



**MOUNT CARMEL COLLEGE, BENGALURUR-560052
CENTRE FOR SCIENTIFIC RESEARCH AND ADVANCED LEARNING
INSTITUTIONAL HUMAN ETHICAL COMMITTEE**

Proposal formats

FORM - A

APPLICATION FOR PERMISSION FOR STUDIES ON HUMAN SUBJECTS

1. Name & Address of the dept :
2. Name & designation of Investigator :
3. Place where study will be conducted :
4. Date of commencement & duration of study :
5. Funding agency / sponsor :

Investigator's Declaration

Certified that

1. The research proposal is not duplicative of previously reported research
2. All investigators working on this proposal are aware of the ICMR ethical guidelines
3. I / we have reviewed the pertinent scientific literature
4. I / we will obtain approval from IEC before initiating any deviation / changes in the study
5. The study shall be initiated only upon review & approval of IEC
6. I / we shall maintain all the records as per format [form B or C]
7. Informed consent will be obtained & confidentiality of the subjects will be maintained

Place:

Date

Chief Investigator:

For Office use only

Proposal number

Date of receipt

Approval date

Date received after revision

Expiry date

Secretary

Chairman



MOUNT CARMEL COLLEGE, BENGALURUR-560052
CENTRE FOR SCIENTIFIC RESEARCH AND ADVANCED LEARNING
INSTITUTIONAL HUMAN ETHICAL COMMITTEE

FORM -B

Proforma for routine PG class work (Practicals) involving Human Subjects.

1. Name of the Department :

2. List of Practical and Nature of each practical in brief.
(Including Objectives and Methods :
to be employed)

3. Specify the method of Subject selection for Practical class work :
 - (a) PG Students
 - (b) Patients
 - (c) Students (from other Institutions.)
 - (d) Any other, specify

4. Specify the source of obtaining :
blood samples

UNDERTAKING

It is certified that,

Work is conducted purely as part of routine curriculum by PG students.

Signature of the Teacher-in-charge.

Chairperson

MOUNT CARMEL COLLEGE, AUTONOMOUS

FORM - A

APPLICATION FOR PERMISSION FOR STUDIES ON HUMAN SUBJECTS

	Name & Designation / Qualification	Address Tel & Fax no Email	Signature
Name of PI/ PhD candidate			
Research Guide			
Co-PI, if any			
Research fellow			
Place where study will be conducted			
Date of commencement & duration of study			
Funding agency / sponsor			

Investigator's Declaration

Certified that

1. The research proposal is not duplicative of previously reported research
2. All investigators working on this proposal are aware of the ICMR ethical guidelines
3. I / we have reviewed the pertinent scientific literature
4. I/we will obtain approval from IEC before initiating any deviation/changes in the study
5. The study shall be initiated only upon review & approval of IEC
6. I /we shall maintain all the records as per format [form B or C]
7. Informed consent will be obtained & confidentiality of the subjects will be maintained

Place:

Date

Chief Investigator

For Office use only

Proposal number

Date of receipt

Approval date

Date received after revision

Expiry date

Secretary

Chairman

**MOUNT CARMEL COLLEGE, AUTONOMOUS
INSTITUTIONAL HUMAN ETHICAL COMMITTEE (IHEC)**

FORM – C

**Proforma for submission to Institutional Ethical Committee, for undertaking
studies involving human subjects**

1. Title:			
Tick one : PhD <input type="checkbox"/> Sponsored project <input type="checkbox"/> PG dissertation <input type="checkbox"/>			
2. Details of Investigating Team :			
	Name & Designation / Qualification	Dept. Address Tel & Fax no Email	Signature
Investigator			
Research Guide			
Any Others			
Name of sponsor			
Expertise of the investigating team			
3. Type of Study : Epidemiological <input type="checkbox"/> Basic Sciences <input type="checkbox"/> Survey <input type="checkbox"/>			
Clinical: Single center <input type="checkbox"/> Multicentric <input type="checkbox"/> Behavioral <input type="checkbox"/>			
(b) Data Collection: From Records <input type="checkbox"/>			
Using Questionnaire <input type="checkbox"/>			
(c) Any other, specify: <input type="checkbox"/>			
4. Duration of the study :			
Probable date of initiation :			
Completion :			
5. Pre-clinical studies done, if any :			
(in brief)			
Publications, if any :			

Note: It is compulsory to provide all the required information, incomplete applications will be rejected.

6. Study design [Brief description of the proposal – Introduction, aim (s) & objectives, justification for study, methodology describing number of subjects, Inclusion / exclusion criteria, dosages of drug, duration of treatment, potential risks & benefits, outcome measures, statistical analysis and whether it is of national significance with rationale. Attach sheet with maximum 500 words. See page 4 for more details]											
7. Will any advertising be done for recruitment of Subjects? (posters, flyers, brochure, websites – if so kindly attach a copy)											
8. Does the study involve (a) Anthropometric Measurements : Yes / No (b) Blood samples : Yes / No (c) Urine analysis : Yes / No (d) Lifestyle modification : Yes / No (e) Other (specify). If answer is Yes to (b) & (c) mention the tests											
9. Intervention Studies- Oral (a) Product evaluation : Yes / No (b) Dietary : Yes / No (c) Synthetic : Yes / No If Yes, is toxicological evaluation carried out. (d) Known medication : Yes / No If yes, give a brief summary of dosage, administration, Contra indications (if any)											
10. Use of biological/hazardous material : Yes <input type="checkbox"/> No <input type="checkbox"/> (If the answer is Yes, give details)											
11. Consent : Written <input type="checkbox"/> Oral <input type="checkbox"/> i. Subject consent form - enclose ii. Who will obtain consent ? <table style="width: 100%; border: none;"> <tr> <td style="width: 35%;"></td> <td style="width: 15%;">PI/Co-PI</td> <td style="width: 10%;"><input type="checkbox"/></td> <td style="width: 15%;">Nurse/Counsellor</td> <td style="width: 10%;"><input type="checkbox"/></td> </tr> <tr> <td></td> <td>Research staff</td> <td><input type="checkbox"/></td> <td>Any other</td> <td><input type="checkbox"/></td> </tr> </table>			PI/Co-PI	<input type="checkbox"/>	Nurse/Counsellor	<input type="checkbox"/>		Research staff	<input type="checkbox"/>	Any other	<input type="checkbox"/>
	PI/Co-PI	<input type="checkbox"/>	Nurse/Counsellor	<input type="checkbox"/>							
	Research staff	<input type="checkbox"/>	Any other	<input type="checkbox"/>							
12. Risks & Benefits: i. Is the risk reasonable compared to the anticipated benefits to subjects / community / country? Yes <input type="checkbox"/> No <input type="checkbox"/> ii. Is there physical / social / psychological risk / discomfort? Yes <input type="checkbox"/> No <input type="checkbox"/> iii. Is there a benefit <table style="width: 100%; border: none;"> <tr> <td style="width: 40%;">a) to the subject ?</td> <td style="width: 15%;">Direct</td> <td style="width: 10%;"><input type="checkbox"/></td> <td style="width: 15%;">Indirect</td> <td style="width: 10%;"><input type="checkbox"/></td> </tr> <tr> <td>b) Benefit to society</td> <td>Direct</td> <td><input type="checkbox"/></td> <td>Indirect</td> <td><input type="checkbox"/></td> </tr> </table> if yes, explain		a) to the subject ?	Direct	<input type="checkbox"/>	Indirect	<input type="checkbox"/>	b) Benefit to society	Direct	<input type="checkbox"/>	Indirect	<input type="checkbox"/>
a) to the subject ?	Direct	<input type="checkbox"/>	Indirect	<input type="checkbox"/>							
b) Benefit to society	Direct	<input type="checkbox"/>	Indirect	<input type="checkbox"/>							
13. i. Are the subjects remunerated for their involvement in the research? Yes <input type="checkbox"/> No <input type="checkbox"/> ii. If yes, is this remuneration provided irrespective of their social and economic conditions? iii. Compensation for travel, Specify amount and type:											

14. Data Monitoring

- i. Is there a data & safety monitoring committee
- ii. Is there a plan for reporting of adverse events?

If Yes, reporting is done to :

Sponsor Ethics Committee

15. Is there any conflict of interest?

(financial/non-financial)

If Yes, specify :

(Signature, Name & Designation of the Applicant)

Place:

Date:

Checklist for attached documents:

1. Form A- 1 copy
2. Project proposal – 2 Copies (Form B or C as applicable)
3. Informed Consent form -1 copy
4. Investigator’s brochure for recruiting subjects, if any
5. Advertisements /Information brochures
6. Copy of clinical trial protocol and/or Questionnaire
7. Ph. D Registration confirmation letter
8. Project sanction copy

Note: one copy each of Items 4, 5 & 6 to be attached only if applicable to the study.