| (ch | UREOMYCIN® |
|---|--|
| narian: | Client: |
| SS: | Business/Home Address: |
| : | Phone: |
| | Fax or Email (optional): |
| | : (select one and specify additional required information) |
| Growing Cattle (over 400 lb): For the reduction of th Drug Concentration: g/ton (to provi Duration of Feeding: days | e incidence of liver abscesses. ide 70 mg/head/day) |
| Beef Cattle: Control of bacterial pneumonia associate chlortetracycline. Drug Concentration: g/ton (to provi Duration of Feeding: days | ed with shipping fever complex caused by <i>Pasteurella spp.</i> susceptible to ide 350 mg/head/day) |
| chlortetracycline. Drug Concentration: g/ton (to prove | n of anaplasmosis caused by <i>Anaplasma marginale</i> susceptible to ide 350 mg/head/day) |
| Beef Cattle (over 700 lb): Control of active infection of chlortetracycline. Drug Concentration: g/ton (to provi | of anaplasmosis caused by <i>Anaplasma marginale</i> susceptible to ide 0.5 mg/lb body weight/day) |
| 5. Beef and Non-lactating Dairy Cattle: As an aid in consusceptible to chlortetracycline when delivered in a final prug Concentration: 8000 g/ton (to provide 0.5 to 2.0 mg/lb body weight/of [Must use a FDA-approved proprietary formulation.] 6000 g/ton (to provide 0.5 to 2.0 mg/lb body weight/of [Must use a FDA-approved proprietary formulation or formulations g/must use a FDA-approved proprietary formulation.] 700 g/ton (to provide 0.5 to 2.0 mg/lb body weight/daff [Must use a FDA-approved proprietary formulation.] Duration of Feeding: days | day) day) ulation in 21 CFR 558.128(e)(6).] day) |
| caused by Pasteurella multocida organisms susceptible Drug Concentration: Complete Feed g/ton (500 to 4,00 to 20 g/ton of Feeding: days (Feed for not g/ton of Feeding: | 00 g/ton to provide 10 mg/lb body weight/day) 0,000 g/ton to provide 10 mg/lb body weight/day) |
| DIRECTED ON THE LABELING | G (EXTRA-LABEL USE) IS NOT PERMITTED. Premises or Location of Cattle: |
| | marian: |

___(dd/mm/yyyy)

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. (List the specific approved combination)

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 withdrawl 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.

Veterinarian's signature: WHITE Original - Veterinarian

Date of VFD Issuance: _

CANARY Copy - Supplier

Date of VFD Expiration: ___

(Cannot exceed 6 months after issuance)

PINK Copy - Client

_(dd/mm/yyyy)