Viral taxonomy needs a spring clean; its exploration era is over

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Abstract

The International Committee on Taxonomy of Viruses has recently changed its approved definition of a viral species, and also discontinued work on its database of virus descriptions. These events indicate that the exploration era of viral taxonomy has ended; over the past century the principles of viral taxonomy have been established, the tools for phylogenetic inference invented, and the ultimate discriminatory data required for taxonomy, namely gene sequences, are now readily available. Further changes would make viral taxonomy more informative. First, the status of a 'taxonomic species' with an italicized name should only be given to viruses that are specifically linked with a single 'type genomic sequence' like those in the NCBI Reference Sequence Database. Secondly all approved taxa should be predominately monophyletic, and uninformative higher taxa disendorsed. These are 'quality assurance' measures and would improve the value of viral nomenclature to its users. The ICTV should also promote the use of a public database, such as Wikipedia, to replace the ICTV database as a store of the primary metadata of individual viruses, and should publish abstracts of the ICTV Reports in that database, so that they are 'Open Access'.
past phylogenetic history, and so provide most of the information required for taxonomy.

**Viral names; names of what?**

The first descriptions of viruses used vernacular names for viruses that were based on the diseases they caused; tobacco mosaic virus, smallpox virus, etc. In the 1930-40s, F.O. Holmes and H.H. McKinney attempted to introduce Latinized binomials for viruses, *Marmor tabaci* for TMV, however they were not adopted because the nature of viruses was still largely unknown and no sensible classification had yet emerged. Consequently throughout the past century the different host-specific branches of virology have devised their own vernacular names; plant and animal viruses from disease, host or provenance names, bacteriophages from isolate codes, etc. It is interesting, therefore, that virologists studying the Pandoraviruses, one of the currently emerging areas of virus discovery, are using classical latinized binomials for them [8].

The ICTV regularized the naming of viruses in 1993 [9], and produced a list of approved taxonomic names for 3,600 of them. Confusingly most of these were already in use as vernacular names of viruses, but they were distinguished by writing the vernacular name of the virus in Roman script and the formal name of the species in italics [10]. Thus *Tobacco mosaic virus* became the name of the species that includes the subcellular organism named tobacco mosaic virus (TMV). This pedantic nomenclatural practice is frequently used incorrectly or ignored, and also conflicts with the frequent use of generic or specific names of plants as part of a viral name [11]; the long-established taxonomies of the larger cellular organisms take a more relaxed view of taxonomic and organisal names, and rely more on the good sense of the reader to infer the status of a name from the context in which it is used.

A very significant unresolved problem with the taxonomic names of viruses published by the ICTV is that there is no formal description of the virus to which each name is attached. By contrast, the formal taxonomic name for each plant, animal or bacterium is attached to one particular specimen, culture, description, or even illustration, kept in a secure collection, museum or herbarium (http://en.wikipedia.org/wiki/Type_(biology)). These are called ‘type’ specimens and each is permanently associated with a particular name, and thus provides total stability to that name. These preserved types can be compared with new specimens, and can be examined by techniques that have been invented since the type was collected, most notably gene sequencing. By contrast there are no comparable viral types; specimens or descriptions. In the early days of viral taxonomy most viruses could not be stored or maintained by serial passaging. Attempts were therefore made to collate viral descriptions based on the best data winnowed from the scientific literature. The earliest compilations were books, such as Smith’s “Textbook of Plant Virus Diseases” (1957) and Andrewes and Pereira’s “Viruses of Vertebrates” (1978). The first computer database of descriptions of viruses was started in the early 1980s as the VIDE (Virus Identification Data Exchange) project [12] using the pioneering DELTA system (http://delta-intkey.com/www/refseq.htm). It aimed to be comprehensive and progressive, and in 1991 became the first component of the ICTVdB [13], which, when work on it was concluded in 2008, contained data on 4949 ‘species’, 286 ‘genera’ or groups and 71 ‘families’, etc. of viruses. However the descriptions in the ICTVdB never became the basis of the formal names as none uniquely described individual viruses, and all were only ever partially complete as the exploratory phase of virology moved ever forward using ever more sophisticated techniques. Nowadays viruses are identified by gene sequence comparisons using the Genbank/EMBL/ DDBJ nucleotide databases, which currently contain over two million viral gene sequences, many of them complete. Therefore unique ‘type genomic sequences’ could and should now be formally linked with the approved names of individual virus species. This was done informally in the 9th ICTV Report [14] using the NCBI Reference Sequence Database [15], as this is a “comprehensive, integrated, non-redundant set of reference sequences”. Work on the ICTVdB is unlikely to be resumed as its proposed functions in viral taxonomy have been largely superseded by genomic sequences.

**Viral classification; polyphyletic or pseudo-monomophyletic?**

The recently approved concept of a viral species, like its predecessor, states that viral species are produced by divergent evolution; a species is now “a monophyletic group of viruses” and, earlier, “a replicating lineage”. However a feature of the ICTV’s Master Species List [16], that is cryptic and may confuse many, is that whereas the lineages that constitute Species and Genera are collections of viruses that share most of their genes, many viral Families and Orders share their genes with more than one Family or Order. Thus the Master Species List is arranged as a series of separate hierarchies, but whereas some are mostly ‘natural,’ others are ‘artificial’; polyphylly masquerading as monophyly.

If, for example, a virus is stated to be from the Order *Tymovirales*, that information only reveals which gene family supplied its replicate, although if that virus was stated to be from the *Tymoviridae* or the *Alphaflexiviridae*, namely Families in the Order, that would be much more informative as two or more genes in the viruses of each Family are related, and are expressed in a range of shared phenotypic characters. Thus there is no
informational gain from linking these two Families into an Order based on a single character. Similarly many species of the Caudovirales share no genes with other species of the Order, but have virions of similar shape, which presumably reflects their mode of infection, therefore it would be better that the formal name of this polyphyletic group be abandoned, and for the traditional vernacular name ‘tailed phages’ to be used when required!

Conclusions; carpe diem

Now is the time for the ICTV to review the results of its considerable efforts over the past half century, and to remove any unhelpful vestiges of that period.

Firstly its Master Species List should be ‘spring-cleaned’. The only viruses to be given approved (italicized) names should be those formally linked with a single properly documented ‘type genomic sequence’. Others that have partial sequences should have the status of ‘tentative species’. All others should be removed from the list. In the 9th ICTV Report, for example, the genus Potyvirus had 143 approved species, yet only 53 of these were listed with NCBI Reference Sequence Accession Codes, and 16 had no sequence data. This change would indicate to a knowledgeable user the informational status of a taxonomic name, and give some idea of how fully the virus has been characterized. These sequences would also provide a series of agreed ‘datum points’ from which sequence relationships of other isolates and viruses could be measured.

Second the ICTV should only recognise and list taxa that are predominantly monophyletic and, where possible, this status should be based on sequence and structural comparisons of viral molecules. The ICTV could produce listings and phylograms, not only of viral species and higher monophyletic groupings of those species, but also lists and phylograms of the genes shared by different polyphyletic viral taxa and those species, but also lists and phylograms of the viral species and higher monophyletic groupings of ports are a significant part of that record. They are an important resource for those identifying and describing novel and emerging viruses. This information has come, mostly pro bono, from a wide variety of sources. The primary metadata, together with abstracts of the information extracted from it by the ICTV and published in its Reports, must be archived in a way that ensures its long term survival, while ensuring that is available to all who wish to use it. This will only be achieved if it is stored in one of the large international ‘Open Access’ databases, as these are probably less susceptible to capricious changes in funding priorities than smaller specialized databases, and because they will promote the decentralized collation, revision and enrichment of the data.

Competing interests

The author declares that he has no competing interests.

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