

FOLD, SEAL, AND RETURN

FOOD AND DRUG ADMINISTRATION
7500 Standish Place (HFV-199), Room N403
Rockville, MD 20855

**VETERINARY ADVERSE DRUG REACTION, LACK OF
EFFECTIVENESS OR PRODUCT DEFECT REPORT**
(For VOLUNTARY Reporting)

Form Approved: OMB 0910-0645
Expiration Date:

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.

Individual Case Safety Report Number (FDA Assigned Number)

Submission Type Initial Follow-up

Report Type Adverse Event Product Problem Both Adverse Event and Product Problem

Date of this Report Month Day Year

Date of Initial Report (If this report is a Follow-up) Month Day Year

Sender Information

First Name Last Name

Street Address City

State Postal/Zip Code

Country Telephone Number

Telephone Number (Other) Fax Number

e-Mail Address

Sender Category Veterinarian Animal Owner Physician Patient
 Other Health Care Professional Other Unknown

Sender Previously Reported to the Manufacturer? Yes No

(If Yes, provide the Manufacturer's Case Number below)

Manufacturer's Case Number

No Identity Disclosure (If you do NOT want your identity disclosed to the manufacturer, place an "X" in this box)

Preferred Method of Contact Telephone e-Mail

Healthcare Professional Information (If different from Sender Information)

First Name Last Name

Street Address City

State Postal/Zip Code

Country Telephone Number

Telephone Number (Other) Fax Number

e-Mail Address

Owner Information (If different from Sender Information)

First Name Last Name

Street Address City

State Postal/Zip Code

Country Telephone Number

Telephone Number (Other) Fax Number

e-Mail Address

Suspected Product Information

Name of Suspected Product

Diagnosis and/or Reason for Use of the Product Dosage Form (Chewable, liquid, tablet, topical, injection, etc.)

Date of First Exposure Month Day Year Date of Last Exposure Month Day Year

Duration of Product Use

Product Use Information for Suspected Product

Dose Administered Interval of Administration (Frequency) Route of Administration

Product Administered By Veterinarian / Veterinary Staff Owner Other

Lot Number Expiration Date Month Day Year

Name of Manufacturer of Suspected Product

Adverse Event Information

Veterinarian's Level of Suspicion that Product Caused the Adverse Event High Medium Low Unknown

Treatment of Adverse Event

Did Adverse Event Abate After Stopping the Product? Yes No Not Applicable

Did Adverse Event Reappear After Reintroduction of the Product? Yes No Not Applicable

Outcome Recovered Died Other

Species

- Budgerigar Cat Cattle Cockatiel Cockatoo
- Dog Ferret Fish Goat Guinea Pig
- Horse Human Parrot Pig Rabbit
- Sheep Other

Breed

Gender Male Male Neutered Female Female Neutered

Age Weight

Overall Health Status When Suspected Product Given

- Excellent Good Fair Poor Critical

Number of Animals Treated Number of Animals Affected

Date of Onset of Adverse Event Month Day Year

Length of Time Between First Exposure to Suspected Product(s) and Onset of Adverse Event

Length of Time Between Last Administration of Suspected Product(s) and Onset of Adverse Event

When the Adverse Event Occurred, Treatment with Suspected Product

- Had Already Been Completed Was Discontinued
- Was Discontinued and Replaced with Another Product Was Discontinued and Reintroduced Later
- Was Continued at an Altered Dose Other (Comment Below)

Attached Document Name (Filename if Electronic)

Attached Document Description

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Concurrent Clinical Problem(s)

Were There Concurrent Clinical Problems? Yes No Do Not Know

List Concurrent Clinical Problem(s)

Concurrent Product Information (Excluding Treatment of Current Event)

Please provide name(s), dose(s), interval(s), date(s) of treatment(s), and other relevant information to describe other products that the patient was taking at the time of the event. Either copy this page as needed to describe the additional products or provide comments in the narrative box at the end of the form.

Were Concurrent Products Given? **Yes** **No** **Do Not Know**

List Names of Concurrent Products Administered

Date of First Exposure

Month

Day

Year

Date of Last Exposure

Month

Day

Year

Duration of Product Use

Adverse Event/Product Problem (Long Narrative)

INSTRUCTIONS FOR COMPLETING AND SUBMITTING FORM FDA 1932a

GENERAL INSTRUCTIONS

- Please either type or print all entries in a font no smaller than 8 point. If filling in the form by hand, please use black ink.
- Please complete all sections that apply.
- For narrative entries, attach additional pages as needed.
- If attaching additional pages, please do the following:
 - Identify all attached pages as Page # of # (e.g., Page 1 of 4);
 - Indicate the appropriate section and block number next to the narrative continuation; and
 - Include the phrase *continued* at the end of each field that has additional information continued onto another page.

Individual Case Safety Report Number: This number will be assigned by the Food and Drug Administration (FDA).

Submission Type: Choose a Submission Type. If this is the first time you have sent FDA information about this, choose "Initial" report. If this is additional information for a previously submitted report, choose "Follow-up" report.

Report Type: Choose a Report Type. If you are reporting something that has affected an animal or a human, including lack of effectiveness, choose "Adverse Event." If you are reporting something associated with a product (such as crumbled tablets or peculiar appearance), choose "Product Problem." If both situations apply, choose "Both."

Date of this Report and Date of Initial Report: Enter dates as mm/dd/yyyy. If exact dates are unknown, provide the best estimate.

Sender Information: Provide the contact information for the person who is filling out this form.

Reporter Category: Choose the appropriate Reporter Category.

Manufacturer's Case Number: Fill in the case number, if applicable or known. If you previously reported to the manufacturer, you can contact the manufacturer for the manufacturer's case number.

Healthcare Professional Information: Please provide the name, mailing address, phone number, and e-mail address of the veterinarian or other health care professional who can be contacted to provide information, if such follow-up is necessary.

If the Health Care Professional is also the Sender, there is no need to repeat the information.

Owner's Name: Please provide the owner's name, mailing address, and phone number. If the Owner is also the Sender, there is no need to repeat the information.

The owner's information is held in strict confidence by FDA and protected to the fullest extent of the law. **The FDA will not disclose the reporter's identity in response to a request from the public, pursuant to the Freedom of Information Act.**

Name of Suspected Product: Provide the brand name of the product.

Diagnosis and/or Reason for Use: Provide the reason or indication for which the product was prescribed or used in the animal.

Dosage Form: Provide the dosage form (e.g., chewable tablet, liquid, tablet, topical, injection, etc.)

Date(s) of First and Last Exposure: Enter the date(s) the product was started and stopped. If actual dates are unknown, enter the approximate time period the product was used in the Duration of Product Use box (e.g., 2 weeks during the summer of 2006). If the product was used less than 1 day, enter the same date in the Date of First Exposure and Date of Last Exposure boxes.

Dose Administered, Interval of Administration (Frequency), and Route of Administration: Describe how the product was administered (e.g., 250 mg), frequency of administration (e.g., every 12 hours for 5 days), and how it was administered (e.g., orally, injection, etc.). Describe how the product was administered, even if it differs from what was prescribed.

Product Administered By: Please check the appropriate box. If given by a member of the veterinarian's staff, please identify (e.g., technician, assistant) in the narrative box at the end of the form. If given by someone other than the owner (e.g., pet sitter, trainer), choose "Owner" but identify in the narrative.

Lot Number and Expiration Date: Please provide the lot number and expiration date from the product, if available.

Name of Manufacturer of Suspected Product: Provide the name of the manufacturer.

Treatment of Adverse Event: If the adverse event was treated, describe the treatment given.

Did Adverse Event Abate after Stopping the Product?

Choose "Yes" if the adverse event lessened or went away when the product was stopped or the dose was decreased.

Choose "Not Applicable" if the product was not stopped or decreased.

Outcome: Choose an Outcome for the adverse event. If "Other" is chosen, describe this outcome in the narrative at the end of the form (e.g., the dog lived but never recovered fully, since it was left with a permanent elevation of liver enzymes).

Species: Choose a box for species (e.g., cat, dog, ferret, horse, human, other, etc.). If "Other" is chosen, identify the species in the narrative box at the end of the form.

Breed: Enter the breed (e.g., Yorkie, Mixed Breed, Lab mix, Siamese/Persian mix). Note: This category is not applicable if the patient is human.

Age: Provide the patient's age at the time of the adverse event, including a time descriptor (e.g., 8 years). Provide the best estimate if exact age is unknown.

Weight: Provide the patient's weight in pounds (lb). Make a best estimate if exact weight is unknown.

Overall Health Status: Check the box that best describes the patient's overall state of health when drug/product was first given.

Number of Animals Treated: If more than one animal was treated with the same drug/product at the same time, please tell us how many were treated (e.g., two kittens received Drug X).

Number of Animals Affected: If more than one animal had an adverse event after the treatment, please tell us how many. If more than one animal had an adverse event, and the reaction was not the same, please submit a separate report for each animal (e.g., two kittens received Drug X, a de-worming medication. One vomited and wouldn't eat for several days, while the other had a seizure).

Date of Onset: Provide the date when the adverse event first started.

Length of Time Between Exposure to Suspected Product and Onset of Adverse Event: Enter the length of time from the first day the product was given to the onset of the adverse event (e.g., 3 days).

Length of Time Between Last Administration of Suspected Product(s) and Onset of Adverse Event: Enter the length of time from the last dose of the product to the onset of the adverse event (e.g., 3 hours).

When the Reaction Occurred, Treatment with Suspected Product:

Check the appropriate box that applies to the reported adverse event. If “Other” is checked, provide comments in narrative box at the end of the form.

Attached Document File Name:

If attaching any supporting documents, such as letters, medical records, or photos, provide the name of the file here.

Examples:

- Documents for Princess.doc.
- Spreadsheet of Princess' laboratory results.xls
- Photographs of Princess before and after treatment.jpg
- Newspaper article about the product.pdf

If you mail your report, these attachments should accompany the paper Form FDA 1932a.

Attached Document Description: If attaching any supporting documents, provide the description of the contents (e.g., medical records, lab tests, photograph, newspaper article, etc.)

Concurrent Clinical Problem(s): Provide information on other known health problems of the patient at the time of exposure to the product (e.g., chronic allergic dermatitis, intermittent vomiting, allergic reaction following vaccination). Check “None” if there are no known concurrent problems.

Concurrent Product Information: Please provide names, doses, and dates of treatments for products that the patient was taking at the time of the event. **Do** include over-the-counter products, such as supplements, vitamins, and homeopathic preparations. **Do not** include products used to treat the event. Check “None” if nothing else was being given at the time of the adverse event.

Adverse Event/Product Defect (Long Narrative): Use this space to describe the event, possible contributing factors, and outcome. Include a description of what happened and a summary of all available clinical information.