EXTERNAL REPORT

September 22, 2014

<NOTE: If the report is an unanticipated problem involving risk to subjects or others unrelated to the local research context, reporting to the DOD is not required.

Send when the organization is notified by any Federal department, agency or national organization that any part of the HRPP is under investigation for cause involving a DoD-supported research protocol.1>

<Attach associated minutes and send to:>
Director, Defense Research and Engineering
ddre@dtic.mil

<If the research is conducted or funded by the Department of the Navy (DOD), attach associated minutes and send to:>
Under Secretary of the Navy
1000 Navy Pentagon
Washington, D.C. 20350-1000

<If the research is also DHHS-regulated or subject to DHHS oversight by virtue of a federalwide assurance (FWA), send to the following address:2>
Kristina Borror
Director of the Division of Compliance Oversight
Office for Human Research Protections
The Department of Health and Human Services
The Tower Building
1101 Wootton Parkway, Suite 200
Rockville, MD 20852
Phone: 301-496-7005
Phone: 866-447-4777 (toll free)
Fax: 240-453-6909
IRPT.OS@hhs.gov

Dear Sir or Madam:

<Name of organization> is submitting this report in fulfillment of its regulatory requirement to follow written procedures for ensuring prompt reporting to the IRB,

2 See: http://www.hhs.gov/ohrp/policy/incidreport_ohrp.html
appropriate institutional officials, and department or agency heads of (i) any 
unanticipated problems involving risks to subjects or others or any serious or continuing 
non-compliance with the regulations or the requirements or determinations of the IRB; 
and (ii) any suspension or termination of IRB approval.

This is a report of: <delete all that do not apply> 
• An unanticipated problem involving risks to subjects or others; 
• Serious or continuing non-compliance with the regulations or the requirements or 
determinations of the IRB; and 
• A suspension or termination of IRB approval.

The institution conducting the research is:

<table>
<thead>
<tr>
<th>Organization:</th>
</tr>
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<tbody>
<tr>
<td>FWA:</td>
</tr>
<tr>
<td>IRB Registration:</td>
</tr>
<tr>
<td>Address:</td>
</tr>
<tr>
<td>Contact Name:</td>
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<tr>
<td>Contact Title:</td>
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<td>Contact Phone:</td>
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<td>Contact Fax:</td>
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<td>Contact Email:</td>
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</tbody>
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This report is in regard to:

<table>
<thead>
<tr>
<th>Type of Review:</th>
<th>&lt;Indicate Initial, Continuing, or Modification&gt;</th>
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<tbody>
<tr>
<td>Title:</td>
<td></td>
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<tr>
<td>Investigator:</td>
<td></td>
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<tr>
<td>IRB ID:</td>
<td></td>
</tr>
<tr>
<td>Funding:</td>
<td>&lt;Indicate “None” if there is none.&gt;</td>
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<tr>
<td>Grant Title:</td>
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<td>Grant ID:</td>
<td>&lt;Indicate “None” if there is none.&gt;</td>
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<tr>
<td>IND, IDE or HDE:</td>
<td>&lt;Indicate “None” if there is none.&gt;</td>
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<tr>
<td>Documents Reviewed:</td>
<td></td>
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<td>IRB Review Date:</td>
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Description of the problem including the findings of the organization and the reasons for 
the IRB’s decision:

<Insert description.>

Actions the organization is taking or plans to take to address the problem:

<Describe actions. For example: Revise the protocol, suspend subject enrollment, 
terminate the research, revise the informed consent document, inform enrolled 
subjects, or increase monitoring of subjects.>
Follow-up Plans

<Delete if none. Otherwise describe plans, if any, to send a follow-up or final report by the earlier of (1) a specific date or (2) when an investigation has been completed or a corrective action plan has been implemented.>

Please let us know if you need additional information.

Sincerely,

Organizational Official <or designee>
cc:  <Convened IRB by inclusion in the agenda materials as an information item>
     <Protocol Contact>
     <Principal Investigator>
     <Sponsor. Delete if none.>
     <Contract Research Organization. Delete if none>
     <Chairman or Supervisor of the Principal Investigator>
     <Sponsored Projects Services (SPS)>
     <Legal Counsel>
     <Risk Management>
     <Others as deemed appropriate by the Organizational Official.>
     <For international or collaborative research, the local research ethics committee or equivalent, as applicable>
     <The Privacy Officer of an organization, if the report involves unauthorized use, loss, or disclosure of the organization’s individually identifiable information>
     <The Information Security Officer of an organization, if the report involves violations of the organization’s information security requirements.>
     <The following regulatory agencies when they conduct, fund, or oversee the research when they want reporting separate from OHRP>
     Agency for International Development (22 CFR 225)
     Central Intelligence Agency (Executive order)
     Consumer Products Safety Commission (16 CFR 1028)
     The Department of Agriculture (7 CFR 1c)
     The Department of Commerce (15 CFR 27)
     The Department of Education (ED) (34 CFR 97)
     The Department of Energy (DOE) (10 CFR 745)
     The Department of Homeland Security (Public law 108-458 Sec. 8306)
     The Department of Justice (DOJ) (28 CFR 46)
     The Department of Transportation (49 CFR 11)
     The Environmental Protection Agency (EPA) (40 CFR 26)
     <send to:>
     The Environmental Protection Agency (EPA) Human Subjects Research Review official
     Housing and Urban Development (24 CFR 60)
National Aeronautics and Space Administration (14 CFR 1230)
National Science Foundation (45 CFR 690)
Office of Science and Technology Policy (Adoption of policy)
Social Security Administration (Public law 7.5.26)