- 1 <u>Title</u> Discovery and Genomic Characterization of a Novel Henipavirus, Angavokely virus, from
- 2 fruit bats in Madagascar.
- 3
- 4 <u>Running Title</u> Novel Henipavirus, AngV, found in Madagascar fruit bat
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35 Abstract

36 37	The genus <i>Henipavirus</i> (family <i>Paramyxoviridae</i>) is currently comprised of seven viruses, four of which have demonstrated prior evidence of zoonotic capacity. These include the biosafety level
38	4 agents Hendra (HeV) and Nipah (NiV) viruses, which circulate naturally in pteropodid fruit
39	bats. Here, we describe and characterize Angavokely virus (AngV), a divergent henipavirus
40	identified in urine samples from wild, Madagascar fruit bats. We report the near-complete
41	16,740 nt genome of AngV, which encodes the six major henipavirus structural proteins
42	(nucleocapsid, phosphoprotein, matrix, fusion, glycoprotein, and L polymerase). Within the
43	phosphoprotein (P) gene, we identify an alternative start codon encoding the AngV C protein
44	and a putative mRNA editing site where the insertion of one or two guanine residues encodes,
45	respectively, additional V and W proteins. In other paramyxovirus systems, C, V, and W are
46	accessory proteins involved in antagonism of host immune responses during infection.
47	Phylogenetic analysis suggests that AngV is ancestral to all four previously described bat
48	henipaviruses—HeV, NiV, Cedar virus (CedV), and Ghanaian bat virus (GhV)—but evolved more
49	recently than rodent- and shrew-derived henipaviruses, Mojiang (MojV), Gamak (GAKV), and
50	Daeryong (DARV) viruses. Predictive structure-based alignments suggest that AngV is unlikely to
51	bind ephrin receptors, which mediate cell entry for all other known bat henipaviruses.
52	Identification of the AngV receptor is needed to clarify the virus's potential host range. The
53	presence of V and W proteins in the AngV genome suggest that the virus could be pathogenic
54	following zoonotic spillover.
55	
56	
57	
58	
59	Importance
60	Henipaviruses include highly pathogenic emerging zoonotic viruses, derived from bat, rodent,
61	and shrew reservoirs. Bat-borne Hendra (HeV) and Nipah (NiV) are the most well-known
62	henipaviruses, for which no effective antivirals or vaccines for humans have been described.
63	Here we report the discovery and characterization of a novel henipavirus, Angavokely virus
64	(AngV), isolated from wild fruit bats in Madagascar. Genomic characterization of AngV reveals
65 66	all major features associated with pathogenicity in other henipaviruses, suggesting that AngV
66 67	could be pathogenic following spillover to human hosts. Our work suggests that AngV is an
67 68	ancestral bat henipavirus which likely uses viral entry pathways distinct from those previously described for HeV and NiV. In Madagascar, bats are consumed as a source of human food,
69	presenting opportunities for cross-species transmission. Characterization of novel henipaviruses
70	and documentation of their pathogenic and zoonotic potential are essential to predicting and
70	preventing the emergence of future zoonoses that cause pandemics.
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72 Introduction

73 Henipaviruses (HNVs) belong to a genus of bat-, rodent-, and shrew-borne viruses within the 74 family Paramyxoviridae with demonstrated zoonotic potential. HNVs can manifest extreme 75 virulence in human hosts, as exemplified by the prototypical HNVs, Hendra virus (HeV), and 76 Nipah virus (NiV), which cause severe acute respiratory distress and/or encephalitis in humans. 77 yielding case fatality rates that can exceed 90% (1–3). This high pathogenicity and the lack of approved HNV therapeutics or vaccines for humans have garnered HeV and NiV classification as 78 79 Biological Safety Level 4 (BSL4) agents and WHO priority diseases. Since their discovery in the 1990s, HeV and NiV have periodically spilled over to humans from their reservoir hosts, 80 pteropodid bats. HeV zoonosis is mediated by spillover to intermediate horse hosts, from which 81 82 humans acquire infection(4). NiV can spillover to humans via intermediate transmission 83 through pig hosts, or directly from bat-to-human, resulting in near-annual outbreaks of fatal 84 encephalitis in South Asia, where subsequent human-to-human transmission also occurs (2, 5– 85 7). 86

87 Novel HNVs continue to emerge from wildlife hosts and represent ongoing threats to human

- 88 health. Initially, the *Henipavirus* genus comprised only HeV and NiV; however, the past two
- 89 decades have witnessed the discovery of five new HNVs: bat-borne Cedar virus (CedV) and
- 90 Ghanaian bat virus (GhV), rodent-borne Mojiang virus (MojV), and shrew-borne Gamak (GAKV)
- 91 and Daeryong viruses (DARV) (8–11). Of these novel HNVs, at least two show evidence of
- 92 zoonotic potential: serological data suggests prior human exposure to GhV in West Africa (12),
- 93 while MojV was first identified following a human outbreak of severe pneumonia in Chinese
- 94 mine workers, all of whom died after infection(9). In addition to their high potential for
- 95 pathogenicity, HNVs possess a broad host range that spans at least seven mammalian orders,
- 96 including bats (10, 13).
- 97

98 Cross-species viral spillover necessitates effective inter-species transmission, which first 99 requires a virus to successfully enter the cells of diverse host species. In general, HNVs use the 100 highly-conserved ephrin family of proteins, both type A and type B, as cell entry receptors (1, 8, 101 14–16). A notable exception to this pattern is MojV, which does not use ephrin proteins—or the 102 sialic acid and CD150 receptors common to non-HNV paramyxoviruses—to gain cell entry (14, 103 17). Indeed, as of yet, the viral entry receptor for MojV—and the closely related GAKV and 104 DARV—remain unknown. In general, viruses in the genus *Henipavirus* have broad host ranges 105 and cause high case fatality rates following human spillover, making the characterization of new 106 HNVs a high public health priority.

107

108 The HNV genome consists of six structural proteins: nucleocapsid (N), phosphoprotein (P), 109 matrix (M), fusion (F), glycoprotein (G), and polymerase (L). In comparison with other members 110 of the family Paramyxoviridae, HNVs have relatively larger genomes (approximately 18kb vs 111 16kb). This extended length is largely due to several, long 3' untranslated regions (UTR) of the 112 N, P, F and G mRNAs (18, 19). The genome length of HNVs, like all paramyxoviruses, adheres to 113 the so-called 'Rule of Six', whereby viral genomes consistently demonstrate polyhexameric 114 length (20). The 'Rule of Six' is believed to be a requirement for efficient genome replication 115 under the unique mRNA editing features of the paramyxovirus genome (20). The paramyxovirus 116 P locus exhibits notable transcription properties that are shared across most members of the 117 Paramyxoviridae family. The P gene permits the translation of additional accessory proteins 118 from either gene editing events within the locus (prior to translation) or an overlapping ORF in 119 the P gene itself. All HNVs, with the exception of CedV, harbor a highly conserved mRNA editing 120 site at which the insertion of additional guanine residues can result in the translation of 121 accessory proteins, V and W, involved in viral antagonism and evasion of the host immune 122 system (1). The HNV P gene also contains an overlapping ORF that allows for the synthesis of a 123 third accessory protein, C, which is also involved in viral host immune evasion (1).

124

125 Our lab has previously demonstrated evidence of exposure to henipa-like viruses in serum

126 collected from three endemic Madagascar fruit bat species (*E. dupreanum, Pteropus rufus,* and

127 *Rousettus madagascariensis*) using a Luminex serological assay which identified cross reactivity

to CedV/NiV/HeV-G and -F proteins (21). The most significant antibody binding previously

129 detected corresponded to the NiV-G antigen for *E. dupreanum* serum and the HeV-F antigen for

130 *P. rufus* and *R. madagascariensis* serum, suggesting the potential circulation of multiple HNVs in

131 the Madagascar fruit bat system (21). Fruit bats, including *E. dupreanum*, are consumed widely

in Madagascar as a source of human food, presenting opportunities for cross-species zoonotic

133 emergence. This underscores the importance of further characterization of the pathogenic and

134 zoonotic potential of AngV and other potential HNVs circulating in the Madagascar fruit bat

135 system. Here, we describe and characterize a novel bat HNV, Angavokely virus (AngV),

136 recovered from urine samples collected from the Madagascar fruit bat, *E. dupreanum*. Our

137 work suggests AngV is part of an ancestral group of HNVs that may rely on a novel, non-ephrin-

- 138 mediated viral entry pathway.
- 139

140 Methods

141 Ethics Statement

142 Animal capture and handling and subsequent collection of biological samples were conducted

143 in strict accordance with the Madagascar Ministry of Forest and the Environment (permit

numbers 019/18, 170/18, 007/19) and guidelines posted by the American Veterinary Medical

145 Association. Field protocols were approved by the UC Berkeley Animal Care and Use Committee

- 146 (ACUC Protocol # AUP-2017-10-10393), as previously described (22).
- 147
- 148 Animal capture, sample collection, and RNA extraction
- 149 Fruit bats were captured and processed in part with a long-term study investigating the
- 150 seasonal dynamics of potentially zoonotic viruses in Madagascar, as has been previously
- 151 described (21–25). Animals were identified morphologically by species, sex, and age class
- 152 (juvenile vs. adult), and urine swabs were collected into viral transport medium from any
- 153 individual that urinated during handling. Urine swabs were flash-frozen in liquid nitrogen in the
- 154 field and delivered to -80°C freezers at Institut Pasteur de Madagascar for long-term storage.
- 155 Urine specimens from 206 bats were randomly selected for total RNA extraction using the Zymo
- 156 Quick RNA/DNA Microprep Plus kit, performed as previously described (22).
- 157
- 158 mNGS library preparation
- 159 Total urine RNAs were diluted with nuclease-free H₂O, and 5uL of each specimen was used as
- 160 input for mNGS library preparation. A 2-fold dilution series of a 25ng/uL stock of HeLa total RNA
- 161 (n=8 samples), along with 5 water samples were included and processed in parallel as positive
- and negative controls, respectively. Additionally, a 25pg aliquot of External RNA Control
- 163 Consortium (ERCC) spike-in mix (Thermo-Fisher) was included in each sample. Dual-indexed
- 164 mNGS library preparations for the samples were miniaturized and performed in 384-well
- 165 format with NEBNext Ultra II RNAseq library preparation kit (New England Biolabs) reagents.
- 166 RNA samples were fragmented for 12 min at 94°C, and 16 cycles of PCR amplification were
- 167 performed. Per sample read yields from a small scale iSeq (Illumina) paired-end 2 x 146bp
- 168 sequencing run on an equivolume pool of the individual libraries were used to normalize
- volumes of the individual mNGS libraries to generate an equimolar pool. Paired-end 2 x 146bp
- sequencing of the resulting equimolar library pool was performed on the NovaSeg6000
- 171 (Illumina) to obtain approximately 50 million reads per sample.
- 172

173 Sequence analysis

- 174 Raw reads from urine sample sequencing were first uploaded to the CZBID (v6.8) platform for
- host and quality filtering and *de novo* assembly (26). In brief, in the CZBID pipeline, adaptor
- sequences were removed with Trimmomatic (v.0.38), and reads were quality filtered (27).
- 177 Reads then underwent host-filtration against the Malagasy fruit bat genome, *E. dupreanum*,
- using STAR (v 2.7.9a) (28) and a second host-removal step using Bowtie2 (29). After host
- 179 filtering, reads were aligned using rapsearch2 (30) and Gsnap (31), and putative pathogen taxa
- 180 were identified. Next, reads were assembled using SPADES (v.3.15.3) (32), and all contigs
- 181 generated were subject to BLAST analysis against the putative taxa previously identified by

- 182 rapsearch2 and Gsnap. We considered samples positive for HNV if the CZBID pipeline produced
- 183 at least one contig with an average read depth of two or more, which yielded a BLAST
- alignment length >100 nt/aa and an e-value < 0.00001 (BLASTn v2.5.0+) or a bit score >100
- 185 (BLASTx v2.5.0+) when queried against an HNV database derived from all HNV genomes
- 186 available in NCBI (Accessed July 2021).
- 187
- 188 Genome Annotation and Comparison
- 189 One urine sample, collected from an adult female *E. dupreanum* fruit bat in March 2019,
- 190 yielded a near full-length HNV genome, which we analyzed in greater depth in subsequent
- analyses and annotated as the novel HNV, AngV (Genbank Accession #: ON613535). Nucleotide
- 192 BLAST of the AngV genome identified NiV (GenBank Accession #: AF212302) as the top hit for
- this novel virus and was subsequently chosen as the reference genome for further analysis. We
- aligned AngV to NiV (GenBank Accession #: AF212302) in the program Geneious Prime
- 195 (v2020.2.4) and annotated all six major HNV structural genes, and the accessory C ORF, within
- the P gene. We identified the putative mRNA editing site within the P gene sequence (spanning
- 197 nucleotides 1,225-1,232 of the P gene) and manually added one or two guanine (G) residues to
- the 3' end of the conserved HNV mRNA editing site to generate V and W ORFs, respectively,
- and their corresponding proteins. We furthered queried all identified transcriptional elements
- against publicly available sequences using NCBI BLAST and BLASTx (33). Resulting BLAST and
- 201 BLASTx hits were used in phylogenetic analyses as described below.
- 202

We used the program pySimPlot to scan the whole genome sequence of AngV for nucleotide
 sequence identity to the NiV genome (GenBank Accession #: AF212302) and the nucleotide and

amino acid sequences of individual Open Reading Frames (ORFs) contained therein.

- 206 Respectively, window size and scanning were specified as 50 and one for nucleotide pairwise
- identity and 50 and five for amino acid pairwise identity. Results were visualized using Prism(9.2.0).
- 209

210 Phylogenetic analyses

- 211 We constructed 10 Maximum Likelihood (ML) phylogenetic trees to analyze the evolutionary
- 212 relatedness of our putative HNV to previously described paramyxoviruses. These included (a)
- 213 one amino acid L protein phylogeny comparing the L protein of AngV to all reference L protein
- 214 paramyxovirus sequences in NCBI, in addition to the newly-described shrew HNV, GAKV and
- 215 DARV (accessed November 2021) and (b) nine amino acid phylogenies comparing each
- 216 individual protein annotated in AngV (N, P, C, V, W, M, F, G, L) against the top 50 BLASTx
- sequence hits for each protein collapsed on 98% sequence similarity. Distinct outgroups were
- applied: (a) Sunshine Coast Virus (GenBank Accession #: YP_009094051.1) for L protein, (b)

- 219 Human orthopneumovirus (HRSV, GenBank Accession #: NC_001781) for N, P, M, F, G, L
- proteins, and (c) Sendai virus (GenBank Accession #: NP_056872) for gene C.

221 For each phylogenetic tree, we aligned sequences via the MUSCLE algorithm (v3.8.1551) (34)

- and determined the best fit nucleotide or amino acid substitution model for using ModelTest-
- NG (35). Phylogenies were then constructed in RAxML-NG (36), using the corresponding best fit
- 224 model: JTT (complete L-protein sequence) or LG+G4+F (individual proteins). In accordance with
- best practices outlined in the RAxML-NG manual, twenty ML inferences were made on each
- original alignment. Bootstrap replicate trees were inferred using Felsenstein's method (37).
- 227 MRE-based bootstopping test was applied after every 200 replicates (38), and bootstrapping
- 228 was terminated once diagnostic statistics dropped below the threshold value. Bootstrap
- support values were drawn on the best-scoring tree.

230 We additionally computed one Bayesian time-resolved phylogeny, using all 77 full-length HNV

- 231 nucleotide sequences available in NCBI, including our newly contributed AngV (GenBank
- Accession #: ON613535). As with ML trees, sequences were first aligned in MUSCLE (v3.8.1551)
- 233 (34), and the best fit nucleotide substitution model was subsequently queried in ModelTest-NG
- 234 (35). We then constructed a Bayesian timetree in the program BEAST2 (39, 40), using the best
- fit GTR+I+G4 model inferred for the whole genome alignment from ModelTest-NG and
- assuming a constant population prior. Sampling dates corresponded to collection data as
- reported in NCBI Virus; we assumed a collection date of 31-July in cases where only year of
- collection was reported. We computed trees using both an uncorrelated exponentially
- distributed relaxed molecular clock (UCED) and a strict clock but here report results from the
- 240 strict clock only as similar results were inferred from both. We ran Markov Chain Monte Carlo
- 241 (MCMC) sample chains for 1 billion iterations, checked convergence using TRACER v1.7 (41) and
- averaged trees after 10% burn-in using TreeAnnotator v2.6.3 (42) to visualize mean posterior
- 243 densities at each node. The resulting phylogeny was visualized in R v.4.0.3 for MacIntosh in the
- 244 'ggtree' package (43).
- 245
- 246 AngV G Protein Structure Modeling

We used the Artificial Intelligence system, AlphaFold, to predict the 3D structure of the AngV
glycoprotein (G) (44). Molecular graphics and analyses of the AngV glycoprotein structure were
performed with UCSF ChimeraX (45). HNV glycoprotein ephrin binding residues were aligned
using the program Geneious Prime (v2020.2.4).

- 251
- 252 Results
- 253 Discovery and prevalence of HNV in Malagasy bats

- 254 Urine swab specimens from 206 bats were collected in 8 roosting sites across the island of
- 255 Madagascar from 2013 to 2019 (Figure 1). Urine samples were collected during wet and dry
- 256 seasons from all three Madagascar fruit bat species, *P. rufus, E. dupreanum*, and *R.*
- 257 madagascariensis (Table 1). Isolated RNA from urine swab specimens generated an average of
- 258 19 million paired-read sequences. In total, 10/206 (4.9%) bats were positive for HNV; all
- positive samples were collected from *E. dupreanum* bats (10/106; 9.4%) at the Angavokely cave
- 260 roosting site (Table 1). Positive samples were collected in wet and dry seasons from both male
- and female adults.
- 262

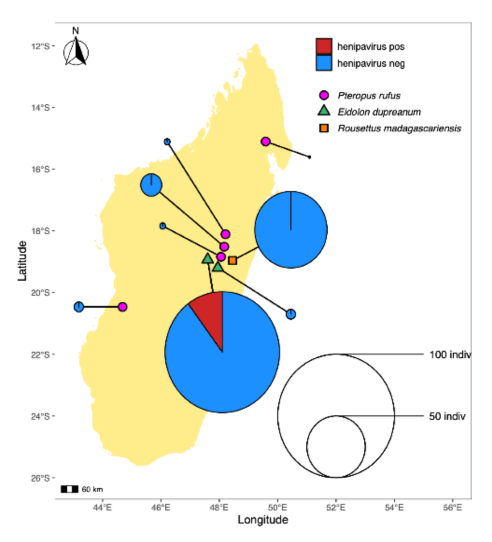




Figure 1. Geographic location of sampling sites used in this study. Sampling sites grouped by bat species found
depicted as follows *P. rufus* (pink circles) Ambakoana (-18.51 S, 48.17 E) / Mahabo (-20.46 S, 44.68 E) /
Mahialambo (-18.11 S, 48.21 E) / Makira (-15.11 S, 49.59 E) / Marovitsika (-18.84 S, 48.06 E) roosts; *E. dupreanum*(green triangles) Angavobe (-18.94 S, 47.95 E) /Angavokely (-18.93 S, 47.76 E) caves; *R. madagascariensis* (orange
squares) Maromizaha cave (-18.96 S, 48.45). Pie charts are size-weighted by total bat population sampled at each
site, corresponding to the legend. The percentage of HNV positive samples is shown for all sampled species and
sites. HNV positive samples were only recovered from the *E. dupreanum* Angavokely site.

271

272 Genomic characterization of AngV

273 We recovered one near-full-length HNV contig (16,740 nt), supported by an average sequencing

depth of 14 reads (Figure 2A), from a urine sample collected from an adult, non-lactating *E*.

275 *dupreanum* female in the 2018-2019 wet season (capture date: 15-March 2019). We focused

276 subsequent genomic analyses on this longest sequence, which we named Angavokely virus

277 (AngV) after the site of *E. dupreanum* capture.

278

As with other members of the *Henipavirus* genus, the genome of AngV is organized into 6 open

280 reading frames (ORF) arranged in the order 3'-N-P-M-F-G-L-5'. AngV shares an average

nucleotide identity of 36% with the NiV reference genome (AF212302) and a varying amino acid

identity that is highest across the ORFs encoding for the nucleocapsid and L polymerase

283 proteins (Figures 2B, 2C).

284

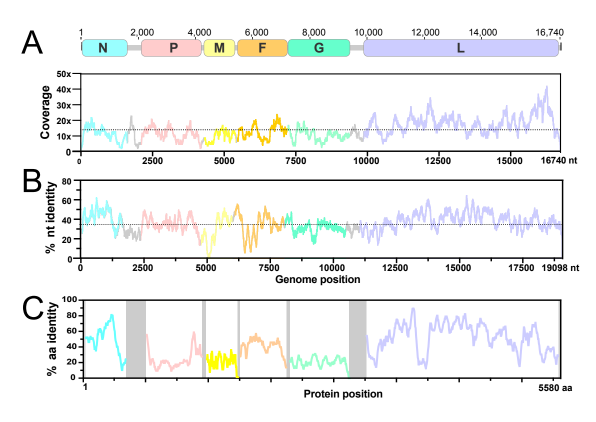




Figure 2. AngV genome organization. A. Coding regions for each gene are shown and depicted in color, non-coding
 intergenic and terminal regions are highlighted in gray. Depicted genes represented as follows: nucleocapsid (N),

RNA polymerase (P and L), matrix (M), fusion (F), and glycoprotein (G). Sequencing read depth supporting each
 position of the recovered genome sequence is plotted below the genomic schematic. Scanning nucleotide (B) and
 amino acid (C) pairwise identity to Nipah virus (GenBank Accession #: AF212302). Dotted horizontal lines represent
 average read depth (14.29) or average nucleotide pairwise identity (36%).

292

293 AngV coding regions

- 294 The P gene of AngV follows an organization similar to most members of the *Henipavirus* genus.
- 295 AngV harbors alternative start sites which, respectively, encode the P and C proteins, as well as
- a conserved putative mRNA editing site common to most paramyxoviruses A⁴⁻⁶G²⁻³ (Figure 3A
- and B). AngV shares an identical putative mRNA editing site with the recently discovered HNVs,
- 298 MojV and GAKV. Pseudotemplated addition of one or two G residues at the conserved putative
- 299 mRNA editing site generates a putative V and W protein, respectively (Figure 3C; W protein in
- 300 Supplemental Figure 1). In congruence with members of the *Henipavirus* genus that encode the
- 301 conserved putative mRNA editing site, the putative V protein of AngV harbors a unique C-
- 302 terminal region that contains a highly conserved cysteine-rich zinc finger domain (Figure 3C).
- 303
- 304

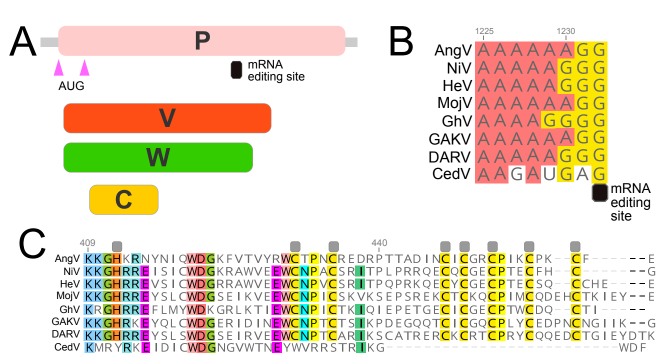


Figure 3. Organization of the P gene of AngV. A. Alternative transcriptional start sites (pink triangle) generate the P and C protein. Pseudotemplated addition of one or two guanine nucleotides at the putative mRNA editing site generates a V and W protein, respectively. B. Sequence alignment of the putative mRNA editing site across
 members of the *Henipavirus* genus (cRNA depicted). C. Amino acid alignment of the unique C terminal region of

310 the V protein following the addition of one guanine nucleotide to the putative mRNA editing site. Gray boxes

311 denote conserved cysteine and histidine residues suggested to directly coordinate bound zinc ions (54). Individual 312 nucleotides or amino acids are color coordinated if at least 75% conserved at the alignment position. Nucleotide or 313 amino acid position numbers displayed represent the position within the AngV gene or protein. Virus name 314 (abbreviation), followed by GenBank Accession #: Angavokely virus (AngV) ON613535; Nipah virus (NiV) AF212302; 315 Hendra virus (HeV) AF017149; Mojiang virus (MojV) KF278639; Ghanaian bat Henipavirus (GhV) HQ660129; 316 Daeryong virus (DARV) MZ574409; Gamak virus (GAKV) MZ574407, Cedar virus (CedV) JQ001776. CedV is shown 317 here only for comparison, as the CedV P protein is not believed to undergo RNA editing or to generate a functional 318 V protein (8, 54).

- 319
- 320 The length of each ORF in the AngV genome resembles those from previously described HNVs,
- with the exception of the gene encoding the glycoprotein (G)—which, at 688 aa, is 56 aa longer
- than the longest previously-characterized HNV glycoprotein from GhV (632 aa; Table 2) (15).
- 323 BLAST analysis indicates that AngV ORFs for genes encoding the nucleocapsid (N), matrix (M),
- 324 and polymerase (L) proteins exhibit the highest nucleotide and amino acid pairwise identity
- with other HNVs, with highest similarity shared with the NiV L protein (nt 74.7%, aa 52.2%,
- Table 2). In contrast to many emerging viruses, AngV largely exhibits higher nucleotide vs.
- 327 amino acid identity with other HNVs (Table 2). The more recently discovered HNVs (MojV,
- 328 CedV, GAKV, DARV, GhV) mirror this pattern, showing higher nucleotide vs. amino acid identity
- 329 when compared to NiV and HeV (data not shown).
- 330

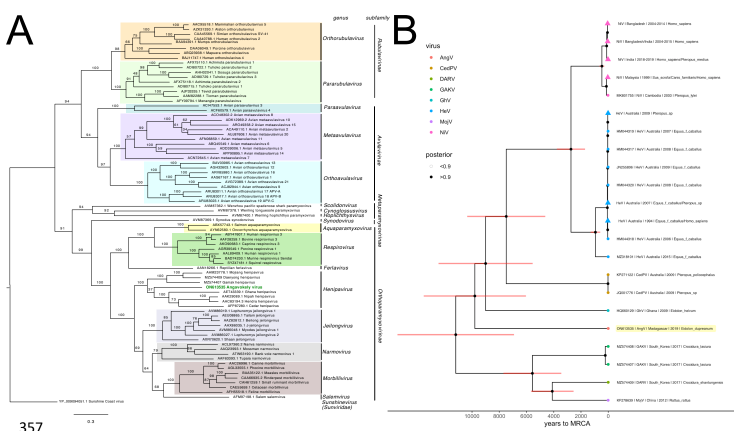
331 AngV non-coding regions

Examination of all viral intergenic regions (in cRNA orientation) reveals that AngV exhibits the 332 highly conserved CTT intergenic junction site characteristic of other HNVs, as well as gene stop 333 and gene start sites with high similarity to those of previously described HNVs (Supplemental 334 335 Table 1). We were unable to locate the intergenic junction site and transcriptional start or stop site in the 5' region of the N ORF for AngV, suggesting that the genomic 3' untranslated regions 336 (UTR) for AngV have not yet been fully recovered. Comparison of the 5' and 3' UTRs for AngV 337 338 with those of other HNVs reveals UTRs of varying lengths within the *Henipavirus* genus 339 (Supplemental Table 2). Nevertheless, AngV exhibits similar lengths and a nucleotide identity of 340 roughly 30-40% for the 5' and 3' UTRs of most HNV genes; however, the P gene 3' UTR, the M 341 gene 5' and 3' UTRs, and the F gene 5' and 3' UTRs, are significantly shorter in AngV compared 342 with previously described HNVs. Correspondingly, nucleotide identity varies when comparing 343 this shorter subset of 5' and 3' UTRs for AngV against other HNVs (Supplemental Table 2).

344

345 Phylogenetic Analyses

346 Phylogenetic analysis of complete L protein amino acid sequences across the Paramyxoviridae 347 family places AngV within the *Henipavirus* genus at <0.82 nucleotide substitutions away from the node distinguishing the family Paramyxoviridae from the Sunviridae (Figure 4A). AngV 348 349 clusters independently within the *Henipavirus* genus and diverges ancestral to all currently known bat-borne HNVs. Our time-resolved Bayesian phylogeny further corroborates this result, 350 placing AngV ancestral to all previously described bat-borne HNVs but more recently diverged 351 352 than the rodent- and shrew-borne HNVs, MojV, GAKV, and DARV (Figure 4B; Supplemental Figure 2). We estimate the divergence of the AngV lineage from the rest of the HNV clade at 353 9,794 years ago (95% HPD 6,519-14,024 years), and the time to the Most Recent Common 354 355 Ancestor (MRCA) for the entire HNV genus as 11,195 years ago (95% HPD 7,351-15,905 years). 356





Tree is rooted with Sunshine Coast Virus (GenBank Accession #: YP_009094051.1) as an outgroup, with outgroup branch length shrunk for ease of viewing. Novel HNV, AngV, is depicted in green. Subfamilies and genera are

361 demarcated, excluding those unassigned to subfamily (genera Scolidonvirus, Cynoglossusvirus, Hoplichthysvirus). 362 Bootstrap support is depicted and GenBank Accession numbers displayed next to virus names. Scale bar represents 363 substitutions per site. B. Time-resolved Bayesian phylogeny computed in BEAST2 incorporating all available 364 Henipavirus whole genome nucleotide sequences, with the addition of newly discovered GAKV, DARV, and AngV. 365 Closely-related sequences are collapsed at triangle nodes for NiV and HeV (phylogeny with un-collapsed branches 366 available in Supplemental Figure 2). 95% HPD intervals around the timing of each branching node are visualized as 367 red horizontal bars. Posterior support >.9 is indicated by black coloring of the corresponding node, and distinct 368 Henipavirus species are indicated by colored tip points, with AngV highlighted in yellow for further emphasis. The 369 estimated time to MRCA for Angavokely virus and the previously-described bat-borne HNVs is 9,794 (95% HPD 370 6,519 – 14,025) years ago.

371

372	In N, P, C, V, W, M and G amino acid phylogenies, the AngV proteins cluster closely with those
373	of other HNVs (Supplemental Figure 3). Interestingly, in the amino acid phylogeny, the AngV F
374	protein, like the F proteins of MojV, GAKV, and DARV, localizes ancestral to non-HNV
375	paramyxoviruses and distinct from the bat-borne HNV clade (Supplemental Figure 3). The AngV
376	L protein shows the highest amino acid identity to the L protein of rodent-borne Mount Mabu
377	Lophuromys virus 2 (MMLV-2), a putative Jeilongvirus (46), but is nonetheless nested between

378 the MojV/GAKV/DARV clade and the bat-borne HNV clade (Supplementary Figure 3).

379

380 AngV glycoprotein

381 We further examined the AngV G protein for conserved structural features and amino acid

residues historically associated with HNV ephrin binding. AlphaFold analysis revealed a six-

383 bladed β-propeller fold that is characteristic of *Paramyxoviridae* glycoproteins, with each blade

largely composed of 4 antiparallel β -strands (Figure 5A). The β -propeller fold is stabilized by

385 seven disulfide bonds that are conserved among HNVs (Supplemental Figure 4). This HNV

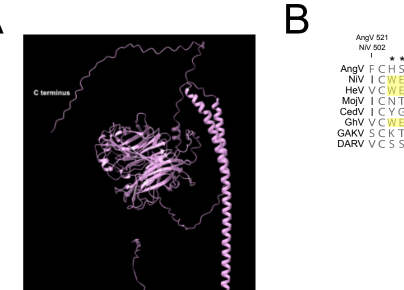
protein G structure-based alignment reveals that the elongated AngV G protein primarily

results from a lengthy C-terminal tail with an additional 67 aa beyond that of NiV G protein

388 (Supplemental Figure 4). Similar to the MojV G protein, the AngV G protein lacks previously

described ephrin binding residues (NiV W504, E505, T531, A532, E533, N557, and Y581) (Figure

- 390 5B and Supplemental Figure 4) (16, 47, 48).
- 391



AngV 521	AngV 548	AngV 575	AngV 599
NiV 502	NiV 529	NiV 556	NiV 580
۱ * *	I	*	' *
Anal/ FCHSSC	$v c \dot{u} c \dot{v} \dot{c}$		
AngV F C H S S G			
Niv I C W E G V	Y N <mark>Q T A E</mark>		Q 🛛 🏹 🛛
HeV V C <mark>W E</mark> G T	Y N <mark>Q T A E</mark>	T N A (Q 🛛 🏹 🛛
MojVICNTRG	Y ··· N N G G T	· Y S I '	T G K
CedV CYGGT	Y … D <mark>Q L A E</mark>	L <mark>N</mark> T I	R T N I
GhV V C <mark>W E</mark> G T	Y E <mark>Q V A E</mark>	S S A I	R T
GAKV SCKTWN	F····KTGNS	QSI	G G V
DARV V C S S Y G	Y … N G E G T	F K I	G Q

392 393

Figure 5. AlphaFold-predicted AngV glycoprotein 3D structure and ephrin binding residue sequence alignment. A. 394 AlphaFold-predicted 3D structure of AngV glycoprotein. N and C termini are indicated in white. B. Alignment of 395 HNV ephrin binding residues. The position of previously-described HNV ephrin binding residues are noted by a star, 396 and residues conserved across most HNVs are highlighted yellow. Amino acid position numbers displayed 397 represent the position within the AngV or NiV glycoproteins. Virus name (abbreviations) followed by GenBank 398 Accession #: Angavokely virus (AngV) ON613535; Nipah virus (NiV) AF212302; Hendra virus (HeV) AF017149; 399 Mojiang virus (MojV) KF278639; Cedar virus (CedV) JQ001776; Ghanaian bat Henipavirus (GhV) HQ660129; Gamak 400 virus (GAKV) MZ574407; Daeryong virus (DARV) MZ574409.

401

402

403 Discussion

404 We describe and characterize a novel HNV, AngV, from a urine sample collected from an E.

405 dupreanum Malagasy fruit bat. In this study, urine samples from 206 unique fruit bats were

406 assessed by metagenomic sequencing, yielding an overall positive HNV detection rate of 4.9%

- (10/206) for all bats studied and a HNV prevalence of 9.4% (10/106) for the E. dupreanum 407
- 408 hosts. Of all the HNV positive samples, only one sample yielded sufficient reads for assembly of
- 409 a complete coding sequence and subsequent genomic analysis. In a 6-year collection period
- 410 spanning multiple wet/dry seasons, HNV positive samples were only recovered from E.
- 411 dupreanum bats in the Angavokely roosting site, despite prior serological evidence of HNV
- 412 infection in P. rufus and R. madagascariensis bats, as well (21). HNV RNA was recovered from E.

413 dupreanum in both wet and dry seasons, though higher sampling intensity throughout the wet 414 season precludes any conclusions regarding underlying seasonal patterns in these data. 415 Previous work in this system has suggested a seasonal increase in fruit bat seroprevalence 416 across the winter low nutrient season, which also overlaps the gestation period for these 417 synchronously breeding fruit bats (21). In fruit bat systems elsewhere, HNVs are also shed in 418 urine at higher rates during the nutrient-poor dry seasons for the localities in question (49–51); 419 in the case of NiV and HeV, these seasonal viral shedding pulses have been linked to zoonotic 420 spillover.

421

422 The recovered genome of AngV exhibits a structural organization characteristic of the 423 Henipavirus genus and a nucleotide and amino acid identity to HeV and NiV that is comparable 424 to those shared with the more distantly related HNVs, MojV, GhV and CedV. A limited quantity 425 of available original sample precluded full genome recovery for AngV (as evidenced by the lack 426 of the 5' UTR region of the N ORF), which prevented analysis of the extent to which the full 427 AngV genome may abide by the 'Rule-of-Six', observed by all other members of the 428 Orthoparamyxovirinae subfamily (52). Phylogenetic analyses of AngV support classification of 429 this virus as a distinct novel bat-borne *Henipavirus* (Lgene amino acid distance <0.82 distance 430 for the subfamily Orthoparamyxoviringe), in accordance to the International Committee on 431 Taxonomy of Viruses (ICTV) criteria (19). This novel HNV is estimated to have diverged 432 approximately 9,800 years ago, prior to the currently known African and Asian bat-borne HNV 433 lineages but considerably more recently than the estimated mid-to late-Miocene divergence of 434 E. dupreanum from its sister species, E. helvum, on the African continent (53). Recent 435 characterization of *Betacoronaviruses* in Madagascar fruit bats demonstrates surprising identity 436 to lineages circulating in West Africa (22), suggesting that, despite their endemism, Malagasy 437 fruit bats likely experience some form of contact with the African continent. Of the 49 bat 438 species that inhabit the island nation of Madagascar, nine species are widely distributed across 439 Africa, Asia, and/or Europe, presenting opportunities for inter-species viral transmission via 440 island-hopping. Intensified viral sampling of Madagascar's insectivorous bat populations for 441 HNVs thus represents an important future research priority.

442

443 As an ancestral bat-borne HNV, AngV may provide important insight into HNV evolution and 444 pathogenesis. Similar to other paramyxoviruses, the encoded AngV P gene is able to produce 445 multiple immunomodulatory protein products (54). One such protein product is the V protein, 446 thought to be involved in immune evasion and considered a significant determinant of viral 447 pathogenicity and lethality (55, 56). AngV harbors the highly conserved mRNA editing site and a 448 predicted ORF that encodes a V protein with a conserved cysteine-rich C-terminus, suggesting 449 that AngV has the capacity to produce a functional V protein. With the exception of the newly 450 discovered HNVs in shrews, GAKV and DARV, all HNVs harboring a V protein have previously 451 demonstrated evidence of human infection, highlighting the potential for AngV to cause 452 productive infection in humans (1, 9, 12). Further studies are needed to ascertain the virulence 453 potential and host breadth of this novel virus.

454

Characterization of the AngV glycoprotein (G) through AlphaFold modeling and structure-based
alignments revealed a similar structural organization to other HNV glycoproteins. Notably, the
AngV glycoprotein surpasses that of GhV as the longest glycoprotein of the *Henipavirus* genus.
Like that of GhV, the AngV glycoprotein harbors a long C terminal extension (Supplemental
Figure 4). It is unclear if the C terminal extension of the AngV glycoprotein has a functional role,
though the C terminal extension of the glycoprotein in GhV is known to play a functional role in
receptor-mediated fusion (15).

462

463 Henipavirus host tropism and virulence rely on a myriad of factors, one of which is the HNV 464 glycoprotein. The previously characterized HNV glycoproteins of NiV, HeV, CedV, and GhV, 465 utilize members of the ephrinA and ephrinB class family as host-cell receptors for viral entry 466 into human cells (15–17, 47, 57). However, like MojV, the AngV glycoprotein lacks these well-467 conserved ephrin binding residues. Structure-based alignments can shed light on potential 468 receptor binding residues when characterizing novel viruses. For instance, sequence-based 469 comparisons of the GhV and NiV glycoproteins were used to predict GhV ephrin binding (12), 470 which was later confirmed by crystallography (15). Structure-based alignment of the AngV

glycoprotein shows a lack of highly conserved ephrin binding residues, including NiV E533 – a seminal residue for ephrinB2 binding that is conserved across all ephrin binding HNVs. This suggests that, like MojV—and probably DARV and GAKV—the AngV glycoprotein may not bind ephrins, pointing to the possible use of an ancestral viral entry pathway. The growing number of novel HNVs that appear not to rely on ephrin binding for cellular entry could warrant re-evaluation of the existing HNV genus to better reflect conserved function and pathobiology. This work presents a novel bat-HNV, AngV, identified from a Malagasy fruit bat. AngV joins a growing group of ancestral HNVs with unknown cell-entry receptors. Discovery of the cell surface receptor for AngV represents an important future research priority that will shed light on the breadth of host range for this virus, including its zoonotic potential.

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- 517 Conflict of Interests
- 518 The authors declare no competing interests.
- 519
- 520 Data Availability
- 521 Raw and assembled sequencing data are deposited in NCBI Bioproject PRJNA837298. The full
- 522 genome of AngV is available in GenBank under Accession # ON613535. All raw data and code
- 523 for figures can be obtained in our open-access GitHub repository:
- 524 https://github.com/brooklabteam/angavokely-virus

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729 **Table 1.** Prevalence of HNV infections in the urine of Malagasy bats captured during 2013-2019

730 collection period.

		By Site			By Season/Sex		
Roost site	Species	Total sampled	Total henipavirus positive	Season	Total sampled # (M,F)	Total henipavirus positive # (M,F)	
				wet '13	2 (1,1)		
				wet '14/'15	3 (3,0)		
Ambakoana	P. rufus	18	0	wet '15/'16	2 (2,0)	0 (0,0)	
				wet '17/'18	2 (2,0)		
				wet '18/'19	9 (6,3)		
•				wet '17/'18	5 (1,4)	0 (0,0)	
Angavobe	E. dupreanum	8	0	dry '18	3 (1,2)	0 (0,0)	
			10	wet '15/'16	2 (1,1)	0 (0,0)	
		98		wet '17/'18	38 (5,33)	4 (0,4)	
Angavokely	E. dupreanum			dry '18	11 (8,3)	2 (1,1)	
				wet '18/'19	47 (20,27)	4 (2,2)	
Mahabo	P. rufus	8	0	dry '14	8 (4,4)	0 (0,0)	
Mahialambo	P. rufus	5	0	wet '18/'19	5 (2,3)	0 (0,0)	
Makira	P. rufus	2	0	dry '15	2 (2,0)	0 (0,0)	
				wet '17/'18	9 (3,6)		
Marovitsika	P. rufus	62	0	dry '18	7 (3,4)	0 (0,0)	
				wet '18/'19	46 (20,26)		
				wet '13	2 (1,1)		
Maromizaha	R.	5	0	dry '14	1 (0,1)	0 (0,0)	
	madagascariensis			wet '14/'15	2 (2,0)		

731

732 733

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734

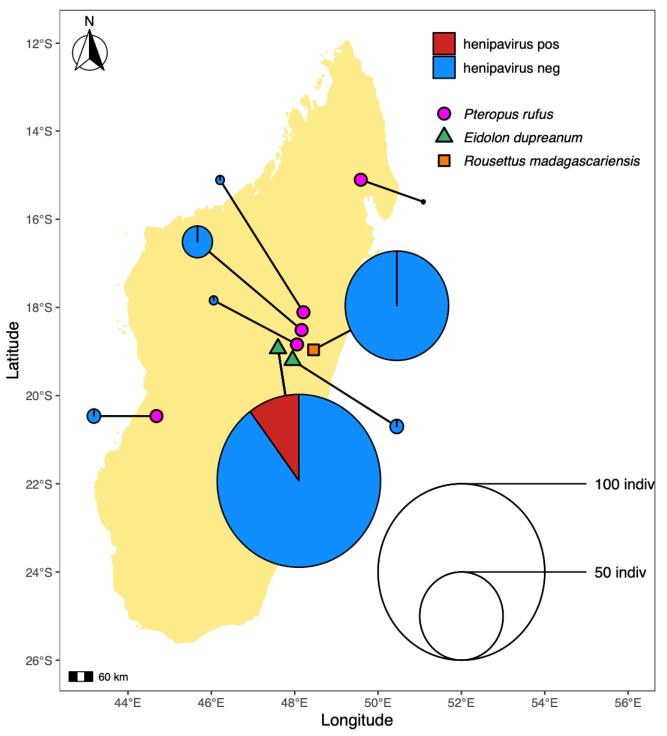
735

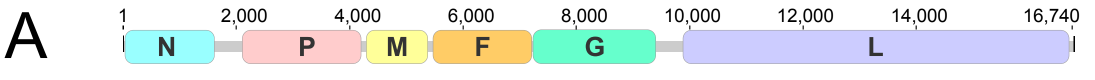
736

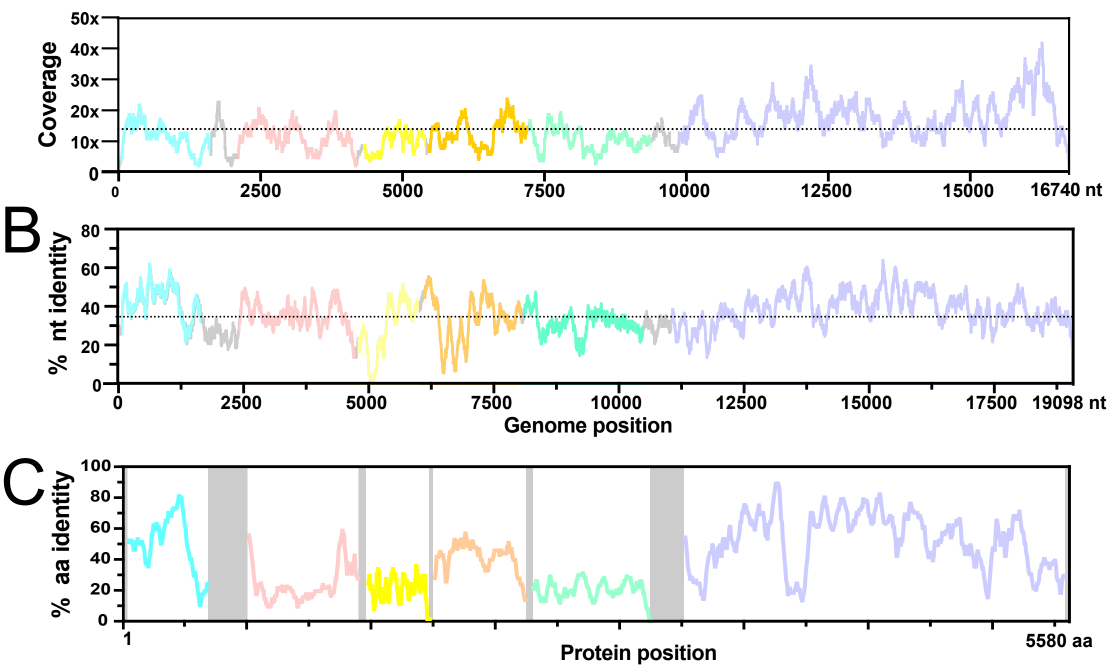
Table 2. Length and pairwise sequence identity of predicted open reading frames of AngV and

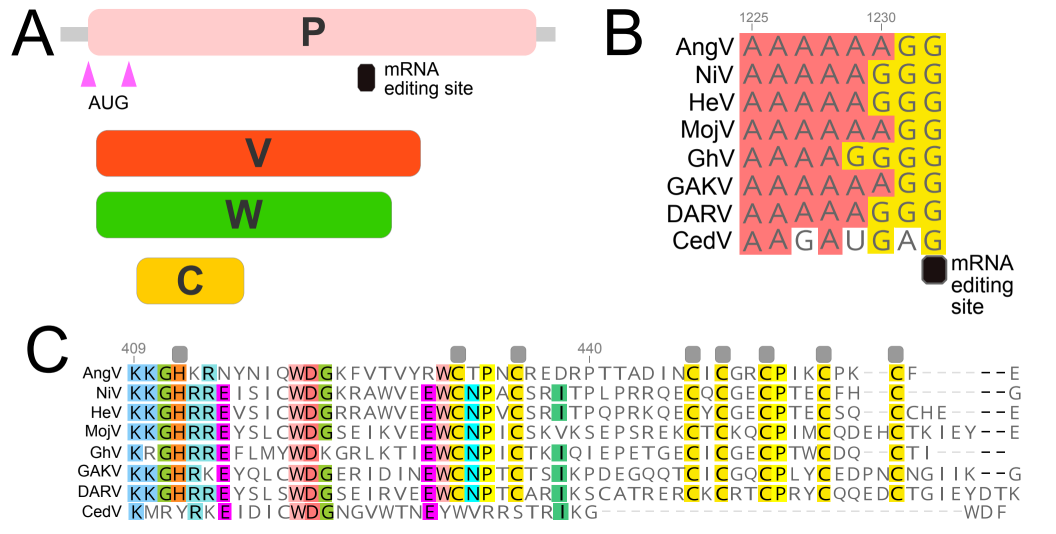
738 other HNV.

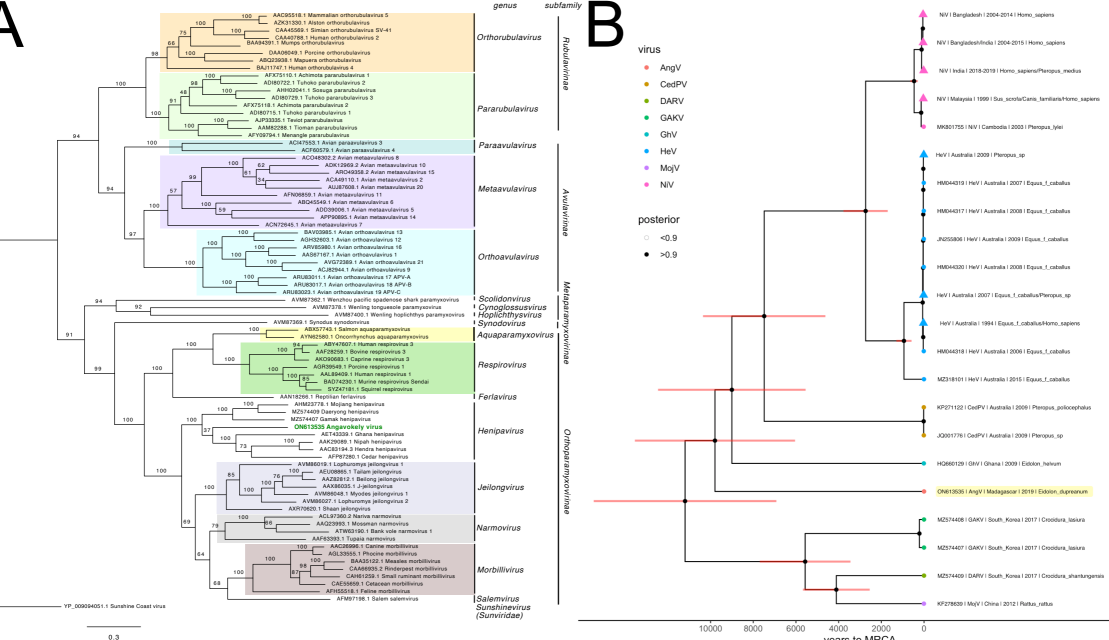
Gene	AngV length aa	NiV length aa (%nt, %aa)	HeV	MojV	CedV	GAKV	DARV	GhV
N	514	532	532	539	510	533	574	514
		(56.4, 48.0)	(55.4, 46.7)	(55.8, 45.3)	(57.5, 48.3)	(58.0, 47.5)	(55.3, 24.0)	(56.2, 48.9)
Р	693	709	707	694	737	586	698	870
•		(47.8, 25.1)	(48.1, 24.1)	(47.6, 23.6)	(42.3, 21.6)	(43.8, 22.5)	(47.3, 23.2)	(42.2, 23.1)
<u> </u>	173	166	166	177	177	184	175	163
с	173	(51.3, 25.7)	(51.5, 25.1)	(47.6, 22.5)	(54.5, 29.8)	(47.4, 27.5)	(48.1, 24.0)	(46.1, 23.1)
v	461	456	457	464		370	468	621
v		(46.9, 22.8)	(48.5, 23.6)	(47.2, 24.5)		(43.0, 22.3)	(47.2, 22.2)	(24.7, 19.8)
w	412	450	448	435		331	437	572
vv		(47.0, 21.3)	(45.0, 20.0)	(46.5, 22.5)		(43.9, 22.7)	(45.5, 21.1)	(39.3, 16.8)
м	354	352	352	340	360	340	345	343
IVI		(60.0, 54.6)	(59.5, 54.0)	(57.1, 53.4)	(56.9, 53.1)	(56.1, 50.8)	(56.5, 52.3)	(56.9, 51.4)
-	539	546	546	545	557	565	545	662
F	239	(53.7, 39.4)	(51.7, 39.4)	(52.2, 40.6)	(51.8, 36.3)	(53.3, 39.4)	(58.8, 40.1)	(45.5, 35.4)
<u> </u>	600	602	604	625	622	635	628	632
G	688	(43.5, 21.4)	(43.8, 20.4)	(43.0, 21.2)	(43.3, 19.8)	(44.9, 17.8)	(44.3, 18.7)	(42.5, 19.8)
	2.250	2,244	2,244	2,277	2,501	2,291	2,271	2,250
L	2,259	(74.7, 52.2)	(57.6, 51.9)	(57.5, 51.8)	(52.4, 44.8)	(57.7, 51.0)	(58.8, 51.3)	(57.0, 49.1)





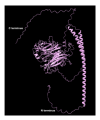






vears to MRCA





В

AngV 521 NV 502	AngV 548 NV 529	Ang/ 575 NIV 555	AngV 599 NIV 580
AngV FCHSSG	i Y S Ĥ Ŝ Ŷ (5 I A A	G I R
NV CWEGV			
HeV V C W E G T	Y-NOTAL	T N A	Q I YI
MOVICNTRG	Y ···· N N G G	F Y S I '	TGK
CedV CYGGT	Y-DOLA	L N T	R ···· T N I
GhV V C WEGT	Y E Q V A I	S S A	R I T I
GAKV SCKTWN	I E … K T G N S	5 Q S I	G G V
DARV VCSSYG	Y N G E G	Г F К I	I G Q