

THE GOVERNMENT PHARMACEUTICAL ORGANIZATION

RAW MATERIAL SPECIFICATION

Title: Fluoxetine Hydrochloride USP

Spec. No. : SP-AK30-F23

(Item No. 41031975)

Reference(s): USP 41 page 1809 - 1810

Rev. No. : 03

Other Requirements: GPO Specification

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USP 41

Test Parameters	Requirement		
Description	White to off-white, crystalline powder.		
Solubility	Freely soluble in alcohol and in methanol; sparingly soluble in water and in		
	dichloromethane; practically insoluble in ether.		
Identification	A. Infrared absorption.		
	B. It responds to the tests for Chloride.		
Organic impurities	Aminomethyl-1-phenylpropanol : Not more than 0.25%.		
	Fluoxetine related compound B : Not more than 0.25%.		
	Fluoxetine related compound A : Not more than 0.15%.		
	4-Trifluoromethylphenol : Not more than 0.1%.		
	Any individual unspecified impurity : Not more than 0.1%.		
	Total impurities : Not more than 0.5%.		
Water	Not more than 0.5%.		
Assay	98.0% - 102.0% of Fluoxetine hydrochloride (C ₁₇ H ₁₈ F ₃ NO·HCl), calculated on the anhydrous basis.		

เอกสารไม่ควบคุม ใช้ในการจัดซื้อ

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Head of Raw Material	Director of Raw Material	Director of Drug Registration and	Director of Quality Assurance	13/03/20
Standard Section 2	Standard Division	Pharmacovigilance Division	Department (Activa)	



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GPO Specification

Test Parameters	Requirement	
Heavy metals	Not more than 30 ppm.	
Tapping volume	5.0-g portion occupies not more than 14.0 mL when tapped down mechanically for 20 drops at 3-mm height and nominal rate of 250 drops/min.	
Fineness	For information.	
[Pass through sieve number 30]		
Tapped density	For information.	

เอกสารไม่ควบคุะ ใช้ในการจัดซื้

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Product Information

Approved source(s)	Refer to current version of Approved Supplier List of Fluoxetine Hydrochloride USP (Item No. 41031975).
Sampling plan	For Identification : 100%. For Other Tests : n plan.
Testing procedure	Tests to be performed as per current version of WI-AK30-F23.
Storage condition	Store at a condition stated on the label from the manufacturer. Preserve in tight containers.
Retest period	1 year after first testing date.

History of changes

Description	Effective Date
ประกาศใช้ครั้งแรก USP 27	25/06/04
Update ข้อกำหนดตาม USP 34	
	29/09/11
	10/05/15
	Description

เอกสารไม่ควบคุม ใช้ในการจัดซื้อ

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J. 1867 padech / 29/01/20	Linerount , antollau	130/05/20	Rumquongroj , 30/0-1/20	15/03/20
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