Section 1932A

FOLD, SEAL, AND RETURN

VETERINARY ADVERSE DRUG REACTION, LACK O EFFECTIVENESS OR PRODUCT DEFECT REPORT			DATE REPORTED	Form Approved: OMB No. 0910-0284 Expiration Date: January 31, 2010		
NOTE: This report is authorized by 21 U.S.C 352(a) and (f). While you are not required to report, your cooperation is needed to assure comprehensive and timely assessment of product labeling.						
If you do NOT want your identity disclosed to the manufacturer, place an "X" in this	1. VETERINARIAN'S NAME AND ADDRESS			2. OWNER'S NAME OR CASE ID (In Confidence)		
box.	TELEPHONE (Include Area Code)			3. NADA NUMBER (For FDA Use)		
4. SUSPECTED DRUG AND DOSAGE FORM				5. MANUFACTURER'S NAME		
6. DIAGNOSIS AND / OR REASON FOR USE OF DRUG				7. ADMINISTERED BY VETERINARIAN OWNER		
8. DOSAGE ADMINISTERED AND ROUTE (Ex. 250 mg. q 12h, 5 days, orally)				9. DATE(S) OF ADMINISTRATION		
10. SPECIES	11. BREED	12. AGE		13. SEX	14. WEIGHT	
15. CONCURRENT CLINICAL PROBLEMS NONE OVERALL STATE OF HEALTH WHEN SUSPECTED DRUG GIVEN: GOOD FAIR POOR CRITICAL			16. CONCURRENT DRUGS ADMINISTERED NONE			
17. REACTION INFORMATION						
a. TIME BETWEEN INITIATION OF THERAPY WITH SUSPECTED DRUG AND ONSET OF REACTION WAS b. TIME BETWEEN LAST ADMINISTRATION OF SUSPECTED DRUG AND ONSET OF REACTION WAS c. OUTCOME: RECOVERED FROM REACTION DIED FROM REACTION OTHER (Comment Below) d. WAS THE REACTION TREATED? NO YES (Comment Below) e. WHEN THE REACTION APPEARED, TREATMENT WITH SUSPECTED DRUG: HAD ALREADY BEEN COMPLETED WAS DISCONTINUED DUE TO REACTION WAS DISCONTINUED AND REPLACED WITH ANOTHER DRUG WAS DISCONTINUED AND REPLACED WITH ANOTHER DRUG REACTION OTHER (Comment Below) f. LEVEL OF SUSPICION THAT DRUG CAUSED THE REACTION: HIGH MEDIUM LOW						
POSSIBLE CONTRIBUTING	ADD DETAILS ABOUT CASE HISTORY AND O FACTORS. DESCRIBE LACK OF EFFECTIVENING TO THE PROPERTY OF THE PROPERTY O	ESS OR PRO	DDUCT DEFECT (Includ			

An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number.

Department of Health and Human Services Food and Drug Administration CVM, HFV-210 (0910-0012) 7500 Standish Place Rockville, MD 20855

Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information for reducing this burden to:

FOLD NO POSTAGE **DEPARTMENT OF NECESSARY HEALTH & HUMAN SERVICES** IF MAILED IN THE **UNITED STATES** Public Health Service Food and Drug Administration Rockville MD 20857 Official Business Penalty for Private use \$300 **BUSINESS REPLY MAIL** FIRST CLASS PERMIT NO. 946 ROCKVILLE MD POSTAGE WILL BE PAID BY FOOD AND DRUG ADMINISTRATION Department of Health and Human Services Food and Drug Administration CVM, HFV-210 (0910-0012) 7500 Standish Place Rockville MD 20855 Intelligated about the latest and the Historia FOI D THANK YOU FOR SHARING YOUR CONCERN ABOUT ANIMAL DRUG EFFECTS 18. (Continued)

FOR FDA USE ONLY

Confidentiality: The owner's identity is

protected to the fullest extent of the law.

held in strict confidence by FDA and

The reporter's identity, including the identity of self-reporter, may be shared

with the manufacturer unless requested otherwise. However, FDA will not disclose the reporter's identity in response to a request from the public, pursuant to

the Freedom of Information Act.

☐ NAI

☐ AI

AP

__ AL

CONT

PR

CR

___ I.L.

COMMENT