

CHIMAK HEALTHCARE

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QUALITY CONTROL DEPARTMENT THE DRUG & COSMETIC ACT. 1940 & THE RULES THERE UNDER

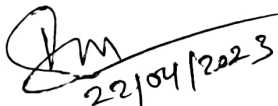


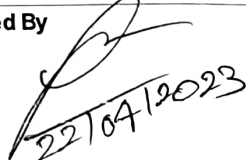
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CERTIFICATE OF ANALYSIS

Product Name : SUPER TADAPOX (10X10)	
Generic Name : Tadalafil & Dapoxetine Tablets. IHS	
Batch No. : LRMB23003	A.R. No. : FGLT04231206
Batch Size : 1,00,000.000 TAB	Qty. of Sample : 60.000 TAB
Mfg. Dt. : APR-2023	Exp. Dt. : MAR-2026
Parent Product Code : SMD000443	Received Dt. : 22/04/2023
Parent Batch No. : LGYR23003	Release Dt. : 22/04/2023
Parent Batch Size : 1,00,000.000 TAB	Specification No. : CHC/QC/FGPS/912
Storage Condition : Store at a temperature not exceeding 25°C in a dry place, protect from light	

Sr.	Test	Result	Specification
1	Description	Yellow coloured, round shaped, onese breakline, biconvex, film coated tablet. contains in Blister pack, imprinted with SUPER TADAPOX	Yellow coloured, round shaped, onese breakline, biconvex, film coated tablet. contains in Blister pack, imprinted with SUPER TADAPOX
2	Identification test	Complies.	Should be positive for Tadalafil & Dapoxetine Hcl.
3	Average weight	326.50 mg	326.0 mg \pm 5.0 %
4	Uniformity of weight	-2.90%, +4.13%	\pm 5 % of average weight
5	Diameter	10.10 mm	10.10 mm \pm 0.2 mm
6	Thickness	3.94 mm	4.00 mm \pm 0.5 mm
7	Hardness in Kg/cm ²	9.4 kg/cm ²	NLT 5 kg/cm ²
8	Dissolution for Tadalafil 40 mg	Min 83.36% & Max 84.61%	NLT 60.0%
9	Dissolution for Dapoxetine HCl eq. to Dapoxetine 60 mg	Min 95.56% & Max 98.60%	NLT 70.0%
10	Assay for Tadalafil USP 40 mg	38.6956 mg (96.74%)	NLT 90.0 % & NMT 110.0%
11	Assay for Dapoxetine HCl eq. to Dapoxetine 60 mg	59.2316 mg (98.72%)	NLT 90.0 % & NMT 110.0%

In the opinion of undersigned the sample referred to above is standard quality / ~~not of standard quality~~, as per IHS specification.

Analysed By  22/04/2023	Reviewed By  22/04/2023	QA Department  22/04/2023	Approved By  22/04/2023
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