

Summary of Studies Supporting USDA Product Licensure

Establishment Name	Boehringer Ingelheim Vetmedica, Inc.
USDA Vet Biologics Establishment Number	124
Product Code	47A1.20
True Name	Canine Parainfluenza Vaccine, Modified Live Virus, Bordetella Bronchiseptica Bacterin
Tradename(s) / Distributor or Subsidiary (if different from manufacturer)	
Date of Compilation Summary	December 06, 2017

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	Bordetella bronchiseptica (Bb)
Study Purpose	Pivotal Efficacy against respiratory disease due to Bb
Product Administration	Two doses administered subcutaneously 21 days apart
Study Animals	30 Dogs, 7 weeks old, randomly divided into 15 vaccinates and 15 placebo controls
Challenge Description	Bb was administered 14 days after the last vaccination
Interval observed after challenge	Dogs were observed for clinical signs 2x daily for 14 days post challenge.
Results	Dogs were considered affected by Bb challenge if coughing was observed for two or more days post challenge. Positive Bb Clinical results: Vaccinates: 4/15 (27%) positive Controls: 14/15 (93%) positive See the attached table with the clinical observations for coughing.
USDA Approval Date	April 4, 2013

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Individual Clinical Scores for Coughing

							Experim	ental Vac	Experimental Vaccine Group	dr							
Dog	-2DPC	-1DPC	ODPC	1DPC	2DPC	3DPC	4DPC	SDPC	6DPC	7DPC	8DPC	9DPC	10DPC	11DPC	12DPC	13DPC	14DPC
1	0	0	0	0	0	0	0	1	0	0	0	0	0	0	0	0	0
2	0	0	0	0	0	1	1	0	1	1	0	0	1	0	0	1	0
3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	1	0	0
4	0	0	0	0	0	0	1	1	1	1	1	1	1	1	1	0	0
5	0	0	0	0	0	0	0	0	1	0	0	0	0	0	0	0	0
9	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
7	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
8	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
6	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
10	0	0	0	0	0	0	0	1	1	0	0	0	0	0	0	0	0
11	0	0	0	0	0	0	0	0	1	0	0	0	0	0	0	0	0
12	0	0	0	0	0	0	0	1	0	0	0	0	0	0	0	0	0
13	0	0	0	0	0	0	0	0	1	0	0	0	0	0	0	0	0
14	0	0	0	0	0	0	0	0	1	0	0	0	0	0	0	0	0
15	0	0	0	0	0	0	0	1	1	0	0	0	0	0	0	0	0
							Place	Placebo Control Group	ol Group								
Dog	-2DPC	-1DPC	ODPC	1DPC	2DPC	3DPC	4DPC	SDPC	6DPC	7DPC	8DPC	9DPC	10DPC	11DPC	12DPC	13DPC	14DPC
16	0	0	0	0	0	0	0	1	1	0	0	0	0	0	0	0	0
17	0	0	0	0	1	1	1	0	1	0	0	0	0	0	1	0	0
18	0	0	0	0	0	1	1	1	1	0	0	0	0	0	0	1	0
19	0	0	0	0	0	0	0	1	1	0	1	0	0	0	0	0	0
20	0	0	0	0	0	1	1	1	1	1	1	1	1	0	0	0	0
21	0	0	0	0	0	0	1	1	1	1	0	0	0	0	0	0	0
22	0	0	0	0	1	1	1	1	1	1	1	1	1	1	1	1	1
23	0	0	0	0	0	1	0	0	0	0	0	0	0	0	0	0	0
24	0	0	0	0	0	1	0	1	0	0	1	0	0	0	0	0	0
25	0	0	0	0	0	0	1	1	1	1	1	1	0	0	0	0	0
26	0	0	0	0	0	0	1	1	1	0	1	1	0	1	1	0	1
27	0	0	0	0	1	1	1	1	1	1	1	1	0	1	1	1	0
28	0	0	0	0	0	1	1	1	1	0	1	1	1	1	1	0	0
29	0	0	0	0	0	0	0	0	0	0	1	1	0	0	0	0	0
30	0	0	0	0	0	0	1	1	1	1	0	0	0	0	0	0	0
1 - cough observed	erved																
0 - no cough observed	perved																
* Dogs were considered affected by Bb challenge if coughing was observed for two or more days post challenge.	onsidere	daffected	by 8b ch	allenge if	coughing	was obs	erved for	two or m	ore days	post cha	lenge.						

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Study Type	Efficacy
Pertaining to	Canine Parainfluenza (CPI)
Study Purpose	Efficacy
Product Administration	
Study Animals	
Challenge Description	
Interval observed after challenge	
Results	Study data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance. Study data, however, are no longer available.

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Study Type	Safety							
Pertaining to	ALL							
Study Purpose	Demonstrate safety of	'product u	ındar tyni	cal usa co	nditions			
Product	2 Doses administered a	_						
Administration	2 Doses administrated a	ii a 3 wccs	inicivai	by the SQ	Toute.			
Study Animals	628 privately owned c	anines we	re includ	ed in the t	final analy	veie Mor	e than	
Study Allillais	one-third of the canine							
	days of age) at the tim				idy were _	o week	, (<u> </u>	
	639 Total dogs were e				ete the sti	ndv		
Challenge	NA	monea oc		iot compi	oto the st	aay.		
Description								
Observation	Canines were observed for 30 min following the first vaccination and daily							
interval after	till the second vaccina			_			•	
last treatment	following the second							
Results	Frequency of adverse		<u> </u>					
		Number	Percent	Number	Percent	Total	Percent	
	Adverse Event	≤ 59	≤ 59	> 59	> 59	number	of all	
	X 1	days old	days old	days old	days old		animals	
	No adverse events	157	73.36	374	90.34	531	84.55	
	Diarrhea*	50	23.36	11	2.66	61	9.71	
	Gastroenteritis*	24	11.21	4	0.97	28	4.46	
	Injection site lump	3	1.40	10	2.42	13	2.07	
	Depression	8	3.74	1	0.24	9	1.43	
	Anorexia	8	3.74	0	0.00	8	1.27	
	Decreased appetite	4	1.87	4	0.97	8	1.27	
	Not drinking	8	3.74	0	0.00	8	1.27	
	Mortality Affirmed by licensee to have probable cause other than vaccination	4	1.87	2	0.48	6	0.96	
	Injection site pain	4	1.87	1	0.24	5	0.80	
	Injection site granuloma	0	0.00	4	0.97	4	0.64	
	Abdominal pain	3	1.40	0	0.00	3	0.48	
	Cough	0	0.00	3	0.72	3	0.48	
	Hypersalivation	3	1.40	0	0.00	3	0.48	
	Hyperactivity	0	0.00	2	0.48	2	0.32	
	Aggression	0	0.00	1	0.24	1	0.16	
	Corneal edema	0	0.00	1	0.24	1	0.16	
	Digestive tract disorder NOS	1	0.47	0	0.00	1	0.16	
	Fever	0	0.00	1	0.24	1	0.16	
	Fungal skin infection NOS	1	0.47	0	0.00	1	0.16	
	Hot spot (pyotraumatic dermatitis)	0	0.00	1	0.24	1	0.16	

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	Injection site abscess	0	0.00	1	0.24	1	0.16
	Joint pain	0	0.00	1	0.24	1	0.16
	Local swelling (not application site)	0	0.00	1	0.24	1	0.16
	Miscellaneous eating disorder NOS	0	0.00	1	0.24	1	0.16
	Nasal discharge	1	0.47	0	0.00	1	0.16
	Ocular discharge	0	0.00	1	0.24	1	0.16
	Polydipsia	0	0.00	1	0.24	1	0.16
	Skin swelling	0	0.00	1	0.24	1	0.16
	Sneezing	0	0.00	1	0.24	1	0.16
	Tremor	0	0.00	1	0.24	1	0.16
	Weakness	0	0.00	1	0.24	1	0.16
	* 78 animals had confirm						
	gastroenteritis not attribut	able to vac	cination (se	veral anima	ıls had mor	e than one c	lisease).
USDA	February 28, 2017						
Approval Date							

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