

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

Establishment Name	Intervet Inc.
USDA Vet Biologics Establishment Number	165A
Product Code	44B1.20
True Name	Bovine Rhinotracheitis-Virus Diarrhea-Parainfluenza 3- Respiratory Syncytial Virus Vaccine, Modified Live Virus, Campylobacter Fetus-Leptospira Canicola-Grippotyphosa- Hardjo-Icterohaemorrhagiae-Pomona Bacterin
Tradename(s) / Distributor or Subsidiary (if different from manufacturer)	Bovilis Vista 5 VL5 SQ - Merck Animal Health Bovilis Vista 5 VL5 SQ - No distributor specified Vista 5 L5 SQ - Merck Animal Health Vista 5 VL5 SQ - Merck Sharp & Dohme Saude Animal Ltda Merck Animal Health Vista 5 VL5 SQ - No distributor specified
Date of Compilation Summary	December 15, 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.

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Study Type	Efficacy
Pertaining to	Bovine Viral Diarrhea Virus Type 1 (BVDV1)
Study Purpose	To demonstrate efficacy against respiratory disease caused by
	BVDV1.
<b>Product Administration</b>	
Study Animals	Bovine
<b>Challenge Description</b>	BVDV Type 1b NY-1 strain
Interval observed after	
challenge	
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.
<b>USDA Approval Date</b>	February 20, 2004

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Study Type	Efficacy
Pertaining to	Bovine Viral Diarrhea Virus Type 1 (BVDV1)
Study Purpose	To demonstrate efficacy against persistent infection of calves
	caused by BVDV1.
Product Administration	
Study Animals	Bovine
Challenge Description	BVDV Type 1b, strain SD02 BVD09
Interval observed after	
challenge	
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.
<b>USDA Approval Date</b>	June 23, 2005

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Study Type	Efficacy
Pertaining to	Bovine Viral Diarrhea Virus Type 1 (BVDV1)
Study Purpose	To demonstrate efficacy against fetal infection caused by BVDV1
	206 days after vaccination.
<b>Product Administration</b>	
Study Animals	Bovine
<b>Challenge Description</b>	BVDV Type 1b strain SD02 BVD09
Interval observed after	
challenge	
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.
<b>USDA Approval Date</b>	October 6, 2005

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Study Type	Efficacy
Pertaining to	Bovine Viral Diarrhea Virus Type 1 (BVDV1)
Study Purpose	To demonstrate efficacy against respiratory disease caused by
	BVDV1 206 days after vaccination.
<b>Product Administration</b>	
Study Animals	Bovine
<b>Challenge Description</b>	
Interval observed after	
challenge	
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.
<b>USDA Approval Date</b>	October 6, 2005

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Study Type	Efficacy
Pertaining to	Bovine Viral Diarrhea Virus Type 2 (BVDV2)
Study Purpose	To demonstrate efficacy against respiratory disease caused by
	BVDV2.
Product Administration	
Study Animals	Bovine
<b>Challenge Description</b>	BVDV2a strain 1373
Interval observed after	
challenge	
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.
<b>USDA Approval Date</b>	December 10, 2003

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Study Type	Efficacy						
Pertaining to	Bovine Viral Diarrhea Virus Type 1 (BVDV2)						
Study Purpose	To demonstrate efficacy against persistent infection of calves						
	caused by BVDV2						
<b>Product Administration</b>							
Study Animals	Bovine						
<b>Challenge Description</b>	BVDV Type 2 strain SD02 BVD05						
Interval observed after							
challenge							
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.						
<b>USDA Approval Date</b>	April 25, 2005						

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Study Type	Eff	Efficacy							
Pertaining to	Boy	Bovine Viral Diarrhea Virus Type 2 (BVDV2)							
Study Purpose	To	demonstrate e	fficacy against fe	etal infection ca	aused by				
		BVDV2 206 days after vaccination.							
Product Administration	1 d	1 dose administered by the subcutaneous route 28 days prior to							
	bre	eding.							
Study Animals			eifers, 28 vaccina						
Challenge Description	All	heifers were c	hallenge with B	VDV2 strain Γ	V809-04 at 164-				
	178	days of gesta	tion.						
Interval observed after		-	ere collected on o	•					
challenge		_	s isolation. Fetu	ses were colle	cted on day 60				
		after challenge.							
Results	Vir	us Isolation or	<u> </u>						
		Group	# of Animals	# Affected	Percent (%)				
		Vaccinates	28	2	7				
		Controls	18	18	100				
			om fetal samples						
		`	/	-	rirus was isolated				
			tissue (lung, sple	een, thymus, k	idney, buffy				
		coat).	T		- (a)				
	Group # of Animals # Affected Percent (%)								
	Vaccinates 28 2 4								
	Controls 18 17 94								
	_	1 . 1	1 1						
	Rav	w data shown (	on attached pages	S.					
HCD A A LD 4	0.54	halan 4, 2007							
USDA Approval Date	Oct	tober 4, 2007							

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## Viremia of Challenged Heifers

(Vaccinate Group)

Number	0	5	6	7	8	9	10
1536	0	0	: 0	0	. 0	0	0
1538	0	0	0	0	0	0	. 0
1541	0 -	- 0	.0	0	0	0	. 0
1544	0	0	- 1	0	0	0	. 0
1545	0 1	- 0	0	0-	0	0	. 0
1556	0	0 -	0	0	0	. 0	. 0
1558	0	0 -	0	0	0	0	0
1562	. 0	- 0	0	0	0	0 0	0
1566	0	0	0	0	0	0 :	0
1567	0	. 0	0	0	0	0	0
1569	0	0	0	0	0	0 1	0
1570	0	0 .	0	0	0	0	- 0
1575	0	0	0	0	0	0	. 0
1581	0	0	0	0	0	0	. 0
1582	0	0 -	0	0	0	0	. 0
1585	0	0	1	0	0 0	0	0.0
1594	0	0	0	0	. 0	0 '	0
1596	0	0	0	0	0	. 0	. 0
1597	0	0	0	0	0	. 0	- 0
1598	0	0	.0	0	0	0 '	0
1599	0 .	0	0	- 0	0	. 0	0
1601	0	0	0	0	0	0	. 0
1605	. 0	0	0 -	0	0	. 0	0
1606	0 -	. 0	0	0	. 0	0	0
1607	0	. 0	0	0	0	0 .	0
1608	0	0	0	0	. 0	0	0
1609	0	0	0	0 '	0	0	0
1614	. 0	0	0	0	0	0	.0

0=negative;

1=positive.

#### (Control Group)

Number	0	5	6	7	8	9	10
1540	0	1	11	1 1	1	0	0
1542	0 '	1 - 1	1.	0	11	0	- 0
1543	0	1	1 .	1.1	1:	0	1
1546	0	0	1.	111	1 :	0	. 0
1549	0	. 1	1.	- 1	1	1 1	0
1553	. 0	1	1 :	1	1.	1	. 0
1557	0	0	1	. 1	. 1	0	0
1571	0	- 1	. 1	15	1 1	0	0
1572	0	1 -	1 1	0 1	. 1	0	0
1573	0	1 1	1 .	1	1	- 0	0
1574	0	1	1	-d -d	.0	- 1 -	0
1577	0	- 1	1.1	1.1	0	0	0
1586	0	-1"	1	1	0	0	. 0
1590	0 -	1	1 1	1	0	0	0
1591	0	1	1 1 1	1	1 1 4	- 1	. 0
1593	0	1	1	1	. 1	1	0
1595	0	1	1	1	1	1	0
1615	0	1	0	0.	0	1 1	. 0

0=negative;

1=positive.

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**Virus Isolation from Fetal Samples** 

Groups		Virus isolations					
	Heifer ID	Thymus	Spleen	Lung	Kidney	Buffy- coats	VI Results
	1536	0	0	0	. 0	0	0
	1538	0	0	0	0	0	0
	1541	0	0	0	0	0	0
- 1	1544	0 .	0	- 0	0	0	0
	1545	1 1	-1 -	1-	0	1 1	1
. [	1556	0	0	0	0	0	0
	1558	- 0	0	0	0	- 0	0
	1562	0	. 0	0	0	0	0
	1566	0	0	0	0 .	0 .	0
	1567	0	0	0	0	0	0
	1569	- 0	0	. 0	0	0 -	0
1, 1	1570	0	0	0	. 0	0	0
-	1575	0	0	. 0	0	0	0
	1581	0	0	0	0	0	0
, .	1582	- 0	0 -	0	0 -	- 0	0 -
	1585	0	0	0	0	0	0.
	1594	0	0	0	0	0	0
	1596	0	0	0	0	0 -	. 0
Vaccinate	1597	0	0	0	0 -	0	0
	1598	0	0	0 - "	0	0	0
1 1	1599	0	0	0	0	0	- 0
	1601	0	0	0	0	1 .	1 ,
	1605	0	0	0	0	0	. 0
	1606	- 0	0	0	0	0	0
	1607	0 '	0	0	0	0	0
	1608	0	. 0	0.0	0	0	0
1	1609	0 .	0 .	0	0	0 .	0
	1614	0	0	0 -	0	0	0

Groups			Virus isolations					
	Heifer	Thymus	Spleen	Lung	Kidney	Buffy- coats	VI Results	
	1540	1 1	11	11	1	1	1	
[	1542	. 1	1	1	1	0 -	1	
	1543	0	0	0	0	1	1 1	
· .	1546	0	- 0	0	0 -	1	1.	
	1549	1 5	1	. 1	1	1	1 1	
	1553	1	1 1	1	1	. 1	1.	
	1557	1 -	1	1 1	- 1	· 1	. 1	
	1571	0,	0	0 0	0	0 -	0	
	1572	0	0	0	0	1 1	. 1	
Controls	1573	0	0	0	0	1	1	
, , ,	1574	-1	1.	1	1	0 -	1	
	1577	0	0	. 0	0	1.	1	
,	1586	1	1	-11	. 1	0 -	1	
	1590	1	1 1	1	1 1	0	1	
	1591	1.	1	1	1 1	0	.1.	
1	1593	0	1 1 1	0	1 ,	1 1	1	
	1595	1	: 1	1	. 1	1 '	- , 1	
	1615	0	0	. 0	0	1	1	

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Study Type	Efficacy		
Pertaining to	Bovine Viral Diarrhea Virus Type 2 (BVDV2)		
Study Purpose	To demonstrate efficacy against respiratory disease caused by		
	BVDV2 200 days after vaccination.		
<b>Product Administration</b>			
Study Animals	Bovine		
<b>Challenge Description</b>			
Interval observed after			
challenge			
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.		
<b>USDA Approval Date</b>	June 22, 2006		

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Study Type	Efficacy
Pertaining to	Campylobacter fetus
Study Purpose	To demonstrate effectiveness against infertility caused by <i>C</i> .
	fetus
<b>Product Administration</b>	Subcutaneous
Study Animals	Bovine
<b>Challenge Description</b>	
Interval observed after	
challenge	
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.
<b>USDA Approval Date</b>	March 26, 2004

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Study Type	Efficacy
Pertaining to	Infectious Bovine Rhinotracheitis (IBR)
Study Purpose	To demonstrate effectiveness against disease caused by IBR
<b>Product Administration</b>	Subcutaneous
Study Animals	Bovine
<b>Challenge Description</b>	
Interval observed after	
challenge	
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.
USDA Approval Date	March 29, 2004

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Study Type	Efficacy		
Pertaining to	Infectious Bovine Rhinotracheitis (IBR)		
Study Purpose	To demonstrate effectiveness against abortions caused by IBR		
<b>Product Administration</b>	Subcutaneous		
Study Animals	Bovine		
<b>Challenge Description</b>			
Interval observed after			
challenge			
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.		
<b>USDA Approval Date</b>	June 27, 2005		

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Study Type	Efficacy
Pertaining to	Infectious Bovine Rhinotracheitis (IBR)
Study Purpose	To demonstrate efficacy against abortions caused by IBR at 217
	days post vaccination.
<b>Product Administration</b>	Subcutaneous
Study Animals	Bovine
<b>Challenge Description</b>	
Interval observed after	
challenge	
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.
USDA Approval Date	March 16, 2006

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Study Type	Efficacy		
Pertaining to	Infectious Bovine Rhinotracheitis (IBR)		
Study Purpose	To demonstrate efficacy against respiratory disease caused by IBR		
	182 days after vaccination.		
<b>Product Administration</b>			
Study Animals	Bovine		
<b>Challenge Description</b>			
Interval observed after			
challenge			
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.		
<b>USDA Approval Date</b>	March 30, 2004		

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Study Type	Efficacy
Pertaining to	Leptospira canicola
Study Purpose	To demonstrate effectiveness against disease caused by <i>L. canicola</i>
<b>Product Administration</b>	Subcutaneous
Study Animals	Bovine
Challenge Description	
Interval observed after	
challenge	
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.
<b>USDA Approval Date</b>	March 10, 2005

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Study Type	Efficacy		
Pertaining to	Leptospira grippotyphosa		
Study Purpose	To demonstrate effectiveness against disease caused by		
_	L. grippotyphosa		
Product Administration	Subcutaneous		
Study Animals	Bovine		
Challenge Description			
Interval observed after			
challenge			
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.		
USDA Approval Date	March 10, 2005		

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Study Type	Efficacy		
Pertaining to	Leptospira hardjo		
Study Purpose	To demonstrate effectiveness against disease caused by <i>L. hardjo</i>		
<b>Product Administration</b>	Subcutaneous		
Study Animals	Bovine		
Challenge Description			
Interval observed after			
challenge			
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.		
<b>USDA Approval Date</b>	May 24, 2004		

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Study Type	Efficacy	Efficacy			
Pertaining to	Leptospira hardjo				
Study Purpose	To demonstrate effectiveness against urinary shedding of $L$ .				
	hardjo				
<b>Product Administration</b>	1 dose administered by the subcutaneous route				
Study Animals	29 calves, > 6 m	onths of age, 14	vaccinates and	15 controls	
Challenge Description	All calves were challenged with <i>L. hardjo</i> 21 days after				
	vaccination.				
Interval observed after	All calves were	monitored daily	for 30 days for	clinical signs of	
challenge	disease. Urine samples were taken on days 16 (Week 2), 23				
	(Week 3) and 30 (Week 4) post challenge for Leptospira				
	isolation.				
Results	Leptospira Isolation: A calf was considered positive if				
	leptospires were isolated from the urine during any of the four				
	post-challenge weeks.				
	Group	# of Animals	# Positive	Percent (%)	
	Vaccinates	14	0	0	
	Controls	15	15	100	
	Raw data shown	on the attached	page.		
USDA Approval Date	March 27, 2012				

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## L. hardjo Urine Isolations

Group	Animal No.	al No. Post-challenge Week		
GIOUP.	Ahiillai Nu.	2	3	4
	507	+	-	-
	509	+	+	-
	510	+	+	+
	517	+	+	+
	518	+	+	-
	519	+	-	-
	520	-	+	-
Control	521	-	+	-
	523	+	-	-
	526	+	-	-
	527	+	+	-
	529	+	-	-
	531	-	+	-
	533	+	+	-
	540	+	+	-
	508	-	-	-
	511	-	-	
	512	-	-	-
	514	-	-	-
	516	-	-	-
	522	-	-	-
Vaccinated	524	-	-	-
vaccinated	525	-	-	•
	532	-	-	-
	536	-	-	-
	537	-	-	-
	541	=		-
	542	-	-	-
	543	-	_	-

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Study Type	Efficacy		
Pertaining to	Leptospira icterohaemorrhagiae		
Study Purpose	To demonstrate effectiveness against disease caused by $L$ .		
	icterohaemorrhagiae		
<b>Product Administration</b>	Subcutaneous		
Study Animals	Bovine		
Challenge Description			
Interval observed after			
challenge			
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.		
<b>USDA Approval Date</b>	June 22, 2004		

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Study Type	Efficacy				
Pertaining to	Leptospira pomona				
Study Purpose	To demonstrate effectiveness against disease caused by L. pomona				
<b>Product Administration</b>	Subcutaneous				
Study Animals	Bovine				
Challenge Description					
Interval observed after					
challenge					
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.				
<b>USDA Approval Date</b>	May 24, 2004				

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Study Type	Efficacy				
Pertaining to	Parainfluenza <sub>3</sub> Virus (PI3)				
Study Purpose	To demonstrate effectiveness against shedding caused by PI3				
<b>Product Administration</b>	Subcutaneous				
Study Animals	Bovine				
<b>Challenge Description</b>					
Interval observed after					
challenge					
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.				
<b>USDA Approval Date</b>	April 12, 2005				

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Study Type	Efficacy				
Pertaining to	Bovine Respiratory Syncytial Virus (BRSV)				
Study Purpose	To demonstrate effectiveness against BRSV				
<b>Product Administration</b>	Subcutaneous				
Study Animals	Bovine				
Challenge Description					
Interval observed after					
challenge					
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.				
<b>USDA Approval Date</b>	June 3, 2004				

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Study Type	Safety							
Pertaining to	NA NA							
Study Purpose	Demonstrate safety in pregnant cows or in calves nursing pregnant cows if							
	previously vaccinated pre-breeding with same product							
Product	One dose of test product was administered subcutaneously 30-60 days before							
Administration	breeding to all study animals. Then the test or control product was given during the							
	targeted trimester of gestation.							
	• <u>Test product:</u> Contains modified live virus fractions • Control product: Product of similar composition except virus fraction(s) are							
	<u>Control product:</u> Product of similar composition except virus fraction(s) are killed							
Study Animals	Pregnant cows: separate groups vaccinated at each trimester of gestation. Similar							
<b>,</b>	sized groups in each trimester were maintained as controls.							
Challenge	NA							
Description								
Interval observed		nge. After vacci		•	_		s. Calves	
after challenge	monitored	for 48 hours (bu	ıll calves) o	r 4 weeks	(heiter calv	ves).		
Results								
			<u> </u>	Removed	gnant Cows	I I		
		Vaccinated During		from	Unrelated			
	Trimester	Gestation With	Vaccinated	studya	Abortionsb	Abortions		
	1	Control Product	233	7	2	2		
	1	Test Product	235	4	3	1		
	2	Control Product	230	0	2	1		
	2	Test Product	231	2	6	1		
	3	Control Product	224	2	5	1		
	3	Test Product	216	1 8		0		
		s affirmed by stu						
		s affirmed by stu bovine rhinotrac					ed to	
							tracheitis or	
	<sup>c</sup> Abortions with unknown causes OR related to infectious bovine rhinotracheitis or bovine virus diarrhea viruses							
			Nur	mber of Calves				
	Trimester	Vaccinated During Gestation With	Live Births <sup>a</sup>	Mortality Prior to End of Observation Period				
	1	Control Product	224	4				
	1	Test Product	229	6				
	2	Control Product	233	5				
	2	Test Product	230	2				
	3	Control Product	223	14				
	3	Test Product	221	5				
	<sup>a</sup> Live births include twins							
TICDAA	M 0. 2012							
USDA Approval	May 9, 2013							
Date								

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Study Type	Safety				
Pertaining to	ALL				
Study Purpose	To demonstrate safety under field conditions.				
<b>Product Administration</b>					
Study Animals	Bovine				
<b>Challenge Description</b>					
Interval observed after					
challenge					
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. No data are published because this study was submitted to USDA-APHIS prior to January 1, 2007, and APHIS only requires publication of data submitted after that date.				
<b>USDA Approval Date</b>	June 16, 2004				

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Study Type	Safety				
Pertaining to	ALL				
Study Purpose	To demonstrate safety in pregnant animals under field conditions				
_	when cows or heifers are vaccinated prior to breeding, within the				
	previous 12 months, with a modified live Infectious Bovine				
	Rhinotracheitis Virus (IBRV) and Bovine Viral Diarrhea Virus				
	(BVDV) product				
Product Administration	Two doses, administered subcutaneously. First vaccination				
	given 14 to 60 days prior to breeding. Second vaccination given				
	during a specified trimester of pregnancy.				
Study Animals	<u>1st Trimester Study</u> : 468 pregnant heifers $(52 - 86)$ days				
	pregnant) 2 years of age and older.				
	2 <sup>nd</sup> Trimester Study: 461 pregnant heifers (100 – 180 days				
	pregnant) 2 – 14 years of age.				
	3 <sup>rd</sup> Trimester Study: 440 pregnant heifers (≥190 days pregnant)				
	2 years of age and older.				
<b>Challenge Description</b>	Not applicable				
Interval observed after	All cows were observed from pre-breeding vaccination through				
challenge	calving.				
Results	Summary of the results listed in the table below				
<b>USDA Approval Date</b>	May 9, 2013				

### Summary of the results as follows:

		No. of Cows			Fetal Loss (%)
Trimester	Group	Entered	Removed*	Fetal Loss (%) related to vaccination	unrelated to vaccination as affirmed by licensee
1 <sup>st</sup>	Vaccinates	235	4	1 (0.4 %)	3 (1.3%)
	Controls	233	7	2 (0.9%)	2 (0.9%)
2 <sup>nd</sup>	Vaccinates	231	2	1 (0.4%)	6 (2.5%)
2	Controls	230	0	1 (0.4%)	2 (0.8%)
3 <sup>rd</sup>	Vaccinates	216	1	0 (0%)	8 (3.7%)
	Controls	224	2	1 (0.5%)	5 (2.2%)

<sup>\*</sup>Number of cows removed from the study results due to death serious illness considered unrelated to vaccination as affirmed by licensee

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