



Summary of Studies Supporting USDA Product Licensure

Establishment Name	Elanco US Inc.
USDA Vet Biologics Establishment Number	196
Product Code	4X59.R1
True Name	Bovine Rhinotracheitis-Virus Diarrhea-Parainfluenza 3-Respiratory Syncytial Virus Vaccine, Modified Live Virus, Mannheimia Haemolytica Bacterial Extract-Toxoid
Tradename(s) / Distributor or Subsidiary (if different from manufacturer)	Nuplura PH + 5 - Elanco US Inc.
Date of Compilation Summary	September 17, 2024

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	Comparative Serology
Pertaining to	Infectious Bovine Rhinotracheitis (IBR) Virus, Bovine Viral Diarrhea (BVD) Virus Type 1, BVD Type 2, and Parainfluenza 3 (PI3) Virus
Study Purpose	Comparative serology to demonstrate lack of interference between the IBR, BVD Type 1, BVD Type 2, and PI3 viral fractions and the Mannheimia Haemolytica bacterial extract when the antigens are combined to show that combining fractions with satisfactory efficacy studies does not reduce the immune response.
Product Administration	One dose of combination test vaccine administered subcutaneously to cattle, followed by a second dose of monovalent Bovine Respiratory Syncytial Virus (BRSV) 21 days later.
Study Animals	9 month old calves were vaccinated twice 21 days apart. Animals were seronegative to BVD Type 1, BVD type 2, IBR and PI3. Forty-five animals were vaccinated with the combination test vaccine (BVD1-BVD2-IBR-PI3-BRSV-M Haem) and 45 animals were vaccinated with the reference vaccine (BVD1-BVD2-IBR- PI3-BRSV).
Challenge Description	NA
Interval observed after challenge	Blood was collected on study days D0, D28, D35
Results	The primary outcome was neutralizing antibody serum titer for the BVD type 1, BVD type 2, PI3, and IBR antigens. The study met the criterion in which the test product Geometric Mean Titer (GMT) must be at least 63% of the reference product GMT. Raw data on pages below
USDA Approval Date	November 18, 2016

BVD Type 1 Serum Neutralization Titers

<i>Combination Test Vaccine</i>		
Calf ID	Study Day 0	Study Day 28
1	<2	362
2	<2	1024
3	<2	362
4	<2	790
5	<2	912
6	<2	1312
7	<2	664
8	<2	1117
9	<2	821
10	<2	664
11	<2	575
12	<2	1149
13	<2	724
14	<2	1448
15	<2	1149
16	<2	2702
17	<2	861
18	<2	1328
19	<2	3158
20	<2	1218
21	<2	724
22	<2	1579
23	<2	1449
24	<2	724
25	<2	821
26	<2	912
27	<2	411
28	<2	1643
29	<2	279
30	<2	1117
31	<2	2435
32	<2	1218
33	<2	2048
34	<2	1825
35	<2	1149
36	<2	323
37	<2	1277
38	<2	319
39	<2	1024
40	<2	512
41	<2	1652
42	<2	1448
43	<2	1117
44	<2	1579
45	<2	1290

<i>Reference Vaccine</i>		
Calf ID	Study Day 0	Study Day 28
1	<2	91
2	<2	724
3	<2	1328
4	<2	2435
5	<2	1328
6	<2	456
7	<2	664
8	<2	342
9	<2	1149
10	<2	575
11	<2	724
12	<2	<2
13	<2	724
14	<2	72
15	<2	790
16	<2	1148
17	<2	638
18	<2	456
19	<2	395
20	<2	664
21	<2	744
22	<2	181
23	<2	1328
24	<2	1448
25	<2	724
26	<2	332
27	<2	458
28	<2	1448
29	<2	456
30	<2	558
31	<2	645
32	<2	664
33	<2	206
34	<2	664
35	<2	724
36	<2	1290
37	<2	1149
38	<2	861
39	<2	889
40	<2	878
41	<2	1579
42	<2	458
43	<2	821
44	<2	2048
45	<2	256

BVD Type 2 Serum Neutralization Titers

<i>Combination Test Vaccine</i>		
Calf ID	Study Day 0	Study Day 28
1	<2	29
2	<2	57
3	<2	14
4	<2	51
5	<2	205
6	<2	197
7	<2	13
8	<2	112
9	<2	114
10	<2	36
11	<2	21
12	<2	203
13	<2	45
14	<2	152
15	<2	160
16	<2	72
17	<2	40
18	<2	41
19	<2	91
20	<2	28
21	<2	10
22	<2	12
23	<2	38
24	<2	166
25	<2	29
26	<2	16
27	<2	36
28	<2	57
29	<2	54
30	<2	45
31	<2	219
32	<2	14
33	<2	51
34	<2	41
35	<2	14
36	<2	20
37	<2	31
38	<2	31
39	<2	83
40	<2	20
41	<2	41
42	<2	45
43	<2	18
44	<2	97
45	<2	37

<i>Reference Vaccine</i>		
Calf ID	Study Day 0	Study Day 28
1	<2	181
2	<2	8
3	<2	128
4	<2	16
5	<2	25
6	<2	51
7	<2	11
8	<2	9
9	<2	5
10	<2	114
11	<2	51
12	<2	40
13	<2	25
14	<2	64
15	<2	11
16	<2	40
17	<2	51
18	<2	27
19	<2	2
20	<2	81
21	<2	40
22	<2	144
23	<2	83
24	<2	10
25	<2	76
26	<2	7
27	<2	128
28	<2	16
29	<2	41
30	<2	102
31	<2	181
32	<2	36
33	<2	3
34	<2	9
35	<2	205
36	<2	11
37	<2	23
38	<2	51
39	<2	80
40	<2	103
41	<2	29
42	<2	140
43	<2	11
44	<2	149
45	<2	29

IBR Serum Neutralization Titers

<i>Combination Test Vaccine</i>		
Calf ID	Study Day 0	Study Day 28
1	<2	14
2	<2	26
3	<2	41
4	<2	83
5	<2	36
6	<2	38
7	<2	45
8	<2	29
9	<2	49
10	<2	41
11	<2	35
12	<2	23
13	<2	29
14	<2	36
15	<2	23
16	<2	72
17	<2	29
18	<2	72
19	<2	26
20	<2	41
21	<2	45
22	<2	80
23	<2	41
24	<2	12
25	<2	80
26	<2	49
27	<2	41
28	<2	45
29	<2	29
30	<2	32
31	<2	23
32	<2	80
33	<2	57
34	<2	55
35	<2	41
36	<2	38
37	<2	49
38	<2	51
39	<2	49
40	<2	26
41	<2	25
42	<2	35
43	<2	41
44	<2	76
45	<2	83

<i>Reference Vaccine</i>		
Calf ID	Study Day 0	Study Day 28
1	<2	20
2	<2	11
3	<2	8
4	<2	11
5	<2	20
6	<2	36
7	<2	49
8	<2	18
9	<2	18
10	<2	29
11	<2	21
12	<2	4
13	<2	18
14	<2	14
15	<2	23
16	<2	13
17	<2	11
18	<2	12
19	<2	29
20	<2	18
21	<2	36
22	<2	7
23	<2	21
24	<2	36
25	<2	25
26	<2	12
27	<2	45
28	<2	20
29	<2	29
30	<2	23
31	<2	45
32	<2	29
33	<2	45
34	<2	26
35	<2	91
36	<2	23
37	<2	45
38	<2	18
39	<2	29
40	<2	36
41	<2	14
42	<2	23
43	<2	38
44	<2	23
45	<2	18

PI3 Serum Neutralization Titers

<i>Combination Test Vaccine</i>		
Calf ID	Study Day 0	Study Day 28
1	<2	29
2	<2	114
3	<2	5
4	<2	197
5	<2	49
6	<2	23
7	<2	45
8	<2	181
9	<2	72
10	<2	49
11	<2	23
12	<2	49
13	<2	41
14	<2	91
15	<2	12
16	<2	99
17	<2	144
18	<2	72
19	<2	144
20	<2	72
21	<2	72
22	<2	83
23	<2	57
24	<2	45
25	<2	57
26	<2	9
27	<2	41
28	<2	25
29	<2	45
30	<2	23
31	<2	72
32	<2	45
33	<2	91
34	<2	36
35	<2	72
36	<2	25
37	<2	45
38	<2	83
39	<2	91
40	<2	29
41	<2	49
42	<2	41
43	<2	51
44	<2	83
45	<2	91

<i>Reference Vaccine</i>		
Calf ID	Study Day 0	Study Day 28
1	<2	41
2	<2	<2
3	<2	72
4	<2	45
5	<2	36
6	<2	3
7	<2	5
8	<2	3
9	<2	26
10	<2	49
11	<2	45
12	<2	<2
13	<2	23
14	<2	45
15	<2	8
16	<2	91
17	<2	29
18	<2	3
19	<2	45
20	<2	83
21	<2	36
22	<2	23
23	<2	32
24	<2	10
25	<2	3
26	<2	6
27	<2	36
28	<2	6
29	<2	36
30	<2	6
31	<2	41
32	<2	3
33	<2	29
34	<2	11
35	<2	3
36	<2	18
37	<2	9
38	<2	3
39	<2	36
40	<2	20
41	<2	5
42	<2	41
43	<2	91
44	<2	72
45	<2	45

Study Type	Efficacy																								
Pertaining to	<i>Mannheimia Haemolytica</i>																								
Study Purpose	Demonstrate the efficacy of a combination product (IBR, BVD Type 1, BVD Type 2, BRSV, and PI3 combined with Mannheimia Haemolytica in Combination test vaccine) in protection against <i>M. haemolytica</i> as a means to demonstrate absence of interference of IBR, BVD Type 1, BVD Type 2, and PI3 to <i>M. Haemolytica</i> immunity to show that combining fractions with satisfactory efficacy studies does not reduce the immune response.																								
Product Administration	One dose of combination vaccine administered subcutaneously																								
Study Animals	63 Calves, 80-82 days of age, randomly distributed into three treatment groups. One group of 21 Vaccinates (Combination test vaccine), one group of 21 reference vaccinates (Reference vaccine – Viral fractions only), and one group of 21 placebo controls.																								
Challenge Description	<i>Mannheimia Haemolytica</i> , administered 10 days after vaccination																								
Interval observed after challenge	Lungs evaluated 4 days after challenge																								
Results	The percent of the lung mass that was abnormal (consolidated) was calculated for every animal. 5-number summary for lung consolidation (%) <table><tr><th><i>Treatment</i></th><th><i>Minimum</i></th><th><i>Q1</i></th><th><i>Median</i></th><th><i>Q3</i></th><th><i>Maximum</i></th></tr><tr><td>Combination Test vaccinates</td><td>0.2</td><td>2.6</td><td>5.6</td><td>8.2</td><td>26</td></tr><tr><td>Reference Vaccinates</td><td>2.1</td><td>7.9</td><td>24.1</td><td>24.6</td><td>90.4</td></tr><tr><td>Placebo Controls</td><td>1.4</td><td>9.5</td><td>17.6</td><td>32.5</td><td>84.8</td></tr></table> Raw data shown on attached page.	<i>Treatment</i>	<i>Minimum</i>	<i>Q1</i>	<i>Median</i>	<i>Q3</i>	<i>Maximum</i>	Combination Test vaccinates	0.2	2.6	5.6	8.2	26	Reference Vaccinates	2.1	7.9	24.1	24.6	90.4	Placebo Controls	1.4	9.5	17.6	32.5	84.8
<i>Treatment</i>	<i>Minimum</i>	<i>Q1</i>	<i>Median</i>	<i>Q3</i>	<i>Maximum</i>																				
Combination Test vaccinates	0.2	2.6	5.6	8.2	26																				
Reference Vaccinates	2.1	7.9	24.1	24.6	90.4																				
Placebo Controls	1.4	9.5	17.6	32.5	84.8																				
USDA Approval Date	September 16, 2016																								

Lung Consolidation (%), in order of rank:

Combination Test vaccinates	Reference Vaccinates	Placebo Controls
0.21	2.14	1.44
1.8	3.1	2.64
1.94	4.47	3.56
2.18	4.54	4.3
2.29	4.99	7.36
2.57	7.94	9.53
2.81	8.39	9.55
3.49	8.87	10.23
3.51	14.65	15.23
5.4	22.95	15.98
5.57	24.05	17.59
5.64	24.36	17.66
6.51	28.45	23.77
6.51	30.95	23.84
6.67	33.95	26.75
8.15	34.55	32.45
10.99	37.3	34.1
11.35	45.9	37.25
13	48.73	38.4
16.57	58.2	39.05
26.01	90.4	84.75

Study Type	Efficacy																		
Pertaining to	Mannheimia haemolytica																		
Study Purpose	To demonstrate effectiveness against respiratory disease caused by M. Haemolytica as early as 10 days after administration.																		
Product Administration	1 dose subcutaneously																		
Study Animals	30 calves, 75-83 days of age. 20 vaccinates and 10 controls.																		
Challenge Description	Mannheimia Haemolytica, administered 10 days after second vaccination																		
Interval observed after challenge	Lungs evaluated 4 days after challenge																		
Results	The percent of the lung mass that was abnormal (consolidated) was calculated for every animal. 5-number summary for lung consolidation (%) <table><tr><td>Treatment</td><td>Minimum</td><td>Q1</td><td>Median</td><td>Q3</td><td>Maximum</td></tr><tr><td>Controls</td><td>4.6</td><td>20.6</td><td>23</td><td>52.3</td><td>57.8</td></tr><tr><td>Vaccinates</td><td>1.0</td><td>6.5</td><td>9.9</td><td>14.7</td><td>41.3</td></tr></table> Raw data shown on attached page.	Treatment	Minimum	Q1	Median	Q3	Maximum	Controls	4.6	20.6	23	52.3	57.8	Vaccinates	1.0	6.5	9.9	14.7	41.3
Treatment	Minimum	Q1	Median	Q3	Maximum														
Controls	4.6	20.6	23	52.3	57.8														
Vaccinates	1.0	6.5	9.9	14.7	41.3														
USDA Approval Date	June 5, 2009																		

Lung Consolidation scores (%), in order of rank:

Vaccinate	Control
0.97	4.61
1.69	13.35
3.66	20.63
4.45	20.65
5.95	20.66
7.13	25.4
7.2	26.35
7.25	52.25
7.35	55.85
9.48	57.75
10.25	
10.93	
11.21	
12.26	
12.86	
16.45	
16.59	
20.4	
38.25	
41.3	

Study Type	Efficacy																		
Pertaining to	Bovine Respiratory Syncytial Virus (BRSV)																		
Study Purpose	Demonstrate the efficacy of a combination product (IBR, BVD Type 1, BVD Type 2, BRSV, and PI3 combined with Mannheimia Haemolytica in Combination test vaccine) in protection against BRSV as a means to demonstrate absence of interference of IBR, BVD Type 1, BVD Type 2, and PI3 to BRSV immunity to show that combining fractions with satisfactory efficacy studies does not reduce the immune response																		
Product Administration	One dose of combination test vaccine administered subcutaneously to cattle, followed by a second dose of monovalent BRSV 21 days later.																		
Study Animals	Twenty-nine (29) 88-to-90-day old colostrum-deprived Holstein cross-calves, steers, and heifers; 14 vaccinates and 15 controls.																		
Challenge Description	Twenty-one (21) and 22 days after 2nd vaccination, calves were challenged with BRSV.																		
Interval observed after challenge	Calves were observed daily for clinical signs for eight days then lung lesions evaluated.																		
Results	The primary outcome was lung lesion scores summarized in Table 1. Table 1: Five Number Summary (Minimum = Min, 25th Percentile = Q1, Median = Med, 75th Percentile = Q3, and Maximum = Max) of Percent of Pneumonic Tissue by Treatment Group <table><tr><th><i>Group (Number of calves)</i></th><th><i>Min</i></th><th><i>Q1</i></th><th><i>Med</i></th><th><i>Q3</i></th><th><i>Max</i></th></tr><tr><td><i>Control group</i></td><td>0</td><td>13</td><td>15</td><td>19</td><td>37</td></tr><tr><td><i>Vaccinate group</i></td><td>0</td><td>0</td><td>1</td><td>1</td><td>5</td></tr></table> 1 calf was removed between the first and second vaccination Raw data found on the next page.	<i>Group (Number of calves)</i>	<i>Min</i>	<i>Q1</i>	<i>Med</i>	<i>Q3</i>	<i>Max</i>	<i>Control group</i>	0	13	15	19	37	<i>Vaccinate group</i>	0	0	1	1	5
<i>Group (Number of calves)</i>	<i>Min</i>	<i>Q1</i>	<i>Med</i>	<i>Q3</i>	<i>Max</i>														
<i>Control group</i>	0	13	15	19	37														
<i>Vaccinate group</i>	0	0	1	1	5														
USDA Approval Date	June 21, 2024																		

Animal ID	Group	Lung Lesion Score							
		Right Cranial Lobe	Right Posterior Cranial Lobe	Right Middle Lobe	Right Caudal Lobe	Left Cranial Lobe	Left Posterior Cranial Lobe	Left Caudal Lobe	Right Accessory Lobe
919	Control	7	0	0	0	0	0	0	0
920	Control	60	10	20	8	10	20	10	25
929	Control	10	3	10	7	10	12	15	10
939	Control	40	15	20	10	10	45	20	20
944	Control	12	20	30	20	7	15	20	25
951	Control	15	12	85	12	15	90	15	15
962	Control	20	15	15	10	15	15	15	15
967	Control	30	15	20	12	10	30	12	10
974	Control	20	15	25	10	10	15	15	25
978	Control	98	40	80	15	10	75	25	10
981	Control	85	15	85	25	20	90	20	85
993	Control	75	20	20	0	15	25	10	0.5
994	Control	15	7	12	15	10	10	12	10
1002	Control	25	10	20	15	12	30	15	15
1011	Control	45	7	10	20	7	15	15	20
922	Vaccinate	10	0	0	0	0	0	0	0
943	Vaccinate	1	0	0	0	0	0	0	0
952	Vaccinate	7	0	0	0	0	3	0	0
958	Vaccinate	11	0	0	0	0	0	0	7
959	Vaccinate	0	0	0	0	0	0	0	0
961	Vaccinate	20	0	5	0	0	0	0	0
973	Vaccinate	10	3	0	0	0	0	0	0.5
975	Vaccinate	NA	0	0	0	NA	NA	NA	3
976	Vaccinate	10	0	0	0	0	0	0	0
980	Vaccinate	10	0	0	0	0	0	0	0
984	Vaccinate	20	0	0	0	5	1	0	0
988	Vaccinate	0	0	0	0	0.5	0	0.5	0
992	Vaccinate	0	0	0	0	0	0	0	0
996	Vaccinate	30	7	20	0	1	10	0	10

Study Type	Safety																																				
Pertaining to	All																																				
Study Purpose	Demonstrate safety of product under typical use conditions																																				
Product Administration	One dose administered subcutaneously, followed by a monovalent bovine respiratory syncytial virus (BRSV) injection 2 weeks later.																																				
Study Animals	627 calves, 18-29 days of age at each of three sites.																																				
Challenge Description	NA																																				
Interval observed after challenge	No challenge. Animals were observed at 2 hours after each injection and daily for a minimum of 42 days after vaccination.																																				
Results	All Adverse Events: <table> <tr> <th>VeDDRA Term</th><th>Number of Calves</th></tr> <tr> <td>No Adverse Events</td><td>329</td></tr> <tr> <td>Injection Site Swelling</td><td>259</td></tr> <tr> <td>Increased Respiratory Rate</td><td>39</td></tr> <tr> <td>Death*</td><td>8</td></tr> <tr> <td>Anorexia</td><td>4</td></tr> <tr> <td>Fever</td><td>13</td></tr> <tr> <td>Hypothermia</td><td>3</td></tr> <tr> <td>Arthritis</td><td>1</td></tr> <tr> <td>Conjunctivitis</td><td>1</td></tr> <tr> <td>Diarrhea</td><td>14</td></tr> <tr> <td>Labored Breathing</td><td>1</td></tr> <tr> <td>Cough Productive</td><td>1</td></tr> <tr> <td>Respiratory Distress</td><td>5</td></tr> <tr> <td>Lameness</td><td>2</td></tr> <tr> <td>Pink Eye</td><td>11</td></tr> <tr> <td>Bloated</td><td>1</td></tr> <tr> <td>Lethargy</td><td>2</td></tr> </table> *Deaths not attributable to vaccine	VeDDRA Term	Number of Calves	No Adverse Events	329	Injection Site Swelling	259	Increased Respiratory Rate	39	Death*	8	Anorexia	4	Fever	13	Hypothermia	3	Arthritis	1	Conjunctivitis	1	Diarrhea	14	Labored Breathing	1	Cough Productive	1	Respiratory Distress	5	Lameness	2	Pink Eye	11	Bloated	1	Lethargy	2
VeDDRA Term	Number of Calves																																				
No Adverse Events	329																																				
Injection Site Swelling	259																																				
Increased Respiratory Rate	39																																				
Death*	8																																				
Anorexia	4																																				
Fever	13																																				
Hypothermia	3																																				
Arthritis	1																																				
Conjunctivitis	1																																				
Diarrhea	14																																				
Labored Breathing	1																																				
Cough Productive	1																																				
Respiratory Distress	5																																				
Lameness	2																																				
Pink Eye	11																																				
Bloated	1																																				
Lethargy	2																																				

	Injection Site Reactions, Size:								
	Site	Total number animals	Vaccination	# of injection site reactions by Size (cm ²)			Injection site reactions not observed		
				≤ 2.0 cm ²	2.0-10.0 cm ²	>10 cm ²			
	1	207	4X59.R1	4	0	0	203		
			BRSV	0	0	0	207		
	2	210	4X59.R1	1	50	69	90		
			BRSV	0	0	0	210		
	3	210	4X59.R1	95	32	0	83		
			BRSV	23	0	0	187		
	Injection Site Reactions, Duration:								
	Sites	Total number animals	Total # Animals with Reactions	Vaccination	Five Number Summary (Days)				
					Min	Q1	Med	Q3	Max
	1	207	4	4X59.R1	6.0	7.0	8.0	8.0	8.0
				BRSV	NA	-	-	-	-
	2	210	120	4X59.R1	1.0	2.0	13.0	21.0	50.0*
				BRSV	NA	-	-	-	-
	3	210	127	4X59.R1	1.0	1.0	8.0	14.0	28.0
				BRSV	1.0	1.0	1.0	1.0	8.0
	* Six reactions were not resolved by study day 50. All were reported to be less than 2 cm ² at the end of the study.								
	USDA Approval Date		April 10, 2020						