

**FATAL-PLUS SOLUTION - pentobarbital sodium injection, solution**  
**Vortech Pharmaceuticals, Ltd.**

*Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.*

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**Description**

**Fatal-Plus Solution**

FOR VETERINARY USE ONLY

For Euthanasia of Animals

**DOSAGE AND ADMINISTRATION:**

Intravenous injection is the preferred route. However, intraperitoneal or intracardiac injections may be made where the intravenous injection is impractical, as in the very small dog and cat, or in the comatose animal with depressed vascular function. Inject rapidly 1 mL for every 10 lbs. body weight Minimum 1 mL.

**WARNING: THIS IS A DENATURED SOLUTION FOR ANIMAL EUTHANASIA ONLY. POISONOUS IF TAKEN INTERNALLY.** Must not be used for therapeutic purposes. Do not use in animals intended for food.

**ENVIRONMENTAL HAZARD:**

This product is toxic to wildlife. Birds and mammals feeding on treated animals may be killed. Birds and mammals feeding on treated animals may be killed. Euthanized animals must be properly disposed of by deep burial, incineration, or other method in compliance with state and local laws, to prevent consumption of carcass material by scavenging wildlife.

**CAUTION:** Federal law restricts this drug to use by or on the order of a licensed veterinarian.

**POISON: KEEP OUT OF THE REACH OF CHILDREN**

**Principle Display Panel**



**Fatal-Plus Solution - NDC: 0298-9373-68**

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<b>FATAL-PLUS SOLUTION</b>				
pentobarbital sodium injection, solution				
<b>Product Information</b>				
<b>Product Type</b>	PRESCRIPTION ANIMAL DRUG	<b>Item Code (Source)</b>	NDC:0298-9373	
<b>Route of Administration</b>	INTRAVENOUS, INTRAPERITONEAL, INTRACARDIAC	<b>DEA Schedule</b>	CII	
<b>Active Ingredient/Active Moiety</b>				
<b>Ingredient Name</b>		<b>Basis of Strength</b>	<b>Strength</b>	
Pentobarbital Sodium (UNII: NJJ0475N0S) (Pentobarbital - UNII:I4744080IR)		Pentobarbital Sodium	390 mg in 1 mL	
<b>Product Characteristics</b>				
<b>Color</b>	blue (DARK BLUE)	<b>Score</b>		
<b>Shape</b>		<b>Size</b>		
<b>Flavor</b>		<b>Imprint Code</b>		
<b>Contains</b>				
<b>Packaging</b>				
<b>#</b>	<b>Item Code</b>	<b>Package Description</b>	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
1	NDC:0298-9373-68	250 mL in 1 VIAL, MULTI-DOSE		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		06/14/1982	

**Labeler** - Vortech Pharmaceuticals, Ltd. (052399276)

**Registrant** - Vortech Pharmaceuticals, Ltd. (052399276)

## Establishment

Name	Address	ID/FEI	Business Operations
Vortech Pharmaceuticals, Ltd		052399276	manufacture

## Establishment

Name	Address	ID/FEI	Business Operations
Siegfried USA, LLC		001213784	api manufacture

Revised: 12/2016

Vortech Pharmaceuticals, Ltd.