

Classification, nomenclature, and database development for hepatitis C virus (HCV) and related viruses: proposals for standardization

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Summary. This paper presents a summary of the recommendations that were formulated for the purposes of unifying the nomenclature for hepatitis C virus (HCV), based upon guidelines of the International Committee on Virus Taxonomy (ICTV), and provides guidelines for the incorporation of sequence data into an HCV database that will be available to researchers through the internet. Based upon the available data, the genus Hepacivirus should be regarded as comprising a single species with HCV-1 as the prototype. All currently known isolates of HCV can be divided into six phylogenetically distinct groups, and we recommend that these groups are described as clades 1 to 6. Whether or not these should be regarded as different species within the Hepacivirus genus requires additional clinical, virological, and immunological information. Clades 1, 2, 4, and 5 would correspond to genotype 1, 2, 4, and 5 while clade 3 would comprise genotype 3 and genotype 10, and clade 6 comprise genotypes 6, 7, 8, 9, and 11. We propose that existing subtype designations are reassigned within these clades based upon publication priority, the existence of a complete genome sequence and prevalence. The assignment of isolates to new clades and subtypes should be confined to isolates characterized from epidemiologically unlinked individuals. Comparisons should be based on nucleotide sequences of at least two coding regions and preferably of complete genome sequences, and should be based on phylogenetic analysis rather than percent identity. A forum for discussion and contributions to these recommendations will be made available at the international HCV database at http://s2as02.genes.nig.ac.jp.

Background

The long-awaited discovery of hepatitis C virus (HCV), the virus responsible for over 90% of non-A, non-B hepatitis, by Chiron Corporation in 1989 [11] and the simultaneous development of the technology of polymerase chain reaction led to a surge of sequence information for this virus (reviewed in [5, 27, 29, 47]). The first indication of the genetic diversity of HCV came from investigators in Japan [12, 19] and subsequent studies have identified at least six major genetic groups [1-4, 7-10, 12, 20, 22, 36, 38-40, 42, 44, 45, 48, 53]. Due to the rapid growth of sequence information, nomenclature describing the genetic variability has been inconsistent, resulting in considerable confusion for researchers and students alike who enter the field. In 1993, two research groups defined the existence of 6 major genotypes of HCV [3, 44]. Using regions within the core, E1, and NS5B, the proportion of identical nucleotides was proposed to distinguish types, subtypes, and isolates [44, 53]. However, the wide range of HCV genetic variability that has been discovered upon sequencing of subsequently identified strains revealed the existence of isolates that did not fit within these defined ranges [18, 28, 55–57, 58]. In light of these problems, a small but internationally representative group of HCV researchers gathered in Santa Fe, New Mexico, USA, in late August 1997 to generate a consensus regarding the taxonomic position of HCV and the nomenclature of HCV isolates.

Framework for discussions

The initial discussion focused upon the guidelines for virus nomenclature and classification provided by the International Committee on Taxonomy of Viruses (ICTV). The ICTV categorizes viruses as orders (-virales), families (-ae), subfamilies (virinae), genera (-virus) and species. A species is recognized as a "polythetic class of viruses that constitutes a replicating lineage and occupies a particular ecological niche". As such, more than one of the following properties have to be present to differentiate individual species: 1) sequence differences; 2) host range; 3) cell and tissue tropism; 4) pathogenicity or cytopathology; 5) physical properties; or 6) antigenic properties. In the case of a virus that cannot be propagated in cell culture (such as HCV), the complete genome sequence must be known and the virus must have further distinguishing characteristics such as virus neutralization [34]. At present, the ICTV recognizes the *Hepacivirus* genus as containing HCV [59]; the question of whether or not the 6 to 11 genotypes that have been reported should be categorized as individual species remains to be determined.

Sequence approaches for classification

Genetic classification can be broadly approached from two perspectives generally identified as either distance-based or character-based [25]. A distance-based approach is based upon numerical values describing the similarity, irrespective of the evolutionary relatedness, with the values usually expressed as measures of distance. These numerical values can be graphically depicted as a frequency distribution or as a tree-like structure; the generation of a "tree" from these data does not necessarily imply an evolutionary correspondence, especially when relatively short sequences are under investigation. Unweighted pair group method analysis (UPGMA) and neighbor-joining (NJ) are commonly used distance-based

methods used to generate a tree; both methods generate a single tree, which can be a starting point for evolutionary analysis.

The other general approach is character-based and evaluates the phylogenetic relatedness of sequences based upon a subset of positions called "informative sites". Parsimony methods (weighted and unweighted) and maximum likelihood are two frequently used approaches. Typically multiple trees (cladograms) are generated and evaluated for accuracy, and so these methods are computationally more demanding.

Bootstrapping is a test that can be applied to trees generated by either character-based or distance-based methods in order to estimate the frequency with which a particular branch (node) occurs. Sequences are repeatedly reshuffled and a tree analysis under the appropriate model (N-J, parsimony, etc.) is performed. The percentage of bootstrap replicates in which a particular set of sequences cluster together gives an indication of the strength of the grouping. Values of greater than 70% out of 1000 replicates are considered as supporting the grouping [16].

Methods used for genetic analysis of HCV

Three coding regions, core, E1, and NS5B were identified initially as being useful for identifying genotypes by distance-based methods [3, 4, 36, 44, 45, 53]. Sequence distances (dissimilarities) have been calculated in several HCV laboratories using the PHYLIP program DNADIST [13, 14] with the Kimura 2-parameter metric for transitions and transversions. In other laboratories, 1-parameter (simple percent dissimilarities) and 6-parameter [15] measures have been employed. Plotting the values obtained from any of these approaches as a frequency distribution (the number of times each pair-wise value is observed in a large data set) usually results in three peaks, each with the characteristics of a random distribution. The troughs between these peaks have been used to demarcate types, subtypes, and isolates [30, 31, 36, 44, 53]. Although these criteria were satisfactory for distinguishing between isolates from Europe, Japan, the Middle East and North America, analysis of strains from Southeast Asia and Africa produced ambiguous results with many distances falling in the troughs. Whether a sequence was considered a type, subtype or isolate might depend upon which particular sequences it was compared with [28, 48]. In contrast, when distances between sequences are used to construct neighbor joining trees, six different clades are observed and are supported by bootstrap analysis [3, 4, 28, 31, 36, 48, 52, 53]. Similar conclusions arise from analysis of complete virus genomes by the character based methods of maximum likelihood [28, 52] or parsimony (Fig. 1). A virus previously described as genotype 10 clusters on the same branch as viruses within genotype 3, together forming clade 3, while a virus described as genotype 11 clusters with a genotype 6a virus, together forming clade 6. This concordance of distance-based and character-based methods is the basis of our proposal that HCV variants be grouped into six clades.

The GB viruses

The issues which relate to the taxonomic status of the GB and HGV viruses, GBV-A [32, 49, 50], GBV-B [32, 49, 50], and GBV-C/HGV [23, 26, 37, 49], are also whether or not these viruses are monophyletic or polyphyletic in relationship to HCV and also to each other. Figure 2 is a reconstruction of the relationships based upon inversely weighted

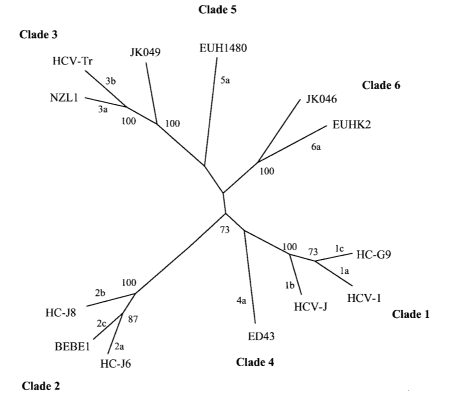


Fig. 1. Parsimony analysis of representative HCV sequences. Inversely weighted parsimony was performed using PAUP 3.1.1 as previously described [17, 21]. To minimize noise, only homologous second base positions from complete genomic sequences were studied: of the 3183 total sites that were aligned, 1216 varied sites were analyzed. A virtually identical topology was obtained using maximum likelihood analysis at all nucleotide positions [52]. Bootstrap analysis was subsequently performed (DNABOOT) and values of 70% or more are indicated at the appropriate nodes; in general, bootstrap values of 70% may correspond to confidence values of 95% [16]. Accession numbers (not identified in the figure) are: HC-G9 (D14853), HCV-1 (M62321), HCV-J (D90208), ED43(Y11604), HC-J6 (D00944), HC-J8 (D10988), NZL1(D17763), HCV-Tr (D49347), JK049 (D63821), EUH1480 (Y13184), JK046 (D63822), and EUHK2 (Y12083)

parsimony using only second base positions within the NS3 helicase region. The inclusion of other flavivirus sequences in this analysis is important as it tends to argue against a monophyletic relationship.

The GBV-A and GBV-C/HGV viruses are closely related to each other phylogenetically, have similar genome structures (with a short or absent potential coding sequence for a homologue of the core gene) and are non-hepatotrophic. Based upon such criteria, an argument could be made for these viruses being considered as members of a new genus of non-hepatotropic *Flaviviridae*. The known sequences of the GBV-C/HGV viruses can be grouped into three geographic variants based upon sequence of the whole genome or sequence of the 5'UTR [33, 41, 51]. However, other subgenomic regions smaller than one-quarter of the complete genome, with the exception of the first 1200 5' terminal nucleotides or a fragment spanning nucleotides 3600–4800, do not reflect the pattern seen within the whole genome [51]. Given these observations, and the recent findings by Takahashi et al. [54], investigators are cautioned against assigning names until these

relationships are clarified. The GBV-A viruses have sequences that reflect the species from which they were derived [6, 24] although experimental studies have demonstrated that cross-species transmission can occur [6, 24].

The GBV-B virus causes hepatitis in New World monkeys [6, 43] and is more closely related to HCV than to GBV-A and GBV-C (Fig. 2); furthermore, it has a genome structure that resembles HCV, specifically with the presence of a reading frame encoding a core polyprotein. Based upon these characteristics, it may be considered as a member of the genus *Hepacivirus* or alternatively as a member distinct from, but sharing some similarities with the hepaciviruses.

Summary of recommendations

Hepacivirus genus

At this time, the genus *Hepacivirus* should be regarded as consisting of a single species, with HCV-1 as the prototype of the species. The future designation of species corresponding to the 6 clades would be appropriate if additional information such as cross-neutralization or cross-protection studies clearly distinguish these viruses. In the absence of such information, however, we conclude that it is difficult to argue for more than a single species.

Type nomenclature

To avoid confusion with any previous terminology and to reflect the inclusive nature of the groupings, we recommend that the term clade be used to identify the 6 genetic groups. Thus, genotypes 1, 2, 4, and 5 would be identified as clades 1, 2, 4, and 5, respectively; genotypes 3 and 10 would become members of clade 3, while genotypes 6, 7, 8, 9, and 11 would be members of clade 6 (Fig. 1).

Subtype nomenclature

The designation of subtype names may be important for the study of clinical differences within different clades of HCV, for the study of new therapies and vaccines, and for epidemiological investigations. A proposal for renaming currently identified subtypes based upon publication priority, number of isolates, and the presence of a complete genome sequence is being developed (Maertens et al., manuscript in prep.). In the future, we recommend that at least two epidemiologically unlinked isolates be identified which cluster together phylogenetically and are clearly distinguished from other subtypes.

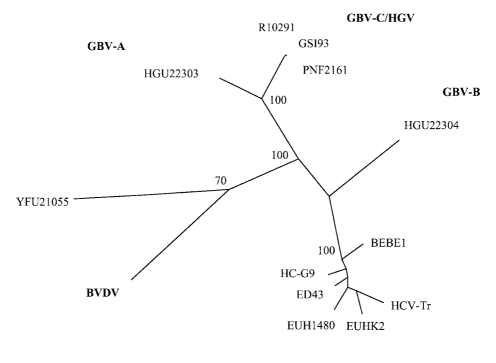
Approach to classify new sequences

In the interest of defining a common approach to classification of clades, subtypes, and isolates the following steps (illustrated in Fig. 3) are recommended : 1) Sequence the C, E1, or NS5B region. 2) Align sequences with representatives of each of the six clades. 3) Determine % similarity; the highest % similarity gives a tentative clade assignment. 4) Calculate DNADIST (Kimura 2 parameter option) using PHYLIP followed by the generation of a neighbor-joining tree using representatives of all known clades; bootstrap to confirm. These steps will result in identifying the clade assignment, or if the sequence appears to be a new clade, indicate that additional (complete genome) sequence information is needed.

To identify a subtype, steps 2–4 should be repeated with representatives of subtypes within the clade. Subtype identification can result in three possible outcomes: 1) The isolate is identified as belonging to an existing subtype. 2) The sequence appears to be a new subtype. However, prior to assigning a new subtype name, the sequence of at least one additional sample which is epidemiologically unrelated is needed. Alternatively, a full length sequence of the proposed new subtype would provide adequate data to support the designation of a new subtype. 3) The analyses are ambiguous. The sequences can be analysed by other phylogenetic methods or sent to one of the HCV research labs capable of performing in-depth analyses. As has been proposed previously [46], the sequence of two sub-genomic regions should be obtained and compared to verify the phylogenetic relationship of new HCV subtypes.

Database policies and development

For clarity in communication, investigators should note that subtypes are referred to with small letters (e.g., "5a"), whereas the protein components of HCV and HGV/GBV-C are characterized by capital letters (e.g., "NS5A"). It was suggested that, for consistency,



HCV

Fig. 2. Parsimony analysis illustrating the relationship of representative hepaciviruses, BVDV (pestivirus), yellow fever virus (flavivirus), GBV-A, GBV-B, and GBV-C. Inversely weighted parsimony was performed using PAUP 3.1.1 as previously described [17, 21]. To minimize noise, only homologous second base positions from NS3 coding sequences were studied: of the 601 total alignable sites, 526 varied sites were analyzed. Bootstrap analysis was subsequently performed (PAUP 3.1.1) and values of 70% or more are indicated at the appropriate nodes; in general, bootstrap values of 70% may correspond to confidence values of 95% [16]. Accession numbers (not identified in the figure) are: HCV-Tr (D49347), EUHK2 (Y12083), EUH1480 (Y13184), ED43 (Y11604), HC-J9 (D14853), BVDV (M31182), YFU21055 (U21055), HGU22303 (U22303), HGU44402 (U44402), R10291(U45966), GSI93 (D87263), and PNF2161 (U22304)

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nucleotide numbering should initiate at the AUG codon, with 5' terminal nucleotides given negative values numbered from $\langle -1$ to $-341 \rangle$. The term 5' UTR should be used to refer to the 5' region of the RNA that is untranslated. To avoid confusion and the possibility that identical names are given to individual samples, it was suggested that samples be identified by 1) the ISO two character country name [35], 2) identification of the laboratory in which the isolation and sequencing was performed, and 3) a sample number, i.e., USCDC10 (meaning United States, Centers for Disease Control, sample number 10). No patient initials should be used for virus identification, and the identifier should be kept to under 10 characters in total. In looking to the future, other useful information which could usefully be incorporated into a database format would include information such as age, sex, year of isolation, year of infection, etiology (transfusion, IVDU, sexual, unknown), disease stage (asymptomatic, CAH, LC, HCC) and ethnic origin.

Currently there is a database of all available HCV sequences (downloaded regularly from DDBJ [DNA DataBase of Japan]) available through the world wide web at http:// s2as02.genes.nig.ac.jp. The master database contains all HCV sequences with an alignment based upon the reference strain HCV HC-J4 (D10750). A reproduction of the database home page containing this graphic alignment is shown in Fig. 4; individual sequences are illustrated by lines under a schematic of the HCV genome indicating the various gene regions and products (adapted from [5]). Genome regions can be selected from this graphic map of the HCV genome and all sequences from that region can then be viewed as fully annotated entries – nucleic acid or amino acid sequences – and compared to each other by tree analysis. In a separate section of this database, there is FASTA similarity search capability.

The purpose of this communication is to solicit feedback on these proposed guidelines, and to generate ideas for future categories that would be useful for database development. A forum for discussion and contributions to these recommendations will be made available at the international HCV database at http://s2as02.genes.nig.ac.jp. One of the services that this database could provide is a source of curatorial information for researchers interested in evaluating sequence information from a wide range of perspectives – molecular, epidemio-

Clade Determination

1. Sequence C, E1, or NS5B region

2. Align sequences with representative sequences from each clade

3. Highest % similarity gives tentative clade assignment

4. Analyze using DNADIST of Phylip (Kimura 2-parameter), bootstrap to confirm

Subtype Determination

5. Repeat steps 2 - 4 with representatives of each subtype within the clade

Fig. 3. Flow chart illustrating the steps recommended to identify HCV clades and subtypes

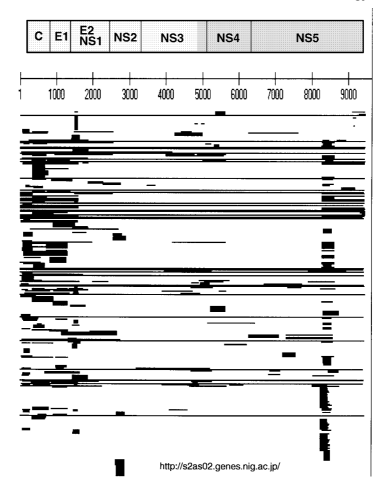


Fig. 4. Graphic of the HCV genome from the HCV Database (http://s2as02.genes.nig.ac.jp) illustrating alignment of complete and partial genome sequences. Each individual line represents an HCV sequence whose location has been aligned relative to the complete genome. The potential cleavage sites resulting in NS4A, NS4B, NS5A, and NS5B are not shown in this graphic

logic, and clinical. The types of information that are useful to these different groups may vary, and the ultimate usefulness will depend upon the inclusion of the maximum amount of appropriate information from each sample.

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