

ZELNATE[®] DNA IMMUNOSTIMULANT JUMP-STARTS THE ANIMAL'S OWN DEFENSE SYSTEM TO HELP FIGHT BRD.

Zelnate[®] is the first licensed immunostimulant that aids in the treatment of bovine respiratory disease (BRD) due to Mannheimia haemolytica. Zelnate jump-starts the animal's own defense system to help fight infectious disease.



- Aids in the treatment of BRD due to Mannheimia haemolytica in cattle 4 months of age or older, when administered at the time of, or within 24 hours after, a perceived stressful event.
- In six studies, Zelnate was shown to significantly reduce mortality due to BRD from 4.45% to 3.5% or a relative change of 20.83%.1
- IM (intramuscular) administration
 - No injection site issues were observed during field safety studies.
- Intranasal administration
 - Spray into one nostril with a syringe using an atomization tip.

- Proprietary DNA liposome complex stimulates the innate immune system in cattle.
 - The innate immune system has been shown to provide a rapid, potent and broad protectective response to infectious agents.
- A novel technology that enhances an animal's natural defenses and contains no antibiotics and no preservatives.

Elanco

- Can be used in natural programs.
- Available in 10 and 50 dose package sizes.

FOR MORE INFORMATION, PLEASE CONTACT YOUR ELANCO SALES REPRESENTATIVE OR CALL US AT 800-364-2014.

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Do not administer within 21 days of slaughter.

^{&#}x27;Nickell, J., Keil, D., Settje, T., et al. 2016. "Efficacy and safety of a novel DNA immunostimulant in cattle." Bov Pract. 50(1):9-20.

This product is based on technology developed by Juvaris BioTherapeutics and is patent protected. Animal health applications are being developed exclusively under the rights of Elanco and are protected by patents.

See productdata.aphis.usda.gov for a summary of the studies approved by the USDA for licensing this product. This package insert may also contain additional information developed by the licensee.

DNA Immunostimulant



For Intramuscular or Intranasal Administration to Cattle

FOR VETERINARY USE ONLY

02320

READ IN FULL

DESCRIPTION

The innate immune system in cattle has been shown to provide a potent, rapid, nonspecific, protective response to infectious agents, such as Mannheimia haemolytica that can lead to Bovine Respiratory Disease (BRD). BRD is a serious condition that commonly causes lung lesions, reduced lung capacity and mortality.

ZELNATE® is a bacterial-produced plasmid DNA with a liposome carrier that stimulates the innate immune system and has been shown to be effective against bovine respiratory disease due to Mannheimia haemolytica.

The freeze-dried (desiccate) product is packaged with two different sterile diluents. The First Sterile Rehydrator (vial 1) is used to reconstitute the desiccate cake (vial 2), and then transferred to the Final Sterile Solution (vial 3) to achieve the proper concentration for administration.

INDICATION

This product has been shown to be effective for the treatment of cattle, 4 months of age or older, against bovine respiratory disease due to *Mannheimia haemolytica*. For more information regarding efficacy and safety data, see productdata.aphis.usda.gov.

This product has been shown to be effective at the time of, or within 24 hours after, a perceived stressful event.

IMPORTANT STORAGE CONDITIONS

Store Refrigerated

2°C to 8°C (35°F to 46°F) DO NOT FREEZE.

Stability has been demonstrated for at least 8 hours after reconstitution if vial is refrigerated and sterility is maintained.



METHOD OF ADMINISTRATION

Inject 2 mL intramuscularly at the time of, or within 24 hours after, a perceived stressful event (for example: weaning, shipping, commingling or adverse environmental conditions). Alternatively, spray 2 mL into one nostril using an atomization tip attached to the syringe; the atomizer should produce a fine mist of particles 30-100 microns in size for delivery to the mucosal membranes. Use entire contents of vial once first opened.

CAUTION

must be

2

In case of human exposure, contact a physician.

| Study Type | | | Efficacy | | | | | | | | | | | | | | | |
|--|----|---|---|---|-------------------------------------|--|---|----------------------------------|-----------------------------------|--------------------------------|--------------------------------|-----------------------------------|-------------------------------|---------|---------|----|--|--|
| Pertaining to Study Purpose Product Administration Study Animals | | | Mannheimia haemolytica Efficacy against bovine respiratory disease | | | | | | | | | | | | | | | |
| | | | | | | | | | | | | | | | | | One dose administered by IM route <u>at the time of challenge</u> . Control group administered diluent only | |
| | | | 64 Holstein steers of 3-4 months of age; randomized into 2 groups of 32 calves each | | | | | | | | | | | | | | | |
| | | | Challenge Description | | | live M. | haem | olytica | a inocu | lum | | | | | | | | |
| Interval observed after challenge | | e | Observed daily for 5 days. Lungs were evaluated 5 days after challenge. | | | | | | | | | | | | | | | |
| | | | includ | ed in t | he ana | | | · | ior to E ation | Jay J, | | uupsy | r rung s | | vas IIU | L | | |
| | | | Treatment | | | | | | | | | | | Maximum | | | | |
| | | | | | | mum | <u> </u> | | /ledian | | | | | | | | | |
| | | | <u>Treatn</u> Contro Treate | ls | | <u>mum</u> 0% 0% | | % | <u>Aedian</u> 10% 4% | 15 | i%)% | Maxim 33% 22% | 6 | | | | | |
| | | | Contro Treate | d data sh | own oi | 0% 0% | 69 19 | % | 10% | 15 | i%)% | 33% 22% | 6 % | y 5 are | marke | ed | | |
| | | | Contro Treate Raw d an aste The de | d ata sh erick (aths p | own o *). prior to | 0% 0% n the ta Day 5 | 69 19 able be were: | % % :low. T 1/32 i | 10% 4% | 15 10 nals th | i%)% nat die oup: 1/ | 33% 22% d prior '32 in (| 6 % r to Day Control | | | | | |
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| USDA Aproval Date | | | Contro Treate Raw d an aste The de was se 28-Feb | ils d ata sh erick (eaths p evere t o-2013 | own oi *). prior to povine | 0% 0% n the ta Day 5 respira | 69 19 able be were: atory d | % klow. T 1/32 i isease | 10% 4% The anir | 15 1(mals th ted gro | i%)% nat die oup: 1/ | 33% 22% d prior '32 in (| 6 % r to Day Control | | | | | |
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death prior to Day 5 DIAMOND

Treated (Cont.)

Control (Cont.)

MANUFACTURED BY: Diamond Animal Health, Inc. Des Moines, IA 50327 U.S. Veterinary License No. 213 PCN 9381.D0 Made in U.S.A November, 2018 85877690 I V1811



4% 5% 5% 6% 8% 9% 10% 10% 10% 11% 12% 13% 13% 15% 18% 22

10% 10% 10% 11% 13% 14% 15% 15% 18% 18% 21% 23% 27% 29% 33% 349

DISTRIBUTED BY: Bayer HealthCare LLC, Animal Health Division P.O. Box 390 Shawnee Mission, KS 66201 U.S.A. 1-800-633-3796

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PRECAUTION

Do not administer within 21 days of slaughter. Do not mix with other products, except as specified on this label. This product has not been tested in pregnant animals.

OTHER INFORMATION

Contains no antibiotics and no preservatives.

HOW SUPPLIEDVials of 10 and 50 doses.

