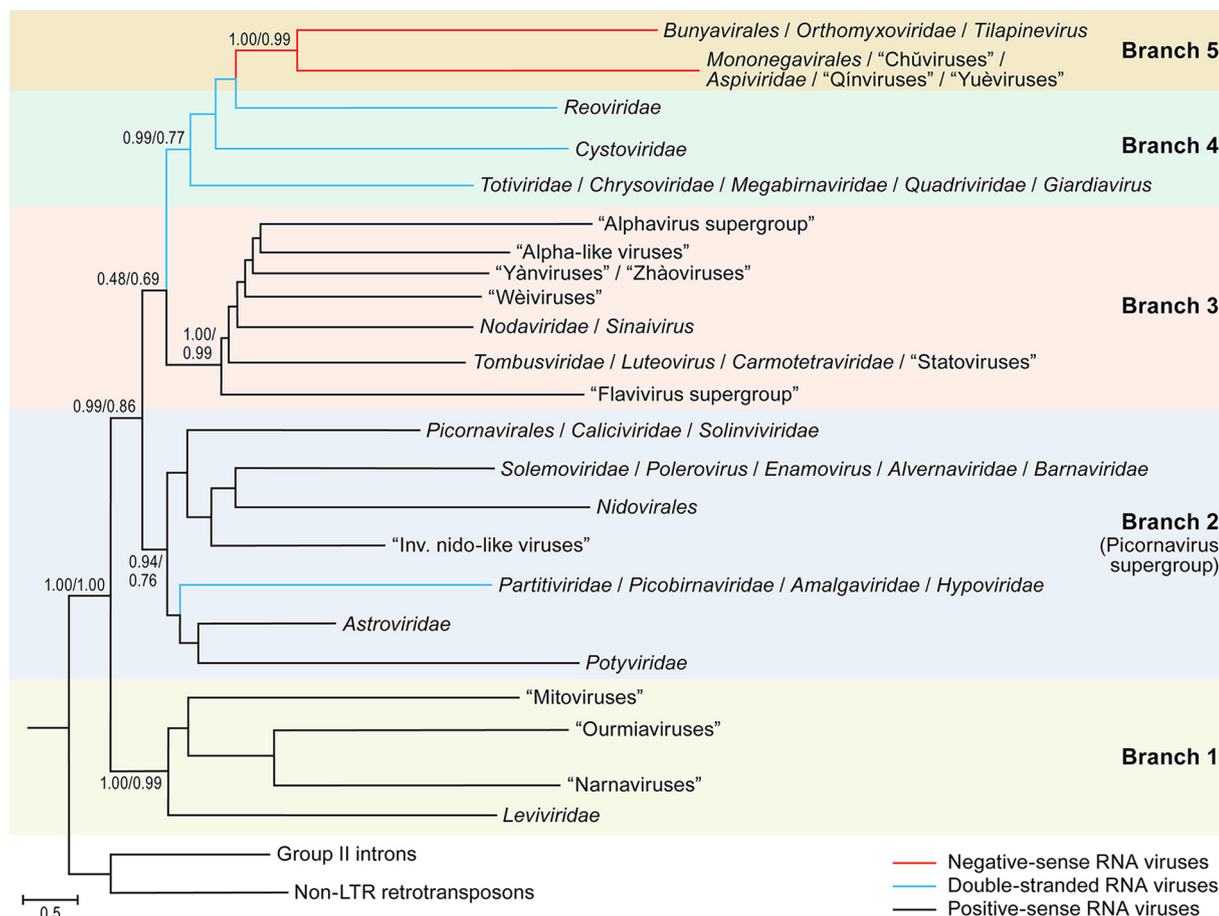


FIG 1 Phylogeny of RNA virus RNA-dependent RNA polymerases (RdRps) and reverse transcriptases (RTs): the main branches (branches 1 to 5). Each branch represents collapsed sequences of the corresponding set of RdRps. The 5 main branches discussed in the text are labeled accordingly. The bootstrap support values obtained by the indicated numerator/denominator calculations are shown for each internal branch. LTR, long-terminal repeat.

via vernacular designations (recognizable through their lack of capitalization and italicization). For instance, the members of the ICTV-approved order *Bunyavirales* are called bunyaviruses, whereas those of the family *Tombusviridae* are called tombusviruses. However, in this work, both taxon and vernacular terms are to be understood *sensu lato*: if our analysis indicates certain viruses to be members of or very closely related to an ICTV-established taxon, we consider them members of that taxon despite the lack of current ICTV recognition. As a result, the order *Bunyavirales* has more members in our analysis than in the official taxonomy. (ii) We use vernacular names in quotation marks for viruses/lineages that are clearly distinct from those covered by the official ICTV framework. Whenever possible, we use names that circulate in the literature (e.g., “hepeliviruses,” “statoviruses”). In the absence of such unofficial names, we name the lineage reminiscent of the next most closely related lineage (e.g., “levi-like viruses” are a clearly distinct sister group to *Leviviridae*/leviviruses). (iii) Monophyletic clusters that transcend the currently highest ICTV-accepted rank (i.e., order) are labeled according to terms circulating in the literature (i.e., “alphavirus supergroup,” “flavivirus supergroup,” and “picornavirus supergroup”). (iv) Lineages represented by a single virus are labeled with the respective virus name.

Rooting the phylogenetic tree, generated using PhyML (48), between the RTs and RdRps resulted in a well-resolved topology of RNA viruses in which the tree splits into 5 major branches, each including a substantial diversity of viruses (Fig. 1).

Branch 1 consists of leviviruses and their eukaryotic relatives, namely, “mitoviruses,” “narnaviruses,” and “ourmiaviruses” (the latter three terms are placed in quotation

marks as our analysis contradicts the current ICTV framework, which classifies mitoviruses and narnaviruses as members of one family, *Narnaviridae*, and ourmiaviruses as members of a free-floating genus, *Ourmiavirus*).

Branch 2 (“picornavirus supergroup”) consists of a large assemblage of +RNA viruses of eukaryotes, in particular, those of the orders *Picornavirales* and *Nidovirales*; the families *Caliciviridae*, *Potyviridae*, *Astroviridae*, and *Solemoviridae*, a lineage of dsRNA viruses that includes partitiviruses and picobirnaviruses; and several other, smaller groups of +RNA and dsRNA viruses.

Branch 3 consists of a distinct subset of +RNA viruses, including the “alphavirus supergroup” along with the “flavivirus supergroup,” nodaviruses, and tombusviruses; the “statovirus,” “wèivirus,” “yànvirus,” and “zhàovirus” groups; and several additional, smaller groups.

Branch 4 consists of dsRNA viruses, including cystoviruses, reoviruses, and totiviruses and several additional families.

Branch 5 consists of –RNA viruses.

Each of these 5 major branches of the tree is strongly supported by bootstrap replications (Fig. 1). Assuming the RT rooting of the tree, branch 1, which consists of leviviruses and their relatives infecting eukaryotes, is a sister group to the rest of RNA viruses; this position is highly robust with respect to the choice of the phylogenetic method and parameters. This tree topology is compatible with the monophyly of the RdRps and, by inference, of RNA viruses and the origin of eukaryotic RNA viruses from a prokaryotic RNA virus ancestor shared with leviviruses. The deeper history remains murky. We have no information on the nature of the common ancestor of retroelements and RNA viruses, let alone on whether the ancestor was an RNA virus or a retroelement. However, parsimony considerations suggest that a retroelement ancestor is more likely given that capsids first appeared in the virus part of the tree rather than having been lost in retroelements.

The next split in the tree occurs between branch 2 and the short stem that formally joins branches 3, 4, and 5. However, the unification of branch 3 with branches 4 and 5 is weakly supported and might not reflect actual common ancestry.

Arguably, the most striking feature of the RNA virus tree topology is the paraphyly of +RNA viruses relative to dsRNA and –RNA viruses. Indeed, according to this phylogeny, –RNA viruses evolved from within dsRNA viruses, whereas dsRNA viruses are polyphyletic (Fig. 1). One major group of dsRNA viruses that includes partitiviruses and picobirnaviruses is firmly embedded within +RNA virus branch 2, whereas another, the members of a larger dsRNA virus group that includes cystoviruses, reoviruses, totiviruses, and viruses from several other families comprise a distinct branch, branch 4, which might be related to +RNA virus branch 3 (Fig. 1). This placement of the two branches of dsRNA viruses is conceptually compatible with the previous evolutionary scenarios of independent origins from +RNA viruses. However, the presence of a strongly supported branch combining 3 lineages of dsRNA viruses that infect both prokaryotes and eukaryotes suggests a lesser extent of polyphyly in the evolution of dsRNA viruses than originally proposed (15, 32).

An alternative phylogenetic analysis of the same RdRp alignment using RAXML yielded the same 5 main branches, albeit some with weak support (Data Set S3B). Furthermore, although the dsRNA viruses are split the same way using RAXML as in the PhyML tree, the nested tree topology, in which branch 4 (the bulk of dsRNA viruses) is lodged deep within +RNA viruses and branch 5 (–RNA viruses) is located inside branch 4, is not reproduced (Data Set S3B). Instead, branches 4 and 5 are separate and positioned deep in the tree, immediately above the split between branch 1 and the rest of the RdRps. Given the poor resolution of the RAXML tree and a strong biological argument, namely, the absence of identified –RNA viruses in prokaryotes or protists (with the exception of the “leishbuviruses” infecting kinetoplastids [49, 50]; see also Discussion below), we believe that the tree topology presented in Fig. 1 carries more credence than that shown in Data Set S3B. Nevertheless, these discrepancies emphasize

that utmost caution is due when biological interpretation of deep branching in trees of highly divergent proteins is attempted.

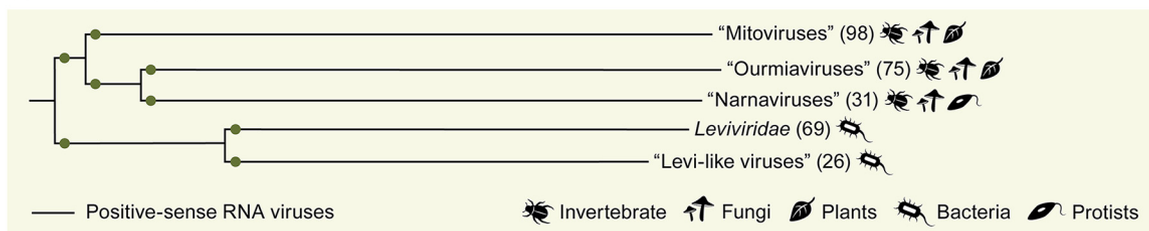
Evolution of the 5 major branches of RNA viruses and reconstruction of gene gain and loss events. (i) Reconstruction of gene gains and losses. The RdRp is the only universal protein of RNA viruses. Accordingly, other viral genes must have been gained and/or lost at different stages of evolution. Thus, after performing the phylogenetic analysis of RdRps, we assigned the proteins and domains shared by diverse viruses to the branches of the RdRp tree. Multiple alignments and hidden Markov model (hmm) profiles were constructed for 16,814 proteins and domains encoded by RNA viruses, and a computational pipeline was developed to map these domains on the viral genomes (Data Set S5). The resulting patterns of domain presence/absence in the branches of the RdRp tree were used to reconstruct the history of the gains and losses of RNA virus genes (or proteins and domains, for simplicity), both formally, using the ML-based Gloome method (51), and informally, from parsimony considerations.

These reconstructions reveal a high level of branch specificity in the evolution of the gene repertoire of RNA viruses. The only protein that is likely to have been gained at the base of the eukaryotic virus subtree (branches 2 to 5) (which also includes the bacterial cystoviruses) is the single jelly-roll capsid protein (SJR-CP) (Fig. S2A). Retroelements lack capsid proteins, and therefore, there is no indication that SJR-CP was present in the hypothetical element that encoded the common ancestor of RdRps and RTs. Furthermore, reconstruction of the evolution of branch 1 (leviviruses and their relatives) argues against the ancestral status of SJR-CP in this branch.

(ii) Branch 1: leviviruses and their descendants. The current RdRp tree topology combined with gene gain-loss reconstruction suggests the following evolutionary scenario for branch 1 (Fig. 2A): a levivirus-like ancestor that, like the extant members of the *Leviviridae*, possessed a capsid protein unrelated to SJR-CP (19, 52) gave rise to naked eukaryotic RNA replicons known as “mitoviruses” and “narnaviruses.” These replicons consist of a single RdRp gene (Fig. 2B) and replicate in mitochondria and in the cytosol of the host cells of fungal and invertebrate hosts, respectively (the latter hosts were identified in metaviromic holobiont analyses) (14, 53). Recently, the existence of plant “mitoviruses” has been reported although it is not known whether these viruses reproduce in the mitochondria (54). The “narnavirus” RdRp is also the ancestor of the RdRp of the expanding group of “ourmiaviruses” (Fig. 2A). “Ourmiaviruses” were originally identified in a narrow range of plants, and genomic analysis revealed the chimeric nature of these viruses, with a “narnavirus”-like RdRp but SJR-CPs and movement proteins (MPs) which were apparently acquired from “picorna-like” and “tombus-like” viruses, respectively (55). Use of metaviromics has led to the identification of numerous related viruses associated with invertebrates, many of which encode distinct SJR-CP variants and some of which acquired an RNA helicase (Fig. 2B and S2) (14). Thus, the evolution of this branch apparently involved the loss of the structural module of leviviruses, which yielded naked RNA replicons that reproduced in the mitochondria of early eukaryotes. A group of these replicons subsequently escaped to the cytosol, which was followed by the reacquisition of unrelated structural modules from distinct lineages of eukaryotic viruses inhabiting the same environment (Fig. 2B). This complex evolutionary scenario emphasizes the key role of modular gene exchange in the evolution of RNA viruses.

(iii) Branch 2: picornavirus supergroup. Expansive branch 2 generally corresponds to the previously described “picornavirus supergroup” (Fig. 1 and 3) (7, 31). Some of the virus groups that were previously considered peripheral members of this supergroup, such as totiviruses and nodaviruses, were relocated to different branches in the present tree (branches 4 and 3, respectively), whereas the viruses of the order *Nidovirales* were moved inside branch 2 from an uncertain position in the tree. Nevertheless, the core of the supergroup remains coherent, suggestive of common ancestry. Within branch 2, 3 major clades are strongly supported (Fig. 3A); however, many of the internal branches are less reliable, so that the relative positions of partitivirus-picobirnavirus, potyvirus-astrovirus, and nidovirus clades within branch 2 remain uncertain.

A Branch 1 (299)



B

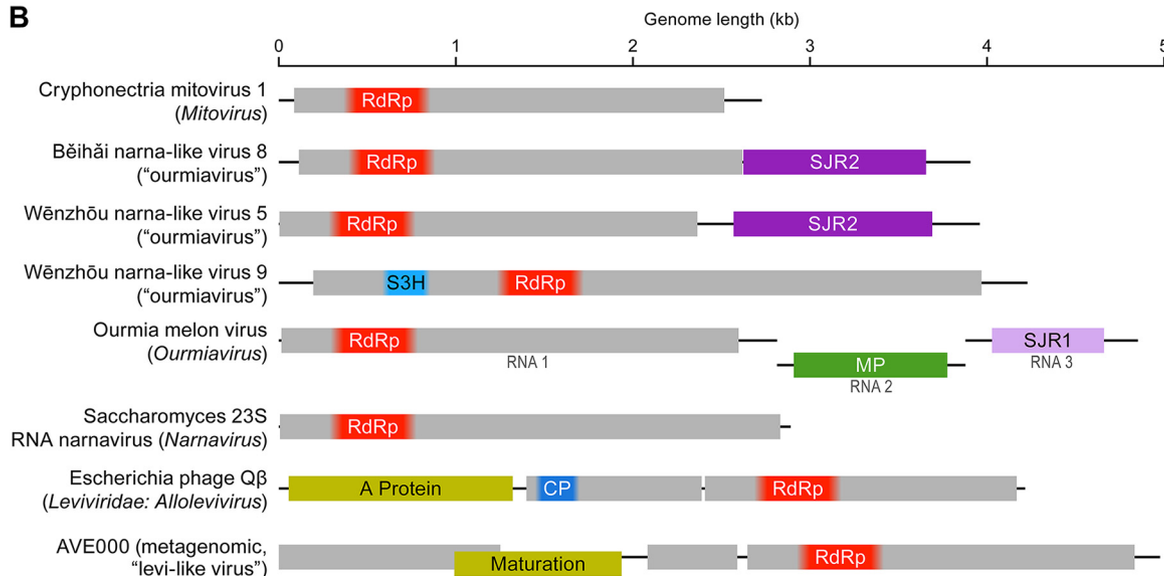
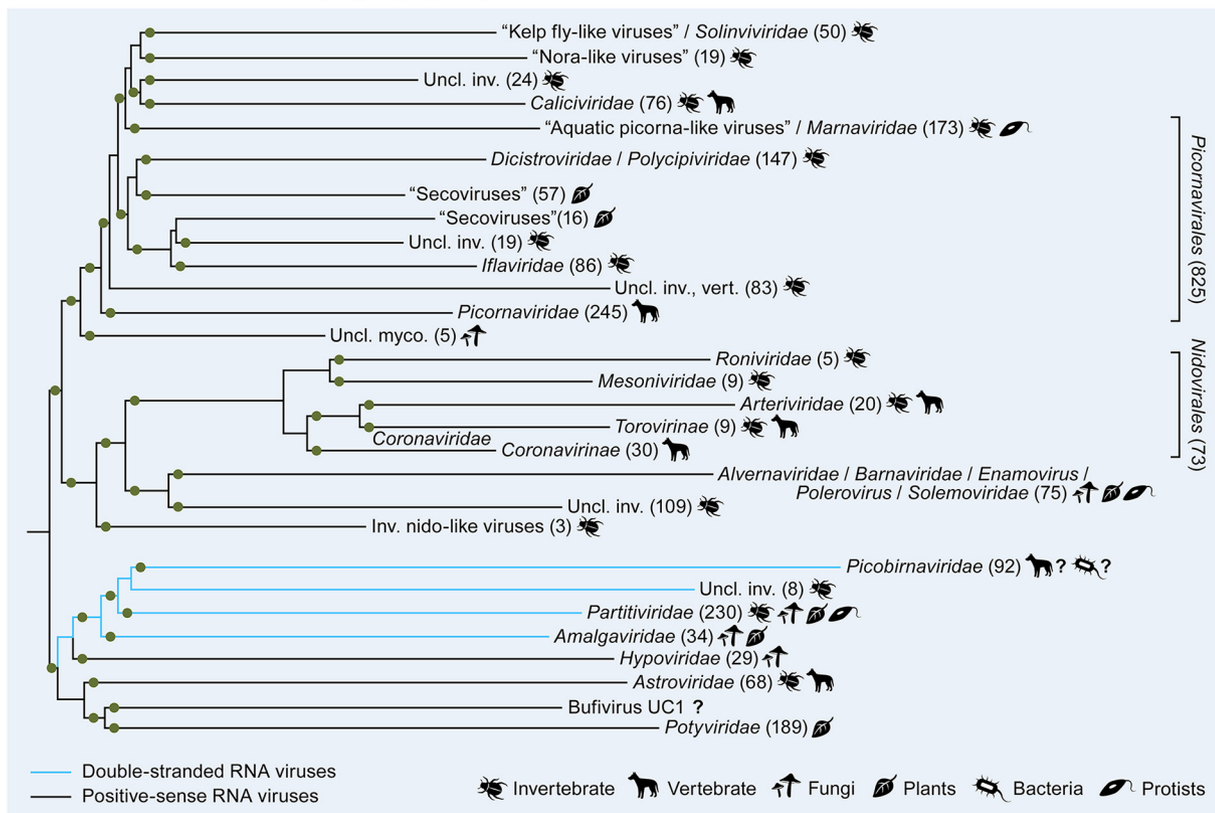


FIG 2 Branch 1 of the RNA virus RNA-dependent RNA polymerases (RdRps): leviviruses and their relatives. (A) Phylogenetic tree of the virus RdRps showing ICTV-accepted virus taxa and other major groups of viruses. Approximate numbers of distinct virus RdRps present in each branch are shown in parentheses. Symbols to the right of the parentheses summarize the presumed virus host spectrum of a lineage. Green dots represent well-supported (≥ 0.7) branches. (B) Genome maps of a representative set of branch 1 viruses (drawn to scale) showing color-coded major conserved domains. Where a conserved domain comprises only a part of the larger protein, the rest of this protein is shown in light gray. The locations of such domains are approximated (indicated by fuzzy boundaries). CP, capsid protein; MP, movement protein; S3H, superfamily 3 helicase; SJR1 and SJR2, single jelly-roll capsid proteins of type 1 and type 2 (see Fig. 7).

The largest and most coherent of the branch 2 clades includes the cornerstone of the picornavirus supergroup, the ~ 826 viruses-strong order *Picornavirales* (56), expanded with caliciviruses, solinviruses, and a multitude of unclassified viruses infecting invertebrates, vertebrates, fungi, protists, and undefined hosts (for viruses discovered by metaviromics) (Fig. 3) (11, 14, 17, 57–59). The second largest, deep-branching clade consists of two lineages that include, respectively, +RNA and dsRNA viruses (Fig. 3). The +RNA virus lineage combines astroviruses and potyviruses, the evolutionary affinity of which is well recognized (31, 60). The dsRNA lineage includes the members of the families *Amalgaviridae*, *Hypoviridae*, *Partitiviridae*, and *Picobirnaviridae*, with each of these families greatly expanded by unclassified affiliates. Finally, the “middle” clade is smaller and less diverse; it encompasses nidoviruses, including the longest of all +RNA virus genomes (61), and solemoviruses with much shorter genomes (Fig. 3). Notably, some members of the family *Luteoviridae* and *Heterocapsa circularisquama* RNA virus, the only known alvernavirus (62), are nested within the solemovirus clade. Given the lack of support beyond the phylogenetic affinity of the RdRps and the dramatic differences in the genomic architectures of nidoviruses and solemoviruses, the possibility that this unification was caused by a tree construction artifact is difficult to rule out (the branch support notwithstanding).

Hypoviruses, representing a group of fungal capsidless RNA replicons, have been traditionally viewed as dsRNA viruses. However, comparisons of genome architectures and phylogenetic analyses suggested that hypoviruses are derivatives of potyviruses

A Branch 2: “Picornavirus supergroup” (1901)



B

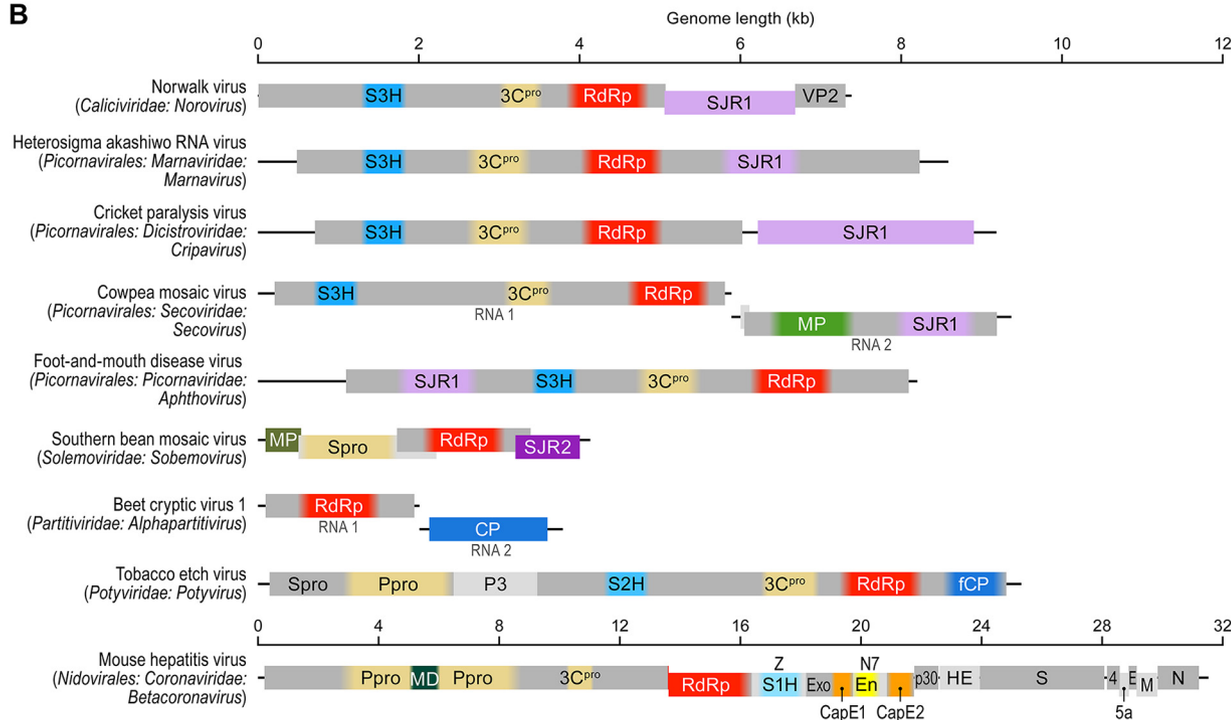


FIG 3 Branch 2 of the RNA virus RNA-dependent RNA polymerases (RdRps): “picornavirus supergroup” of the +RNA viruses expanded to include nidoviruses and two groups of dsRNA viruses, partitiviruses, and picobirnaviruses. (A) Phylogenetic tree of the virus RdRps showing ICTV accepted virus taxa and other major groups of viruses. Approximate numbers of distinct virus RdRps present in each branch are shown in parentheses. Symbols to the right of the parentheses summarize the presumed virus host spectrum of a lineage. Green dots represent well-supported (≥ 0.7) branches. Inv., viruses of invertebrates (many found in holobionts, making host assignment uncertain); myco., mycoviruses; Uncl., unclassified; vert., vertebrate. (B) Genome maps of a representative set of branch 2 viruses (drawn to scale) showing color-coded major conserved domains.

(Continued on next page)

that have lost the capsid protein (63, 64). In the current RdRp tree, hypoviruses cluster with the dsRNA viruses of the partitivirus-picobirnavirus clade rather than with potyviruses (Fig. 3). Whether this position is an artifact of tree construction or whether hypoviruses actually share the RdRps with dsRNA viruses is unclear.

The partitivirus-picobirnavirus clade within branch 2 represents a transition to the *bona fide* dsRNA Baltimore class (Fig. 3). Typical partitiviruses and picobirnaviruses have minimalist genomes that consist of two dsRNA segments encapsidated separately into distinct 120-subunit T=1 capsids (65–68). These genome segments encode, respectively, RdRps and CPs that are clearly homologous between the two families. The CPs of the partitivirus-picobirnavirus clade have been suggested to be distantly related to those of other dsRNA viruses that belong to branch 4 (33, 69). Notably, this clade also includes some naked RNA replicons that reproduce in algal mitochondria or chloroplasts, use a mitochondrial genetic code, and, in terms of lifestyle, resemble “mitoviruses” (14, 70, 71). By analogy, the origin of the partitivirus-picobirnavirus group in an as-yet-undiscovered lineage of prokaryotic RNA viruses seems likely. More specifically, this group of dsRNA viruses could have evolved through reassortment of genomic segments encoding, respectively, a +RNA virus RdRp of branch 2 (possibly a naked RNA replicon) and a dsRNA virus capsid protein related to those of branch 4 viruses. The most recently evolved branch of partitiviruses is characterized by larger, 4- to 6-partite genomes, in contrast to the mono- or bipartite genomes in the deeper branches (14). This observation emphasizes a major tendency in virus evolution: increases in genome complexity via gradual acquisition of accessory genes (72).

Apart from the SJR-CP, an apparently ancestral protein that is likely to represent a shared derived character (synapomorphy) of branch 2 is a serine protease that is present in members of the order *Picornavirales* (with the diagnostic substitution of cysteine for the catalytic serine), members of the potyvirus-astrovirus clade, solemoviruses, alvernavirus, and nidoviruses (Fig. 3B; see also Fig. S2B). As demonstrated previously, this viral protease derives from a distinct bacterial protease, probably of mitochondrial origin, which is compatible with an early origin of branch 2 in eukaryotic evolution (31).

The reconstruction of protein gain and loss, together with the comparison of genome architectures in this branch, revealed extensive rearrangements as well as gene and module displacement (Fig. 3B; see also Fig. S2B). Branch 2 includes viruses with relatively long genomes and complex gene repertoires (nidoviruses, potyviruses, and many members of *Picornavirales*) along with viruses with much shorter genomes and minimal sets of genes (astroviruses and solemoviruses). Clearly, evolution of branch 2 viruses involved multiple gene gains. Of special note is the gain of 3 distinct helicases in 3 clades within this branch: superfamily 3 helicases (S3H) in members of *Picornavirales*, superfamily 2 helicases (S2H) in potyviruses, and superfamily 1 helicases (S1H) in nidoviruses (Fig. 3B; see also Fig. S2B). This independent, convergent gain of distinct helicases reflects the trend noticed early in the study of RNA virus evolution, namely, that most viruses with genomes longer than ~6 kb encode helicases, whereas smaller ones do not. This difference conceivably exists because helicase activity is required for the replication of longer RNA genomes (73). Another notable feature is the change of virion morphologies among potyviruses (replacement of ancestral SJR-CP by an unrelated CP forming filamentous virions) and nidoviruses (displacement by a distinct nucleocapsid protein). The dramatic change in virion morphology and mode of ge-

FIG 3 Legend (Continued)

Where a conserved domain comprises only a part of the larger protein, the rest of this protein is shown in light gray. The locations of such domains are approximated (indicated by fuzzy boundaries). 3C^{Pro}, 3C chymotrypsin-like protease; CP, capsid protein; E, envelope protein; En, nidoviral uridylylate-specific endoribonuclease (NendoU); Exo, 3'-to-5' exoribonuclease domain; fCP, capsid protein forming filamentous virions; M, membrane protein; MD, macrodomain; MP, movement protein; MT, ribose-2-O-methyltransferase domain; N, nucleocapsid protein; N7, guanine-N7-methyltransferase; Ppro, papain-like protease; SJR1 and SJR2, single jelly-roll capsid proteins of type 1 and type 2; spike, spike protein; S1H, superfamily 1 helicase; S2H, superfamily 2 helicase; S3H, superfamily 3 helicase; VP2, virion protein 2; Z, Zn-finger domain; Spro, serine protease; P3, protein 3. Distinct hues of same color (e.g., green for MPs) are used to indicate cases where proteins that share analogous function are not homologous.

nome encapsidation might have been necessitated by the inability of SJR-CP-based icosahedral capsids to accommodate the larger genomes of the ancestral potyviruses and nidoviruses. In addition, nidoviruses gained capping enzymes (CapE) (Fig. 3B; see also Fig. S2B) that most likely were acquired independently of the capping enzymes of other RNA viruses. Nidoviruses also gained ribonucleases and other accessory proteins that are involved in genome replication, virulence, and other aspects of the infection cycle of these largest known RNA viruses (61, 74–76).

(iv) Branch 3: “alphavirus supergroup” and “flavivirus supergroup” and the extensive diversity of “tombus-like viruses.” Branch 3 is the part of the +RNA virus RdRp tree that underwent the most dramatic rearrangements compared to previous versions. This branch consists of two strongly supported clades of +RNA viruses: (i) the assemblage that originally was defined as the “alphavirus supergroup” (7, 15) joined by several additional groups of viruses and (ii) flaviviruses and related viruses (“flavivirus supergroup”; Fig. 1 and 4). In the former clade, the alphavirus supergroup encompasses an enormous diversity of plant, fungal, and animal +RNA viruses and consists of 3 well-supported lineages, namely, tymoviruses, virgaviruses/alphaviruses/endornaviruses, and hepeviruses/benyviruses, each accompanied by related viruses that often form as-yet-unclassified lineages (Fig. 4A). Within the “alphavirus supergroup” alone, the genome lengths range from ~6 to ~20 kb. Despite this length variation, all supergroup members harbor a conserved RNA replication gene module encoding a CapE, S1H, and RdRp; the conservation of this module attests to the monophyly of the supergroup (Fig. 4B).

In contrast, virion architectures differ dramatically even within each of the three lineages of the “alphavirus supergroup.” The major structural themes include variants of icosahedral capsids formed by SJR-CP (e.g., bromoviruses and tymoviruses); unrelated icosahedral capsids enveloped in a lipoprotein bilayer (togaviruses); flexuous filamentous capsids formed by a distinct type of CP (alphaflexiviruses, betaflexiviruses, gammaflexiviruses, and closteroviruses); and rigid rod-shaped capsids assembled from another distinct CP (benyviruses and virgaviruses). It was traditionally thought that the latter capsid type is specific to viruses of flowering plants (20). However, the recent discovery of a virgavirus-like CP in invertebrate viruses (e.g., Běihǎi charybdis crab virus 1 [Fig. 4B]) (14) suggests that the emergence of this unique CP fold antedates land colonization by plants at ~100 million years ago (Mya). Yet another “structural” theme is offered by endornaviruses, naked RNA replicons which, similarly to the hypoviruses in branch 2 (see above), originally were classified as dsRNA viruses. However, endornaviruses possess all the hallmarks of the alphavirus supergroup and are clearly derived from +RNA viruses of this group. They seem to have been mislabeled dsRNA viruses due to the accumulation of dsRNA replication intermediates in infected cells (18, 77). A parallel loss of the CP genes apparently occurred in the deltaflexiviruses, which, in RdRp phylogenies, form a sister group to the flexible filamentous gammaflexiviruses (78), and in the umbraviruses that are included in the family *Tombusviridae* based on the RdRp phylogeny. Notably, unlike most other capsidless viruses that are vertically inherited, umbraviruses can hijack capsids of coinfecting luteoviruses for aphid transmission (79).

Within branch 3, the phylogenetically compact alphavirus supergroup is embedded within the radiation of diverse virus groups, including the well-known tombusviruses and nodaviruses, along with several newcomers discovered via metaviromics, such as the “statovirus,” “wèivirus,” “yànvirus,” and “zhàovirus” groups (14, 80, 81) (Fig. 4A). Our RdRp analysis revealed remarkable phylogenetic heterogeneity within and among these groups and split the “tombus-like viruses” into 5 lineages with distinct evolutionary affinities (groups “Uncl. [unclassified] inv. [invertebrates]” and subsets of “tombus-like viruses” and “nodaviruses” in Fig. 4A). This subdivision is also supported by the analysis of the CPs of these viruses (see section on SJR-CP evolution) (Fig. 7). Therefore, in contrast to the alphavirus supergroup, nodaviruses, and the flavivirus supergroup, the term “tombus-like” loses its evolutionary and taxonomic coherence. Accordingly, we use the term “tombusviruses” (without quotation marks) only for one

A Branch 3 (1,208)

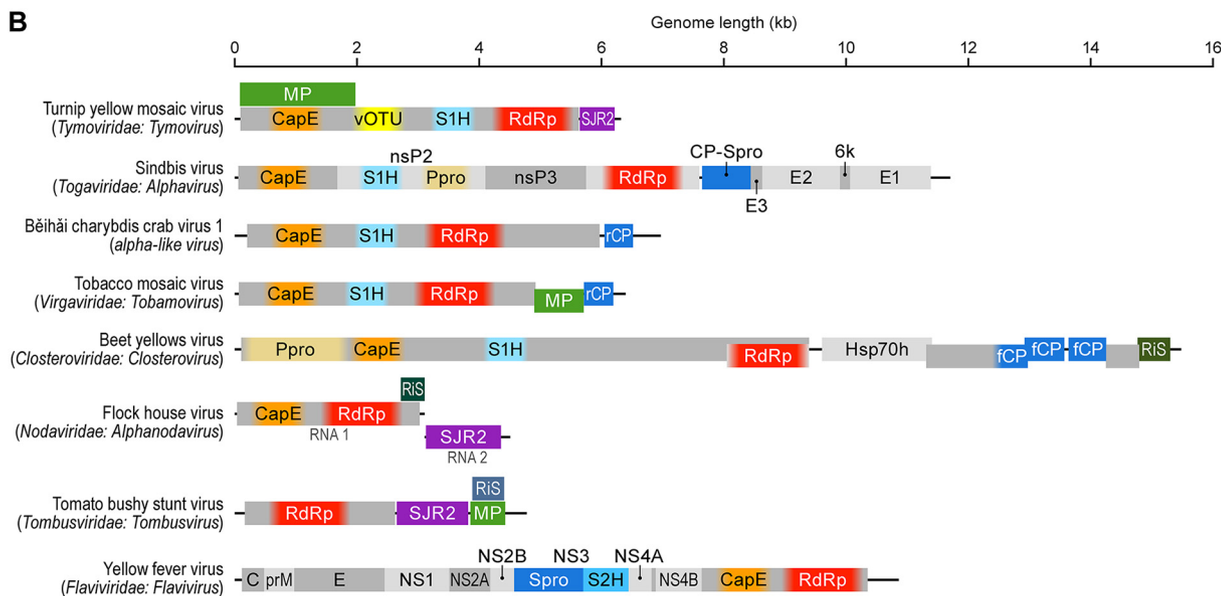
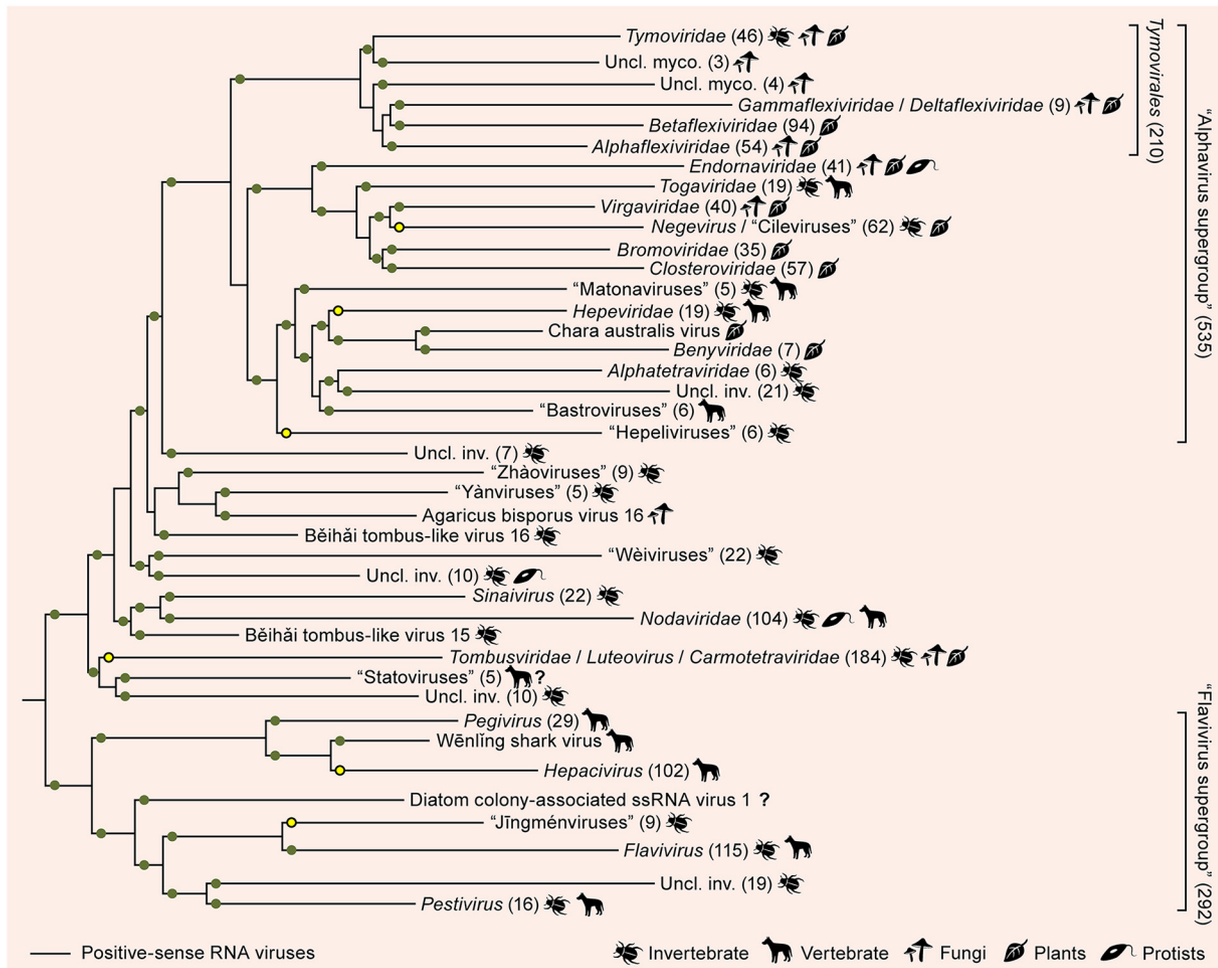


FIG 4 Branch 3 of the RNA virus RNA-dependent RNA polymerases (RdRps): alphavirus superfamily and radiation of related tombusviruses, nodaviruses, and unclassified viruses and flavivirus supergroup. (A) Phylogenetic tree of the virus RdRps showing ICTV-accepted virus taxa and other major groups of viruses. Approximate numbers of distinct virus RdRps present in each branch are shown in parentheses. Symbols to the right of the parentheses summarize the presumed virus host spectrum of a lineage. Green dots represent well-supported (≥ 0.7) branches, whereas yellow dots correspond to weakly supported branches. Inv., viruses of invertebrates (many found in holobionts, making host assignment

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lineage that includes the members of the current family *Tombusviridae* along with a broad variety of related plant and invertebrate holobiont viruses (14).

The previously suggested, tenuous *Flaviviridae-Tombusviridae* affinity is gone in the present tree, although members of both families belong to the same major branch, branch 3. Plant tombusviruses (and members of closely related plant virus genera), representing the only group of “tombus-like viruses” that was available at the time of previous analyses (7, 21), now form but a small twig deep within the large assemblage that we refer to as tombusviruses. Tombusviruses are affiliated with “statoviruses” (80) and with a subset of unclassified viruses from invertebrate holobionts rather than with flaviviruses (Fig. 4A). Flaviviruses now form a separate clade within branch 3, the flavivirus supergroup that includes members of four recognized flaviviral genera (*Pegivirus*, *Hepacivirus*, *Flavivirus*, and *Pestivirus*), the newly discovered “jīngménviruses” with segmented genomes (38), and a variety of unclassified, extremely divergent “flavi-like viruses” of animals and plants. This clade is split into two well-supported lineages; one includes pegiviruses and hepaciviruses, and the other consists of the rest of flaviviruses (Fig. 4A). Flaviviral virions are enveloped, with the envelope proteins forming an external icosahedral shell, whereas the core nucleocapsid is apparently disordered; the evolutionary provenance of the core protein, with its distinct fold, is unclear (19, 82, 83). Notably, flaviviral envelope proteins are class II fusion proteins that are closely related to alphavirus envelope E1 proteins (84). The theme of gene swapping between these distantly related virus groups of branch 3 is further emphasized by the homology between the alphavirus CPs that form icosahedral capsids under the lipid envelopes and the flavivirus nonstructural NS3 proteases that share a chymotrypsin-like fold (84). Because the RdRp tree topology implies that the alphavirus ancestor is more recent than the ancestor of flaviviruses (Fig. 1), such adoption of the NS3 protease for a structural role is suggestive of emerging alphaviruses borrowing their structural module from preexisting flaviviruses (19).

The hallmark of branch 3 is the capping enzyme (CapE), which is present in the entire “alphavirus supergroup” and in flaviviruses (Fig. 4; see also Fig. S2B). A highly divergent version of CapE has been identified in nodaviruses (85) and, in our present analysis, in the additional subset of viruses that grouped with nodaviruses, as well as in a few viruses scattered throughout the clade. Formally, CapE is inferred to be ancestral in the entire branch 3. However, CapEs of “alphavirus supergroup” members, nodaviruses, and flaviviruses are related only distantly to one another, and at least the latter have closer eukaryotic homologs, namely, the FtsJ family methyltransferases (86, 87). Furthermore, tombusviruses, statoviruses, yànviruses, zhàoviruses, wèiviruses, and members of the *Pegivirus-Hepacivirus* lineage of flaviviruses lack CapE, putting into question its presence in the ancestor of this branch (Fig. 4B; see also Fig. S2B). The most credible evolutionary scenario seems to involve convergent acquisition of CapEs on at least 3 independent occasions, recapitulating the apparent history of helicases in branch 2 (see above) (Fig. 3B; see also Fig. S2B). The trend of the capture of helicases by +RNA viruses with longer genomes also holds in branch 3 and includes the acquisition of S1H at the base of the alphavirus supergroup and of S2H by the ancestral flavivirus (Fig. 4B; see also Fig. S2B).

To an even greater extent than in branch 2, the apparent routes of virus evolution in branch 3 involve lineage-specific gene capture that resulted in evolution of complex

FIG 4 Legend (Continued)

uncertain); myco., mycoviruses; uncl., unclassified. (B) Genome maps of a representative set of branch 3 viruses (drawn to scale) showing color-coded major conserved domains. Where a conserved domain comprises only a part of the larger protein, the rest of this protein is shown in light gray. The locations of such domains are approximated (indicated by fuzzy boundaries). C, nucleocapsid protein; CapE, capping enzyme; CP-Spro, capsid protein-serine protease; E, envelope protein; fCP, divergent copies of the capsid protein forming filamentous virions; Hsp70h, Hsp70 homolog; MP, movement protein; NS, nonstructural protein; nsP2 to nsP3, nonstructural proteins; Ppro, papain-like protease; prM, precursor of membrane protein; rCP, capsid protein forming rod-shaped virions; RiS, RNA interference suppressor; S1H, superfamily 1 helicase; S2H, superfamily 2 helicase; SJR2, single jelly-roll capsid proteins of type 2; Spro, serine protease; vOTU, virus OTU-like protease; NS, nonstructural protein. Distinct hues of same color (e.g., green for MPs) are used to indicate the cases when proteins that share analogous function are not homologous.

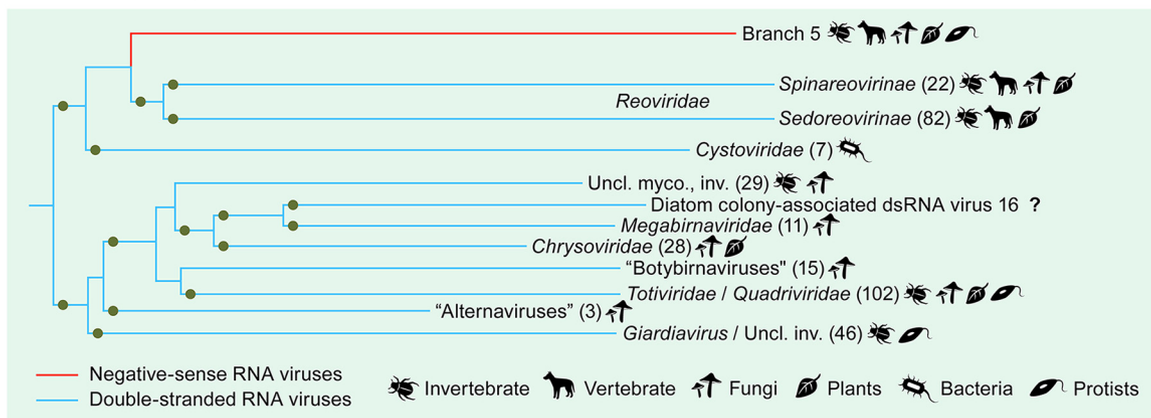
genome architectures (Fig. 4B). The most notable cases are the closteroviruses and divergent flaviviruses that have genomes of up to 20 to 26 kb, rivalling coronaviruses in terms of genome length and the complexity of the gene repertoire (38, 88–90).

The lack of genes assigned to the common ancestor of branch 3 (with the obvious exception of the RdRp) prevents development of a coherent evolutionary scenario for the entire branch. In the case of the clade encompassing the “alphavirus supergroup” and related viruses, a potential common ancestor could be a simple virus that encoded only an RdRp and a SJR-CP, a CP fold most broadly represented in this clade, including diverse tombusviruses, nodaviruses and members of *Bromoviridae*, and *Tymoviridae* within the “alphavirus supergroup.” Proposal of such an ancestor for the flavivirus clade is challenged by the lack of viruses with short and simple genomes among flaviviruses. Indeed, the lengths of the genomes in this clade range from ~9 kb to ~26 kb, with even the shortest ones encoding at least three of the flavivirus signature genes (serine protease [Spro], S2H, and RdRp). One potential clue, however, is provided by “jīngménviruses,” with tetrapartite genomes in which the protease-helicase modules and the RdRps are encoded by separate genome segments; two other segments apparently encode structural proteins of unclear provenance (38). This genome architecture could hint at an ancestral flavivirus genome that was assembled from genes borrowed from preexisting viruses, one of which possessed a divergent “tombus-like virus” RdRp. Although the origins of branch 3 are murky, major trends in its subsequent evolution clearly included lineage-specific gene capture, starting with helicases and CapEs in the ancestors of the major lineages and followed by diverse genes in smaller groups (Fig. 4B).

(v) Branch 4: dsRNA viruses. Branch 4, which joins branch 3 with weak support, includes the bulk of the dsRNA viruses (Fig. 1 and 5). All dsRNA viruses in this branch share a unique virion organization and encode homologous CPs. In particular, the specialized icosahedral capsids of these viruses, involved in transcription and replication of the dsRNA genome inside cells, are constructed from 60 homo- or heterodimers of CP subunits organized on an unusual T=1 (also known as pseudo-T=2) lattice (69, 91). The only exceptions are the chrysovirus, which encode large CPs corresponding to the CP dimers of other dsRNA viruses and form genuine T=1 capsids (92). The icosahedral capsids of partitiviruses and picobirnaviruses, which encode RdRps belonging to branch 2, are also constructed from 60 homodimers (66, 67, 93) and have been suggested to be evolutionarily related to those of the dsRNA viruses from RdRp branch 4 (94) despite little structural similarity between the corresponding CPs. Totiviruses, many of which have “minimal” genomes encoding only RdRps and CPs, comprise one of the two major clades in branch 4, whereas cystoviruses, the only known prokaryotic dsRNA viruses, together with members of the vast family *Reoviridae*, which consists of multisegmented dsRNA viruses infecting diverse eukaryotes, comprise the second clade (Fig. 5). The concept of the closer phylogenetic affinity between cystoviruses and reoviruses appears to be corroborated by the fact that the inner T=1 icosahedral capsid is uniquely encased by the outer icosahedral shell constructed on a T=13 lattice in both families (34). Both cystoviruses and reoviruses appear to have gained many clade-specific genes, in particular, RecA-like packaging ATPases of the former (95) and the CapEs of the latter that are only distantly related to CapEs of other RNA viruses and likely were acquired independently (96, 97) (Fig. 5; see also Fig. S2B).

(vi) Branch 5: –RNA viruses. Branch 5, the 100% supported lineage combining all –RNA viruses, is lodged within branch 4 as the sister group of reoviruses, and this position is upheld by two strongly supported internal branches in the RdRp tree (Fig. 1 and 6). The –RNA branch splits into 2 strongly supported clades. The first clade encompasses the 348 viruses-strong membership of the order *Mononegavirales* (98), along with the members of the distantly related family *Aspiviridae* (99), 3 groups of –RNA viruses discovered through metaviromics (“chūviruses,” “qínviruses,” and “yuèviruses”) (14, 39), and a group of unclassified fungal viruses (Fig. 6A) (42, 100). In contrast to the members of the *Mononegavirales*, most of which possess unsegmented genomes, the remainder of this clade is characterized by bi-, tri-, or even tetraseg-

A Branch 4 (346)



B

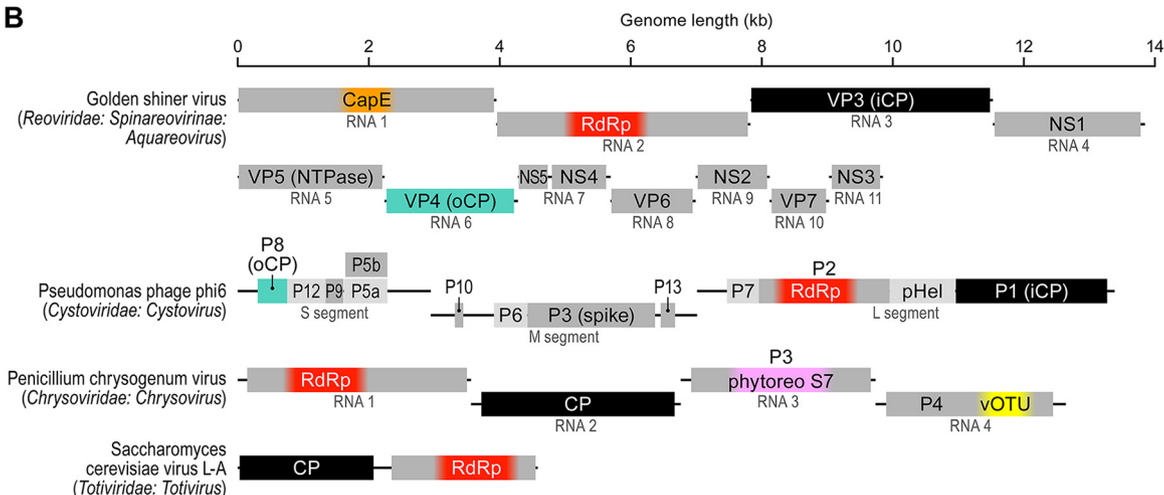
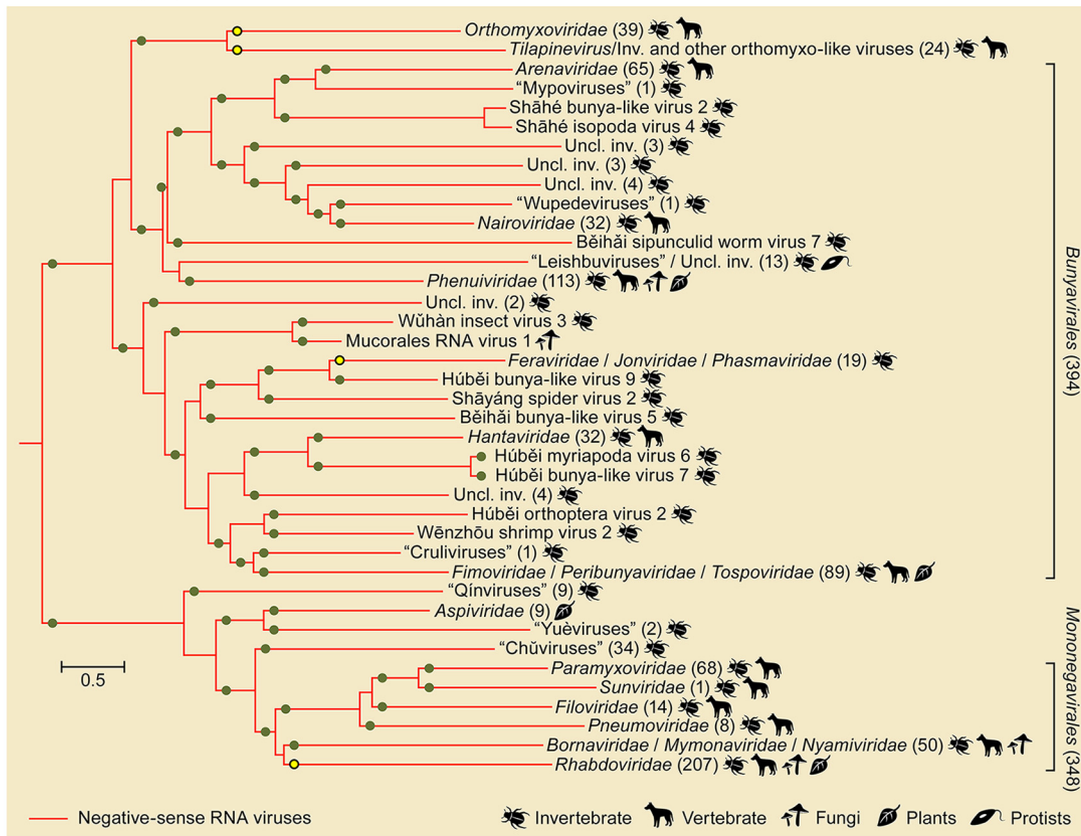


FIG 5 Branch 4 of the RNA virus RNA-dependent RNA polymerases (RdRps): dsRNA viruses of eukaryotes and prokaryotes. (A) Phylogenetic tree of the virus RdRps showing ICTV-accepted virus taxa and other major groups of viruses. Approximate numbers of distinct virus RdRps present in each branch are shown in parentheses. Symbols to the right of the parentheses summarize the presumed virus host spectrum of a lineage. Inv., viruses of invertebrates (many found in holobionts, making host assignment uncertain); myco., mycoviruses; uncl., unclassified. Green dots represent well-supported (≥ 0.7) branches. (B) Genome maps of a representative set of branch 4 viruses (drawn to scale) showing color-coded major conserved domains. Where a conserved domain comprises only a part of the larger protein, the rest of this protein is shown in light gray. The locations of such domains are approximated (indicated by fuzzy boundaries). CapE, capping enzyme; CP, capsid protein; iCP, internal capsid protein; NS, nonstructural protein; NTPase, nucleotide triphosphatase; oCP, outer capsid protein; P, protein; phytoreo57, homolog of S7 domain of phytoreoviruses; pHel, packaging helicase; vOTU, virus OTU-like protease; VP, viral protein. The CPs of totiviruses and chrysoviruses are homologous to iCPs of reoviruses and cystoviruses (black rectangles).

mented genomes (Fig. 6B). The second clade combines the family *Orthomyxoviridae*, the genus *Tilapinevirus* (101), and the large order *Bunyavirales* (394 viruses) (102). The latter order consists of two branches, one of which is the sister group to the orthomyxovirus/tilapinevirus clade (albeit with weak support). The numbers of negative-sense or ambisense genome segments in this clade range from 2 to 3 in most of bunyaviruses to 8 in viruses of the genus *Emaravirus* and the orthomyxovirus/tilapinevirus group (10, 99, 101, 103). A notable acquisition in the first clade is a CapE, whereas members of the second clade share “cap-snatching” endonucleases (En) (10).

Patterns of the single jelly-roll capsid protein evolution. The SJR-CP is the dominant type of CP among +RNA viruses and is also found in members of one family of dsRNA viruses (*Birnaviridae*). Structural comparisons indicate that SJR-CPs of RNA viruses form a monophyletic group and likely were recruited from cellular SJR proteins on a single occasion during the evolution of RNA viruses (19). The short length and high divergence of SJR-CPs preclude adequate resolution in phylogenetic analysis; thus, we

A Branch 5 (859)



B

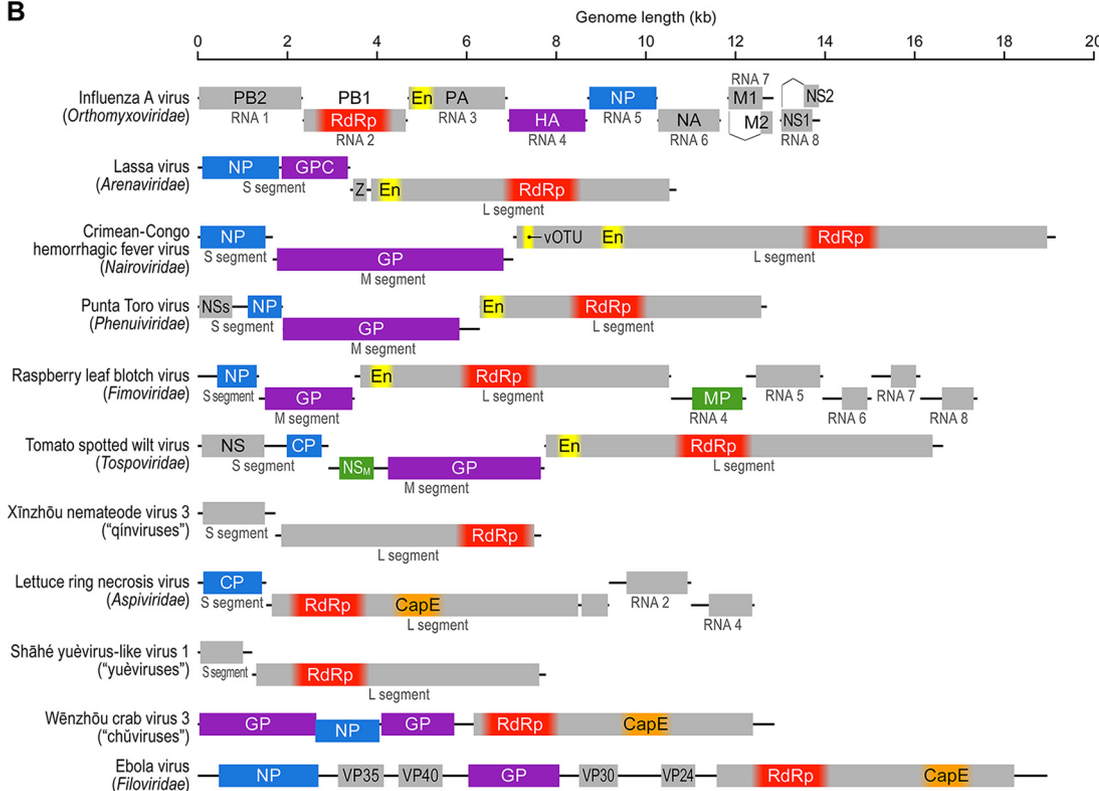


FIG 6 Branch 5 of the RNA virus RNA-dependent RNA polymerases (RdRps): –RNA viruses. (A) Phylogenetic tree of the virus RdRps showing ICTV-accepted virus taxa and other major groups of viruses. Approximate numbers of distinct virus RdRps present in each (Continued on next page)

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performed profile-profile sequence comparisons and clustering of all viral SJR-CP sequences in our data set (see Materials and Methods for details). Analysis of the resulting network reveals patterns that are generally congruent with the RdRp phylogeny and provides further insights into the evolution of different branches of RNA viruses.

At conservative P value thresholds ($P < 1e^{-10}$), the majority of SJR-CPs segregated into two large clusters, both of which contained representatives from RdRp branch 2. Cluster 1 included the members of *Picornavirales* and *Caliciviridae* and diverse “picorna-like viruses” of invertebrates, whereas cluster 2 consisted of the members of the families *Astroviridae*, *Luteoviridae*, and *Solemoviridae* and “sobemo-like viruses” (Fig. S3). In addition, cluster 2 contained members of several families from RdRp branch 3, namely, *Tombusviridae* (and diverse “tombus-like viruses”), *Hepeviridae*, a subgroup of *Nodaviridae*, and “statoviruses.”

At less-restrictive P value thresholds ($P < 1e^{-03}$), all SJR-CPs were interconnected, largely through making contacts to the core of cluster 2. Only “ourmiaviruses” had stronger affinity to picornaviruses in cluster 1 (Fig. 7). This pattern of connectivity is consistent with the radiation of SJR-CPs from a common ancestor, likely resembling sequences from cluster 2 of branch 2. This analysis also reveals high CP sequence divergence among members of some families (e.g., *Bromoviridae*) and numerous cases of apparent CP gene replacement. For instance, the CPs of nodaviruses fall into two groups; one is related to the turreted CPs of tetraviruses, and the other is similar to CPs of tombusviruses, mirroring the RdRp phylogeny (Fig. 7). At a greater phylogenetic distance, CPs of astroviruses and hepeviruses are closely related despite their affiliation with branches 2 and 3, respectively, suggesting CP gene replacement in the ancestor of one of the two families. Given that the CPs of hepeviruses connect to SJR-CPs of other viruses through astroviruses, CP gene replacement most likely occurred in the ancestor of hepeviruses (Fig. S3). Notably, the CPs of “zhàoviruses,” “wèiviruses,” and “tombus-like” and “sobemo-like” viruses (representing a diverse virus assemblage within the solemovirus branch, to the exclusion of bona fide *Solemoviridae*) did not form discrete clusters but rather were affiliated with diverse virus groups, suggesting extensive recombination in these viruses, with multiple CP gene exchanges (Fig. 7). In the case of unclassified “narnaviruses” and “ourmiaviruses,” the CP genes apparently were acquired on more than 3 independent occasions from different groups of viruses, emphasizing the impact of recombination and gene shuffling in the evolution of RNA viruses. Previously, a similar extent of chimerism was also observed among single-stranded DNA (ssDNA) viruses (104, 105), highlighting the evolutionary and functional plasticity of short viral genomes.

The modular gene-sharing network of RNA viruses: gene transfer and module shuffling. The pronounced structural and functional modularity of virus proteomes and the pervasive shuffling of the genomic regions encoding distinct protein modules are key features of virus evolution (11, 15, 17). Therefore, a productive approach to the study of the virosphere that complements phylogenetics is the construction and analysis of networks of gene sharing. Bipartite networks, in which one type of node corresponds to genes and another to genomes, have been employed to investigate the double-stranded DNA (dsDNA) domain of the virosphere (45). This analysis revealed hierarchical modular organization of the network, with several modules that included

FIG 6 Legend (Continued)

branch are shown in parentheses. Symbols to the right of the parentheses summarize the presumed virus host spectrum of a lineage. Green dots represent well-supported (≥ 0.7) branches, whereas yellow dots correspond to weakly supported branches. Inv., viruses of invertebrates (many found in holobionts, making host assignment uncertain); uncl., unclassified. (B) Genome maps of a representative set of branch 5 viruses (drawn to scale) showing color-coded major conserved domains. Where a conserved domain comprises only a part of the larger protein, the rest of this protein is shown in light gray. The locations of such domains are approximated (indicated by fuzzy boundaries). CapE, capping enzyme; CP, capsid protein; EN, “cap-snatching” endonuclease; GP, glycoprotein; GPC, glycoprotein precursor; HA, hemagglutinin; M, matrix protein; MP, movement protein; NA, neuraminidase; NP, nucleoprotein; NS, nonstructural protein; NS_M, medium nonstructural protein; NS_S, small nonstructural protein; PA, polymerase acidic protein; PB, polymerase basic protein; vOTU, virus OTU-like protease; VP, viral protein; Z, zinc finger protein.

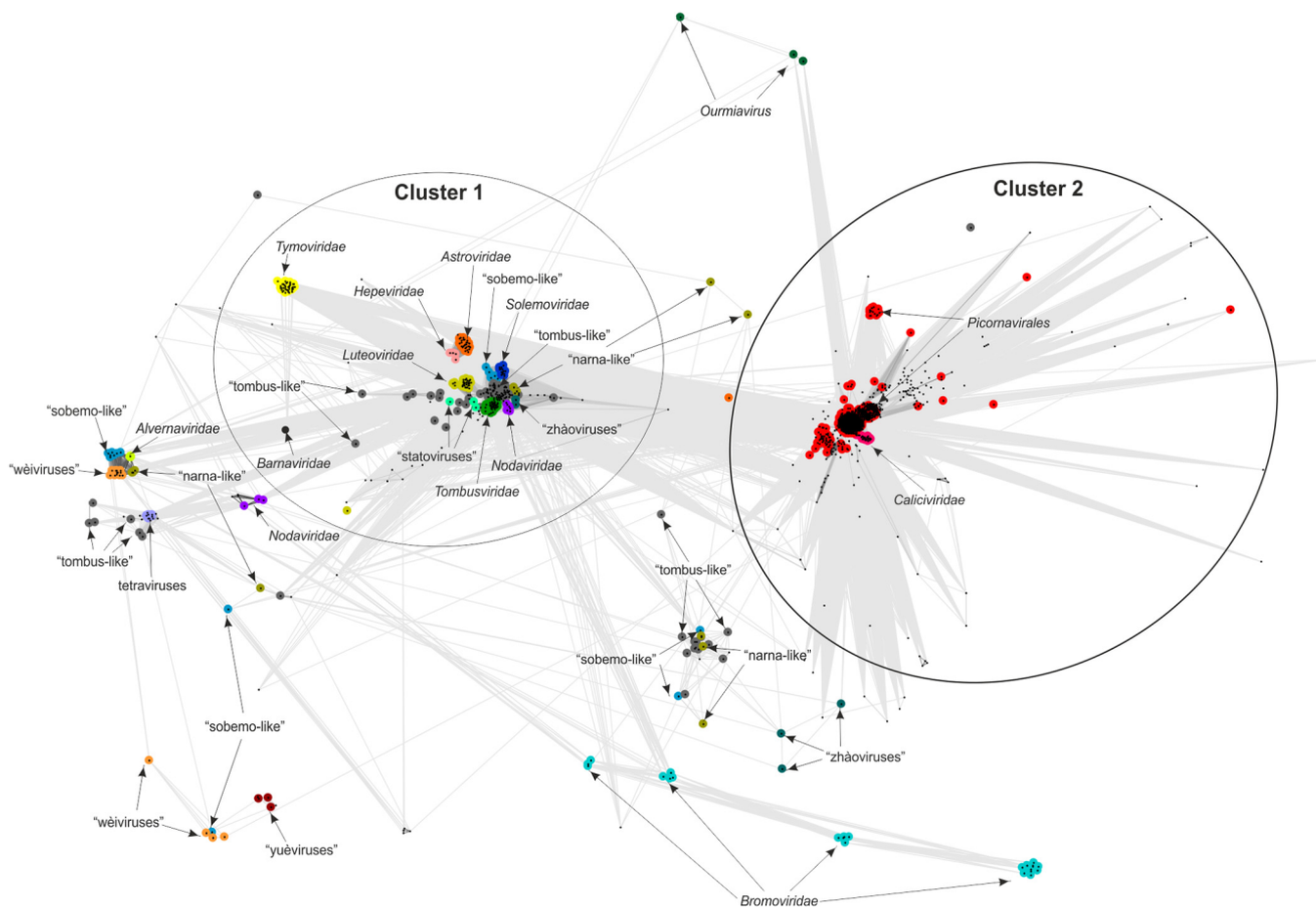


FIG 7 Sequence similarity networks of SJR-CPs. Protein sequences were clustered by the pairwise similarity of their hmm profiles using CLANS. Different groups of SJR-CPs are shown as clouds of differentially colored circles, with the corresponding subgroups labeled as indicated in the figure. Edges connect sequences with CLANS P values of $\leq 1e-03$.

nonobvious connections between disparate groups of viruses (44, 106). Although this type of analysis is less informative for RNA viruses due to the small number of proteins encoded in each viral genome, the “pan-proteome” of RNA viruses is large (Data Set S5), prompting us to experiment with bipartite gene sharing networks for RNA viruses. The initial search for statistically significant modularity identified 54 distinct modules, most of which included a single virus family (Fig. 8A and B). Remarkably, the family *Reoviridae* has been split into 5 modules, highlighting the vast diversity of this family, comparable to that seen in order-level taxa. Among the exceptions, the most expansive module included the viruses of the order *Picornavirales* (module 29), together with those of the family *Caliciviridae* (module 47), which are linked through the conserved suite of genes that includes those encoding SJR-CP, chymotrypsin-like protease, S3H, and the RdRp (Fig. 8A and C). Viruses of the order *Bunyavirales* were also recovered in a single module characterized by the presence of a conserved nucleocapsid (with the exception of the families *Nairoviridae* and *Arenaviridae*) and the cap-snatching endonuclease (module 51; Fig. 8A and C).

The next stage of the network analysis aimed at detecting the supermodules that are formed from the primary modules via connecting genes. The results of the analyses of the supermodules of RNA viruses failed to attain statistical significance due to the small number of shared genes; nevertheless, some notable connections were revealed by this analysis. Specifically, 8 overlapping supermodules were identified (Fig. 8C). The largest and, arguably, most remarkable is a supermodule that combines +RNA, dsRNA, and -RNA viruses that share the capping enzymes, S1H (with the exception of

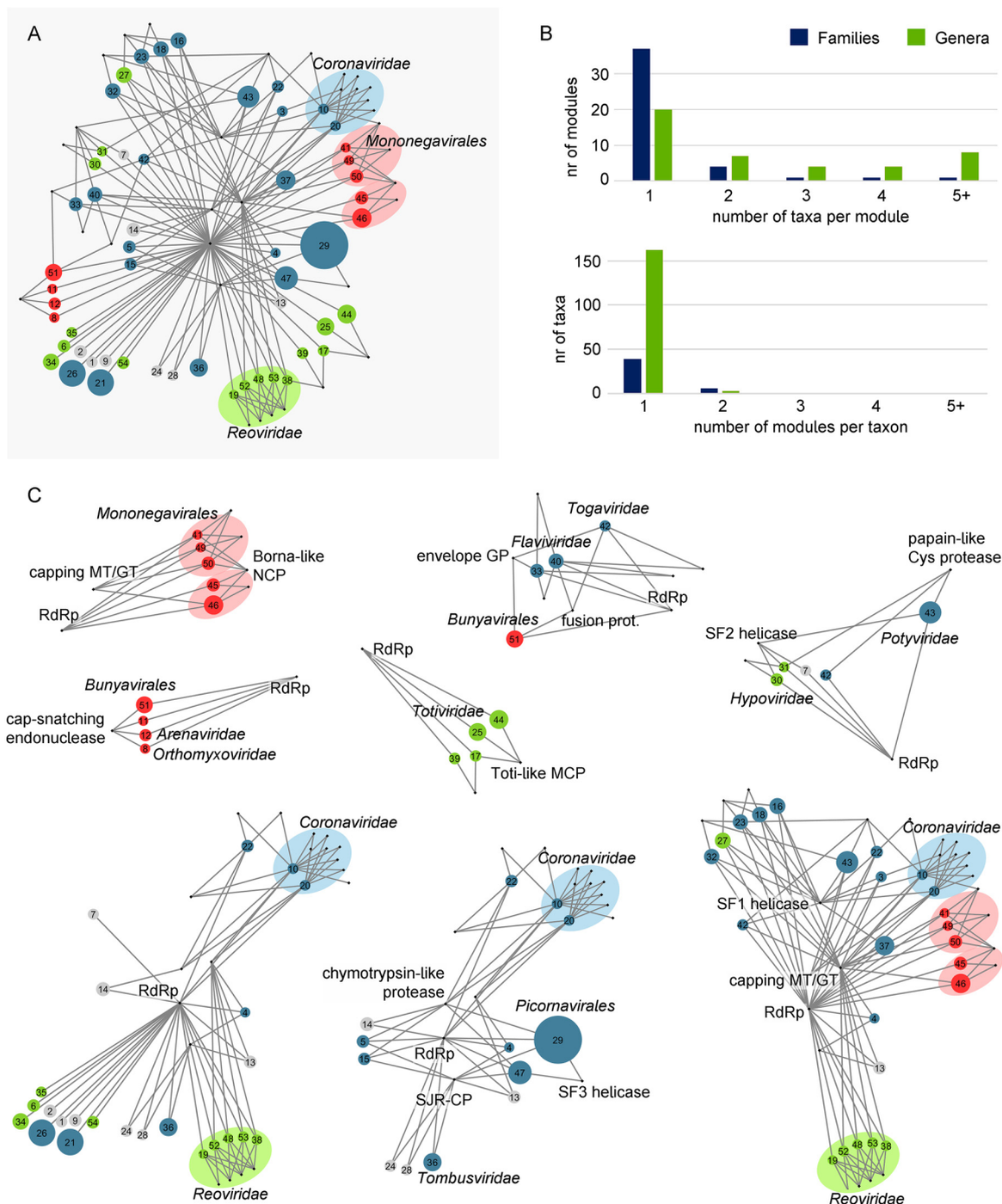


FIG 8 Bipartite network of gene sharing in RNA viruses. (A) Groups of related viruses were identified as the modules of the bipartite genome-gene network (not shown), whereas connector genes were defined as those genes present in two or more modules with prevalence greater than 65%. The network presented in panel A shows viral modules as colored circles (blue, +RNA viruses; green, dsRNA viruses; red, -RNA viruses) linked to the connector genes (black dots) that are present in each module. The size of the circles is proportional to the number of genomes in each module. Shaded ovals indicate statistically significant, first-order supermodules that join modules from taxonomically related groups. (B) Taxonomic analysis of the network modules confirms that most modules contain viruses from a single family and that families do not tend to split among modules. (C) High-order supermodules of the RNA virus network, obtained by iteratively applying a community detection algorithm on the bipartite network of (super)modules and connector genes. GP, glycoprotein; GT, guanylyltransferase; MT, methyltransferase; NCP, nucleocapsid; RdRp, RNA-dependent RNA polymerase; SF, superfamily; SJR-CP, single jelly-roll capsid protein.

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Reoviridae and *Mononegavirales*), and additional connector genes (e.g., the OTU family protease) that link some of the constituent modules. The second largest supermodule combines large subsets of viruses from RdRp branches 2 and 3 that are connected through the SJR-CP and the chymotrypsin-like protease. Another supermodule encompasses enveloped +RNA viruses of the families *Flaviviridae* and *Togaviridae* and –RNA viruses of the order *Bunyavirales* (except for *Arenaviridae*) that share homologous envelope glycoproteins (except for flaviviruses) and class II fusion proteins.

Information from gene sharing is inherently limited for RNA viruses due to the small number of genes in each genome. Nevertheless, the bipartite network analysis revealed prominent “horizontal” connections that are underlain either by actual gene exchange or by parallel acquisition of homologous genes by distinct RNA viruses.

Host ranges of RNA viruses: evolutionary implications and horizontal virus transfer (HVT). RNA viruses have been identified in representatives of all major divisions of eukaryotes, whereas in prokaryotes, members of two families of RNA viruses are known to infect only a limited range of hosts (11, 13, 15, 31). For branch 1 in our phylogenetic tree of RdRps, the route of evolution from leviviruses infecting prokaryotes to eukaryotic “ourmiaviruses” of plants and invertebrates is readily traceable and involves a merger between a levivirus-derived naked RNA replicon that eukaryotes most likely inherited from the mitochondrial endosymbiont with the SJR-CP of a eukaryotic “picorna-like virus.” Notably, such a merger seems to have occurred on at least three other independent occasions in branch 1 because several groups of invertebrate holobiont “narnaviruses” and some “ourmiaviruses” encode distantly related SJR-CPs that apparently were acquired from different groups of plant and animal viruses (Fig. 7).

The case of cystoviruses is less clear given that this clade is sandwiched between eukaryotic viruses in branch 4 and therefore does not seem to be a good candidate for classification as the ancestor of this branch. It appears more likely that the ancestor was a toti-like virus, whereas cystoviruses were derived forms, which would imply virus transfer from eukaryotes to prokaryotes. However, an alternative scenario might be considered. No known prokaryotic viruses are classified in branch 2, but it has been proposed that the picobirnaviruses, for which no hosts have been reliably identified, actually are prokaryotic viruses. This proposal is based on the conspicuous conservation of functional, bacterium-type, ribosome-binding sites (Shine-Dalgarno sequences) in picobirnavirus genomes (107, 108). Should that be the case, viruses of prokaryotes might be lurking among totiviruses as well. Then, branch 4 would stem from a prokaryotic ancestor, obviating the need to invoke virus transfer from eukaryotes to prokaryotes to explain the origin of the cystoviruses.

We made an attempt to quantify the potential horizontal virus transfer (HVT) events in the RNA viruses that represent the 5 major branches of the RdRp tree. The leaves of the tree were labeled with the known hosts, the entropy of the host ranges for each subtree was calculated, and the resulting values were plotted against the distance from the root (Fig. 9). By design, for all branches, entropy (host diversity) drops from the maximum values at the root to zero at the leaves. All branches show substantial host range diversity such that, for example, at the half-distance from the root to the leaves, all branches, except for branch 1, retain at least half of the diversity (Fig. 9). Furthermore, the differences between the branches are substantial, with the highest entropy observed in branch 4 (dsRNA viruses) and branch 5 (–RNA viruses). With all the caveats due to potential errors and ambiguities in host assignment, this analysis strongly suggests that HVT played an important role in the evolution of all major groups of RNA viruses.

DISCUSSION

RNA virus evolution coming into focus. This work was prompted by the advances of metaviromics, which have dramatically increased the known diversity of RNA viruses (11, 13, 17, 37, 57). We reasoned that this expansion of the RNA virosphere could provide an improved understanding of virus evolution. Although further progress of

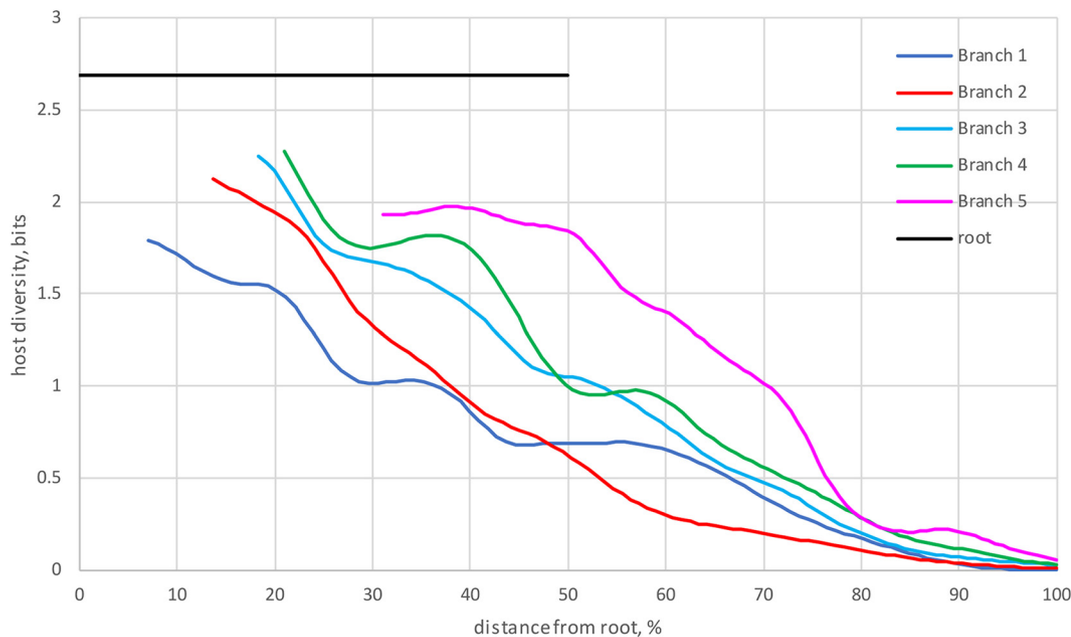


FIG 9 Quantitative analysis of the host range diversity of RNA viruses. The entropy of host ranges is plotted against the ultrameterized tree depth for the 5 main branches of the RdRp phylogeny (see Fig. 1).

metaviromics and enhanced phylogenomic methods will undoubtedly change current ideas, we believe that some key aspects of RNA virus evolution are indeed coming into focus.

The expanded diversity of RNA viruses, combined with the iterative procedure for phylogenetic analysis, allowed us to obtain a tree of all RdRps and the most closely related RTs in which the main branches are strongly supported and thus appear to be reliable (Fig. 1). To our knowledge, the picture of RNA virus evolution emerging from the tree has not been presented previously. The tree seems to clarify the relationships between the 3 Baltimore classes of RNA viruses by revealing the nested tree structure in which dsRNA viruses evolved, on at least two occasions, from +RNA viruses, whereas –RNA viruses evolved from a distinct group of dsRNA viruses.

The derivation of –RNA viruses from dsRNA viruses is, arguably, the most unexpected outcome of the present analysis, considering the lack of genes (other than the RdRp gene) shared by these virus classes. Clearly, given that the primary evidence behind the derivation of –RNA viruses from within dsRNA viruses comes from deep phylogeny, extreme caution is due in the interpretation of this observation. However, the pronounced similarity between the three-dimensional (3D) structures of the RdRps of –RNA influenza virus A and bacteriophage $\phi 6$ dsRNA cystovirus (35) is compatible with our findings. Further, because virtually no –RNA viruses are known in prokaryotes or unicellular eukaryotes (with the single exception of the “leischbuviruses” in parasitic trypanosomatids [50], which were likely acquired from the animal hosts of these protists), their later origin from a preexisting group of +RNA or dsRNA viruses appears most likely.

The +RNA to dsRNA to –RNA scenario of RNA virus genome evolution also makes sense in terms of the molecular logic of genome replication-expression strategies. Indeed, +RNA viruses use the simplest genomic strategy and, in all likelihood, represent the primary pool of RNA viruses. The dsRNA viruses conceivably evolved when a +RNA virus switched to encapsidating a replicative intermediate (dsRNA) together with the RdRp. Naked replicons similar to those of “mitoviruses,” hypoviruses, and endornaviruses might have been evolutionary intermediates in this process. This switch does not seem to be as “easy” and common as previously suspected (15, 32) but, nevertheless, appears to have occurred at least twice during the evolution of RNA viruses. The

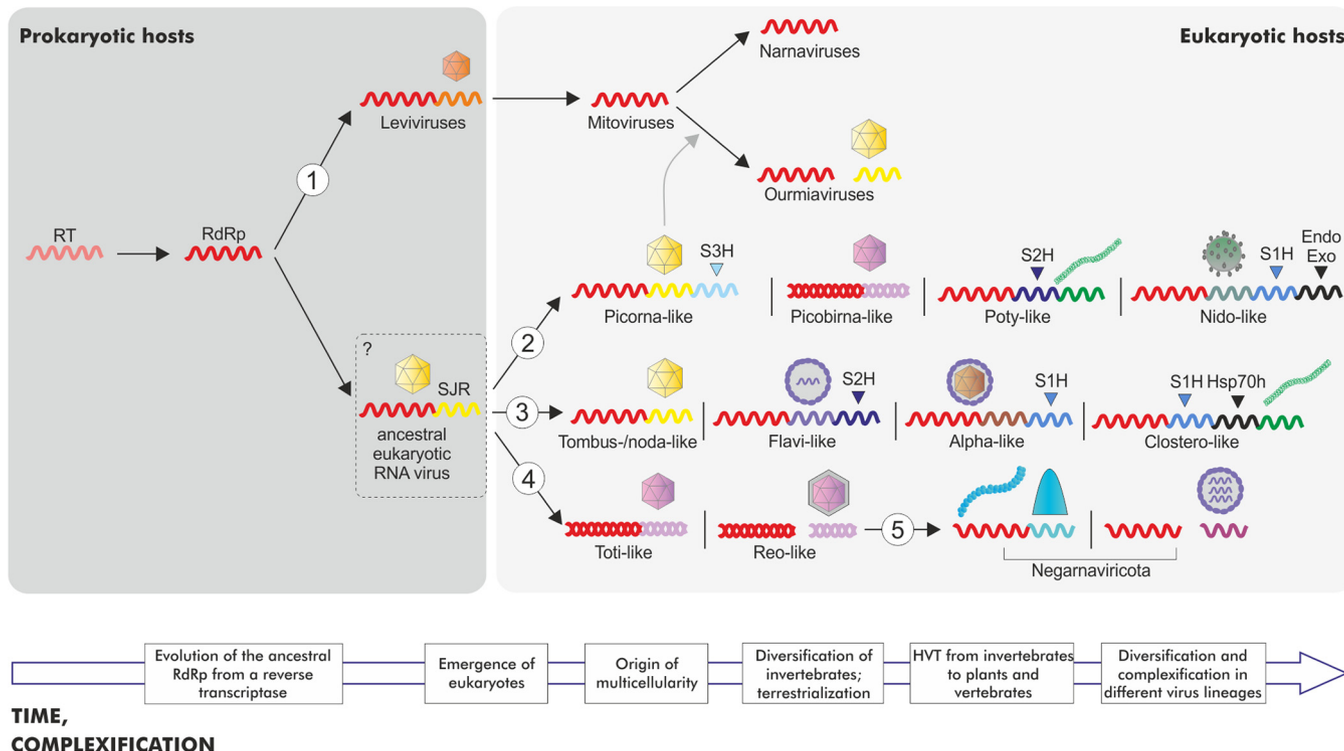


FIG 10 A general scenario of RNA virus evolution. The figure is a rough scheme of the key steps of RNA virus evolution inferred in this work. The main branches from the phylogenetic tree of the RdRps are denoted with the numbers 1 to 5 as described in the Fig. 1 legend. Only the genes corresponding to RdRp, CP, and helicases (S1H, S2H, and S3H for the helicases of superfamilies 1, 2, and 3, respectively) are shown systematically. The helicases appear to have been captured independently and in parallel in 3 branches of +RNA viruses, facilitating the evolution of larger, more complex genomes. Additional genes, namely, the Endo and Exo genes (for endonuclease and exonuclease, respectively) and the Hsp70h gene (heat shock protein 70 homolog), are shown selectively, to emphasize the increased genome complexity, respectively, in *Nidovirales* and *Closteroviridae*. The virion architecture is shown schematically for each included group of viruses. Icosahedral capsids composed of unrelated CPs are indicated by different colors (see the text for details). The question mark at the hypothetical ancestral eukaryotic RNA virus indicates the uncertainty with regard to the nature of the host (prokaryotic or eukaryotic) of this ancestral form. The block arrow at the bottom indicates the time flow and the complexification trend in RNA virus evolution.

origin of –RNA viruses is the next step during which the plus strand is discarded from the virions, perhaps simplifying the processes of transcription and replication. Conceivably, the evolution of dsRNA and –RNA viruses, in which transcription and replication of the viral genomes were confined to the interior of virions or nucleocapsid transcription/replication complexes and no dsRNA accumulated in the infected cells, was driven by the advantage of escaping some of the host defense mechanisms, in particular, RNA interference (RNAi) (109, 110). The membrane-associated replication complexes of +RNA viruses could represent an initial step in this direction (9).

Obviously, evolution of the RdRp does not equal evolution of viruses; other genes, in particular, those encoding capsid and other structural proteins, are crucial for virus reproduction, and these genes often have different histories. The reconstruction of gene gain and loss sheds some light on these aspects of RNA virus evolution. The ancestors of each of the major branches of RNA viruses (except for branch 1) appear to have been simply organized +RNA viruses resembling tombusviruses (Fig. 10). Thus, these types of viruses encoding RdRps and SJR-CPs might have been ancestral to the bulk of eukaryotic RNA viruses (apart from those in branch 1 that derive directly from prokaryotic leviviruses). The subsequent parallel capture of different helicases enabled evolution of increasingly complex genomes via accumulation of additional genes (Fig. 10). Notably, similar levels of complexity, with a complete coding capacity of 20 to 40 kb, were reached independently in 4 branches of the RdRp tree, namely, branch 2 (coronaviruses), branch 3 (closteroviruses and flaviviruses), branch 4 (reoviruses), and branch 5 (filoviruses and paramyxoviruses) (Fig. 3 and 6).

Gene exchange and shuffling of gene modules are important factors of RNA virus evolution. For example, it appears that all dsRNA viruses in the two major clades within

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branches 2 and 4 share homologous structural modules that combine with distinct RdRps. At least in the case of partiti-picobirnaviruses in branch 2, the dsRNA virus (“toti-like virus”) CP apparently displaced the ancestral SJR-CP. However, this particular protein structure does not seem to be essential to encapsidate a dsRNA genome; birnaviruses, whose provenance is uncertain due to the permutation in their RdRps, retain SJR-CP, which is most closely related to SJR-CP of nodaviruses and tetraviruses (19). An even more striking example of module shuffling is presented by amalgaviruses, dsRNA viruses that group with partitiviruses in the RdRp tree but encode a distant homolog of the nucleocapsid protein of $-$ RNA bunyaviruses (111–114). More generally, structural and replication modules have been repeatedly shuffled during the evolution of $+$ RNA viruses. Examples include displacement of the ancestral SJR-CP by a filamentous CP in potyviruses and by a helical nucleocapsid protein in nidoviruses and multiple cases of displacement with rod-shaped-like CP and unique nucleocapsid proteins in branch 3. Thus, exchange of genes and gene modules among RNA viruses is pervasive and can cross the boundaries of Baltimore classes.

Another recurring trend in RNA virus evolution is the loss of the structural module, resulting in the emergence of naked RNA replicons such as “narnaviruses” and “mitoviruses” in branch 1, hypoviruses in branch 2, and endornaviruses, umbraviruses, and deltaflexiviruses in branch 3 (18). On some occasions, broad horizontal spread of a gene led to a major shift in the lifestyle of viruses, such as adaptation of viruses to new types of hosts. The primary cases in point are the movement proteins (MPs) of plant viruses that are represented in all 5 branches of the RdRp tree and, outside the RNA part of the virosphere, in plant caulimoviruses and badnaviruses and in ssDNA viruses (115).

The prevalence of HVT and the overall course of RNA virus evolution. Arguably, the most striking realization brought about by metaviromics analyses concerns the diverse host range of numerous groups of viruses, even tight ones that occupy positions near the tips of the RdRp tree. Extending early observations on the highly unexpected similarities between viruses of animals and plants, recent metaviromic analyses revealed numerous clusters of indisputably related viruses infecting animals and plants or plants and fungi or, in some cases, animals or plants and protists. The results of the analyses of evolutionary relationships between viruses with distinct host ranges are supported not only by the phylogeny of the RdRp but also by the fact that these viruses share additional conserved domains, such as, for example, SJR1, S3H, and 3C-Pro in the case of *Picornavirales* members infecting protists, plants, fungi, invertebrates, and vertebrates (Fig. 3).

Invertebrates are particularly promiscuous hosts for viruses, often sharing the same virus group with distantly related organisms. Certainly, much caution is due in the interpretation of host range assignments from metaviromics, especially those resulting from holobiont studies. Viruses identified in holobiont samples of (for example) invertebrates could actually infect protists associated with these animals (50) or could represent contamination from fungal or plant or even prokaryotic sources. These uncertainties notwithstanding, the extensive diversity of hosts even within small branches of the RdRp tree is undeniable. A coevolution scenario in which the ancestors of all these viruses originated from the common ancestor of the respective groups of eukaryotes and coevolved with the hosts implies an enormous diversity of RNA viruses in early eukaryotes. This scenario appears to be highly unlikely given the apparent paucity of RNA viruses in the extant protists (although new metaviromic studies might substantially expand the range of protist viruses). The pervasive HVT alternative seems much more plausible, especially given that arthropods, nematodes, and other invertebrates are well known as virus vectors and thus fit the role of RNA virus reservoirs (11).

In addition to invertebrates, which appear to be the dominant HVT agents, fungi could also play an important role in HVT within the global RNA virome. Indeed, fungi that are tightly associated with plants and insects often share closely related viruses with these organisms (116–118). Furthermore, an indisputable case of cross-kingdom

transfer of an insect iflavivirus to an entomopathogenic fungus has been recently described (119).

These findings appear to be most extensively compatible with a grand evolutionary scenario (Fig. 10) in which the ancestor of the eukaryotic RNA virome was a levi/narnavirus-like naked RNA replicon that originally reproduced in mitochondria and then combined with a host carbohydrate-binding SJR protein (19) or a preexisting SJR-CP from a DNA virus to form a simple ancestral virus. Given that viruses of branches 2, 3, and 4 are present in modern protists, it appears likely that these branches emerged in early eukaryotes. However, because of the apparent dominance of the viruses of RdRp branch 2 (members of the “picornavirus supergroup” in general and of the “aquatic picorna-like viruses” clade in particular) in protists, this branch likely diversified first, whereas the diversification of branches 3 and 4 occurred later, after ancestral protist viruses were transferred to marine invertebrates during the Cambrian explosion. Results of the recent analysis of the viromes of ctenophores, sponges, and cnidarians suggest that substantial diversification of RNA viruses had already occurred in these deeply branching metazoa (120). Invertebrates brought their already highly diverse RNA virome to land at the time of terrestrialization and subsequently inoculated land plants. In land plants, RNA viruses, particularly those of branch 3, dramatically expanded, perhaps in part because of the exclusion of competing large DNA viruses. Finally, it seems plausible that, given the high prevalence of –RNA viruses in metazoa and their virtual absence in protists (with the exception of the recently discovered “leishbuviruses” that likely invaded their parasitic protist hosts via HVT from an animal host [49, 50]), the viruses that comprise RdRp branch 5 evolved in animals via mixing and matching genes from reovirus-like and flavivirus-like ancestors.

The impending overhaul of RNA virus taxonomy. The expansion of the global RNA virome thanks to the advances of metaviromics, combined with the phylogenomics results, seems to call for an overhaul of the current virus taxonomy on multiple levels. Most importantly, creation of a coherent, hierarchical system with multiple taxonomic ranks seems to be imminent. This process has already started with the proposal of a phylum rank for –RNA viruses, for which monophyly is unequivocally supported by the present analysis (Fig. 1 and 6). This phylum could consist of two subphyla with multiple classes and orders. At least 4 additional phyla of RNA viruses can be confidently predicted to emerge, including the dsRNA viruses of branch 4 and the +RNA viruses of branches 1, 2, and 3. Each of these phyla will undoubtedly have a rich internal structure. In addition, some of the current families do not seem to be compatible with the expansive RdRp trees presented here and in a previous analysis (14). For instance, the families *Coronaviridae*, *Togaviridae*, and *Rhabdoviridae* are likely to be split into two families each.

While the present study was in preparation, a major attempt at a comprehensive virus classification was published (121). This work analyzed a dendrogram that was produced from matrices of distances between viruses derived from sequence similarity scores combined with measures of gene composition similarity. Unlike our present analysis, this approach presupposes monophyly of each of the Baltimore classes. Furthermore, given that their analysis was based on measures of similarity rather than on phylogenetic analysis proper, this approach is best regarded as producing a phenetic classification of viruses rather than an evolutionary reconstruction as such. Some of the groups delineated by this method, particularly those among +RNA viruses, are closely similar to those reported here. Others, however, are extensively different; for instance, the order *Mononegavirales* is not classified as monophyletic in their dendrograms. We did not attempt a complete comparison; such an exercise could be useful in the future for better understanding the routes of RNA virus evolution.

Concluding remarks. Through metaviromics, many aspects of the global RNA virome evolution can be clarified. Certainly, reconstruction of the deepest events in this evolutionary history is bound to remain tentative, especially because the RdRp gene is the only universal gene of the RNA viruses and is hence the only one that can serve as

the template for evolutionary reconstructions. At the depth of divergence characteristic of RdRps, the relationship between the major branches in the tree cannot be established with confidence. Nevertheless, monophyly of several expansive groups, in particular, the 5 main branches in the RdRp tree, is strongly supported. Because of the stability of these branches, biologically plausible scenarios of evolution emerge under which dsRNA viruses evolved from different groups of +RNA viruses whereas –RNA viruses evolved from dsRNA viruses.

Evolutionary reconstructions suggest that the last common ancestors of each major lineage of eukaryotic +RNA viruses were simple viruses that encoded only the RdRp and a CP, most likely of the SJR fold. Subsequent evolution involved independent capture of distinct helicases which apparently facilitate replication of larger, more complex +RNA genomes. The helicase-assisted replication of +RNA genomes created the opportunities for parallel acquisition of additional genes encoding proteins involved in polyprotein processing and virus genome expression, such as proteases and capping enzymes, respectively, and proteins involved in virus-host interactions, such as MPs or RNAi suppressors of plant viruses. In addition to these processes of vertical evolution of RNA viruses, the results from phylogenomic analyses reveal multiple cases of gene module exchange among diverse viruses and pervasive HVT, often between distantly related hosts, such as animals and plants. Together, these processes have shaped a complex network of evolutionary relationships among RNA viruses.

The much-anticipated comprehensive exploration of the RNA viromes of prokaryotes and unicellular eukaryotes, such as free-living excavates, chromalveolates, rhizaria, amoebozoa, and choanoflagellates, as well as deeply rooted metazoa, will undoubtedly help in developing better-supported evolutionary scenarios for each of the 5 major branches of the RNA virus tree. Nevertheless, it is already clear that the current taxonomy of RNA viruses is due for a complete overhaul.

MATERIALS AND METHODS

Phylogeny of RNA-dependent RNA polymerases. Protein sequences belonging to RNA viruses (excluding retroviruses) and to unclassified viruses were downloaded from the NCBI GenBank database in April 2017 (122). Initial screening for RdRp domains was performed using PSI-BLAST (123) (E value of 0.01, effective database size of 2×10^8 sequences) with position-specific scoring matrices (PSSMs) produced from the available RdRp alignments. The sources included group-specific alignments for +RNA viruses and dsRNA viruses (12, 31) and the 4 PFAM alignments for the –RNA viruses (pfam00602, pfam00946, pfam04196, and pfam06317) from the NCBI conserved domain database (CDD) (124). Additionally, a set of RTs from group II introns and non-long-terminal-repeat (non-LTR) retrotransposons was extracted from GenBank as an outgroup.

Extracted RdRp footprints were filtered for the fraction of unresolved amino acids (at most 10%) and clustered using UCLUST (125) with a similarity threshold of 0.9. One representative from each cluster was selected for further analysis. The resulting set contained 4,640 virus RdRps. This set went through several rounds of semimanual curation whereby sequences were clustered using UCLUST, aligned using MUSCLE (126), and cross-searched against each other and their parent sequences (often representing complete viral polyproteins) using PSI-BLAST and HHSEARCH (127). Upon obtaining the results of these searches, the boundaries of the RdRp domain were expanded or trimmed to improve their compatibility with each other.

The RdRp and RT sequences were subjected to an iterative clustering and aligning procedure, organized as follows. Initially, sequences were clustered using UCLUST with a similarity threshold of 0.5. The clustered sequences were aligned using MUSCLE, and singletons were converted to pseudoalignments consisting of just one sequence. Sites containing more than 67% gaps were temporarily removed from alignments, and pairwise similarity scores were obtained for clusters by the use of HHSEARCH. Scores for a pair of clusters were converted to distances as follows: the $d_{A,B} = -\log[s_{A,B}/\min(s_{A,A}, s_{B,B})]$ formula, in which $d_{A,B}$ is the distance between cluster A and cluster B and $s_{A,B}$ is the HHSEARCH score for the comparison of these clusters, was used to convert scores s to distances d . The matrix of pairwise distances was used to calculate the mean by the unweighted pair group method using average linkages (UPGMA) (128). Shallow tips of the tree were used as the guide tree for a progressive pairwise alignment of the clusters at the tree leaves using HHALIGN (127), resulting in larger clusters. This procedure was reiterative, ultimately resulting in the single alignment of the whole set of 4,640 virus RdRp sequences and 1,028 RT sequences.

During the clustering procedure, 50 virus RdRp clusters, consisting of 1 to 545 sequences, were defined. These clusters represented either well-established groups of related viruses (roughly comparable to the ICTV family rank) or, in case of poorly characterized and unclassified viruses, groups of well-aligned RdRps that were clearly distinct from others. In all cases, uncertainties were treated conservatively, i.e., favoring placing sequences with questionable relatedness into separate clusters.

Additionally, RT sequences were placed into two clusters consisting of group II intron RTs and non-LTR retrotransposon RTs.

For each cluster consisting of three or more sequences, an approximate maximum likelihood (ML) phylogenetic tree was constructed from the cluster-specific alignment using the FastTree program (129) (Whelan and Goldman [WAG] evolutionary model with gamma-distributed site rates) with sites that contained more than 50% gaps removed from the alignment. Trees were rooted using a variant of the midpoint rooting procedure such that the differences between the (weighted) average root-to-tip distances in the subtrees on the opposite sides of the root were minimized.

To resolve the structure of the global relationships, up to five representatives from each cluster were selected using the within-cluster trees to ensure the diversity of the selected sequences. This procedure resulted in a set of 228 virus RdRps and 10 RTs. The alignments of the selected sequences were extracted from the master alignment and filtered for sites containing more than 50% gaps. A ML phylogenetic tree was reconstructed for the resulting alignment using PhyML (48) (Le Gascuel [LG] evolutionary model with gamma-distributed site rates and empirical amino acid frequencies; support values were calculated using aBayes method implemented in PhyML). Another form of branch support, i.e., bootstrap support provided by the transfer (BOOSTER) phylogenetic bootstrap method (130), was also used to assess the reliability of the major tree divisions. Alternatively, the same RdRp alignment was used as the input for ML phylogenetic analysis using RAXML (LG evolutionary model with gamma-distributed site rates and empirical amino acid frequencies).

RdRps in the global tree were divided into 5 major branches (supergroups). Up to 15 representatives from each cluster were selected to form supergroup-level alignments. The respective trees were reconstructed from these alignments using the same procedure (PhyML tree with LG evolutionary model with gamma-distributed site rates and empirical amino acid frequencies).

The overall tree was assembled manually by first replacing the supergroup representatives in the global tree with the supergroup trees and then replacing the cluster representatives with the cluster trees. The lower-level trees were rooted according to the arrangement of the representatives in the upper-level tree.

Identification of protein domains. Protein domains were identified using a representative set of RNA virus genomes, including representative members of ICTV-approved virus families and unclassified virus groups. This set was annotated manually using sensitive profile-profile comparisons with the HHsuite package (127), and hmm profiles for annotated proteins or their domains were generated by running one iteration of HHblits against the latest (October 2017) unclust30 database (131). Each annotated profile was assigned to a functional category (e.g., "capsid protein_jelly-roll," "chymotrypsin-like protease"). These profiles were used to annotate the genomes of all of the viruses that were included in the RdRp phylogenetic analysis and for which complete (or near-complete) genome sequences were available. Profiles for the latter proteins were generated by running one iteration of Jackhmmer (132) against the UniRef50 database. Protein regions that did not have significant hits were extracted and clustered with cluster analysis of sequences (CLANS) (133), and groups containing at least three members were identified, annotated (if possible), and added to the manually annotated profile database. The last step of the domain identification procedure was then repeated using the updated RNA virus profile database. Highly similar viruses (with identical domain organizations and >94% identical RdRps) were removed from the data set. In addition, only one representative genome was retained for some members of overrepresented species (e.g., *Hepacivirus C*). Many genomes (e.g., alphaviruses, tombusviruses, and nidoviruses) encoded readthrough proteins, resulting in domains from the shorter protein contained within the longer readthrough protein. Such redundancies were removed by filtering out the shorter of the two proteins sharing >80% identity. The final set of annotated genomes included 2,839 viruses.

Reconstruction of the history of gene gain and loss. The protein domains identified in the 2,839 virus genomes were mapped onto the composite tree of virus RdRps. A set of 500 representatives was chosen among the (mostly complete) genomes. The domain complement data and the respective tree were analyzed using the GLOOME program (51). Domain gains and losses were inferred at each tree branch using the difference of posterior probabilities for the domain occurrence in the nodes at the proximal and the distal ends of the branch (differences with values above 0.5 imply gain; differences with values below -0.5 imply loss).

Clustering analysis. The SJR CP network was generated by performing all-against-all comparisons of the SJR CP profiles. To generate hmm profiles, SJR CP sequences were extracted from the total proteome of RNA viruses and two iterations of HHblits were performed against the uniprot20_2016_02 database. The resulting profiles were compared to each other with HHsearch (127). Their similarity *P* values were extracted from the result files and used as an input for CLANS program (133). Clusters were identified using a network-based algorithm implemented in CLANS. Resulting clusters were manually inspected and refined.

Analysis of bipartite gene-genome networks. A bipartite network was built to study the patterns of gene sharing among viral genomes (44, 106). After removal of genomes with fewer than 2 domains and of domains that appeared in less than 3 genomes, the network consisted of 2,829 nodes, of which 2,515 corresponded to genomes and 314 to domains. Genome and domain nodes were connected by links wherever a domain was present in a genome. A consensus community detection approach was used to identify the modules of the network (134, 135). First, we ran 500 replicas with Infomap (136) (bipartite setting, using domains as factors) and built a similarity matrix by assigning to each pair of nodes a similarity value equal to the fraction of replicas in which both nodes were placed in the same module. Then, hierarchical clustering was performed on the similarity matrix, setting the number of

clusters equal to the median number of modules obtained in the 500 replicas. Order statistics local optimization method (Oslo) software (137) was subsequently used to filter significant modules with a *P* value threshold equal to 0.05. To detect higher-order (super)modules, we first identified connector domains as those present in at least 2 modules with prevalence greater than 0.65. The second-order network, composed of 54 modules and 34 connector domains, was searched for second-order modules with Infomap (500 replicas, bipartite setting with connector domains as factors). Due to the small size of the second-order network, consensus community detection did not qualitatively improve the results of the search, and therefore we took the replica with the best Infomap score and skipped hierarchical clustering for this and subsequent steps of the higher-order module search. After assessing statistical significance with Oslo, 4 second-order modules were recovered, encompassing 12 of the original modules associated with closely related virus families. A third-order network was built by pooling the 4 second-order modules with the 46 modules that remained unmerged. The third-order network was analyzed in the same manner to obtain the 9 supermodules. None of these supermodules was assessed by Oslo as significant.

Quantification of virus host range diversity. Known hosts or, in case of viruses isolated from holobionts, virus sources were identified for 3,456 viruses. The host taxonomy (analyzed to the phylum level) was mapped to the leaves of the combined tree. The tree was ultrametrized, and the weights were computed for all leaves as described in reference 138. Each internal node of the tree therefore defines a set of leaves with assigned hosts; the distribution of (weighted) relative frequencies of hosts can be characterized by its Shannon entropy. Each leaf has one host defined such that the entropy of the host range distribution is 0 whereas the host diversity is maximal at the root. The entropy generally declines along the tree because viruses that belong to relatively shallow branches tend to share their host ranges. The node depth versus node host range entropy data were averaged for each major branch separately using a Gaussian kernel with a bandwidth equal to 10% of the total tree depth.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at <https://doi.org/10.1128/mBio.02329-18>.

FIG S1, PDF file, 0.05 MB.

FIG S2, PDF file, 0.1 MB.

FIG S3, PDF file, 0.4 MB.

DATASET S1, XLS file, 1.1 MB.

DATASET S2, TXT file, 0.3 MB.

DATASET S3, TXT file, 0.04 MB.

DATASET S4, TXT file, 0.04 MB.

DATASET S5, XLS file, 1.5 MB.

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