Clinical Research Submission

1. Administration

Name: shanmuka sagar

Department: computer science

Submission Date: 6/10/2025

Review Type: Full Committee Review

Study Title: Artificial Intellengence study

Short Title: Al

Protocol: 4521ABC Version: 254KJH

Date: 2025-06-07T18:30:00.000Z

2. Investigators

Type: principal

Name: shanmuka sagar
Designation: software engineer

Qualification: B.Tech

Department: computer science

Email: shanmukasagar2019@gmail.com

Contact: 8688345501

3. Investigators Count

PI Count: 21

Co-Pl Count: 21

Duration: 13

4. Funding Details

Estimated Budget: 50000

Funding Source: Institutional funding

Other Details:

5. Overview Research

Summary: Developing Ai models

Study Type: Academic clinical trials

Other Type:

6. Methodology

Sample Size: 250

Participants per Site: 12
Lab Outsourcing: No

Lab Details:

7. Participants

Type: Patient

Vulnerable Groups:

Safeguards:

Other Participant:

Reimbursement: No

Details:

Additional Safeguards:

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8. Benefits

Any Risk: No

Risk Details:

Risk Strategy:

Participant Benefits: Indirect
Social Benefits: Indirect
Scientific Benefits: Indirect

9. Consent

Waiver Consent: Yes

Translated Languages:

Other Reason: hskfhslkfahl fhffhlshfslfhksfh jfhfklafhlkafhsflshfk

10. Payment

Injury Treatment: No SAE Compensation: No

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

protocol)

Cover letter View File Brief CV of all Investigators- Updated, signed View File and dated Good Clinical Practice (GCP) training of View File investigators in last 3 years EC clearance of other centers File: null Agreement between collaborating partners File: null File: null MTA between collaborating partners View File Insurance policy / certificate View File **Copy of CTA signed with the sponsor** Provide all significant previous decisions (e.g. those leading to a negative decision or modified protocol) by other ECs / Regulatory authorities File: null for proposed study (whether in same location or elsewhere) and modification(s) to protocol Copy of the detailed protocol (clearly identified numbered and dated) and synopsis (summary View File as far as possible in non-technical language, flowchart, diagrammatic representation of the

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	
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)	
Advertisement / material to recruit participants (fliers, posters, etc.)	File: null
DCGI Approval letter	View File
Others specify	File: null