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High Variety of Known and New RNA and DNA Viruses of Diverse Origins in Untreated Sewage

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Deep sequencing of untreated sewage provides an opportunity to monitor enteric infections in large populations and for high-throughput viral discovery. A metagenomics analysis of purified viral particles in untreated sewage from the United States (San Francisco, CA), Nigeria (Maiduguri), Thailand (Bangkok), and Nepal (Kathmandu) revealed sequences related to 29 eukaryotic viral families infecting vertebrates, invertebrates, and plants (BLASTx E score, <10−4), including known pathogens (>90% protein identities) in numerous viral families infecting humans (Adenoviridae, Astroviridae, Caliciviridae, Hepeviridae, Paroviridae, Picornaviridae, Picobirnaviridae, and Reoviridae), plants (Alphaflexiviridae, Betaflexiviridae, Partitiviridae, Sobemovirus, Secoviridae, Tombusviridae, Tyurnoviridae, Virgaviridae), and insects (Dicistroviridae, Nodaviridae, and Paroviridae). The full and partial genomes of a novel kobuvirus, salivirus, and saposivirus are described. A novel astrovirus (casa astrovirus) basal to those infecting mammals and birds, potentially representing a third astrovirus genus, was partially characterized. Potential new genera and families of viruses distantly related to members of the single-stranded RNA picorna-like virus superfamly were genetically characterized and named Picalivirus, Secalivirus, Hepelivirus, Nedicistrovirus, Cadicistrovirus, and Niflaviru. Phylogenetic analysis placed these highly divergent genomes near the root of the picorna-like virus superfamly, with possible vertebrate, plant, or arthropod hosts inferred from nucleotide composition analysis. Circular DNA genomes distantly related to the plant-infecting Geminiviridae family were named Baminivirus, Nimivirus, and Nimivirus. These results highlight the utility of analyzing sewage to monitor shedding of viral pathogens and the high viral diversity found in this common pollutant and provide genetic information to facilitate future studies of these newly characterized viruses.

The characterization of previously unknown viral genomes has accelerated following the introduction of high-throughput sequencing technologies. We reasoned that untreated sewage would provide a rich source of both known and previously uncharacterized viruses with which to expand the reach of the existing viral taxonomy and also provide candidate viral genomes for future disease association studies, particularly of idiopathic human enteric diseases. Enteric infections cause more than 2 million deaths each year (93), primarily among infants in developing countries (59, 93). In the United States, greater than 40% of cases of diarrhea are caused by unknown agents (33). Viruses in sewage, which reflect in part ongoing enteric infections in the sampled human population (79), may therefore include still unknown human pathogens. Studies of sewage have frequently been used to monitor known viral pathogens, most frequently, polioviruses (6–9, 36, 88, 94). Viruses in untreated sewage may disseminate through rivers, plant irrigation, and fish and shellfish production and affect downstream human, animal, and plant health (27).

Sewage includes feces and urine from humans as well as from domesticated and wild animals, such as pets and rodents. Numerous bacteria, archaea, unicellular eukaryotes, plants, and insects and their associated viruses are also expected within sewage systems. The large number of potential host species contributing to sewage provides an opportunity to sample the viral diversity infecting cellular organisms from all kingdoms of life.

Recent studies have shown that besides human viruses, untreated sewage also contains a wide diversity of other animal, plant, insect, and bacterial viruses (11, 76, 82, 89). We performed a metagenomic deep-sequencing analysis of viral particles in sewage from four countries on three continents. We also acquired the full or partial genomes of several selected viruses and phylogenetically compared them to previously known viruses. We report here on the wide diversity of eukaryotic viruses found in sewage and on multiple novel viral genomes, significantly increasing the known diversity of viruses, especially those related to the single-stranded RNA (ssRNA) picorna-like virus superfamly.

MATERIALS AND METHODS

Sample collection. Untreated sewage waters were collected from a polluted canal in Khlong Maha Nak, Yommarat, Pom Prap Sattru Phai, Bangkok, Thailand, on 18 June 2009; from a junction of a main sewage line and river at the Kalimati Bridge in Kathmandu, Nepal, on 14 August 2009; from the city of Maiduguri (Gombaru Ward) in Nigeria on 15 April
2008; and from the southeast water pollution control plant in San Francisco, CA, on 15 May 2009.

Viral particle purification and sequence-independent nucleic acid amplifications. Following collection, sewage samples were shipped on dry ice and stored at −80°C. Six hundred to 800 ml from each location was thawed in the dark at 4°C over 72 h. Sewage samples were centrifuged at 4,000 × g for 15 min at 4°C followed by a second centrifugation at 10,000 × g for 15 min in a Beckman SW55Ti rotor at 4°C to remove large particulates and bacteria. The resulting supernatant was subjected to 0.22-μm pore-size tangential-flow filtration, and the collected viral fraction was concentrated to 12 ml using a 30-kDa tangential-flow filter (PKGVFPC30 and PXCO30C50; Millipore). Viruses were enumerated using SYBR gold epifluorescence microscopy (73). Six milliliters of the viral fraction were further concentrated by sucrose cushion centrifugation (38%, wt/vol) in a Beckman SW28 rotor. Pelleted viruses were resuspended in 500 μl SM buffer (100 mM NaCl, 8 mM MgSO4 · 7H2O, 50 mM Tris-HCl [pH 7.5]). The resuspended virus particles were treated with a cocktail of DNases (Turbo DNase from Ambion, Baseline-ZERO from Epicentre, and Benzonase from Novagen) and RNase (Fermentas) to digest unprotected nucleic acids (10, 69, 72, 92). Viral nucleic acids were then extracted using QIAamp spin columns (Qiagen).

Viral RNA and DNA were used to construct libraries by random PCR amplification as previously described (92). Twelve different primers (an arbitrarily designed 20-base oligonucleotide followed by a randomized octamer sequence at the 3′ end) were used in separate reverse transcription (RT) reactions for each sample (Invitrogen). After denaturation, the cDNA was then subjected to a round of DNA synthesis using Klenow fragment polymerase (New England BioLabs), followed by PCR amplification using a primer consisting of only the 20-base fixed portion of the random primer. The amplification from each primer was then quantified and pooled in an equal amount, and DNA libraries were prepared and sequenced using a 454 GS FLX titanium platform.

Bioinformatics. Pyrosequencing reads were trimmed and parsed according to their primer tags. For each sample, sequences sharing more than 95% nucleotide sequence identities over 35 bp were assembled into contigs. Metagenomic reads, assembled contigs, and nonassembled singlets were compared to the sequences in the GenBank nonredundant nucleic acid database using BLASTn and BLASTx and an E-value cutoff of 10−3. On the basis of the best BLAST result, sequences were classified into their likely taxonomic groups of origin. Sequence identity distribution analysis was performed by parsing the BLASTx result for the taxonomy, host, and percent protein identities of the local pairwise alignment provided by BLASTx.

Acquisition of novel virus genomes. Total nucleic acid was extracted from the purified virus preparation by using QIAamp spin columns. In order to amplify the extremities of the viral genomes, both 3′ and 5′ rapid amplification of cDNA ends (RACE) amplification kits (Invitrogen and Clontech) were used according to the manufacturer’s instructions. In short, three primers were designed for each virus genome. For 3′ RACE, the first primer was used in a reverse transcription reaction to produce cDNA from the poly(A) 3′ ends. For 5′ RACE, two additional steps were performed: cDNA launched from a primer complementary to the viral sequence was purified using spin columns, and poly(C) was added to the cDNA 5′ end using terminal deoxynucleotidyltransferase and dCTP. For the first round of PCR amplification, cDNA was amplified using a virus-specific primer and a RACE-specific primer ending with poly(G) or poly(A). The second round of PCR was performed using the second downstream primer, together with another RACE-specific primer without the poly(G) or poly(A) supplied by the RACE kit. Amplicons were analyzed using gel electrophoresis and Sanger sequenced by primer walking until the extremities were obtained. The genome sequence fragments originally derived from 454 pyrosequence data were confirmed using Sanger sequencing.

To complete the circular DNA genomes, we used rolling-circle and inverse PCR amplification as described previously (68, 69). Viral nucleic acids were first nonspecifically amplified using random hexamers by rolling-circle amplification (Genomiphi; GE Healthcare) and then further amplified by specific primers designed to amplify the whole circular genome using inverse PCR (70, 71). PCR amplicons were then Sanger sequenced.

Phylogenetic analysis. Phylogenetic analyses were performed using novel virus sequences, their closest BLAST hits, and other type species from related viral genera or families. Due to the divergent nature of the virus genomes, all sequence alignments and phylogenetic analyses were performed on the translated amino acid sequences. Multiple-amino-acid-sequence alignments were performed using the MUSCLE (version 3.8) (19) and MAFFT (50) programs, and the best overall alignments were selected for further analysis. Pairwise distance analyses and conserved amino acid analyses were performed over the alignments produced. Maximum likelihood (ML) trees were generated from translated protein sequences using RAxML and PROTGAMMA and Dayhoff similarity matrix parameters (87). These specify a general time-reversible model with a gamma distribution for rates over sites. All model parameters were estimated by RAxML. ML trees were run with 100 bootstrap replications; branches with 60% or greater bootstrap support are labeled. Resulting trees were examined for consistency with published phylogenetic trees. For trees without an outgroup, midpoint rooting was conducted using the program MEGA (90). Sliding-window analyses were performed using the same alignment, with protein identities between translated sequential segments of 32 in-frame codons, incrementing by 8 codons, being calculated (66, 85).

NCA. Nucleotide composition analysis (NCA) was performed as previously described (46, 84) using sequences infecting mammals (n = 117), insects (n = 63), and plants (n = 167) for classification. The frequencies of each mononucleotide and dinucleotide were used for discriminant analysis to maximize discrimination between control sequences; these canonical factors were then used to infer the host origin of the RNA virus sequences obtained in the current study.

Nucleotide sequence accession numbers. All sequenced genomes were deposited in GenBank under accession numbers JQ898331 to JQ898345. Pyrosequences were deposited in GenBank under short-read archive accession number SRA054852. The GenBank accession numbers of all viral taxa used in the phylogenetic trees in Fig. 3 to 9 are listed (see Table S1 in the supplemental data).

RESULTS

Untreated raw sewage was collected from four locations: San Francisco (United States), Bangkok (Thailand), Kathmandu (Nepal), and Maiduguri (Nigeria) (see Materials and Methods). Epifluorescence microscopy showed 1.46 ± 1010, 9.26 ± 1010, 4.0 ± 108, and 6.01 ± 109 virus particles per ml, respectively. Viral particles were purified using tangential flow filtration, sucrose cushion centrifugation, and nuclease treatment (see Materials and Methods). Nucleic acids were then extracted and amplified by random RT-PCR using primers with degenerate 3′ ends (92). The amplified nucleic acids were prepared into DNA libraries for pyrosequencing in two 454 FLX runs. A total of 304,498, 392,638, 217,383, and 161,798 sequence reads were generated from the U.S., Thai, Nepalese, and Nigerian libraries, respectively (Fig. 1). Overlapping sequence reads were assembled into contigs, and both contigs and singlets of >100 bp were analyzed for translated protein similarities to sequences in the nonredundant database of GenBank using the best BLASTx with a threshold E score of 10−4 (see Materials and Methods). The resulting classification (see Fig. S1 and S2 in the supplemental material) indicates that sequences other than those of viruses remained in our virus concentrates. Recognizable phage sequences were found at a frequency of 13.5% relative to the 11% of sequences showing similarity to eukaryotic viruses. A large
fraction (37%) of the sequences was unrecognizable on the basis of the BLASTx criterion of an E value of $<10^{-4}$. In contrast to other environmental metagenomes, these results indicated that in these sewage samples, eukaryotic viruses and phages were present in roughly equivalent numbers. The large number of unclassifiable sequences, within the range reported by other studies of purified viral particles, is likely to consist, at least in part, of highly divergent prokaryotic and eukaryotic viral sequences unrecognizable by BLASTx against the current viral database.

Twenty-nine eukaryotic viral families represented in sewage viromes. Approximately 110,000 sequences derived from the four sewage libraries exhibited sequence similarity to eukaryotic viruses (BLASTx E score, $<10^{-4}$), including matches to at least 29 different eukaryotic viral families (Fig. 1 and 2; Table S13 in the supplemental material contains the assembled sequences and sequence identity information). The most frequently amplified viral sequences belonged to families infecting invertebrates, followed by viruses found in diverse environmental samples but without defined hosts, plant viruses, viruses of vertebrates, and finally, human viruses at 6% of all identified eukaryotic viral sequences (see Fig. S1 and S2 in the supplemental material). When the threshold of sequence similarity was increased to $>90\%$ amino acid sequence identities, we detected members of 18 eukaryotic viral families (Table 1; see Tables S2 to S4 in the supplemental material). All viruses identified in sewage were nonenveloped viruses. Sequence matches to large-genome lipid-enveloped double-stranded DNA (dsDNA) viruses, including asfaviruses, poxviruses, and herpesviruses, were also detected but upon visual inspection of the alignments were excluded, as these consisted of repeat sequences that were possibly the result of amplification or sequencing artifacts.

**Enteric human viral pathogens.** The human viruses detected included members of seven distinct families (Table 1). Many recently described human viruses were detected (cosavirus [34, 49], cardiovirus [1, 12, 17, 42], salivirus/klassevirus [31, 35, 65], bacoviruses 2 and 3 [4, 47, 48]), underlining sewage’s potential for further human virus discovery. Californian and Nepalese sewage contained the greatest diversity of human viruses. Astroviruses and caliciviruses (noroviruses and sapoviruses) were detected in California, while Nepal’s human viruses consisted mostly of picornaviruses. Poliovirus type 2 vaccine strain (Sabin-2) was detected in the Nepalese sample as a pyrosequence with 100% nucleotide sequence identities to poliovirus 2 (reference genome positions 4329 to 4428). Human Saffold viruses (family Picornaviridae, genus Cardiovirus) were found in both the United States and Nigeria. Aichi viruses (family Picornaviridae, genus Kobuvirus) were found in Nepal, Thailand, and the United States. Bacoviruses 2 and 3 were found in Nepal and the United States, respec-

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**FIG 1** Taxonomic distributions of the eukaryotic virus-related sequences from four sewage samples. Assembled sequences were compared to the sequences in the nonredundant protein database using BLASTx (E score, $<10^{-4}$). The number of sequences with identities to eukaryotic viruses is shown. Twenty-nine eukaryotic virus families were detected from human, vertebrate, plant, and insect hosts (as well as unclassified viruses and satellites).

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**Sewage Viromes from Four Countries**

**San Francisco, CA, USA n=11530**

**Maiduguri, Nigeria n=24438**

**Kathmandu, Nepal n=56747**

**Bangkok, Thailand n=21709**
Human hepatitis E virus (HEV) was detected in the Nepalese sample only. In terms of sequence read numbers, the most strongly represented human viruses were all positive ssRNA viruses: Aichi viruses, tenuivirus, koaviruses, saliviruses, astroviruses, coronaviruses, neproviruses, and parvoviruses. Overall, sequences related to human viruses by best BLASTx comparisons in U.S. sewage showed higher percent identities to database viruses (86%) than similar sequences from the other three countries (74% to 78%). This greater degree of similarity may reflect the larger contribution to GenBank of viral sequences from the United States and other developed countries relative to viral sequences from less developed countries.

**Virology in sewages**

High diversity of vertebrate, plant, and insect viruses in sewage viromes. Viruses infecting other eukaryotes besides humans were also abundant. Overall, at least 29 eukaryotic viral families were found in the sewage samples (Fig. 1 and 2).

**Vertebrate viruses**

<table>
<thead>
<tr>
<th>Virus Family</th>
<th>Vertebrate Viruses</th>
<th>Percentage Identities %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Astroviridae</td>
<td>Mammal mamastrovirus</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Caliciviridae</td>
<td>Mammoth reovirus</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Picornaviridae</td>
<td>Mammoth picornavirus</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Reoviridae</td>
<td>Human enterovirus</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Adenoviridae</td>
<td>Human adenovirus</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Anelloviridae</td>
<td>Human anellovirus</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Paroviridae</td>
<td>Human parovirus</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
</tbody>
</table>

**Plant viruses**

<table>
<thead>
<tr>
<th>Virus Family</th>
<th>Plant Viruses</th>
<th>Percentage Identities %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alphaflexiviridae</td>
<td>Tenuivirus</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Betaflexiviridae</td>
<td>Koavirus</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Luteoviridae</td>
<td>Betaflexiviridae</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Potyviridae</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Secoviridae</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Tombusviridae</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Tombusviridae</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Tyomoviridae</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Virgaviridae</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Unclassified</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Gemmiviridae</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Nanoviridae</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Partitiviridae</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
</tbody>
</table>

**Invertebrate viruses**

<table>
<thead>
<tr>
<th>Virus Family</th>
<th>Invertebrate Viruses</th>
<th>Percentage Identities %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacillariomaviridae</td>
<td>Bacillariomaviruses</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Dicistroviridae</td>
<td>Dicistroviruses</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Picornaviridae</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Novadaviridae</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Tetraviridae</td>
<td>Tetaviruses</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Totiviridae</td>
<td>Totiviruses</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Unclassified</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Birnaviridae</td>
<td>Birnaviruses</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Circoviridae</td>
<td>Circoviruses</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>Paroviridae</td>
<td>Paroviruses</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
<tr>
<td>As coviridae</td>
<td>Unclassified</td>
<td>0.4 0.6 0.8 1.0</td>
</tr>
</tbody>
</table>

These divergent viral sequences showed sequence similarities to the Astroviridae, Caliciviridae, Picornaviridae, Novadaviridae, Adenoviridae, Circoviridae, and Paroviridae families (Fig. 2), reflecting an abundance of novel animal viruses in sewage.

Plant viruses accounted for 21% of the total viral reads, including both known plant viruses and more divergent viral sequences (Fig. 2 and see Fig. S1 in the supplemental material). Known plant pathogens included alphaflexiviruses, betaflexiviruses, parviviruses, secoviruses, sobemovirus, tombusviruses, tymoviruses, and virgaviruses (see Table S4 in the supplemental material). An even larger proportion of the plant virus-like sequences were from more divergent plant viruses (Fig. 2; <90% identities). Sewage therefore contained 12 out of the total 21 known plant viral families, dominated by major groups of nonenveloped plant viruses.

Known insect viruses, including dicistroviruses, nodaviruses, and densovirus (>90% amino acid sequence identities; see Table S3 in the supplemental material) that infect bees, mosquitoes, and other insects, were found in sewage. The majority of the insect virus sequences belonged to more divergent insect viruses that...
shared <90% amino acid similarities with known insect viruses (Fig. 2).

In summary, the sewage contained a very high diversity of known (Table 1; see Tables S2 to S4 in the supplemental material) and divergent (Fig. 2) viruses infecting multiple kingdoms of eukaryotic hosts. Several divergent viral sequences were selected for further directed genome sequencing (Fig. 2, numbered dots), resulting in the characterization of the novel viral genera and families described below. The large number of divergent sequences that were not extended indicates that numerous other novel viral genomes remained only minimally characterized.

**Novel picornaviruses in the kobuvirus and salivirus genera.** Using sequences showing similarities to human picornaviruses as starting points for genome extension, we acquired near complete genomes of new viruses using RT-PCR and 5’ and 3’ RACE amplifications, followed by Sanger sequencing.

From the Nepalese sample, a near complete genome of a novel kobuvirus in the *Picornaviridae* family was characterized and was named kobuvirus sewage Kathmandu (KoV-SewKTM) after the location of its discovery. Related human kobuviruses, the pathogenic Aichi viruses, have been associated with gastroenteritis (3, 30, 63, 77, 83, 95–98), while other kobuviruses have been detected worldwide in pigs and cows with diarrhea (51, 81) as well as in mouse and canine feces (45, 64, 78). KoV-SewKTM (GenBank accession number JQ898342) was 6,939 nucleotides (nt) long and shared ~82% nucleotide sequence identities to human Aichi virus, canine kobuvirus, and murine kobuvirus over its genome (see Fig. S3A in the supplemental material). Pairwise distance analysis indicated that KoV-SewKTM shared the highest amino acid sequence identities in the P1 region with mouse kobuvirus (87%) and was equidistant to other kobuviruses in the 2C plus 3CD region (86% to 88%) (see Table S5 in the supplemental material). Phylogenetic analysis of the P1 region confirmed these relationships (Fig. 3). Since members of each kobuvirus species share >70% amino acid sequence identities in P1 plus >80% amino acid sequence identities in the 2C plus 3CD region (54), KoV-SewKTM, together with murine and canine kobuviruses, all fit into the human Aichi virus species on the basis of the species demarcation criterion used for enteroviruses (54) and recently substantiated by a larger analysis of picornavirus diversity (60). Since the VP1 of KoV-SewKTM shared less than 84% amino acid sequence identities to the Aichi virus-related viruses, a threshold functionally determined to differentiate enterovirus genotypes corresponding to serotypes (74), KoV-SewKTM may qualify as a candidate for a novel kobuvirus genotype.

A near complete genome of a novel salivirus was also sequenced from the Bangkok sample and was named salivirus sewage Bangkok (SaliV-SewBKK; GenBank accession number JQ898343). The near complete genome of SaliV-SewBKK was 6,397 nt long, containing the full-length polyprotein except for only a partial VP0 and no 5’ untranslated region (UTR) at the 5’ end. Currently, human saliviruses (also called klasseviruses) have been associated with diarrhea and detected in feces from both

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**TABLE 1 Distribution and sequence identities of sequence reads in sewage with >90% amino acid identities to known human viruses**

<table>
<thead>
<tr>
<th>Viral family</th>
<th>Virus</th>
<th>% amino acid identity to known human viruses</th>
<th>Nigeria Minimum</th>
<th>Nigeria Maximum</th>
<th>Nepal Minimum</th>
<th>Nepal Maximum</th>
<th>Bangkok Minimum</th>
<th>Bangkok Maximum</th>
<th>California Minimum</th>
<th>California Maximum</th>
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<tbody>
<tr>
<td><em>Astroviridae</em></td>
<td>Human astrovirus</td>
<td>91</td>
<td>91</td>
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<td></td>
<td></td>
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<td></td>
<td>Human astrovirus 1</td>
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<td>100</td>
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<tr>
<td></td>
<td>Human astrovirus 3</td>
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<td>100</td>
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</tr>
<tr>
<td></td>
<td>Human astrovirus 8</td>
<td>97</td>
<td>97</td>
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<tr>
<td><em>Caliciviridae</em></td>
<td>Norovirus Hu/GI/Ototoke/1979/JP</td>
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<td>91</td>
<td></td>
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<tr>
<td></td>
<td>Sapovirus SaKaeo-15/Thailand</td>
<td>98</td>
<td>98</td>
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<td><em>Hepeviridae</em></td>
<td>Hepatitis E virus</td>
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<td>100</td>
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</tr>
<tr>
<td><em>Parvoviridae</em></td>
<td>Adeno-associated virus</td>
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gastroenteritis patients and healthy subjects from Nigeria, Tunisia, Nepal, Australia, and the United States, as well as in sewage from Spain, suggesting a widespread geographic distribution (31, 35, 65). Phylogenetically, the salivirus from sewage from Bangkok (SalIV-SewBKK) branches out at a basal position to the known saliviruses that have recently been proposed to form their own genus in the family Picornaviridae (54, 60). Saliv-SewBKK was equidistant to other saliviruses in the P1 region (85% to 86%), 2C plus 3CD region (90%), and VP1 region (82% to 85%), as shown in both similarity plot and pairwise distance analysis (see Tables S5A to C in the supplemental material). Comparing such percent identity with criteria for enterovirus serotypes, SaliV-SewBKK may be considered the second serotype of the salivirus species since the prior saliviruses/klasseviruses from human feces were all equidistant to other saliviruses in the P1 region (85% to 86%), 2C plus 3CD region (90%), and VP1 region (82% to 85%), as shown in both similarity plot and pairwise distance analysis (see Tables S5A to C in the supplemental material). Comparing such percent identity with criteria for enterovirus serotypes, SaliV-SewBKK may be considered the second serotype of the salivirus species since the prior saliviruses/klasseviruses from human feces were all closely related (see Table S5 in the supplemental material).

New sapovirus in California. A new sapovirus, namely, sapovirus sewage/California/2009 (SaV-SewSFO; GenBank accession number JQ898334-6), was tentatively named from the California sewage sample. A fraction of its genome (1,385 nt long), including half of the capsid, was sequenced and phylogenetically compared to the other sapoviruses within the Sapovirus genus (see Fig. S5 in the supplemental material). On the basis of genetic distance criteria and clustering, it appears that this virus may be classified either as a divergent member of genogroup 2 or as a member of a new genogroup.

A divergent astrovirus related to mamma- and avastroviruses. Nearly half of the genome of a highly divergent astrovirus was acquired from the California sewage sample and was tentatively named casa (for California sewage-associated) astrovirus (AstV-casa; GenBank accession number JQ898338), was characterized from the California sewage sample. A fraction of its genome (3,206 nt) consists of a highly divergent astrovirus, placing it at the root of the Astroviridae family (Fig. 4). The 3’ UTR of AstV-casa lacked the stem-loop-2-like motif described in a subset of known astroviruses (40). Phylogenetic analyses indicated that AstV-casa was highly divergent from the other astroviruses, placing it at the root of the Astroviridae family (Fig. 4). Pairwise distance analysis showed that AstV-casa was equidistant with mamastrovirus and avastrovirus in both the RdRP (<20%) and capsid (<11%) regions (see Table S7 in the supplemental material). Its basal position makes it difficult to suggest either birds or mammals as likely hosts.

To classify AstV-casa, we compared the amino acid sequence identities throughout its coding region using sliding-window analyses. When AstV-casa was compared to known astroviruses, it showed greater divergence from them than the intragenus variations within mamastroviruses and avastroviruses (see Fig. S5 in the supplemental material). Together with the results of phylogenetic analysis and pairwise distance analysis (see Table S7 in the supplemental material), the sequence identities suggested that AstV-casa reflects the existence of a third Astroviridae genus, provisionally named Casastrovirus.

Picalivirus, a highly divergent virus in the picorna-like virus superfamily. Positive-strand RNA virus-related sequences showing even lower-level sequence similarities to both Picornaviridae and Caliciviridae families were abundant. We acquired the full-length genome of one such virus from Nepalese sewage (GenBank accession number JQ898334-6), and provisionally named it picalivirus (picornavirus- and calicivirus-related virus [PicaV]). The picalivirus RNA genome consisted of 8,996 nucleotides beginning with a 2,025-amino-acid-long ORF encoding a nonstructural polyprotein, followed by a second ORF encoding a capsid protein and a 3’ UTR ending in a poly(A) tail (Fig. 5). As characterized by 5’ RACE amplification, the 5’ end of the picalivirus genome contained a very short 5’ UTR in a manner reminiscent of caliciviruses. The genomic organization of picalivirus resembled most closely that of the Lagovirus and Sapovirus genera in the Caliciviridae family (15), with their very short 5’ UTR and a long nonstructural polyprotein containing the helicase and RdRP regions, followed by a second ORF encoding the capsid protein (Fig. 5A; see FIG 3 Genome organization and phylogenetic relationships of new kobovirus and salivirus. (A) Genome organization of the kobovirus sewage Kathmandu (KoV-SewKTM) and salivirus sewage Bangkok (SalIV-SewBKK). (B) Phylogenetic analysis of the two novel picornaviruses with known members of the Kobovirus and Salivirus genera in the family Picornaviridae, using translated protein sequence alignments of P1 capsid and 3D polymerase regions and the maximum likelihood method. Aligned regions are shown by black bars.
Amino acid sequence identities in the helicase and RdRP regions of picalivirus and members of the Picornavirales order (families Picornaviridae, Iflaviridae, Dicistroviridae, Marnaviridae, and Secoviridae) (61) and Caliciviridae (133) were calculated through pairwise analysis (see Tables S8A and B in the supplemental material). The most closely related genome was that of a calicivirus with 25% and 20% amino acid sequence identities in the helicase and RdRP regions, respectively. Except for several short conserved motifs, including the helicase nucleotide-binding motif [GXXGXGK(T/S)] and several RdRP motifs (KDEX, YGDD, and FLRR) (43, 57, 61), the picalivirus polyprotein was highly divergent from the other RNA viruses (Fig. 5A and C; see Tables S8A and B in the supplemental material). Conserved motifs were slightly closer to those of the Caliciviridae than to other members of the Picornavirales order (Fig. 5C). Protein structure prediction (86) showed that the structures closest to the PicaV RdRP were the poliovirus three-dimensional (3D) RdRP (E value, 4e−103) and Norwalk virus RdRP (E value, 7e−96), while the structures closest to the PicaV capsid were the cripavirus capsid (E value, 7e−15) and the cardiovirus capsid (E value, 2e−9). Phyllogenetic analyses using the RdRP and the helicase regions placed picalivirus close to the root of the Picornavirales order, indicating the early divergence of this viral group and suggesting picalivirus as a candidate prototype of a novel viral family provisionally named Picaliviridae (Fig. 5B). Previous genetic analyses have described a large grouping of positive-strand RNA viruses which includes the Picornavirales order (58, 61), the Caliciviridae, plus other positive single-stranded RNA viral families into a picorna-like virus superfamily (58). The picorna-like virus superfamily, due to its wide distribution in all forms of nucleated cells, is theorized to have emerged early in the evolution of eukaryotes (58). A subset of the viruses in the picorna-like virus superfamily encodes a superfamily 3 helicase, including all members of the Picornavirales order plus the Caliciviridae (58). Picalivirus also contains a homolog of this enzyme. On the basis of its high level of divergence relative to other members of the picorna-like virus superfamily, picalivirus may therefore be a prototype of a new viral family related to the Picornavirales order.

Fig. S6 in the supplemental material for a comparison of ORF organizations.

Amino acid sequence identities in the helicase and RdRP regions of picalivirus and members of the Picornavirales order (families Picornaviridae, Iflaviridae, Dicistroviridae, Marnaviridae, and Secoviridae) (61) and Caliciviridae (133) were calculated through pairwise analysis (see Tables S8A and B in the supplemental material). The most closely related genome was that of a calicivirus with 25% and 20% amino acid sequence identities in the helicase and RdRP regions, respectively. Except for several short conserved motifs, including the helicase nucleotide-binding motif [GXXGXGK(T/S)] and several RdRP motifs (KDEX, YGDD, and FLRR) (43, 57, 61), the picalivirus polyprotein was highly divergent from the other RNA viruses (Fig. 5A and C; see Tables S8A and B in the supplemental material). Conserved motifs were slightly closer to those of the Caliciviridae than to other members of the Picornavirales order (Fig. 5C). Protein structure prediction (86) showed that the structures closest to the PicaV RdRP were the poliovirus three-dimensional (3D) RdRP (E value, 4e−103) and Norwalk virus RdRP (E value, 7e−96), while the structures closest to the PicaV capsid were the cripavirus capsid (E value, 7e−15) and the cardiovirus capsid (E value, 2e−9). Phyllogenetic analyses using the RdRP and the helicase regions placed picalivirus close to the root of the Picornavirales order, indicating the early divergence of this viral group and suggesting picalivirus as a candidate prototype of a novel viral family provisionally named Picaliviridae (Fig. 5B). Previous genetic analyses have described a large grouping of positive-strand RNA viruses which includes the Picornavirales order (58, 61), the Caliciviridae, plus other positive single-stranded RNA viral families into a picorna-like virus superfamily (58). The picorna-like virus superfamily, due to its wide distribution in all forms of nucleated cells, is theorized to have emerged early in the evolution of eukaryotes (58). A subset of the viruses in the picorna-like virus superfamily encodes a superfamily 3 helicase, including all members of the Picornavirales order plus the Caliciviridae (58). Picalivirus also contains a homolog of this enzyme. On the basis of its high level of divergence relative to other members of the picorna-like virus superfamily, picalivirus may therefore be a prototype of a new viral family related to the Picornavirales order.

FIG 4 Genome organization and phylogenetic relationship of the casa astrovirus (AstV-casa). (A) Genome organization of the casa astrovirus compared with human and turkey astroviruses; (B) phylogenetic analysis of the translated partial RdRP and the full capsid protein sequences of AstV-casa and representatives of the Mamastrovirus and Avastrovirus of the Astroviridae family using the maximum likelihood method.
High level of picalivirus diversity and geographic distribution. Metagenomics-derived sequences were analyzed for the presence of sequences closely related to the first picalivirus genome. Two other picalivirus genomes were identified and partially sequenced using RACE amplifications (GenBank accession numbers JQ898335 and JQ898336). Phylogenetic analysis of their RdRP regions showed that they shared a common root with the original picalivirus genome (Fig. 5B) with amino acid sequence identities of 30% to 34% (see Table S8 in the supplemental material), suggesting that these picaliviruses constitute prototypes of three Picaliviridae genera (using as criteria the genetic distances between genera within the Picornaviridae and Caliciviridae families) (52, 60). Picaliviruses were found in both San Francisco and Kathmandu, suggesting a wide geographic distribution.

Secalivirus, a virus distantly related to Caliciviridae. The family Caliciviridae consists of five genera, Norovirus, Sapovirus, Vesivirus, Lagovirus, and Nebovirus, as well as a recently described group, recovirus (14, 15, 20, 75). Caliciviruses are known to infect only vertebrate hosts, causing a wide range of diseases, including respiratory infections, vesicular lesions, gastroenteritis, and hemorrhagic disease (14, 16). The partial genome of a highly divergent calicivirus was acquired from the Nepal sewage sample and preliminarily named secalivirus (sewage-associated calici-like virus [SecaliV]; GenBank accession number JQ898339). The partial genome consisted of 4,068 nucleotides, including a partial capsid gene, followed by two ORFs of unknown function, and a 553-nt-long 3’ UTR ending with a poly(A) tail (Fig. 6). Pairwise identity analysis suggested that the SecaliV capsid protein was highly divergent from the capsid proteins of the Caliciviridae (10.5 to 17.5% identity) (see Table S9 and Fig. S7 in the supplemental material).

The capsid of SecaliV did contain a conserved capsid motif (calicivirus coat protein pfam00915), with certain amino acid positions being more like those of caliciviruses than other positive
**ssRNA viruses** (see Fig. S7 in the supplemental material). Similarly, protein structure prediction (86) showed that the structure closest to the SecaliV capsid was the calicivirus coat protein (E value, $2 \times 10^{-57}$). Comparing the capsid protein of SecaliV, caliciviruses, and other members of the *Picornavirales*, the phylogenetic analysis also placed SecaliV near the root of the *Caliciviridae* (Fig. 6C). Lastly, sliding-window analysis showed that SecaliV shared a degree of divergence from known caliciviruses similar to the intergenus variations of known calicivirus genera (Fig. 6B). Collectively, these results indicate that secaliavirus may be tentatively classified as the prototype of a new genus in the *Caliciviridae*.

**A highly divergent HEV-like genome.** A sequence from Nepal shared low-level similarity to HEV in the family *Hepeviridae*. This genome was tentatively named hepevirus-like virus (hepelivirus [HepeV]; GenBank accession number JQ898340). The partial genome of hepelivirus acquired was 2,721 nt long, consisted of half of ORF1 (RdRP), a complete ORF2 (capsid), as well as the 3′ UTR (Fig. 7). Nepal was the only sampled region where human HEV sequences were detected (i.e., >90% protein identity) (Table 1). HEV can cause self-limited or fulminant hepatitis in humans, as well as infect a range of mammalian species as a zoonotic agent. Recently, other HEV-related viral species have been characterized in rodent, avian, bat, and fish host species (2, 5, 18, 38, 52).

Pairwise distance analysis of the RdRP amino acid suggested that HepeV was equidistant to HEV, rodent HEV, avian HEV, and cutthroat trout virus (~20% identity), while showing weaker identity to members of other ssRNA families (7% to 12%) (see Table S10 in the supplemental material). HepeV contained conserved features of RdRP of the *Hepeviridae*. Like other HEVs, HepeV contained FKGDDS (underscore highlights a conserved RdRP motif) (5, 28) and DVXR motifs, compared to the XY GDDX and FL(K/R)R motifs common to members of the *Picornavirales* (see Fig. S8 in the supplemental material) (43, 57, 61).

**HepeV, like other genomes in *Hepeviridae*, lacked the KDEX motif that is consistently found among members of the order *Picornavirales* (see Fig. S8 in the supplemental material).** Phylogenetic analysis of the RdRP region also showed HepeV’s closer relationship to the family *Hepeviridae*, placing HepeV at the root of HEV-related viruses and suggesting an early divergence from other known *Hepeviridae* species (Fig. 7).

All three analyses, i.e., conserved motifs, pairwise distances, and phylogenetics, suggested that HepeV belongs to the *Hepeviridae* family. When HepeV was compared to other members of the *Hepeviridae*, it showed divergence greater than their interspecies variations (Fig. 7B). HepeV may therefore represent a prototype for a new genus within the family *Hepeviridae*.

**Genomes of other highly divergent ssRNA viruses.** We also characterized the RdRP regions together with their 3′ ends of three other highly divergent viruses, provisionally called nedicistrovirus (Nepal sewage dicistro-like virus [NediV]; GenBank accession numbers JQ898341, JQ898344, and JQ898345, respectively). NediV contained conserved features of RdRP of the *Hepeviridae*. Like other HEVs, NediV contained FKGDDS (underscore highlights a conserved RdRP motif) (5, 28) and DVXR motifs, compared to the XY GDDX and FL(K/R)R motifs common to members of the *Picornavirales* (see Fig. S8 in the supplemental material) (43, 57, 61).
the RdRP, followed by the capsid protein in the same reading frame (see Fig. S9 in the supplemental material). NiflaV (partial genome of 1,614 nt) resembled iflaviruses in terms of a polyprotein containing RdRP, without a capsid-encoding gene at the 3’ end of the genome. Phylogenetic analysis of the RdRP region suggested that CadiV and NiflaV branched out near the root of the tree, suggesting an early divergence from the picorna-like virus superfamily (Fig. 8).

NediV, CadiV, and NiflaV are three examples of divergent ssRNA viruses among many likely present in the sewage viromes (Fig. 2). Given the number of sequences showing low-level similarities to known viruses, we postulate that many more divergent ssRNA genomes remain uncharacterized in these sewage samples.

**Genomes distantly related to plant geminiviruses.** Sewage viromes contained plant viral sequences with various degrees of divergence from known plant pathogens, especially from the single-stranded DNA (ssDNA) family Geminiviridae. The family Geminiviridae contains four genera, Begomovirus, Curtovirus, Topocuvirus, and Mastrevirus, which are responsible for various types of plant diseases (21, 23, 52). Geminiviruses often limit the production of tomato, pepper, squash, melon, and cotton in the subtropics and tropics (67, 80), causing famines in the developing world (62). We describe three highly divergent geminivirus-like genomes (GenBank accession number JQ898331-3).

Baminivirus (Bangkok gemini-like virus [BamiV]), niminivirus (Nigeria gemini-like virus [NimiV]), and nepavirus (Nepal gemini-like virus virus [NepaV]) genomes consisted of circular DNA 2.3 to 2.8 kb long. Similar to other geminiviruses, these genomes contained two major ORFs encoding Rep and capsid proteins in an opposite orientation (Fig. 9A). Notably, BamiV and NimiV also contained a stem-loop with the 15-nt sequence CTATTATAACATTGC. Similar to geminiviruses, RCR motifs 1, 2, 3 and Walker A and B motifs were identified in the Rep protein of the three gemini-like viruses (Arguello-Astorga G., personal communication). ORFs related to movement protein (MP) were also identified in BamiV and NimiV, with ~35% protein identities to an array of geminivirus MP, suggesting that BamiV and NimiV genomes may be monopartite.

Phylogenetic analysis showed that BamiV, NimiV, and NepaV, while highly divergent from known geminiviruses, still clustered in the same clade as other ssDNA viral families (Fig. 9B). Baminivirus and niminivirus Rep proteins shared less than 30% pairwise amino acid sequence identities with other geminiviruses (see Table S12 in the supplemental material). Pairwise distance analysis of BamiV and NimiV with the Rep protein of other circular ssDNA genomes showed >19% Rep protein identities to Rep proteins in the Geminiviridae versus <15% identities to those in the Circoviridae and Nanoviridae (see Table S12 in the supplemental material). Sliding-window analysis of the replication gene amino acid sequence alignment showed that the identities between baminivirus and different geminiviruses were greater than the intragenus variations of geminiviruses (Fig. 9C), suggesting that baminivirus and niminivirus may be prototypes for two new genera within the Geminiviridae. Formal classification of these viruses in the Geminiviridae will require particle structure analysis and confirmation of their ability to infect plants.

Nepavirus lacked the exact stem-loop nonanucleotide signature of geminiviruses; however, it was still more closely related to Geminiviridae than to other known ssDNA viral families in pairwise distance analysis (>11 to 19% Rep protein identities to the Rep proteins of the Geminiviridae versus <7 to 12% identities to the Rep of a circovirus and nanovirus (see Table S12 in the supplemental material). Nepavirus is therefore related to the Gemini-
viridae family, but due to its sequence divergence and lack of conserved nonanucleotide, it may represent a new viral family.

Inference of cellular hosts using nucleotide composition analysis. We previously showed that ssRNA viruses from vertebrates, invertebrates, and plants could be broadly differentiated on the basis of a discriminant analysis of their di- and trinucleotide composition (46, 84). The complete and partial genome sequences generated here were therefore analyzed using nucleotide composition analysis (Fig. 10). The kobuvirus (KoV-SewKTM), salivirus (SaliV-SewBKK), and sapovirus (SaV-SewSFO) showed a nucleotide composition consistent with their expected vertebrate origins based on phylogenetic affinities described in previous sections (Fig. 3 and 4; see Fig. S4 in the supplemental material). NCA also inferred a vertebrate host for the new astrovirus AstV-casa.

The full or partial genomes of picaliviruses (PicaVs A, B, and C), secalivirus (SecaliV), hepelivirus (HepeV), and cadicistrovirus (CadiV) showed nucleotide composition properties comparable to those of invertebrate (insect/nematode)-infecting viruses. Currently, all known members of the Hepeviridae and Caliciviridae infect vertebrates. If the NCA predictions of invertebrate hosts for hepelivirus and secalivirus are correct, these two deep clades may represent insect-infecting viruses that diverged early in the evolution of Hepeviridae and Caliciviridae, respectively, expanding the host range for these two families. More experimental data addressing the tropism of these viruses are required to confirm this prediction.

The niflavirus (NiflaV) clustered with plant viruses, whereas the nedicistrovirus (NediV) did not group with any of these three kingdoms of eukaryotic hosts. Further discriminant analysis using more nucleotide composition components yielded the same inferred hosts for all viruses and a vertebrate host for NediV (data not shown).

DISCUSSION
Several trends were evident on the basis of our metagenomic analyses of viruses in untreated sewage from four countries. Each sample contained distinct subsets of human viral pathogens (Table 1). Numerous novel viral genotypes, species, genera, and, possibly, families of nonenveloped eukaryotic RNA and DNA viruses were identified. Sewage therefore harbors a very high viral diversity of known and previously uncharacterized viruses that allows for...
large-scale viral discovery and for monitoring of the presence of pathogens from humans and other cellular hosts.

While a large body of literature analyzing wastewater effluents for the presence of a specific virus(es) using PCR exists, unbiased studies of viruses in wastewaters using metagenomics approaches remain few. A metagenomics analysis of reclaimed water (for non-potable public uses) using pyrosequencing of viral particle-associated DNA indicated 98% prokaryotic versus 2% eukaryotic viral sequences, while the viral RNA population showed a more equally mixed distribution of viruses from a variety of inferred hosts more akin to that reported here (82). A metagenomics analysis of DNA viruses purified from activated sludge (a product of wastewater treatment) also showed a predominance (95%) of prokaryotic DNA viruses (76). A pyrosequencing analysis of the DNA viruses in the influent, activated sludge, effluent, and anaerobic digester of a wastewater treatment plant showed >90% of viral reads to be of likely bacterial origin (89). Lastly, a recent metagenomic analysis using pyrosequencing of DNA and RNA in viral particles from untreated wastewater from Pittsburgh, PA, Barcelona, Spain, and Addis Ababa, Ethiopia, found >80% of virus-like sequences likely originating from prokaryotes (11). The last study also revealed a great diversity of DNA and RNA viruses in which 85% of the eukaryotic virus-like sequences had a likely plant origin and generated the full viral genome of a phage-like genome previously thought to be a eukaryotic virus called non-A non-B hepatitis virus (11).

The sewage samples analyzed here were also rich in plant viruses, possibly reflecting the diversity of local plants as well as those plants consumed by local residents and animals. Some plant viruses (pepper mild mottle virus) are known to pass through the human gastrointestinal tract and remain infectious (99). Sewage may therefore provide another means for the dissemination of plant viruses if used untreated as fertilizer. Metagenomics studies of DNA viruses in insect vectors (whiteflies) also showed the majority of virus-like sequences to be closely related to known plant geminiviruses, including novel begomoviruses (68). Mosquito viromes also contained sequences related to Circoviridae and Geminiviridae (72). Sequences with best BLASTx E score to human viruses showed higher percentage identities when derived from U.S. sewage than from that of other countries (average of 84% versus 72 to 74%, respectively). Such a result may reflect the greater contribution to GenBank of viral genomes from developed relative to more resource-constrained countries. The ease of recognition of diver-

FIG 9 (A) Genome organization of the novel geminivirus-related viruses (baminivirus, niminivirus, and nepavirus) and two geminiviruses (reference species of begomoviruses and mastreviruses). Baminivirus and niminivirus both contained the nonanucleotide TAATATTAC within a stem-loop. (B) Phylogenetic analysis of the translated Rep protein sequences among niminivirus, baminivirus, and nepavirus members and representatives of Geminiviridae and other ssDNA viruses with circular ssDNA genomes using the maximum likelihood method. (C) Sequence identity comparisons of baminivirus and geminivirus diversity through sliding-window analysis of pairwise translated protein p distances in the Rep gene alignment. Niminivirus showed a similar result in sliding-window analysis (data not shown).
ent viruses, in the form of better E scores and higher percent identity, may therefore be improved as further studies contribute more viral genomes from less exhaustively sampled geographic regions.

The latest (ninth) release of the International Committee on Taxonomy of Viruses reports a total of 86 recognized viral families infecting prokaryotes and eukaryotes, plus other unassigned genera and numerous unclassified sequences (52). In this report, we identified, in highly variable proportions, sequences related to members of 29 eukaryotic viral families (BLASTx E score, < 10^-4). Specific extension and genome sequencing of selected viral sequences led to the genetic characterization of multiple novel viruses, broadening the diversity of the known virosphere. The cellular hosts of many sewage-derived viruses remain unknown. On the basis of their genetic similarities to viruses with known tropism or their nucleotide composition, broad categories of hosts such as mammals, vertebrates, insects/nematodes, or plants may be inferred. Inoculation of a complex viral mixture into model animals or plants may help to determine the host range of such highly divergent viruses. The addition of highly divergent viral genomes to public databases will also facilitate the identification of previously unrecognized viruses distantly related to these new genomes. It is also conceivable that the novel vertebrate viruses in the Picornaviridae, Astroviridae, and Caliciviridae families are of human origin. Knowledge of their genomes will facilitate further studies of their tropism, seroprevalence, and disease associations, as well as improve the design of consensus PCR-based assays for further viral discovery.

Sewage contains the fecal input of large numbers of local residents, some of which will be shedding diarrhea-causing viruses. Future metagenomic efforts may be conducted on larger scales, at greater frequencies, and with greater sequencing depths or targetting specific viral populations (such as polioviruses [37]) to expand the long tradition of analyzing sewage for enteric viruses. Metagenomic analysis combined with genomic characterization of viruses in sewage also has the potential to help monitor seasonal trends of enteric infections, better detect epidemic outbreaks, measure the extinction of wild-type polioviruses, as well as assist in optimizing vaccination strategies by characterizing recently replicated viral genotypes.

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