

Whole-proteome phylogeny of large dsDNA virus families by an alignment-free method

Guohong Albert Wu^{a,b}, Se-Ran Jun^a, Gregory E. Sims^{a,b}, and Sung-Hou Kim^{a,b,1}

^aDepartment of Chemistry, University of California, Berkeley, CA 94720; and ^bPhysical Biosciences Division, Lawrence Berkeley National Laboratory, 1 Cyclotron Road, Berkeley, CA 94720

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The vast sequence divergence among different virus groups has presented a great challenge to alignment-based sequence comparison among different virus families. Using an alignment-free comparison method, we construct the whole-proteome phylogeny for a population of viruses from 11 viral families comprising 142 large dsDNA eukaryote viruses. The method is based on the feature frequency profiles (FFP), where the length of the feature (*l*-mer) is selected to be optimal for phylogenomic inference. We observe that (i) the FFP phylogeny segregates the population into clades, the membership of each has remarkable agreement with current classification by the International Committee on the Taxonomy of Viruses, with one exception that the mimivirus joins the phycodnavirus family; (ii) the FFP tree detects potential evolutionary relationships among some viral families; (iii) the relative position of the 3 herpesvirus subfamilies in the FFP tree differs from gene alignment-based analysis; (iv) the FFP tree suggests the taxonomic positions of certain "unclassified" viruses; and (v) the FFP method identifies candidates for horizontal gene transfer between virus families.

alignment-free genome comparison | feature frequency profile | horizontal gene transfer | whole-genome phylogeny | virus phylogeny

Phylogenetic and taxonomic studies of viruses have become increasingly important as more and more whole viral genomes are sequenced (1–4). Knowledge of viral taxonomy and phylogeny is useful for understanding the diversity and evolution of viruses not only within a viral family, but also among different viral families that may have a common origin (5). They also provide useful information in drug design against virally induced diseases (6).

One of the unusual aspects of viral genomes is that they exhibit high sequence divergence due to high mutation rate, genetic recombination, reassortment, horizontal gene transfer (HGT), gene duplication, and gene gain/loss (7, 8). A direct consequence of the high sequence divergence and relatively small number of genes in viruses is that the number of highly conserved genes among different viral families is very small or, sometimes, undetectable. For example, the relationship among different families of eukaryote large DNA viruses (LDV) has often been studied based on multiple sequence alignment of a single gene, the DNA polymerase gene (9). Whether this single-gene based analysis can be used to properly infer viral species phylogeny is debatable.

Due to this and other limitations (10) of multiple sequence alignment comparison of 1 or a few selected viral genes, there has been a growing interest in alignment-free methods for whole-genome comparison and phylogenomic studies (11, 12). Alignment-free approaches have been used in the reconstruction of virus genome trees for individual virus families (13, 14) and across virus families. Examples of the latter include the composition vector method used to construct a genome tree for large dsDNA viruses (15), the average common substring approach used for phylogenomic analysis of the reverse-transcribing viruses and the negative-sense ssRNA viruses (16), and tetranucleotide usage patterns that have been found useful for inferring host-virus coevolution among bacteriophages and eukaryotic viruses (17). Besides genome trees,

self-organizing maps (18) have also been used to understand the grouping of viruses.

In the previous alignment-free phylogenomic studies using *l*-mer profiles, 3 important issues were not properly addressed: (i) the selection of the feature length, *l*, appears to be without logical basis; (ii) no statistical assessment of the tree branching support was provided; and (iii) the effect of HGT on phylogenomic relationship was not considered. HGT in LDVs has been documented by alignment-based methods (19–22), but these studies have mostly searched for HGT from host to a single family of viruses, and there has not been a study of interviral family HGT among LDVs.

To address these issues, we have developed an alignment-free method using feature frequency profiles (FFPs) (23). In this work, we use the FFP method, supplemented by an HGT detection technique, to study the taxonomic grouping and phylogenomic relationship among subfamilies within each family, and phylogenomic relationship among 11 LDV families and 4 dsDNA insect viruses that have not yet been assigned to any virus family by the International Committee on the Taxonomy of Viruses (ICTV). Altogether, we analyze 142 complete LDV proteomes from National Center for Biotechnology Information's non-redundant Ref-Seq database (24).

Results and Discussion

We first present results on the whole-proteome tree reconstruction, including the choice of optimal feature length, and the identification of interviral-family HGT genes. To increase the sensitivity of the FFP method, we have applied 2 filtering schemes: the filtering of HGT candidate genes and the filtering of low-complexity features. Next, we describe the overall features of the LDV proteome tree, possible evolutionary relationship among families, and the differences between the FFP phylogeny and existing alignment-based phylogenies of several individual viral families. Finally, we compare the FFP tree to a previously published alignment-free analysis.

Optimal Feature Length. When whole proteomes are compared using *l*-mer FFP, different feature (*l*-mer) lengths can lead to different tree topologies. Thus, determining the optimal feature length is critical for phylogeny inference. Based on both cumulative relative entropy (CRE) and relative sequence divergence (RSD) analyses, the optimal feature length for LDV proteomes is determined to be 8 aa (see *Materials and Methods*). This estimate depends on the range of proteome sizes and the sequence divergence properties of the viruses (Fig. 1).

Horizontal Gene Transfer Between Viral Families. We use the Jensen-Shannon divergence (JS) (25) of pairwise FFPs to estimate the dissimilarity of 2 proteomes. JS provides a summary statistic of

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¹To whom correspondence should be addressed. E-mail: shkim@cchem.berkeley.edu.

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Herpesvirales. Herpesviruses are morphologically distinct from other viruses and they divide into 3 families under the recently established order Herpesvirales (30, 31), namely Herpesviridae, Alloherpesviridae, and Malacoherpesviridae. In the FFP tree, each family forms a clade, but the 3 families do not cluster to form a monophyletic group, indicating a lack of interfamily phylogenetic relationship at the sequence level despite of morphological similarities. The Herpesviridae clade further divides into 3 monophyletic subgroups corresponding to the α , β , and γ subfamilies with high statistical support. Of the 3 subfamilies, the β subfamily branches off first. This branching order is at variance with alignment-based analysis (31). The 4-member clade of the Alloherpesviridae shows moderate statistical support as a result of its great sequence divergence among the 4 viral proteomes, of which all but ICHV1 are currently not assigned at the genus level.

At the genus level, all except the rhadinovirus genus of the γ subfamily (shown in blue in inner ring of Fig. 3) are monophyletic. Within the rhadinovirus genus, the murid herpesvirus 4 (MHV4) proteome shows great sequence divergence and is separated from other members of the genus. Sequence alignment-based analysis also found that MHV4 has a particularly high level of sequence divergence, causing difficulties in determining its phylogenetic position unambiguously (32). The unclassified Tupaiid herpesvirus 1 (TuHV1) clusters with the cytomegalovirus genus of the β subfamily (shown in light green in the inner ring) in the FFP tree, although it may or may not be assigned to the same genus.

Phycodnaviridae and Mimiviridae. There are 9 phycodnaviruses and 1 mimivirus with complete proteomes in our dataset. Each multimeric genus forms its own clade with high branch support. The recently sequenced marine green algae virus OtV5 (33) is not yet included in the ICTV 2008 Official Taxonomy, although sequence comparison of the DNA polymerase gene suggested that it belong to the genus prasinovirus (33). In the FFP tree, OtV5 is positioned next to the chlorovirus genus (shown in red in inner ring of Fig. 3), as is also the case with the DNA polymerase-based analysis.

The 9 phycodnaviruses do not form a monophyletic group in the FFP tree, because mimivirus (APMV) nests within them. However, all phycodnaviruses and the mimivirus together form a monophyletic group with moderate statistical support. Sequence alignment using the major capsid protein (34) or the DNA polymerase gene (35) found similar mixing between the mimivirus and phycodnaviruses. This is at variance with an earlier phylogenetic analysis suggesting that the mimivirus form a separate family (36). In the FFP tree, the mimivirus, OtV5, and the chlorovirus genus form a highly supported clade. Both the FFP tree and the recent sequence-alignment analyses show the high sequence divergences among the genera of Phycodnaviridae (37), suggesting a possible taxonomic revision of the Phycodnaviridae family (34, 38) and the mimivirus (35).

Poxviridae. The grouping of poxviruses in the proteome tree is consistent with the ICTV classification. The highly supported poxvirus clade falls into 2 monophyletic groups corresponding to the entomopoxvirinae and chordopoxvirinae subfamilies (middle ring, purple and green respectively), and the latter further divides into 3 monophyletic groups associated with reptilian, avian and mammalian hosts, respectively. Each genus forms a clade in the FFP tree. The branching order of different genera mostly agrees with an analysis based on alignment of a core set of 35 genes common to the chordopoxvirinae (39), although minor discrepancies also exist, for example, in the relative position of cervipoxvirus (DPV) and capripoxvirus (SHPV, GTPV, LSDV). In the FFP tree, the unclassified crocodile poxvirus (CRV) is the outgroup of the chordopoxvirinae clade and positioned next to the avipoxvirus genus (FWPV, CNPV). This suggests that CRV could be assigned to a new genus within the chordopoxvirinae subfamily.

Other viruses. There are 4 insect viruses that are not assigned to any viral family. Two (HzNV1 and GbNV) are nudiviruses, and they form a clade and cluster with the baculovirus family in the FFP tree,

consistent with an analysis based on alignment of the DNA polymerase gene (40). The other two insect viruses causing salivary gland hypertrophy (MdSGHV and GpSGHV) form a clade with strong support, corroborating a recent finding that the two are related and form a distinct clade based on analysis of gene trees (41). They cluster with WSSV. The FFP tree also suggests that the 2 nudiviruses and the 2 SGHVs be separately assigned to 2 new viral families.

Comparison with Another Alignment-Free Method. In a previous report on the reconstruction of the whole-proteome phylogeny of large dsDNA viruses (15), the authors used an *l*-mer-based composition vector (CV) method with subtracted background “noise” modeled by a Markov chain estimator. Notable differences between the FFP tree and the CV tree are (i) the CV tree was based on *l*-mers of length 5, but the optimal feature length for FFP tree is 8; (ii) the CV tree did not explicitly deal with HGT among LDV families; (iii) the authors did not provide statistical assessment of branch support in the CV tree; (iv) neither baculoviruses nor iridoviruses are monophyletic in the CV tree; (v) the phycodnaviruses do not form a monophyletic group, with or without the mimivirus in the CV tree; and (vi) ascoviruses were not included in the CV tree, which could further distort the CV tree topology due to the extensive HGT between ascovirus and baculovirus.

FFP Method vs. Multiple Sequence Alignment (MSA) Method. MSA method has to select a set of highly conserved genes for alignment, and assumes that phylogeny of those selected genes represents species phylogeny. Thus, MSA can be applied only within individual families or for closely related families, and cannot be used for comparing diverse multiple families of LDVs. For inferring phylogeny of diverse families, FFP method has at least 3 advantages: (i) the whole genome/proteome is used to represent each species, (ii) it does not require selection of highly conserved genes common to all families, and (iii) it is not very sensitive to large-scale genome rearrangement and other changes including gene gain and loss.

On a more technical note, the presence of a common 8-mer between two proteins does not in general imply that the two proteins are homologous, and vice versa. This is illustrated in Fig. 2D, which shows that random sequences can have common 8-mers, and in Fig. 2C, which shows that there may be no common 8-mer between many protein pairs of DNA polymerase from different viral families. To make the distinction between distant and closely related viral species, we use 50 type species representing all of the LDV genera and find that only 53% of the interfamily 8-mer-sharing protein pairs are homologous, after excluding HGT genes and low complexity features and using a blast E-value cutoff 0.01. In contrast, for intrafamily protein pairs, 8-mer conservation implies gene homology 95% of the time. However, even for the latter case, FFP and MSA, which use the whole proteome and a fraction of the proteome respectively, can give different phylogenies as exemplified by the branching order of the α , β , and γ subfamilies of the herpesviridae. These observations suggest that 8-mer conservation is not a useful measure for phylogenetic inference, but the profile of all 8-mers determines the FFP tree topology.

Conclusion

Using the alignment-free FFP method, we have studied the molecular phylogeny and horizontal gene transfer (HGT) between families for a broad population of large dsDNA eukaryote viruses consisting of 11 viral families. The unique aspects of this study include: (i) the selection of optimal feature length for phylogeny inference, (ii) a modified bootstrap support analysis of the branching orders in the FFP tree, and (iii) identification of interfamily HGT candidate genes and exclusion of the genes from the FFP tree reconstruction. The analysis of the FFP tree for the broad population of LDVs suggests that the method is suitable for grouping diverse families of viruses, subgrouping within individual families,

finding possible evolutionary relationship among the families, and assigning “unclassified” species, even when there are no or few common genes among the broad population.

Materials and Methods

Dataset. The viral sequences were downloaded from National Center for Biotechnology Information’s REFSEQ database (September 2008 release) (24). Protein sequences for large eukaryote dsDNA viruses are extracted from the .faa file. Polydnviruses are excluded from consideration because they are a distinct group and hardly share any common genes with other virus families. The final dataset of 142 LDVs consists of 11 viral families and 4 insect viruses unassigned to any family. The list of viruses is included in Table S2.

Feature Frequency Profile (FFP) and Distance Matrix. A general description of FFP method is published in ref. 23. The feature frequency profile of a given sequence is obtained by counting all overlapping features of length l by sliding a window of width l along the sequence, advancing 1 letter at a time. The FFP of a proteome is the total sum of the FFPs for each protein sequence contained therein. In this work, we use the normalized FFP, i.e., the probability of occurrence of each word in a proteome. The dissimilarity between 2 FFPs can be estimated from the Jensen–Shannon divergence (JS) (25). For 2 probability distributions $P = (p_1, p_2, \dots)$ and $Q = (q_1, q_2, \dots)$, JS is given by

$$JS(P, Q) = \frac{1}{2} KL\left(P, \frac{P+Q}{2}\right) + \frac{1}{2} KL\left(Q, \frac{P+Q}{2}\right), \quad [1]$$

where $KL(P, Q)$ is the Kullback–Leibler divergence (42) or relative entropy

$$KL(P, Q) = \sum_i p_i \log_2 \frac{p_i}{q_i}, \quad [2]$$

and the summation is over all features. Note that JS is bounded between 0 and 1. Strictly speaking, JS is not a distance metric, because it does not satisfy the triangle inequality. However, this violation happens only for short feature lengths and is of no concern to us. For a given feature length l , the distance matrix for a collection of proteomes is constructed from all pairwise JSs.

Relative Sequence Divergence (RSD), Cumulative Relative Entropy (CRE), and Optimal Feature Length. Two methods exist for estimating the optimal feature length for whole-genome phylogeny. The first is related to information theory and makes use of cumulative relative entropy (CRE) of individual proteomes. By contrast, the second method estimates the relative sequence divergence (RSD) of a proteome relative to a random sequence of the same size by comparing their relatedness (in terms of FFP) to a group of proteomes. Both methods give the same estimate for LDVs.

CRE. This method estimates the minimal feature length for which the information content of a proteome can be approximated by its FFP. This is done by requiring the CRE between the FFP of a proteome and that of a Markov chain estimator to be small. Under a Markov chain model of order $l-2$, the expected l -mer frequencies of a sequence or proteome is given by frequencies of features of lengths $l-1$ and $l-2$ as follows (43),

$$\tilde{f}_{a_1 \dots a_l} = \frac{f_{a_2 \dots a_l} * f_{a_1 \dots a_{l-1}}}{f_{a_2 \dots a_{l-1}}}, \quad [3]$$

where f denotes observed feature-frequencies of a proteome, a_i denotes amino acid type at position i of a feature. The difference between the estimated and observed l -mer frequencies can be measured by the relative entropy $KL(\tilde{P}_l, \hat{P}_l)$, where \tilde{P}_l and \hat{P}_l are estimated and observed probability vectors of l -mers respectively. This difference as a function of feature length exhibits a peak, whose position can be estimated using random sequences (zero-order Markov chains) and is well approximated by

$$l_{peak} \cong \log_{20} N + 1, \quad [4]$$

where the base 20 is the number of amino acid types and N is the proteome size.

A monotonically decreasing function can be constructed for the cumulative relative entropy (CRE),

$$CRE(l) = \sum_{i \geq l} KL(P_i, \tilde{P}_i). \quad [5]$$

The minimal feature length at which $CRE(l)$ approaches zero can be used iteratively to infer approximate frequencies of increasingly longer features, and is defined as the optimal feature length for phylogeny inference. For a group of divergent sequences like LDVs, this is approximately given by

$$l_{CRE} \sim 2 \log_{20} N, \quad [6]$$

where N denotes the largest proteome size. For LDVs, the largest proteomes (i.e., mimivirus and phycodnaviruses) give $l_{CRE} \approx 8$. This estimate is confirmed in Fig. 1A, where CRE values from Eq. 5 are plotted for all LDVs against feature length, and they all approach zero at feature length 8, with the largest proteome of the mimivirus (APMV) as the main determining factor.

RSD. This method requires that, on average, a biological sequence shares more features than a random sequence of the same length with a group of bio-sequences. For a group of n related biological sequences, the relative sequence divergence (RSD) for a biological sequence s_i at feature length l with $i = 1..n$ can be defined as

$$RSD(s_i, l) = \frac{\sum_{j \neq i} c(r_i, s_j, l)}{\sum_{j \neq i} c(s_i, s_j, l)}, \quad [7]$$

where $c(s_i, s_j, l)$ denotes the number of common feature of length l between sequences s_i and s_j . r_i denotes a random sequence of zero-order Markov chain with the same length as s_i . For short feature lengths ($l < l_{peak}$), nearly all possible features are used by both the random sequence and viral proteomes, and the RSD is approximately 1. For longer feature lengths ($l > l_{peak}$), the feature space is sparsely sampled, with all of the viral proteomes sampling one region and the random sequence a different region. As feature length increases, the overlap in feature space between the viral proteomes and random sequence becomes smaller and the RSD decreases to zero. Optimal feature length for phylogeny inference is obtained when RSD becomes much smaller than 1.

In Fig. 1B, the RSD’s are plotted for 4 representative LDV proteomes including the smallest (NeleNPV), the largest (APMV), and intermediate (SHFV and CNPV), and they all fall < 0.05 at feature length 8 and longer. Thus, both RSD and CRE analyses give $l = 8$ as the optimal feature length of the LDV proteomes. With longer feature lengths, RSD and CRE become even smaller, but the average number of shared features between viral proteomes (especially distantly related ones) becomes fewer and the resulting tree topology is less robust.

Interfamily HGT Candidates. HGT between viral families can cause some distortion of the tree topology, because JS can be biased by the few highly similar genes shared between 2 viruses as measured by the number of common 8-mers. For LDV proteomes at the optimal feature length $l = 8$, the distribution of common 8-mers in a protein pair is illustrated in Fig. 2. In particular, Fig. 2B shows the results from pairwise comparison of all proteins from different viral families, and Fig. 2D shows the same comparison after the amino acids in each protein sequence are randomly permuted. From Fig. 2D we infer that a protein-pair from our dataset can share up to 4 different 8-mers by chance. Fig. 2C plots the number of common 8-mers from DNA polymerase pairs between viral families, and the maximum number of shared 8-mers is 8. Thus, a protein pair from different viral families that share unusually high number of 8-mers relative to the DNA polymerase protein, which is common to all members, are candidates for HGT. For example, as shown in Fig. 2A, the unusually large number of common 8-mers present in protein pairs from the ascovirus HVAV3e and the baculovirus HZSNPV suggests direct or indirect HGT events between the 2 viruses.

To see the effect of using different HGT cutoffs (i.e., number of shared 8-mers) on LDV phylogeny, we compare tree topologies with cutoffs ranging from 6 to 40. We observe that the tree topology remains stable for a HGT cutoff in the range 13–31 (Fig. S1). For this work, we use a conservative HGT cutoff of 20, and identified 164 HGT instances consisting of 8 genes (Table S1).

Filtering Out Low Complexity Features. Features with low complexity generally bear no or little phylogenetic signal and could distort the tree topology if enough of them are present in the viral proteomes. One measure of feature complexity is the Shannon entropy

$$K_2 = - \sum_i \frac{n_i}{l} \log_2 \frac{n_i}{l}, \quad [8]$$

where i runs over the 20 aa types, n_i is the occurrence frequency of amino acid type i in a given feature, and l is the feature length. This and another closely related

complexity measure K_1 were used to detect and exclude regions of low complexity in amino acid sequences (44) during sequence alignment. For 8-mers, K_2 takes on values between 0 and 3, corresponding to using 1 and 8 aa types respectively.

The effect of using different low complexity cutoffs on phylogenetic tree reconstruction is illustrated in Fig. S2. Note that even excluding only the least complex features (i.e., homo 8-mers) causes appreciable change in the tree topology. For K_2 between 0 and 1.5, we observe that the tree topology is most stable for cutoffs 0.9, 1.1, and 1.3. Based on this analysis, we filter out 8-mer features with $K_2 < 1.1$ for this study. These features account for 0.3% of the viral proteomes on average, and up to a maximum of 2% for the EhV86 proteome. By way of comparison, for random sequences with equal usage of different amino acid types, the fraction of 8-mers with $K_2 < 1.1$ is $< 10^{-5}$. The compositional types of these low complexity features include A_8 , $A_x B_{8-x}$ ($x = 1-4$), and $A_6 B_1 C_1$, where A, B, and C denote different amino acid types.

Phylogenetic Tree Reconstruction and Robustness Test. Phylogenetic trees are constructed from distance matrices using BIONJ (45). Robustness of the tree topology is estimated using a modified version of the bootstrap method (46),

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which works as follows. A table is first constructed with each row representing 1 viral proteome and each column representing 1 feature present in a viral proteome. Each table element indicates the feature frequency in a proteome (zero if absent). The bootstrap is applied to the columns of the table except that columns that are redrawn are treated as drawn only once (i.e., each column is either present or absent in the bootstrapped table). Thus, the resampled table has fewer columns but each feature maintains the same frequency as in the original table. This procedure is equivalent to a jackknife test deleting $1/e$ (i.e., 37%) of the features. A new distance matrix is then calculated for the resampled table. We use 200 replicates to estimate the branch support for the un-bootstrapped tree. For the LDV dataset, a significant proportion of the features are unique to only 1 proteome, thus the resampling is expected to underestimate the branch support. We have taken this and other factors (47) into consideration when making phylogenetic inferences.

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