

# Senior Scientific/Regulatory Writer

Job ID REQ-10021041 Sep 17, 2024 Irlanda

#### Resumen

-To write, review and/or manage the production of high quality clinical and safety documentation for submission to regulatory authorities in support of marketing applications. To provide documentation related consultancy to other line functions.

#### **About the Role**

### Major accountabilities:

- Core member of clinical Trial Team/participate in safety Management Team -Actively participate in planning of data analyses and presentation used in CSRs.
- Act as documentation consultant in CTTs and SMTs To ensure compliance of documentation To internal company standards and external regulatory guidelines.
- May Act as Program Writer ensuring adequate medical writing resources are available for assigned Program and consistency between documents.
- Reporting of technical complaints / adverse events / special case scenarios related to Novartis products within 24 hours of receipt -Distribution of marketing samples (where applicable)

## Key performance indicators:

 Delivery of high quality clinical and safety documents in time and in compliance with internal and external standards -Customer / partner/ project feedback and satisfaction -Adherence to Novartis policy and guidelines

### **Minimum Requirements:**

#### Work Experience:

- · Functional Breadth.
- Cross Cultural Experience.
- · Collaborating across boundaries.
- Operations Management and Execution.

#### Skills:

- Clinical Research.
- · Clinical Trials.
- · Detail Oriented.
- · Medical Writing.
- · Regulatory Compliance.

Safety.

#### Languages:

• English.

**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? <a href="https://www.novartis.com/about/strategy/people-and-culture">https://www.novartis.com/about/strategy/people-and-culture</a>

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División

Operations

**Business Unit** 

**CTS** 

Ubicación

Irlanda

Sitio

Dublin (Novartis Corporate Center (NOCC))

Company / Legal Entity

IE02 (FCRS = IE002) Novartis Ireland Ltd

Alternative Location 1

Home Worker, Reino Unido

Alternative Location 2

Hyderabad (Office), India

**Functional Area** 

Research & Development

Job Type

Full time

**Employment Type** 

Regular

Shift Work

No

Apply to Job

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

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- 1. https://www.novartis.com/about/strategy/people-and-culture
- 2. https://talentnetwork.novartis.com/network
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- 4. https://novartis.wd3.myworkdayjobs.com/en-US/Novartis\_Careers/job/Dublin-Novartis-Corporate-Center-NOCC/Senior-Scientific-Regulatory-Writer\_REQ-10021041-1
- 5. https://novartis.wd3.myworkdayjobs.com/en-US/Novartis\_Careers/job/Dublin-Novartis-Corporate-Center-NOCC/Senior-Scientific-Regulatory-Writer\_REQ-10021041-1