

ANTI DANDRUFF SHAMPOOS- pyrithione zinc liquid
Guangdong Quaker Daily Chemical Co., LTD

anti dandruff SHAMPOOS

Pyrithione zinc 1%

Anti-dandruff

helps prevent recurrence of flaking and itching associated with dandruff

For external use only.

In eyes and mucous membranes. If contact occurs, rinse thoroughly with water.

you have a condition that covers a large area of the body.

condition worsens or does not improve after regular use of this product as directed.

Irritation, dryness, or an allergic reaction occurs and consult a healthcare professional.

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

for best results, use at least twice a week or as directed by a doctor.

for maximum dandruff control, use every time you shampoo.

shake before use

wet hair, massage onto scalp, rinse, repeat if desired.

water

sodium lauryl sulfate

cocamidopropyl betaine

guar hydroxypropyltrimonium chloride

acrylates/c10-30 alkyl acrylate crosspolymer

zinc chloride

sodium chloride

methylchloroisothiazolinone

dimethicone

trideceth-3

trideceth-10

citric acid

prunus persica (peach) leaf extract

trehalose

aloe barbadensis leaf juice

ammonium lauryl sulfate

sodium benzoate

fragrance

blue 1 CI 42090

红色尺寸部分无需印刷

qb

QUEST
BASICS

ANTI DANDRUFF

SKU: LOT#: FCCID#:

INCB:
C M Y K

2746 C

COMMENTS:

DIELINE
CUT FOLD

DESIGNER: Milagros Santos
DATE: 10/23/2024

COMPARE TO THE ACTIVE INGREDIENT IN
HEAD & SHOULDERS® CLASSIC CLEAN

biopure.

anti
dandruff
SHAMPOOS

FOR DRYNESS & ITCH RELIEF

EFFECTIVE DANDRUFF CONTROL
HYDRATES DRY SCALP
SOOTHES ITCHINESS
GENTLE CLEANSING
REDUCES INFLAMMATION

16 FL OZ (473ML)

biopure.
12mm

Drug Facts

Active Ingredient

Purpose

Pyrrithione zinc 1% Anti-dandruff

Uses

helps prevent recurrence of flaking and itching associated with dandruff.

Warnings

For external use only.

Do not use

In eyes and mucous membranes. If contact occurs, rinse thoroughly with water.

Ask a doctor before use if

you have a condition that covers a large area of the body.

Stop use and ask a doctor if

condition worsens or does not improve after regular use of this product as directed.

irritation, dryness, or an allergic reaction occurs and consult a healthcare professional.

Keep out of reach of children. If swallowed, get medical help or contact a Poison Control Center right away.

Directions

for best results, use at least twice a week or as directed by a doctor.

for maximum dandruff control, use every time you shampoo.

shake before use.

wet hair, massage onto scalp, rinse, repeat if desired.

Other information

close and store at room temperature.

Inactive Ingredients

water, sodium laureth sulfate, cocamidopropyl betaine, sodium chloride, trehalose, sodium benzoate, fragrance, dimethicone, acrylates/c10-30 alkyl acrylate crosspolymer, aloe barbadensis leaf juice, ammonium lauryl sulfate, guar hydroxypropyltrimonium chloride, zinc chloride, methylchlorosulfathiazoline, citric acid, prunus persica(peach) leaf extract, trideceth-3, trideceth-10, blue 1 (CI 42090).

Questions? Call 718-875-2586

BP2773-DT

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Made in China
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CAP: 10249 C



BOTTLE: WHITE

ANTI DANDRUFF SHAMPOOS

pyrrithione zinc liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:85434-001
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PYRITHIONE ZINC (UNII: R953O2RHZ5) (PYRITHIONE ZINC - UNII:R953O2RHZ5)	PYRITHIONE ZINC	1 g in 100 mL

Inactive Ingredients

Ingredient Name	Strength
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
ZINC CHLORIDE (UNII: 86Q357L16B)	

METHYLCHLOROISOTHIAZOLINONE (UNII: DEL7T5QRPN)	
TRIDECETH-10 (UNII: G624N6MSBA)	
WATER (UNII: 059QF0KO0R)	
SODIUM LAURETH SULFATE (UNII: BPV390UAP0)	
PRUNUS PERSICA LEAF (UNII: VN3501T41P)	
ALOE BARBADENSIS LEAF POWDER (UNII: ZY81Z83H0X)	
AMMONIUM LAURYL SULFATE (UNII: Q7AO2R1M0B)	
FRAGRANCE 13576 (UNII: 5EM498GW35)	
COCAMIDOPROPYL BETAINE (UNII: 5OCF3O11KX)	
SODIUM BENZOATE (UNII: OJ245FE5EU)	
DIMETHICONE (UNII: 92RU3N3Y1O)	
CITRIC ACID (UNII: 2968PHW8QP)	
GUAR HYDROXYPROPYLTRIMONIUM CHLORIDE (UNII: B16G315W7A)	
ACRYLATES/C10-30 ALKYL ACRYLATE CROSSPOLYMER (60000 MPA.S) (UNII: 8Z5ZAL5H3V)	
TRIDECETH-3 (UNII: 438A02204X)	
TREHALOSE (UNII: B8WCK70T7I)	
CI 42090 (UNII: H3R47K3TBD)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:85434-001-01	1 in 1 BOX	03/26/2025	
1		473 mL in 1 BOTTLE; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M032	03/26/2025	

Labeler - Guangdong Quaker Daily Chemical Co., LTD (548075342)

Establishment

Name	Address	ID/FEI	Business Operations
Guangdong Quaker Daily Chemical Co., LTD		548075342	manufacture(85434-001)

Revised: 3/2025

Guangdong Quaker Daily Chemical Co., LTD