The Ohio State University (Ohio State) is committed to providing a safe and healthy working environment for its employees. To meet this commitment, Ohio State has developed and implemented Safety, Health, and Environmental (SHE) Practices that address safety and environmental concerns for all employees. Additionally, Ohio State is subject to strict local, state, and federal regulations promulgated by such agencies as the Nuclear Regulatory Commission (NRC), the Environmental Protection Agency (EPA), and the Public Employment Risk Reduction Program, which has adopted Occupational Safety and Health Administration (OSHA) standards. Ohio State is also committed to complying with current safety regulations and guidelines as issued by the United States Departments of Health and Human Services and Agriculture, the National Institutes of Health (NIH) and the Centers for Disease Control and Prevention (CDC).

This biosafety manual provides safety guidelines, policies, and procedures for the use and manipulation of biohazards and recombinant and synthetic nucleic acid molecules. Although the implementation of these procedures is mainly the responsibility of the Principal Investigator (PI), success in biosafety depends upon the combined efforts of everyone in the laboratory. Planning for and implementation of biological safety must be part of every laboratory activity in which biohazard materials are used. The purpose of the overall biological safety plan is to ensure the safe handling of biohazardous materials in any work performed under Ohio State aegis and to thereby protect personnel, research outcomes, and the environment.

The biosafety program consolidates the compliance programs for the Public Employment Risk Reduction Program adoption of the OSHA Hazard Communication Standard (29 CFR 1910.1200), the OSHA Occupational Exposure to Hazardous Chemicals in the Laboratory (29 CFR 1910.1450), the OSHA Occupational Exposure to Bloodborne Pathogens Standard (29 CFR 1910. 1030), the NIH’s NIH Guidelines for Research Involving Recombinant and Synthetic Nucleic Acid Molecules (NIH Guidelines) and the CDC/NIH publication Biosafety in Microbiological and Biomedical Laboratories (6th edition). Additionally, the Institution’s Safety Management Guidebook provides Ohio State procedures for medical surveillance, sharps injuries, and working alone that are to be followed by principal investigators and laboratory personnel.

This Institutional Biosafety Manual applies to all Ohio State faculty, staff, hosted visitors, students, participating guests and volunteers, contract laborers, supplemental personnel and employees of firms working at locations where the Ohio State has management control of specific biohazards.

Biohazards at Ohio State are defined as infectious agents (i.e., pathogens) or materials produced by living organisms that may cause disease in other living organisms. This definition encompasses not only human pathogens, but also materials that may contain such pathogens (human-, nonhuman-primate- and other animal- and plant-sourced materials) that can cause disease in humans, animals or plants. Work with some experimental animals and arthropods also constitutes potential exposure to biohazardous materials since these animals may harbor infectious agents and/or proteins in their dander, urine, saliva, serum, etc., to which personnel may react or may become allergic.
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Section II
Responsibilities

The basic safety principle is that all injuries are preventable. Management, from the Ohio State President to the Principal Investigator/Supervisor, has a responsibility to encourage a safety effort in a sustained and consistent manner by establishing safety goals, promoting a positive safety culture, demanding accountability for safety performance, and providing the resources to the safety program.

II.1 Office of Environmental Health and Safety (EHS)

EHS has an institutional responsibility to help promote the safety and health of Ohio State employees. EHS personnel serve as safety consultants to the departments and other units of Ohio State and provide information on applicable safety-related regulations or guidelines. EHS personnel are involved in the development of safety practices and procedures. They also provide guidance to personnel in safety matters through consultation. The unit also has the responsibility of ensuring, through the auditing of laboratory facilities and work practices, that the work is completed in a safe and environmentally sound manner.

EHS has institutional responsibility for the disposal of radioactive materials, hazardous chemicals, and infectious waste. Training in specific areas of safety is provided by EHS personnel, including those associated with the collection and disposal of biological wastes.

Use, possession, and transfer of select agents and toxins must be in accordance with federal regulations. Select agents and toxins are biological agents and toxins that have the potential to pose a severe threat to the public, animal or plant health, or a threat to animal or plant products. EHS oversees the use, possession, and transfers of select agents and toxins to ensure compliance with federal regulations. Appendix B contains additional information on select agents and toxins. Questions about select agents and toxins should be directed to the Institutional Biosafety Officer (IBO) at 614-247-4867.

II.2 Office of Responsible Research Practices (ORRP)

The Office of Responsible Research Practices (ORRP), a unit of the Office of Research, provides support for review committees, many of which are federally mandated, including a Biomedical Institutional Review Board (IRB), a Cancer IRB, a Behavioral and Social Sciences IRB, the Institutional Animal Care and Use Committee (IACUC), and the Institutional Biosafety Committee (IBC). The Office of Responsible Research Practices also supports the institution by promoting ethical conduct of research and educating Ohio State students, faculty, and staff regarding research regulations. Access to eProtocol, information on policies and procedures, and Statements of Assurance can be obtained through the ORRP website (http://orrp.osu.edu).
II.3 Institutional Biosafety Officer (IBO)

The Institutional Biosafety Officer (IBO) is responsible for the development, implementation, and direction of the comprehensive biological safety program at the institution. The biosafety program includes work with human tissues, fluids, cells, or cell cultures, recombinant or synthetic nucleic acids, transgenic plants/animals, infectious agents, work with animals known to be vectors of zoonotic diseases, gene transfer, xenotransplantation, etc. This individual serves as the Institutional Biosafety Officer as defined by the NIH Guidelines and thereby serves on the Institutional Biosafety Committee. The membership of this committee reviews and approves protocols involving the use of biohazards, recombinant or synthetic nucleic acid, and gene transfer. The IBO assists investigators and staff with all matters related to biosafety. The IBO audits laboratories and work practices for compliance with Ohio State policies and procedures.

II.4 Institutional Biosafety Committee (IBC)

Ohio State is committed to the safe, legal, and ethical use of biologically derived hazardous materials. Acting as the agent for Ohio State in such matters, the Institutional Biosafety Committee acts to assure that activities involving recombinant or synthetic nucleic acid molecules and biohazards meet the legal and ethical requirements for the responsible use of these agents, that safety levels are appropriately classified, and that work is performed in accordance with good safety practices. Additionally, the committee membership works to establish policies and make recommendations regarding such activities, maintain and promote an open and cooperative relationship with investigators and other committees, and educate concerning the regulatory requirements of biosafety.

The IBC reviews all aspects of research involving recombinant or synthetic nucleic acid molecules, vectors, and host cells that cannot be classified as human or animal biohazards. The committee also has responsibility for traditional research activities that utilize biohazards and for recombinant or synthetic nucleic acid projects that include biohazards (including mammalian viral vectors, pathogenic organisms, and the use of human blood, tissues, body fluids, cells, blood products, human stem cells and other potentially infectious materials). All human gene transfer protocols, those animal gene transfer protocols not exempted from review by the NIH Guidelines, and potential dual use research of concern is also reviewed by the IBC.

The IBC retains the authority to refuse permission for a principal investigator to work with specific biological agents if, in the opinion of the committee, public health or the environment would be compromised by granting such use.

II.5 President and Vice Presidents

The Ohio State President and Vice Presidents encourage a climate of compliance with federal, state, and local regulations and support an ongoing commitment to this compliance.
As needed, an ad-hoc committee may be formed to consider policy implications of emerging technologies and processes that have a biosafety component. This committee will work with the standing Ohio State committees and administrative units with responsibilities for biosafety issues in research and teaching to provide advice to the Vice President for Research.

II.6 Deans

Deans encourage compliance with safety, health, and environmental practices by departments within their jurisdiction. All academic and non-academic departments, schools, and divisions shall participate in all applicable required programs.

II.7 Department Chairs, Center Directors, and Other Facility Directors

Department Chairs/Directors shall:

- Develop building emergency and evacuation plans (BEAPs), appoint building safety committees, departmental biosafety officers, and appoint building safety managers and alternates (in some cases with the associate dean for research and research officers);

- Maintain discipline, enforce rules and regulations, and take prompt, effective corrective action when necessary. The department chair shall also provide assistance to EHS and ORRP staff when investigations arise involving the conduct or work practices of PIs and/or other personnel in the department.

- Ensure the compliance of principal investigators and other supervisory personnel with federal, state, and local regulations and Ohio State policies applicable to the department’s work, including enrollment of individuals in the Occupational Health program. Regulatory and policy documents are available from EHS and ORRP. The department chair may delegate safety- and health-related responsibilities to principal investigators or other supervisors, but it is the department chair’s responsibility to understand the regulations and to see that the requirements are met.

- Take corrective actions to halt any violations should violations of Ohio State biohazard policies occur, in concert with the IBO, EHS, the department safety officer, and the appropriate Ohio State standing committee.

II.8 Principal Investigators (PIs) and Supervisors

Direct responsibility for compliance with Ohio State’s safety and health programs is assigned to the Principal Investigator. This means that the PI shall provide a safe workplace and shall implement health and safety programs. This includes ensuring that personnel are adequately trained, research protocols/safety plans are prepared and submitted to the IBC for review, and laboratories are submitted to periodic inspections. PIs are responsible for maintaining good working order of equipment in their laboratories (including the appropriate certification of biological safety cabinets).
Principal Investigators shall:

- Communicate to those in the laboratory Ohio State’s high priority regarding health and safety and concern for the environment and shall ensure that environmental, health and safety obligations are fulfilled by all personnel in the laboratory.

- Analyze work procedures for hazard identification and implementation of measures to eliminate or control workplace hazards prior to beginning new experiments.

- Ensure that all laboratory personnel, maintenance personnel and visitors who may be exposed to any biohazard are informed in advance of their potential risk and the behavior required to minimize that risk.

- Correct deficiencies noted during the periodic laboratory inspection and respond in writing with the corrective action and date of implementation, to EHS within the required time.

- Submit a research protocol (Safety Plan) covering the use of biohazardous agents for review and approval by the IBC before laboratory work commences and submit the laboratory to periodic inspections by a representative of EHS.

- Submit any significant changes in the research protocol to the IBC for review and approval.

- Ensure any research projects covered by the NIH Guidelines that require prior agency approval before initiation, are reviewed by the IBC before seeking or obtaining agency approval. Experiments that require IBC approval prior to initiation are reviewed and approved before beginning work.

- Encourage regular self-assessment inspections by employees in order to review work habits and correct deficiencies. Prompt reporting of health and safety problems by project personnel is to be encouraged. Persons who file reports concerning laboratory shortcomings in good faith will be protected from retaliatory actions based on such filings.

- Ensure that all individuals in the laboratory know how to access the Institutional Biosafety Manual available on the EHS website and maintain a written acknowledgment of understanding by these individuals.

- Ensure training of all individuals involved in the handling and disposal of biohazard agents and that all training records are maintained as directed by the standards; The PI is responsible for taking all safety training required by his/her laboratory staff.

- Create and foster an environment in the laboratory that encourages open discussion of biosafety issues, problems, and modifications of procedures.
- Ensure that Personal Protective Equipment (PPE) appropriate to the biohazardous agent(s) is available, in good condition, and utilized appropriately.

- Ensure the participation of all personnel in a Medical Surveillance Program (i.e., Online Risk Assessment Tool (ORAT)). Personnel should identify all use of biohazardous agents as part of the ORAT completion.

- Ensure that all accidents and biohazard exposures are reported as required under Ohio State policy in a timely manner to the IBO and the Chair of the IBC. The Biohazard Incident Reporting policy and forms can be found at http://orrp.osu.edu/ibc/osuibcpolicies/incidentreporting/.

- Notify the IBO if a laboratory-acquired infection is known or suspected.

- Stop any work posing imminent danger. Prudent practices are to be employed by those working in the laboratory.

- Ensure that appropriate signage is used at the entrance(s) to and within the laboratory. Animal Hazard Safety Protocol (AHSP) signage must be in place in the vivarium before beginning animal experiments which include hazardous materials (consult with the animal vivarium supervisor for more information).

- Develop (with the IBO) plans for handling emergencies (accidental spills, fires, riots, etc.).

- Ensure that the animal vivarium team leader and ULAR hazard liaison is notified via the eProtocol system at least three working days before animals under the care of ULAR staff are treated with biohazardous agents. Consultation with ULAR and EHS personnel may be needed to ensure that risks and required PPE (personal protective equipment) are understood by all individuals involved. Post Animal Hazard Safety Protocols (AHSP) on applicable door when using hazardous agents for the duration specified on the form.

II.9 The Individual

YOU ARE RESPONSIBLE FOR YOUR OWN SAFETY!

The health and safety of each employee is extremely important. Employees should bring their concerns to their supervisor, the departmental safety officer, department chair, the IBO, the IBC, or EHS.

Each employee is expected to be conscientious in assuming personal safety responsibility from the first day on the job. Each employee must understand that he or she is responsible for working safely.
The individual shall:

- Comply with Ohio State’s safety policies and rules and follow both oral and written instructions from the principal investigator or supervisor. The individual shall report to the principal investigator any unsafe conditions and/or any accident or exposure to chemicals or biological agents. If the individual receives no response or an unsatisfactory response, he/she should contact the department chair, EHS or the chair of the IBC.

- Know the hazards of the chemicals and biological agents in the workplace as well as proper handling and disposal procedures. Training shall be provided by the principal investigator or designee prior to the commencement of work. The individual must minimize all potential exposures to infectious materials or contaminated items. He/she will learn what precautions and protective equipment are needed for specific jobs and practice good hygiene.

II.10 Students, Visitors and Guests

Ohio State is committed to providing a safe and healthy work environment to its employees that, in turn, fosters a safe learning environment for students. Ohio State requires students, visitors, and guests to abide by applicable safety guidelines when using campus facilities. It is the policy of Ohio State to ensure that all students who might be exposed to hazardous materials in the course of their activities are adequately protected.
Section III
Risk Assessment and Research Protocols

III.1. Risk Assessment

New research or development initiatives are evaluated by the Principal Investigator in the early planning stages for the hazards that can be posed by the biological, toxic, flammable, reactive, explosive, and/or radioactive materials required by the proposed work. New or inexperienced investigators are encouraged to read the Institutional Laboratory Biosafety Manual and attachments, to seek consultation with the departmental safety officers, the chair of the Institutional Biosafety Committee, the Institutional Biosafety Officer, and/or Environmental Health and Safety or industrial hygiene personnel, as well as to review published expert opinion regarding regulatory requirements.

The assessment of infection risks associated with the laboratory use of biohazard materials requires the proper risk classification of the material, the risk modification of the manipulations of the material, the environmental risks associated with the material, and the laboratory requirements for containment associated with those risks. Agents of Risk Group 1 or 2 can usually be assessed in consultation with the departmental safety officer but may still require the submission of a research protocol to the IBC. It is required that PIs submit a research protocol for approval by the Institutional Biosafety Committee for the use of Risk Group 2 or higher agents prior to the beginning of the research. Agents that require BSL3 containment require submissions for approval of research protocols and are best managed by early consultation with the BSL-3 Operations Group, the Institutional Biosafety Committee, the Institutional Biosafety Officer, and Environmental Health and Safety (EHS) personnel.

Established, ongoing research or development initiatives should be evaluated annually by the Principal Investigator to assure that risks have not changed and that the established safety program is in compliance with the current regulatory requirements (Institutional Biosafety Manual, latest edition). The PI has the responsibility for auditing facilities and work practices to help ensure that the work is completed in a safe and environmentally sound manner. New initiatives that increase the risk profile of the laboratory must be accompanied by appropriate changes in the laboratory’s research protocol (such as engineering controls, PPE, training requirements, spill containment, etc.) to reflect the increase in risk. Work at the new risk level must not begin until the appropriate research protocol is approved and in place. New investigation initiatives in the same laboratory that do not change the risk profile of that laboratory but significantly change the scope of work, also requires an update to the laboratory’s research protocol. Beginning in 2015, all IBC protocols will be approved for a 5-year time period. At the end of the approval period, a renewal protocol must be submitted for approval, or the protocol and the approved work will be terminated.
The training of personnel must be documented in writing and the records kept by the Principal Investigator. All personnel must be made aware of the potential hazards associated with their work and must be trained in the designated safety procedures as well as the appropriate use of the safety equipment (including personal protective equipment and engineering controls) required and the appropriate handling of spills. It is the Principal Investigator's responsibility to make sure that all training is completed at the required intervals by their laboratory personnel. Be aware that while allowing unpaid volunteers (both minors and adults) to work in the laboratory is a department level policy, if allowed all volunteers must take all the required safety training prior to beginning work in the laboratory. Ohio State’s policy regarding minors on campus must also be followed.

III.2. Risk Group Assessment

There are considerable variances in the assignment of risk to infectious agents. In order to standardize the approach to this important consideration, Risk Groups will be selected according to the information in Table III.1.

Information pertaining to the assignment of agents to risk groups may be found in Appendix B of the NIH Guidelines for Research Involving Recombinant and Synthetic Nucleic Acid Molecules (NIH Guidelines) and Section VIII of the CDC/NIH publication Biosafety in Microbiological and Biomedical Laboratories, 6th Ed.

This method of Risk Group (RG) assignment is the most consistent with that of the World Health Organization (WHO).

Table III.1 Basis for the Classification of Biohazardous Agents by Risk Groups (RG).

<table>
<thead>
<tr>
<th>Risk Group 1 (RG1)</th>
<th>Agents not associated with disease in healthy human adults</th>
</tr>
</thead>
<tbody>
<tr>
<td>Risk Group 2 (RG2)</td>
<td>Agents associated with human disease that is rarely serious and for which preventative or therapeutic interventions are <em>often</em> available. All human source material (including blood, human cell lines and other potentially infectious materials) are considered RG2.</td>
</tr>
<tr>
<td>Risk Group 3 (RG3)</td>
<td>Agents associated with serious or lethal human disease for which preventive or therapeutic interventions <em>may</em> be available (high individual risk but low community risk).</td>
</tr>
<tr>
<td>Risk Group 4 (RG4)</td>
<td>Agents likely to cause serious or lethal human disease for which preventive or therapeutic interventions <em>not usually</em> available (high individual risk and high community risk).</td>
</tr>
</tbody>
</table>
III.3. Research Protocol

The research protocol is a project-specific biological safety plan for research and teaching laboratories, or common facilities shared by more than one of these activities. In each laboratory/facility, the PI/supervisor must specify the safety practices/procedures to be used in the laboratory and is responsible for the implementation of the plan. Approved practices must be based on the risk assessment for that laboratory/project and must comply with accepted national standards. A research protocol for the containment of Risk Group 2 and 3 infectious agents and any work with recombinant or synthetic nucleic acid molecules must be submitted to the IBC prior to implementation (work with agents of RG 4 is not permitted). The research protocol must address the following issues as appropriate to the individual laboratory risks:

- **Risk Group of infectious hazard:** the characteristics of infectious agents, the laboratory requirements and primary laboratory hazards of working with the agents can be found in *Biosafety in Microbiological and Biomedical Laboratories* (6th Edition), in *Physical and Biological Hazards of the Workplace*, in similar references, or by contacting the Institutional Biosafety Officer, the IBC, or EHS personnel.

- **Bloodborne Pathogens Standard (OSHA) [29 CFR 1910.1030]:** laboratory use of human or animal blood (intentionally infected with HIV or HBV), blood components, or tissues.

- **Containment requirements/engineering controls:** for biosafety level (BSL) 2 or 3, biosafety cabinets, storage requirements, transport containers, personal protective equipment (PPE) (i.e., hoods, eye protection, gloves, gowns, respirators, etc.), spill management and waste disposal.

- **University Laboratory Animal Resources:** animal husbandry matters and safety of researchers and animal facility (ULAR) personnel.

- **Exposure and Post-Exposure Follow-up:** exposure definition, prophylaxis if available (e.g., Hepatitis B vaccine), and exposure follow-up coordinated through Occupational Health and Wellness (Employee Health).

- **Specific safety training requirements:** all laboratory personnel (including the Principal Investigator) must be trained on the specific standard operating procedures to be used in the research; personnel must also be informed according to Public Employment Risk Reduction Program regulations (i.e. OSHA regulations) of the potential hazards associated with their work and must be trained in the designated safety procedures, use of safety equipment and Personal Protective Equipment, appropriate waste disposal, and availability of preventive measures such as vaccines. Examples of annually required education: standard operating procedures, bloodborne pathogens training for blood or body fluid exposure prevention, respirator training, safe animal handling, hearing conservation, laboratory chemical hazards, etc.
Training records: protocols for recording and maintaining records of initial personnel training and the annually required yearly exposure-prevention education programs must be established and in place.

III.4. Safety Documentation and References

Laboratory safety documents should be easily recognized and accessible for laboratory/facility personnel and safety personnel. As applicable, safety documentation should include the following:

- Hazard Communication Program
- Chemical Hygiene Plan
- Departmental Emergency Plans
- Radiation Safety Information
- Annual Chemical Inventory
- Safety Data Sheets (Safety Data Sheets) or access thereto
- Bloodborne Pathogens Exposure Control Plan
- Approved Research Protocols (IBC, IRB and IACUC) and Annual Reviews
- Personal Protective Equipment (PPE) hazard assessment
- Respirator Records (Respirator Plan) plus records of examinations and Fit Test Reports
- Training Documentation

Boilerplates of some of the elements of the documents listed above may be found at the EHS website at www.ehs.osu.edu.

III.5. Select Agents

The use, possession or transfer of select agents requires prior approval by the Responsible Official. For additional information on select agents please see Appendix B of this document or contact the IBO at 614-247-4867.
Section IV
Recombinant and Synthetic Nucleic Acids, Gene Transfer, and Transgenic Organisms

Recombinant and synthetic nucleic acids are defined as (1) molecules that a) are constructed by joining nucleic acid molecules and b) that can replicate in a living cell, i.e., recombinant nucleic acids, (2) nucleic acid molecules that are chemically or by other means synthesized or amplified, including those that are chemically or otherwise modified but can base pair with naturally occurring nucleic acid molecules, i.e., synthetic nucleic acids, or (3) molecules that result from the replication of those described in (1) or (2) above.

All work with recombinant and synthetic nucleic acid molecules requires an IBC protocol and, if the research is not exempt from full committee review under the NIH Guidelines for Research Involving Recombinant or Synthetic Nucleic Acid Molecules (NIH Guidelines), it must be sent for full committee review and be approved by the IBC. Good laboratory practices and containment procedures are to be followed by all individuals using recombinant and synthetic nucleic acids at Ohio State. Exempted research is also subject to inspection by the Institutional Biosafety Officer or other EHS personnel.

All Principal Investigators intending to use recombinant and synthetic nucleic acid molecules shall notify the IBC by submitting an electronic research protocol (e-Protocol) according to the nature of the research. All recombinant and synthetic nucleic acid research falls into the six classes below:

- Experiments that require IBC approval and NIH director approval before initiation
- Experiments that require NIH/OSP and IBC approval before initiation
- Experiments involving Human Gene Transfer that require IBC approval before initiation
- Experiments that require IBC approval before initiation
- Experiments that require IBC notice simultaneous with initiation
- Exempt experiments

Experiments involving the generation of recombinant and synthetic nucleic acids require, at a minimum, registration with the Institutional Biosafety Committee (IBC) through the submission of an IBC protocol. Some experiments also require approval of the IBC. The NIH Guidelines are applicable to all research conducted at or sponsored by an institution that receives funds from the National Institutes of Health for recombinant and synthetic nucleic acid research. Ohio State is required to monitor all recombinant and synthetic nucleic acid research. Compliance is required to protect employees, the community, and the environment from the creation and release of any novel organisms that might be pathogenic to man, animals, and plants or harmful to the environment.
The NIH Guidelines deal primarily with laboratory research and human gene-therapy protocols. A few types of experiments are prohibited by the NIH Guidelines and several others require prior approval by NIH. The rest come under the jurisdiction of the Ohio State IBC, which reviews the research and sets the level of biosafety and containment necessary to safely conduct the experiments. In recent years, the Guidelines have been relaxed considerably and much of the recombinant and synthetic nucleic acid research is exempt from full committee review. But even the research that is exempted from full committee review must be registered through the submission of an IBC protocol and carried out using prudent laboratory practices.

Most regulated experiments involve hosts and/or genes that are derived from etiologic agents or have a known biohazard associated with them. The release of genetically engineered organisms into the environment, their use as drugs or food products, and human gene-therapy protocols may be regulated by the USDA, EPA and FDA. The IBC may not have the sole authority to approve research in these areas and researchers may need to obtain permission from the federal agencies in addition to the IBC.

All protocols involving the generation of recombinant and synthetic nucleic acid molecules for human gene transfer must be approved by the IBC prior to the enrollment of the first human subject.

Investigators who use transgenic plants or animals must submit a recombinant and synthetic nucleic acid protocol and submit it via e-Protocol to the IBC. The IBC will review the form to determine if IBC approval of the experiments is necessary.

**IV.1. Institutional Biosafety Committee (IBC)**

An Institutional Biosafety Committee (IBC) is required at all institutions that receive funding from the National Institutes of Health (NIH). The IBC is composed of researchers, administrators, Ohio State staff, and community members. The committee meets as needed to review research protocols and advise on matters of recombinant and synthetic nucleic acid safety or biosafety matters. Different members of the IBC review recombinant and synthetic nucleic acid experimentation, biohazard research, and gene transfer experiments based on their areas of expertise. All non-exempt recombinant and synthetic nucleic acid and gene transfer protocols are approved by the full committee. If the protocol is considered exempt by the NIH Guidelines, it is approved administratively without being reviewed by the full committee. All recombinant and synthetic nucleic acid research at Ohio State, regardless of funding source, must be conducted in accordance with the NIH Guidelines for Research Involving Recombinant and Synthetic Nucleic Acid Molecules and must be registered with the Ohio State IBC.

The Ohio State IBC is further charged with reviewing and approving research conducted with human, plant, or animal pathogens. This review is conducted pursuant to the Centers for Disease Control and Prevention (CDC)/NIH publication, Biosafety in Microbiological and Biomedical Laboratories (6th edition).
The Ohio State IBC in conjunction with the Office of Environmental Health and Safety also provides guidance to the Ohio State research community regarding proper acquisition, handling, transfer, and disposal of potentially hazardous or regulated biological materials. The Office of Responsible Research Practices (ORRP) will also provide assistance with IBC registrations and applications. Additional information is provided at the ORRP Biosafety web page (https://orrp.osu.edu/ibc/) under the Investigator Guidance link. This site provides detailed descriptions of IBC registration requirements, including a comprehensive Guidance Document, and the IBC review process.

**IV.2. Principal Investigator’s Responsibilities**

Under the *NIH Guidelines* a PI has the responsibility to:

1. Evaluate the proposed research and establish appropriate containment conditions for that research.
2. Inform all laboratory personnel of the potential hazards associated with the work.
3. Develop an appropriate safety plan and procedures to minimize potential personnel exposure to hazardous materials.
4. Ensure that the host/vector systems used in all research projects are safe.
5. Ensure that all recombinant and synthetic nucleic acid waste is appropriately decontaminated prior to disposal.
Section V
Biohazard Signs and Tags

The United States Occupational Safety and Health Administration (OSHA) regulations (29 CFR 1910.145, *Specifications for accident prevention signs and tags*) require that warning signs and/or symbols be used to warn personnel and visitors of the potential hazards in the workplace. Specifically, with regard to biohazards, the universal biohazard sign must be used to “…signify the actual or potential presence of a biohazard and to identify equipment, containers, rooms, materials, experimental animals or combinations thereof, which contain or are contaminated with, viable hazardous agents.” OSHA recommends that biohazard signs be fluorescent orange or orange-red with the lettering and symbols a contrasting color. An example of a sign is given at the end of this section (Figure 5.1).

- Ohio State requires that the universal biohazard symbol be used to designate the presence of materials defined as biohazard (see Section I of this document)
- All laboratories must display room signs signifying the biohazards present, an emergency contact and phone number, and the necessary precautions to be taken when entering or working in the area (room signs [Fig 5.2] may be requested through EHS by using the “Room Sign Request Form” found on the EHS website)
- PIs are responsible for ensuring that hazard signs are posted and are current and accurate.
- When using experimental animals cared for by ULAR staff, the PI must give a minimum of three days’ notice via the e-Protocol online system, to the animal vivarium team leader and ULAR hazard liaison before exposing or treating the animals with biohazardous agents. This will trigger an email to the study team outlining animal room requirements and allow ULAR staff to prepare for appropriate animal husbandry. A working day is defined as a “day” during which Ohio State offices are open and excludes weekends and holidays. The PI or laboratory staff should confirm the appropriate signage is posted at the animal room level as indicated on the Ohio State Animal Hazard Safety Protocols (AHSP) prior to initiating hazardous work. The AHSP is provided to the PI and ULAR supervisors have access to the signage. A research protocol for the use of the particular agent as prepared by the PI must be submitted to the IBC and the research protocol must receive IBC approval before the research can begin.
OSU Stadium

**PI**
Dr. Brutus Buckeye

**Emergency Contacts**
Rose Bowl: 614-xxx-xxx
Ryan Day: 614-xxx-xxx

**IN EMERGENCY:** Firemen may enter: Call office of Radiation Safety 2-1284 or University Police 2-2121. Radiation Safety Emergency Pager: 614-240-0705

**Designated Area Lab**
Access limited to authorized personnel only

**No Food or Drink**
Section VI
Containment

The term containment is used to describe safe methods for managing biohazard agents in the laboratory environment. The three essential elements of containment are (1) laboratory practice and technique, (2) safety equipment, and (3) facility design. The purpose of containment is to reduce exposure of laboratory workers and others to potentially hazard agents and prevent the escape of these agents into the outside environment.

Research activities involving biohazard agents of RG 2 or higher can only be conducted with prior approval of the IBC. The elements of a safety plan have been discussed previously. The NIH/CDC manual *Biosafety in Microbiological and Biomedical Laboratories (BMBL)* (6th Edition) provides guidance for the appropriate containment of biohazard work. The biosafety levels are based on the probability of occupationally-acquired infections resulting from the handling of specific agents in the laboratory.

CDC describes four biosafety levels (BSLs) which consist of combinations of laboratory practices and techniques, safety equipment, and laboratory facilities. Each combination is specifically appropriate for the operations conducted, the documented or suspected routes of transmission of the infectious agents, and for the laboratory function or activity. The recommended biosafety level for an organism represents the conditions under which the agent can be ordinarily handled safely. When specific information is available to suggest that virulence, pathogenicity, antibiotic resistance patterns, vaccine and treatment availability, or other factors are significantly altered, more (or less) stringent practices may be specified.

**Biosafety Level 1** is appropriate for work done with defined and characterized strains of viable microorganisms not known to cause disease in healthy adult humans. It represents a basic level of containment that relies on standard microbiological practices with no special primary or secondary barriers recommended, other than a sink for hand washing.

**Biosafety Level 2** is applicable to work done with a broad spectrum of indigenous moderate-risk agents present in the community and associated with human disease of varying severity. Agents can be used safely on the open bench, provided the potential for producing splashes or aerosols is low. Primary hazards to personnel working with these agents relate to accidental percutaneous or mucous membrane exposures or ingestion of infectious materials. Procedures with high aerosol or splash potential must be conducted in primary containment equipment such as biosafety cabinets. Personal Protective Equipment (PPE) such as splash shields, face protection, gowns and gloves should be used as appropriate. Hand washing and waste decontamination facilities must also be available.
**Biosafety Level 3** is applicable to work done with indigenous or exotic agents with a potential for respiratory transmission and which may cause serious and potentially lethal infection. Primary hazards to personnel working with these agents (i.e., *Mycobacterium tuberculosis*, *St. Louis encephalitis virus* and *Coxiella burnetii*) include auto-inoculation, ingestion and exposure to infectious aerosols. Greater emphasis is placed on primary and secondary barriers to protect personnel in adjoining areas, the community and the environment from exposure to infectious aerosols. For example, all laboratory manipulations should be performed in biological safety cabinets or other enclosed equipment. Secondary barriers include controlled access to the laboratory and a specialized ventilation system to prevent the release of infectious agents in the event an accidental release occurs in the laboratory.

**Biosafety Level 4** is applicable for work with dangerous and exotic agents that pose a high individual risk of life-threatening disease, which may be transmitted via the aerosol route and for which there is no available vaccine or therapy. Agents with close or identical antigenic relationship to Biosafety Level 4 agents should also be handled at this level. Primary hazards to workers include respiratory exposure to infectious aerosols, mucous membrane exposure to infectious droplets and auto-inoculation. The facility is generally a separate building or a completely isolated zone within a complex with specialized ventilation and waste management systems to prevent the release of viable agents to the environment. All manipulations of potentially infected materials and isolates pose a high risk of exposure and infection to personnel, the community and the environment. Isolation of aerosolized infectious materials is accomplished primarily by working in a Class III biological safety cabinet and a full-body, air-supplied positive pressure personnel suit.

**Vertebrate Animal Biosafety Levels**

There are four animal biosafety levels (ABSLs), designated Animal Biosafety Level 1 through 4, for work with infectious agents in mammals. The levels are combinations of practices, safety equipment and facilities for experiments on animals infected with agents that produce or may produce human infection. In general, the biosafety level recommended for working with an infectious agent in vivo and in vitro is comparable.

**Animal Biosafety Level 1** is suitable for work involving well characterized agents that are not known to cause disease in healthy human adults, and that are of minimal potential hazard to laboratory personnel and the environment.

**Animal Biosafety Level 2** is suitable for work with those agents associated with human disease. It addresses hazards from ingestion as well as from percutaneous and mucous membrane exposure.

**Animal Biosafety Level 3** is suitable for work with animals infected with indigenous or exotic agents that present the potential of aerosol transmission and of causing serious or potentially lethal disease.

**Animal Biosafety Level 4** is suitable for addressing dangerous and exotic agents that pose high risk of like threatening disease, aerosol transmission, or related agents with unknown risk of transmission.
Complete descriptions of all **Biosafety Levels and Animal Biosafety Levels** are outlined in the NIH/CDC manual *Biosafety in Microbiological and Biomedical Laboratories (BMBL)* (6th Edition). The *BMBL* provides minimum guidelines for containment of biohazards. Ohio State containment requirements and laboratory practices may be more stringent. When in doubt, contact the Institutional Biosafety Officer for confirmation of Ohio State requirements.

Research at the Ohio State is limited to containment at Biosafety Levels 1, 2, or 3. Infectious materials must be clearly identified and stored in such a manner as to preclude accidental exposure. This normally includes double containment and labeling of the storage freezer/refrigerator/liquid nitrogen tank.

**VI.1. Laboratory-acquired Infections**

A number of infectious agents have been documented as causes of laboratory-acquired infections. Included in the list are bacterial, viral, chlamidial and rickettsial, and parasitic organisms.

**VI.2. Laboratory Practices**

The most important element of containment is strict adherence to standard microbiological practice and techniques. Persons working with biohazard agents or infected materials shall be aware of potential hazards and shall be trained and proficient in the practices and techniques required for safe handling. When standard laboratory practices are not sufficient to control the hazard associated with a particular agent or laboratory procedure, additional measures such as safety equipment and facility design must be used.

**VI.3. Safety Equipment**

Safety equipment includes personal protective equipment, biological safety cabinets, sealed, leak proof containers, and other engineering controls designed to prevent or minimize exposures to hazardous biological materials. The use of vaccines, if available, is encouraged or in some instances specified to provide an increased level of personal protection.

**VI.3.1. Biological Safety Cabinets (BSC)**

The biological safety cabinet is the principal device used to provide containment of infectious splashes or aerosols. Biological Safety Cabinets are designed to protect the worker, the integrity of the experiment, and the environment. There are three types of biological safety cabinets: Class I, Class II, and Class III.
CDC and NIH have published a document entitled *Primary Containment for Biohazards: Selection, Installation and Use of Biological Safety Cabinets* that is available for reference concerning the specifics of BSC use, including a section on appropriate risk assessment. This document is available as Appendix A of the NIH/CDC manual *Biosafety in Microbiological and Biomedical Laboratories (BMBL)* (6th Edition).

VI.3.1.1. Class I BSCs

The Class I BSC provides personnel and environmental protection but no product protection. It is similar in airflow to a chemical fume hood, but has a High Efficiency Particulate Air (HEPA) filter in the exhaust system to protect the environment.

VI.3.1.2. Class II (Types A1, A2, B1, and B2) BSCs

Class II BSCs provide personnel, environmental and product protection. Airflow is drawn around the operator into the front grille of the cabinet providing personnel protection. In addition, the downward laminar flow of HEPA-filtered air provides product protection by minimizing the chance of cross-contamination along the work surface of the cabinet. Because cabinet air has passed through the exhaust HEPA filter, it is contaminant-free (environment protection), and may be recirculated to the cabinet workspace, back into the laboratory (Class II Type A1 and A2 BSCs) or ducted out of the building (Class II Type B1, B2, and A2 BSCs). A Class II Type A2 BSC may optionally be installed such that air re-circulates back into the room or is ducted outdoors. Under the new NSF/ANSI 49 Standard, no newly installed Type A cabinet may be directly ducted to the building’s exhaust system; thimble exhaust connections should be used to connect all new installations of Type A BSCs to the building exhaust system.

HEPA filters are effective at trapping particulates and infectious agents, but not at capturing volatile chemicals or gases. Only Class II Type B2 BSCs that have 100% of air ducted to the outside should be used when working with volatile chemicals. Class II Type B1 BSCs recirculate 30% of exhaust air to the work area and should ONLY be used with minute amounts of volatile chemicals as long as the re-circulating vapors do not present a problem in the work. The same is true of Class II Type A2 cabinets that are vented to the outdoors (with the exception that this type of cabinet re-circulates 70% of the air back to the cabinet); when the Type A2 cabinet is vented back into the room, it should NOT be used with toxic chemicals.

All Class II cabinets are designed for work with microorganisms assigned to Risk Groups 1, 2 and 3. Class II cabinets provide the microbe-free work environment necessary for cell culture propagation, and also may be used for the formulation of nonvolatile antineoplastics or chemotherapeutic drugs.
VI.3.1.3. Class III BSCs

Class III cabinets provide the highest level of protection. A Class III BSC is a totally enclosed glove-box cabinet of gas-tight construction. The cabinet is maintained under negative air pressure of at least 0.5 inches of water gauge. Supply air is drawn into the cabinet through HEPA filters, and the exhaust air is filtered by two HEPA filters in series before discharge to the outside. Generally, the ventilation system is separate from the facility’s ventilation system. Class III cabinets are available for high-risk biological agents.

VI.3.1.4. Horizontal Laminar Flow “Clean Benches”

Horizontal Laminar Flow “Clean Benches” are not BSCs. HEPA-filtered air flows across the work surface and toward the user. These devices provide product protection ONLY. They can be used for certain clean activities, such as dust-free assembly of sterile equipment or electronic devices. These benches should not be used when handling cell culture materials or drug formulations, or when manipulating potentially infectious materials. The worker is exposed to materials (including proteinaceous antigens) being manipulated on the clean bench and can experience hypersensitivity reactions. Horizontal clean air benches should never be used as a substitute for a biological safety cabinet in research, biomedical or veterinary laboratories or as a substitute for a chemical hood.

VI.3.1.5. Vertical Laminar Flow “Clean Benches”

Vertical Laminar Flow “Clean Benches” are also not BSCs. They may be useful in hospital pharmacies when a clean area is needed for preparation of intravenous drugs. While these units usually have a sash, the air is discharged into the room under the sash, resulting in the same potential problems as the horizontal laminar flow clean benches.

VI.3.1.6 Biological Safety Cabinets vs. Chemical Fume Hoods

Biological Safety Cabinets (BSCs) and Chemical Fume Hoods (CFHs) are not interchangeable. BSCs and CFHs are different equipment designed for different applications. BSCs are for working with biological materials and CFHs are for use when working with hazardous chemicals.

There are two main differences between CFHs and BSCs. CFHs have inward airflow, offering personnel protection only. BSCs have both inward and downward airflow, allowing for both personnel and product protection (clean work environment). The second difference is a CFH has no HEPA filtration offering no environmental protection; a BSC has HEPA filters on both the supply and the exhaust which provides both product and environmental protection.

It is important to understand the differences between these two types of equipment. If you have questions as to which type of equipment you should be using for your research, please contact Environmental Health and Safety for assistance.
VI.3.2. Use of Biological Safety Cabinets

Biological safety cabinets with the potential to be used to protect workers from hazardous biological agents shall be tested and certified after installation and before use, any time they are moved, when major repairs are performed and at least annually. According to NSF/ANSI Standard 49, prior to repair or replacement of components located in contaminated plenums, prior to relocation and in some cases prior to recertification, BSCs must be gas decontaminated by a qualified contractor. The PI shall provide annual certification records for each biosafety cabinet under that individual’s control. Testing shall meet the criteria in NSF/ANSI Standard 49 - 2012 Biosafety Cabinetry: Design, Construction, Performance, and Field Certification, Annex F. Call EHS for information on the standard and a list of companies qualified to certify biological safety cabinets.

- A BSC is required in Biosafety Level 2 laboratories whenever a laboratory procedure results in the formation of an aerosol (below are a list of activities that are prone to aerosol formation).
  - Centrifugation
  - Vigorous shaking and mixing
  - Pipetting
  - Grinding
  - Aspiration/washing
  - Injection
  - Sonication
  - Working with materials under pressure

A BSC is required for all pathogen manipulations performed in a Biosafety Level 3 laboratory.

Biological safety cabinets are only effective when personnel operate them properly:

- Understand the function and use of the biological safety cabinet before beginning work.
- Demonstrate proficiency in working in the BSC.
- No modifications may be made to any BSC without first contacting the IBO.

Open flames are not permitted to be used in BSCs. The flame creates turbulence which disrupts the pattern of HEPA-filtered air being supplied to the work surface. In addition, the heat from the continuous flame may damage the supply and/or exhaust HEPA filters, requiring replacement of the filters.

Biosafety cabinets are designed for a single operator. Never work with two or more people at a time in any BSC, regardless of manufacturer, model, or size. Multiple users will cause air disruptions and potentially destroy the containment capabilities of the BSC, possibly creating personnel, product, or environmental protection issues.
Any procedure specific exemption or waiver from this policy must be submitted to the Institutional Biosafety Officer and/or the Institutional Biosafety Committee Chair for review and approval prior to commencement.

A thorough evaluation of the proposed work (including the biological and chemical agents to be used and the procedures to be performed) must be executed before selecting the appropriate biological safety cabinet. Contact the Institutional Biosafety Officer for assistance when selecting a new biosafety cabinet.

Biosafety cabinets should be located away from:

- Laboratory equipment that creates air movement
- Portable fans
- Air supply registers
- Chemical fume hoods
- Open windows
- Foot traffic

Contact the Institutional Biosafety Officer for assistance when selecting a location for a new biosafety cabinet.

If a biosafety cabinet is posted for radioactive sample work, it must be cleared by EHS prior to its annual recertification. Please reach out to your EHS representative to obtain a maintenance letter prior to recertifying any biosafety cabinet that is used for radioactive materials.

Additional information on BSCs can be found in the “Biological Safety Cabinets” online training available on the EHS training website.

VI.3.3. Other Safety Equipment

Leak proof containers for the processing, transporting or storage of etiologic agents are also safety equipment. An example of a leak proof container is the safety centrifuge cup/rotor that is designed to prevent the release of aerosols during centrifugation.

A secondary container is required when transporting etiological agents between rooms, floors, or buildings. These secondary containers should be leak-proof, durable containers that must be marked with the universal biohazard symbol and contain enough absorbent material to contain the liquid samples if they were to leak during transit.
A secondary container is also required when transporting ABSL2 animals between room, floors and buildings. Secondary containment is necessary to contain cage contents and prevent a release of potentially infectious material if a cage is upset or toppled while in transit. Secondary containment options could include biohazard bags, Rubbermaid containers on carts or enclosed cabinets on wheels. It is also important to remember not to overload a cart with too many cages even if the cages are in secondary containment.

Personal protective equipment (PPE) (e.g., gloves, coats, gowns, shoe covers, boots, respirators, face shields, and safety glasses or goggles) is clothing and equipment generally used in combination with BSCs and other devices to contain the agents, animals, or materials during manipulation. PPE is covered in more detail in Chapter VII of this manual.

In situations where it is impractical to work in BSCs, personal protective devices may form the primary barrier between personnel and the infectious materials. Examples of such situations include certain animal studies, animal necropsy, and activities relating to maintenance, service, or support of the laboratory facility.

Appropriate safety equipment must be considered when performing a risk assessment for a particular project. The Institutional Biosafety Officer, EHS, and/or the IBC must be consulted when additional containment devices are determined to be necessary.

**WARNING:** Chemical fume hoods and laminar-flow clean-air benches (both vertical and horizontal) are *not* to be used for work with biohazard materials.

### VI.4. Facility Design

Secondary barriers not only protect the environment within the facility, but also outside the laboratory (and the community outside the facility) from exposure to infectious materials. The design of the facility provides the secondary barrier. The three facility designs are the basic laboratory, the containment laboratory, and the maximum containment laboratory. Laboratories are typically visited by EHS staff on an annual basis and should be found to be in compliance with the appropriate biosafety-level containment for the biohazards in use as defined by the NIH/CDC and Ohio State guidelines.

In addition to IBC approval, further approvals are required to work with agents classified as RG3 prior to initiation of work.
VI.4.1. The Basic Laboratory

The Basic Laboratory provides general space where work is done with viable agents that are not associated with disease in healthy adults; it may include Biosafety Levels 1 and 2 facilities. This laboratory is also appropriate for work with infectious agents or potentially infectious materials when the hazard levels are low and laboratory personnel can be adequately protected by standard laboratory practice. While work is commonly conducted on the open bench, certain operations are confined to BSCs (especially those that produce aerosols). Conventional laboratory designs are adequate.

VI.4.2. The Containment Laboratory

The Containment Laboratory has specialized engineering features that enable laboratory workers to handle hazardous materials without endangering themselves, the community, or the environment. The containment laboratory is described as a Biosafety Level 3 facility. The features that distinguish this laboratory from the basic laboratory are the provisions for access control and a specialized ventilation system. In all cases, a controlled access zone separates the laboratory from areas open to the public.

VI.4.3. The Maximum Containment Laboratory

The Maximum Containment Laboratory has special engineering and containment features that allow laboratory workers to safely conduct activities involving infectious agents that are extremely hazardous to humans or capable of causing serious epidemic disease. The maximum containment laboratory is described as a Biosafety Level 4 facility. Containment requirements at this level will not be approved.

VI.5. Recombinant and Synthetic Nucleic Acid Biosafety Levels

Laboratory-scale recombinant and synthetic nucleic acid research and development (i.e., <10 liters) must be carried out at the biosafety level determined to be appropriate by review of the NIH Guidelines for Research Involving Recombinant or Synthetic Nucleic Acid Molecules. Although some experiments are found to be exempt from IBC review under the NIH Guidelines, the containment necessary for performing these experiments is dependent upon the biosafety level assigned to the host/vector system.

Large-scale recombinant and synthetic nucleic acid production (≥ 10 liters) must be approved by the IBC. The appropriate level of containment will be determined by the IBC at the time of review of the protocol.
Section VII
Personal Protective Equipment

VII.1. Regulations

Working safely in a laboratory requires having the proper containment equipment and engineering controls, wearing appropriate Personal Protective Equipment (PPE), using proper work practices, knowing safety information for the materials and equipment used, and following safety instructions and laboratory protocols. Some labs contain more than one type or category of hazardous material. The protective equipment and work practices in such a lab are those that provide protection against the most hazardous agent.

The appropriate use of personal protective equipment (equipment for eyes, face, head, and extremities, protective clothing, respiratory devices, and protective shields and barriers) minimizes the potential for exposure to biohazard, toxic, and corrosive agents.

The Ohio Public Employment Risk Reduction Program has adopted the Occupational Safety and Health Administration standard (29 CFR 1910.132) covering the availability and use of PPE. According to the standard, the PI shall assess the workplace to determine if hazards are present, or are likely to be present, that necessitate the use of PPE. If such hazards are present, or are likely to be present, the PI shall:

- Offer the employee with the potential to exposure, types of PPE that will protect them from the hazards identified in the hazard assessment.
- Communicate the selection decisions to each employee.
- Select PPE that properly fits each employee.
- Educate the employee on proper use of PPE.

The PI is responsible for ensuring that each person using PPE knows when PPE is necessary, what PPE is necessary, how to properly don, doff, adjust and wear the PPE, what the limitations of the PPE are, and the proper care, maintenance, useful life and disposal of the PPE. The PI shall verify that each affected person has received and understood the required training through a written certification that contains the person’s name, the date(s) of the training, and that identifies the subject of the certification.

VII.2. General Comments

Some protection is provided by ordinary clothing and glasses. However, one must dress sensibly for laboratory work. Laboratory-provided clothing protects the clothing underneath. It is the responsibility of the lab worker to use special protective clothing and equipment when they are required for safety. Protective wear may include laboratory coats, wrap around gowns, masks, coveralls, aprons, gloves, shoe covers, eye protection, and respirators. It is necessary to select the
garments and fabric used based upon the nature of the hazardous agent. The PI must provide or ensure provision of appropriate PPE to each employee who is subject to occupational exposure to human blood or other potentially biohazard material. The PPE is provided at no cost to the employee.

The PI must either directly or through delegation ensure that each employee uses PPE when warranted. Aprons, laboratory coats, gloves, and other protective clothing, preferably made of chemically inert material, shall be readily available. Laboratory coats are essential to protect street clothing from biological aerosols or chemical splashes and spills, vapors, or dusts. PPE shall be provided in a sanitary and reliable condition and shall be cleaned regularly to avoid spreading contamination.

Protective equipment in appropriate sizes must be available in the work area or issued to employees. Hypoallergenic gloves or similar alternatives must be readily available to those allergic to the latex or vinyl gloves normally provided. Additionally, the type of glove used must be compatible with the usage: some gloves are permeable to certain compounds. Check the Safety Data Sheet for incompatibility.

PPE must be replaced if damaged. Staff should not attempt to repair PPE under any circumstances.

Eyes are very vascular and can quickly absorb many chemicals. Regulations require the use of protective eye and face equipment where there is a reasonable probability that their use can prevent injury.

Safety glasses with clear side shields are adequate protection for general lab use. Goggles shall be worn when there is danger of splashing chemicals or biologicals or flying particles (such as when chemicals are poured, or glassware is used under elevated or reduced pressure). A face shield (or face shield with goggles) offers maximum protection (for example, with macaque monkeys, with a vacuum system that may implode or when emptying the liquid from vacuum traps).

Corrective lenses in spectacles do not in themselves provide sufficient protection for working in the lab. Regulations require that persons whose vision requires corrective lenses, and who are required to wear eye protection, shall wear goggles over their eyeglasses, prescription safety glasses, or goggles with prescription lenses. Persons who wear contact lenses in laboratories must also wear appropriate eye protection.

Unprotected skin should be protected whenever possible. Suitable clothing shall be worn in the laboratory. Street clothing may absorb liquid spills that might otherwise contact skin. Shorts and other garments that do not provide sufficient skin protection are prohibited in the laboratory. Long sleeves protect arms; long sleeves shall fit snugly when working around moving machinery. Wool affords more protection from flash burns or corrosives than cotton or synthetic fabrics. Some synthetic fabrics may increase the severity of injury in the case of fire. Cotton is less prone to static electricity build up than nylon or other synthetics.
The wearing of substantial leather shoes in the lab protects against chemical splashes or broken glass. The wearing of sandals or other open-toed footwear is prohibited. Cleaning up spills on floors may require extra protection such as rubber boots or plastic shoe covers. Steel-toed shoes should be used when handling heavy items such as gas cylinders or heavy equipment components. Gloves must be worn when it is reasonably anticipated that hand contact with blood or other potentially infectious materials, mucous membranes, or non-intact skin might occur as well as when employees perform vascular-access procedures and handle or touch contaminated items or surfaces.

Disposable gloves must be replaced as soon as practical when contaminated or when torn, punctured, or otherwise compromised in their ability to function as a barrier. They must not be reused.

Utility gloves (non-disposable gloves) may be decontaminated for reuse provided the integrity of the glove is not compromised. They must be discarded if cracked, peeling, torn or punctured or exhibit other signs of deterioration.

Gloves must be removed, and hands washed when exposure is no longer anticipated and prior to leaving the work area.

For certain protocols and projects, additional PPE such as respiratory protection may be required. Respirator selection is based on the hazard and protection factor required. Personnel who require respiratory protection must contact Occupational Health and Wellness (Employee Health) for medical evaluation and clearance, and EHS for fit testing and training, prior to using a respirator.

Personal hygiene is extremely important to individuals working in a lab. Contamination of food, beverages, or smoking materials is a potential route of exposure to toxic chemicals or biological agents through ingestion. Laboratory personnel shall not prepare, store, or consume food or beverages, pipette by mouth, smoke, apply cosmetics, or handle contact lenses in the lab.

Hand washing is a primary safeguard against inadvertent exposure to toxic chemicals or biological agents. Individuals should always wash their hands before leaving the lab, even if using gloves. Wash hands after removing protective clothing, before leaving the lab, and before eating, drinking, smoking, or using the restroom. Individuals should wash their hands periodically during the day at intervals dictated by the nature of the work being completed. Wash with soap and running water, with hands held downwards to flush the contaminants off the hands. Turn the tap off with a clean paper towel to prevent recontamination and dry hands with clean towels.

Confine long hair and loose clothing when in the lab to keep them from catching fire, dipping into hazards, or becoming entangled in moving machinery. Avoid the wearing of finger rings and wristwatches that may become contaminated, react with chemicals, puncture, or tear gloves, or be caught in moving parts or equipment.

Remove laboratory coats and gloves before leaving the lab and entering public spaces (i.e., elevators and restrooms) to prevent spreading contamination to other areas.
Section VIII
Biosafety Laboratory Practices and Equipment

Guidelines for Good Practices at all biosafety levels can be found in the Biosafety in Microbiological and Biomedical Laboratories 6th edition. Ohio State does not conduct research at BSL4.

All laboratory personnel shall engage in good microbiological laboratory practices at all times. The following practices incorporate minimal practices and provide guidance for ensuring the protection of personnel, research, and the environment for the level of containment used in academic and research laboratories.

Hands should be washed frequently during the day. Wash hands after removing gloves, before leaving the laboratory, before and after contact with patients or animals, and before eating, smoking, handling contacts or applying cosmetics.

Hands must also be washed immediately after accidental contact with blood, body fluids, and contaminated materials. Refrigerators, freezers, water baths, and centrifuges should be cleaned and disinfected periodically (the frequency to be established by the PI/laboratory director) and when gross contamination occurs. Wear gloves, gown, and appropriate PPE during cleaning.

Exits and aisles must not be obstructed in any way. No trash, supplies, equipment, or furniture should be permitted in exit routes or aisles.

Exit doors must not be obstructed, bolted, or blocked in any way. Smoke doors must not be obstructed in any way that prevents automatic closing in case of fire.

Do not cover or block access to fire extinguishers, fire alarm boxes, electric panels, emergency blankets, safety showers, eyewashes or exits at any time, for any reason.

All hazardous materials, including biological, chemical and radioactive materials, should be secured when unattended, to protect from unauthorized access, misuse or removal.

VIII.1. Biosafety Level 1 (BSL-1)

Biosafety Level 1 is suitable for work involving well-characterized agents not known to consistently cause disease in healthy adult humans, and of minimal potential hazard to laboratory personnel and the environment. The laboratory is not necessarily separated from the general traffic patterns in the building. Work is generally conducted on open bench tops using standard microbiological practices. Special containment equipment or facility design is neither required nor generally used. Laboratory personnel have specific training in the procedures conducted in the laboratory and are supervised by a scientist with general training in microbiology or a related science.
VIII.2. **Biosafety Level 2 (BSL-2)**

**Biosafety Level 2** is similar to Biosafety Level 1 and is suitable for work involving agents of moderate potential hazard to personnel and the environment. It differs from BSL-1 in that (1) laboratory personnel have specific training in handling pathogenic agents and are directed by competent scientists; (2) access to the laboratory is limited when work is being conducted; (3) extreme precautions are taken with contaminated sharp items; and (4) certain procedures in which infectious aerosols or splashes may be created are conducted in biological safety cabinets or other physical containment equipment.

VIII.3. **Work with Human or Animal Tissues**

Human blood, blood products, cells and cell lines, body fluids and tissues are listed as potentially hazardous biological materials. Animal tissues may also be contaminated with biohazardous materials. Biosafety Level 2 practices and procedures must be followed when handling human blood, blood products, cells and cell lines (including established cell lines), body fluids and tissues and animal tissues because of the infectious agents that they may contain. Biosafety Level 2 practices and procedures are consistent with the concept of “Universal Precautions” that require all specimens of human blood, blood products, body fluids and tissues to be treated as if they are infectious. The federal regulation, the Occupational Exposure to Bloodborne Pathogens (29 CFR 1910.1030), adopted by the Ohio Public Employment Risk Reduction Program, mandates a series of engineering and work practice controls, training, and Hepatitis B vaccination to control the health risk to employees resulting from occupational exposure to human blood and other potentially infectious materials that may contain human pathogens.

VIII.4. **Biosafety Level 3 (BSL-3)**

Specific BSL-3 practices and procedures are described in the Ohio State BSL-3 facility biosafety manuals.
Section IX
Decontamination and Spills

IX.1. Definitions

- **Sterilization:** the act or process, physical or chemical, which destroys or eliminates all forms of life, especially microorganisms.

- **Decontamination:** reduction of all organisms and the destruction of pathogenic organisms in or on a material so that material is no longer considered to be capable of transmitting disease.

- **Disinfection:** the act of destroying or irreversibly inactivating specific viruses, bacteria, or pathogenic fungi, but not necessarily their spores, on inanimate surfaces. Most disinfectants are not effective sterilizers.
  - High Level Disinfectants kill all viruses and vegetative cells, but they may not kill endospores reliably.
  - Intermediate Level Disinfectants destroy all vegetative cells including Mycobacteria, fungi and most, but not all viruses. They cannot kill endospores.
  - Low Level (General Purpose) Disinfectants destroy vegetative bacteria, except Mycobacteria, fungi, and non-enveloped viruses.

- **Antiseptic:** a substance that prevents or arrests the growth or action of microorganisms either by inhibiting their activity or by destroying them. The term is used especially for preparations applied to living tissue.

IX.2. Evaluation

The initial risk assessment for any project should include an evaluation of the processes and/or agents to be used to ensure that the biohazardous materials involved in the research are inactivated during spill cleanup, before cleaning equipment for re-use, and for final disposal.

The OSHA Bloodborne Pathogens Standard requires that all equipment and environmental and working surfaces shall be cleaned and decontaminated after contact with blood or other potentially infectious materials. The standard also requires decontamination of contaminated work surfaces after completion of procedures, immediately or as soon as feasible after any overt contamination of surfaces or any spill of potentially infectious material, and at the end of the work shift if the work surface has become contaminated. All reusable equipment shall be decontaminated immediately or as soon as feasible after visible contamination.
For any infectious material adequate pre-cleaning of surfaces is important for any disinfection or sterilization procedure. Ten minutes of exposure to a disinfectant is not adequate to disinfect objects that have narrow channels or other areas that can harbor microorganisms. Alcohols, 70%, for example, are effective for killing HBV but are not recommended for this purpose because of their rapid evaporation and the consequent difficulty of maintaining proper contact times. Alcohols have been removed from many laboratories because they are flammable. Alcohols should be maintained only in small volumes and may be desirable as an adjunct to skin disinfection.

Chlorine compounds are widely used disinfectants in the laboratory. An inexpensive, broad-spectrum disinfectant for use on bench tops and similar surfaces can be prepared by diluting common household bleach (5.25 % sodium hypochlorite solution [some cut-rate brands might not contain this much hypochlorite]) to obtain at least 500 ppm of free available chlorine. (Some bleach solutions available now contain about 1/3 more hypochlorite than the solutions mentioned above. Check to see what the concentration of chlorine in the bleach solution you are using.) A 1:10 dilution of commercial bleach (10%) that produces a solution containing 5000 ppm of free chlorine can be used to disinfect spills. The use of higher concentrations of bleach in chemical fume hoods should be reserved for significant contamination.

High concentrations of bleach solutions should not go into an autoclave.

Prepare a fresh solution of bleach every two weeks; discard unused portions down the sink drain and then flush with fresh water. Be aware that chlorine compounds may corrode metals, especially aluminum. To help prevent corrosion after using the bleach solution, rewipe the surfaces with 70% ethanol.

Chlorine dioxide, either as a liquid solution or as a gas, can also be used for decontamination purposes. The liquid form (i.e. Clidox) is most commonly used as a 1:5:1 or 1:18:1 dilution (base:water:activator). Chlorine dioxide gas can be used for large equipment or space decontaminations.

Iodophors that are registered with the EPA may be effective hard-surface decontaminants when used per the manufacturer’s instructions, but iodophors formulated as antiseptics are not suitable for use as disinfectants (i.e., Wescodyne).

Phenolics that are registered with the EPA may be effective hard-surface decontaminants when used per the manufacturer’s instructions (i.e.,VESPHENE, Hil-Phene).

Quaternary ammonium compounds are low-level disinfectants and are not recommended for spills of human blood, blood products, and other potentially infectious materials (i.e., Conflikt, End-BacII).
The use of such chemicals requires that the laboratory have a current Chemical Hygiene Plan (29 CFR 1910.1450, Occupational Exposure to Hazardous Chemicals in Laboratories). Safety Data Sheets (SDSs) for the chemicals in use must be made available to the individuals in the lab, as well as training on special procedures for handling the chemicals.

IX.3. Sterilization

Unless the facility is permitted by the Ohio EPA to treat infectious waste, all terminal treatment is incineration. Consequently, all pre-treated and untreated biohazards (i.e., infectious waste) shall be placed in a burn box. See Appendix C. Consult EHS for assistance.

According to the OEPA, for terminal sterilization to be allowed, the sterilization process (steam autoclaving, dry heat, etc.) must be validated, and the validation documented. Liquid "cold" sterilants may be used to sterilize equipment that will not withstand the heat of steam or the chemical reactivity of ethylene oxide processing.

Additionally, the sterilization process must also be monitored at least weekly (or a quality-control run completed if the autoclave is used less often than weekly) with biological indicators (spore strips, time/temperature charts, etc.), and records of monitoring kept for review.

Since Ohio State has no autoclave licensed for terminal sterilization by the OEPA, individuals using autoclaves must still have their autoclaved waste prepared and sent off-site for incineration per the Ohio State contract. If an autoclave is being used for infectious waste pre-treatment, periodic monitoring of the effectiveness of the sterilization process is still recommended.

IX.3.1. Steam Sterilization

Steam sterilization (autoclaving) is the primary means of sterilization. The following points must be kept in mind when steam sterilization is to be used:

- Materials affected (e.g., denatured or melted) by heat will be destroyed by this method of sterilization.

- Steam must reach the material for a prescribed period of time (adequate sterilization time) to ensure sterilization. Containers must be open to allow steam penetration, or water must be placed in the container before placing in the sterilizer.

- Use extreme caution when opening the autoclave following the sterilization cycle. Steam can cause serious injury. Additionally, malfunctioning autoclaves can fill with superheated water that will be released when the autoclave is opened.
- **Suggested sterilization cycle times:**
  - 60 minutes @ 121°C & 15 PSI for decontaminating waste
  - Lengthen time for large or dense loads
  - 30 minutes @ 121°C & 15 PSI for sterilizing clean materials (i.e., glassware)
  - Use slow exhaust for liquids and fast exhaust for glassware

Additional information can be found in the “Autoclave Safety” training, available on the EHS website.

**IX.4. Disinfection**

An integral part of the biosafety program is the identification of appropriate disinfectants or decontaminating agents. Such materials are to be kept readily available in the use-dilution required. The disinfectant and the disinfection process must be validated, and the validation documented. Personnel must be trained in the appropriate use of the approved disinfectant. EHS personnel can assist in the development of an appropriate validation and monitoring process.

Disinfectants must always be used in accordance with the manufacturer’s recommendations. Failure to follow the manufacturer’s recommendations can result in the failure of the disinfectant to perform as expected.

**Disinfection Hazards**

Disinfectants are potentially hazardous chemicals and should be handled with care. Check the manufacturer’s Safety Data Sheet (SDS) before use.

Personnel should be informed of the hazards associated with disinfectant use and provided with appropriate PPE to minimize exposure under use conditions.

Appropriate disposal requirements must be specified for each disinfectant used.

**IX.5. Spills and Spill Cleanup**

Spills of biohazardous materials may constitute a significant health hazard if not handled appropriately. All personnel working with biohazardous materials must be trained in the specific cleanup and disinfectant procedures to be used for their particular laboratory. Personnel must also be informed of the handling and disposal of contaminated clothing and personal protective devices. All of this information should be included in a Standard Operating Procedure developed by the PI.

A biological spill shall be followed by prompt action to contain and clean up the spill. When a spill occurs, warn everyone in the area and call for assistance as needed. The degree of the risk involved in a spill depends on the volume of the material spilled, the creation of infectious aerosols, the concentration of organisms in the material spilled, the hazard of the organisms involved, the route of infection of the organisms, and the diseases caused by the organisms.
Spills of biological agents can contaminate areas and lead to infection of laboratory workers. Prevention of exposure is the primary goal in spill containment and cleanup, exactly as in chemical spills. In evaluating the risks of spill response, generation of aerosols and droplets is a major consideration.

**IX.5.1. Generic Spill Cleanup Plans**

As part of the laboratory Safety Plan, each laboratory must have a biological spill kit and a spill cleanup plan detailing specific disinfectants and procedures for agents used in that laboratory. The biological spill kit should contain supplies to clean up any spill of biological origin, including plant, animal or human material and recombinant or synthetic nucleic acids, both infectious and non-infectious. Cleanup of any spill requires the use of appropriate personal protective equipment (i.e., laboratory coat, shoe covers, gloves, and possible respiratory protection). To comply with OEPA regulations, all spills of infectious materials greater than one gallon or one cubic foot must be reported to EHS. The following procedures should serve as a guide for the development of specific procedures for the laboratory Safety Plan.

A copy of the EHS “Infectious Waste Spill Containment & Clean Up Procedure” can be obtained by contacting your EHS Safety Representative. This procedure should be posted in all labs where working with or storing potentially infectious materials, including recombinant or synthetic nucleic acids.

**IX.5.1.1. Spill Cleanup Procedures**

The following procedures are to assist lab personnel with containment and cleanup of spills under specific circumstances.

**Spill Contained Within a Biological Safety Cabinet (BSC)**

- BSC must run during cleanup to contain aerosols & HEPA-filter exhaust air.
- Don appropriate personal protective gear before initiating cleanup.
- Initiate clean up as soon as possible using a 10% bleach solution or other EPA approved tuberculocidal disinfectant.
- If the spill is contained on a bench diaper, remove the contaminated bench diaper & discard as infectious waste.
- If the spill is on the work area surface, cover spilled material with disinfectant-soaked towels. Allow the appropriate contact time (30 minutes for bleach; other EPA approved tuberculocidal disinfectants follow manufacturer’s recommendations) then remove the contaminated towels & discard as infectious waste.
• Wipe down the interior of the cabinet & any splatter on items within the cabinet with a disinfectant-soaked towel.

• Wipe down non-autoclavable materials with disinfectant. Allow the appropriate contact time (30 minutes for bleach; other EPA approved tuberculocidal disinfectants follow manufacturer’s recommendations) with disinfectant before any items are removed from cabinet.

• Place items designated as contaminated used sharps in an appropriate infectious waste sharps container using tongs/forceps. Place other contaminated disposable materials used in the cleanup process in a biohazard bag. Process as infectious waste.

• Place contaminated re-usable items in autoclave bags or autoclavable pans with lids. Sterilize, preferably by autoclaving, and clean for re-use.

• If the cabinet has a catch basin beneath the work surface & the spill resulted in liquids flowing into this area, more extensive decontamination is required.
  1) Ensure the drain valve under the cabinet is closed.
  2) Pour disinfectant onto the work surface & through the front and rear grilles into the drain pan. Allow 30 minutes contact time.
  3) Absorb spilled fluid-disinfectant from work surface with paper towels & discard in biohazard bag.
  4) Prepare to empty drain pan. Place disinfectant solution in a collection vessel. Attach flexible tubing to the drain valve. The tube should be of sufficient length to allow the open end to be submerged in the collection vessel to minimize aerosol generation.
  5) Open the drain valve & empty the drain pan into the collection vessel containing disinfectant. Flush the drain pan with water & remove the flexible tubing. Manage contaminated materials as if they are infectious.
  6) Remove protective clothing used during cleanup & dispose of as infectious waste. Wash hands when gloves are removed.
  7) Notify Principal Investigator or supervisor. Consult with EHS (614-292-1284) to determine whether gas decontamination of the cabinet and filters is necessary, especially if a high-risk agent or a major spill of a moderate-risk agent occurred.
  8) Run BSC at least 10 minutes after cleanup, before resuming activity in the cabinet.

Spill Outside a Cabinet, Inside the Laboratory

• If a spill occurs in a Biosafety Level 2 facility, outside the BSC, notify other individuals in the laboratory to evacuate.

• Exit the laboratory to the hallway, closing the door behind you.
- Remove any contaminated clothing (turn contaminated portion inward) & place it in an autoclave bag.

- Wash all exposed skin.

- Place signs on door(s) to the laboratory warning individuals who may want to enter that a spill occurred & access is denied.

- Allow aerosols to settle for at least 30 minutes before re-entering the laboratory.

- Assemble supplies (disinfectant, sharps containers, towels, tongs, biohazard bags, etc.) before entering the laboratory.

- Don appropriate personal protective equipment (i.e., disposable gown, protective eyewear, gloves, shoe coverings & respiratory protection [if necessary]).

- Clean up spill with a 10% bleach solution or other EPA approved tuberculocidal disinfectant as follows:

  1) Surround spill area with disinfectant or diking material that is soaked in disinfectant.
  2) Place items designated as contaminated used sharps in an appropriate infectious waste sharps container. Place other disposable materials used in the cleanup process in a biohazard bag. Process as infectious waste.
  3) Place paper towels over the entire spill area to absorb the spill. Clean the area and dispose of the material as infectious waste.
  4) Apply disinfectant and allow the appropriate contact time (30 minutes for bleach; other EPA approved tuberculocidal disinfectants follow manufacturer’s recommendations) with the disinfectant to ensure adequate germicidal action.
  5) Wipe down non-autoclavable materials with disinfectant.
  6) Place contaminated re-usable items in autoclave bags or autoclavable pans with lids. Sterilize, preferably by autoclaving, and clean for re-use.
  7) Remove protective clothing used during cleanup then place in a biohazard bag for autoclaving.

- Wash hands when gloves are removed.

- Notify Principal Investigator or supervisor & EHS (614-292-1284)

**If spill assistance is needed, contact EHS by calling 614-292-1284 and choosing Option 1.**
Spill Inside a Centrifuge

*The potential for multiple infections from a single centrifuge accident is high. Aerosols are created when fluid escapes from the rotor or cup while the centrifuge is operating at high speed. All opening of centrifuges must be performed slowly.*

Unsealed buckets:

- If a centrifuge tube breaks while the centrifuge is running, turn off motor. Allow the machine to be at rest for 30 minutes before opening. If breakage is discovered after the machine has stopped, re-close the lid immediately & allow the unit to be at rest for 30 minutes.

- Unplug centrifuge before initiating clean up.

- Don strong, thick, rubber gloves & other PPE before proceeding with clean up.

- Flood centrifuge bowl with a 10% bleach solution or other EPA approved tuberculocidal disinfectant. Place paper towels soaked in a disinfectant over the entire spill area. Allow the appropriate contact time (30 minutes for bleach; other EPA approved tuberculocidal disinfectants follow manufacturer’s recommendations) with the disinfectant. Use mechanical means (such as forceps) to remove broken tubes & glass fragments. Place them in a sharps container for disposal as infectious waste.

- Remove buckets, trunnions & rotor then place in disinfectant for 30 minutes or autoclave.

- Unbroken, capped tubes may be placed in disinfectant & recovered after appropriate contact time.

- Use mechanical means to remove remaining disinfectant soaked materials from centrifuge bowl & discard as infectious waste.

- Place paper towels soaked in a disinfectant in the centrifuge bowl & allow it to soak for 30 minutes, wipe down again with disinfectant, wash with water & dry. Discard disinfectant soaked materials as infectious waste.

- Remove protective clothing used during cleanup & place in a biohazard bag for autoclaving. Wash hands whenever gloves are removed.

Sealed buckets (safety cups):

- Remove the sealed bucket to a biological safety cabinet before opening.

- If breakage occurred, replace the cap on the safety cup loosely and autoclave.
Notify Principal Investigator or supervisor & EHS (614-292-1284).

Spill Outside the Laboratory; during Transport (on Ohio State Campus)

_The major emphasis should be on preventing spills during transport. All transport of infectious materials must be in a rigid, securely sealed, watertight primary container, which is contained within a second rigid, leak proof sealed container. Sufficient absorbent should be added to the second container to absorb contents in case of leakage from the primary container. The outer container must be labeled with the universal biohazard symbol._

If a spill occurs during transport, don gloves and initiate cleanup immediately with a 10% bleach solution or other EPA approved tuberculocidal disinfectant as follows:

- Surround spill area with disinfectant or diking material that is soaked in disinfectant.
- Place contaminated used sharps in an appropriate infectious waste sharps container.
- Place paper towels over the entire spill area to absorb the spill. Clean the area and dispose of the material as infectious waste.
- Apply disinfectant and allow the appropriate contact time (30 minutes for bleach; other EPA approved tuberculocidal disinfectants follow manufacturer’s recommendations) with the disinfectant to ensure adequate germicidal action.
- Place all materials used in the cleanup process (including contaminated gloves) in a biohazard bag and process as infectious waste.
- Wash hands as soon as possible.

**IX.5.1.2. Biological Spill on a Person**

If a biological material is spilled onto a person, emergency response is based on the hazard of the biological agent spilled (including the ability of the organism to penetrate intact skin), the amount of material spilled, and whether significant aerosols were generated. If aerosol formation is believed to have been associated with the spill, a contaminated person should leave the contaminated area immediately. If possible, he or she should go to another laboratory so that hallways and other public areas do not become contaminated.

Contaminated clothing is removed and segregated as biohazard laundry for disinfecting. Contaminated skin shall be thoroughly flushed with water and washed with a disinfectant soap. Showering may be appropriate, depending on the extent of the spill. For Risk Group 2 and Risk Group 3 pathogens, the employee must report to Occupational Health and Wellness (Employee Health) immediately for evaluation.
Section X
Biological Waste Disposal

Laboratory waste may be potentially hazardous (infectious, radioactive, or toxic chemicals) and must be handled appropriately to prevent possible harm to personnel and/or environmental contamination. Certain wastes are regulated and must be handled according to approved methods.

All applicable rules and regulations of local, state, and federal agencies are to be followed in the handling, treatment, and disposal of biomedical waste.

All biohazard waste must be packaged, contained, and stored in a manner that protects and prevents the waste from release at any time. If storage is necessary, putrefaction and the release of infectious agents into the air must be prevented.

X.1. Responsibility

It is the responsibility of the PI/laboratory supervisor to identify the classes of wastes that are generated in the laboratory and to ensure that the appropriate methods of waste disposal are followed. Information regarding infectious waste disposal is covered under the Principal Investigator Assurances section of the IBC e-Protocol submission.

It is the responsibility of each laboratory employee to ensure that he/she follows the proper method of waste disposal.

X.2. Requirements

Waste must be segregated on the basis of potential hazard.

All infectious waste will be handled, and disposed of in accordance with Appendix C, Infectious Waste Guidelines.
Section XI
Ordering, Receiving, Shipping and Movement of Biohazard Materials

The transport of biohazard material is regulated by a number of government agencies. It is imperative that personnel are aware of the applicable regulations and comply with them. The shipper of biohazard material is responsible for the proper classification, identification, packaging, labeling, and documentation of the shipped material. Failure to comply could result in fines, the confiscation and destruction of the material and loss of valuable research time.

XI.1. Applicable Regulations

The following is a list of regulations that control the shipping of hazardous materials:

1. U. S. Department of Health and Human Services (HHS) and United States Department of Agriculture (USDA): 42 CFR 73, 7 CFR 331, and 9 CFR 121 for select agents or toxins; 42 CFR 73 is the implementation of the Public Health Security and Bioterrorism Preparedness and Response Act of 2002; 7 CFR 331 and 9 CFR 121 are implementations of the Agricultural Bioterrorism Protection Act of 2002 (Part of Title II of the PHSBPRA). The select agent list may be found in Appendix B of this manual.


6. USPHS 42 CFR - Part 71 Foreign Quarantine. Part 71.54 Etiologic agents, hosts, and vectors

XI.2 Shipments within the United States

The United States Department of Transportation (DOT) has issued regulations covering the intrastate, interstate, and foreign shipment of hazardous materials.

Shippers of dangerous goods must be trained (49 CFR 172.700 et seq.) on the DOT regulations. Training must be documented and is required every three years. Individuals must comply with shipping regulations and certify that the shipped materials will arrive at their destination in good
condition and will not present any hazards to humans or animals during the shipment. Commercial carriers will refuse to accept any packages that do not meet the regulations. Substantial fines, for both the individual and Ohio State, may be incurred if an individual is not compliant.

Shipment or transfer of exempt amounts of select agent toxins (see Appendix B) falls under the DOT regulations for hazardous materials. If shipping or transferring a non-exempt amount of select agent toxin, contact your Select Agent Program’s Responsible Official.

XI.3 International Shipments

If hazardous or infectious materials are being shipped internationally, the shipper must complete training on the Dangerous Goods Regulations, International Air Transport Association, and (IATA). This training must be completed every two years, or whenever the regulations change, whichever comes first.

The U.S. Government actively regulates the use, import, export, and interstate transport of many microorganisms, toxins, vectors and other infectious substances and biological materials. In many cases a permit or license from the Department of Commerce, the Centers for Disease Control and Prevention (CDC) and/or the Animal and Plant Health Inspection Service (APHIS) will be required. For specific information detailing current permit and licensing requirements by type of material see the “Import, Export and Transfer of Biological Materials Guide” on the EHS Biosafety webpage.

Material containing etiologic agents being imported into the United States must be accompanied by a U.S. Public Health Service importation permit. Importation permits are issued only to the importer, who must be located in the United States. The importation permit, with the proper packaging and labeling, will expedite clearance of the package through the United States Public Health Service Division of Quarantine and release by U.S. Customs.

The USDA Animal and Plant Health Inspection Service (APHIS) also regulates the importation of certain plants, animals and animal products into the United States. Please visit the APHIS website (https://www.aphis.usda.gov/aphis/ourfocus/importexport) for more information on obtaining an APHIS permit.

The importer is legally responsible for assuring that the foreign personnel package, label, and ship the infectious materials according to Federal and International regulations. Shipping labels with the universal biohazard symbol, the address of the importer, the permit number, and the expiration date, are also issued to the importer with the permit. The importer must send the labels and one or more copies of the permit to the shipper. The permit and labels inform the U.S. Customs Service and U.S. Division of Quarantine Personnel of the package contents.
XI.4 Intracampus transport of biohazardous materials

Transportation of biohazard material on campus (including between laboratories) must be completed in a manner that takes into account the potential risk of the agent being moved. The biohazard material must:

- be enclosed in a primary vessel contained within a secondary vessel.
- have a closed, leak proof secondary vessel, marked with the biohazard symbol, and marked with the name of the agent contained within the primary vessel.
- be completely absorbed by an absorbent material packed into the secondary vessel should the primary vessel be broken.
- have a secondary vessel constructed in such a manner that there will be no release into the environment of the agent in the case that the primary vessel becomes broken or leaks.

Inappropriate transport of biohazard material constitutes a violation of Ohio State’s Biohazard Policy and will be dealt with accordingly.
Section XII
Accident and Incident Reporting

Rapid and accurate reporting of accidents and incidents involving occupational exposures to biohazard material is important in identifying potentially hazardous operations and procedures. Furthermore, it allows personnel to be treated appropriately and minimizes the potential for actually contracting a disease associated with the infectious agent.

- Report all accidents involving potential exposures to biohazard material and occupational illnesses to supervisory personnel, the appropriate administrative unit (department, division, etc.), the Institutional Biosafety Officer and Occupational Health and Wellness (Employee Health) or Student Health Services. An accident report form must be completed.

- An investigation of any incident or accident may be performed in accordance with Ohio State and EHS policy (see Appendix D for more information).

- Additional information relating to biohazard/rDNA incident reporting requirements is available at: http://orrp.osu.edu/ibc/osuibcpolicies/incidentreporting/

Ohio State Recombinant DNA/Biohazard Research Incident Reporting Policy and Process

Ohio State is required to report certain incidents involving recombinant DNA or biohazard research to the National Institutes of Health. This policy outlines the information necessary to determine the nature and extent of the incident, as well as the appropriate reporting requirements and process.

Reporting Responsibilities

1. Personnel involved will immediately report the incident to the Institutional Biosafety Officer, who will contact the Senior Director of Environmental Health and Safety, the Director of ULAR, the Chair of the Institutional Biosafety Committee and the Offices of Responsible Research Practices and Research Compliance as needed.

2. The Institutional Biosafety Officer and the PI will collectively complete a rDNA / Biohazard Incident Report Form or an Animal Bite/Exposure Report Form in a timely manner as determined by the nature of the incident and agency reporting timelines. The report will be provided to the Institutional Biosafety Committee (IBC) for review.

3. Following review by the IBC and the Office of Research Compliance, the Chair of the IBC and Office of Responsible Research Practices will submit the final incident report with the Chair’s signature to the respective federal agency on behalf of Ohio State. Copies of the incident report will be provided to the Office of Legal Affairs, the Associate Dean for Research of the College involved, and the Chair of the Department involved.
Reportable Incidents and Timelines

1. The following incidents should be reported by immediately to the Principal Investigator, the Institutional Biosafety Officer, and the Chair of the Institutional Biosafety Committee:

   Spills or accidents in a BSL2 laboratory resulting in an overt exposure, injury or illness of personnel, including bites/exposures to animals intentionally infected with RG2 agents or potential zoonotic diseases.

   Spills or accidents in a BSL3 laboratory resulting in an overt potential exposure, injury or illness of personnel, including bites/exposures to animals intentionally infected with RG3 agents or potential zoonotic diseases.

   Release of a Risk Group 2 or 3 agent / genetic material from a primary containment device (e.g., biological safety cabinet, centrifuge, or primary container into the laboratory)

   Spills or accidents that lead to personal injury or illness or breach of containment (e.g., aerosols released outside of containment, skin punctures with needles containing Risk Group 2 or 3 agents or genetic material from these agents).

   Failure to adhere to the containment and biosafety practices described in the NIH Guidelines.

2. The following should be reported by Ohio State to NIH OSP:

   Section IV-B-2-b-(7) of the NIH Guidelines for Research Involving Recombinant or Synthetic DNA Molecules (NIH Guidelines) states that “...any significant problems with or violations of the NIH Guidelines and any significant research-related accidents or illnesses” must be reported to NIH OSP within 30 days.

   For work conducted in BSL2 or BSL3 laboratories, Appendix G of the NIH Guidelines specifies that “spills and accidents which result in overt exposures to organisms containing recombinant or synthetic nucleic acid molecules are immediately reported to the IBC and NIH OSP” (i.e., 1.a. and 1.b. above).
rDNA / BIOHAZARD INCIDENT REPORT FORM

1. List the name(s) of the employee(s) involved:

2. List the date, time, and location (building and room) in which the accidental exposure occurred:

3. As thoroughly as possible, describe the circumstances of the exposure incident:

4. List the biological agent / genetic material and route(s) of possible exposure (e.g., inhalation, subcutaneous, etc.):

5. What is the nature of the organism strain to which the employee has been exposed? (strain name and history, complete drug-resistance/susceptibility profile, genetic modifications, any other information that might be pertinent to treatment):

6. List steps taken to evaluate employee’s health, and action taken to prevent recurrence of a similar incident:

(Attach additional pages as necessary)

Employee Signature ____________________________

Date ____________________________

Principal Investigator Signature ____________________________

Submit to Chair of the Institutional Biosafety Committee for review via the Office of Responsible Research Practices (IBCinfo@osu.edu).
ANIMAL BITE / EXPOSURE REPORT FORM WILD CAUGHT & USDA SPECIES
(To be completed by employee and supervisor)

Employee’s Name:

Principal Investigator’s Name:

Date of incident:

Time of incident:

Location of incident:

Animal species involved:

Describe the circumstances of the exposure incident:

Possible risks of exposure:

What first aid / medical attention was given to employee following exposure:

What action has been taken to prevent recurrence of a similar incident (if any):

(Attach additional pages as necessary)

Employee Signature ____________________________

Date ____________________________

Principal Investigator Signature ____________________________

Submit to Chair of the Institutional Biosafety Committee for review via the Office of Responsible Research Practices (IBCinfo@osu.edu).
Section XIII
Occupational Health Program

Overview

All individuals (i.e., faculty, staff, students, visiting scientists and volunteers) who work in laboratories must participate in Ohio State's Occupational Health Program. This program includes identification and enrollment of personnel, hazard evaluations/ risk assessments, exposure controls, medical evaluations, and occupational health and safety training. The purpose and goal of the Occupational Health Program is to identify, evaluate, manage, and reduce potential health risks associated with work involving animals, biohazards, and hazardous chemicals.

Hazard assessments and medical surveillance are critical components of an effective occupational health program and involve the evaluation of health risks associated with an individual's occupational exposures, as well as an individual's current health status. A comprehensive occupational medicine program is provided through Occupational Health and Wellness (Employee Health), Ohio State Wexner Medical Center, Ohio State Student Health Services, or a designated contractor for regional sites. After reviewing an individual’s information, a member of the Occupational Health team determines if a medical evaluation is necessary. During the medical evaluation, employees are counseled about exposure to hazardous agents that they may encounter in the course of their employment. Sources for additional information are given to the employee at that time. In many cases, an initial evaluation and risk assessment is all that is necessary. For some individuals, a clinical examination, vaccinations, and medical monitoring may be required as well.

Enrollment

Personnel shall enroll in the Occupational Health Program by the Online Risk Assessment Tool (ORAT) at http://orrp.osu.edu/iacuc/occhealth/. If an individual experiences a change in their health status or a change in their occupational exposure, he/she must update their information by updating relevant information at http://orrp.osu.edu/iacuc/occhealth/.

Medical Disclosures

Personnel that work with biohazards and have chronic medical conditions are asked to disclose these conditions to Occupational Health and Wellness (Employee Health) (unpaid students to Student Health Services). These conditions will be evaluated as part of the health evaluation. Some medical conditions and treatments can increase the severity or risk of an adverse health outcome resulting from occupational exposures.
Research Risk Assessment

Use of hazardous agents requires an approved Biosafety Research Protocol, Chemical Hygiene Plan and/or Radioactive Materials Permit, which are reviewed and approved by the Institutional Biosafety committee (IBC), the Office of Environmental Health and Safety (EHS), and/or the Radiation Safety Committee (URSC) respectively. When completing the initial risk assessment for a biohazard research project, the PI should include an evaluation of the appropriate medical surveillance, prophylactic measures (e.g., immunizations), possible treatment options, and post exposure follow-up requirements for the biohazard agent(s). This assessment is performed by the supervisor/PI in conjunction with Occupational Health and Wellness (Employee Health). The requirements for routine medical surveillance, prophylaxis, and post-exposure treatment and follow-up for work with biohazards are dictated by the risk assessment.

Vaccinations

Hepatitis B virus (HBV) vaccine is available free of charge to all employees who could reasonably anticipate exposures to human blood or other potentially infectious materials (e.g., human tissues, human cell lines, blood products, etc.) while performing their job duties. Employees with a potential for exposure to human blood or potentially infectious materials must be offered HBV immunization (cf., OSHA Bloodborne Pathogen Standard, 29 CFR 1910.1030).

When vaccinations are deemed necessary in the biohazard risk assessment, personnel will be offered the vaccine. In some cases, personnel can decline but must sign a declination form. Personnel should consult with Occupational Health and Wellness (Employee Health) for specific information.

Routes of Exposure

Exposures can occur in research personnel via:

- Injections, including, but not limited to cuts, abrasions, puncture wounds or via contamination of an existing skin injury.
- Absorption through skin (failure to wear PPE/ properly use PPE), as well as splashes to mucous membranes (e.g., eyes, nose, mouth).
- Ingestion resulting from improper lab practices (e.g., eating/drinking in lab, failure to wash hands prior to exiting lab).
- Inhalation.
Reporting Exposures, Illnesses, or Injuries

Personnel experiencing any injury or illnesses related to occupational exposures must report the event to their supervisor or PI and Occupational Health and Wellness (Employee Health) (614-293-8146), as well as submit an Employee Accident Report. Concerns or symptoms of allergies to lab animals should be reported as soon as they are noted. Known exposures to infectious agents or other biologically hazardous material (e.g., recombinant or synthetic nucleic acid molecules) must be reported to PI/supervisor; Occupational Health and Wellness (Employee Health); and to the Institutional Biosafety Officer (Environmental Health & Safety at 614-292-1284). If medical treatment is needed, personnel should go to Occupational Health and Wellness (Employee Health) or Student Health Services. Student employees shall go to Occupational Health and Wellness (Employee Health Services) and non-paid students to Student Health Services. Occupational Health and Wellness is located on the 2nd floor, McCampbell Hall at 1581 Dodd Dr. Student Health Services (614-292-4321) is located at 1875 Millikin Road. If medical treatment is needed after hours, personnel should report to the Wexner Medical Center Emergency Department. Personnel working at regional campuses shall go to the nearest Emergency Department for medical treatment. If personnel seek medical treatment in an Emergency Department, they must have an evaluation and complete an Employee Accident Report at Occupational Health and Wellness, Student Health Services or the regional site’s designated contractor prior to returning to work.

Reproductive Hazards

Persons capable of and considering reproduction should consider the ramifications of working with chemical, biological, or radiological agents, or animals. Information that should be reviewed while considering whether precautions will be necessary include Safety Data Sheets (SDS), the laboratory’s Chemical Hygiene Plan and/or the biohazard risk assessment. Individuals who are concerned about potential reproductive hazards in the workplace may contact the Office of Environmental Health and Safety (614-292-1284) with questions on locating relevant safety information.
Section XIV
Animal Research Safety

XIV.1 General

A laboratory animal facility (vivarium) is an extension of the research laboratory, and all requirements for work with biohazardous agents and toxic chemicals in the research laboratory are applicable to work in the animal facility. The Biosafety Level (facilities, practices, and operational requirements) recommended for working with biohazard agents in vivo and in vitro are comparable. All animal work shall be in compliance with all applicable standards and regulations as noted earlier in this Manual as well as the Guide for the Care and Use of Laboratory Animals (8th Edition) and the Laboratory Animal Welfare Regulations [Animal Welfare Act] (9 CFR Subchapter A, Parts 1, 2 and 3). All research involving animals is subject to prior review by the Institutional Animal Care and Use Committee (IACUC).

The PI, in consultation with ULAR and EHS, is responsible for developing a research protocol to be submitted to the IBC when conducting animal research involving biohazard agents. This research protocol must include appropriate engineering controls, work practices and personal protective equipment to protect all personnel from the recognized hazards associated with the work.

All animal research involving biohazard agents will be completed at the appropriate animal biosafety level indicated for the biohazard agent being used as assigned by the Principal Investigator and approved by the Institutional Biosafety Officer and the IBC.

Supervisors and PIs must evaluate work done with animals and, in addition to ensuring compliance with applicable animal research regulations, ensure all personnel (research, as well as ULAR) will be adequately protected from exposure to hazardous agents associated with the animal research.

Animal satellite housing areas must be appropriate for the biosafety level of the work being performed in the space. For example, ABSL2 work cannot be performed or ABSL2 animals cannot be housed in a satellite housing spaces that are only rated or approved as ABSL1. Contact the Institutional Biosafety Officer for assistance with questions regarding animal satellite housing.

The PI must notify the animal vivarium team leader and ULAR hazard liaison via the e-Protocol system at least three working days before animals under the care of ULAR staff are treated with hazardous agents. For agents that do not require any additional handling or processing by ULAR, notification in advance is not required. The PI is responsible for posting the Animal Hazard Safety Protocol (AHSP) at the lab, housing, or procedure space for the duration of the hazard. A working day is defined as a day during which Ohio State offices are open and excludes weekends and holidays. In the interest of safety, ULAR reserves the right to euthanize those animals exposed to biohazard agents or toxic chemicals if ULAR has not received the appropriate notification.
All animal carcasses, whether infectious or not, must be disposed of as infectious waste in accordance with Appendix C. Only bedding and waste from infected animals must be disposed of in accordance with the same section of this *Manual*.

Individuals working in vivaria must recognize that conscientious personal hygiene practices establish an important barrier to infection. All individuals handling animals must wear gloves. After handling animals, their secretions or excretions, individuals shall remove their gloves, wash their hands with disinfectant soap and water and then dry their hands. Protective clothing (lab coat, uniform or surgical gown) and other safety devices such as hearing protection, facemasks and safety glasses may be required when working with animals. Eating, drinking, storing food and/or drink, smoking, or applying cosmetics in animal rooms are prohibited. Individuals should keep their hands away from their mouths, eyes, noses, and hair after handling animals. Inadvertent self-contamination with pathogens is the primary cause of reported illnesses among laboratory workers. Individuals who have open wounds should also take additional care when working with animals.

### XIV.2 Laboratory Animal Allergies

It is important to minimize exposures that could result in sensitization of animal-care and laboratory-research personnel to animal allergens. Engineering controls of animal facilities, adequate work procedures and the use of appropriate personal protective equipment can minimize exposure to laboratory animal urine, dander, fur, saliva, serum, etc.

- Allergic reactions to laboratory animals are common among personnel working with laboratory animals.
- Personnel should be made aware of the signs and symptoms of laboratory animal allergies.
- Personnel exhibiting any signs of hypersensitivity to laboratory animals (contact dermatitis, allergic conjunctivitis, allergic rhinitis, asthma) must report to Occupational Health and Wellness (Employee Health) Services for further evaluation and/or treatment.

### XIV.3 Zoonoses and Arthropodoses

Researchers working with lab animals must recognize the possibility of naturally infected animals capable of transmitting those infections (zoonoses) to lab personnel. This is particularly true of non-human primates and farm animals, but it is possible with other lab-animal species. Research work also may involve exposure to arthropods (members of the phylum *Arthropoda*, which includes the classes *Insecta, Arachnida, Pentastomida, and Crustacea*), and laboratory workers should be aware of the risks in working with these species. It is important to understand the extent to which the arthropods may or may not have been infected with agents, and to which exposure to both infected or uninfected arthropods can impact health and well-being.
Personnel working with lab animals and arthropods must be made aware of the diseases that may infect these animals and that may be transmitted to humans as well as the methods of transmission (i.e., aerosol for Tuberculosis, bites/scratches/intimate contact for Cercopithecine herpesvirus 1 [Herpes virus simiae], Rocky Mountain Spotted Fever from tick bites, skin contact for ringworm and orf, etc.).

Guidelines for Arthropod Containment at all Biosafety Levels can be found in the Arthropod Containment Guidelines, Version 3.2. Ohio State does not conduct research at ACL4.

PIs working with animals or species that can cause injury or disease in humans due to bites (snakes, ticks, mites, etc.), touch (some amphibians and fish) or other methods of transfer of venoms, poisons, etc., must work with Occupational Health and Wellness (Employee Health) to develop an appropriate plan to ensure reporting of possible exposures and provide for medical evaluation, treatment, and follow-up of personnel who are exposed to such agents.

For more information on zoonotic diseases associated with laboratory animals, click on the Animal Research tab on the EHS Research-Biosafety page.

**XIV.4 Vertebrate Animal Biosafety Level Criteria**

Institutional management must provide appropriate facilities and staff and must establish practices that reasonably assure appropriate levels of environmental quality, safety, and care for experimental animals. As a general principle, the biosafety level (facilities, practices, and operational requirements) recommended for working with biohazard agents in vivo and in vitro are comparable. The animal room, however, is not the laboratory, and can present unique problems. In the laboratory, hazardous conditions are caused by personnel or the equipment that is being used. In the animal room, the animals themselves can introduce new hazards. Animals may produce aerosols, and they may also infect and traumatize animal handlers by biting and scratching.

Guidelines for Good Practices at all Animal Biosafety Levels can be found in the Biosafety in Microbiological and Biomedical Laboratories 6th edition. Ohio State does not conduct research at ABSL4.

Specific ABSL-3 practices and procedures are described in the Ohio State ABSL-3 facility biosafety manuals.
Appendix A
Selected References

Arthropod Containment Guidelines, 2019, American Committee of Medical Entomology; American Society of Tropical Medicine and Hygiene, Vector Borne and Zoonotic Diseases; 19(3):152-173.


Biosafety in the Laboratory, 1989; National Academy Press, Washington, D.C.


Classification of Etiologic Agents on the Basis of Hazard, 1979, Department of Health, Education, and Welfare, Public Health Service, Center for Disease Control, Atlanta, Georgia.


CRC Handbook of Toxicology, 1995, Michael J. Derelanko and Mannfred A. Hollinger (Eds.), CRC Press, Boca Raton, Florida.


Protection of Laboratory Workers from Instrument Biohazards (Proposed Guideline), 1991, National


Subcommittee on Arbovirus Laboratory Safety for Arboviruses and Certain Other Viruses of Vertebrates, 1980, American Journal of Tropical Medicine Hyg, 29(6), 1359-1381.


Appendix B
Select Agents and Toxins

The Centers for Disease Control and Prevention (CDC) and the Animal and Plant Health Inspection Service (APHIS) oversee the possession, use and transfer of select agents and toxins in the United States. Select agents and toxins are biological agents and toxins that have the potential to pose a severe threat to public, animal or plant health, or to animal or plant products. Select agents and toxins are listed in Table 1 of this appendix.

Select agent regulations require entities to implement many provisions, including without limitation, Department of Justice security risk assessments of all individuals who will access select agents; providing training to all staff with access to ensure safety and security while conducting research involving select agents or toxins; developing biosecurity, biosafety, and incident response plans; and maintaining records, e.g. select agent inventory and access records and training documentation. Certain individuals, referred to as restricted persons, cannot possess or have access to select agents or toxins. Personnel are subject to significant criminal and civil penalties for the inappropriate use, possession, or transfers of select agents and toxins.

Some attenuated strains of select biological agents and toxins are excluded from the requirements of the select agent regulations. Select agents and toxins exclusions can be found at: https://www.selectagents.gov/SelectAgentsandToxinsExclusions.html

Table 2 of this appendix lists the toxins, which are excluded from select agent regulations when the amount under the control of a principal investigator does not exceed, at any time, the amounts indicated.

For additional information, contact the Institutional Biosafety Officer at the Office of Environmental Health and Safety at 614-292-1284. All select agent and toxin shipments to or from Ohio State require prior approval from the Responsible Official (RO). Ohio State has designated the RO the authority and control to ensure compliance with Select Agent Regulations.

Dual Use Research of Concern (DURC) Program. The NIH OSP defines DURC as “life science research that, based on current understanding, can be reasonably anticipated to provide knowledge, information, products, or technologies that could be directly misapplied to pose a significant threat with broad potential consequences to public health and safety, agricultural crops and other plants, animals, the environment, or national security. The United States Government’s oversight of DURC is aimed at preserving the benefits of life sciences research while minimizing the risk of misuse of knowledge, information, products, or technologies provided by such research”.

Specific information regarding the DURC Program at Ohio State can be found at https://orc.osu.edu/regulations-policies/dual-use-research-of-concern/. DURC applies to research involving the 15 select agents and toxins listed in Table 3.

<p>| TABLE 1: SELECT AGENTS AND TOXINS LIST |</p>
<table>
<thead>
<tr>
<th>HHS SELECT AGENTS AND TOXINS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abrin</td>
</tr>
<tr>
<td>Bacillus cereus Biovar anthracis*</td>
</tr>
<tr>
<td>Botulinum neurotoxins*</td>
</tr>
<tr>
<td>Botulinum neurotoxins producing species of Clostridium*</td>
</tr>
<tr>
<td>Conotoxins (Short, paralytic alpha conotoxins containing the following amino acid sequence X1CCX2PACGX3X4X5X6X7)1</td>
</tr>
<tr>
<td>Coxiella burnetti</td>
</tr>
<tr>
<td>Crimean-Congo haemorrhagic fever virus</td>
</tr>
<tr>
<td>Diacetoxyscripenol</td>
</tr>
<tr>
<td>Eastern Equine Encephalitis virus3</td>
</tr>
<tr>
<td>Ebola virus*</td>
</tr>
<tr>
<td>Francisella tularensis*</td>
</tr>
<tr>
<td>Lassa fever virus</td>
</tr>
<tr>
<td>Lujo virus</td>
</tr>
<tr>
<td>Marburg virus*</td>
</tr>
<tr>
<td>Monkeypox virus3</td>
</tr>
<tr>
<td>Reconstructed replication competent forms of the 1918 pandemic influenza virus containing any portion of the coding regions of all eight gene segments (Reconstructed 1918 Influenza virus)</td>
</tr>
<tr>
<td>Ricin</td>
</tr>
<tr>
<td>Rickettsia prowazekii</td>
</tr>
<tr>
<td>SARS-associated coronavirus (SARS-CoV)</td>
</tr>
<tr>
<td>Saxitoxin</td>
</tr>
<tr>
<td><strong>South American Haemorrhagic Fever viruses:</strong></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Chapare</td>
</tr>
<tr>
<td>Guanarito</td>
</tr>
<tr>
<td>Junin</td>
</tr>
<tr>
<td>Machupo</td>
</tr>
<tr>
<td>Sabia</td>
</tr>
<tr>
<td>Staphylococcal enterotoxins A, B, C, D, E subtypes</td>
</tr>
<tr>
<td>T-2 toxin</td>
</tr>
<tr>
<td>Tetrodotoxin</td>
</tr>
<tr>
<td><strong>Tick-borne encephalitic complex (flavi) viruses:</strong></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Far Eastern subtype</td>
</tr>
<tr>
<td>Siberiansubtype</td>
</tr>
<tr>
<td>Kyasanur Forest disease virus</td>
</tr>
<tr>
<td>Omsk hemorrhagic fever virus</td>
</tr>
<tr>
<td>Variola major virus (Smallpox virus)*</td>
</tr>
<tr>
<td>Variola minor virus (Alastrim)*</td>
</tr>
<tr>
<td>Yersinia pestis*</td>
</tr>
</tbody>
</table>
## OVERLAP SELECT AGENTS AND TOXINS

<table>
<thead>
<tr>
<th>Agent/Pathogen</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacillus anthracis*</td>
</tr>
<tr>
<td>Bacillus anthracis Pasteur strain</td>
</tr>
<tr>
<td>Brucella abortus</td>
</tr>
<tr>
<td>Brucella melitensis</td>
</tr>
<tr>
<td>Brucella suis</td>
</tr>
<tr>
<td>Burkholderia mallei*</td>
</tr>
<tr>
<td>Burkholderia pseudomallei*</td>
</tr>
<tr>
<td>Hendra virus</td>
</tr>
<tr>
<td>Nipah virus</td>
</tr>
<tr>
<td>Rift Valley fever virus</td>
</tr>
<tr>
<td>Venezuelan equine encephalitis virus3</td>
</tr>
</tbody>
</table>

## USDA SELECT AGENTS AND TOXINS

<table>
<thead>
<tr>
<th>Agent/Pathogen</th>
</tr>
</thead>
<tbody>
<tr>
<td>African horse sickness virus</td>
</tr>
<tr>
<td>African swine fever virus</td>
</tr>
<tr>
<td>Avian influenza virus3</td>
</tr>
<tr>
<td>Classical swine fever virus</td>
</tr>
<tr>
<td>Foot-and-mouth disease virus*</td>
</tr>
<tr>
<td>Goat pox virus</td>
</tr>
<tr>
<td>Lumpy skin disease virus</td>
</tr>
<tr>
<td>Mycoplasma capricolum3</td>
</tr>
<tr>
<td>Mycoplasma mycoides3</td>
</tr>
<tr>
<td>Newcastle disease virus2,3</td>
</tr>
<tr>
<td>Peste des pesteis ruminants virus</td>
</tr>
<tr>
<td>Rinderpest virus*</td>
</tr>
<tr>
<td>Sheep pox virus</td>
</tr>
<tr>
<td>Swine vesicular disease virus</td>
</tr>
</tbody>
</table>

## USDA PLANT PROTECTION AND QUARANTINE (PPQ) SELECT AGENTS AND TOXINS

<table>
<thead>
<tr>
<th>Agent/Pathogen</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coniothyrium glycines (formerly Phoma glycinicola and Pyrenochaeta glycines)</td>
</tr>
<tr>
<td>Peronosclerospora philippinensis (Peronosclerospora sacchari)</td>
</tr>
<tr>
<td>Ralstonia solanacearum</td>
</tr>
<tr>
<td>Rathayibacter toxicus</td>
</tr>
<tr>
<td>Sclerophthora rayssiae</td>
</tr>
<tr>
<td>Synchytrium endobioticum</td>
</tr>
<tr>
<td>Xanthomonas oryzae</td>
</tr>
</tbody>
</table>

*Denotes Tier 1 Agent

1 C = Cysteine residues are all present as disulfides, with the 1st and 3rd Cysteine, and the 2nd and 4th Cysteine forming specific disulfide bridges; The consensus sequence includes known toxins α-MI and α-GI (shown above) as well as α-GIA, α-CnIA, α-CnIB; X1 = any amino acid(s) or Des-X; X2 = Asparagine or Histidine; P = Proline; A = Alanine; G = Glycine; X3 = Arginine or Lysine; X4 = Asparagine, Histidine, Lysine, Arginine, Tyrosine, Phenylalanine or Tryptophan; X5 = Tyrosine, Phenylalanine, or Tryptophan; X6 = Serine, Threonine, Glutamate, Aspartate, Glutamine, or Asparagine; X7 = Any amino acid(s) or Des-X and; “Des X” = “an amino acid does not have to be present at this position.” For example if a peptide sequence were XCCHPA then the related peptide CCHPA would be designated as Des-X.
A virulent Newcastle disease virus (avian paramyxovirus serotype 1) has an intracerebral pathogenicity index in day-old chicks (Gallus gallus) of 0.7 or greater or has an amino acid sequence at the fusion (F) protein cleavage site that is consistent with virulent strains of Newcastle disease virus. A failure to detect a cleavage site that is consistent with virulent strains does not confirm the absence of a virulent virus.

Select agents that meet any of the following criteria are excluded from the requirements of this part: Any low pathogenic strains of avian influenza virus, South American genotype of eastern equine encephalitis virus, west African clade of Monkeypox viruses, any strain of Newcastle disease virus which does not meet the criteria for virulent Newcastle disease virus, all subspecies Mycoplasma capricolum except subspecies capripneumoniae (contagious caprine pleuropneumonia), all subspecies Mycoplasma mycoides except subspecies mycoides small colony (Mmm SC) (contagious bovine pleuropneumonia), and any subtypes of Venezuelan equine encephalitis virus except for Subtypes IAB or IC, provided that the individual or entity can verify that the agent is within the exclusion category.

**TABLE 2: HHS Toxins**

<table>
<thead>
<tr>
<th>Toxin</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abrin</td>
<td>1000 mg</td>
</tr>
<tr>
<td>Botulinum neurotoxins</td>
<td>1 mg</td>
</tr>
<tr>
<td>Short, paralytic alpha conotoxins</td>
<td>100 mg</td>
</tr>
<tr>
<td>Diacetoxyripenol (DAS)</td>
<td>10,000 mg</td>
</tr>
<tr>
<td>Ricin</td>
<td>1000 mg</td>
</tr>
<tr>
<td>Saxitoxin</td>
<td>500 mg</td>
</tr>
<tr>
<td>Staphylococcal Enterotoxins (Subtypes A, B, C, D, and E)</td>
<td>100 mg</td>
</tr>
<tr>
<td>T-2 toxin</td>
<td>10,000 mg</td>
</tr>
<tr>
<td>Tetrodotoxin</td>
<td>500 mg</td>
</tr>
</tbody>
</table>

**TABLE 3: DURC AGENTS**

<table>
<thead>
<tr>
<th>Agent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Highly Pathogenic Avian Influenza virus</td>
</tr>
<tr>
<td><em>Bacillus anthracis</em></td>
</tr>
<tr>
<td>Botulinum neurotoxin</td>
</tr>
<tr>
<td><em>Burkholderia mallei</em></td>
</tr>
<tr>
<td><em>Burkholderia pseudomallei</em></td>
</tr>
<tr>
<td>Ebola virus</td>
</tr>
<tr>
<td>Foot-and-mouth disease virus</td>
</tr>
<tr>
<td><em>Francisella tularensis</em></td>
</tr>
<tr>
<td>Marburg virus</td>
</tr>
<tr>
<td>Reconstructed 1918 Influenza virus</td>
</tr>
<tr>
<td>Rinderpest virus</td>
</tr>
<tr>
<td>Toxin-producing strains of <em>Clostridium botulinum</em></td>
</tr>
<tr>
<td>Variola major virus</td>
</tr>
<tr>
<td>Variola minor virus</td>
</tr>
<tr>
<td><em>Yersinia pestis</em></td>
</tr>
</tbody>
</table>
Appendix C
Infectious Waste Guidelines

Infectious Waste Generation and Treatment

The Ohio State University, as required by Ohio Administrative Code (OAC) Section 3745-27, is registered with the Ohio Environmental Protection Agency (OEPA) as a large-quantity generator of infectious waste. Faculty and staff who generate infectious waste must comply with OEPA regulations. For generators of infectious waste (faculty, staff, students, etc.) the following pages contain information dealing with these regulations. Individual PI’s/Supervisors are responsible for assuring compliance with infectious waste regulations including:

1. Identification and segregation
2. Proper packaging
3. Proper treatment
4. Personnel treatment
5. Spill and containment plans
6. Spill response
7. Spill reporting
8. Contingency plans
9. Storage

It is the PI’s/Supervisor’s responsibility to notify the Institutional Biosafety Committee (IBC), or EHS of their activities and to comply with OEPA regulations. Assistance is available from EHS to help develop and implement procedures consistent with the regulations.

Individuals who wish to treat their own infectious waste must register with EHS at 614-292-1284, obtain a treatment facility permit from OEPA and undergo quarterly laboratory audits by EHS and a representative of OEPA. Records must be kept of all waste treatment and disposal. This includes treating liquid waste with bleach and disposing in the sanitary sewer.

C.2 Definitions of Infectious Waste

1. Cultures and stocks of infectious agents (human pathogens) and associated biologicals, including without limitation, specimen cultures, cultures of stocks of infectious agents, wastes from production of biologicals and discarded live and attenuated vaccines.

2. Laboratory wastes that were, or are likely to have been, in contact with infectious agents that may present a substantial threat to public health if improperly managed.
3. Pathological wastes, including, without limitation, human and animal tissues, organs, and body parts, and body fluids and excreta that are contaminated with or are likely to be contaminated with infectious agents, removed or obtained during surgery or autopsy or for diagnostic evaluation provided that, with regard to pathological waste from animals, the animals have or likely to have been exposed to a zoonotic or infectious agent.

4. Waste materials from the rooms of humans, or the enclosures of animals, that have been isolated because of diagnosed communicable diseases that are likely to transmit infectious agents. Also included are waste materials from the rooms of patients who have been placed on blood and body fluid precautions under the Universal Precaution System established by the Centers for Disease Control and Prevention in the Public Health Service of the United States Department of Health and Human Services, if specific wastes generated under the Universal Precaution System have been identified as infectious wastes by rules referred to in C.2.8 below.

5. Human and animal blood specimens and blood products are being disposed of provided that with regard to “blood specimens and blood products” from animals, the animals were or were likely to have been exposed to a zoonotic or infectious agent. Blood products do not include patient care waste such as bandages or disposable gowns that are lightly soiled with blood or bodily fluids unless such wastes are soiled to the extent that the generator of the wastes determines that they should be managed as infectious wastes.

6. Contaminated carcasses, body parts, and bedding of animals that were intentionally exposed to infectious agents from zoonotic or human diseases during research, production of biologicals, or testing pharmaceuticals, and carcasses and bedding of animals otherwise infected by zoonotic of infectious agents that may present a substantial threat to public health if improperly handled.

7. Sharp wastes used in the treatment, diagnosis, or inoculation of human beings or animals or that have, or likely to have, come in contact with infectious agents in medical, research, or industrial laboratories, including, without limitation, hypodermic needles and syringes, scalpel blades, and glass articles that have been broken. Such wastes are hereinafter in this rule referred to as “sharp infectious waste” or “sharps”.

8. Any other waste material generated in the diagnosis, treatment, or immunization of humans or animals, in research pertaining thereto, or in the production or testing of biologicals that the Public Health Council created in Section 3701.33 of the Revised Code, by rules adopted in accordance with Chapter 119 of the Revised Code, identifies as infectious waste after determining that the wastes represent a substantial threat to public health when improperly managed because they are contaminated, or likely to be contaminated with infectious agents.
C.3 Packaging, Storage and Disposal of Untreated Infectious Waste

To meet Ohio Administrative Code Section 2745-27-30 for the packaging, storage and disposal of infectious waste, Ohio State requires the following:

C.3.1 Material

1. Red bags or biohazard bags, biohazard shipping boxes, and sharps containers.

2. All material in C.3.1.1 except sharps containers are available at no charge from EHS, excluding the Medical Center operations. Contact Environmental Services in the Hospital for additional information. Sharps containers are available through the eStores. For Ohio State locations, delivery of waste supplies can be requested by logging into the EHS online secure web application.

C.3.2 Packaging

1. Assemble the infectious waste box provided by EHS and ensure that all markings are oriented correctly with the “Up Arrows” pointed upward.

2. Tape all seams with sturdy packaging tape. NOTE: masking tape is not acceptable.

3. Line the infectious waste box with the EHS provided red plastic infectious waste bag prior to placement of infectious waste materials into the container.

4. Place only infectious material or infectious contaminated materials in the infectious waste bags used to line infectious waste boxes.

5. Store liquid infectious waste in Department of Transportation (DOT) approved plastic containers or carboys prior to packaging for pickup. NOTE: Total liquid volume is not to exceed four gallons.

6. Place liquid infectious waste containers in the bottom of a double-lined infectious waste box to facilitate pickup and storage. NOTE: Total liquid volume is not to exceed four gallons.

7. If not being immediately sealed, infectious waste containers must be closed when waste is not being actively added to the container.

8. Limit the total weight in the infectious waste boxes to 30 pounds.

9. Seal the bag prior to sealing the box.

10. Seal the box securely with packaging tape.
11. Include the building name, room number, name of the principal investigator or lab supervisor, and the waste request number on the top of the box.

12. Arrange for pickup of packaged infectious waste or to request storage containers or packaging materials via the EHS website at https://ehs.osu.edu/service-requests.

C3.3 Storage

1. Lock outside storage areas containing infectious waste containers to prevent unauthorized access.

2. Designate infectious waste storage areas. Those storage areas that are not locked, shall be visibly labeled with a sign stating “Warning: Infectious Waste” and/or displaying the international biohazard symbol on all points of access.

C3.4 Disposal

1. Request pick-up of packaged infectious waste via the EHS website at https://ehs.osu.edu/service-requests. Additional information on proper disposal, packaging requirements, and service request processes are available at EHS website.

2. Generators of infectious waste may discharge untreated liquid or semi-liquid infectious wastes consisting of blood, blood products, body fluids, and excreta into the sanitary sewer system as defined in Section 6111.01 of the Revised Code, unless the discharge is inconsistent with the terms and conditions of any permit for the system involved under Chapter 6111 of the Revised Code (OAC 3745-27-30-C).

C3.5 Spills

1. All individuals who use biohazard substances must record in a log all spills or accidents involving infectious waste. For spills in quantities greater than one gallon or which involve exposure of laboratory personnel, EHS must be notified.

2. All individuals who use biohazard substances must develop and implement a spill-containment and clean-up procedure. The procedure must be readily available to persons likely to handle infectious waste.

3. The following sections on Spill Containment and Cleanup Procedures and the Ohio State Contingency Plan are provided to meet these requirements. Modifications of procedures must be forwarded to EHS for review and comments.
C.4 Treatment by Incineration

Those who wish to treat infectious waste onsite by incineration must comply with OAC 3745-27-32. In the past, infectious waste (primarily animal carcasses and bedding) had been incinerated at Wiseman Hall, Biological Sciences or Goss Laboratory. These incinerators are not permitted by the OEPA to burn infectious waste. Administrative units responsible for these incinerators have been notified that all incineration of infectious waste must cease immediately. Infectious waste currently treated at one of these locations should be packaged according to instructions provided above in C.3. Contact EHS at 614-292-1284 for further assistance.

C.5 Treatment by Steam Sterilization

Those who wish to treat infectious waste onsite using steam sterilization must also comply with OAC 3745-27-32 and be permitted by OEPA. Contact Environmental Health and Safety for more information.

C.6 Chemical Treatment

Chemical treatment of infectious waste also requires complying with OAC 3745-27-32 and be permitted by OEPA. Contact EHS for more information.

The OEPA has only approved chemical treatment of infectious waste categorized as cultures. Therefore, chemical treatment of any other category of infectious waste must be approved by the Director of OEPA or an alternate approved-treatment method used.

C.7 Spill Containment and Clean-up Procedures

According to OAC 3745-27-30, spill containment and a clean-up kit shall be available in those areas designated in the Spill Containment and Cleanup Procedures. The location of the kits shall provide for rapid and efficient cleanup of spills anywhere within these areas.

C.7.1 Spill Kit Materials

The kit shall include but is not limited to:
1. Absorbent material
2. One gallon approved chemical disinfectant (bleach)
3. Red bags or bags labeled with the biohazard symbol
4. Impermeable and disposable overalls (preferably Tyvek total body coveralls)
5. Gloves (heavy neoprene or latex)
6. Goggles (can be reusable)
7. Rigid plastic container for sharps
8. first aid kit unless emergency care is available on the premises
9. Boundary tape and other appropriate safety equipment
C.7.2 Cleanup Procedures

1. A copy of the cleanup procedures is provided later in this section (See Appendix C.9).
2. More specific or detailed cleanup procedures can be prepared by the generator.

C.7.3 Spill Log

1. A copy of the spill log is also provided.
2. Spill logs must be maintained for 5 years.
3. All spills greater than one gallon or which involve exposure of laboratory personnel must be reported to EHS immediately and those spills of volume greater than one cubic foot must be reported to EHS and to the Director of OEPA within 48 hours.

C.8 Contingency Plan

In accordance with OAC 3745-27-32 and 35, a contingency plan for treatment facilities must be available at treatment sites. In the event that sites which treat infectious waste cannot meet the storage requirements described below or are experiencing a malfunction in treatment possess, the contingency plan shall be implemented.

C.8.1 Storage

1. Store infectious waste in a manner that maintains the integrity of packing.
2. Maintain waste in a non-putrescent state, using refrigeration or freezing if necessary
3. Lock outside storage to prevent unauthorized access.
4. Designate and label storage areas by posting biohazard warning signs.
5. Store infectious waste in a manner that affords protection from animals.
6. No infectious waste may be stored more than 14 days.
7. No more than seven times the treatment facility’s total maximum daily throughput capacity shall be stored for treatment.
8. Contain and clean up any spills of infectious waste within a storage area using approved methods.
CONTINGENCY PLAN

Emergency Coordinator: Tom Novotny, EHS
Telephone: 614-292-1284
Alternate Coordinator: On-call Representative (EHS Emergency Response Team): 614-292-1284

1. If you cannot comply with the storage requirements set forth, the following contingency plan shall be implemented:
   a. Notify your Emergency Coordinator.
   b. Call EHS and request red bags, biohazard boxes, and sharps containers as needed for packing infectious waste at your treatment location.
   c. Following packaging of infectious waste, EHS will arrange for offsite incineration.

2. Listing of emergency telephone numbers in addition to the Emergency Coordinator.
   a. Ohio State Police Dispatcher: 911
   b. EHS Chemical/Infectious Waste Management: 614-292-1284 -option 2
   c. EHS Main Office: 614-292-1284
   d. OEPA Central Distract Office: 614-728-3778
   e. Emergency Number: 911
   f. Columbus Health Department: 614-645-7417

CONTACT: Tom Novotny, 1314 Kinnear Rd, Tel: 614-292-1284

Infectious Waste Spill Containment and Cleanup Procedure

In accordance with OAC 3745-27-30, the following containment and cleanup procedures are to be implemented in the event of an infectious waste spill.

Infectious waste spills must be contained and cleaned up immediately.

I. A spill kit containing absorbent material, bleach or other USEPA registered tuberculocidal disinfectant, biohazard bags, gloves, eye protection, and a biohazard sharp container must be accessible in the laboratory.

II. To use bleach as a disinfectant, a 1:10 dilution (minimum 10% sodium hypochlorite solution) of household bleach should be prepared immediately prior to use, with a minimum of 30 minutes contact time with the waste. If another USEPA registered tuberculocidal disinfectant is used, the manufacturer’s recommendations for concentration and contact time should be followed.
a. Limit access to area to authorized personnel.
b. Open the spill kit.
c. Put on appropriate PPE (i.e. gloves, eye protection, coveralls).
d. Contain liquid spills by covering with absorbent pads. Place contaminated absorbent pads and other contaminated solids into a biohazard bag. Seal the bag by tying in a knot and place into a second biohazard bag. Sharps (i.e. needles, blades, or broken glassware) associated with the spill should be placed in a biohazard sharps container.
e. Clean the spill and cover contaminated surfaces with absorbent pads and soak with appropriate disinfectant (See II above). Allow the disinfectant to stand on the contaminated material for the minimum recommended contact time.
f. Place all materials used during the cleanup process in a biohazard bag. Seal the bag by tying in a knot and place into a second bag. Place all into a biohazard burn box.
g. Disinfect all re-usable materials from the spill kit. Replenish disposable items from the spill kit.

**Infectious Waste Spill Report**

A spill report is required under OAC 3745-27-30(A) (10) for any spills that are greater than or equal to one cubic foot in volume. Complete this report and return to the address listed below.

Date and Time of Spill: __________________________ Date of Report: _______________

Location of Spill: _______________________________________________________________

Employee(s) Involved in Cleanup: _______________________________________________

Waste Spilled: __________________________ Quantity (estimated):__________________

Describe Cleanup Procedure: ____________________________________________________

____________________________________________________________________________

Summary of Events Causing Spill (if known): ______________________________________

____________________________________________________________________________

Printed Name: __________________________ Date: __________________________

Signature: ______________________________

Mail Completed Report to:
  David Puskas
  Office of Environmental Health and Safety
  1314 Kinnear Road, CAMPUS
Appendix D
Accident/Incident Follow-up Information

For severe or life-threatening injuries to employees of Ohio State, immediately call 911. Also, promptly contact the EHS at 614-292-1284 to report the injury.

For minor occupational injuries, the supervisor should direct the employee to Occupational Health and Wellness (Employee Health) during business hours (McCaw Hall 2nd Floor, 1581 Dodd Dr. 614-293-8146) or the Ohio State Wexner Medical Center Emergency Department after hours (410 West 10th Ave.).

For all injuries, the supervisor shall:

- Ensure the injured employee completes an Ohio State Employee Accident Report.
- Sign the Employee Accident Report.
- Complete the Supervisor Accident Analysis Report after investigating the incident to determine a root cause; and as appropriate, implement measures to prevent recurrence.
- Ensure the accident report and supporting documentation is submitted to the Office of Human Resources, Absence Management and Vocational Services by email at accidentreport@osu.edu or fax 614-688-8120.

For assistance or if you have questions, contact EHS at 614-292-1284.

D.1 Accident Investigation Purpose

The intent of the investigation is prevention and correction (i.e., identify the root cause(s), not to assign blame.

D.2 Definitions

Accident: an undesired event that results in personal injury or property damage.

Incident: an unplanned, undesired event that adversely affects the completion of a task.

Near Miss: incidents where no property was damaged and no personal injury was sustained, but where, given a slight shift in time or position, damage and/or injury easily could have occurred.

D.3 When to Conduct an Investigation

All incidents whether a near miss or an actual injury-related event should be investigated. Near miss reporting and investigation enable identification and control of hazards before they cause a more serious incident. Accident/incident investigations are tools for uncovering hazards that were either...
missed during earlier job hazard analyses or have managed to slip away from the controls planned for them. To be useful an investigation needs to be done with the aim of discovering every contributing factor to accident/incident in order to “fail-safe” the condition and/or activity to prevent recurrence.

While all accidents should be investigated, including accidents involving property damage only; the extent of the investigation shall be reflective of the seriousness of the accident.

When an accident results in a fatality or hospitalization of employees, promptly contact EHS. In these situations, (except to the extent necessary to protect employees and the public) evidence at the scene of an accident shall be left untouched until the site has been inspected by health and safety officials.

D.4 Who Should Investigate

The usual investigator for incidents is the supervisor in charge of the involved area and/or activity. The supervisor should be accountable for accidents in his/her area, should know the situation and the people involved best, has a personal interest in cause identification and can take immediate corrective action. The injured or impacted employee should be involved as the individual can clarify any uncertainties by providing details on what happened and why it occurred. Employee involvement in investigations will not only provide additional expertise and insight, but in the eyes of the workers, will lend credibility to the results. Employee involvement can enhance employee knowledge of potential hazards and the experience can make employees advocates of the importance of safety thus strengthening the safety culture of the organization.

EHS will assist with investigation of accidents involving fatalities, serious injuries or extensive property damage. Upon request, EHS will provide assistance investigating any accident, incident or near miss. EHS will review the investigation finding and recommendations.

D.5 Key Questions

Six key questions should be answered: who, what, when, where, why, and how. Facts should be distinguished from opinion, and both should be presented carefully and clearly. The report should include thorough interviews with everyone with any knowledge of the incident. A good investigation is likely to reveal several contributing factors, and it probably will recommend several preventative actions.

Avoid the trap of laying sole blame on the injured employee. Even if an injured worker openly blames his/herself for making a mistake or not following a prescribed process; the accident investigator should not be satisfied that all contributing causes have been identified. The error made by the employee may not be the most important contributing cause. The employee who has not followed prescribed procedures may have been encouraged directly or indirectly by a supervisor or expected workload to “cut corners”. The prescribed procedures may not be practical or even safe in the eyes of the employee. Sometimes where elaborate and difficult procedures are required,
engineering redesign might be a better answer. In such cases, management error—not employee error—may be the most contributing cause.

All supervisors and others who investigate accidents/incidents should be held accountable for describing causes carefully and clearly. Investigation reports should not include catch-phrases, such as “Employee did not plan job properly.” While such a statement may suggest an underlying problem with this worker, it is not conducive to identifying all possible causes, preventions, and controls. Certainly, it is too late to plan a job when the employee is about to do it. Further it is unlikely that safe work will always result when each employee is expected to plan procedures alone.

D.6 Implications of Accident Investigations

Recommended preventative actions should make it very difficult, if not impossible, for the incident to recur. The investigative report should list all the ways to “fail-safe” the condition or activity. Considerations of cost or engineering should not enter at this stage. The primary purpose of accident investigations is to prevent future occurrences. Beyond this immediate purpose, the information obtained through the investigation should be used to update and revise controls used to reduce hazards to employees. For example, the Job Safety Analysis should be revised, and employees retrained to the extent that it fully reflects the recommendations made by an accident/incident investigation report. Implications from the root causes of the accident need to be analyzed for their impact on all other operations and procedures.

D.7 References

1. Occupational Safety and Health Administration Fact Sheets: Accidents/Incident Investigations.

2. Occupational Safety and Health Administration: Accident/Incident Investigation Tools and Tips.
Appendix E
Plant Biosafety

The NIH Guidelines for Research Involving Recombinant or Synthetic Nucleic Acid Molecules specifies the physical and biological containment conditions and practices suitable to the greenhouse conduct of experiments involving recombinant or synthetic nucleic acid molecule-containing plants, plant associated microorganisms, and small animals.

Information regarding standard practices, greenhouse design and access, records, and decontamination and inactivation requirements for BSL1-P through BSL4-P can be found at https://osp.od.nih.gov/wp-content/uploads/NIH_Guidelines.pdf.

Plant containment levels BSL1-P through BSL4-P are designed to provide differential levels of biosafety for plants in the absence or presence of other experimental organisms that contain recombinant or synthetic nucleic acid molecules.

For experiments in which plants are grown at BSL1 through BSL4 laboratory settings, containment practices shall be followed as described in Section VI. These containment practices include the use of plant tissue culture rooms, growth chambers within laboratories, or experiments performed on open benches.
Appendix F
Website Addresses

Ohio State Environmental Health and Safety. https://ehs.osu.edu/

Office of Responsible Research Practices. https://orrp.osu.edu/

College of Medicine. https://medicine.osu.edu/

ABSA International. https://absa.org/

American Society for Microbiology. https://www.asm.org/


Centers for Disease Control and Prevention. https://www.cdc.gov/

U.S. Food and Drug Administration. https://www.fda.gov/home


Occupational Safety and Health Administration. https://www.osha.gov/


USDA APHIS. https://www.aphis.usda.gov/aphis/home/
Appendix G
Revisions

2023 Institutional Biosafety Manual Revisions
- Throughout entire document, CDC’s *Biosafety in Microbiological and Biomedical Laboratories* 5th edition updated to 6th edition.
- Throughout the entire document, Institutional Biosafety contact information updated.
- Throughout the entire document, hyperlinks verified and updated when necessary.
- Page 22, proper location of a Biological Safety Cabinet (BSC) was updated.
- Entire document was reformatted, branded, and made accessible.

Biosafety cabinets should be located away from:
- Laboratory equipment that creates air movement
- Portable fans
- Air supply registers
- Chemical fume hoods
- Open windows
- Foot traffic

Contact the Institutional Biosafety Officer for assistance when selecting a location for a new biosafety cabinet.

- Page 22, secondary container requirement added for transporting etiological agents.
  A secondary container is required when transporting etiological agents between rooms, floors, or buildings. These secondary containers should be leak-proof, durable containers that must be marked with the universal biohazard symbol and contain enough absorbent material to contain the liquid samples if they were to leak during transit.
- Page 22, secondary container requirement added for transporting ABSL2 animals.
  A secondary container is also required when transporting ABSL2 animals between rooms, floors and buildings. Secondary containment is necessary to contain cage contents and prevent a release of potentially infectious material if a cage is upset or toppled while in transit. Secondary containment options could include biohazard bags, Rubbermaid containers on carts or enclosed cabinets on wheels. It is also important to remember not to overload a cart with too many cages even if the cages are in secondary containment.
- Page 22, clearing a RAD posted BSC for recertification. If a biosafety cabinet is posted for radioactive sample work, it must be cleared by EHS prior to its annual recertification. Please reach out to your EHS representative to obtain a maintenance letter prior to recertifying any biosafety cabinet that is used for radioactive materials.
- Page 50, satellite housing statement added.
  Animal satellite housing areas must be appropriate for the biosafety level of the work being performed in the space. For example, ABSL2 work cannot be performed or ABSL2 animals cannot be housed in a satellite housing spaces that are only rated or approved as ABSL1. Contact the Institutional Biosafety Officer for assistance with questions regarding animal satellite housing.