







**ZENITH DRUGS LIMITED**  
(Formerly - Zenith Drugs Private Limited)

**CONTACT US:**

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 **PlantAdd.-72/5&72/1,72/3,74/1/1,75/1/1.Muradpura(Orangpura),  
Dhar Road, Near Kalaria, INDORE, PIN-453001(MP)INDIA**



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**Registered Office Address - K. No. 72/5, Village Muradpura, Depalpur, Indore, Madhya Pradesh - 453001**

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Date: February 10, 2026

To,  
Listing Compliance Department,  
National Stock Exchange of India Limited,  
Exchange Plaza, Bandra Kurla Complex,  
Bandra (East), Mumbai – 400051, Maharashtra, India.

**NSE Symbol: ZENITHDRUG; ISIN- INE0QWN01013**

**Subject: Intimation of receiving GMP Certificate under Revised Schedule 'M'**

Regulation: Regulation 30 of SEBI (Listing Obligations and Disclosure Requirements), Regulations, 2015

Dear Sir/Madam,

Pursuant to Regulation 30 of the SEBI (Listing Obligations and Disclosure Requirements) Regulations, 2015, **Zenith Drugs Limited** is pleased to inform that the Company has been granted a **Good Manufacturing Practices (GMP) Certificate** under **Revised Schedule 'M' of the Drugs Rules, 1945** by the **Food & Drugs Administration, Government of Madhya Pradesh**.

The GMP Certificate has been issued following a detailed inspection conducted on **02 February 2026** by the Drugs Inspector, District Indore, at the Company's manufacturing facility situated at **Muradpura (Orangpura), Dhar Road, Near Kalaria, Indore (M.P.)**.

This certification confirms that the Company's manufacturing operations are fully compliant with the stringent **quality, safety, and regulatory standards** prescribed under Revised Schedule 'M'. The GMP Certificate shall remain valid up to **12 September 2028**, aligning with the validity of the Company's drug manufacturing licences.

The receipt of this certification is a significant milestone and reflects **Zenith Drugs Limited's continued commitment to operational excellence, robust quality systems, and adherence to global manufacturing standards**. The Company believes that this achievement further enhances confidence among regulators, customers, investors, and other stakeholders.

The above information is being disclosed for the information of the exchanges and stakeholders.

Thanking you.

Yours faithfully,  
**For Zenith Drugs Limited**

**Sakshi Bhawsar**  
**Company Secretary & Compliance Officer**