NORTH EASTERN INDIRA GANDHI REGIONAL INSTITUTE OF HEALTH & MEDICAL SCIENCES, SHILLONG, MEGHALAYA
(An Autonomous Institute, Ministry of Health & Family Welfare, Govt. of India)

Combined Format for Submitting Research Proposal of Faculty for Consideration

By
NEIGRIHMS SCIENTIFIC ADVISORY COMMITTEE (NSAC)
INSTITUTE ETHICS COMMITTEE (IEC) (HUMAN STUDIES) INTRAMURAL RESEARCH GRANT COMMITTEE
Version 1.0
SECTION 1
(For NSAC)

PART A – GENERAL INFORMATION

1. Title of the Project :

2. Name, Designation & Address of the Principal Investigator with mobile number, e-mail ID & Number of ongoing projects as Principal Investigator :

3. Name(s), Designation(s) & Address(es) of the Co-Investigator(s) with mobile numbers & e-mail IDs :

4. Duration of study :

5. A. If the study is institutional, state whether it is intra-departmental or inter-departmental. :

   B. If the study is inter-departmental,

   (i) State the names of collaborating departments :

   (ii) State whether consent has been obtained from them :

6. A. If the study is inter-institutional, state whether it is national or international. :

   B. State the name of coordinating institution :

   C. State the names of collaborating institutions. :

   D. State whether consent has been obtained from collaborating institutions. Enclose copies of the same.

   E. State whether you have enclosed a copy of the original research protocol submitted by the coordinating institution.

   F. State the responsibilities of each collaborating Institution. :

7. Details of foreign collaboration with supporting evidence :

8. Details of foreign extramural funding with supporting evidence:

   A. Details of source(s) of funding
   B. Details of overall funding
   C. Details of funding to NEIGRIHMS with breakup

9. Details of Indian extramural funding with supportive evidence:

   A. Details of source(s) of funding
   B. Details of overall funding
   C. Details of funding to NEIGRIHMS with breakup
### PART B – TECHNICAL DETAILS

1. Title of the project

2. Background

   A. Rationale
   B. Novelty
   C. Expected outcome & application

3. Research question(s)

4. Research hypothesis (es), if any

5. Aim and objectives: Primary objective(s) & secondary objective(s)

6. Brief review of literature

7. Study participants
   (humans, animals or both)

8. Study design / type

9. for participants, mention

   A. Inclusion criteria
   B. Exclusion criteria

   C. Withdrawal criteria, if any (trial-related therapy, follow-up and documentation are terminated prematurely as it is indicated to ensure safety of the participants)

   D. Rescue criteria, if applicable (starting symptomatic therapy either to control symptoms of disease or to overcome lack of adequate efficacy of the study drug or placebo)

10. Number of groups to be studied, their names and definitions

11. Sampling

   A. Population
   B. Sampling method
   C. Sample size in each group and sample size calculation method(s)
12. Randomization details
A. Selection of participants
B. Allocation to groups

13. Methods
A. Intervention details with standardization techniques (drugs / devices / invasive procedures / noninvasive procedures / others)
B. Are the drugs/devices to be used approved for these indications by Drug Controller General of India (DCGI)? (Enclose the approval letter from DCGI for trial on humans or give undertaking to get the approval from DCGI; For all drugs and devices submit documents showing DCGI approval for the proposed indication of the study)
C. Are all procedures to be used professionally acceptable?
D. List of variables and their measurement methods with standardization techniques
   (i) Independent variables
   (ii) Dependent variables
   (iii) Confounding & interacting variables
E. Data collection methods including settings & periodicity
F. List variable-wise statistical tests to be used for data analysis

14. Relevant references for the project
(Maximum 20) (in Vancouver style, to be cited sequentially in the text of project)

15. Enclosures
A. Brief CV of all investigators
B. Data collection proforma
C. Questionnaire(s)
D. Copy of signed original protocol in multicentric Studies
E. Copy of signed consent letter from coordinator in multicentric studies
F. Others

16. Undertakings (please retain what is applicable)
A. The principal investigator hereby gives undertaking to obtain required DCG-I approval and submit its copies to NSAC and IEC.
B. The principal investigator hereby gives undertaking to obtain HMSC approval and submit its copies to NSAC and IEC.
C. The principal investigator hereby gives undertaking to follow official guidelines for exchange of human biological material.

D. The principal investigator hereby gives undertaking to get the required MoU signed and submit its copies to NSAC and IEC.

<table>
<thead>
<tr>
<th>A. Signature of the Investigator</th>
<th>Signature of Head of the Department of the Investigator</th>
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<td>(Name, Designation, Department, Seal and Date)</td>
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<tr>
<th>B. Signature(s) of the Co-Investigator(s)</th>
<th>Signature(s) of Head(s) of the Department of the co-investigator(s)</th>
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<tr>
<td>(Name, Designation, Department, Seal and Date)</td>
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SECTION – 2
(For Institute Ethics Committee (IEC)-Human Studies)

Proforma to be submitted to the Institute Ethics Committee (Human Studies) for faculty projects

1. Title of the project:

2. Ethical issues involved in the study:
   - less than minimal risk / minimal risk / more than minimal risk to the study subjects (for guidance please consult ICMR guidelines for biomedical research in human participants, 2006)
   [Along with level of risk, the risks should be written in detail. If you feel there will be no risk, give justification]

3. Benefit of the study:

4. Details of Informed Consent Process:
   i. Who will take the informed consent?
   ii. When will the informed consent be taken?
   iii. How will the informed consent be taken?
   iv. Where will the informed consent be taken?

5. Do you need exemption from obtaining Informed Consent from study subjects - if so give justifications.

6. Whether Consent forms in English and in local language are enclosed?
   (if the consent form in local language is not applicable, appropriate explanations must be provided)
   a. Documents attached
   b. Review Exemption Application Form (if applicable)
   c. Brief CV of investigators (including no. of projects with him/her) - Needed for all Investigators for each project separately
   d. Investigator’s Brochure
   e. For student projects, the guide should give a signed statement on a separate sheet with details of the project proposal that “I take full responsibility and accountability for planning, execution and adverse events occurring during the study. The data collected and records will be retained by me for a period of three years”.
   f. Others

7. Conflict of interest for any other investigator(s) (if yes, please explain in brief)

8. We, the undersigned, have read and understood this protocol and hereby agree to conduct the study in accordance with this protocol and to comply with all requirements of the ICMR guidelines (2006)

   Signature of the Investigators: Date:

   Signature of the Head of the Department Date:

   Signature of the Co- Investigators: Date:
Signature of the Heads of the Department of Co-Investigators  Date:

(Note: The proforma must be accompanied by Informed Consent Document (ICD) in Khasi, English & Hindi. Informed Consent Document should comprise Patient Information Sheet and the consent form. The investigator must provide information to the subjects in a simple language, and it should address the subjects, in a dialogue format. Studies involving children below 7 years should include parent / LAR consent form while studies involving children above 7 years and below 18 years of age should include assent form in addition to parent / LAR consent form)
INFORMED CONSENT DOCUMENT (ICD)

Patient / Participant information sheet

INFORMATION FOR PARTICIPANTS OF THE STUDY

Instructions - This is the patient information sheet. It should address the participant of this study. Depending upon the nature of the individual project, the details provided to the participant may vary. A separate consent form for the patient/test group and control (drug/procedure or placebo) should be provided as applicable. While formulating this sheet, the investigator must provide the following information as applicable in a simple language in English, Khasi and Hindi which can be understood by the participant. (Do not copy & paste from the study protocol submitted to NSAC).

- Title of the project
- Name of the investigator/guide
- Purpose of this project/study
- Procedure/methods of the study including withdrawal criteria
- Expected duration of the subject participation
- The benefits to be expected from the research to the participant or to others and the post trial responsibilities of the investigator
- Any risks expected from the study to the participant
- Maintenance of confidentiality of records
- Provision of free treatment for research related injury
- Reimbursement for participating in the study
- Compensation to the participants for foreseeable risks and unforeseeable risks related to research study leading to disability or death.
- Freedom to withdraw from the study at any time during the study period without the loss of benefits that the participant would otherwise be entitled
- Possible current and future uses of the biological material to be generated from the research and if the material is likely to be used for secondary purposes or would be shared with others, this should be mentioned
- Possible current and future uses of the data to be generated from the research and if the data is likely to be used for secondary purposes or would be shared with others, this should be mentioned
- Address and mobile number of the Principal investigator (PI) and Co-PI, if any:

    Signature of the investigator:

    Signature of the participant:

Place:
Date:
CONSENT FORM

Title of the project:

Participant’s name: Address:

The details of the study have been provided to me in writing and explained to me in my own language. I confirm that I have understood the above study and had the opportunity to ask questions. I understand that my participation in the study is voluntary and that I am free to withdraw at any time, without giving any reason, without the medical care that will normally be provided by the hospital being affected. I agree not to restrict the use of any data or results that arise from this study provided such a use is only for scientific purpose(s). I have been given an information sheet giving details of the study. **Risk and benefit of this project has been explained to me.** I fully consent to participate in the above study.

(I also consent / do not consent to use my stored biological samples for future scientific purposes: Yes/ No – if applicable)

Signature/thumb impression of the participant: ______________________ Date: _____________

Signature of the witness: ______________________ Date: _____________

Name and address of the witness:

Signature of the investigator: ______________________ Date: _____________
CONSENT FORM (for participants less than 18 years of age)

Parent/Legally acceptable representative (LAR)

Title of the project:

Participant’s name: 
Address: 

Parent/LAR’s name:

The details of the study have been provided to me in writing and explained to me in my own language. I confirm that I have understood the above study and had the opportunity to ask questions. I understand that my child/ward’s participation in the study is voluntary and that I am free to withdraw my child/ward at any time, without giving any reason, without the medical care that will normally be provided by the hospital being affected. I agree not to restrict the use of any data or results that arise from this study provided such a use is only for scientific purpose(s). I have been given an information sheet giving details of the study. **Risk and benefit of this project has been explained to me.** I fully consent for the participation of my child/ward in the above study.

Assent of child/ward obtained (for participants 7 to 18 years of age)

(I also consent / do not consent to use my child/ward’s stored biological samples for future scientific purposes: Yes/No – if applicable)

Signature/ thumb impression of the parent/ LAR: ______________________ Date: _____________

Signature of the witness: ______________________ Date: _____________

Name and address of the witness:

Signature of the investigator: ______________________ Date: _____________
ASSENT FORM
(for children above 7 years and below 18 years of age)

Assent form to participate in a clinical research

Child Participant’s name: Date of birth/Age:

Parent/LAR’s name: Address:

Title of the project:

The details of the study have been provided to me in writing and explained to me in my own language. I confirm that I have understood the above study and had the opportunity to ask questions. I understand that my participation in the study is voluntary and that I am free to withdraw at any time, without giving any reason, without the medical care that will normally be provided by the hospital being affected. I agree not to restrict the use of any data or results that arise from this study provided such a use is only for scientific purpose(s). I understand that following completion of study as well as during publication of the results, confidentiality of my identity will be maintained. I have been given an information sheet giving details of the study. Risk and benefit of this project has been explained to me. I fully assent to participate in the above study.

(I also assent / do not assent to use my stored biological samples for future scientific purposes: Yes/No – if applicable)

Signature of the child participant: Date:
(If child knows to sign/Thumb impression)

Signature of the parent or guardian: Date:

Name and address of the witness:

Signature of the witness: Date:

Signature of the Investigator: Date:

(Asent form should be accompanied by patient / participant information sheet for children in a simple language comprehensible to a child of 7-18 years; Language used should be simpler for children in the age group 7-12 years compared to children in the age group >12-18 years)
**CHECK LIST**  
*(To be filled and duly signed by the principal investigator)*

Title of the study:

Name of the Investigator:

Designation & Department:

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<tr>
<th>S.No</th>
<th>Items</th>
<th>Yes/No</th>
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<tbody>
<tr>
<td>1</td>
<td>Exact title as approved by NSAC</td>
<td></td>
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<tr>
<td>2</td>
<td>Date of NSAC approval mentioned in proper format (dd/mm/yyyy)</td>
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<tr>
<td>2</td>
<td>Source of funding mentioned</td>
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<td>3</td>
<td>Adequate literature review with justification for the study mentioned</td>
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<td>4</td>
<td>Detailed description about methodology (Study design, number of groups, sample size etc)</td>
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<tr>
<td>5</td>
<td>No mirror statement in Inclusion/Exclusion criteria (Ex: Age &lt;18 in inclusion &amp; Age &gt;18 in exclusion)</td>
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<tr>
<td>6a</td>
<td>Permission from DCGI <em>(if applicable)</em>.</td>
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<tr>
<td>6b</td>
<td>DCGI approval for the mentioned indication in the study (for drugs, devices, cosmetics etc)</td>
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<tr>
<td>7</td>
<td>Adequate justification for exemption from obtaining informed consent given <em>(if applicable)</em>.</td>
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<td>8</td>
<td>Informed Consent Document in Khasi, English and Hindi attached as per NEIGRIHMS SOP format.</td>
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<td>9</td>
<td>Information to the participant/ parent/guardian in layman (simple) language.</td>
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<td>10</td>
<td>Validated questionnaire both in Khasi, English &amp; Hindi attached <em>(if study involves interview/ questioning)</em></td>
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<td>11</td>
<td>Signature of all investigators (Principal &amp; Co-investigator) and Head of corresponding department obtained with date</td>
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<td>12</td>
<td>Compensation mentioned as per NEIGRIHMS guidelines in consent form part 1</td>
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<td>13</td>
<td>Confidentiality mentioned as per NEIGRIHMS guidelines in consent form part 1</td>
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<tr>
<td>14a</td>
<td>Separate consent form for subjects &lt; 7 yrs attached <em>(if applicable)</em></td>
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<tr>
<td>14b</td>
<td>Separate assent form for subjects &gt; 7 yrs &lt; 18 yrs attached <em>(if applicable)</em></td>
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<td>15</td>
<td>Separate consent form for cases and controls attached <em>(if applicable)</em></td>
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<tr>
<td>16</td>
<td>Ethical issues explained in detail with level of risk</td>
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<td>17</td>
<td>No discrepancy between Khasi, English &amp; Hindi consent form</td>
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<tr>
<td>18a</td>
<td>Declaration form from Guide (for all UG/PG/PhD/DM,MCh projects) regarding overall responsibility for the research</td>
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<tr>
<td>18b</td>
<td>Declaration form from principal investigators / Guide stating that all procedures used in the study are standard and professionally acceptable (for faculty projects/ for all UG/PG/PhD/DM,MCh)</td>
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</table>

Date: 

Signature of principal investigator

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12
(It is mandatory to submit this form along with proforma)

REVIEW EXEMPTION APPLICATION FORM

1 Principal Investigator’s Name:

_____________________________________________________

2 Department:

________________________________________________________

3 Title of Project:

________________________________________

4 Names of other participating staff and students:

___________________________________________________

5 Brief description of the project:
Please give a brief summary (approx. 300 words) of the nature of the proposal, including the aims/objectives/hypotheses of the project, rationale, participants’ description, and procedures/methods to be used in the project:-

6 State reasons why exemption from ethics review is requested?
✓ Audits of educational practices
✓ Research on microbes cultured in the laboratory
✓ Research on immortalized cell lines
✓ Research on cadavers or death certificates provided such research reveals no identifying personal data
✓ Analysis of data freely available in public domain
✓ Any other

(This should include justification for exemption e.g. study does not involve human participants. If exemption is being requested on the basis of low risk involved in the study please refer to the backside of this annexure.)

Principal Investigator’s signature: __________________________

Date __________
Forwarded by the Head of the department:

Name: ______________________________ Signature: ____________
Date__________

Recommendations by the IEC Member Secretary:

Exemption
Cannot be exempted
Reasons______________________________________________
Discussion at full board
Signature of the Member Secretary: ______________________
Date ______________

Final Decision:

Exemption
Cannot be exempted
Reasons______________________________________________
Discussion at full board
Signature of the Chairperson: __________________________
Date ______________

Final Decision at Full Board meeting held on
_________________________________________________________________
Signature of the Chairperson: __________________________
Date ______________

No research can be counted as low risk if it involves:

(i) Invasive physical procedures or potential for physical harm
(ii) Procedures which might cause mental/emotional stress or distress, moral or cultural offence
(iii) Personal or sensitive issues
(iv) Vulnerable groups
(v) Cross cultural research
(vi) Investigation of illegal behaviour(s)
(vii) Invasion of privacy
(viii) Collection of information that might be disadvantageous to the participant
(ix) Use of information already collected that is not in the public arena which might be disadvantageous to the participant
(x) Use of information already collected which was collected under agreement of confidentiality
(xi) Participants who are unable to give informed consent
(xii) Conflict of interest e.g. the researcher is also the lecturer, teacher, treatment-provider, colleague or employer of the research participants, or there is any other power relationship between the researcher and the research participants.
(xiii) Deception
(xiv) Audio or visual recording without consent
(xv) Withholding benefits from “control” groups
(xvi) Inducements
(xvii) Risks to the researcher

This list is not definitive but is intended to sensitize the researcher to the types of issues to be considered. Low risk research would involve the same risk as might be encountered in normal daily life.

Please check that your application / summary has discussed:

- Procedures for voluntary, informed consent
- Privacy & confidentiality
- Risk to participants
- Needs of dependent persons
- Conflict of interest
- Permission for access to participants from other institutions or bodies
- Inducements

In some circumstances research which appears to meet low risk criteria may need to be reviewed by the IEC. This might be because of requirements of:

- The publisher of the research
- An organization which is providing funding resources, existing data, access to participants etc.
SECTION – 3
FOR INTRAMURAL RESEARCH GRANT COMMITTEE

BUDGET DETAILS

1. Title of the Project:

2. Total amount required:

3. Year wise break-up of the amount:

4. Budget requirement:
   a. Consumable (Provide the list of items required with all relevant details)
   b. Non-consumable (Detailed justification required)
   c. Travel (Not for attending conference) – field work etc.

5. Justification for the budget:

6. For Faculty project:
   a. No. of intramural grants received in last five years:
   b. Enclose order copy of last intramural grant:
   c. Enclose copy of UC, SOE and progress report of last intramural grant:
   d. No. of extramural grants received in last five years:
   e. Enclose order copy of last extramural grant:
   f. Enclose copy of UC, SOE and progress report of last extramural grant:

7. For projects where faculty as a guide:
   a. Name of the Candidate:
   b. Study course:
   c. Year of the study:
   d. No. of previous intramural grant received:
   e. Enclose order copy of last intramural grant:
   f. Year of receiving the last intramural grant:
   g. Amount of receiving the last intramural grant:
   h. Enclose copy of UC, SOE and progress report of last intramural grant:
Declaration:

A) I/we declare that the infrastructure necessary for carrying out the above mentioned research scheme are available with me/us.
B) I/we agree to submit within, one month of termination of the scheme a final report on the work and an annual report within one month of expiry of a year if the project goes for more than one year. Extension of the project will be subject to approval of the report by the expert committee.
C) The faculty members those who have not submitted the final reports in respect of earlier projects granted by the Institute, are not entitled for the Institute Grant in future till they submit the report.

Principal Investigator

Co-Investigator (S)

Forwarded with remarks from Head of the Department
(in which The Principal Investigator is working)