Guidance on the use, testing and maintenance of laboratory and animal isolators for the containment of biological agents
1. Introduction

1. This guidance gives advice on factors to consider when selecting, using, testing and maintaining negative pressure isolators for experiments involving biological agents and replaces previous guidance issued in 1985. The main scenarios covered by this guidance are laboratory isolators for housing equipment/large experiments etc and animal husbandry isolators for animals infected with biological agents. Pharmaceutical isolators, particularly those operated at positive pressures are covered by separate publications.

2. It is aimed at employers, laboratory managers, safety advisors and users. Other groups such as manufacturers and companies which offer testing and maintenance services may also find it useful.

3. It applies in situations where isolators are selected as the mechanism of primary containment, usually because the task is unsuitable to carry out in a microbiological safety cabinet. It does not cover other laboratory or animal containment strategies such as individually ventilated cages (IVCs) or downdraft tables. The main reason for this is current limited knowledge on the effectiveness of these alternatives for adequately containing aerosols of biological agents, and the range and complexity of testing required to validate this containment. This is not to say that these alternatives cannot be used for work with biological agents, but that it will fall to dutyholders to undertake the relevant testing and monitoring to demonstrate ongoing containment. Given that the use of IVCs is expanding in this area, it is likely that more data will become available to hopefully enable supplementary guidance to be made available in the future.

2. Legal requirements

Health & Safety Issues

Biological Agents

4. The Control of Substances Hazardous to Health Regulations 2002 (as amended) require employers to prevent, or if this is not reasonably practicable, adequately control exposure of employees to biological agents using measures other than personal protective equipment. Where exposure cannot be prevented employers must take steps to ensure that exposure is adequately controlled by using a combination of control measures specified in Schedule 3 of which states that infected material, including any animal is to be handled in a safety cabinet or isolator or other suitable containment.

5. The Genetically Modified Organisms (Contained Use) Regulations 2000 require suitable containment and control measures be used to adequately control against exposure to biological agents which are genetically modified and the Genetically Modified Organisms (Contained Use) (Amendment) Regulations 2005 specifically state that work with infected animals at containment levels 3 and 4 must be undertaken in isolators.

6. For small scale in vitro laboratory activities a microbiological safety cabinet (MSC) is widely recognised as offering a practicable engineering solution for controlling aerosols. However for non-standard laboratory activities (such as those involving large pieces of equipment) or animals infected with biological agents, it may not be practicable to use an MSC.

7. Isolators that are used, tested and maintained appropriately are recognised as being able to provide effective engineering control against aerosols from biological agents associated with such activities. However, fundamental to the whole process is a robust risk assessment addressing the specifics of the activities to be undertaken and taking into account relevant guidance, for example working safely with research animals.

Other Health & Safety issues

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*A biological agent is defined in 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’.*
8. Whilst the primary use of isolators is to control exposure of operators to aerosols of biological agents, it is important to take into account other health and safety issues, including human factors such as ergonomic considerations, in the risk assessments. Further information on this subject can be found in Annex A.

Animal Welfare Issues

9. Animal welfare requirements are specified by the Animals (Scientific Procedures) Act 1986. The associated Home Office Code of Practice\(^6\) specifies, amongst other things, the minimum environmental parameters that must be met in relation to space, noise, temperature and humidity etc for different animal species.

Environmental Issues

10. The GMO Contained Use Regulations require that appropriate measures be in place to protect the environment as well as human health. In addition, the Specified Animal Pathogen Order 1998\(^7\) requires the containment of exotic animal pathogens so that the environment is protected. Isolators may be a useful form of primary containment to reduce the likelihood of an environmental pathogen escaping.

3. Types of isolators

11. During selection of an isolator consideration of the main task(s) to be carried out along with related activities (for example, the movement of samples, removal of waste etc) is essential. Further information regarding isolator selection can be found in Annex B.

Flexible Film Isolators

12. Flexible Film Isolators (FFIs) are tented enclosures of flexible plastic built on a metal frame. They are routinely accessed from the side through gauntlets or sleeves with cuffed gloves. Ports allow material to both enter and exit the isolator.

13. The size of these units and therefore the amount of equipment within is limited by the degree of access afforded by the gauntlets.

*Flexible Film Isolators (photo Copyright Health Protection Agency)*
Rigid Isolators

14. These are essentially the same as FFI’s except their canopies are made of a rigid material.

Flexible half-suit isolators

15. These are larger than FFIs and are accessed by the operator by half suit(s) in the base of the isolator and also by gauntlets/sleeves and gloves on the side. Operators enter the half suit by ducking under the base and then standing up within the positively pressurised suit.

16. This combination gives the potential for large work surfaces within the isolator due to the increase in accessibility.

Flexible half-suit isolators (photo Copyright Health Protection Agency)
Rigid half-suit isolators

17. These are essentially the same as flexible half suits isolators except their canopies are made of a rigid material. This makes them more resistant to damage, but less able to absorb pressure fluctuations, for example when entering or exiting the half suits.

Transfer isolators

18. These are FFIs or rigid isolators used specifically for transferring materials from one isolator/MSC to another. They are usually battery powered and have docking ports to enable attachment and transfer.
4. Commissioning tests for isolators

19. Whilst MSCs have a defined British Standard\(^8\), isolators currently have no such equivalent (although general guidance is given in Annex A of the Laboratory Operations Standard\(^9\)). From a containment perspective there are many similarities between an isolator and a Class III MSCs and wherever possible, isolators should aim to provide a similar level of protection as afforded by the MSC. Following the testing and maintenance regimes in this document will usually ensure that the expected requirements are met.

20. Commissioning of an isolator should be when it is in its final position within the laboratory/animal room. Factory commissioning only is not acceptable. Commissioning will also be required after any major changes to the isolator such as following dismantling and/or movement and additions/alterations.

21. The following 5 tests are considered essential at commissioning:

- air leaktightness
- leak detection
- filters
- negative pressure
- air change rates

Air Leaktightness

22. The isolator should be leak-tight in order to control exposure to aerosols of biological agents and also gaseous agents (e.g. formaldehyde) used in the disinfection or task process. The MSC British Standard requires a positive pressure hold test be undertaken to determine the air leaktightness of a Class III MSC. The specified overpressure of 500 Pa is unlikely to be practicable in an isolator, particularly film isolators. Instead a practicable alternative could be considered to be a minimum overpressure of 150 Pa. If it was not practicable to overpressure to 150 Pa then an appropriate level as determined by risk assessment and advice from the supplier/manufacturer should be used.

23. A suitable pass criteria would be no more than a 10% loss of pressure when the internal isolator pressure is raised to 150 Pa and held for 30 minutes. One way of assessing this is described in the information box.

Information Box

One method for undertaking positive pressure hold test: the isolator should be assembled with all connections (gloves, gauntlets, half suits etc) made. The extract air duct should be blanked off and the internal pressure raised to 150 Pa by the supply air, which should then be blanked off. For flexible isolators the canopy should be allowed to stretch fully (e.g. for 30-60 minutes) before the 150 Pa pressure is re-established. The supply air should be finally blanked off and the pressure noted at 5 minute intervals over a 30 minute period. Over this time the pressure of 150 Pa should be maintained within a 10% margin.

Leak detection

24. Leak detection should still be undertaken whether or not the isolator passes its pressure hold test. This is because the hold test relates to leak rates, which allows for a certain amount of breach in the containment before the test fails. Leak detection is important in order to identify minor breaches, which could deteriorate during use.

25. The particle tracer test is the recommended method for leak testing given that it is sensitive, relatively easy to perform and uses the same type of equipment as for filter testing. In addition it can be undertaken at the same time as the positive pressure hold test. One way of assessing this is described in the information box.
Information Box

One method for undertaking particle tracer test: place a hot or cold smoke generator capable of generating aerosols of small (e.g. 0.3 micron) particles (e.g. food oil Ondina DL) within the isolator. Raise the internal pressure to minimum of 150 Pa and blank off the supply and extract air. Use a photometer to scan the whole canopy, filter housings etc.

Filters

26. H14 HEPA filters, meeting the relevant standard\textsuperscript{10}, must be in place for the extract air; double HEPA filters must be used if the exhaust air is to be re-circulated into the laboratory and the supply and extract fans must be interlocked to prevent positive pressurisation. In addition, filtration may also be required for supply air, if animal protection is needed.

27. The filters must be seated appropriately and all must be fitted with access ports to allow independent, full face scan testing. The test should meet the standard listed above, with particulate concentrations on both sides of the individual filters measurable.

Negative pressure

28. There will be a broad range of acceptable working pressures, influenced by airflow rates, type of isolator and activity. However, a minimum pressure of 30 Pa below the air pressure of the laboratory should be maintained.

29. As a minimum, positive pressurisation must not occur during normal working practices, which includes when operators enter and leave the isolator (e.g. through half-suits). In addition, the pressure differential should be such that negative pressure is maintained in the event of a foreseeable breach in the isolator envelope.

Air change rates

30. As for negative pressures, the air change rate selected will depend on the type of isolator and the activity. The rapid removal of air potentially contaminated with aerosols reduces the likelihood of accidental exposure and the higher the air change rate the more rapidly aerosols can be removed. In addition, the higher the rate, the lower the waiting time required for disinfection between tasks and following any spills within the isolator.

31. High levels of air change rates are considered inappropriate when housing experimental animals. The Code of Practice for the Animals (Scientific Procedures) Act requires that animals are not kept in draughts with controlled ventilation, noise, humidity and temperature. For example, it is recommended that rodents should have 15-20 changes of fresh air per hour. It is however, generally considered that animal husbandry with infected animals is less likely to generate aerosols of biological agents of high concentration and as such lower levels of air change rates may be appropriate. A minimum of 13 air changes per hour (ach) should be in place.

32. Activities involving laboratory work and pieces of equipment are likely to have the potential to generate aerosols of high concentration and as such high levels (e.g. 40 ach) may be required. In addition, equipment housed within isolators are likely to generate heat hence higher air change rates will assist in dispersing this and controlling heat stress to the canopy.

33. One way of assessing air change rates is described in the information box.

Information Box

One method for calculating air change rate: measure the mean airflow velocity at the exhaust with an anemometer, multiply this by 3600 and divide by the isolator volume.

34. Summary of requirements of tests required at commissioning of isolator
<table>
<thead>
<tr>
<th>Requirement</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air leaktightness</td>
<td>No more than 10% loss of pressure when held at +150 Pa for 30 mins</td>
</tr>
<tr>
<td>Leak detection</td>
<td>Pressure held at +150Pa with no leaks detected</td>
</tr>
<tr>
<td>HEPA filters</td>
<td>Following BS EN 1822:2000</td>
</tr>
<tr>
<td>Negative pressure</td>
<td>At least –30Pa below laboratory pressure</td>
</tr>
<tr>
<td>Air change rate</td>
<td>Minimum of 13 air changes per hour</td>
</tr>
</tbody>
</table>

**5. Maintenance testing of isolators**

35. Isolators are considered as local exhaust ventilation systems under COSHH and as such are required to have regular examination and testing at intervals of not more than 14 months. At containment level 3 and 4 it is recommended that this be done every 6 months, in line with the recommendations for MSCs\(^1\).

36. If it is not practicable to test regularly, for example because an animal study runs for longer than 6 months, then robust ‘in use’ testing should be carried out and documented to ensure the isolator continues to provide an adequate control system. In addition, because isolators tend to be more fragile than MSC’s and more prone to damage during normal operation regular in use testing is essential.

37. The following is a non-exhaustive list of maintenance tests along with a recommended frequency. However, the frequency of testing on an isolator should be individually assessed according to the activity it is being used for.

<table>
<thead>
<tr>
<th>Test</th>
<th>Recommended Minimum frequency</th>
<th>Reasons</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inspection of the canopy, gauntlets/ sleeves and gloves and/or half suit.</td>
<td>Visual In use</td>
<td>Any damage should be dealt with immediately</td>
</tr>
<tr>
<td>Monitoring of manometers</td>
<td>Visual In use</td>
<td>for pressure and air flow rate.</td>
</tr>
<tr>
<td>Housekeeping inside isolator</td>
<td>In use</td>
<td>Should be kept clean and free of unnecessary equipment.</td>
</tr>
<tr>
<td>Isolator surfaces, including the extract ducting.</td>
<td>Visual 6 monthly</td>
<td>examined for defects, cracks and any other damage.</td>
</tr>
<tr>
<td>Integrity of canopy tested</td>
<td>Soft soap, smoke pencil, careful visualisation (see information box) 6 monthly</td>
<td>Repair breaches in the canopy for example, using silicone sealant and/or patching</td>
</tr>
<tr>
<td>Integrity of gauntlet/gloves tested</td>
<td>Soft soap, smoke pencil, careful visualisation 6 monthly</td>
<td>Replace and test new before starting next study</td>
</tr>
<tr>
<td>Test</td>
<td>Recommended Minimum frequency</td>
<td>Reasons</td>
</tr>
<tr>
<td>-------------------------------------------</td>
<td>-------------------------------</td>
<td>-------------------------------------</td>
</tr>
<tr>
<td>Alarm testing</td>
<td>6 monthly</td>
<td>Confirm functioning correctly</td>
</tr>
<tr>
<td>Anti blowback valve</td>
<td>6 monthly</td>
<td>Confirm functioning correctly</td>
</tr>
<tr>
<td>Integrity of filter and housing confirmed.</td>
<td>Full face scanning</td>
<td>6 monthly</td>
</tr>
<tr>
<td>Inlet filter</td>
<td>Anemometer readings</td>
<td>6 monthly</td>
</tr>
<tr>
<td>Calibration of manometer(s).</td>
<td>Calibration</td>
<td>annually</td>
</tr>
</tbody>
</table>

**Information Box**

One or more of the following tests could be used to detect damage and leaks in the canopy, half suits, gauntlets/gloves.

**Soft soap**

Raise the internal pressure to minimum of 150 Pa and apply 2.5% solution of soft soap in tepid, distilled water to all welds, gaskets & joints using a paintbrush. Any leaks should be identifiable by the creation of bubbles.

**Smoke pencil**

Raise the internal pressure to 150 Pa and monitor all welds, gaskets, joints filter housings etc using a smoke pencil. Any leaks should be visualised by deflections to the smoke plume.

**Careful visualisation**

Use a plate to seal off the glove across the port/shoulder ring. Monitor the pressure change within the glove system. If it equilibrates, rate of change can provide direct assessment of leakage.

Place a flexible diaphragm across a glove port/shoulder ring to trap air within the glove system. If there is leakage the air within the glove will be drawn into isolator (and then out through filter), causing the diaphragm to adopt a visible concave shape.
Annex A

Human Factors

It is important that human factors along with other health and safety issues are taken into account when working with isolators. Human factors can be described as the interaction between the user, task and equipment including behaviours and ergonomics. Selection and use of an isolator needs to be driven by the requirements of the operator and not solely by the task. For example, lack of consideration to good ergonomic practice can result in reduced work efficiency, user discomfort, injury and compromised safety.

For example, animal husbandry is a physically demanding occupation involving a great deal of lifting and carrying with the majority of the work time spent standing. Isolators used for animal work need to be selected to meet the requirements of the animals and to achieve the best ergonomic performance for the user.

An assessment of the manual handling risks should be undertaken referring to relevant guidance\textsuperscript{12} to avoid unacceptable strain due to the type of work and/or amount of equipment. It may be necessary to select a different type of isolator or adapt/reduce the cages/equipment designated for the study.

Ergonomic issues to consider before selecting an isolator:

- The physical and psychological characteristics of the users.
- Training and supervision.
- The procedures to be undertaken i.e. whether manual, gross or fine finger dexterity is required.
- The reduction of manual dexterity when working in an isolator.
- Access and egress, especially with half suit isolators where the user has to duck down under the equipment to enter the suit.
- Positioning of equipment inside the isolator so that they are within a zone of convenient reach.
- The best ergonomic performance for example, half suit isolators provide the user with the ability to twist and rotate, whereas FFI's do not.
- The frequency of breaks or changes in task to reduce fatigue.
- Means of communication e.g. if working in half-suit isolators

Before final selection it may be advantageous to put together a ‘mock up’ to ensure all the ports, gloves, half suits etc are situated in the best positions for the task to be carried out and to provide hands-on training of staff.

The nature of research, diagnostic and animal work means that it may be necessary to work out of hours and/or alone. A local risk assessment will identify any specific hazards for lone working and the use of isolators in the containment laboratory.
Annex B
Factors to consider when selecting an isolator to ensure that it is fit for purpose

<table>
<thead>
<tr>
<th>Feature</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Canopy</td>
<td>Sealable without the use of excess sealant; able to fumigate</td>
</tr>
<tr>
<td>Flexible canopy</td>
<td>Flexible and resistant to tear and disinfectant that will be used during procedures and cleaning</td>
</tr>
<tr>
<td>Floor</td>
<td>Hard, resilient surface</td>
</tr>
<tr>
<td>Half suit</td>
<td>Connected to port using double rubber seals (not silicone sealant)</td>
</tr>
<tr>
<td>Services</td>
<td>Supplied through a metal plate (using gas tight sockets) to prevent damage to canopy – electricity required for fumigation</td>
</tr>
<tr>
<td>Filters</td>
<td>H14 HEPA filters on extract; may be needed on supply for animal protection.</td>
</tr>
<tr>
<td>Battery back up</td>
<td>Necessary if housing animals, useful for long-term studies.</td>
</tr>
<tr>
<td>Location of alarms (e.g. negative pressure)</td>
<td>Good visibility and access</td>
</tr>
<tr>
<td>Fans</td>
<td>Supply (if present) and extract interlocked to prevent positive pressurisation.</td>
</tr>
<tr>
<td>Pass box</td>
<td>Separate fan and filters (usually with higher air change rate than main isolator) to assist in passing out samples and fumigating out waste. Usually only on larger half suit isolators.</td>
</tr>
<tr>
<td>Communication</td>
<td>In half suit isolators may require e.g. headsets to allow communication between users</td>
</tr>
<tr>
<td>Waste port</td>
<td>If waste removal required and no pass box present</td>
</tr>
<tr>
<td>Docking port</td>
<td>Double ended for attaching transfer isolator or another isolator/MSC</td>
</tr>
<tr>
<td>Dunk tank</td>
<td>Could be used for sample movement</td>
</tr>
<tr>
<td>Entry and exit to half suit isolators</td>
<td>Thought given to where the various ports are sited and ease of access will prevent injuries to users and damage to the isolator.</td>
</tr>
</tbody>
</table>

References

1 ‘Guidance on the Use, Testing and Maintenance of Laboratory and Animal Flexible Film Isolators’ Advisory Committee on Dangerous Pathogens (ACDP)


3 Control of Substances hazardous to Health Regulations 2002 Approved Code of Practice L5 HSE Books 2005 ISBN 0 7176 2981 3


5 Working Safely with research animals: management of infection risks (1997) HSE Books 1997 ISBN 0 7176 1377 1

6 Code of practice for the housing and care of animals used in scientific procedures http://www.homeoffice.gov.uk/docs/cop_hcasp.html
8 Biotechnology. Performance criteria for microbiological safety cabinets BS EN 12469:2000
9 Biotechnology. Laboratories for research, development and analysis. Guidance for biotechnology laboratory operations BS EN 12741:1999
10 High efficiency air filters (HEPA and ULP) classification, performance, testing and marketing. BS EN 1822:2000
11 The management, design and operation of microbiological containment laboratories Guidance, HSE Books 2001, ISBN 0 7176 2034 4

FURTHER INFORMATION

HSE produces a wide range of documents. Some are available as printed publications, both priced and free, and others are only accessible via the HSE website, www.hse.gov.uk.

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