

Summary of Studies Supporting USDA Product Licensure

Establishment Name	Boehringer Ingelheim Animal Health USA Inc.
USDA Vet Biologics Establishment Number	124
Product Code	4845.21
True Name	Encephalomyelitis-Rhinopneumonitis-Influenza Vaccine, Eastern & Western, Killed Virus, Tetanus Toxoid
Tradename(s) / Distributor or Subsidiary (if different from manufacturer)	Vetera 5xp - Boehringer Ingelheim (Canada) Ltd. Vetera 5xp - No distributor specified
Date of Compilation Summary	February 06, 2019

Disclaimer: Do not use the following studies to compare one product to another. Slight differences in study design and execution can render the comparisons meaningless.

Study Type	Efficacy
Study Type	
Pertaining to	Clostridium tetanus
Study Purpose	Demonstration of efficacy against Clostridium tetanus
Product Administration	One dose, administered intramuscularly
Study Animals	10 guinea pigs (10 vaccinates)
Challenge Description	Not applicable
Interval observed after	Not applicable
challenge	
Results	6 weeks after the injection, vaccinate serum samples were collected and pooled, then tested for antitoxin content by indirect Enzyme-Linked Immunosorbent Assay.A satisfactory value which met the requirements per 9 CFR 113.114(c) was achieved.
USDA Approval Date	February 15, 2011

Study Type	Efficacy									
Pertaining to	Eastern equine encephalomyelitis									
Study Purpose	Demonstration of efficacy against Eastern equine									
	encephalomyelitis									
Product Administration	Two doses, administered intramuscularly, 14 to 21 days apart									
Study Animals	12 guinea pigs (10 vaccinates, 2 controls)									
Challenge Description	Not applicable									
Interval observed after	Not applicable									
challenge										
Results	Serum samples were tested by a plaque reduction, serum neutralization test, 14 to 21 days after the second injection. Vaccinates and controls were evaluated in terms of Eastern equine encephalomyelitis per the criteria in 9 CFR 113.207(b) and the requirements were met.									
USDA Approval Date	February 15, 2011									

Study Type	Efficacy
Pertaining to	Western equine encephalomyelitis
Study Purpose	Demonstration of efficacy against Western equine
	encephalomyelitis
Product Administration	Two doses, administered intramuscularly, 14-21 days apart
Study Animals	12 guinea pigs (10 vaccinates, 2 controls)
Challenge Description	Not applicable
Interval observed after	Not applicable
challenge	
Results	Serum samples were tested by a plaque reduction, serum neutralization test, 14 days after the second injection. Vaccinates and controls were evaluated in terms of Western equine encephalomyelitis per the criteria in 9 CFR 113.207(b) and the requirements were met.
USDA Approval Date	February 15, 2011

Study Type	Efficacy												
Pertaining to	Equine herpesvirus type 1 (EHV-1)												
Study Purpose	Demonstration of efficacy against respiratory disease caused by EHV-1												
Product Administration	Two doses, adn	wo doses, administered intramuscularly, 21 days apart											
Study Animals	40 horses (20 v	0 horses (20 vaccinates, 20 controls), 4-5 months of age											
Challenge Description	Equine herpesv	Equine herpesvirus type 1 administered 15 days post-final											
	vaccination												
Interval observed after	Horses were ob	Iorses were observed daily for 14 days post-challenge											
challenge Results	Saa rayy data ar	ee raw data on following pages.											
Kesuits	See law uata of	ee raw data on following pages.											
	signs of respira classified as "n following class Disease status Normal Mild Moderate	tory disease formal", "m fication of th 0 or 1 1.5 or 2 4 or 6	Nasal Score	sal discharge was									
	The number of	horses in eac	h category were:										
	Norr	nal Mild	Moderate										
	Control 0	10	10										
	Vaccine 6	11	3										
USDA Approval Date	January 28, 200	9											

Nasal Discharge:

								tchall								
Treatment	ID	0	1	2	3	4	5	6	7	8	9	10	11	12	13	14
	1					1.5			1.5	1.5	1	1.5				
	2						1.5		1.5	1.5	1	1.5	1.5	1		
	3						1.5			1.5	2			1.5		
	4			1		2	1.5		1.5	1.5	1.5	1.5				1.5
	5				2	2	2	1	4	2	2	1.5	1.5		1.5	
	6			1		4	6	4	4	4	4	2	2	2		
	7					1.5	1.5	1.5	1.5	1.5	2	4	1		1	1
	8								1.5	2	2	4	1.5	4	2	1.5
	9				1	1.5	1.5	1.5	1.5	1.5	2	1.5				
Controls	10			1			1		1.5	1.5	2	4	4		1.5	1.5
(20 horses)	11						1.5	1.5	1.5		2		1.5	1.5	1.5	
	12						1.5	1.5		2						1.5
	13		l				2	1.5	1.5	2	2	2	1.5	1.5	1.5	4
	14				1.5	2		1.5		1.5	1.5	1.5			2	2
	15				1	2	1.5	1	1.5		4		1		4	1.5
	16					1.5	1.5	2	2	2	2	1.5	1	1	4	2
	17					1.5		1			1.5	2		1.5	1.5	
	18						1	1.5	1.5	4	4	2	1.5	4	1.5	2
	19				1	2	1.5		1.5	2	4	1	1.5		1	
	20						1.5	1.5	2	1.5	2				1.5	
	1					1		1				1.5				
	2				1											
	3						1	1.5	4		1.5	1.5			1	
	4				1						2	1				
	5				1				1	1						
	6				1	1.5						1.5	2	2	2	1.5
	7							2					1.5			
	8															
	9					2	1.5	2	2	6	2	1.5		1.5	4	2
Vaccinates	10								1				1	1.5		
(20 horses)	11				1		1.5		2	2	1	1.5				
,	12				1		1.5	2	1.5	2	2	2		2	2	1.5
	13				1.5						1.5	1.5			1.5	1.5
	14		İ				l	1	1		l	1	l		1.5	
	15		İ		1		l		l		l		l			
	16				1		1.5	1.5	1			1.5				
	17															
	18						1			1.5		1.5				
	19														6	2
	20		l				l		l		l		l			

Day Postchallenge

Scoring:

Blank is 0 =none;

1 = slight serous, as may be observed in both normal and diseased horses;

- 1.5 = very slight mucopurulent discharge;
- 2 = moderate clear serous discharge, or slight mucopurulent discharge;
- 3 = abundant serous discharge;
- 4 = moderate mucopurulent discharge;
- 6 = heavy mucopurulent discharge

Study Type	Efficacy												
Pertaining to	Equine herpesvirus ty	vpe 4 (EHV-4)											
Study Purpose	Demonstration of efficacy against respiratory disease caused by EHV-4 Two doses, administered intramuscularly, 21 days apart 40 horses (20 vaccinates, 20 controls), 4 months of age												
Product Administration	Two doses, administe	red intramuscularly	, 21 days apart										
Study Animals													
Challenge Description	Equine herpresvirus type 4 administered 14 days post-final vaccination Horses were observed daily for 14 days post-challenge												
Interval observed after challenge	Horses were observed daily for 14 days post-challenge												
Results	See raw data on follo	wing pages.											
	The horses were assessed for the presence of nasal and ocular discharge as signs of respiratory disease. The severity of the combined findings (nasal and ocular discharge) were classified as "mild" or "moderate" according to the following classification:Disease statusNasal scoreOcular scoreNormal = 00 or 10 or 1												
	score Normal = 0 0 or 1 0 or 1 Mild = 1 0 or 1 2 Mild = 1 1.5, 2, or 3 any												
	Moderate $= 2$	4 or 6	any										
	Moderate respiratory controls and 1/20 vac in 12/20 placebo cont None of the placebo c challenge, whereas 2 disease.	cinated horse, and m rols and 17/20 vacci controls remained he	nild disease was inated horses. ealthy following	observed									
USDA Approval Date	May 31, 2011												

Ocular Discharge:

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		19															

Day Postchallenge

Scoring:

Blank is 0=none 1=mild or moderate 2=severe

Nasal Discharge:

Day Postchallenge

				1		ostch										
Treatment	Animal	0	1	2	3	4	5	6	7	8	9	10	11	12	13	14
	1				1	1			1	2	3		3		3	
	2				2	3	3	2	2	3	3	2	4	3	3	2
	3				3	3		2	4			3	3	2	2	
Controls	4					4	4	3	3	4	3	3			2	2
	5					2	3	3	3		3	2	2		2	3
	6						3		2	4	3	3	2	3	2	
	7				1	2	1	2	2	2	2	3	2		2	2
	8								2		2					
	9							2	2	3	2	2	2	3		
	10				3	4	3	3	3	2		2	2	2	2	2
	11															1
	12						3		2	2	2				3	3
	13					3	2	2	2	2	1	2	2			
	14				2	3	4	4	2	4	2	4	3	4	3	
	15				1		3	3	3	3		3	3			2
	16				3	3	3	4	2	4	4	3	4	2	2	2
	17					1		2	2	3	2		3	3		
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	19						1	4	2	3		3			2	3
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	4				1											
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	6										3					
	7					1										
	8							2	3	1	3					
	9											1				2
Vaccinates	10										3		2			
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	12								3	2	3	1	3			2
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	18									4	2		2		2	
	19															
	20								2			3	3			

Dav Postchallenge

Scoring:

Blank is 0 = none

1 = slight clear serous, as may be observed in both normal and diseased horses;

1.5 = very slight mucopurulent discharge, one or both nostrils;

2 = moderate clear serous discharge, easily seen in one or both nostrils;

3 = abundant clear serous discharge typically seen only in diseased horses;

4 = moderately mucopurulent, in large quantities in both nostrils;

5 = heavy mucopurulent discharge in large amounts in both nostrils

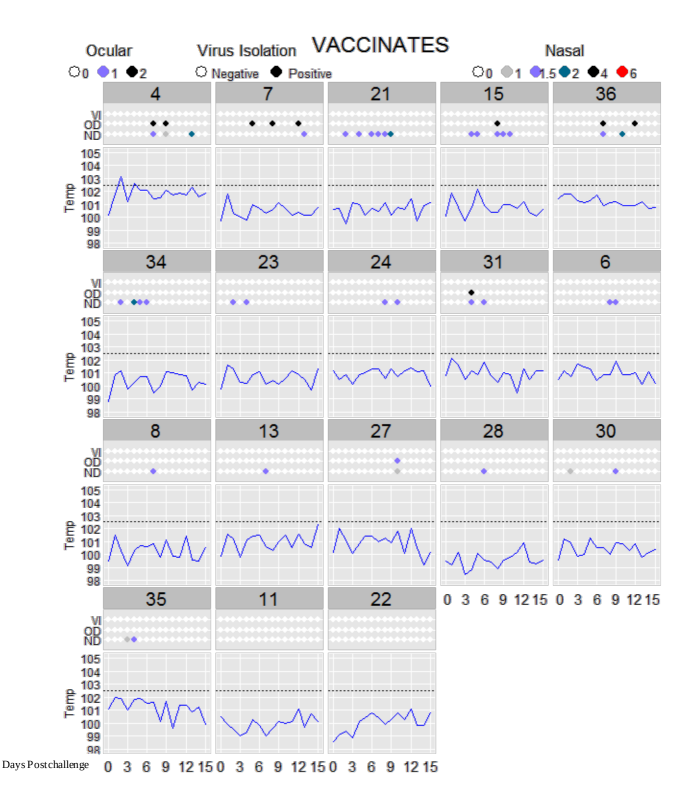
Study Type	Efficacy
Pertaining to	Equine influenza virus
Study Purpose	Demonstration of 6-month duration of immunity against
,	respiratory disease caused by equine influenza
Product Administration	Two doses, administered intramuscularly, 21 days apart.
	Vaccinates received test product, and controls received
	adjuvanted diluent.
Study Animals	30 horses (20 vaccinates, 10 controls), 5-6 months of age
Challenge Description	Influenza A/eq/Ohio/2003 administered 184 days post-final
	vaccination
Interval observed after	Horses were observed daily for 10 days post-challenge
challenge	
Results	See tables at the end of document for data.
	 Clinical Signs: An animal was considered positive (affected by challenge) if the animal exhibited: Fever (temperature >102.5°F), OR Nasal discharge (moderate serous discharge or mucopurulent discharge), OR Ocular discharge A total of 9/10 (90%) controls were positive as compared to only 9/20 (45%) vaccinates. There were no adverse reactions to vaccine administration at any timepoint.
USDA Approval Date	September 7, 2010

					D	ays Po	ost-ch	alleng	ge			
Treatment	Clinical Sign	0	1	2	3	4	5	6	7	8	9	10
Controls												
	Fever											
1	Nasal discharge						+	+	+	+		
	Ocular discharge						+			+		+
	Fever											
2	Nasal discharge			+			+		+	+	+	
	Ocular discharge						+	+			+	+
	Fever											
3	Nasal discharge							+		+		
	Ocular discharge			+			+			+		+
	Fever											
4	Nasal discharge											
	Ocular discharge						+	+	+			+
	Fever											
5	Nasal discharge					+	+	+	+	+	+	
	Ocular discharge											
	Fever											
6	Nasal discharge					+			+		+	+
	Ocular discharge											+
	Fever											
7	Nasal discharge			+			+		+			+
	Ocular discharge			+				+				
	Fever								+			
8	Nasal discharge						+	+	+			+
	Ocular discharge			+	+		+	+				+
	Fever											
9	Nasal discharge											
	Ocular discharge											
	Fever											
10	Nasal discharge						+	+	+	+	+	
	Ocular discharge					+	+		+	+	+	

					D	ays P	ost-ch	allen	ge			
Treatment	Clinical Sign	0	1	2	3	4	5	6	7	8	9	10
Vaccinates												
	Fever											
1	Nasal discharge											
	Ocular discharge											
	Fever											
2	Nasal discharge											
	Ocular discharge											
	Fever											
3	Nasal discharge											
	Ocular discharge						+			+	+	
	Fever											
4	Nasal discharge								+			
	Ocular discharge											
	Fever											
5	Nasal discharge											
	Ocular discharge											
	Fever											
6	Nasal discharge											
	Ocular discharge											
	Fever											
7	Nasal discharge											
	Ocular discharge											
	Fever											
8	Nasal discharge											
	Ocular discharge											
	Fever											
9	Nasal discharge											
	Ocular discharge											
	Fever						+					
10	Nasal discharge							+	+			
	Ocular discharge									+		+
	Fever											
11	Nasal discharge						+			+	+	+
	Ocular discharge											
	Fever											
12	Nasal discharge									+		
	Ocular discharge						1		1	1		

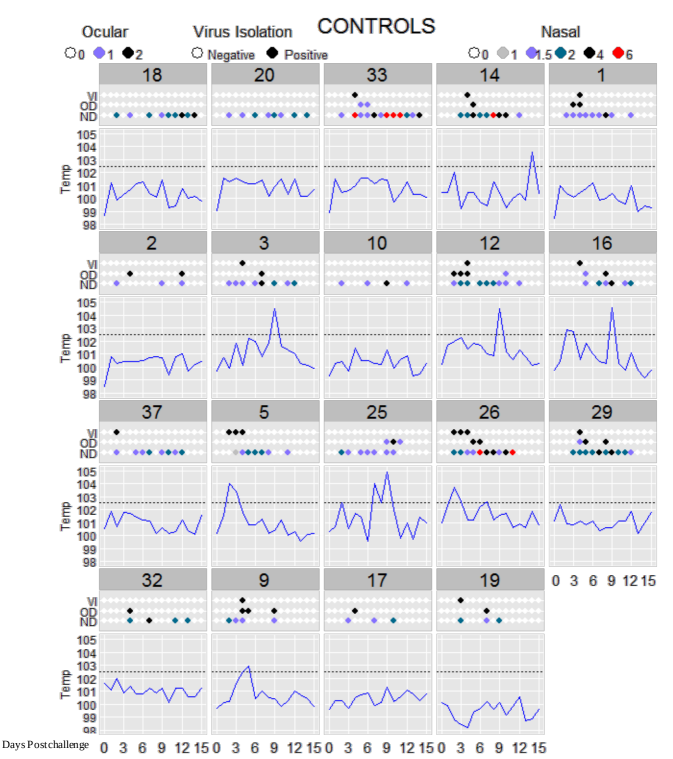
					D	ays P	ost-ch	allen	ge			
Treatment	Clinical Sign	0	1	2	3	4	5	6	7	8	9	10
Vaccinates												
	Fever											
13	Nasal discharge					+						+
	Ocular discharge											
	Fever											
14	Nasal discharge											
	Ocular discharge											
	Fever											
15	Nasal discharge											
	Ocular discharge						+		+			
	Fever											
16	Nasal discharge							+				
	Ocular discharge											
	Fever											
17	Nasal discharge											
	Ocular discharge											
	Fever											
18	Nasal discharge											
	Ocular discharge											
	Fever											
19	Nasal discharge							+		+		
	Ocular discharge											
	Fever											
20	Nasal discharge											
	Ocular discharge											

Study Type	Efficacy
Pertaining to	Equine influenza virus
Study Purpose	Demonstration of efficacy against respiratory disease and shedding caused
	by equine influenza
Product	Two doses, administered intramuscularly, 21 days apart.
Administration	
Study Animals	37 horses (18 vaccinates, 19 controls), approximately 9-10 months of age
Challenge	Influenza A/eq/Ohio/2003 administered 3 weeks post-final vaccination
Description	
Interval	Horses were observed, and nasal swabs were collected, daily for 15 days
observed after	post-challenge.
challenge	
Results	See tables at the end of document for data.
	Clinical Signs:
	An animal was considered positive (affected by challenge) if the animal
	exhibited the following at any post-challenge observation point:
	• Fever (temperature $\geq 102.5^{\circ}$ F), OR
	Ocular discharge, OR
	• Nasal discharge (very slight mucopurulent discharge, or worse)
	Duration of disease was calculated from the date the animal was first observed to be positive to the date of last positive observation for that animal. Based on this calculation, the median duration of disease for the controls was determined to be 11 days as compared to 3 days for the vaccinates.
	Nasal shedding of influenza virus was evaluated through nasal swab virus isolation results. An animal was considered positive if virus was isolated from nasal swabs on one or more occasions following challenge.
	0/18 vaccinates shed virus and 12/19 controls shed virus.
	There were no adverse reactions to vaccine administration at any timepoint.
USDA Approval Date	April 8, 2013



Ocular Discharge: 0=none; 1=mild to moderate; 2=severe

Nasal Discharge: 0=none; 1=slight clear serous, as may be observed in both normal and diseased horses; 1.5=very slight mucopurulent discharge, one or both nostrils; 2=moderate clear serous discharge, easily seen in one or both nostrils; 3=Abundant clear serous discharge typically seen only in diseased horses; 4=moderately mucopurulent, in large quantities in both nostrils; 5=heavy mucopurulent discharge in large amounts in both nostrils



Ocular Discharge: 0=none; 1=mild to moderate; 2=severe

Nasal Discharge: 0=none; 1=slight clear serous, as may be observed in both normal and diseased horses; 1.5=very slight mucopurulent discharge, one or both nostrils; 2=moderate clear serous discharge, easily seen in one or both nostrils; 3=Abundant clear serous discharge typically seen only in diseased horses; 4=moderately mucopurulent, in large quantities in both nostrils; 5=heavy mucopurulent discharge in large amounts in both nostrils

C4d. True o	Efficient
Study Type	Efficacy
Pertaining to	Equine influenza
Study Purpose	Demonstration of efficacy against respiratory disease caused by
	equine influenza A2 strain Richmond 07
Product Administration	Two doses, administered intramuscularly, 21 days apart
Study Animals	20 horses (20 vaccinates), 12 months of age
Challenge Description	Not applicable
Interval observed after	Not applicable
challenge	
Results	This product class allows the manufacturer to update micro- organisms in this vaccine under expedited procedures to respond to emerging needs. Abbreviated data to support influenza strain updates to the product composition were evaluated by USDA- APHIS and found to be acceptable based on regulations and policies at the time of approval. Full vaccination-challenge studies may not have been required for these updates.
USDA Approval Date	February 2, 2012

Study Type	Efficacy
Pertaining to	Equine influenza
Study Purpose	Demonstration of efficacy against respiratory disease caused by
	equine influenza A2 strain Kentucky 95
Product Administration	Two doses, administered intramuscularly, 21 days apart
Study Animals	20 horses (20 vaccinates), 12 months of age
Challenge Description	Not applicable
Interval observed after	Not applicable
challenge	
Results	This product class allows the manufacturer to update micro- organisms in this vaccine under expedited procedures to respond to emerging needs. Abbreviated data to support influenza strain updates to the product composition were evaluated by USDA- APHIS and found to be acceptable based on regulations and policies at the time of approval. Full vaccination-challenge studies may not have been required for these updates.
USDA Approval Date	February 2, 2012

Study Type	Safety							
Pertaining to	All fractions							
Study Purpose	To demonstr	ate safety u	nder field condi	itions at th	ree differ	ent test sit	tes	
Product	2 doses given intramuscularly 21 days apart							
Administration								
Study Animals	622 horses vaccinated with two doses including:							
			nonth-old foals					
			month-old foals					
Challange		1 year or ol	der norses					
Challenge Description	Not Applicat	ble						
Interval	Horses were	observed or	n Days 0, 1 and	3 followi	ng the firs	t vaccinat	ion and	
observed after			wing the second		-			
vaccination	injection site							
Results			reactions obser	ved at any	of the th	ree sites.	Local	
			re summarized					
	N 1 S 1	a.						
	North Dakot	a Site:		Tuon	sient			
	Summary	Total	Number		on Site	Number	Normal	
	j samma y	Number	with 2 doses	Swelling		rumber ruma		
	Age			1 st dose	2 nd dose	1 st dose	2 nd dose	
	2-4 mo	149	149	0	0	149	149	
	5-7 mo	0	0	n/a	n/a	n/a	n/a	
	8-11 mo	0	0	n/a	n/a	n/a	n/a	
	1 yr-5yr	23	23	0	0	23	23	
	6-15 yr	121	121	0	0	121	121	
	>16 yr	3	3	0	0	3	3	
	Total	296	296	0	0	296	296	
	California Si	te:						
		Total	Number		sient			
	Summary	Number	with 2 doses		on Site	Number	Normal	
	Age			1 st dose	lling 2 nd dose	1 st dose	2 nd dose	
	2-4 mo	0	0	n/a	n/a	n/a	n/a	
	5-7 mo	5	5	0	0	5	5	
	8-11 mo	0	0	n/a	n/a	n/a	n/a	
	1 yr-5yr	25	25	0	4	25	21	
	6-15 yr	15	15	0	3	15	12	
	>16 yr	6	6	0	1	6	5	
	Total	51	51	0	8*	51	43	
			were minimal.	The reported	d reactions	were mild,	transient,	
	non-painful in	njection swell	lings.					

	Missouri Site	•								
	Summary	Total Number	Number with 2 doses	Transient Injection Site Swelling		Number Normal				
	Age			1 st dose	2 nd dose	1 st dose	2 nd dose			
	2-4 mo	55	54	0	0	55	54			
	5-7 mo	15	14	0	0	15	14			
	8-11 mo	0	0	n/a	n/a	n/a	n/a			
	1 yr-5yr	134	132	0	0	134	132			
	6-15 yr	68	68	0	0	68	68			
	>16 yr	7	7	0	0	7	7			
	Total	279	275	0	0	279	275			
	Total Across Three Sites:									
	Site	TotalNumberNumberwith 2 doses		Injecti Swe	on Site lling	Number Normal				
				1 st dose	2 nd dose	1 st dose	2 nd dose			
	North Dakota	296	296	0	0	296	296			
	California	51	51	0	8*	51	43			
	Missouri	279	275	0	0	279	275			
	Total	626	622	0	8*	626	614			
	swellings afte	er the second	were minimal an vaccination in eig actions observed.	ght (8) olde			1			
USDA	February 14, 2012									
Approval Date	reordary ri,	2012								

Study Type	Safety
Pertaining to	All fractions
Study Purpose	To demonstrate safety in pregnant mares under field conditions at
	two different test sites
Product	Two intramuscular doses, given 16-28 days apart. 54 pregnant mares
Administration	were injected with placebo and 325 pregnant mares were vaccinated
	with test product.
Study Animals	Three hundred seventy-nine pregnant mares at two locations were
	included in the study. The mares were confirmed to be pregnant by
	serum hormonal evaluation on the day of the first vaccination.
Challenge	Not applicable
Description	
Interval observed	1 st and 2 nd trimester: Mares observed immediately after vaccination
after vaccination	and daily for overall health and for abortion. Resulting foals were
	observed daily for 7 days following birth.
	3 rd trimester: Mares observed immediately after vaccination and
	daily for overall health and for abortion. Resulting foals were
	observed daily for 30 days following birth.
Results	Results shown on next page

Results	Study 2013 North Daka									
	Group	Vaccin		Confirmed Pregnant	Foals		Parturition Rate			
	1 st trimester product	c/ 143		27			90%			
	1st trimeste placebo	r/ 59	5	54	49	9	91%			
	2 nd trimeste product	r/ 6	6	5	6		100%			
	3 rd trimester product	r/ 140	1	.17	117		100%			
	Total – all animals	348	3	304	286	9	94%			
	Total – product on	289	2	250	237	9	95%			
	Total – placebo on	59	5	54	49	9	91%			
		Study 2013-PM-1009								
	Group	Vaccin		onfirmed regnant			Parturition Rate			
	2011 3 rd trimester	5	5	- Built	5		0%			
	2012 1 st trimester	1	1		1	10	0%			
	2012 2 nd trimester	53	43	3	39	91	%			
	2012 3 rd trimester	26	20	5	25	96	%			
	Total – product	85	75	5	70 93		3%			
	Study 2014-PM-1009 North Dakota Site:									
		Vaccinated	Confirm Pregnan		d Parturi Rate	ition	Foals Survived to End of Observation Period			
	2 nd trimester	52	52	52	100%		51*			
	vaccinated 3 rd trimester vaccinated	69	69	67**	97.1%		67			
	*Lost foal af **One mare cooperator.	*Lost foal affirmed by study cooperator to be due to causes other than vaccination **One mare died due to causes other than vaccination, as affirmed by study								
	AII UIIEI 10a	is were normal	i anu nealtí	LY						