

## WALLEN-BRIDGE™, warfarin sodium tablet

Warfarin Sodium Tablets, USP

THIS INFORMATION IS FOR PROFESSIONALS ONLY.  
These highlights do not include all of the information needed to use WARLEN-BRIDGE safely and effectively. See full prescribing information for WARLEN-BRIDGE.

**1.1. Approved 2016** **INDICATIONS AND USAGE**  
WARLEN-BRIDGE is indicated for the treatment of thromboembolic disorders. See full prescribing information for complete details.  
**1.2. Important Warnings and Precautions**  
• **Bleeding risk:** WARLEN-BRIDGE increases the risk of bleeding. Patients should be monitored for signs of bleeding. See full prescribing information for complete details.  
• **Interactions:** WARLEN-BRIDGE has numerous drug interactions. See full prescribing information for complete details.  
**2. DOSAGE AND ADMINISTRATION**  
See full prescribing information for complete details.  
**3. WARNINGS AND PRECAUTIONS**  
**3.1. Bleeding Risk:** WARLEN-BRIDGE increases the risk of bleeding. Patients should be monitored for signs of bleeding. See full prescribing information for complete details.  
**3.2. Laboratory Monitoring:** Regular monitoring of INR is necessary. See full prescribing information for complete details.  
**3.3. Drug Interactions:** WARLEN-BRIDGE has numerous drug interactions. See full prescribing information for complete details.  
**3.4. Contraindications:** WARLEN-BRIDGE is contraindicated in patients with active bleeding or other conditions that increase the risk of bleeding. See full prescribing information for complete details.  
**4. HOW TO USE YOUR TABLETS**  
See full prescribing information for complete details.  
**5. STORAGE AND HANDLING**  
See full prescribing information for complete details.

**6. HOW TO USE YOUR TABLETS**  
See full prescribing information for complete details.  
**7. SIDE EFFECTS**  
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Walfarin Sodium Tablets, USP, 4 mg  
NDA 021-130



WALFARIN SODIUM			
NDA 021-130			
<b>Product Information</b>			
Product Type	ANDA (New Active Ingredient) - New Drug (sNDA)	Drug Code (sNDA)	Drug Code (ANDA/BLA/OTC)
Route of Administration			
<b>Active Ingredient/Inactive Ingredients</b>			
Active Ingredient Name	Supplemental Name	Strength	Strength
<b>Manufacturer Information</b>			
<b>Product Characteristics</b>			
Color	Shape	Size	Imprint
<b>Packaging</b>			
4 (1) Unit Code	Package Description	Marketing Start Date	Marketing End Date
<b>Marketing Information</b>			
Marketing Category	Application Number or Marketing Application	Marketing Start Date	Marketing End Date

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