

## Senior Global Process Owner - Study & Site Management

Job ID  
REQ-10075540

4月 16, 2026

India

### 摘要

TAs Senior Global Process Owner for Study & Site Management you will own the end to end, regulatory compliant study and site process, strengthening its health and maturity through continuous improvement, and powering faster, quality driven clinical delivery that gets transformative medicines to patients sooner.

The Senior Global Process Owner for Study & Site Management acts as a single point of ownership that drives process health and continuous improvement for sustained process maturity.

### About the Role

The Sr GPO will be responsible for overall governance and oversight of a process by setting appropriate strategy, coordinating process mapping activities, overseeing the development the various procedural documents related to a process, ensuring efficiency and effectiveness of the process and managing risks. In addition, the Sr GPO would also be responsible to monitor process

performance via KPIs/KQIs aligned with regulatory and organizational strategies.

Key Responsibilities :

### 1. End to End Process Ownership & Strategy

- Accountable for the overall design, delivery, maintenance, and continuous improvement of the designated process(es).
- Lead long term process strategy, ensuring alignment with regulatory expectations and business needs.
- Anticipate internal/external changes and assess their impact on processes and supporting systems.

### 2. Cross Functional Collaboration & Process Improvement

- Lead and support cross functional process improvement and change management initiatives.
- Drive simplification, automation, and standardization across functions.
- Ensure transformed processes can be executed globally by responsible line functions.

### 3. Governance, Documentation Oversight & Compliance

- Ensure oversight and lifecycle management of controlled documents (SOPs, WPs, manuals) for the process.
- Ensure coherence and harmonization across procedural documents within the process.
- Oversee process related risks and ensure appropriate mitigation strategies.
- Monitor performance trends, conduct root cause analysis/FMEAs when needed, and ensure appropriate risk management.

Minimum Requirements:

- Education: University degree in Life Science, quantitative science or business. Desirable qualifications in shared services, outsourcing, global sourcing, project management/Coaching, 6-Sigma, Lean education/training, Master of Business Administration or equivalent
- Extensive knowledge of end-to-end processes within clinical development, including supporting systems, regulations, and awareness of business changes.
- 5 years' Site Management, Clinical Trial Monitoring, CRA Management and/or Clinical Project Management (Country level) domain experience essential.
- Ability to anticipate and assess the impact of external and internal changes on the end-to-end process, supporting systems (and vice-versa), and associated training requirements.
- Experience in effective process improvement.
- Strategic thinker with the ability to contribute to long-term process improvements and operational planning.
- Experience with process simplification and optimization, including improvements to quality documentation.
- Demonstrated ability to collaborate effectively across functions, supporting performance improvements within the end-to-end clinical development value chain.

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部门  
Development

Business Unit  
Development

地点  
India

站点  
Hyderabad (Office)

Company / Legal Entity  
IN10 (FCRS = IN010) Novartis Healthcare Private Limited

Functional Area  
Research & Development

Job Type  
Full time

Employment Type  
Regular

Shift Work

No

```
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showSpeedMenu: false }, css : "/modules/custom/arcticnckalturaaddon/css/kalturavideo.css",
components: { fullscreen: { disableDoubleClick: false } }, uiComponents: [ { presets: ['Playback',
'Live'], area: 'BottomBarRightControls', replaceComponent: 'Fullscreen', get:
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config.plugins.googleTagManager.customEventsTracking.custom = [];
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false, playlistEvents: false, castEvents: false } }; }
```

```
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kalturaPlayerVideos = []; kalturaPlayerVideos.push(kalturaPlayer); } // Load the Player for other
media. kalturaPlayer.loadMedia({entryId: "1dgfvmafo"}); } catch (e) { console.error(e.message) }
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