Section 9a

VETERINARY ADVERSE DRUG REACTION, LACK OF EFFECTIVENESS, PRODUCT DEFECT REPORT (Forward to address at left. Attach all correspondence that pertains to this reaction)

Form Approved: OMB No. 0910-0284 Expiration Date: January 31, 2010

Public reporting burden for this collection of information is estimated to average 2 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to:

Food and Drug Administration

An agency may not conduct or sponsor, and a person is not required

Rockville, MD 20855	210), ROOM N403		MB control numbe	r.	it displays a currently	
).300). Failure t			pproval of the application.
1. REPORT SOURCE AND	ADDRESS (Mfr., Dis	tr.)		b. DATE SENT TO F		
4. NAME, ADDRESS AND PHONE NO. OF ATTENDING VETERINARIAN (In confidence) Name:				5. NAME OR CASE IDENTIFICATION OF OWNER (In confidence)		
Street Address:						
City:	State:	ZIP:				
Phone No. ()				7- NAME OF MANU	IEACTURER	
(Include dosage form and strength - Ex., tab, 500 mg.)				7a. NAME OF MANUFACTURER		
				b. NADA NO.		
8. LOT NUMBER(S) 9. DOSAGE ADMINISTERED AND ROUTE (Ex. 250 mg., q 12 h, p.o.)				10. DATE(S) OF ADMINISTRATION		
11. ILLNESS/REASON FOR USE OF THIS DRUG				12. DRUG WAS ADMINISTERED BY		
				UVETERINARIAN, STAFF OWNER, OTHER		
13. NUMBER OF ANIMALS IN THIS INCIDENT			14. REACTING ANIMALS			
a. TREATED WITH DRUG	b. REACTED)	c. DIED	a. SPECIES	b	. BREED
15. CONCOMITANT MEDICAL PROBLEMS				c. AGE	d	. WEIGHT
				e. SEX		PREGNANT NEUTERED
				Y NEW ILLNESS DEV CT DRUG STARTED?		DIAGNOSIS CHANGE AFTER
GOOD FAIR POOR CRITICAL NO					Explain)	
18. CONC NAME OF DRUG ROUTE			NCOMITANT DRU	DOSAGE REGIMEN DATE(S) OF ADMINISTRATION		
			FOR FDA			
		COM	IMENT	USE UNLT		
1	PR	NAI AI AP AL	-			
□ I.L. □ C	CR CONT					

REACTION DATA 19. DESCRIBE SUSPECTED ADVERSE REACTION: INCLUDE ALL SIGNS, RESULTS OF PERTINENT LAB TESTS, NECROPSY RESULTS, POSSIBLE CONTRIBUTING FACTORS, ETC. ALSO, INCLUDE IN THIS SECTION PRODUCT INEFFECTIVENESS AND PRODUCT DEFECTS SUCH AS CRACKED TABLETS, CLOUDY SOLUTION, ETC. 20a. ATTENDING VETERINARIAN'S LEVEL OF SUSPICION THAT DRUG 20b. WAS THERE EXTRA LABEL USE (ELU) INVOLVED? CAUSED REACTION HIGH ■ MEDIUM LOW NO ATTENDING VET. ■ NO YES (Explain) 21. LENGTH OF TIME BETWEEN LAST ADMINISTRATION OF SUSPECT DRUG AND 22. DATE OF ONSET 23. DURATION OF REACTION ONSET OF REACT (Hrs., days, etc.) (Mo., day, yr.) 24. WAS THE ADVERSE REACTION TREATED? 25. OUTCOME OF REACTION TO DATE ☐ NO YES (Describe treatment) ☐ DIED (Give date) ☐ REMAINS UNDER TREATMENT ALIVE WITH SEQUELAE RECOVERED UNKNOWN 26. WHEN REACTION APPEARED, TREATMENT WITH SUSPECT DRUG: ☐ HAD ALREADY BEEN COMPLETED ☐ CONTINUED ☐ DISCONTINUED DUE TO THE REACTION STOPPED AND THE ☐ DISCONTINUED, REPLACE WITH ANOTHER DRUG **REACTION** ☐ DISCONTINUED, REINTRODUCED LATER RECURRED OTHER (Explain) ☐ CONTINUED AT ALTERED DOSE OTHER (Explain) _ 27. HAD ANIMAL(S) BEEN PREVIOUSLY EXPOSED TO THIS DRUG? YES NO UNKNOWN 28. DID ANIMAL(S) PREVIOUSLY REACT TO THIS DRUG? ■ NO YES UNKNOWN 29. HAD ANIMAL(S) PREVIOUSLY REACTED TO OTHER DRUGS? □NO YES UNKNOWN (If yes, give drug(s) and reaction if known) 30. HAS THE ATTENDING VETERINARIAN SEEN SIMILAR REACTIONS TO THIS DRUG IN ANY OTHER ANIMALS? □ NO YES (Describe treatment) 31. NAME AND TITLE OF INDIVIDUAL RESPONSIBLE FOR ACCURACY 32. SIGNATURE OF INDIVIDUAL RESPONSIBLE FOR ACCURACY OF REPORTED OF REPORTED INFORMATION (Type or print) INFORMATION