# BAD BREATH KILLER- propolis strip TECOZYME INC

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

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### **ACTIVE INGREDIENT**

Propolis extract 8.29%

### **PURPOSE**

**Antibacterial** 

### Uses

- Helps prevent and reduce oral bacterial that leads to bad breath, gingivitis, periodontitis
- For the dry mouth relief overnight

### WARNINGS

Use with caution if allergic to bee products Keep out of reach of children under 3 years age

### **KEEP OUT OF REACH OF CHILDREN**

Keep out of reach of children under 3 years age

### **Directions**

- Use 1 strip per day or per night.
- Fold and apply to upper molar gum surface
- Do not push the strips with tongue for 10 second once applied.
- Safe to use overnight

### Other Information

Store away from high temperatures and humidity or direct sunlight.

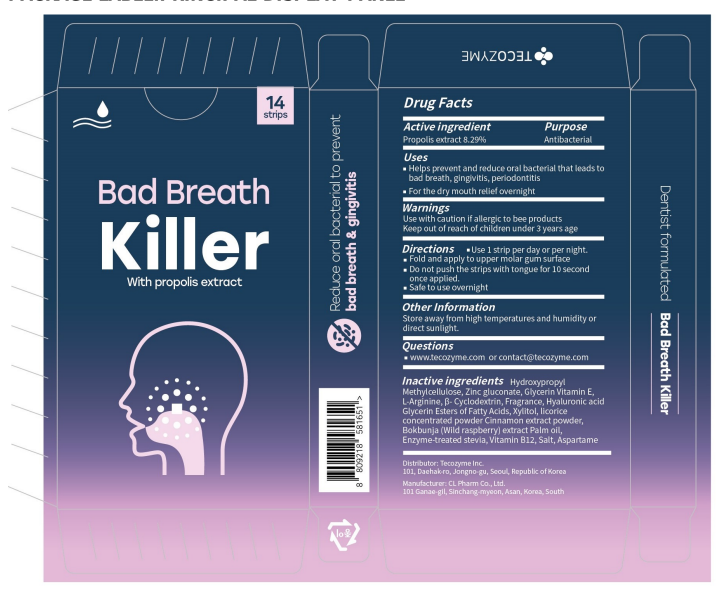
### **QUESTIONS**

■ www.tecozyme.com or contact@tecozyme.com

### **INACTIVE INGREDIENTS**

Hydroxypropyl Methylcellulose, Zinc gluconate, Glycerin Vitamin E, L-Arginine, β-Cyclodextrin, Fragrance, Hyaluronic acid, Glycerin Esters of Fatty Acids, Xylitol, licorice concentrated powder, Cinnamon extract powder, Bokbunja (Wild raspberry) extract, Palm oil, Enzyme-treated stevia, Vitamin B12, Salt, Aspartame

### PACKAGE LABEL.PRINCIPAL DISPLAY PANEL



# Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:79717-0011 Route of Administration DENTAL

Active Ingredient/Active Moiety			
Ingredient Name	<b>Basis of Strength</b>	Strength	
PROPOLIS WAX (UNII: 6Y8XYV2NOF) (PROPOLIS WAX - UNII:6Y8XYV2NOF)	PROPOLIS WAX	8.29 mg in 100 mg	

Inactive Ingredients		
Ingredient Name	Strength	
HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29V3WO)		
Zinc gluconate (UNII: U6WSN5SQ1Z)		
Xylitol (UNII: VCQ006KQ1E)		

l	Packaging					
	# Item Code	Package Description	Marketing Start Date	Marketing End Date		
	1 NDC:79717- 0011-2	14 in 1 CARTON	08/01/2021			
	1 NDC:79717- 0011-1	84.5 mg in 1 POUCH; Type 0: Not a Combination Product				

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		08/01/2021	

## Labeler - TECOZYME INC (694613703)

# Registrant - TECOZYME INC (694613703)

Establishment				
Name	Address	ID/FEI	Business Operations	
CL Pharm Co.,Ltd.		694759867	manufacture(79717-0011)	

Revised: 8/2021 TECOZYME INC