
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## **1. Purpose**

To provide instructions for taking action and maintaining records that identify investigators/institutes who fail to follow the procedures written in the approved protocol or to comply with national / international guidelines for the conduct of human and animal research, including those who fail to respond to the MRIN EC's requests.

## **2. Scope**


This SOP applies to all MRIN EC approved research protocols involving human and animal subjects.

## **3. Responsibility**

The responsibility of the Board is to identify, decide the deviation and take appropriate action. The designated member of the MRIN EC members is responsible for collecting and recording the non-compliance list (AF/01-016/2020/01.4). Protocol Deviation Report must be reviewed by the Primary Reviewer

## **4. Flow chart**

<b><u>No.</u></b>	<b><u>Activity</u></b>	<b><u>Responsibility</u></b>
1	Noting protocol deviation / non-compliance / violation.	EC members and Chairperson
	↓	
2	Board discussion and decision	EC members and Chairperson
	↓	
3	Notify the investigator	EC Secretariat,
	↓	
4	Keep records and follow up	EC Secretariat

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## **5. Detailed instructions**

### **5.1 Noting Protocol Deviation/ Non-Compliance/ Violation**

EC members may identify protocol deviation/violation from progress report or site monitoring

- Ensure that the issues as well as the details of deviation /non-compliance/violation involving research investigators are included in the agenda of the MRIN EC meeting.
- Maintain a file that identifies investigators who are found to be non-compliance with national/international regulations or who fail to follow protocol approval stipulations or fail to respond to the MRIN EC's request for information/action.

### **5.2 The MRIN EC's Decision**


- The Board may suspend or terminate approval of current studies or refuse subsequent applications from the investigators cited. Such decisions are recorded in the minutes.
- The chairperson notifies the investigator of the MRIN EC's action in writing, when the Board
  - Suspend or terminate approval
  - Corrective action required
  - Site visit needed
  - refuse subsequent applications from the investigators cited

### **5.3 Notify the investigator**

- The MRIN EC Secretariat members record the MRIN EC's decision.
- Draft and type a notification letter.
- Get the Chairperson to sign and date the letter.
- Make copies of the notification letter.
- Send the original copy of the notification to the investigator.
- Send a copy of the notification to the relevant national authorities and institute
- Send the third copy to the sponsor or the sponsor's representative of the study

### **5.4 Keep records and follow up**

- Keep the last copy of the notification letter in the "non-compliance" folder of protocol file.
- Store the file in the shelf with an appropriate label.
- Follow up the action after a reasonable time.

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## **6. Glossary**

Deviation / Non-compliance / Violation      The MRIN EC monitors whether investigators do not perform the study in compliance with the approved protocol, ICH GCP, FDA regulations and/or fail to respond to the MRIN EC's request for information/action.


## **7. Annex**

Annex 1      AF/01-016/2020/01.4      Deviation/Non-Compliance/Violation Record

Annex 2      AF/02-016/2020/01.4      Document History

## **8. References**


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
**Annex 1**  
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**Deviation / Non-Compliance / Violation Record**

Application Number:	Date:
Study Title:	
Investigator	Contact No.:
Institution:	Contact No.:
Sponsor:	Contact No.:
<input type="checkbox"/> Deviation from protocol <input type="checkbox"/> Non-Compliance <input type="radio"/> Major <input type="radio"/> Minor <input type="checkbox"/> Violation	
Description:	
Found by:.....                      Reported by:..... Date:.....                      Date:.....	
<b>SIGNATURES:</b>	
Protocol Reviewer	Date : .....

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
<p><b>APPROVAL:</b></p> <p style="text-align: center;">_____ Date: .....</p> <p style="text-align: center;">Chairperson, MRIN EC</p> <p><b>COMPLETION:</b></p> <p style="text-align: center;">_____ Date:.....</p> <p style="text-align: center;">Secretary, MRIN EC</p>
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Form AF/02-016/2020/01.4

Document History

Author	Version	Date	Description
EC Members	01.0	2 January 2013	<b>Final version</b>
Ivet, Lia, Mona	01.1	11 October 2014	<ol style="list-style-type: none"> <li>1. Synchronize the topic number and SOP number</li> <li>2. Format Document History : Author, Version, Date and Description of the main change</li> <li>3. Section 5.1. Post approval documents should be reviewed by the primary reviewers</li> <li>4. Annex 1 : addition of section of reviewer's signature</li> </ol>
Ivet, Lia, Mona	01.2	15 November 2014	<ol style="list-style-type: none"> <li>1. Section 5.1. : Replace "Post Approval Documents" with "Protocol Deviation Report"</li> <li>2. "Protocol Deviation Report must be reviewed by the Primary Reviewer" under Sec 3</li> <li>3. Addition of "No further Action" and "Request Information" under Section 5.2.</li> </ol>
Budhianto Suhadi, Debbie S. Retnoningrum	01.3	1 April 2017	<ol style="list-style-type: none"> <li>1. Delete Komite Etik Penelitian Kesehatan (KEPK) on the header</li> </ol>
MW,IS,GE,LSH	01.4	2 January 2020	<ol style="list-style-type: none"> <li>1. 5.1 Replace the statement of "whenever protocol deviation.... Has been observed with "EC members may identify protocol deviation.... Or site monitoring"</li> <li>2. 5.2. Delete " Requires further information, Needs no further action " and replace with "suspend or terminate approval, corrective action required, site visit needed., etc."</li> </ol>

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			3. References : <ul style="list-style-type: none"> <li>• Delete link WHO</li> <li>• Delete FERCAP SOP</li> <li>• Add ICH Nov 2016</li> </ul> 4. Revise Annex
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