

Summary of Studies Supporting USDA Product Licensure

| Establishment Name | Intervet Inc. |
|---------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| USDA Vet Biologics Establishment Number | 165A |
| Product Code | 12D5.30 |
| True Name | Bursal Disease-Reovirus Vaccine, Standard & Variant, Killed Virus |
| Tradename(s) / Distributor or Subsidiary (if different from manufacturer) | Breedervac-Reo-Plus - Intervet Mexico S.A. de C.V. Breedervac-Reo-Plus - Intervet Mexico S.A. de C.V Merck Sharpe and Dohme (MSD) Breedervac-Reo-Plus - Intervet, Inc. Breedervac-Reo-Plus - Merck Animal Health |
| Date of Compilation Summary | October 21, 2021 |

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 | Efficacy |
|-------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Infectious Bursal Disease (IBD), Standard Strain |
| Study Purpose | To demonstrate effectiveness in chickens to provide passive |
| | immunity against IBD, Standard strain to progeny. |
| Product Administration | Subcutaneous, Intramuscular |
| Study Animals | Chickens |
| 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 | September 15, 1987 |

| Study Type | Efficacy |
|-------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Infectious Bursal Disease (IBD), Delaware Variant A Strain |
| Study Purpose | To demonstrate effectiveness in chickens to provide passive |
| | immunity against IBD, Delaware Variant A strain, to progeny. |
| Product Administration | Subcutaneous, Intramuscular |
| Study Animals | Chickens |
| 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 | September 15, 1987 |

| Study Type | Efficacy |
|-------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Infectious Bursal Disease (IBD), Delaware Variant E Strain |
| Study Purpose | To demonstrate effectiveness in chickens to provide passive |
| | immunity against IBD, Delaware Variant E strain, in progeny. |
| Product Administration | Subcutaneous, Intramuscular |
| Study Animals | Chickens |
| 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 | July 29, 1988 |

| Study Type | Efficacy |
|-------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Infectious Bursal Disease (IBD), GLS-5 Strain |
| Study Purpose | To demonstrate effectiveness in chickens to provide passive |
| | immunity against IBD, GLS-5 strain, to progeny. |
| Product Administration | Subcutaneous, Intramuscular |
| Study Animals | Chickens |
| 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 8, 1991 |

| Study Type | Efficiency |
|-------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Study Type | |
| Pertaining to | Infectious Bursal Disease (IBD), Standard & Variant Strains |
| Study Purpose | To demonstrate effectiveness in chickens to provide passive |
| · · · | immunity against IBD, standard and variant strains, to progeny. |
| Product Administration | Subautanaous Intramuscular |
| Troduct Administration | |
| Study Animals | Chickens |
| 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 | May 8, 1992 |

| Study Type | Efficacy |
|-------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Avian Reovirus |
| Study Purpose | To demonstrate effectiveness in chickens to provide passive |
| | immunity against malabsorption caused by Avian Reovirus to |
| | progeny. |
| Product Administration | Subcutaneous, Intramuscular |
| Study Animals | Chickens |
| 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 | November 17, 1983 |

| Study Type | Efficacy |
|-------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | Avian Reovirus |
| Study Purpose | To demonstrate effectiveness against tenosynovitis caused by |
| | avian reovirus |
| Product Administration | Intramuscular |
| Study Animals | Chickens |
| 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 | November 17, 1983 |

| Study Type | Efficacy |
|-------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Doutoining to | Avior Booving |
| Pertaining to | Avian Reovirus |
| Study Purpose | To demonstrate effectiveness against tenosynovitis caused by |
| | Avian Reovirus. |
| Product Administration | Subcutaneous |
| Study Animals | Chickens |
| 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. Study data, however, are no longer available. |
| USDA Approval Date | March 23, 1994 |

| Study Type | Safety |
|-------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Pertaining to | ALL |
| Study Purpose | To demonstrate safety of product under typical field conditions |
| Product Administration | Subcutaneous, Intramuscular |
| Study Animals | Chickens |
| 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 | January 3, 1985 |