

## Summary of Studies Supporting USDA Product Licensure

Establishment Name	Boehringer Ingelheim Animal Health USA Inc.
USDA Vet Biologics Establishment Number	124
Product Code	4865.A1
True Name	Encephalomyelitis Vaccine, Eastern & Western, Killed Virus, Tetanus Toxoid
Tradename(s) / Distributor or Subsidiary (if different from manufacturer)	Vetera EWT - No distributor specified
Date of Compilation Summary	October 28, 2020

## 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					
<b>Product Administration</b>	One dose, administered intramuscularly					
Study Animals	10 guinea pigs (10 vaccinates) Not applicable					
Challenge Description	Not applicable					
Interval observed after	Not applicable					
challenge						
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				
<b>Product Administration</b>	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	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	Study 2013-PM-1009						
	North Dako	ota Site:						
	Group	Vaccin	nated	Cont Preg	firmed nant	Foals		Parturition Rate
	1 <sup>st</sup> trimester product	/ 143		127		114	1	90%
	1st trimester	r/ 59		54		49		91%
	2 <sup>nd</sup> trimester	r/ 6		6		6		100%
	3 <sup>rd</sup> trimester	./ 140		117		117		100%
	Total –	348		304		286		94%
	Total –	289		250		237		95%
	Total –	1y 59		54		49		91%
	placebo on	ly						
	Study 2013 Misssouri S	-PM-1009 /ite:						
	Group	Vaccina	ated	Confi Pregr	rmed ant	Foals	Pa R	arturition ate
	2011 3 <sup>rd</sup>	5		5		5	10	00%
	2012 1 <sup>st</sup>	1		1		1	10	00%
	2012 2 <sup>nd</sup>	53		43		39	91	%
	2012 3 <sup>rd</sup>	26		26		25	96	5%
	Total –	85		75		70	93	%
	Study 2014 North Dako	-PM-1009 ta Site:	1					
	Group	Vaccinated	Conf Pregi	irmed nant	Foaled	l Parturi Rate	ition	Foals Survived to End of Observation Period
	2 <sup>nd</sup> trimester	52	52		52	100%		51*
	3 <sup>rd</sup> trimester vaccinated	69	69		67**	97.1%		67
	*Lost foal aff **One mare cooperator.	firmed by study died due to cau	y coope ises oth	erator to er than	be due vaccina	to causes o tion, as affi	ther t rmed	han vaccination. by study
LISDA Annuaral	All other foal	$\frac{15 \text{ were normal}}{12 2014}$	and he	aitny				
USDA Approval	Date September	12, 2014						

Study Type	Safety	Safety					
Pertaining to	All fractions	All fractions					
Study Purpose	To demonst	To demonstrate safety under field conditions					
Product	Two doses,	wo doses, administered intramuscularly approximately 3 – 4 weeks apart					
Administration							
Study Animals	880 horses,	including 21	8 foals 3 month	s of age a	and 52 foa	als 5 mont	hs of age
Challenge	Not application	ble					
Description							
Interval	Not application	ble					
observed after							
challenge							
Results	Horses were	e observed at	least daily follo	owing eac	h vaccina	ation, until	
	resolution o	f any observe	ed reactions.				
	There were	no systemic	reactions observ	ved at any	of the si	tes. Two f	oals and
	one horse di	one horse died from causes affirmed by licensee not attributed to vaccination.					
	Adverse eve	Adverse events were limited to transient non-painful swellings at the injection					
	site that resolved without treatment						
	Local inject	ion site react	ions are summa	rized belo	ow across	s the four s	sites:
				Vacci With Tr	nates	Numb	oer Of
		Total	Number Of	Injecti	on Site	Nor	mal
	Site	Number	Vaccinates	Swe	lling	Vacci	inates
		Of	Administered	After	After	A.C. 1 St	A.C.
		vaccinates	2 doses	1 <sup>st</sup>	$2^{nd}$	After 1 <sup>st</sup>	After $2^{nd}$ doso
				dose	dose	uose	2 0080
	North Dakota	378	378	4	0	374	378
	California	43	43	4	3	39	40
	Missouri	292	290	0	0	292	290
	Texas	170	169	6	1	164	168
	Total	883	880	14 (1.6%)	4 (0.5%)	869 (98.4%)	876 (99.5%)
	Results from	n each site ar	e summarized o	on the foll	owing pa	iges.	

Summary	Number Of	Number Of Vaccinates	Vaccina Transient Site Sv	tes With t Injection welling	Number Vacc	Of Normal inates
Age	vaccinates	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	Vaccinates With Transient Injection Site Swelling		Number Vacc	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	hary Number Of Number Of Vaccinates With Number Of Vaccinates Site Swelling		Number Of Norma Vaccinates			
Age	vaccinates	2 doses	After 1 <sup>st</sup>	After 2 <sup>nd</sup>	After 1st	After 2 <sup>nd</sup>
Age		2 00505	dose	dose	dose	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	Number Of Vaccinates	Vaccinates With Transient Injection Site Swelling		Number Vacc	Of Normal inates
Age	vaccinates	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
*Swellings were **Swelling was	e <1.5cm were ol 5cm observed 1	bserved 4-7 days day post vaccinat	post vaccination that res	ation and res	olved within 2 days.	n 6 days.

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