Veterinary Feed Directive for Cattle Aureomycin® (chlortetracycline)

Sequential VFD ID Number, if appropriate

Veterinarian	ian: Client:		
Address:	Business/Hom	e Address:	
Phone #:	Phone #:	ntional	
	nail: (optional) FAX or email: (o		
Indications information	ns, Drug Level in Medicated Feed, and Duration of Use: (select ion)	one and specify additional required	
1)			
	Drug Concentration: g/ton (to provide 70 mg/head	d/day)	
	1 ,		
	spp. susceptible to chlortetracycline. Drug Concentration:g/ton (to provide 350 mg/hea	ad/day)	
	Duration of Feeding:days 3) Beef Cattle (under 700 lb): Control of active infection of anaplasmosis caused by <i>Anaplasr</i> susceptible to chlortetracycline.		
	Drug Concentration:g/ton (to provide 350 mg/heat Duration of Feeding:days	ad/day)	
		nosis caused by Anaplasma marginale	
	Drug Concentration:g/ton (to provide 0.5 mg/lb b Duration of Feeding:days	ody weight/day)	
	8000 g/ton (to provide 0.5 to 2.0 mg/lb body weight/day)		
	[Must use an FDA-approved proprietary formulation.]		
	6000 g/ton (to provide 0.5 to 2.0 mg/lb body weight/day) [Must use an FDA-approved proprietary formulation or the FDA	A-approved formulation in 21 CFR 558.128(e)(6).]	
	5000 g/ton (to provide 0.5 to 2.0 mg/lb body weight/day) [Must use an FDA-approved proprietary formulation.]		
	700 g/ton (to provide 0.5 to 2.0 mg/lb body weight/day) [Must use an FDA-approved proprietary formulation.]		
	Duration of Feeding:days		
6)	Calves, Beef and Non-Lactating Dairy Cattle: Treatment of bact bacterial pneumonia caused by Pasteurella multocida organism Drug Concentration:		
		ovide 10 mg/lb body weight/day) ²	
	Top Dressg/ton (4000 to 20,000 g/ton to provi	de 10 mg/lb body weight/day)	
	Duration of Feeding: days (Feed for not more than 5	3 /	
^{1,2} W indi *******	. ² When used in combination with 30 to 181.8 g/ton lasalocid to proviondication is only approved for use in cattle weighing up to 800 lbs.	le 1 mg/2.2 lb body weight per day, this	
	USE OF FEED CONTAINING THIS VETERINARY FEED DIRECTIVE (VFD DIRECTED ON THE LABELING (EXTRA-LABEL USE)	DRUG IN A MANNER OTHER THAN AS	
Approximate number of <i>Cattle</i> to be treated:			
	s or Location of cattle:		
	nstructions and/or other animal identifications:		

Affirmation	of Intent (for combination VFD drugs): check the appropriate box	:		
	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 with any other animal drugs.			
☐ 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.				
	Drug(s) and Dose Range(s) [all listed doses are 90% DM basis]			
	10 to 30 g/ton lasalocid to provide 100 to 360 mg per head per day (BOVATEC®) [NADA 141-250]	Cattle fed in confinement for slaughter.		
	25 to 30 g/ton lasalocid to provide 250 to 360 mg per head per day (BOVATEC®) [NADA 141-250]	Cattle fed in confinement for slaughter.		
	30 to 600 g/ton lasalocid to provide 60 to 360 mg per head per day (BOVATEC®) [NADA 141-250]	Pasture cattle (slaughter, stocker, feeder cattle, dair and beef replacement heifers.)		
	30 to 181.8 g/ton lasalocid to provide 1 mg per 2.2 lb bodyweight per day (maximum 360 mg lasalocid daily) (BOVATEC®) [NADA 141-250]			
	5 g/ton laidlomycin propionate potassium to provide 30 to 75 mg per head per day (CATTLYST®) [NADA 141-201]	Cattle fed in confinement for slaughter.		
	5 to 10 g/ton laidlomycin propionate potassium to provide 30 to 150 mg per head per day (CATTLYST®) [NADA 141-201]	Cattle fed in confinement for slaughter.		
	12.9 to 90.8 g/ton decoquinate to provide 22.7 mg per 100 lb body weight per day (DECCOX®) [NADA 141-185]	Calves, beef and non-lactating dairy cattle.		
	90.9 to 535.7 g/ton decoquinate to provide 22.7 mg per 100 lb body weight per day (DECCOX®) [NADA 141-185]	Calves, beef and non-lactating dairy cattle.		
	Other FDA-approved, conditionally approved, or indexed combination:			
	*for complete information see the approved Type C medicated f	eed label		
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.				
Warning: No withdrawal period is required. A withdrawal period has not been established for this product in pre-ruminating calves. Do not use in calves to be processed for veal.				
Date of VFD Issuance:(dd/mm/yyyy)				

n Color X Copy – Supplier
All parties must retain a copy of this VFD for 2 years after issuance

Color Y Copy - Client

Veterinarian's signature:

Color Z Original – Veterinarian