

MINISTRY OF HEALTH PHARMACY AND POISONS BOARD

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Pharmacy and Poisons Board Building Lenana Road P.O Box 27663-00506 Nairobi, Kenya

GMP CERTIFICATE No: PPB/INS/GMP/CERT/029/23

CERTIFICATE OF GOOD MANUFACTURING PRACTICES (GMP) COMPLIANCE OF A MANUFACTURER

PART 1

This Certificate is issued in accordance with Section 35B of the Pharmacy and Poisons Act (Cap 244) of the Laws of Kenya. The Pharmacy and Poisons Board, The National Medicines Regulatory Authority of Kenya, confirms the following:

The manufacturer:	Bafna Pharmaceuticals Ltd
Site address:	No. 147, Madhavaram Redhills High Rd, Grantlyon Village, Vadakarai Post, Chenai-
	600052, India.

Has been inspected in connection with Marketing Authorization(s) listing manufacturers located outside Kenya.

From the knowledge gained during the inspection of this manufacturer, the latest of which was conducted on 6th - 7th March 2023, GMP Report No. **PPB/INS/GMP/RPT/029/23**, the site complies with the prescribed Good Manufacturing Practices as per the relevant WHO Technical Report Series and other internationally acceptable guidelines.

This certificate reflects the status of the manufacturing site at the time of the inspection noted above and should not be relied upon to reflect the compliance status if more than three years have elapsed since the date of that inspection, after which time the Kenya Pharmacy and Poisons Board should be consulted.

PART 2

1.1	Non-sterile products				
	DOSAGE FORM	CATEGORY	PRODUCT TYPE	ACTIVITIES	
	Oral Solids	General	Tablets & hard	All manufacturing activities	
			Gelatin Capsules	including all operations of	
				purchase of materials and	
				products, production,	
			and the second sec	quality control testing	
				and/or batch release	
				storage and distribution of	
		1000	pharmaceutical products		
				and the related controls.	

The compliance status shall be deemed valid unless it is invalidated under any of the following conditions;

- 1. The activities and/or categories certified herewith are changed.
- 2. The site is no longer considered to be in compliance with WHO cGMP.
- 3. The manufacturing site is changed.

The authenticity of this certificate may be verified with the Kenya Pharmacy and Poisons Board.

REGISTRAR PHARMACY AND POISONS BOARD MINISTRY OF HEALTH P. O. Box 27663 - 00506, NAIROBI Date: 31st October 2023

DR. F. M. SIYOI REGISTRAR/CHIEF EXECUTIVE OFFICER PHARMACY AND POISONS BOARD