Sequential VFD ID Number, if appropriate

Veterinary Feed Directive for Swine Aureomycin® (chlortetracycline)

/eterin	naria	n:	Client:				
Addres	ss:		Business or Home Address:				
Phone	щ.	<u> </u>	Phone #:				
		ail: (optional)					

ndicat			n of Use: (select one and specify additional required				
	1)	Swine: Control of porcine proliferative enteropathies (ileitis) caused by <i>Lawsonia ntracellularis</i> susceptible to chlortetracycline.					
			vide 10 mg/lb body weight/day, which is equivalent to				
		Duration of use: days (Feed for no	ot more than 14 days)				
П	2)	Swine: Treatment of bacterial enteritis caused by <i>Escherichia coli</i> and <i>Salmonella</i>					
Ш	,	choleraesuis and bacterial pneumonia caused by Pasteurella multocida susceptible to					
		chlortetracycline.	·				
		Drug Concentration:g/ton (to pro	vide 10 mg/lb body weight/day, which is equivalent to				
		approximately 400 g/ton)	t more than 14 days)				
	Duration of use:days (feed for not more than 14 days) 3) Swine: Reduction in the incidence of cervical lymphadenitis (jowl abscesses) caused						
Ш	3)	Broup E <i>Streptococci</i> susceptible to chlortetracycline.					
		Drug Concentration:g/ton (50 to					
		Duration of use: days	100 grany				
	۸)	Breeding Swine: Control of lentoenirosi	s (reducing the incidence of abortion and shedding				
Ш	4)	of <i>leptospirea</i>) caused by <i>Leptospira po</i>	,				
		Drug Concentration: 400 g/ton	miona susceptible to chlorietracycline.				
		Duration of use: days (feed continuous)	ly for not more than 14 days)				
*****	****	***************************************	*******************				
USE	OF	FEED CONTAINING THIS VETERINARY FEE	D DIRECTIVE (VFD) DRUG IN A MANNER OTHER THAN AS				
			XTRA-LABEL USE) IS NOT PERMITTED.				
Prem	ises	or Location of animals:					
Speci	ial In	structions and/or other animal identifications:					
Affirn		on of Intent (for combination VFD drugs): check					
		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, condition						
			d feed that contains the VFD drug(s) as a component.				
		Drug(s) and Dose Range(s)	Specifications*				
		10 to 30 g/ton bacitracin methylene disalicylate;					
	_	(BMD®) [NADA 200-242]	This combination has a zero day withdrawal.				
		Other FDA-approved, conditionally approved, or	indexed				
		combination:					
		*for complete information see the approved Type	a C medicated feed lahel				

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.

for complete information see the approved Type C medicated feed label

Withdrawal Period: No withdrawal period is required.						
Date of VFD Issuance:	(dd/mm/yyyy)	Date of VFD Expira (Cannot exceed 6 r	ation: (dd/mm/yyyy) months after issuance)			
Veterinarian's signature:		<u> </u>	,			
Color 7 Original	_ Votorinarian	Color X Cony - Supplier	Color V Cony - Client			

Color Z Original – Veterinarian Color X Copy – Supplier

All parties must retain a copy of this VFD for 2 years after issuance

Color Y Copy - Client