# UNIVERSITY OF ESSEX BIOLOGICAL SAFETY POLICY

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<tr>
<td>2</td>
<td>C. Smith</td>
<td>16/09/2021</td>
<td>Update to section 4.7 ethical approval and section 4.8 on donors.</td>
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SECTION 1
POLICY STATEMENT

The University of Essex recognises its responsibility to comply with current legislation and appropriate guidelines in the management of hazardous biological agents. This document and the departmental local rules for biological agents form part of the University requirements to comply with the Control of Substances Hazardous to Health Regulations.

This policy applies to all intentional work which involves the handling, storage, transport and waste disposal of biological agents at the University. Work placements in external health care facilities are not covered by this policy. This also does not include contact with blood/bodily fluid through first aid/cleaning activities.

The use of Genetically Modified Organisms (GMOs) is not covered in detail in this policy. This work is only carried out in the School of Life Sciences and is therefore covered by departmental local rules.

The policy will be reviewed and updated by Workplace Wellbeing every 5 years.
SECTION 2
RESPONSIBILITIES

Full details of health and safety responsibilities of all employees can be found in the University’s Health and Safety Policy. The additional responsibilities for biological safety are as follows:

2.1 UNIVERSITY STRATEGIC RESPONSIBILITIES

- Full strategic responsibilities for health and safety are in the University Health and Safety Policy.
- The University Steering Group (USG) has delegated authority from Council for the effective implementation of the University’s Health and Safety Policy. All members of USG are collectively and individually responsible for the overall health and safety management of the University.

2.2 DESIGNATED HEAD OF DEPARTMENT (DHoD)

- Overall responsibility for ensuring that there are effective arrangements in place and that there are adequate resources allocated to the control of biological safety within their department.
- Appointment of a Departmental Biological Safety Officer (DBSO).
- Approval of all biological schemes of work/risk assessments.

2.3 UNIVERSITY BIOLOGICAL SAFETY ADVISER (UBSA)

- Advises on technical requirements and procedures to enable the University to meet statutory requirements, relevant industry/sector guidance and Approved Codes of Practice.
- Ensures recording systems meet requirements of statutory bodies.
- Advises on the technical implications of new and revised legislation and codes of practice.
- Chairs the Biological Hazards and Genetic Modification Sub-Group (BHGMSG) and gives final signature for schemes of work on behalf of the committee.
- Oversees the content of relevant Local Rules and the University Biological Safety Policy.
- Maintains a list of current schemes of work/workers lists.
- Submits documentation as required to the Health and Safety Executive (HSE), including notification of first use of a substance (refer to section 4.5).
- May obtain expert competent technical advice where appropriate.

2.4 DEPARTMENTAL BIOLOGICAL SAFETY OFFICER (DBSO)

- Appointed by the DHoD in consultation with the UBSA.
- Advises the DHoD on all matters relating to biological hazards.
- Represents the department on the BHGMSG.
- Appraises all schemes of work for the department before they are submitted to the BHGMSG for approval.
- Co-ordinates day to day operational activities and pre-screens local issues before they are sent to the UBSA or BHGMSG.
- Advises the DHoD on whether Class 1 projects can be approved or should be escalated to the BHGMSG.
- May obtain expert competent technical advice where appropriate.
2.5 BIOLOGICAL HAZARDS AND GENETIC MODIFICATION SUB-GROUP (BHGMG)

- Considers changes in legislation and best practice that significantly affect the management of biological agents at the University and recommend to HSG any changes in policy or practice that might be required.
- Examines risk assessments and other relevant documentation for all projects involving:
  - genetic modification (GM) work
  - the use of biological agents
  - any proposed use of specified animal pathogens
  - the use of plants, plant products and plant pests requiring a licence or other permission
  - the use of pathogens and toxins subject to Schedule 5 of the Anti-terrorism, Crime and Security Act 2001
- Gives final approval for schemes of work for Hazard Group 2 pathogens (refer to section 3.1).
- To monitor and report, as appropriate, to Health & Safety Group on compliance with statutory and University requirements.
- Provides reports, gives advice and makes recommendations to and on behalf of the Health & Safety Group on any matter involving biosafety for the possession, use and control/containment of GMOs and other hazardous biological agents/materials, as relates to this Policy.

The Sub-group must be satisfied that:

a) schemes relating to undergraduate practicals are justified academically and there is no viable alternative; and
b) all reasonable measures have been taken to minimise any risk to researchers and students.

2.6 PRINCIPAL INVESTIGATOR (PI)

- Ensure that the risks associated with their work are fully assessed as required by current legislation.
- Maintains an up to date copy of the scheme of work for their project and provide copies of these for the UBSA.
- Maintain a list of all researchers (staff, post-graduates, undergraduates and visiting workers) who are currently working (or have worked on) their projects that include biological agents and provide a copy for the UBSA.
- Regularly review the risks and worker’s lists.
- Where there is no approved classification for a biological agent, provisionally classify the agent according its nature and properties.
- Ensure that workers understand and comply with departmental local rules and work to the correct containment level.
- Ensure that adequate training is given and that changes are notified.
- Provide safety information for Permits to Work which are required by maintenance staff and contractors before they may enter any area where work with biological agents is taking place.
- Inform the UBSA of any new schemes of work which may need notification to the HSE (refer to section 4.5).
SECTION 3
DEFINITIONS

3.1 HAZARD GROUPINGS OF BIOLOGICAL AGENTS

The term biological agent is defined under the Control of Substances Hazardous to Health Regulations (COSHH) as “a microorganism, cell culture or human endoparasite, whether or not genetically modified, which may cause infection, allergy, toxicity or otherwise create a hazard to human health.” Biological agents include bacteria, viruses, fungi, agents which cause Transmissible Spongiform Encephalopathies (TSEs), microscopic endoparasites and microscopic forms of larger endoparasites.

Biological agents are classified into four Hazard Groups, based on their ability to infect healthy humans:

<table>
<thead>
<tr>
<th>Hazard Group</th>
<th>Description</th>
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<tbody>
<tr>
<td>1</td>
<td>Unlikely to cause human disease</td>
</tr>
<tr>
<td>2</td>
<td>Can cause human disease and may be a hazard to employees; it is unlikely to spread to the community and there is usually effective prophylaxis or treatment available.</td>
</tr>
<tr>
<td>3</td>
<td>Can cause severe human disease and may be a serious hazard to employees; it may spread to the community, but there is usually effective prophylaxis or treatment available.</td>
</tr>
<tr>
<td>4</td>
<td>Causes severe human disease and is a serious hazard to employees; it is likely to spread to the community and there is usually no effective prophylaxis or treatment available.</td>
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Source: ACDP Approved List of Biological Agents

The Advisory Committee on Dangerous Pathogens (ACDP) categorises biological agents that are human pathogens and publishes these in the Approved List of Biological Agents.

Only agents in Hazard Groups (HG) 2, 3 and 4 are listed in this publication. Those not listed in these groups are not necessarily classified in Group 1. If there is no approved classification for an agent, the Principal Investigator must provisionally classify the agent according its nature and properties (refer to Schedule 3 Part 1 of the COSHH Regulations). A risk assessment should be carried out to determine the containment level required for working with any organisms. The risks of allergenicity and toxicity should also be considered.

3.2 CONTAINMENT LEVELS

Working with HG2 biological agents requires a minimum of containment level 2 (CL2). For laboratory areas, the ACDP publication Management and operation of microbiological containment laboratories describes the requirements for containment levels.

**Note:** The University of Essex does not have facilities to work at CL3 or 4; therefore research at these levels is not permitted under any circumstances.

If work is being carried out on unscreened blood where the presence of HG 3 or 4 is unlikely, the sample is presumed to be HG2 and therefore requires containment level 2.
3.3 GENETICALLY MODIFIED ORGANISMS

Genetic modification is defined as any alteration of the genetic material (DNA or RNA) of an organism using a method that does not occur naturally by mating and/or natural recombination. As a result of this modification, a novel product will be formed that may have an altered characteristic to that of the original organism. This is known as a genetically modified organism (GMO). GMOs which are plants or animals are called larger GMOs (LGMOs); and genetically modified micro-organisms (such as bacteria, viruses, parasites or fungi) are called genetically modified micro-organisms (GMMs).

Work with GMOs and GMMs at the University is covered by the Genetically Modified Organisms (Contained Use) Regulations 2014. “Contained Use” covers any activity in which organisms are genetically modified, or in which GMOs are cultured, stored, transported, destroyed, disposed of or used in any other way and for which physical, chemical or biological barriers, or any combination of such barriers, are used to limit their contact with people and the environment. There is a requirement to notify the HSE of GM work under certain circumstances.

Work with genetically modified organisms must have a risk assessment where appropriate and will need to be approved by the Biological Hazards and Genetic Modification Sub-Group (BHGMSG). Departmental local rules must be in place before any GMO work commences.

The School of Life Sciences is currently the only department permitted to carry out genetic modification research. Any other department considering GMO research MUST contact the UBSA for advice before commencing any work.
SECTION 4
ARRANGEMENTS

4.1 APPOINTMENT OF A DBSO
All departments which handle biological samples must appoint a DBSO. This should be a member of staff (usually academic) who is knowledgeable and competent in work with biological agents.

4.2 DEPARTMENT LOCAL RULES
All departments which handle biological samples must have documented Local Rules detailing their arrangements for controlling the risks. This must include:

- The scope of the activities covered in the department
- Identification and description of the containment levels and areas.
- Local responsibilities (in addition to those in Section 2 of this policy and the University Health and Safety Policy).
- Arrangements for visitors, visiting workers and contractors.
- Risk assessment/scheme of work requirements (such as justification for grouping projects).
- Training and supervision requirements.
- Recordkeeping.
- Equipment maintenance and inspection regimes (including fumigation if applicable).
- Control measures, such as use of microbiological safety cabinets, designation of work areas, signage and personal protective equipment.
- Disinfection procedures.
- Emergency procedures including identifying how accidental exposures may occur and the appropriate action to take.
- Transport, receipt, labelling and packaging of samples (including action to take if packaging is damaged).
- Storage and inventory management.
- Lone working and out of hours policy.
- Waste disposal arrangements.

Local rules should be reviewed at least every 3 years.

4.3 RISK ASSESSMENT
All work with biological agents (or material which may contain biological agents, such as human tissue) must have an up to date risk assessment, which is covered under a scheme of work. Refer to Section 5.

4.4 WORKERS LISTS
An up to date workers list must be kept for every scheme of work involving biological agents or genetically modified organisms. This is maintained by the Principal Investigator and a copy provided to the UBSA.

4.5 NOTIFICATION TO THE HSE
The first use (i.e. deliberate work) of a biological agent at a premises must be notified to the HSE prior to the work commencing. This is carried out by the UBSA, therefore they should be informed of all new scheme proposals and the revision of any existing schemes at CL2 before work commences. Allow at least 20 days for the assessment by the HSE before work starts.
There is also a requirement to notify subsequent use of specific agents listed in Part V of Schedule 3 of the COSHH Regulations.

The formal acknowledgement letter from the HSE will be held by the UBSA.

4.6 APPROVAL PROCESS

All work with HG2 agents must be approved by the BHGMSG before the work commences. Refer to Section 5.

4.7 ETHICS APPROVAL

All research with human participants (including work with human tissue and bodily fluids) must follow the University ethics approval process. Any research involving human tissue or bodily fluids which are not immediately rendered acellular may require ethical approval from an NHS Research Ethics Committee flagged to review such research in order to ensure compliance with the Human Tissue Act. In such instances, further advice should be sought from the REO Research Governance team.

A valid risk assessment must accompany the application for ethical approval in the ERAMS.

4.8 RISKS FROM BLOOD-BORNE INFECTIONS

Blood-borne viruses (BBVs) are found in blood, semen and vaginal secretions and breast milk. There is minimal risk associated with other body fluids unless contaminated with blood. Carriers of these viruses (including HIV, Hepatitis B and Hepatitis C) may not be ill or have any symptoms. Therefore all human blood, body fluids and tissue samples should be considered to be potentially infectious.

In the laboratory -

- Potential transmission of BBVs is most commonly by accidental puncture injury through the skin, by a contaminated needle or other sharp object.
- BBVs may also be spread by spillage or splashing on to mucous membranes, eyes or open wounds, abrasions/damaged skin.

All work carried out on materials that may contain unidentified biological agents must be carried out at a minimum of CL2. ACDP has advised that all such specimens must be treated as though they are infected with BBVs and allocated to HG2 as a minimum. Processes that generate large drops or aerosols of blood or similar infectious fluids must be carried out in a Class II microbiological safety cabinet.

Collection of samples

Only trained phlebotomists may take whole blood samples of up to 100ml for research purposes. A certificate of qualification for phlebotomy must be seen and validated by the supervising academic before blood can be taken. Samples taken using an autolet do not require a qualified phlebotomist. However, staff/students taking samples by this method or by or equivalent medical device must receive training (either by someone competent/experienced in the technique, or a formal course) before commencing their research. The use of manual lancets will only be considered in exceptional circumstances following completion and authorisation of a specific risk assessment.

It is the University policy that phlebotomists must be referred to Occupational Health for a course of Hepatitis B vaccinations.

Donors

Blood may be taken from volunteer donors for research purposes, provided that the informed and written consent of the donor is obtained. The department should define a pre-screening form for potential donors with advice from Occupational Health.
The designated person taking blood must maintain a record of the volumes taken from each donor in a blood sample collection record book. The volume of blood donated by any individual in a 12-week period should not exceed 200 ml, which is the safe level for healthy donors to the National Blood Service. It is the responsibility of the designated person to ensure that this amount is not exceeded and that those donors, who suffer symptoms of ill health, cease participation in the research and consult their medical practitioner. The DBSO must keep a central record of blood donations available for inspection.

**Needlestick injuries**

Anyone who sustains a needle stick injury is required to seek first aid, submit an incident report and be assessed by Occupational Health.

### SECTION 5

**RISK ASSESSMENT AND APPROVAL**

#### 5.1 SCHEME OF WORK

All work with biological agents (or material which may contain biological agents, such as human tissue) must have an up to date scheme of work. This forms the risk assessment for the project and must specify the containment level for the work. (Note that there may also be other associated risk assessments required for the project, such as for specific laboratory techniques). It may be appropriate to combine more than one project on the same scheme of work (such as for undergraduate project work) but this must be agreed with the DBSO or UBSA in advance.

Organisms that produce toxins or allergens harmful to humans (e.g. toxin-producing cyanobacteria or toxin-producing corals) must also have a biological scheme of work.

Schemes of work and risk assessments should be reviewed regularly, at an interval defined by the department.

**No work with HG3 or HG4 agents is permitted at the University of Essex under any circumstances.**

#### 5.2 APPROVAL PROCESS FOR WORK WITH BIOLOGICAL AGENTS

All work with HG2 agents must be approved by the BHGMSG before the work is carried out. The approval process for HG1 and HG2 agents is shown is as follows:
New or revised biological risk assessment

Complete section 1 of RA

Hazard group 1

DBSO for approval

HoD for approval

UBSA for record

Summary to BHGMSG

Hazard group 2

Complete section 2 of RA

DBSO for approval

HoD for approval

UBSA for circulation to BHGMSG

BHGMSG for approval

UBSA for signoff and notifications as required
SECTION 6
RECORD KEEPING

6.1 BHGMSG DOCUMENTATION
The UBSA holds the following documents:

- Copies of all schemes of work, associated risk assessments and workers lists.
- Correspondence with the HSE.
- Notifications to the HSE.

The BHGMSG Secretary holds the records of approvals and minutes from the BHGMSG meetings.

The DBSO holds donor lists and local records.

6.2 RISK ASSESSMENTS
The department must keep copies of all relevant documentation for work with biological hazards; this may be electronic or hard copy.

SECTION 7
TRAINING AND SUPERVISION

All departments must ensure that personnel are adequately trained in safe handling of biological agents before being allowed to work with them.

They must ensure that relevant staff and students work in accordance with the local rules, risk assessments and procedures for use of laboratory facilities and equipment.

Students must be effectively supervised by senior staff who are competent and experienced in the procedures being supervised.

There are specific training and competence requirements for the DBSO and UBSA roles, arranged by the Health and Safety team.

SECTION 8
REVIEW

Compliance with this policy and local rules is checked annually as part of the routine departmental health and safety inspections. More detailed audits of the system are carried out periodically by the UBSA/DBSO.

Incidents and near misses must be reported through the normal University procedures and will be followed up by the UBSA/DBSO.
## Appendix 1 - Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>ACDP</td>
<td>Advisory Committee on Dangerous Pathogens</td>
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<td>BBV</td>
<td>Blood-borne viruses</td>
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<tr>
<td>BHGMSG</td>
<td>Biological Hazard and Genetic Modification Sub Group</td>
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<tr>
<td>CL</td>
<td>Containment Level</td>
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<tr>
<td>COSHH</td>
<td>Control of Substances Hazardous to Health</td>
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<td>DBSO</td>
<td>Department Biological Safety Officer</td>
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<td>DHoD</td>
<td>Designated Head of Department</td>
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<td>GMOs</td>
<td>Genetically modified organisms</td>
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<td>HG</td>
<td>Hazard Group</td>
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<td>HSE</td>
<td>Health and Safety Executive</td>
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<td>HSG</td>
<td>Health and Safety Group</td>
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<td>MSC</td>
<td>Microbiological Safety Cabinet</td>
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<td>PI</td>
<td>Principal Investigator</td>
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<tr>
<td>PPE</td>
<td>Personal Protective Equipment</td>
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<tr>
<td>UBSA</td>
<td>University Biological Safety Adviser</td>
</tr>
<tr>
<td>USG</td>
<td>University Steering Group</td>
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