

## Facilities Deviation Writer

Job ID  
REQ-10079705  
Июн. 23, 2026  
Мексика

### Сводка

#LI-Hybrid

Location: Mexico City, Mexico

Relocation Support: This role is based in Mexico City, Mexico. Novartis is unable to offer relocation support: please only apply if accessible.

Take ownership of quality where it truly matters - at the intersection of facilities, engineering, and patient impact. In this role, you will play a key part in ensuring deviations are thoroughly investigated, understood, and resolved, helping to maintain high standards in a complex GMP environment. You'll work closely with cross-functional teams to drive robust investigations and meaningful corrective actions, while shaping a culture of continuous improvement and quality excellence. If you enjoy combining analytical thinking with hands-on collaboration, this role offers a chance to make a real difference.

### About the Role

#### Key Responsibilities

- Open and assess deviations within required timelines, ensuring accurate classification and documentation
- Evaluate product impact of deviations, aligning with batch release requirements and quality standards
- Author, own, and drive investigations through to timely and compliant closure; update SOPs as relevant to Deviations/CAPAs
- Apply structured root cause analysis methods to identify causes of process and product deviations
- Ensure investigations are complete, accurate, and fully supported with robust documentation
- Design and execute experiments or studies to support investigation outcomes
- Collaborate cross-functionally to assess deviation impact and maintain compliant operations
- Develop, document, and implement effective corrective and preventive actions
- Monitor CAPA effectiveness and ensure execution through Good Manufacturing Practice systems and training
- Deliver training and communication to reinforce quality practices and maintain compliance

#### Essential Requirements

- Bachelor's degree in a relevant field with pharmaceutical industry experience
- Fluency in written and spoken English
- Two to five years of experience in a pharmaceutical, facilities, engineering, or quality environment
- Hands-on experience with deviation management and writing investigations in a GMP environment
- Proven experience in CAPA management, including creation, implementation, and effectiveness tracking
- Strong root cause analysis skills using structured investigation methodologies
- Strong understanding of current good manufacturing practices and regulatory expectations for biologics manufacturing
- Excellent technical writing skills, with the ability to clearly structure investigations and present findings

#### Commitment to Diversity and Inclusion

Novartis is committed to building an outstanding, inclusive work environment and diverse team representative of the patients and communities we serve.

#### Accessibility and accommodation

Novartis is committed to work with and provide reasonable accommodation to individuals with disabilities. If, because of a medical condition or disability, you need a reasonable accommodation for any part of the recruitment process, or in order to perform the essential functions of a position, please send an e-mail to [tas.mexico@novartis.com](mailto:tas.mexico@novartis.com) and let us know the nature of your request and your contact information. Please include the job requisition number in your message.

**Why Novartis:** Helping people with disease and their families takes more than innovative science. It takes a community of smart, passionate people like you. Collaborating, supporting and inspiring each other. Combining to achieve breakthroughs that change patients' lives. Ready to create a brighter future together? <https://www.novartis.com/about/strategy/people-and-culture>

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Дивизион  
Operations  
Business Unit  
Production / Manufacturing

Место

Мексика

Сайт

INSURGENTES

Company / Legal Entity

MX06 (FCRS = MX006) Novartis Farmacéutica S.A. de C.V.

Functional Area

Technical Operations

Job Type

Full time  
Employment Type  
Regular  
Shift Work  
No

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