

# Equine Influenza A(H3N8) Virus Isolated from Bactrian Camel, Mongolia

## Technical Appendix

Technical Appendix Table 1. Virus name, subtype, host species, collection date, and GenBank accession number (PB2 segment) for the 36 H3N8 viruses used in the whole-genome phylogenetic analysis conducted for all eight viral genome segments.

Virus name	Subtype	Host species	Collection date	Accession no.
A_camel_Mongolia_335_2012	H3N8	Camel	Nov-2012	CY164127.1
A_avian_Japan_8KI0102_2008	H3N8	Avian	Oct-08-08	CY079266
A_avian_Japan_8KI0129_2008	H3N8	Avian	Oct-08-08	CY079258
A_avian_Japan_8KI0150_2008	H3N8	Avian	Oct-08-08	CY079242
A_avian_Japan_8KI0162_2008	H3N8	Avian	Oct-08-08	CY079234
A_chicken_Laos_A0573_2007	H3N8	Avian	2007	CY040963
A_chicken_Vietnam_G14_2008	H3N8	Avian	Jan-2008	AB593452
A_donkey_Xinjiang_5_2007	H3N8	Equine	Dec-2007	EU794572
A_duck_Beijing_40_04	H3N8	Avian	2004	EU492488
A_duck_Beijing_61_05	H3N8	Avian	2005	EU492492
A_duck_Hokkaido_8_1980	H3N8	Avian	1980	AB274963
A_duck_Hunan_S1256_2012	H3N8	Avian	Mar-23-12	CY146601
A_duck_Hunan_S1824_2012	H3N8	Avian	Mar-24-12	CY146625
A_duck_Nanchang_1681_1992	H3N8	Avian	Dec-01-92	CY005475
A_duck_Vietnam_G119_2006	H3N8	Avian	Nov-2006	AB593428
A_environment_Hunan_S4350_2011	H3N8	Avian	Nov-13-11	CY146753
A_equine_Gansu_7_2008	H3N8	Equine	Jan-2008	EU794492
A_equine_Heilongjiang_1_2010_	H3N8	Equine	Apr-23-10	KF309031
A_equine_Heilongjiang_10_2008	H3N8	Equine	Apr-2008	EU794508
A_equine_Huabei_1_2007	H3N8	Equine	Dec-03-07	GU571147
A_equine_Inner_Mongolia_8_2008	H3N8	Equine	Feb-2008	EU794524
A_equine_Kyonggi_SA1_2011	H3N8	Equine	Jul-01-11	JX844143
A_equine_Liaoning_9_2008	H3N8	Equine	Apr-2008	EU794516
A_equine_Qinghai_1_1994	H3N8	Equine	1994	EU794532
A_equine_Sachiyama_1_1971	H3N8	Equine	1971	CY034941
A_equine_Tokyo_2_1971	H3N8	Equine	1971	CY096922
A_equine_Tottori_1_07	H3N8	Equine	2007	AB591847
A_equine_Xinjiang_1_2007	H3N8	Equine	Nov-2007	EU794540
A_equine_Xinjiang_2_2007	H3N8	Equine	Nov-2007	EU794548
A_equine_Xinjiang_3_2007	H3N8	Equine	Nov-2007	EU794556
A_equine_Xinjiang_4_2007	H3N8	Equine	Dec-2007	EU794564

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<b>Virus name</b>	<b>Subtype</b>	<b>Host species</b>	<b>Collection date</b>	<b>Accession no.</b>
A_equine_Xuzhou_01_2013	H3N8	Equine	Aug-27-13	KF806992
A_Mallard_SanJiang_90_2006_2006	H3N8	Avian	2006	CY100631
A_muscovy_duck_Vietnam_LBM240_2012	H3N8	Avian	2012	AB786912
A_swine_Anhui_01_2006	H3N8	Swine	Jan-06-06	FJ200417
A_swine_Chibi_01_2005	H3N8	Swine	Dec-15-05	FJ200425

Technical Appendix Table 2. Virus names for the hemagglutinin sequences of the 155 equine A/H3N8 viruses used in Figure 1. Viruses containing the two amino acid insertion near the beginning of the hemagglutinin are specified.

<b>Virus name</b>	<b>Subtype</b>	<b>Insertion</b>
A_camel_Mongolia_335_2012	H3N8	yes
A_donkey_Xinjiang_5_2007	H3N8	
A_equine_Alaska_29759_1991	H3N8	
A_equine_Algiers_1_1972	H3N8	
A_equine_Almaty_26_2007	H3N8	yes
A_equine_Argentina_1_93	H3N8	
A_equine_Austria_421_1992	H3N8	
A_equine_Bari_2005	H3N8	yes
A_equine_Berlin_1_1989	H3N8	
A_equine_California_1_1980	H3N8	
A_equine_California_103_1982	H3N8	
A_equine_California_191_2003	H3N8	
A_equine_California_4537_1997	H3N8	
A_equine_California_83_1982	H3N8	
A_equine_California_8560_2002	H3N8	
A_equine_Carlow_1_2009	H3N8	
A_equine_Colorado_10_2007	H3N8	
A_equine_Cordoba_18_1985	H3N8	
A_equine_Donegal_1_2007	H3N8	yes
A_equine_Donegal_1_2009	H3N8	
A_equine_Down_1_2008	H3N8	yes
A_equine_Egypt_6066NAMRU3-VSVRI_2008	H3N8	
A_equine_Florida_1_93	H3N8	
A_equine_Florida_612_2004	H3N8	
A_equine_Florida_779_2004	H3N8	
A_equine_Fontainbleu_1_1979	H3N8	
A_equine_Fontainebleau_1976	H3N8	
A_equine_France_1_1976	H3N8	
A_equine_Gansu_7_2008	H3N8	
A_equine_Georgia_1_1981	H3N8	
A_equine_Georgia_10_1981	H3N8	

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<b>Virus name</b>	<b>Subtype</b>	<b>Insertion</b>
A_camel_Mongolia_335_2012	H3N8	yes
A_equine_Georgia_13_1981	H3N8	
A_equine_Georgia_3_1981	H3N8	
A_equine_Georgia_9_1981	H3N8	
A_equine_Guelph_06-28865_2006	H3N8	
A_equine_Guelph_G03-0250_2003	H3N8	
A_equine_Guelph_G03-55399_2003	H3N8	
A_equine_Guelph_G04-54701_2004	H3N8	
A_equine_Heilongjiang_1_2010	H3N8	yes
A_equine_Heilongjiang_10_2008	H3N8	
A_equine_Hokkaido_I828_2008	H3N8	
A_equine_Hong_Kong_J_1992	H3N8	
A_equine_Huabei_1_2007	H3N8	yes
A_equine_Hubei_6_2008	H3N8	
A_equine_Ibadan_6_91	H3N8	
A_equine_Ibadan_9_91	H3N8	
A_equine_Ibaraki_1_07	H3N8	
A_equine_Idaho_37875_1991	H3N8	
A_equine_Inner_Mongolia_8_2008	H3N8	
A_equine_Italy_1062_1991	H3N8	
A_equine_Italy_1199_1992	H3N8	
A_equine_Italy_824_1991	H3N8	
A_equine_Johannesburg_1_1986	H3N8	
A_equine_Kanazawa_1_2007	H3N8	
A_equine_Kascakew_1_1978	H3N8	
A_equine_Katra-Jammu_6_2008	H3N8	yes
A_equine_Kentucky_1_1981	H3N8	
A_equine_Kentucky_1_1986	H3N8	
A_equine_Kentucky_1_1987	H3N8	
A_equine_Kentucky_1_1990	H3N8	
A_equine_Kentucky_1_1991	H3N8	
A_equine_Kentucky_1_1992	H3N8	
A_equine_Kentucky_1_1994	H3N8	
A_equine_Kentucky_1277_1990	H3N8	
A_equine_Kentucky_2_1980	H3N8	
A_equine_Kentucky_2_1981	H3N8	
A_equine_Kentucky_2_1986	H3N8	
A_equine_Kentucky_2_1987	H3N8	
A_equine_Kentucky_211_1987	H3N8	
A_equine_Kentucky_3_1981	H3N8	

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<b>Virus name</b>	<b>Subtype</b>	<b>Insertion</b>
A_camel_Mongolia_335_2012	H3N8	yes
A_equine_Kentucky_3_1986	H3N8	
A_equine_Kentucky_4_1980	H3N8	
A_equine_Kentucky_5_2002	H3N8	
A_equine_Kentucky_692_1988	H3N8	
A_equine_Kentucky_694_1988	H3N8	
A_equine_Kentucky_698_1988	H3N8	
A_equine_Kentucky_8_1994	H3N8	
A_equine_Kentucky_bitter_boredom5_1976	H3N8	
A_equine_Kentucky_magnificent_genius1_1981	H3N8	
A_equine_Kentucky_pass_the_pepper1_1976	H3N8	
A_equine_Kentucky_Rosie100_1981	H3N8	
A_equine_Kildare_1_2007	H3N8	yes
A_equine_Kyonggi_SA1_2011	H3N8	
A_equine_Liaoning_9_2008	H3N8	
A_equine_Limerick_1_2010	H3N8	
A_equine_Lincolnshire_1_2007	H3N8	
A_equine_Lonquen_1_2006	H3N8	
A_equine_Massachussetts_213_2003	H3N8	
A_equine_Miami_1_1963	H3N8	
A_equine_Mongolia_20_2011	H3N8	yes
A_equine_Mongolia_3_2011	H3N8	yes
A_equine_Mongolia_56_2011	H3N8	yes
A_equine_Mongolia_6_2011	H3N8	yes
A_equine_Montana_9233_2007	H3N8	
A_equine_Mysore_1_2008	H3N8	
A_equine_New_Market_1_1979	H3N8	
A_equine_New_Market_1976	H3N8	
A_equine_New_Market_nasalwash1_1979	H3N8	
A_equine_New_York_1_1975	H3N8	
A_equine_New_York_1_1999	H3N8	
A_equine_New_York_146066_2007	H3N8	
A_equine_New_York_452_2003	H3N8	
A_equine_New_York_VR-297_1983	H3N8	
A_equine_Newmarket_5_2003	H3N8	
A_equine_Ohio_1_2003	H3N8	
A_equine_Ohio_113461-1_2005	H3N8	
A_equine_Ohio_113461-2_2005	H3N8	
A_equine_Ohio_113461-3_2005	H3N8	
A_equine_Otar_764_2007	H3N8	yes

Technical Appendix Table 2. Virus names for the hemagglutinin sequences of the 155 equine A/H3N8 viruses used in Figure 1. Viruses containing the two amino acid insertion near the beginning of the hemagglutinin are specified.

<b>Virus name</b>	<b>Subtype</b>	<b>Insertion</b>
A_camel_Mongolia_335_2012	H3N8	yes
A_equine_Qinghai_1_1994	H3N8	
A_equine_Richmond_1_2007	H3N8	yes
A_equine_Romania_1_1980	H3N8	
A_equine_Rome_5_1991	H3N8	
A_equine_Rook_93753_1989	H3N8	
A_equine_Sachiyama_1_1971	H3N8	
A_equine_Santa_Fe_1_1985	H3N8	
A_equine_Sao_Paulo_1_1969	H3N8	
A_equine_Sao_Paulo_6_1963	H3N8	
A_equine_Spain_1_2007	H3N8	yes
A_equine_Suffolk_89	H3N8	
A_equine_Sussex_1_1989	H3N8	
A_equine_Switzerland_1118_1979	H3N8	
A_equine_Switzerland_173_1993	H3N8	
A_equine_Sydney_6085_2007	H3N8	
A_equine_Taby_1991	H3N8	
A_equine_Tennessee_5_1985	H3N8	
A_equine_Tennessee_5_1986	H3N8	
A_equine_Texas_117793_2005	H3N8	
A_equine_Texas_39655_1991	H3N8	
A_equine_Tiaret_1_2011	H3N8	yes
A_equine_Tiaret_10_2011	H3N8	yes
A_equine_Tiaret_2_2011	H3N8	yes
A_equine_Tiaret_3_2011	H3N8	yes
A_equine_Tiaret_4_2011	H3N8	yes
A_equine_Tiaret_5_2011	H3N8	yes
A_equine_Tiaret_6_2011	H3N8	yes
A_equine_Tiaret_7_2011	H3N8	yes
A_equine_Tiaret_8_2011	H3N8	yes
A_equine_Tiaret_9_2011	H3N8	yes
A_equine_Tokyo_1971	H3N8	
A_equine_Tokyo_2_1971	H3N8	
A_equine_Tottori_1_07	H3N8	
A_equine_Uruguay_1_1963	H3N8	
A_equine_Virginia_131054-3_2005	H3N8	
A_equine_Wisconsin_1_03	H3N8	
A_equine_Xinjiang_1_2007	H3N8	
A_equine_Xinjiang_2_2007	H3N8	
A_equine_Xinjiang_3_2007	H3N8	

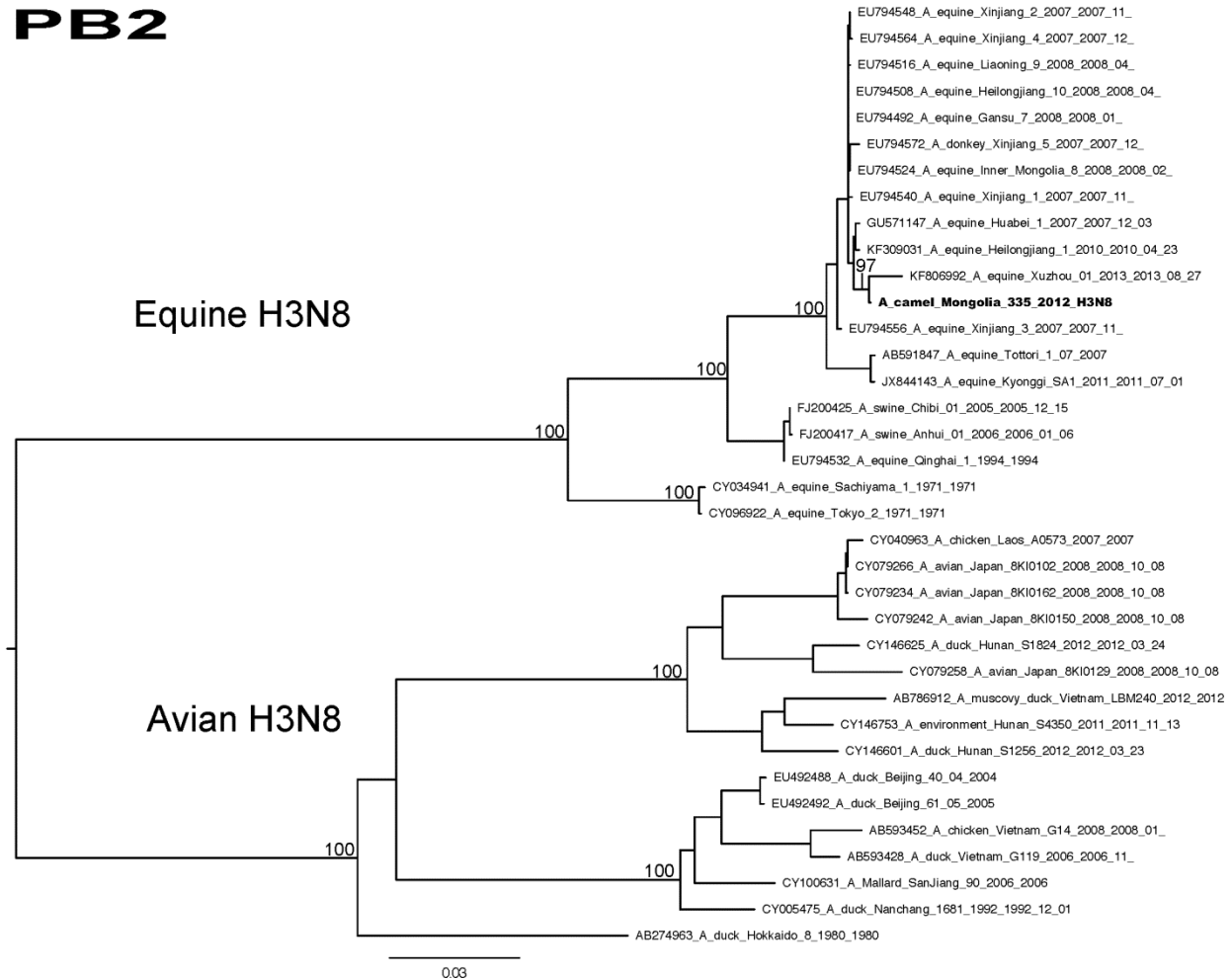
Technical Appendix Table 2. Virus names for the hemagglutinin sequences of the 155 equine A/H3N8 viruses used in Figure 1. Viruses containing the two amino acid insertion near the beginning of the hemagglutinin are specified.

<b>Virus name</b>	<b>Subtype</b>	<b>Insertion</b>
A_camel_Mongolia_335_2012	H3N8	yes
A_equine_Xinjiang_4_2007	H3N8	
A_equine_Xuzhou_01_2013	H3N8	yes
A_equine_Yokohama_aq13_2010	H3N8	yes
A_equine_Yokohama_aq19_2009	H3N8	
A_equine_Yokohama_aq29_2011	H3N8	
A_equine_Yokohama_aq5_2011	H3N8	
A_equine_Yokohama_aq53_2011	H3N8	
A_equine_Yokohama_aq79_2011	H3N8	

Technical Appendix Table 3. Position of the two amino acid insertion at the beginning of the hemagglutinin sequence.

Example of virus with insertion:	
A/equine/Almaty/26/2007/H3N8	MKTTIIFIFILLTHW
Example of virus without insertion:	
A/equine/Alaska/29759/1991/H3N8	MKTTIIL--ILLTHW

# PB2

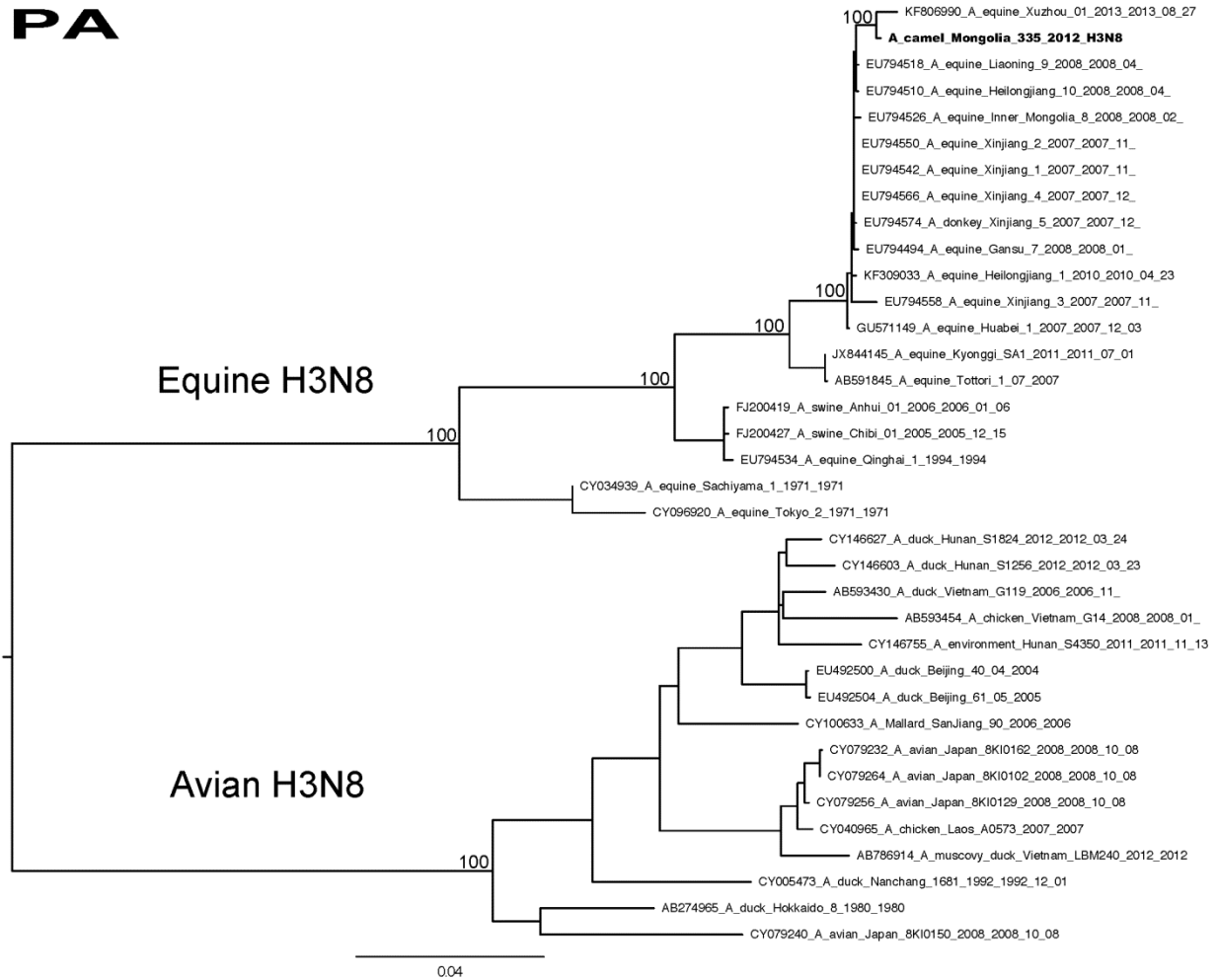


Technical Appendix Figure 1. Evolutionary relationships of the PB2 segment of 36 influenza A viruses of the H3N8 subtype collected in Asia from horses (n = 17), avian species (n = 16), swine (n = 2), and a camel (*A/camel/Mongolia/335/2012(H3N8)*, highlighted in bold). The tree is midpoint rooted for clarity, and all branch lengths are drawn to scale. High bootstrap values (> 70) are provided for key nodes. Scale bar indicates nucleotide substitutions per site.



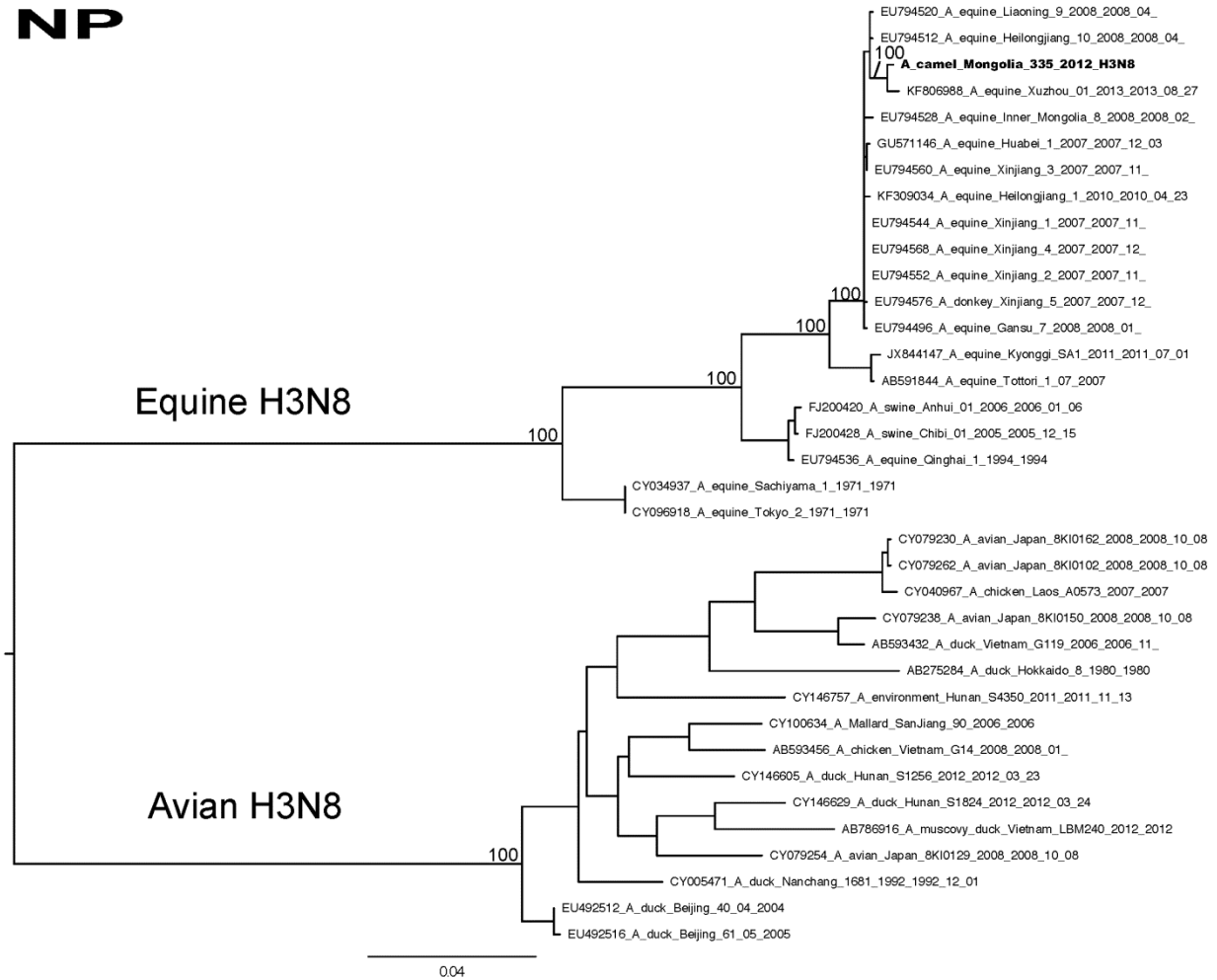


PA

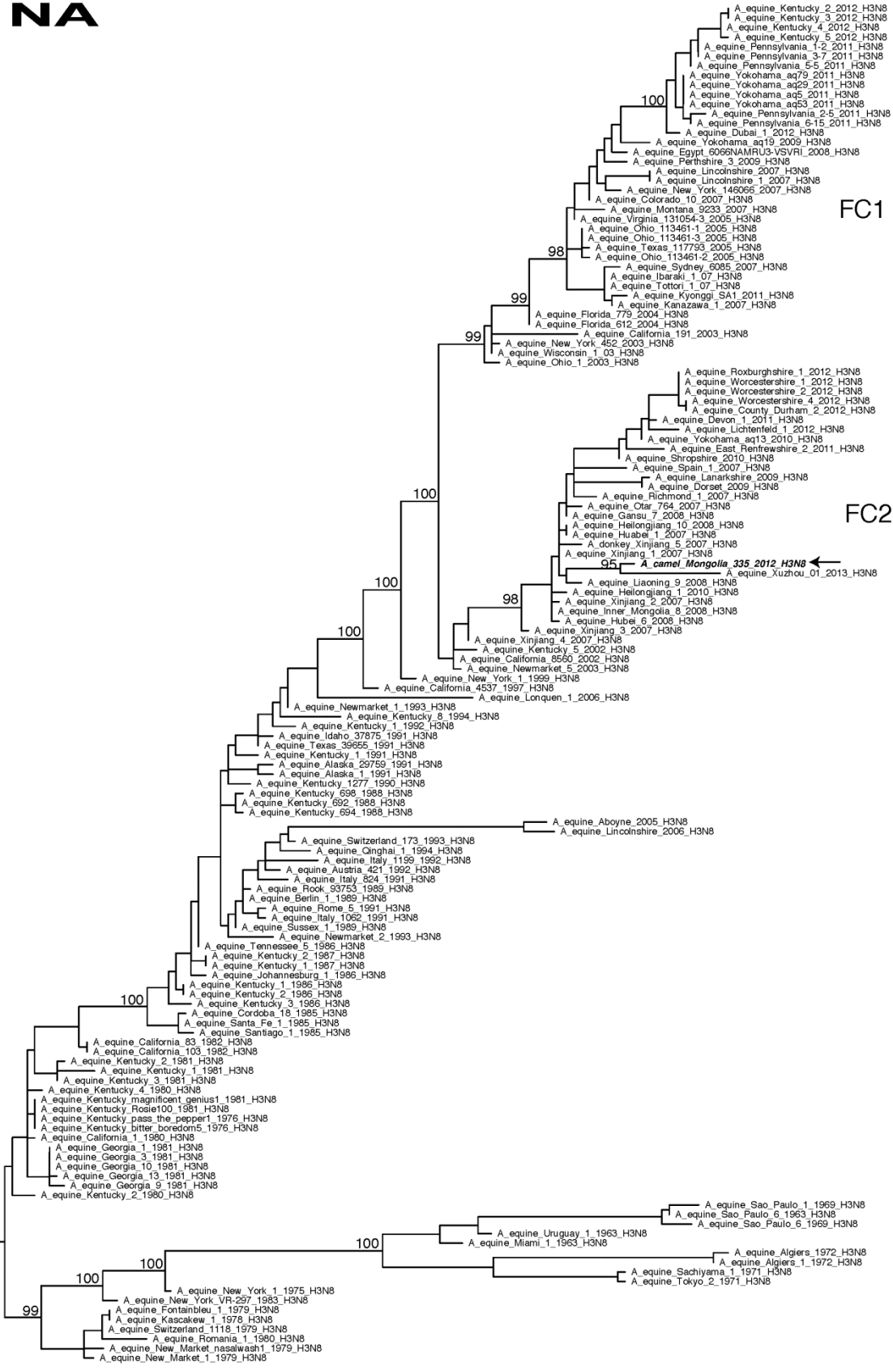


Technical Appendix Figure 3. Evolutionary relationships of the PA segment of 36 influenza A viruses of the H3N8 subtype collected in Asia from horses (n = 17), avian species (n = 16), swine (n = 2), and a camel (*A/camel/Mongolia/335/2012(H3N8)*, highlighted in bold). The tree is midpoint rooted for clarity, and all branch lengths are drawn to scale. High bootstrap values (> 70) are provided for key nodes. Scale bar indicates nucleotide substitutions per site.

# NP



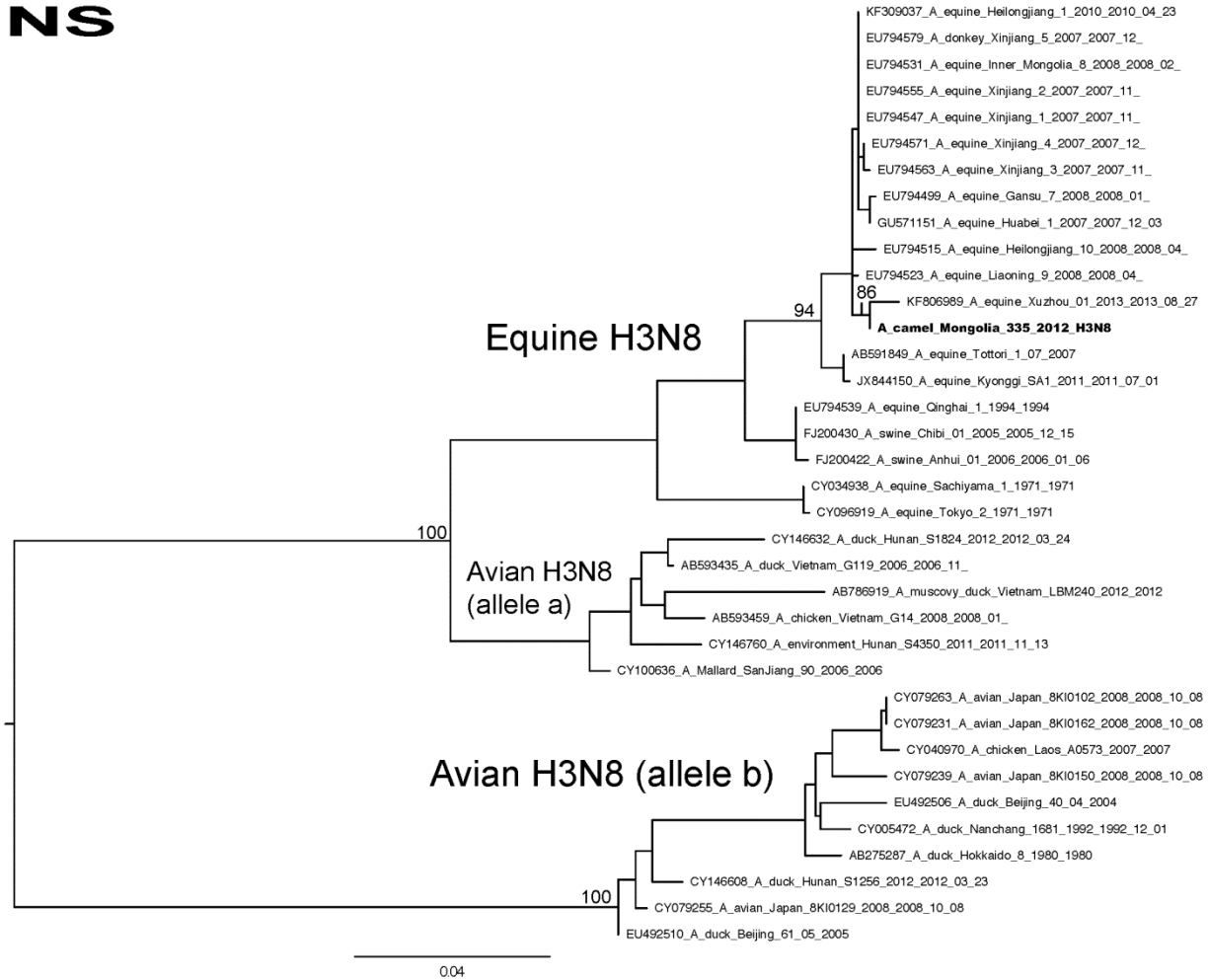
Technical Appendix Figure 4. Evolutionary relationships of the NP segment of 36 influenza A viruses of the H3N8 subtype collected in Asia from horses (n = 17), avian species (n = 16), swine (n = 2), and a camel (*A/camel/Mongolia/335/2012(H3N8)*, highlighted in bold). The tree is midpoint rooted for clarity, and all branch lengths are drawn to scale. High bootstrap values (> 70) are provided for key nodes. Scale bar indicates nucleotide substitutions per site.



0.02



NS



Technical Appendix Figure 7. Evolutionary relationships of the NS segment of 36 influenza A viruses of the H3N8 subtype collected in Asia from horses (n = 17), avian species (n = 16), swine (n = 2), and a camel (A/camel/Mongolia/335/2012(H3N8), highlighted in bold). The tree is midpoint rooted for clarity, and all branch lengths are drawn to scale. High bootstrap values (> 70) are provided for key nodes.