



SURVEILLANCE REPORT

Influenza virus characterisation

Summary Europe, June 2016

Summary

Since week 40/2015, almost 139 000 influenza detections across the Region have been reported. Influenza type A viruses have prevailed over type B with A(H1N1)pdm09 viruses, greatly outnumbering A(H3N2) and B/Victoria-lineage detections representing over 91% of the type B viruses assigned to a lineage.

Since 1 January 2016, EU/EEA countries have shared 498 influenza-positive specimens with the Francis Crick Institute, London, for detailed characterisation. Since the May report, 160 viruses have been characterised antigenically, and genetic analyses are ongoing.

Of the 63 A(H1N1)pdm09 viruses characterised antigenically, 60 (95%) were similar to the vaccine virus A/California/7/2009. Worldwide new genetic subclusters of viruses within the 6B clade have emerged, with two being designated as subclades: 6B.1 defined by HA1 amino acid substitutions S162N and I216T, and 6B.2 defined by HA1 amino acid substitutions V152T and V173I. Of the 381 viruses characterised genetically for the 2015–2016 season, 29 (8%) were clade 6B, 344 (90%) were subclade 6B.1, and eight (2%) were subclade 6B.2.

The 25 A(H3N2) test viruses characterised by haemagglutination inhibition (HI) assay were poorly recognised by reference antiserum raised against egg-propagated A/Switzerland/9715293/2013, the vaccine virus recommended for use in the 2015–2016 northern hemisphere influenza season. The test viruses were recognised somewhat better by antisera raised against egg-propagated A/Hong Kong/4801/2014, the virus recommended for use in 2016 southern hemisphere and 2016–2017 northern hemisphere influenza vaccines. Of 114 A(H3N2) viruses characterised genetically for the 2015–2016 season: two (2%) were clade 3C.3, 84 (74%) were subclade 3C.2a, and 28 (24%) were subclade 3C.3a.

The 63 B/Victoria-lineage viruses were antigenically similar to tissue culture-propagated surrogates of B/Brisbane/60/2008. All 159 viruses characterised genetically for the 2015–2016 season fell in genetic clade 1A, as do recently collected viruses worldwide.

Nine B/Yamagata viruses have been characterised since the previous report; they reacted well with post-infection ferret antiserum raised against egg-propagated B/Phuket/3073/2013, the recommended vaccine virus for the northern hemisphere 2015–2016 influenza season and for quadrivalent vaccines in the 2016 southern hemisphere and 2016–2017 northern hemisphere seasons. Of the 25 viruses characterised genetically for the 2015–2016 season, 24 fell in genetic clade 3 and one in clade 2.

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Table 1 shows a summary of influenza virus detections in the WHO European Region reported to TESSy for the weekly reporting period (weeks 40/2015–20/2016) of the 2015–2016 season. A total of over 138 000 detections had been made with type A viruses prevailing over type B at a ratio of 2.3:1; this compares to a ratio of 5.8:1 to week 7/2016, indicating a surge in influenza type B circulation over the subsequent 13 weeks. As of week 20/2016, of the type A viruses subtyped ($n = 66\,707$) and the type B viruses ascribed to lineage ($n = 7\,834$), A(H1N1)pdm09 have prevailed over A(H3N2) and B/Victoria over B/Yamagata by ratios of 10.2:1 and 11.1:1, respectively. While relatively few influenza detections have been reported for weeks 21–25/2016, it is notable that the ratios for type A:type B, A(H1N1)pdm09:A(H3N2) and B/Victoria:B/Yamagata have dropped to 0.4:1, 1.1:1 and 2.0:1, respectively.

Since 1 January 2016, 47 shipments of specimens have been received at the Crick Worldwide Influenza Centre (WIC), from 26 countries in the EU/EEA. These packages contained 498 specimens, a mix of clinical samples and virus isolates originating from 21 countries, with collection dates after 31 December 2015 (Table 2). The majority (71%) were type A viruses, and A(H1N1)pdm09 outnumbered A(H3N2) at a ratio of 4.4:1. Of the 146 type B specimens received (29% of the specimens), 125 were B/Victoria-lineage and 19 B/Yamagata-lineage. A number of specimens are still in the process of being characterised. The antigenic and genetic properties of influenza virus isolates characterised since the May 2016 report¹ are presented and discussed in this report.

Table 1. Influenza virus detections in the WHO European Region from the start of reporting for the 2015–2016 season (week 40/2015)

Virus type/subtype/lineage	Cumulative number of detections						Totals*			
	Sentinel sources		Non-sentinel sources		Totals		%		Ratios	
	Weeks 40/2015-20/2016	Weeks 21-25/2016	Weeks 40/2015-20/2016	Weeks 21-25/2016	Weeks 40/2015-20/2016	Weeks 21-25/2016	Weeks 40/2015-20/2016	Weeks 21-25/2016	Weeks 40/2015-20/2016	Weeks 21-25/2016
Influenza A	10496	3	85919	167	96415	170	69.7	26.9	2.3:1	0.4:1
A(H1N1)pdm09	8665	1	52083	39	60748	40	91.1	51.3	10.2:1	1.1:1
A(H3N2)	1365	1	4594	37	5959	38	8.9	48.7		
A not subtyped	466	1	29242	91	29708	92				
Influenza B	8144	11	33791	452	41935	463	30.3	73.1		
Victoria lineage	3974	2	3210	46	7184	48	91.7	66.7	11.1:1	2.0:1
Yamagata lineage	145	0	505	24	650	24	8.3	33.3		
Lineage not ascribed	4025	9	30076	382	34101	391				
Total detections (total tested)	18 640 (50 861)	14 (510)	119 710 (536 625)	619 (17 575)	138 350 (587 486)	633 (18 085)				

* Percentages are shown for total detections (types A & B [in bold type], and for viruses ascribed to influenza A subtype and influenza B lineage). Ratios are given for type A:B [in bold type], A(H1N1)pdm09: A(H3N2) and Victoria:Yamagata lineages.

¹ European Centre for Disease Prevention and Control. Influenza virus characterisation, summary Europe, May 2016. Stockholm: ECDC; 2016. Available from: <http://ecdc.europa.eu/en/publications/Publications/influenza-virus-characterisation-may-2016.pdf>

Table 2. Summary of clinical samples and virus isolates, with collection dates after 31 December 2015, received from EU/EEA Member States

MONTH*	TOTAL RECEIVED	A		H1N1pdm09		H3N2		B		B Victoria lineage		B Yamagata lineage		
		Number received	Number propagated	Number received	Number propagated	Number received	Number propagated ²	Number received	Number propagated	Number received	Number propagated	Number received	Number propagated	
2016														
JANUARY														
Bulgaria	18			18	18									
Cyprus	15			9	5	1	1	0		5	3			
Czech Republic	3			3	3									
Estonia	3			2	0					1	1			
Finland	1			1	1									
Germany	26			11	11	5	5	0		8	8	2	2	
Greece	27			27	17									
Hungary	7			4	4					3	3			
Iceland	6			5	5							1	1	
Ireland	10			9	9					1	1			
Italy	2			1	1	1	1	0						
Latvia	8			6	6					2	2			
Netherlands	2			2	2									
Portugal	6			6	6									
Romania	8			7	7	1	0	1						
Slovenia	8			3	3	3	0	3	2	0				
Spain	19			16	15	1	0	1		2	2			
2016														
FEBRUARY														
Bulgaria	47			34	33	1	0	1		12	12			
Cyprus	9			8	8					1	1			
Czech Republic	8			8	8									
Finland	5			4	4	1	1	0						
Germany	13			6	6	2	2	0				5	5	
Greece	4			4	2									
Iceland	3			2	2							1	1	
Italy	22			7	7	10	7	3		4	4	1	1	
Latvia	2			2	2									
Lithuania	12			10	10					2	2			
Netherlands	1			1	1									
Portugal	1			1	1									
Romania	6			3	3	2	2	0		1	1			
Slovakia	9			3	3	1	1	0		5	5			
Slovenia	17			5	5	9	4	4		3	3			
Spain	17			15	14					2	2			
Sweden	5			1	1	2	2	0		2	2			
2016														
MARCH														
Bulgaria	16			6	6	2	0	2		8	8			
Czech Republic	11			2	2	1	1	0		8	8			
Finland	5			2	2					3	3			
Germany	9					1	0	1		7	7	1	1	
Iceland	3			1	1	1	0	1		1	1			
Italy	10			1	1	5	4	1		4	4			
Norway	2				in process									
Portugal	10			6	6					4	3			
Romania	9			6	6	1	1	0		1	1	1	1	
Slovakia	10			3	3					5	5	2	2	
Slovenia	14			3	3	6	3	3		5	5			
Sweden	5			1	1	1	1	0		2	2	1	1	
2016														
APRIL														
Iceland	8			2	2	2	0	1		3	3	1	1	
Italy	3			1	1	1	0	1		1	1			
Portugal	6			1	1					5	5			
Romania	8			3	3					5	5			
Slovakia	2			1	1							1	1	
Slovenia	4			1	1					2	2	1	1	
Sweden	1					1	0	1						
2016														
MAY														
Iceland	4					2	1	1		2	2			
Norway	5			1	in process	1	in process			2	in process	1	in process	
Slovenia	2									2	2			
2016														
JUNE														
Iceland	1									1	1			
21 Countries	498	0	0	287	263	65	37	25	2	0	125	120	19	18
				57.6%		13.1%					25.1%		3.8%	
				70.7%							29.3%			

* Month indicates the months in which the clinical specimens were collected

1. Propagated to sufficient titre to perform HI assay

2. Propagated to sufficient titre to perform HI assay in presence of 20nM oseltamivir; numbers in red indicate viruses recovered but with insufficient HA titre to permit HI assay

Influenza A(H1N1)pdm09 virus analyses

Haemagglutination inhibition (HI) analyses of viruses that have been performed since the May 2016 report are shown in Tables 3-1 and 3-2. Of the 63 A(H1N1)pdm09 viruses from EU/EEA countries antigenically characterised, 60 were similar to the vaccine virus, A/California/7/2009, reacting with antiserum raised against the vaccine virus at titres within fourfold of the homologous titre. Generally, the test viruses were recognised by the panel of antisera at titres within fourfold of the titres for the homologous viruses, with the exception of the antiserum raised against A/Christchurch/16/2010. This antiserum recognised only 5/63 (8%) test viruses at a titre within fourfold of the titre for the homologous virus. In addition, antiserum raised against A/Lviv/N6/2009 showed eightfold reduced titres with 17 of the test viruses, compared to the homologous titre. Reference viruses carrying HA1 G155E amino acid substitutions, A/Bayern/69/2009 and A/Lviv/N6/2009, showed reduced recognition by the antisera raised against A/California/7/2009 and reference viruses in genetic clades 4, 5, 6, 7 and subclades 6A, 6B, 6B.1 and 6B.2.

HA gene sequencing was performed for 62 of the test viruses: all fell in subclade 6B.1 (Tables 3-1 and 3-2). Since 2009, the HA genes have evolved, and nine clades have been designated. For well over a year, viruses in clade 6, represented by A/St Petersburg/27/2011 and carrying amino acid substitutions of D97N, S185T and S203T in HA1 and E47K and S124N in HA2 compared with A/California/7/2009, have predominated worldwide, with a number of subclades emerging. All EU/EEA viruses characterised since the September 2014 report² carry HA genes in subclade 6B, which is characterised by additional amino acid substitutions of K163Q, A256T and K283E in HA1 and E172K in HA2 compared with A/California/7/2009, e.g. A/South Africa/3626/2013. A number of virus clusters have emerged within clade 6B and two of these have been designated as subclades: viruses in subclade 6B.1 are defined by HA1 amino acid substitutions S84N, S162N (which results in the formation of a new potential glycosylation motif at residues 162-164 of HA1) and I216T, while those in subclade 6B.2 are defined by HA1 amino acid substitutions V152T and V173I (Figure 1).

² European Centre for Disease Prevention and Control. Influenza virus characterisation, summary Europe, September 2014. Stockholm: ECDC; 2014. Available from: <http://ecdc.europa.eu/en/publications/Publications/Influenza-ERLI-Net-report-Sept-2014.pdf>

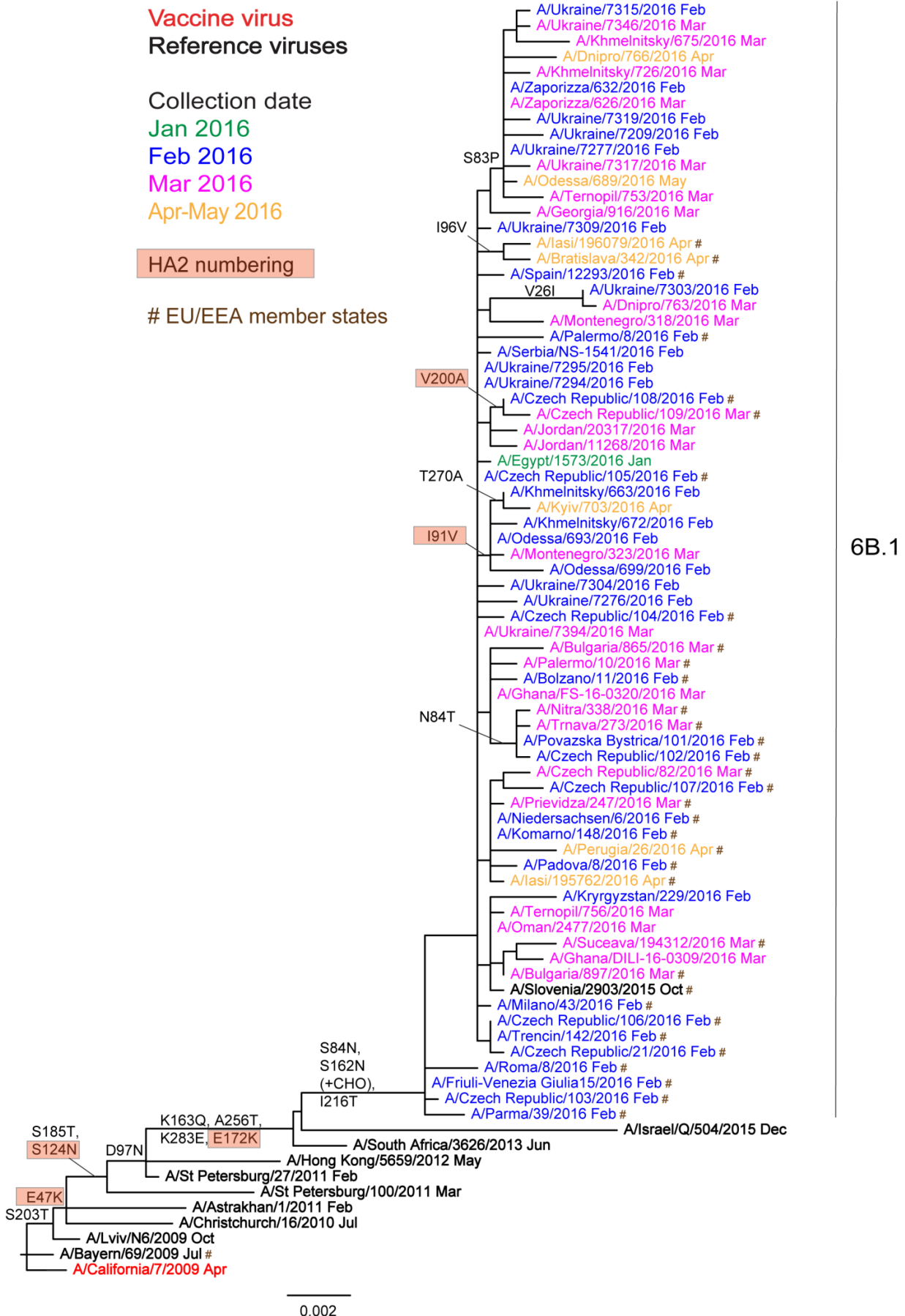
Table 3-1. Antigenic analysis of A(H1N1)pdm09 viruses by HI

Viruses	Other information	Collection date	Passage history	Haemagglutination inhibition titre													
				Post-infection ferret antisera													
				A/Cat 7/09 Egg F06/16 ¹	A/Bayem 69/09 MDCK F09/15 ¹	A/Lviv N6/09 MDCK F14/13 ¹	A/Chch 16/10 Egg F15/14 ¹	A/Astrak 1/11 MDCK F22/13 ¹	A/St. P 27/11 Egg F26/14 ¹	A/St. P 100/11 Egg F24/11 ¹	A/HK 5659/12 MDCK F30/12 ¹	A/St. Afr 3626/13 Egg F03/14 ¹	A/Sbv 2903/2015 Egg F02/16 ²	A/Israel Q-504/15 MDCK F08/16 ²			
REFERENCE VIRUSES																	
A/California/7/2009	clone 38-32	2009-04-09	E3/E3	1280	640	640	640	1280	1280	640	2560	2560	1280	2560	2560	2560	2560
A/Bayem/69/2009		2009-07-01	MDCK5/MDCK1	40	320	80	80	40	40	40	40	40	40	80	80	40	40
A/Lviv/N6/2009		2009-10-27	MDCK4/SIAT1/MDCK3	80	1280	320	320	80	80	80	80	80	80	320	320	160	160
A/Christchurch/16/2010	4	2010-07-12	E1/E3	1280	2560	1280	5120	2560	640	5120	2560	2560	2560	2560	2560	2560	2560
A/Astrakhan/1/2011	5	2011-02-28	MDCK1/MDCK5	1280	640	1280	1280	2560	640	5120	2560	2560	2560	2560	2560	2560	2560
A/St. Petersburg/27/2011	6	2011-02-14	E1/E4	1280	640	1280	1280	1280	640	2560	2560	2560	2560	2560	2560	2560	2560
A/St. Petersburg/100/2011	7	2011-03-14	E1/E4	1280	640	1280	1280	1280	640	2560	2560	2560	2560	2560	2560	2560	2560
A/Hong Kong/5659/2012	6A	2012-05-21	MDCK4/MDCK2	320	160	160	320	320	160	1280	1280	1280	320	640	640	640	640
A/South Africa/3628/2013	6B	2013-06-06	E1/E3	640	640	320	320	640	640	1280	1280	1280	1280	2560	2560	1280	1280
A/Slovenia/2903/2015	clone 37	2015-10-26	E1/E1	2560	1280	1280	1280	2560	1280	5120	2560	2560	2560	5120	2560	5120	5120
A/Israel/Q-504/2015	6B.2	2015-12-15	C1/MDCK2	1280	640	320	320	1280	320	2560	2560	2560	2560	2560	2560	2560	2560
TEST VIRUSES																	
A/Bulgaria/033/2016	6B.1	2016-01-14	SIAT1/MDCK1	640	320	320	320	640	640	1280	1280	1280	640	1280	1280	1280	1280
A/Bulgaria/031/2016	6B.1	2016-01-14	SIAT1/MDCK1	1280	640	640	640	1280	640	2560	2560	2560	1280	2560	2560	2560	2560
A/Bulgaria/037/2016	6B.1	2016-01-15	SIAT1/MDCK1	640	1280	320	320	1280	320	1280	1280	1280	1280	2560	2560	1280	1280
A/Bulgaria/043/2016	6B.1	2016-01-18	SIAT1/MDCK1	320	320	320	320	640	640	1280	1280	1280	640	1280	1280	640	640
A/Bulgaria/042/2016	6B.1	2016-01-18	SIAT1/MDCK1	640	320	320	320	640	320	1280	1280	1280	640	1280	1280	640	640
A/Bulgaria/052/2016	6B.1	2016-01-19	SIAT1/MDCK1	1280	640	640	640	1280	640	2560	2560	2560	1280	2560	2560	1280	1280
A/Bulgaria/099/2016	6B.1	2016-01-22	SIAT2/MDCK1	1280	1280	640	640	1280	640	2560	2560	2560	1280	2560	2560	1280	1280
A/Bulgaria/093/2016	6B.1	2016-01-22	SIAT2/MDCK1	640	320	320	320	640	320	1280	1280	1280	640	2560	2560	1280	1280
A/Bulgaria/129/2016	6B.1	2016-01-25	SIAT2/MDCK1	1280	640	640	640	1280	640	2560	2560	2560	1280	2560	2560	1280	1280
A/Bulgaria/110/2016	6B.1	2016-01-25	SIAT2/MDCK1	640	320	320	320	640	640	1280	1280	1280	640	2560	2560	1280	1280
A/Bulgaria/132/2016	6B.1	2016-01-25	SIAT1/MDCK1	640	640	640	640	1280	640	2560	2560	2560	1280	2560	2560	1280	1280
A/Bulgaria/132/2016	6B.1	2016-01-27	SIAT2/MDCK1	640	640	320	320	640	320	1280	1280	1280	640	2560	2560	1280	1280
A/Bulgaria/127/2016	6B.1	2016-01-27	SIAT2/MDCK1	640	320	320	320	640	640	1280	1280	1280	640	2560	2560	1280	1280
A/Bulgaria/126/2016	6B.1	2016-01-27	SIAT2/MDCK1	1280	640	640	640	1280	640	2560	2560	2560	1280	2560	2560	1280	1280
A/Povazska Bystrica/101/2016	6B.1	2016-02-03	MDCK2/MDCK1	640	320	320	320	640	640	1280	1280	1280	640	2560	2560	1280	1280
A/Trencin/142/2016	6B.1	2016-02-16	MDCKx/MDCK1	1280	640	640	640	1280	640	2560	2560	2560	1280	2560	2560	1280	1280
A/Komarno/148/2016	6B.1	2016-02-16	MDCKx/MDCK1	640	640	640	640	1280	640	1280	1280	1280	640	2560	2560	1280	1280
A/Prievidza/247/2016	6B.1	2016-03-03	MDCKx/MDCK1	320	320	320	320	640	320	1280	1280	1280	640	2560	2560	1280	1280
A/Tinava/273/2016	6B.1	2016-03-08	MDCKx/MDCK1	640	640	320	320	640	320	1280	1280	1280	640	2560	2560	1280	1280
A/Nitra/338/2016	6B.1	2016-03-31	MDCKx/MDCK1	1280	640	640	640	1280	640	2560	2560	2560	1280	2560	2560	1280	1280
A/Bratislava/342/2016	6B.1	2016-04-01	MDCKx/MDCK1	80	320	320	320	640	160	1280	1280	1280	320	640	640	640	1280
Vaccine																	

* Superscripts refer to antiserum properties (< relates to the lowest dilution of antiserum used)

1 <= <40
2 <= <80

Figure 1. Phylogenetic comparison of influenza A(H1N1)pdm09 HA genes



Influenza A(H3N2) virus analyses

As described in many previous reports³, influenza A(H3N2) viruses continue to be difficult to characterise antigenically by HI assay due to variable agglutination of red blood cells (RBCs) from guinea pigs, turkeys and humans or the loss of the ability of viruses to agglutinate any of these RBCs. This is a particular problem for most viruses that fall in genetic subgroup 3C.2a, as was highlighted first in the November 2014 report⁴.

Results of HI tests performed with guinea pig RBCs in the presence of 20nM oseltamivir, added to circumvent NA-mediated binding of A(H3N2) viruses to the RBCs, are shown in Table 4. Twenty-five test viruses retained sufficient HA titre to be analysed by HI assay and 22 of these were characterised genetically, with all falling in subclade 3C.2a.

The test viruses, propagated in MDCK-SIAT1 cells, reacted poorly in HI assays, with the panel of post-infection ferret antisera relative to the titres of the antisera with their respective homologous viruses (shown in red: Table 4). However, the antisera raised against A/Stockholm/6/2014 (3C.3a: tissue culture- and egg-propagated), tissue culture-propagated A/Hong Kong/5738/2014 (3C.2a), tissue culture-propagated A/Georgia/532/2015 and A/Hong Kong/4801/2014 (3C.2a: tissue culture- and egg-propagated), gave reactivity with the vast majority of test viruses. The antiserum raised against egg-propagated A/Switzerland/9715293/2013 (3C.3a), the northern hemisphere 2015–2016 vaccine component, reacted with 10 test viruses at a titre of 40 (32-fold reduced compared to the homologous titre), while antiserum raised against egg-propagated A/Hong Kong/4801/2014, the virus recommended for use in vaccines for the southern hemisphere 2016 and northern hemisphere 2016–2017 influenza seasons, yielded titres reduced by two-to-eightfold with 22 test viruses compared to the homologous titre and gave absolute titres of 40 or greater.

Since 2009, seven genetic groups based on the HA gene have been defined for A(H3N2) viruses. Phylogenetic analysis of the HA genes of representative A(H3N2) viruses with recent collection dates is shown in Figure 2. The HA genes fall within clade 3C. This clade has three subdivisions: 3C.1 (represented by A/Texas/50/2012, the vaccine virus recommended for use in the 2014–15 northern hemisphere season), 3C.2 and 3C.3. Viruses in these three subdivisions had been antigenically similar. In 2014 three new subclades emerged, one in subdivision 3C.2, 3C.2a, and two in 3C.3, 3C.3a and 3C.3b, with subclade 3C.2a viruses dominating in recent months (Figure 2). While viruses in subclades 3C.2a and 3C.3a are antigenic drift variants, those in 3C.3b (e.g. A/Netherlands/525/2014) have remained antigenically similar to previously circulating viruses in the 3C.3 subdivision. Amino acid substitutions that define these subdivisions and subclades are:

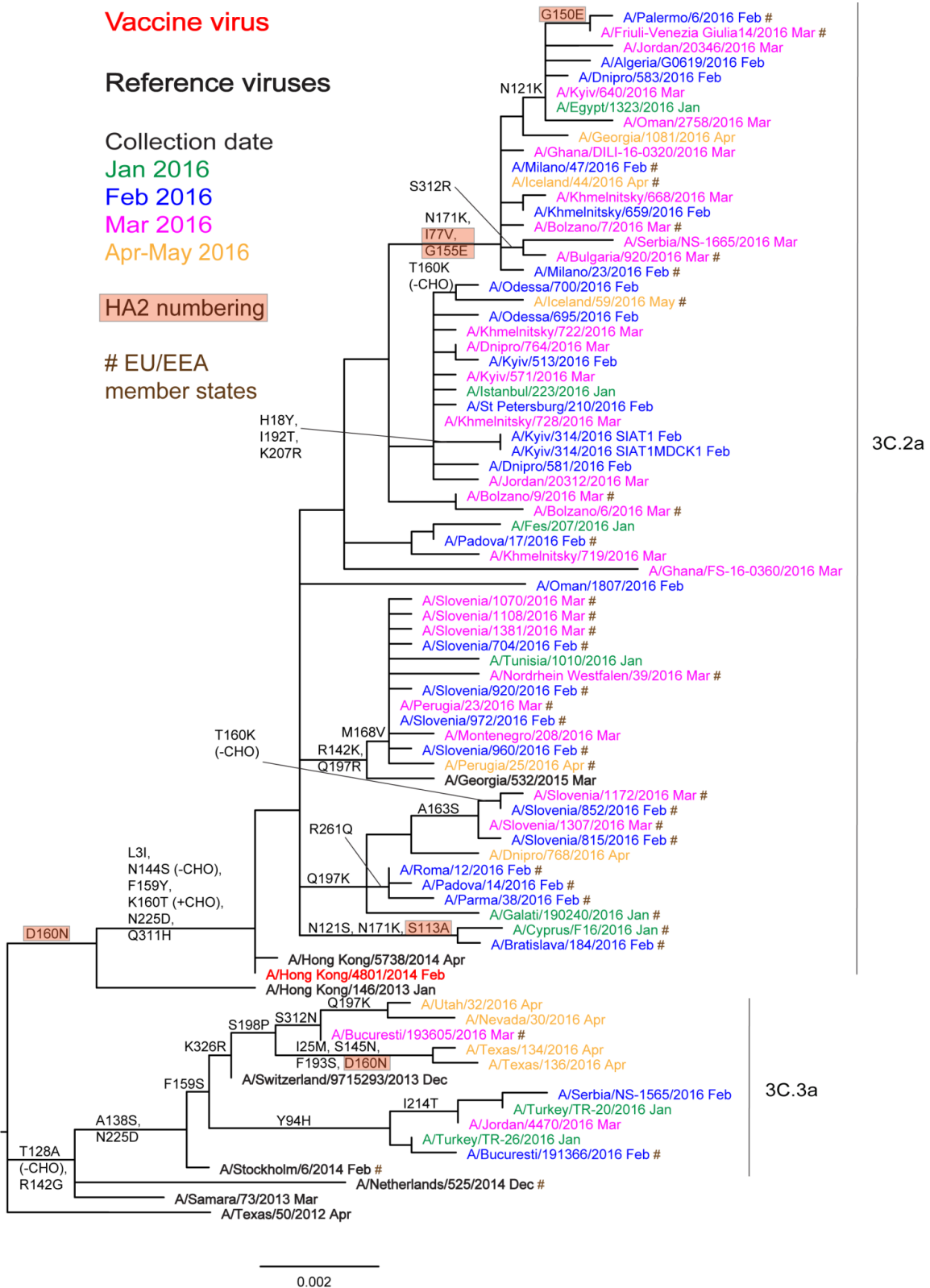
- (3C.2) **N145S** in **HA1**, and **D160N** in **HA2**, e.g. A/Hong Kong/146/2013
- (3C.2a) Those in 3C.2 plus **L3I**, **N144S** (resulting in the loss of a potential glycosylation site), **F159Y**, **K160T** (in the majority of viruses, resulting in the gain of a potential glycosylation site), **N225D** and **Q311H** in **HA1**, e.g. A/Hong Kong/5738/2014
- (3C.3) **T128A** (resulting in the loss of a potential glycosylation site), **R142G** and **N145S** in **HA1**, e.g. A/Samara/73/2013
- (3C.3a) those in 3C.3 plus **A138S**, **F159S** and **N225D** in **HA1**, many with **K326R**, e.g. A/Switzerland/9715293/2013
- (3C.3b) those in 3C.3 plus **E62K**, **K83R**, **N122D** (resulting in the loss of a potential glycosylation site), **L157S** and **R261Q** in **HA1** with **M18K** in **HA2**, e.g. A/Netherlands/525/2014

Based on results available at the time of the February 2015 vaccine composition meeting showing cross-reactivity of antisera raised against subclade 3C.3a and 3C.2a viruses, but with changes acquired on egg-adaptation of genetic subgroup 3C.2a viruses and, at that time, the lack of a suitable 3C.2a vaccine candidate, the World Health Organization recommendation was to use an A/Switzerland/9715293/2013-like virus as the A(H3N2) component of vaccines for the northern hemisphere 2015–16 influenza season [1]. After February 2015, a new subclade designated 3C.3b emerged, these three subclades being antigenically distinguishable, but subclade 3C.2a viruses became prevalent and have remained so. While ferret antisera raised against 3C.3a and 3C.2a subclade viruses showed some cross-reactivity with viruses in all three subclades, antisera raised against 3C.3b viruses were subclade specific. With the availability of new subclade 3C.2a vaccine candidates and the continued cross-reactivity of antisera raised against viruses in subclades 3C.3a and 3C.2a viruses, the World Health Organization recommendation for the A(H3N2) component of influenza vaccines for the southern hemisphere 2016 [2] and northern hemisphere 2016-17 [3] influenza seasons was for an A/Hong Kong/4801/2014-like (3C.2a) virus.

³ For example, the September 2013 report: European Centre for Disease Prevention and Control. Influenza virus characterisation, summary Europe, September 2013. Stockholm: ECDC; 2013. Available from: <http://www.ecdc.europa.eu/en/publications/Publications/influenza-virus-characterisation-sep-2013.pdf>

⁴ European Centre for Disease Prevention and Control. Influenza virus characterisation, summary Europe, November 2014. Stockholm: ECDC; 2014. Available from: http://www.ecdc.europa.eu/en/publications/Publications/ERLI-Net_report_November_2014.pdf

Figure 2. Phylogenetic comparison of influenza A(H3N2) HA genes



Influenza B virus analyses

EU/EEA countries have provided 146 influenza type B-positive specimens with collection dates after 31 December 2015; 144 were ascribed to a lineage: 125 B/Victoria-lineage and 19 B/Yamagata-lineage (Table 2).

Influenza B – Victoria lineage

Since the May 2016 report, 63 viruses of this lineage have been characterised antigenically. HI results are shown in Tables 5-1 to 5-3 with the 59 viruses characterised genetically all falling in clade 1A.

The test viruses showed similar HI reactivity patterns to those from the 2014–2015 influenza season: only 15 (24%) test viruses showed HI titres reduced within fourfold of the titre for the homologous virus with post-infection ferret antisera raised against the recommended vaccine virus for quadrivalent live and inactivated vaccines for the northern hemisphere 2015–2016 influenza season, B/Brisbane/60/2008. The number increased to 43 (68%) when eightfold reductions in HI titre were considered. Similarly, the test viruses were poorly recognised by post-infection ferret antisera raised against reference viruses propagated in eggs (B/Malta/636714/2011, B/Johannesburg/3964/2012 and B/South Australia/81/2012) and cell-propagated B/Formosa/V2367/2013. In contrast, all test viruses showed reactivity within fourfold, the majority within twofold, of the titres for the corresponding homologous viruses with antisera raised against viruses that are considered to be surrogate tissue culture-propagated antigens representing the egg-propagated B/Brisbane/60/2008 prototype virus; these antisera were raised against tissue culture propagated viruses B/Hong Kong/514/2009 (clade 1B), and recently circulating viruses B/Ireland/3154/2016 and B/Nordrhein-Westfalen/1/2016 (both clade 1A).

Phylogenetic analysis of the HA gene of representative B/Victoria lineage viruses is shown in Figure 3. Throughout the previous season and this season to date, viruses from Europe, and elsewhere, have HA genes that fall into the B/Brisbane/60/2008 clade (clade 1A) and remain antigenically similar to the vaccine virus B/Brisbane/60/2008. The great majority of viruses, with collection dates since October 2015, fall in a major subcluster defined by amino acid substitutions I117V, N129D and V146I within clade 1A.

These results, linked with the rise in the proportion of B/Victoria-lineage viruses seen in the 2015 southern hemisphere and 2015–2016 northern hemisphere influenza seasons, support the recommendations made to include B/Brisbane/60/2008 in trivalent influenza vaccines for the southern hemisphere 2016 [2] and northern hemisphere 2016–2017 [3] influenza seasons and in the quadrivalent vaccines.

Influenza B – Yamagata lineage

HI results for nine B/Yamagata-lineage test virus analysed since the May 2016 report are shown in Table 6. Eight of the test viruses were characterised genetically with seven falling in clade 3 and one (B/Poprad/320/2016) in clade 2.

The homologous titres of the 10 post-infection ferret antisera, shown in red, ranged from 80–1280, and all test viruses showed reactivity with nine of the antisera (Table 6).

Antisera raised against egg-propagated clade 3 viruses B/Phuket/3073/2013 (the virus recommended for inclusion in trivalent influenza vaccines for the northern hemisphere 2014–2015 season) and B/Hong Kong/3417/2014 recognised the test viruses at titres within twofold of their respective homologous titres. Only four of the test viruses showed HI reactivity within fourfold, compared to the homologous titre, with antiserum raised against egg-propagated B/Massachusetts/02/2012, the clade 2 vaccine virus recommended for use in the 2014–2015 northern hemisphere influenza season. Two viruses, B/Poprad/320/2016 (clade 2) and B/Milano/101/2016 (clade 3 but carrying a B/Victoria-lineage NA), showed high reactivity with antisera raised against two cell-propagated clade 2 viruses, B/Estonia/55669/2011 and B/Massachusetts/02/2012.

Figure 4 shows a phylogenetic analysis of the HA genes of representative B/Yamagata-lineage viruses. Worldwide, the vast majority of HA genes from recently collected viruses have fallen in the B/Wisconsin/1/2010–B/Phuket/3073/2013 clade (clade 3), with the great majority falling in a subgroup defined by HA1 L172Q amino acid substitution. A few viruses, annotated in the phylogenetic tree and including B/Milano/101/2016, are reassortants carrying NA genes normally associated with the B/Victoria-lineage.

Based on such results, a B/Phuket/3073/2013-like virus has been recommended for inclusion in quadrivalent vaccines for the 2016 southern hemisphere [2] and 2016–2017 northern hemisphere [3] influenza seasons.

Table 5-1. Antigenic analysis of influenza B/Victoria-lineage viruses by HI

Viruses	Haemagglutination inhibition titre											
	B/Bris	B/Mal	B/Bris	B/Mal	B/Jhb	B/For	B/Sh Aus	B/HK	B/Ireland	B/Nord-West	Passage history	Collection date
REFERENCE VIRUSES												
B/Malaysia/2506/2004	2560	640	20	80	160	80	160	20	<	<	E3/E7	2004-12-06
B/Brisbane/60/2008	5120	80	160	80	640	320	1280	160	40	40	E4/E4	2008-08-04
B/Mal/636714/2011	2560	80	80	80	320	320	640	80	40	40	E4/E1	2011-03-07
B/Johannesburg/3964/2012	5120	640	640	1280	1280	1280	1280	320	320	320	E1/E2	2012-08-03
B/Formosa/V2367/2012	5120	40	80	320	320	320	640	80	80	80	MDCK1/MDCK3	2012-08-06
B/South Australia/81/2012	5120	80	160	320	320	320	1280	80	40	40	E4/E2	2012-11-28
B/Hong Kong/514/2009	2560	<	<	40	20	40	80	80	80	80	MDCK3	2009-10-11
B/Ireland/3154/2016	5120	<	<	40	20	80	80	80	80	80	MDCK1/MDCK1	2016-01-14
B/Nordrhein-Westfalen/1/2016	5120	<	<	40	40	80	80	80	80	80	C2/MDCK1	2016-01-04
TEST VIRUSES												
B/Nira/1/25/2016	5120	<	<	40	40	80	80	80	80	80	MDCK1/MDCK1	2016-02-09
B/Levice/134/2016	5120	<	<	40	40	80	80	80	80	80	MDCKx/MDCK1	2016-02-11
B/Kosice/150/2016	5120	<	<	40	20	40	80	80	80	80	MDCKx/MDCK1	2016-02-11
B/Tmava/192/2016	5120	<	<	40	40	80	80	80	80	80	MDCKx/MDCK1	2016-02-22
B/Breitslava/226/2016	2560	<	<	20	20	40	80	40	40	40	MDCKx/MDCK1	2016-02-29
B/Topolcan/240/2016	5120	<	<	40	40	80	80	80	80	80	MDCKx/MDCK1	2016-03-02
B/Banska Bystrica/306/2016	5120	<	<	40	40	80	80	80	80	80	MDCKx/MDCK1	2016-03-04
B/Trencin/266/2016	5120	<	<	40	40	80	80	80	80	80	MDCKx/MDCK1	2016-03-10
B/Piestany/296/2016	5120	<	<	40	40	80	80	80	80	80	MDCKx/MDCK1	2016-03-14
B/Banska Bystrica/336/2016	5120	<	<	20	20	80	80	80	80	80	MDCKx/MDCK1	2016-03-15

Vaccine
NH 2015-16[#]
SH 2016
NH 2016-17

¹ < =

* Superscripts refer to antiserum properties (< relates to the lowest dilution of antiserum used) <40; ² < = <10; ³ hyperimmune sheep serum

[#] B/Victoria-lineage virus recommended for use in quadrivalent vaccines

Table 5-2. Antigenic analysis of influenza B/Victoria-lineage viruses by HI

Viruses	Hemagglutination inhibition titre											
	Post-infection ferret antisera											
	B/Bris 6008 Egg Sh 539, 540, 543, 544, 570, 571, 574 ^{1,3}	B/Mal 250604 Egg F37/11 ²	B/Bris 6008 Egg F26/13 ³	B/Mal 630714/11 Egg F29/13 ²	B/Jhb 3964/12 Egg F04/16 ²	B/For V2367/12 MDCK F04/13 ²	B/SH Aus 8/12 Egg F4/113 ²	B/HK 514/09 MDCK F09/13 ²	B/Ireland 3154/16 MDCK F15/16 ²	B/Nord-West 1/16 MDCK F16/16 ²		
Passage history	Collection date	Passage history	Collection date	Passage history	Collection date	Passage history	Collection date	Passage history	Collection date	Passage history	Collection date	Passage history
REFERENCE VIRUSES												
B/Malaysia/2506/2004	2560	640	80	80	40	80	160	20	20	80	20	<
B/Brisbane/60/2008	2560	80	640	320	320	320	1280	80	80	40	40	<
B/Mal/63/71/2011	2560	80	640	640	320	320	1280	80	80	40	40	<
B/Johannesburg/9864/2012	5120	640	1280	1280	1280	1280	1280	320	320	160	160	160
B/Formosa/V2367/2012	5120	40	640	160	160	320	640	80	80	40	40	40
B/South Australia/81/2012	2012-11-28	80	80	320	320	320	1280	80	80	40	40	40
B/Hong Kong/514/2009	2009-10-11	2560	<	80	40	40	40	80	80	80	80	40
B/Ireland/3154/2016	2016-01-14	5120	10	80	40	40	80	80	80	80	80	40
B/Nordrhein-Westfalen/1/2016	2016-01-04	2560	<	80	40	80	80	80	80	80	80	80
TEST VIRUSES												
B/Paderborn/2/2016	2016-02-02	MDCK3/MDCK1	2560	10	160	80	80	80	80	40	40	80
B/France/2/2016	2016-02-02	MDCK2/MDCK1	2560	<	40	40	40	40	40	40	40	40
B/Lithuania/3/663/2016	2016-02-02	MDCK3/MDCK1	2560	<	160	20	40	40	40	80	40	20
B/Lithuania/4/516/2016	2016-02-07	MDCK3/MDCK1	2560	<	80	20	40	40	40	40	40	20
B/Romania/6/2016	2016-02-08	C2/MDCK1	5120	20	320	40	80	80	80	80	80	40
B/Panama/14/2016	2016-02-24	MDCK1/MDCK1	2560	10	160	20	80	40	40	40	40	40
B/Palermo/4/2016	2016-03-02	MDCK3/MDCK1	2560	10	160	20	80	40	40	40	40	40
B/Milano/14/5/2016	2016-03-07	MDCK1/MDCK1	5120	10	160	20	80	40	40	40	40	40
B/Fiji/Venetsia Giulis22/2016	2016-03-11	MDCK1/MDCK1	5120	10	80	40	80	80	80	80	80	40
B/Palermo/6/2016	2016-03-11	MDCK3/MDCK1	5120	10	80	40	80	80	80	80	80	40
B/Czech Republic/11/2016	2016-03-15	C2/MDCK1	5120	20	80	20	80	40	40	40	40	20
B/Czech Republic/11/2016	2016-03-15	C2/MDCK1	2560	10	80	20	20	20	20	40	40	20
B/Czech Republic/11/2016	2016-03-17	C2/MDCK1	5120	10	160	40	80	80	80	80	80	40
B/Czech Republic/11/2016	2016-03-17	MDCK1/MDCK1	2560	10	40	20	40	40	40	40	40	20
B/Czech Republic/11/2016	2016-03-18	C2/MDCK1	2560	<	80	20	40	40	40	40	40	20
B/Czech Republic/11/2016	2016-03-21	C3/MDCK1	2560	40	80	40	80	80	80	80	80	40
B/Czech Republic/11/2016	2016-03-21	C2/MDCK1	2560	<	80	20	40	40	40	40	40	40
B/Czech Republic/11/2016	2016-03-22	SIAT1/MDCK1	2560	10	160	20	40	40	40	40	40	40
B/Czech Republic/11/2016	2016-03-23	SIAT1/MDCK1	2560	<	160	20	40	40	40	40	40	20
B/Czech Republic/11/2016	2016-03-23	SIAT1/MDCK1	2560	<	160	20	40	40	40	40	40	40
B/Czech Republic/11/2016	2016-03-24	C2/MDCK1	2560	20	80	20	80	80	80	80	80	40
B/Czech Republic/11/2016	2016-03-29	C2/MDCK1	2560	10	80	20	80	80	80	80	80	40
B/Czech Republic/11/2016	2016-04-06	SIAT2/MDCK1	1280	1280	80	20	40	40	40	40	40	20
B/Czech Republic/11/2016	2016-04-11	MDCK1/MDCK1	2560	10	80	40	80	80	80	80	80	40
B/Czech Republic/11/2016	2016-04-11	SIAT2/MDCK1	2560	10	160	20	40	40	40	40	40	20
B/Czech Republic/11/2016	2016-04-11	SIAT2/MDCK1	2560	<	160	20	40	40	40	40	40	20
B/Czech Republic/11/2016	2016-04-12	MDCK1/MDCK1	2560	<	80	40	80	80	80	80	80	40
B/Czech Republic/11/2016	2016-04-15	MDCK1/MDCK1	2560	10	80	20	40	40	40	40	40	40
B/Czech Republic/11/2016	2016-04-18	SIAT1/MDCK1	2560	<	40	10	40	40	40	40	40	20
B/Czech Republic/11/2016	2016-04-26	MDCK1/MDCK1	2560	<	80	20	40	40	40	40	40	20
B/Czech Republic/11/2016	2016-05-02	MDCK1/MDCK1	2560	10	160	40	80	80	80	80	80	40
B/Czech Republic/11/2016	2016-05-11	MDCK1/MDCK1	2560	10	80	20	40	40	40	40	40	40
B/Czech Republic/11/2016	2016-06-05	MDCK1	2560	<	160	20	40	40	40	40	40	80

Vaccine NH
2015-16¹ SH 2016
NH 2016-17

¹ < = <40; ² < = <10; ³

* Superscripts refer to antiserum properties (< relates to the lowest dilution of antiserum used)
Hyperimmune sheep serum

[†] B/Victoria-lineage virus recommended for use in quadrivalent vaccines

Table 5-3. Antigenic analysis of influenza B/Victoria-lineage viruses by HI

Viruses	Passage history	Collection date	Haemagglutination inhibition titre														
			B/Bris 60/08 Egg	B/Mal 2506/04 Egg	B/Bris 6008 Egg	B/Mal 636714/11 Egg	B/Jhb 3964/12 Egg	B/For V2387/12 MDCK	B/Sh Aus 81/12 Egg	B/HK 514/09 MDCK	B/Ireland 3154/16 MDCK	B/NordWest 1/16 MDCK	Genetic group				
REFERENCE VIRUSES																	
B/Malaysia/2506/2004		2004-12-06	1280	640	40	80	40	80	40	160	20	<	<				
B/Brisbane/602/08		2008-08-04	2560	80	640	320	160	320	160	1280	80	40	40	20			
B/Mal/636714/2011		2011-03-07	2560	80	640	320	320	320	1280	1280	80	40	40	40			
B/Johannesburg/3964/2012		2012-08-03	5120	640	1280	1280	1280	1280	1280	1280	320	160	160	160			
B/Formosa/V2387/2012		2012-08-06	5120	40	640	160	320	320	320	640	80	40	40	20			
B/South Australia/812/012		2012-11-28	2560	80	160	320	320	40	40	80	80	40	40	20			
B/Hong Kong/514/2009		2009-10-11	2560	<	80	40	40	40	40	80	80	40	40	40			
B/Ireland/3154/2016		2016-01-14	5120	<	80	40	40	40	40	80	40	80	80	40			
B/Nordrhein-Westfalen/1/2016		2016-01-04	5120	<	80	40	40	40	80	80	80	80	80	40			
TEST VIRUSES																	
B/Stockholm/1/1/2016		2016-02-08	5120	10	160	80	80	80	80	160	40	20	20	40			
B/Stockholm/12/2016		2016-02-08	5120	10	40	20	40	40	40	40	40	40	40	40			
B/Slovenia/701/2016		2016-02-09	5120	<	40	40	40	40	40	80	40	40	40	40			
B/Slovenia/914/2016		2016-02-21	5120	<	40	40	40	40	40	80	80	80	80	40			
B/Slovenia/1038/2016		2016-02-26	5120	<	40	40	40	40	40	80	80	80	80	40			
B/Stockholm/15/2016		2016-03-01	5120	<	80	40	40	40	40	80	40	40	40	80			
B/Finland/588/2016		2016-03-01	5120	<	80	40	40	40	40	80	40	40	40	40			
B/Slovenia/1171/2016		2016-03-09	5120	<	80	40	40	40	40	80	80	80	80	40			
B/Slovenia/1186/2016		2016-03-09	5120	<	80	40	40	40	40	80	80	80	80	40			
B/Finland/592/2016		2016-03-11	5120	10	80	40	40	40	40	80	80	80	80	40			
B/Slovenia/1228/2016		2016-03-14	5120	<	40	40	40	40	40	80	80	80	80	40			
B/Slovenia/1237/2016		2016-03-15	5120	<	40	40	40	40	40	80	80	80	80	40			
B/Stockholm/16/2016		2016-03-16	5120	<	80	40	40	40	40	80	40	40	40	40			
B/Slovenia/1311/2016		2016-03-21	5120	10	80	40	40	40	40	160	80	80	80	40			
B/Finland/623/2016		2016-03-21	5120	<	80	40	40	40	40	80	80	80	80	40			
B/Lisboa/13/2016		2016-04-06	5120	<	80	40	40	40	40	80	80	80	80	40			
B/Slovenia/1459/2016		2016-04-06	5120	<	80	40	40	40	40	160	80	80	80	40			
B/Slovenia/1523/2016		2016-04-16	5120	<	80	40	40	40	40	160	80	80	80	40			
B/Slovenia/1604/2016		2016-05-01	2560	<	80	40	40	40	40	80	80	80	80	40			
B/Slovenia/1601/2016		2016-05-01	5120	<	80	40	40	40	40	80	80	80	80	40			

Vaccine NH
2015-16¹ SH 2016
NH 2016-17

¹ < = <40, ² < = <10, ³

* Superscripts refer to antiserum properties (< relates to the lowest dilution of antiserum used)
hyperimmune sheep serum

³ B/Victoria-lineage virus recommended for use in quadrivalent vaccines

Figure 3. Phylogenetic comparison of influenza B/Victoria-lineage HA genes

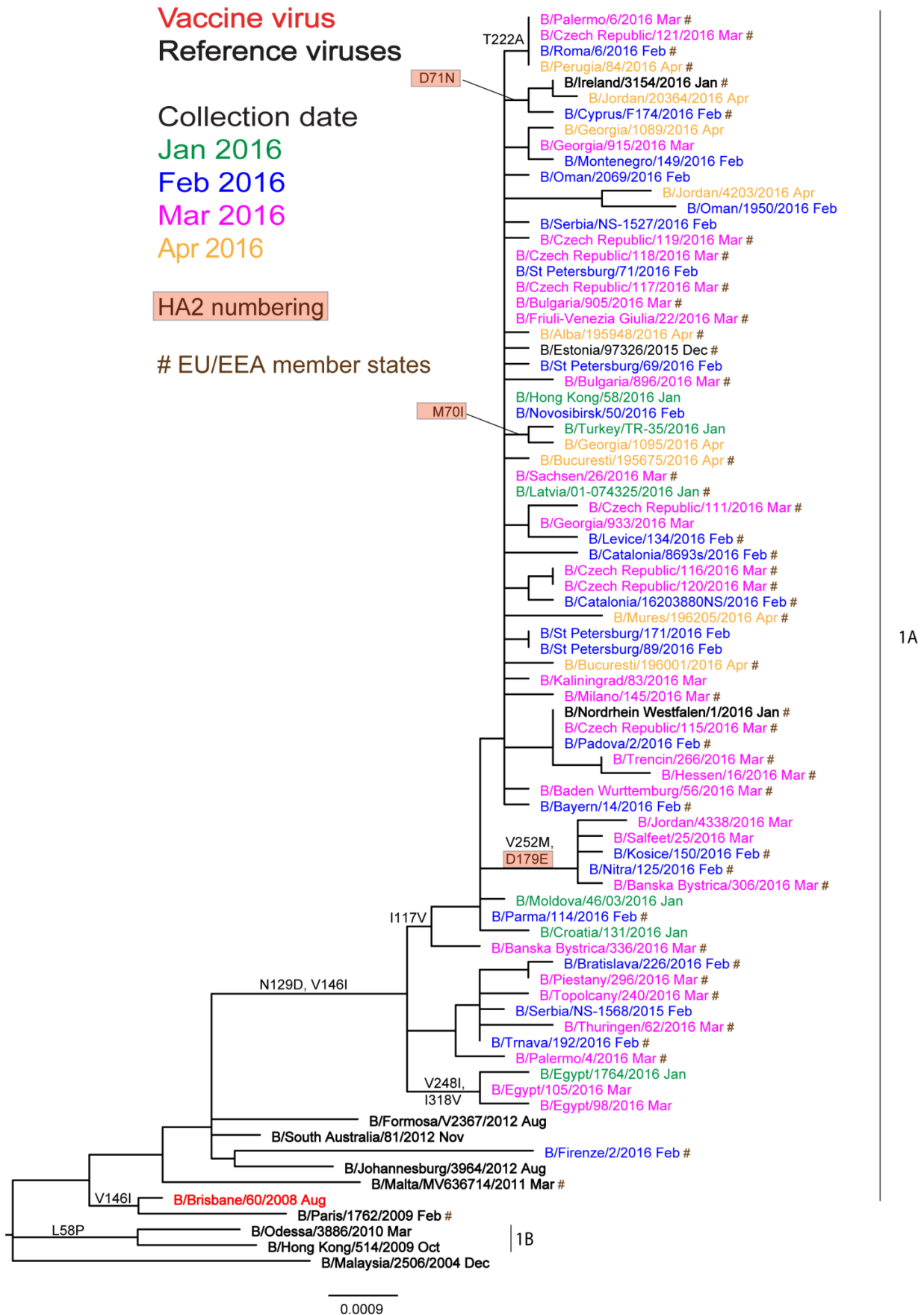


Table 6. Antigenic analysis of influenza B/Yamagata-lineage viruses by HI

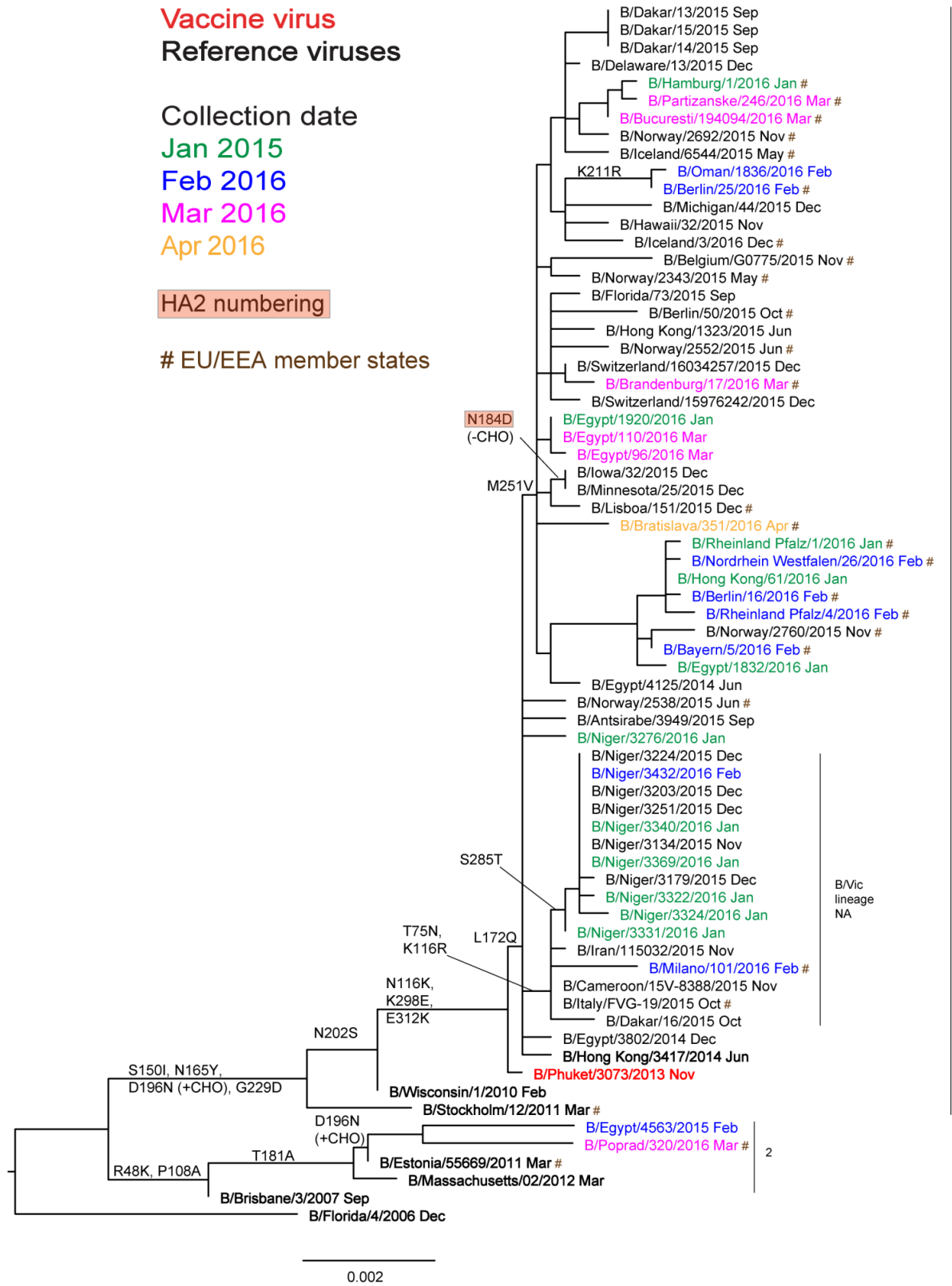
Viruses	Haemagglutination inhibition titre											
	Post-infection ferret antisera											
	B/Phuket	B/FI	B/Bris	B/Estonia	B/Mass	B/Mass	B/Wis	B/Stock	B/Phuket	B/Phuket	B/HK	
	3073/13	4/06	3/07	55669/11	02/12	02/12	1/10	12/11	3073/13	3073/13	3417/14	
Passage history	Egg	Egg	Egg	MDCK	MDCK	Egg	Egg	Egg	MDCK	Egg	Egg	
Ferret number	SH614 ^{1,3}	F17/13 ¹	F38/14 ²	F32/12 ²	F05/15 ²	F42/14 ²	F10/13 ²	F06/15 ¹	F35/14 ²	F36/14 ²	St-Judes	
Genetic Group	3	1	2	2	2	2	3	3	3	3	F715/14 ^{2,4}	3
REFERENCE VIRUSES												
B/Florida/4/2006	1280	1280	640	80	80	1280	160	160	20	160	160	160
B/Brisbane/3/2007	1280	640	320	40	80	640	80	80	10	80	80	160
B/Estonia/65669/2011	640	80	40	80	160	80	40	10	40	40	80	80
B/Massachusetts/02/2012	1280	640	320	320	320	640	160	80	80	160	160	160
B/Massachusetts/02/2012	1280	640	320	40	80	640	80	80	10	80	160	160
B/Wisconsin/1/2010	2560	320	160	10	10	160	160	80	20	160	160	160
B/Stockholm/12/2011	1280	160	80	10	<	80	80	80	20	80	80	80
B/Phuket/3073/2013	5120	160	160	160	320	320	160	80	320	320	160	160
B/Phuket/3073/2013	5120	320	160	20	<	320	320	160	40	160	320	320
B/Hong Kong/3417/2014	1280	80	40	<	<	40	40	20	10	40	160	160
TEST VIRUSES												
B/Iceland/25/2016	2560	160	80	80	40	160	160	40	160	160	160	160
B/Milano/101/2016	5120	320	160	320	640	320	320	160	1280	640	160	160
B/Partizanske/246/2016	2560	80	40	40	20	80	80	40	80	80	160	160
B/Poprad/320/2016	2560	80	160	320	320	160	80	40	160	80	320	320
B/Stockholm/14/2016	5120	160	160	40	20	320	320	80	160	160	320	320
B/Bratislava/351/2016	2560	80	80	40	<	80	160	40	160	80	160	160
B/Iceland/49/2016	2560	80	80	80	80	80	160	40	160	160	160	160
B/Slovenia/1471/2016	5120	80	80	40	<	80	160	40	80	80	160	160
B/Norway/3694/2016	2560	80	80	40	<	80	160	40	80	80	160	160

Vaccine
NH 2015-16
SH2016[#]
NH 2016-17[#]

* Superscripts refer to antiserum properties (< relates to the lowest dilution of antiserum used)
1 < + <40; 2 < + <10; 3 hyperimmune sheep serum; 4 RDE serum pre-absorbed with TRBC

B/Yamagata-lineage virus recommended for use in quadrivalent vaccines

Figure 4. Phylogenetic comparison of influenza B/Yamagata-lineage HA genes



Summary of genetic data submitted to TESSy

For the period covering weeks 40/2015–20/2016, 2601 viruses have been characterised genetically: 1770 A(H1N1)pdm09 clade 6B represented by A/South Africa/3626/2013 (6B.1 and 6B.2 subclade designations were not available as reporting categories at the start of the 2015–2016 influenza season); 211 A(H3N2) subclade 3C.2a represented by A/Hong Kong/4801/2014, 65 subclade 3C.3a represented by A/Switzerland/9715293/2013, two subclade 3C.3b represented by A/Stockholm/28/2014, and two subclade 3C.3 represented by A/Samara/73/2013; 496 B/Victoria-lineage clade 1A represented by B/Brisbane/60/2008; and 55 B/Yamagata-lineage clade 3 represented by B/Phuket/3073/2013.

Antiviral susceptibility

For weeks 40/2015–20/2016 of the 2015–2016 influenza season, countries reported on the antiviral susceptibility of 2700 A(H1N1)pdm09 viruses, 172 A(H3N2) viruses and 523 influenza type B viruses from sentinel and non-sentinel sources. All but 31 showed no molecular or phenotypic evidence of reduced inhibition (RI) by neuraminidase inhibitors (oseltamivir and zanamivir). Twenty-six A(H1N1)pdm09 viruses carried NA H275Y amino acid substitution associated with highly reduced inhibition (HRI) by oseltamivir, one A(H3N2) virus showed RI by oseltamivir associated with NA-E119V amino acid substitution and four B/Victoria-lineage viruses showed HRI by both drugs due to NA-R374K amino acid substitution.

Phenotypic testing for susceptibility to oseltamivir and zanamivir has been conducted on 673 viruses at the WIC: 404 A(H1N1)pdm09, 110 A(H3N2), 136 B/Victoria-lineage and 23 B/Yamagata-lineage viruses. All but one A(H1N1)pdm09 virus showed normal inhibition (NI) by these neuraminidase inhibitors: A/Bayern/151/2015 showed reduced inhibition (RI) by zanamivir and carried NA I117R amino acid substitution.

Influenza A(H7N9) virus

On 1 April 2013, the World Health Organization (WHO) Global Alert and Response [4] reported that the China Health and Family Planning Commission notified the WHO of three cases of human infection with influenza A(H7N9). The cases were confirmed by laboratory testing on 29 March 2013 by the Chinese CDC. A description of the characteristics of H7N9 viruses can be found on the WHO website [5]. Increased numbers of cases were reported over the course of the 2013–14, 2014–15 and 2015–2016 seasons and cases have been reported recently [6]. A revised Rapid Risk Assessment [7] for these A(H7N9) viruses was carried out by ECDC and posted on 11 February 2015. WHO posted a summary of human infection on 31 January 2014 [8], updated on 13 June 2016 [9] with 11 new cases since the report of 09 May 2016, and conducted a risk assessment on 23 February 2015 [10]. In light of the assessment, WHO advised that countries continue to strengthen influenza surveillance. WHO last summarised the numbers of cases of human infection related to their geographic location on 14 July 2014 [11] and has provided subsequent situation updates, with the latest being on 01 July 2016 [6].

Influenza A(H5) virus

The most recent monthly risk assessment of influenza at the human-animal interface was published by WHO on 13 June 2016 [9]. Since the last WHO Influenza update on 09 May 2016, one laboratory-confirmed human case of avian influenza A(H5N6) virus infection in China has been reported to WHO. One human case of A(H5N1) infection in Egypt has been reported for the same period. ECDC published an updated rapid risk assessment on the situation in Egypt on 13 March 2015 [12] and an epidemiological update 10 April 2015 [13]. On 02 December 2015 ECDC published a rapid risk assessment related to identification highly pathogenic H5 viruses in poultry in France [14].

WHO CC reports

A description of results generated by the WHO Collaborating Centre for Reference and Research on Influenza at the Crick Worldwide Influenza Centre, the Francis Crick Institute, Mill Hill Laboratory and used at the WHO Vaccine Composition Meetings held in Memphis, USA, 21–23 September 2015, and at WHO Geneva, 22–24 February 2016, can be found at:

https://www.crick.ac.uk/media/273950/crick_sep2015_vcm_report_to_post.pdf and
https://www.crick.ac.uk/media/286458/crick_feb2016_vcm_report_to_post.pdf

Note on the figures

The phylogenetic trees were constructed using [RAxML](#), drawn using [FigTree](#) and annotated using Adobe Illustrator. The bars indicate the proportion of nucleotide changes between sequences. Reference strains are viruses to which post-infection ferret antisera have been raised. The colours indicate the month of sample collection. Isolates from WHO NICs in EU/EEA countries are marked (#). Sequences for some viruses from non-EU/EEA countries were recovered from GISAID. We gratefully acknowledge the authors, originating and submitting laboratories of the sequences from GISAID's EpiFlu database which were downloaded for use in the preparation of this report (all submitters of data may be contacted directly via the [GISAID website](#)), along with all laboratories who submitted sequences directly to the London WHO Collaborating Centre.

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