

GUJARAT UNIVERSITY
Institutional Ethics Committee (IEC), School of Sciences,
Gujarat University, Ahmedabad

Form to be filled by the Principal Investigator (PI) for submission to Institutional Ethics Committee (IEC), Gujarat University, Ahmedabad.

Serial No of IEC Management Office:
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Proposal Title:

	Name, Designation & Qualifications	Address Tel & Fax Nos. Email ID	Signature
PI			
Co-PI / Collaborators			
1.			
2.			
3.			

Please attach detailed Curriculum Vitae of all Investigators (with subject specific publications limited to previous 5 years).

Tick appropriately

Sponsor Information :			
1. Indian	a) Government	<input type="checkbox"/>	Central <input type="checkbox"/> State <input type="checkbox"/> Institutional <input type="checkbox"/>
	b) Private	<input type="checkbox"/>	
2. International	Government	<input type="checkbox"/>	Private <input type="checkbox"/> UN agencies <input type="checkbox"/>
3. Industry	National	<input type="checkbox"/>	Multinational <input type="checkbox"/>
Contact Address of Sponsor:			

Total Budget :

1.Type of Study : Epidemiological Basic Sciences Animal studies
 Clinical: Single center Multicentric Behavioral

2. Status of Review: New Revised

**3. Clinical Trials:
 Drug /Vaccines/Device/Herbal Remedies :**

i. Does the study involve use of :
 Drug Devices Vaccines
 Indian Systems of Medicine/
 Alternate System of Medicine Any other NA

ii. Is it approved and marketed
 In India UK & Europe USA
 Other countries, specify

iii. Does it involve a change in use, dosage, route of administration?	Yes	No
If yes, whether DCGI's /Any other Regulatory authority's Permission is obtained?	Yes	No
If yes, Date of permission :		

iv. Is it an Investigational New Drug? If yes, IND No:	Yes	No
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a). Investigator's Brochure submitted	Yes	No
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b). <i>In vitro</i> studies data	Yes	No
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c). Preclinical Studies done	Yes	No
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d). Clinical Study is : Phase I <input type="checkbox"/> Phase II <input type="checkbox"/> Phase III <input type="checkbox"/> Phase IV <input type="checkbox"/>		
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e). Are you aware if this study/similar study is being done elsewhere ? If Yes, attach details	Yes	No
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4. Brief description of the proposal – Introduction, review of literature, aim(s) & objectives, justification for study, methodology describing the potential risks & benefits, outcome measures, statistical analysis and whether it is of national significance with rationale (Attach sheet with maximum 500 words):

5. Subject selection:
 i. Number of Subjects :

ii. Duration of study :

iii. Will subjects from both sexes be recruited	Yes	No
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iv.	Inclusion / exclusion criteria given			Yes	No	
v.	Type of subjects	Volunteers	<input type="checkbox"/>	Patients	<input type="checkbox"/>	
vi.	Vulnerable subjects (Tick the appropriate boxes)		Yes	No	<input type="checkbox"/>	
	pregnant women	<input type="checkbox"/>	children	<input type="checkbox"/>	elderly	<input type="checkbox"/>
	fetus	<input type="checkbox"/>	illiterate	<input type="checkbox"/>	handicapped	<input type="checkbox"/>
	terminally ill	<input type="checkbox"/>	seriously ill	<input type="checkbox"/>	mentally challenged	<input type="checkbox"/>
	economically & socially backward	<input type="checkbox"/>	any other	<input type="checkbox"/>		
vii.	Special group subjects (Tick the appropriate boxes)		Yes	No	<input type="checkbox"/>	
	captives	<input type="checkbox"/>	institutionalized	<input type="checkbox"/>	employees	<input type="checkbox"/>
	students	<input type="checkbox"/>	nurses/dependent	<input type="checkbox"/>	armed	<input type="checkbox"/>
	any other	<input type="checkbox"/>	staff		forces	<input type="checkbox"/>
6. Privacy and confidentiality						
i.	Study involves -		Direct Identifiers		<input type="checkbox"/>	
			Indirect Identifiers/coded		<input type="checkbox"/>	
			Completely anonymised/ delinked		<input type="checkbox"/>	
ii.	Confidential handling of data by staff			Yes	No	
7. Use of biological/ hazardous materials				Yes	No	
i.	Use of fetal tissue or abortus					
ii.	Use of organs or body fluids			Yes	No	
iii.	Use of recombinant/gene therapy			Yes	No	
	If yes, has Department of Biotechnology (DBT) approval for rDNA products been obtained?			Yes	No	
iv.	Use of pre-existing/stored/left over samples			Yes	No	
v.	Collection for banking/future research			Yes	No	
vi.	Use of ionising radiation/radioisotopes			Yes	No	
	If yes, has Bhaba Atomic Research Centre (BARC) approval for Radioactive Isotopes been obtained?			Yes	No	
vii.	Use of Infectious/biohazardous specimens			Yes	No	
viii.	Proper disposal of material			Yes	No	
ix.	Will any sample collected from the patients be sent abroad ?			Yes	No	
If Yes, justify with details of collaborators						
	a) Is the proposal being submitted for clearance from Health Ministry's Screening Committee (HMSC) for International collaboration?			Yes	No	

b) Sample will be sent abroad because (Tick appropriate box):		
	<input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/>	
Facility not available in India Facility in India inaccessible Facility available but not being accessed If so, reasons...		
8. Consent : *Written <input type="checkbox"/> Oral <input type="checkbox"/> Audio-visual <input type="checkbox"/>		
i. Consent form : (tick the included elements)		
Understandable language Statement that study involves research Sponsor of study Purpose and procedures Risks & Discomforts Benefits Compensation for participation Compensation for study related injury	<input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/>	Alternatives to participation Confidentiality of records Contact information Statement that consent is voluntary Right to withdraw Consent for future use of biological material Benefits if any on future commercialization eg. genetic basis for drug development
*If written consent is not obtained, give reasons:		
ii. Who will obtain consent ?		
	PI/Co-PI <input type="checkbox"/> Research staff <input type="checkbox"/>	Nurse/Counsellor <input type="checkbox"/> Any other <input type="checkbox"/>
9. Will any advertising be done for recruitment of Subjects ? (posters, flyers, brochure, websites – if so kindly attach a copy)		
	Yes	No
10. Risks & Benefits:		
i. Is the risk reasonable compared to the anticipated benefits to subjects / community / country?		
	Yes	No
ii. Is there physical / social / psychological risk / discomfort?		
If Yes, Minimal or no risk More than minimum risk High risk	<input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/>	Yes No
iii. Is there a benefit a) to the subject ?		
	Direct <input type="checkbox"/> Indirect <input type="checkbox"/>	
b) Benefit to society <input type="checkbox"/>		
11. Data Monitoring		
i. Is there a data & safety monitoring committee/ Board (DSMB)?		
	Yes	No
ii. Is there a plan for reporting of adverse events ?		
If Yes, reporting is done to :		
Sponsor <input type="checkbox"/> Ethics Committee <input type="checkbox"/> DSMB <input type="checkbox"/>		
iii. Is there a plan for interim analysis of data?		
	Yes	No

vi. Are there plans for storage and maintenance of all trial database? If Yes, for how long ?	Yes	No
12. Is there compensation for participation? If Yes, Monetary <input type="checkbox"/> In kind <input type="checkbox"/> Specify amount and type:	Yes	No
13. Is there compensation for injury? If Yes, by Sponsor <input type="checkbox"/> by Investigator <input type="checkbox"/> by insurance <input type="checkbox"/> by any other <input type="checkbox"/> company	Yes	No
14. Do you have conflict of interest? (financial/nonfinancial) If Yes, specify :	Yes	No
Checklist for attached documents:		
Project proposal – 20 Copies	<input type="checkbox"/>	
Curriculum Vitae of Investigators	<input type="checkbox"/>	
Brief description of proposal	<input type="checkbox"/>	
Patient information sheet	<input type="checkbox"/>	
Informed Consent form	<input type="checkbox"/>	
Investigator's brochure for recruiting subjects	<input type="checkbox"/>	
Copy of advertisements/Information brochures	<input type="checkbox"/>	
Copy of clinical trial protocol and/or questionnaire	<input type="checkbox"/>	
Institutional Ethics Committee clearance	<input type="checkbox"/>	
Institutional Animal Ethics Committee clearance	<input type="checkbox"/>	
CPCSEA clearance, if any	<input type="checkbox"/>	
HMSC/DCGI/DBT/BARC clearance if obtained	<input type="checkbox"/>	

Place:
Date:

Signature & Designation of PI/Co-PI/Collaborator

GUJARAT UNIVERSITY
Institutional Ethics Committee (IEC)

Model Form to be filled by Reviewers

Serial No of IEC Management Office:

Proposal Title:

Principal Investigator:

Co-investigator: 1.

2.

3.

Supporting Agency: ICMR/ non ICMR

If non ICMR, name of agency:

Project Status: New

Revised

Review: Regular

Interim

Date of Review:

1. Research Design

i. Scientifically sound enough to expose subjects to risk Yes No

ii. Relevant to contribute to further knowledge Yes No

iii. Of national importance Yes No

2 Risks

a. Is there physical/social/psychological risk/discomfort? Yes No

b. Is the overall risk/benefit ratio Acceptable Unacceptable

3 Benefits

Direct: Reasonable Undue None

Indirect: Improvement in science/knowledge Any other

4 Subject selection :

- i Inclusion / exclusion criteria addressed? Yes No
- ii Vulnerable subjects (woman, child, mentally challenged, seriously or terminally ill, foetus, economically or socially backward and healthy volunteers) adequately protected ? Yes No
- iii. Special group subjects (captives, students, nurses & dependant staff) adequately protected? Yes No

5 Privacy & Confidentiality maintained? Yes No

6 Patient Information Sheet: Adequate Inadequate

7. Consent form components addressed adequately? Yes No

8. Compensation, (if applicable) addressed adequately? Yes No

9. Is there a Conflict of Interest? Yes No

If yes, Acceptable Unacceptable

10. Budget: Appropriate Inappropriate

11. Decision of review
Recommended Recommended with suggestions
Revision Rejected

Any other remarks/suggestions:

Reviewer's name and Signature

**Communication of Decision of the Institutional Ethics Committee(IEC)/
Institutional Review Board(IRB)**

IEC/IRB No:

Protocol title:
Principal Investigator:
Name & Address of Institution:
<input type="checkbox"/> New review <input type="checkbox"/> Revised review <input type="checkbox"/> Expedited review
Date of review (D/M/Y): Date of previous review, if revised application:
Decision of the IEC/ IRB: <input type="checkbox"/> Recommended <input type="checkbox"/> Recommended with suggestions <input type="checkbox"/> Revision <input type="checkbox"/> Rejected
Suggestions/ Reasons/ Remarks:
Recommended for a period of :

Please note *

- **Inform IEC/IRB immediately in case of any Adverse events and Serious adverse events.**
- **Inform IEC/IRB in case of any change of study procedure, site and investigator**
- **This permission is only for period mentioned above. Annual report to be submitted to IEC/IRB.**
- **Members of IEC/IRB have right to monitor the trial with prior intimation.**

Signature of Member Secretary
IEC/IRB