MAXIMUM STRENGTH 4% LIDOCAINE PATCH GOOD NEIGHBOR PHARMACY-lidocaine patch

InterMed Laboratories Private Limited

Maximum Strength 4% Lidocaine Patch

Active Ingredient

Lidocaine 4%

Purpose

External Analgesic

Uses

For temporary relief of pain

Warnings

For external use only

Do Not Use

- more than 1 patch on your body at a time or on cut, irritated or swollen skin
- for more than one week without consulting a doctor
- over raw surfaces or blistered areas
- if pouch is damaged or opened

When Using This Product

- use only as directed. Read and follow all directions and warnings on this label.
- do not allow contact with the eyes

Stop Use and Ask a Doctor if

- condition worsens
- skin irritation develops
- symptoms persist for more than 7 days or clear up and occur again within a few days

If Pregnant or Breast Feeding

ask a health professional before use.

Keep Out of Reach of Children and Pets

If swallowed, get medical help or contact a Poison Control Center right away.

Directions

Adults and children over 2 years:

- clean and dry affected area free of lotions, ointments and creams.
- carefully remove backing from patch starting at a corner. Apply sticky side of patch to affected area.
- Do not use more than one patch in a 12 hour period. Maximum 2 patches per day. Discar patch after single use.

children 2 years or younger:ask a physician.

Inactive Ingredients

dihydroxy aluminium aminoacetate, disodium EDTA, glycerin, methyl paraben, polyvinyl alcohol, propyl paraben, propylene glycol, sodium carboxy methyl cellulose, sodium polyacrylate, tartaric acid, titanium dioxide, water

Bulk Package Label



MAXIMUM STRENGTH 4% LIDOCAINE PATCH GOOD NEIGHBOR PHARMACY

lidocaine patch

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Prod	UCT	Inform	ation

Product Type HUMAN OTC DRUG Item Code (Source) NDC:71811-008

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name Basis of Strength Strength

LIDOCAINE (UNII: 98PI200987) (LIDOCAINE - UNII:98PI200987) LIDOCAINE 40 mg

Inactive Ingredients

Ingredient Name	Strength
ingregient Name	Strength

TARTARIC ACID (UNII: W4888I119H)
TITANIUM DIOXIDE (UNII: 15FIX9V2|P)

CARBOXYMETHYLCELLULOSE SODIUM (UNII: K6790BS311)

DIHYDROXYALUMINUM AMINOACETATE (UNII: DO250MG0W6)

GLYCERIN (UNII: PDC6A3C0OX)	
EDETATE DISODIUM ANHYDROUS (UNII: 8NLQ36F6MM)	
METHYLPARABEN (UNII: A2I8C7HI9T)	
SODIUM POLYACRYLATE (2500000 MW) (UNII: 05115JN12J)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
POLYVINYL ALCOHOL, UNSPECIFIED (UNII: 532B59J990)	
PROPYLPARABEN (UNII: Z8IX2SC1OH)	
WATER (UNII: 059QF0KO0R)	

P	Packaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date			
1	NDC:71811-008- 05	5 in 1 CARTON	11/06/2025				
1		1 in 1 POUCH; Type 0: Not a Combination Product					

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC Monograph Drug	M017	11/06/2025			

Labeler - InterMed Laboratories Private Limited (676244169)

Revised: 11/2025 InterMed Laboratories Private Limited