Febrile Respiratory Illness (FRI) Surveillance Update

Operational Infectious Diseases, Naval Health Research Center, San Diego, CA





2016 Week 38 (through 24 September 2016)

Contact Information

Address: Commanding Officer Attn: Code 166 Naval Health Research Center 140 Sylvester Rd. San Diego, CA 92106

Web: Click here

Email: Click here

Items of Note

- FRI rates at all basic training centers are at/below expected values
- NHRC is able to test for novel influenza strains, MERS coronavirus, enterovirus EV-68, Ebola virus, and Zika virus

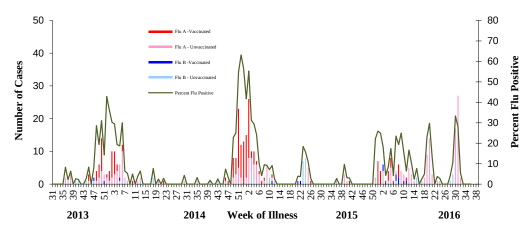
Influenza

NHRC Laboratory-Confirmed Influenza Cases, US Military Basic Trainees

		Curre	nt Week*			Since	e Oct.1, 201	5	
	A /				A /				No.
Site	Untyp.	A/H3	A/H1	В	Untyp.	A/H3	A/H1	В	Tested
Ft. Benning							1	5	207
Ft. Jackson							3		197
Ft. Leonard Wood						1	7	4	172
NRTC Great Lakes							3	1	162
Lackland AFB							1	1	26
MCRD Parris Island						30	6		186
MCRD San Diego						18	1	4	458
CGTC Cape May					2	47	3	9	195
Total	0	0	0	0	2	96	25	24	1603

- NHRC also conducts shipboard FRI surveillance that includes large-deck ships of the US Fleet Forces (Atlantic), 3rd (Pacific), and 7th (Far East) Fleets.
- NHRC conducts FRI surveillance among (non-active duty) DoD beneficiaries in Illinois and San Diego (details on page 7).
- NHRC-CDC-BIDS collaborative border FRI/SARI surveillance program is ongoing at 7 US-Mexico border sites in San Diego (2) and Imperial (5) counties in California. Details on page 8.
- For more information about NHRC FRI surveillance programs, please contact NHRC.

Vaccination Status of Confirmed Influenza Cases Among US Military Basic Trainees, 2013-16

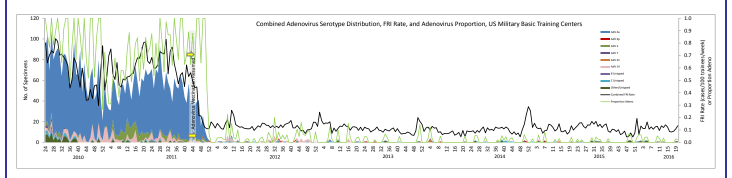


Other Items of Interest

NHRC is conducting laboratory-based surveillance for meningococcal disease. The program's purpose
is to track and characterize meningococcal cases among DoD medical beneficiaries. For more
information and the most recent data, click here.

Adenovirus

- Vaccination against types 4 and 7 adenovirus was instituted at all basic training centers by mid-November 2011.
- FRI rates and the proportion of FRI cases positive for adenovirus have decreased markedly since vaccine was reintroduced.
- Sporadic adenovirus cases at basic training centers 2012-16. FRI rates remain low in general.



FRI Rates

- FRI surveillance is ongoing at 8 U.S. military basic training centers, representing all service branches. As each week's FRI count is reported, FRI Rate Status is classified into one of 3 categories:
 - At or below expected value (expected value shown as dashed line)
 - Moderately elevated
 - Substantially elevated

Week ending 24 September 2016:

- At or below expected value:

Fort Benning

Fort Jackson

Fort Leonard Wood

Naval Recruit Training Command, Great Lakes

Marine Corps Recruit Depot, San Diego

Marine Corps Recruit Depot, Parris Island

Lackland Air Force Base (data through 10 Sept.)

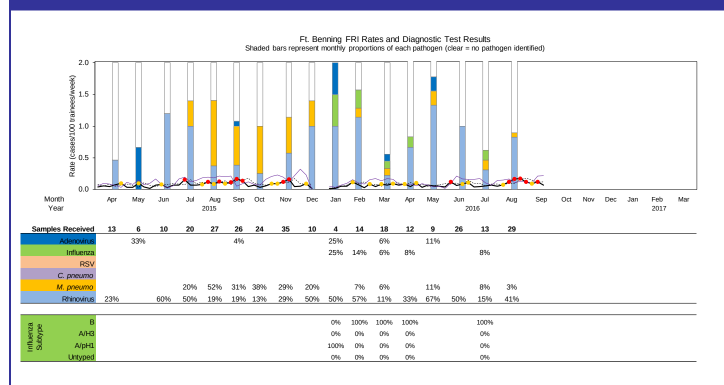
Coast Guard Training Center, Cape May

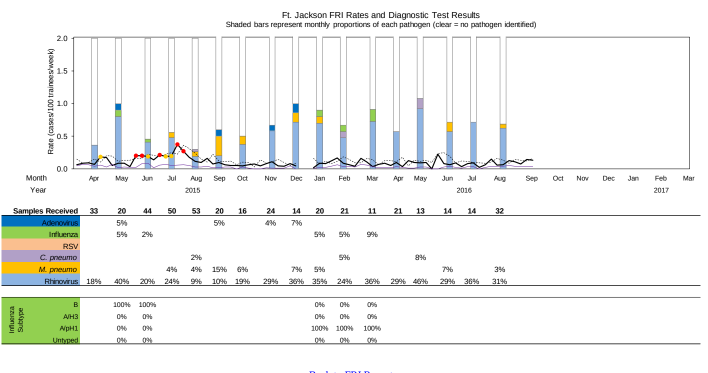
Moderately elevated:

None

• Substantially elevated:

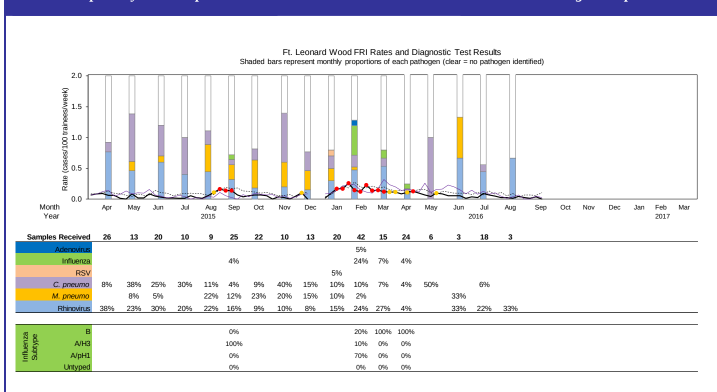
None

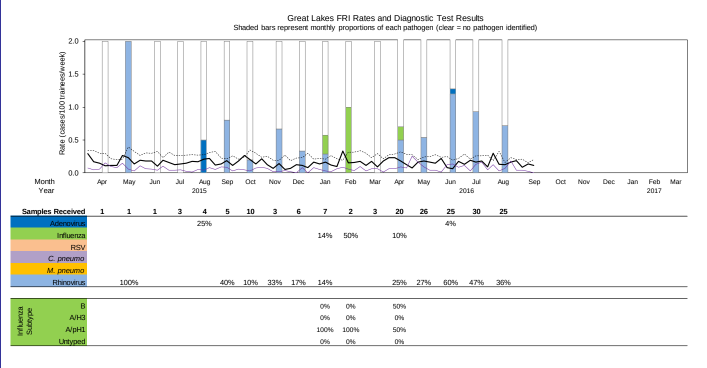




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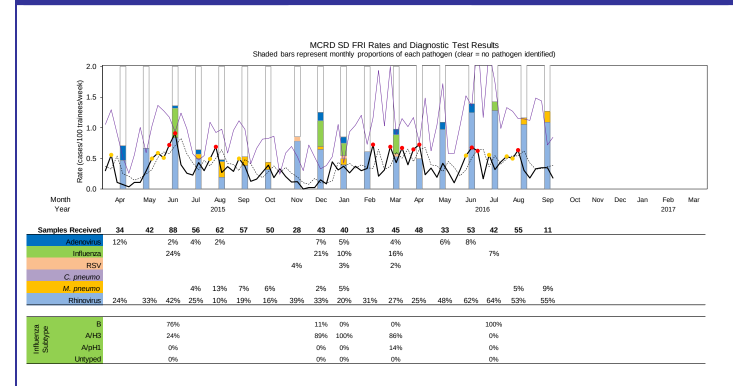
- Observed FRI rate (expected rate = dashed line) • Moderately elevated • Substantially elevated — Pneumonia rate (incl. afebrile)

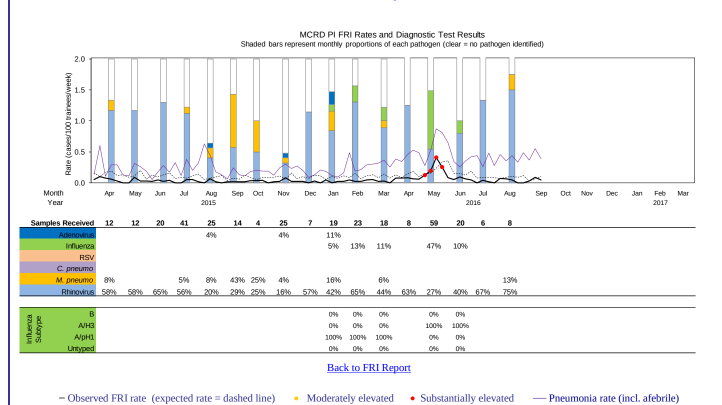


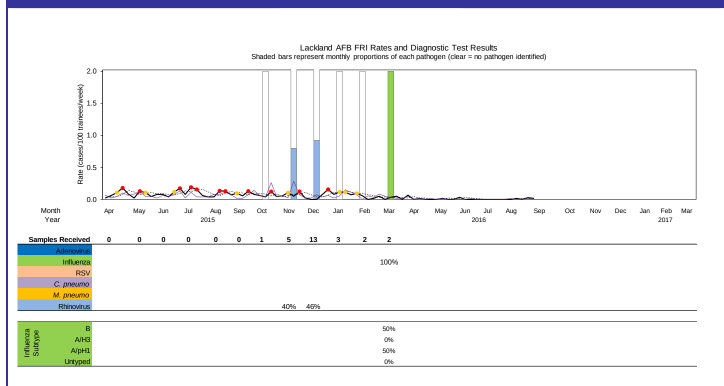


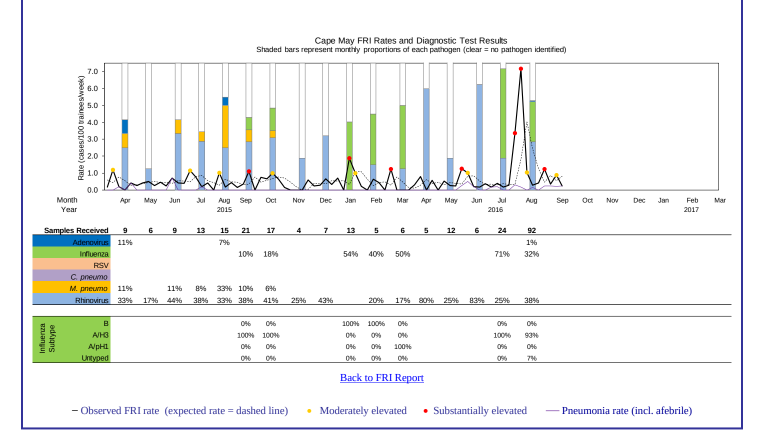
Back to FRI Report

- Observed FRI rate (expected rate = dashed line) • Moderately elevated • Substantially elevated — Pneumonia rate (incl. afebrile)



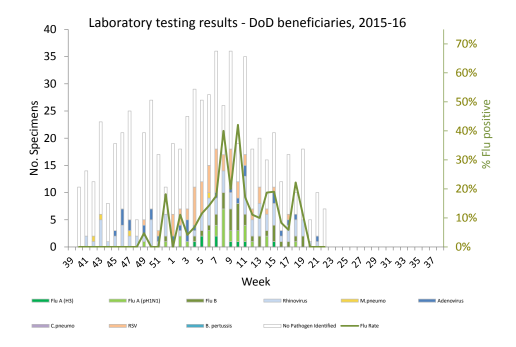






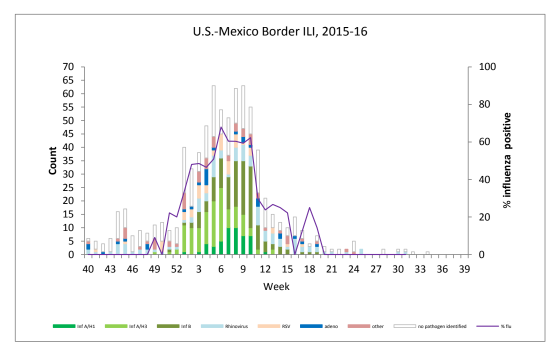
DoD Beneficiary Surveillance

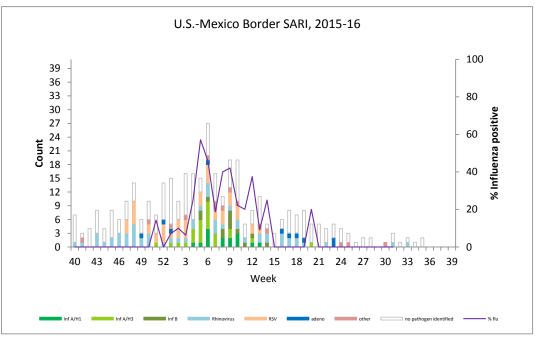
- NHRC conducts FRI surveillance among (non-active duty) DoD beneficiaries at Lovell FHCC (Great Lakes, IL) and in 4 NMC San Diego facilities: the Emergency Department, Pediatric Department, Naval Branch Health Clinic Kearny Mesa, and NH Camp Pendleton.
 - For questions regarding surveillance in this population, please contact the principal investigator (Chris Myers, christopher.a.myers48.civ@mail.mil) or the study coordinator (Erin Hansen, erin.a.hansen.ctr@mail.mil).



US-Mexico Border Surveillance

• In collaboration with the CDC Border Infectious Disease Surveillance program and San Diego/Imperial Counties, NHRC performs laboratory testing for FRI cases among civilians near the US-Mexico border. Both outpatient (FRI) and inpatient (Severe Acute Respiratory Illness; SARI) cases are included in the program. Current sites are located in San Diego (2) and Imperial (5) counties in California.





Phylogenetic Comparison of Influenza A-pandemic 2009 H1N1 (pH1N1) HA and NA Protein Sequences

- 47 analyzed Influenza A pH1N1 HA sequences were derived from MDCK isolates and 4 belonged to clade 6B while 43 belonged to the subclade 6B.1, defined by mutations S84N, S162T (ADD GLY) and I216T.
- 46 analyzed Influenza A pH1N1 NA sequences were derived from MDCK isolates.
- Phylogenetic trees for both HA and NA protein sequences were generated by Clustal V method using DNASTAR® Lasergene Megalign software.
- Amino acid changes shown (both HA and NA sequences) are with respect to A/California/07/2009-like virus.

Summary of Influenza A (pH1N1) Protein Homology When Compared to 2015-2016 Vaccine Strain

Segment	No. Isolates	2015-2016 Vaccine Strain	Protein Homology
A/pH1N1 HA	47	A/California/07/2009	97.0-98.0%
A/pH1N1 NA	46	A/California/07/2009	96.1-97.3%

Summary of Influenza A (pH1N1) N-Linked Glycosylation Mutations

Loss or gain of N-linked glycosylation sites affect host innate immune system recognition and the ability to
induce adaptive immune response thus altering its viral antigenicity.¹ Predicted loss or gain of N-linked
glycosylation of protein sequences were calculated using CBS NetNGlyc 1.0 Server
http://www.cbs.dtu.dk/services/NetNGlyc/

	Mutation		
A/pH1N1 Segment	ADD GLY	LOSS GLY	
HA	S162N		
NA	N44S, S70N	N386K	

Sun X. Jayaraman A. Mainprasad P. Raman R. Houser K. Pappas C. ... Tumpey T. (2013).
 N-Linked Glycosylation of the Hemagglutin Protein Influences Virulence and Antigenicity of the 1918 Pandemic and Seasonal H1N1 Influenza A Viruses. Journal of Virology 8756-8766.

^{2.} Lee H. Tang J. Kong D. Loh T. Chiang D. Lam T. Koay E. (2013). Comparison of Mutation Patterns in Full-Genome A/H3N2 Influenza Sequences Obtained Directly from Clinical Samples and the Same Samples after a Single MDCK Passage. *PLoS One*.

Phylogenetic Comparison of Influenza A (H3N2) HA and NA Protein Sequences

- **33** analyzed H3N2 HA sequences were derived from MDCK isolates and **5** were derived from clinical specimen.
- 28 analyzed H3N2 NA sequences were derived from MDCK isolates and 5 were derived from clinical specimen.
- Phylogenetic trees for both HA and NA protein sequences were generated by Clustal V method using DNASTAR® Lasergene Megalign software.
- Amino acid changes shown are with respect to 2010 vaccine strain
 A/Perth/16/2009-like virus for HA sequences and A/Norway/1186/2011 for NA sequences

Summary of Influenza A (H3N2) Protein Homology When Compared to 2015-2016 Vaccine Strain

Segment	No.	2015-2016 Vaccine Strain	Protein Homology
H3N2 HA	38	A/Switzerland/9715293/2013	97.0-99.6%
H3N2 NA	33	A/Switzerland/9715293/2013	97.9-98.6%

Summary of Influenza A (H3N2) N-Linked Glycosylation Mutations

Loss or gain of N-linked glycosylation sites affect host innate immune system recognition and the ability to
induce adaptive immune response thus altering its viral antigenicity.¹ Predicted loss or gain of N-linked
glycosylation of protein sequences were calculated using CBS NetNGlyc 1.0 Server
http://www.cbs.dtu.dk/services/NetNGlyc/

	Mutation		
A/H3N2 Segment	ADD GLY	LOSS GLY	
НА	K160T	S47P N122D T128A N144S N158K N158H T160K T160I	
NA	S245N N329I S331R	S247T N329T	

Sun X. Jayaraman A. Mainprasad P. Raman R. Houser K. Pappas C. ... Tumpey T. (2013).
 N-Linked Glycosylation of the Hemagglutin Protein Influences Virulence and Antigenicity of the 1918 Pandemic and Seasonal H1N1 Influenza A Viruses. Journal of Virology 8756-8766.

Phylogenetic Comparison of Influenza B (Victoria and Yamagata) HA and NA Protein Sequences

- 50 analyzed Influenza B HA sequences were derived from MDCK isolates. 26 (52%) isolates belonged to the V1A clade of the Victoria Lineage and 24 (48%) belonged to the Y3 clade of the Yamagata lineage.
- 50 analyzed Influenza B NA sequences were derived from MDCK isolates. 26 isolates belonged to the V1A clade of the Victoria Lineage and 24 belonged to the Y3 clade of the Yamagata lineage.
- Phylogenetic trees for both HA and NA protein sequences were generated by Clustal V method using DNASTAR® Lasergene Megalign software.
- Amino acid changes shown (both HA and NA sequences) are with respect to previous vaccine strains B/Ohio/01/2005-like virus for B/Victoria specimens and B/Florida/04/2006-like virus for B/Yamagata specimens.

Summary of Influenza B Protein Homology When Compared to 2015-2016 Vaccine Strain

Segment	No. Isolates	2015-2016 Vaccine Strain	Protein Homology
B/Victoria HA	26	B/Brisbane/60/2008	99.1-99.5%
B/Victoria NA	26	B/Brisbane/60/2008	98.3-98.54%
B/Yamagata HA	24	B/Phuket/3073/2013	99.1-99.5%
B/Yamagata NA	24	B/Phuket/3073/2013	98.7-100.0%

Summary of Influenza B N-Linked Glycosylation Mutations

Loss or gain of N-linked glycosylation sites affect host innate immune system recognition and the ability to
induce adaptive immune response thus altering its viral antigenicity.¹ Predicted loss or gain of N-linked
glycosylation of protein sequences were calculated using CBS NetNGlyc 1.0 Server
http://www.cbs.dtu.dk/services/NetNGlyc/

	Mutation		
INF B Segment	ADD GLY LOSS GLY		
HA	A198T	N196D T198N	
NA	D463N		

Sun X. Jayaraman A. Mainprasad P. Raman R. Houser K. Pappas C. ... Tumpey T. (2013).

N-Linked Glycosylation of the Hemagglutin Protein Influences Virulence and Antigenicity of the 1918 Pandemic and Seasonal H1N1 Influenza A Viruses. Journal of Virology 8756-8766.

^{2.} Lee H. Tang J. Kong D. Loh T. Chiang D. Lam T. Koay E. (2013). Comparison of Mutation Patterns in Full-Genome A/H3N2 Influenza Sequences Obtained Directly from Clinical Samples and the Same Samples after a Single MDCK Passage. *PLoS One*.

Summary of Influenza A/pH1N1, A/H3N2 and Influenza B Hemagglutinin (HA) Genetic Groups

