# NATRUM SALICYLICUM- natrum salicylicum pellet Rxhomeo Private Limited d.b.a. Rxhomeo, 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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#### **ACTIVE INGREDIENT**

NATRUM SALICYLICUM HPUS 2X and higher

#### **USES**

Post-Flu Tonic

#### **INDICATIONS**

Condition listed above or as directed by the physician

#### **DOSAGE**

Adults- Take 4 or 6 Pellets by mouth, three times daily or as suggested by physician. Children 2 years and older- take 1/2 the adult dose.

#### **WARNINGS**

This product is to be used for self-limiting conditions

If symptoms do not improve in 4 days, or worsen, discontinue use and seek assistance of health professional

As with any drug, if you are preganant, or nursing a baby, seek professional advice before taking this product

Keep this and all medication out of reach of children

#### **INACTIVE INGREDIENTS**

Sucrose

#### **STORAGE**

Store in a cool dark place

## **QUESTIONS OR COMMENTS**

www.Rxhomeo.com | 1.888.2796642 | info@rxhomeo.com

Rxhomeo, Inc 9415 Burnet Road, Suite 312, Austin, TX 78758



# **NATRUM SALICYLICUM 2X**

Active Ingredients: As above Inactive Ingredients: Sucrose USES: Post-Flu Tonic





Jactured according to the pathic Pharmacopoeia of the ad States Est. # 30052969310 fo@rxhomeo.com m 1-888-BRYONIA ( 2796642 )





Distributed in the US by Rxhomeo, Inc 9415 Burnet Road, Suite 312, Austin, TX 78758 Manufactured by: Rxhomeo Private Limited "Indradhanush", 4-1-424 to 426, Bank Street, Abids, Hyderabad #500001 India.

NDC: 15631-0311-0 Batch No: XXXXXXXX Contents 100 Pellets

## NATRUM SALICYLICUM

natrum salicylicum pellet

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:15631-0311
Route of Administration	ORAL		

**Active Ingredient/Active Moiety Ingredient Name Basis of Strength** Strength SODIUM SALICYLATE (UNII: WIQ 1H8 5 SYP) (SALICYLIC ACID - UNII: O414PZ4LPZ) SODIUM SALICYLATE 2 [hp\_X]

Inactive Ingredients	
Ingredient Name	Strength
SUCROSE (UNII: C151H8M554)	

1	Packaging				
#	# Item Code	Package Description	<b>Marketing Start Date</b>	<b>Marketing End Date</b>	
1	NDC:15631-0311-0	100 in 1 PACKAGE; Type 0: Not a Combination Product	0 1/0 1/20 18		
2	NDC:15631-0311-1	200 in 1 PACKAGE; Type 0: Not a Combination Product	0 1/0 1/20 18		
3	NDC:15631-0311-2	400 in 1 PACKAGE; Type 0: Not a Combination Product	0 1/0 1/20 18		
4	NDC:15631-0311-3	750 in 1 PACKAGE; Type 0: Not a Combination Product	0 1/0 1/20 18		
5	NDC:15631-0311-4	2500 in 1 PACKAGE; Type 0: Not a Combination Product	0 1/0 1/20 18		
6	NDC:15631-0311-5	12500 in 1 PACKAGE; Type 0: Not a Combination Product	0 1/0 1/20 18		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
unapproved homeopathic		09/10/2015		

Labeler - Rxhomeo Private Limited d.b.a. Rxhomeo, Inc (650833994)

Establishment				
Name	Address	ID/FEI	Business Operations	
Rxhomeo Private Limited d.b.a. Rxhomeo, Inc		650833994	manufacture(15631-0311), label(15631-0311)	

Revised: 3/2020

Rxhomeo Private Limited d.b.a. Rxhomeo, Inc