Expansion of the Metazoan Virosphere: Progress, Pitfalls, and Prospects

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Abstract:
Metagenomic sequencing has led to a recent and rapid expansion of the animal virome. It has uncovered a multitude of new virus lineages from under-sampled host groups, including many that break up long branches in the virus tree, and many that display unexpected genome sizes and structures. Although there are challenges to inferring the existence of a virus from a ‘virus-like sequence’, in the absence of an isolate the analysis of nucleic acid (including small RNAs) and sequence data can provide considerable confidence. As a consequence, this period of molecular natural history is helping to re-shape our views of deep virus evolution.

Highlights:
- Metagenomic discovery now contributes substantially to our view of virus evolution
- Serendipitous virus sequences from transcriptomes and genomes are under-utilised
- Small-RNA and strand-specific sequencing aid interpretation of viral metagenomes
- Barcode-switching and cryptic/unintended host material hamper robust host assignment
- Systematic host sampling is required to estimate the total number of animal viruses
Explosive metagenomic growth

It is 120 years since the word ‘virus’ was first applied specifically to a viral pathogen [1], but the number of known viruses is growing faster than ever (figure 1A; [2]). Much of this growth is through metagenomic discovery: the undirected large-scale sequencing of nucleic acids sampled from potential hosts or their environment [2-4]. Pioneered by studies of bacteriophage in the marine environment [5], recent years have witnessed an explosion in metagenomic sampling of the metazoan virosphere. This boom has focussed first on viruses likely to infect us and our livestock, particularly the virome of mammalian faeces [e.g. 6], on putative disease reservoirs such as bats [e.g. 7,8], and on arbovirus vectors [e.g. 9]. Subsequently, the focus has expanded to include neglected animal lineages, identifying hundreds of new RNA viruses in arthropods and other invertebrates [10-13], and recently in divergent and under-sampled chordates [14,15].

Compared to the isolation of new virus cultures, metagenomic discovery seems (relatively) cheap, easy, and (virtually) guaranteed—sequences often appear ‘for free’ when sequencing genomes and transcriptomes (Figure 1B-E) [10,16-18]. Nevertheless, there are clearly limitations to metagenomic discovery—especially for important applied questions such as “Where is the pandemic coming from?” [2]. With an isolate in hand we would have more than just a ‘virus-like sequence’: we could unambiguously confirm the host, be confident we hadn’t been misled by a computational artefact, and study viral replication, host range and immunity [19-21]. However, our catalogue of the virosphere is in its infancy, and there are still great gains to be made from simple ‘molecular natural history’. Fewer than 5 thousand viruses have received formal taxonomic recognition [22] and only around 15 thousand have even been named informally (Figure 1A). This is less comprehensive than the 17th century view of plant diversity, even in absolute terms [ca. 18 thousand species, 23], but few biologists today would claim the naturalists of subsequent centuries wasted their effort when making herbarium collections. And a modern evolutionary virologist can probably learn more from a virus genome than a 17th century botanist could from a dried specimen.

Metagenomic discovery has already had a huge impact on our knowledge of virus diversity. It has ‘filled in’ shallower parts of the tree, finding close relatives of iconic human pathogens, such as new influenzastrains in toads and eels [14]. It has also discovered new deep branches, such as clades of insect-infecting Partitiviruses [10,11] and Lu- teo/Sobemo-like viruses [10,24], and whole new families, such as the Chuviruses [25]. This in turn has led to renewed interest in inferring deep viral phylogenies [11,26], and has prompted proposals for large-scale updates of higher-level virus taxon- omy [27]. More importantly, metagenomics now contributes to our thinking on virus evolution. It has provided a better perspective on host-association and host-switching [14,28,29], found familiar virus lineages with unexpected genome sizes and structures [11,25,30], and uncovered an unexpectedly dynamic history of ‘modular’ protein swapping [11,26]. Finally, merely having a PCR product from a metagenomic sample can provide an experimental route to the functional biology of an uncultured virus [31].

Potential pitfalls

The recent viral bonanza partly reflects advances in nucleic acid sequencing, a technology that has left Moore’s Law—that computational power doubles every 2 years—far behind [32]. But sequencing is just one of the challenges to exploring the virosphere. The lack of a viable meta-bar-coding sequence means that virus discovery often takes a full metagenomic approach, sequencing total (or virus-enriched) nucleic acid, and subse- quently assigning sequences through inferred ho- mology [e.g. 3,33,34]. This is challenging because high divergence means that only the most conserved sequences are recognisable (e.g. RNA virus polyme-rases), and even then, only at the protein level. Sensitive surveys therefore benefit from assembled contigs rather than raw reads (so that divergent genes are linked to recognisable ones) and protein rather than nucleic-acid similarity searches (because divergence is high). This can be done using off-the-shelf assemblers and search al gorithms such as SPAdes [35] or Trinity [36], and Diamond [37], but there is also a growing ecosys- tem of virus-specific metagenomic packages and pipelines available [34].
As with any field in rapid development, best practice is uncertain and fluid, and there are pitfalls for the unwary [3]. For example, although the assembly of virus (especially RNA virus) genomes is facilitated by their small size and largely unrepeatitive nature, the high complexity of metagenomic pools tends to promote artefactual and chimeric contigs [4,38]. These can unite viral sequences with non-viral ones, especially high-copy-number host sequences such as those from mitochondria and ribosomes. Such ‘wide’ chimeras are partly mitigated by the use of paired-end and strand-specific reads, ensuring effective adaptor removal, and (when possible) removing host reads before assembly (although this can introduce problems if virus reads can cross-map to the host). Chimeric mis-assemblies among divergent viruses or viral segments are also possible, especially when they share identical stretches of sequence, such as structural RNA motifs or terminal repeats. These are harder to diagnose, and may ultimately require PCR verification, but can often be flagged by comparison with close relatives (if available), unexpected local variation in read-depth, and comparison across metagenomic samples.

These challenges aside, discovering a ‘virus-like sequence’ remains easier than confirming its status as an infectious agent of the targeted host. First, even if a sequence is ultimately virus-derived in an evolutionary sense, its immediate origin may have been an Endogenous Viral Element (EVE) [39]. If expressed and/or ‘domesticated’ by the recipient genome, EVEs may be represented at high levels and retain open reading frames [39,40]. Equivalently, host sequences—especially transposable elements (TEs)—are often incorporated into large DNA viruses and can move freely between hosts and viruses [41], allowing these host sequences to be misclassified as viral in origin. Second, the host can be misassigned if samples contain multiple hosts, either naturally or through contamination. Although nucleic acid contamination is minimised by good laboratory practice, externally contaminated reagents [e.g. 42], nucleic acids involved in reagent production (e.g. reads from Murine leukaemia virus [43]) and library miss-assignment at the point of sequencing, can all be harder to identify and to exclude. In particular, ‘barcode switching’ (or ‘hopping’) in some Illumina platforms can misattribute reads among libraries at rates of up to 1% [44], and while this is reduced by incorporating barcodes in both adaptors (‘dual indexing’), it is not always completely mitigated. Multi-host samples are often explicitly recognised as such, for example those from ‘holobionts’ such as anemones [45]. However, the multi-host nature of other samples is sometimes downplayed. For example, faecal samples are often dominated by viruses infecting the gut microbiota and/or organisms in the host’s diet [46,47], but virus-like sequences are sometimes reported (at least in the headline) as if they were viruses of the faecal donor itself. And, if nucleic acids or viruses are prepared from whole host individuals, viruses in faecal matter and viral infections of parasites (notably nematodes, platyhelminthes, and microscopic arthropods) and pathogens (fungi, trypanosomatids, apicomplexans, amoebae, and many others) will also be represented among the sequences. Pre-screening of samples for specific parasites by PCR [such as nematodes, e.g. 13]—can mitigate against this, as can tissue dissection [14] (although at the potential risk of biasing discovery toward viruses with a strong tissue tropisms). However, the potential for viral infections of eukaryotic parasites within the metazoan host means that even dissected tissue may be cryptically multi-host. For example, the only dimarhabdovirus recorded from a plant sample derives from RNA contaminated with thrips [16].

**Going beyond ‘virus-like sequences’**

Such pitfalls make some authors (justifiably) hesitant to proclaim a new virus from metagenomic sequencing alone, and many instead choose to report ‘virus-like sequences’—providing an implicit *caveat emptor*. But even in the absence of an isolate, sequence data and nucleic acid analysis can be used to support the existence of a free/replicating virus. First, the nature and quantity of the nucleic acid provides useful clues. Endogenous DNA copies can be identified by a comparison of PCR and RT-PCR (or direct DNA and RNA sequencing) [10,11,13,48]. For example, functional DNA viruses must express their proteins, so that the absence of viral mRNAs argues against active replication. Active replication also affects strand-bias in RNA viruses, so that strand-specific PCR [49] or RNA sequencing can identify the negative-sense replication intermediates of positive-
sense single-stranded (+ss) RNA viruses, and quantitative analyses can detect the presence of coding products from +ssRNA and dsRNA viruses [48]. And, for both DNA and RNA viruses, contaminating sequences are likely to be at relatively low titre, whereas the copy-number of inherited EVEs will match the host genome. This means that high copy-number itself provides an argument in favour of viral status [11], especially when viruses can contribute more than 10%, and sometimes in excess of 50%, of total (non-ribosomal) RNA in some species [50].

Second, contigs that encode complete viral genomes with intact open reading frames are more consistent with functional viruses than with EVEs. And, although whole viruses can be (retro)-copied into a host’s genome, there is rarely selective pressure to maintain the virus genome intact—resulting in segregating frameshift and nonsense mutations. Even expressed and functional (i.e. ‘domesticated’) EVEs generally only provide the host with one or two beneficial sequences [39,40]. Complete or near-complete virus genomes can also rule out the misattribution of host TEs as viral sequences, as the larger DNA viruses that carry TEs are unlikely to be represented by their TE sequences alone. Third, the distribution of virus-like sequences across metagenomic pools and host individuals (e.g. surveyed by PCR) can help to confirm a genuine viral origin [10,13,50]. Presence/absence patterns can help to weed out EVEs, as—unless it is very recent in origin—an EVE insertion is likely to be present in all host genomes, whereas virus prevalence is likely below 100% and variable among populations and over time [10,13]. The co-occurrence of virus-like and other sequences across host individuals can be used to correctly infer hosts, as viruses that infect a contaminating microparasite will co-occur with it. Similarly, patterns of co-occurrence can also help to identify missing parts of the viral genome, such as fragments of incompletely assembled genomes and components of segmented viruses that are not recognisable using sequence similarity [10,50].

Finally, perhaps the ultimate evidence of infection is recognition by the host antiviral immune system [10,51,52]. In vertebrates, the presence antibodies can be used to corroborate infection [51]. In nematodes and arthropods, the distinctive small RNAs (viRNAs) generated from viral genomes by antiviral RNA-interference (RNAi) [53,54] can be used in a similar way. Because Dicer-mediated viRNA biogenesis targets dsRNA such as replication intermediates, viRNAs can demonstrate both an antiviral response and viral replication. Importantly, viRNAs usually have a tight and characteristic length distribution (e.g. 20nt in Lepidoptera, 21nt in Drosophila, 22nt in C. elegans) [53,54] and a 3’ 2-O-methyl group, making them distinguishable from degradation products. Their size distribution and base composition also distinguish them from TE- and EVE-derived piwi-associated RNAs [13,40,54]. Notably, and unlike antibodies, viRNAs can identified directly from the metagenomic discovery RNA pool, allowing confirmation concomitantly with metagenomic discovery [10,52].

How many animal viruses are there, and what are they doing?

Our expanded view of the animal virosphere has already started to answer old questions and provoke new ones, but these two stand out. What prospect is there of answering them? Given any definition of ‘different virus’ [4,21], whether based on an operational taxonomic unit or a functional biological definition, virus lineages are countable. Sampling of nine virus families to near-saturation from one bat species in Bangladesh identified 55 different viruses and implied an estimate of 320 thousand viruses infecting mammals [55]. However, if a substantial proportion of these viruses were either multi-host or represented recent spillover from other hosts (i.e. without onward transmission) the estimate would be very different, and the estimate might also be biased by the particular choice of virus families and geographic region. A more confident estimate could be made from unbiased metagenomic samples of the joint distribution of prevalence across host and virus lineages, sampled across their geographic range. For example, near-saturation sampling of multiple taxonomic groups within a single ecosystem-type across a geographic region would not only allow the virus diversity within host lineages to be assessed, but would also allow an assessment of host range and—from sequence analysis—the timescale of host switching.

Such metagenomic surveys may soon be possible for a few carefully-considered host groups,
but they would still miss those virus sequences that we cannot see because they lack detectable homology with known viruses, the so-called viral ‘dark matter’ [56,57]. Many of these ‘dark matter’ sequences, perhaps the majority, are likely to represent the poorly-conserved regions of otherwise recognisable viruses [57,58]. However, some completely new and/or highly divergent virus lineages, which cannot be detected using the de facto default choice of search tools and parameters, probably remain to be discovered. Indeed, several recently-identified viruses were initially detected from contig size and nucleic acid abundance, and only subsequently attributed as viral using higher-sensitivity similarity searches [11]. In the future, more powerful search tools, such as those based on protein profile-profile comparisons [57,59], and a search for deeper homologies, such as those provided by protein structure [60], may prove useful. Where no remaining protein similarity exists, a complementary approach to is to consider sequences that are flagged as potentially viral in origin by the antiviral RNAi immune response of plants, insects, and fungi. Webster et al [10] proposed around 60 such viRNA-based ‘candidate virus’ contigs based on metagenomic sequencing from Drosophila, and approximately half of these have since been identified as fragments from known virus lineages by the subsequent discovery of related viruses, or by an analysis of co-occurrence across samples [e.g. 50]. This leaves open the possibility that some do represent genuinely new viruses (e.g. Figure 1E), but the ultimate confirmation of genuinely novel virus lineages probably represents a case in which viral isolates are unequivocally necessary.

What are these viruses doing to their hosts? It is almost axiomatic that viruses are parasites, but micro-organisms are often mutualist or commensal, and although viruses necessarily use host resources, their impact on host fitness may be negligible and/or outweighed by provision of some unknown benefit [61]. It might initially seem that elucidating the fitness consequences of infection must also require isolates for experimentation. However, experimental studies are rarely useful for inferring real-world fitness. First, most studies measure traits such as survival or reproduction in place of fitness. This can misinterpret life-history tradeoffs, such as mistaken a host response to mitigate cost (e.g. terminal investment) for a virus-derived benefit (increased early-life reproduction). Second, such studies tend to be under-powered: an absence of detectable harm does not imply costs are absent, only that they are small. But at what point is a cost so small that the virus is effectively commensal? The ultimate arbiter of costliness must be natural selection: if the presence of the virus selects for host resistance, then the virus imposes a net fitness cost, by definition. A resistance mutation is expected to spread if its fitness benefit substantially exceeds the impact of genetic drift (i.e. \( N_e s >> 1 \) where \( N_e \) is effective population size and \( s \) is the selective benefit). Very conservatively, an infection cost of 0.1% in Drosophila (or many other small invertebrates with large effective population size) would select strongly for host resistance. However, this cost is probably an order of magnitude too small to measure experimentally in a multicellular organism [62], meaning that it is effectively impossible to experimentally distinguish between a low-cost virus and a commensal one. Far from requiring more isolates, the best solution to understanding fitness consequences of infection could also be a metagenomic one, by adding metagenomic screens to fitness studies of animals in the wild [e.g. 63].

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**Figure 1**

**Panel A:** The number of distinct names for viruses (excluding phage) in the GenBank nucleotide database, by year (colours provide a scale for Panels B-D). Counts were obtained by finding the record creation date and GenBank ‘species’ (collapsing strain identifiers) for each of 2.6 million virus sequences. Exclusion of unrecognised species names and the merging of divergent strains are likely to make this an underestimate. **Panel B:** Midpoint-rooted maximum likelihood phylogeny of picorna-like viruses and caliciviruses, inferred from approximately 250 amino acids of the polymerase. Branches are coloured by the year in which the lineage was first recorded in GenBank (colours provided by panel A). Approximately 8000 picorna-like polymerase sequences from the NCBI non-redundant protein (nr) and transcriptome shotgun assembly (tsa_nt) databases were identified by blastp and tblastn. These were collapsed into 1140 clusters at a threshold of 96% identity, with one representative of each cluster used to infer the tree. Around 10% of the represented picorna-like lineages are known only as unannotated virus-like sequences from transcriptomes (pale yellow; viruses from transcriptome datasets are treated as unpublished and given a more recent date). Note that the short conserved-sequence length leads to poor resolution and fails to recover some named genera, and that similarity criteria for inclusion means that some picornavirus groups were excluded. **Panels C and D:** To illustrate with ease with which new virus-like sequences can be found in public datasets, I obtained the most recently deposited *Drosophila* RNAseq dataset (PRJNA414017 [64]), performed a *de novo* assembly using Trinity [36], and identified virus-like sequences using Diamond [37]. I found complete genomes for two picorna-like viruses (red labels; MH320557 and MH320558): a divergent sequence of Kilifi virus from *D. bipectinata* (previously known from *D. melanogaster*) and a Dicistro-like virus from *D. kikawai*, related to Hubei diptera virus 1 [11]. Maximum-likelihood phylogenies for these two sequences were inferred from around 700 amino acids of the polymerase, mid-point rooted, and coloured as in panel B. These trees illustrate the dominance of recent discoveries, including the many virus-like sequences in transcriptome assemblies (blue taxon labels). They also illustrate the potential confusion introduced by naming faecal-sample viruses after the faecal donor (all close relatives of Goose Dicistrovirus infect invertebrates). **Panel E:** Phylogeny of two putative ‘dark matter’ viruses from *Drosophila*, including related transcriptome sequences. These putative viruses each comprise four 1.5Kb segments encoding a single long open reading frame (the most conserved of which was used for phylogenetic inference), but they lack detectable homology with any known virus lineage and were inferred to be viral on the basis of viRNA profiles and co-occurrence across samples [10]. Data associated with this figure are available from via FigShare https://dx.doi.org/10.6084/m9.figshare.6272066.