

# **Summary of Studies Supporting USDA Product Licensure**

Establishment Name	Boehringer Ingelheim Animal Health USA Inc.
USDA Vet Biologics Establishment Number	124
Product Code	48W5.21
True Name	Encephalomyelitis-West Nile Virus Vaccine, Eastern & Western, Killed Virus, Tetanus Toxoid
Tradename(s) / Distributor or Subsidiary (if different from manufacturer)	Equi-Jec WNV+EWT - No distributor specified  Vetera EWT + WNV - Boehringer Ingelheim (Canada) Ltd.  Vetera EWT + WNV - 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.

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Study Type	Efficacy
Pertaining to	Clostridium tetanus
<b>Study Purpose</b>	Demonstration of efficacy against Clostridium tetanus
<b>Product Administration</b>	One dose, administered intramuscularly
Study Animals	10 guinea pigs (10 vaccinates)
<b>Challenge Description</b>	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.
<b>USDA Approval Date</b>	April 18, 2008

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Study Type	Efficacy
Pertaining to	Eastern equine encephalomyelitis
Study Purpose	Demonstration of efficacy against Eastern equine
	encephalomyelitis
<b>Product Administration</b>	Two doses, administered intramuscularly, 14 to 21 days apart
Study Animals	12 guinea pigs (10 vaccinates, 2 controls)
<b>Challenge Description</b>	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 Eastern equine encephalomyelitis per the criteria in 9 CFR 113.207(b) and the requirements were met.
<b>USDA Approval Date</b>	April 18, 2008

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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 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.
<b>USDA Approval Date</b>	April 18, 2008

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Study Type	Efficacy		
Pertaining to	West Nile Virus (WNV)		
Study Purpose	Demonstration of twelve me	onth duration of	immunity against disease
	caused by WNV		
<b>Product Administration</b>	Two doses, administered intr	amuscularly, 25 da	ays apart
Study Animals	30 horses (20 vaccinates, 10	placebo controls) 4	4-5 months of age
<b>Challenge Description</b>	West Nile Virus was admin	istered at 380 day	ys (10 vaccinated and 5
	placebo control animals) or	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 neurological disease, as me evidence of virus-induced  Animals were also monitor the blood).  Results are summarized as	easured by morta brain disease (his red for viremia (c	lity and microscopic stopathology).
	Outcome	Controls	Vaccinates
	Mortality	7/10 (70%)	1/20 (5%)
	Viremia at least one day	10/10 (100%)	2/20 (10%)
	See raw data on following	pages.	
<b>USDA Approval Date</b>	September 3, 2010		

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Treatment	#	Died or Euthanized due	Severity Histopat	hological lesions
1 reatment	#	to disease severity	Medulla	Pons
	1	Yes	3	3
	2	Yes	3	3
	3	Yes	3	3
	4	Yes	3	3
<b>Controls</b>	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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M AM PM 7 8 9 10 D D D D D D D D D D D D D D D D D D D	Viremia:								Da	ys Post	Days Post-challenge	ge								
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	Treatment	•	Ψ	Н	Н	Н	Н	PM	Н	PM	AM	PM	ΑM	PM	,	•	,	Π	ŧ	7.
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	Controls	v		4	$\vdash$	$\vdash$	⊢	20	2	r								Δ	A	h
	(10 horses)	9	L	L	$\vdash$	$\vdash$	$\vdash$	55	10										А	<u>P</u>
		7	L	F	$\vdash$	$\vdash$	⊢	225	13	121									z	h
		œ	L	H	55	⊬	33	105	20	2										L
		6	L	<u></u>	$\vdash$	$\vdash$	25	135	240	9	20									
		10		L		8	6	40	2	r										
2   4		ī																	А	h
3   4   6   6   6   6   6   6   6   6   6		2																		
Vaccinates         4         40         1 <t< td=""><td></td><td>3</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></t<>		3																		
S   95   95   95   95   95   95   95		4																		
Vaccinates (20 horses)         11         40         9           (20 horses)         11         12         13           13         16         17         18         19           Actual value in plaque-forming units per milliliter equivalents (PEUeg/mL) = Positive for virus isolation (<5 PEUeg/mL)		S																		
7		9			95															
Vaccinates         10         40           (20 horses)         11         12           13         14         15           16         17         18           19         19         10           Actual value in plaque-forming units per milliliter equivalents (PEUcq/mL) = Positive for virus isolation (<5 PEUcq/mL)		7																		
Vaccinates         9           (20 horses)         11           12         13           14         15           16         17           18         19           Actual value in plaque-forming units per milliliter equivalents (PEUcg/mL) = Positive for virus isolation         ≤ DEUcg/mL)           D = Dead         N = Not recorded; horse was circling with sporadic head / neck tremors.		<b>20</b>		-	40															
Vaccinates         10           (20 horses)         11           12         13           13         14           16         10           17         18           18         19           Actual value in plaque-forming units per milliliter equivalents (PFUeg/mL) = Positive for virus isolation (<5 PFUeg/mL)		٥ )	4	$\dashv$	+	$\perp$														
11   12   13   14   15   16   17   17   18   19   19   19   19   19   19   19	Vaccinates	01	$\perp$	$\dashv$	_															
12	(20 horses)	11																		
13		12		-																
14   15   16   17   18   19   19   19   19   19   19   19		I3		-																
15   16   17   18   19   19   19   19   19   19   19		14	4	$\dashv$	$\downarrow$	$\downarrow$			1	1										
Actual value in plaque-forming units per milliliter equivalents (PFUeg/mL) = Positive for virus isolation (<5 PEUeg/mL)  D = Dead  N = Not recorded; horse was circling with sporadic head / neck tremors.		IS	4	+	+	4			1	1										
Actual value in plaque-forming units per milliliter equivalents (PEUeg/mL) = Positive for virus isolation  Blank = Negative for virus isolation (<5 PEUeg/mL)  D = Dead  N = Not recorded; horse was circling with sporadic head / neck tremors.		0 1	4	+	+	4			1	1										
Actual value in plaque-forming units per milliliter equivalents (PFUeg/mL) = Positive for virus isolation  Blank = Negative for virus isolation (<5 PFUeg/mL)  D = Dead  N = Not recorded; horse was circling with sporadic head / neck tremors.		1	_	+	+	-				1										
Actual value in plaque-forming units per milliliter equivalents (PFUeg/mL) = Positive for virus isolation  Blank = Negative for virus isolation (<5 PFUeg/mL)  D = Dead  N = Not recorded; horse was circling with sporadic head / neck tremors.		e	4	$\dashv$	4	4			1	1										
Actual value in plaque-forming units per milliliter equivalents (PFUeq/mL) = Positive for virus isolation  Blank = Negative for virus isolation (<5 PFUeg/mL)  D = Dead  N = Not recorded; horse was circling with sporadic head / neck tremors.		19	4	+	_	_				1										
Actual value in plaque-forming units per milliliter equivalents (PFUeg/mL) = Positive for virus isolation  Blank = Negative for virus isolation (<5 PFUeg/mL)  D = Dead  N = Not recorded; horse was circling with sporadic head / neck tremors.		0.7		_																
D = Dead N = Not recorded; horse was circling with sporadic head / neck tremors.	Actual value in Blank = Negati	plaque-form ive for virus is	ing un solatio	its per n (<5.]	millilite PEUeg/	er equiv: mL)	alents (E	FUeg/n	nL) = Pc	ositive i	for virus	s isolat	non							
14 - IVOLICCOLOCCI, INCISC WAS CILCIME WITH SPOTAGOL INCAR ULTIMOTS.	D = Dead N = Not record	ed: horse me	i oiroli	to the	h enors/	fir head	t doest	remore												
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Study Type	Efficacy
Pertaining to	West Nile Virus
Study Purpose	Demonstration of efficacy against WNV
<b>Product Administration</b>	Two doses, administered intramuscularly 21 days apart
Study Animals	28 horses (19 vaccinates, 9 placebo controls) 4-5 months of age
Challenge Description	West Nile Virus was administered intrathecally at 14 days (to 10
	vaccinated and 5 placebo control animals) and 28 days (to 9
	vaccinated and 4 placebo control animals) after the second
	vaccination
Interval observed after	Horses were bled on the day of challenge, twice daily for 6 days
challenge	post-challenge, once daily for an additional 4 days post-challenge,
	and on day 14 post-challenge
Results	The primary outcome was viremia (detection of WNV in the
	blood). While the test method was quantitative, an animal was
	considered to be positive (affected by challenge) if any virus was
	detected in the blood on one or more occasions post-challenge.
	The number of animals positive for (affected by) viremia at least
	once is summarized as follows:
	Controls Vaccinates
	8/9 (89%)   1/19 (5%)
	See raw data on the following page.
<b>USDA Approval Date</b>	August 25, 2008

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hallenge	5 6 7 8	PM AM PM AM PM / 8 9 10 14	20	20 65 45	15	80 15		70	D	15 10   10	10 5																		
Days Post-Challenge	4	AM		5	120			120		20	15																		
Day	3	PM			495	235		70		15																			
		AM	135		355	140		110		30	20																		
		PM	280	40	645	235		675		09	15																		
	2	AM	390		1475	85	165	110		5	10																		
		PM	20	5	125	20																							
		AM	15																										
		>																											
	Horse	П	13	19	20	45	29	71	72	74	79	14	16	21	22	23	24	25	56	27	37	73	75	77	80	81	82	83	70
Viremia:		l reatment					Controls	(Sagnon 6)											I	77.00.00	Vaccinates (10 benegation)								

Actual value in plaque-forming units per milliliter equivalents (PFUeq/mL) = Positive for virus isolation Blank = Negative for virus isolation (<5 PFUeq/mL)

D = Dead (euthanized on Day 11 due to West Nile Virus)

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Study Type	Efficacy			
Pertaining to	West Nile Virus	(WNV)		
Study Purpose	Demonstration of	of six month durat	ion of immuni	ty against WNV
<b>Product Administration</b>	Two doses, adm	inistered intramus	cularly 21 day	ys apart
Study Animals		ccinates, 10 place		
Challenge Description		est Nile Virus was		
				o control animals)
		allenge Group 2:		and 5 placebo
		after second vacc		
Interval observed after		d on the day of ch		
challenge	-	<u>-</u>	additional 4 da	ys post-challenge,
	and on day 14 p			
Results		come was viremia		
		ne test method wa	•	
		positive (affected	•	•
	detected in the b	lood on one or mo	ore occasions	post-challenge.
		nimals positive fo		east once
	_ <u> </u>	marized as follow		_
	Challeng	e Controls	Vaccinates	
	Group	5/5 (1000/)	2/10 (200/)	-
	2	5/5 (100%) 5/5 (100%)	2/10 (20%) 4/10 (40%)	-
	Combine	` '	6/20 (30%)	-
	Comone	1 10/10 (100/0)	0/20 (30/0)	_
	See raw data on	the following pag	e	
	See Taw Gata Off	and following pag	<b>.</b> .	
USDA Approval Date	October 21, 200	9		

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	22			425	20	30	40	85	09	65							Z		Positive
	810			10	300	125	125	80	45									Pe	Positive
	S11				50	30	40	40	25									Pe	Positive
	S13				410	110	135	110	55	15							Z		Positive
$\vdash$	S1														Г	$\vdash$	H	ž	Negative
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	6S																Z		Negative
<u> </u>	S12																	ž	Negative
	S14																	Ne	Negative
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	S53		70	320	380	100	135	45	10									P	Positive
	S54							5		5	5	5					Z		Positive
Challenge 2	S55	10	5	95	70	30	25	40										Pe	Positive
	S29			90	265	20	70	45	45	5								Pe	Positive
H	S46															$\vdash$	_	Ž 	Negative
	S47							5											Positive
	S48																	Ž	Negative
	S49			15														Pc	Positive
vaccinates (10 benge)	S51																	Ž —	Negative
Chellenge 2	S52																	ž	Negative
<b>」</b>	92S																	Ne Ne	Negative
	S57			2	5													P(	Positive
	S58		5															Pe	Positive
L	0,0																		Nogotivo

Actual value in plaque-forming units per milliliter equivalents (PFU eq/mL) = Positive for virus isolation Blank = Negative for virus isolation (<5 PFU eq/mL) N = Not recorded Positive = affected by challenge if virus was detected in the blood on one or more occasions post-challenge. Negative = virus was detected in the blood on zero occasions post-challenge.

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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.
<b>Study Animals</b>	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

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# Results

## Study 2013-PM-1009

North Dakota Site:

Group	Group Vaccinated Confirmed Foals		Foals	Parturition
		Pregnant		Rate
1 <sup>st</sup> trimester/ product	143	127	114	90%
1st trimester/ placebo	59	54	49	91%
2 <sup>nd</sup> trimester/ product	6	6	6	100%
3 <sup>rd</sup> trimester/ product	140	117	117	100%
Total – all animals	348	304	286	94%
Total – product only	289	250	237	95%
Total – placebo only	59	54	49	91%

## Study 2013-PM-1009

#### Misssouri Site:

Group	Vaccinated	Confirmed Pregnant	Foals	Parturition Rate
2011 3 <sup>rd</sup>	5	5	5	100%
trimester				
2012 1st	1	1	1	100%
trimester				
2012 2 <sup>nd</sup>	53	43	39	91%
trimester				
2012 3 <sup>rd</sup>	26	26	25	96%
trimester				
Total –	85	75	70	93%
product				

# Study 2014-PM-1009

## North Dakota Site:

Group	Vaccinated	Confirmed Pregnant	Foaled	Parturition Rate	Foals Survived to End of Observation Period
2 <sup>nd</sup> trimester vaccinated	52	52	52	100%	51*
3 <sup>rd</sup> trimester vaccinated	69	69	67**	97.1%	67

<sup>\*</sup>Lost foal affirmed by study cooperator to be due to causes other than vaccination.

All other foals were normal and healthy

**USDA Approval Date** 

September 12, 2014

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<sup>\*\*</sup>One mare died due to causes other than vaccination, as affirmed by study cooperator.

Study Type	Safety							
Pertaining to	All fractions							
Study Purpose	To demonstra	ate safety u	nder field cond	itions				
Product	Two doses, a	Two doses, administered intramuscularly 3 – 4 weeks apart						
Administration								
<b>Study Animals</b>	556 horses, i	ncluding 43	88 foals betwee	n 2 month	s and app	roximatel	y 1 year	
	of age							
Challenge	Not applicab	le						
Description								
Interval	Not applicab	le						
observed after								
challenge								
Results		any observ	t least daily followed reactions. O	_				
		•	reactions observable licensee not as		•		norse died	
	Scoring Method for Injection Site Reactions:  0 = No reaction  1 = Localized swelling at or near the injection site which is not visible; detectable only by palpation. Not painful.  2 = Localized visible swelling at or near the injection site. Not painful.  3 = Localized visible swelling at or near the injection site. Raised, circumscribed and painful when palpated.							
	Site	Total Number Of	Number Of Vaccinates Administere	Vaccina Tran Injecti	tes With asient on Site lling	Numl Nor	per Of rmal inates	
		Vaccinat es	d 2 doses	After	After	After	After	
				1 <sup>st</sup> dose	2 <sup>nd</sup> dose	1 <sup>st</sup> dose	2 <sup>nd</sup> dose	
	Missouri	315	314	(1.0%)	9 (2.9%)	312 (99.0%)	305 (97.1%)	
				1	2	109	108	
	Oklahoma	110	110	(0.9%)	(1.8%)	(99.1%)	(98.2%)	
	Texas	131	131	0 (0%)	0 (0%)	131 (100%)	131 (100%)	
	Total	556	555	4 (0.7%)	11 (2.0%)	552 (99.3%)	544 (98.0%)	
	Results from	each site an	re summarized	on the fol	lowing pa	ige.		

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Summary	Number Of Vaccinates	Number Of Vaccinates Administered	Transien	ites With t Injection welling		Of Normal inates
Age	vaccinates	2 doses	After 1st dose	After 2 <sup>nd</sup> dose	After 1st dose	After 2 <sup>nd</sup> dose
2-4 months	55	55	0	0	55	55
5-7 months	8	8	0	0	8	8
8-11 months	1	1	0	0	1	1
1 year	170	170	1	2	169	168
≥ 2 years	81	80	2	7	79	73
Total	315	314	3	9	312	305

Horse No.	Age	Reaction Description	Injection#	Day	Score	Resolution Day
10	11 y	Swelling on day 3, 5.5 cm x 2.25 cm x 5mm	2	3	2	7
22	8 y	Swelling on day 3, 12 cm circle, raised 1.5 cm, painful,	2	3	3	7
129	1 y	Swelling on day 3, 2.3 cm circle, raised 4 mm, painful but no heat	2	3	3	7
183	1 y	Swelling on day 7, raised lesion 1.5 cm circle, height 0.2 cm	1	7	2	14
183	1 y	Swelling on day 1, 3 cm lesion, not raised but palpalble	2	1	1	3
222	9 y	Swelling on day 3, 6 cm x 7 cm x 1.2 cm, raised lesion hard and painful	2	3	3	7
266	10 y	Swelling localized in several places unsure if related to vaccine	1	1	2	3
266	10 y	Swelling small palpapable mass ~ 2 cm size, still present day 3 no worse	2	1	2	3
271	13 y	Swelling 5cm circle, raised 5 mm, solid and painful	2	3	3	7
288	8 y	Swelling < 2 cm, raised lesion ~ 1 mm deep	1	3	2	7
288	8 y	Swelling ~ 8.5 cm circle raised ~ 1.3 cm, painful, not hot to touch	2	3	3	7
300	10 y	Swelling 6 cm circle, solid swelling not painful	2	3	2	7

## Oklahoma Site:

Summary	Number Of Vaccinates	Number Of Vaccinates Administered	Transien	ites With t Injection welling		Of Normal inates
Age	vaccinates	2 doses	After 1st dose	After 2 <sup>nd</sup> dose	After 1st dose	After 2 <sup>nd</sup> dose
4-6 months	49	49	1	1	48	48
1 year	25	25	0	1	25	24
≥ 2 years	36	36	0	0	36	36
Total	110	110	1	2	109	108

Horse No. 19-A	Age 4 m	Reaction Description Swelling redness painful injection area 6 cm in diameter, reaction subsided in	Injection#	Day	Score	Resolution Day
		10 days	1	7	3	17
33-A	5 m	Small swelling, 3 cm diameter, subsided in 3 days	2	1	2	4
43	1 y	Mid-sized swelling, 5 cm diameter, reduced to 2.5 cm in 6 days; small, hard 2 cm at 10 days, probable subcutaneous leakage	2	1	2	Study End

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Texas Site:  Summary	Number Of Vaccinates	Number Of Vaccinates Administered	Tra Inject	ates With nsient ion Site elling		Of Normal inates
Age		2 doses	After 1st dose	After 2 <sup>nd</sup> dose	After 1st dose	After 2 <sup>nd</sup> dose
7-9 months	130	130	0	0	130	130
≥ 2 years	1	1	0	0	1	1
Total	131	131	0	0	131	131

USDA Approval	September 14, 2009
Date	

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