

UC Davis

UC Davis Previously Published Works

Title

Enhanced arbovirus surveillance with deep sequencing: Identification of novel rhabdoviruses and bunyaviruses in Australian mosquitoes

Permalink

<https://escholarship.org/uc/item/3k8570q3>

Authors

Coffey, Lark L
Page, Brady L
Greninger, Alexander L
et al.

Publication Date

2014

DOI

10.1016/j.virol.2013.09.026

Copyright Information

This work is made available under the terms of a Creative Commons Attribution-NonCommercial-NoDerivatives License, available at <https://creativecommons.org/licenses/by-nc-nd/4.0/>

Peer reviewed

Published in final edited form as:

Virology. 2014 January 5; 448: . doi:10.1016/j.virol.2013.09.026.

Enhanced arbovirus surveillance with deep sequencing: identification of novel rhabdoviruses and bunyaviruses in Australian mosquitoes

Lark L. Coffey^{1,2}, Brady L. Page^{1,2}, Alexander L. Greninger², Belinda L. Herring³, Richard C. Russell^{4,5}, Stephen L. Doggett⁵, John Haniotis⁵, Chunlin Wang⁶, Xutao Deng^{1,2}, and Eric L. Delwart^{1,2,#}

¹Blood Systems Research Institute, University of California San Francisco, San Francisco, California, USA

²Department of Laboratory Medicine, University of California San Francisco, San Francisco, California, USA

³Griffith University, Gold Coast Campus, Queensland, Australia

⁴Sydney Medical School, University of Sydney

⁵Centre for Infectious Diseases and Microbiology, Westmead Hospital, New South Wales, Australia

⁶Genome Technology Center, Stanford University School of Medicine, Stanford, California, USA

Abstract

Viral metagenomics characterizes known and identifies unknown viruses based on sequence similarities to any previously sequenced viral genomes. A metagenomics approach was used to identify virus sequences in Australian mosquitoes causing cytopathic effects in inoculated mammalian cell cultures. Sequence comparisons revealed strains of Liao Ning virus (*Reovirus*, *Seadornavirus*), previously detected only in China, livestock-infecting Stretch Lagoon virus (*Reovirus*, *Orbivirus*), two novel dimarhabdoviruses, named Beaumont and North Creek viruses, and two novel orthobunyaviruses, named Murrumbidgee and Salt Ash viruses. The novel virus proteomes diverged by ~50% relative to their closest previously genetically characterized viral relatives. Deep sequencing also generated genomes of Warrego and Wallal viruses, orbiviruses linked to kangaroo blindness, whose genomes had not been fully characterized. This study highlights viral metagenomics in concert with traditional arbovirus surveillance to characterize known and new arboviruses in field-collected mosquitoes. Follow-up epidemiological studies are required to determine whether the novel viruses infect humans.

© 2013 Elsevier Inc. All rights reserved.

#Corresponding author: Eric Delwart, Blood Systems Research Institute, University of California, San Francisco, Department of Laboratory Medicine, 270 Masonic Avenue, San Francisco, California, 94118, delwarte@medicine.ucsf.edu.

Competing Interests

None.

Authors' contributions

LLC conducted project, interpreted results and wrote the paper. BLP constructed libraries for deep sequencing. ALG, CW and XD performed bioinformatics and contig assembly. BLH, RCR, SLD, and JH collected mosquito pools, screened samples for known arboviruses, and provided samples to LLC and ELD for deep sequencing. ELD designed the project and edited the manuscript.

Publisher's Disclaimer: This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final citable form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

Keywords

virus discovery; deep sequencing; arbovirus surveillance; bunyavirus; rhabdovirus; reovirus; novel virus; mosquito; Australia

Background

Numerous arboviruses that cause significant human disease are endemic to Australia. To address the risks posed by arboviruses to human and veterinary health in Australia, annual arbovirus surveillance is conducted by a network of regional diagnostic laboratories that trap mosquitoes and other insects and test them for known viruses. Viruses tested include members of the *Flaviviridae* and *Togaviridae* families-Barmah Forest virus (BFV), Edge Hill virus (EHV), Kunjin virus (KUNV), Kokobera virus (KOKV), Murray Valley encephalitis virus (MVEV), Ross River virus (RRV), Sindbis virus (SINV), and Stratford virus (STRV)-detected using antigenic tests on inoculated mosquito and vertebrate cell monocultures showing cytopathic effects (CPE) (1). Many of these arboviruses cause human disease, ranging from mild febrile illness to encephalitis and death. Identifying these circulating viruses, as well as other novel viruses, in anthropophilic mosquitoes is therefore important to implement strategies to detect and mitigate arbovirus transmission to humans and other animals. However, despite extensive diagnostic testing, some CPE-causing viral isolates cannot be identified using current assays specific for known arboviruses.

An increasing number of viruses are being discovered in arthropods (reviewed in (2)). Viral metagenomics, so-called 'deep sequencing', has been used in mosquitoes to survey viral diversity (3) and to sequence arboviruses previously broadly classified using antigenic cross-reactivity (4–9). However, its use for identifying viruses in surveillance settings has so far been limited to detection of an insect-restricted densovirus in Chinese mosquitoes (10).

In this study, deep sequencing was used to non-specifically amplify and sequence enriched viral nucleic acids from CPE-positive supernatants in which arboviruses were not identified by conventional antigenic tests. Viral sequences were identified by similarities of their *in silico* translated protein sequences with all previously sequenced viral proteins. We detected sequences of known reoviruses including Liao Ning virus (LNV), Stretch Lagoon orbivirus (SLOV), Wallal virus (WALV), and Warrego virus (WARV), as well as two novel rhabdoviruses, and two novel bunyaviruses. RRV, EHV and KOKV sequences were also identified in some pools. Detection of novel and known arboviruses not previously recognized in Australia highlights the use of complementing conventional arbovirus surveillance with viral metagenomics approaches.

Materials & Methods

Insect Collection and Virus Culture

Mosquitoes and midges were collected in dry-ice baited traps at various sites in NSW, Australia between 1992 and 2010 (Table 1). As part of routine arbovirus surveillance in the region, individual mosquitoes and midges were identified using morphological criteria (11) and pooled into groups of up to 50 insects. Pools were mechanically homogenized with glass beads and inoculated onto porcine stable equine kidney (PSEK) or baby hamster kidney (BHK) cells that are highly susceptible to flavivirus and alphavirus infection, respectively, and monitored by microscopic examination for CPE, including cell rounding and death characteristic of arbovirus infections. CPE positive supernatants were transferred to new cultures of PSEK, BHK or *Aedes albopictus* larval (C6/36) cells for a total of 2 to 6 passages (only sample 934 represented unpassaged mosquito homogenate) until a strong CPE was

elicited, suggesting virus infection. The lowest remaining available passage was used in this study. Virus identification using supernatants from passages was attempted by enzyme immunoassay with a panel of monoclonal antibodies to Australian arboviruses including BFV, EHV, KOKV, KUNV, MVEV, RRV, SINV, and STRV. Prior to 1994, viruses were identified by micro-neutralization tests using specific antisera to the same viruses as well as Alfuy (ALFV), Gan Gan (GGV), and Trubanaman viruses (TRUV) (12).

Viral Particle Purification

Supernatants from low passage insect pools were clarified from cell debris by 12,000 g centrifugation for 2 minutes and then passed through a 0.45- μ m filter (Millipore) to remove large particulates including bacteria. The filtrate was treated with a nuclease cocktail consisting of DNases (Turbo DNase from Ambion, Baseline-Zero from Epicentre and Benzonase from Novagen) and with RNase (Fermentas) to digest host nucleic acids and non-encapsulated viral nucleic acids. Nucleic acids protected from nuclease digestion were then extracted using the QIAamp viral RNA mini kit (Qiagen) according to manufacturer's recommendations with a five minutes incubation step at room temperature prior to elution from the column.

Sequence-Independent Nucleic Acid Amplification for Deep Sequencing

Deep sequencing was performed using Roche 454 and Illumina MiSeq platforms. For Roche 454 libraries, RNA-only and DNA-plus-RNA sequence-independent amplifications were combined before sequencing. For RNA-only amplification, 10 μ l of extracted nucleic acids were treated with DNase (Ambion) and were used as a template for cDNA synthesis using Superscript III reverse transcriptase (Invitrogen). cDNAs were primed with an arbitrary sequence and eight fixed nucleotides at the 3' end and six fixed nucleotides at the 5' end that served as a barcode (13). For the DNA-plus-RNA amplification, the DNase step prior to reverse transcription was omitted. After reverse transcription, cDNAs were heat denatured and the primers were allowed to re-anneal for a single round of second strand DNA synthesis using Klenow DNA polymerase (New England Biolabs). Thirty-five PCR cycles were performed on double stranded cDNAs using primers consisting of arbitrary sequences also used as molecular tags. Each PCR reaction was performed in duplicate to increase sampling of viral nucleic acids, and then the four resulting PCR products were pooled and purified using the QIAquick PCR purification kit (Qiagen). Randomly amplified nucleic acids from each insect pool derived culture supernatant were normalized using Nanodrop (Thermo Scientific) and mixed at equimolar ratios to obtain 2.6 μ g of DNA/sample. Amplicons from 300–1000 base pairs in this mixture were size selected by extraction from a 2% agarose gel and purified using the QIAquick gel extraction kit (Qiagen). PCR ends were polished using T4 polynucleotide kinase (New England Biolabs) and Roche 454 adaptors were ligated to the ends. Small fragments were removed using AmpPure XP beads (Agencourt Biosciences) according to the manufacturer's protocol (GS FLX Titanium Library Preparation Kit, Roche).

To characterize sequences in selected samples more deeply to fill in gaps in genomes, libraries were also made and sequenced using MiSeq (Illumina). Nucleic acids were extracted from the same filtered and nuclease treated culture supernatants and the ScriptSeq v2 RNA-Seq Library Preparation Kit (Epicentre) was used following the manufacturer's protocol with 1X purifications performed using AmpPure XP beads (Agencourt Biosciences). Up to 35 PCR cycles were used when amplicons were not observed in 1% agarose gels after fewer cycles. Size distributions of libraries were assessed using an Agilent 2100 Bioanalyzer and concentrations of cluster-competent DNAs were determined using the SYBR FAST LightCycler 480 qPCR Kit (Kapa) according to manufacturer's instructions. Libraries were diluted to 7 pM for sequencing on the MiSeq.

For both sequencing platforms, barcode sequences were included in primers during non-specific amplifications to multiplex multiple insect pool derived culture supernatants in the same run.

Deep Sequencing Data Processing and Sequence Assembly

Sequences were binned according to barcode and then the barcode and primer were removed. Clonal reads were eliminated and sequences were de novo assembled into contigs using Mira (Roche 454), SOAP (Illumina), PRICE (14) or manually using Sequencher (Gene Codes; where alignments were made when 90% of at least 10 overlapping base pair reads were identical). Assembled contigs and single sequences that did not collapse into contigs were then analyzed by tBLASTx, querying the translated nucleotide database using a translated nucleotide query. To measure sequence similarity, expectation (E) scores were used, with a cutoff of $E < 10^{-5}$. E scores as high as 10^{-3} are used for virus discovery (13); here 10^{-5} was selected to increase stringency but still allow for detection of divergent sequences. Contigs and singlets were classified as eukaryotic, bacterial, phage, or viral based on best E score matches; sequences with tBLASTx E scores $> 10^{-5}$ were deemed unclassifiable. Contigs and singlets with best matches to annotated viruses in GenBank were then sorted by virus taxon.

Genome Acquisitions by Reverse Transcription-Polymerase Chain Reactions

Contigs and singlets showing strong tBLASTx E scores to viruses in GenBank were joined together by RT-PCRs using specific primers designed from deep sequencing reads and directly Sanger sequenced. Areas where deep sequencing coverage was lower than 5X or regions with sequencing discrepancies were also re-sequenced using Sanger technology directly on PCR products. The Titan One Tube RT-PCR kit (Roche) was used according to the manufacturer's protocol and cycling conditions were: cDNA synthesis at 50°C for 30 m followed by denaturation at 94°C for 2 minutes (m), then 35 cycles of: 94°C for 30 s, 58°C for 30 s, and 68°C for 2 m, followed by a final extension of 68°C for 10 m. Full genomes, when obtained, were assembled using Sequencher (Gene Codes).

Phylogenetic Analyses, Similarity and Identity Comparisons

Reference viral amino acid sequences representing bunyavirus, reovirus and rhabdovirus families were obtained from GenBank. Sequence alignments were performed using CLUSTALX (15) on the Mobylye portal website (<http://mobylye.pasteur.fr>). Phylogenies were generated using the maximum likelihood (Jones-Taylor-Thornton model) and neighbor-joining (maximum composite-likelihood) methods with MEGA software (16). Statistical significance of tree topologies was evaluated by 1000 bootstrap re-sampling iterations. Sequence identities between species were calculated using the SIAS server (<http://imed.med.ucm.es/Tools/sias.html>) and the equation: percent identity = $100 * (\text{identical positions} / \text{length of alignment})$ where gaps in alignments were included in analyses. Similarity plots, calculated in % identities, were generated by aligning nucleotide sequences and calculating scanning pairwise identities using a window size of 100 bp using mVISTA (<http://genome.lbl.gov/vista/mvista/submit.shtml>).

Results

Detection of Virus Genomes and Sequence Determinations

Virus genomes were detected by deep sequencing in 21 of the 47 culture supernatants from insect pools (Table 1, Table S1, only pools with identified virus genomes are listed; raw sequence reads are available upon request). Pools contained from 0 to 3 viruses. At least 17 pools contained sequences that matched with 90% nucleotide identity to known

arboviruses in GenBank. These viruses included EHV, KOKV, LNV, RRV, SLOV, WALV and WARV. Since these viral sequences were highly similar to published sequences, we consider that they represent strains of known viruses. Since only partial genome data was available in GenBank for WALV and WARV, we sequenced the genomes from 1 pool each for both viruses.

Five pools contained sequences with ~50% amino acid (aa) identity to different rhabdovirus or bunyavirus species in GenBank. Two of these pools (932 and 934) contained sequences that were ~99% identical, reflecting variants of the same viral species. Sequences from the other 3 pools were dissimilar. Complete genomes of 2 novel bunyaviruses were obtained. Partial genome sequences comprising 75% of the L gene for one virus and 90% of the complete genome of the other virus for 2 novel rhabdoviruses were obtained.

Novel Rhabdoviruses

Two mosquito pools, 6 and 954, contained genomes consistent with rhabdovirus genome organization (17). Sample 6 (GenBank accession number KF310911) was isolated from *Anopheles annulipes* in Griffith, NSW near Beaumont; we propose the name 'Beaumont virus' (BEAUV) for this rhabdovirus. Sample 954 (GenBank accession numbers KF360970-3) was isolated from *Culex sitiens* in Ballina, NSW near North Creek. We propose the name 'North Creek virus' (NORCV) for this rhabdovirus. BEAUV and NORCV both exhibit similar genome organization, genome length and %GC in the regions studied to representative rhabdovirus species (Table 2, Figure S2). A maximum likelihood phylogeny (Figure 1) was constructed using L gene, the RNA dependent RNA polymerase (RdRp), aa sequences of BEAUV and NORCV and other rhabdoviruses, including members of all *Rhabdoviridae* genera. GenBank sequences for selected viruses, including most Australian rhabdoviruses, were often short (133 aa, e.g.), and were therefore excluded from phylogenies. Despite the exclusion of these sequences from trees, BEAUV and NORCV shared <70% aa identity in the 133 aa region available for the Australian rhabdoviruses in GenBank, indicating that the novel viruses do not constitute known Australian rhabdovirus species that are poorly genetically characterized. Both novel viruses are located in the **dipteran-mammal associated rhabdovirus** (dimarhabdovirus) supergroup that includes several unassigned clades, tentative groups Tibrogargan (as well as Almpiwar, Hart Park, and Le Dantec that are not included in Figure 1 since no full L gene sequence data is available in GenBank), and two established genera, Vesiculovirus and Ephemerovirus. BEAUV and NORCV are positioned at two deeply rooted branches in the dimarhabdovirus supergroup and neither virus clusters with any other known viral species, including any rhabdoviruses previously identified in Australia. BEAUV and NORCV share 37% aa identity in the L gene, a level similar to their identities with representative species in other dimarhabdovirus clades (Table 3). Nucleotide similarity plots (Figure S2) show that both novel viruses share 65% L gene nucleotide (nt) identity with species from different dimarhabdovirus clades, supporting the placement of both viruses outside of established dimarhabdovirus clades. Compared to other species that cluster together (shaded boxes in Table 3), BEAUV and NORCV are less similar to each other (Table 3), supporting their placement on two separate deeply rooted dimarhabdovirus branches.

Novel Bunyaviruses

Three mosquito pools, 931, 932, and 934, contained genomes consistent with orthobunyavirus genome organization (18) including three RNA segments: a large (L) segment that encodes the RdRp, a medium (M) segment that encodes a polyprotein, and a small (S) segment that encodes a nucleocapsid protein (NP) and a non-structural protein (NSs). Samples 934 (GenBank accession numbers KF234253-5) and 932 contained sequences that were ~99% identical to each other. Both originated from *Anopheles*

annulipes mosquitoes from Griffith, NSW, near the Murrumbidgee River. We therefore propose the name ‘Murrumbidgee virus’ (MURBV) for the novel virus. Sample 931 (GenBank accession numbers KF234256-8) was isolated from a pool of *Aedes vigilax* from Port Stephens, NSW, near the town of Salt Ash. We propose the name Salt Ash virus (SASHV) for this virus. MURBV and SASHV both exhibit similar genome organization, gene length and %GC to other orthobunyavirus species (Table 2, Figure S3). Examining nucleotide similarity plots and segment-wide identity with representative orthobunyaviruses from different clades, SASHV exhibits 54–56% nt identity and MURBV shows 53–63% nt identity (L segment, Figure S3). The two viruses are more similar to each other (63% in L, 57% nt identity in M, 74% in S, Figure S3) than they are to other orthobunyaviruses. Considering aa identity, the two novel viruses exhibit 51% aa identity (L segment) and 33% (M) with representative members of the orthobunyavirus genus (Table 4A,B), consistent with similarities observed between orthobunyavirus species that belong to different antigenic groups (19). Phylogenetic comparisons based on maximum-likelihood algorithms of MURBV and SASHV with L (Figure 2A) and M (Figure 2B) segments of published bunyavirus genomes show that, despite their relatively low aa identity to each other (61% in the L segment and 52% in M), the two viruses group together in a deeply rooted branch that is distinct from any of the established orthobunyavirus groups (Bunyamwera, Wyeomyia, California Encephalitis, Simbu). The M segment phylogeny (Figure 2B) places MURBV and SASHV in a sister clade to the orthobunyaviruses and the L segment phylogeny (Figure 2A) positions the novel viruses within the orthobunyavirus clade. Neighbor joining algorithms and phylogenies based on S segments (data not shown) yielded similar tree topologies. Maximum likelihood trees containing all *Bunyavirus* genera (Supplemental Figure 1A,1B) also show that MURBV and SASHV represent a distinct and deeply rooted orthobunyavirus clade. Notably, MURBV and SASHV cluster separately from the Simbu group viruses, many of which are borne by midges in Australia.

Reoviruses

Four reovirus genomes were detected in mosquito pools. LNV (*Seadornavirus* genus) was present in 8 pools from multiple species, and SLOV, WALV and WARV (*Orbivirus* genus) were detected in 2 pools each (pools 4 and 929, 922 and 927, 948 and 949, respectively). Reovirus genomes consist of 10–12 double-stranded RNA segments. The same viral species in different pools shared 95% nt identity. While WALV and WARV have been detected in NSW previously, only a portion of one genome segment is available in GenBank. We therefore assembled complete sequences for each of the 10 segments for WALV (GenBank accession numbers KF23459-68) and for 9 of 10 segments for WARV (GenBank accession numbers KF310902-10). The genome organization, segment lengths and GC% for each segment are similar to related orbiviruses (Table 2). A phylogenetic tree based on segment 1, the RdRp gene (Figure 3) shows the genetic relatedness of genomes detected here to other reoviruses. Although all 4 detected viruses have been isolated from mosquitoes previously, LNV has only been reported in China. The 8 LNV isolates sequenced here were between ~1–4% different from each other (nt level) in the RdRp gene and were at least 5% different from previously published Chinese LNV (data not shown).

Togaviruses and Flaviviruses

In addition to our detection of novel viruses and known viruses not previously observed in NSW, deep sequencing also detected the *Togavirus* RRV in 10 pools and the flaviviruses KOKV and EHV in one pool each. Virus sequences from all of these pools shared ~95% aa identity to published sequences from Australian isolates in GenBank (data not shown). Based on this high identity, these sequences represent strains of known flaviviruses and togaviruses endemic to Australia that were undetected during antigen testing, possibly due to low viral loads.

Discussion

Summary

We used deep sequencing to detect viral sequences in mammalian cell culture supernatants from insect pools from NSW, Australia. This approach identified 4 novel virus genomes and 4 genomes of known viruses that had not been observed extensively in NSW, recognized in Australia, or fully sequenced. These findings highlight the utility of deep sequencing for identifying viruses that would otherwise be overlooked by conventional serologic testing and virus-specific RT-PCR, although until recently molecular viral detection was not standard in diagnostic platforms employed in NSW. Identification of these virus genomes in anthropophilic mosquitoes collected in NSW, coupled with their ability to replicate in mammalian cells, indicates that they may be capable of infecting humans or other vertebrates. The absence of viral sequences in 26 of the CPE-positive insect pools tested here indicates that some viral sequences went undetected, perhaps because of distant homology to viruses in the GenBank database, or that other microbes, such as bacteria or fungi, could have caused the CPE but were excluded from deep sequencing results via our selective filtration and bioinformatics steps.

Geographic Distributions of Australian Arboviruses

The locations of arbovirus isolations in this study, as well as areas where closely related bunyaviruses, rhabdoviruses, and reoviruses have been isolated previously, are shown in Figure 4. The alphavirus and flaviviruses RRV, EHV and KOKV are not shown since they exhibit diffuse continental distributions and are not in the same families as the viruses detected here. We detected LNV at multiple sites in NSW. To our knowledge, this is the first detection of this virus outside China (20, 21). Although both WALV and WARV have been isolated in NSW as recently as 1995–6 (22, 23), detection of these viruses in Griffith and Hawkesbury (near Sydney) extends their distributions farther south. None of the Almpiwar group Australian rhabdoviruses (Almpiwar (ALMV), Charleville (CHVV) and Humpty doo (HDOOV) viruses) have been detected in NSW; however, knowledge of their distributions relies on data based solely on single isolates from the Northern Territory and Queensland. The Australian orthobunyavirus AKAV exhibits a diffuse distribution along the northern and eastern coast and overlaps with GGV, including in the Port Stephens area where SASHV was recovered. BEAUV and MURBV were both isolated in Griffith, in a region where other orthobunyaviruses have not been previously isolated. NORCV was isolated in Ballina near Port Stephens at the northern coastal edge of NSW, outside the known distribution of Australian rhabdoviruses. SLOV has previously been isolated in Western Australia and the Northern Territory, and once in Queensland and Sydney. Here we detected SLOV for the first time in Griffith, NSW.

Novel Rhabdoviruses

Rhabdoviruses are single-stranded, negative-sense RNA enveloped viruses that comprise 6 recognized genera and more than 130 unassigned viruses with a worldwide distribution (17) that impose a significant cost to human and veterinary health and to the agriculture industry. In Australia, there are at least 12 known arthropod-borne rhabdoviruses, including Adelaide river, Kimberley and Berrimah viruses (genus *Ephemerovirus*) that circulate among livestock and dipterans, Parry Creek and Wongabel viruses (Hart Park group) that use mosquitoes and midges as vectors, ALMV, HDOOV, and CHVV (Almpiwar group), Tibrogargan (TIBV) and Coastal Plains (CPV) viruses (Tibrogargan group) that use lizards, mosquitoes and midges as hosts, as well as the mosquito-borne Oak Vale virus (OVRV) and midge-borne Ngaingan virus (unassigned groups) (reviewed in (17)). None of these Australian rhabdoviruses have been linked to human disease; this may reflect their inability

or lack of present ecological opportunities to infect humans or infrequent attempts at detection in patients.

Here, we describe two novel rhabdovirus genomes in mosquito pools, BEAUV and NORCV, which diverge by 60% at the amino acid level in the RdRp from other rhabdoviruses. Strong bootstrap support leads us to classify these viruses in 2 deeply rooted branches in the dimarhabdovirus supergroup. Although rhabdoviruses in the same genera and dimarhabdovirus subgroups occur across different continents, genetically related virus species share similar hosts (Figure 1). Most rhabdoviruses, lyssaviruses excluded, cycle between vertebrate and arthropod hosts (17). To date, rhabdoviruses from Australia all belong to the Tibrogargan, Hart Park and Ampliwar groups. Given that BEAUV and NORCV do not cluster with any of these groups, they may constitute 2 novel mosquito-borne Australian dimarhabdovirus clades.

Novel Bunyaviruses

Bunyaviruses comprise >350 known segmented negative-stranded RNA viruses that infect a variety of plants and animals sometimes causing hemorrhagic fever (18). Of the five known *Bunyaviridae* genera (*Hantavirus*, *Nairovirus*, *Orthobunyavirus*, *Phlebovirus* and *Tospovirus*), all but the hantaviruses are vectored by mosquitoes, midges, sandflies, ticks and thrips. The largest genus is the *Orthobunyavirus*, which contains >220 virus species historically differentiated by serologic relatedness (24). Although many orthobunyaviruses cause disease in humans and animals, molecular characterization has been limited to selected species (19, 24). In Australia, all known orthobunyaviruses are members of the Simbu group and include Aino (AINOV), Akabane (AKAV), Douglas (DOUV), Facey's Paddock (FPV), GGV, Peaton (PEAV), Tinaroo (TINV), TRUV, and Thimiri (THIV) viruses (25), as well as recently identified Leanyer virus (LEAV) that may represent a distinct genetic and antigenic complex (19). Here we characterize the genomes of two novel orthobunyaviruses, MURBV and SASHV, that form a deeply rooted clade separate from any of the established orthobunyavirus groups (Figures 2A,B, S1A,B), possibly reflecting a sixth group. Orthobunyaviruses show little geographic clustering (Figure 2A), excepting *Wyeomyia* group species that have been reported exclusively in South America. Here we identify a new viral clade in Australia comprising MURBV and SASHV that likely use mosquitoes as vectors.

Novel Detection of Reoviruses in Australia

The family *Reoviridae* contains 9 genera including the arthropod-borne seadornaviruses and orbiviruses. Seadornaviruses are transmitted by anopheline and culicine mosquitoes and their genomes consist of 12 enveloped double-stranded RNA segments. There are three species of seadornaviruses: Banna virus (BAV), Kadiporo virus (KDV) and LNV. BAV has been isolated from encephalitic humans (26, 27), pigs, cattle (28), and mosquitoes in China (29, 30), Indonesia (31) and Vietnam (32). Serosurveys of patients with encephalitis across China revealed a BAV seroprevalence of about 10% (33). Unlike BAV, KDV and LNV have not been associated with human disease, although LNV replicates in vertebrate cells and produces viremia and nasal hemorrhages in mice (34). LNV was first isolated in *Aedes dorsalis* mosquitoes from Liao Ning province in northeastern China in 2006 (34) and has subsequently been isolated from *Culex* spp. and *Ae. dorsalis* throughout northern China (20, 21, 35). Here we report the first detection of LNV in multiple pools of different species of mosquitoes and midges from NSW collected from 1995–2005 (Table 1) indicating that LNV is geographically diffuse in NSW, Australia, and may use multiple vector species there, similar to LNV circulation in China. Detection of LNV in pools of mosquitoes collected in 1995 indicates that the virus was present in Australia before it was recognized in China in 2006. No samples from China were ever present in our laboratories in Australia or the

United States, and the level of sequence divergence (ca. 5%) between Australian LNV and Chinese LNV from GenBank is higher than would be expected if the sequences detected here represent contamination by Chinese LNV. Furthermore, a negative control hamster kidney cell culture that was passaged with insect pool extracts did not contain any arbovirus genome segments (Table 1), confirming the lack of cross-contamination between virus isolates or passages.

Orbiviruses, a genus within the Reovirus family, consist of 22 recognized viral species and 13 unassigned viruses, each with 10 double stranded RNA segments (36). Many are transmitted by ticks, *Culicoides*, mosquitoes, and sand flies, and use cows, goats, sheep, equids, camelids, marsupials, sloths, bats, large carnivores, and humans as hosts (37). Among these, SLOV was first isolated in 2002 from a pool of *Culex annulirostris* mosquitoes collected in Kimberley, Western Australia, and a subsequent serologic survey revealed SLOV antibody in horses, goats and donkeys in the Northern Territory (37). The potential for human infection with SLOV is unknown, and SLOV has only once been isolated outside of Queensland and the Northern Territory, in Sydney (Figure 4) (38). Here we report the detection of SLOV genomes in two pools from Griffith, NSW: *Cx. annulirostris* collected in 2006 and a mixed pool of *Cx. australicus* and *Cx. molestus* collected in 1995. These detections extend the distribution of SLOV in NSW and also indicate that SLOV was probably circulating in mosquitoes in Australia for at least 6 years before it was first isolated in 2002 (39).

Midge- and mosquito-borne orbiviruses WALV and WARV are associated with epidemics of blindness in kangaroos and WALV-inoculated kangaroos developed chorioretinitis (23, 40, 41). Full genome sequences of these 2 viruses have not been previously published. WALV and WARV were first isolated in the 1970s in the Northern Territory from the two vector species implicated here, *Cx. annulirostris* and *An. annulipes* (41). Standfast et al. (41) also detected these two viruses in other insects, including *An. farauti* and various *Culicoides* spp., suggesting that WALV and WARV exhibit broad vector ranges. Nine park workers sampled after a 1994–5 outbreak of kangaroo blindness lacked neutralizing antibody to WALV (42); in the absence of widespread serosurveys or clinical surveillance of patients, the potential for either of these viruses to cause human infections and disease remains unknown.

Vector Use Patterns

Two of the viruses characterized here, BEAUV and MURBV, were isolated from freshwater *Anopheles annulipes* s.l. mosquitoes, members of the Australasian *Annulipes* complex that comprises at least 15 sibling species, some of which vector malaria and myxomatosis in rabbits (43). *An. annulipes* exhibits a diffuse distribution throughout Australia and predominantly lays its eggs in freshwater streams, marshes and lakes (44, 45). The arboviruses OVRV (*Rhabdoviridae*, unassigned), (46) RRV (*Togaviridae*, alphavirus) and the bunyavirus Mapputta virus (MAPV, unassigned) (41) have all been isolated from *An. annulipes*. SASHV, by contrast, was isolated from *Aedes vigilax*, a significant vector of RRV, BFV and GGV (47, 48), that is highly abundant in salt marshes (45). Blood meal identification studies show that *Ae. vigilax* exhibits broad host-feeding patterns, imbibing from dogs, birds, humans, horses and possums in Brisbane (49). *Culex sitiens*, a mosquito that oviposits in brackish or saline coastal lagoons (50), from which NORCV was isolated, has been implicated as a vector of numerous alphaviruses and flaviviruses in Australia, including RRV, BFV, SINV, MVEV, KUNV, KOKV, ALFV, and Japanese encephalitis virus (JEV) (51–53). The 4 newly genetically characterized viruses reported here therefore appear to use the same highly abundant vectors as other arboviruses known to pose significant threats to human health in Australia.

Future Directions and Use of Deep Sequencing in Arbovirus Surveillance

Serologic surveys using either neutralization of viral isolates or detection of antibody to viral antigens will be required to determine whether the 4 novel viruses identified here, as well as LNV and SLOV, infect humans or other animals in Australia. Serologic cross-reactivity of these novel viruses with lesser-described rhabdoviruses and bunyaviruses for which little or no sequence data is available (e.g. MAPV, GGV and TRUV,) while unlikely due to large expected genetic distances, may also help define their serologic relatedness to other members of the rhabdovirus and bunyavirus families.

Recent outbreaks of pathogenic arboviruses belonging to numerous viral families including the *Togaviridae* (chikungunya virus (CHIKV)) (54), *Bunyaviridae* (severe fever with thrombocytopenia syndrome virus (SFTSV)) (55), *Flaviviridae* (West Nile virus (WNV) (56), dengue virus (DENV)) (57), *Rhabdoviridae* (Bas-Congo virus (BASV)) (58), and *Reoviridae* (BANV) (26) attest to their potential to cause significant human disease. Arbovirus epidemics may be potentiated by changing viral ecology and point mutations that enhance transmission. Arbovirus surveillance therefore plays an important role as an early warning system to mitigate consequences of infections that cause human and animal disease. This study provides an important step by identifying circulating viruses of potential concern in Australian mosquitoes. Future studies are warranted to address the possibility of human or animal infection by the novel viruses detected here, as well as LNV. This study also highlights the use of deep sequencing for identifying arboviruses in mosquitoes. Most vector surveillance programs screen pools only for known endemic and enzootic viruses, precluding detection of novel or newly emerging viruses. Here we show that deep sequencing can identify novel viruses without prior knowledge of virus genomes present, making this approach a valuable alternative to designing and optimizing family- or genus-level degenerate primers for PCR-based detection. Deep sequencing is rapidly becoming cheaper and widespread, and barcoding many samples for pooling in a single run can further reduce cost without significantly compromising data quantity, providing a possible new tool for virus detection and characterization in arbovirus surveillance settings. The focus of this study was to detect viruses that caused CPE but which were not identified by conventional serology or antigen tests used in arbovirus surveillance. For this reason, we did not test pools where an arbovirus was identified by serology, potentially precluding the detection of co-infecting viruses. Furthermore, serologic tests were not 100% sensitive, as evidenced by detection of tested viruses RRV, EHV and KOKV in selected pools. Some pools also contained multiple viruses (Table 1). These results suggest that if surveillance platforms begin using deep sequencing for virus detection, testing of some known arbovirus-positive samples may be warranted, even at added expense. Although the majority of the viruses detected in this study were from 3rd or 4th passages, we also detected MURBV and sequenced its entire genome directly from a homogenized pool of mosquitoes (sample 934), indicating that this approach will also work directly on unpassaged arthropod pools.

Conclusions

We used viral metagenomics in concert with traditional arbovirus surveillance to characterize known and new arboviruses in field-collected mosquitoes. This study exemplifies the application of deep sequencing technologies for the genetic characterization of arboviruses following their amplification in cell culture, as this both demonstrates their ability to replicate in mammalian and insect cells, and increases viral titer to facilitate full genome sequencing. We identified 2 novel rhabdoviruses, 2 novel bunyaviruses, and we detected the arbovirus LNV not previously recognized in Australia. Further deployment of deep sequencing to characterize arboviruses in anthropophilic vectors will enable a more complete description of potential arbovirus threats to human and animal health. Whether the

novel and newly recognized arboviruses detected here infect animals or humans and cause disease will require serological studies and the further analysis of unexplained disease in exposed animal and human populations.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

The NSW Arbovirus Surveillance Program was funded by the NSW Ministry of Health. The sequencing work was supported by the Blood Systems Research Institute and National Institutes of Health R01 HL105770.

List of Virus Abbreviations

ABLV	Australian bat lyssavirus
AHSV	African horse sickness virus
AINOV	Aino virus
AKAV	Akabane virus
ALFV	Alfuy virus
ALMV	Almpiwar virus
BANV	Banna virus
BASV	Bas-Congo virus
BEAUV	Beaumont virus
BEFV	bovine ephemeral fever virus
BFV	Barmah Forest virus
BYSM	barley yellow striate mosaic
CCHFV	Crimean-Congo hemorrhagic fever virus
CHIKV	chikungunya virus
CHVV	Charleville virus
CPV	Coastal Plains virus
DAff sigma virus	<i>Drosophila affinis</i> sigma virus
DENV	dengue virus
DMel sigma virus	<i>Drosophila melanogaster</i> sigma virus
DObs sigma virus	<i>Drosophila obscura</i> sigma virus
DOUV	Douglas virus
EBLV	European bat lyssavirus
EHDV	epizootic hemorrhagic disease virus
EHV	Edge Hill virus
EUBV	Eubenangee virus
FPV	Facey's Paddock virus

GGV	Gan Gan virus
HDOOV	Humpty Doo virus
IHNV	infectious hematopoietic necrosis virus
JEV	Japanese encephalitis virus
KDV	Kadiporo virus
KOKV	Kokobera virus
KUNV	Kunjin virus
LEAV	Leanyer virus
LNV	Liao Ning virus
LYMoV	lettuce yellow mottle virus
LNyV	lettuce necrotic yellows virus
MAPV	Mapputta virus
MFSV	maize fine streak virus
MURBV	Murrumbidgee virus
MVEV	Murray Valley encephalitis virus
NCMV	northern cereal mosaic virus
NORCV	North Creek virus
PEAV	Peaton virus
PHSV	Peruvian horse sickness virus
RRV	Ross River virus
RVFV	Rift Valley fever virus
SASHV	Salt Ash virus
SFTSV	severe fever with thrombocytopenia syndrome virus
SINV	Sindbis virus
SLOV	Stretch Lagoon orbivirus
STRV	Stratford virus
SVCV	spring viremia of carp virus
TIBV	Tibrogargan virus
TILV	Tilligerry virus
TINV	Tinaroo virus
THIV	Thimiri virus
TRUV	Trubanaman virus
VHSV	viral hemorrhagic septicemia virus
VSV	vesicular stomatitis virus
WALV	Wallal virus
WARV	Warrego virus

WNV

West Nile virus

References

1. Knope K, Whelan P, Smith D, Nicholson J, Moran R, Doggett S, Sly A, Hobby M, Wright P. Arboviral diseases and malaria in Australia, 2010–11: Annual report of the National Arbovirus and Malaria Advisory Committee. *Commun Dis Intell Q Rep*. 2013; 37:E1–E20. [PubMed: 23692155]
2. Junglen S, Drosten C. Virus discovery and recent insights into virus diversity in arthropods. *Curr Opin Microbiol*. 2013; 16:507–513. [PubMed: 23850098]
3. Ng TF, Willner DL, Lim YW, Schmieder R, Chau B, Nilsson C, Anthony S, Ruan Y, Rohwer F, Breitbart M. Broad surveys of DNA viral diversity obtained through viral metagenomics of mosquitoes. *PLoS One*. 2011; 6:e20579. [PubMed: 21674005]
4. Bishop-Lilly KA, Turell MJ, Willner KM, Butani A, Nolan NM, Lentz SM, Akmal A, Mateczun A, Brahmabhatt TN, Sozhamannan S, Whitehouse CA, Read TD. Arbovirus detection in insect vectors by rapid, high-throughput pyrosequencing. *PLoS Negl Trop Dis*. 2010; 4:e878. [PubMed: 21085471]
5. Chowdhary R, Street C, Travassos da Rosa A, Nunes MR, Tee KK, Hutchison SK, Vasconcelos PF, Tesh RB, Lipkin WI, Briese T. Genetic characterization of the Wyeomyia group of orthobunyaviruses and their phylogenetic relationships. *J Gen Virol*. 2012; 93:1023–1034. [PubMed: 22278828]
6. Hall-Mendelin S, Allcock R, Kresoje N, van den Hurk AF, Warrilow D. Detection of arboviruses and other micro-organisms in experimentally infected mosquitoes using massively parallel sequencing. *PLoS One*. 2013; 8:e58026. [PubMed: 23460918]
7. Quan PL, Junglen S, Tashmukhamedova A, Conlan S, Hutchison SK, Kurth A, Ellerbrok H, Egholm M, Briese T, Leendertz FH, Lipkin WI. Moussa virus: a new member of the *Rhabdoviridae* family isolated from *Culex decens* mosquitoes in Cote d'Ivoire. *Virus Res*. 2010; 147:17–24. [PubMed: 19804801]
8. Swei A, Russell BJ, Naccache SN, Kabre B, Veeraraghavan N, Pilgard MA, Johnson BJ, Chiu CY. The genome sequence of Lone Star virus, a highly divergent bunyavirus found in the *Amblyomma americanum* tick. *PLoS One*. 2013; 8:e62083. [PubMed: 23637969]
9. Vasilakis N, Widen S, Travassos da Rosa AP, Wood TG, Walker PJ, Holmes EC, Tesh RB. Malpais spring virus is a new species in the genus vesiculovirus. *Virology*. 2013; 447:10–19. [PubMed: 23497016]
10. Ma M, Huang Y, Gong Z, Zhuang L, Li C, Yang H, Tong Y, Liu W, Cao W. Discovery of DNA viruses in wild-caught mosquitoes using small RNA high throughput sequencing. *PLoS One*. 2011; 6:e24758. [PubMed: 21949749]
11. Russell, RC. *A Colour Photo Atlas of Mosquitoes of Southeastern Australia*. Department of Medical Entomology; Westmead, New South Wales: 1996.
12. Russell RC, Doggett SL, Clancy J, Haniotis J, Patsouris K, Hueston L, Marchetti M, Dwyer DE. Arbovirus and vector surveillance in NSW, 1997–2000. *Arbovirus Research in Australia*. 2001; 8:304–313.
13. Victoria JG, Kapoor A, Li L, Blinkova O, Slikas B, Wang C, Naeem A, Zaidi S, Delwart E. Metagenomic analyses of viruses in stool samples from children with acute flaccid paralysis. *Journal of Virology*. 2009; 83:4642–4651. [PubMed: 19211756]
14. Ruby JG, Bellare P, Derisi JL. PRICE: software for the targeted assembly of components of (Meta) genomic sequence data. *G3 (Bethesda)*. 2013; 3:865–880. [PubMed: 23550143]
15. Thompson JD, Higgins DG, Gibson TJ. CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. *Nucleic Acids Res*. 1994; 22:4673–4680. [PubMed: 7984417]
16. Tamura K, Peterson D, Peterson N, Stecher G, Nei M, Kumar S. MEGA5: molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. *Mol Biol Evol*. 2011; 28:2731–2739. [PubMed: 21546353]
17. Kuzmin IV, Novella IS, Dietzgen RG, Padhi A, Rupprecht CE. The rhabdoviruses: biodiversity, phylogenetics, and evolution. *Infect Genet Evol*. 2009; 9:541–553. [PubMed: 19460320]

18. Walter CT, Barr JN. Recent advances in the molecular and cellular biology of bunyaviruses. *J Gen Virol.* 2011; 92:2467–2484. [PubMed: 21865443]
19. Savji N, Palacios G, Travassos da Rosa A, Hutchison S, Celone C, Hui J, Briese T, Calisher CH, Tesh RB, Lipkin WI. Genomic and phylogenetic characterization of Leanyer virus, a novel orthobunyavirus isolated in northern Australia. *J Gen Virol.* 2011; 92:1676–1687. [PubMed: 21402599]
20. Lu Z, Liu H, Fu S, Lu X, Dong Q, Zhang S, Tong S, Li M, Li W, Tang Q, Liang G. Liao ning virus in China. *Virol J.* 2011; 8:282. [PubMed: 21649929]
21. Lv X, Mohd Jaafar F, Sun X, Belhouchet M, Fu S, Zhang S, Tong SX, Lv Z, Mertens PP, Liang G, Attoui H. Isolates of Liao ning virus from wild-caught mosquitoes in the Xinjiang province of China in 2005. *PLoS One.* 2012; 7:e37732. [PubMed: 22649554]
22. Hooper PT, Lunt RA, Gould AR, Hyatt AD, Russell GM, Kattenbelt JA, Blacksell SD, Reddacliff LA, Kirkland PD, Davis RJ, Durham PJ, Bishop AL, Waddington J. Epidemic of blindness in kangaroos—evidence of a viral aetiology. *Aust Vet J.* 1999; 77:529–536. [PubMed: 10494400]
23. Hooper P. Kangaroo blindness and some other new viral diseases in Australia. *Aust Vet J.* 1999; 77:514–515. [PubMed: 10494397]
24. Mores CN, Turell MJ, Dyer J, Rossi CA. Phylogenetic relationships among orthobunyaviruses isolated from mosquitoes captured in Peru. *Vector Borne Zoonotic Dis.* 2009; 9:25–32. [PubMed: 18759638]
25. Blacksell SD, Lunt RA, White JR. Rapid identification of Australian bunyavirus isolates belonging to the Simbu serogroup using indirect ELISA formats. *J Virol Methods.* 1997; 66:123–133. [PubMed: 9220398]
26. Li Q. First isolation of eight strains of new orbivirus (Banna) from patients with innominate fever in Xinjiang. *Endemic Diseases Bulletin.* 1992; 7:77–82.
27. Xu P, Wang Y, Zuo J, Lin J, Xu P. Isolation of orbiviruses from sera of patients with unknown fever and encephalitis collected from Yunnan Province. *Chinese Journal of Virology.* 1990; 6:27–33.
28. Liu H, Li MH, Zhai YG, Meng WS, Sun XH, Cao YX, Fu SH, Wang HY, Xu LH, Tang Q, Liang GD. Banna virus, China, 1987–2007. *Emerg Infect Dis.* 2010; 16:514–517. [PubMed: 20202434]
29. Chen B, Tao S. Arbovirus survey in China in recent ten years. *Chin Med J (Engl).* 1996; 109:13–15. [PubMed: 8758350]
30. Liting S, Chen B, Chou Z. Isolation and identification of new members of coltivirus from mosquitoes collected in China. *Zhonghua Shi Yan He Lin Chuang Bing Du Xue Za Zhi.* 1995; 9:7–10.
31. Brown SE, Gorman BM, Tesh RB, Knudson DL. Coltiviruses isolated from mosquitoes collected in Indonesia. *Virology.* 1993; 196:363–367. [PubMed: 8102827]
32. Nabeshima T, Thi Nga P, Guillermo P, Parquet Mdel C, Yu F, Thanh Thuy N, Minh Trang B, Tran Hien N, Sinh Nam V, Inoue S, Hasebe F, Morita K. Isolation and molecular characterization of Banna virus from mosquitoes, Vietnam. *Emerg Infect Dis.* 2008; 14:1276–1279. [PubMed: 18680655]
33. Tao SJ, Chen BQ. Studies of coltivirus in China. *Chin Med J (Engl).* 2005; 118:581–586. [PubMed: 15820089]
34. Attoui H, Mohd Jaafar F, Belhouchet M, Tao S, Chen B, Liang G, Tesh RB, de Micco P, de Lamballerie X. Liao ning virus, a new Chinese seadornavirus that replicates in transformed and embryonic mammalian cells. *J Gen Virol.* 2006; 87:199–208. [PubMed: 16361432]
35. Li WJ, Wang JL, Li MH, Fu SH, Wang HY, Wang ZY, Jiang SY, Wang XW, Guo P, Zhao SC, Shi Y, Lu NN, Nasci RS, Tang Q, Liang GD. Mosquitoes and mosquito-borne arboviruses in the Qinghai-Tibet Plateau—focused on the Qinghai area, China. *Am J Trop Med Hyg.* 2010; 82:705–711. [PubMed: 20348523]
36. Mertens, P.; Maan, S.; Samuel, A.; Attoui, H. Orbiviruses, *Reoviridae*. In: Fauquet, C.; Mayo, M.; Maniloff, J.; Desselberger, U.; Ball, L., editors. *Virus taxonomy: either report of the international committee on taxonomy of viruses*. Elsevier Academic Press; London: 2005. p. 466-483.

37. Belaganahalli MN, Maan S, Maan NS, Tesh R, Attoui H, Mertens PP. Umatilla virus genome sequencing and phylogenetic analysis: identification of stretch lagoon orbivirus as a new member of the Umatilla virus species. *PLoS One*. 2011; 6:e23605. [PubMed: 21897849]
38. Jansen CC, Prow NA, Webb CE, Hall RA, Pyke AT, Harrower BJ, Pritchard IL, Zborowski P, Ritchie SA, Russell RC, Van Den Hurk AF. Arboviruses isolated from mosquitoes collected from urban and peri-urban areas of eastern Australia. *J Am Mosq Control Assoc*. 2009; 25:272–278. [PubMed: 19852216]
39. Cowled C, Palacios G, Melville L, Weir R, Walsh S, Davis S, Gubala A, Lipkin WI, Briese T, Boyle D. Genetic and epidemiological characterization of Stretch Lagoon orbivirus, a novel orbivirus isolated from *Culex* and *Aedes* mosquitoes in northern Australia. *J Gen Virol*. 2009; 90:1433–1439. [PubMed: 19282430]
40. Reddacliff L, Kirkland P, Philbey A, Davis R, Vogelnest L, Hulst F, Blyde D, Deykin A, Smith J, Hooper P, Gould A, Hyatt A. Experimental reproduction of viral chorioretinitis in kangaroos. *Aust Vet J*. 1999; 77:522–528. [PubMed: 10494399]
41. Standfast HA, Dyce AL, St George TD, Muller MJ, Doherty RL, Carley JG, Filippich C. Isolation of arboviruses from insects collected at Beatrice Hill, Northern Territory of Australia, 1974–1976. *Aust J Biol Sci*. 1984; 37:351–366. [PubMed: 6152599]
42. Tallis G, Ng S, Azuolas J. A sero-prevalence study for ‘wallal-type’ virus infection among park workers. *Aust N Z J Public Health*. 1998; 22:515. [PubMed: 9659784]
43. Foley DH, Wilkerson RC, Cooper RD, Volovsek ME, Bryan JH. A molecular phylogeny of *Anopheles annulipes* (Diptera: Culicidae) *sensu lato*: the most species-rich anopheline complex. *Mol Phylogenet Evol*. 2007; 43:283–297. [PubMed: 17126567]
44. Cooper RD, Frances SP, Waterson DG, Piper RG, Sweeney AW. Distribution of anopheline mosquitoes in northern Australia. *J Am Mosq Control Assoc*. 1996; 12:656–663. [PubMed: 9046472]
45. Russell RC, Cloonan MJ, Wells PJ, Vale TG. Mosquito (Diptera: *Culicidae*) and arbovirus activity on the south coast of New South Wales, Australia, in 1985–1988. *J Med Entomol*. 1991; 28:796–804. [PubMed: 1685192]
46. Quan PL, Williams DT, Johansen CA, Jain K, Petrosov A, Diviney SM, Tashmukhamedova A, Hutchison SK, Tesh RB, Mackenzie JS, Briese T, Lipkin WI. Genetic characterization of K13965, a strain of Oak Vale virus from Western Australia. *Virus Res*. 2011; 160:206–213. [PubMed: 21740935]
47. Azuolas JK, Wishart E, Bibby S, Ainsworth C. Isolation of Ross River virus from mosquitoes and from horses with signs of musculo-skeletal disease. *Aust Vet J*. 2003; 81:344–347. [PubMed: 15080456]
48. Harley D, Sleigh A, Ritchie S. Ross River virus transmission, infection, and disease: a cross-disciplinary review. *Clin Microbiol Rev*. 2001; 14:909–932. [PubMed: 11585790]
49. Kay BH, Boyd AM, Ryan PA, Hall RA. Mosquito feeding patterns and natural infection of vertebrates with Ross River and Barmah Forest viruses in Brisbane, Australia. *Am J Trop Med Hyg*. 2007; 76:417–423. [PubMed: 17360861]
50. Clements, AN. *The Biology of Mosquitoes*. Vol. 1. CABI Publishing; New York: 2000. Osmotic and Ionic Regulation; p. 125
51. Van Den Hurk AF, Montgomery BL, Northill JA, Smith IL, Zborowski P, Ritchie SA, Mackenzie JS, Smith GA. Short report: the first isolation of Japanese encephalitis virus from mosquitoes collected from mainland Australia. *Am J Trop Med Hyg*. 2006; 75:21–25. [PubMed: 16837702]
52. Johansen CA, Nisbet DJ, Foley PN, Van Den Hurk AF, Hall RA, Mackenzie JS, Ritchie SA. Flavivirus isolations from mosquitoes collected from Saibai Island in the Torres Strait, Australia, during an incursion of Japanese encephalitis virus. *Med Vet Entomol*. 2004; 18:281–287. [PubMed: 15347396]
53. Johansen CA, Nisbet DJ, Zborowski P, van den Hurk AF, Ritchie SA, Mackenzie JS. Flavivirus isolations from mosquitoes collected from western Cape York Peninsula, Australia, 1999–2000. *J Am Mosq Control Assoc*. 2003; 19:392–396. [PubMed: 14710742]
54. Powers AM, Logue CH. Changing patterns of chikungunya virus: re-emergence of a zoonotic arbovirus. *J Gen Virol*. 2007; 88:2363–2377. [PubMed: 17698645]

55. Yu XJ, Liang MF, Zhang SY, Liu Y, Li JD, Sun YL, Zhang L, Zhang QF, Popov VL, Li C, Qu J, Li Q, Zhang YP, Hai R, Wu W, Wang Q, Zhan FX, Wang XJ, Kan B, Wang SW, Wan KL, Jing HQ, Lu JX, Yin WW, Zhou H, Guan XH, Liu JF, Bi ZQ, Liu GH, Ren J, Wang H, Zhao Z, Song JD, He JR, Wan T, Zhang JS, Fu XP, Sun LN, Dong XP, Feng ZJ, Yang WZ, Hong T, Zhang Y, Walker DH, Wang Y, Li DX. Fever with thrombocytopenia associated with a novel bunyavirus in China. *N Engl J Med*. 2011; 364:1523–1532. [PubMed: 21410387]
56. Suthar MS, Diamond MS, Gale M Jr. West Nile virus infection and immunity. *Nat Rev Microbiol*. 2013; 11:115–128. [PubMed: 23321534]
57. Simmons CP, Farrar JJ, van Vinh Chau N, Wills B. Dengue. *New England Journal of Medicine*. 2012; 366:1423–1432. [PubMed: 22494122]
58. Grard G, Fair JN, Lee D, Slikas E, Steffen I, Muyembe JJ, Sittler T, Veeraraghavan N, Ruby JG, Wang C, Makuwa M, Mulembakani P, Tesh RB, Mazet J, Rimoin AW, Taylor T, Schneider BS, Simmons G, Delwart E, Wolfe ND, Chiu CY, Leroy EM. A novel rhabdovirus associated with acute hemorrhagic fever in central Africa. *PLoS Pathog*. 2012; 8:e1002924. [PubMed: 23028323]
59. Monath TP, Cropp CB, Frazier CL, Murphy FA, Whitfield SG. Viruses isolated from reptiles: identification of three new members of the family *Rhabdoviridae*. *Arch Virol*. 1979; 60:1–12. [PubMed: 90494]
60. St George TD, Standfast HA, Cybinski DH, Filippich C, Carley JG. Peaton virus: a new Simbu group arbovirus isolated from cattle and *Culicoides brevitarsis* in Australia. *Aust J Biol Sci*. 1980; 33:235–243. [PubMed: 7436869]
61. Murray MD, Kirkland PD. Bluetongue and Douglas virus activity in New South Wales in 1989: further evidence for long-distance dispersal of the biting midge *Culicoides brevitarsis*. *Aust Vet J*. 1995; 72:56–57. [PubMed: 7779035]
62. Boughton CR, Hawkes RA, Naim HM. Arbovirus infection in humans in NSW: seroprevalence and pathogenicity of certain Australian bunyaviruses. *Aust N Z J Med*. 1990; 20:51–55. [PubMed: 2108660]
63. Animal Health Australia. 2009–2010 Report National Arbovirus Monitoring Program. 2010.

Research Highlights

- We used metagenomics to identify virus sequences in Australian mosquitoes
- We detected two novel dimarhabdoviruses and two novel orthobunyaviruses
- Liao Ning virus (*Reovirus*), previously only observed in China, was also detected
- Viral metagenomics combined with surveillance can identify novel arboviruses

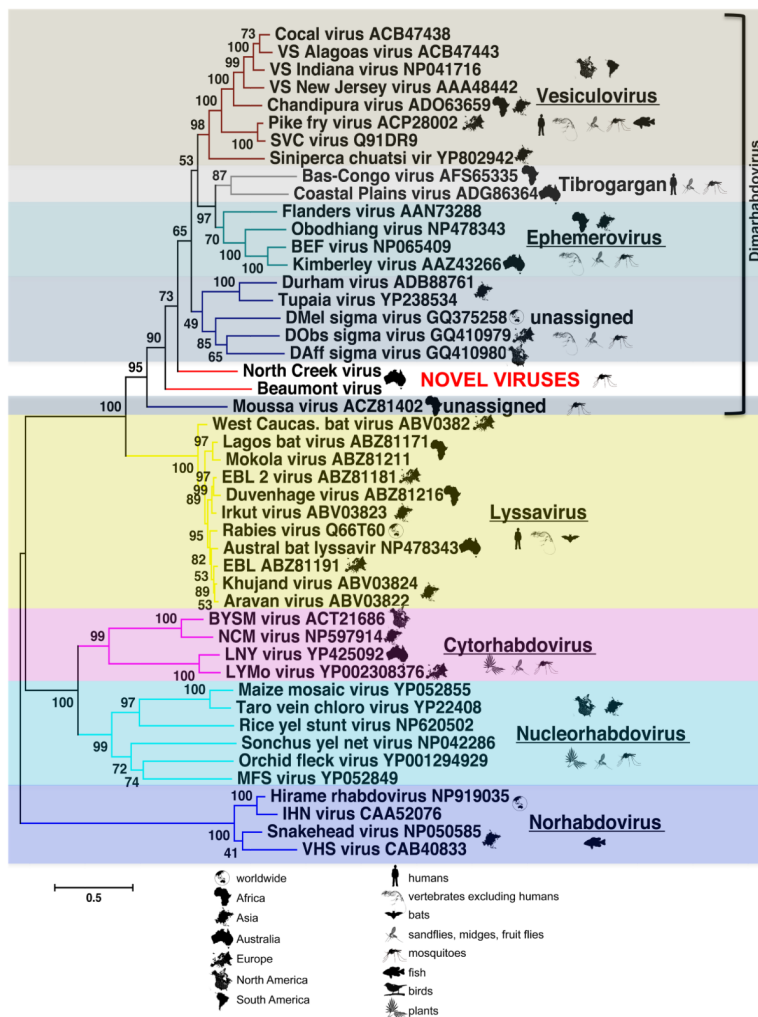
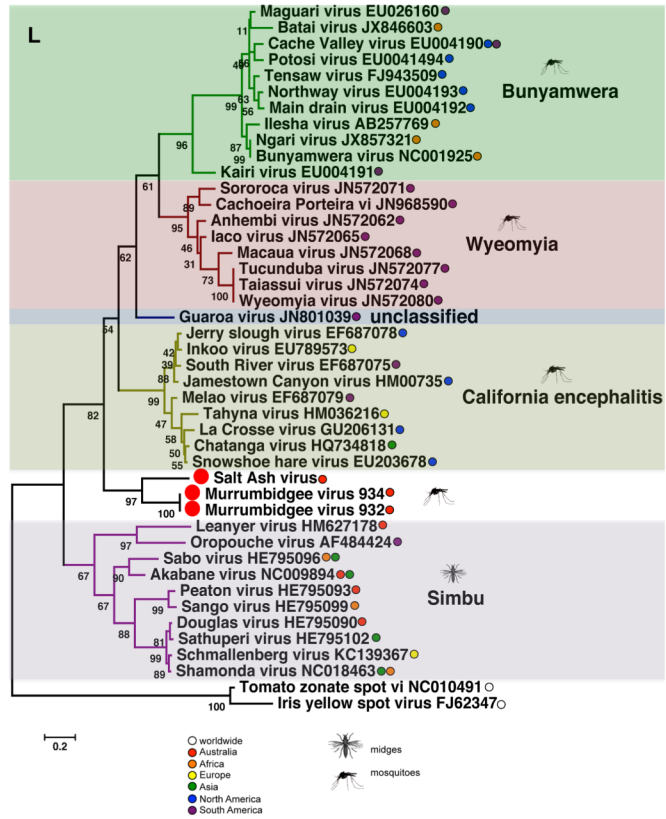


Figure 1. *Rhabdovirus* maximum likelihood L gene open reading frame phylogeny with novel viruses highlighted in red. Numbers left of branches show statistical significance of tree topologies based on 1000 bootstrap re-sampling iterations. Symbols indicate geographic distributions and organism pictures show host ranges. Colored boxes show clade groupings; underlined groups denote established genera. L gene sequences used were 1568 aa (BEAUV), 2120 aa (NORCV, complete gene), ≈2100 aa (complete gene) for GenBank sequences.

Figure 2A



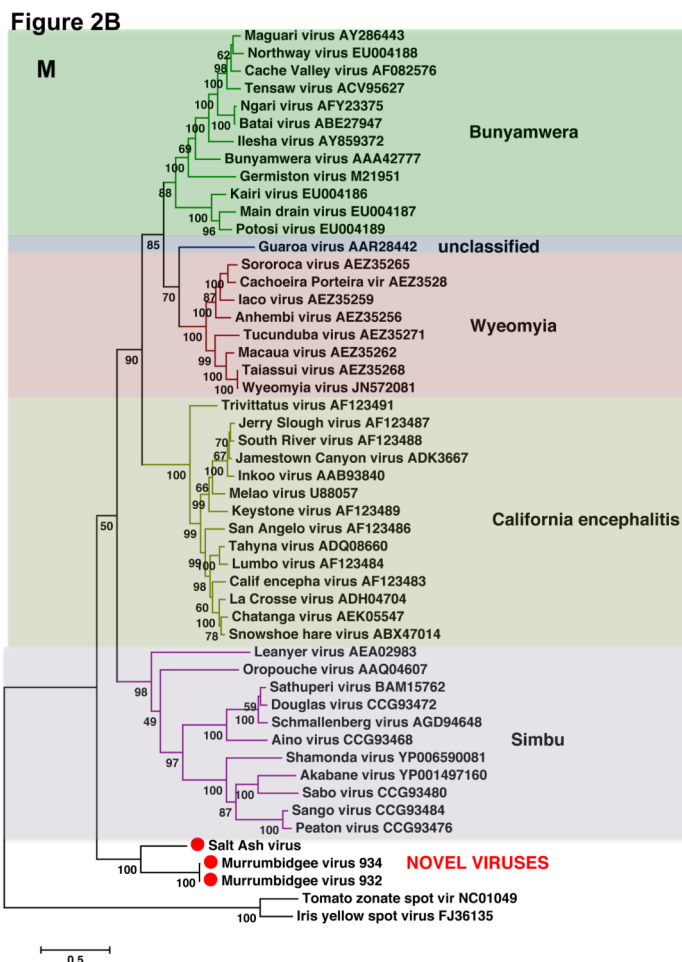


Figure 2.
 Figure 2A, B: Orthobunyavirus maximum likelihood phylogenies of the L (A) and M (B) segment open reading frames with novel viruses highlighted in red. Sequences for M and L from GenBank were derived from the same isolate, when possible. Numbers to the left of branches show statistical significance of tree topologies based on 1000 bootstrap re-sampling iterations. Neighbor joining trees (not shown) exhibited similar topologies. Colored boxes show established orthobunyavirus groups defined based on genetic or serologic relatedness. Insect pictures show primary vectors. Gene sequences used were L: 2212 aa for SASHV, complete 2242 aa for MURBV, complete \approx 2200 aa for GenBank sequences, M: complete 1377 aa for SASHV, complete 1371 aa for MURBV, complete \approx 1400 aa for GenBank sequences.

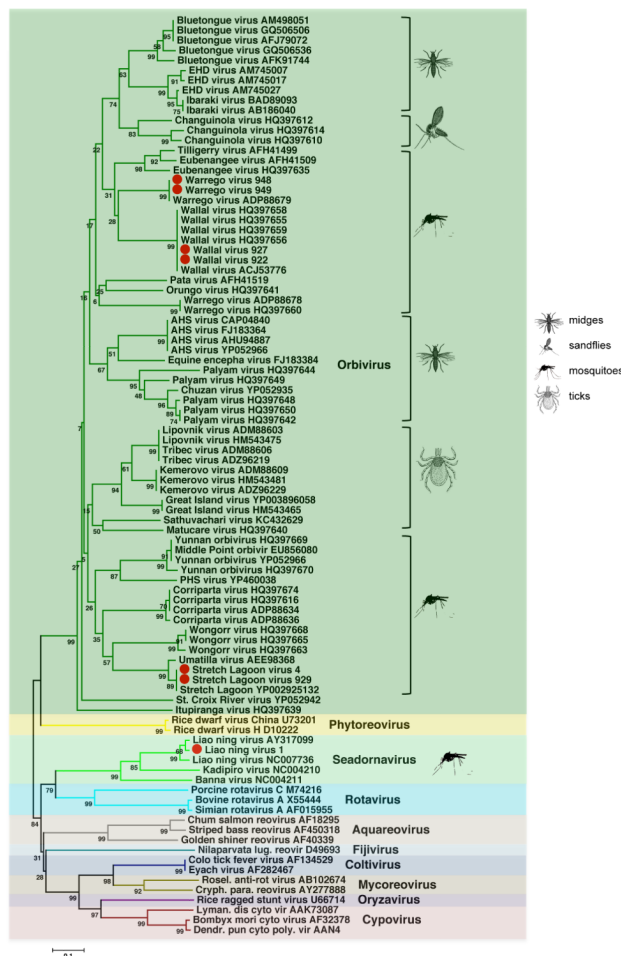


Figure 3. *Reoviridae* RdRp gene maximum likelihood phylogeny with viruses detected here highlighted in red. Numbers to the left of branches show statistical significance of tree topologies based on 1000 bootstrap re-sampling iterations. Colored boxes show established reovirus genera. Pictures show primary vector types. Full ≈ 1250 aa RdRp gene sequences we used for viruses sequenced in this study, some shorter ≈ 130 aa sequences from GenBank were used when full sequences were not available.

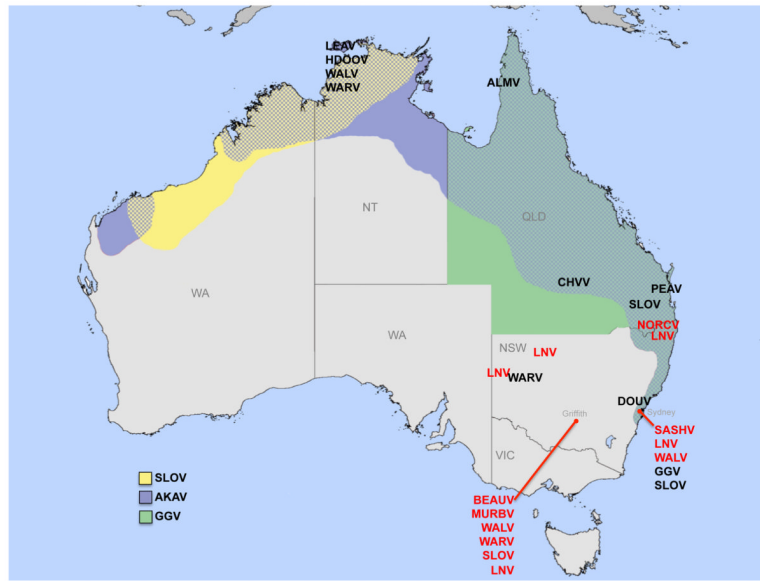


Figure 4. Map of Australia showing mosquito collection locations for arboviruses detected in this study (red) as well as sites where other Australian arboviruses have been isolated previously (black and shaded regions) (59–62); Akabane virus (AKAV, purple) (63), SLOV (yellow) (19, 62), and GGV (green) (62). EHV, KOKV, and RRV are not shown since they exhibit diffuse continental distributions. LNV has not been previously isolated in Australia. NSW, New South Wales; NT, Northern Territory; QLD, Queensland; SA, South Australia; VIC, Victoria, WA, Western Australia.

Table 1

Virus genomes detected by deep sequencing insect pools passaged in cell culture. Bold denotes novel genomes detected for the first time.

Sample	Collection year	Collection source	Collection location in New South Wales, Australia	Passage history	Virus genome detected		
					virus 1	virus 2	virus 3
1	2004	<i>Anopheles amictus</i>	Bourke	ppcc	LNV	RRV	
2	1998	biting midges	Wentworth	bbc	LNV	RRV	
3	1996	<i>Aedes camptorhynchus</i>	Sydney	bbcc	RRV		
4	1995	<i>Culex australicus</i> , <i>Cx. molestus</i>	Griffith	bbbc	SLOV	LNV	RRV
5	2005	<i>Anopheles amictus</i>	Boggabilla	bbc	LNV	RRV	
6	2005	<i>Anopheles annulipes</i>	Griffith	ppp	BEAUV (<i>Rhabdoviridae</i>)	LNV	RRV
7	1995	<i>Culex quinquefasciatus</i>	Sydney	ppcc	RRV		
8	1997	<i>Culex sitiens</i>	Ballina	pppcc	LNV	RRV	
9	1995	<i>Aedes vigilax</i>	Batemans Bay	pppcc	LNV	RRV	
10	1995	<i>Culex sitiens</i>	Tweed Heads	ppcc	LNV	RRV	
922	2005	<i>Culex annulirostris</i>	Hawkesbury	ppcc	WALV		
927	2010	<i>Anopheles annulipes</i>	Griffith	ppc	WALV		
929	2006	<i>Culex annulirostris</i>	Griffith	ppc	SLOV		
931	1992	<i>Aedes vigilax</i>	Port Stephens	bbc	SASHV (<i>Bunyaviridae</i>)		
932	1996	<i>Anopheles annulipes</i>	Griffith	bb	MURBY (<i>Bunyaviridae</i>)		
933	1999	<i>Culex annulirostris</i>	Griffith	ppppcc	KOKV		
934	1997	<i>Anopheles annulipes</i>	Griffith	none	MURBY (<i>Bunyaviridae</i>)		
935	1998	<i>Aedes vigilax</i>	Batemans Bay	pppcc	EHV		
948	1995	<i>Culex annulirostris</i>	Griffith	bbbc	WARV		
949	1995	<i>Anopheles annulipes</i>	Griffith	bbc	WARV		
954	1997	<i>Culex sitiens</i>	Ballina	pppcc	NORCV (<i>Rhabdoviridae</i>)		
955	n/a	hamster cell culture	n/a	b			

Key to abbreviations: b: baby hamster kidney cell passage, p: porcine stable equine kidney cell passage, c: C6/36 *Aedes albopictus* larval cell passage, BEAUV: Beaumont virus, EHV: Edge Hill virus, KOKV: Kobobera virus, LNV: Liao Ning virus, MURBV: Murrumbidgee virus, NORCV: North Creek virus, RRV: Ross River virus, SASH: Salt Ash virus, SLOV: Stretch Lagoon orbivirus, WALV: Wallal virus, WARV: Warrego virus.

Table 3

Percent L gene amino acid identities for selected dimarhabdoviruses, with GenBank accession numbers noted.

Beaumont virus	100	Beaumont virus	North Creek virus	Tupaia virus YP238534	DMel sigma virus GQ375258	Cocal virus ACB47438	Chandipura virus ADO63659	Bas-Congo virus AFS65335	Coastal Plains virus ADG86364	Obodhiang virus NP478343	Kimberley virus AAZ43266	Moussa virus ACZ81402
Beaumont virus	100											
North Creek virus	37	100										
Tupaia virus YP238534	33	37	100									
DMel sigma virus GQ375258	31	37	41	100								
Cocal virus ACB47438	34	39	45	42	100							
Chandipura virus ADO63659	35	39	44	41	61	100						
Bas-Congo virus AFS65335	39	34	36	34	38	39	100					
Coastal Plains virus ADG86364	31	35	38	36	41	40	42	100				
Obodhiang virus NP478343	33	35	37	34	39	39	38	40	100			
Kimberley virus AAZ43266	31	35	37	36	41	40	39	39	42	50	100	
Moussa virus ACZ81402	27	31	33	30	33	33	31	31	31	30	30	100

Colors denote species that cluster (Figure 1) and shaded boxes highlight identities for species in the same group.

Table 4

L (A) and M (B) segment amino acid identities for the novel Murrumbidgee and Salt Ash viruses versus representative orthobunyavirus species with GenBank accession numbers noted.

L segment									
	Murrumbidgee virus	Salt Ash virus	Oropouche virus AAQ04607	Peaton virus CCG93476	Guaroa virus JN801039	Jamestown Canyon ADK3667	La Crosse virus ADH04704	Bunyamwera virus AAA42777	Wyomyia virus JNS72081
Murrumbidgee virus	100								
Salt Ash virus	61	100							
Oropouche virus AAQ04607	46	47	100						
Peaton virus CCG93476	45	48	59	100					
Guaroa virus JN801039	47	47	47	46	100				
Jamestown Canyon ADK3667	50	51	51	50	49	100			
La Crosse virus ADH04704	50	50	51	49	49	83	100		
Bunyamwera virus AAA42777	49	49	50	48	48	55	56	100	
Wyomyia virus JNS72081	48	50	49	47	48	55	55	54	100

M segment									
	Murrumbidgee virus	Salt Ash virus	Oropouche virus AAQ04607	Peaton virus CCG93476	Guaroa virus JN801039	Jamestown Canyon ADK3667	La Crosse virus ADH04704	Bunyamwera virus AAA42777	Wyomyia virus JNS72081
Murrumbidgee virus	100								
Salt Ash virus	52	100							
Oropouche virus AAQ04607	29	30	100						
Peaton virus CCG93476	30	31	40	100					
Guaroa virus JN801039	33	33	32	30	100				
Jamestown Canyon ADK3667	32	33	33	33	32	100			
La Crosse virus ADH04704	32	33	33	33	32	75	100		
Bunyamwera virus AAA42777	32	33	35	32	32	44	44	100	
Wyomyia virus JNS72081	33	33	34	32	33	44	44	54	100

Colors denote orthobunyavirus groups and correspond to patterns in Figure 2. Shaded boxes highlight identities for species in the same group.