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Gene duplication and phylogeography of North American members of the Hart Park serogroup of avian rhabdoviruses

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\textbf{A B S T R A C T}

\textit{Flanders virus} (FLAV) and Hart Park virus (HPV) are rhabdoviruses that circulate in mosquito–bird cycles in the eastern and western United States, respectively, and constitute the only two North American representatives of the Hart Park serogroup. Previously, it was suggested that FLAV is unique among the rhabdoviruses in that it contains two pseudogenes located between the P and M genes, while the cognate sequence for HPV has been lacking. Herein, we demonstrate that FLAV and HPV do not contain pseudogenes in this region, but encode three small functional proteins designated as U1–U3 that apparently arose by gene duplication. To further investigate the U1–U3 region, we conducted the first large-scale evolutionary analysis of a member of the Hart Park serogroup by analyzing over 100 spatially and temporally distinct FLAV isolates. Our phylogeographic analysis demonstrates that although FLAV appears to be slowly evolving, phylogenetically divergent lineages co-circulate sympatriically.

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\section*{Introduction}

\textit{Flanders virus} (FLAV) and Hart Park virus (HPV) are two closely-related members of the Hart Park serogroup of the family \textit{Rhabdoviridae} that are maintained in mosquito–passerine bird transmission cycles in the eastern and western United States, respectively (Whitney, 1964; Johnson, 1965; Kokernot et al., 1969; Crane et al., 1970; Main et al., 1979; Main, 1981). Viruses in the Hart Park serogroup were initially classified together based on antigenic cross-reactivity in complement fixation, neutralization, immunodiffusion and/or immunofluorescence assays (Boyd, 1972; Frazier and Shope, 1979). In addition to FLAV and HPV, other members in earlier classifications of the serogroup included Mosquito virus (MQOV), a virus first isolated in Brazil, and two African viruses, Mosaic virus (MOSV) and Kamese virus (KAMV) (Tesh et al., 1983; Calisher et al., 1989). Besides their antigenic relatedness, these five geographically disparate viruses appear to share a similar mechanism of transmission, as virus isolation data indicated that they were predominately associated with birds and/or culicine (e.g., \textit{Culex}, \textit{Culiseta}) mosquitoes (Karabatsos, 1985).

More recently, \textit{Wongabel virus} (WONV), Parry Creek virus (PCRV), and \textit{Ngaoiangan virus} (NGAV) have also been provisionally included into the serogroup based on genetic and phylogenetic (rather than antigenic) relationships (Bourhy et al., 2003; Gubala et al., 2008, 2010). These three viruses were originally isolated in Australia, and besides the serological observation that the natural host range of NGAV may include macropods, they also appear to be predominantly associated with birds and culicine mosquitoes or other hematophagous insects such as Culicoids biting midges (Humphrey-Smith et al., 1991; Bourhy et al., 2008; Gubala et al., 2010). Additionally, two recently described (but historically isolated) Australian viruses recovered from \textit{Culex annulirostris} – Holmes Jungle virus (HQV) and Orchid River virus (ORRV) – appear to be new members of the serogroup (Gubala, 2012), as do Bangoran virus (BGNV) and Porton’s virus (PORV) (Dacheux et al., 2010). Whether these twelve potential Hart Park serogroup
members will eventually be designated as a new genus within the Rhabdoviridae will likely entail a more comprehensive phylogenetic analysis (such as full genome studies) of these and other unclassified rhabdoviruses of the Dimarhabdovirus supergroup.

FLAV is unique among the rhabdoviruses in that it purportedly contains a 19 kDa protein gene flanked on either side by putative pseudogenes (GenBank accession AH012179). No comparative sequence for HPV has previously been available. These three consecutive genes, originally termed pseudogene 1, 19 kDa protein gene, and pseudogene 2, are located between the phosphoprotein (P) and matrix (M) genes, such that the FLAV genome is currently represented as 3′-nucleoprotein(N)-P-pseudogene1–19K-pseudogene2–M-glycoprotein(G)-polymerase(L)-5′ (Dietzgen et al., 2011). However, given the constraints on genome size that seem surprising that FLAV would apparently carry two sequences that have no functional role. As Australian Hart Park serogroup viruses (i.e., WONV and NGAV) contain three complete intact ORFs between their P and M genes (Gubala et al., 2008, 2010), we sought to analyze this region in the two North American members of the serogroup, FLAV and HPV, and clarify this apparent genomic complexity. Additionally, we investigated the potential encoding of a viroporin-like small hydrophobic (SH) protein located between the G and L proteins and undertook the first comprehensive evolutionary study of a Hart Park serogroup virus by analyzing more than 100 pseudogene region sequences of FLAV isolates collected over a 50-year period.

Results and discussion

Gene, mRNA, and protein analysis of the pseudogene region and SH ORF

Our genetic analysis of multiple FLAV isolates indicated that the two putative pseudogenes located between the P and M genes contained complete uninterrupted ORFs flanked by conserved transcriptional start (UCCUCMKIAG) and stop/polyadenylation (CU3) sequences, suggesting that they in fact encode functional proteins (GenBank accessions KF028661–KF028670). The predicted proteins associated with pseudogene 1, the 19 kDa protein gene, and pseudogene 2 ORFs in FLAV were very similar in size, with lengths of 161, 165, and 160 amino acids, respectively. Similar results were found with HPV (GenBank accession KF028764), indicating both viruses had three complete ORFs between the P and M genes. Cloning of RT-PCR products generated from RNA extracted from FLAV-infected Vero cells demonstrated that polyadenylated transcripts of the two putative pseudogene sequences (as well as the 19 kDa protein gene) were being produced, again indicating that they are functional ORFs. Functionality was further supported as an analysis of the pseudogene 1, 19 kDa protein gene, and pseudogene 2 sequences of 10 FLAV isolates produced d\(d_0/d_s\) ratios of 0.07, 0.02 and 0.09, respectively, indicative of strong selective (i.e., functional) constraints rather than the selective neutrality expected of pseudogenes (in which \(d_0/d_s\) ratios would tend to be a value of ~1.0). Similarly, a \(d_0/d_s\) of 0.07 was observed in 103 pseudogene 1 (U1) sequences (see below), again revealing strong selective constraints.

In addition to the predicted N, P, M, G, and L proteins, we detected three small viral protein bands when we probed FLAV-infected Vero cell lysates in a Western blot using FLAV-specific antisera (Fig. 1). Based on their respective molecular weights, the L (238.54 kDa), G (71.05 kDa), N (50.40 kDa), and M (25.83 kDa) proteins were identified by their approximate size in the immunoblot (Fig. 1). Although the predicted P protein (25.78 kDa) is very similar in size to the M protein, the former is known to migrate in SDS-PAGE gels at between 40 and 50 kDa (Dietzgen et al., 2011), suggesting P is the band around 40 kDa (size known from additional blots) beneath N. As the predicted molecular weights of the products of pseudogene 1, the 19 kDa protein gene, and pseudogene 2 are essentially identical to one another (18.58, 18.98, and 18.93 kDa, respectively), this suggests that the band just beneath the 20 kDa marker (which is as immunoreactive as the N or M bands) might be the co-migration of the three protein products, provided that their migration is not affected by any post-translational modifications or physiochemical differences. Similarly, the slightly larger band of ~23 kDa might represent a modified form (e.g., phosphorylated) of one of the pseudogene region proteins or an in vivo cleavage product as suggested by Boyd and Whitaker-Dowling (1988). Finally, the lowest band could represent an additional cleavage product, a faster migrating form of one of the pseudogene region proteins (e.g., the acidic pseudogene 1), or the putative SH protein, a predicted 10.37 kDa viroporin-like protein lying between the G and L genes (see below).

To determine if the lower viral protein bands detected in the immunoblot were the pseudogene region products (and/or SH protein) or proteolytic truncated forms of the five major structural proteins, FLAV was purified by sucrose density gradient ultracentrifugation and select SDS-PAGE protein bands were further analyzed by nano-scale high performance liquid chromatography coupled to tandem mass spectrometry (nano HPLC-MS/MS). Although the same or similarly-sized viral bands seen in the infected cell lysates (Fig. 1) were also present (but at a lower intensity) in purified virions by immunoblotting, they were not clearly observed in the SYPRO Ruby-stained gels, suggesting that these proteins/peptides may be incorporated into virions at low concentrations, either selectively or randomly. However, a bright band(s) approximately 10–20 kDa was demonstrated to be abundantly present in purified viruses and was the only distinct band(s) present beneath the putative M protein in the fluorescent gel (not shown). In-gel tryptic digestion of this band followed by nano HPLC–MS/MS analysis identified peptides corresponding to both pseudogene 1 and pseudogene 2 products (Table 1), conclusively demonstrating that proteins of these reported pseudogenes are being expressed; whether they are normal structural components
of the virus or are incorporated into particles by chance during morphogenesis is uncertain. Additionally, peptides corresponding to N, and to a lesser extent P, M, and G, were also detected (Table 1), suggesting that cleavage products of the major structural proteins may also contribute to the observed immunoreactivity in Western blots. However, the vast majority (96 M percent) of the peptides (and hence, the major component of the 10–20 kDa band intensity) identified in the MS/MS spectra were derived from two cellular proteins, histone H4 (~11.4 kDa) and cyclophilin A (~17.9 kDa) (Table 1), both of which have been previously identified as being incorporated into rhabdovirus virions. In vesicular stomatitis New Jersey virus (VSNJV), cyclophilin A (a chaperone protein involved in protein folding) has been shown to bind to N and is required for VSNJV replication (Bose et al., 2003). Histone H4 has been observed in vesicular stomatitis Indiana virus particles (Moerdyk-Schauwecker et al., 2009), as well as other viruses such as retroviruses (Chertova et al., 2006; Segura et al., 2008) and coronaviruses (Neuman et al., 2008). Although contamination of chromat on the viral surface could be the source of histone H4, the complete absence of other similarly sized core histone proteins (i.e., H2A, H2B, H3), despite the very high abundance of histone H4, suggests its incorporation into virions may be selective and that FLAV infection may entail a tentative nuclear phase as observed in other rhabdoviruses (Godowski et al., 2002), including the related WONV (see below). Additional cellular proteins of interest found in FLAV particles (but at a much lower concentration than either histone H4 or cyclophilin A) were CD59 and heat shock protein 70 (Hsp70) (Table 1). CD59 is a complement regulatory protein which inhibits the membrane attack complex and has previously been found embedded in the outer membrane of a number of different viruses, thus providing a unique mechanism to avoid complement-mediated lysis (Vanderplaschen et al., 1998; Hu et al., 2010; Amet et al., 2012). Like cyclophilin A, Hsp70 is a protein chaperone that has been demonstrated to associate with N in rhabdovirus particles (Lahaye et al., 2012), and is often subverted from the host by viruses for a variety of functions (Gurer et al., 2002; Mayer, 2005; Nagy et al., 2011). The biological significance of these cellular proteins within FLAV particles and their potential role in the viral life cycle, if any, remains to be determined (e.g., Colpitts et al., 2011).

Based on these results, we suggest that the pseudogene 1 and 2 sequences be renamed as U1 and U3, respectively, to conform to the standard nomenclature first set forth by Gubala et al. (2008) with WONV. We also suggest that the 19 kDa protein gene be renamed as U2 for clarity among related viruses. Additionally, a putative ORF between the G and L genes in FLAV (GenBank accession KF028861), denoted as the SH ORF by Walker et al. (2011), encodes a viroporin-like protein which contains a hydrophobic transmembrane domain (WIGTLGIGLGGYVIK), similar to the transmembrane domain of the G protein (WISGILIVSLUC), and a highly basic C-terminus. Although the putative SH protein (120 aa) could be translated by mechanisms such as leaky ribosomal scanning, similar to that observed with the C proteins of the vesiculoviruses (Spiropoulou and Nichol, 1993), or by ribosomal frameshifting (~1) to produce a G-SH polyprotein (Liston and Briedis, 1995), conserved motifs found in FLAV strongly suggest that the SH protein is expressed by coupled translation. In addition to the pentanucleotide UAAUG junction between the G and SH proteins (where UAA is the termination codon for G and AUG is the start codon for SH), which has previously been demonstrated to be a common sequence for translational termination–reinitiation in a number of viruses (Horvath et al., 1990; Powell et al., 2008; Gue et al., 2009), FLAV contains sequences (motifs 1, 2, and 2′) that constitute the termination upstream ribosome-binding site (TURBS) essential for coupled translation that are very similar to those seen in members of the Norovirus genus within the family Caliciviridae, as well as Influenza B virus (Fig. 2) (Meyers, 2007; Powell, 2010). This represents the first, albeit tentative, recognition of a rhabdovirus utilizing coupled translation for protein expression and demonstrates that convergent evolution of this expression strategy has occurred in a diverse range of viral families.

Although direct evidence that the U2 and SH proteins are expressed is still lacking and will likely require more specific immunological analysis with protein-specific antibodies and/or
further mass spectrometry analysis, the fact that U1 and U3 proteins were detected and that other accessory proteins in related rhabdoviruses have been shown to be expressed (Walker et al., 2011), suggests that the U2 and SH proteins are also likely being produced in FLAV. Based of these results, we propose that the genomic organization of FLAV be demonstrated as 3′-N-P-U1-U2-U3-M-G-SH-L-5′ (Fig. 3). Whether other small putative ORFs within the FLAV genome, such as overlapping ORFs found within the N gene (Walker et al., 2011), may also be functional and express proteins remains to be determined.

**Gene duplication of the U proteins in FLAV and HPV**

The degree of amino acid sequence similarity between the U1–U3 proteins in FLAV (Fig. 4) and HPV was particularly evident and strongly suggests that they arose through gene duplication in an ancestral rhabdovirus, similar to that observed with WONV and suggested for FLAV (Walker et al., 2011; Simon-Loriere and Holmes, 2013). Additionally, the presence of identical motifs present in U1 and U3, but not U2 (e.g., YDFVWP in WONV), is of interest, and means that the order in which duplication of the genes occurred is uncertain. Previously, gene duplication has been described in a number of other rhabdoviruses, including *Bovine ephemeral fever virus* (BEFV) and *Adelaide River virus* (ARV) (Walker et al., 1992; Wang and Walker, 1993), illustrating that this particular mechanism of virus evolution appears to have occurred multiple independent times among different members of the *Rhabdoviridae*, thereby facilitating the noted complexity of their genomes. As rhabdovirus genomes contain similar initiation and termination sequences within each gene, this repetitive genetic feature may facilitate the occurrence of homologous gene duplication and novel gene evolution within the family. In the case of BEFV and ARV, a nonstructural G protein (*Gns*), which lies directly downstream of the G protein, is believed to have been generated by homologous gene duplication of the G protein in an ancestral rhabdovirus (Walker et al., 1992; Wang and Walker, 1993). As the G and Gns proteins exhibit low levels of amino acid identity, and the Gns protein does not share characteristics of the G protein such as being incorporated into virions or inducing neutralizing antibodies in the host (Hertig et al., 1996; Johal et al., 2008), it is likely that Gns has undergone adaptive evolution and functional divergence after duplication, although its role in viral infection is unknown. While recent functional analysis of WONV has demonstrated that U3 is required for efficient viral replication, it is unclear whether U1 and U2 have similar roles to U3, and whether the three proteins may act synergistically. Similarly, whether the functions of U1–U3 are conserved throughout the viruses of the Hart Park serogroup remains to be determined.

**Molecular evolution of FLAV in the United States**

To explore the evolution of U1 in more detail, we analyzed 103 FLAV isolates from mosquitoes and birds collected annually over a 9-year period (2002–2010) in Georgia and over a 6-year period in Texas (2005–2010), as well as additional isolates from other states and older archived viruses dating back to the prototype FLAV isolate from Flanders, New York in 1961 (Table 2). Although our phylogenetic analysis revealed a low level of evolutionary change, with the vast majority of viruses falling into a single large clade (denoted as lineage A) (Fig. 5), the most notable result was the identification of a unique FLAV variant (termed lineage B), which demonstrated ~15% nucleotide divergence in U1 to lineage A. This variant lineage, which was first identified in 2005, appears to be localized to the lower coastal plain region of Georgia (Lowndes Co., Chatham Co.), and despite longitudinal in-state surveillance has never been found outside of this two-county area. Interestingly, both the prototypical FLAV (lineage A) and the variant (lineage B) appear to circulate sympatrically (Fig. 5), as they have been repeatedly isolated together from the same county (i.e., Lowndes) over a 6-year period (2005–2010). Despite such co-circulation in Georgia, it is also notable that all viruses sampled outside of Georgia fell into lineage A, as did all viruses sampled from 1961 to 1999. Although available data suggests that lineage B is transmitted primarily by the same mosquito species as other FLAV isolates (Table 2), the evolutionary factors that have driven this phylogenetic divergence in a sympatric set of viruses, such as switching to non-avian (more sedentary) hosts or to a mosquito-only cycle, are unknown and would require additional virus surveillance and serological surveying in the region. In this context, it is important to note that some *Culex* species (e.g., *Cx.

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**Fig. 3.** Newly proposed genomic configuration of FLAV demonstrating the U1–U3 region between the P and M genes and the SH ORF between the G and L genes. The genomic organization of the related *Wongabel* virus (WONV) is shown for comparison.

**Fig. 4.** Alignment of the U1–U3 proteins of the prototype isolate of FLAV (61-7484) showing amino acid identity indicative of gene duplication. Residues of identity are highlighted in blue. Asterisks, colons, and periods indicate identical, conserved, and semi-conserved residues, respectively, among the three proteins. Tryptic peptides of U1 and U3 identified by mass spectrometry are shown in bold italic.

| Isolate       | Host             | County     | State     | Year | Lineage | GenBank |
|--------------|------------------|------------|-----------|------|---------|---------|
| 61-7484      | Culiseta (Cs.) melanura | Suffolk    | New York  | 1961 | A       | KT028651 |
| C182         | Aegeains phoenicus | Unknown    | Unknown   | 1963 | A       | KT028675 |
| Ar 228-74    | Culex (Cx.) restuans | Unknown    | Connecticut | 1974 | A       | KT028663 |
| Ar 274-74    | Cs. melanura      | Unknown    | Connecticut | 1974 | A       | KT028676 |
| Ar 46-84     | Cs. restuans/Cs. melanuraa | Unknown    | Connecticut | 1984 | A       | KT028674 |
| Ar 77-84     | Cs. restuans/Cs. melanurab | Unknown    | Connecticut | 1984 | A       | KT028673 |
| RI 907-36    | Cs. melanura      | Westernly | Rhode Island | 1999 | A       | KT028677 |
| CLA 31-02    | Cx. spp.         | Clayton    | Georgia   | 2002 | A       | KT028662 |
| WV 382-02    | Sparrow spp.     | Jefferson  | West Virginia | 2002 | A       | KT028667 |
| CHC 1015-03  | Cs. melanura     | Chatham    | Georgia   | 2003 | A       | KT028665 |
| DN 133-03    | Cx. spp.         | DeKalb     | Georgia   | 2003 | A       | KT028678 |
| FTN 724-03   | Cx. quinquefasciatus | Fulton      | Georgia   | 2003 | A       | KT028715 |
| FTN 787-03   | Cx. quinquefasciatus | Fulton      | Georgia   | 2003 | A       | KT028730 |
| GWI 41-03    | Cx. spp.         | Gwinnett   | Georgia   | 2003 | A       | KT028724 |
| RCK 2-03     | Cx. spp.         | Rockdale   | Georgia   | 2003 | A       | KT028719 |
| WV 376-03    | Turdus migratorius | Raleigh    | West Virginia | 2003 | A       | KT028668 |
| CHC 948-04   | Cx. quinquefasciatus | Chatham    | Georgia   | 2004 | A       | KT028691 |
| CHC 1014-04  | Cx. quinquefasciatus | Chatham    | Georgia   | 2004 | A       | KT028731 |
| CHC 1216-04  | Cx. spp.         | Chatham    | Georgia   | 2004 | A       | KT028762 |
| CHC 1256-04  | Cx. quinquefasciatus | Chatham    | Georgia   | 2004 | A       | KT028762 |
| CHC 1315-04  | Cx. quinquefasciatus | Chatham    | Georgia   | 2004 | A       | KT028755 |
| CHC 1591-04  | Cx. quinquefasciatus | Chatham    | Georgia   | 2004 | A       | KT028743 |
| FTN 133-04   | Cx. quinquefasciatus | Fulton     | Georgia   | 2004 | A       | KT028692 |
| FTN 251-04   | Cx. quinquefasciatus | Fulton     | Georgia   | 2004 | A       | KT028742 |
| FTN 252-04   | Cx. quinquefasciatus | Fulton     | Georgia   | 2004 | A       | KT028757 |
| FTN 281-04   | Cx. quinquefasciatus | Fulton     | Georgia   | 2004 | A       | KT028704 |
| FTN 283-04   | Cx. quinquefasciatus | Fulton     | Georgia   | 2004 | A       | KT028761 |
| FTN 323-04   | Cx. quinquefasciatus | Fulton     | Georgia   | 2004 | A       | KT028783 |
| M 10567      | Cx. quinquefasciatus | Harris     | Texas     | 2005 | A       | KT028716 |
| M 11750      | Cx. quinquefasciatus | Harris     | Texas     | 2005 | A       | KT028664 |
| M 13419      | Cx. quinquefasciatus | Harris     | Texas     | 2005 | A       | KT028723 |
| CHC 302-05   | Cx. quinquefasciatus | Chatham    | Georgia   | 2006 | A       | KT028696 |
| CHC 3088-05  | Cs. melanura      | Chatham    | Georgia   | 2005 | A       | KT028709 |
| CLA 19-05    | Cx. quinquefasciatus | Clayton    | Georgia   | 2005 | A       | KT028694 |
| DKB 532-05   | Cx. restuans      | DeKalb     | Georgia   | 2005 | A       | KT028666 |
| FTN 178-05   | Cx. spp.         | Fulton     | Georgia   | 2005 | A       | KT028700 |
| GWI 78-05    | Cx. quinquefasciatus | Gwinnett  | Georgia   | 2005 | A       | KT028693 |
| LWN 1608-05  | Cs. salinarius    | Lowndes    | Georgia   | 2005 | B       | KT028720 |
| M 2876-06    | Cx. quinquefasciatus | Harris     | Texas     | 2006 | A       | KT028722 |
| M 3028-06    | Cx. quinquefasciatus | Harris     | Texas     | 2006 | A       | KT028721 |
| CHC 121-07   | Cx. quinquefasciatus | Chatham    | Georgia   | 2007 | A       | KT028687 |
| CHC 452-07   | Cx. quinquefasciatus | Chatham    | Georgia   | 2007 | A       | KT028687 |
| CHC 522-07   | Cx. quinquefasciatus | Chatham    | Georgia   | 2007 | A       | KT028708 |
| CHC 576-07   | Cx. quinquefasciatus | Chatham    | Georgia   | 2007 | A       | KT028701 |
| FTN 4-07     | Cx. quinquefasciatus | Fulton     | Georgia   | 2007 | A       | KT028707 |
| FTN 13-07    | Cx. quinquefasciatus | Fulton     | Georgia   | 2007 | A       | KT028686 |
| LWN 74-07    | Cx. quinquefasciatus | Lowndes    | Georgia   | 2007 | A       | KT028699 |
| LWN 81-07    | Cx. quinquefasciatus | Lowndes    | Georgia   | 2007 | A       | KT028698 |
| LWN 205-07   | Cx. quinquefasciatus | Lowndes    | Georgia   | 2007 | B       | KT028734 |
| M 16631      | Cx. quinquefasciatus | Harris     | Texas     | 2007 | A       | KT028712 |
| M 18684      | Cx. quinquefasciatus | Harris     | Texas     | 2007 | A       | KT028729 |
| CHC 306-08   | Cx. spp.         | Chatham    | Georgia   | 2008 | A       | KT028706 |
| CHC 363-08   | Cx. quinquefasciatus | Chatham    | Georgia   | 2008 | A       | KT028705 |
| CHC 441-08   | Cx. quinquefasciatus | Chatham    | Georgia   | 2008 | A       | KT028738 |
| CHC 561-08   | Cx. quinquefasciatus | Chatham    | Georgia   | 2008 | A       | KT028740 |
| CHC 1663-08  | Cx. spp.         | Chatham    | Georgia   | 2008 | A       | KT028735 |
| DKB 105-08   | Cx. quinquefasciatus | DeKalb   | Georgia   | 2008 | A       | KT028758 |
| DKB 270-08   | Cx. quinquefasciatus | DeKalb   | Georgia   | 2008 | A       | KT028732 |
| LWN 167-08   | Cx. quinquefasciatus | Lowndes    | Georgia   | 2008 | A       | KT028739 |
| LWN 171-08   | Cx. quinquefasciatus | Lowndes    | Georgia   | 2008 | B       | KT028733 |
| LWN 241-08   | Cx. quinquefasciatus | Lowndes    | Georgia   | 2008 | B       | KT028736 |
| LWN 325-08   | Cx. quinquefasciatus | Lowndes    | Georgia   | 2008 | A       | KT028744 |

Table 2: FLAV isolates recovered from 1961 to 2010 in the eastern United States that were analyzed during the study.
Materials and methods

Mosquito collection and virus isolation

Mosquitoes in Georgia, USA, were collected as part of a statewide arbovirus surveillance program using a variety of methods (CDC light traps, gravid traps), identified to the species level (when possible), and stored at −80 °C until further processing. Mosquito pools were mechanically homogenized in BA-1 media (Lanciotti et al., 2000), clarified by centrifugation (6700 × g for 10 min), and an aliquot (100 μl) was inoculated into confluent 2-day-old 4.0 cm² cultures of Vero E6 cells. Wells exhibiting cytopathology were harvested and RNA was extracted using a QiAamp Viral RNA Mini kit (Qiagen, Valencia, CA) and virus isolates were identified as FLAV by RT-PCR targeting the N gene (Nasci et al., 2001) using an AMV reverse transcriptase/Gotaq® Flexi DNA polymerase system (Promega, Madison, WI). Arbovirus surveillance in Texas, USA, was performed as described previously (Lillibridge et al., 2004). A small number of avian isolates of FLAV were included in the analysis (Table 2), and these were recovered from homogenized brain tissue of dead bird submissions using the methods described above. Archived FLAV isolates from 1961 to 1999 (Table 2) and the prototype strain of HPV from 1955 (Ar70, Culex tarsalis, Hart Park, Kern County, California, USA) were obtained from the World Reference Center for Emerging Viruses and Arboviruses (WRCVEA) at the University of Texas Medical Branch (UTMB).

U1–U3 gene and mRNA analysis

The pseudogene 1, 19 kDa gene, and pseudogene 2 sequences in a representative set of spatially and temporally discrete FLAV isolates, including the original prototype strain 61-7484 (GenBank accession KF028661–KF028670), were amplified by RT-PCR using primers designed from the original FLAV sequences (GenBank accession AH012179). The analogous region in HPV (GenBank accession KF028764) was amplified by designing primers based on highly conserved regions in FLAV. All pseudogene 1 (U1) sequences used in phylogenetic analysis (see below) have been submitted to GenBank under the accession numbers KF028671–KF028763. cDNA products of the transcripts of the pseudogene region were generated using an oligo(dT) primer and gene-specific primers based on the 5′-terminal mRNA sequence and cloned using a PCR Cloning kit (Qiagen, Valencia, CA). Primer sequences are available from the authors upon request.

Table 2 (continued)

| Isolate | Host | County | State | Year | Lineage | GenBank |
|---------|------|--------|-------|------|---------|---------|
| LWN 504-08 | Cs. melanura | Lowndes | Georgia | 2008 | B | KF028737 |
| M 27056 | Cs. quinquefasciatus | Harris | Texas | 2008 | A | KF028713 |
| M 29588 | Cs. quinquefasciatus | Harris | Texas | 2008 | A | KF028727 |
| CHC 109-09 | Cs. spp. | Chatham | Georgia | 2009 | A | KF028745 |
| CHC 368-09 | Cs. spp. | Chatham | Georgia | 2009 | A | KF028751 |
| CHC 622-09 | Cs. quinquefasciatus | Chatham | Georgia | 2009 | B | KF028684 |
| CHC 624-09 | Cs. quinquefasciatus | Chatham | Georgia | 2009 | B | KF028672 |
| DKB 160-09 | Cs. restuans | DeKalb | Georgia | 2009 | A | KF028747 |
| DKB 166-09 | Cs. quinquefasciatus | DeKalb | Georgia | 2009 | A | KF028749 |
| DKB 238-09 | Cs. restuans | DeKalb | Georgia | 2009 | A | KF028753 |
| DKB 252-09 | Cs. quinquefasciatus | DeKalb | Georgia | 2009 | A | KF028750 |
| DKB 74-09 | Cs. restuans | DeKalb | Georgia | 2009 | A | KF028746 |
| FTN 11-09 | Cs. spp. | Fulton | Georgia | 2009 | A | KF028754 |
| LWN 414-09 | Cs. quinquefasciatus | Lowndes | Georgia | 2009 | A | KF028748 |
| LWN 524-09 | Cs. quinquefasciatus | Lowndes | Georgia | 2009 | A | KF028752 |
| LWN 681-09 | Cs. quinquefasciatus | Lowndes | Georgia | 2009 | B | KF028678 |
| M 38933 | Cs. quinquefasciatus | Harris | Texas | 2009 | A | KF028726 |
| M 39509 | Cs. quinquefasciatus | Harris | Texas | 2009 | A | KF028714 |
| CHC 1169-10 | Cs. quinquefasciatus | Chatham | Georgia | 2010 | A | KF028670 |
| CHC 1217-10 | Cs. spp. | Chatham | Georgia | 2010 | A | KF028683 |
| DKB 318-10 | Cs. restuans | DeKalb | Georgia | 2010 | A | KF028685 |
| LWN 52-10 | Cs. restuans | Lowndes | Georgia | 2010 | B | KF028680 |
| LWN 47-10 | Cs. restuans | Lowndes | Georgia | 2010 | B | KF028679 |
| LWN 713-10 | Cs. quinquefasciatus | Lowndes | Georgia | 2010 | B | KF028682 |
| LWN 903-10 | Cs. quinquefasciatus | Lowndes | Georgia | 2010 | B | KF028681 |
| M 2735 | Cs. quinquefasciatus | Harris | Texas | 2010 | A | KF028728 |
| M 28263 | Cs. quinquefasciatus | Harris | Texas | 2010 | A | KF028717 |
| LOU 026-22 | Cs. pipiens/restuans | Loudoun | Virginia | 2010 | A | KF028669 |

* Precise identification not determined; information derived from Yale Arbovirus Research Unit annual reports.
**SDS-PAGE and immunoblotting**

FLAV mouse hyperimmune ascites fluid (MHIAF) was generated as described previously (Tesh et al., 1983) and carried out under an animal use protocol approved by the UTMB. Immunoblots were performed according to standard methods (Harlow and Lane, 1999). Vero cells were infected with FLAV at a multiplicity of infection (M.O.I.) of 1, trypsinized at day 3 post-infection, and pelleted by light centrifugation (4300 g for 15 min). The cell pellet was washed 2X in PBS and then lysed in RIPA buffer (50 mM Tris–HCl, pH 7.4, 150 mM NaCl, 1% NP-40, 0.1% SDS, 0.5% sodium deoxycholate, 1 mM EDTA). Insoluble protein was removed by centrifugation (6700 g for 10 min) and the lysate was mixed with 5X Laemmli sample buffer (250 mM Tris–HCl, pH 6.8, 25% β-mercaptoethanol, 10% SDS, 50% glycerol, 0.05% bromophenol blue) and boiled for 5 min. Proteins were electrophoresed by SDS-PAGE in a 10% or 12% polyacrylamide gel and transferred to 0.45 μm nitrocellulose. The membrane was blocked with 5% dry milk in TBS-0.05% Tween and probed using a 1:100 dilution of FLAV MHIAF and a 1:2000 dilution of a goat anti-mouse IgG (H+L).

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**Fig. 5.** Evolutionary relationships among 103 U1 nucleotide sequences of FLAV depicting the two main viral lineages (A) and (B). To illustrate the sympatric co-circulation of the two lineages, those viruses sampled from Lowndes County, Georgia, are shown in bold italics. All horizontal branches are scaled according to the number of nucleotide substitutions per site, and bootstrap support values are shown for key nodes.
HRP conjugate (Jackson ImmunoResearch Laboratories, West Grove, PA). Viral protein sizes were estimated against a SuperSignal™ Molecular Weight Protein Ladder (ThermoScientific, Waltham, MA) and protein–antibody complexes were detected using a SuperSignal™ West Pico Chemiluminescent Substrate Kit (ThermoScientific). Blots were analyzed using a ChemiDoc™ MP imaging system (BioRad, Hercules, CA).

Virus purification and tandem mass spectrometry

To obtain viral proteins for mass spectrometry, large-scale purification of FLAV was performed. Briefly, confluent Vero MARU cell cultures were grown in 850 cm² roller bottles (Corning Inc., Corning, NY) and infected with FLAV at an M.O.I. of ~1. Supernatant was harvested at day 4 post-infection, clarified by low-speed centrifugation at 4400 × g for 30 min, and virus was precipitated overnight at 4°C with 7% polyethylene glycol (PEG) and 2.3% NaCl. Virus was pelleted by centrifugation at 13,000 × g for 1 h and the pelleted was resuspended in TES buffer (10 mM Tris-Cl, pH 7.4, 2 mM EDTA, 150 mM NaCl) and centrifuged (13,000 × g, 15 min) to remove the PEG. Virus was then purified on a 20% sucrose cushion followed by a 20–60% sucrose gradient in a Beckman SW 32 Ti rotor at 134,000 × g for 2 h at 4°C using an Optima™ L-100K Ultracentrifuge (Beckman Coulter, Brea, CA). The virus band was recovered, loaded on an Amicon™ Ultra-15 100K centrifugal filter unit for concentration and to remove low molecular weight proteins (Millipore, Billerica, MA), and subjected to SDS-PAGE as previously noted except that the gel was stained with a SYPRO Ruby Protein Gel Stain (Molecular Probes, Invitrogen, Carlsbad, CA).

Proteins in the gel were visualized using an UV transilluminator and a band corresponding to the approximate size of the accessory proteins of interest (U1–U3, SH; ~10–20 kDa) was cut from the gel. Nano-scale high performance liquid chromatography coupled to tandem mass spectrometry (nano HPLC–MS/MS) was performed as described previously (Hochrainer et al., 2012). Briefly, SYPRO Ruby-stained proteins were destained, reduced using dithiothreitol (10 mM), alkylated with iodoacetamide (55 mM), and digested overnight with trypsin (0.5 μg). Tryptic peptides were collected by centrifugation (4000 × g, 2 min) and the remaining peptides in the gel were sonicated in 50% acetonitrile-5% formic acid and collected. Tryptic peptides were pooled, evaporated in a Speedvac SC110 (Thermo Savant, Milford, MA, USA), reconstituted in 2% acetonitrile–0.1% trifluoroacetic acid, and analyzed with nano HPLC–MS/MS using an LTQ–Orbitrap Elite mass spectrometer (Thermo-Fisher Scientific, San Jose, CA). Proteins were identified by searching MS/MS spectra using the Mascot Daemon search engine (version 2.3.02, Matrix Science, Boston, MA). Proteins identiﬁed by searching MS/MS were subjected to Mascot search in Mascot.

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