

Summary of Studies Supporting USDA Product Licensure

Establishment Name	Zoetis Inc.
USDA Vet Biologics Establishment Number	190
Product Code	7430.00
True Name	Clostridium Chauvoei-Septicum-Novyi-Sordellii-Perfringens Types C & D-Mannheimia Haemolytica Bacterin-Toxoid
Tradename(s) / Distributor or Subsidiary (if different from manufacturer)	One Shot Ultra 7 - No distributor specified One Shot Ultra 7 - Zoetis South Africa Ltd
Date of Compilation Summary	July 05, 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	Clostridium chauvoei
Study Purpose	Demonstrate effectiveness against Clostridium chauvoei
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	February 3, 1999

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Study Type	Efficacy
Pertaining to	Clostridium novyi
Study Purpose	Demonstrate effectiveness against Clostridium novyi
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	February 3, 1999

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Study Type	Efficacy
Pertaining to	Clostridium perfringens Type C
Study Purpose	Demonstrate effectiveness against Clostridium perfringens Type
	C
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	February 3, 1999

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Study Type	Efficacy
Pertaining to	Clostridium perfringens Type D
Study Purpose	Demonstrate effectiveness against Clostridium perfringens Type
-	D
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	February 3, 1999

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Study Type	Efficacy
Pertaining to	Clostridium septicum
Study Purpose	Demonstrate effectiveness against Clostridium septicum
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	February 3, 1999

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Study Type	Efficacy
Pertaining to	Clostridium sordelli
Study Purpose	Demonstrate effectiveness against Clostridium sordelli
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	February 3, 1999

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Study Type	Efficacy
Pertaining to	Mannheimia haemolytica
Study Purpose	Demonstrates effectiveness against Mannheimia haemolytica
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	June 21, 1991

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Study Type	Efficacy							
Pertaining to	Mannheimia haemolytica							
Study Purpose	Demonstrate efficacy against respiratory disease caused by							
_	Mannheimia haemolytica							
Product Administration	One dose administered subcutaneously							
Study Animals	9 vaccinates	s and 11 co	ntrols calve	es				
Challenge Description	M. haemoly	<i>tica</i> admin	istered 14 d	lays post-v	accination			
Interval observed after	Animals were observed daily from days 12 through 18 post-							
challenge	challenge then lung lesions were evaluated.							
Results	Lung lesions (%) was the efficacy variable.							
	Lung lesion							
		Minimum	25 th	Median	75 th	Maximum		
			Percentile		Percentile			
	Placebo	1.6	5.6	9.5	17.1	92.9		
	Vaccinates	0.4	1.0	1.7	2.6	14.8		
	Mortality:							
	Controls: 2/	/11 (18.2%))					
	Vaccinates:	0/9 (0%)						
		` ′						
	See individ	ual data att	ached					
USDA Approval Date	08/20/1999	-	-	-	-			

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Lung Lesions and Mortality by Treatment and Animal

Controls

Mortality Animal Lung Lesions (%) Post-Challenge 281 9.7 No 600 No 10.1 603 No 4.9 17.1 605 No 606 27.0 Yes 610 92.9 Yes 615 7.7 No 616 9.2 No 618 **5.6** No **627** 1.6 No 628 9.5 No

Vaccinates

Animal	Lung	Mortality
	Lesions (%)	Post-
		Challenge
269	0.4	No
279	1.0	No
287	2.1	No
297	1.7	No
601	0.4	No
602	2.6	No
604	1.0	No
609	9.5	No
625	14.8	No

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Study Type	Safety							
Pertaining to	ALL							
Study Purpose	Demonstrate safety in cattle under field conditions							
Product Administration	Two doses admir	nistered su	ıbcutan	eously	28 day	ys apar	t	
Study Animals	200 crossbred be	ef heifers						
Challenge Description	NA	NA						
Interval observed after	Calves were obse	erved for o	ne hou	ır after	vaccin	ation a	and dai	ly
challenge	throughout the st	tudy for ac	lverse e	events.	Anima	als wer	e exam	nined
	for injection site	reactions	on day	s 14, 2	8, and	42 pos	t-vacci	nation
	(left side) and da	y 14 post-	vaccin	ation (right si	de).		
Results	Number of Animals with Adverse Events							
						•		-
	a g s							
			sal ıarg	rhea	atic	nes	esce	
	Nasal Discharge Diarrhea Respiration Rate Lameness Leg Abscess							
	Vaccinate	es	55	1	1	1	1	-
	All observations were	affirmed by 1	icensee to	be due	to causes	other tha	an vaccin	ation.
	Number of Anim	nals with I	njectio	n Site	Reaction	<u>ons</u>		
	Vaccinates	1.4	Da	ys Post	t-Vaccin	<u>ation</u>	42	
	Left Side	14 129			28 41		42 13	
	Left Side 129 41 13 Right Side 153 - -							
	All injection site reactions were transient and under 4 cm in diameter.							
USDA Approval Date	09/08/1999							

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Study Type	Safety						
Pertaining to	ALL						
Study Purpose	Demonstrate safety in cattle under field conditions						
Product Administration	Two doses adm	inistered su	ıbcutaı	neously	28 days	s apart	
Study Animals	196 crossbred b	196 crossbred beef heifers					
Challenge Description	NA						
Interval observed after	Calves were obs	served for o	one ho	ur after	vaccina	tion and daily	
challenge	throughout the s	study for ac	lverse	events.	Animal	s were examined	
	for injection site	e reactions	on day	s 14, 2	8, and 42	2 post-	
	vaccination (lef	t side) and	day 14	post-v	accinati	on (right side).	
Results	Number of Anii	mals with A	Advers	e Event	<u>s</u>		
			1	1			
			SS		ion		
			ene	eye	ess		
			Tameness	Pink eye	Depression		
	Vaccinates		15	1	1		
		e affirmed by 1	icensee t	to be due	to causes o	ther than vaccination.	
	Number of Anii	mals with I	njectio	n Site l	Reaction	<u>1S</u>	
	Vaccinates	Days Post	-Vaccin			T.,	
	Left Side	14 74		28 7		42	
	Right Side	94		-		3	
	All injection site reactions were transient and under 2 cm in diameter.						
USDA Approval Date	09/08/1999					_	
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Study Type	Safety			
Pertaining to	ALL			
Study Purpose	Demonstrate safety in cattle under field conditions			
Product Administration	Two doses administered 28 days apart			
Study Animals	200 crossbred beef heifers			
Challenge Description	NA			
Interval observed after	Calves were observed for one hour after vaccination and daily			
challenge	throughout the study for adverse events. Animals were examined			
	for injection site reactions on days 14, 28, and 42 post-			
	vaccination (left side) and day 14 post-vaccination (right side).			
Results	Number of Animals with Adverse Events			
		1		
		SSS	Ring worm	
		lene) N	
		Lameness	Sing	
	Vaccinates 1 1			
	All observations were affirmed by licensee to be due to causes other than vaccination.			
	Number of Animals with Injection Site Reactions			
	Vaccinates	Days Post-Vacci	128	42
	Left Side	80	22	7
	Right Side	95	-	-
	All injection site reactions were transient and under 4 cm in diameter.			
	00/00/4000			
USDA Approval Date	09/08/1999			

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