



WINDLAS BIOTECH LIMITED

40/1 Mohabewala Industrial Area Dehradun-248110 (Uttarakhand)

Approved by: FDA Uttarakhand, GLP certified Lab

FORM 39 [See rule 150E(f)]

Report of Analysis by Approved Institution

Lic. No.: 1/UA/CTL*/2024

CERTIFICATE OF ANALYSIS FOR FINISHED PRODUCT

Product Name	ATORVAHEAL 10 TABLET (SALE)		
Generic Name/ Dosage Form Strength (mg)	Atorvastatin Tablets IP 10 mg		
Batch No.	AVU26001	Booking ID/A.R. No.	QC/FG/26/03874
Batch Size	200000 STRIP	Pack Size	01 X 10 TABLET
Mfg. Date	03/2026	Exp. Date	02/2028
Reference	I.P	Sample Qty.	1x125 TABLET
Sample Received on	21/03/2026	Manufacturer Lic. No	34/UA/2013,55/UA/SC/P-2013
Protocol/STP No.	WBL/STP/FG/0418	Ref. SPS No.	WBL/FPS/FG/8473
Date of Analysis started on	29/03/2026	Date of Analysis completed on	06/04/2026
Sample Send by and Address	WINDLAS BIOTECH LIMITED (PLANT-2) ,Khasra No. 141 to 143 & 145, Mohabewala Industrial Area, Dehradun-248110 UK		
Manufactured By	WINDLAS BIOTECH LIMITED (PLANT-2)	Page No.	01 of 03

S.No.	Test	Specification	Result
1	Description	White to off white coloured, round, biconvex, film coated tablets, plain on both sides.	White coloured, round, biconvex, film coated tablets, plain on both sides.
2	Identification	In the Assay, the principal peak in the chromatogram obtained with the test solution corresponds to the peak in the chromatogram obtained with the reference solution.	Complies
3	Disintegration Time	Not more than 30 minutes	01 Min 12 Sec
4	Average weight	93.10 to 102.90 mg	98.08 mg
5	Uniformity of weight	Not more than 2 tablets in 20 deviates from the average weight by more than 10%. No tablet deviates from the average weight by more than 15%.	-3.54% to +4.42%

	Prepared By-QC	Checked By-QC	Reviewed By-Lab.-QA	Approved By- Person-in-charge
Name	Aman Kamboj	Ravi Kumar	Manoj Gautam	Jogendra Singh
Designation	Trainee	Assistant Manager	Assistant Manager	G. M.
Date	06/04/2026	06/04/2026	06/04/2026	06/04/2026

Printed By: Vanshita Singh

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Manufactured By	WINDLAS BIOTECH LIMITED (PLANT-2)	Page No.	02 of 03

S.No.	Test	Specification	Result
6	Dissolution	Not less than 70% (Q) of the stated amount in 30 minutes. Stage: S1 Number tested: 6 Each unit is not less than Q+5 percent of the labelled content. Stage: S2 Number tested: 6 Average of 12 units (S1+S2) is equal to or greater than Q, and no unit is less than Q-15 percent of the labelled content. Stage: S3 Number tested: 12 Average of 24 units (S1+S2+S3) is equal to or greater than Q, not more than 2 units are less than Q-15 percent of the labelled content and no unit is less than Q-25 percent of the labelled content.	Min-90.2% Max-95.3% Avg-92.4%
7	Uniformity of Dosage Unit (By Content uniformity)	Acceptance value should be less than 15	3.52
8	Related substances		
8a.	Any secondary peak	Not more than 1.0%	0.095%

	Prepared By-QC	Checked By-QC	Reviewed By-Lab.-QA	Approved By- Person-in-charge
Name	Aman Kamboj	Ravi Kumar	Manoj Gautam	Jogendra Singh
Designation	Trainee	Assistant Manager	Assistant Manager	G. M.
Date	06/04/2026	06/04/2026	06/04/2026	06/04/2026

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S.No.	Test	Specification	Result
8b.	Sum of all secondary peak	Not more than 4.0%	0.095%
9	Assay Each film coated tablet contains:		
9a.	Atorvastatin Calcium IP eq. to Atorvastatin...10mg	90.0 to 110.0%	99.6%

Remarks : The results of analysis relate only to the analysis applied.

In the opinion of the undersigned, the sample referred to above is of standard Quality as defined in the Act and the rules made thereunder for the reason given below.

	Prepared By-QC	Checked By-QC	Reviewed By-Lab.-QA	Approved By- Person-in-charge
Name	Aman Kamboj	Ravi Kumar	Manoj Gautam	Jogendra Singh
Designation	Trainee	Assistant Manager	Assistant Manager	G. M.
Date	06/04/2026	06/04/2026	06/04/2026	06/04/2026

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