

Summary of Studies Supporting USDA Product Licensure

Establishment Name	Diamond Animal Health, Inc.			
USDA Vet Biologics Establishment Number	213			
Product Code	1155.20			
True Name	Bovine Rhinotracheitis-Virus Diarrhea Vaccine, Killed Virus			
Tradename(s) / Distributor or Subsidiary (if different from manufacturer)	Diamond Animal Health, Inc. Master Guard 3 - Elanco Animal Health - Diamond Animal Health, Inc.			
Date of Compilation Summary	December 20, 2023			

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 Virus Diarrhea (BVD) Type 1			
Study Purpose	To demonstrate efficacy of against BVD virus Type 1			
Product Administration	Two doses by subcutaneous or intramuscular injection to healthy,			
	susceptible cattle			
Study Animals	Bovine			
Challenge Description	BVD Type 1, NY-1 Strain (CVB-L lot 97-12)			
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 27, 1999			

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Study Type	Efficacy				
Pertaining to	Bovine Virus Diarrhea (BVD) Type 2				
Study Purpose	To demonstrate efficacy of product for BVD virus Type 2				
Product Administration	Two doses by subcutaneous or intramuscular injection				
Study Animals	Bovine				
Challenge Description	BVD virus Type 2, 890 Strain (lot 1444-57, from CVB-L lot 91-3)				
Interval observed after	14 days				
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 27, 1999				

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Study Type	Efficacy			
Pertaining to	Infectious Bovine Rhinotracheitis (IBR)			
Study Purpose	To demonstrate efficacy of product for IBR virus			
Product Administration	Two doses by intramuscular or subcutaneous injection			
Study Animals	Bovine			
Challenge Description	Cooper Challenge Strain IBR Virus Lot 50980			
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 15, 1986			

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Study Type	Safety				
Pertaining to	ALL				
Study Purpose	Demonstrate safety of product under typical use conditions				
Product Administration	Two doses 14 days apart subcutaneously or intramuscularly				
Study Animals	Bovine				
Challenge Description	NA				
Interval observed after	NA				
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 12, 1999				

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Study Type	Safety					
Pertaining to	ALL					
Study Purpose	To demonstrate safety in pregnant cows					
Product	Two doses administered	Two doses administered subcutaneously to heifers and cows, 1				
Administration	dose prior to breeding a	nd 1 dose during preg	nancy at different			
	trimesters. Heifers and o	cows were confirmed	to be pregnant at			
	administration during pr	administration during pregnancy.				
Study Animals	Heifers and cows-separate groups vaccinated during each trimester.					
	Similar sized groups in each trimester were maintained as controls.					
Challenge Description	NA					
Interval observed after	Heifers and cows observ	ved from the pre-breed	ling vaccination to			
challenge	post-breeding vaccination	post-breeding vaccination.				
Results						
	First Trimester (<93 day	First Trimester (<93 days of gestation)				
		Vaccinate	Control			
	Enrolled	209	213			
	Excluded (not related	1	0			
	to vaccination)					
	Second Trimester (94-1	87 days of gestation)				
		Vaccinate	Control			
	Enrolled	315	310			
	Excluded (not related	2	2			
	to vaccination)					
	Third Trimester (188-250 days of gestation)					
		Vaccinate	Control			
	Enrolled	205	208			
	No adverse events were reported.					
	N. 1.5.2012					
USDA Approval Date	March 5, 2013					

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