

CERTIFICATE OF FINISHED PRODUCT ANALYSIS

Product Name:	Frasix-4ml Injection (Furosemide IP 10 mg/ml)		
Batch No.	25AH146B	A.R. No.	FP250235
Batch Size	10200 Ampoule	Sample Quantity	60 Nos
Mfg. Date	08/2025	Date of Testing	08/08/2025
Exp. Date	07/2027	Date of Release	23/08/2025
Reference Batch No	25AH146	MFG. Lic. No.	N-MB-2023/263

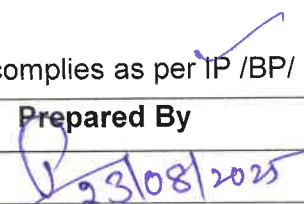
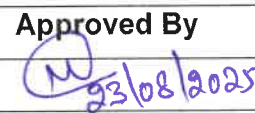
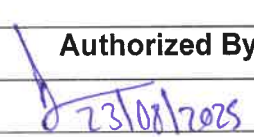
Sr. No.	Test	Specification	Observation
1.	Description	A clear, colourless or almost colourless solution.	Clear and colorless solution.
2.	Identification	A. When examined in the range 220 nm to 360 nm the final solution obtained in the Assay shows three absorption maxima at about 228 nm, 271 nm and 333 nm. B. A red-violet color is produced	Complies
3.	pH	Between 8.0 to 9.3.	8.66
4.	Extractable Volume	NLT 4.0 ml	4.0 ml
5.	Particulate Matter	Injections that are solutions, when examined under suitable conditions of visibility, are clear and practically free from particles that can be observed on visual inspection by the unaided eye.	Complies
6.	Related substance	In the chromatogram obtained with the test solution the area of any secondary peak is not more than the area of the principal peak obtained with the reference solution (1per cent), and sum of areas of all the secondary peaks is not more than 1.5 times the area of the peak in the chromatogram obtained with the reference solution (1.5 per cent).	Complies

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7.	Bacterial Endotoxins	Not more than 3.5 Endotoxin Units per mg of frusemide.	Less than 3.5 Endotoxin Units per mg of frusemide.
8.	Sterility	Should be complies	Sterile
9.	Assay: Each ml contains Furosemide IP 10 mg	95.0 % to 105.0 % (9.5 mg to 10.5 mg)	102.4 % (10.24 mg)

Remark: The sample is complies as per IP /BP/ USP/IH specification.

	Prepared By	Approved By	Authorized By
Sign/Date	 23/08/2025	 23/08/2025	 23/08/2025
Name	Prayash	Manisha Thakur	Bhupinder Navet
Designation	Officer QC	Executive-QC	Head-QC