

Triamcinolone acetoneide

CODE ARTICLE: 96882	BATCH: 0124881	Active pharmaceutical ingredient
TEST DATE: 24/01/2024	TEST: QC-00260678	
EXPIRY DATE: 30/07/2028	FABRICATION DATE: 30/08/2023	MR: 434,50
CAS: [76-25-5]	FORMULA: C ₂₄ H ₃₁ FO ₆	
STORAGE: Store in tightly closed containers in a cool, dry place.		

TEST	RESULTS	Units	MIN	MAX	METHOD
APPEARANCE	CONFORM	-		Crystal powder	PH.EUR.*
COLOUR	CONFORM	-		White	PH.EUR.*
<u>SOLUBILITY</u>					
SOLUBLE	SPARINGLY SOLUBLE IN ETHANOL				PH.EUR.*
INSOLUBLE	PRACTICALLY INSOLUBLE IN WATER				PH.EUR.*
<u>IDENTIFICATION</u>					
IR IDENTIFICATION	CONFORM				PH.EUR.*
IDENTIFICATION C	CONFORM				PH.EUR.*
ROTACION OPTICA (SUST. ANHIDRA)	111	-	110	117	PH.EUR.*
<u>RELATED SUBSTANCES</u>					
TOTAL IMPURITIES	0,2% (PASS)	-		<= 0.5% (<= 5xarea Triamcinolone RS b)	PH.EUR.*
UNSPECIFIED IMPURITIES	< DL=0,05% (PASS)	-		<= 0.10% (<= Triamcinolone Area RS B)	PH.EUR.*
IMPURITY B	0,1% (PASS)	-		<= 0.2% (<= 2xarea triamcinolone RS b)	PH.EUR.*
IMPURITY C	0,06% (PASS)	-		<= 0.15% (<= 1.5xarea triamcinolone RS b)	PH.EUR.*
WATER	1,5 %		0,0	2	PH.EUR.*
ASSAY (ANHIDROUS BASIS)	98,7 %		97,5	102	PH.EUR.*
<u>RESIDUAL SOLVENTS</u>					
ACETONE	88 ppm		0,0	5.000	Manufacturer's Standard
METHANOL	115 ppm		0,0	3.000	Manufacturer's Standard
DICHLOROMETHANE	N.D. MG/KG			600	Manufacturer's Standard
ACETIC ACID	N.D. ppm			5000	Manufacturer's Standard
DIISOPROPYLETHER	N.D. ppm			5000	Manufacturer's Standard
PARTICLE SIZE <10 MICRON	CONFORM	%	99		Manufacturer's Standard
CORRECT BATCH	YES				Manufacturer's Standard
SAMPLING	COLLECT				Manufacturer's Standard

The data expressed in this certificate of analysis is facilitated by our supplier and/or obtained in our control laboratory, in no case being exempted of the controls demanded by each sector. Tests marked with (*) are verified in Guinama.

Manufacturer: 53686 M. P. C. PVT. LTD. (India) Manufacturer Batch: TRAFP23012 Manufacturer original CoA available under request

GUINAMA S.L. Calle Oslo, 3 46185 La Pobla de Vallbona (Valencia) Spain

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STORAGE: Store in tightly closed containers in a cool, dry place.		

TEST	RESULTS	Units	MIN	MAX	METHOD
SPECIFICATION	PH.EUR 11				Manufacture's Standard
Elemental Impurities ICHQ3D	PASS				Manufacture's Standard
Risk Assessment Evaluation for Nitrosamines	PASS				Manufacture's Standard

NON-sterile



This product has been handled in a NON-sterile room. For batches suitable for sterile use, check availability.

Silvia Sancho-Tello Ripoll
Technical Director.

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