Biosafety Level 3 (BSL-3) Laboratory Design Standards

January 2020
Introduction

The University of California (UC) is home to some of the world’s most advanced and pioneering biomedical science and clinical research. Consistently ranked among the leaders in the field of Infectious Disease research, the UC is often called upon to provide critical expertise in response to worldwide public health crises. Infectious Disease research involving highly pathogenic and primarily aerosol transmissible agents (i.e., Risk Group 3 agents) has the potential to present significant risk to individuals, the community, and the environment. The ability to safely conduct these research activities is largely dependent upon the highly engineered Biosafety Level 3 (BSL-3) laboratory, the highest-level containment facilities currently operated by the UC. The primary objective of these laboratories is to provide the best possible physical containment of Risk Group 3 agents. Hence, the design and engineering of these laboratories must be maintained at the highest attainable standards.

Several authorities have published standards for the design of BSL-3 laboratories. Development of the UC Biosafety Level 3 Design Standards has incorporated input from several of these sources, including the following:

- CDC Biosafety in Microbiological and Biomedical Laboratories (BMBL), 5th Edition, 2009
- NIH Design Requirements Manual for Biomedical Laboratories and Animal Research Facilities (DRM), 2019
- NIH Guidelines for Research Involving Recombinant or Synthetic Nucleic Acid Molecules (NIH Guidelines), 2016
- National Institutes of Health Biosafety Level 3 Laboratory Certification Requirements, 2006
- Industry Standards and Best Practices

The information presented in these Standards addresses facility design and engineering systems for BSL-3 containment laboratories and incorporates additional design elements to assist campuses in conducting facility risk assessments. It is important to note that an effective BSL-3 program is not only reliant upon the facility and engineering, but also operation and maintenance (O&M) protocols, robust training programs, administrative controls, and standard operating procedures (SOPs).

Application

The BSL-3 Laboratory Design Standards represent the UC requirements for constructing BSL-3 laboratories. These Standards need to be referenced and used by UC design teams including
researchers, architects and engineers, as well as outside contractors. They provide necessary tools and information for BSL-3 planning, construction, and commissioning on UC campuses. While incorporating all applicable regulatory requirements, the standards identify UC requirements, provide explanations of those standards, and document best practices for consideration during the facility risk assessment. This range of acceptable parameters will allow campuses to build each laboratory with a level of features commensurate with the facility risk assessment. These Standards pertain to new facility builds and/or major renovations or retrofits of existing BSL-3 laboratory space. It is not the intent of these Standards to require upgrades to existing BSL-3 spaces, but to provide design standards and guidance when embarking on a new construction or major renovation project.

For the purposes of these Design Standards, major renovations/retrofits are defined as any of the following: 1. Changes to fixed primary containment equipment (e.g. ducted BSCs or any other devices connected to the BSL-3 exhaust system); 2. Changes to major HVAC systems/components (e.g. exhaust fans, air handling units, airflow control valves, isolation dampers, Building Management Systems / Building Automation Systems (BMS/BAS), ductwork, etc.); 3. Changes to floorplans or structural components of the secondary containment boundaries (e.g. walls, ceilings, or attached fixtures); 4. Any other changes to the facility that could impact overall safety, operations, or ventilation system performance. This Standard is applicable to the specific retrofit and not meant to apply to the entire facility. As plans to renovate specific features of an existing facility evolve, it is important to address any identified issues that may affect the safe operations of a BSL-3 laboratory, potentially presenting a risk to the UC community or the environment. These Standards have been approved by the UC systemwide High Containment Laboratory Oversight Committee (HCLOC), and shall be incorporated or referenced in Campus Building Standards.

Note: Facilities required to comply with the Federal Select Agent Program and/or Dual Use Research of Concern may have additional biosafety or biosecurity design features beyond these UC Standards. In addition to the UC Standards, campuses must reference and comply with the applicable Federal regulatory standards when designing Federal Select Agent or Dual Use Research high-containment facilities.

**Campus Involvement**

At the onset of a new BSL-3 laboratory or existing laboratory renovation design project, it is important to consider the involvement of specific campus experts. Consultation with these key campus members may provide insight into critical design flaws that may affect specific aspects of the laboratory’s functionality. These contributions could prove to be exceptionally cost-effective in
terms of both time and resources. Involvement from the following campus members is recommended:

- PI (researcher)/Department Chair
- EH&S Biosafety
- High Containment Laboratory Director
- Campus Design Management Group
- Capital Programs
- UC Building Officials
- Campus ADA Reviewer
- Campus Security or Campus Police Department
- Campus Emergency Response
- Campus Fire Marshal
- Campus Facilities Operations & Management Group

Deviations

Campuses must comply with the UC Biosafety Level 3 Design Standards and Best Practices should be considered when appropriate. When justified by a documented risk assessment, a deviation may be considered. A campus seeking a deviation approval must first complete the **UC BSL3 Design Standards Deviation Request Form, Appendix I**. The completed form and requested documentation shall be presented to the campus High Containment Laboratory Oversight Group (HCLOG) for approval. With campus HCLOG approval, the deviation request will be presented to the UC systemwide HCLOC, for final approval.

Revisions to the Standards

A full review and revision of the UC BSL3 Design Standards shall be conducted every two years, on even years. Between full revisions, the Design Standards may be revised on an as needed basis via continual evaluation and application. Through the use of these Design Standards, any identified gaps, errors, or sections requiring further clarification will be noted and highlighted by local EH&S and Design professionals. Corrections, clarifications and all identified issues shall be presented to the UC systemwide HCLOC for confirmation and approval. Once approved by the HCLOC, updates will be completed and distributed to each location. Revised versions of the Standards must receive approval by the UC systemwide HCLOC prior to implementation.
Plumbing References ....................................................................................................................... 44

**Telecom, Security & Documentation** ................................................................. 47
Telecom .......................................................................................................................... 47
Security .......................................................................................................................... 47
Signage ............................................................................................................................ 49
Enhanced Facilities ........................................................................................................... 50
Verification & Documentation ......................................................................................... 50
Seismic ............................................................................................................................. 51
Telecom, Security & Documentation References ......................................................... 52

**Safety and Decontamination** .............................................................................. 53
Safety .............................................................................................................................. 53
Decontamination ........................................................................................................... 55
Safety and Decontamination References ..................................................................... 57

**BSL-3 Standard Equipment** ................................................................................. 59
Autoclaves ..................................................................................................................... 59
Biosafety Cabinets ......................................................................................................... 60
Flow Cytometry ............................................................................................................ 61
Centrifuge ...................................................................................................................... 61
BSL-3 Standard Equipment References ...................................................................... 62
List of Codes, Standards and Guidelines

- Biosafety in Microbiological and Biomedical Laboratories (BMBL) 5th Edition, 2009
  https://www.cdc.gov/labs/pdf/CDC-BiosafetyMicrobiologicalBiomedicalLaboratories-2009-P.PDF

  The BMBL is published jointly by the U.S. Department of Health and Human Services, the Centers for Disease Control and the National Institutes of Health. The document describes Biological risk assessment, lab practices, safety equipment, and facilities requirements required for designing and operating laboratories at Biosafety Levels (BSL) 1 through 4. Although not a code, certification of BSL facilities is dependent on following the recommendations in this document.

- NIH Design Requirements Manual (DRM) 2019 Rev 1.4: 4/24/19

- National Institutes of Health Biosafety Level 3 Laboratory Certification Requirements, July 2006

- ANSI/ASSE Z9.14-2014 Testing and Performance-Verification Methodologies for Ventilation Systems for Biosafety Level 3 (BSL-3) and Animal Biosafety Level 3 (ABSL-3) Facilities - This document can be purchased at: https://store.assp.org/PersonifyEbusiness/Store/ProductDetails/productId/11562952

- NIH Guidelines for Research Involving Recombinant or Synthetic Nucleic Acid Molecules (NIH Guidelines), 2016

# BSL-3 Architectural

## General

<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. New UC BSL-3 laboratories and designated retrofits must be reviewed for compliance with applicable provisions of the American with Disabilities Act (ADA) and related codes and regulations.</td>
<td>Access to the BSL-3 laboratory and work activities conducted in the facility must be evaluated on a case-by-case basis to determine appropriate accommodations and accessibility. Work with the appropriate units on campus to determine specific ADA design requirements and compliance with Title 24 California Building Code Chapter 11 Accessibility.</td>
<td>1. N/A</td>
</tr>
<tr>
<td>2. Design the laboratory to be easily cleaned and decontaminated.</td>
<td>Smooth walls, floors, and ceilings with finish that is resistant to repeated chemical decontamination.</td>
<td>2. N/A</td>
</tr>
<tr>
<td>3. Keep the BSL-3 laboratory separate from unsecured high traffic areas.</td>
<td></td>
<td>3. Consider locating the facility on an upper floor. Consider location of BSL-3 laboratories in relation to other zones such as BSL-2 support laboratories, offices and break rooms, elevators, loading docks, etc. for effects on laboratory pressurization and airflow.</td>
</tr>
<tr>
<td>4. Provide dedicated anteroom to include space for personal protective equipment (PPE) and waste. Doors are self-closing.</td>
<td>Dedicated anteroom is accessed through self-closing doors that should be physically interlocked with over-ride for emergency use.</td>
<td>4. N/A</td>
</tr>
</tbody>
</table>
5. Provide access to non-containment laboratory support spaces in close proximity to the BSL-3 laboratory, including but not limited to BSL-2 workspace, storage, and break rooms. 13, 14, 15, 16

6. Sealed penetrations (through partitions, walls, floors and ceilings) are required. 17, 18, 19, 20

7. Primary containment devices must be placed in such a way as to minimize impact of room airflow on function/containment. This must be determined in conjunction with the High Containment Laboratory Director (HCLD). 21, 22, 23, 24

5. The needs of the research program and research staff must be considered in design and location of support spaces.

6. Sealed penetrations should be visible for ease of inspection and verification.

7. Primary containment equipment such as BSCs, shall be placed in the room so that containment is not impacted by supply diffusers, exhaust grilles, doors, traffic flow, or other equipment.

6. N/A

7. Consider Appendix A of the DRM and Appendix A of the BMBL for exact distances from features and for placement in a space.

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**Containment Barrier**

<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
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<tbody>
<tr>
<td>8.</td>
<td>Partitions must be designed to resist damage from differential air pressure surges during HVAC testing, and to withstand water and moisture. Interior partitions (walls) must extend slab to slab. 25</td>
<td>N/A</td>
</tr>
<tr>
<td>9.</td>
<td>Construct BSL-3 containment barrier with steel studs, sheet rock and appropriate backing material. 26, 27</td>
<td>Frame partition assemblies shall include gypsum board that is selected and detailed to be appropriately impact, moisture and water resistant. Standard gypsum board is not acceptable. Light-gauge steel studs used for partition framing shall be 18-gauge minimum thickness.</td>
</tr>
</tbody>
</table>
Screw spacing in gypsum wallboard assemblies shall not exceed 152 mm (6 in.). Structural adequacy shall be verified, and additional lateral reinforcement provided if required. All partitions that do not extend to the underside of the structure shall be capped. Concrete masonry units (CMU) walls shall include masonry units utilizing fine sand aggregate or ground face to provide an appropriate substrate for block filler and epoxy paint. Voids in CMU partitions shall be sealed above the ceiling.

### Anterooms

<table>
<thead>
<tr>
<th>Standard</th>
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</tr>
</thead>
<tbody>
<tr>
<td>10. Anteroom doors must be self-closing and have a mechanism for interlocking. Mechanically interlocked doors must have an emergency override.</td>
<td>10. N/A</td>
<td>10. N/A</td>
</tr>
<tr>
<td>11. Anterooms must be appropriately sized for program requirements based on risk assessment and considerations for future expansion of research focus. Anterooms must be sized to meet local Fire Code requirements.</td>
<td>11. Provide anteroom large enough to allow for a change bench, storage shelves for clean PPE, laundry receptacle and waste container.</td>
<td>11. Anteroom should also include space for a logbook and wall calendar. If the anteroom will be the main entryway for equipment, size appropriately for largest piece of equipment, including door sizes.</td>
</tr>
<tr>
<td>Standard</td>
<td>Explanation</td>
<td>Best Design Practice</td>
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</tr>
<tr>
<td>12. Doors must have door locks access controls that allows for fast and efficient removal of access clearance, such as card keys. Hard key access must be off master and provided as a backup to electronic lock systems. Electronic locks must fail secure with a bypass for egress.</td>
<td>12. Electronic lock systems shall be linked to restricted remote access management software. Adhere to any local/UC policy (UCOP IS-3 policy)</td>
<td>12. Consider including biometric or PIN devices for locks on doors access control based on risk assessment.</td>
</tr>
<tr>
<td>14. Door must be welded steel or fiberglass constructed of sufficiently durable material (e.g.: stainless steel, FRP, or epoxy-painted welded steel) with welded steel frames, and large enough for passage of equipment.</td>
<td>14. Wooden doors and frames are not permitted.</td>
<td>14. Provide 42 in. single leaf doors.</td>
</tr>
<tr>
<td>15. Doors shall be seamless with no top or bottom recesses. Cutouts in doors and frames shall be sealed to enhance sanitation and resist air infiltration. Doors and door hardware shall be free of sharp edges. Door hardware shall be easily operated with PPE.</td>
<td>15. N/A</td>
<td>15. N/A</td>
</tr>
<tr>
<td>16. Door sweeps must be provided for pest control, but must be adjustable to facilitate inward airflow movement.</td>
<td>16. Broom sweeps, no rubber sweeps.</td>
<td>16. N/A</td>
</tr>
<tr>
<td>17. Provide stainless steel kick plates.</td>
<td>17. N/A</td>
<td>17. N/A</td>
</tr>
<tr>
<td>18. Vision panels must be in all interior doors</td>
<td>18. Vision panels in exterior doors should be</td>
<td>18. N/A</td>
</tr>
</tbody>
</table>
unless prohibited for programmatic reasons. 44, 45

sealed and rated for fifteen-minute forced entry break resistance.

19. Access to the laboratory must be through two interlocked doors; provide interlock over-ride for emergency use. 46, 47, 48

19. Egress only doors may be single door configuration. No hardware on outside of the door.

19. N/A

20. Door swing direction must be determined in discussion with EH&S and Fire Marshal. 49

20. Doors should swing in the direction of airflow (clean to dirty).

20. N/A

21. No pocket doors, bifold or sliding doors. 50, 51

21. N/A

22. Doors serving locations that require directional airflow shall be configured to allow for sufficient air movement at the undercut to achieve proper operation without excessive pull or closure pressure. 52

Airflow through undercut doors allows for containment airflow and plays a role in establishing operating pressure differential between spaces, which requires verification of height and airflow to ensure proper cascading pressurization control of the facility.

22. N/A

Windows

<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td>23. All windows must be appropriately sealed and non-operable. 53, 54, 55, 56, 57</td>
<td>23. N/A</td>
<td>23. Windows on perimeter walls or doors should be considered only after a security assessment.</td>
</tr>
<tr>
<td>24. Window frames must be fully welded and sealed. 58 • Provide laminated and tempered glass at all windows</td>
<td>24. N/A</td>
<td>24. If windows are located on perimeter walls, they should be appropriately glazed, as described in the DRM.</td>
</tr>
<tr>
<td>25. Sills must be sloped for ease of cleaning. 59</td>
<td>25. N/A</td>
<td>25. N/A</td>
</tr>
<tr>
<td>Floors</td>
<td>Standard</td>
<td>Explanation</td>
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</tbody>
</table>
| 26. Carpets and rugs are not permitted.  
63, 64 | 26. N/A | 26. N/A |
| 27. Provide monolithic seamless welded sheet vinyl or troweled-on epoxy floors with integral coved base. Floor must be chemical resistant and impermeable to liquids.  
65, 66, 67, 68, 69  
• Floors shall have a slip resistant finish. | 27. N/A | 27. N/A |
| 28. Floors of mechanical rooms and interstitial levels located above biocontainment areas shall be designed to prevent leaks. Penetrations through the floor shall be protected by raised curbs or sleeves.  
70, 71 | 28. N/A | 28. Floors of mechanical rooms and interstitial levels located directly above biocontainment areas should be marked to indicate the location of rooms and utility distribution below where possible. |

<table>
<thead>
<tr>
<th>Base</th>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
</tr>
</thead>
</table>
| 29. Floors shall be monolithic with an integral coved base 6 inches’ minimum height, sealed to the wall finish.  
72, 73 | 29. N/A | 29. N/A |

<table>
<thead>
<tr>
<th>Walls</th>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>30. Walls shall be durable, monolithic, and resistant to chemicals and disinfectants. Walls shall be sealed to the base, ceiling, doorframes, cover plates, and all other</td>
<td>30. N/A</td>
<td>30. Corner guards and bumper rails should be provided to protect wall surface in high traffic/impact areas.</td>
<td></td>
</tr>
</tbody>
</table>
openings and penetrations. Epoxy paint or panelized composite systems are standard. Wall construction and materials must be selected to ensure compatibility with finish systems, and to provide a smooth, void-free substrate. 74, 75, 76

31. Suite walls must be full height, extending to the structural deck above. 77, 78

### Ceilings

<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>32. Finish monolithic ceiling with a durable moisture resistant substrate such as epoxy paint that is resistant to all chemical and cleaning agents that may be used. 79, 80, 81, 82, 83</td>
<td>32. Ceilings shall be monolithic, seamless construction. Ceiling systems shall be designed to resist damage from deflection caused by differential air pressure surges that may occur during HVAC fan-failure testing. Ceiling material and support systems shall be designed to be moisture and sag resistant. • Ceilings shall be durable and resistant to moisture, wash-downs and pressurization. Ceilings shall be monolithic, seal to the walls, and with sealed access panel, lights, diffusers, and other ceiling-mounted devices. Epoxy painted gypsum board or panelized composite systems are standard. Gypsum board shall not be standard wall board, but must be specified and detailed to be appropriately moisture and sag resistant. Acoustical tile ceiling systems are not acceptable.</td>
<td>32. N/A</td>
</tr>
</tbody>
</table>
33. The ceiling must be high enough over BSCs to allow appropriate duct or thimble connections. Allowances for maintenance access must be included. Clearances must be per NSF49. 84

<table>
<thead>
<tr>
<th>Access Panels</th>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>34. Minimize access panels. Any access panels must be gasketed and latched to allow for room decontamination. Access panels must be piano type hinged. 85, 86</td>
<td>34. N/A</td>
<td>34. Locate access panels outside the BSL-3 suite.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Finishes</th>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
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<tbody>
<tr>
<td>35. The interior finishes of a BSL-3 shall form a durable, monolithic, impermeable enclosure. Mock-ups of all finishes shall be provided for review and approval, and as a basis for acceptance of the final installation. The mock-ups shall be constructed in the same conditions, and using the same materials and techniques as the final installation. The mock-ups shall include all typical conditions, including sealants, transitions between materials, inside and outside corners. 87, 88, 89</td>
<td>35. N/A</td>
<td>35. BSL-3 wall and ceiling finishes, unless factory finished, should have high performance reinforced multi-coat resinous paint finish (reinforced epoxy paint or equivalent). Paint applicators should be Society for Protective Coatings (SSPC) Coating Application Specialist (CAS Level II) Certified, and should be trained and approved by the paint manufacturer. All high performance resinous paint applications should be inspected by an independent third party Coating Inspector Program (CIP) level 3 certified inspector.</td>
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</table>
36. All finishes in the BSL-3 laboratory shall be chemical resistant and resistant to decontamination agents.

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<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>37.</td>
<td>Laboratory furniture must be sturdy and capable of supporting anticipated loads and uses. 90</td>
<td>37. N/A</td>
</tr>
<tr>
<td>38.</td>
<td>Spaces between benches, cabinets and equipment must be accessible for cleaning. 91, 92</td>
<td>38. Fixed millwork, casework, and countertops shall either be sealed to walls and floors or positioned with sufficient spacing to facilitate cleaning and minimize harborage of pests.</td>
</tr>
<tr>
<td>39.</td>
<td>Bench tops must be impervious to water and resistant to heat, organic solvents acids, alkalis, and other chemicals. 60, 61, 62, 93, 94</td>
<td>39. Bench tops should be epoxy, stainless steel, or phenolic resin.</td>
</tr>
<tr>
<td>40.</td>
<td>Chairs must be covered with non-porous chemical and decontaminant resistant material. 95, 96</td>
<td>40. Consider ergonomic needs when choosing chairs. Adhere to any local/UC policy.</td>
</tr>
<tr>
<td>41.</td>
<td>Shelving standards shall be open fronted or otherwise detailed to allow for full sanitation. Slotted standards with inaccessible concealed areas are not allowed. 97</td>
<td>41. N/A</td>
</tr>
<tr>
<td>42.</td>
<td>Cabinets shall be designed without inaccessible areas that cannot be easily sanitized or disinfected. Joints, holes, and</td>
<td>42. N/A</td>
</tr>
</tbody>
</table>
anchors shall be sealed.  

43. Casework finishes shall be resistant to agents used for decontamination. Stainless steel, phenolic resin, or other corrosion-resistant materials shall be used in damp or corrosive environments.  

44. Minimize cabinet space in the BSL-3 laboratory. Movable tables, mobile base units, and cantilevered bench tops shall be used whenever possible. Where fixed casework components must be used, they must be set on monolithic bases to facilitate installation of integral cover flooring base. Void areas behind fixed case work must be sealed to walls. All fixed items must be sealed to the floor, wall, and adjacent items. 

Sealing

<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td>45. Refer to NIH DRM Appendix L, Sealant Table for areas to be sealed and sealant type.</td>
<td>45. N/A</td>
<td>45. N/A</td>
</tr>
<tr>
<td>46. Doorframes, hinges and latches must be sealed.</td>
<td>46. N/A</td>
<td>46. N/A</td>
</tr>
<tr>
<td>47. Ductwork must be sealed where it penetrates the barrier. Supply and exhaust grills must be sealed where it meets barrier surface.</td>
<td>47. N/A</td>
<td>47. N/A</td>
</tr>
</tbody>
</table>
48. All windows in the laboratory must be sealed. 103, 104

49. All penetrations into and through partitions, floors, and ceilings shall be sealed to enhance sanitation, facilitate gas and vapor decontamination, and resist air infiltration. Piping, ductwork, electrical boxes, and other penetrating items shall be firmly anchored to resist movement that could damage seals. Penetrations shall be visible for inspection and maintenance. 105, 106, 107

50. Seams between walls, floors, and ceilings and between all dissimilar materials shall be fully sealed. Sealant at movement joints shall be applied after installation of high-performance finishes to resist cracking. 108, 109

51. Sealant shall be applied in a uniform, smooth, and continuous manner, resulting in a finish free of voids, pinholes, sharp edges, or excess sealant. Sealant must be compatible with all material that comes in contact with it, including other sealant. Sealant must have chemical resistance, flexibility, durability, adherence, and other characteristics appropriate for its use. 110
   • Where required, apply sealant before application of fire stopping.
BSL-3 Architectural References

1. The laboratory must be designed so that it can be easily cleaned and decontaminated. BMBL 5th ed. Section IV. Biosafety Level 3.D.3
2. The interior finishes of a BSL-3 facility shall form a durable, monolithic, impermeable enclosure. NIH DRM Section 4.9.5
3. Clean-ability of all surfaces including furniture. NIH BSL-3 Certification Section I.2.A
4. Visual inspection of the Architectural features of the laboratory spaces. ANSI Z9.14 Section 8.3.2
5. The laboratory must be separated from areas that are open to unrestricted traffic flow within the building. BMBL 5th ed. Section IV. Biosafety Level 3.D.1
6. A biocontainment laboratory shall not be accessed from the exterior, but through a lower-risk laboratory area (generally BSL-2) with locks, card readers, and other appropriate security devices. NIH DRM Section 4.9.2.1
7. The laboratory is separated from areas which are open to unrestricted traffic flow within the building. NIH Guidelines Appendix G-II-C-4-a
8. Access to the laboratory is through two self-closing doors. BMBL 5th ed. Section IV. Biosafety Level 3.D.1
9. Movement of staff, materials, equipment, and waste in and out of the containment zone requires the use of a directionally pressurized anteroom to provide separation from areas with unrestricted traffic flow. Anterooms shall be designed to accommodate the storage and donning of required PPE. This Passage shall be arranged with two sets of self-closing doors. The anteroom doors shall be interlocks to prevent simultaneous opening of doors between the outside corridor and containment areas. Entrance interlocks, when present, shall be provided with a manual override for use in case of emergency. NIH DRM Section 2.5.3.5
10. Visual inspection: Access to the laboratory should be through a series of two self-closing interlocked doors. ANSI Z9.14 Section 8.3.2
11. Inspect two self-closing doors. Doors should not open simultaneously. Ensure that doors automatically close and latch. Check function of door interlock systems or facility SOPs as appropriate. Check the distance between entrance door from the anteroom for adequate door operation. Check emergency overrides of door interlocks. ANSI Z9.14 Section 8.3.3
12. Passage through two sets of doors is the basic requirement for entry into the laboratory from access corridors or other contiguous areas. Physical separation of the high containment laboratory from access corridors or other laboratories or activities may be provided by a double-door clothes change room (showers may be included), airlock, or other access facility which requires passage through two sets of doors before entering the laboratory. NIH Guidelines Appendix G-II-C-4-a
13. Food must be stored outside the laboratory area in cabinets or refrigerators designated and used for this purpose. BMBL 5th ed. Section IV. Biosafety Level 3.A.3
14. Personnel support areas required for the safe conduct of laboratory work shall be provided at appropriate locations. NIH DRM Section 2.1.3.3
15. Assess location of BSL-3 laboratories in relation to BSL-2 support laboratories, offices and break rooms, elevators, loading docks, etc. for effects on laboratory pressurization and airflow. NIH BSL-3 Certification Section I.3
16. Appropriate directional airflow between containment support spaces and adjacent areas shall be maintained and verified in accordance with the risk assessment. ANSI Z9.14 Section 8.4.10
17. Seams, walls, floors, and ceiling surfaces should be sealed. BMBL 5th ed. Section IV. Biosafety Level 3.D.3
18. All penetrations into and through partitions, floors, and ceiling shall be sealed to enhance sanitation, facilitate gas and vapor decontamination, and resist air infiltration. Penetrations shall be visible for inspection and maintenance. NIH DRM Section 4.9.6
19. Inspect, & Evaluate Architectural Features for Maintenance, Operations (Finishes, penetrations & caulking integrity such as doors, around
20. Properly sealed laboratory surfaces are essential to maintain controlled directional airflow and ventilation system performance. Room leakage or tightness is also critical when gaseous fumigants are used for decontamination.  

**ANSI Z9.14 Section 8.3.2**

21. BSCs must be installed so that fluctuations of the room air supply and exhaust do not interfere with proper operations. BSCs should be located away from doors, heavily traveled laboratory areas, and other possible airflow disruptions.  

**BMBL 5th ed. Section IV.Biosafety Level 3.D.6**

22. Primary containment equipment such as BSCs and individual ventilated caging (IVC) shall be placed in the room so that containment is not impacted by supply diffusers or exhaust grilles or doors or traffic flow.  

**NIH DRM Section 6.6.1.F**

23. Verify correct placement of biological safety cabinets with respect to supply and exhaust diffusers, doors and traffic patterns.  

**NIH BSL-3 Certification Section II.26**

24. Verify correct placement of primary containment equipment (fume hoods, BSCs, etc.) with respect to air devices, doors, and traffic patterns.  

**ANSI Z9.14 Section 8.3.7**

25. Partitions in a biocontainment facility shall be selected to withstand pressurization, impacts and water or moisture. Partition material and detailing shall minimize differential movement. Interior partitions which are part of the secondary barrier shall extend to and be sealed to the underside of the structure. Additionally, partitions shall comply with all physical security requirements as dictated by a threat risk assessment.  

**NIH DRM Section 4.9.4**

26. Frame partition assemblies shall include gypsum board that is selected and detailed to be appropriately impact and moisture or water resistant. Standard gypsum board is not acceptable. Light-gauge steel studs used for partition framing shall be 18-gauge minimum thickness.  

**NIH DRM Section 4.9.4**

27. Screw spacing in gypsum wallboard assemblies shall not exceed 152 mm (6 in.). Structural adequacy shall be verified, and additional lateral reinforcement provided if required. All partitions that do not extend to the underside of the structure shall be capped. Concrete masonry units (CMU) walls shall include masonry units utilizing fine sand aggregate or ground face to provide an appropriate substrate for block filler and epoxy paint. Voids in CMU partitions shall be sealed above the ceiling.  

**NIH DRM Section 4.9.4**

28. A clothing change room (anteroom) may be included in the passageway between the two self-closing doors.  

**BMBL 5th ed. Section IV.Biosafety Level 3.D.1**

29. An anteroom for clean storage of equipment and supplies with dress-in, shower-out capabilities may be required based on agent summary statement, risk assessment, or applicable local, state, or federal regulations.  

**BMBL 5th ed. Section IV.Biosafety Level 3.D.14**

30. Movement of staff, materials, equipment, and waste in and out of the containment zone requires the use of a directionally pressurized anteroom to provide separation from areas with unrestricted traffic flow. Anterooms shall be designed to accommodate the storage and donning of required PPE. This Passage shall be arranged with two sets of self-closing doors. The anteroom doors shall be interlocks to prevent simultaneous opening of doors between the outside corridor and containment areas. Entrance interlocks, when present, shall be provided with a manual override for use in case of emergency.  

**NIH DRM Section 2.5.3.5**

31. Anteroom doors are far enough apart that large pieces of equipment can be moved in or out.  

**ANSI Z9.14 Section 8.4.3**

32. Passage through two sets of doors is the basic requirement for entry into the laboratory from access corridors or other contiguous areas. Physical separation of the high containment laboratory from access corridors or other laboratories or activities may be provided by a double-door clothes change room (showers may be included), airlock, or other access facility which requires passage through two sets of doors before entering the laboratory.  

**NIH Guidelines Appendix G-II-C-4-a.**

33. Doors must have locks in accordance with the institutional policy.  

**BMBL 5th ed. Section IV.Biosafety Level 3.D.1**
34. A biocontainment laboratory shall not be accessed from the exterior, but through a lower-risk laboratory area, with locks, card readers, and other appropriate security devices. NIH DRM Section 4.9.2.1.A
35. Laboratory doors are kept closed when experiments are in progress. NIH Guidelines Appendix G-II-C-2-a
36. External facility doors must be self-closing and self-locking. BMBL 5th ed. Section IV.Biosafety Level 3.D.1
37. The BMBL requires directional airflow into the containment zone through two self-closing doors. NIH DRM Section 2.5.3.7.E
38. Access doors to the laboratory or containment module are self-closing. NIH Guidelines Appendix G-II-C-4-g
39. Doors shall be stainless steel, fiberglass-reinforced polymer (FRP), epoxy –painted welded steel or another durable material. Frames shall be fully welded, sized for the passage of large equipment, and shall have stainless steel protection plates. Knock-down frames are not permitted. NIH DRM Section 4.9.3.2
40. Doors shall be seamless with no top or bottom recesses. Cutouts in doors and frames shall be sealed to enhance sanitation and resist air infiltration. Doors and door hardware shall be free of sharp edges. Door hardware shall be easily operated with PPE. NIH DRM Section 2.6.2.6
41. An effective integrated pest management program is required. BMBL 5th ed. Section IV.Biosafety Level 3.A.10
42. Door sweeps shall be adjustable to facilitate air movement. NIH DRM Section 4.9.3.2
43. Provide stainless steel armor, kick, mop, and stretcher plates on doors based on door location and room use. Center plates horizontally on the door at a width 2 inches less than the door width. Door plates shall maintain UL rating, and must be specified in the door schedule. NIH DRM Section 4.2.1.6.E
44. Vision panels shall be in all doors unless prohibited for programmatic reasons. NIH DRM Section 4.9.3.2
45. Select Agent Laboratories: Vision panels shall be sealed and rated for fifteen minute forced entry break resistance. NIH DRM Section 4.2.2.8.D
46. Laboratory doors must be self-closing and have locks in accordance with the institutional polices. Access to the laboratory is through two self-closing doors. A clothing change room (anteroom) may be included in the passageway between the two self-closing doors. BMBL 5th ed. Section IV.Biosafety Level 3.D.1
47. Required entrances and exits must be configured in vestibules with interlocking doors and directional airflow to maintain the integrity of the barrier. NIH DRM Section 2.5.0.2.a
48. Access to the laboratory should be through a series of two self-closing interlocked doors. ANSI Z9.14 Section 8.3.2
49. Doors serving biocontainment laboratories ordinarily swing in the direction of air movement (clean to dirty). NIH DRM Section 4.2.2.3.A
50. Pocket doors, bifold doors, and accordion doors are not permitted in NIH biomedical laboratories or animal research facilities. NIH DRM Section 4.2.1
51. Manual surface mounted sliding doors may be allowed under the following conditions: Serving rooms which are low hazard and low occupancy. Serving a room with a non-laboratory function. Not required for fire rating or pressurization. NIH DRM Section 4.2.1
52. Doors serving locations that require directional airflow shall be configured to allow for sufficient air movement at the undercut to achieve proper operation without excessive pull or closure pressure. NIH DRM Section 4.9.3.2
53. All windows in the laboratory must be sealed. BMBL 5th ed. Section IV.Biosafety Level 3.D.5
54. Operable windows are not permitted in NIH research laboratories and ARFs (animal research facilities). NIH DRM Section 4.1.4.A
55. Inspect caulking integrity for non-operable windows. NIH BSL-3 Certification Section I.2
56. Inspect: Windows on containment perimeter wall are non-operable and sealed-in-place type construction. ANSI Z9.14 Section 8.3.2
57. Windows in the laboratory are closed and sealed. NIH Guidelines Appendix G-II-C-4-f
58. All window frames shall be fully welded and sealed. Frames shall be foam filled or otherwise seal to prevent infiltration. When windows are located in the barrier wall, removable glazing stops shall be located on the non-containment side of the barrier. Glazing shall be tempered safety or laminated glass. NIH DRM Section 2.6.2.7

59. All interior window sill shall be sloped, and all windows shall be sealed to ensure ease of cleaning and decontamination. NIH DRM Section 4.1.4.E

60. Bench tops must be impervious to water and resistant to heat, organic solvents acids, alkalis, and other chemicals. BMBL 5th ed. Section IV.Biosafety Level 3.D.4.a

61. Bench tops shall be epoxy, stainless steel, or phenolic resin. NIH DRM Section 4.9.8.E

62. Bench tops are impervious to water and resistant to acids, alkalis, organic solvents, and moderate heat. NIH Guidelines Appendix G-II-C-4-c.

63. Carpets and rugs not permitted. BMBL 5th ed. Section IV.Biosafety Level 3.D.3

64. Carpeting is not permitted in any area of the laboratory, including office areas that can only be accessed by passing through a laboratory. NIH DRM Section 4.4.3.2.C

65. Floors must be slip resistant. BMBL 5th ed. Section IV.Biosafety Level 3.D.3.a

66. Floors must be impervious to liquids. BMBL 5th ed. Section IV.Biosafety Level 3.D.3.a

67. Consideration should be given to the installation of seamless, sealed, resilient, or poured floors, with integral coves. BMBL 5th ed. Section IV.Biosafety Level 3.D.3.a

68. Floors must be resistant to chemicals. BMBL 5th ed. Section IV.Biosafety Level 3.D.3.a

69. Floors shall be durable, slip-resistant and resistant to degradation from chemicals, disinfectants, and decontaminants. Floors shall be monolithic with an integral coved base 6 in. minimum height, sealed to the wall finish. The floor shall extend under all equipment and casework. Heat-welded sheet vinyl is most typically used in BSL-3 laboratories, and epoxy is standard where high durability or load capacity is required. NIH DRM Section 4.9.5.1

70. Floors of mechanical rooms and interstitial levels located above biocontainment areas shall be designed to prevent leaks. Penetrations through the floor shall be protected by raised curbs or sleeves. NIH DRM Section 4.9.5.1

71. Floors of mechanical rooms and interstitial levels located directly above biocontainment areas shall be marked to indicate the location of rooms and utility distribution below where possible. NIH DRM Section 4.9.5.1

72. Consideration should be given to the installation of seamless, sealed, resilient, or poured floors, with integral coves. BMBL 5th ed. Section IV.Biosafety Level 3.D.3.a

73. Floors shall be monolithic with an integral coved base 6 inches minimum height, sealed to the wall finish. NIH DRM Section 4.9.5.1

74. Walls should be constructed to produce a sealed smooth finish that can be easily cleaned and decontaminated. BMBL 5th ed. Section IV.Biosafety Level 3.D.3.b

75. Walls shall be durable, monolithic, and resistant to chemicals and disinfectants. Walls shall be sealed to the base, ceiling, door frames, cover plates, and all other openings and penetrations. Epoxy paint or panelized composite systems are standard. Wall construction and materials must be selected to ensure compatibility with finish systems, and to provide a smooth, void-free substrate. NIH DRM Section 4.9.5.2

76. The integrity of all surfaces, penetrations, and seals on the containment perimeter shall be visually inspected. ANSI Z9.14 Section 8.3.2

77. Interior partitions which are part of the secondary barrier shall extend to and be sealed to the underside of the structure. NIH DRM Section 4.9.4
78. Wall finishes shall be protected from impact and wear utilizing corner guards, crash rails, FRP panels or other methods in vulnerable areas. **NIH DRM Section 4.9.5.2**

79. Ceilings should be constructed, sealed, and finished in the same general manner as walls. **BMBL 5th ed. Section IV.Biosafety Level 3.D.3.c**

80. Ceilings shall be monolithic, seamless construction. Ceiling systems shall be designed to resist damage from deflection caused by differential air pressure surges that may occur during HVAC fan-failure testing. Ceiling material and support systems shall be designed to be moisture and sag resistant. **NIH DRM Section 2.6.2.4**

81. Ceilings shall be durable and resistant to moisture, wash-downs and pressurization. Ceilings shall be monolithic, seal to the walls, and with sealed access panel. Lights, diffusers, and other ceiling-mounted devices. Epoxy painted gypsum board or panelized composite systems are standard. Gypsum board shall not be standard wall board, but must be specified and detailed to be appropriately moisture and sag resistant. Acoustical tile ceiling systems are not acceptable. **NIH DRM Section 4.9.5.3**

82. Verify structural capability of wall and ceiling systems prior to determining leak test pressures. **ANSI Z9.14 Section 8.4.8.2**

83. The interior surfaces of walls, floors, and ceilings are water resistant so that they can be easily cleaned. Penetrations in these surfaces are sealed or capable of being sealed to facilitate decontaminating the area. **NIH Guidelines Appendix G-II-C-4-b**

84. Ceiling height to be coordinated with cabinet requirements to ensure proper airflow and containment within the cabinet. **NIH DRM Section 4.6.1.11.A.8**

85. All utility systems shall be configured to minimize the need for access panels within the containment barrier. **NIH DRM Section 4.9.7**

86. Access panel frames shall be sealed to the wall or ceiling, and fitted with a continuous, gastight door gasket. Panel design shall ensure compression of the gasket around the entire panel door perimeter without discontinuity at the hinge or latch when in the latched position. Access door assemblies shall be stainless steel or another non-coated, corrosion-resistant material. **NIH DRM Section 4.9.7**

87. The interior finishes of a BSL-3 shall form a durable, monolithic, impermeable enclosure. Mock-ups of all finishes shall be provided for review and approval, and as a basis for acceptance of the final installation. The mock-ups shall be constructed in the same conditions, and using the same materials and techniques as the final installation. The mock-ups shall include all typical conditions, including sealants, transitions between materials, inside and outside corners. **NIH DRM Section 4.9.5**

88. BSL-3 wall and ceiling finishes, unless factory finished, shall have high performance reinforced multi-coat resinous paint finish (reinforced epoxy paint or equivalent). Paint applicators shall be Society for Protective Coatings (SSPC) Coating Application Specialist (CAS Level II) Certified, and must be trained and approved by the paint manufacturer. All high performance resinous paint applications must be inspected by an independent third party Coating Inspector Program (CIP) level 3 certified inspector. **NIH DRM Section 4.9.5**

89. Inspect and evaluate finishes. **NIH BSL-3 Certification Section I.2**

90. Laboratory furniture must be capable of supporting anticipated loads and uses. **BMBL 5th ed. Section IV.Biosafety Level 3.D.4**

91. Spaces between benches, cabinets, and equipment must be accessible for cleaning. **BMBL 5th ed. Section IV.Biosafety Level 3.D.4**

92. Fixed millwork, casework, and countertops shall either be sealed to walls and floors or positioned with sufficient spacing to facilitate cleaning and minimize harborage of pests. **NIH DRM Section 4.5.1.4**

93. Bench tops must be impervious to water and resistant to heat, organic solvents acids, alkalis, and other chemicals. **BMBL 5th ed. Section IV.Biosafety Level 3.D.4.a**

94. Bench tops shall be epoxy, stainless steel, or phenolic resin. **NIH DRM Section 4.9.8.E**

95. Chairs must be covered with a non-porous material that can be easily cleaned and decontaminated with appropriate disinfectant. **BMBL 5th ed. Section IV.Biosafety Level 3.D.4.b**

96. Chairs and other laboratory furnishings shall be covered with a non-porous material that is easily cleaned and decontaminated with
appropriate disinfectant. BMBL Section IV.BiosafetyLevel3.D.4.b.

97. Shelving standards shall be open fronted or otherwise detailed to allow for full sanitation. Slotted standards with inaccessible concealed areas are not allowed. NIH DRM Section 4.9.8.A

98. Cabinets shall be designed without inaccessible areas that cannot be easily sanitized or disinfected. Joints, holes, and anchors shall be sealed. NIH DRM Section 4.9.8.B

99. Casework finishes shall be resistant to agents used for decontamination. Stainless steel, phenolic resin, or other corrosion-resistant materials shall be used in damp or corrosive environments. NIH DRM Section 4.9.8.F

100. Movable tables, mobile base units, and cantilevered bench tops shall be used whenever possible. Where fixed casework components must be used, they shall be set on monolithic bases to facilitate installation of integral cover flooring base. Void areas behind fixed case work shall be sealed to walls. All items shall have smooth corners and edges and be free of open joints and voids. All fixed items shall be sealed to the floor, wall, and adjacent items. Welded stainless steel is standard in BSL-3 laboratories where moisture or frequent cleaning is required. NIH DRM Section 4.9.8

101. Spaces around doors should be capable of being sealed for space decontamination. BMBL 5th ed. Section IV.Biosafety Level 3.D.3

102. Spaces around ventilation openings should be capable of being sealed for space decontamination BMBL 5th ed. Section IV.Biosafety Level 3.D.3

103. All windows in the laboratory must be sealed. BMBL 5th ed. Section IV.Biosafety Level 3.D.5

104. Window frames shall be fully welded and sealed. Frames shall be foam filled or otherwise sealed to prevent infiltration. When windows are located in the barrier wall, removable glazing stops shall be located on the non-containment side of the barrier. Glazing shall be tempered safety or laminated glass. NIH DRM Section 2.6.2.7

105. Seams, floors, walls, and ceilings should be sealed. BMBL 5th ed. Section IV.Biosafety Level 3.D.3

106. All penetrations into and through partitions, floors, and ceilings shall be sealed to enhance sanitation, facilitate gas and vapor decontamination, and resist air infiltration. Piping, ductwork, electrical boxes, and other penetrating items shall be firmly anchored to resist movement that could damage seals. Penetrations shall be visible for inspection and maintenance. NIH DRM Section 4.9.6

107. Visual Inspection: Continuous seal between ductwork and room exhaust grille. Verify during initial installation and maintain written/photo documentation for records. ANSI Z9.14 Section 8.3.4

108. Seams, floors, walls, and ceiling surfaces should be sealed. BMBL 5th ed. Section IV.Biosafety Level 3.D.3

109. Seams between walls, floors, and ceilings and between all dissimilar materials shall be fully sealed. Sealant at movement joints shall be applied after installation of high-performance finishes to resist cracking. NIH DRM Section 4.9.6

110. Sealant shall be applied in a uniform, smooth, and continuous manner, resulting in a finish free of voids, pinholes, sharp edges, or excess sealant. Sealant must be compatible with all material that it is in contact with, including other sealant. Sealant must have chemical resistance, flexibility, durability, adherence, and other characteristics appropriate for its use. NIH DRM Section 4.9.6
### Heating, Ventilation & Air Conditioning

<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
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</table>
| 1.       | The laboratory must be designed for ease of maintenance, so that access to critical mechanical equipment is outside containment. | 1. Equipment should be accessible for maintenance, repairs and annual verification.  
• Shutoff valves should be in non-containment spaces.  
• Mechanical HVAC spaces should be on full size interstitial space/floor. |
| 2.       | A BSL-3 environment must have a minimum of 6 air changes per hour. | 2. N/A |
| 3.       | A BSL-3 environment must have supply/exhaust fan interlock. | 3. Consider redundancy by providing both Software and Hardware Interlocks |

1, 2

3, 4

5, 6, 7, 8
4. Gas tight damper and/or low leakage airflow control valves are required to facilitate isolation of individual laboratory rooms/space based on agent summary statement, risk assessment, or applicable local, state or federal regulations. 9, 10


6. Provide a ducted dedicated, single pass exhaust ventilation system. The exhaust fan system shall be redundant (N+1 minimum) and provide directional airflow, from least hazardous area to most potentially hazardous. 12, 13, 14, 15, 16
   • The laboratory exhaust air must not re-circulate to any other area of the building. 30, 31, 32, 33

7. Under normal operations, system shall provide directional airflow and shall be exhausted to outside. 17, 18, 19, 20

4. Gas tight isolation dampers or low leakage airflow control valves allow the facility or laboratory room to become isolated from the rest of the HVAC system without affecting other areas. This feature can be used when performing area decontamination of the room or facility or when removing a large piece of equipment.

5. Adherence to the latest version of energy codes will ensure compliance with local, state and federal requirements.
   • Include local and UC standards

6. BSL-3 exhaust ductwork shall not be shared with non-BSL-3 spaces.

4. On supply and exhaust systems (see DRM reference).

5. N/A

6. Dedicated ventilation system (supply and exhaust) will allow a faster response of the management system by being able to isolate BSL-3 facility without affecting the rest of the building.

7. N/A
8. 100 CFM doorway infiltration. 8. 100 CFM will maintain an average pressure difference of ~ -0.05 in.w.g. Differential pressure range of -0.04 to -0.15 in.w.g. shall be maintained between each pressure zone. In the event that multiple containment zones exist within a laboratory or laboratory suite, sequentially more negative pressure differentials must be established so that the more contaminated spaces are maintained at a negative pressure with respect to less contaminated areas.
   • Smoke test to verify directional airflow

9. A N+1 redundancy is required for High-Containment laboratory rooms. Capacity and size of the make-up air system serving containment devices/equipment shall correspond to 100% of the programmed containment devices/equipment plus future capacity as defined by the campus personnel. 21, 22

9. The air-handling unit must provide enough capacity for future upgrades. Capacity of the cooling system shall include the program cooling demand plus an allowance for future expansion of internal heat gain requirements as defined by the campus personnel.

9. A dedicated air handing system is highly recommended. Consider designs with multiple fans, isolation dampers and controls maybe designed to satisfy the N+1 requirements.

Considerations should be made for design of the HVAC system with regard to constant volume or variable volume.

10. Provide 100% outdoor air supplied to the BSL-3 laboratory that is then exhausted directly. 23

10. Providing outside air will minimize cross-contamination between spaces.

10. Provide dedicated supply (air handling unit and ductwork).

11. Provide digital monitor or analog magnehelic gauge at each door leading to a new pressure zone. 24, 25, 26

11. Pressure monitor must allow laboratory personnel to verify that laboratory is under negative pressure.

11. N/A
• “Ball-In-tube” units are not acceptable

12. Provide audible and visual alarms to notify personnel of air flow disruption. 27, 28

12. Local alarms shall notify laboratory personnel of issues with the facility.

12. Alarm shall report to appropriate personnel and Facilities.

12. Alarm should report to Facilities personnel and Building Management System (BMS) after the appropriate alarm delay set point.

13. A BSL-3 environment shall have air pressure differential of -.05" in.w.g. at each barrier door and laboratory. 29

13. Having a differential pressure of -0.05 in.w.g. provides the containment of pathogens in the case of a release outside the BSC

13. N/A

14. Exhaust air must be HEPA filtered. The Laboratory building exhaust air should be dispersed away from occupied areas and from building air intake locations. 34, 35, 43, 44, 45

14. HEPA filter sterilizes the air before it is discharged into the environment

14. N/A

• Consider other regulatory requirements when volatile chemicals and radioactive materials will be used in B2 cabinets.

15. Exhaust air systems shall be arranged with single fans or multiple manifolded fans designed to achieve N+1 redundancy and maintain the exhaust air system fully operational, at all times. Each fan shall be designed to be capable of being fully isolated while the overall system remains fully operational. 37, 42

15. N+1 configuration provides redundancy in the system in case of a failure.

15. N/A

16. Provide a dedicated exhaust system with double HEPA filters in series for a Class III BSC. 38

16. Class III BSC are commonly used for aerosol procedures involving biological materials with hazardous chemicals or radioactive materials.

16. N/A
17. Exhaust hoods on the "dirty" side of double door autoclaves shall be connected to the BSL-3 exhaust air system. 39

18. HEPA filter housings shall have gas-tight isolation dampers, decontamination ports, and/or bag-in/bag-out (with appropriate decontamination procedures) capability. 46, 47

19. The HEPA filter housing shall allow for leak testing of each filter and assembly. 48, 49, 50

17. Connecting exhaust hood to BSL-3 exhaust system will minimize contamination of non-containment areas in case of failure.

18. This allows for HEPA filter housing decontamination before any repairs and/or maintenance is required.
   - Isolation dampers, decontamination ports and bag in-bag out should be located in areas that are easily accessible with enough room for maintenance/filter change-outs.
   - HEPA filter housing shall be welded stainless steel construction.
   - Each HEPA filter shall be capable of in situ decontamination and full face filter scanning.

19. HEPA filters must be checked for leaks on an annual basis.

17. N/A

18. N/A

19. N/A
HVAC References

1. Laboratory must be designed so that it can be easily cleaned and decontaminated. [BMBL 5th ed. Section IV. Biosafety Level 3.D.3]
2. The interior of walls, floors and ceilings are water resistant so that they can be easily cleaned. [NIH Guidelines Appendix G-II-C-4-b]
3. A BSL-3 environment shall have 6 air changes per hour minimum. [NIH DRM Section 6.1.11.1.A]
4. Table 1. Airborne contamination removal and air change rates. [CDC Infection Control. Environmental Infection Control Guidelines. Part IV appendix]
5. A BSL-3 environment shall have: Supply/Exhaust fan interlock. [NIH DRM Section 7.7.1.B]
6. Ensure that interlock between supply and exhaust is operational. [NIH BSL-3 Certification Section II.6]
7. Verify Control Systems/Fail Safe Operation [ANSI Z9.14 Section 8.3.9]
8. The air handling system shall be designed such that under failure conditions, the airflow will not be reversed and periodic verification. [NIH DNA Guidelines Appendix G-II-C-4-i]
9. Gas tight dampers to facilitate laboratory isolation may be required based on agent summary statement, risk assessment, or applicable local, state, or federal regulations. [BMBL 5th ed. Section IV. Biosafety Level 3.D.14]
10. Bubble-tight isolation damper shall be provided between the room supply air terminal, the room supply air diffuser, and between the room exhaust grille and room exhaust air terminal. [NIH DMR Section 6.6.9.B]
12. A ducted air ventilation system is required. This system must provide sustained directional airflow by drawing air into the laboratory from "clean" areas toward "potentially contaminated" areas. [BMBL 5th ed. Section IV. Biosafety Level 3.D.9]
13. A BSL-3 environment shall have single pass airflow. [NIH DRM Section 2.5.3.7.E, 6.6.7]
14. Ensure single pass airflow. Directional airflow must be established from clean areas into contaminated areas. In the event that multiple containment zones exist within a laboratory or laboratory suite, sequentially more negative pressure differentials must be established so that the more contaminated spaces are maintained at a negative pressure with respect to less contaminated areas. [NIH BSL-3 Certification Section II.14]
15. The directional airflow shall be verified in normal and failure modes as well as equipment such as BSC's, fume hoods, filtration systems, etc. connected to the ventilation system. [ANSI Z9.14 Section 8.4.1, 8.4.2]
16. A ducted exhaust air ventilation system is provided. The system creates directional airflow that draws air into the laboratory from uncontaminated spaces surrounding the laboratory. [NIH Guidelines Appendix G-II-C-4-i]
17. At BSL-3..., exhaust laboratory air must be directly exhausted to the outside since it is considered potentially contaminated. This concept is referred to as a dedicated, single-pass exhaust system. [BMBL 5th ed. Appendix A Section VI. Building Exhaust]
18. A BSL-3 environment shall have a supply system independent and separate from the remainder of the building. [NIH DRM Section 6.1.8.1]
19. BSL-3 laboratory spaces shall be provided with dedicated supply air systems, which do not serve any other laboratory spaces outside the containment laboratory. [NIH DRM Section 6.6.2.A]
20. Minimal acceptable conditions for ventilation system performance in BSL-3 facilities include: 1) during normal facility operations, the maintenance of air movement from areas that are not contaminated by any biological hazards towards areas that may be progressively
more contaminated inside the laboratory. **ANSI Z9.14 Section 5.2**

21. Air-handling units (AHU) shall be designed to provide N+1 reliability and maintain 100% capacity in the event of a lead component failure. Multiple parallel air-handling units shall be provided to operate simultaneously to meet full load conditions. Each AHU and its related components shall be capable of total isolation by the use of isolation dampers located upstream and downstream of each air-handling unit. **NIH DRM Section 6.1.8.1.B, 6.6.9.A**

22. Capacity and size of the make-up air system to serve containment devices/equipment shall correspond to 120% of the programmed containment devices/equipment. Capacity of the cooling system shall include the program cooling demand plus an allowance for 20% future expansion of internal heat gain requirements. **NIH DRM Section 6.2.1**

23. Use of 100% outdoor air to provide all the room air to be exhausted through laboratory spaces and laboratory containment equipment; (2) Size the exhaust air system to handle the simultaneous operation of all laboratory spaces and all laboratory containment equipment, and (3) Directing airflow from low hazard areas to high hazard areas at all times. Air supplied to the corridor and adjacent clean spaces shall be exhausted through the laboratory to achieve effective negative pressurization. **NIH DRM Section 6.1.13.1.A**

24. Laboratory personnel must be able to verify directional airflow. A visual monitoring device, which confirms directional airflow, must be provided at the laboratory entry. **BMBL 5th ed. Section IV.Biosafety Level 3.D.9.a**

25. A BSL-3 environment shall have pressure differential monitors. **NIH DRM Section 7.7.1.B, 7.7.2.D**

26. Visual Inspection: A means for users to verify airflow direction should be present. Monitor should be provided, both at the outer entry of the BSL-3 suite and at interior doors, based on the risk assessment. **ANSI Z9.14 Section 8.3.3**

27. Audible alarms should be considered to notify personnel of airflow disruption. **BMBL 5th ed. Section IV.Biosafety Level 3.D.9.a**

28. The BAS shall provide differential pressure monitors on classified spaces to indicate the room differential pressure and shall alarm when the pressure goes beyond adjustable thresholds and time durations. **NIH DRM Section 13.9.4**

29. A BSL-3 environment shall have air pressure differential of -0.05" w.g. at each barrier door and laboratory. **NIH DRM Section 6.1.15.C**

30. The laboratory exhaust air must not re-circulate to any other area of the building. **BMBL 5th ed. Section IV.Biosafety Level 3.D.9.b**

31. Laboratory and animal facility air shall not be recirculated to another space or facility to prevent migration of chemical fumes or airborne pathogens and prevent cross-contamination between spaces. **NIH DRM Section 6.1.8**

32. Visual inspection of the exhaust air systems including exhaust stack location and discharge velocities, code distance from exhaust stack and air intakes operable windows, etc., the exhaust stack height, the presence and operation of the interlock between the supply and exhaust, and the exhaust air filtration. **ANSI Z9.14 Section 8.3.4**

33. The exhaust air is not recirculated to any other area of the building, is discharged outside, and is dispersed away from the occupied areas and air intakes. **NIH Guidelines Appendix G-II-C-4-i**

34. The laboratory building exhaust air should be dispersed away from occupied areas and from building air intake locations or the exhaust air must be HEPA filtered. **BMBL 5th ed. Section IV.Biosafety Level 3.D.9.c**

35. Construction documents shall include the design of exhaust stack height and discharge air velocity characteristics to overcome the building cavity boundary and avoid re-entrainment of exhaust. Stacks shall be shown as part of the architectural design and the design rationale shall be described in the early design reports. In general, exhaust stacks shall be designed to meet the following requirements - discharge shall be a minimum of 3 m (10 ft.) above the roofline and any roof element within a horizontal distance of a 4 m (13 ft.) radius, upward velocity shall be a minimum of 15 m/s (3,000 fpm) at the point of discharge. Reentry calculations may dictate higher discharge velocities, safety concerns shall always take precedence over aesthetics, and manifolds for multiple exhaust fans shall have separate exhaust stacks for each fan to avoid having positive pressure ductwork on the discharge side of fans not operating. **NIH DRM Section 6.2.3.C**
36. Ensure that discharge of local exhaust ventilation (LEV) devices is removed from air intakes to prevent re-entrainment. NIH BSL-3 Certification Section II.40

37. Exhaust air systems shall be arranged with multiple manifolded fans designed to achieve N+1 redundancy and maintain the exhaust air system fully operational, at all times. Each manifolded fan shall be designed to be fully isolated while the overall system remains fully operational. In the case of single fan systems, in addition to the main fan, a standby fan shall be provided. The A/E shall review redundancy requirements for each particular system with the program user and the NIH/DOHS. NIH DRM Section 6.1.22.2

38. Exhaust air from a Type B BSC shall be a dedicated system with double HEPA filters in series. NIH DRM Section 7.5.15.A

39. Exhaust hoods on the "dirty" side of pass through autoclaves are connected to the BSL-3 exhaust air system. NIH DRM Section 4.9.10

40. At BSL-3…, exhaust laboratory air must be directly exhausted to the outside since it is considered potentially contaminated. This concept is referred to as a dedicated, single-pass exhaust system. BMBL 5th ed. Appendix A Section VI.Building Exhaust

41. A BSL-3 environment shall have an exhaust system independent and separate from the remainder of the building. NIH DRM Section 6.6.7.A

42. A BSL-3 environment shall have: Redundant exhaust fans (N+1). NIH DRM Section 6.6.1.D

43. The laboratory building exhaust air should be dispersed away from occupied areas and from building air intake locations or the exhaust air must be HEPA filtered. BMBL 5th ed. Section IV.Biosafety Level 3.D.9.c

44. Exhaust air HEPA filtration is recommended to eliminate the possibility of re-entrainment of BSL-3 exhaust air into the intake air and to filter highly infectious agents and pathogens that may cause risk to the environment. NIH DRM Section 6.6.8.B

45. The exhaust air is not recirculated to any other area of the building, is discharged outside, and is dispersed away from the occupied areas and air intakes. NIH Guidelines Appendix G-II-C-4-i

46. HEPA filter housings should have gas-tight isolation dampers, decontamination ports, and/or bag-in/bag-out (with appropriate decontamination procedures) capability. BMBL 5th ed. Section IV.Biosafety Level 3.D.9

47. HEPA filter housing shall be welded stainless steel construction. Each HEPA filter shall be capable of in situ decontamination and full face scanning. NIH DRM Section 6.6.8.D

48. The HEPA filter housing should allow for leak testing of each filter and assembly. BMBL 5th ed. Section IV.Biosafety Level 3.D.9

49. The housing shall be accessible and space provided for filter change-outs. NIH DRM Section 6.6.8.D

50. HEPA filter installations shall be visually inspected. HEPA filtration shall be tested at least annually. ANSI Z9.14 Section 8.3.4

51. Final HEPA filtration of the laboratory exhaust air may be required by the agent summary statement, risk assessment, or applicable local state or federal regulations. BMBL 5th ed. Section IV.Biosafety Level 3.D.14

Exhaust air HEPA filtration is recommended and each particular system/application shall be reviewed with the user and EHS. NIH DRM Section 6.6.8.B
### Electrical

<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>1.</strong> All electrical penetrations shall be sealed airtight internally at the containment boundary.</td>
<td>Sealed penetrations are required for decontamination and directional airflow</td>
<td>1. N/A</td>
</tr>
<tr>
<td>• Refer to NIH DRM for areas to be sealed and sealant type.</td>
<td>• J box is sealed and in wet areas consider using an outlet cover</td>
<td></td>
</tr>
<tr>
<td><strong>2.</strong> Electrical faceplates and devices shall be able to withstand routine decontamination, as well as, after spills, splashes, or other potential contamination.</td>
<td>BSL-3 laboratories must be able to be area decontaminated without affecting the electrical components</td>
<td>2. Utilize stainless steel cover plates for switches and receptacles in rooms that will utilize vaporized paraformaldehyde or chlorine dioxide gaseous decontamination protocols.</td>
</tr>
<tr>
<td>• Use of foam and rubber shall be minimized.</td>
<td></td>
<td>• Based on decontamination methods, consider providing electric relays for control of power outlets for disinfecting agent generation equipment, mixing equipment, and neutralizing agent generation equipment (if required by disinfecting agent), to be controlled from outside the room(s). Provide power to dehumidification equipment if required by disinfection agent protocol.</td>
</tr>
<tr>
<td>• Wall-mounted electrical receptacles and switches and ceiling-mounted receptacles shall be hinged, self-closing, weatherproof and stainless steel SS-316 with rounded edge cover plates with neoprene closed-cell gasket material for all wall devices.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>3.</strong> Conduits for all systems - Conduit applications in BSL-3 facilities are as follows:</td>
<td>Threaded rigid conduit with sealed penetrations at containment barrier provides a long term and durable pathway that meets the airtight</td>
<td>3. N/A</td>
</tr>
<tr>
<td>• Conduit type: use rigid galvanized steel (RGS) conduit with threaded</td>
<td></td>
<td></td>
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</tbody>
</table>
fittings in all BSL-3 areas

- Seal-off: provide seal-off fittings when conduits exit defined BSL-3 perimeter
- SMR: use of surface metal raceway systems (SMRs) is not allowed in BSL-3 areas

4. BSL-3 facilities - All components of the HVAC system (supply and exhaust), alarms, emergency lighting and laboratory outlets for essential equipment (Biological Safety Cabinets, freezers, autoclave, etc.) shall be on a backup power system. 12, 13, 14, 15
- All BSL-3 equipment and controls on backup power shall be supplied backup power through a closed transition transfer switch. BAS controllers and BSCs shall have uninterruptible power supply (UPS) with a run time of at least 120 seconds
- UPS shall be installed such that it is easy to monitor and replace on a regular basis. Backup power to the laboratory and associated systems shall be on dedicated circuits from the generator.
- All HVAC equipment must be capable of auto-restart after a power failure.

5. Provide emergency power for the following essential equipment: 16, 17
   1. HVAC

4. BSL-3 laboratories shall have emergency power for critical components.

5. Emergency Power shall connect the following loads to the standby electrical systems:

4. Emergency Generator - Provide a local generator dedicated to the facility to provide emergency/standby power. Consider providing 100% generator backup.
2. Alarms
3. Emergency lighting
4. BSCs
5. Freezers
6. Incubators
   • During the design phase, review equipment with facility users to determine emergency power backup needs.

6. The emergency power system shall be designed to meet applicable codes and standards. \(^{18, 19, 20}\)

7. Life safety loads shall be wired separately from normal powered, legally required and optional standby loads. \(^ {21}\)

8. Provide UPS power to all alarms and electronic key access systems. During the design phase, review with facility user’s additional equipment that may require UPS power backup. \(^ {22, 23, 24}\)
   • Biological Safety Cabinets, refrigerators and freezers should be

   • Heating and cooling units provided for critical support function
   • Receptacles serving selected equipment identified as critical by the program
   • General lighting in laboratories
   • Critical control and containment equipment compressed air system
electrical system:
   • Sterilizers
   • Cage wash equipment for remote locations
   • Other loads as required by the program based on risk assessment

6. Emergency electrical service size shall be adequate to meet the current and future emergency electrical demand, and applicable codes and standards

6. N/A

7. Life Safety Loads may include all or some of the following:
   • Emergency egress lighting
   • Egress signage
   • Communications systems (including PA systems
   • Fire alarm and mass notification systems
   • Self-contained battery-powered lighting at generator set location
   • Fire-suppression systems (fire pumps, jockey pumps, compressors, valves, etc.)

7. N/A

8. A central UPS system or a number of local UPSs may be required to backup all building wide low voltage systems that are essential for containment operation, safe shutdown of the facility and for critical BSL-3 specific loads.

8. N/A
9. J boxes shall be cast and/or sealed airtight. 25, 26, 27

9. Provide cast boxes with external mounting provisions, external hub, and gasketed device cover plates
   • Sealing: Provide a 25 mm (1 in.) barrier of silicone caulk around the wire within a device box hub.
   Provide a continuous bead of caulk between the device box and the adjacent surface. Provide a continuous bead of caulk around the device cover plate and the adjacent surface.

9. Consider double gang J boxes
   • Type and Depth: All boxes should be double gang type; the box depth should be at least the next larger than minimum size required per code.

10. Panel boards must be located outside the contained space. Circuit breakers must be labeled. 28

10. Electrical equipment that requires service shall not be installed within a containment area.

10. N/A

11. Provide GFCI outlets at sinks for electrically operated pre-mixing valves.

11. N/A

11. N/A

12. Power Wiring Insulation shall be compatible with sealing compound (sealing compound non-deleterious to insulation), using THW, THWN, THHN/THWN, or XHHW, with minimum size #12 wire, except #10 wire for special purpose receptacles. The A/E shall coordinate requirements for security wiring, on a per project basis with Facilities and EH&S. 29, 30, 31

12. Verify that manufacturers wiring jacketing is chemically compatible with sealing compounds.

12. N/A

13. The electrical equipment shall mirror the N+1 configuration of the critical

13. The separate distribution panels shall be powered from separate transfer

13. N/A
mechanical systems. Where redundant mechanical system components are provided, the electrical system shall be designed to limit the impact of the loss of a single electrical component. (e.g. if 2 fans are provided in a N+1 configuration, each fan should be provided from a separate distribution panels so that the N units of the equipment can continue to operate during failure of any single electrical system component.)

14. Confirm electrical infrastructure minimum requirements for reliability and redundancy with approving authorities.

14. Redundant electrical services eliminate single points of failure and correspond to critical mechanical system redundancies. Coordinate with campus and utility provider for service infrastructure for high reliability. Review with grant officers, principal investigators, facility operations and authorities having jurisdiction to confirm requirements.

• Downstream distribution for BSL-3 projects from switchgear for critical areas, such as mechanical support rooms with redundant motors in each set, shall consist of pairs of distribution switchboards/panel boards, each fed from a separate side of the switchgear, to supply approximately half of each set of motors. This would include supplying packaged units with multiple motors with separate switches. The A/E shall evaluate and minimize single points of failure for all projects.
feeders for each motor, where possible.

15. Wall penetrations shall be prepared and sleeved. Wall penetrations through fire rated walls shall be avoided. Where necessary, penetrations shall be in accordance with NFPA and NIH regulations. All fire-rated features of the contract design must be approved by Fire Marshal. 35

15. Submission and Mock-up - Penetrations into the containment barrier (including mounting of electrical boxes) shall be detailed in the construction documents and require mock-ups to be constructed and tested prior to installation. Penetration details for equipment shall be coordinated with the equipment manufacturer.

16. Provide gasketed, sealed, weather tight surface mounted light fixtures. 36

16. N/A

17. The BSL-3 lighting systems shall be designed to ensure biohazards are contained within the area. Lighting fixtures shall be easily opened to permit full decontamination. 37, 38

17. N/A

18. Provide emergency lighting in each room of the BSL-3 suite. 39

18. Emergency lighting installation shall comply with the following requirements:
- Connect at least 50% of lighting fixtures in the laboratories to the emergency power source.
- Provide at least one lighting fixture per room in laboratory areas with self-testing emergency battery ballast. Connect the lighting fixtures with emergency ballasts to unswitched local emergency generator circuits.
19. Lighting levels must be sufficient for tasks.

20. Conventional light switching (voltage toggle) is acceptable.  

20. Occupancy sensors may be used in laboratory support and other areas, but not in containment spaces where BSCs are to be located. In addition, specialized lighting controls (such as dimming systems and DC lighting controls) shall be provided per specific program requirements.
Electrical References

1. The laboratory must be designed so that it can be easily cleaned and decontaminated. Carpets and rugs are not permitted. Seams, floors, walls, and ceiling surfaces should be sealed. BMBL 5th ed. Section IV. Biosafety Level 3.D

2. All penetrations shall be durable, sealed, and tested to meet room tightness criteria for BSL-3 laboratories. NIH DRM Section 10.8.2.A.2

3. Inspect and Evaluate – Finishes, penetrations & caulking integrity for architectural elements such as doors, around the ceilings, lighting fixtures, electrical devices, etc. within containment to meet requirements for: Sealed seams and penetrations. NIH BSL-3 Certification Section I.2

4. Visual inspection – The integrity of all surfaces, penetrations, and seals on the containment perimeter shall be visually inspected. ANSI Z9.14 Section 8.3.2

5. Decontaminate work surfaces after completion of work and after any spill or splash of potentially infectious material with appropriate disinfectant. BMBL 5th ed. Section IV. Biosafety Level 3.A.7

6. All surface finishes shall be selected to be compatible with the anticipated agents and methods used for cleaning, disinfection, or sterilization and protocols used by program without damage or degradation, including discoloration. NIH DRM Section 4.4.5

7. Cover plates for building interior receptacles, switches, and boxes shall be stainless steel, brushed aluminum, or hospital-grade impact resistant nylon. Cover plates for cast boxes shall be gasketed and weatherproof. NIH DRM Section 10.5.3.8

8. Provide weatherproof covers for receptacles and switches where they are exposed to water. Utilize stainless steel cover plates for switches and receptacles in rooms that will utilize vaporized paraformaldehyde or chlorine dioxide gaseous decontamination protocols. NIH DRM Section 10.5.5.H.4

9. Properly sealed laboratory surfaces (walls, floors, and ceilings) are essential to maintain controlled directional airflow and ventilation system performance. Room leakage or tightness is also critical when gaseous fumigants are used for decontamination. ANSI Z9.14 Section 8.3.2

10. Conduits for all systems: Conduit applications in BSL-3 facilities are as follows: 1) Conduit Type: Use Rigid Galvanized Steel (RGS) conduit with threaded fittings in all BSL-3 areas. 2) Seal-off: Provide seal-off fittings when conduits exit defined BSL-3 perimeter. SMR: Use of surface metal raceway systems (SMRs) is not allowed in BSL-3 areas. NIH DRM Section 10.8.3.A

11. Inspect and Evaluate – Finishes, penetrations & caulking integrity for architectural elements such as doors, around the ceilings, lighting fixtures, electrical devices, etc. within containment to meet requirements for: Surface impermeability to liquids; Resistance of surfaces to chemical (organic solvents, acids, alkalis), disinfectants and moderate heat; Gas tightness for decontamination; Pest management requirements. NIH BSL-3 Certification Section I.2

12. Develop HVAC system and electrical systems failure tests consistent with laboratory design parameters. Perform tests and record data. To verify correct operations these tests should include at a minimum: 1) Normal operations to emergency power. 2) Emergency power to normal operations NIH BSL-3 Certification Section II.17

13. Pneumatic control air system and control power system shall be assessed and verified to be in good operating condition by visual inspection so as to maintain any required critical control or secondary containment features during normal operation and in the event of loss of power, failure of a generator, or failure of primary compressors and equipment. ANSI Z9.14 Section 8.3.6

14. Visual inspection of control power system should include control power for all heating, ventilating, and air-conditioning (HVAC) controls should be served by standby power system. ANSI Z9.14 Section 8.3.6

15. HVAC systems shall be tested for directional airflow during system failures. During system failures, momentary changes in airflow patterns can be acceptable based on the facility risk assessment. Failure of the normal/preferred source power supporting supply and exhaust fan systems...
components and transition to the emergency or alternate source. The ability to transition from normal to the backup system should be verified. Return from power outage or emergency or alternate power source to normal or preferred power source should be verified. ANSI Z9.14 Section 8.4.2

16. Standby Power Requirements: In addition to the loads listed in Section 10.3, Emergency Power, connect the following loads to the standby electrical systems: 1) Heating and cooling units provided for critical support function. 2) Receptacles serving selected equipment identified as critical by the program. 3) General lighting in laboratories. 4) Critical control and containment equipment compressed air systems. NIH DRM Section 10.8.1.B

17. As determined per program basis, recommend connecting the following loads to the standby electrical system: 1) Freezers/refrigerators, 2) Sterilizers, 3) Cage wash equipment for remote locations, where there are no other options available to accommodate an extended outage, 4) Other loads as required by the program based on the risk assessment. NIH DRM Section 10.8.1.B

18. The emergency distribution panel and all emergency gear up to the load end of the automatic transfer switches shall be located in a separate dedicated electrical room away from the normal power electrical room. NIH DRM Section 10.3.3

19. Wiring for the emergency power shall be separate from the normal power. NIH DRM Section 10.3.4.A

20. This is to ensure that emergency electrical service will meet current and future demand of the facility. NIH DRM Section 10.3.1

21. Life Safety Loads may include all or some of the following: 1) Emergency egress lighting, 2) Egress signage, 3) Communications systems (including PA systems), 4) Fire alarm and mass notification systems, 5) Self-contained battery-powered lighting at generator set location, 6) Fire-suppression systems (fire pumps, jockey pumps, compressors, valves, etc.) NIH DRM Section 10.3.1.B

22. A central UPS system or a number of local UPSs may be required to backup all building wide low voltage systems that are essential for containment operation, safe shutdown of the facility and for critical BSL-3 specific loads. Central UPS shall be of the double-conversion online type; wet cell type batteries are recommended. NIH DRM Section 10.8.1.C

23. Develop HVAC system and electrical systems failure tests consistent with laboratory design parameters. Perform tests and record data. To verify correct operations these tests should include at a minimum: 1) If a UPS is installed, verify operation of relays, 2) Provide UPS for BAS, 3) Assess if UPS is operational. NIH BSL-3 Certification Section II.17

24. Containment systems should respond in such ways as to avoid operating conditions that present risks to workers and the environment. Electrical systems including backup generators for normal power outages provide effective strategies to maintain containment including minimization of flow reversals. Emergency generators can normally respond within 10-15 seconds to restore power, whereas UPS systems (where installed) provide continuity of power between transitions of normal to emergency power systems. Based on the risk assessment for the facility, each supporting system should have a documented sequence of operation/strategy for power failure and restoration. ANSI Z9.14 Section 8.4.6

25. Boxes for All Systems: General requirements of device boxes are as follows: 1) Type and Depth: All boxes shall be double gang type; the box depth shall be at least the next larger than minimum size required per code. 2) Cast Boxes: Provide cast boxes with external mounting provisions, external hub, and gasketed device cover plates. 3) Sealing: Provide a 25 mm (1 in.) barrier of silicone caulk around the wire within a device box hub. Provide a continuous bead of caulk between the device box and the adjacent surface. Provide a continuous bead of caulk around the device cover plate and the adjacent surface. NIH DRM Section 10.8.3.D

26. Inspect and Evaluate – Finishes, penetrations & caulking integrity for architectural elements such as doors, around the ceilings, lighting fixtures, electrical devices, etc. within containment to meet requirements for: Sealed seams and penetrations; Surface impermeability to liquids; Resistance of surfaces to chemical (organic solvents, acids, alkalis), disinfectants and moderate heat; Gas tightness for decontamination; Pest management requirements. NIH BSL-3 Certification Section I.2
27. Properly sealed laboratory surfaces (walls, floors, and ceilings) are essential to maintain controlled directional airflow and ventilation system performance. Room leakage or tightness is also critical when gaseous fumigants are used for decontamination. Examples of elements to inspect include: Piping and electrical wall or ceiling penetrations including fire sprinklers; Seals around light fixtures, receptacles, diffusers, and grilles; Sealing of electrical and communication/data wires within conduit; Components mounted to walls and ceilings. ANSI Z9.14 Section 8.3.2

28. Electrical Installation in Containment Areas: Avoid installing electrical equipment which requires service within a containment area. NIH DRM Section 10.8.2.A

29. The A/E shall coordinate requirements for security wiring, on a per project basis with the Project Officer and Division of Physical Security Management (DPSM). NIH DRM Section 10.8.3.C

30. Inspect and Evaluate – Finishes, penetrations & caulking integrity for architectural elements such as doors, around the ceilings, lighting fixtures, electrical devices, etc. within containment to meet requirements for: Sealed seams and penetrations. NIH BSL-3 Certification Section I.2

31. Properly sealed laboratory surfaces (walls, floors, and ceilings) are essential to maintain controlled directional airflow and ventilation system performance. Room leakage or tightness is also critical when gaseous fumigants are used for decontamination. Examples of elements to inspect include: Piping and electrical wall or ceiling penetrations including fire sprinklers; Seals around light fixtures, receptacles, diffusers, and grilles; Sealing of electrical and communication/data wires within conduit; Components mounted to walls and ceilings. ANSI Z9.14 Section 8.3.2

32. RGS conduit with threaded fittings shall be used in all BSL-3 areas. NIH DRM Section 10.8.3.A.1

33. Facilities shall have a minimum of two dedicated utility services, physically separated in different duct-banks and different manholes. These dedicated services shall be fed by different primary substations or by one double-ended utility substation, which is fed by two dedicated utility service lines. Each required electric service to the facility shall be sized to handle 100% of the design load (i.e., N + 1 redundancy). NIH DRM Section 10.8.1.A

34. Evaluate and minimize single points of failure for all systems including power supplies, electrical distribution, grounding, equipment, and controls for all projects. Downstream electrical distribution from switchgear to critical areas, such as mechanical support rooms with redundant motors in each set, shall comprise pairs of distribution switchboards/panel boards, each fed from a separate side of the switchgear, to supply approximately half of each set of motors. Where possible, provide separate feeders to packaged units with multiple motors, i.e., a separate feeder for each motor. NIH DRM Section 10.8.1.A.6, 10.8.1.A.7

35. Electrical Installation in Containment Areas: Avoid installing electrical equipment which requires service within a containment area. Electrical systems and equipment not serving the BSL-3 area shall not be located within the containment area. Containment Barrier Penetrations: Penetrations through the containment barriers shall comply with the following requirements: 1) Penetrations through the containment barriers shall be gas-tight, non-porous, smooth and cleanable; and readily visible for routine inspection, cleaning, and maintenance. Penetrating components shall be sufficiently rigid in construction and adequately braced to structure to maintain the long-term integrity of the penetration. The result shall be free of sharp edges or similar hazards. 2) All penetrations shall be durable, sealed, and tested to meet the room tightness criteria for BSL-3 laboratories. Submission and Mock-up: Penetrations into the containment barrier (including mounting of electrical boxes) shall be detailed in the construction documents and shall require mock-ups to be constructed and tested prior to installation. Penetration details for equipment shall be coordinated with the equipment manufacturer. NIH DRM Section 10.8.2.A, 10.8.2.B

36. Lighting fixture installation shall comply with the following requirements: 1) Fixtures shall be UL listed with a minimum IP65 rating, 2) Lighting fixtures shall allow full decontamination with ease of effort, and permit easy re-lamping and access to ballasts. 3) Lighting fixtures shall be
provided with stainless steel housings, glass or heavy duty acrylic prismatic lens, and stainless steel door with tool-less fasteners or captive, flush, stainless steel screws. 4) Use surface mounted, fully sealed, enclosed, and gasketed fluorescent or LED fixtures. Seal surface mounted fixtures with a continuous bead of sealant around its perimeter to seal housing to ceiling. Lighting fixture must have a sealed conduit entrance to housing. Lighting fixtures may be pendant mounted only in an open ceiling. Pendant-mounted lighting fixtures shall be fully sealed and gasketed with same features as those of surface mounted fixtures including both a sealed conduit entrance to housing and a sealed conduit entrance a ceiling canopy. 5) Install fixtures in continuous rows and aligned with edge of laboratory bench in laboratories and laboratory support areas. Install lighting fixtures in a symmetrical pattern. Fixtures mounted in continuous rows should have a minimum of 152 mm (6 in.) between fixtures to allow for disinfection. 6) In specific imaging modalities, such as MRI, incandescent or LED fixtures and conduit shall be made of non-ferrous materials. Utilize EMI filters and DC dimming methods that are compatible with imaging equipment. All special requirements shall be coordinated with the manufacturer of equipment in each modality. NIH DRM Section 10.8.4.A, 10.8.4.B

37. The BSL-3 lighting systems shall be designed to ensure biohazards are contained within the area. Lighting fixtures shall be easily opened to permit full decontamination. NIH DRM Section 10.8.4.A

38. BSL-3 laboratory lighting fixtures and systems shall be designed with the containment barrier at the outlet box. NIH DRM Section 10.8.4

39. Emergency lighting installation shall comply with the following requirements: 1) Connect at least 50% of lighting fixtures in the laboratories to the emergency power source. 2) Provide at least one lighting fixture per room in laboratory areas with self-testing emergency battery ballast. Connect the lighting fixtures with emergency ballasts to unswitched local emergency generator circuits. NIH DRM Section 10.8.4.C

40. Laboratories shall utilize line voltage toggle switches. Occupancy sensors may be used in laboratory support and other areas. In addition, specialized lighting controls (such as dimming systems and DC lighting controls) shall be provided per specific program requirements. If occupancy sensors are used, specify infrared sensors with sealed enclosures. NIH DRM Section 10.8.4.D
<table>
<thead>
<tr>
<th>Standard</th>
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</tr>
</thead>
<tbody>
<tr>
<td>1. Provide labeled shut-off valves to utilities servicing the BSL-3 area(s) and locate them outside the containment barrier.</td>
<td>To maintain the integrity of the containment envelope there should be no access panels in the BSL-3 laboratory. Laboratory Mechanical, Electrical, Plumbing (MEP) items that may need maintenance shall be outside the containment zone.</td>
<td>Access to shut-off valves should be restricted to trained individuals whenever possible. • Labels should be understandable to trained individuals, but not compromise security.</td>
</tr>
<tr>
<td>2. Cylinder gases shall be piped from adjacent rooms outside the BSL-3 suite</td>
<td>N/A</td>
<td>Provide a locked closet for cylinder gases.</td>
</tr>
<tr>
<td>3. Provide back-flow prevention devices at each water service. • Provide back-flow prevention devices at each water faucet</td>
<td>Backflow device shall be Reduced Pressure Principle type for highest risk level noted in the California Plumbing Code Table 603.2.</td>
<td>N/A</td>
</tr>
<tr>
<td>4. Minimize surface mounted utilities.</td>
<td>Exposed piping shall be chemically compatible, non-porous, smooth, with sanitary surfaces, and shall utilize sanitary type piping clamps and supports. Hangers/clamps and all associated attachments shall be corrosion resistant (typically stainless steel with plastic grommets), free of sharps.</td>
<td>N/A</td>
</tr>
<tr>
<td>5. Do not provide compressed air to the BSL-3 suite</td>
<td>The use of compressed air increases the risk of aerosolizing microorganisms. Inhalation of infectious aerosols is the primary route of laboratory acquired infections.</td>
<td>N/A</td>
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</table>
6. All piping into the BSL-3 shall be secured to prevent movement that may break the barrier and the outside.  

Water Distribution

<table>
<thead>
<tr>
<th>Standard</th>
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<tbody>
<tr>
<td>7. The water distribution system shall be designed to provide the required flow and pressure for the most hydraulically demanding fixture/equipment. System must comply with local regulations and industry standards.</td>
<td>7. Pipe mains shall be designed to achieve the maximum calculated flow rate and to provide a 20% allowance for future expansion.</td>
<td>7. N/A</td>
</tr>
<tr>
<td>8. Piping systems shall be properly insulated.</td>
<td>8. N/A</td>
<td>8. N/A</td>
</tr>
<tr>
<td>9. Comply with all local plumbing codes.</td>
<td>9. N/A</td>
<td>9. N/A</td>
</tr>
<tr>
<td>10. If an Effluent Decontamination System (EDS) is provided, plumbing must be installed within the containment zone.</td>
<td>10. The requirement for EDS in the laboratory will be based on a risk assessment</td>
<td>10. N/A</td>
</tr>
</tbody>
</table>

Vacuum Line

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<th>Standard</th>
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<tbody>
<tr>
<td>11. Provide easily replaceable HEPA filters where vacuum lines connect to BSCs.</td>
<td>11. N/A</td>
<td>11. Installed central vacuum systems should be dedicated to the BSL-3 laboratory, should not serve other areas, and should be located in close proximity to the containment space. Central</td>
</tr>
<tr>
<td>• HEPA filters on the vacuum lines must be easily replaceable and appropriately decontaminated prior to</td>
<td></td>
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</tbody>
</table>
disposal. Vacuum pumps should be located in a negatively pressurized room (negative to adjacent spaces).

### Sinks

<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>12. Provide wall mounted, hands-free type 304 stainless steel (16 gauge) hand wash sink with electrically operated hot/cold premixing valves near main entrance/exit to each room in a laboratory suite. Seal all openings in sink. Sink to have coved backsplash.</td>
<td>12. N/A</td>
<td>12. N/A</td>
</tr>
<tr>
<td>• Provide hands-free paper towel dispenser and soap dispenser at sink in anteroom</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13. Provide hands-free hand wash sink with electrically operated valves at each zone within the BSL-3 laboratory.</td>
<td>13. Provide additional sinks as required by risk assessment. • Risk assessment of agents and laboratory procedures may recommend additional sinks in an effort to minimize exposures.</td>
<td>13. N/A</td>
</tr>
<tr>
<td>14. Faucets must be designed such that the splash is minimized during handwashing and other procedures.</td>
<td>14. N/A</td>
<td>14. Faucets within containment should have atmospheric vacuum breaker and laminar flow, non-aerating, non-splash outlet.</td>
</tr>
<tr>
<td>15. Flow rates shall be at least 1.8 gal/min.</td>
<td>15. Laboratory sinks are not regulated by CalGreen (California code) or LEED. UC Sustainability policy calls for water use reduction in facilities. Adopting this</td>
<td>15. Suggest that Calgreen kitchen sink requirement be adopted: 1.8 GPM normal maximum with intermittent maximum of 2.2</td>
</tr>
</tbody>
</table>
standard would be consistent with UC GPM.

16. Where stainless steel sinks are used, surface undercoating and sound-deadening pads shall be omitted. 32

16. N/A

Shower

<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>17. Shower out requirements are based on risk assessment. 33, 34, 35, 36, 37</td>
<td>17. A shower out requirement will be based on risk assessment. If required, pass-through design shower should be considered to allow traffic in one direction only. Dirty clothing/PPE shall not contaminate the clean area.</td>
<td>17. When not required, consideration should be given to the future addition of showers with regards to laboratory design layout and plumbing demands.</td>
</tr>
<tr>
<td>• Hand-held showers shall not be utilized except where specifically required for barrier-free compliance and shall include a vacuum breaker.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>18. Fixtures requiring potable supply direct from the domestic potable water system (e.g., emergency eyewash, showers, and toilet room/shower fixtures located in containment) shall be isolated from other functions with an ASSE 1013 backflow preventer. 38, 39</td>
<td>18. Backflow device shall be Reduced Pressure Principle type for highest risk level noted in the California Plumbing Code Table 603.2</td>
<td>18. N/A</td>
</tr>
</tbody>
</table>

Drains

<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>19. If laboratory intends to dispose of</td>
<td>19. Requirement for EDS will be based on a</td>
<td>19. N/A</td>
</tr>
</tbody>
</table>
contaminated liquid waste down the drains or disposal of such could possibly occur unintentionally, it must be discharged to a dedicated waste system prior to decontamination. 40

risk assessment and best practices guidelines.

• Laboratory personnel may eliminate the need for EDS by decontaminating liquid waste with appropriate disinfectant and contact time prior to disposal.

20. Floor sinks provided for receiving the condensate from the sterilizer chamber shall be on the clean side of the autoclave bioseal. 41, 42

20. N/A

20. Floor drains/floor sinks should be avoided in containment.

21. Drains pipes must be resistant to liquid decontaminant used in the facility.

21. N/A

21. Metal pipes should be avoided in areas where decontaminated liquid waste will be disposed.

### Traps

<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
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</tr>
</thead>
</table>
| 22. 6-inch P traps (minimum) must be provided if significant changes in pressure could occur.  
• Traps shall be selected to exceed the exhaust fan static pressure to maintain at least a 37-50 mm (1.5 -2 in.) seal depth under HVAC operating and fan failure modes, as well as extra depth as required for control of pressure transients. 43 | 22. N/A | 22. Deep traps provide safer protection by maintaining trap seal when sudden changes in space pressure occur. |
<p>| 23. Traps shall be resistant to common decontamination agents such as bleach. | 23. N/A | 23. N/A |</p>
<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>24. Provide cleanouts in such a way as to maintain integrity of the containment barrier.</td>
<td>24. N/A</td>
<td>24. Threaded cleanout plugs should be coated with threaded joint sealer for an airtight seal.</td>
</tr>
</tbody>
</table>
**Plumbing References**

1. Each pressurized piping penetration from outside the barrier into containment shall be provided with a shut-off valve located outside the containment barrier serving only the BSL-3 area(s). Each equipment connection shall include dedicated isolation valve. Fire-sprinkler piping need not be provided with a shut-off valve at each penetration; however, the sprinkler zone serving BSL-3 should be capable of independent isolation.  **NIH DRM Section 8.6.1.D**

2. All piping systems shall be identified using system-nomenclature-specific pipe labels. **NIH DRM Section 8.1.3.4.B**

3. Gas cylinders shall not be located inside containment spaces and shut-off valves should be provided for each penetration into containment to permit independent isolation of each service. **NIH DRM Section 8.6.6.B**

4. Water supplies to BSL-3 spaces shall be isolated from other functions with an approved back-flow preventer (BFP) installed outside containment prior to serving BSL-3 areas. **NIH DRM Section 8.6.2.A**

5. Each water service shall be provided with a service entrance backflow preventer sized for 100% demand load, ASSE 1013 type for domestic water and ASSE 1015/1048 or ASSE 1013/1047 type for fire service as appropriate. On facilities with single water service is utilized to supply each system, backflow preventers on the individual service shall be arranged in parallel, sized to provide N+1 redundancy. **NIH DRM Section 8.3.2.E**

6. Exposed piping shall be chemically compatible, non-porous, smooth, with sanitary surfaces, and shall utilize sanitary type piping clamps and supports. Hangers/clamps and all associated attachments shall be corrosion resistant (typically stainless steel with plastic grommets), free of sharps, and where applied in ABSL-3 areas exposed piping shall be in conformance with ASME BPE-2002 and properly sealed with an approved sealant. **NIH DRM Section 1.15.4.F**

7. Each laboratory, animal, and other critical facilities shall be provided with two normally operating (online) water services, each sized for total demand and appropriately connected to the campus loop. Water supplies and distribution systems shall be arranged to minimize potential for single point failure or service loss. **NIH DRM Section 8.3.2.A**

8. The use of compressed air within a BSC must be carefully considered and controlled to prevent aerosol production and reduce the potential for vessel pressurization. **BMBL 5th ed. Appendix A**

9. Pressurized gas systems shall not be connected to primary containment devices, including BSCs... Only where pressurized gas services are necessary for required equipment operation or procedures in primary containment equipment (e.g., aerosol inhalation chambers) may such services be provided... the service line shall have backflow preventer check valve arrangement at the point of use. **NIH DRM Section 8.6.6.C**

10. Each pressurized piping penetration from outside into containment shall be provided with a shutoff valve. Each equipment connection shall include isolation valves. **NIH DRM Section 8.6.1.D**

11. A separate and distinct central laboratory/non-potable water subsystem shall be provided and distributed throughout the building, sourced and isolated from the domestic water system with parallel ASSE 1013 backflow preventers, which have been sized and arranged to N+1 redundancy. **NIH DRM Section 8.3.4.B**

12. The water distribution system shall be designed to provide the required flow and pressure for the most hydraulically demanding fixture/equipment. Systems shall be designed to provide at least 280 kPa (40 psi) residual (flowing) pressure at the most remote outlet on the laboratory water system and not less than 240 kPa (35 psi) residual pressure for the hydraulically remote fixture on the domestic plumbing system. **NIH DRM Section 8.3.4.1.B**

13. A pressure-reducing station shall be provided if required to limit maximum water pressure to 550 kPa (80 psi) at any service outlet. Major
pressure reducing station valves shall be of the hydraulically-operated pilot-type automatic control valve, municipal grade with stainless steel trim. NIH DRM Section 8.3.4.1.C

14. Piping systems shall be properly insulated. NIH DRM Section 8.3.5.C

15. Pipe mains shall be designed for the maximum calculated flow at the design stage and to provide a 20% allowance for future expansion. NIH DRM Section 8.3.5 F

16. The pipe sizing criteria shall comply with requirements of Table 8.4.5, Minimum Waste Piping Diameters. NIH DRM Section 8.4.5.B

17. Vacuum lines must be protected with HEPA filters, or their equivalent. BMBL 5th ed. Section IV.Biosafety Level 3.D.7

18. Filters must be replaced as needed. BMBL 5th ed. Section IV.Biosafety Level 3.D.7

19. Liquid disinfectant traps may be required. BMBL 5th ed. Section IV.Biosafety Level 3.D.7

20. Utility services needed within a BSC must be planned carefully. Protection of vacuum systems must be addressed (Figure 12). BMBL 5th ed. Appendix I

21. If vacuum lines are used with toxin, they should be protected with a HEPA filter to prevent entry of toxins into the line. BMBL 5th ed. Appendix I

22. The use of disinfectant traps and hydrophobic filters are required at each point of use, including biological safety cabinets and aerosol chambers. Filters utilized shall be at least HEPA efficiency for liquid and gas streams; and permanent type pipe-line filters should be sterilizing grade for repeated usage. NIH DRM Section 8.6.5.A

23. Laboratories must have a sink for hand washing. BMBL 5th ed. Section IV.Biosafety Level 3.D.2

24. Sink must be hands free or automatically operated. BMBL 5th ed. Section IV.Biosafety Level 3.D.2

25. Locate sink near the exit door. BMBL 5th ed. Section IV.Biosafety Level 3.D.2

26. All laboratories require a hands-free or automated hand washing sink located near the exit door. NIH DRM Section 4.9.8.G

27. Sink faucets shall be hands-free that are either electric sensor operated and hard-wired to AC power, or foot pedal actuated with slow-close off-the-floor mounted valves with flip-up pedals to permit cleaning. Battery actuated faucets and faucets which operate with hands, wrist or elbow are not acceptable. Knee operated valves shall include suitable anchorage, or shall be knee-panel type with pneumatic control valves (scrub-sink type). NIH DRM Section 8.6.11.1.B

28. Sinks must be available for handwashing in each zone within a BSL-3 suite. BMBL 5th ed. Section IV.Biosafety Level 3.D.2

29. Additional sinks may be required by risk assessment. BMBL 5th ed. Section IV.Biosafety Level 3.D.2

30. Faucets within containment shall have gooseneck-type spouts and be fitted with integral ASSE 1001 atmospheric vacuum breaker and laminar flow, non-aerating, non-splash outlet. NIH DRM Section 8.6.11.1

31. Flow rates shall be at least 2 gal/min. NIH DRM Section 8.6.11.1

32. Where stainless steel sinks are used, surface undercoating and sound-deadening pads shall be omitted. NIH DRM Section 8.6.11.4.A

33. Shower-out capability may be required based on agent summary statement, risk assessment, or applicable local, state, or federal regulations. BMBL 5th ed. Section IV.Biosafety Level 3.D.14

34. Risk assessment may require the use of pass through showers in the exiting sequence. When not required, consideration should be given to the addition of showers for future flexibility. NIH DRM Section 2.5.3.5.C

35. Maximum outlet temperature at showers shall be limited to 45 degrees C (112 deg. F) at the individual fixture limit stop. NIH DRM Section 8.2.9.B

36. Showers shall provide a minimum flow rate of 2.5 gal/min. NIH DRM Section 8.6.11.2.A

37. Hand-held shower shall not be utilized except where specifically required for barrier-free compliance and shall include a vacuum breaker. NIH
**DRM Section 8.6.11.2.A**

38. Outlets requiring potable supply direct from the domestic potable water system (e.g., emergency eyewash, showers, and toilet room/shower fixtures located in containment) shall be isolated from other functions with an ASSE 1013 backflow preventer. **NIH DRM Section 8.6.2.C**

39. Indirect waste (including but not limited to fire sprinkler drains) shall not terminate into other plumbing fixtures including janitor mop sinks, showers, or other fixtures not dedicated specifically for receipt of indirect waste. **NIH DRM Section 8.4.10.D**

40. All drain inlets within containment shall discharge to the dedicated waste system serving BSL-3 facilities. **NIH DRM Section 8.6.9.1.D**

41. The sterilized effluent from BSL-3 autoclave chambers fitted with decontamination of all chamber effluent may discharge through the sanitary system or general building laboratories waste system as an indirect connection through a floor sink. The drain receptor shall be located on the clean (non-contained) side of the bioseal (typically within the clean sterilizer service access area). **NIH DRM Section 8.6.9.1.P**

42. Floor drains/floor sinks shall be avoided in containment. **NIH DRM Section 8.6.9.1.H**

43. Waste systems shall be atmospherically vented to the building exterior and shall include deep seal traps for all drain inlets within containment. **NIH DRM Section 8.6.9.1.A**

44. Waste pipe clean-out access shall be arranged similar to other piping penetrations to maintain integrity of the containment barrier, typically arranged as a threaded capped or plugged pipe extension through a wall. **NIH DRM Section 8.6.9.1.J**
### Telecom

<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Exposed conduit, piping or pathways are to be resistant to decontamination chemicals and easily cleaned behind, if surface mounted.</td>
<td>1. Any material exposed in the BSL-3 laboratory shall be resistant to decontamination methods.</td>
<td>1. N/A</td>
</tr>
<tr>
<td>2. Provision for landline telephone must be made available.</td>
<td>2. Landline makes communication with dispatch easier</td>
<td>2. Provide an intercom or hands-free telephone in each room, including anteroom¹ &lt;li&gt;System must be connected to a location that has personnel available for emergency response at all times work is being performed in a BSL-3 laboratory.&lt;/li&gt;</td>
</tr>
<tr>
<td>3. Provide a method for electronic transfer of information to outside of containment (e.g.: scanner, fax, iPad, etc.).²</td>
<td>3. Electronic transfer method eliminates the practice of transferring documents from the BSL-3 to areas outside of containment.</td>
<td>3. N/A</td>
</tr>
</tbody>
</table>

### Security

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<tr>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
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</thead>
<tbody>
<tr>
<td>4. Anteroom doors must be self-closing and interlocked. Interlocked doors must have an emergency override.³ ⁴</td>
<td>4. Self-closing doors are required to both maintain the pressure differential and the security of the space. Locks are required to prevent unauthorized access to the containment facility.</td>
<td>4. N/A</td>
</tr>
</tbody>
</table>
5. Electronic access control is required at Anteroom for access to BSL-3 suite. System must provide access history. 5, 6, 7.

5. Due to the use of biohazardous materials, access to the biocontainment facility must be strictly limited to trained authorized personnel.
   - Provide card key, biometric, or PIN security access. Link system to EH&S and/or building/campus security.

5. Video surveillance cameras should be installed to provide live and recorded video activity outside of the secured space, in interior activity spaces, and covering materials of interest. IP-based, high resolution (minimally 3 megapixel), powered using Power-over-Ethernet (PoE) and should be Wide Dynamic Range and Low Light (if not infrared) capable.
   - The access control and video surveillance system devices should be coordinated to allow for recording and monitoring of entry and exit events. Due to the sensitivity of research, the physical security systems should be integrated to provide for real time monitoring and post evening auditing of the video surveillance and electronic access control systems. Please utilize any existing enterprise systems in coordination with the Campus Police Department.
   - Emergency/duress notification systems should be installed in the laboratory spaces where work is being
performed. A two way, single button push emergency phone with direct connection to the Campus Police Department should be installed in a highly visible and easily accessible location in the laboratory. Please utilize any existing enterprise systems in coordination with the Campus Police Department.

6. Consider a method for maintaining an access control record for mechanical/support spaces.

6. Access to mechanical and support spaces must be designed to limit unauthorized users.

6. N/A

<table>
<thead>
<tr>
<th>Signage</th>
<th>Explanation</th>
<th>Best Design Practice</th>
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</thead>
<tbody>
<tr>
<td>7. Provide space on or near the anteroom door for appropriate signage, coordinated with EH&amp;S and the Fire Marshal.</td>
<td>Accurate signage information is important for many reasons, including health and safety. Conversely, for security reasons detailed signage may be a concern. To determine actual signage requirements, the design team should meet with the University's EH&amp;S and the Fire Marshal.</td>
<td>7. N/A</td>
</tr>
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</table>
## Enhanced Facilities

<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
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</thead>
<tbody>
<tr>
<td>8. Enhanced facility requirements shall be determined by risk assessment study. Enhancements may be required for facility and security based on Select Agent Requirements. Example includes shower-out capabilities for projects involving work with Highly Pathogenic Avian Influenza. 8, 9, 10</td>
<td>In some cases, risk assessment may call for facility designs that exceed those recommended in the BMBL BSL-3 guidelines. This may be due to the agents and/or volumes in use, the experimental procedures or other guidelines that may be followed. The enhancements noted have all been noted in other parts of this document and may be required to further protect the users and environment, to better be able to decontaminate the facility or to provide better security. • Laboratory effluent decontamination</td>
<td>8. N/A</td>
</tr>
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</table>

## Verification & Documentation

<table>
<thead>
<tr>
<th>Standard</th>
<th>Explanation</th>
<th>Best Design Practice</th>
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<tbody>
<tr>
<td>9. The BSL-3 facility design, operational parameters, and procedures must be verified and documented prior to operation. Facilities must be re-verified and documented at least annually. 11</td>
<td>BSL-3 facilities are some of the most complicated facilities to design, construct and operate. The complex systems must be thoroughly documented and facilities personnel should be trained to maintaining the various systems.</td>
<td>9. N/A</td>
</tr>
<tr>
<td>10. An independent third-party commissioning agent (CA) is required for design consultation and certification of the</td>
<td>Documentation of initial testing must be provided and must comply with ANSI Z9.14 standards.</td>
<td>10. Commissioning agent should be involved from the beginning of the project.</td>
</tr>
</tbody>
</table>
BSL-3 Facility. The Biosafety Officer and High-Containment Director must be included in the BSL-3 facility design and certification. BSL-3 laboratories must be certified by an experienced team of engineers and biosafety professionals before initial operation. 12, 13

- Certification must be done in the presence of the Biosafety officer and/or High-Containment Director.
- The facility must be tested and verified at least annually in coordination with the HCLD to provide assurance that the facility is in proper operational condition.

<table>
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<tr>
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</thead>
<tbody>
<tr>
<td>11. All tall (over 42 inches) and/or heavy equipment must be provided with seismic restraints designed to withstand shaking equivalent to 7.0 on the Richter scale. 14</td>
<td>11. This is to prevent damage of equipment in the event of an earthquake.</td>
<td>11. N/A</td>
</tr>
<tr>
<td>12. Tall and/or movable cabinets must be seismically anchored. Anchor points that penetrate the containment barrier must be properly sealed.</td>
<td>12. Penetrations made to the containment envelop need to be sealed to maintain proper directions airflow.</td>
<td>12. N/A</td>
</tr>
</tbody>
</table>
Telecom, Security & Documentation References

1. Evaluate availability of emergency two-way communication system. NIH BSL-3 Certification Section I.3
2. Evaluate availability of system provided for electronic transfer of information to outside of containment. NIH BSL-3 Certification Section I.3
3. Laboratory doors must be self-closing and have locks in accordance with the institutional policies. BMBL 5th ed. Section IV.Biosafety Level 3.D.1
4. Physical separation of the high containment laboratory from access corridors or other laboratories or activities may be provided by a double-door clothes change room (showers may be included), airlock, or other access facility which requires passage through two sets of doors before entering the laboratory. NIH Guidelines Appendix G-II-C-4-a
5. Laboratory access is restricted. BMBL 5th ed. Section IV.Biosafety Level 3.D.1
6. At BSL-3, more emphasis is placed on primary and secondary barriers. Secondary barriers for this level include controlled access to the laboratory. BMBL 5th ed. Section III.Principles of Biosafety.Biosafety Levels
7. Advanced access control devices, such as biometrics, may be required by the agent summary statement, risk assessment, or applicable local, state or federal regulations. BMBL 5th ed. Section IV.Biosafety Level 3.D.14
8. Biosafety Level 3 Enhanced for Research Involving Risk Group 3 Influenza Viruses. (See Appendices G-II-C-2-n, G-II-C-2-r, and G-II-C-4-i for additional guidance for facilities, waste handling, and serum collection for research involving mammalian-transmissible HPAI H5N1 virus. NIH Guidelines Appendix G-II-C-5
9. Enhanced environmental and personal protection may be required by the agent summary statement, risk assessment, or applicable local, state or federal regulations. Laboratory enhancements may include, one or more of the following: BMBL 5th ed. Section IV.Biosafety Level 3.D.14
10. An anteroom for clean storage of equipment and supplies with dress-in, shower-out capabilities BMBL 5th ed. Section IV.Biosafety Level 3.D.14
11. The BSL-3 facility design, operational parameters, and procedures must be verified and documented prior to operation. Facilities must be re-verified and documented at least annually. BMBL 5th ed. Section IV.Biosafety Level 3.D.15
12. An independent third-party commissioning agent (CA) is required. NIH DRM Section 1.10.1.E
13. Personnel shall verify that the direction of the airflow (into the laboratory) is proper. The exhaust air from the laboratory room may be discharged to the outside without being filtered or otherwise treated unless research is being conducted with mammalian transmissible HPAI H5N1 virus. For research with mammalian-transmissible HPAI H5N1 virus, exhaust air must be HEPA filtered and there must be sealed ductwork from the containment barrier to the filter. In addition, the air handling system shall be designed such that under failure conditions, the airflow will not be reversed and periodic verification, with annual verification of the HEPA filters, shall be performed. NIH Guidelines Appendix G-II-C-4-i
14. Any equipment, including but not limited to, appliances and shelving to be installed by the contractor, which is 42 inches or higher and has the potential for falling over during an earthquake, or moving and blocking corridors or doors, shall be permanently braced or anchored to wall-studs, structural columns and/or the floor. UC Laboratory Design Manual, Seismic Hazard Abatement #3.
## Safety and Decontamination

### Safety

<table>
<thead>
<tr>
<th>Standard</th>
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<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Provide adequate storage for safety equipment and supplies in the BSL-3 suite, including biological and chemical spill kits.</td>
<td>Readily available safety equipment will facilitate response in an emergency.</td>
<td>N/A</td>
</tr>
<tr>
<td>2. Provide binocular, hands-free eyewash.</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>3. Provide a combination emergency shower/eyewash unit. Coordinate exact location(s) with EH&amp;S. Locate unit in a central location within the BSL-3 laboratory/suite. Emergency shower/eyewash unit location must be compliant with ANSI 358.1-2014 and Cal-OSHA regulations.</td>
<td>N/A</td>
<td>Flow alarms should be provided on emergency showers in lab spaces to notify the staff of possible issue(s).</td>
</tr>
<tr>
<td>4. Provide mounted type ABC fire extinguisher at contained side of anteroom door. Risk assessment and Fire Marshal requirements shall determine need for additional fire extinguishers. The maximum travel distance to an extinguisher shall be 15 m (50 ft.). For open labs, the fire extinguisher shall be located as closely as possible to the exit access doors.</td>
<td>Provide type ABC fire extinguisher at contained side of anteroom door. Risk assessment and Fire Marshal requirements shall determine need for additional fire extinguishers.</td>
<td>Provide in gasket, recessed cabinet.</td>
</tr>
</tbody>
</table>
5. BSL-3 facilities shall be fully protected with an automatic sprinkler system based on local regulations. Fire sprinkler heads, if in canopy hoods, should be rated higher than the steam temperature. ⁹

6. Sprinkler heads shall be pendent type and not be recessed or concealed. The piping drop shall extend through the penetration sufficient to allow for application of a visible seal. Escutcheons shall not be provided. A flat solid stainless steel plate or washer that is tight fitting against the pipe may be utilized bedded in sealant and sealed to the pipe circumference. Pipes shall be braced to prevent movement that may damage the barrier seal. ¹⁰

7. Provide space in anteroom for storage of clean PPE. Provide areas for laundry and/or waste bin for disposal of dirty PPE as necessary. Provide space and outlets for PAPR charging station and adequate space for donning/doffing procedures.

8. Provide local audible and visual alarms for the following: ¹¹
   - Fire hazard
   - HVAC failure
   - Differential pressure at any door out

5. Fire sprinkler sensors for placement near autoclaves or other high heat/steam generating equipment should be designed to prevent nuisance triggering and floods

6. N/A

7. Considerations for privacy and changing areas should be made

8. Consider also reporting alarm to PI, facilities or campus security/police, as appropriate for a given alarm condition.
   - Freezer and incubator alarms
9. Alarms must be on UPS power.  

<table>
<thead>
<tr>
<th>Standard</th>
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</tr>
</thead>
</table>
| 10. Provide pass-through autoclave in the BSL-3 suite  
  - Provide exhaust hoods on the clean and dirty sides of double door autoclaves. The dirty side canopy hood must connect to the BSL-3 exhaust. The clean side canopy hood is connected to the general lab exhaust. 
  - Provide space adjacent to the autoclave on the dirty and clean side of pass-through autoclave for waste collection and disposal. | 10. N/A | 10. N/A |
| 11. Rooms shall be capable of being sealed for protocols that require gaseous and/or vapor decontamination. | 11. The facility should be capable of being fully sealed for decontamination purposes. | 11. Consider placement of decontamination ports that are accessible from the outside of the facility. Decontamination corridor integrated into the design for... |
12. Liquid effluent from BSL-3 autoclaves may be discharged to sanitary sewer or reclaimed as appropriate.  

13. Air evacuated from the autoclave chamber prior to decontamination cycle shall be considered potentially contaminated and requires HEPA filtration or other validated method.

14. Class II Biological Safety Cabinets must meet the NSF/ANSI 49 standard.  

- Class III BSC may be required based on a risk assessment.

12. Based on risk assessment, liquid from the BSL-3 autoclave may need to be treated prior to disposal to municipal sewer.

13. Provide HEPA filter at the autoclave vacuum system to minimize release of infectious material during purging phase.

14. Primary containment is required for manipulations of infectious materials.

Facility should be designed to allow for individual rooms or zones to be isolated for decontamination.
Safety and Decontamination References

1. Evaluate availability of emergency equipment. **NIH BSL-3 Certification Section I.3**
2. An eyewash station must be readily available in the laboratory. **BMBL 5th ed. Section IV.Biosafety Level 3.D.8**
3. A BSL-3 environment shall have: Hands free hand washing sink with eyewash. **NIH DRM 8.2.10.1**
4. Available hands-free sink near laboratory exit. **NIH BSL-3 Certification Section I.3**
5. In spaces where a significant hazard exists and it is likely a user may be present without supervision, a flow alarm shall be provided to indicate emergency shower operation. The alarm shall provide local audible alert and remote alert. **NIH DRM Section 8.2.10.D**
6. Validate availability of emergency shower. **NIH BSL-3 Certification Section II.13**
7. All fire extinguisher cabinets shall be sized to contain the approved fire extinguisher. Cabinet doors shall not have locks. The maximum travel distance to an extinguisher shall be 15 m (50 ft.). For open labs, the fire extinguisher shall be located as closely as possible to the exit access doors. **NIH DRM Section 9.4.4**
8. Evaluate availability of working fire extinguisher. **NIH BSL-3 Certification Section I.3**
9. BSL-3 facilities shall be full protected with an automatic sprinkler system. **NIH DRM Section 9.5.1.1.A**
10. Sprinkler heads shall be pendent type and not be recessed or concealed. The piping drop shall extend through the penetration sufficient to allow for application of a visible seal. Escutcheons shall not be provided. A flat solid stainless steel plate or washer that is tight fitting against the pipe may be utilized bedded in sealant and sealed to the pipe circumference. Brace pipes to prevent movement that may damage the barrier seal. **NIH DRM Section 9.5.1.1.B, 9.5.1.1.D**
11. In the event of loss of directional airflow, personnel in the containment space shall be notified of such condition via audible and/or visual alarms. Alarm indication should be located inside the laboratory space(s), at the entrance of the BSL-3. **ANSI Z9.14 Section 8.3.3**
12. Power to BMS and other control systems for monitoring, controlling, and alarming HVAC normal and failure modes should be on UPS. **ANSI Z9.14 Sec. 8.4.6**
13. A method for decontaminating all laboratory wastes should be available in the facility, preferably within the laboratory (e.g., autoclave, chemical disinfection, or other validated decontamination method). **BMBL 5th ed. Section IV.Biosafety Level 3.D.11**
14. Provide stainless steel exhaust canopy hoods over door to capture steam (both sides). “Dirty” side canopy exhaust should be tied into contaminant exhaust. “Clean” side canopy exhaust may be tied into non-contaminant exhaust. **NIH DRM Section 6.6.10.A**
15. Autoclave for decontaminating laboratory wastes is available. **NIH Guidelines Appendix G-II-C-4-h**
16. Equipment that penetrates secondary barrier shall be provided with a manufacturer-supplied bioseal. Bioseal shall form a continuous, airtight seal. **NIH DRM Section 4.9.11.B**
17. Selection of the appropriate materials and methods used to decontaminate the laboratory must be based on the risk assessment. **BMBL 5th ed. Section IV.Biosafety Level 3.D.3**
18. All penetrations into and through partitions, floors, and ceilings shall be sealed to enhance sanitation, facilitate gas and vapor decontamination, and resist air infiltration. **NIH DRM Section 2.6.2.5.**
19. Gas tightness for decontamination **NIH BSL-3 Certification Section I.2**
20. The sterilized effluent from BSL-3 autoclaves may discharge through sanitary system or general building laboratory waste system as an indirect connection through a floor sink. The drain receptor shall be located on the clean side of the bioseal (including with the clean utility access area.) **NIH DRM Section 8.6.9.P**
21. All procedures involving the manipulation of infectious materials must be conducted within a BSC, or other physical containment devices. No work with open vessels is conducted on the bench. When a procedure cannot be performed within a BSC, a combination of personal protective equipment and other containment devices, such as a centrifuge safety cup or sealed rotor must be used. BMBL 5th ed. Section IV. Biosafety Level 3.B.10

22. Refer to Appendix F "Biological Safety Cabinet (BSC) Placement Requirements for New Buildings and Renovations" for BSC placement guidelines. NIH DRM Appendix F

23. Verify correct placement of biological safety cabinets with respect to supply and exhaust diffusers, doors and traffic patterns. NIH BSL-3 Certification Section II.6
### Autoclaves

<table>
<thead>
<tr>
<th>Minimum Standards</th>
<th>Explanation</th>
<th>Best Design Practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Provide an autoclave within the laboratory.</td>
<td>1. N/A</td>
<td>1. Provide a pass-through autoclave opening to a secure room outside the BSL-3 suite.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Pass through autoclaves penetrating the BSL-3 barrier should have an approved bioseal. The body of the autoclave should be on the outside of the containment barrier.</td>
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<tr>
<td></td>
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<td>• Autoclave doors should be interlocked.</td>
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<td>• Floor sink for the autoclave should be on the outside of the containment barrier.</td>
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<td></td>
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<td>• Clean side autoclave doors should not open directly onto a public corridor but should open into a vestibule with doors separating it from the public corridor.</td>
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<td></td>
<td></td>
<td>• Provide exhaust hoods on the clean and dirty sides of double door autoclaves. The dirty side canopy hood should connect to the BSL-3 exhaust. The clean side canopy hood can be connected to the general lab exhaust.</td>
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<td></td>
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<td>• Provide space adjacent to the</td>
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</table>
2. Autoclave size, accessories and decontamination cycles shall be determined during the design phase in coordination with EH&S.  

### Biosafety Cabinets

<table>
<thead>
<tr>
<th>Minimum Standards</th>
<th>Explanation</th>
<th>Best Design Practice</th>
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</thead>
<tbody>
<tr>
<td>3. BSCs must meet the NSF/ANSI 49 standard. Include space for seismic bracing. 18, 19, 20</td>
<td>Locate BSCs to minimize air current effects of passing traffic and door openings.</td>
<td>BSCs should be placed according to the NIH DRM Appendix A. Adhere to any UC ergonomics policies. • Perform a risk assessment to determine if the BSCs can be safely exhausted back into the room or if they require a thimble or hard connection. Hard ducted cabinets (B1 or B2) cannot be installed without consultation with EH&amp;S.</td>
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4. For labs that will be performing aerosol challenges, consideration must be given to Class III BSC with enhancements (e.g.: double door sterilizer or dunk tank) or other primary containment devices. Work with HCLD and EH&S for risk assessment of specific aerosol challenges. 21, 22, 23 • Provide HEPA filtration for supply and exhaust of Class III BSCs. | Aerosol challenges shall be conducted within a class III biological safety cabinet, other primary containment device, or other approved inhalation - exposure system capable of withstanding rigorous decontamination by disinfectants required by the program. | N/A |
• Class III BSCs must be directly (hard) connected up through the second exhaust HEPA filter of the cabinet. Supply air must be provided in such a manner that prevents positive pressurization of the cabinet.

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<tr>
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<th>Explanation</th>
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<tbody>
<tr>
<td>Flow Cytometry</td>
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<tr>
<td>5. If flow cytometers are used in a BSL-3 facility, they must be installed within a biological safety cabinet or other primary containment device specifically designed for the purpose of containing any aerosols produced.</td>
<td>5. N/A</td>
<td>5. N/A</td>
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</tbody>
</table>

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<thead>
<tr>
<th>Minimum Standards</th>
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<th>Best Design Practice</th>
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<tr>
<td>Centrifuge</td>
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<td>6. Centrifuges shall be equipped with gasketed safety cups, sealed rotors or operated only within a BSC.</td>
<td>6. N/A</td>
<td>6. N/A</td>
</tr>
</tbody>
</table>
BSL-3 Standard Equipment References

1. A method for decontaminating all laboratory wastes should be available in the facility, preferably within the laboratory (e.g., autoclave, chemical disinfection, or other validated decontamination method). **BMBL 5th ed. Section IV.Biosafety Level 3.A.8, Section IV.Biosafety Level 3.D.11**

2. Solid waste shall be decontaminated by use of an autoclave. Autoclaves with a double door pass through function are preferable, but single door autoclaves are acceptable. **NIH DRM Section 4.9.10**

3. An autoclave for decontaminating laboratory wastes is available preferably within the laboratory. **NIH Guidelines Appendix G-II-C-4-h**

4. All equipment integral to the containment barrier requires a manufacturer supplied biological seal in the containment wall. The biological seal shall be a structural stable, mechanically fastened gasketed seal capable of containing decontaminating gas and allowing for differential movement and remaining intact for gaseous decontamination protocols. **NIH DRM Section 2.6.2.5**

5. Design documents shall include large scale penetration details for each type of barrier wall, floor and ceiling penetration including bioseals. **NIH DRM Section 2.6.4.1**

6. Penetration seals shall be readily visible (when applicable) for inspection, cleaning, and maintenance. **ANSI Z9.14 Section 8.3.2**

7. Pass through autoclaves shall not permit door opening on the clean side prior to completion of a validated decontamination cycle. **NIH DRM Section 4.9.10**

8. Interlocks shall be connected to the Building Automation System. **NIH DRM Section 4.9.3.2**

9. A sink shall be located in the clean side of autoclaves for disposal of sterilized liquid waste. **NIH DRM Section 4.9.10**

10. The floor sink for receiving the sterilized chamber condensate shall be located on the clean side of the bioseal, typically in the utility service access space. **NIH DRM Section 4.9.10**

11. A pass-through autoclave from containment to an anteroom outside of the containment barrier eliminates the need for transport of contaminated material outside of containment. **NIH DRM Section 2.5.4.2**

12. Clean side autoclave doors shall not open directly onto a public corridor but shall open into a vestibule with doors separating it from the laboratory. **NIH DRM Section 4.9.10**

13. The room shall have adequate exhaust capacity to remove heat, steam, and odors generated by the use of the autoclave. **NIH DRM Section 4.6.1.12.C**

14. A canopy exhaust hood shall be provided above the autoclave door to capture steam and odors. In two-door pass through configurations, a canopy hood is required above the non-containment side door. A second exhaust hood may be required over the containment side door if the program calls for autoclaving materials into the containment barrier. Canopy hoods on the containment side shall be connected to the HEPA-filtered containment zone exhaust system. Canopy hoods on the non-containment side shall be connected to the non-containment exhaust system. **NIH DRM Section 4.9.10**

15. Each laboratory area shall be provided with designated space for the safe storage of biological. **NIH DRM Section 2.1.3.7.4**

16. Laboratories shall be designed with a room for temporary storage of hazardous waste. **NIH DRM Section 1.11.3.4**

17. Risk Assessment, Systems Failure & Disaster Mitigation **NIH DRM Section 1.15.6**

18. All procedures involving the manipulation of infectious materials must be conducted within BSCs or other physical containment devices. **BMBL 5th ed. Section IV.Biosafety Level 3**

19. Placement of fume hoods and BSCs in laboratories shall be in accordance with the Biological Safety Cabinet Placement Guide. **NIH DRM**
Appendix A

20. BSC’s should be located away from doors, heavily traveled laboratory areas, and other possible airflow disruptions. **BMBL 5th ed. Section IV.Biosafety Level 3.D.6**

21. Class III BSCs must be directly and independently exhausted through two HEPA filters in series. **BMBL 5th ed. Section IV.Biosafety Level 3.D.10**

22. Class III BSCs must be directly (hard) connected up through the second exhaust HEPA filter of the cabinet. Supply air must be provided in such a manner that prevents positive pressurization of the cabinet. **BMBL 5th ed. –Section IV.Biosafety Level 3.D.10**

23. Aerosol challenges shall be conducted within a class III biological safety cabinet or other approved inhalation- exposure equipment capable of withstanding rigorous decontamination using chemicals required by the program. All materials in the containment cabinet shall be decontaminated prior to exiting the cabinet. An in-line double-door sterilizer and a dunk tank or fumigation integral to the BSC may be required based on risk assessment and program requirements. **NIH DRM Section 4.9.11.C**

24. Equipment that may produce infectious aerosols must be contained in primary barrier devices that exhaust air through HEPA filtration or other equivalent technology before being discharged into the laboratory. These HEPA filters should be tested and/or replaced at least annually. **BMBL 5th ed. Section IV.Biosafety Level 3.D.12**

25. Ensure that continuous flow centrifuges or other equipment that may produce aerosols are contained in devices that exhaust air through HEPA filters before discharge into the laboratory. **NIH BSL-3 Certification Section II-10**

26. Equipment that may produce infectious aerosols must be contained in primary barrier devices that exhaust air through HEPA filtration or other equivalent technology before being discharged into the laboratory. These HEPA filters should be tested and/or replaced at least annually. **BMBL 5th ed. Section IV.Biosafety Level 3.D.12**

Procedures and equipment used routinely for handling infectious agents in laboratories, such as pipetting, blenders, non-self-contained centrifuges, sonicators, and vortex mixers are proven sources of aerosols. **BMBL 5th ed. Section II.Hazardous Characteristics of Laboratory Procedures**
Appendix I

UC BSL3 Design Standards Deviation Request Form

To initiate a deviation request, present this completed form to your campus High Containment Laboratory Oversight Group (HCLOG) for review. Once approved by the campus HCLOG, the request will be provided to the systemwide High Containment Laboratory Oversight Committee (HCLOC) for final review.

As part of the review process, the HCLOC Chair will select a subcommittee of five designated reviewers. One member shall be an HCLOG chair from a location other than the submitting campus, one member shall be the HCLD from the submitting campus, and the remaining three members will be chosen by the HCLOC Chair or Vice Chair. The subcommittee will strive to provide the submitting campus a decision within 10 working days of receiving all the necessary information.

Please note financial restrictions do not outweigh safety requirements. Requests for deviations cannot be based solely on financial constraint.

---

**Campus:**

**High Containment Lab Director:**

**Campus BSL3 Building Location Details** (see request for relevant documentation below):

**Principle Investigator(s):**

**Date:** Click or tap to enter a date.

---

**UC BSL3 Design Standard** – provide the standard section or sections you are seeking to deviate from

- **Section:**
- **Number:**
- **State the standard:**

**Justification for Deviation**

- Explain why you are unable to adhere to the standard:
- Discuss any alternate plans for mitigation:
- Discuss any possible risks associated with the deviation:

**Proposed research**

- Overview of research projects to be conducted in the facility:
- List proposed agents/toxins in use:
Select Agents:
Select Toxins:
Tier 1 Select Agents/Toxins:
List any other applicable Federal Agency Requirements:

List experimental procedures:
Describe standard laboratory procedures:
Describe any proposed high-risk procedures:
Discuss any future projects or plans for this facility:

Personnel
How many BSL3 users:
Support staff:

Facility Layout – provide a full description of the space
Describe the laboratory space:
Is there an ante room:
Will there be ABSL3 space:

Relevant documentation – attach the following:
1. Laboratory Floor Plans
2. Building Floor Plans
3. Facility Specifications
4. Mechanical Plans
5. Approved BUAs or Research Summaries

Submitting Campus Signatures:
HCLOG Chair
HCLD
HCLOG Review Date: Click or tap to enter a date.

HCLOC Review Notes:
Date Received: Click or tap to enter a date.
HCLOC review notes and decision:
Date Returned to submitting campus: Click or tap to enter a date.