

(THE DRUGS & COSMETICS ACT 1940 & RULES THEREUNDER)

Sample Name CIPROFLOXACIN HCL TABLETS-IP.500mg. Mfg Lic No [REDACTED]  
Submitted By Quality Control Advisor. (MMDSL) Shillong. Report No [REDACTED]  
Address DHS LaitumkhrahNHM, EastKhasi Hills Meghalaya. Receipt Date 19-Mar-24  
Mfg By N.S. Report Date 29-Mar-24  
Supplied By N.S. Ref No N.S.  
Batch No Mfg. Date Exp Date Batch Size Sample Qty  
MMDSL/QC-0140 03/2023 02/2026 N.S. 60 TABS.

RESULT OF ANALYSIS

Date / Period of Performance of test 19/03/2024 to 29/03/2024.

Reference to protocol :- I.P-2022.

Description :- White elongated biconvex film coated tablet having scored on one side.

Identification A (by HPLC) :- Complies.  
B (by TLC) :- Complies.

Average weight :- 783.8mg.

Uniformity of weight :- Within limit.

Dissolution (by U.V) :- Complies(88.64% to 94.26%) Limit:- NLT 80%+5%

Assay (by HPLC) :- Each film coated tablet contains:-

Content of	Obtd./Avg.wt.	Claim	LIMIT		Method
			Lower	Upper	
Ciprofloxacin Hydrochloride eq. to Ciprofloxacin	:- 499.13mg	500.0mg	450.0mg	550.0mg	I.P.

NOTE:- SAMPLE CONSUMED IN TESTING.

Report: In Opinion of the undersigned, The sample referred to above is of Standard Quality as defined in the Act and the rules made there under for the reason given: Complies as per IP.

- NOTE :
1. The result listed refer only to the tested samples and applicable parameter. Endorsement of products is neither inferred nor implied.
  2. Sample not drawn by us. Total liability of this laboratory limited to the invoice amount.
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