

Summary of Studies Supporting USDA Product Licensure

Establishment Name	Boehringer Ingelheim Animal Health USA Inc.
USDA Vet Biologics Establishment Number	124
Product Code	4469.24
True Name	Bovine Rhinotracheitis-Virus Diarrhea-Parainfluenza 3- Respiratory Syncytial Virus Vaccine, Modified Live Virus, Leptospira Canicola-Grippotyphosa-Hardjo- Icterohaemorrhagiae-Pomona Bacterin
Tradename(s) / Distributor or Subsidiary (if different from manufacturer)	Express FP 10 - Agricultural Machinery and Materials Co. Ltd Express FP 10 - Boehringer Ingelheim (Canada) Ltd. Express FP 10 - No distributor specified Lepto 5 - No distributor specified
Date of Compilation Summary	November 02, 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	Bovine Virus Diarrhea (BVD)						
Study Purpose	Demonstration of efficacy against persistent infection of calve						
	with BVD Type 1						
Product Administration	Pregnant heifers						
Study Animals	Bovine						
Challenge Description	BVD Type 1a isolate BJ						
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	September 19, 2003						

Study Type	Efficacy
Pertaining to	Bovine Virus Diarrhea (BVD)
Study Purpose	Demonstration of efficacy against BVD Type 1 (respiratory
	disease)
Product Administration	
Study Animals	Bovine
Challenge Description	BVD Type 1 isolate NY-1
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	November 9, 1998

Study Type	Efficacy
Pertaining to	Bovine Virus Diarrhea (BVD)
Study Purpose	Demonstration of efficacy against persistent infection of calves with BVD Type 1
Product Administration	
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	January 2, 2002; September 25, 2002

Study Type	Efficacy						
Pertaining to	Bovine Virus Diarrhea Type 1 (BVD1)						
Study Purpose	Demonstration of 12 month duration of immunity against BVD1						
	(respiratory and reproductive)						
Product Administration	One dose, subcut	aneously					
Study Animals	Heifers (22 Vacc	inates and 23 Co	ontrols), 12-15 m	nonths of age			
Challenge Description	Challenged with	noncytopathic B	VD1b BJ strain,	12 months			
	(368 days) after v	vaccination and a	approximately 93	3 days of			
	gestation.						
Interval observed after	Observed for 14	days after challe	nge. Blood colle	ected on days 0,			
challenge	2, 4, 6, 8, 10, 12	and 14 after chal	lenge to evaluat	e viremia and			
	leukopenia. Fetu	ses collected on	day 72 after cha	llenge.			
Results	Results of the stu	dy are summariz	zed as follows:				
	Blood was evalua	ated for viremia	(the presence of	virus) and			
	leukopenia (at lea	ast one white blo	od cell count no	greater than			
	40% of pre-challe	enge baseline co	unt).				
	Dositivo for J	Viromia and La	ukononio.				
	r ositive for	Viromio	ukopema:	1			
	Vacinator			-			
	Vaccinates	0/22(0%)	$\frac{6}{22} (30\%)$	-			
	Controls	19/23 (0370)	21/23 (9170)				
	Calvas (fatusas)	wara considered	positivo for por	vistont BVD			
	infaction if at loa	st one fotal tissu	positive for pers	r if haifars wara			
	open when the fe	si one retar tissu	e was positive of	tissues spleen			
	thymus hoart blo	ad and coroball	um wore evaluat	tad for the			
	presence of BVD	by virus isolat	ion				
	presence of DVD	of by virus isolat	1011.				
	Positive for l	RVD Persistent	Infection				
	Vaccinates	1/22 (5%)					
	Controls	20/23 (87%)					
		20/23 (07/0)]				
	See tables on the	following pages	for data.				
USDA Annuaval Data	October 3 2011						

Viremia

Vaccinates (22 bovine)

Animal	Days Post-Challenge									
ID	0	2	4	6	8	10	12	14		
5	-	-	-	-	-	-	-	-		
11	-	1	1	-	-	-	-	-		
29	-	1	1	-	-	-	-	-		
37	-	-	-	-	-	-	-	-		
56	-	1	1	-	-	-	-	-		
64	-	I	1	-	-	-	-	-		
77	-	-	-	-	-	-	-	-		
92	-	-	-	-	-	-	-	-		
120	-	-	-	-	-	-	-	-		
125	-	-	-	-	-	-	-	-		
149	-	-	-	-	-	-	-	-		
152	-	-	-	-	-	-	-	-		
156	-	-	-	-	-	-	-	-		
181	-	-	-	-	-	-	-	-		
185	-	-	-	-	-	-	-	-		
201	-	-	-	-	-	-	-	-		
223	-	-	-	-	-	-	-	-		
250	-	-	-	-	-	-	-	-		
260	-	-	-	-	-	-	-	-		
263	-	-	-	-	-	-	-	-		
277	-	-	-	-	-	-	-	-		
300	-	-	-	-	-	-	-	-		

Controls (23 bovine)

Animal	Days Post-Challenge									
ID	0	2	4	6	8	10	12	14		
17	-	-	-	-	+	-	-	-		
22	-	-	-	-	-	-	-	-		
51	-	-	-	-	+	-	-	-		
53	-	-	-	-	-	-	-	-		
58	-	-	-	-	+	-	-	-		
66	-	-	-	+	-	-	-	-		
94	-	-	-	-	+	+	-	-		
103	-	-	-	-	+	-	-	-		
111	-	-	-	+	+	-	-	-		
134	-	-	-	-	+	-	-	-		
135	-	-	-	+	-	-	-	-		
136	-	-	-	-	+	-	-	-		
141	-	-	-	-	-	-	-	-		
179	-	-	-	+	+	-	-	-		
198	-	-	-	-	+	-	-	-		
225	-	-	-	+	+	-	-	-		
230	-	-	-	-	+	-	-	-		
236	-	-	-	-	+	-	-	-		
241	-	-	-	+	+	-	-	-		
243	-	-	+	+	+	-	-	-		
259	-	-	-	-	+	-	-	-		
262	-	-	-	-	+	-	-	-		
283	-	-	-	-	-	-	-	-		

+ = positive for virus (highlighted yellow)

- = negative for virus

Animal	White Blood Cell Count per each Day Post-Challenge									
ID	Baseline (-2)	0	2	4	6	8	10	12	14	Result
5	6.8	7.5	6	8.9	7	10	6.6	7.7	6.7	-
11	8.8	8.1	4.9	7.3	5.4	5.6	4.9	4	5.9	+
29	7.2	6.2	3.6	4.6	5	4.5	4.7	5	4.3	+
37	8.9	7.3	6.2	6.9	6.1	4.4	7.1	4.1	3.9	+
56	6.5	5.3	6.4	4.9	5	5.3	6.8	4.8	5.1	-
64	8.8	7	7.2	8.9	8.1	6.7	5.3	7.4	6.6	-
77	5	7.1	3.3	5	5.9	5.8	3.1	3.9	2.7	+
92	6	7.1	4.6	4	5.7	5.6	4.8	5.2	4.4	-
120	8.1	4.8	5.6	7.5	5	4.9	4.7	4.7	6.9	+
125	4.6	7.8	8.3	5.3	5.3	5.7	4.8	5.1	4.1	-
149	9.1	8.3	7.4	6.7	6.6	6.2	8.4	5.9	5.5	-
152	9.4	12	8.6	5.4	6.9	6.8	5.3	7.9	7.8	+
156	8.1	10	7.3	7.1	7.3	7.9	6.6	7.1	6.6	-
181	6.8	5.7	4.9	4.6	4.2	4.5	6	3.9	2.8	+
185	7.2	4.9	5.8	5.4	7.7	7.8	5.8	5.6	5.1	-
201	7.2	8.2	6.2	5.6	6.2	6.5	5.9	6.3	8.1	-
223	7.8	7.8	8	7.6	6	5.1	5.7	5	6	-
250	7.9	6	6.2	5	5	8.5	5.6	6	5.7	-
260	9.3	6.3	5.9	6.1	7.3	5.9	6.1	6.2	6.1	-
263	7	9.7	7.2	6.8	7.6	7	7.7	6.3	8.3	-
277	11.3	5.6	6	5.9	4.9	4.5	6.8	4.1	4.1	+
300	6.2	7.2	5.2	4.4	6.3	6.7	5.5	6.5	6.9	-

Leukopenia in Vaccinates (22 bovine)

White Blood Cell Count:

• Numerical values = white blood cell count in thousands of cells per cubic millimeter of blood (i.e. 6.8 equals 6,800 cells/mL³)

Overall Result:

+ = positive for leukopenia at least one day (highlighted yellow)

Animal	White Blood Cell Count per each Day Post-Challenge											
ID	Baseline (-2)	0	2	4	6	8	10	12	14	Result		
17	7.4	7.6	5.3	2.4	3.5	2.7	6.6	8.2	5.7	+		
22	7.2	7.3	6.4	2.5	3.6	3.2	5.1	5.2	5.9	+		
51	4	6.6	5.4	2.7	4.1	4	4.5	4.8	3.9	-		
53	7.3	8.1	6.9	2.5	3.2	5.9	8.8	4.2	4.5	+		
58	11.5	10.2	8	4	4.7	4.7	9.5	5	7.5	+		
66	8.3	6.2	3.5	3.7	2.7	5	6.5	3.8	5.6	+		
94	8.3	6.8	6.7	4.5	6.2	6.3	9.5	9.4	8.4	+		
103	12.5	7.2	5.8	3.5	4.3	4.9	5	4.2	6.6	+		
111	8.9	5.6	6.8	3.6	5.5	5.6	6.8	6.5	3.8	+		
134	8.6	9.6	6.1	4.5	3.8	6.3	5.7	7.4	11.4	+		
135	6.4	6.1	5.8	3.4	4	3.7	3.9	5.3	6.7	+		
136	7.1	6.4	5.9	3.4	4.8	5.5	6.8	7.4	7.1	+		
141	19.3	16	10.5	4.9	5.8	6.7	5.9	6.7	6.3	+		
179	5.5	6.8	6.7	2	2.5	5	2.9	3.2	5.8	+		
198	12.8	9	8.1	2.6	2.8	3.5	4.5	4.4	4.6	+		
225	8.3	8.4	9.6	4.7	4.4	5.3	4.3	3.6	5.1	+		
230	7.7	9.2	7.9	3.5	4.9	6.4	5.9	6	8	+		
236	11.4	11.8	9.2	3.6	4.4	5.5	7.8	7.2	5.5	+		
241	9.4	7.7	5.9	3.2	4	5.7	8.7	5.4	6.4	+		
243	5	5	5.7	2.9	3.4	8.1	3.5	3.4	5.4	+		
259	5.2	6.4	8.4	3.7	5.6	10.1	8.5	8.9	6.8	-		
262	8.6	8.1	7.7	3.7	6.3	6.8	3.9	8.5	4.8	+		
283	4.8	5.5	5.5	1.5	3	4.6	2.9	2.8	2.5	+		

Leukopenia in Controls (23 bovine)

White Blood Cell Count:

• Numerical values = white blood cell count in thousands of cells per cubic millimeter of blood (i.e. 6.8 equals 6,800 cells/mL³)

Overall Result:

+ = positive for leukopenia at least one day (highlighted yellow)

Persistent Infection of Calves

Vaccinates (22 bovine)						Controls (23 bovine)						
		V	irus Is BVD 1	olatior Fype 1	n:			Virus Isolation: BVD Type 1				
Animal ID	Result	Spleen	Thymus	Heart Blood	Brain	Animal ID	Result	Spleen	Thymus	Heart Blood	Brain	
5	Negative	-	-	-	-	17	Positive	+	+	+	+	
11	Negative	-	-	-	-	22	Negative	-	-	-	-	
29	Negative	-	-	-	-	51	Positive	+	+	+	+	
37	Negative	-	-	-	-	53	Negative	-	-	-	-	
56	Negative	-	-	-	-	58	Positive	+	+	+	+	
64	Negative	-	-	-	-	66	Positive	+	+	+	+	
77	Negative	-	-	-	-	94	Positive	+	+	+	+	
92	Negative	-	-	-	-	103	Positive	-	+	+	+	
120	Negative	-	-	-	-	111	Positive	+	+	+	+	
125	Negative	-	-	-	-	134	Positive	+	+	+	+	
149	Negative	-	-	-	-	135	Positive	NA	NA	NA	NA	
152	Negative	-	-	-	-	136	Positive	+	+	+	+	
156	Negative	-	-	-	-	141	Positive	NA	NA	NA	NA	
181	Positive	NA	NA	NA	NA	179	Positive	+	+	+	+	
185	Negative	-	-	-	-	198	Negative	-	-	-	-	
201	Negative	-	-	-	-	225	Positive	NA	NA	NA	NA	
223	Negative	-	-	-	-	230	Positive	NA	NA	NA	NA	
250	Negative	-	-	-	-	236	Positive	+	+	+	+	
260	Negative	-	-	-	-	241	Positive	-	+	+	+	
263	Negative	-	-	-	-	243	Positive	+	+	+	+	
277	Negative	-	-	-	-	259	Positive	+	+	+	+	
300	Negative	-	-	-	-	262	Positive	NA	NA	NA	NA	
						283	Positive	+	+	+	+	

Vaccinatos (?? hovina)

Result:

Positive = positive for BVD persistent infection because at least one fetal tissue was positive, or heifer was open Negative = negative for BVD persistent infection because all fetal tissues were negative

Virus Isolation:

+ = fetal tissue positive for BVD1 by virus isolation

- = fetal tissue negative for BVD1 by virus isolation

NA = not applicable because heifer was open

Study Type	Efficacy
Pertaining to	Bovine Virus Diarrhea (BVD)
Study Purpose	Demonstration of efficacy against BVD Type 2 (persistently
	infected calves)
Product Administration	Pregnant heifers
Study Animals	Bovine
Challenge Description	BVD Type 2a isolate PA131
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	August 24, 2006

Study Type	Efficacy							
	Dervice View Director (DVD)							
Pertaining to	Bovine Virus Diarrhea (BVD)							
Study Purpose	Demonstration of efficacy against BVD Type 2 (respiratory							
	disease)							
Product Administration								
Study Animals	Bovine							
Challenge Description	BVD Type 2a isolate BVD 890							
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	November 9, 1998							

Study Type	Efficacy
Pertaining to	Bovine Virus Diarrhea (BVD)
Study Purpose	Demonstration of efficacy against persistent infection of calves
	with BVD Type 2
Product Administration	Pregnant heifers
Study Animals	Bovine
Challenge Description	BVD Type 2 isolate PA131
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	September 19, 2003

Study Type	Efficacy
Pertaining to	Bovine Virus Diarrhea (BVD)
Study Purpose	Demonstration of efficacy against persistent infection of calves
	with BVD Type 2
Product Administration	Pregnant heifers
Study Animals	Bovine
Challenge Description	BVD Type 2a isolate NY-93
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	January 2, 2002; September 25, 2002

Study Type	Efficacy	Efficacy					
Pertaining to	Bovine Virus Dia	Bovine Virus Diarrhea Type 2 (BVD2)					
Study Purpose	Demonstration of 12 month duration of immunity against BVD2						
	(respiratory and p	(respiratory and persistent infection of calves)					
Product Administration	One dose, subcut	aneously					
Study Animals	Heifers (18 Vacc	inates and 22 Co	ontrols), 13-16 m	onths of age			
Challenge Description	Challenged with	noncytopathic B	VD2 PA131 stra	in, 12 months			
	(374 days) after v	vaccination and a	at approximately	90 days of			
	gestation						
Interval observed after	Observed for 14	days after challe	nge. Blood colle	ected on days 0,			
challenge	2, 4, 6, 8, 10, 12	and 14 after cha	llenge to evaluate	e viremia and			
	leukopenia. Fetu	ses collected on	day 65 after cha	llenge.			
Results	Results of the stu	dy are summarized	zed as follows:				
	D1 1 1	. 1.0	(4)	• \ 1			
	Blood was evalua	ated for viremia	(the presence of	virus) and			
	100/ of any oball	ast one white blo	ood cell count no	greater than			
	40% of pre-chang	enge basenne co	unt).				
	Positive for V	Viremia and Le	ukopenia:				
		Viremia	Leukopenia				
	Vaccinatos	ViremiaLeukopeniaVariation $1/19$ ($0/2$)					
	$\begin{array}{ c c c c c c c c c c c c c c c c c c c$						
	Controls	20/22 (91%)	1/18 (6%)				
	Controls	20/22 (91%)	14/22 (50%)				
	Calves (fetuses)	20/22 (91%) were considered	1718 (6%) 14/22 (50%) positive for pers	istent BVD			
	Calves (fetuses) vinfection if at lea	were considered st one fetal tissu	positive for pers was positive.	istent BVD The fetal tissues			
	Calves (fetuses) infection if at lea spleen, thymus, h	were considered st one fetal tissu	positive for pers was positive.	istent BVD The fetal tissues evaluated for			
	Calves (fetuses) infection if at lea spleen, thymus, h the presence of B	were considered st one fetal tissu heart blood, and SVD2 by virus is	positive for pers e was positive. T cerebellum were olation.	istent BVD The fetal tissues evaluated for			
	Calves (fetuses) infection if at lea spleen, thymus, h the presence of B	were considered st one fetal tissu heart blood, and VD2 by virus is	positive for pers e was positive. The cerebellum were olation.	istent BVD The fetal tissues evaluated for			
	Calves (fetuses) infection if at lea spleen, thymus, h the presence of B Positive for 1	were considered st one fetal tissu heart blood, and BVD2 by virus is	positive for pers e was positive. The cerebellum were olation.	istent BVD The fetal tissues evaluated for			
	Calves (fetuses) v infection if at lea spleen, thymus, h the presence of B Positive for I Vaccinates	were considered st one fetal tissu heart blood, and VD2 by virus is BVD Persistent 0/18 (0%)	positive for pers e was positive. The cerebellum were olation.	istent BVD The fetal tissues evaluated for			
	Calves (fetuses) infection if at lea spleen, thymus, h the presence of B Positive for 1 Vaccinates Controls	were considered st one fetal tissu heart blood, and BVD2 by virus is BVD Persistent 0/18 (0%) 21/22 (95%)	positive for pers e was positive. The cerebellum were olation.	istent BVD The fetal tissues evaluated for			
	Calves (fetuses) v infection if at lea spleen, thymus, h the presence of B Positive for 1 Vaccinates Controls	1718 (0%)20/22 (91%)were consideredst one fetal tissuheart blood, and bVD2 by virus isBVD Persistent0/18 (0%)21/22 (95%)	positive for pers e was positive. The cerebellum were olation.	istent BVD The fetal tissues evaluated for			
	Calves (fetuses) infection if at lea spleen, thymus, h the presence of B Positive for 1 Vaccinates Controls See tables on the	were considered st one fetal tissu heart blood, and VD2 by virus is BVD Persistent 0/18 (0%) 21/22 (95%) following pages	positive for pers e was positive. The construction of the construc	istent BVD The fetal tissues evaluated for			

Viremia

Vaccinates (18 bovine)

Animal	Days Post-Challenge								
ID	0	2	4	6	8	10	12	14	
9	-	-	-	-	-	-	-	-	
21	-	-	-	-	-	-	-	-	
43	-	1	-	-	-	-	-	-	
57	-	-	-	-	-	-	-	-	
70	-	I	-	-	-	-	-	-	
82	-	I	-	-	-	-	-	-	
90	-	1	-	-	-	-	-	-	
97	-	1	-	-	-	-	-	-	
106	-	-	-	-	-	-	-	-	
114	-	-	-	-	-	-	-	-	
130	-	1	-	+	-	-	-	-	
137	-	-	-	-	-	-	-	-	
191	-	-	-	-	-	-	-	-	
196	-	-	-	-	-	-	-	-	
227	-	-	-	-	-	-	-	-	
242	-	-	-	-	-	-	-	-	
271	-	-	-	-	-	-	-	-	
272	-	-	-	-	-	-	-	-	

Controls (22 bovine)

Animal		Days Post-Challenge							
ID	0	2	4	6	8	10	12	14	
2	-	-	-	-	+	-	-	-	
3	-	-	-	+	+	+	-	-	
7	-	-	-	+	+	-	-	-	
12	-	-	-	-	-	-	-	-	
20	-	-	-	1	+	+	-	-	
24	-	-	-	+	+	-	-	-	
27	-	-	-	-	-	-	-	-	
81	-	-	-	+	+	-	-	-	
88	-	-	-	+	+	+	+	-	
91	-	-	-	+	-	-	-	-	
145	-	-	-	+	+	+	-	-	
157	-	-	-	I	+	-	-	-	
159	-	-	+	+	+	+	-	+	
168	-	-	-	+	+	-	-	-	
170	-	-	-	+	+	-	-	-	
199	-	-	-	+	+	-	-	-	
202	-	-	-	+	+	-	-	-	
211	-	-	-	I	+	+	-	-	
224	-	-	+	+	+	+	-	-	
248	-	-	-	+	+	-	-	-	
269	-	-	-	+	+	+	-	-	
279	-	-	-	+	+	-	-	-	

+ = positive for virus (highlighted yellow)

- = negative for virus

A		White Blood Cell Count per each								Orverall
ID	Baseline (-2)	0	2	4	6	8	10	12	14	Result
9	4.9	4.3	6.5	8.8	6.8	7.4	8.6	8.5	6.4	-
21	7	6.6	9	5.4	7.2	7	5.4	5.9	6.7	-
43	4	5.2	9.8	9.1	3.7	7.9	6.5	6.3	7.5	-
57	5.4	6.6	6.8	5.2	5	6.6	8.3	6.2	7.6	-
70	4.6	8.6	6.2	6	4.5	6.4	6.1	9.6	5.4	-
82	6	6.7	8.4	10.2	5.5	9.9	6.8	7.3	6.9	-
90	5	6.5	5.9	9.6	4.9	6.3	6.6	6	7.1	-
97	5.2	6.7	6.8	7.7	6.4	5.5	6.8	7	4.6	-
106	3.8	8.1	6.3	6.5	4.7	8	7.2	5.8	6.6	-
114	6.4	6.9	6.1	5.1	6.3	6.9	6.9	5.3	5.3	-
130	5.4	4.5	7.2	5.5	3.6	9.4	6.4	5.2	6.6	-
137	8	5.1	4.9	6.5	10	7.4	9.1	6.4	4.7	+
191	3.6	7.6	7.9	7.2	4.3	7.2	5.7	6.5	5.5	-
196	4.2	8.2	4.3	6.5	8.4	8.3	5	4.9	4.8	-
227	3.7	4.8	5.8	4.8	5.3	8.2	6.2	9.2	6.5	-
242	8.7	10	9	7.1	5.6	8.8	7.4	5.4	5.5	-
271	5.4	6.1	5.6	5.9	5.8	10.9	6.7	7.1	10.1	-
272	5.9	5.3	4.2	5.8	4.7	5.1	5.9	5.7	7.1	-

Leukopenia in Vaccinates (18 bovine)

White Blood Cell Count:

Overall Result: + = positive for leukopenia at least one day (highlighted yellow)

Numerical values = white blood cell count in thousands of cells per cubic millimeter of blood (i.e. 4.9 equals • 4,900 cells/mL³)

Animal		White Blood Cell Count per each Day Post-Challenge								
ID	Baseline (-2)	0	2	4	6	8	10	12	14	Result
2	4.5	6.4	8.1	2.6	4.8	3.3	4.5	7.3	8.2	+
3	10.4	6	8.9	4.8	6.7	6.1	8.3	6.4	6.7	+
7	3.7	3.8	5.4	3.9	3.6	5.2	4.3	5.1	5.1	-
12	3.4	4.6	6.2	3.7	3.7	5.4	4.5	4.9	5.5	-
20	6.3	3.4	7.5	3.5	4.3	4.8	5.5	4.3	4.5	-
24	4.5	6	7	3.1	3.6	4.4	6.2	5.1	6.9	+
27	1.5	4.3	6	2.5	1.4	2.5	5	2.8	2.9	+
81	3.2	4	7.6	2.7	3.2	4	5.6	4.4	4.3	-
88	7.2	7.9	7.3	5.9	7.5	5.4	5	6.3	5.6	-
91	5.9	7.8	9.7	3.7	4.3	7.3	12.7	9.9	11	+
145	4.3	4.2	5.5	2.9	4.5	3.6	4.3	5.7	6.7	-
157	6.4	8.6	12.1	3.1	4.9	6.9	3.8	4.5	6.5	+
159	4.8	5.6	8.3	4.4	7	4.7	4	5.1	4.1	-
168	7.7	7	5.6	4	3.4	3.8	3.7	3.9	6.6	+
170	5.2	5.5	7	3	3.7	3.8	4	4.3	4.8	+
199	3.5	4.6	4.3	3.6	3.2	5.7	3.2	3.7	5.6	-
202	5.6	5.6	6	2	5.2	2.5	1.1	3.3	4.8	+
211	5.2	8.6	7.3	3.4	3.3	2.8	6.1	4.1	5.9	+
224	2.7	6.9	5	5	6.3	3	2.2	5.3	4.8	+
248	5.9	6.4	8.2	3.4	6.2	5.5	4.2	4.1	6.6	+
269	7.2	8.1	6.4	4.8	2.6	ND	4.2	3.9	4.7	+
279	5.7	7.4	6	3	4.2	3.1	4.7	3.3	3.2	+

Leukopenia in Controls (22 bovine)

White Blood Cell Count:

• Numerical values = white blood cell count in thousands of cells per cubic millimeter of blood (i.e. 4.9 equals 4,900 cells/mL³)

Overall Result:

+ = positive for leukopenia at least one day (highlighted yellow)

Persistent Infection of Calves

	Vaccinates (18 bovine)						
		Virus Isolat BVD Type					
		BVD Type 2					
Animal ID	Result	Spleen	Thymus	Heart Blood	Brain		
9	Negative	-	-	-	-		
21	Negative	-	-	I	-		
43	Negative	-	-	-	-		
57	Negative	-	-	-	-		
70	Negative	-	-	-	-		
82	Negative	-	-	-	-		
90	Negative	-	-	-	-		
97	Negative	-	-	-	-		
106	Negative	-	-	-	-		
114	Negative	-	-	-	-		
130	Negative	-	-	-	-		
137	Negative	-	-	-	-		
191	Negative	-	-	-	-		
196	Negative	-	-	-	-		
227	Negative	-	-	-	-		
242	Negative	-	-	-	-		
271	Negative	-	-	-	-		
272	Negative	-	_	_	-		

Virus Isolation: **BVD Type 2** Animal Thymus Result Spleen Heart Blood Brain ID + + + 2 Positive -3 + + Positive + + 7 + + + + Positive 12 Positive + + + + 20 + + + + Positive 24 + Positive ++ + + + + + 27 Positive 81 Negative ----88 Positive + + + + 91 Positive + + + + 145 Positive + + + + 157 Positive _ + + + 159 + + + Positive ++ 168 Positive + + + 170 + + + + Positive 199 Positive + + + + + + + + 202 Positive 211 Positive + + + + + + + + 224 Positive 248 + + + + Positive 269 Positive + + + +

Controls (22 bovine)

Result:

Positive = positive for BVD persistent infection because at least one fetal tissue was positive, or heifer was open Negative = negative for BVD persistent infection because all fetal tissues were negative

279

Positive

+

+

+

+

Virus Isolation:

+ = fetal tissue positive for BVD2 by virus isolation

- = fetal tissue negative for BVD2 by virus isolation

Study Type	Efficacy
Pertaining to	Infectious Bovine Rhinotracheitis (IBR)
Study Purpose	Demonstration of efficacy against IBR (respiratory disease)
Product Administration	
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	May 4, 1994

Study Type	Efficacy	Efficacy					
Pertaining to	Infectious Boy	vine Rhinotracheit	is (IBR)				
Study Purpose	Demonstration	n of efficacy again	st IBR (reproductiv	ve disease) 12			
	months after v	accination					
Product Administration	One dose, sub	cutaneously appro	ximately five mon	ths prior to			
	breeding						
Study Animals	32 bovine (13	vaccinates and 19	controls), 7 - 9 mc	onths of age			
Challenge Description	Challenged wi	th IBR Cooper str	ain 386 days after	vaccination at			
	approximately	7 months of gesta	ntion				
Interval observed after	Cattle were ob	served daily after	challenge and unti	l calving for			
challenge	signs of aborti	on. Fetal tissues v	were evaluated for	the presence of			
	IBR and other	causes of abortion	1.				
Results	Cattle were co	nsidered affected	if the fetus was abo	orted and testing			
	results of the f	etus were negative	e for other causes of	of abortion			
	(Bovine viral c	diarrhea virus (BV	DV) and abortifac	ient bacteria).			
	Results of the	study are summar	ized as follows:				
			_				
	Abortions in v	vaccinates and co	ontrols:	1			
		Non-Aborted	Aborted				
	Vaccinates	11/13 (84.6%)	2/13 (15.4%)				
	Controls 1/19 (5.3%) 18/19 (94.7%)						
	See table on th	e following page	for data.				
USDA Approval Date	October 5, 201	1					

Treatment	Animal	Abortion	IBR by PCR		IBR by V	Virus Isola	ation (VI))	BVDV by VI
				Brain	Kidney	Liver	Lung	Thymus	Same tissues
	6	No	NA	NA	NA	NA	NA	NA	NA
	10	Yes	Negative	-	-	-	-	-	-
	34	No	NA	NA	NA	NA	NA	NA	NA
	45	No	NA	NA	NA	NA	NA	NA	NA
	89	No	NA	NA	NA	NA	NA	NA	NA
Vaccinatas	117	No	NA	NA	NA	NA	NA	NA	NA
(13 howing)	155	No	NA	NA	NA	NA	NA	NA	NA
(15 00vine)	176	Yes	Positive	-	-	-	-	+	-
	180	No	NA	NA	NA	NA	NA	NA	NA
	206	No	NA	NA	NA	NA	NA	NA	NA
	209	No	NA	NA	NA	NA	NA	NA	NA
	228	No	NA	NA	NA	NA	NA	NA	NA
	276	No	NA	NA	NA	NA	NA	NA	NA
	18	Yes	Positive	+	-	-	-	-	-
	26	Yes	Positive	-	-	-	-	-	-
	30	Yes	Positive	-	-	-	-	-	-
	41	Yes	Positive	-	-	-	-	-	-
	42	Yes	Positive	-	-	-	-	-	-
	47	Yes	Positive	-	-	-	-	-	-
	48	Yes	Positive	-	-	-	-	-	-
	62	Yes	Positive	-	-	-	+	-	-
Controla	119	Yes	Positive	-	-	-	+	-	-
(10 howine)	128	No	NA	NA	NA	NA	NA	NA	NA
(19 00vine)	154	Yes	Positive	-	-	-	-	-	-
	161	Yes	Positive	-	-	-	-	-	-
	174	Yes	Positive	-	-	-	-	-	-
	187	Yes	Positive	-	-	-	+	-	-
	194	Yes	Positive	-	-	-	-	-	-
	210	Yes	Positive	-	-	-	-	-	-
	219	Yes	Positive	-	+	-	-	-	-
	257	Yes	Positive	+	-	-	-	-	-
	282	Yes	Positive	+	-	-	+	-	-

Abortion status and evaluation of fetal tissues:

NA = Not applicable since calf was not aborted.

Positive = Positive for the presence of IBR virus by PCR in all fetal tissues examined. **Negative** = Negative for the presence of IBR virus by PCR in all fetal tissues (brain, kidney, liver, lung, and thymus).

+ = Positive for the presence of IBR or BVDV by virus isolation.

- = Negative for the presence of IBR or BVDV by virus isolation.

The same tissues were assessed for BVDV (brain, kidney, liver, lung, thymus).

Tissues were negative for abortifacient bacteria. Data not shown.

Study Type	Efficacy
Pertaining to	Leptospira canicola
Study Purpose	Demonstration of efficacy against leptospirosis caused by
	Leptospira canicola
Product Administration	
Study Animals	Bovine and Porcine
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	July 14, 1981

Study Type	Efficacy
Pertaining to	Leptospira grippotyphosa
Study Purpose	Demonstration of efficacy against leptospirosis caused by
	Leptospira grippotyphosa
Product Administration	
Study Animals	Bovine and Porcine
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	July 14, 1981

Study Type	Efficacy
Pertaining to	Leptospira hardjo
Study Purpose	Demonstration of efficacy against leptospirosis caused by
	Leptospira hardjo
Product Administration	
Study Animals	Bovine and Porcine
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	July 14, 1981

Study Type	Efficacy									
Pertaining to	Leptospira hardjo									
Study Purpose	Demonstration	Demonstration of efficacy against Leptospira borgpetersenii								
	serovar hardje	p-bovis								
Product Administration	Two doses, 2	Two doses, 21 days apart, Subcutaneously								
Study Animals	32 bovine (21	32 bovine (21 vaccinates, 11 controls), 6 months of age								
Challenge Description	Challenged wa	ith <i>Leptospira borg</i>	<i>petersenii</i> sero ^s	var <i>hardjo-bovis</i>						
	on 84, 85 and	86 days after the se	cond vaccinati	on						
Interval observed after	Cattle were observed daily after challenge. Urine samples were									
challenge	taken weekly for 8 weeks. On day 56 and 57 after challenge,									
	kidneys, ovaries, and uterine tissues were cultured for <i>Leptospira</i>									
	isolation.									
Results	An animal was considered affected if urine cultures were									
	positive at one	e or more points afte	er challenge.							
	Results of the	study are summariz	ed as follows:							
	TT • 1/			. 1 . 1						
	Urine cultures	were positive for L	<i>eptospira</i> on a	t least one day:						
	Group	# Positive / Total	% Affected	-						
	Vaccinates	0/21	100%	-						
	Controis	11/11	10070	J						
	Kidney cultur	es were positive for	Lentonsira at	necronsv						
	Group	# Positive / Total	% Affected							
	Vaccinates	0 / 21	0%							
	Controls	10 / 11	91%							
				-						
	Ovary culture	s were positive for <i>I</i>	<i>Leptopsira</i> at n	ecropsy:						
	Group	# Positive / Total	% Affected							
	Vaccinates	0 / 21	0%							
	Controls	2 / 11	18%							
		1. 1.0		0 0 1						
	No Leptospire	<i>i</i> was cultured from	the uterine fiss	sue of any of the						
	vaccinated or	control heiters at ne	cropsy.							
	Saa tablaa an	the following pages	for data							
	See tables on 1	the following pages	tor data.							
USDA Approval Date	April 5, 2010									

Urine, Kidney and Ovary Cultures:

Vaccinates:

Animal #	Weekly Urine Observations						tions		Overall Urine	Kidney	Ovary
Allillal #	1	2	3	4	5	6	7	8	Outcome	Outcome	Outcome
2	-	-	-	-	-	-	-	-	Negative	Negative	Negative
7	-	-	-	-	I	-	-	I	Negative	Negative	Negative
10	-	-	-	-	I	-	-	I	Negative	Negative	Negative
11	-	-	-	-	I	-	-	I	Negative	Negative	Negative
12	-	-	-	-	I	-	-	I	Negative	Negative	Negative
13	-	-	-	-	I	-	-	I	Negative	Negative	Negative
14	-	-	-	-	I	-	-	I	Negative	Negative	Negative
15	-	-	-	-	I	-	-	I	Negative Negative		Negative
16	-	-	-	-	1	-	-	I	Negative Negative		Negative
17	-	-	-	-	I	-	-	I	Negative	Negative	Negative
30	-	-	-	-	1	-	-	I	Negative	Negative	Negative
32	-	-	-	-	-	-	-	-	Negative	Negative	Negative
37	-	-	-	-	-	-	-	-	Negative Negative		Negative
41	-	-	-	-	-	-	-	-	Negative Negative		Negative
42	-	-	-	-	I	-	-	I	Negative	Negative	Negative
43	-	-	-	-	I	-	-	I	Negative	Negative	Negative
49	-	-	-	-	I	-	-	I	Negative	Negative	Negative
50	-	-	-	-	-	-	-	-	Negative	Negative	Negative
51	-	-	-	-	-	-	-	-	Negative	Negative	Negative
53	-	-	-	-	-	-	-	-	Negative	Negative	Negative
54	-	-	-	-	-	-	-	-	Negative	Negative	Negative

Controls:

Animal #	Weekly Urine Observations		Overall Urine	Kidney	Ovary						
Allillal #	1	2	3	4	5	6	7	8	Outcome	Outcome	Outcome
4	-	-	+	+	+	+	+	-	Positive Negative		Negative
5	-	-	+	+	+	+	+	+	Positive	Positive	Positive
6	-	-	+	+	+	+	+	+	Positive	Positive Positive	
9	-	-	-	+	+	+	+	+	Positive Positive		Negative
23	-	-	-	+	+	-	+	+	Positive	Positive	Negative
27	-	-	+	+	+	-	-	+	Positive	Positive	Negative
28	-	-	+	+	+	+	+	I	Positive Positive		Positive
31	-	-	-	-	-	+	+	-	Positive Positive		Negative
34	-	-	+	-	+	+	+	+	Positive	Positive	Negative
35	-	-	-	+	+	-	+	+	Positive	Positive	Negative
52	-	-	-	+	+	+	+	-	Positive	Positive	Negative

Weekly Urine Observations:

- = Urine sample was negative for *Leptospira*

+ = Urine sample was positive for *Leptospira* (highlighted yellow)

Overall Urine / Kidney / Ovary Outcome:

Negative = All urine samples / kidney / ovary were negative for *Leptospira*

Positive = At least one urine sample / kidney / ovary was positive for *Leptospira* (highlighted yellow)

~	7 97
Study Type	Efficacy
Pertaining to	Leptospira icterohaemorrhagiae
Study Purpose	Demonstration of efficacy against leptospirosis caused by
	Leptospira icterohaemorrhagiae
Product Administration	
Study Animals	Bovine and Porcine
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	July 14, 1981

Study Type	Efficacy
Pertaining to	Leptospira pomona
Study Purpose	Demonstration of efficacy against leptospirosis caused by
	Leptospira pomona
Product Administration	
Study Animals	Bovine and Porcine
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	July 14, 1981

Study Type	Efficacy
Pertaining to	Bovine Parainfluenza Type 3 (PI ₃)
Study Purpose	Demonstration of efficacy against PI ₃
Product Administration	
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	September 8, 1994; January 16, 2001

Study Type	Efficacy
Pertaining to	Bovine Parainfluenza Type 3 (PI ₃)
Study Purpose	Demonstration of efficacy against PI ₃
Product Administration	
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	November 3, 2000

Study Type	Efficacy
Pertaining to	Bovine Respiratory Syncytial Virus (BRSV)
Study Purpose	Demonstration of efficacy against BRSV
Product Administration	
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	September 19, 2003

Study Type	Efficacy								
Pertaining to	Bovine Respiratory Syncytial Virus (BRSV)								
Study Purpose	Demonstration of efficacy against BRSV								
Product Administration	Two doses, 26 days apart, subcutaneously								
Study Animals	29 bovine (14 vaccinates, 15 controls), 29 – 37 days old								
Challenge Description	Challenged wit	th BRSV at 40 - 4	1 days after final	vaccination					
Interval observed after	Observed daily	for 9 days after o	challenge. Nasal	swabs were					
challenge	collected from	cattle on days 3,	4, 5, 6, 7, 8 and 9	after challenge.					
_	The lungs of ca	attle were examin	ed on 9 days after	the second					
	challenge.								
Results	Results of the s	study are summar	ized as follows:						
	Nasal swabs w	ere evaluated for	BRSV shedding.	An animal was					
	considered pos	itive if shedding	was detected on a	t least one day:					
	Group	Positive	Negative						
	Vaccinates	s 2/14 (14%)	12/14 (86%)						
	Controls	13/15 (87%)	2/15 (13%)						
	Lung lesions were evaluated visually and by palpation. An animal was considered positive if its lungs had any visual or palpable abnormality:								
	Group	Positive	Negative						
	Vaccinates	5/13 (38%)	8/13 (62%)						
	Controls	15/15 (100%)	0/15 (0%)						
	Note: calf #17 wa	as removed from ana	lysis of the lungs due	to a difference in					
	treatment during h	umane euthanasia th	at potentially affected	d the gross					
	appearance of the	lungs.							
	PDSV was isol	lated from the lun	as using virus iso	lation and lung					
	tissue was eval	usted by fluoresc	igs usilig vilus iso ent antibody testi	ng					
	ussue was eval	ualed by Hubbese	on annouy testi	пд.					
	See tables on th	he following page	es for data						
USDA Approval Data	February 12 20	010	.5 101 u ula.						
USDA Appioval Date	1 COluary 12, 20	010							

Crown	Animal	Outcomo		Day	ys Po	st-C	halle	nge	
Group	ID	Outcome	3	4	5	6	7	8	9
	2	Negative	-	-	-	-	-	-	-
	4	Negative	-	-	-	-	-	-	-
	8	Negative	-	-	-	-	-	-	-
	9	Negative	-	-	-	-	-	-	-
	10	Positive	-	-	-	+	-	+	-
Vaccinatas	15	Negative	-	-	-	-	-	nge 8 8 - - - - - - - - - - - - - - - - -	-
v accinates	16	Negative	-	-	-	-	-	-	-
(14 boying)	17	Negative	-	-	-	-	-	-	-
bovine)	26	Negative	-	-	-	-	-	-	-
	27	Positive	-	-	-	-	+	-	-
Group Vaccinates (14 bovine) Controls (15 bovine)	29	Negative	-	-	-	-	-	-	-
	33	Negative	-	-	-	-	-	-	-
	39	Negative	-	-	-	-	-	-	-
	41	Negative	-	-	-	-	-	-	-
	1	Positive	-	-	-	+	+	-	-
	3	Positive	-	-	-	-	+	-	-
	5	Negative	-	-	-	-	-	-	-
	6	Positive	-	-	-	-	+	-	-
	7	Positive	-	-	-	-	+	-	-
	12	Positive	-	-	-	-	+	+	-
Controls	14	Positive	-	-	-	+	, , , , , , , , , , , , , , , , , , ,	-	
(15	18	Positive	-	-	-	-	+	+	-
bovine)	19	Positive	-	-	-	+	+	-	-
	20	Positive	-	-	-	-	+	-	-
Group Vaccinates (14 bovine)	22	Positive	-	-	-	+	+	-	-
	28	Positive	-	-	+	+	+	-	-
Vaccinates (14 bovine) Controls (15 bovine)	31	Positive	-	-	-	+	+	-	-
	35	Positive	-	-	+	-	+	-	-
	37	Negative	-	-	-	-	-	-	-

Nasal Swab Results for BRSV by Virus Isolation:

Outcome =

- Positive if any day was positive (+) for BRSV virus isolation
- Negative if all days were negative (-) for BRSV virus isolation

Nasal swab results =

- + if BRSV was detected by virus isolation
- - if BRSV was not detected by virus isolation

	Animal	Outcome	Total S Lu	Score for Ings	BRSV from Lungs		
Group	Animai ID	(Overall) for Lungs			Virus	Fluorescent	
	12		Visual	Palpable	Isolation	Antibody	
					(VI)	(FA) Testing	
	2	Negative	0	0	Negative	Negative	
	4	Negative	0	0	Negative	Negative	
	8	Negative	0	0	Negative	Negative	
	9	Positive	1	0	Negative	Negative	
	10	Positive	2	0	Negative	Negative	
Vaccinates	15	Negative	0	0	Negative	Negative	
(13	16	Negative	0	0	Negative	Negative	
bovine)	26	Positive	1	0	Negative	Negative	
	27	Negative	0	0	Negative	Negative	
	29	Negative	0	0	Negative	Negative	
	33	Positive	0	5	Negative	Negative	
	39	Positive	3	0	Negative	Negative	
	41	Negative	0	0	Negative	Negative	
	1	Positive	6	2	Positive	Negative	
	3	Positive	6	0	Negative	Negative	
	5	Positive	8	0	Negative	Negative	
	6	Positive	8	3	Negative	Negative	
	7	Positive	19	3	Negative	Negative	
	12	Positive	8	0	Negative	Negative	
Controls	14	Positive	9	0	Positive	Positive	
(15	18	Positive	6	0	Positive	Negative	
bovine)	19	Positive	6	2	Negative	Negative	
	20	Positive	4	0	Negative	Negative	
	22	Positive	4	0	Positive	Negative	
	28	Positive	6	3	Positive	Negative	
	31	Positive	7	4	Positive	Negative	
	35	Positive	17	11	Negative	Positive	
	37	Positive	6	1	Negative	Negative	

Summary of Results for Lung Lesions and Virus Isolation

Outcome (Overall) for Lungs =

- Positive if any parameter is positive (visual lesions, palpable lesions, VI, FA)
- Negative if all parameters are negative (visual lesions, palpable lesions, VI, FA)

Total Score for Lungs =

• Sum of scores for all lung lobes (see following pages for each lung lobe).

• Scores range from 0 (negative / normal) to 32. Any score of 1 or higher is considered positive. **BRSV Virus Isolation (VI) =**

- Positive if BRSV was isolated from lung tissue
- Negative if BRSV was not isolated from lung tissue

Fluorescent Antibody (FA) Testing =

- Positive if BRSV specific staining was observed in lung tissue
- Negative if BRSV specific staining was not observed in lung tissue

						Visual				
Group	Animal ID	Total Score (Sum)	Left cranial	Left middle	Left caudal	Right cranial	Right posterior cranial	Right middle	Right caudal	Inter- mediate
	2	0	0	0	0	0	0	0	0	0
	4	0	0	0	0	0	0	0	0	0
	8	0	0	0	0	0	0	0	0	0
	9	1	0	0	0	0	1	0	0	0
	10	2	0	0	2	0	0	0	0	0
Maarinataa	15	0	0	0	0	0	0	0	0	0
(12 howing)	16	0	0	0	0	0	0	0	0	0
(15 bovine)	26	1	0	0	0	0	0	1	0	0
	27	0	0	0	0	0	0	0	0	0
	29	0	0	0	0	0	0	0	0	0
	33	0	0	0	0	0	0	0	0	0
	39	3	0	1	0	1	0	1	0	0
	41	0	0	0	0	0	0	0	0	0
	1	6	0	1	1	1	1	0	1	1
	3	6	1	1	1	1	0	1	0	1
	5	8	1	1	1	1	1	1	1	1
	6	8	1	2	1	0	1	2	0	1
	7	19	2	2	3	2	2	2	3	3
	12	8	1	1	1	1	0	1	1	2
Controlo	14	9	1	1	1	2	0	1	1	2
(15 bovine)	18	6	0	1	1	2	0	1	1	0
	19	6	0	1	0	2	1	1	0	1
	20	4	0	1	0	0	1	1	0	1
	22	4	0	1	0	2	0	1	0	0
	28	6	1	2	0	1	1	1	0	0
	31	7	0	0	1	3	0	1	1	1
	35	17	2	2	2	2	2	2	3	2
	37	6	0	1	1	1	0	1	1	1

Visual Lung Lesions for Each Lung Lobe:

Scoring System for Lung Lobes (Visual)

	Description
0	Normal
1	Slight multifocal or diffuse congestion
2	Moderate congestion with visible lobular pattern (+/- mild edema)
3	Multiple consolidated lobules; minimal to mild pleuritis
4	Most of all of the lobe consolidated; moderate to severe pleuritis

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_					le					
Group	Animal ID	Total Score (Sum)	Left cranial	Left middle	Left caudal	Right cranial	Right posterior cranial	Right middle	Right caudal	Inter- mediate
	2	0	0	0	0	0	0	0	0	0
	4	0	0	0	0	0	0	0	0	0
	8	0	0	0	0	0	0	0	0	0
	9	0	0	0	0	0	0	0	0	0
	10	0	0	0	0	0	0	0	0	0
X7	15	0	0	0	0	0	0	0	0	0
vaccinates	16	0	0	0	0	0	0	0	0	0
(15 bovine)	26	0	0	0	0	0	0	0	0	0
	27	0	0	0	0	0	0	0	0	0
	29	0	0	0	0	0	0	0	0	0
	33	5	0	1	1	0	0	1	1	1
	39	0	0	0	0	0	0	0	0	0
	41	0	0	0	0	0	0	0	0	0
	1	2	0	0	0	1	0	0	1	0
	3	0	0	0	0	0	0	0	0	0
	5	0	0	0	0	0	0	0	0	0
	6	3	0	1	0	0	1	1	0	0
	7	3	0	0	0	0	0	1	1	1
	12	0	0	0	0	0	0	0	0	0
Controlo	14	0	0	0	0	0	0	0	0	0
(15 hoving)	18	0	0	0	0	0	0	0	0	0
(15 bovine)	19	2	0	0	1	0	0	0	1	0
	20	0	0	0	0	0	0	0	0	0
	22	0	0	0	0	0	0	0	0	0
	28	3	0	0	1	0	0	1	1	0
	31	4	0	1	1	0	0	0	1	1
	35	11	0	1	1	2	1	3	2	1
	37	1	0	0	0	0	0	1	0	0

Palpable Lung Lesions for Each Lung Lobe:

Scoring System for Lung Lobes (Palpable)

	Description
0	Normal
1	Slight or mild diffuse firmness within lobe
2	Moderate diffuse firmness within lobe
3	Non-homogeneous firmness throughout lobe, with palpable solid areas
4	Most or all of lobe palpably solid

Study Type	Safety					
Pertaining to	All fractions					
Study Purpose	To demonstrate safety under field conditions					
Product Administration						
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	July 6, 1999					

Study Type	Safety						
Pertaining to	All fractions						
Study Purpose	To demonstrate safety in pregnant heifers/cows and nursing calves						
Product Administration	Two doses, administered subcutaneously. First vaccination given						
	1-2 months prior to breeding. Second vaccination given during a						
	specified trimester of pregnancy.						
Study Animals	Site 1:						
	2,063 cows a	nd heifers r	eceived vaccine	e prior to bre	eding.		
	1,586 cows	and heifer	s received vac	cine or a p	placebo during		
	pregnancy and are included in this summary.						
	Site 2:						
	120 calves fro	$\frac{m}{1}$ dams that	received vaccine	in the 2 nd or	3 rd trimester.		
Challenge Description	Not applicab	le					
Interval observed after	Not applicab	le					
Challenge Degulte	A 11	haifarr	no obcoment f		na waasin - 4' -		
Kesuits	All cows and	i neifers we	re observed Irol	in pre-breed	ling vaccination		
	Docults of the	ng and carv	es were observe	follows:	ks postpartum.		
	Results of the	e study are s	summarized as i	tonows.			
	Fetal Loss (S	Site 1):					
		Vac	cinates	Control	s (Placebo)		
			Fetal Loss		Fetal Loss		
	Trimester	Enrolled	(%)	Enrolled	(%)		
	1 st	306	7 (2.3%)	274	6 (2.2%)		
	2 nd 237 1 (0.4%) 235 3 (1.3%)						
	3 rd	267	5 (1.9%)	267	6 (2.2%)		
	The number of animals during pregnancy was reduced due to						
	normal losses including dystocia, lameness, and non-study related						
	causes (as af	firmed by li	censee).				
	Fetal loss wa	s due to abo	ortion or open (1	non-pregnan	t). For all three		
	trimesters, r	10 COWS O	r heifers (0.0%	%) in eithe	r group were		
	diagnosed	as having	aborted due	to Infec	ctious Bovine		
	Rhinotracheitis (IBR) or Bovine Virus Diarrhea Virus (BVDV).						
	All tests for viral detection and isolation of IBR and BVDV on all						
	fetal tissues were negative.						
	Eatal Infaction (Site 2).						
	Serum same	oles were o	collected from	calves prio	r to receiving		
	colostrum 6	51 calves	were from cox	vs vaccinat	ed in the 2^{nd}		
	trimester and	d 59 calves	s were from co	ows vaccing	ated in the 3 rd		
	trimester 6 serum samples were removed from the study due to						
	equipment malfunction or concerns that colostrum was received						
	All valid sar	nples tested	l negative for a	antibodies to	BR. BVD1		
	and BVD2.	Serum sam	ples were also	negative for	IBR by virus		
	isolation and	negative fo	r BVD1 and BV	/D2 by PCR			

USDA Approval Date	January 11, 2008