

**VANILLA HAND SANITIZER- benzalkonium chloride gel**  
**Shenzhen Lantern Science 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.*

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**☐Active Ingredient**

Benzalkonium Chloride 0.1%

Antibacterial

Keep out of reach of children.

**☐Warnings**

- for external use only
- if irritation occurs discontinue use
- keep out of eyes
- avoid contact with broken skin
- do not inhale or ingest
- stop use and ask if doctor if skin irritation develops
- keep out of reach of children
- if swallowed, get medical help or contact a poison control center right away.

**☐Directions**

- Rub a dime sized drop into hands.
- For children under 6 use under adult supervision.

**☐Inactive Ingredients**

water/aqua, hydroxyethylcellulose, phenoxyethanol, fragrance/parfum, disodium edta

To decrease bacteria on hands

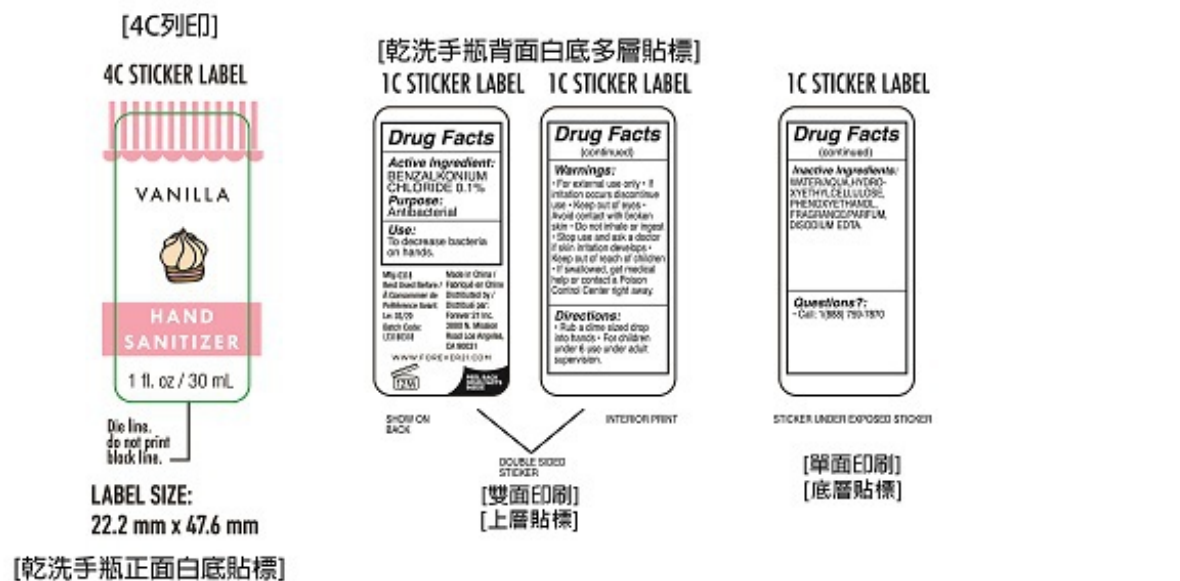
# TO:蘭亭

## #TB01905 (Forever 21 特別版)

Forever 21 版不需改稿，只需加上製造日期標及確認資訊是否正確

\*不需成分

1. 一般英文版 乾洗手 成分(字高 1/32")
2. DIRECTION FOR SAFE USE (英文) (字高 1/32")
3. 品名字高 1/16"
4. 正確重量: 乾洗手: 1FL OZ (30 ml) PER UNIT (字高 1/16")
5. 產地: Made in China
6. 請加上保值期圖案
7. 正確地址為: Distributed by / Distribué par: Forever 21 Inc. 3880 N, Mission Road, Los Angeles, CA 90031
6. 請加上 Batch Code: LT31B0318
7. Mfg: 03/18
8. Best used before : 03/20



# TO:蘭亭





[乾洗手瓶正面白底貼標]

#TB01905 (Forever 21 特別版)

UPC:

Forever 21 系列 1FL OZ (30 mL) Hand Sanitizer with holder 乾洗手含軟膠套繩組

內容物: (乾洗手聞起來要有香甜味，並沒有含酒精成分，但是成分為“苯扎氯胺:比率 0.1-0.15%(殺菌作用))

1 個 Forever 21 1FL OZ (30 ML)橢圓瓶 Hand Sanitizer with holder 乾洗手:

Vanilla 香草香味

透明 Hand Sanitizer with holder 乾洗手內料

橢圓瓶上蓋:白色

透明橢圓瓶尺寸: 8.2(H) x 3.5(W) x 2.3 (D) CM

瓶身套上一粉紅色(PMS 7422C)全包開窗軟膠套+含 3 顆調整珠扣軟膠套繩(蘭亭全包)

以上 1 個成一套。

**VANILLA HAND SANITIZER**

benzalkonium chloride gel

**Product Information**

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:54860-089
<b>Route of Administration</b>	TOPICAL		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>BENZALKONIUM CHLORIDE</b> (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII:7N6JUD5X6Y)	BENZALKONIUM CHLORIDE	0.1 g in 100 mL

## Inactive Ingredients

Ingredient Name	Strength
HYDROXYETHYL ETHYLCELLULOSE (UNII: ZDN57Z154K)	
EDETATE DISODIUM (UNII: 7FLD91C86K)	
PHENOXYETHANOL (UNII: HIE492ZZ3T)	
WATER (UNII: 059QF0K00R)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:54860-089-01	30 mL in 1 BOTTLE; Type 0: Not a Combination Product	01/30/2018	

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333E	01/30/2018	

**Labeler** - Shenzhen Lantern Science Co., Ltd. (421222423)

## Establishment

Name	Address	ID/FEI	Business Operations
Shenzhen Lantern Science Co., Ltd.		421222423	manufacture(54860-089)

Revised: 2/2018

Shenzhen Lantern Science Co., Ltd.