

# Summary of Studies Supporting USDA Product Licensure

Establishment Name	Boehringer Ingelheim Animal Health USA Inc.
USDA Vet Biologics Establishment Number	124
Product Code	4855.24
True Name	Encephalomyelitis-Rhinopneumonitis-Influenza-West Nile Virus Vaccine, Eastern & Western, Killed Virus, Tetanus Toxoid
Tradename(s) / Distributor or Subsidiary (if different from manufacturer)	Equi-Jec 6 - 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									
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	11									
Results	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	May 1, 2008									

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	May 1, 2008

Study Type	Efficacy
Pertaining to	Western equine encephalomyelitis
Study Purpose	Demonstration of efficacy against Western 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 Western equine encephalomyelitis per the criteria in 9 CFR 113.207(b) and the requirements were met.
USDA Approval Date	May 1, 2008

Study Type	Efficacy													
Pertaining to	Equine herpesvirus type 1 (EHV-1)													
Study Purpose	Demonstration of efficacy against respiratory disease caused by EHV-1 Two doses administered intramuscularly, 21 days apart													
Product Administration	Two doses, adn	Two 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 herpesvirus type 1 administered 15 days post-final													
	vaccination													
Interval observed after	Horses were ob	Horses were observed daily for 14 days post-challenge												
challenge Results	Saa rayy data ar	See 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											
	<b>Control</b> 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		İ				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		l				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		İ				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	Efficacy Equine herpesvirus (EHV) type 1											
Study Purpose	Equine herpesvirus (EHV) type 1 Demonstration of efficacy against EHV-4											
Product Administration	Two doses administered intramuscularly 21 days apart											
	Two doses administered intramuscularly 21 days apart 37 horses (24 vaccinates, 13 controls), 4 to 5 months of age											
Study Animals							Ų					
Challenge Description	Equine herpesv second vaccinat	•	ype 4 wa	is adi	ninistered	15 da	lys following					
Interval observed after challenge	Horses were observed daily for 14 days following challenge											
Results	The horses were assessed for the presence of nasal and ocular discharge as signs of respiratory disease. The severity of the nasal and ocular discharge resulted in the classification of the observed clinical signs as "mild" or "moderate" according to the following											
	table:											
	Disease Status Nasal Score Ocular Score											
	Normal $= 0$		$\frac{0 \text{ or } 1}{0 1}$		0 or 1 2							
	Mild = 1		$\frac{0 \text{ or } 1}{1.5 2 \text{ or }}$	2	_							
	Moderate = 2		1.5, 2, or 4 or 6	3	Any							
	Iviouerate – 2	2 2	4 01 0		Any							
	Respiratory dise	ease w	was obse	rved	as follows	s:						
		Contr	rols	Vac	cinates							
	Normal	0 out			t of 24							
	Mild		t of 13		out of 24							
	Moderate 1 out of 13 1 out of 24											
	See raw data on	n the f	following	g pag	es.							
USDA Approval Date	September 17, 2	2009										

# **Ocular Discharge:**

	Horse						Γ	<b>D</b> avs	Post	t-cha	allen	ge				
Treatment	ID	0	1	2	3	4	5	6	7	8	9	10	11	12	13	14
	8300															
	8861								1		1	2	1	1	1	1
	0327										1	1				1
	1029								1	1	1		1		1	1
	2603											1				
	8822															
Controls	5542												1	1	1	2
(13 horses)	3280				1				1				1	1	1	1
	5597														1	2
	9331								2	2	2	2	1		2	1
	9339											1	1			1
	5103															
	6528															
	1278							1					1		1	
	1602							1	1	1	1	1	1	2		
	8026															
	3857									1	1	1				1
	5560															
	5636															
	0261											1	1			
	0285						1	1								1
	6051															
	6311							1			1					
	1310															
Vaccinates	5381															
(24 horses)	8023												1			
	8881				1	1		1					1			
	0019											1				
	1381		1			1										
	2333		1			1										1
	3086		1			1										
	3347		1			1										
	5379		1	1		1		Ì	Ì	Ì			1			
	7297		1			1										
	7580		1	1		1		1	Ì	Ì			1			
	7806		1			1		1	1							
	8004	1	1			1								1		
Saaring	000	I	I	<u> </u>	1	I	1	I	I	I	I	I	I	L	I	I

Scoring: No entry indicates 0 = horse is normal

1 = normal

2 = mild

#### **Nasal Discharge:**

	Horse						]	Days ]	Post-	challe	nge					
Treatment	#	0	1	2	3	4	5	6	7	8	9	10	11	12	13	14
	8300							1.5	1.5	1.5	1.5			1.5		
	8861							1.5	1	1			1.5			1.5
	0327							1	1		1.5	1.5		2		1
	1029				1.5						1.5			1.5	1.5	
	2603				1.5			1.5			1.5					
Controla	8822					2	1.5	1.5		2						
Controls (13 horses)	5542								1.5	1.5	2	1.5	1		1.5	2
(15 norses)	3280					1		2								
	5597					1.5		1	1.5	2	1.5		1		1.5	1.5
	9331						1.5	1.5	1.5	4	1.5	1			1.5	1.5
	9339							1.5	1.5	1.5	1.5	1.5	2	1		
	5103					1	1.5	2	1.5	2	1.5	2				
	6528							1.5	1.5			1.5		1		
	1278					1.5		1								
	1602							1.5	1.5	2	2	2	2	4		
	8026					2	1.5	1				2	1.5	2		
	3857									1.5	1.5					
	5560															1.5
	5636									1.5						
	0261										1.5					
	0285										1.5					
	6051				1.5					1						1.5
	6311							1.5			2					
	1310															
Vaccinates	5381															
(24 horses)	8023															
	8881				1.5	1.5					1.5		1.5			
	0019															
	1381							1.5			1.5					
	2333															
	3086															
	3347				1.5											
	5379														2	
	7297															
	7580					1										1.5
	7806															
Securing	8004										1					

## Scoring:

No entry is 0 = horse is normal

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 or slightly mucopurulent discharge, in one or both nostrils;

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

- 4 = moderately mucopurulent discharge, in large quantities in both nostrils;
- 6 = heavy mucopurulent discharge in large amounts filling both nostrils

Study Type	Efficacy
Pertaining to	Equine influenza virus
Study Purpose	Demonstration of 6-month duration of immunity against
<b>,</b>	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.
	<ul> <li>Clinical Signs: An animal was considered positive (affected by challenge) if the animal exhibited: <ul> <li>Fever (temperature &gt;102.5°F), OR</li> <li>Nasal discharge (moderate serous discharge or mucopurulent discharge), OR</li> <li>Ocular discharge</li> </ul> </li> <li>A total of 9/10 (90%) controls were positive as compared to only 9/20 (45%) vaccinates.</li> <li>There were no adverse reactions to vaccine administration at any timepoint.</li> </ul>
USDA Approval Date	September 7, 2010

					D	ays Po	ost-ch	allen	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
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	Efficacy
Pertaining to	Equine influenza
Study Purpose	Demonstration of efficacy against respiratory disease caused by
	equine influenza A2 strain New Market 2/93
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	West Nile Virus (WNV)		
Study Purpose	Demonstration of twelve m	onth duration of	immunity against disease
	caused by WNV		
Product Administration	Two doses, administered intr	amuscularly, 25 da	ays apart
Study Animals	30 horses (20 vaccinates, 10	placebo controls) 4	4-5 months of age
Challenge Description	West Nile Virus was admin	istered at 380 day	ys (10 vaccinated and 5
	placebo control animals) or	r 408 days (10 v	accinated and 5 placebo
	control animals) post-final va	accination.	
Interval observed after	Horses were observed twice	e daily for 14 da	ys post-challenge and
challenge	once daily for an additiona	l 7 days post-cha	llenge.
Results	An animal was considered	affected by chall	lenge if it developed
	neurological disease, as me	easured by morta	lity and microscopic
	evidence of virus-induced	brain disease (his	stopathology).
		1.6 • • • /	
	Animals were also monitor	red for viremia (c	letection of WNV in
	the blood).		
	Results are summarized as	follows	
	Outcome	Controls	Vaccinates
	Mortality	7/10 (70%)	1/20 (5%)
	Viremia at least one day	10/10 (100%)	2/20 (10%)
	- I china at reast one day		
	See raw data on following	nages	
		P.9.9.	
USDA Approval Date	September 3, 2010		

Treatment	#	Died or Euthanized due	Severity Histopat	hological lesions
Treatment	#	to disease severity	Medulla	Pons
	1	Yes	3	3
	2	Yes	3	3
	3	Yes	3	3
	4	Yes	3	3
Controls	5	Yes	3	3
(10 horses)	6	Yes	2	2
	7	Yes	1	1
	8	No	1	1
	9	No	1	1
	10	No	1	0.5
	1	Yes	3	3
	2	No	2	0.5
	3	No	1	1
	4	No	1	0.5
	5	No	1	0.5
	6	No	1	0.5
	7	No	0.5	0.5
	8	No	0.5	0.5
	9	No	0.5	0
Vaccinates	10	No	0	0.5
(20 horses)	11	No	0	0
	12	No	0	0
	13	No	0	0
	14	No	0	0
	15	No	0	0
	16	No	0	0
	17	No	0	0
	18	No	0	0
	19	No	0	0
	20	No	0	0

Scoring of hi	stopathological lesions:
0 =	No significant lesions.
0.5 =	Rare, small, multifocal glial nodules scattered throughout the parenchyma.
1 =	Mild, nonsuppurative encephalitis characterized by mild multifocal perivascular cuffs with lymphocytes and plasma cells and a rare neutrophil and scattered multifocal glial nodules composed of glial cells with a few mononuclear inflammatory cells. Occasionally within this grade, there may be minimval perivascular cuffing and more moderate scattered glial nodules.
2 =	Moderate nonsuppurative encephalitis characterized by moderate lymphoplasmacytic perivascular cuffs around many vessels and multifocal accumulations of glial nodules scattered throughout the parenchyma.
3 =	Severe nonsuppurative encephalitis characterized by severe and thick lymphoplasmacytic perivascular cuffing with multiple scattered glial nodules throughout the parenchyma.

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	_			$\vdash$	+	30	25	50	10									P	þ	þ
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3       1	2		$\vdash$																	
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8       40       9       9       9         9       10       10       10       10       10       10         11       10       10       10       10       10       10       10         12       13       12       10       1																				
9         1	~				40															
Vaccinates       10       10       10         11       11       11       11       11         12       11       11       11       11       11         13       11       11       11       11       11       11         14       11 <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>																				
(20 horse)       11       (20 horse)         12       1       1         13       1       1         14       1       1         13       1       1         14       1       1         15       1       1         16       1       1         17       1       1         18       1       1         19       1       1         10       1       1         11       1       1         12       1       1         13       1       1         14       1       1         15       1       1         16       1       1         17       1       1         18       1       1         19       1       1         10       1       1         11       1       1         12       1       1         13       1       1         14       1       1         15       1       1         16       1       1         17																				
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14         12         15         10         11         11         12         13         14         15         16         17         18         19         11         11         12         13         14         15         16         17         18         19         11         11         12         13         14          15          16          17          18          19         11          10         11          12          13          14          15          16          17          18          19         10          11          11         14        15        16        17 <td< td=""><td>13</td><td></td><td><math>\vdash</math></td><td><math>\vdash</math></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></td<>	13		$\vdash$	$\vdash$																
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	N = Not recorded; horse was circling with sporadic head / neck tremors.	was circ	ling w	rith sp(	oradic	head /	neck tr	emors.												

Study Type	Efficient				
Study Type	Efficacy				
Pertaining to		virus (WNV)			
Study Purpose				of immunity against V	VNV
<b>Product Administration</b>				ly 22 days apart	
Study Animals				ntrols) 4-5 months of a	ge
Challenge Description				l days (Group 1:	
		-		mals) or 222 days	
	· •		nd 5 placebo	control animals) after the	he
	second vacc				
Interval observed after				ge, twice daily for 6 day	
challenge	post-challen	ge, once daily	for an addition	onal 4 days post-challer	nge,
	and on day 1	14 post-challe	nge		
Results	The primary	outcome was	s viremia (dete	ection of WNV in the	
	blood). An	animal was co	onsidered to be	e positive if virus was	
	detected in t	he blood on o	ne or more oc	casions post-challenge	•
	The number	of animals po	ositive for vire	emia at least once is	
	summarized	for as follow	s:		
	Challenge	Controls	Vaccinates		
	Group				
	1	5/5 (100%)	1/10 (10%)		
	2				
		5/5 (100%)	3/10 (30%)		
	The outcom	e for <b>viremia</b>	is as follows	for the first group of ho	orses
	The outcom	e for <b>viremia</b> 201 days follo	is as follows t wing the seco	nd vaccination:	orses
	The outcom	e for <b>viremia</b> 201 days follo Horse ID	is as follows twing the seco	nd vaccination:	orses
	The outcom	e for <b>viremia</b> 201 days follo Horse ID S16	is as follows t wing the seco Challenge Posi	nd vaccination: e Group 1 tive	orses
	The outcom challenged 2	e for <b>viremia</b> 201 days follo Horse ID S16 S21	is as follows t wing the seco Challenge Posi Posi	nd vaccination: <b>Group 1</b> tive tive	orses
	The outcom challenged 2 Controls	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23	is as follows t wing the seco Challenge Posi Posi Posi	nd vaccination: <b>Group 1</b> tive tive tive	orses
	The outcom challenged 2	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26	is as follows t wing the seco Challenge Posi Posi Posi Posi	nd vaccination: <b>Group 1</b> tive tive tive tive	orses
	The outcom challenged 2 Controls	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30	is as follows t wing the seco Challenge Posi Posi Posi Posi Posi	nd vaccination: <b>Group 1</b> tive tive tive tive tive tive	orses
	The outcom challenged 2 Controls	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30 S17	is as follows t wing the seco Challenge Posi Posi Posi Posi Posi Nega	nd vaccination: <b>Group 1</b> tive tive tive tive tive tive tive	orses
	The outcom challenged 2 Controls	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30 S17 S18	is as follows t wing the seco Challenge Posi Posi Posi Posi Posi Nega Nega	nd vaccination: <b>Group 1</b> tive tive tive tive tive tive tive tive tive tive	orses
	The outcom challenged 2 Controls	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30 S17 S18 S19	is as follows t wing the seco Challenge Posi Posi Posi Posi Posi Nega Nega	nd vaccination: <b>Group 1</b> tive tive tive tive tive tive tive tive tive tive tive tive tive tive	orses
	The outcom challenged 2 Controls (5 horses)	e for <b>viremia</b> 201 days follo <b>Horse ID</b> S16 S21 S23 S26 S30 S17 S18 S19 S20	is as follows t wing the seco Challenge Posi Posi Posi Posi Nega Nega Nega Nega	nd vaccination: <b>Group 1</b> tive tive tive tive tive tive tive tive tive tive tive tive tive tive tive tive tive tive	orses
	The outcom challenged 2 Controls (5 horses) Vaccinates	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30 S17 S18 S19 S20 S20 S22	is as follows t wing the seco Challenge Posi Posi Posi Posi Nega Nega Nega Nega Nega Nega	nd vaccination: <b>Group 1</b> tive tive tive tive tive tive ative ative ative ative tive	orses
	The outcom challenged 2 Controls (5 horses)	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30 S17 S18 S19 S20 s S22 S20 s S22 S20 s S22 S24	is as follows t wing the seco Challenge Posi Posi Posi Posi Nega Nega Nega Nega Nega Nega Nega	nd vaccination: <b>Group 1</b> tive tive tive tive tive tive ative ative ative ative ative ative ative ative	prses
	The outcom challenged 2 Controls (5 horses) Vaccinates	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30 S17 S18 S19 S20 S20 S22 S22 S24 S25	is as follows t wing the seco Challenge Posi Posi Posi Posi Nega Nega Nega Nega Nega Nega Nega Nega	nd vaccination: Group 1 tive tive tive tive tive tive ative ative ative ative ative ative ative ative ative ative ative ative	prses
	The outcom challenged 2 Controls (5 horses) Vaccinates	e for <b>viremia</b> 201 days follo <b>Horse ID</b> S16 S21 S23 S26 S30 S17 S18 S19 S20 S S22 S22 S22 S24 S25 S27	is as follows f wing the seco Challenge Posi Posi Posi Posi Nega Nega Nega Nega Nega Nega Nega Nega	nd vaccination: <b>Group 1</b> tive tive tive tive tive tive ative ative ative ative ative ative ative ative ative ative ative	orses
	The outcom challenged 2 Controls (5 horses) Vaccinates	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30 S17 S18 S19 S20 s S22 S20 s S22 S22 S24 S25 S27 S28	is as follows t wing the seco Challenge Posi Posi Posi Posi Nega Nega Nega Nega Nega Nega Nega Nega	nd vaccination: <b>Group 1</b> tive tive tive tive tive tive ative ative ative ative ative ative ative ative ative ative ative ative ative ative	prses
	The outcom challenged 2 Controls (5 horses) Vaccinates (10 horses	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30 S17 S18 S19 S20 S S22 S22 S24 S25 S27 S28 S29	is as follows t wing the seco Challenge Posi Posi Posi Posi Nega Nega Nega Nega Nega Nega Nega Nega	nd vaccination: <b>Group 1</b> tive tive tive tive tive tive ative ative ative ative ative ative ative ative ative ative ative ative ative ative ative ative ative ative	
	The outcom challenged 2 Controls (5 horses) Vaccinates (10 horses Positive = W	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30 S17 S18 S19 S20 S S22 S22 S24 S25 S27 S28 S29 NV detected in b	is as follows t wing the seco Challenge Posi Posi Posi Posi Nega Nega Nega Nega Nega Nega Nega Nega	nd vaccination: Group 1 tive tive tive tive tive tive ati	
	The outcom challenged 2 Controls (5 horses) Vaccinates (10 horses Positive = W	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30 S17 S18 S19 S20 S S22 S22 S24 S25 S27 S28 S29 NV detected in b	is as follows t wing the seco Challenge Posi Posi Posi Posi Nega Nega Nega Nega Nega Nega Nega Nega	nd vaccination: <b>Group 1</b> tive tive tive tive tive tive ative ative ative ative ative ative ative ative ative ative ative ative ative ative ative ative ative ative	
	The outcom challenged 2 Controls (5 horses) Vaccinates (10 horses Positive = W	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30 S17 S18 S19 S20 S S22 S22 S24 S25 S27 S28 S29 NV detected in b	is as follows t wing the seco Challenge Posi Posi Posi Posi Nega Nega Nega Nega Nega Nega Nega Nega	nd vaccination: Group 1 tive tive tive tive tive tive ati	
	The outcom challenged 2 Controls (5 horses) Vaccinates (10 horses Positive = W	e for <b>viremia</b> 201 days follo Horse ID S16 S21 S23 S26 S30 S17 S18 S19 S20 S S22 S22 S24 S25 S27 S28 S29 NV detected in b	is as follows t wing the seco Challenge Posi Posi Posi Posi Nega Nega Nega Nega Nega Nega Nega Nega	nd vaccination: Group 1 tive tive tive tive tive tive ati	

1				
			as follows for the seco	
	horses challeng		following the second v	accination:
		Horse ID	Challenge Group 2	
		S32	Positive	
	Controls	S36	Positive	
	Controls	S39	Positive	
	(5 horses)	S40	Positive	]
		S43	Positive	]
		S31	Negative	
		S33	Positive	
		S34	Negative	
		S35	Positive	
	Vaccinates	S37	Negative	
	(10 horses)	S38	Negative	
		S41	Negative	
		S42	Negative	]
		S44	Negative	]
		S45	Positive	]
			od on one or more occasion	
	Negative = WNV	V detected in blo	ood on zero occasions post-	challenge
USDA Approval Date	November 2, 2	009		

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 <sup>st</sup> and 2 <sup>nd</sup> 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 <sup>rd</sup> 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 Dake							
	Group	Vaccin		Confirm Pregnant		Foals		Parturition Rate
	1 <sup>st</sup> trimester product	:/ 143		127		114		90%
	1st trimeste placebo	r/ 59		54		49	9	91%
	2 <sup>nd</sup> trimeste product	r/ 6	,	6		6		100%
	3 <sup>rd</sup> trimester product	r/ 140		117		117		100%
	Total – all animals	348		304		286	9	94%
	Total – product on	289		250		237	9	95%
	Total – placebo on	59		54		49	9	91%
	Study 2013 Misssouri S	-PM-1009				_		
	Group	Vaccin		Confirme Pregnant	I I	Foals		arturition ate
	2011 3 <sup>rd</sup> trimester	5	5		5	5	_	00%
	2012 1 <sup>st</sup> trimester	1	1		1	-	10	00%
	2012 2 <sup>nd</sup> trimester	53	4	3	3	89	91	%
	2012 3 <sup>rd</sup> trimester	26	2	6	2	25	96	5%
	Total – product	85	7	5	7	/0	93	5%
	Study 2014 North Daka		·					
		Vaccinated	Confirm Pregnan		aled	Parturit Rate	ion	Foals Survived to End of Observation Period
	2 <sup>nd</sup> trimester	52	52	52		100%		51*
	vaccinated 3 <sup>rd</sup> trimester vaccinated	69	69	67*	**	97.1%		67
	*Lost foal af **One mare cooperator.	firmed by stud died due to cau ls were normal	uses other	than vacc				han vaccination by study
	All other 10a	is were normal	i anu nealt	пу				

Study PurposeTo deStudy PurposeTo deProductTwo deAdministrationStudy AnimalsStudy Animals880 hChallengeNot ajDescriptionNot ajIntervalNot ajObserved afterHorseChallengeHorseChallengeAdver	doses, orses, pplical pplical es were ution o e were orse di rse even nat reso	including 21 including 21 ble ble e observed at f any observe no systemic ied from caus	reactions observ ses affirmed by ited to transient	y approxi s of age a owing eac yed at any licensee i	and 52 foa the vaccina of the sinot attribute	als 5 mont ation, until tes. Two fe uted to vac	hs of age
ProductTwo ofAdministrationTwo ofStudy Animals880 heChallengeNot apDescriptionNot apIntervalNot apObserved afterHorseChallengeHorseChallengeHorseChallengeHorseChallengeHorseChallengeHorseStatistic threeHorseStatistic threeHorseStatistic threeHorseStatistic threeHorseStatistic threeHorse	doses, orses, pplical pplical es were ution o e were orse di rse even nat reso	administered including 21 ble ble e observed at of any observed no systemic ied from cause	l intramuscularly 8 foals 3 month least daily follo ed reactions. reactions observ ses affirmed by ited to transient	y approxi s of age a owing eac yed at any licensee i	and 52 foa the vaccina of the sinot attribute	als 5 mont ation, until tes. Two fe uted to vac	hs of age
AdministrationStudy Animals880 hmStudy Animals880 hmChallengeNot apDescriptionIntervalIntervalNot apobserved afterHorsechallengeThereResultsHorseresoluThereone hmAdversite th	pplical pplical pplical es were ution o e were orse di rse even nat reso	including 21 ble ble e observed at f any observe no systemic ied from cause	8 foals 3 month least daily follo ed reactions. reactions observ ses affirmed by ited to transient	s of age a owing eac yed at any licensee	and 52 foa the vaccina of the sinot attribute	als 5 mont ation, until tes. Two fe uted to vac	hs of age
Study Animals880 hChallengeNot ajDescriptionIntervalIntervalNot ajobserved afterIntervalchallengeHorseResultsHorseresoluThereone heAdversite th	pplical pplical es were ation o e were orse di rse even nat reso	ble ble e observed at of any observe no systemic ied from cause ents were lim	least daily follo ed reactions. reactions observ ses affirmed by ited to transient	owing eac yed at any licensee	h vaccina of the si not attribu	ation, until tes. Two f uted to vac	oals and ecination.
Challenge DescriptionNot applicationInterval observed after challengeNot applicationResultsHorse resoluThere one how site the	pplical pplical es were ation o e were orse di rse even nat reso	ble ble e observed at of any observe no systemic ied from cause ents were lim	least daily follo ed reactions. reactions observ ses affirmed by ited to transient	owing eac yed at any licensee	h vaccina of the si not attribu	ation, until tes. Two f uted to vac	oals and ecination.
DescriptionIntervalNot apolyticalobserved afterIntervalchallengeHorseResultsHorseThereone hopeAdversite th	pplical pplical es were ation o e were orse di rse even nat reso	ble e observed at of any observe no systemic ied from cause ents were lim	ed reactions. reactions observ ses affirmed by ited to transient	ved at any licensee 1	of the si tot attribu	tes. Two fo uted to vac	oals and cination.
IntervalNot applicationobserved afterNot applicationchallengeHorseResultsHorseresoluThereone hoAdversite th	es were ution o e were orse di rse eve nat reso	e observed at f any observe no systemic ied from cause ents were lim	ed reactions. reactions observ ses affirmed by ited to transient	ved at any licensee 1	of the si tot attribu	tes. Two fo uted to vac	oals and cination.
bserved after challenge Results Horse resolu There one ho Adver site th	es were ution o e were orse di rse eve nat reso	e observed at f any observe no systemic ied from cause ents were lim	ed reactions. reactions observ ses affirmed by ited to transient	ved at any licensee 1	of the si tot attribu	tes. Two fo uted to vac	oals and cination.
challenge     Horse       Results     Horse       resolut     There       one hor     Adver       site th	ution o e were orse di rse eve nat reso	of any observe no systemic ied from cause ents were lim	ed reactions. reactions observ ses affirmed by ited to transient	ved at any licensee 1	of the si tot attribu	tes. Two fo uted to vac	oals and cination.
Results Horse resolu There one ho Adven site th	ution o e were orse di rse eve nat reso	of any observe no systemic ied from cause ents were lim	ed reactions. reactions observ ses affirmed by ited to transient	ved at any licensee 1	of the si tot attribu	tes. Two fo uted to vac	oals and cination.
resolu There one he Adver site th	ution o e were orse di rse eve nat reso	of any observe no systemic ied from cause ents were lim	ed reactions. reactions observ ses affirmed by ited to transient	ved at any licensee 1	of the si tot attribu	tes. Two fo uted to vac	oals and cination.
There one he Adver site th	e were orse di rse eve nat reso	no systemic ied from caus ents were lim	reactions observ ses affirmed by ited to transient	licensee	not attribu	uted to vac	cination.
one he Adver site th	orse di rse eve nat reso	ied from cause ents were lim	ses affirmed by ited to transient	licensee	not attribu	uted to vac	cination.
Adver site th	rse eve nat reso	ents were lim	ited to transient				
site th	hat reso			, non-pai	nful swel	lings at the	e injection
site th	hat reso			, p			•
Local	iniaat						
	Inneci	ion site react	ions are summa	rized bel	ow across	s the four s	sites.
	injeet				inates		
		T-4-1	Namel an Of		ransient	Numt Nor	oer Of
		Total Number	Number Of Vaccinates	Injecti	on Site		inates
Si	ite	Of	Administered		lling	vacci	mates
		Vaccinates	2 doses	After	After	After 1st	After
				1 <sup>st</sup>	2 <sup>nd</sup>	dose	2 <sup>nd</sup> dose
	orth			dose	dose		
	kota	378	378	4	0	374	378
	fornia	43	43	4	3	39	40
Miss	souri	292	290	0	0	292	290
Те	exas	170	169	6	1	164	168
То	otal	883	880	14 (1.6%)	4 (0.5%)	869 (98.4%)	876 (99.5%)

Summary	Number Of Vaccinates	Number Of Vaccinates		tes With t Injection welling	Number Of Normal Vaccinates	
Age		Administered 2 doses	After 1 <sup>st</sup> dose	After 2 <sup>nd</sup> dose	After 1 <sup>st</sup> dose	After 2 <sup>nd</sup> dose
2-4 months	179	179	0	0	179	179
5-7 months	0	0	n/a	n/a	n/a	n/a
8-11 months	0	0	n/a	n/a	n/a	n/a
1-5 years*	121	121	2	0	119	121
6-15 years*	78	78	2	0	76	78
>16 years	0	0	n/a	n/a	n/a	n/a
Total	378	378	4	0	374	378

\*Swellings were 3cm in size observed 1-3 days post vaccination that resolved within 3 days.

#### California Site:

Summary	Number Of Vaccinates	Number Of Vaccinates Administered	Transien	ites With t Injection welling		Of Normal inates
Age		2 doses	After 1 <sup>st</sup> dose	After 2 <sup>nd</sup> dose	After 1 <sup>st</sup> dose	After 2 <sup>nd</sup> dose
2-4 months*	7	7	0	2	7	5
5-7 months**	1	1	1	0	0	1
8-11 months	0	0	n/a	n/a	n/a	n/a
1-5 years***	19	19	2	0	17	19
6-15 years****	15	15	1	1	14	14
>16 years	1	1	0	0	1	1
Total	43	43	4	3	39	40

\*Swellings were 3cm in size observed within hours post vaccination that resolved within several hours. \*\*Swelling was 3cm in size observed immediately post vaccination that resolved within several hours. \*\*\*1 horse had a swelling 1cm in size observed immediately post vaccination that resolved within several hours. 1 horse had a swelling observed on day 1 that increased in size to 9cm on day 3 post vaccination and resolved by day 5.

\*\*\*\*Same horse had a swelling after each vaccination that resolved within 3 weeks. Size after the first vaccination was 24cm. Size after the second vaccination was 10cm.

Missouri Site: Summary	Number Of Number Of		Vaccinates With Transient Injection Site Swelling		Number Of Normal Vaccinates	
Age	Vaccinates	Administered 2 doses	After 1 <sup>st</sup> After 2 <sup>nd</sup> dose dose	After 1 <sup>st</sup> dose	After 2 <sup>nd</sup> dose	
2-4 months	33	32	0	0	33	32
5-7 months	0	0	n/a	n/a	n/a	n/a
8-11 months	0	0	n/a	n/a	n/a	n/a
1-5 years	225	224	0	0	225	224
6-15 years	32	32	0	0	32	32
>16 years	2	2	0	0	2	2
Total	292	290	0	0	292	290

## Texas Site:

Summary	Number Of Vaccinates	Number Of Vaccinates Administered	Transien	ites With t Injection welling	n Number Of Normal Vaccinates	
Age		2 doses	After 1 <sup>st</sup> dose	After 2 <sup>nd</sup> dose	After 1 <sup>st</sup> dose	After 2 <sup>nd</sup> dose
2-4 months	0	0	n/a	n/a	n/a	n/a
5-7 months	52	51	1	1	51	50
8-11 months	0	0	n/a	n/a	n/a	n/a
1-5 years	114	114	5	0	109	114
6-15 years	0	0	n/a	n/a	n/a	n/a
>16 years	4	4	0	0	4	4
Total	170	169	6*	1**	164	168
		bserved 4-7 days day post vaccinat				n 6 days.

USDA	November 1, 2010
<b>Approval Date</b>	