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BL3-N	APPENDIX Q. PHYSICAL AND BIOLOGICAL CONTAINMENT FOR RECOMBINANT DNA RESEARCH INVOLVING ANIMALS March 2013	N/A	Conform	Non- Conform	Comments
	Appendix Q-II-C-1. Standard Practices (BL3-N)				
	Appendix Q-II-C-1-a. Animal Facility Access (BL3-N)				
	Appendix Q-II-C-1-a-(1). The containment area shall be locked.				
	Appendix Q-II-C-1-a-(2). The containment area shall be patrolled or monitored at frequent intervals.				
	Appendix Q-II-C-1-a-(3). The containment building shall be controlled and have a locking access.				
	Appendix Q-II-C-1-a-(4). The Animal Facility Director shall establish policies and procedures whereby only persons who have been advised of the potential hazard and who meet any specific entry requirements (e.g., vaccination) shall enter the laboratory or animal rooms.				
	Appendix Q-II-C-1-a-(5). Animal room doors, gates, or other closures shall be kept closed when experiments are in progress.				
	Appendix Q-II-C-1-b. Decontamination and Inactivation (BL3-N)				
	Appendix Q-II-C-1-b-(1). The work surfaces of containment equipment shall be decontaminated when work with organisms containing recombinant DNA molecules is finished. Where feasible, plastic-backed paper toweling shall be used on nonporous work surfaces to facilitate clean-up.				
	Appendix Q-II-C-1-b-(2). All animals shall be euthanized at the end of their experimental usefulness and the carcasses decontaminated before disposal in an approved manner.				
	Appendix Q-II-C-1-b-(3). Needles and syringes shall be promptly placed in a puncture-resistant container and decontaminated, preferably by autoclaving, before discard or reuse.				
	Appendix Q-II-C-1-b-(4). Special safety testing, decontamination procedures, and Institutional Biosafety Committee approval shall be required to transfer agents or tissue/organ specimens from a BL3-N animal facility to a facility with a lower containment classification.				
	Appendix Q-II-C-1-b-(5). Liquid effluent from containment equipment, sinks, biological safety cabinets, animal rooms, primary barriers, floor drains, and sterilizers shall be decontaminated by heat treatment before being released into the sanitary system. The procedure used for heat decontamination of liquid wastes shall be monitored with a recording thermometer. The effectiveness of the heat decontamination process system shall be revalidated every 30 days with an indicator organism.				
	Appendix Q-II-C-1-c. Signs (BL3-N)				

Appendix Q-II-C-1-c-(1). When the animal research requires special provisions for entry (e.g., vaccination), a warning sign incorporating the	
universal biosafety symbol shall be posted on all access doors to the animal work area. The sign shall indicate: (i) the agent, (ii) the animal species, (iii) the name and telephone number of the Animal Facility Director or other responsible individual, and (iv) any special requirements for entering the laboratory.	
Appendix Q-II-C-1-d. Protective Clothing (BL3-N)	
Appendix Q-II-C-1-d-(1). Full protective clothing that protects the individual (e.g., scrub suits, coveralls, uniforms) shall be worn in the animal area. Clothing shall not be worn outside the animal containment area and shall be decontaminated before laundering or disposal. Personnel shall be required to shower before exiting the BL3-N area and wearing of personal clothing.	
Appendix Q-II-C-1-d-(2). Special care shall be taken to avoid skin contamination with microorganisms containing recombinant DNA. Impervious and/or protective gloves shall be worn when handling experimental animals and when skin contact with an infectious agent is unavoidable.	
Appendix Q-II-C-1-d-(3). Appropriate respiratory protection shall be worn in rooms containing experimental animals.	
Appendix Q-II-C-1-e. Records (BL3-N)	
Appendix Q-II-C-1-e-(1). Documents regarding experimental animal use and disposal shall be maintained in a permanent record book.	
Appendix Q-II-C-1-e-(2). Any incident involving spills and accidents that result in environmental release or exposure of animals or laboratory workers to organisms containing recombinant DNA shall be reported immediately to the Biological Safety Office, Animal Facility Director, Institutional Biosafety Committee, NIH/OBA, and other appropriate authorities (if applicable). Reports to the NIH/OBA shall be sent to the Office of Biotechnology Activities, National Institutes of Health, 6705 Rockledge Drive, Suite 750, MSC 7985, Bethesda, MD 20892-7985 (20817 for non-USPS mail), 301-496-9838, 301-496-9839 (fax). Medical evaluation, surveillance, and treatment shall be provided as appropriate and written records maintained. If necessary, the area shall be appropriately decontaminated.	
Appendix Q-II-C-1-e-(3). When appropriate and giving consideration to the agent handled, baseline serum samples shall be collected and stored for animal care and other at-risk personnel. Additional serum specimens may be collected periodically depending on the agent handled or the function of the facility.	
Appendix Q-II-C-1-f. Transfer of Materials (BL3-N)	

 Appendix Q-II-C-1-f-(1). Biological materials removed from the animal containment laboratory in a viable or intact state shall be transferred to a non-breakable sealed primary container and then enclosed in a non-breakable sealed secondary container. All containers, primary and secondary, shall be disinfected before removal from the animal facility. Advance approval for transfer of material shall be obtained from the Animal Facility Director. Packages containing viable agents may be opened only in a facility having an equivalent or higher level of physical containment unless the agent is biologically inactivated or incapable of reproduction. Appendix Q-II-C-1-f-(2). Special safety testing, decontamination 	
procedures, and Institutional Biosafety Committee approval shall be required to transfer agents or tissue/organ specimens from a BL3-N animal facility to a facility with a lower containment classification.	
Appendix Q-II-C-1-g. Other (BL3-N)	
Appendix Q-II-C-1-g-(1). All genetically engineered neonates shall be permanently marked within 72 hours after birth, if their size permits. If their size does not permit marking, their containers should be marked. In addition, transgenic animals should contain distinct and biochemically assayable DNA sequences that allow identification of transgenic animals from among non-transgenic animals.	
Appendix Q-II-C-1-g-(2). Appropriate steps should be taken to prevent horizontal transmission or exposure of laboratory personnel. If the agent used as the vector is known to be transmitted by a particular route (e.g., arthropods), special attention should be given to preventing spread by that route. In the absence of specific knowledge of a particular route of transmission, all potential means of horizontal transmission (e.g., arthropods, contaminated bedding, or animal waste) should be prevented.	
Appendix Q-II-C-1-g-(3). Eating, drinking, smoking, and applying cosmetics shall not be permitted in the work area.	
Appendix Q-II-C-1-g-(4). Individuals who handle materials and animals containing recombinant DNA molecules shall be required to wash their hands before exiting the containment area.	
Appendix Q-II-C-1-g-(5). Experiments involving other organisms that require containment levels lower than BL3-N may be conducted in the same area concurrently with experiments requiring BL3-N containment provided that they are conducted in accordance with BL3-N practices.	
Appendix Q-II-C-1-g-(6). Animal holding areas shall be cleaned at least once a day and decontaminated immediately following any spill of viable materials. Appendix Q-II-C-1-g-(7). All procedures shall be performed carefully to	
minimize the creation of aerosols.	
Appendix Q-II-C-1-g-(8). A double barrier shall be provided to separate male and female animals unless reproductive studies are part of the experiment or other measures are taken to avoid reproductive transmission. Reproductive incapacitation may be used.	

	Appendix Q-II-C-1-g-(9). The containment area shall be in accordance		
	with state and Federal laws and animal care requirements. Appendix Q-II-C-1-g-(10). All animals shall be euthanized at the end of	 	
	their experimental usefulness and the carcasses decontaminated		
	before disposal in an approved manner.		
	Appendix Q-II-C-1-g-(11). Personnel shall be required to shower before		
	exiting the BL3-N area and wearing personal clothing.		
	Appendix Q-II-C-1-g-(12). Animals of the same or different species,		
	which are not involved in the work being performed, shall not be		
	permitted in the animal area.		
	Appendix Q-II-C-1-g-(13). Needles and syringes shall be used only for		
	parenteral injection and aspiration of fluids from laboratory animals and		
	diaphragm bottles. Only needle-locking syringes or disposable syringe-		
	needle units (i.e., needle is integral to the syringe) shall be used for the		
	injection or aspiration of fluids containing organisms that contain		
	recombinant DNA. Extreme caution shall be used when handling		
	needles and syringes to avoid autoinoculation and the generation of aerosols during use and disposal. Following use, needles shall not be		
	bent, sheared, replaced in the needle sheath or guard or removed from		
	the syringe. The needles and syringes shall be promptly placed in a		
	puncture-resistant container and decontaminated, preferably by		
	autoclaving, before discard or reuse.		
	Appendix Q-II-C-1-g-(14). A biosafety manual shall be prepared or	 	
	adopted. Personnel shall be advised of special hazards and required to		
	read and follow instructions on practices and procedures.		
	Appendix Q-II-C-2. Animal Facilities (BL3-N)		
	Appendix Q-II-C-2-a. Animals shall be contained within an enclosed		
	structure (animal room or equivalent) to minimize the possibility of theft		
	or unintentional release and avoid arthropod access. The special		
	provision to avoid the entry or escape of arthropods from the animal		
	areas may be waived if the agent in use is not known to be transmitted		
	by arthropods.		
	Appendix Q-II-C-2-b. The interior walls, floors, and ceilings shall be		
	impervious to water and resistant to acids, alkalis, organic solvents,		
	and moderate heat, to facilitate cleaning. Penetrations in these		
<u> </u>	structures and surfaces (e.g., plumbing and utilities) shall be sealed. Appendix Q-II-C-2-c. Windows in the animal facility shall be closed,		
	sealed, and breakage resistant (e.g., double-pane tempered glass or		
	equivalent). The need to maintain negative pressure should be		
	considered when constructing or renovating the animal facility.		
	Appendix Q-II-C-2-d. An autoclave, incinerator, or other effective means		
	to decontaminate animals and waste shall be available, preferably		
	within the containment area. If feasible, a double-door autoclave is		
	preferred and should be positioned to allow removal of material from		
	the containment area.		
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ag are op us	opendix Q-II-C-2-e. If arthropods are used in the experiment or the ent under study can be transmitted by an arthropod, the interior work ea shall be appropriately screened (52 mesh). All perimeter joints and enings shall be sealed, and additional arthropod control mechanisms ed to minimize arthropod entry and propagation, including propriate screening, or the equivalent of access doors.		
Ар	ppendix Q-II-C-2-f. Access doors to the containment area shall be self- osing.		
Ap are oth ph are sh	opendix Q-II-C-2-g. The animal area shall be separated from all other eas. Passage through two sets of doors shall be the basic quirement for entry into the animal area from access corridors or her contiguous areas. The animal containment area shall be ysically separated from access corridors and other laboratories or eas by a double-door clothes change room, equipped with integral owers and airlock.		
sin dra be he rec pro	opendix Q-II-C-2-h. Liquid effluent from containment equipment, hks, biological safety cabinets, animal rooms, primary barriers, floor ains, and sterilizers shall be decontaminated by heat treatment fore being released into the sanitary system. The procedure used for eat decontamination of liquid wastes shall be monitored with a cording thermometer. The effectiveness of the heat decontamination occess system shall be revalidated every 30 days with an indicator ganism.		
pro the ex if t aw	opendix Q-II-C-2-i. An exhaust air ventilation system shall be ovided. This system shall create directional airflow that draws air into e animal room through the entry area. The building exhaust, or the haust from primary containment units, may be used for this purpose he exhaust air is discharged to the outside and shall be dispersed vay from occupied areas and air intakes. Personnel shall verify that e direction of the airflow (into the animal room) is proper.		
Ар	ppendix Q-II-C-2-j. If the agent is transmitted by aerosol, then the haust air shall pass through a high efficiency particulate air/HEPA		
	ppendix Q-II-C-2-k. Vacuum lines shall be protected with high iciency particulate air/HEPA filters and liquid disinfectant traps.		
Ap an ca sp ex an	ppendix Q-II-C-2-I. In lieu of open housing in the special animal room, imals held in a BL3-N area may be housed in partial-containment ging systems (e.g., Horsfall units or gnotobiotic systems, or other ecial containment primary barriers). Prudent judgment must be ercised to implement this ventilation system (e.g., animal species) d its discharge location.		
au ne	opendix Q-II-C-2-m. Each animal area shall contain a foot, elbow, or tomatically operated sink for hand washing. The sink shall be located ar the exit door.		
	opendix Q-II-C-2-n. Restraining devices for animals may be required avoid damage to the integrity of the animal containment facility.		