

**PROPRANOLOL SCOPOLAMINE- propranolol scopolamine tablet  
TPS**

*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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Take one tablet orally as directed by your physician as needed for symptoms of panic or anxiety.

Do not exceed two tablets per day.

**TPS LLC**

3524 DECATUR HWY FULTONDALE, AL 35068

877-608-4995 1-877-608-4995 BT9752747

Caution: Federal law prohibits transfer of this drug to any other person than patient for whom prescribed

**Rx 263095 Jack Doe/Dr. Jane Doe MD**

**JOHN DOE**

123 MAIN ST AUBURN, AL 12345

PROPRANOLOL SCOPOLAMINE TABLET TRITURATE

40 MG/0.5 MG

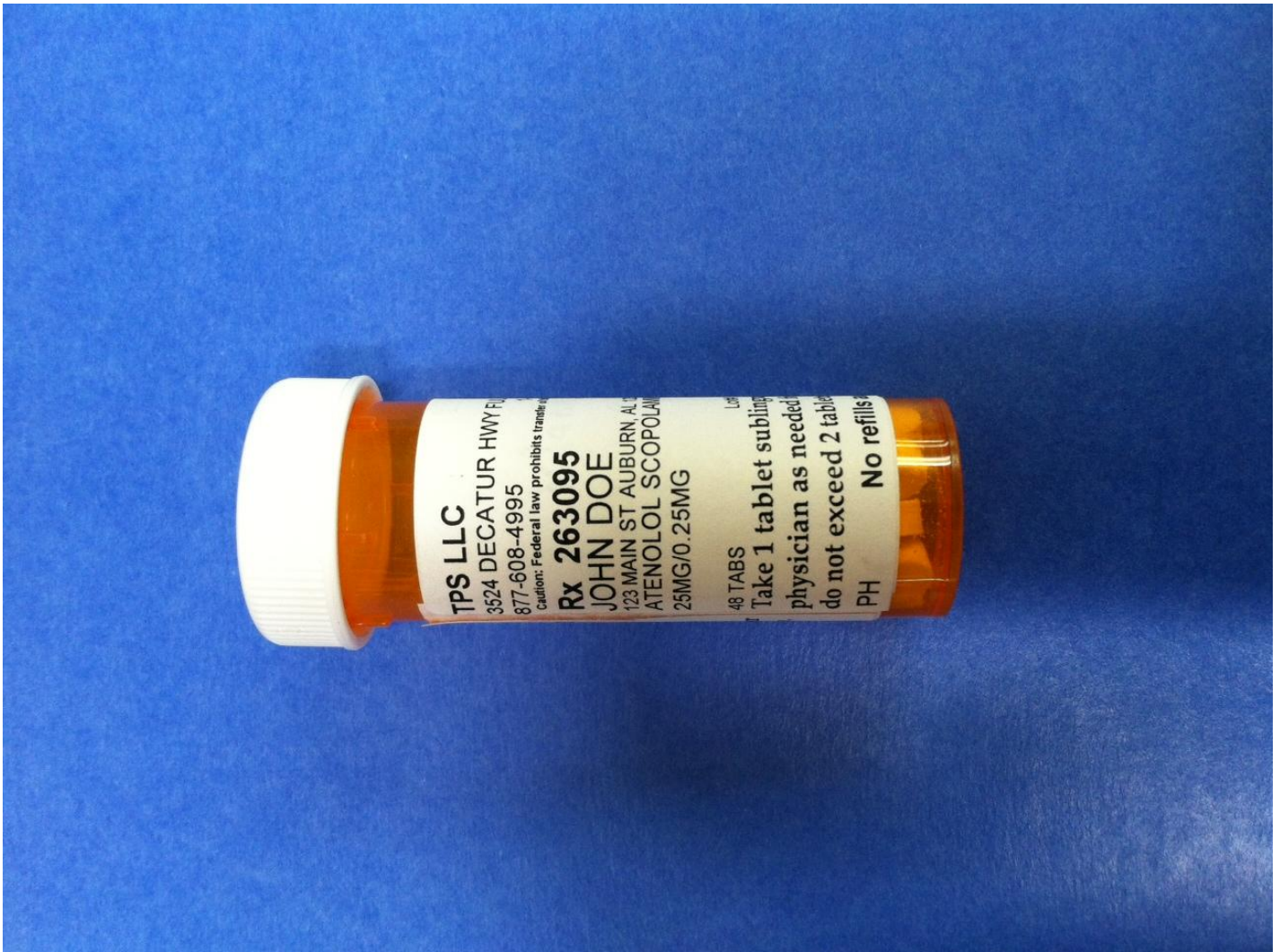
48 TABS Lot# Exp

Take 1 tablet sublingually or orally as directed by your physician as needed for symptoms of panic or anxiety.

Do not exceed 2 tablets per day

PH No refills authorized 10/8/2014

Pill bottle low res.jpg



<b>PROPRANOLOL SCOPOLAMINE</b>			
propranolol scopolamine tablet			
<b>Product Information</b>			
<b>Product Type</b>	HUMAN PRESCRIPTION DRUG	<b>Item Code (Source)</b>	NDC:69267-202
<b>Route of Administration</b>	ORAL		
<b>Active Ingredient/Active Moiety</b>			
	<b>Ingredient Name</b>	<b>Basis of Strength</b>	<b>Strength</b>
	PROPRANOLOL HYDROCHLORIDE (UNII: F8A3652H1V) (PROPRANOLOL - UNII:9Y8NXQ24VQ)	PROPRANOLOL HYDROCHLORIDE	40 mg in 40.5 mg
	SCOPOLAMINE HYDROBROMIDE (UNII: 451IFR0GX8) (SCOPOLAMINE - UNII:DL48G20X8X)	SCOPOLAMINE HYDROBROMIDE	.5 mg in 40.5 mg
<b>Product Characteristics</b>			
<b>Color</b>	white	<b>Score</b>	no score
<b>Shape</b>	ROUND (THE DIAMETER DEPENDS ON DIES)	<b>Size</b>	5mm
<b>Flavor</b>		<b>Imprint Code</b>	

**Contains****Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:69267-202-06	243 mg in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	10/01/2014	
2	NDC:69267-202-12	486 mg in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	10/01/2014	
3	NDC:69267-202-24	972 mg in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	10/01/2014	
4	NDC:69267-202-48	1944 mg in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	10/01/2014	

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		10/01/2014	

**Labeler** - TPS (044805267)**Establishment**

Name	Address	ID/FEI	Business Operations
TPS		044805267	manufacture(69267-202)

Revised: 10/2014

TPS