VETERINARY ADVERSE DRUG REACTION, LACK OF EFFECTIVENESS, PRODUCT DEFECT REPORT

Food and Drug Administration 7500 Standish Place (HFV-240) Rockville, MD 20855-9921

(Forward to address at right. Attach all correspondence that pertains to this reaction.)

NOTE: This report is required by law (21 CFR 514.80 and 512 (I) of the Federal Food, Drug, and Cosmetic Act (FDCA)). Failure to report can result in withdrawal of approval of the application (21 CFR 514.80 (h) and 512 (e) of the FDCA).

The data elements marked with an asterisk [*] require a value or text to be entered. An asterisk at the section level applies to all fields within that section. An asterisk at the subsection level applies to all fields within that subsection. Otherwise, asterisks apply to individual fields.

	A	Pa dministrative and Ide	rt A entificatio	on In	nformation	
		Regulatory Aut	nority - RA	(A.1	1)	
RA Name (A.1.1)*			Street Ade	-		
City (A.1.3)* State/County or Province		ce (A.1.4)		Mail/Zip Code (A.1.5)*	3-Character Country Code (A.1.6)*	
		Marketing Authorizati	on Holde	- M/	 AH (A.2)	
		MAH Inform	ation (A.2.	1)		
Business Name (A.2.1.1)*			Street Add	dress	(A.2.1.2)*	
City (A.2.1.3)*		State/County or Provinc	 xe (A.2.1.4)		Mail/Zip Code (A.2.1.5)*	3-Character Country Code (A.2.1.6)*
		Person Acting on Beh	alf of the l	ЛАН	(A.2.2)	
Title (e.g., Mr., Ms., Dr.) First (A.2.2.1) First	Name (A	.2.2.2)		Last	Name (A.2.2.3)	
Telephone Number (A.2.2.4)	Fax Nur	nber (A.2.2.5)	Email Add	lress	(A.2.2.6)	
		Person(s) Involve	d in the A	ER (A.3)	
		Primary Rep	oorter (A.3.	1)		
Primary Reporter Category (A.3.1	.1)* (Sele	ct One)				
Veterinarian Animal Owr	ner 🗌 F	Physician 🗌 Patient	Other I	lealth	n Care Professional 🗌 Othe	er 🗌 Unknown
Last Name (A.3.1.2)*		First Name (A.3.1.3)				
Telephone Number (A.3.1.4) Fax Number (A.3.1.5)		Email Address (A.3.1.6)				
Business Name (A.3.1.7)			Street Ad	dress	(A.3.1.8)	
City (A.3.1.9)		State/County or Provinc	⊥ ≿e (A.3.1.10))	Mail/Zip Code (A.3.1.11)	3-Character Country Code (A.3.1.12)*

Part	A - Adm	ninistrative and Iden	tification Info	ormation (Continued)	
		erson(s) Involved in tl		, ,	
			orter (A.3.2)		
Other Reporter Category (A.3.2.1)* (require	d if any of the A.3.2 informat	ion is provided) (S	Select One)	
Veterinarian Animal Ow	ner	Physician 🗌 Patient	Other Hea	alth Care Professional 🗌 Otl	her 🗌 Unknown
Last Name (A.3.2.2)			First Name (A.3	3.2.3)	
Telephone Number (A.3.2.4)Fax Number (A.3.2.5)Email Address (A.3.2.6)					
Business Name (A.3.2.7)			Street Address	(A.3.2.8)	
City (A.3.2.9)		State/County or Provinc	e (A.3.2.10)	Mail/Zip Code (A.3.2.11)	3-Character Country Code (A.3.2.12)
		AER Inform	nation (A.4)		
Unique AER Identification Numbe	r (A.4.1)*		. ,		
Original Receive Date (A.4.2)* (dd	l/mm/yyy	y)	Date of Current	t Submission (A.4.3)* (dd/mm/y	<i>ууу)</i>
Day Month Year Day Month Year				ır	
		Type of Re	eport (A.4.4)		
Type of Submission (A.4.4.1)* (Se	lect One)				
Expedited Periodic	E Fo	llow-up 🗌 Nullification	n 🗌 3-Day Fie	eld Alert	
Reason for Nullification Report (A	,	ovide if nullification is select	ted from A.4.4.1)		
	- /				
			rt B		
		Descriptio	on of the AE		
		nin this section (B.1) are		y if an animal is associated wi	th the report.)
Number of Animals Treated (B.1.)		Number of Anir	mals Affected (B.1.2)*	
Attending Veterinarian's Assessm	ent of An	imal Health Status Prior	to VMP Use (B.1	l.2.1)	

Species (B.1.3)*:

Breed (B.1.4)			
Purebred Information (B.1.4.1)			
Breed (B.1.4.1.1) of Animal 1	Breed (B.1.4.1.1) of Animal 2	Breed (B.1.4.1.1) of Animal 3	

	Part B - Description	of the AE (Continue	ed)
	Animal Data (B.1) (Continued)	
Crossbred Information (B.1.4.2)			
Breed (B.1.4.2.1)	Breed (B.1.4.2.1)		Breed (B.1.4.2.1)
Gender (B.1.5) (Select One)		Reproductive Status ((B.1.6) (Select One)
Female Male Mixed	Unknown	Intact Ne	eutered 🗌 Mixed 🗌 Unknown
Female Physiological Status (B.1.7) (Select	t One)		
Nonpregnant Lactating	pregnant Nonlactating	Pregnant Lactating	g 🗌 Pregnant Nonlactating
Mixed Not	Applicable	Unknown	
	Weigh	t (B.1.8)	
Measured, Estimated, Unknown Weights (B.1.8.1)*		nt in Kilograms (B.1.8.2) red or Estimated selected	
Measured Estimated Ur	Iknown		
	Age	(B.1.9)	
Measured, Estimated, Unknown Age (B.1.	9.1)* known		
Minimum Age (B.1.9.2) (provide if Measured	or Estimated Minimum A	ge Units (B.1.9.2.1) (pro	ovide if B.1.9.2 is given) (Select One)
selected from B.1.9.1)			Hour 🗌 Day 🗌 Month 🗌 Year
Maximum Age (B.1.9.3)			ovide if B.1.9.3 is given) (Select One)
			Hour Day Month Year
		and Usage (B.2)	
	appropriate B.2.1-B.2.6	-	onding pages of additional forms.)
Registered or Brand Name (B.2.1)*		Product Code (B.2.1.1	1)
Registration Identifier (B.2.1.2)*		ATCvet Code (B.2.1.3	3)*
Company or MAH (B.2.1.4)			
The following fields (B.2.1.5-B.2.1.7.1.3.3)	are applicable only if an	animal is associated wi	th the report
MAH Assessment (B.2.1.5)			
× ,			
RA Assessment (B.2.1.6)			
RA Assessment Term (B.2.1.6.1)			
Explanation Relating to Assessment (B.2.1.6.1.1)		
Route of Exposure (B.2.1.7)			
	Dose Per Admin	istration (B.2.1.7.1)	
Numeric Value for Dose (Numerator) (B.2		• •	r) (B.2.1.7.1.1.1) (provide if B.2.1.7.1.1 is given)
Numeric Value for Dose (Denominator) (B	21712) Units of Va	lue for Dose (Denomina	tor) (B 2 1 7 1 2 1) (provide if B 2 1 7 1 2 is given)

Part B - Description of the AE (Continued)				
VMP(s) Data and Usage (B.2) (Continued)				
Interval of Admir	nistration (B.2.1.7.1.3)			
	s of Value for Interval of Administration (B.2.1.7.1.3.1.1) vide if B.2.1.7.1.3.1 is given) <i>(Select One)</i>			
	Second 🗌 Minute 🗌 Hour 🗌 Day 📄 Month 🗌 Year			
Date of First Exposure (B.2.1.7.1.3.2) (dd/mm/yyyy)	Date of Last Exposure (B.2.1.7.1.3.3) (dd/mm/yyyy)			
Day Month Year	Day Month Year			
Active Ingre	edient(s) (B.2.2)			
1st Entry				
Active Ingredient(s) (B.2.2.1)*				

Numeric Value for Strength (Numerator) (B.2.2.1.1)*	Units for Numeric Value for Strength (Numerator) (B.2.2.1.1.1)*
Numeric Value for Strength (Denominator) (B.2.2.1.2)*	Units for Numeric Value for Strength (Denominator) (B.2.2.1.2.1)*
Active Ingredient Code (B.2.2.1.3):	
2nd Entry	
Active Ingredient(s) (B.2.2.1)*	

Numeric Value for Strength (Numerator) (B.2.2.1.1)*	Units for Numeric Value for Strength (Numerator) (B.2.2.1.1.1)*
Numeric Value for Strength (Denominator) (B.2.2.1.2)*	Units for Numeric Value for Strength (Denominator) (B.2.2.1.2.1)*
Active Ingredient Code (B.2.2.1.3):	
3rd Entry	
Active Ingredient(s) (B.2.2.1)*	

Numeric Value for Strength (Numerator) (B.2.2.1.1)*	Units for Numeric Value for Strength (Numerator) (B.2.2.1.1.1)*
Numeric Value for Strength (Denominator) (B.2.2.1.2)*	Units for Numeric Value for Strength (Denominator) (B.2.2.1.2.1)*
Active Ingredient Code (B.2.2.1.3):	

Dosage Form (B.2.2.2)

Lot Number (B.2.3)	Expiration Date (B.2.3.1) (dd/mm/yyyy)		
	Day Month Year		

Par	t B - Description of the AE (Continued)		
V	MP(s) Data and Usage (B.2) (Continued)		
The following fields (B.2.4-B.2.5.1) are applical	ble only if an animal is associated with the report.		
Who Administered the VMP? (B.2.4) (Select On	e)		
Veterinarian Animal Owner	Physician Patient Multiple Administrators		
Other Health Care Professional] Other 🔄 Unknown		
Use According to Label (B.2.5) (Select One)			
Yes No No Information			
Explanation for	or the Off-Label Use Code (B.2.5.1) (Select All That Apply)		
Was the target species Off-Label (B.2.5.1.1)	Was the indication Off-Label (B.2.5.1.6)		
Yes No Information	Yes No No Information		
Was the route of administration Off-Label (B.2.	5.1.2) Was the storage condition Off-Label (B.2.5.1.7)		
Yes No No Information	Yes No No Information		
Was the animal overdosed (B.2.5.1.3)	Was the product expired (B.2.5.1.8)		
Yes No No Information	Yes No No Information		
Was the animal underdosed (B.2.5.1.4)	Was there any other Off-Label issue (B.2.5.1.9)		
Yes No No Information	Yes No No Information		
Was the treatment regime Off-Label (B.2.5.1.5)			
Yes No No Information			
Proa	uct/Manufacturing Defect Information (B.2.6)		
The fields within this subsection (B.2	2.6.1-B.2.6.5) are applicable only if reporting a product/manufacturing defect.		
Manufacturing Site Identifier Number (B.2.6.1)	Manufacturer's Identifier Type (B.2.6.1.1) (select one if B.2.6.1 is given)		
	FEI Number DUNS Number		
Manufacturing Date (B.2.6.2) (dd/mm/yyyy)			
Day Month Year			
Number of Defective Items (B.2.6.3)	Defective Item Units (B.2.6.3.1)		
Number of Items Returned (B.2.6.4)	Returned Item Units (B.2.6.4.1)		
ORA District Field Office (B.2.6.5)			

Narrative of AE (B.3.1)*

AE Data (B.3)

Part B - Description of the AE (Continued)

AE Data (B.3) (Continued)

Narrative of AE (B.3.1)* (Continue, if needed)

Adverse Clinical Manifestations (B.3.2)*	Number of Animals (B.3.2.1)		Number of Animals 3.2.1.1)
		Actual	Estimated

Part B - Description	of the AE (Continued)
AE Data (B.3) (Continued)	
Date of Onset of AE/PP Found Date (B.3.3)* (dd/mm/yyyy)	
Day Month Year	
The following fields (B.3.4-B.5.1) are applicable only if an animal is associated with the report.	
Length of Time Between Exposure to VMP(s) and Onset of AE (B.3.4) (Select One)	
□ <2 Minutes □ <24 Hours □ <7 Days	>30 Days and <6 Months
<1 Hour	
<12 Hours	
Duration of AE (B.3.5)	
	on Time Units (B.3.5.1.1) (provide if B.3.5.1 is given) (Select One)
	Second Minute Hour Day Month Year
Serious AE (B.3.6)* (Select One)	Treatment of AE (B.3.7) (Select One)
	Yes No Unknown No Information
Outcome to Date (B.3.8) (Enter appropriate numbers where applicable)	
Ongoing (B.3.8.1) Recovered/Normal (B.3.8.2)	Recovered with Sequela (B.3.8.3)
Died (B.3.8.4) Euthanized (B.3.8.5)	Unknown (B.3.8.6)
Previous Exposure to the VMP? (B.3.9) (Select One)	Previous AE to the VMP? (B.3.10) (Select One)
Yes No Unknown No Information	Yes No Unknown No Information
Dechallenge - Rechallenge Information (B.4)	
Did AE Abate After Stopping the VMP? (B.4.1) (Select One)	
Yes No Unknown No Information Not Applicable	
Did AE Reappear After Re-introduction of the VMP? (B.4.2) (Select One)	
Yes No Unknown No Information Not Applicable	
Assessment of AE (B.5)	
Attending Veterinarian's Assessment (B.5.1) (Select One)	
🗌 Probable 📄 Possible 📄 Unlikely 📄 Unknown 📄 No Assessment 📄 No Attending Veterinarian	
Report Number(s) of Linked Report(s) (B.6)	
Unique AER Identification Number (B.6.1)	
Explanation for Linkage (B.6.1.1) (provide if B.6.1 is given) <i>(Select One)</i>	
Parent - Offspring Same patient Duplicate report Similar reports from same reporter (cluster) Other link type	
Supplemental Documents (B.7)	
Attached Document Name(s) (Filename(s) if Electronic) (B.7.1)	Attached Document Type(s) (B.7.1.1) (provide if B.7.1 is given)

Part B - Description of the AE (Continued)

HL7 ICSR Wrapper Data Elements (B.8)

Only sections B.8.2.2.3-B.8.2.2.8, B.8.2.5, and B.8.2.6 are relevant for submission of the paper form.

	rapper (B.8.1) (Identifier (B.8.1.1))
	/Identifier (B.8.1.1)*
Batch Number/Identifier - Root (B.8.1.1.1) Not Applicable for Paper Form	Batch Number/Identifier - Extension (B.8.1.1.2) Not Applicable for Paper Form
Batch Se	nder (B.8.1.2)
Batch Sender - Root (B.8.1.2.1)* Not Applicable for Paper Form	Batch Sender - Extension (B.8.1.2.2)* Not Applicable for Paper Form
Batch Sender - Title (B.8.1.2.3) Not Applicable for Paper Form	
Batch Sender - Last Name (B.8.1.2.4)* Not Applicable for Paper Form	Batch Sender - First Name (B.8.1.2.5)* Not Applicable for Paper Form
Batch Sender - Telephone (B.8.1.2.6)* Not Applicable for Paper Form	Batch Sender - Fax (B.8.1.2.7) Not Applicable for Paper Form
Batch Sender - Email (B.8.1.2.8)* Not Applicable for Paper Form	
Batch Receiver (B.8.1.3)	
Batch Receiver - Root (B.8.1.3.1)* USFDA	Batch Receiver - Extension (B.8.1.3.2) US Food and Drug Administration
Date of Batch Creation (B.8.1.4)* Not Applicable for Paper Form	VICH AER Version Number (B.8.1.5)* VICH AER 1.0.0
	n Wrapper (B.8.2)
	umber (B.8.2.1)*
Message Number - Root (B.8.2.1.1) Not Applicable for Paper Form	Message Number - Extension (B.8.2.1.2) Not Applicable for Paper Form
Pharmacovigilance Contact Person	for the MAH (Message Sender) (B.8.2.2)
Message Sender - Root (B.8.2.2.1) Not Applicable for Paper Form	Message Sender - Extension (B.8.2.2.2) Not Applicable for Paper Form
Title (Message Sender - Title) (B.8.2.2.3)	
Last Name (Message Sender - Last Name) (B.8.2.2.4)*	First Name (Message Sender - First Name) (B.8.2.2.5)*
Telephone (Message Sender - Telephone) (B.8.2.2.6)*	Fax (Message Sender - Fax) (B.8.2.2.7)
Email (Message Sender - Email) (B.8.2.2.8)*	
 Message R	eceiver (B.8.2.3)
Message Receiver - Root (B.8.2.3.1)*	Date of Message Creation (B.8.2.4)* Not Applicable for Paper Form
	Day Month Year
Report Identifier (B.8.2.5)* Domes	tic vs. Foreign Report Category (B.8.2.6)* <i>(Select One)</i> Domestic Soreign - Same Other Soreign - Similar
Profile Identifier (B.8.2.7)* Not Applicable for Paper Form (Select One)	
Adverse Event Adverse Event and Product Prob	lem Product Problem

This section applies only to requirements of the Paperwork Reduction Act of 1995.

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Department of Health and Human Services Food and Drug Administration Office of Operations Paperwork Reduction Act (PRA) Staff *PRAStaff@fda.hhs.gov*

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