

Australian Government

Department of Health

Therapeutic Goods Administration

Public Summary

Summary for ARTG Entry:	260025	Prospan Drops
ARTG entry for	Medicine Listed	
Sponsor	SFI Australasia	
Postal Address	PO Box 1027, Cl Australia	ROWS NEST, NSW, 1585
ARTG Start Date	16/09/2015	
Product Category	Medicine	
Status	Active	
Approval Area	Listed Medicines	

Conditions

Colouring agents used in listed medicine for ingestion, other than those listed for export only under section 25 of the Act, shall be only those included in the list of 'Colourings permitted in medicines for oral use'.

The sponsor shall keep records relating to this listed medicine as are necessary to: (a) Expedite recall if necessary of any batch of the listed medicine, (b) Identify the manufacturer(s) of each batch of the listed medicine. Where any part of or step in manufacture in Australia of the listed medicine is sub-contracted to a third party who is not the sponsor, copies of relevant Good Manufacturing Practice agreements relation to such manufacture shall be kept.

The sponsor shall retain records of the distribution of the listed medicine for a period of five years and shall provide the records or copies of the records to the Complementary Medicines Branch, Therapeutic Goods Administration, upon request.

The sponsor of the listed medicine must not, by any means, intentionally or recklessly advertise the medicine for an indication other than those accepted in relation to the inclusion of the medicine in the Register.

All reports of adverse reactions or similar experiences associated with the use or administration of the listed medicine shall be notified to the Head, Office of Product Review, Therapeutic Goods Administration, as soon as practicable after the sponsor of the goods becomes aware of those reports. Sponsors of listed medicines must retain records of such reports for a period of not less than 18 months from the day the Head, Office of Product Review is notified of the report or reports.

The sponsor shall not supply the listed medicine after the expiry date of the goods.

Where a listed medicine is distributed overseas as well as in Australia, product recall or any other regulatory action taken in relation to the medicine outside Australia which has or may have relevance to the quality, safety or efficacy of the goods distributed in Australia, must be notified to the National Manager Therapeutic Goods Administration, immediately the action or information is known to the sponsor.

Products

1 . Prospan Drops						
Product Type	Single Medicine Product	Effective Date	30/08/2019			
Permitted Indications						
Anti-inflammatory/relieve inflammation						
Decrease/reduce/reli	ieve mild bronchial irritation					
Decrease/reduce/reli	ieve bronchial mucous congestion					
Soothe/calm the che	st					
Decrease/reduce excess chest phlegm						
Loosen chest phlegn	n					
Decrease/reduce exe	cess mucous					
Decrease/reduce/reli	ieve mild upper respiratory tract congestion					
Expectorant/clear res	spiratory tract mucous					
Loosen respiratory tr	act mucous					
Decrease/reduce/reli	ieve mild bronchial cough					
Decrease/reduce/reli	ieve cough					
Enhance/improve/promote/increase cough productivity						
Soothe respiratory tr	act mucous membranes/mucous tissue					
Indication Requiren	nents					
Product presentation must not imply or refer to serious forms of respiratory disorders/diseases, such as: asthma, pneumonia, COAD, COPD, influenza.						
Label statement: If s	symptoms persist, talk to your health profession	nal.				

Respiratory tract infections must be qualified by 'mild'.

Label statement: Adults only, OR Not to be used in children under 2 years of age without medical advice (or words to that effect).

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Product presentation must only refer to mild bronchitis.

Standard Indications

No Standard Indications included on Record

Specific Indications

No Specific Indications included on Record

Warnings

Adults only. OR Not to be used in children under two years of age without medical advice (or words to that effect).

If coughing persists consult your doctor (or a healthcare professional) (or words to that effect).

Adults only. OR Not to be used in children under two years of age without medical advice (or words to that effect).

Do not use if pregnant or likely to become pregnant (or words to that effect)

Use in children under 12 years is not recommended.

Contains saccharin sodium (or words to that effect).

Contains ethanol or contains alcohol.

Do not use while breastfeeding.

Keep out of reach of children (or words to that effect).

Additional Product information

Pack Size/Poison information	on			
Pack Size		Poison Schedule		
Components				
1. Formulation 1				
Dosage Form	Oral Liquid			
Route of Administration	Oral			
Visual Identification				C
Active Ingredients				
Hedera helix leaf Extract of	dry concentrate		20 mg/mL	
Equivalent: Hedera helix ((Dry)		125 mg/mL	~
Other Ingredients (Excipie	ents)			
Anise Oil				
ethanol				
Fennel Oil				
Peppermint Oil				
purified water				
saccharin sodium				1

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