

**PSORINOHEEL - sulfur, thuja occidentalis leafy twig, vaccinia virus strain new york city board, gonorrhoeal urethral secretion human, sodium chloride, nerium oleander leaf, cicutu virosa root, solution**  
**Heel Inc**

*Disclaimer: This homeopathic product has not been evaluated by the Food and Drug Administration for safety or efficacy. FDA is not aware of scientific evidence to support homeopathy as effective.*

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**Psorinoheel oral vial**

**DESCRIPTION**

**Each 1.1 ml ampule contains:**

Ingredient Name	Potency	Quantity
<b>Bacillinum</b>	<b>12x</b>	0.55µ
<b>Bufo rana</b>	<b>10x</b>	1.1µ
<b>Cictuta virosa</b>	<b>5x</b>	0.55µ
<b>Kali bismuthum iodatum</b>	<b>5x</b>	0.55 µ
<b>Medorrhinum</b>	<b>12x</b>	1.1µ
<b>Natrum muriaticum</b>	<b>12x</b>	1.1µ
<b>Oleander</b>	<b>4x</b>	0.55µ
<b>Psorinum</b>	<b>10x</b>	1.1µ
<b>Sulphur</b>	<b>6x</b>	1.1µ
<b>Thuja occidentalis</b>	<b>6x</b>	1.1µ
<b>Vaccinotoxinum</b>	<b>8x</b>	1.1µ

**Inactive Ingredient:** Isotonic Sodium Chloride solution

**INDICATION AND USAGE**

Psorinoheel® Oral Vials is a homeopathic drug product indicated for the stimulation of the defense mechanism in chronic skin disorders, hepatic damage , and chronic illness

**DOSAGE AND ADMINISTRATION**

Dosage:

**Adults and children above 6 years:** 1 vial orally 1-3 times daily

**Children up to 6 years:** ½ vial orally 1-3 times daily

**CONTRAINDICATIONS**

Contraindications

Psorinoheel® Oral Vials are contraindicated in patients with known hypersensitivity to Psorinoheel® or any of its ingredients.

**WARNINGS AND PRECAUTIONS**

## **Warnings and Precautions**

None

## **ADVERSE REACTIONS**

No adverse events have been reported with a causal relationship Psorinoheel® Oral Vials.

## **OVERDOSAGE**

**Overdosage:** No negative effects of an overdose have been reported and none are expected due to the homeopathic dilutions

## **CLINICAL PHARMACOLOGY**

### **Mechanism of Action**

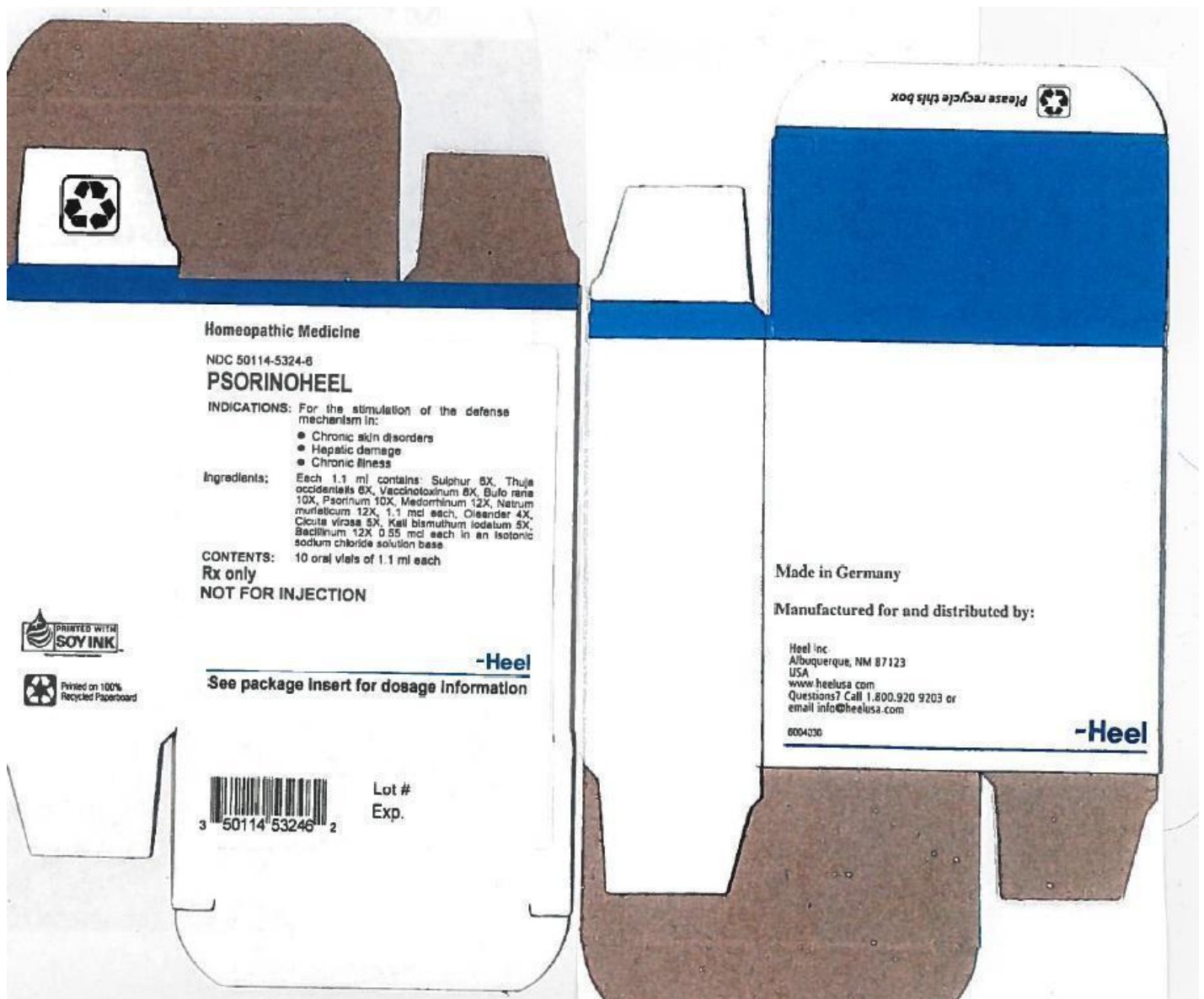
The exact mechanism of Psorinoheel® Oral Vials is not fully understood.

### **Pharmacodynamics**

Not applicable for homeopathic medicinal products.

## **DOSAGE**

1 oral vial containing 1.1 ml solution for oral administration



## PSORINOHEEL

sulfur, thuja occidentalis leafy twig, vaccinia virus strain new york city board, gonorrheal urethral secretion human, sodium chloride, nerium oleander leaf, cicuta virosa root, solution

### Product Information

<b>Product Type</b>	HUMAN PRESCRIPTION DRUG	<b>Item Code (Source)</b>	NDC:50114-5324
<b>Route of Administration</b>	ORAL		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SULFUR (UNII: 70FD1KFU70) (SULFUR - UNII:70FD1KFU70)	SULFUR	6 [hp_X] in 1.1 mL
THUJA OCCIDENTALIS LEAFY TWIG (UNII: 1NT28V9397) (THUJA OCCIDENTALIS LEAFY TWIG - UNII:1NT28V9397)	THUJA OCCIDENTALIS LEAFY TWIG	6 [hp_X] in 1.1 mL
VACCINIA VIRUS STRAIN NEW YORK CITY BOARD OF HEALTH LIVE ANTIGEN (UNII: 4SV59689SK) (VACCINIA VIRUS STRAIN NEW YORK CITY BOARD OF HEALTH LIVE ANTIGEN - UNII:4SV59689SK)	VACCINIA VIRUS STRAIN NEW YORK CITY BOARD OF HEALTH LIVE ANTIGEN	8 [hp_X] in 1.1 mL

<b>BUFO BUFO CUTANEOUS GLAND</b> (UNII: Q59QU6N72Q) (BUFO BUFO CUTANEOUS GLAND - UNII:Q59QU6N72Q)	BUFO BUFO CUTANEOUS GLAND	10 [hp_X] in 1.1 mL
<b>SCABIES LESION LYSATE (HUMAN)</b> (UNII: 5UAU16Z1U4) (SCABIES LESION LYSATE (HUMAN) - UNII:5UAU16Z1U4)	SCABIES LESION LYSATE (HUMAN)	10 [hp_X] in 1.1 mL
<b>GONORRHEAL URETHRAL SECRETION HUMAN</b> (UNII: 9BZG9E38F) (GONORRHEAL URETHRAL SECRETION HUMAN - UNII:9BZG9E38F)	GONORRHEAL URETHRAL SECRETION HUMAN	12 [hp_X] in 1.1 mL
<b>NERIUM OLEANDER LEAF</b> (UNII: 7KV510R6H6) (NERIUM OLEANDER LEAF - UNII:7KV510R6H6)	NERIUM OLEANDER LEAF	4 [hp_X] in 1.1 mL
<b>CICUTA VIROSA ROOT</b> (UNII: YEA9P21S8N) (CICUTA VIROSA ROOT - UNII:YEA9P21S8N)	CICUTA VIROSA ROOT	5 [hp_X] in 1.1 mL
<b>POTASSIUM DICHROMATE</b> (UNII: T4423S18FM) (DICHROMATE ION - UNII:9LKY4BFN2V)	POTASSIUM DICHROMATE	5 [hp_X] in 1.1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:50 114-5324-6	10 in 1 CARTON		
1		1.1 mL in 1 AMPULE		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved homeopathic		01/31/1993	

**Labeler** - Heel Inc (102783016)

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Heel Inc