Rapid Identification of Known and New RNA Viruses from Animal Tissues

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Abstract

Viral surveillance programs or diagnostic labs occasionally obtain infectious samples that fail to be typed by available cell culture, serological, or nucleic acid tests. Five such samples, originating from insect pools, skunk brain, human feces and sewer effluent, collected between 1955 and 1980, resulted in pathology when inoculated into suckling mice. In this study, sequence-independent amplification of partially purified viral nucleic acids and small scale shotgun sequencing was used on mouse brain and muscle tissues. A single viral agent was identified in each sample. For each virus, between 16% to 57% of the viral genome was acquired by sequencing only 42–108 plasmid inserts. Viruses derived from human feces or sewer effluent belonged to the Picornaviridae family and showed between 80% to 91% amino acid identities to known picornaviruses. The complete polyprotein sequence of one virus showed strong similarity to a simian picornavirus sequence in the provisional Sapelovirus genus. Insects and skunk derived viral sequences exhibited amino acid identities ranging from 25% to 98% to the segmented genomes of viruses within the Reoviridae family. Two isolates were highly divergent: one is potentially a new species within the orthoreovirus genus, and the other is a new species within the orbivirus genus. We demonstrate that a simple, inexpensive, and rapid metagenomics approach is effective for identifying known and highly divergent new viruses in homogenized tissues of acutely infected mice.

Introduction

Clinical laboratories are rapidly adopting viral species-specific nucleic acid amplification for virus identification, thereby increasing the sensitivity of detection and reducing the time needed for diagnosis. Although widely successful, these methods are limited for detecting divergent viruses due to their high specificity. Failure rates in determining the etiological cause of disease are varied. For example, the rate for encephalitis is between 30–85% [reviewed in [1]], approximately 12% for acute flaccid paralysis [2], and for non A-E hepatitis between 18–62% [reviewed in [3]]. In cases where routine screening fails, newer technologies are now being employed. Prominent among these are microarrays and sequence-independent amplification and sequencing of viral nucleic acids [4,5,6,7,8,9,10,11,12,13,14].

Viral microarrays can be used to screen for all viral families simultaneously and have been used successfully to detect novel human rhinoviruses [10], human coronaviruses [10], and a human gamma retrovirus closely related to mouse retroviruses [14]. Microarrays require sufficient sequence similarities between virus and array oligonucleotides for hybridization to occur, making the detection of highly divergent viruses problematic. Sequence-independent amplification of nuclease protected viral particles [15] abrogates the need for a prior sequence information, allowing the detection of viruses recognizable through their protein sequence homologies to known viruses and has successfully been used to identify novel human and bovine paroviruses [4,8,15], polyomaviruses [7,16,17], anelloviruses [5], an arenavirus [18], a dicistrovirus associated with honey bee colony collapse disorder [19], and a seal picornavirus [9].

In this study we utilized sequence-independent amplification of partially purified viral nucleic acid from mouse tissue followed by low-scale shotgun sequencing to quickly identify the viral agents in five samples negative by tests available at the time of inoculations. Of the five viruses identified, two belonged to the Picornaviridae family, and three to the Reoviridae family.

Materials and Methods

Sample preparation and viral culture

VRDL1 isolate was originally derived from sewer effluent from Hamilton, Montana in 1959 and passaged five times through intracerebral inoculation of suckling mice. Subsequently, the viral stock was adapted to growth in primary rhesus monkey kidney cells, and mice were re-inoculated by intramuscular injection. Moribund mice were sacrificed; muscle tissue was collected. VRDL2 was isolated from the muscle of a moribund mouse inoculated with a fecal suspension of a 17 year old female suffering flu like symptoms in 1960. VRDL3 and VRDL4 were isolated from the brains of sucking mice paralyzed after inoculation of ground tissue from a pool of C. tarsalis mosquitoes or a pool of mosquitoes from multiple species, respectively. VRDL5 was isolated from the brain of a sucking mouse paralyzed after inoculation of filtered, ground skunk brain tissue in 1974. For all
**Author Summary**

Viral surveillance programs aim to identify circulating viruses to safeguard the public and livestock from viral outbreaks. Occasionally, samples suspected of harboring a virus cause severe disease in laboratory animals, but the identity of the virus eludes researchers. Here, we applied a simple viral discovery technique to identify viruses directly from the tissues of inoculated symptomatic mice and found a single virus in each sample using a rapid viral particle purification and random nucleic acid amplification method. Two viruses appear to be closely related to the members of the *Picornaviridae* family. In three other samples, originally collected from pools of crushed mosquitoes and the brain of a sick skunk, a known and novel viruses related to members of the *Reoviridae* family were identified. Reoviruses are considered major pathogens of livestock. Our studies provide the groundwork for further analysis of the prevalence and pathogenesis of these divergent viruses and illustrate the ease with which new viral species can be identified in tissues of acutely infected animals.

**Isolation of viral nucleic acid**

Cellular debris and bacteria were removed from mouse tissue homogenates by filtration through 0.45 μm filter (Millipore). Filtered viral particles were then pelleted by centrifugation at 22,000 g for 2 hr at 8°C and resuspended in Hanks buffered saline solution (Gibco, BRL). Non-particle protected (naked) DNA and RNA was removed by digestion with a cocktail of DNase enzymes consisting of 14 U of turbo DNase (Ambion), 20 U benzonase (Novagen) and 20 U of RNase One (Promega) at 37°C for 90 minutes in 1× DNase buffer (Ambion). Remaining total nucleic acid was then isolated using Qiamp Viral RNA isolation kit (Qiagen) according to manufacturer’s protocol. Purified viral RNA was protected from degradation by addition of 40 U of RNase inhibitor (Invitrogen) and stored at −70°C.

**Reverse transcriptase priming and amplification of nucleic acids**

Viral cDNA synthesis was performed by incubation of the extracted viral RNA/DNA with 100 pmol of primer K-8N (GAC CAT CTA GCC ACC TCC TGC NNN NNN N) [13], and 0.5 mM of each deoxynucleoside triphosphate (dNTP) at 75°C for 5 min. Subsequently, 5 U of RNase inhibitor, 10 mM dithiothreitol, 1× first-strand extension buffer, and 200 U of SuperScript II (Invitrogen) were added to the mixture and incubated at 25°C for 5 min, followed by 37°C incubation for 1 hr. In order to generate products with the fixed portion of primer K-8N on the opposite strand of the cDNA, single round priming and extension was performed using Klenow fragment (New England Biolabs). 20 μl of cDNA was heated to 95°C for 2 min and then cooled to 4°C in the presence of 20 pmol of primer K-8N in 1× klenow reaction buffer. 3 U of Klenow fragment were added and incubated at 37°C for 60 min.

PCRamplification was performed using 5 μl of the reaction described above in a total reaction volume of 50 μl containing 2.5 mM MgCl₂, 0.2 mM dNTPs, 1× PCR Gold buffer, 0.8 μM primer K (GAC CAT CTA GCC ACC TCC TGC AC), and 0.94 U of AmpliTaq Gold. Temperature cycling was performed as follows: 1 cycle of 95°C for 5 min, 5 cycles of denaturing at 95°C for 1 min, annealing at 59°C for 1 min, extension at 72°C for 1 min, 33 cycles of denaturing at 95°C for 20 sec, 59°C for 20 sec, 72°C for 1 min + 2 sec each cycle. An additional extension for 7 min at 72°C was added to the end of the run. Products were distinguished as a smear by agarose gel electrophoresis. Fragments larger than 500 bp were isolated (Qiagen) and subcloned into pGem T-easy vector (Invitrogen) for sequencing.

**Comparative sequence analysis**

Reference sequences were obtained from NCBI and edited using GENEDOC software. Accession numbers for sequences used in analyses are as follows: Picornaviruses: Simian picornavirus 1 (AY064708), Simian picornaviruses VPI (AAL69622-AAL69637), Human enteroviruses 89 (AY697459), 91 (AY697461), 83 (AY343301), 94 (EF107097), and 70 (DQ201177), 94 (EF107097), 70 (DQ201177), Coxsackie A viruses 5 (AF081296), 10 (AF081300), 3 (AF41342), 11 (AF499636), 15 (AF463512), 19 (AF08138) and 1 (AF499635), Coxsackie B4 virus (AF311939), Human echoviruses 30 (AF311938) and 2 (AY302545). Bovine Enteroviruses (AF129433, D00214), Porcine Enteroviruses (AF363453, AF406813), Duck Picornavirus TW09A (VP164335). Reoviruses: Eyach virus (NC_003696-NC_003707), Colorado Tick and Fever Virus (NC_004180_004191). Sequence alignments were generated using the CLUSTAL_W package with the default settings. CLUSTAL_W alignments were used to generate phylogenetic trees in MEGA 4 using either neighbor-joining, maximum likelihood or maximum parsimony with bootstrap values calculated from 1000 replicates. The generated phylogenetic trees were visualized using the program MEGA 4.

**Results**

**Summary of Viruses**

As part of infectious agent surveillance programs, sewer effluent, ground mosquito pools and skunk brain tissue collected between 1959 and 1980 by the California Department of Public Health were used to inoculate suckling mice intramuscularly or intracerebrally. In five samples, infectious agents could not be indentified in brain or muscle homogenates using available techniques, although the inoculated mice exhibited severe pathology. In this study, we used sequence-independent amplification of viral particle protected nucleic acids following nuclease digestion [8,13,15] of infected mouse tissue to identify a viral agent in each of these samples. These viruses are summarized in Table 1. In total, 47% of all plasmid subclones were viral in origin, indicative of a high viral prevalence in tissues of infected animals.

**Picornaviruses**

**Enterovirus.** According to California Department of Public Health records, sample VRDL2 produced flaccid paralysis in...
injected suckling mice with diffuse myositis of trunk and limb muscles. These symptoms are consistent with Coxsackie A virus infection. However, all serological tests were negative for enteroviruses [20]. In the current study, purified viral nucleic acids randomly amplified from VRDL2-infected mouse muscle confirmed infection with a moderately divergent Coxsackie A virus (CoxA). Pair-wise sequence comparisons of VRDL2 revealed between 74–86% nucleotide identity and 72–98% amino acid identity to CoxA1, CoxA19 and CoxA22, which comprise a monophyletic cluster within the human enterovirus C species (HEV-C). Coxsackie virus serotypes have been shown to correlate with their phylogenetic groupings based on sequence similarity, with approximately >75% nucleotide identity and >88% amino acid identity in the VP1 corresponding to the same serotype [21]. To determine the phylogenetic relationships between VRDL2 and serotypes of HEV-C several phylogenetic reconstruction algorithms (maximum likelihood, maximum parsimony and neighbor-joining) were used to compare HEV-C VP1 sequences (Fig. 1). At least three representatives, when available, from each member of HEV-C were used; a single member of human enterovirus B was included as an outgroup. As suggested by amino acid identity, VRDL2 segregated with CoxA22 with bootstrap values between 71–98%, as a basal member of the CoxA22 serotype.

**Sapelovirus.** Viral sequences obtained from sample VRDL1 represent a highly divergent picornavirus most closely related to simian picornavirus 1 (SV2), a member of the putative “sapelovirus” genus [22], with only 63% deduced amino acid identity within the originally sequenced VP2, VP3 and VP4 regions. Sub-genomic fragments were linked by PCR and both 5’- and 3’-RACE were utilized to acquire the complete VRDL1 polyprotein sequence which exhibited 80% amino acid identity to SV2. Except for the complete genome of SV2, sequence of simian sapeloviruses polyproteins currently are limited to VP1 and 3D regions [23,24]. To characterize the relation of VRDL1 within the sapeloviruses, the deduced amino acid sequence for VP1 was used for phylogenetic reconstruction (Fig. 2). VRDL1 VP1 clustered tightly to SV49 VP1, a divergent member within the sapelovirus cluster; the two sequences showing 91% amino acid identity (Fig. 2). Similar results were seen for partial sequence data within 3D (data not shown) [24]. We therefore report the complete polyprotein sequence of SV49 strain VRDL1 (SV49-VRDL1) which is sufficiently divergent from SV2, with amino acid identities ranging from 52.3% in VP1 to 98.9% within the conserved 3D polymerase, to be considered the prototypic genome sequence for a second simian sapelovirus genotype.

**Reoviruses**

**Eyach virus isolated in United States.** The VRDL4 sample was originally derived from a pool of California mosquitoes collected in 1980 as part of the arbovirus surveillance program. Viral sequences from VRDL4 shared strong amino acid identity to members of the *Coltivirus* genus. Eyach virus (EYAV), Colorado tick fever virus (CTFV), and CTFV-related viruses are the sole members of the *Coltivirus* genus. Partial sequencing of segments 1, 2, 3, 4, 5, 7, 8, and 9 all shared greater than 94% nucleotide identity to the prototype strain of EYAV, originally identified in *Ixodes ricinus* ticks in Germany in 1976 [25] (Fig. 3a). To date, the reported geographical distribution of EYAV has been limited to Europe, while CTFV (and related strains) have been isolated in the United States [26]. In a phylogenetic tree based on concatenated VP1 amino acid sequence, VRDL4 paired with EYAV, with strong bootstrap support (data not shown). Overall amino acid and nucleotide
identities across all segments sequenced suggest VRDL4 represents the first isolation of EYAV (EYAV strain VRDL4) from the United States. (Fig. 3b).

**Novel Orthoreovirus.** VRDL5, isolated from skunk brain tissue intracerebrally inoculated into suckling mice, displayed significant sequence identity to several fragments of mammalian orthoreoviruses. Sequence from VRDL5 was obtained for 5 of the 9 segments, with amino acid identities to known mammalian orthoreoviruses ranging from 23% to 79%. The greatest amount of coverage (979bp of 1150bp) was obtained for the S3 fragment encoding the sigma NS protein. This fragment shared approximately 78% amino acid identity with baboon reovirus (Fig. 4). For the S4 segment, VRDL sequences also paired with baboon reovirus. To date, no baboon reovirus class L or M segments have been published. VRDL5 fragment M2 shared closest sequence identity to avian orthoreovirus, while both L2 and M3 were weakly similar to mammalian orthoreoviruses. We propose VRDL5 represents a new orthoreovirus, tentatively named “skunk orthoreovirus”, which shares greatest similarity to baboon reovirus.

**Novel Orbivirus.** Sample VRDL3 consisted of brain tissue from suckling mice that had been inoculated intracerebrally with pooled *Culex tarsalis* homogenate. Viral sequences from VRDL3 displayed closest similarity to members of the Orbivirus genus in the Reoviridae family. Sequence from six of the ten total segments of Orbiviruses displayed weak amino acid identities (19–36%) to subcloned VRDL3 fragments (Fig. 5). Amino acid sequence from VP7, a structural protein, which along with VP3 forms the viral capsid [27], is the protein primarily used for phylogenetic analysis of Orbiviruses and exhibits the highest level of conservation within members of the Orbivirus genus with intraspecies amino acid
Figure 3. Eyach Virus. (A) Positions of VRDL4 fragments amplified relative to segmented Coltivirus genome. (B) Pairwise amino acid (nucleotide) percent identities between VRDL4 and CTFV and VRDL4 and EYAV for each segment.
doi:10.1371/journal.ppat.1000163.g003

Figure 4. Novel Orthoreovirus. Alignment of six VRDL5 fragments amplified through viral discovery approach to ten genomic fragments of model orthoreovirus with amino acid identities (bottom). Neighbor-joining tree from orthoreovirus sigmaNS protein with bootstrap values from 1000 pseudo-replicates (top).
doi:10.1371/journal.ppat.1000163.g004
identities between 83–99% [28,29,30]. VRDL3 VP7 exhibited only 19% amino acid identity to the closest Orbiviruses, and phylogenetic analysis revealed a deep, weakly bootstrap-supported pairing to the Great Island Broadhaven virus (Fig. 5). We propose VRDL3 is a novel virus and propose the name California Mosquito Pool Virus (CMPV), in accordance with traditional nomenclature of Orbiviruses.

Several members of the Orbivirus genus, including Bluetongue virus (BTV) and epizootic hemorrhagic disease virus (EPHDV), contain a conserved RGD motif located between positions 168 to 170 of VP7 shown to be necessary for attachment to Culicoides cells [31]. This motif is missing in Orbiviruses transmitted by mosquitoes, (e.g. Yunnan virus), or ticks [e.g. Broadhaven (BRDV) and St. Croix River virus (SCRV)]. CMPV VP7 contained a RHD motif, similar to the conserved RGD motif of BTV and EPHDV, despite being isolated from Culex mosquitoes. BTV-10 VP7 crystal structure [32] and functional analysis also have revealed several conserved residues within nine alpha helices involved in formation of the icosahedral viral core. Predicted secondary structure from CMPV VP7 partial sequence reveals the presence of the four C-terminal alpha helices involved in formation of the viral core (data not shown) and includes conserved residues corresponding to BTV-10 D318 and Y271, the latter shown to be necessary for stable trimerization and particle formation [27].

**Discussion**

Genetic and antigenic diversity and the presence of uncharacterized viruses can confound virus identification using species or serotype specific reagents. PCR tests generally exhibit increased sensitivity relative to antibody based antigen detection. However, viral genetic diversity can prevent the detection of viral variants. Use of degenerate consensus primers targeting conserved viral regions can improve detection of variants, but often at the cost of decreased sensitivity.

In this study, we circumvent the need for prior knowledge of viral sequence using sequence independent amplification of nuclease protected viral particles [15]. We tested samples of ground mouse muscle or brain that had been inoculated with suspected infectious material including human feces, ground mosquito pools, and sewer effluent. Processing of samples and identification of five previously undetected viruses was achieved in under a week. From each of the samples tested, a single viral species was identified. Typically, the majority of clones sequenced contained viral sequence. Of the five viruses amplified two were picornaviruses. One picornavirus was a borderline new serotype of HEV-C, most closely related to CoxA22, and the other was a member of the putative sapelovirus genus, SV49 strain VRDL1. Three reoviruses were identified: one was a very close match to Eyach virus, while the other two were sufficiently divergent from members of the *Orbivirus* and *Orthoreovirus* genera to be candidates for new species with the *Reoviridae* family. The Eyach virus, derived from pooled mosquitoes collected in 1980 is the first reported case of Eyach detection in the US. Eyach virus previously was thought to be restricted to Europe, while the related coltivirus species CTFV was found only in the US [26]. Current prevalence in US mosquitoes remains to be determined.

**Identification of novel viruses**

We consider it unlikely that SV49-VRDL1, a simian sapelovirus, was derived from the sewer effluent used as inoculum, but...
instead was likely a contamination of lengthy in vitro manipulation in primary monkey kidney cells prior to mouse inoculation. Isolation of several simian enteroviruses in the early development and testing of poliovirus vaccine in primary monkey kidney cells illustrates the hazard of cell culture introducing viral contaminants [33,34,35,36]. Regardless of its origin, we report the complete genome sequencing of SV49-VRDL1, a second simian sapelovirus exhibiting an overall amino acid identity of 80% and 77% nucleotide identity compared to SV2.

Orthropiruses currently consist of four species: mammalian orthopirivirus (MRV), avian orthopirivirus (ARV), Nelson Bay orthopirivirus (NBV), and Baboon reovirus (BRV), although complete genetic sequence of BRV has not been described. Additionally, incomplete genomes from three tentative species have been described recently in reptiles [37]. Depending on the host and viral species, orthopiruses can cause respiratory illness, diarrhea, and severe neurological disorders. Infection in humans is generally benign, but can cause respiratory or gastrointestinal illness, with increased severity in infants [38,39]. Demarcation of orthopirivirus species is based on sequence divergence, host species, and pathogenesis.

To determine if the skunk orthopirivirus identified represents a new species of orthopirivirus or a novel variant of the baboon reovirus species, intraspecies and interspecies sequence diversity within the orthopirivirus genus were examined. Deduced sigma non-structural protein sequences share 83–98% amino acid identity within avian orthopiruses (ARV) and 73%–94% identity within mammalian orthopiruses (MRV) [40,41]. Pair-wise amino acid comparisons between AVR, BRV, and MRV species exhibit identities between 18–53% [37,40]. The skunk orthopirivirus S3 fragment shares 78% amino acid identity to baboon reovirus, falling into a gray zone with regard to species demarcation. Similarly, the BRV and skunk orthopirivirus S4 segment encoding the fusion-associated small transmembrane protein exhibit 79% amino acid identity. Due to the paucity of known baboon reovirus sequences in Genbank we cannot make direct comparisons with class L and M orthopiruses.

Comparisons between AVR, BRV, and MRV species exhibit identities below 36% to all described Orthopiruses, qualifying it as a new Orthopivirus species. Transmission of Orthopiruses involves three primary vectors; midges (BTV, AHSV), ticks (BRDV, SCRV) and mosquitoes (Yunnan virus). While phylogenetic analysis of VP7 indicated CMPV branches with very low bootstrap support with BRDV, CMPV was isolated from C. tarsalis mosquito pools while BRDV is tick-borne [40]. Taken together these data indicate CMPV may be the founding member of a new species within the Orthopirivirus genus.

These results indicate that known and novel viruses can be readily characterized using limited sequencing, provided that they are in the high concentration expected from affected tissues of recently inoculated mice showing acute pathologies. Application of such techniques to hard-to-type isolates may therefore accelerate the identification of new viral species.

Acknowledgments

We thank Dr. Michael P. Busch and Blood Systems Research Institute for sustained support.

Author Contributions

Conceived and designed the experiments: JGV AK ELD. Performed the experiments: JGV AK. Analyzed the data: JGV AK. Contributed reagents/materials/analysis tools: JGV KD DPS. Wrote the paper: JGV ELD.

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