Sequential VFD ID Number

Veterinary Feed Directive for use in Calves, Beef Cattle and Nonlactating Dairy Cattle Chlortetracycline

Client: Business or Home Address: Phone Number:		JOE PRODUCER	Veterinarian:	DR JD EXAMPLE		
		222 SECOND DRIVE	Address: Phone Number:	111 FIRST LANE		
		XXX-XXX-XXXX		XXX-XXX-XXXX		
Location	of animals:	r of animals to be treated: 100 222 SECOND LANE nd/or other animal identifications:				
FEED 4 O	UNCES PER	HEAD DAILY TO PROVIDE 750 MG PER 150	00 ANIMAL			
Indication	ns, Drug Lev	vel in Medicated Feed, and Duration	of Use (select one and spec	ify the additional required inf	ormation):	
	A. For Gro	owing Cattle (over 400 lbs): For the re Drug level: g/ Duration of use:	ton in order to provide 70 n			
	B. For Beef Cattle: For the control of bacterial pneumonia associated with shipping fever complex caused by <i>Pasteurella spp</i> . susceptible to chlortetracycline. Drug level:g/ton in order to provide 350 mg / head/ day					
	Duration of use: days C. For Beef Cattle (under 700 lbs.): Control of active infection of anaplasmosis caused by <i>Anaplasma marginale</i> susceptible to chlortetracycline. Drug level: g/ton in order to provide 350 mg / head/ day Duration of use: days					
Ø	D. For Beef Cattle (over 700 lbs.): Control of active infection of anaplasmosis caused by <i>Anaplasma marginale</i> susceptible to chlortetracycline. Drug level: 6000 g/ton in order to provide 0.5 mg/lb body weight / day Duration of use: 180 days					
	E. For Calves, Beef, and Nonlactating Dairy Cattle; For treatment of bacterial enteritis caused by <i>Escherichia coli</i> and bacterial pneumonia caused by <i>Pasteurella multocida</i> susceptible to chlortetracycline. Drug level: g/ton in order to provide 10 mg/lb body weight / day Duration of use: days (1 to 5 days)					
Caution:	Use of feed	containing this Veterinary Feed Dire	ctive (VFD) drug in a manne	er other than as directed on th	ne labeling (extra-label use) is not permitted.	
		For use i	n Dry Feeds Only. Not for U	se in Liquid feed Supplements	i.	
	Residue	e Warnings: Zero-day withdrawal pe	riod. A withdrawal period h Do not use in calves to be		ois product in pre-ruminating calves.	
Combina	ation Use:					
	This VFD only authorizes the use of the VFD drug(s) cited in this order and is not intended to authorize the use of such drug(s) in combination wit other animal drugs.				e use of such drug(s) in combination with any	
	This VFD authorizes the use of the VFD drug(s) cited in this order in the following FDA-approved, conditionally approved, or indexed combination(s) in medicated feed that contains the VFD drug(s) as a component. (List the specific approved combination(s))					
Ø		This VFD authorizes the use of the VFD drug(s) cited in this order in any FDA-approved, conditionally approved, or indexed combination(s) in medicated feed that contains the VFD drug(s) as a component.				
VFD Issua	ince Date:	01-01-xxxx	VFD Expiration Date	e: 6-01-XXXX		
Veterinar	ian's Signat	ure:		Mont (Not to exceed 6 months	:h/Day/Year s from issuance date)	
	– Veterina		Copy – Supplier		Copy – Client	
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All parties must retain a copy of this veterinary feed directive for 2 years after issuance