

Summary of Studies Supporting USDA Product Licensure

| Establishment Name | Elanco US Inc. |
|---------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| USDA Vet Biologics Establishment Number | 196 |
| Product Code | 44D7.22 |
| True Name | Bovine Rhinotracheitis-Virus Diarrhea-Parainfluenza 3- Respiratory Syncytial Virus Vaccine, Killed Virus, Haemophilus Somnus-Leptospira Canicola-Grippotyphosa- Hardjo-Icterohaemorrhagiae-Pomona Bacterin |
| Tradename(s) / Distributor or Subsidiary (if different from manufacturer) | Vira Shield 6 + L5 HB Somnus - Elanco US Inc. Vira Shield 6 + VL5 HB Somnus - No distributor specified |
| Date of Compilation Summary | April 24, 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.

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| Study Type | Efficacy |
|---------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Bovine Virus Diarrhea (BVD) |
| Study Purpose | To demonstrate effectiveness against Bovine Virus Diarrhea |
| | Type 1b |
| Product Administration | |
| Study Animals | Bovine |
| Challenge Description | BVD isolate NY-1, BVD1b |
| 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 require publication of data submitted after that date. |
| USDA Approval Date | July 16, 2003 |

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| Study Type | Efficacy |
|-------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Bovine Virus Diarrhea (BVD) |
| Study Purpose | To demonstrate effectiveness against Bovine Virus Diarrhea |
| | Type 2a |
| Product Administration | |
| Study Animals | Bovine |
| Challenge Description | BVD isolate 890, Type BVD2a |
| 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 require publication of data submitted after that date. |
| USDA Approval Date | November 3, 1998 |

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| Study Type | Efficacy |
|-------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Bovine Rhinotracheitis |
| Study Purpose | To demonstrate effectiveness against Bovine Rhinotracheitis. |
| 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 require publication of data submitted after that date. |
| USDA Approval Date | October 22, 1986 |

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| Study Type | Efficacy |
|-------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Haemophilus somnus |
| Study Purpose | To demonstrate effectiveness against <i>Haemophilus somnus</i> |
| 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 require publication of data submitted after that date. |
| USDA Approval Date | August 2, 1988 |

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| Study Type | Efficacy |
|-------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Leptospira canicola, Leptospira grippotyphosa, Leptospira |
| Ct. I. D | icterohaemorrhagiae, Leptospira pomona |
| Study Purpose | To demonstrate effectiveness against <i>Leptospira spp</i> . |
| Product Administration | |
| Study Animals | Bovine, swine |
| 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 require publication of data submitted after that date. |
| USDA Approval Date | March 18, 1983 |

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| Study Type | Efficacy |
|-------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Leptospira hardjo type hardjo-bovis |
| Study Purpose | To demonstrate effectiveness against <i>Leptospira hardjo</i> type |
| _ | hardjo-bovis |
| 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 require publication of data submitted after that date. |
| USDA Approval Date | July 24, 2001 / November 9, 2001 |

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| Study Type | Efficacy |
|-------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Bovine Parainfluenza Type 3 |
| Study Purpose | To demonstrate effectiveness against Bovine Parainfluenza Type |
| | 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 require publication of data submitted after that date. |
| USDA Approval Date | March 12, 1987 (License Issued) |

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| Study Type | Efficacy |
|-------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Bovine Respiratory Syncytial Virus |
| Study Purpose | To demonstrate effectiveness against Bovine Respiratory |
| | Syncytial Virus |
| 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 require publication of data submitted after that date. |
| USDA Approval Date | December 18, 1987 |

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| Study Type | Efficacy |
|-------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Bovine Virus Diarrhea |
| Study Purpose | To demonstrate effectiveness against Bovine Virus Diarrhea |
| | Type 2a at 11 months post-vaccination |
| Product Administration | |
| Study Animals | Bovine |
| Challenge Description | BVD Isolate 890, Type BVD2a |
| 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 require publication of data submitted after that date. |
| USDA Approval Date | September 24, 1996 |

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| Study Type | Safety |
|-------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | All fractions |
| Study Purpose | Safety by IM route in bovine |
| Product Administration | |
| Study Animals | |
| Challenge Description | |
| Interval observed after | |
| challenge | |
| Results | Scientific data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. |

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| Study Type | Safety |
|-------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | All fractions |
| Study Purpose | To demonstrate safety under field conditions |
| Product Administration | |
| Study Animals | Bovine, including pregnant cows and heifers |
| Challenge Description | |
| Interval observed after | |
| challenge | |
| Results | Scientific data were evaluated by USDA-APHIS prior to product licensure and met regulatory standards for acceptance at the time of submission. However, study data are not available. |

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