# FERROUS SULFATE- ferrous sulfate tablet Richmond Pharmaceuticals, Inc.

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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#### FERROUS SULFATE TABLETS 325 mg

#### Supplement Facts [/S]

Serving Size: 1 Tablet Amount per Tablet	% Daily Value
Iron (as ferrous sulfate) 65 mg	360 %

#### SUGGESTED USE

**Adults:** One (1) tablet daily, preferably after meals or as directed by the doctor. **Children:** As directed by the doctor.

#### **Active Ingredient**

## EACH TABLET CONTAINS: % U.S. RDA\*

Elemental Iron 65 mg 360

(Equivalent to 325 mg of Ferrous Sulfate)

\* U.S. Recommended Daily Allowance

#### **Inactive Ingredients**

Croscarmellose sodium, dicalcium phosphate, FD&C RED#40 (Al-lake), FD&C yellow #6 (Al-lake), hypromellose, magnesium stearate, microcrystalline cellulose, PEG 400, titanium dioxide

#### Purpose

Iron Supplement

#### WARNING:

Accidental overdose of iron containing products is a leading cause of fatal poisoning in children under 6. **Keep this product out of reach of children**. In case of accidental overdose, call a doctor or Poison Control Center immediately.

**WARNINGS: Do not exceed recommended dosage.** The treatment of any anemic condition should be under the advice and supervision of doctor. Occasional gastrointestinal discomfort (such as nausea) may be minimized by taking with meals. Iron-containing medication may occasionally cause constipation or diarrhea.

As with any drug, if you are pregnant or nursing baby, seek the advice of a health professional before using this product.

# **DRUG INTERACTION PRECAUTION**

Since oral iron products interfere with absorption of oral tetracycline antibiotics , these products should not be taken within two hours of each other.

# DOSAGE AND ADMINISTRATION

**Each tablet contains:** Calcium 20 mg (2% daily value) Store in a dry place at controlled room temperature at 15-30 °C (59°-86° F). Do not expose to excessive heat or moisture.

Questions or Comments DO NOT USE IF IMPRINTED SAFETY SEAL UNDER CAP IS MISSING OR DAMAGED Call 804-270-4498 Monday-Friday, 8.30 am – 4.30 pm ET

Package Label

FERROUS SULFATE TABLETS 325 mg (5 gr)

**Iron Supplement** 

NDC: 54738-963-01 - 100 Tablets

## NDC: 54738-963-03 - 1000 TABLETS



# FERROUS SULFATE

ferrous sulfate tablet

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Product Type		HUMAN OTC DR	UG	Ite m C	Code (Source) NDC:54738-9		-963		
Route of Administra	tion	ORAL							
Active Ingredient	t/Active M	biety							
	Ingredient Name				Basis of	Basis of Strength Str			
-						FERROUS	CATION	325 mg	
Inactive Ingredie	nts								
		Ingrediei	nt Name				Strength		
CROSCARMELLOSE	SODIUM (UN	II: M280L1HH48)							
DIBASIC CALCIUM PI	HO SPHATE D	IHYDRATE (UNII: O	7TSZ97GEP)						
FD&C RED NO.40 (U	NII: WZB91272	KOA)							
FD&C YELLOW NO.6 (UNII: H77VEI93A8)									
HYPROMELLOSES (UNII: 3NXW29V3WO)									
MAGNESIUM STEARATE (UNII: 70097M6I30)									
CELLULOSE, MICRO	CRYSTALLI	NE (UNII: OP1R32D61	lU)						
POLYETHYLENE GLYCOL 400 (UNII: B697894SGQ)									
TITANIUM DIO XIDE (									
TTANIOW DIO AIDE (	UNII: 15FIX9V	2JP)							
TTAINOW DIOAIDE (	UNII: 15F1X9 V	2JP)							
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Labeler - Richmond Pharmaceuticals, Inc. (043569607)

Registrant - Advance Pharmaceutical Inc. (078301063)

# Establishment

Name	Address	ID/FEI	Business Operations
Advance Pharmaceutical Inc.		078301063	manufacture(54738-963)

Revised: 12/2019

Richmond Pharmaceuticals, Inc.