

BABY WIPES WITH ALLANTOIN- allantoin liquid
Yinjing Medical Technology (Shanghai) Co., Ltd.

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

Baby Wipes with Allantoin

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Allantoin 0.5%(w/w)

Skin Protectant

Uses:

Temporarily protects and helps relieve minor skin irritations

Warnings:

For external use only.

When using this product, do not get into eyes.

Stop use and ask a doctor if the rash persists for more than 7 days, or tends to reoccur. Keep out of reach of children

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Directions

- Open lid, remove tab and pull wipe through
- Take wipe and rub thoroughly over the affected area
- Use as many wipes as needed
- DO NOT FLUSH, dispose of used wipes in trash

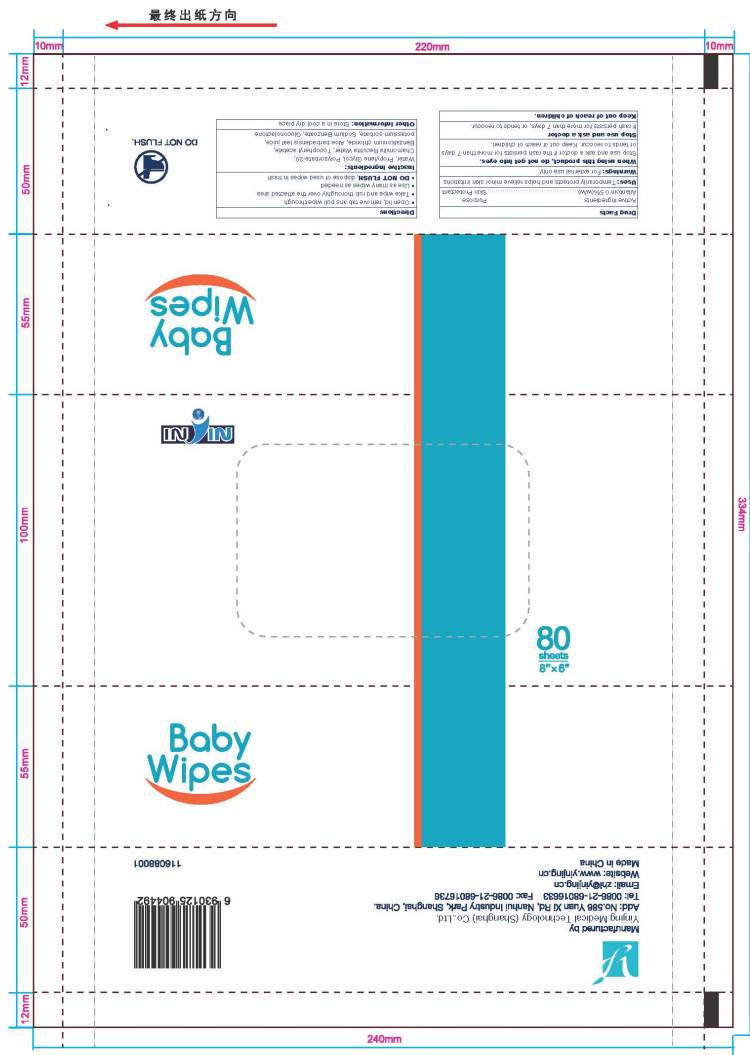
Inactive Ingredients:

Water, Propylene Glycol, Polysorbate-20, Chamomilla Recutita Water, Tocopheryl acetate, Benzalkonium chloride, Aloe barbadensis leaf juice, potassium sorbate, Sodium Benzoate, Gluconolactone.

Other Information:

Store in a cool dry place

Baby Wipes with Allantoin 80 Count (44019-400-80)



- 专色亮橙
- 银京蓝



彩色盖贴 (89*48mm)



白色盖贴 (100*60mm)

BABY WIPES WITH ALLANTOIN

allanto in liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:440 19-400
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALLANTOIN (UNII: 344S277G0Z) (ALLANTOIN - UNII:344S277G0Z)	ALLANTOIN	0.5 g in 100 g

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0K00R)	

PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
POLYSORBATE 20 (UNII: 7T1F30V5YH)	
.ALPHA.-TOCOPHEROL ACETATE (UNII: 9E8X80D2L0)	
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7)	
ALOE VERA LEAF (UNII: ZY81Z83H0X)	
POTASSIUM SORBATE (UNII: 1VPU26JZZ4)	
SODIUM BENZOATE (UNII: OJ245FE5EU)	
GLUCONOLACTONE (UNII: WQ29KQ9POT)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:44019-400-80	360 g in 1 BAG; Type 0: Not a Combination Product	08/03/2016	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	part347	08/03/2016	

Labeler - Yinjing Medical Technology (Shanghai) Co., Ltd. (530501535)

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