

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type/ Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
Altimmune (UK) (Immune Targeting Systems Ltd)	FP-01.1 (Flunisyn)	Nucleoprotein (NP), Matrix protein (M1), RNA polymerase PB1, RNA polymerase PB2, HA head domain, conserved epitopes	Synthetic peptide-based	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None	Inactive, no longer in development	NCT01265914	Completed	Immune Targeting Systems Ltd	Industry	Phase 1	8/1/2010	3/1/2011	8/1/2011	49 Adults (18 to 55 years)	London, United Kingdom
Altimmune (UK) (Immune Targeting Systems Ltd)	FP-01.1 (Flunisyn)	Nucleoprotein (NP), Matrix protein (M1), RNA polymerase PB1, RNA polymerase PB2, HA head domain, conserved epitopes	Synthetic peptide-based	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None	Inactive, no longer in development	NCT02071329	Completed	Immune Targeting Systems Ltd	Industry	Phase 1	1/1/2014	12/1/2014	12/1/2014	111 Adults (18 to 45 years)	London, United Kingdom
Altimmune (UK) (Immune Targeting Systems Ltd)	FP-01.1 (Flunisyn)	Nucleoprotein (NP), Matrix protein (M1), RNA polymerase PB1, RNA polymerase PB2, HA head domain, conserved epitopes	Synthetic peptide-based	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Other: Unspecified	Inactive, no longer in development	NCT01677676	Completed	Immune Targeting Systems Ltd	Industry	Phase 1	1/1/2012	5/1/2012	9/1/2012	48 Adults (18 to 55 years)	Brisbane, Queensland, Australia
Altimmune (UK) (Immune Targeting Systems Ltd)	FP-01.1 (Flunisyn)	Nucleoprotein (NP), Matrix protein (M1), RNA polymerase PB1, RNA polymerase PB2, HA head domain, conserved epitopes	Synthetic peptide-based	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Other: Unspecified	Inactive, no longer in development	NCT01701752	Completed	Immune Targeting Systems Ltd	Industry	Phase 1	9/1/2012	4/1/2013	4/1/2013	120 Older Adults (65 to 74 years)	Unspecified
Altimmune (UK) (Immune Targeting Systems Ltd)	FP-01.1 (Flunisyn)	Nucleoprotein (NP), Matrix protein (M1), RNA polymerase PB1, RNA polymerase PB2, HA head domain, conserved epitopes	Synthetic peptide-based	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular		Inactive, no longer in development	See preclinical information		Immune Targeting Systems Ltd	Industry						
Altimmune (UK) (Immune Targeting Systems Ltd)	FP-01.1 (Flunisyn)	Nucleoprotein (NP), Matrix protein (M1), RNA polymerase PB1, RNA polymerase PB2, HA head domain, conserved epitopes	Synthetic peptide-based	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular		Inactive, no longer in development			Immune Targeting Systems Ltd	Industry						
BiondVax Pharmaceuticals Ltd (Israel)	Multimeric-001 (M-001)	HA head domain, conserved epitopes, Nucleoprotein (NP), Matrix protein (M1)	Peptide-based, Recombinant protein	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT03450915	Open, recruiting	BiondVax Pharmaceuticals Ltd.	Industry	Phase 3	8/1/2019	Estimated May 2020	Estimated December 2020	Estimated 9630 Adults and Older Adults (60 years and older)	Poland
BiondVax Pharmaceuticals Ltd (Israel)	Multimeric-001 (M-001)	HA head domain, conserved epitopes, Nucleoprotein (NP), Matrix protein (M1)	Peptide-based, Recombinant protein	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT03058692	Completed	National Institute of Allergy and Infectious Disease (NIAID)	Government	Phase 2	4/9/2018	1/14/2019	1/14/2019	120 Adults (18 to 49 years)	United States: Iowa, Ohio, Texas
BiondVax Pharmaceuticals Ltd (Israel)	Multimeric-001 (M-001)	HA head domain, conserved epitopes, Nucleoprotein (NP), Matrix protein (M1)	Peptide-based, Recombinant protein	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT02691130	Completed	(a) BiondVax Pharmaceuticals Ltd. (b) Seventh Framework Program	Industry, Government	Phase 2	11/1/2015	10/1/2016	1/1/2017	224 Adults (18 to 60 years)	Budapest, Hungary

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> 6/10/2014 Francis 2015 PMID: 24928790							[1] Francis 2015 (PMID: 24928790) [2] Gottlieb 2014 (PMID: 25172355) [3] <a href="https://clinicaltrials.gov/ct2/show/NCT01265914">https://clinicaltrials.gov/ct2/show/NCT01265914</a>
Results not yet reported									[1] Francis 2015 (PMID: 24928790) [2] Gottlieb 2014 (PMID: 25172355) [3] <a href="https://clinicaltrials.gov/ct2/show/NCT02071329">https://clinicaltrials.gov/ct2/show/NCT02071329</a>
Results not yet reported									[1] Francis 2015 (PMID: 24928790) [2] Gottlieb 2014 (PMID: 25172355) [3] <a href="https://clinicaltrials.gov/ct2/show/NCT01677676">https://clinicaltrials.gov/ct2/show/NCT01677676</a>
Results not yet reported									[1] Francis 2015 (PMID: 24928790) [2] Gottlieb 2014 (PMID: 25172355) [3] <a href="https://clinicaltrials.gov/ct2/show/NCT01701752">https://clinicaltrials.gov/ct2/show/NCT01701752</a>
				Unknown	Unknown				[1] Francis 2015 (PMID: 24928790) [2] Gottlieb 2014 (PMID: 25172355)
									[1] Francis 2015 (PMID: 24928790) [2] Gottlieb 2014 (PMID: 25172355)
Results not yet reported									[1] Rudolph 2014 (PMID: 21285533) [2] Gottlieb 2014 (PMID: 25172355) [3] Astmon 2012 (PMID: 22318394) [4] Astmon 2014 (PMID: 25173483) [5] Van Doorn 2017 (PMID: 28296763) [6] <a href="https://clinicaltrials.gov/ct2/show/NCT03450915">https://clinicaltrials.gov/ct2/show/NCT03450915</a>
Results not yet reported									[1] Rudolph 2014 (PMID: 21285533) [2] Gottlieb 2014 (PMID: 25172355) [3] Astmon 2012 (PMID: 22318394) [4] Astmon 2014 (PMID: 25173483) [5] Van Doorn 2017 (PMID: 28296763) [6] <a href="https://clinicaltrials.gov/ct2/show/NCT03058692">https://clinicaltrials.gov/ct2/show/NCT03058692</a>
Interim results reported	<a href="#">Sponsor press release</a> 07/20/17 <a href="http://www.biondavax.com/2017/07/biondavax-reports-positive-phase-2b-clinical-trial-results-for-its-universal-flu-vaccine/">http://www.biondavax.com/2017/07/biondavax-reports-positive-phase-2b-clinical-trial-results-for-its-universal-flu-vaccine/</a>								[1] Rudolph 2014 (PMID: 21285533) [2] Gottlieb 2014 (PMID: 25172355) [3] Astmon 2012 (PMID: 22318394) [4] Astmon 2014 (PMID: 25173483)

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Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type/ Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
BiondVax Pharmaceuticals Ltd (Israel)	Multimeric-001 (M-001)	HA head domain, conserved epitopes, Nucleoprotein (NP), Matrix protein (M1)	Peptide-based, Recombinant protein	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT02293317	Completed	BiondVax Pharmaceuticals Ltd.	Industry	Phase 2	11/1/2014	3/1/2015	6/1/2015	37 Adults and Older Adults (50 to 65 years)	Tel Aviv, Israel
BiondVax Pharmaceuticals Ltd (Israel)	Multimeric-001 (M-001)	HA head domain, conserved epitopes, Nucleoprotein (NP), Matrix protein (M1)	Peptide-based, Recombinant protein	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	Aluminum salts; Alum		NCT01419925	Completed	BiondVax Pharmaceuticals Ltd.	Industry	Phase 2	8/1/2011	1/1/2012	1/1/2012	120 Older Adults (65 years and older)	Jerusalem, Israel
BiondVax Pharmaceuticals Ltd (Israel)	Multimeric-001 (M-001)	HA head domain, conserved epitopes, Nucleoprotein (NP), Matrix protein (M1)	Peptide-based, Recombinant protein	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	Other: unspecified		NCT01146119	Completed	BiondVax Pharmaceuticals Ltd.	Industry	Phase 2	7/1/2010	5/1/2011	6/1/2011	200 Adults (18 to 49 years)	Israel: Jerusalem, Tel Aviv
BiondVax Pharmaceuticals Ltd (Israel)	Multimeric-001 (M-001)	HA head domain, conserved epitopes, Nucleoprotein (NP), Matrix protein (M1)	Peptide-based, Recombinant protein	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	Oil-in-water: Montanide ISA VG51		NCT01010737	Completed	BiondVax Pharmaceuticals Ltd.	Industry	Phase 1	9/1/2009	3/1/2010	3/1/2010	60 Adults and Older Adults (55 to 75 years)	Tel Aviv, Israel
BiondVax Pharmaceuticals Ltd (Israel)	Multimeric-001 (M-001)	HA head domain, conserved epitopes, Nucleoprotein (NP), Matrix protein (M1)	Peptide-based, Recombinant protein	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	Oil-in-water: Montanide ISA VG51		NCT00877448	Completed	BiondVax Pharmaceuticals Ltd.	Industry	Phase 1	6/1/2009	10/1/2009	11/1/2009	63 Adults (18 to 49 years)	Tel Aviv, Israel

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
									[5] Van Doorn 2017 (PMID: 28296763)
									[6] <a href="https://clinicaltrials.gov/ct2/show/NCT02691130">https://clinicaltrials.gov/ct2/show/NCT02691130</a>
Interim results reported	<a href="http://www.biondvax.com/clinical-trials/">Sponsor website. http://www.biondvax.com/clinical-trials/</a>								[1] Rudolph 2014 (PMID: 21285533)
									[2] Gottlieb 2014 (PMID: 25172355)
									[3] Astmon 2012 (PMID: 22318394)
									[4] Astmon 2014 (PMID: 25173483)
									[5] Van Doorn 2017 (PMID: 28296763)
									[6] <a href="https://clinicaltrials.gov/ct2/show/NCT02293317">https://clinicaltrials.gov/ct2/show/NCT02293317</a>
Interim results reported. Results reported in peer-reviewed journal	<a href="http://www.biondvax.com/clinical-trials/">Sponsor website. http://www.biondvax.com/clinical-trials/</a>	<a href="#">Peer-reviewed publication or journal. Astmon 2014 10/7/2014 PMID: 25173483</a>							[1] Rudolph 2014 (PMID: 21285533)
		<a href="#">Peer-reviewed publication or journal. Lowell 2017 2/1/2017 PMID: 28065476</a>							[2] Gottlieb 2014 (PMID: 25172355)
									[3] Astmon 2012 (PMID: 22318394)
									[4] Astmon 2014 (PMID: 25173483)
									[5] Van Doorn 2017 (PMID: 28296763)
									[6] Lowell 2017 (PMID: 28065476)
									[7] <a href="https://clinicaltrials.gov/ct2/show/NCT01419925">https://clinicaltrials.gov/ct2/show/NCT01419925</a>
Interim results reported	<a href="http://www.biondvax.com/clinical-trials/">Sponsor website. http://www.biondvax.com/clinical-trials/</a>								[1] Rudolph 2014 (PMID: 21285533)
									[2] Gottlieb 2014 (PMID: 25172355)
									[3] Astmon 2012 (PMID: 22318394)
									[4] Astmon 2014 (PMID: 25173483)
									[5] Van Doorn 2017 (PMID: 28296763)
									[6] <a href="https://clinicaltrials.gov/ct2/show/NCT01146118">https://clinicaltrials.gov/ct2/show/NCT01146118</a>
Interim results reported	<a href="http://www.biondvax.com/clinical-trials/">Sponsor website. http://www.biondvax.com/clinical-trials/</a>								[1] Rudolph 2014 (PMID: 21285533)
									[2] Gottlieb 2014 (PMID: 25172355)
									[3] Astmon 2012 (PMID: 22318394)
									[4] Astmon 2014 (PMID: 25173483)
									[5] Van Doorn 2017 (PMID: 28296763)
									[6] <a href="https://clinicaltrials.gov/ct2/show/NCT01010737">https://clinicaltrials.gov/ct2/show/NCT01010737</a>
Interim results reported. Results reported in both peer reviewed journal and registry	<a href="http://www.biondvax.com/clinical-trials/">Sponsor website. http://www.biondvax.com/clinical-trials/</a>	<a href="#">Peer-reviewed publication or journal. 2/9/2012 Astmon 2012 PMID: 22318394</a>	<a href="https://clinicaltrials.gov">https://clinicaltrials.gov</a> 3/27/2013 <a href="https://clinicaltrials.gov/ct2/show/results/NCT00877448">https://clinicaltrials.gov/ct2/show/results/NCT00877448</a>						[1] Rudolph 2014 (PMID: 21285533)
									[2] Gottlieb 2014 (PMID: 25172355)
									[3] Astmon 2012 (PMID: 22318394)
									[4] Astmon 2014 (PMID: 25173483)
									[5] Van Doorn 2017 (PMID: 28296763)

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<b>Recombinant Antigens/Proteins</b>																	
BiondVax Pharmaceuticals Ltd (Israel)	Multimeric-001 (M-001)	HA head domain, conserved epitopes, Nucleoprotein (NP), Matrix protein (M1)	Peptide-based, Recombinant protein	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		See preclinical information		BiondVax Pharmaceuticals Ltd.	Industry						
BiondVax Pharmaceuticals Ltd (Israel)	Multimeric-001 (M-001)	HA head domain, conserved epitopes, Nucleoprotein (NP), Matrix protein (M1)	Peptide-based, Recombinant protein	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular												
Bionor Pharma AS (Norway) Univ of Groningen (Netherlands)	Vacc-FLU	Nucleoprotein (NP), Membrane protein, IAV (M2)	Peptide-based	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Subcutaneous	ISA51		See preclinical information		Bionor Pharma	Industry						
Calder BioSciences (US) (previously Avilar Medical, LLC) Vanderbilt Univ (US)	DT-headless HA	Hemagglutinin (HA), conserved stalk domain	Other: headless HA	Strain-specific immunity	Unknown	Unknown		See preclinical information		Calder BioSciences (previously Avilar Medical, LLC) James Crowe, Vanderbilt University	Industry, Academic						
Dynavax (US)	N8295	Membrane protein ion channel ectodomain (MZe), Nucleoprotein (NP)	Fusion protein	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Unknown	Inactive, no longer in development	Unknown	Completed?	Dynavax	Industry	Phase 1	7/1/2010			54 Adults (18 to 40 years)	
Dynavax (US)	N8295: Preclinical	Membrane protein ion channel ectodomain (MZe), Nucleoprotein (NP)	Fusion protein	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Unknown	Inactive, no longer in development	See preclinical information		Dynavax	Industry						
Dynavax (US)	N8295	Membrane protein ion channel ectodomain (MZe), Nucleoprotein (NP)	Fusion protein	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Unknown											

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									<a href="https://clinicaltrials.gov/ct2/show/NCT00877448">[6] https://clinicaltrials.gov/ct2/show/NCT00877448</a>
				[1] Aged Mice: Proof of Concept [2] Mice: Model to examine human epitopes	<a href="#">[1-2] Astmon 2012</a>				<a href="#">[1] Rudolph 2014 (PMID: 21285533)</a> <a href="#">[2] Gottlieb 2014 (PMID: 25172355)</a> <a href="#">[3] Astmon 2012 (PMID: 22318394)</a> <a href="#">[4] Astmon 2014 (PMID: 25173483)</a> <a href="#">[5] Van Doorn 2017 (PMID: 28296763)</a>
									<a href="#">[1] Rudolph 2014 (PMID: 21285533)</a> <a href="#">[2] Gottlieb 2014 (PMID: 25172355)</a> <a href="#">[3] Astmon 2012 (PMID: 22318394)</a> <a href="#">[4] Astmon 2014 (PMID: 25173483)</a> <a href="#">[5] Van Doorn 2017 (PMID: 28296763)</a>
				[1] Mice: Determine the cellular and humoral immune responses; and to assess protective potential of induced immune response [2] Mice: Evaluate efficacy	[1] Herrera-Rodriguez 2018 [2] Bionor press release 2011				<a href="#">[1] Herrera-Rodriguez 2018 (PMID: 29223787)</a> <a href="#">[2] Bionor press release 2011</a>
				[1] Mice: evaluate efficacy [2] Ferrets and mice: test heterologous protection	[1] Grant proposal [2] Grant proposal				<a href="#">[1] Calder UV program description</a> <a href="#">[2] Grant Summary</a>
									<a href="#">[1] Zheng 2014 (PMID: 24178189)</a> <a href="#">[2] Press release 2011</a> <a href="#">[3] Press release Sep 2010</a> <a href="#">[4] Press release Apr 2010</a> <a href="#">[5] News Summary 2010</a>
				Unknown	Unknown				<a href="#">[1] Zheng 2014 (PMID: 24178189)</a> <a href="#">[2] Press release 2011</a> <a href="#">[3] Press release Sep 2010</a> <a href="#">[4] Press release Apr 2010</a> <a href="#">[5] News Summary 2010</a>
									<a href="#">[1] Zheng 2014 (PMID: 24178189)</a> <a href="#">[2] Press release 2011</a> <a href="#">[3] Press release Sep 2010</a> <a href="#">[4] Press release Apr 2010</a> <a href="#">[5] News Summary 2010</a>

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<b>Recombinant Antigens/Proteins</b>																	
Ghent Univ (Belgium) Sanofi (US)	CBC NA rNA antigens	Neuraminidase (NA): 3 recombinant NA proteins (NA5200, NA7900, NA9100)	Recombinant NA proteins	B cell response (e.g., neutralizing antibodies)	Intranasal	Sigma Adjuvant System (SAS), containing the immunostimulants monophosphoryl Lipid A and synthetic trehalose dicorymycolate		See preclinical information		Ghent University (Belgium) Sanofi	Academic, Industry						
Icahn School of Medicine at Mount Sinai (US)	mHA (IBV)	HA head domain, conserved epitopes, Hemagglutinin (HA), conserved stalk domain	Recombinant proteins: mosaic HA	Cross-reactive immune response	Intranasal	None		See preclinical information		Icahn School of Medicine at Mount Sinai	Academic						
Icahn School of Medicine at Mount Sinai (US)	Hyperglycosylated HA	Hemagglutinin (HA), conserved stalk domain	Other: Hyperglycosylation of globular head domain	Humoral response	Intramuscular	polyinosinic-polycyidylic acid (poly I:C)		See preclinical information		Icahn School of Medicine at Mount Sinai	Academic						
Icahn School of Medicine at Mount Sinai (US)	rNA proteins	Neuraminidase (NA)	Recombinant NA proteins	Mucosal immune response	Intramuscular, intranasal	polyinosinic-polycyidylic acid (poly I:C)		See preclinical information		Icahn School of Medicine at Mount Sinai	Academic						
Imutex Ltd (SEEK/HVVO) (UK)	FLU-v	Nucleoprotein (NP), Matrix protein (M1), Membrane protein, IAV (M2)	Synthetic peptide-based	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Other: Subcutaneous	Oil-in-water: Montanide ISA VG51		NCT02962908 2015-001932-38	Completed	(a) PepTcell Limited (b) Seventh Framework Program (c) University of Groningen (d) University Medical Center Groningen (e) Robert Koch Institute (f) Norwegian Institute of Public Health	Industry, Government, Academic, Academic, Government, Government	Phase 2	8/1/2016	7/18/2017	7/18/2017	170 Adults (18 to 60 years)	Zwolle, Netherlands
Imutex Ltd (SEEK/HVVO) (UK)	FLU-v	Nucleoprotein (NP), Matrix protein (M1), Membrane protein, IAV (M2)	Synthetic peptide-based	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Other: Subcutaneous	Oil-in-water: Montanide ISA VG51		NCT03180801 FLU-v-004 2016-002134-74 2015-25472	Completed	(a) PepTcell Limited (b) NIAID	Industry, Government	Phase 2	8/18/2016	3/31/2017	5/25/2017	153 Adults (18 to 55 years)	London, United Kingdom
Imutex Ltd (SEEK/HVVO) (UK)	FLU-v	Nucleoprotein (NP), Matrix protein (M1), Membrane protein, IAV (M2)	Synthetic peptide-based	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Other: Subcutaneous	Oil-in-water: Montanide ISA VG51		NCT01226758	Completed	PepTcell Limited	Industry	Phase 1	6/1/2010	12/1/2010	12/1/2010	32 Adults (18 to 45 years)	London, United Kingdom
Imutex Ltd (SEEK/HVVO) (UK)	FLU-v	Nucleoprotein (NP), Matrix protein (M1), Membrane protein, IAV (M2)	Synthetic peptide-based	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Other: Subcutaneous	Oil-in-water: Montanide ISA VG51		NCT01181336	Completed	PepTcell Limited	Industry	Phase 1	4/1/2010	7/1/2010	7/1/2010	48 Adults (18 to 40 years)	London, United Kingdom
Imutex Ltd (SEEK/HVVO) (UK)	FLU-v	Nucleoprotein (NP), Matrix protein (M1), Membrane protein, IAV (M2)	Synthetic peptide-based	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Other: Subcutaneous	Oil-in-water: Montanide ISA VG51		See preclinical information		PepTcell Limited	Industry						

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				[1] Mice: Evaluate protective capability [2] Mice: evaluate if vaccination could reduce viral lung load [3] Mice: examine if antibodies were the major mediators of protection induced by vaccination with CBC-NAs [4] Mice: evaluate scope of protection [5] Mice: evaluate if CBC designs could mediate NI against an HA variant	[1-5] Job 2018				[1] Job 2018 (PMID: 30510776)
				[1] Mice: Evaluate the pathogenicity of mosaic viruses, evaluate immunogenicity and efficacy; evaluate virus clearance in lungs by cross-reactive antibodies [2] Mice: Evaluate immunogenicity	[1] Sun 2019 [2] Broecker 2019				[1] Sun 2019 (PMID: 30944178) [2] Broecker 2019 (PMID: 31341648)
				[1] Mice: test reactivity to parental strain; assess binding to stalk without interference from head reactivity; evaluate immunogenicity	[1] Eggink 2014				[1] Eggink 2014 (PMID: 24155380)
				[1] Guinea pigs: evaluate immunogenicity; determine delivery method; determine breadth of immunity	[1] McMahon 2019				[1] McMahon 2019 (PMID: 31113896)
Interim results reported. Results reported in registry	<a href="#">Sponsor press release</a> 6/18/2018 <a href="https://www.innovaplc.com/media/portfolio-news/2018/2018-06-18">https://www.innovaplc.com/media/portfolio-news/2018/2018-06-18</a>		<a href="#">clinicaltrials.gov</a> 04/08/19 <a href="https://clinicaltrials.gov/ct2/show/results/NCT02962908">https://clinicaltrials.gov/ct2/show/results/NCT02962908</a>						[1] Van Doorn 2017 (PMID: 28376743) [2] Piquezuelos 2015 (PMID: 25994549) [3] Gottlieb 2014 (PMID: 25172355) [4] Piquezuelos 2012 (PMID: 22575166) [5] <a href="https://clinicaltrials.gov/ct2/show/NCT02962908">https://clinicaltrials.gov/ct2/show/NCT02962908</a>
Interim results reported. Results reported in registry	<a href="#">Sponsor press release</a> 03/26/18 <a href="https://www.innovaplc.com/media/portfolio-news/2018/2018-03-26">https://www.innovaplc.com/media/portfolio-news/2018/2018-03-26</a>		<a href="#">clinicaltrials.gov</a> 04/01/19 <a href="https://clinicaltrials.gov/ct2/show/results/NCT03180801">https://clinicaltrials.gov/ct2/show/results/NCT03180801</a>						[1] Van Doorn 2017 (PMID: 28376743) [2] Piquezuelos 2015 (PMID: 25994549) [3] Gottlieb 2014 (PMID: 25172355) [4] Piquezuelos 2012 (PMID: 22575166) [5] <a href="https://clinicaltrials.gov/ct2/show/NCT03180801">https://clinicaltrials.gov/ct2/show/NCT03180801</a>
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> 6/23/2012 Piquezuelos 2015 PMID: 25994549							[1] Van Doorn 2017 (PMID: 28376743) [2] Piquezuelos 2015 (PMID: 25994549) [3] Gottlieb 2014 (PMID: 25172355) [4] Piquezuelos 2012 (PMID: 22575166) [5] Piquezuelos 2015 (PMID: 26084515) [6] <a href="https://clinicaltrials.gov/ct2/show/NCT01226758">https://clinicaltrials.gov/ct2/show/NCT01226758</a>
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> 8/2015 Piquezuelos 2015 PMID: 26084515							[1] Van Doorn 2017 (PMID: 28376743) [2] Piquezuelos 2015 (PMID: 25994549) [3] Gottlieb 2014 (PMID: 25172355) [4] Piquezuelos 2012 (PMID: 22575166) [5] <a href="https://clinicaltrials.gov/ct2/show/NCT01181336">https://clinicaltrials.gov/ct2/show/NCT01181336</a>
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> 6/29/2012 Piquezuelos 2012 PMID: 22575166							[1] Van Doorn 2017 (PMID: 28376743) [2] Piquezuelos 2015 (PMID: 25994549) [3] Gottlieb 2014 (PMID: 25172355)
				[1] Mice: Evaluate Immunogenicity	[1] Stoloff 2007				[1] Van Doorn 2017 (PMID: 28376743) [2] Piquezuelos 2015 (PMID: 25994549) [3] Gottlieb 2014 (PMID: 25172355)



Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Imutex Ltd (SEEK/nVIVO) (UK)	FLU-v	Nucleoprotein (NP), Matrix protein (M1), Membrane protein, IAV (M2)	Synthetic peptide-based	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Other: Subcutaneous	Oil-in-water: Montanide ISA VG51				PepTcell Limited	Industry						
Janssen Pharmaceuticals (The Netherlands) Cruell Vaccine Institute and Scripps Research Institute	Mini-HA	Hemagglutinin (HA), conserved stalk domain	Other: Headless HA	Humoral response	Intramuscular	Aluminum salts: Alum		See preclinical information		Janssen Pharmaceuticals, Cruell Vaccine Institute and Scripps Research Institute	Industry, Industry, Academic						
Korea Univ College of Pharmacy (Korea)	nM2PR	Membrane protein ion channel ectodomain (M2e)	Peptide-based	Immunogen-specific response	Intraperitoneal	Freund's adjuvant		See preclinical information		Korea Univ College of Pharmacy	Academic						
NIAID Indian Institute of Science	H3- and H7-SI	Hemagglutinin (HA), conserved stalk domain	2 HA stem-immunogen (SI) vaccine	Non-neutralizing antibody response	Intramuscular; Intraperitoneal; Intranasally	Squalene-based oil-in-water adjuvants		See preclinical information		NIAID	Government						
National Yang-Ming Univ (Taiwan)	HA stem <sub>103</sub>	Hemagglutinin (HA), conserved stalk domain	Recombinant proteins	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Glycolipid C34		See preclinical information		National Yang-Ming University (Taiwan)	Academic						
Pasteur Institute of Iran	3M2e-HSP	Membrane protein ion channel ectodomain (M2e)	Chimer protein	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Subcutaneous	HSP70		See preclinical information		Pasteur Institute of IRAN	Government						
SutroVax, Inc. (US)	Xpress CF platform	Hemagglutinin (HA), conserved stalk domain	Xpress CF platform, Cell-free/novo	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Unknown	Unknown		See preclinical information		SutroVax, INC.	Industry						
Texas Tech Univ (US)	Efn-3xM2e-HA2+PA	Three tandem M2e repeats plus HA2	Recombinant antigens	T cell response (e.g., cytotoxic T-lymphocytes), Antigen-specific IgG response	Intranasal	Detoxified anthrax toxin system		See preclinical information		Texas Tech University	Academic						
Univ Autonoma del Estado de Morelos (Mexico)	q-DEC-205-M2e conjugate	Membrane protein ion channel ectodomain (M2e)	Peptide-based	T cell response (e.g., cytotoxic T-lymphocytes)	Subcutaneous	Polyinosinic-polycyidylic acid (poly I:C)		See preclinical information		Univ Autonoma del Estado de Morelos	Academic						
University of Hong Kong	HA-mini-stems	Hemagglutinin (HA), conserved stalk domain	Recombinant proteins with pre-fusion headless HA mini-stem	Humoral response	Intramuscular	Addavax (Invivogen)		See preclinical information		University of Hong Kong	Academic						
University of Oxford Blue Water Vaccines (UK)	OREO epitope	HA head domain, variable epitopes OREO epitope	Epitope-based	Cross-reactive immune response	Intramuscular	Alum: Alhydrogel		See preclinical information		University of Oxford Blue Water Vaccines	Academic, Industry						
University of Rochester Medical Center (US)	Chimeric vaccine construct (CH7/3)	H7 globular head, H3 stem	Recombinant proteins: Chimeric	T cell response (e.g., cytotoxic T-lymphocytes)	Subcutaneous	Aluminum salts: Alum		See preclinical information		University of Rochester	Academic						
VA Pharma LLC (Russia) Russian Federation Ministry of Health	Unifu	Membrane protein ion channel ectodomain (M2e), HA2 stalk epitopes	Recombinant proteins	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Other: Flagellin		NCT03789539	Open, not recruiting	VA Pharma Limited Liability Company	Industry	Phase 1	6/2/2018	12/2/2018	Estimated 12/31/18	54 Adults (18 to 60 years)	Saint-Petersburg, Russia

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
									<a href="#">[4] Piequozuelos 2012 (PMID: 22575166)</a>
									<a href="#">[5] Stoloff 2007 (PMID: 17688898)</a>
									<a href="#">[1] Van Doorn 2017 (PMID: 28376743)</a>
									<a href="#">[2] Piequozuelos 2015 (PMID: 25994549)</a>
									<a href="#">[3] Gottlieb 2014 (PMID: 25172355)</a>
									<a href="#">[4] Piequozuelos 2012 (PMID: 22575166)</a>
									<a href="#">[5] Stoloff 2007 (PMID: 17688898)</a>
				[1] Mice: evaluate breadth of protection [2] Mice: evaluate whether serum antibodies are responsible for in vivo protection [3] Cynomolgus monkeys: evaluate immunogenicity and protective efficacy [4] Mice: evaluate the impact of previous exposure to influenza on the induction of broadly influenza reactive antibodies by mini-HA antigen	[1] Impagliazzo 2015 [2] Impagliazzo 2015 [3] Impagliazzo 2015 [4] Van der Lubbe 2018				<a href="#">[1] Impagliazzo 2015 (PMID: 26303961)</a>
									<a href="#">[2] Van der Lubbe 2018 (PMID: 29977611)</a>
				[1] Mice: investigate production and specificity of the anti-m2e Abs in mice; evaluate efficacy	[1] Kim 2019				<a href="#">[1] Kim 2019 (PMID: 31501717)</a>
				[1] Mice: evaluate immunogenicity [2] Mice: evaluate efficacy [3] Mice: passive transfer experiment to determine whether protection was antibody-mediated [4] Ferrets: evaluate antibody response [5] Ferrets: evaluate efficacy [6] Mice: evaluate immunogenicity and efficacy [7] Mice: evaluate immunogenicity [8] Mice: evaluate efficacy	[1-5] Sutton 2017 [6] Mallajosyula 2015 [7] Mallajosyula 2014 [8] Bommakanti 2012				<a href="#">[1] Sutton 2017 (PMID: 29263880)</a>
									<a href="#">[2] Mallajosyula 2014 (PMID: 24927560)</a>
									<a href="#">[3] Mallajosyula 2015 (PMID: 26167164)</a>
									<a href="#">[4] Angelelli 2019 (PMID: 31213541)</a>
									<a href="#">[5] Bommakanti 2012 (PMID: 23015722)</a>
				[1] Mice: Evaluate immunogenicity	[1] Wang 2019				<a href="#">[1] Wang 2019 (PMID: 30388628)</a>
				[1] Mice: evaluate immunogenicity	[1] Farahmand 2018				<a href="#">[1] Farahmand 2018 (PMID: 30382564)</a>
				[1] Mice and ferrets: Determine immunogenicity and protective efficacy	[1] Grant summary				<a href="#">[1] Grant summary</a>
									<a href="#">[2] SutroVax Website</a>
				[1] Mice: Evaluate systemic antibody responses [2] Mice: Test if specific immunity against anthrax toxins was also developed	[1] Arevalo 2017				<a href="#">[1] Arevalo 2017 (PMID: 27775159)</a>
				[1] Mice: evaluate immunogenicity; evaluate role of effector CD4+ T cells on protection	[1] Padilla-Courate 2019				<a href="#">[1] Padilla-Courate 2019 (PMID: 30955979)</a>
				[1] Mice: evaluate use of group 1 HA min-stem to induce protection against both group 1 and group 2 viruses	[1] Valkenburg 2016				<a href="#">[1] Valkenburg 2016 (PMID: 26947245)</a>
				[1] Mice: evaluate immunogenicity	[1] Thompson 2018				<a href="#">[1] Thompson 2018 (PMID: 30242149)</a>
									<a href="#">[2] Recker 2007 (PMID: 17460037)</a>
									<a href="#">[3] Blue Water Vaccines website</a>
				[1] Mice: Evaluate immunogenicity	[1] DiPiazza 2019				<a href="#">[1] DiPiazza 2019 (PMID: 31399519)</a>
									<a href="#">[1] Taybalova 2015 (PMID: 26976546)</a>
									<a href="#">[2] Taybalova 2018 (PMID: 30138320)</a>
									<a href="#">[3] Stepanova 2015 (PMID: 25799221)</a>
									<a href="#">[4] Stepanova 2018 (PMID: 29631629)</a>

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
VA Pharma LLC (Russia) Russian Federation Ministry of Health	Unifu	Membrane protein ion channel ectodomain (M2e), HA2 stalk epitopes	Recombinant proteins	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Other: Flagellin		See preclinical information		VA Pharma Limited Liability Company	Industry						
VA Pharma LLC (Russia) Russian Federation Ministry of Health	Unifu	Membrane protein ion channel ectodomain (M2e), HA2 stalk epitopes	Recombinant proteins	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Other: Flagellin				VA Pharma Limited Liability Company	Industry						
Vaxinnate Corp (US)	TIV+VAX102 STF2.4M2e Vax102 is a recombinant influenza M2e-flagellin vaccine fused to the TLR5 agonist, flagellin	Membrane protein ion channel ectodomain (M2e)	Recombinant fusion protein	B cell response (e.g., neutralizing antibodies)	Intramuscular	Other: TLR5 agonist: Flagellin		NCT00921973	Completed	Vaxinnate Corporation	Industry	Phase 1	6/1/2009	7/1/2009	9/1/2009	80 Adults (18 to 49 years)	Lenexa, Kansas Nashville, Tennessee
Vaxinnate Corp (US)	TIV+VAX102 STF2.4M2e Vax102 is a recombinant influenza M2e-flagellin vaccine fused to the TLR5 agonist, flagellin	Membrane protein ion channel ectodomain (M2e)	Recombinant fusion protein	B cell response (e.g., neutralizing antibodies)	Intramuscular	Other: TLR5 agonist: Flagellin		NCT00921947	Completed	Vaxinnate Corporation	Industry	Phase 1	6/1/2009	7/1/2009	8/1/2009	60 Adults (18 to 49 years)	Salt Lake City, Utah
Vaxinnate Corp (US)	TIV+VAX102 STF2.4M2e Vax102 is a recombinant influenza M2e-flagellin vaccine fused to the TLR5 agonist, flagellin	Membrane protein ion channel ectodomain (M2e)	Recombinant fusion protein	B cell response (e.g., neutralizing antibodies)	Intramuscular	Other: TLR5 agonist: Flagellin		NCT00921206	Completed	Vaxinnate Corporation	Industry	Phase 1	6/1/2009	10/1/2009	12/1/2009	21 Adults (18 to 49 years)	Denver, Colorado
Vaxinnate Corp (US)	TIV+VAX102 STF2.4M2e Vax102 is a recombinant influenza M2e-flagellin vaccine fused to the TLR5 agonist, flagellin	Membrane protein ion channel ectodomain (M2e)	Recombinant fusion protein	B cell response (e.g., neutralizing antibodies)	Intramuscular	Other: TLR5 agonist: Flagellin		NCT00603811	Completed	Vaxinnate Corporation	Industry	Phase 1	9/1/2007	10/1/2008	10/1/2008	60 Adults (18 to 49 years)	Lenexa, Kansas Galveston, Texas
Vaxinnate Corp (US)	VAX102	Membrane protein ion channel ectodomain (M2e)	Recombinant fusion protein	B cell response (e.g., neutralizing antibodies)	Intramuscular	Other: TLR5 agonist: Flagellin		See preclinical information		Vaxinnate Corporation	Industry						
Vaxinnate Corp (US)	VAX102	Membrane protein ion channel ectodomain (M2e)	Recombinant fusion protein	B cell response (e.g., neutralizing antibodies)	Intramuscular	Other: TLR5 agonist: Flagellin				Vaxinnate Corporation	Industry						
<b>Reassortant/Recombinant Influenza Virus-Based Vaccines</b>																	
Codagenix, Inc. (US)	Codavax	Neuraminidase (NA), HA head domain, conserved epitopes	Live-attenuated influenza virus (e.g., single-replication viruses)	Humoral and cellular	Intranasal	None		NCT03926416 CODA01-001	Completed	Codagenix	Industry	Phase 1	2/21/2017	5/29/2018	9/14/2018	125 Adults (18 to 45 years)	Queensland, Australia

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Data and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
									[5] Stepanova 2018 (PMID: 29713522)
									[6] <a href="https://clinicaltrials.gov/ct2/show/NCT03789539">https://clinicaltrials.gov/ct2/show/NCT03789539</a>
				[1] Mice: Evaluate immunogenicity and protective properties of vaccine preparations [2] Mice: evaluate immunogenicity of enhanced vaccine candidate [3] Mice: evaluate immunogenicity and efficacy [4] Mice: compare immunogenicity and protective action of two recombinant proteins which feature different designs which target different antigens [5] Mice: compare the effect of different insertion points of the target antigens into flagellin on the structure, stability and immunogenicity of the recombinant proteins	[1] Tsybalova 2015 [2] Tsybalova 2018 [3] Stepanova 2015 [4] Stepanova 2018 [5] Stepanova 2018				[1] Tsybalova 2015 (PMID: 25976546) [2] Tsybalova 2018 (PMID: 30138320) [3] Stepanova 2015 (PMID: 25799221) [4] Stepanova 2018 (PMID: 29631629) [5] Stepanova 2018 (PMID: 29713522)
									[1] Tsybalova 2015 (PMID: 25976546) [2] Tsybalova 2018 (PMID: 30138320) [3] Stepanova 2015 (PMID: 25799221) [4] Stepanova 2018 (PMID: 29631629) [5] Stepanova 2018 (PMID: 29713522)
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> 12/28/2010 Talbot 2010 PMID: 21203437							[1] Turley 2011 (PMID: 21624416) [2] Talbot 2010 (PMID: 21203437) [3] <a href="https://clinicaltrials.gov/ct2/show/NCT00921973">https://clinicaltrials.gov/ct2/show/NCT00921973</a>
Results reported in registry			<a href="http://clinicaltrials.gov">clinicaltrials.gov</a> 08/22/11 <a href="https://clinicaltrials.gov/ct2/show/results/NCT00921947">https://clinicaltrials.gov/ct2/show/results/NCT00921947</a>						[1] Turley 2011 (PMID: 21624416) [2] Talbot 2010 (PMID: 21203437) [3] <a href="https://clinicaltrials.gov/ct2/show/NCT00921947">https://clinicaltrials.gov/ct2/show/NCT00921947</a>
Results not yet reported									[1] Turley 2011 (PMID: 21624416) [2] Talbot 2010 (PMID: 21203437) [3] <a href="https://clinicaltrials.gov/ct2/show/NCT00921206">https://clinicaltrials.gov/ct2/show/NCT00921206</a>
Results not yet reported									[1] Turley 2011 (PMID: 21624416) [2] Talbot 2010 (PMID: 21203437) [3] <a href="https://clinicaltrials.gov/ct2/show/NCT00603811">https://clinicaltrials.gov/ct2/show/NCT00603811</a>
				[1] Mice: Evaluate immunogenicity and protective efficacy	[1] Talbot 2010; Turley 2011				[1] Turley 2011 (PMID: 21624416) [2] Talbot 2010 (PMID: 21203437)
									[1] Turley 2011 (PMID: 21624416) [2] Talbot 2010 (PMID: 21203437)
Results not yet reported									[1] Yang 2013 (PMID: 23600603) [2] <a href="https://clinicaltrials.gov/ct2/show/NCT03926416">https://clinicaltrials.gov/ct2/show/NCT03926416</a>

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type/ Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Codagenix, Inc. (US)	CodaVax	Neuraminidase (NA), HA head domain, conserved epitopes	Live-attenuated influenza virus (e.g., single-replication viruses)	Humoral and cellular	Intranasal	None		See preclinical information		Codagenix	Industry						
Codagenix, Inc. (US)	CodaVax	Neuraminidase (NA), HA head domain, conserved epitopes	Live-attenuated influenza virus (e.g., single-replication viruses)	Humoral and cellular	Intranasal	None				Codagenix	Industry						
Flugen, Inc (US)	Redeeflu	Membrane protein, IAV (M2)	Live-attenuated influenza virus (e.g., single-replication viruses)	T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	None		2017-004971-30	Completed	FluGen Inc	Industry	Phase 2	2/27/2018	3/6/2019	3/6/2019	120 Adults (18 to 55 years)	Madison, Wisconsin
Flugen, Inc (US)	Redeeflu H3N2 M2SR coadministered with QIV boost	Membrane protein, IAV (M2)	Live-attenuated influenza virus (e.g., single-replication viruses)	T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal, Intramuscular	None		NCT03553940	Open, not recruiting	National Institute of Allergy and Infectious Disease (NIAID)	Government	Phase 1	8/15/2018	Estimated 10/31/20	Estimated 10/31/20	43 Adolescents and Children (9 to 17 years)	Saint Louis, Missouri
Flugen, Inc (US)	Redeeflu H3N2 M2SR coadministered with QIV boost	Membrane protein, IAV (M2)	Live-attenuated influenza virus (e.g., single-replication viruses)	T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal, Intramuscular	None		NCT02822105	Open, not recruiting	FluGen Inc	Industry	Phase 1	6/1/2016	Estimated 12/1/17	Estimated 6/1/18	96 Adults (18 to 49 years)	Lenexa, Kansas
Flugen, Inc (US)	Redeeflu	Membrane protein, IAV (M2)	Live-attenuated influenza virus (e.g., single-replication viruses)	T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	None		See preclinical information		FluGen Inc NIAID	Industry, Government						
Flugen, Inc (US)	Redeeflu	Membrane protein, IAV (M2)	Live-attenuated influenza virus (e.g., single-replication viruses)	T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	None				FluGen Inc NIAID	Industry, Government						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
				[1] Mice: Evaluate efficacy	[1] Yang 2013				[1] Yang 2013 (PMID: 23690603) [2] Mueller 2010 (PMID: 20543832)
									[1] Yang 2013 (PMID: 23690603) [2] Mueller 2010 (PMID: 20543832)
Interim results reported	[1] Sponsor Press Release 02/12/19 <a href="https://www.businesswire.com/news/home/20190212005173/en/FluGen%E2%80%99s-MCSR-Influenza-Vaccine-Succeeds-Phase-2">https://www.businesswire.com/news/home/20190212005173/en/FluGen%E2%80%99s-MCSR-Influenza-Vaccine-Succeeds-Phase-2</a>  [2] Press Release 02/12/19 <a href="https://www.fiercepharma.com/vaccines/flu-gen-plans-further-development-universal-flu-shot-after-phase-2-win">https://www.fiercepharma.com/vaccines/flu-gen-plans-further-development-universal-flu-shot-after-phase-2-win</a>								[1] Hatta 2018 (PMID: 30007825) [2] Hatta 2017 (PMID: 28668565) [3] Sarawar 2016 (PMID: 27595896) [4] Hatta 2011 (PMID: 21272601) [5] Watanabe 2009 (PMID: 19321619) [6] <a href="https://www.clinicaltrialsregister.eu/cti-search.html/2017-004971-30/BE">https://www.clinicaltrialsregister.eu/cti-search.html/2017-004971-30/BE</a>
Results not yet reported									[1] Hatta 2018 (PMID: 30007825) [2] Hatta 2017 (PMID: 28668565) [3] Sarawar 2016 (PMID: 27595896) [4] Hatta 2011 (PMID: 21272601) [5] Watanabe 2009 (PMID: 19321619) [6] <a href="https://clinicaltrials.gov/ct2/show/NCT03553940">https://clinicaltrials.gov/ct2/show/NCT03553940</a>
Results not yet reported									[1] Hatta 2018 (PMID: 30007825) [2] Hatta 2017 (PMID: 28668565) [3] Sarawar 2016 (PMID: 27595896) [4] Hatta 2011 (PMID: 21272601) [5] Watanabe 2009 (PMID: 19321619) [6] <a href="https://clinicaltrials.gov/ct2/show/NCT02822105">https://clinicaltrials.gov/ct2/show/NCT02822105</a>
				[1] Ferrets: Evaluate efficacy with preexisting immunity to influenza [2] Mice and Ferrets: Evaluate efficacy against H5N1 highly pathogenic avian influenza viruses [3] Mice: Evaluate immunogenicity [4] Mice: Assess attenuation and protective efficacy [5] Mice: Evaluate potential of MZKO influenza A virus as a live attenuated vaccine [6] Mice: Evaluate efficacy against heterologous influenza B virus challenge	[1] Hatta 2018 [2] Hata 2017 [3] Sarawar 2016 [4] Hatta 2011 [5] Watanabe 2009 [6] Moser 2019				[1] Hatta 2018 (PMID: 30007825) [2] Hatta 2017 (PMID: 28668565) [3] Sarawar 2016 (PMID: 27595896) [4] Hatta 2011 (PMID: 21272601) [5] Watanabe 2009 (PMID: 19321619) [6] Moser 2019 (PMID: 31280945)
									[1] Hatta 2018 (PMID: 30007825) [2] Hatta 2017 (PMID: 28668565) [3] Sarawar 2016 (PMID: 27595896) [4] Hatta 2011 (PMID: 21272601)

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Gamma Vaccines Pty Ltd (Australia)	GammaFlu	Other: Whole virus	Whole virion gamma-irradiated virus	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	Self-adjuvanted		See preclinical information		Gamma Vaccines	Industry						
Icahn School of Medicine at Mount Sinai (US)	cHA Ca2 M2 virus vaccine	Membrane protein, IAV (M2), Hemagglutinin (HA), conserved stalk domain, cHA	Inactivated influenza virus	Antibody specific response	Intramuscular	Squalene-based oil-in-water adjuvants		See preclinical information		Icahn School of Medicine at Mount Sinai	Academic						
Icahn School of Medicine at Mount Sinai (US) GSK (US)	Chimeric HA (cHA)-based vaccines: cH8/1N1 LAIV and cH5/1N1 IV with adjuvant Flu D-SUIV (GSK3816302A)	Hemagglutinin (HA), conserved stalk domain, HA head domain, conserved epitopes, Novel Antigen: cHA, Neuraminidase (NA)	Headless HA; Functional chimeric HA (cHA)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal, Intramuscular	Aluminum salts: ASO3A		NCT03300050	Open, not recruiting	PATH	Industry	Phase 1	10/10/2017	4/24/2018	Estimated 9/5/19	65 Adults (18 to 39 years)	Durham, North Carolina
Icahn School of Medicine at Mount Sinai (US) GSK (US)	Chimeric HA (cHA)-based vaccines: cH8/1N1 LAIV and cH5/1N1 IV with adjuvant Flu D-SUIV (GSK3816302A)	Hemagglutinin (HA), conserved stalk domain, HA head domain, conserved epitopes, Novel Antigen: cHA, Neuraminidase (NA)	Headless HA; Functional chimeric HA (cHA)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Aluminum salts: ASO3A	Inactive, no longer in development	NCT03275389 EUCTR2017-001584-20-BE	Open, not recruiting	GlaxoSmithKline	Industry	Phase 1	9/18/2017	5/7/2020	5/7/2020	471 Adults (18 to 39 years)	South Miami, Florida Wichita, Kansas Rochester, New York Austin, Texas Norfolk, Virginia Wilrijk, Belgium
Icahn School of Medicine at Mount Sinai (US) GSK (US)	Chimeric HA (cHA)-based vaccines: cH8/1N1 LAIV and cH5/1N1 IV with adjuvant Flu D-SUIV (GSK3816302A)	Hemagglutinin (HA), conserved stalk domain, HA head domain, conserved epitopes, Novel Antigen: cHA, Neuraminidase (NA)	Headless HA; Functional chimeric HA (cHA)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Aluminum salts: ASO3A		See preclinical information		GlaxoSmithKline Path	Industry, Industry						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
									<a href="#">[5] Watanabe 2009 (PMID: 19321619)</a>
									<a href="#">[6] Moser 2019 (PMID: 31280945)</a>
				[1] Mice: investigate mechanism behind cross-protection	[1] Fuyura 2010				<a href="#">[1] Fuyura 2010 (PMID: 20164241)</a>
				[1] Mice: evaluate vaccine efficacy	[1] Sun 2019				<a href="#">[1] Sun 2019 (PMID: 31560436)</a>
Results reported in peer-reviewed journal	<a href="#">Sponsor Press Release</a> 05/01/19 <a href="https://www.fiercebitech.com/biotech/ask-dumps-universal-flu-vaccine-after-interim-data-readout">https://www.fiercebitech.com/biotech/ask-dumps-universal-flu-vaccine-after-interim-data-readout</a>	<a href="#">Peer-reviewed journal</a> <a href="#">Bernstein 2019</a> 10/17/19 <a href="#">PMID: 31630990</a>							<a href="#">[1] Liu 2019 (PMID: 31105689)</a> <a href="#">[2] Nachbagauer 2017 (PMID: 29263881)</a> <a href="#">[3] Kramer 2019 (PMID: 30715353)</a> <a href="#">[4] Wang 2010 (PMID: 20956293)</a> <a href="#">[5] Ermier 2017 (PMID: 28356526)</a> <a href="#">[6] Nachbagauer 2018 (PMID: 30044403)</a> <a href="#">[7] Sunwoo 2018 (PMID: 30223475)</a> <a href="#">[8] Choi 2019 (PMID: 31032479)</a> <a href="#">[9] https://clinicaltrials.gov/ct2/show/NCT03300050</a> <a href="#">[10] Bernstein 2019 (PMID: 31630990)</a>
Interim results reported	<a href="#">Sponsor Press Release</a> 05/01/19 <a href="https://www.fiercebitech.com/biotech/ask-dumps-universal-flu-vaccine-after-interim-data-readout">https://www.fiercebitech.com/biotech/ask-dumps-universal-flu-vaccine-after-interim-data-readout</a>								<a href="#">[1] Liu 2019 (PMID: 31105689)</a> <a href="#">[2] Nachbagauer 2017 (PMID: 29263881)</a> <a href="#">[3] Kramer 2019 (PMID: 30715353)</a> <a href="#">[4] Wang 2010 (PMID: 20956293)</a> <a href="#">[5] Ermier 2017 (PMID: 28356526)</a> <a href="#">[6] Nachbagauer 2018 (PMID: 30044403)</a> <a href="#">[7] Sunwoo 2018 (PMID: 30223475)</a> <a href="#">[8] Choi 2019 (PMID: 31032479)</a> <a href="#">[9] https://clinicaltrials.gov/ct2/show/NCT03275389</a> <a href="#">[10] Development discontinued</a>
				[1] Mice: Evaluate ability of the conjugate to elicit the production of antibodies, evaluate efficacy, evaluate role of anti-LAH antibody in protection [2] Ferrets: Evaluate immunogenicity; compare one and two dose regimen, and examine breadth of protective immunity [3] Mice: Evaluate protective effect of sequential vaccination, evaluate efficacy [4] Ferrets: Evaluate efficacy in male ferrets to determine if ferret model recapitulates gender differences in immune responses observed in humans [5] Pigs: Assess immune responses and efficacy [6] Mice: explore the cHA strategy in mice by comparing use of two adjuvants [7] Mice: Evaluate immunogenicity and efficacy [8] Ferrets: Evaluate efficacy	[1] Wang 2010 [2] Nachbagauer 2017 [3] Ermier 2017 [4] Nachbagauer 2018 [5] Sunwoo 2018 [6] Choi 2019 [7] Nachbagauer 2016 [8] Nachbagauer 2016				<a href="#">[1] Liu 2019 (PMID: 31105689)</a> <a href="#">[2] Nachbagauer 2017 (PMID: 29263881)</a> <a href="#">[3] Kramer 2019 (PMID: 30715353)</a> <a href="#">[4] Wang 2010 (PMID: 20956293)</a> <a href="#">[5] Ermier 2017 (PMID: 28356526)</a> <a href="#">[6] Nachbagauer 2018 (PMID: 30044403)</a>



Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Icahn School of Medicine at Mount Sinai (US) GSK (US)	Chimeric HA (cHA)-based vaccines: cH8/1N1 LAIV and cH5/1N1 IV with adjuvant Flu D-SURV (GSK3816302A)	Hemagglutinin (HA), conserved stalk domain, HA head domain, conserved epitopes, Novel Antigen: cHA, Neuraminidase (NA)	Headless HA; Functional chimeric HA (cHA)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Aluminum salts: AS03A				GlaxoSmithKline	Industry						
Institute of Experimental Medicine (Russia)	cHA LAIV LAIV+NP	Nucleoprotein (NP), cHA	Recombinant proteins (e.g., chimeric HA, COBRA, mosaic HA), Live attenuated influenza virus (e.g., single-replication viruses)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	None		See preclinical information		Institute of Experimental Medicine (Russia) Icahn School of Medicine at Mount Sinai	Academic, Academic						
KJ Biosciences LLC (US)	Flu-ECP	HA head domain, conserved epitopes, HAZ	Inactivated influenza virus	Cross-reactive immune response	Intramuscular	None		See preclinical information		KJ Biosciences LLC Texas A&M University	Industry, Academic						
Univ of Georgia (US)	COBRA-based LAIV	HA head domain, variable epitopes, rHA	LAIV, Recombinant proteins: COBRA	Cellular immune response	Intramuscular	[1] IFA or alum hydroxide [2] AF03 squalene-in-water emulsion [3] Inject alum		See preclinical information		University of Georgia	Academic						
Univ of Oxford (UK)	S-FLU	HA head domain, conserved epitopes	Single replication (signal minus) virus	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	TS6		See preclinical information		University of Oxford	Academic						
Vacchera BioTech GmbH (Austria)	VTH201 UniFluVec	Nonstructural protein (NS1), Nuclear export protein (NEP)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	Unknown		See preclinical information		Vacchera Biotech	Industry						
Vivaldi Biosciences (US) Icahn School of Medicine at Mount Sinai (US) AVIR Green Hills Biotechnology AG (Austria)	deltaFLU LAIV deIN51-inactivated vaccine	Nonstructural protein (NS1)	Replication-deficient LAIV	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	Self-adjuvanted		NCT01369862 GHB-CS08	Completed	AVIR Green Hills Biotechnology AG Vivaldi Biosciences Icahn School of Medicine at Mount Sinai (US)	Industry, Industry, Academic	Phase 1	1/1/2011	8/1/2011	8/1/2011	80 Adults (18 to 60 years)	Vienna, Austria

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
									<a href="#">[7] Sunwoo 2018 (PMID: 30223475)</a> <a href="#">[8] Choi 2019 (PMID: 31032479)</a> <a href="#">[9] Development discontinued</a> <a href="#">[10] Nachbagauer 2016 (PMID: 29250436)</a> <a href="#">[11] Nachbagauer 2016 (PMID: 26719251)</a>
									<a href="#">[1] Liu 2019 (PMID: 31105686)</a> <a href="#">[2] Nachbagauer 2017 (PMID: 29263881)</a> <a href="#">[3] Krammer 2019 (PMID: 30715353)</a> <a href="#">[4] Wang 2010 (PMID: 20956283)</a> <a href="#">[5] Ermler 2017 (PMID: 28356526)</a> <a href="#">[6] Nachbagauer 2018 (PMID: 30044403)</a> <a href="#">[7] Sunwoo 2018 (PMID: 30223475)</a> <a href="#">[8] Choi 2019 (PMID: 31032479)</a> <a href="#">[9] Development discontinued</a> <a href="#">[10] Nachbagauer 2016 (PMID: 29250436)</a> <a href="#">[11] Nachbagauer 2016 (PMID: 26719251)</a>
				[1] Mice: evaluate efficacy [2] Ferrets: evaluate immunogenicity	[1] Isakova-Sivak 2018 [2] Korenkov 2018				<a href="#">[1] Isakova-Sivak 2018 (PMID: 29574336)</a> <a href="#">[2] Korenkov 2018 (PMID: 29520000)</a> <a href="#">[3] Korenkov 2018 (PMID: 29252117)</a>
				[1] Mice: evaluate the potential of low pH-treated antigens for increased cross-reactive immune response and cross protection	[1] Ni 2018				<a href="#">[1] KJ Biosciences Website</a> <a href="#">[2] Ni 2018 (PMID: 30140267)</a>
				[1] Mice: assess viral replication; evaluate efficacy [2] Ferrets: evaluate efficacy [3] Mice and ferrets: evaluate efficacy	[1] Sautto 2018 [2] Allen 2018 [3] Giles 2012				<a href="#">[1] Sautto 2018 (PMID: 31022693)</a> <a href="#">[2] Allen 2018 (PMID: 30265682)</a> <a href="#">[3] Giles 2012 (PMID: 22180399)</a>
				[1] Mice and ferrets: evaluate efficacy, and investigate cellular immune response [2] Ferrets and pigs: evaluate efficacy [3] Mice: Evaluate immunogenicity	[1] Baz 2015 [2] Holzer 2018 [3] Powell 2019				<a href="#">[1] Baz 2015 (PMID: 26488862)</a> <a href="#">[2] Holzer 2018 (PMID: 29703861)</a> <a href="#">[3] Powell 2019 (PMID: 30714836)</a>
				[1] Mice and ferrets: evaluate immunogenicity	[1] Vachera Website				<a href="#">[1] Vachera Website</a>
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> <a href="#">12/16/2013</a> <a href="#">Mosler 2013</a> <a href="#">PMID: 24183981</a>							<a href="#">[1] Morokutti 2014 (PMID: 24560674)</a> <a href="#">[2] Mosler 2013 (PMID: 24183981)</a> <a href="#">[3] Wachek 2010 (PMID: 20039806)</a> <a href="#">[4] <a href="https://clinicaltrials.gov/ct2/show/NCT01369862">https://clinicaltrials.gov/ct2/show/NCT01369862</a></a>

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Vivaldi Biosciences (US) Icahn School of Medicine at Mount Sinai (US) AVIR Green Hills Biotechnology AG (Austria)	deltaFLU LAIV deIN51-trivalent vaccine	Nonstructural protein (NS1)	Replication-deficient LAIV	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	Self-adjuvanted		NCT00724907 2006-001176-20	Completed	AVIR Green Hills Biotechnology AG Vivaldi Biosciences Icahn School of Medicine at Mount Sinai (US)	Industry, Industry, Academic	Phase 1	3/1/2007	7/1/2008	8/1/2008	48 Adults (18 to 50 years)	Vienna, Austria
Vivaldi Biosciences (US) Icahn School of Medicine at Mount Sinai (US) AVIR Green Hills Biotechnology AG (Austria)	deltaFLU LAIV deIN51-trivalent vaccine	Nonstructural protein (NS1)	Replication-deficient LAIV	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	Self-adjuvanted		See preclinical information		AVIR Green Hills Biotechnology AG Vivaldi Biosciences Icahn School of Medicine at Mount Sinai (US)	Industry, Industry, Academic						
Vivaldi Biosciences (US) Icahn School of Medicine at Mount Sinai (US) AVIR Green Hills Biotechnology AG (Austria)	deltaFLU LAIV deIN51-trivalent vaccine	Nonstructural protein (NS1)	Replication-deficient LAIV	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	Self-adjuvanted				AVIR Green Hills Biotechnology AG Vivaldi Biosciences Icahn School of Medicine at Mount Sinai (US)	Industry, Industry, Academic						
<b>Virus-Vectored Vaccines</b>																	
Altimmune, Inc (US)	NasoVAX	Other: targets respiratory tract without directly attacking the IFV	Other: Replication-deficient Ad5-vectored HA	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	None	Static (not progressing or terminated), looking for additional research collaborators	NCT03760549	Completed	Altimmune, Inc.	Industry	Phase 2	1/21/2019	2/18/2019	2/18/2019	8 Adults (18 to 49 years)	Rockville, Maryland
Altimmune, Inc (US)	NasoVAX	Other: targets respiratory tract without directly attacking the IFV	Other: Replication-deficient Ad5-vectored HA	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	None	Static (not progressing or terminated), looking for additional research collaborators	NCT03232567	Completed	Altimmune, Inc.	Industry	Phase 2	9/18/2017	3/7/2018	6/15/2018	60 Adults (18 to 49 years)	Rockville, Maryland
Altimmune, Inc (US)	NasoVAX	Other: targets respiratory tract without directly attacking the IFV	Other: Replication-deficient Ad5-vectored HA	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	None	Static (not progressing or terminated), looking for additional research collaborators	See preclinical information		Altimmune, Inc.	Industry						
Altimmune, Inc (US)	NasoVAX	Other: targets respiratory tract without directly attacking the IFV	Other: Replication-deficient Ad5-vectored HA	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	None	Static (not progressing or terminated), looking for additional research collaborators			Altimmune, Inc.	Industry						
China Center for Disease Control & Prevention	RVJ-4M2eNP	Nucleoprotein (NP), Matrix protein (M1), Membrane protein, IAV (M2), RNA polymerase PB1	Viral vector	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None		See preclinical information		China Center for Disease Control & Prevention	Government						
Eubetics (US)	Ad5-InflA-HA/M2e and Ad5-InflB-HA	HA head domain, conserved epitopes, Membrane protein, BBV (BM2), Membrane protein, IAV (M2), Nucleoprotein (NP)	Ad-vectored Vaccine	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal; Intradermal	None		See preclinical information		Eubetics Corporation	Industry						
Ewha Womans University (South Korea)	rAdB-NP	Nucleoprotein (NP)	rAd-vectored	T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	None		See preclinical information		Ewha Womans University (South Korea)	Academic						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal 2/22/14</a> <a href="#">Morokutti 2014</a> PMID: 24560674 <a href="#">Peer-reviewed publication or journal 2/01/10</a> <a href="#">Wachek 2010</a> PMID: 20038806							<a href="#">[1] Morokutti 2014 (PMID: 24560674)</a> <a href="#">[2] Mosler 2013 (PMID: 24183881)</a> <a href="#">[3] Wachek 2010 (PMID: 20038806)</a> <a href="#">[4] https://clinicaltrials.gov/ct2/show/NCT00724997</a>
				[1] Unknown: generate supporting information to advance these viruses into the clinic	[1] Grant Summary				<a href="#">[1] Morokutti 2014 (PMID: 24560674)</a> <a href="#">[2] Mosler 2013 (PMID: 24183881)</a> <a href="#">[3] Wachek 2010 (PMID: 20038806)</a> <a href="#">[4] Grant Summary</a>
									<a href="#">[1] Morokutti 2014 (PMID: 24560674)</a> <a href="#">[2] Mosler 2013 (PMID: 24183881)</a> <a href="#">[3] Wachek 2010 (PMID: 20038806)</a> <a href="#">[4] Grant Summary</a>
Results not yet reported									<a href="#">[1] Zhang 2011 (PMID: 21818346)</a> <a href="#">[2] https://clinicaltrials.gov/ct2/show/NCT03760549</a>
Results reported in registry			<a href="#">clinicaltrials.gov</a> <a href="#">04/11/19</a> <a href="#">https://clinicaltrials.gov/ct2/show/study/NCT03232567</a>						<a href="#">[1] Zhang 2011 (PMID: 21818346)</a> <a href="#">[2] https://clinicaltrials.gov/ct2/show/NCT03232567</a>
				[1] Mice: Evaluate protective efficacy and immunogenicity	[1] Zhang 2011				<a href="#">[1] Zhang 2011 (PMID: 21818346)</a> <a href="#">[1] Zhang 2011 (PMID: 21818346)</a>
									<a href="#">[1] Wang 2019 (PMID: 31240620)</a>
				[1] Mice: Characterize immunogenicity of fusion antigens expressed by the recombinant vaccinia viruses; evaluate protective efficacy	[1] Wang 2019				<a href="#">[1] Grant Summary</a> <a href="#">[2] Jones 2011 (PMID: 21821082)</a> <a href="#">[3] Gabibersch 2012 (PMID: 23041546)</a>
				[1] Mice and Ferrets: Evaluate efficacy against various influenza A and B viruses [2] Mice: determine immunologic effect of immunizations with increasing doses and determine efficacy [3] Ferrets: evaluate efficacy [4] Rhesus macaques: evaluate ability of vaccine to overcome pre-existing Ad5 immunity	[1] Grant summary [2,3] Jones 2011 [4] Gabibersch 2012				<a href="#">[1] Kim 2019 (PMID: 30639307)</a> <a href="#">[2] Lee 2019 (PMID: 30775351)</a>
				[1] Mice: evaluate immunization route; evaluate efficacy; identify epitope in the NP to determine specific T-cell responses [2] Mice: examine efficacy, antibody responses, CTL responses, and morbidity/mortality after challenge were measured	[1] Kim 2019 [2] Lee 2019				

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type/ Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Food and Drug Administration (US)	A/NP+M2-Ad	Nucleoprotein (NP), Membrane protein, IAV (M2)	rAd-vectored	T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	None		See preclinical information		FDA	Government						
Icahn School of Medicine at Mount Sinai (US)	ChA, NP and M1 delivered by ChAdOx1 and MVA viral-vectored vaccines	Nucleoprotein (NP), Matrix protein (M1)	Viral vector; chimeric HA	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None		See preclinical information		Icahn School of Medicine at Mount Sinai	Academic						
Jenner Institute/ Univ of Oxford (UK)	MVA-NP+M1 and ChAdOx1 NP+M1	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None		NCT01818362	Completed	University of Oxford	Academic	Phase 1	4/1/2013	11/1/2015	11/1/2015	72 Adults (18 to 50 years)	Oxford, Guildford and Southampton, United Kingdom
Jenner Institute/ Univ of Oxford (UK)	ChAdOx1 NP+M1	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None		NCT01623518	Completed	University of Oxford	Academic	Phase 1	6/1/2012	3/1/2013	3/1/2013	15 Adults (18 to 50 years)	Oxford, United Kingdom
Jenner Institute/ Univ of Oxford (UK)	ChAdOx1 NP+M1	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None		See preclinical information		University of Oxford	Academic						
Jenner Institute/ Univ of Oxford (UK)	ChAdOx1 NP+M1	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None				University of Oxford	Academic						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
				[1] Mice: Evaluate immunization route and efficacy [2] Mice: Evaluate immunogenicity [3] Mice: Evaluate efficacy and immunization route [4] Mice: Evaluate efficacy with diverse prior histories [5] Mice: evaluate impact of prior influenza infection on vaccine performance; examine effect of RSV-A2 and RV18 on performance	[1] Price 2010 [2] Soboleski 2011 [3] Price 2018 [4] Rowell 2018 [5] Rowell 2019				[1] Price 2010 (PMID: 20976273) [2] Soboleski 2011 (PMID: 21789196) [3] Price 2018 (PMID: 30037481) [4] Rowell 2018 (PMID: 29249542) [5] Rowell 2019 (PMID: 30986224)
				[1] Mice: Determine protective effect against 2 HA expressing viruses [2] Ferrets: evaluate efficacy and assess the impact of the cHA-NP+M1 bivalent viral vectors on inducing both cellular and humoral immunity	[1] Anunkumar 2019 [2] McMahon 2019				[1] Anunkumar 2019 (PMID: 31399277) [2] McMahon 2019 (PMID: 31497020)
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal 2/15/2018 Coughlan 2018 PMID: 29519670</a>							[1] Antrobus 2014 (PMID: 24374965) [2] Coughlan 2018 (PMID: 29519670) [3] Coughlan 2018 erratum (PMID: 29735416) [4] Lambe 2013 (PMID: 23485942) [5] Altenburg 2014 (PMID: 25036462) [6] Tully 2017 (PMID: 28724579) [7] <a href="https://clinicaltrials.gov/ct2/show/NCT01818362">https://clinicaltrials.gov/ct2/show/NCT01818362</a>
Results not yet reported									[1] Antrobus 2014 (PMID: 24374965) [2] Coughlan 2018 (PMID: 29519670) [3] Coughlan 2018 erratum (PMID: 29735416) [4] Lambe 2013 (PMID: 23485942) [5] Altenburg 2014 (PMID: 25036462) [6] Tully 2017 (PMID: 28724579) [7] <a href="https://clinicaltrials.gov/ct2/show/NCT01623518">https://clinicaltrials.gov/ct2/show/NCT01623518</a>
				[1] Mice: Evaluate efficacy [2] Mice: Evaluate immunogenicity [3] Ferrets: Evaluate efficacy	[1] Tully 2017 [2] Lambe 2013 [3] Altenburg 2014				[1] Antrobus 2014 (PMID: 24374965) [2] Coughlan 2018 (PMID: 29519670) [3] Coughlan 2018 erratum (PMID: 29735416) [4] Lambe 2013 (PMID: 23485942) [5] Altenburg 2014 (PMID: 25036462) [6] Tully 2017 (PMID: 28724579)
									[1] Antrobus 2014 (PMID: 24374965) [2] Coughlan 2018 (PMID: 29519670) [3] Coughlan 2018 erratum (PMID: 29735416) [4] Lambe 2013 (PMID: 23485942) [5] Altenburg 2014 (PMID: 25036462)

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Purdue Univ (US)	Multi-epitope Ad-based vaccine	Neuraminidase (NA), Nucleoprotein (NP), Hemagglutinin (HA)	Ad-vectored	Humoral and cell-mediated immune responses	Intramuscular	None		See preclinical information		Purdue University	Academic						
Univ of Hong Kong	WyethIL-15/56u	HA, NA, M1, M2, NP	Live multivalent-influenza vaccine	T cell response (e.g., cytotoxic T-lymphocytes)	Subcutaneous	IL-15		See preclinical information		University of Hong Kong	Academic						
Univ of Nebraska-Lincoln (US)	Consensus HA gene based Ad-vectored vaccine	HA head domain, conserved epitopes: H1, H2, H3, H5	Viral vector	Cross-protective immune response; potential for shared T and B cell responses	Intramuscular	None		See preclinical information		University of Nebraska-Lincoln	Academic						
Univ of Ottawa (Canada) National Institutes for Food and Drug Control (China)	rAd-SHA2	HA2 subunit	Viral vector	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	Targeting ligand/molecular adjuvant: CD40L		See preclinical information		University of Ottawa National Institutes for Food and Drug Control (China)	Industry, Government						
Vaccltech (UK)	VTP-100 MVA-NP+M1 (co-administered with QIV)	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT03300362	Withdrawn	(a) Vaccltech Ltd. (b) University of Oxford	Industry, Academic	Phase 2	10/13/2017	10/31/2018	10/31/2018	862 Older Adults (65 years and older)	United Kingdom: Bicester, Oxford, Pangbourne, Witney, Wokingham
Vaccltech (UK)	VTP-100 MVA-NP+M1	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT03277456	Completed	(a) Vaccltech Ltd. (b) University of Oxford	Industry, Academic	Phase 1	9/18/2017	11/2/2017	11/2/2017	6 Adults (18 to 50 years)	Oxford, United Kingdom

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
									<a href="#">[6] Tully 2017 (PMID: 28724578)</a>
				[1] Mice: Evaluate immunogenicity and protective efficacy [2] Ferrets: Evaluate immunogenicity and protective efficacy	[1] Hassan 2017 [2] Grant Summary				<a href="#">[1] Hassan 2017 (PMID: 29023601)</a> <a href="#">[2] Grant Summary</a>
				[1] Mice: Evaluate protective efficacy; evaluate immunogenicity; evaluate vaccine efficacy [2] Mice: Evaluate mechanisms of T cell protection and the universality of the vaccine	[1] Valkenburg 2014 [2] Valkenburg 2016				<a href="#">[1] Valkenburg 2014 (PMID: 24708798)</a> <a href="#">[2] Valkenburg 2018 (PMID: 29887326)</a>
				[1] Mice: Evaluate efficacy [2] Mice: Evaluate immunogenicity	[1] Lingel 2017 [2] Webby 2015				<a href="#">[1] Lingel 2017 (PMID: 29097783)</a> <a href="#">[2] Webby 2015 (PMID: 26469190)</a>
				[1] Mice: Evaluate the effects of using CD40L as an adjuvant and targeting molecule on the induction of HA2-specific immune response [2] Mice: Investigate the potential of vaccines to provide cross-protection against influenza viruses from different HA subtypes	[1,2] Fan 2014				<a href="#">[1] Fan 2015 (PMID: 25052763)</a>
Results not yet reported									<a href="#">[1] 2019 press release</a> <a href="#">[2] Folegatti 2019 (PMID: 30909516)</a> <a href="#">[3] De Vries 2018 (PMID: 29912453)</a> <a href="#">[4] Coughlan 2018 (PMID: 29519670)</a> <a href="#">[5] Altenburg 2018 (PMID: 29692427)</a> <a href="#">[6] Mullin 2016 (PMID: 26902548)</a> <a href="#">[7] Antrobus 2012 (PMID: 23831594)</a> <a href="#">[8] Mullarkey 2013 (PMID: 23589155)</a> <a href="#">[9] Powell 2013 (PMID: 23658773)</a> <a href="#">[10] Lillie 2012 (PMID: 22441650)</a> <a href="#">[11] Antrobus 2012 (PMID: 23118984)</a> <a href="#">[12] Berthoud 2011 (PMID: 21148512)</a> <a href="#">[13] <u>https://clinicaltrials.gov/ct2/show/NCT03300362</u></a>
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> <a href="#">3/22/2019</a> <a href="#">Folegatti 2019</a> <a href="#">PMID: 30909516</a>							<a href="#">[1] 2019 press release</a> <a href="#">[2] Folegatti 2019 (PMID: 30909516)</a> <a href="#">[3] De Vries 2018 (PMID: 29912453)</a> <a href="#">[4] Coughlan 2018 (PMID: 29519670)</a> <a href="#">[5] Altenburg 2018 (PMID: 29692427)</a> <a href="#">[6] Mullin 2016 (PMID: 26902548)</a> <a href="#">[7] Antrobus 2012 (PMID: 23831594)</a> <a href="#">[8] Mullarkey 2013 (PMID: 23589155)</a> <a href="#">[9] Powell 2013 (PMID: 23658773)</a> <a href="#">[10] Lillie 2012 (PMID: 22441650)</a>



Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Vaccitech (UK)	VTP-100 MVA-NP+M1 (co-administered with Viroflu6)	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT02014168	Withdrawn	University of Oxford	Academic	Phase 1	1/1/2014	4/1/2014	4/1/2014	3 Adults and Older Adults (18 years and older)	Oxford, United Kingdom
Vaccitech (UK)	VTP-100 MVA-NP+M1 (co-administered with TV)	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT01465035	Completed	University of Oxford	Academic	Phase 1	10/1/2011	11/1/2012	11/1/2012	24 Adults and Older Adults (50 years and older)	Oxford, United Kingdom
Vaccitech (UK)	VTP-100 MVA-NP+M1	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT00993063	Completed	(a) University of Oxford (b) Wellcome Trust	Academic; Other: specify (note developer if different from sponsor)	Phase 2	6/1/2009	3/1/2010	3/1/2010	27 Adults (18 to 45 years)	United Kingdom: Southampton, Oxford

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
									<a href="#">[1] Antrobus 2012 (PMID: 23118984)</a> <a href="#">[12] Berthoud 2011 (PMID: 21148512)</a> <a href="#">[13] https://clinicaltrials.gov/ct2/show/NCT03277456</a>
Results not yet reported									<a href="#">[1] 2019 press release</a> <a href="#">[2] Foleqaath 2019 (PMID: 30909516)</a> <a href="#">[3] De Vries 2018 (PMID: 29912453)</a> <a href="#">[4] Coughlan 2018 (PMID: 29518670)</a> <a href="#">[5] Altenburg 2018 (PMID: 29692427)</a> <a href="#">[6] Mullin 2016 (PMID: 26902548)</a> <a href="#">[7] Antrobus 2012 (PMID: 23831594)</a> <a href="#">[8] Mullarkey 2013 (PMID: 23589156)</a> <a href="#">[9] Powell 2013 (PMID: 23658773)</a> <a href="#">[10] Lillie 2012 (PMID: 22441650)</a> <a href="#">[11] Antrobus 2012 (PMID: 23118984)</a> <a href="#">[12] Berthoud 2011 (PMID: 21148512)</a> <a href="#">[13] https://clinicaltrials.gov/ct2/show/NCT02014168</a>
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> <a href="#">2/6/2013</a> <a href="#">Antrobus 2014</a> <a href="#">PMID: 23831594</a>							<a href="#">[1] 2019 press release</a> <a href="#">[2] Foleqaath 2019 (PMID: 30909516)</a> <a href="#">[3] De Vries 2018 (PMID: 29912453)</a> <a href="#">[4] Coughlan 2018 (PMID: 29518670)</a> <a href="#">[5] Altenburg 2018 (PMID: 29692427)</a> <a href="#">[6] Mullin 2016 (PMID: 26902548)</a> <a href="#">[7] Antrobus 2012 (PMID: 23831594)</a> <a href="#">[8] Mullarkey 2013 (PMID: 23589156)</a> <a href="#">[9] Powell 2013 (PMID: 23658773)</a> <a href="#">[10] Lillie 2012 (PMID: 22441650)</a> <a href="#">[11] Antrobus 2012 (PMID: 23118984)</a> <a href="#">[12] Berthoud 2011 (PMID: 21148512)</a> <a href="#">[13] https://clinicaltrials.gov/ct2/show/NCT01466036</a>
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> <a href="#">3/22/12</a> <a href="#">Lillie 2012</a> <a href="#">PMID: 22441650</a>							<a href="#">[1] 2019 press release</a> <a href="#">[2] Foleqaath 2019 (PMID: 30909516)</a> <a href="#">[3] De Vries 2018 (PMID: 29912453)</a>

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Vacotec (UK)	VTP-100 MVA-NP+M1	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT00942071	Completed	(a) University of Oxford (b) Wellcome Trust	Academic, Other: specify (note developer if different from sponsor)	Phase 1	8/1/2008	11/1/2012	11/1/2012	58 Adults and Older Adults (18 years and older)	Oxford, United Kingdom
Vacotec (UK)	VTP-100 MVA-NP+M1 (co-administered with QIV)	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT03880474	Open, not recruiting	(a) Vacotec Limited (b) Clinical Network Services (CNS) Pty Ltd	Industry, Industry	Phase 2	3/18/2019	Estimated 10/15/2019	Estimated 10/1/2021	6000 Adults and Older Adults (18 years and older)	Australia (New South Wales, Queensland, South Australia, Victoria)

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
									<a href="#">[4] Coughlan 2018 (PMID: 29519670)</a> <a href="#">[5] Altenburg 2018 (PMID: 29692427)</a> <a href="#">[6] Mullin 2016 (PMID: 26902548)</a> <a href="#">[7] Antrobus 2012 (PMID: 23831594)</a> <a href="#">[8] Mullarkey 2013 (PMID: 23588156)</a> <a href="#">[9] Powell 2013 (PMID: 23658773)</a> <a href="#">[10] Lillie 2012 (PMID: 22441650)</a> <a href="#">[11] Antrobus 2012 (PMID: 23118984)</a> <a href="#">[12] Berthoud 2011 (PMID: 21148512)</a> <a href="#">[13] <a href="https://clinicaltrials.gov/ct2/show/NCT00993083">https://clinicaltrials.gov/ct2/show/NCT00993083</a></a>
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> <a href="#">5/3/13</a> <a href="#">Powell 2013</a> <a href="#">PMID: 23658773</a>							<a href="#">[1] 2019 press release</a> <a href="#">[2] Colegaath 2019 (PMID: 30909516)</a> <a href="#">[3] De Vries 2018 (PMID: 29912453)</a> <a href="#">[4] Coughlan 2018 (PMID: 29519670)</a> <a href="#">[5] Altenburg 2018 (PMID: 29692427)</a> <a href="#">[6] Mullin 2016 (PMID: 26902548)</a> <a href="#">[7] Antrobus 2012 (PMID: 23831594)</a> <a href="#">[8] Mullarkey 2013 (PMID: 23588156)</a> <a href="#">[9] Powell 2013 (PMID: 23658773)</a> <a href="#">[10] Lillie 2012 (PMID: 22441650)</a> <a href="#">[11] Antrobus 2012 (PMID: 23118984)</a> <a href="#">[12] Berthoud 2011 (PMID: 21148512)</a> <a href="#">[13] <a href="https://clinicaltrials.gov/ct2/show/NCT00942071">https://clinicaltrials.gov/ct2/show/NCT00942071</a></a>
Results not yet reported		<a href="#">Peer-reviewed publication or journal</a> <a href="#">10/3/12</a> <a href="#">Antrobus 2012</a> <a href="#">PMID: 23118984</a>							<a href="#">[1] 2019 press release</a> <a href="#">[2] Colegaath 2019 (PMID: 30909516)</a> <a href="#">[3] De Vries 2018 (PMID: 29912453)</a> <a href="#">[4] Coughlan 2018 (PMID: 29519670)</a> <a href="#">[5] Altenburg 2018 (PMID: 29692427)</a> <a href="#">[6] Mullin 2016 (PMID: 26902548)</a> <a href="#">[7] Antrobus 2012 (PMID: 23831594)</a> <a href="#">[8] Mullarkey 2013 (PMID: 23588156)</a> <a href="#">[9] Powell 2013 (PMID: 23658773)</a>

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Vacotec (UK)	VTP-100 MVA-NP+M1	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT03883113	Open, not recruiting	Vacotec Limited	Industry	Phase 2	5/3/2019	Estimated 12/16/2019	Estimated 3/17/20	155 Adults (18 to 55 years)	Antwerp, Belgium
Vacotec (UK)	VTP-100 MVA-NP+M1	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None		See preclinical information		Vacotec Limited	Industry						
Vacotec (UK)	VTP-100 MVA-NP+M1	Nucleoprotein (NP), Matrix protein (M1)	Viral vector	T cell response (e.g., cytotoxic T-lymphocytes), B cell response (e.g., neutralizing antibodies)	Intramuscular	None				Vacotec Limited	Industry						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
									<a href="#">[10] Lillie 2012 (PMID: 22441650)</a> <a href="#">[11] Antrobus 2012 (PMID: 23118984)</a> <a href="#">[12] Berthoud 2011 (PMID: 21148512)</a> <a href="#">[13] <a href="https://clinicaltrials.gov/ct2/show/NCT03880474">https://clinicaltrials.gov/ct2/show/NCT03880474</a></a>
Results not yet reported									<a href="#">[1] 2019 press release</a> <a href="#">[2] Folegatti 2019 (PMID: 30909516)</a> <a href="#">[3] De Vries 2018 (PMID: 29912453)</a> <a href="#">[4] Coughlan 2018 (PMID: 29519670)</a> <a href="#">[5] Altenburg 2018 (PMID: 29692427)</a> <a href="#">[6] Mullin 2016 (PMID: 26902548)</a> <a href="#">[7] Antrobus 2012 (PMID: 23831594)</a> <a href="#">[8] Mullarkey 2013 (PMID: 23589155)</a> <a href="#">[9] Powell 2013 (PMID: 23558773)</a> <a href="#">[10] Lillie 2012 (PMID: 22441650)</a> <a href="#">[11] Antrobus 2012 (PMID: 23118984)</a> <a href="#">[12] Berthoud 2011 (PMID: 21148512)</a> <a href="#">[13] <a href="https://clinicaltrials.gov/ct2/show/NCT03883113">https://clinicaltrials.gov/ct2/show/NCT03883113</a></a>
				[1] Porses: Evaluate immunogenicity [2] Mice, chickens and pigs: evaluate use of candidate as adjuvant [3] Mice: evaluate effect of pre-existing immunity to MVA	<a href="#">[1] Brethnach 2006</a> <a href="#">[2] Mullarkey 2012</a> <a href="#">[3] Altenburg 2018</a>				<a href="#">[1] 2019 press release</a> <a href="#">[2] Folegatti 2019 (PMID: 30909516)</a> <a href="#">[3] De Vries 2018 (PMID: 29912453)</a> <a href="#">[4] Coughlan 2018 (PMID: 29519670)</a> <a href="#">[5] Altenburg 2018 (PMID: 29692427)</a> <a href="#">[6] Mullin 2016 (PMID: 26902548)</a> <a href="#">[7] Antrobus 2012 (PMID: 23831594)</a> <a href="#">[8] Mullarkey 2013 (PMID: 23589155)</a> <a href="#">[9] Powell 2013 (PMID: 23558773)</a> <a href="#">[10] Lillie 2012 (PMID: 22441650)</a> <a href="#">[11] Antrobus 2012 (PMID: 23118984)</a> <a href="#">[12] Berthoud 2011 (PMID: 21148512)</a> <a href="#">[13] Brethnach 2006 (PMID: 16194586)</a> <a href="#">[14] Fuksaitawong 2018</a>
									<a href="#">[1] 2019 press release</a>

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type/ Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
				(Neutralizing antibodies)													
Vaxart, Inc (US)	Oral Vaccine: VXA-A1.1	Hemagglutinin (HA)	Ad-based viral vector	B cell response (e.g., neutralizing antibodies)	Mucosal	dsRNA		NCT02918006	Completed	Vaxart Partner: BARDA	Industry	Phase 2, Challenge study	8/31/2016	5/19/2017	1/19/2018	179 Adults (18 to 49 years)	Costal Mesa, California
Vaxart, Inc (US)	Oral Vaccine: VXA-A1.1	Hemagglutinin (HA)	Ad-based viral vector	B cell response (e.g., neutralizing antibodies)	Mucosal	dsRNA		NCT03121339	Completed	Vaxart	Industry	Phase 1	3/31/2017	5/5/2017	4/3/2018	8 Adult Males (18 to 49 years)	Lexington, Kentucky
Vaxart, Inc (US)	Oral Vaccine: VXA-A1.1	Hemagglutinin (HA)	Ad-based viral vector	B cell response (e.g., neutralizing antibodies)	Mucosal	dsRNA		See preclinical information		Vaxart	Industry						
Vaxart, Inc (US)	Oral Vaccine: VXA-A1.1	Hemagglutinin (HA)	Ad-based viral vector	B cell response (e.g., neutralizing antibodies)	Mucosal	dsRNA				Vaxart	Industry						
Wistar Institute, Univ of Pennsylvania (US)	AdC68M2e(3)-NP vector E1-deleted adenovirus (Ad) vectors from chimpanzee serotypes expressing three M2e sequences fused to H1N1 NP	Membrane protein ion channel ectodomain (M2e), Nucleoprotein (NP)	Viral vector based on AdC vector	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None		See preclinical information		Wistar Institute	Industry						
<b>Virus-Like Particle (VLP)</b>																	
Beijing Institute of Microbiology and Epidemiology (China)	3M2e-NP-Hbc	Membrane protein ion channel ectodomain (M2e), Nucleoprotein (NP)	Recombinant subunit VLP (fusion)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intraperitoneal	Oil-in-water: SP01		See preclinical information		Beijing Institute of Microbiology and Epidemiology (China)	Academic						
Cytos Biotechnology AG (Switzerland)	Cytos M2 protein TLR7 vaccine	Membrane protein ion channel ectodomain (M2e)	Recombinant subunit VLP (fusion)	Innate immune response; M2e-specific response	Subcutaneous	None		See preclinical information		Cytos Biotechnology	Industry						
Georgia State Univ (US) Kyung Hee Univ (S Korea) Recently formed small biotech	M2e5x VLP	Novel/Enhanced Antigen: 4/M2e-HA (PR8 backbone)+LAI/IV	Recombinant subunit VLP (fusion)	Humoral and cellular	Intramuscular; Intranasal; Microneedle patch	5-m2e-VLPs		See preclinical information		Georgia State University ADVAC LLC	Academic, Industry						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
									<a href="#">[2] Folegatti 2019 (PMID: 30909516)</a> <a href="#">[3] De Vries 2018 (PMID: 29912453)</a> <a href="#">[4] Coughlan 2018 (PMID: 29519670)</a> <a href="#">[5] Altenburg 2018 (PMID: 29692427)</a> <a href="#">[6] Mullin 2016 (PMID: 26902548)</a> <a href="#">[7] Antrobus 2012 (PMID: 23831594)</a> <a href="#">[8] Mullarkey 2013 (PMID: 23589156)</a> <a href="#">[9] Powell 2013 (PMID: 23658773)</a> <a href="#">[10] Little 2012 (PMID: 22441650)</a> <a href="#">[11] Antrobus 2012 (PMID: 23118984)</a> <a href="#">[12] Berthoud 2011 (PMID: 21148512)</a> <a href="#">[13] Breathnach 2006 (PMID: 16194586)</a> <a href="#">[14] Fuksurtsova 2019</a>
Interim results reported	<a href="#">Poster</a> <a href="https://idsa.confex.com/idsa/2018/webprogram/Paper22242.html">https://idsa.confex.com/idsa/2018/webprogram/Paper22242.html</a>								<a href="#">[1] Scallan 2013 (PMID: 23155123)</a> <a href="#">[2] Scallan 2016 (PMID: 27071663)</a> <a href="https://clinicaltrials.gov/ct2/show/NCT02918006">[3] https://clinicaltrials.gov/ct2/show/NCT02918006</a>
Results not yet reported									<a href="#">[1] Scallan 2013 (PMID: 23155123)</a> <a href="#">[2] Scallan 2016 (PMID: 27071663)</a> <a href="https://clinicaltrials.gov/ct2/show/NCT03121338">[3] https://clinicaltrials.gov/ct2/show/NCT03121338</a>
				[1] Mice, Ferrets: Evaluate immunogenicity and delivery method [2] Evaluate efficacy of mono-, bi-, tri- and quadrivalent vaccine combinations	<a href="#">[1] Scallan 2013</a> <a href="#">[2] Scallan 2016</a>				<a href="#">[1] Scallan 2013 (PMID: 23155123)</a> <a href="#">[2] Scallan 2016 (PMID: 27071663)</a>
									<a href="#">[1] Scallan 2013 (PMID: 23155123)</a> <a href="#">[2] Scallan 2016 (PMID: 27071663)</a>
				[1] Mice: Evaluate antibody response; evaluate immunogenicity; elucidate the immune mechanisms that contribute to protection in vaccinated mice; evaluate the role of antibodies in adoptive transfer studies [2] Mice: test whether V1T or V1K hexon modifications perturb binding of neutralizing antibodies to native hexon; measure M2e-specific antibody responses; evaluate vaccine efficacy	<a href="#">[1] Zhou 2010</a> <a href="#">[2] Zhou 2013</a>				<a href="#">[1] Zhou 2010 (PMID: 20877342)</a> <a href="#">[2] Zhou 2013 (PMID: 23229092)</a> <a href="#">[3] Patent</a>
				[1] Mice: Evaluate immunogenicity and vaccine efficacy	<a href="#">[1] Gao 2013</a>				<a href="#">[1] Gao 2013 (PMID: 23416215)</a>
				[1] Mice: determine importance of the IgG2a/c isotype; determine importance of RNA for protective potency of M2e-AP205	<a href="#">[1] Schmitz 2012</a>				<a href="#">[1] Schmitz 2012 (PMID: 22531913)</a>
				[1] Mice: evaluate humoral and cellular immunogenicity and cross-protective efficacy [2] Mice: evaluate efficacy when immunizing	<a href="#">[1] Kim 2013</a> <a href="#">[2] Kim 2014</a> <a href="#">[3] Kim 2017</a>				<a href="#">[1] Kim 2013 (PMID: 23247101)</a>



Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Georgia State Univ (US)	HAM1 VLPs	H1, H8 and H13 from HA Group 1 H3, H4, and H10 from HA Group 2	Recombinant subunit VLP (fusion)	B cell response (e.g., neutralizing antibodies)	Intranasal	None		See preclinical information		Georgia State University	Academic						
Indiana Univ Bloomington (US)	NP-P22 VLP nanoparticle Vaccine	Hemagglutinin (HA), conserved stalk domain, Nucleoprotein (NP)	Recombinant subunit VLP (fusion)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	None		See preclinical information		Indiana University Bloomington	Academic						
iQur Ltd (UK)	Tandiflu VLP	HA head domain, conserved epitopes, Hemagglutinin (HA), conserved stalk domain	Chimeric VLP	Cellular immune response	Intraperitoneal	Addavax Invivogen	Active, currently in development	See preclinical information		iQur LTD	Industry						
Medicago, Inc (Canada)	Quadrivalent VLP (QVLP)	Hemagglutinin (HA)	HA-bearing quadrivalent virus-like particle (QVLP)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None	Active, currently in development	NCT03739112	Completed	Medicago	Industry	Phase 3	9/18/2018	5/17/2019	6/14/2019	12,793 Elderly Adults (65 years and older)	104 Locations: United States, Canada, Finland, Germany, Thailand
Medicago, Inc (Canada)	Quadrivalent VLP (QVLP)	Hemagglutinin (HA)	HA-bearing quadrivalent virus-like particle (QVLP)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None	Active, currently in development	NCT03301051	Completed	Medicago	Industry	Phase 3	8/21/2017	6/12/2018	6/12/2018	10,137 Adults (18 to 64 years)	74 Locations: United States, Canada, Finland, Germany, Philippines, Thailand, United Kingdom
Medicago, Inc (Canada)	Quadrivalent VLP (QVLP)	Hemagglutinin (HA)	HA-bearing quadrivalent virus-like particle (QVLP)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None	Active, currently in development	NCT03321968	Completed	Medicago	Industry	Phase 3	9/29/2017	12/1/2017	12/1/2017	1200 Adults (18 to 49 years)	10 Locations in Canada: Nova Scotia, Ontario and Quebec
Medicago, Inc (Canada)	Quadrivalent VLP (QVLP)	Hemagglutinin (HA)	HA-bearing quadrivalent virus-like particle (QVLP)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None	Active, currently in development	NCT02831751	Completed	Medicago	Industry	Phase 2	4/1/2016	7/1/2016	1/1/2017	1001 Elderly Adults (64 years and older)	15 Locations: United States and Canada
Medicago, Inc (Canada)	Quadrivalent VLP (QVLP)	Hemagglutinin (HA)	HA-bearing quadrivalent virus-like particle (QVLP)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None	Active, currently in development	NCT02768805	Completed	Medicago	Industry	Phase 2	3/2/2016	5/17/2016	11/26/2016	900 Adults (18 to 64 years)	9 Locations: United States and Canada

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
				<p>Mice: evaluate pre-existing immunity and then by subsequent following vaccination of these previously split vaccine-induced mice with the VLP vaccine</p> <p>[3] Mice: evaluate efficacy, compare efficacy of cross protection with wild-type and recombinant viruses</p> <p>[4] Mice: Evaluate immunogenicity; evaluate cross protective immune correlates</p> <p>[5] Mice: evaluate possible mechanisms of immune response; evaluate differences between VLP and proteins in stimulating innate immune response</p> <p>[6] Mice: determine whether Flg VLP exhibit adjuvant effects on eliciting Th1 type immune responses and improving efficacy</p> <p>[7] Mice: evaluate efficacy of delivery via microneedle patch</p> <p>[8] Mice: evaluate efficacy of supplementing LAIV with M2e5x VLP</p> <p>[9] Mice: evaluate vaccine efficacy induced by combinatorial VLPs</p>	<p>[4] Lee 2018</p> <p>[5] Kim 2018</p> <p>[6] Kim 2018</p> <p>[7] Kim 2019</p> <p>[8] Lee 2019</p> <p>[9] Kang 2019</p>				<p>[2] Kim 2014 (PMID: 25171841)</p> <p>[3] Kim 2017 (PMID: 29107058)</p> <p>[4] Lee 2018 (PMID: 29324805)</p> <p>[5] Kim 2018 (PMID: 30241300)</p> <p>[6] Kim 2018 (PMID: 30199754)</p> <p>[7] Kim 2019 (PMID: 31003421)</p> <p>[8] Lee 2019 (PMID: 30685658)</p> <p>[9] Kang 2019 (PMID: 31246961)</p> <p>[10] Grant Summary</p>
				<p>[1] Mice: Evaluate optimal immunogen designs and iterative antigen exposure</p> <p>[2] Mice: Evaluate enhanced immune protection against drifted viruses</p> <p>[3] Guinea pigs: Determine enhanced protection</p>	<p>[1] Luo 2018</p> <p>[2-3] Grant proposal</p>				<p>[1] Luo 2018 (PMID: 29545521)</p> <p>[2] Grant summary</p>
				<p>[1] Mice: Evaluate immunogenicity and efficacy</p>	<p>[1] Patterson 2013</p>				<p>[1] Patterson 2013 (PMID: 23540530)</p> <p>[2] Grant summary</p>
				<p>[1] Mice: Evaluate immunogenicity</p> <p>[2] Mice: Evaluate efficacy</p>	<p>[1] Kazaks 2017</p> <p>[2] Ramirez 2018</p>				<p>[1] Qur Ltd Website</p> <p>[2] FLUTCORE project site</p> <p>[3] Kazaks 2017 (PMID: 29126399)</p> <p>[4] Ramirez 2018 (PMID: 29306508)</p>
Results not yet reported									<p>[1] Won 2018 (PMID: 30448064)</p> <p>[2] Pillet 2019 (PMID: 31166967)</p> <p>[3] <a href="https://clinicaltrials.gov/ct2/show/NCT03739112">https://clinicaltrials.gov/ct2/show/NCT03739112</a></p>
Results not yet reported									<p>[1] Won 2018 (PMID: 30448064)</p> <p>[2] Pillet 2019 (PMID: 31166967)</p> <p>[3] <a href="https://clinicaltrials.gov/ct2/show/NCT03301061">https://clinicaltrials.gov/ct2/show/NCT03301061</a></p>
Results not yet reported									<p>[1] Won 2018 (PMID: 30448064)</p> <p>[2] Pillet 2019 (PMID: 31166967)</p> <p>[3] <a href="https://clinicaltrials.gov/ct2/show/NCT03321968">https://clinicaltrials.gov/ct2/show/NCT03321968</a></p>
Results not yet reported									<p>[1] Won 2018 (PMID: 30448064)</p> <p>[2] Pillet 2019 (PMID: 31166967)</p> <p>[3] <a href="https://clinicaltrials.gov/ct2/show/NCT02831751">https://clinicaltrials.gov/ct2/show/NCT02831751</a></p>
Results not yet reported									<p>[1] Won 2018 (PMID: 30448064)</p> <p>[2] Pillet 2019 (PMID: 31166967)</p> <p>[3] <a href="https://clinicaltrials.gov/ct2/show/NCT02768805">https://clinicaltrials.gov/ct2/show/NCT02768805</a></p>

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type/ Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Medicago, Inc (Canada)	Quadrivalent VLP (QVLP)	Hemagglutinin (HA)	HA-bearing quadrivalent virus-like particle (QVLP)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None	Active, currently in development	NCT02236052	Completed	Medicago	Industry	Phase 2	7/16/2014	6/17/2015	6/17/2015	450 Older Adults (50 years and older)	3 Locations: Quebec, Canada
Medicago, Inc (Canada)	Quadrivalent VLP (QVLP)	Hemagglutinin (HA)	HA-bearing quadrivalent virus-like particle (QVLP)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None	Active, currently in development	NCT0223816	Completed	Medicago	Industry	Phase 2	8/1/2014	Estimated 5/1/16	Estimated 5/1/16	300 Adults (18 to 49 years)	Florida, United States
Medicago, Inc (Canada)	Quadrivalent VLP (QVLP)	Hemagglutinin (HA)	HA-bearing quadrivalent virus-like particle (QVLP)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None	Active, currently in development	NCT01991587	Completed	Medicago	Industry	Phase 1	10/1/2013	8/1/2014	9/1/2014	120 Adults (18 to 49 years)	Florida, United States
Medicago, Inc (Canada)	Quadrivalent VLP (QVLP)	Hemagglutinin (HA)	HA-bearing quadrivalent virus-like particle (QVLP)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Aluminum salts: Alhydrogel	Active, currently in development	See preclinical information		Medicago	Industry						
Medicago, Inc (Canada)	Quadrivalent VLP (QVLP)	Hemagglutinin (HA)	HA-bearing quadrivalent virus-like particle (QVLP)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None	Active, currently in development			Medicago	Industry						
Medigen, Inc (Canada)	Medigen multivalent VLP vaccine	H1, H2, H3, H5, H7, H9, H10	Novel multivalent VLPs	Neutralizing immune response	Intramuscular	None		See preclinical information		Industry	Industry						
Merck Research Laboratories (US)	Merck M2 based vaccines	M2 extracellular domain	Recombinant subunit VLP (fusion)	M2 peptide-specific antibody response	Intrapentoneal	Aluminum salts: amorphous aluminum hydroxide sulfate		See preclinical information		Merck Research Laboratories	Industry						
National Tsing Hua Univ (Taiwan)	BAFF-VLPs	HA, NA, M1, M2	Recombinant subunit VLP (fusion)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Aluminum salts: Alum		See preclinical information		National Tsing Hua University (Taiwan)	Academic						
NIAID (US)	VLP cocktail	H1, H3, H5 and H7 HA's	Recombinant subunit VLP (fusion)	B cell response (e.g., neutralizing antibodies)	Intranasal	None		See preclinical information		NIAID	Government						
Sanofi Pasteur (US)	ACAM FLU-A	Membrane protein ion channel ectodomain (M2e)	Recombinant subunit VLP (fusion)	B cell response (e.g., neutralizing antibodies)	Intramuscular	Aluminum salts: Alhydrogel; Stimulon QS-21		NCT00819013	Completed	Sanofi	Industry	Phase 1	7/1/2007	1/1/2009	2/1/2009	87 Adults (18 to 40 years)	Miami, Florida Lenexa, Kansas Tacoma, Washington
Sanofi Pasteur (US)	ACAM FLU-A	Membrane protein ion channel ectodomain (M2e)	Recombinant subunit VLP (fusion)	B cell response (e.g., neutralizing antibodies)	Intramuscular	Aluminum salts: Alhydrogel; Stimulon QS-21		See preclinical information		Sanofi	Industry						
Sanofi Pasteur (US)	ACAM FLU-A	Membrane protein ion channel ectodomain (M2e)	Recombinant subunit VLP (fusion)	B cell response (e.g., neutralizing antibodies)	Intramuscular	Aluminum salts: Alhydrogel; Stimulon QS-21				Sanofi	Industry						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> 6/5/19 <a href="#">Pillet 2019</a> PMID: 31166987							<a href="#">[1] Won 2018 (PMID: 30448064)</a> <a href="#">[2] Pillet 2019 (PMID: 31166987)</a> <a href="#">[3] Pillet 2019 (PMID: 31166987)</a> <a href="#">[4] https://clinicaltrials.gov/ct2/show/NCT02236052</a>
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> 6/5/19 <a href="#">Pillet 2019</a> PMID: 31166987							<a href="#">[1] Won 2018 (PMID: 30448064)</a> <a href="#">[2] Pillet 2019 (PMID: 31166987)</a> <a href="#">[3] Pillet 2019 (PMID: 31166987)</a> <a href="#">[4] https://clinicaltrials.gov/ct2/show/NCT02233816</a>
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed journal</a> July 2016 <a href="#">Pillet 2016</a> PMID: 26987887							<a href="#">[1] Won 2018 (PMID: 30448064)</a> <a href="#">[2] Pillet 2019 (PMID: 31166987)</a> <a href="#">[3] Pillet 2016 (PMID: 26987887)</a> <a href="#">[4] https://clinicaltrials.gov/ct2/show/NCT01991587</a>
				[1] Mice: determine wheter H1 and H5-VLPs stimulated DCs in vivo	[1] Won 2018				<a href="#">[1] Won 2018 (PMID: 30448064)</a> <a href="#">[2] Pillet 2019 (PMID: 31166987)</a> <a href="#">[3] Pillet 2016 (PMID: 26987887)</a>
									<a href="#">[1] Won 2018 (PMID: 30448064)</a> <a href="#">[2] Pillet 2019 (PMID: 31166987)</a> <a href="#">[3] Pillet 2016 (PMID: 26987887)</a>
				[1] Ferrets: Evaluate the H1- and quadri-subtype H5/H7/H9/H10 VLPs containing H10 protein for safety and immunogenicity; and evaluate protective efficacy of mono- and quadri- subtype VLPs	[1] Pushko 2016				<a href="#">[1] Pushko 2016 (PMID: 27663671)</a> <a href="#">[2] Tret'yakova 2016 (PMID: 26529299)</a> <a href="#">[3] Grant Summary</a>
				[1] Mice and Rhesus Monkeys: Compare immunogenicity of M2 peptide conjugated to OMPC and M2 peptide expressed on the surface of HBVc antigen basedon dosatration responses	<a href="#">[1] Fu 2009</a>				<a href="#">[1] Fu 2009 (PMID: 19146898)</a>
				[1] Mice: evaluate immunogenicity with and without alum adjuvant; evaluate protective immunity	<a href="#">[1] Hong 2019</a>				<a href="#">[1] Hong 2019 (PMID: 30738080)</a>
				[1] Mice: Assess protection afforded by immunization	[1] Schwartzman 2015				<a href="#">[1] Schwartzman 2015 (PMID: 26199334)</a> <a href="#">[2] Schultz-Cherry 2015 (PMID: 26443464)</a> <a href="#">[3] NIAID Press release</a>
Results reported in both peer reviewed journal and registry		<a href="#">Peer-reviewed publication or journal</a> 3/18/2013 <a href="#">Ibanez 2013</a> PMID: 23527091	<a href="#">clinicaltrials.gov</a> 1/16/2012 <a href="#">https://clinicaltrials.gov/ct2/show/results/NCT00819013</a>						<a href="#">[1] De Filette 2008 (PMID: 18833315)</a> <a href="#">[2] Ibanez 2013 (PMID: 23527091)</a> <a href="#">[3] https://clinicaltrials.gov/ct2/show/NCT00819013</a>
				[1] Mice: Evaluate immunogenicity [2] Mice: Evaluate immunogenicity and protective efficacy	[1] Mice: De Filette 2008 [2] Ibanez 2013				<a href="#">[1] De Filette 2008 (PMID: 18833315)</a> <a href="#">[2] Ibanez 2013 (PMID: 23527091)</a> <a href="#">[1] De Filette 2008 (PMID: 18833315)</a>

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type/ Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Sanofi Pasteur (US) Univ of Georgia (US)	COBRA-VLP	HA head domain, conserved epitopes	COBRA based VLP	B cell response (e.g., neutralizing antibodies)	Intramuscular	Aluminum salts; Inject		See preclinical information		Sanofi Pasteur University of Georgia	Industry, Academic						
Technovax, Inc. (US)	Technovax's VLP vaccine	HA, NA, M1, NP, M2. Novel/enhanced antigen: remodeled HA	Recombinant subunit VLP (fusion)	Broadly neutralizing antibody response	Unknown	Unknown		See preclinical information		TechnoVax NIAID NIH	Industry, Government, Government						
Univ Putra Malaysia (Malaysia)	InvC-M2e3	Membrane protein ion channel ectodomain (M2e)	Recombinant subunit VLP (fusion)	M2e specific antibody response	Subcutaneous	None		See preclinical information		Univ Putra Malaysia (Malaysia)	Academic						
VA Pharma LLC (Russia) Russian Federation Ministry of Health	Plant-produced Flg-4M protein	Membrane protein ion channel ectodomain (M2e)	Recombinant subunit VLP (fusion)	B cell response (e.g., neutralizing antibodies)	Intranasal	Other: Flagellin		See preclinical information		VA Pharma Limited Liability Company	Industry						
<b>Nanoparticle-Based Vaccines</b>																	
Chinese Academy of Sciences (China)	3M2e-rHF	Membrane protein ion channel ectodomain (M2e)	Self-assembling nanoparticle	B cell response (e.g., neutralizing antibodies), humoral, cellular and mucosal immune responses	Intranasal	None		See preclinical information		Chinese Academy of Sciences (China)	Academic						
Georgia State Univ (US)	Multivalent layered nanocluster vaccine	Novel/Enhanced Antigen: 4xM2e+ conformation stabilized HA stem	Nanoparticles	B cell response (e.g., neutralizing antibodies)	Intramuscular, Dissolvable microneedle patch	None		See preclinical information		Georgia State University	Academic						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources	
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References	
										<a href="#">[2] Ibanez 2013 (PMID: 23527091)</a>
				[1] Mice and Ferrets: Evaluate protective efficacy and immunogenicity [2] Mice and Ferrets: Evaluate immunogenicities and efficacies of two strategies [3] Mice: evaluate protective efficacy of three different H5N1 COBRA vaccines: expand the breadth of antibody recognition; to stimulate the broadest breadth of HAI activity against each of the vaccines [4] Mice: determine protective efficacy and breadth of vaccine-elicited antibodies and efficacy of cocktail mixtures [5] Ferrets: evaluate efficacy with preexisting immune status, assess the enhancement of stem-specific antibody titers [6] Mice and Ferrets: Determine specific HAI antibody response; evaluate ability of elicited antibody response to block live virus infection [7] Mice: Evaluate immunogenicity with emulsion-based adjuvant [8] Chickens: evaluate immunogenicity [9] Mice: assess the ability of a set of H1 COBRA HA vaccines to elicit protective antibodies with HAI activity against both human and swine H1 influenza viruses	[1] Giles 2011 [2] Giles 2012 [3] Crevar 2015 [4] Carter 2016 [5] Carter 2017 [6] Wong 2017 [7] Allen 2017 [8] Ross 2019 [9] Skarlupka 2019				<a href="#">[1] Giles 2011 (PMID: 21320540)</a>  <a href="#">[2] Giles 2012 (PMID: 22190399)</a>  <a href="#">[3] Crevar 2015 (PMID: 25671661)</a>  <a href="#">[4] Carter 2016 (PMID: 26912624)</a>  <a href="#">[5] Carter 2017 (PMID: 28978709)</a>  <a href="#">[6] Wong 2017 (PMID: 28978710)</a>  <a href="#">[7] Allen 2017 (PMID: 28789850)</a>  <a href="#">[8] Bar-Peled 2019 (PMID: 31481254)</a>  <a href="#">[9] Ross 2019 (PMID: 30905528)</a>  <a href="#">[10] Skarlupka 2019 (PMID: 31448974)</a>	
				[1] Mice: Evaluate immunogenicity and efficacy of a single or combined candidate vaccine	[1] Grant summary				<a href="#">[1] Grant Summary 1</a>  <a href="#">[2] Grant Summary 2</a>  <a href="#">[3] Press Release 2013</a>  <a href="#">[4] TechnoVax Pipeline</a>	
				[1] Mice: Evaluate protective efficacy and immune responses induced without an adjuvant	[1] Ong 2019				<a href="#">[1] Ong 2019 (PMID: 31430985)</a>	
				[1] Mice: evaluate immunogenicity and efficacy of vaccine linked to flagellin in plants [2] Mice: evaluate strength of immune response and direction of response	[1] Mardanova 2015 [2] Mardanova 2016				<a href="#">[1] Mardanova 2015 (PMID: 26029390)</a>  <a href="#">[2] Mardanova 2016 (PMID: 26710263)</a>  <a href="#">[3] Mardanova 2018 (PMID: 29521217)</a>	
				[1] Mice: Evaluate immunogenicity and protective efficacy	[1] Qi 2018				<a href="#">[1] Qi 2018 (PMID: 29426819)</a>	
				[1] Mice: evaluate protective efficacy; investigate antibody-mediated effector mechanisms [2] Mice: explore protection mechanisms [3] Mice: Examine the potential of crosslinked protein nanoparticles to maintain immunogenicity after cold-chain-independent storage [4] Mice: evaluate immunogenicity	[1] Deng 2018 [2] Deng 2018 [3] Chang 2018 [4] Wang 2014				<a href="#">[1] Deng 2018 (PMID: 29367722)</a>  <a href="#">[2] Deng 2018 (PMID: 30065113)</a>  <a href="#">[3] Chang 2018 (PMID: 30365905)</a>  <a href="#">[4] Wang 2018 (PMID: 30394725)</a>	

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Immvention Therapeutx, Inc (US) Univ North Carolina- Chapel Hill (US)	Ace-DEX polymeric microparticle vaccine	Membrane protein ion channel ectodomain (M2e), HA	Nanoparticles	B cell response (e.g., neutralizing antibodies)	Intramuscular	cGAM0		See preclinical information		Immvention Therapeutx	Industry						
KJ Biosciences LLC (US)	Bfluagen	Membrane protein ion channel ectodomain (M2e), HA2	Dual-domain nanoparticle fusion protein	Cross-reactive immune response	Intramuscular	Oil-in-water; Squalene oil-in-water emulsion containing MPL and trehalose dicorynomycolate		See preclinical information		KJ Biosciences LLC	Industry						
Laval University (Canada)	PapMV and PapMV-sM2e	Membrane protein ion channel ectodomain (M2e), Nucleoprotein (NP)	Nanoparticles	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	PapMV-sM2e nanoparticles possess an adjuvant property		See preclinical information		Laval University (Canada)	Academic						
NIAID VRC (US)	Mosaic receptor-binding domain (RBD) nanoparticle	Other: Receptor-binding domain (RBD) of viral HA on a nanoparticle	Nanoparticles	B cell response (e.g., neutralizing antibodies)	Intramuscular	SAS adjuvant (Sigma)		See preclinical information		National Institute of Allergy and Infectious Disease (NIAID)	Government						
NIAID VRC (US)	H1ssF_3928	Hemagglutinin (HA), conserved stalk domain	Nanoparticles	B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT03814720	Open, recruiting	National Institute of Allergy and Infectious Disease (NIAID)	Government	Phase 1	4/1/2019	Estimated 6/30/21	Estimated 12/31/21	70 Adults and Older Adults (18 to 70 years)	Bethesda, Maryland
NIAID VRC (US)	H1ssF_3928	Hemagglutinin (HA), conserved stalk domain	Nanoparticles	B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT03186761	Open, not recruiting	National Institute of Allergy and Infectious Disease (NIAID)	Government	Phase 1	10/25/2017	9/3/2019	Estimated 9/27/20	50 participants Part 1: Adults at least 18 and born after 1959; Part 2: Adults 18-70 (not born in 1966-1969)	Bethesda, Maryland
NIAID VRC (US)	H1ssF_3928	Hemagglutinin (HA), conserved stalk domain	Nanoparticles	B cell response (e.g., neutralizing antibodies)	Intramuscular	None		See preclinical information		National Institute of Allergy and Infectious Disease (NIAID)	Government						
NIAID VRC (US)	H1ssF_3928	Hemagglutinin (HA), conserved stalk domain	Nanoparticles	B cell response (e.g., neutralizing antibodies)	Intramuscular	None				National Institute of Allergy and Infectious Disease (NIAID)	Government						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
									<a href="#">[5] Chang 2018 (PMID: 30292080)</a>
									<a href="#">[6] Grant Summary</a>
									<a href="#">[7] Wang 2014 (PMID: 23988715)</a>
				[1] Mice: Evaluate efficacy of adjuvant; evaluate immunogenicity and long-term vaccine efficacy [2] Mice: Identify optimal degradation rate of vaccine; evaluate the effect of controlled antigen or adjuvant delivery on immune activation kinetics; evaluate protective efficacy; and evaluate potential for cross-reactivity	[1] Junkins 2018 [2] Chen 2018				<a href="#">[1] Junkins 2018 (PMID: 29170142)</a>
									<a href="#">[2] Chen 2018 (PMID: 30261204)</a>
									<a href="#">[3] Grant Summary</a>
				[1] Mice: Evaluate immunogenicity and efficacy	[1] Ni 2017				<a href="#">[1] Ni 2017 (PMID: 28102171)</a>
									<a href="#">[2] Biofugaen UIV</a>
				[1] Mice: Evaluate immunogenicity and protective efficacy; evaluate adjuvant	[1] Bolduc 2018				<a href="#">[1] Bolduc 2018 (PMID: 30193813)</a>
				[1] Mice: Evaluate immunogenicity	[1] Kanekiyo 2019				<a href="#">[1] Kanekiyo 2019 (PMID: 30742090)</a>
Results not yet reported									<a href="#">[1] Kramer 2019 comment (PMID: 30742079)</a>
									<a href="#">[2] Corbett 2019 (PMID: 30808695)</a>
									<a href="#">[3] Yassine 2015 (PMID: 26301691)</a>
									<a href="#">[4] Kanekiyo 2013 (PMID: 23698367)</a>
									<a href="#">[5] NIH press release</a>
									<a href="https://clinicaltrials.gov/ct2/show/NCT03814720">[6] https://clinicaltrials.gov/ct2/show/NCT03814720</a>
Results not yet reported									<a href="#">[1] Kramer 2019 comment (PMID: 30742079)</a>
									<a href="#">[2] Corbett 2019 (PMID: 30808695)</a>
									<a href="#">[3] Yassine 2015 (PMID: 26301691)</a>
									<a href="#">[4] Kanekiyo 2013 (PMID: 23698367)</a>
									<a href="#">[5] NIH press release</a>
									<a href="https://clinicaltrials.gov/ct2/show/NCT03186781">[6] https://clinicaltrials.gov/ct2/show/NCT03186781</a>
				[1] Mice and Ferrets: Evaluate protective efficacy [2] Mice: Evaluate immunogenicity	[1] Yassine 2015 [2] Corbett 2019				<a href="#">[1] Kramer 2019 comment (PMID: 30742079)</a>
									<a href="#">[2] Corbett 2019 (PMID: 30808695)</a>
									<a href="#">[3] Yassine 2015 (PMID: 26301691)</a>
									<a href="#">[4] Kanekiyo 2013 (PMID: 23698367)</a>
									<a href="#">[5] NIH press release</a>
									<a href="#">[1] Kramer 2019 comment (PMID: 30742079)</a>
									<a href="#">[2] Corbett 2019 (PMID: 30808695)</a>
									<a href="#">[3] Yassine 2015 (PMID: 26301691)</a>



Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Novavax (US)	Nano-Flu	Hemagglutinin (HA), conserved stalk domain, HA head domain, conserved epitopes	Recombinant HA, INIV	B cell response (e.g., neutralizing antibodies)	Intramuscular	Other: Matrix-M		NCT03293498	Completed	Novavax	Industry	Phase 1	8/18/2017	3/14/2018	10/29/2018	330 Adults and Older Adults (50 years and older)	North Carolina, United States
Novavax (US)	Nano-Flu	Hemagglutinin (HA), conserved stalk domain, HA head domain, conserved epitopes	Recombinant HA, INIV	B cell response (e.g., neutralizing antibodies)	Intramuscular	Other: Matrix-M		NCT02078674	Completed	Novavax Department of Health and Human Services	Industry, Government	Phase 1	3/1/2014	7/1/2015	7/1/2015	610 Adults (18 to 64 years)	United States: California, Florida, Idaho, New York, South Carolina
Novavax (US)	Nano-Flu	Hemagglutinin (HA), conserved stalk domain, HA head domain, conserved epitopes	Recombinant HA, INIV	B cell response (e.g., neutralizing antibodies)	Intramuscular	Other: Matrix-M		See preclinical information		Novavax	Industry						
Novavax (US)	Nano-Flu	Hemagglutinin (HA), conserved stalk domain, HA head domain, conserved epitopes	Recombinant HA, INIV	B cell response (e.g., neutralizing antibodies)	Intramuscular	Other: Matrix-M				Novavax	Industry						
Osvax SAS (France)	OVX836	Nucleoprotein (NP)	Self-assembling nanoparticle	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular, Intranasal	None		NCT03594890	Completed	Osvax S.A.S	Industry	Phase 1	6/12/2018	7/7/2019	7/7/2019	72 Adults (18 to 49 years)	Antwerp, Belgium
Osvax SAS (France)	OVX836	Nucleoprotein (NP)	Self-assembling nanoparticle	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular, Intranasal	None		See preclinical information		Osvax S.A.S	Industry						
Osvax SAS (France)	OVX836	Nucleoprotein (NP)	Self-assembling nanoparticle	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular, Intranasal	None				Osvax S.A.S	Industry						
Texas Tech Univ (US)	AuNP-based vaccine AuNP-M2e+sCpG	Novel/Enhanced Antigen: M2 consensus peptides (human M2e, swine M2e, and avian M2e) NA conserved epitope	Nanoparticles	Humoral and cellular: M2e-specific response	Intranasal	CpG		See preclinical information		Texas Tech University	Academic						
TRIA Bioscience Corp (US)	Peptide containing highly conserved Helix A epitope	HA head domain, conserved epitopes	Nanoparticles	B cell response (e.g., neutralizing antibodies)	Intranasal	None		See preclinical information		TRIA Bioscience Corp	Industry						
Univ of Gothenburg (Sweden) Ghent Univ (Belgium)	HA-FFM2e:NPL	Membrane protein ion channel ectodomain (M2e)	Nanoparticles, Fusion protein	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	CTA1-DD		See preclinical information		Ghent University, University of Gothenburg	Academic, Academic						
Univ of Iowa Iowa State Univ (US)	IAV-nanovax	rHA (seasonal and novel immunogen based on equine recombinant HA3 (rHA3)), NP	Nanoparticles	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	CpG		See preclinical information		University of Iowa, Iowa State University	Academic, Academic						
Vanderbilt Univ (US)	pH-responsive NP vaccine	Nucleoprotein (NP)	Nanoparticles	Tissue-resident memory T cells response	Intranasal	CpG		See preclinical information		Vanderbilt Univ (US)	Academic						
Versatope Therapeutics, Inc (US)	Versatope M2e-rOMVs	M2e, Conserved domains from HA, NA and NP	Nanoparticles: Exosome-like particles (extracellular vesicles)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	CC rOMVs		See preclinical information		Versatope Therapeutics, Inc.	Industry						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources	
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References	
									<a href="#">[4] Kanehiko 2013 (PMID: 23696367)</a>	
									<a href="#">[5] NIH press release</a>	
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal, 6/14/2018, Shinde 2018, PMID: 29897849</a>							<a href="#">[1] Smith 2017 (PMID: 28844407)</a>	
									<a href="#">[2] Shinde 2018 (PMID: 29897849)</a>	
									<a href="#">[3] https://clinicaltrials.gov/ct2/show/NCT03293498</a>	
Results not yet reported									<a href="#">[1] Smith 2017 (PMID: 28844407)</a>	
									<a href="#">[2] Shinde 2018 (PMID: 29897849)</a>	
									<a href="#">[3] https://clinicaltrials.gov/ct2/show/NCT02078674</a>	
				[1] Ferrets: Evaluate immunogenicity	[1] Smith 2017				<a href="#">[1] Smith 2017 (PMID: 28844407)</a>	
									<a href="#">[2] Shinde 2018 (PMID: 29897849)</a>	
									<a href="#">[1] Smith 2017 (PMID: 28844407)</a>	
									<a href="#">[2] Shinde 2018 (PMID: 29897849)</a>	
Results not yet reported									<a href="#">[1] Del Campo 2019 (PMID: 30701093)</a>	
									<a href="#">[2] https://clinicaltrials.gov/ct2/show/NCT03594890</a>	
				[1] Mice: Evaluate immunogenicity	[1] Del Campo 2019				<a href="#">[1] Del Campo 2019 (PMID: 30701093)</a>	
									<a href="#">[1] Del Campo 2019 (PMID: 30701093)</a>	
				[1] Mice: Evaluate protective ability against influenza A; evaluate immunogenicity [2] Mice: evaluate the role of free M2e not immobilized on AuNPs towards induction of protective immunity; evaluate the longevity of vaccine-induced immunity [3] Mice: Evaluate broad protection of vaccine; characterize the mucosal immune response generated by the vaccine [4] Mice: Evaluate vaccine efficacy with age and determine if it might require re-administration during a lifetime	[1] Tao 2014 [2] Tao 2015 [3] Tao 2017 [4] Bimler 2019				<a href="#">[1] Tao 2014 (PMID: 23829488)</a>	
									<a href="#">[2] Tao 2015 (PMID: 25842219)</a>	
									<a href="#">[3] Tao 2017 (PMID: 28161578)</a>	
									<a href="#">[4] Bimler 2019 (PMID: 31507643)</a>	
									<a href="#">[4] Grant Summary</a>	
				[1] Mice: Evaluate immunogenicity and efficacy	[1] Zeigler 2019				<a href="#">[1] Zeigler 2019 (PMID: 31341647)</a>	
				[1] Mice: map the M2e T-cell recognition epitope and elucidate its possible mechanisms for protection [2] Mice: Evaluate whether combined HA-PM2e-NFL vaccine vector, hosting the CTA1-SM2e-DD and recombinant HA, stimulated enhanced protective immunity against influenza virus infections	[1] Eliasson 2018 [2] Bernasconi 2018				<a href="#">[1] Eliasson 2018 (PMID: 29295018)</a>	
									<a href="#">[2] Bernasconi 2018 (PMID: 30271406)</a>	
									<a href="#">[3] Eliasson 2011 (PMID: 21481325)</a>	
									<a href="#">[4] Eliasson 2008 (PMID: 18243429)</a>	
				[1] Mice: Evaluate immune capabilities [2] Mice: Evaluate protective efficacy [3] Unknown: Determine optimal vaccine formulation for immunity and protection	[1] Zacharias 2018 [2] Ross 2019 [3] Grant Summary				<a href="#">[1] Zacharias 2018 (PMID: 30233573)</a>	
									<a href="#">[2] Ross 2019 (PMID: 30663733)</a>	
									<a href="#">[3] Grant summary</a>	
				[1] Mice: evaluate immunogenicity and efficacy and demonstrate that clinically relevant antigen, NP, can be delivered via	[1] Knight 2019				<a href="#">[1] Knight 2019</a>	
				[1] Mice: Assess the role LPS play in eliciting humoral response against ROMV displayed proteins; evaluate safety of OMVs; assess	[1-2] Watkins 2017 [3] Rappazzo 2016				<a href="#">[1] Watkins 2017 (PMID: 28215994)</a>	

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
				T lymphocyte, HIV-specific immune response													
<b>DNA/RNA-Based Vaccines</b>																	
BioNTech (Germany) Pfizer (US)	sa-RNA	Hemagglutinin (HA)	mRNA (e.g., self-amplifying RNA)	B cell response (e.g., neutralizing antibodies)	Intramuscular	None		See preclinical information		BioNTech (Germany) Pfizer (US)	Industry, Industry						
Chinese Academy of Sciences	p-PPA-p3M2e and p-p3M2e	Membrane protein ion channel ectodomain (M2e)	DNA	B cell response (e.g., neutralizing antibodies)	Other: Electroporation	None		See preclinical information		Chinese Academy of Sciences	Academic						
CureVac (Germany)	mRNA vaccines based on RNAActive platform	HA, NP	Synthetic mRNA (e.g., self-amplifying RNA)	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	Self-adjuvanted		See preclinical information		CureVac (Germany)	Industry						
GeneOne Life Sciences, Inc. (S Korea) Inovio Pharmaceuticals (US)	VGX-3400	Membrane protein ion channel ectodomain (M2e), Nucleoprotein (NP)	DNA	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular; Other: Electroporation (EP)	None		NCT01184976	Completed	GeneOne Life Sciences, Inc. Inovio Pharmaceuticals	Industry	Phase 1	8/1/2010	4/1/2012	4/1/2012	30 Adults (18 to 39 years)	Seoul, South Korea
GeneOne Life Sciences, Inc. (S Korea) Inovio Pharmaceuticals (US)	VGX-3400	Membrane protein ion channel ectodomain (M2e), Nucleoprotein (NP)	DNA	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Other: Electroporation	None		NCT01142362	Completed	GeneOne Life Sciences, Inc. Inovio Pharmaceuticals	Industry	Phase 1	6/1/2010	11/1/2011	11/1/2011	32 Adults (18 to 50 years)	Overland Park, Kansas Rockville, Maryland
GeneOne Life Sciences, Inc. (S Korea) Inovio Pharmaceuticals (US)	VGX-3400: Preclinical	Membrane protein ion channel ectodomain (M2e), Nucleoprotein (NP)	DNA	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Other: Electroporation	None		See preclinical information		GeneOne Life Sciences, Inc. Inovio Pharmaceuticals	Industry						
GeneOne Life Sciences, Inc. (S Korea) Inovio Pharmaceuticals (US)	VGX-3400	Membrane protein ion channel ectodomain (M2e), Nucleoprotein (NP)	DNA	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Other: Electroporation	None				GeneOne Life Sciences, Inc. Inovio Pharmaceuticals	Industry						
Ghent Univ (Belgium)	mRNA encoding NP	Nucleoprotein (NP)	mRNA (e.g., self-amplifying RNA)	T cell response (e.g., cytotoxic T-lymphocytes)	Other: Intranasal	None		See preclinical information		Ghent University	Academic						
GSK (US)	SAM(HA) vaccine	Nucleoprotein (NP), Matrix protein (M1)	mRNA (e.g., self-amplifying RNA)	T cell response (e.g., cytotoxic T-lymphocytes)	Intramuscular	None; Oil-in-water cationic nanoemulsion (CNE)		See preclinical information		GSK (US)	Industry						
Inovio Pharmaceuticals (US)	INO-3401	HA, Neuraminidase (NA), Nucleoprotein (NP)	Synthetic DNA	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Other: Electroporation	None		NCT01403155 FLU-001	Completed	Inovio	Industry	Phase 1	5/1/2011	8/1/2012	8/1/2012	22 Adults (18 to 50 years)	Rockville, Maryland
Inovio Pharmaceuticals (US)	INO-3401	HA, Neuraminidase (NA), Nucleoprotein (NP)	Synthetic DNA	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Other: Electroporation	None		See preclinical information		Inovio	Industry						
Inovio Pharmaceuticals (US)	INO-3401	HA, Neuraminidase (NA), Nucleoprotein (NP)	Synthetic DNA	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Other: Electroporation	None				Inovio	Industry						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Data and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
				[2] Ferrets: Evaluate safety and efficacy [3] Mice: Evaluate efficacy and immunogenicity [4] Mice: evaluate whether rOMBs could be released in a controlled fashion; to determine whether controlled release of rOMVs could lead to immune protection; assess longevity of a single dose rOMV vs. traditional prime/boost regime					[2] Watkins 2017 (PMID: 2886291) [3] Rappazzo 2016 (PMID: 26827663) [4] Grant summary [5] Grant summary [6] Verstoppe Website
				[1] Mice: determine whether combining antigens in an RNA vaccine affects efficacy; compare sa-RNA vaccines to DNA vaccines and mRNA vaccines [2]	[1] Vogel 2018				[1] Vogel 2018 (PMID: 29275847) [2] Press release [3] Beissant 2019 (PMID: )
				[1] Mice: Evaluate immunogenicity and protective efficacy	[1] Yao 2019				[1] Yao 2019 (PMID: 3086750)
				[1] Mice: Evaluate immunogenicity of RNA active vaccine; evaluate whether a more conserved antigen could mediate protection against homologous and heterologous viral challenge [2] Pigs: investigate whether RNA active vaccines are immunogenic to animals with a weight more similar to humans [3] Mice: compare mRNA vaccines to licensed vaccines based on inactivated virus; evaluate immunogenicity; evaluate vaccine efficacy	[1,2] Kallen 2013 [3] Lutz 2017				[1] Kallen 2013 (PMID: 23921513) [2] Lutz 2017 (PMID: 29263884)
Results not yet reported									[1] GeneOne Life Science [2] https://clinicaltrials.gov/ct2/show/NCT01184976
Results not yet reported									[1] GeneOne Life Science [2] https://clinicaltrials.gov/ct2/show/NCT01142362
				Unknown	Unknown				[1] GeneOne Life Science
				[1] Mice: Evaluate whether intranasally delivered naked mRNA can elicit robust T-cell responses against NP of H3N2 strain compared to DNA vaccination	[1] Joe 2019				[1] Joe 2019 (PMID: 31345237)
				[1] Mice: Evaluate immunogenicity and efficacy [2] Mice and ferrets: evaluate protective immune responses after vaccination with novel HA-based vaccine; evaluate efficacy [3] Mice: Investigate possibility to enhance the immune response induced by a single immunization with SAM by increasing recruitment of APCs at the site of injection	[1] Maghni 2016 [2] Brazzoli 2016 [3] Manara 2019				[1] Maghni 2016 (PMID: 27525409) [2] Brazzoli 2016 (PMID: 26468547) [3] Marana 2019 (PMID: 31227353)
Results not yet reported									[1] Yan 2018 (PMID: 29100705) [2] Yan 2014 (PMID: 24631084) [3] https://clinicaltrials.gov/ct2/show/study/NCT01403155
				[1] Mice, guinea pigs, non human primates and ferrets: Evaluate efficacy	[1] Ferrets: Yan 2018				[1] Yan 2018 (PMID: 29100705) [2] Yan 2014 (PMID: 24631084) [1] Yan 2018 (PMID: 29100705) [2] Yan 2014 (PMID: 24631084)

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type; Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Inovio Pharmaceuticals (US) WinStar Institute, Univ of Pennsylvania (US)	pH3HA	Contemporary H3N2 antigens	Synthetic DNA	B cell response (e.g., neutralizing antibodies)	Other: Intramuscular electroporation (EP)	None		See preclinical information									
Moderna, Inc. (US)	mRNA-1440 (VAL-506440)	HA head domain, conserved epitopes, HA head domain, variable epitopes	mRNA (e.g., self-amplifying RNA)	B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT03076385 2015-003452-48	Completed	ModernaTX, Inc.	Industry	Phase 1	12/1/2015	10/1/2018	10/1/2018	201 Adults (18 to 64 years)	Results reported in peer-reviewed journal
Moderna, Inc. (US)	mRNA-1851 (VAL-339851)	HA head domain, conserved epitopes, HA head domain, variable epitopes	mRNA (e.g., self-amplifying RNA)	B cell response (e.g., neutralizing antibodies)	Intramuscular	None		NCT03345043	Open, not recruiting	ModernaTX, Inc.	Industry	Phase 1	5/25/2016	Estimated 12/1/19	Estimated 12/1/19	156 Adults (18 to 49 years)	Miami, Florida
Moderna, Inc. (US)	mRNA-1851	HA head domain, conserved epitopes, HA head domain, variable epitopes	mRNA (e.g., self-amplifying RNA)	B cell response (e.g., neutralizing antibodies)	Intramuscular	None		See preclinical information		ModernaTX, Inc.	Industry						
Moderna, Inc. (US)	mRNA-1440 mRNA-1851	HA head domain, conserved epitopes, HA head domain, variable epitopes	mRNA (e.g., self-amplifying RNA)	B cell response (e.g., neutralizing antibodies)	Intramuscular	None				ModernaTX, Inc.	Industry						
Profectus Biosciences (US) University of Washington	Novel DNA prime/subunit boost (LT-adjuvanted multi-antigen DNA vaccine)	Hemagglutinin (HA), conserved stalk domain, Membrane protein ion channel ectodomain (M2e), Nucleoprotein (NP), Matrix protein (M1)	DNA, Viral vector boost	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Gene gun (epidermal)	Other: LT adjuvant		See preclinical information		Profectus Biosciences (US) University of Washington	Industry, Academic						
Saint Louis Univ (US)	Multiple pan-DR- and HLA-A2 restricted, highly conserved influenza epitopes	Matrix protein (M1), Membrane protein, IAV (M2), Nucleoprotein (NP)	DNA	T cell response (e.g., cytotoxic T-lymphocytes)	Intranasal	None		See preclinical information		Saint Louis Univ (US)	Academic						
Statens Serum Institute (Denmark) UNIFLUSECURE consortium	Polyvalent DNA vaccine	Internally expressed matrix and nucleoprotein and externally expressed hemagglutinin and neuraminidase	DNA	B cell response (e.g., neutralizing antibodies), T cell response (e.g., cytotoxic T-lymphocytes)	Needle-free intradermal	None		See preclinical information		Statens Serum Institute UNIFLUSECURE consortium	Industry						
Univ of Oslo (Norway)	Combined HA vaccines (MHCII-targeted DNA vaccine)	H5, H6, H8, H9, H11, H13	DNA	B cell response (e.g., neutralizing antibodies)	Other: Electroporation	None		See preclinical information		University of Oslo	Academic						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References
				[1] Mice: Evaluate immunogenicity and protective efficacy	[1] Elliott 2018				[1] Elliott 2018 (PMID: 30062926)
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> 5/10/2019 Feldman 2019 PMID: 310798949							[1] Feldman 2019 (PMID: 31079849) [2] Bahi 2017 (PMID: 28457665) [3] Liang 2017 (PMID: 28958578) [4] Lindgren 2017 (PMID: 29181005) [5] <a href="https://clinicaltrials.gov/ct2/show/NCT03076385">https://clinicaltrials.gov/ct2/show/NCT03076385</a>
Results reported in peer-reviewed journal		<a href="#">Peer-reviewed publication or journal</a> 5/10/2019 Feldman 2019 PMID: 310798949							[1] Feldman 2019 (PMID: 31079849) [2] Bahi 2017 (PMID: 28457665) [3] Liang 2017 (PMID: 28958578) [4] Lindgren 2017 (PMID: 29181005) [5] <a href="https://clinicaltrials.gov/ct2/show/NCT03076385">https://clinicaltrials.gov/ct2/show/NCT03076385</a>
				[1] Mouse, Ferret, and Cynomolgus macaques: Evaluate immunogenicity [2] Cynomolgus macaques: Evaluate immunogenicity [3] Chinese rhesus macaques: Evaluate whether adjuvant could further enhance immune responses; evaluate immunogenicity with different delivery methods	[1] Bahi 2017 [2] Liang 2017 [3] Lindgren 2017				[1] Feldman 2019 (PMID: 31079849) [2] Bahi 2017 (PMID: 28457665) [3] Liang 2017 (PMID: 28958578) [4] Lindgren 2017 (PMID: 29181005)
									[1] Feldman 2019 (PMID: 31079849) [2] Bahi 2017 (PMID: 28457665) [3] Liang 2017 (PMID: 28958578) [4] Lindgren 2017 (PMID: 29181005)
				[1] Cynomolgus macaques: Investigate immunogenicity and protective efficacy	[1] Koday 2017				[1] Koday 2017 (PMID: 29267331) [2] Press release 2016 [3] Grant summary
				[1] Mice: Evaluate protective efficacy	[1] Eickhoff 2019				[1] Eickhoff 2019 (PMID: 31331771)
				[1] Pigs: Investigate immunogenicity of an optimized version of polyvalent DNA vaccine, characterized by a next-generation expression vector without antibiotic resistance markers [2] Rabbits: evaluate immunogenicity when preciously used genes were mixed and administered; evaluate potential of this polyvalent influenza DNA in an optimized setting with codon-optimized influenza DNA in an optimized setting with codon-optimized influenza genes inserted into next generation vectors and delivered with the needle-free jet-injector	[1] Borggren 2016 [2] Borggren 2015				[1] Borggren 2016 (PMID: 27211039) [2] Borggren 2015 (PMID: 25746201)
				[1] Mice: evaluate immunogenicity, assess potential enhancement of immune responses resulting from targeting of HA to MHCII molecules	[1] Anderson 2018				[1] Anderson 2018 (PMID: 29427414)

Vaccine Candidate								Clinical Trial Information									
Developer Name (Specify)	Candidate Name (Specify)	Target Antigen(s)	Vaccine Platform	Proposed Immune Mechanism of Action	Delivery Method	Adjuvant (Type, Specify name)	R&D Status	Registry ID number(s)	Trial Status	Sponsor Name	Sponsor Type	Phase	Study Start Date	Primary Completion Date	Study Completion Date	Sample Size, Enrollment and Age	Location
<b>Recombinant Antigens/Proteins</b>																	
Univ of Pennsylvania (US)	HA mRNA-LPNs	Hemagglutinin (HA), conserved stalk domain	mRNA (e.g., self-amplifying RNA)	Adaptive immune response; Stalk-specific response	Intramuscular, Intranasal	None		See preclinical information		University of Pennsylvania	Academic						
<b>Multiple Platforms</b>																	
Sanofi (US)	HA-ferritin nanoparticles	Divergent H1 sequences, COBRA HA antigens	Nanoparticles, Recombinant proteins (COBRA)	Strain-specific immunity	Intramuscular	SAS, AF03		See preclinical information		Sanofi Pasteur	Industry						
Shanghai Public Health Clinical Center and Institutes of Biomedical Sciences	Three vaccines expressing immunogen sequences PAPB1M1 and PB2NPM2	Highly conserved internal viral epitopes	DNA; adenovirus based; TTV vaccinia-based	T cell response (e.g. cytotoxic T-lymphocytes)	Intranasal, Intramuscular	None		See preclinical information		Shanghai Public Health Clinical Center and Institutes of Biomedical Sciences	Academic						

Results Reporting Information				Preclinical Studies		Key Partners		Other	Data Sources	
Results Reporting Status	Interim Publication Type, Date, and Link	Full Publication Type, Date, and Link	Clinical Trials Registry, Publication Date and Link	Study Intent	Publication Link	Name(s)	Contact Information	Notes	References	
				[1] Mice: Evaluate immunogenicity, determine if HA stalk-specific antibodies could be elicited with a different influenza HA immunogen; evaluate protective efficacy; evaluate protective immune response against antigenically distant subtypes [2] Rabbits and Ferrets: evaluate potency of nucleoside-modified mRNA-LNP influenza virus vaccine [3] Mice: Evaluate efficacy of nasal delivery; evaluate the potential of AAV9.F16 as a vaccine for pandemic influenza [4] Ferrets: evaluate efficacy of intranasal delivery [5] Rhesus macaques: assess feasibility of translating this intranasal delivery strategy into primates	[1-2] Pardi 2018 [3-5] Limberis 2015				<a href="#">[1] Pardi 2018 (PMID: 30135514)</a>  <a href="#">[2] Limberis 2013 (PMID: 23720583)</a>	
				[1] Mice and Ferrets: Evaluate immunogenicity; Evaluate the HAI cross-reactivity elicited by combinations of select HA-Nps; evaluate efficacy	<a href="#">[1] Danicarrere 2018</a>				<a href="#">[1] Danicarrere 2018 (PMID: 30185504)</a>	
				[1] Mice: Evaluate immunogenicity, efficacy and combined administration of vaccines	[1] Xie 2019				<a href="#">[1] Xie 2019 (PMID: 31379782)</a>	