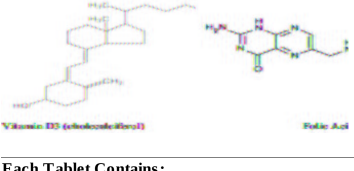
# FOLVIK-D- folic acid and cholecalciferol tablet, coated SLV PHARMACEUTICALS LLC DBA AUM PHARMACEUTICALS

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FOLVIK-D Tablets Rx Only

#### DESCRIPTION

**FOLVIK-D** tablet is an orally administered prescription strength folate product for the dietary management of patients with unique nutritional needs requiring increased folate levels, Vitamin D levels, or used in improving the nutritional status of patients with folic acid and vitamin D deficiency. **FOLVIK-D** should be administered under the supervision of a licensed medical practitioner. Vitamin D3 (cholecalciferol) is a white, crystalline powder, very soluble in water. Folic acid occurs as a yellow or yellowishorange crystalline powder and is very soluble in water and insoluble in alcohol. The structural formula of Vitamin D3 and folic acid are as follows:



<u>Each Tablet Contains:</u>	
Folic Acid	1000 MCG
Vitamin D3 (Cholecalciferol)	94.375 MCG

Each tablet contains the following inactive ingredients: Microcrystalline Cellulose, Dicalcium Phosphate, Croscarmellose Sodium, Magnesium Stearate, Silicon Dioxide, Stearic acid, HPMC E15, HPMC E5/E6, PEG 8000.

## INDICATIONS AND USAGE

**FOLVIK-D** is indicated for dietary management of patients with unique nutritional needs requiring increased folate levels, Vitamin D levels, or used in improving the nutritional status of patients with folic acid and vitamin D deficiency.

## **DOSAGE:**

Usual adult dose is 1 tablet once or twice daily or as prescribed by a licensed medical practitioner.

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This statement has not been evaluated by the FDA. This product is not intended to diagnose, treat, cure, or prevent disease.

# CLINICAL PHARMACOLOGY

The *in vivo* synthesis of the major biologically active metabolites of vitamin D occurs in two steps. The first hydroxylation of Vitamin D takes place in the liver (to 25-hydroxyvitamin D) and the second in the kidneys (to 1, 25-dihydroxyvitamin D). Vitamin D metabolites promote the active absorption of calcium and phosphorus by the small intestine, thus elevating serum calcium and phosphate levels sufficiently to permit bone mineralization. Vitamin D metabolites also mobilize calcium and phosphate from bone and

probably increase the reabsorption of calcium and perhaps also of phosphate by the renal tubules.

There is a time lag of 10 to 24 hours between the administration of vitamin D and the initiation of its action in the body due to the necessity of synthesis of the active metabolites in the liver and kidneys. Parathyroid hormone is responsible for the regulation of this metabolism in the kidneys.

Folic acid is the precursor of tetrahydrofolic acid, which is involved as a cofactor for transformylation reactions in the biosynthesis of purines and thymidylates of nucleic acids. Impairment of thymidylate synthesis in patients with folic acid deficiency is thought to account for the defective deoxyribonucleic acid (DNA) synthesis that leads to megaloblast formation and megaloblastic and macrocytic anemias.

Folic acid is absorbed rapidly from the small intestine, primarily from the proximal portion. Naturally occurring conjugated folates are reduced enzymatically to folic acid in the gastrointestinal tract prior to absorption. Folic acid appears in the plasma approximately 15 to 30 minutes after an oral dose; peak levels are generally reached within 1 hour.

# CONTRAINDCATIONS

This product is contraindicated in patients with a known hypersensitivity to any of the ingredients. **FOLVIK-D** is contraindicated in patients with hypercalcemia, malabsorption syndrome, abnormal sensitivity to the toxic effects of vitamin D, and hypervitaminosis D.

# WARNINGS AND PRECAUTIONS

KEEP OUT OF THE REACH OF CHILDREN. In case of an accidental overdose, call a doctor or a poison control center immediately.

Tell your doctor if you have: kidney problems or thyroid disease.

This medication should be used as directed by your physician during pregnancy or while breastfeeding. Consult your doctor about the risks and benefits.

Folic acid alone is improper therapy in the treatment of pernicious anemia and other megaloblastic anemias where vitamin B12 is deficient. Folic acid in doses above 0.1 mg daily may obscure pernicious anemia in that hematologic remission can occur while neurological manifestations progress.

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There is evidence that the anticonvulsant action of phenytoin is antagonized by folic acid. A patient whose epilepsy is completely controlled

by phenytoin may require increased doses to prevent convulsions if folic acid is given.

# ADVERSE REACTIONS

Allergic sensitization has been reported following both oral and parenteral administration of folic acid.

Call your doctor if you experience any of the following rare but possible signs of hypervitaminosis D: nausea, vomiting, constipation, loss of appetite, increased thirst, increased urination, mental/mood changes or unusual tiredness.

You should call your doctor for medical advice about serious adverse events. To report adverse side effects or to obtain product information, contact Aum pharmaceuticals, at 1-866-760-6565

# HOW SUPPLIED

**FOLVIK - D** tablets are dispensed in bottles of 90 ct (73317-4090-9) **FOLVIK - D** tablets are clear coated and caplet shaped tablets

All prescriptions using this product shall be pursuant to state statutes as applicable. This product is an

Rx only and may be administered only under a physician's supervision. There are no implied or explicit claims on therapeutic equivalence.

#### STORAGE

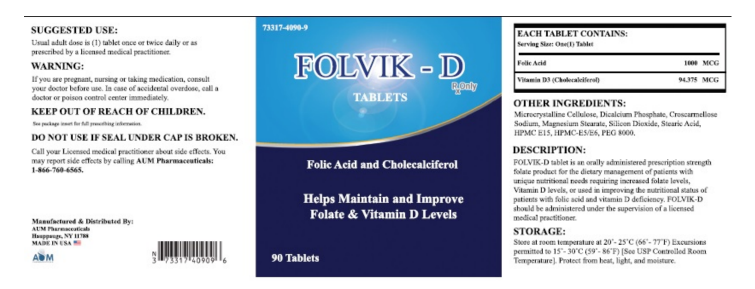
KEEP OUT OF THE REACH OF CHILDREN

Store at 20°-25°C (66°-77°F); excursions permitted to 15°-30°C (59°-86°F) [See USP Controlled Room Temperature.] Protect from heat, light and moisture.

#### **Manufactured for:**

AUM Pharmaceuticals

Hauppauge, NY 11788



# FOLVIK-D

folic acid and cholecalciferol tablet, coated

Product Information					
Product Type	DIETARY SUPPLEMENT	Item Code (Source	NHRIC:73317	2:73317-4090	
Route of Administration	ORAL				
Active Ingredient/Active M	oiety				
	Ingredient Name		Basis o	of Strength	Strength
FOLIC ACID (UNII: 935E97BOY8) (I	D (UNII: 935E97BOY8) (FOLIC ACID - UNII:935E97BOY8) FOLIC ACID		CID	1000 ug	
CHOLECAL CHERDOL (UNIL 100317		1C6V77QF41) CHOLECALCIFERO			
CHOLECALCIFEROL (UNII: 1C6V/	7QF41) (CHOLECALCIFEROL - UN	I:1C6V77QF41)	CHOLEC	ALCIFEROL	94.375 ug
CHOLECALCIFEROL (UNII: 1C6 V/	/QF41) (CHOLECALCIFEROL - UN	I:1C6 V77QF41)	CHOLEC	ALCIFEROL	94.375 ug
	/QF41) (CHOLECALCIFEROL - UN	I:1C6V77QF41)	CHOLEC	ALCIFEROL	94.375 ug
Inactive Ingredients	/QF41) (CHOLECALCIFEROL - UN	I:1C6V77QF41)	CHOLEC	ALCIFEROL	94.375 ug
	/QF41) (CHOLECALCIFEROL - UN Ingredient Name	I:1C6V77QF41)	CHOLEC		94.375 ug strength
	Ingredient Name	I:1C6V77QF41)	CHOLEC		
Inactive Ingredients	Ingredient Name SE (UNII: OP1R32D6 1U)	I:1C6V77QF41)	CHOLEC		
Inactive Ingredients MICROCRYSTALLINE CELLULOS	Ingredient Name SE (UNII: OP1R32D6 1U) ANHYDRO US (UNII: L11K75P92J)	I:1C6V77QF41)	CHOLEC		
Inactive Ingredients MICRO CRYSTALLINE CELLULO S CALCIUM PHO SPHATE, DIBASIC,	Ingredient Name SE (UNII: OP1R32D61U) ANHYDROUS (UNII: L11K75P92J) II: M28OL1HH48)	I:1C6V77QF41)	CHOLEC		

YPROMELLOSES (UNI	I: 3NXW29V3WO)				
OLYETHYLENE GLYC	<b>DL 8000</b> (UNII: Q662QK8M3B)				
Packaging					
# Item Code	Package Description	Marketin	ng Start Date	Ma	arketing End Date
NHRIC:73317-4090-9	90 in 1 BOTTLE				
Marketing Information					
	Application Number or Monograp	h Citation	Marketing Start	Date	Marketing End Dat
Marketing Category					

Supplement F	acts	
Serving Size :		Serving per Container :
	Amount Per Serving	% Daily Value
color		
scoring	1	
shape		
size (solid drugs)	3 mm	

Labeler - SLV PHARMACEUTICALS LLC DBA AUM PHARMACEUTICALS (081225162)

**Registrant** - SLV PHARMACEUTICALS LLC DBA AUM PHARMACEUTICALS (081225162)

# Establishment

Name	Address	ID/FEI	<b>Business Operations</b>
SLV PHARMACEUTICALS LLC DBA AUM PHARMACEUTICALS		081225162	

Revised: 4/2020

SLV PHARMACEUTICALS LLC DBA AUM PHARMACEUTICALS