#### **Clinical Research Submission**

1/7/2025, 2:40:50 pm

#### 1. Administration

Name: sagar

**Department:** computer science

**Submission Date:** 26/6/2025

**Review Type:** Expedited Review **Study Title:** medical research

Short Title: MR
Protocol: 987897
Version: 8798879

**Date:** 2025-06-06T18:30:00.000Z

#### 2. Investigators

**Type:** principal

Name: shanmuka sagar

Designation: software developer

**Qualification:** MD, PhD

**Department:** Clinical Research

**Email:** shanmukasagar2019@gmail.com

**Contact:** 9876543210

**Type:** hod

Name:shanmuka sagarDesignation:Co-Investigator

**Qualification:** MD, PhD

**Department:** Clinical Research

**Email:** shanmukasagar2021@gmail.com

**Contact:** 9876543210

### 3. Investigators Count

PI Count: 12 Co-PI Count: 25 Duration: 32

## 4. Funding Details

**Estimated Budget:** 50000

**Funding Source:** Self-funding

false <>

Proposed Budget: ₹ hfkshksf Cost Per Patient: ₹ fsfsfhskf Total Project Cost: ₹ 650 NIMS Investigations:

**Name:** CHT **Cost:** 250

1

**Name:** BGT **Cost:** 400

Is Outsourced: Yes

Outsourced Investigations:

Name: shanmuka sagar

**Cost:** 600

**Lab:** ameerpet

**NABL:** yes

false false

### 5. Overview Research

kjadhkahd kh **Summary:** 

Academic clinical trials **Study Type:** 

12

**Other Type:** 

### 6. Methodology

**Sample Size:** 250

**Participants per Site: Lab Outsourcing:** No

**Lab Details:** 

### 7. Participants

**Patient** Type:

**Vulnerable Groups: Other Participant:** 

**Reimbursement:** No

**Details:** 

**Additional Safeguards:** 

Justification:

#### 8. Benefits

**Any Risk:** No

**Risk Details:** 

**Risk Strategy:** 

**Participant Benefits:** Indirect **Social Benefits:** Direct **Scientific Benefits:** Indirect

### 9. Consent

Waiver Consent:

No

**Translated Languages:** 

**Other Reason:** 

### 10. Payment

**Injury Treatment:** No **SAE Compensation:** NA

### 11. Storage

**Docs Control:** No **Drugs Control:** No

### 12. Additional Info

**Any Additional:** No

**Details:** 

### 13. Declaration

PI Name: shanmuka sagar

Guide Name: HOD Name:

# 14. Checklist

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Cover letter	View File
Brief CV of all Investigators- Updated, signed and dated	View File
Good Clinical Practice (GCP) training of investigators in last 3 years	View File
EC clearance of other centers	File: null
Agreement between collaborating partners	File: null
MTA between collaborating partners	File: null
Insurance policy / certificate	View File
Copy of CTA signed with the sponsor	View File
Provide all significant previous decisions (e.g. those leading to a negative decision or modified protocol) by other ECs / Regulatory authorities for proposed study (whether in same location or elsewhere) and modification(s) to protocol	File: null
Copy of the detailed protocol (clearly identified numbered and dated) and synopsis (summary as far as possible in non-technical language, flowchart, diagrammatic representation of the protocol)	ि View File
Investigators Brochure (If applicable for drug / biologicals / device trials)	View File
Participant Information Sheet (PIS) and Informed Consent Form (ICF) (English and translated) with version number and dated	ि View File
Assent form for minors (12-18 years) (English and Translated)	File: null
Proforma / Questionnaire / Case Report Forms (CRF) / Interview guides / Guides for Focused Group Discussions (FGDs) (English and translated)	ଐ View File

Advertisement / material to recruit participants (fliers, posters, etc.)

DCGI Approval letter

Others specify

File: null

File: null