Exposure Incident
Report exposure immediately; you may need immediate therapy.

- **Needlesticks/puncture wounds:**
  Wash the affected area with antiseptic soap and warm water for 15 minutes

- **Mucous membrane exposure:**
  Flush the affected area for 15 minutes using an eyewash.

For all exposure incidents:

- Notify Principal Investigator, manager or supervisor (if available) to initiate accident or exposure incident report.

- Seek medical assistance immediately (within 1-2 hours) from Yale Health Center, Acute (203-432-0123), Employee Health (203-432-7978) 8:30 a.m. to 5:00 p.m. Monday-Friday. Medical Area employees may also go to the Yale-New Haven Hospital (Y-NHH) Occupational Health Services (203-688-2462), East Pavilion room 40 (behind cafeteria) from 7:30 a.m. to 4:00 p.m. Monday-Friday (excluding holidays) or the Y-NHH Emergency Room (203-688-2222) from 4:00 p.m. to 7:30 a.m.

All employees should receive follow up care through Yale Employee Health (203-432-7978)

Emergency Phone Numbers

Police and Fire, on campus: 911

Yale Health Center: Acute Care 203-432-0123

Yale Employee Health Monday – Friday 8:30am to 5:00pm 203-432-7978

EHS Emergency Numbers 203-785-3555
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SECTION 1.0 INTRODUCTION

Healthcare workers' concern for their health and safety increased as the AIDS epidemic was recognized. The Centers for Disease Control (CDC) responded to these concerns and developed a new approach to infection control - Universal Precautions.

Universal Precautions are protective measures employees use to eliminate or minimize exposure to infectious agents that may be present in human blood, tissues or certain body fluids. Universal Precautions are based upon the premise that all human blood, tissues and certain body fluids are treated as if known to be infectious for HIV, HBV and other bloodborne pathogens. Individuals who handle blood or these body fluids must wear appropriate personal protective equipment to prevent contact with potentially infectious materials.

The Occupational Safety and Health Administration (OSHA) incorporated many of CDC's Universal Precautions guidelines into a standard titled "Occupational Exposure to Bloodborne Pathogens". The standard was published on December 6, 1991 and a copy can be found in Appendix A of this manual. The OSHA standard outlines: (1) the criteria employers are to use to determine who is potentially exposed to bloodborne pathogens; (2) the elements of an acceptable exposure control plan; (3) the acceptable methods for reducing or eliminating potential exposure (engineering and work practice controls, personal protection equipment and housekeeping); (4) the information on bloodborne pathogen hazards that must be communicated to all potentially exposed employees; (5) the special training and work practice requirements in HIV and HBV research laboratories and production facilities; (6) the employer's obligation to provide Hepatitis B vaccination to all potentially exposed employees and post-exposure follow-up to employees exposed during incidents; and (7) the records that must be maintained.

The bloodborne pathogen standard applies to all employers with "occupationally exposed" employees. OSHA's regulatory definition of "occupational exposure" in this standard departs from its use in other standards. In other standards, occupational exposure has referred to an actual exposure event. In the bloodborne pathogen standard, OSHA defined "occupational exposure" as reasonably anticipated skin, eye, mucous membrane, or parenteral contact with human blood or "other potentially infectious materials" that may result from performance of an employee's duties. OSHA uses the term "exposure incident" to refer to an actual exposure. Therefore an individual is considered occupationally exposed even if he/she does not have direct contact with blood or other potentially infectious material.

In the regulation, OSHA defines "other potentially infectious materials" as:

"...semen, vaginal secretions, cerebrospinal fluids, synovial fluids, pleural fluid, pericardial fluid, peritoneal fluid, amniotic fluid, saliva in dental procedures, any body fluid that is visibly contaminated with blood, and all body fluids in situations where it is difficult or impossible to differentiate between body fluids; any unfixed tissue or organ (other than intact skin) from a human (living or dead); and human immunodeficiency virus (HIV) - containing cell or tissue cultures, organ cultures, and HIV- or Hepatitis B virus (HBV) - containing culture medium or other solutions; and body organs, or other tissues from experimental animals infected with HIV or HBV." OSHA also considers all primary and continuous cell cultures as potentially infectious if not screened and shown negative for the presence of all bloodborne pathogens.

SECTION 2.0 YALE UNIVERSITY'S EXPOSURE CONTROL PLAN

The information in the Exposure Control Plan is intended to eliminate or minimize employee exposure to bloodborne pathogens and outlines how Yale University will comply with all provisions of the OSHA bloodborne standard. The Yale University Exposure Control Plan also identifies the job classifications of all "occupationally exposed" employees. All employers with occupationally exposed employees must prepare a written exposure determination that provides the job classifications or titles and the tasks and procedures that may involve exposure. Employees are required to perform their exposure determination by completing the Training Requirement Assessment through the Training Management System (TMS) http://www.yale.edu/training/. The Training Requirement Assessment must be completed at time of hire by new employees and thereafter annually.
This Exposure Control Plan is reviewed annually and updated at least annually or when necessary to reflect new or modified tasks and procedures, or new and revised employee positions with occupational exposure.

Yale's Exposure Control Plan is available at the following locations to assure employee access to this document:

- Chemistry Department Instrument Room
- Kline Chemistry Laboratory
- 255 Prospect Street
- Engineering Library
- Becton Engineering & Applied Science Ctr
- 15 Prospect Street
- Law School Library
- 127 Wall Street
- Yale Environmental Health & Safety
- 135 College Street, Suite 100

- Divinity School Library
- 409 Prospect Street
- Kline Science Library
- Kline Biology Tower
- 219 Prospect Street
- Medical School Library
- 333 Cedar Street
- Yale Health Center
- Acute Care
- 55 Lock Street

The Exposure Control Plan can be retrieved on the World Wide Web on the Yale Environmental Health and Safety website (http://ehs.yale.edu/) in the Policies and Program Section.

SECTION 3.0 TRAINING REQUIRED BY OSHA'S BLOODBORNE PATHOGEN STANDARD

The bloodborne pathogens training required by OSHA's standard must be performed at the time of employment and at least annually thereafter. The specific elements of an acceptable training program are outlined in the OSHA standard [1910.1030 (g)(2)], which can be found in Appendix A. This training manual contains all of the required training information.

Additional training is required for individuals working in HIV and HBV research laboratories. HIV and HBV research laboratories are laboratories engaged in the culture, production, concentration, experimentation and manipulation of HIV and HBV. Clinical, diagnostic or other research laboratories engaged solely in the analysis of human blood, tissues or organs are not HIV or HBV research laboratories for the purposes of this standard. The additional training requirements for HIV and HBV research laboratories are outlined in the OSHA standard [1910.1030 (g)(2)(ix)] which can be found in Appendix A. This "laboratory specific" training must be performed by the Principal Investigator or laboratory supervisor.

If you have any questions regarding the training information in this manual, or Yale University's plan to control employee exposure to bloodborne pathogens, you may contact a member of the Occupational Safety and Health staff at 203-785-3550.

SECTION 4.0 EXPOSURE CONTROL RESPONSIBILITIES

Each of us is partially responsible for our health and safety on the job. We share responsibility for the welfare of other people in our work environment. Responsibilities identified in the Exposure Control Plan are highlighted below for your reference.

4.1 Occupationally Exposed Employees

Occupationally exposed employees are in a position to exert enormous control over situations in their workplaces. Their actions can prevent or create exposure risks for other employees. Occupationally exposed employees shall:
• attend training seminars
• learn the information presented and apply it in the workplace
• ensure personal protective equipment and engineering controls are inspected periodically and function properly
• implement sign and label requirements, work practice controls and housekeeping duties
• correct deficiencies in control equipment if possible and report all safety deficiencies to their immediate supervisor
• participate in the University's Medical Surveillance Program
• follow emergency action, exposure incident and post-exposure procedures

4.2 Immediate Supervisor

Immediate supervisors provide a crucial link between the employees and the University and function as advocates of both. Immediate supervisors shall assure:
• all occupationally exposed employees are identified to the Office of Environmental Health and Safety and the Department of Employee Health within 10 days of hire
• all employees participate in a Bloodborne Pathogens "new hire" training seminar before they are assigned tasks where occupational exposure occurs and receive annual "refresher" training thereafter
• employees receive additional training when changes (modification of tasks or institution of new tasks) affect the employee's occupational exposure
• engineering controls are used whenever possible to eliminate or minimize employee exposure
• engineering controls, signs and labels are examined, maintained and replaced as necessary
• employees implement work practice controls
• personal protective equipment worn is appropriate for the task, is inspected before use, and functions properly
• emergency action procedures are understood and followed by all employees

4.3 Area Supervisor / Principal Investigators

Area supervisors and principal investigators are responsible for entire work units. They also provide a link between employees and the University. Area supervisors and principal investigators shall assure:
• all exposed employees and supervisors fulfill their responsibilities
• all shipping requirements are met
• cleaning schedules and decontamination methods are documented
• all equipment and work surfaces are cleaned and properly decontaminated
• special requirements for HIV and HBV research laboratories are fulfilled
4.4 Principal Investigators/Lab Supervisor Responsibilities Checklist

A. Responsibilities with new occupationally exposed employees

Principal Investigators/Area Supervisors must ensure:

- that an exposure determination has been completed for each new employee with occupational exposure to human blood or other potentially infectious materials (OPIM*);
- new occupationally exposed employees receive training prior to initiation of work with human blood or OPIM;
- new occupationally exposed employees have been offered the Hepatitis B vaccine within 10 working days of assignment.

New Employees with Occupational Exposure to Bloodborne Pathogens (include name and relevant dates)

<table>
<thead>
<tr>
<th>Name</th>
<th>Date of Assignment</th>
<th>Exposure Determination Completed</th>
<th>Initial Bloodborne Pathogen Training</th>
<th>HBV Vaccine Offered</th>
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</thead>
<tbody>
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</tbody>
</table>

*OPIM: Other potentially infectious materials include semen, vaginal secretions, cerebrospinal fluid, synovial fluid, pleural fluid, pericardial fluid, peritoneal fluid, amniotic fluid, any body fluid that is visibly contaminated with blood, any unfixed human tissue or organ, HIV and HBV containing cell culture solutions, and blood, unfixed organs or tissues from experimental animals infected with HIV or HBV, and sharp items contaminated with any of the materials listed above.

B. Use of Personal Protective Equipment

Identify tasks and procedures that may result in skin, eye, mucous membrane or parenteral contact with blood or OPIM, and list the personal protective equipment that will be utilized to minimize the exposure potential.

List Procedure and Check Required Protective Clothing

<table>
<thead>
<tr>
<th>Task or Procedure with Blood/OPIM Exposure</th>
<th>Lab Coat</th>
<th>Surgical Gloves</th>
<th>Face Shield</th>
<th>Eye Wear &amp; Surgical mask</th>
<th>Other: Solid Front Gown, Tyvek Jump Suit, Sleeve Covers, Booties, Head Cover, Respiratory Protection, etc.</th>
</tr>
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</tbody>
</table>
Identify the location where supplies of personal protective equipment are kept: _____________________________
____________________________________________________________________________________________

Identify the person responsible for maintaining the supply of personal protective equipment: ________________
____________________________________________________________________________________________

List the location where employees store personal protective equipment before leaving the work area (personal protective equipment must be removed before leaving the laboratory):
____________________________________________________________________________________________
____________________________________________________________________________________________

C. Engineering Controls

List the engineering controls that will be utilized to eliminate or minimize occupational exposure, and identify personnel responsible for maintaining or replacing on a regular schedule to ensure their effectiveness.

<table>
<thead>
<tr>
<th>Engineering Control</th>
<th>Personnel Responsible</th>
<th>Maintenance Schedule</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sharps Container</td>
<td>All lab members</td>
<td>Replace when 2/3 - 3/4 full</td>
</tr>
<tr>
<td>Plexiglass bench shield</td>
<td>All users</td>
<td>Decontaminate after use</td>
</tr>
<tr>
<td>Biowaste bags</td>
<td>All lab members</td>
<td>Replace when 2/3 - 3/4 full</td>
</tr>
<tr>
<td>Biosafety Cabinet</td>
<td>All users</td>
<td>Decontaminate before and after use, and immediately following a spill of blood or OPIM</td>
</tr>
<tr>
<td>Vacuum filters</td>
<td>All users</td>
<td>Replace when contaminated, wet or damaged</td>
</tr>
<tr>
<td>Plastic transport bins</td>
<td>All users</td>
<td>Decontaminate after use</td>
</tr>
<tr>
<td>Forceps and other mechanical means of sharps collection</td>
<td>All personnel</td>
<td>Decontaminate after use</td>
</tr>
</tbody>
</table>

D. Disinfection and Decontamination

The Principal Investigator/Lab Supervisor is responsible for ensuring that laboratory is kept neat and clean. Work surfaces and lab equipment must be decontaminated with a suitable disinfectant (such as 1-10% household bleach or an EPA registered tuberculocidal disinfectant) after use and immediately after a spill of human blood or OPIM.

Provide a schedule for general disinfection in the lab: _________________________________________________
____________________________________________________________________________________________
____________________________________________________________________________________________

Provide an outline of the decontamination and spill response procedure: ________________________________
____________________________________________________________________________________________
____________________________________________________________________________________________
List disinfectants that will be used for these purposes: _______________________________________________________

____________________________________________________________________________________________

Identify responsible personnel: ________________________________________________________________

____________________________________________________________________________________________

____________________________________________________________________________________________

E. Additional Responsibilities of the Principal Investigator/Lab Supervisor (Check the following has been completed)

- Antiseptic towels have been provided for pre-handwashing in areas without hand washing stations. Researchers have been instructed to wash hands as soon as feasible after the pre-wash.

- Equipment scheduled for moving, repair, or disposal has been appropriately decontaminated and labeled with a Biosafety Notice. All components that could not be decontaminated have been labeled with the biohazard symbol.

- Contaminated laundry is placed in a biohazard bag and sent to a service that practices Universal Precautions.

- Personnel understand the appropriate post-exposure response and follow-up procedures.

SECTION 5.0 BLOODBORNE PATHOGEN EPIDEMIOLOGY, SYMPTOMATOLOGY, TRANSMISSION

OSHA has determined that occupational exposure to human blood, tissues and body fluids poses a significant health risk because these may contain bloodborne pathogens such as:

- Human Immunodeficiency Virus
- Bloodborne Hepatitis Viruses (Hepatitis B Virus, Hepatitis D Virus, Hepatitis C Virus)
- Plasmodium species
- Treponema species
- Babesia species
- Brucella species
- Leptospira species
- Francisella species
- Streptobacillus moniliformis
- Spirillum minus
- Colorado Tick Fever Viruses (arboviruses- Colorado Tick fever viruses, Kemerovo, Lipovnik, Quarantin, Bhanja, Ganjam, Thogoto and Dugbe viruses)
- Borrelia species
- Creutzfeldt-Jakob agent
- Human T-lymphotropic Virus Type I

The greatest occupational exposure potential for the clinical and laboratory worker is a puncture wound from a sharp (such as needles, cutting instruments, broken glassware) contaminated with human blood, tissue or body fluid. Handling human specimens and direct contact (person to person) are also potential exposure routes. Direct and indirect contact with bloodborne pathogens enables them to enter the body through broken skin (parenteral entry) and through the mucous membranes of the eyes, nose, mouth and urogenital tract.
The bloodborne pathogens that cause malaria, tularemia, leptospirosis, relapsing fever, Lyme disease, brucellosis, viral hemorrhagic fever and Colorado tick fever can also be transmitted by insect bites or animal contacts. Clinicians and laboratorians whose work involves handling animals and/or insects should be aware of the potential occupational hazards associated with their work.

The table in Appendix B, lists the most common bloodborne pathogens, initial symptoms, routes of entry and transmission information. Contact Office of Environmental Health and Safety for more information concerning bloodborne pathogens. The following two sections review in detail the epidemiology, symptomatology and mode of transmission of AIDS and Viral Hepatitis.

5.1 AIDS: Human Immunodeficiency Virus (HIV)

HIV is a retrovirus that causes the Acquired Immune Deficiency Syndrome (AIDS) - a severe life-threatening illness which suppresses the body's ability to fight infection and can impede neurological function. There are two known strains, HIV-I and HIV-II. HIV-I is the etiologic agent of AIDS in North and South America, Europe and Central and East Africa. HIV-II is endemic only in West Africa.

HIV replicates primarily in human macrophages and T4 lymphocytes. Invasion of these two vital components of the immune system gradually depletes the number of cells necessary for normal immune function. As a result, an infected individual's susceptibility to opportunistic infections is increased.

AIDS was first described in 1981 in New York and Los Angeles during an unusual incidence of Pneumocystis carinii pneumonia and Kaposi's sarcoma in homosexual men who had no known underlying immunodeficiency. In the United States, 51% of all AIDS cases have been men who reported sexual contact with other men. In central Africa and in some areas of the Caribbean, sexual transmission appears to be primarily heterosexual. Other sexually transmitted diseases such as herpes, syphilis, and chancroid may facilitate transmission of HIV through ulcerations in the mucous membranes.

Bloodborne pathogen transmission has occurred: (1) by transfusion of blood from HIV infected donors; (2) through receipt of clotting factors for treatment of hemophilia; (3) through the sharing of needles for injection of drugs; (4) through unprotected sexual intercourse with an HIV-infected person (5) through accidents in health care settings with needles or other sharps contam inated with HIV infected blood; and (6) accidental blood splashes on mucous membranes. Up to 30% of infants born to HIV infected mothers may be infected with HIV themselves. The exposure occurs either in utero or during labor and delivery. There are also reports of HIV virus transmission during breast-feeding.

Post-exposure prophylaxis is available for occupational exposure to HIV. The CDC recommends that workers who have an exposure incident must be evaluated within 1-2 hour for a risk assessment and possible prophylactic treatment with antiviral drugs. All potential HIV exposures must be reported and evaluated within 1-2 hour to insure optimal treatment.

From 2003 through 2011, the estimated numbers of newly diagnosed AIDS cases in the United States decreased. In 2011, the estimated rate of AIDS cases in the United States was 15.8 per 100,000 populations. The number of persons living with AIDS (AIDS prevalence) increased steadily from 2003 through 2011. At the end of 2010, an estimated 891,857persons in the United States were living with AIDS. From 2003 through 2007, the estimated number of deaths of persons with AIDS in the United States decreased 17%. From 2008 through 2010, in the United States, the annual estimated number and rate of deaths of persons with diagnosed HIV infection remained stable. In 2010, the estimated rate of deaths of persons with diagnosed HIV infection was 6.3. From 2008 through 2010, the estimated number of deaths of persons living with diagnosed HIV infection increased. At the end of 2010, an estimated 872,990 persons in the United States were living with diagnosed HIV infection. Persons with AIDS are surviving longer, thus, they are contributing to steady increases in the number of persons living with AIDS. The number of AIDS cases reported in 2011 (33, 698), 2010 (33,980), 2009 (35,498), 2008 (37,818), 2007 (37,041), 2006 (36, 791), 2005 (37,256), 2004 (38,695), 2003 (40,054), 2002 (41,289), 2001 (40, 833), 2000 (41,267) has increased from the numbers reported in 1999 (21,419). The number of AIDS cases
Infection with HIV appears to be lifelong. The disease is characterized by a very long incubation period, which is the time of infection to the onset of life-threatening opportunistic infections, and malignancies that signal the development of full-blown AIDS. Disease progression with HIV is divided into several stages according to types of infections or symptoms reported and are associated with progressive decline in CD4+ cells.

- **Acute Infection**: Within 1 to 4 weeks after infection, an individual may experience acute retroviral syndrome, which will manifest itself in a mononucleosis-like or flu-like illness. Unexplained fever, lymphadenopathy, myalgia (muscle pain), arthralgia (joint pain), headache, sore throat, unexplained diarrhea, fatigue, loss of appetite and rash are signs and symptoms of this usually self-limiting stage.

- **Asymptomatic Infection**: During this stage which occurs 4 to 12 weeks after infection, most individuals will develop HIV antibodies (seroconversion). These individuals may be asymptomatic for months to years but can transmit the virus to others.

- **Symptomatic HIV infection/AIDS Related Complex (ARC)**: During this stage which may last from months to several years after infection, many of the symptoms of acute infection reemerge. Oral candidiasis (oral thrush), oral hairy leukoplakia, Herpes zoster in individuals younger than 60 years and ulcerative Herpes simplex may also occur.

- **AIDS**: Many HIV infected individuals go on to develop AIDS months to several years after initial infection. Their immune systems become severely weakened turning normally mild, opportunistic or rare infections into potentially fatal diseases. Indicator diseases of AIDS include: Pneumocystis carinii pneumonia, the most common cause of death; fungal diseases of the esophagus, trachea, bronchi or lungs especially with *Candida* or *Cryptococcus*; Cytomegalovirus (CMV) retinitis; Kaposi's sarcoma; primary brain lymphoma; extrapulmonary disseminated mycobacteria tuberculosis; any mycobacterial disease caused by mycobacteria other than *Mycobacterium tuberculosis*, especially *Mycobacterium avium* complex or *Mycobacterium kansasii*; pulmonary tuberculosis; recurrent pneumonia; invasive cervical cancer; brain toxoplasmosis; disseminated fungal diseases, especially coccidioidomycosis and histoplasmosis; HIV associated encephalopathy (AIDS dementia); HIV associated wasting syndrome (slim disease); HIV disease progression is often associated with decrease of CD4+ cell count throughout.

5.2 **VIRAL HEPATITIS: Hepatitis B, Hepatitis D, and Hepatitis C Viruses**

Viral hepatitis is an ancient disease. The liver, the major target organ in viral hepatitis, plays an essential role in metabolism and degradation and detoxification mechanisms. Cellular damage to the liver, either directly or indirectly, results in a broad range of individual symptoms.

Despite widespread diagnosis and knowledge of infection routes, hepatitis remains one of the leading viral diseases requiring hospitalization in the United States. Over 200,000 cases are reported annually to the CDC. The clinical manifestations of viral hepatitis vary widely from asymptomatic clinical inapparent disease to severe acute symptoms of fatigue, nausea, anorexia, vomiting and fever to chronic and debilitating liver disease followed by cirrhosis and hepatocellular carcinoma.

5.2.1 **Hepatitis B Virus (HBV)**

HBV is a hepadnavirus. As the name indicates, it is a DNA virus, which infects the liver and replicates in liver cells (hepatocytes). HBV is released into the bloodstream from infected hepatocytes. HBV infection may result in a long term carrier state with either mild or severe chronic liver disease including primary hepatocellular carcinoma. The virus is found all over the world with over 400 million carriers worldwide. There are estimated to be about 1,000,000 to 1,400,000 carriers among "healthy" adults in the United States. Total infection with HBV is 5 to 10 times the carrier rate since most infections of adults result in clearance of viremia followed by immunity. Of those people who develop chronic HBV infections (6% of persons infected as adults), 15-25% will eventually die of liver disease. More than 50% of foreign-born persons in the United States from central and southeast Asia, the Middle
East, and Africa are HBV antigen positive, with 5-15% chronically infected. It is estimated that 60,000 new infections occur in the US annually. 4,000 - 5,000 deaths due to chronic liver disease occur each year, including about 200 deaths of health care workers with occupationally contracted HBV.

Blood and blood products are the most effective vehicles for the transmission of HBV. Hepatitis B surface antigen (HBsAg) has been found in virtually all body secretions and excretions, however, only blood, saliva, breast milk, semen and vaginal fluids have shown to be infectious.

Accidental direct percutaneous inoculation (needlestick or other sharp) is the most efficient HBV transmission method. Percutaneous transfer of HBV infected serum or plasma without direct puncture can occur through minute cutaneous scratches or abrasions and through contamination of mucous membranes. Indirect transfer of infective material to skin or mucous membranes can occur by way of contaminated medical devices, gloves or other environmental surfaces.

The efficiency of HBV transmission by the various methods is due to the extraordinary amount of circulating infectious HBV in the blood of infected individuals who are either in the acute phase of infection or who are HBAg carriers and are positive for Hepatitis B "e" antigen (HBeAg). The presence of HBsAg and HBeAg and of HBV viral DNA in an individual's serum is a sign of relatively high infectivity. HBsAg and HBeAg positive human serum can be diluted a hundred million times and still induce HBV infection in experimental animals.

HBV is an extremely stable virus in the environment. Studies at the CDC demonstrated HBV in HBsAg and HBeAg positive sera remained infectious after drying on a surface at 42% relative humidity for at least a week.

5.2.2 Hepatitis D Virus (HDV)

First described in Italy in 1977, HDV (delta agent) is a defective RNA containing virus with HBsAg as a surface coating. HDV requires HBV for replication. It appears to be worldwide in distribution and is endemic in areas of Southern Europe, Africa, the Middle East and South America. The virus is found in HBsAg positive individuals from the same risk groups as in HBV infection. Three types of infection may result from HDV:

- Acute HDV infection superimposed on chronic HBV infection is the most serious category. HDV can replicate indefinitely in the presence of HBV resulting in fulminate disease with high morbidity and mortality. If an individual survives this serious complication of HBV infection, the end result may be chronic hepatitis and cirrhosis.
- Chronic HDV superimposed on chronic HBV infection is often asymptomatic.
- Simultaneous HBV and HDV infection manifests itself as an acute viral hepatitis and usually resolves without complications.

5.2.3 Hepatitis C Virus (HCV)

HCV or parenterally transmitted hepatitis nonA-nonB (NANB) virus is an enveloped RNA virus. HCV infection is the most common chronic bloodborne infection and is a major cause of liver disease in the United States. An estimated 4.1 million Americans have been infected with HCV. There are 26,000 documented cases of HCV in US annually. 55% - 85% of persons with HCV infection become chronically infected, and chronic liver disease with persistently elevated liver enzymes develops in approximately 70% of all HCV infected persons. Persons with chronic hepatitis C are at risk for cirrhosis and primary hepatocellular carcinoma.

Most HCV transmission is associated with direct percutaneous exposure to blood. Health care workers (HCW) are at occupational risk for acquiring this viral infection. However, no vaccine is available to prevent hepatitis C, and immune globulin and antiviral agents are not recommended for postexposure prophylaxis.

The severity of HCV infection ranges from inapparent cases to, in rare cases, fulminating fatal disease. While 80% of HCV infections progress to chronic liver disease over a period of years. Symptomatic chronic HCV infection often improves within 3 to 5 years after infection but may progress to cirrhosis.
In infected individuals, HCV antibody is not detected until an average of 15 weeks after the onset of hepatitis (22 weeks) and may even remain undetectable for a year. There exists a "healthy" carrier state since HCV viremia may persist throughout the course of disease and is not always cleared.

SECTION 6.0 DOOR SIGNS, LABELS AND COLOR CODING

6.1 Wall and Door Signs

Biosafety Level Wall Sign
Entry ways to research and clinical areas that handle human blood and other potentially infectious materials must be posted with a Biosafety Level wall sign that contains the universal biohazard symbol, bears the legend “Biohazard” and BL2 or BL3.

Laboratory Safety Information Card
A Laboratory Safety Information card must be completed and posted with the sign at the laboratory entrance to provide information on the materials handled inside the laboratory, as well as the name and phone numbers of the principal investigator or other responsible person.

Biohazard Door Sign
HIV and HBV research laboratories and production facilities, laboratories working with certain infectious agents that require special provisions for entry (e.g.; vaccination), BL2+, and BL3 laboratories, must have a biohazard door sign posted on all laboratory access doors. The sign includes the international biohazard symbol, bears the legend "Biohazard", and identifies the name of the infectious agent, any special entrance requirements, and the name and phone numbers of the principal investigator or any other responsible persons. The following elements must be included on the door sign:

BIOHAZARD

(Name of infectious agent)
(Special entrance requirements)
(Name, telephone number of the principal investigator or other responsible person)

The door signs shall be fluorescent orange-red (or predominantly so) with lettering or symbols in a contrasting color.

6.2 Labels and Color-coding

Inside the facility, biohazard labels shall be affixed to containers of medical waste, refrigerators, freezers, incubators, waterbaths, sonicators, biological safety cabinets, and centrifuges containing BL2 or BL3 agents, human blood or "other potentially infectious material". Temporary biohazard labels may be placed on equipment that is only sporadically used for work with human blood or other potentially infectious material and decontaminated after each use. For example, a temporary sign incorporating the biohazard symbol, the identity of the material, and the name of the employee who will be operating the centrifuge may be placed on a centrifuge just prior to a run with human blood, other potentially infectious material or other BL2 material. The sign should remain in place until the employee has disinfected the interior bowl of the centrifuge and rotor after use. Once decontaminated, the sign can be removed, returning the equipment back to standard BL1 lab use.
Biohazard labels shall also be affixed to other containers used to store, transport or ship BL2 or BL3 agents, human blood or "other potentially infectious material". Labels required must have the international biohazard symbol and bear the legend "Biohazard" (see figure below).

The labels shall be fluorescent orange-red (or predominantly so) with lettering or symbols in a contrasting color. Labels shall be affixed as close as feasible to the container by string, wire, adhesive, or any other method that prevents their loss or unintentional removal.

The use of warning labels may be waived if: (1) waste is placed in red bags or red containers; (2) containers of blood, blood components, or blood products are labeled as to their contents and have been released for transfusion or other clinical use; or (3) individual containers of blood or "other potentially infectious materials" are placed in a labeled secondary container during storage, transport, shipment or disposal.

### 6.3 Labeling Equipment for Repair, Relocation, or Discard

Potentially contaminated and contaminated equipment sent out for repair or disposal must be decontaminated as thoroughly as possible before sending it out for repair, scheduling on-site service, relocation or disposal. Once the equipment has been decontaminated with an appropriate disinfectant (10% household chlorine bleach, or EPA registered tuberculocidal disinfectant) for an adequate contact time (at least 15 minutes), affix a Biosafety Notice Tag to the equipment. The completed biosafety notice tag will indicate when the equipment was decontaminated, what disinfectant was used, and the name of the person who performed the decontamination. Thorough decontamination of highly technical or sensitive equipment or equipment with limited access to contaminated areas may not be possible. Decontaminate the equipment to the degree possible (flushing lines or wiping down the exterior) and affix a biohazard label indicating which areas of the equipment remain contaminated before sending it out for repair. The biohazard label must include the biohazard symbol as well as the term "biohazard". The label must convey this information to all affected workers (service representatives, manufacturer, etc.). Equipment tags can be obtained from the Yale Environmental Health and Safety (EHS) at 203-785-3550 or copy of tag is available on EHS website (http://ehs.yale.edu/). A sample of this tag appears in Appendix C.

### SECTION 7.0 ENGINEERING CONTROLS

Engineering controls include equipment, devices or supplies that reduce the risk of employee exposure by removing the hazard or isolating the worker from the hazard. Examples of engineering controls include biological safety cabinets, autoclaves, safety centrifuges, splash guards, mechanical pipetting devices, and self-sheathing needles.

Utilize appropriate engineering controls whenever possible. Good work practices are necessary to assure that engineering controls work effectively. Engineering controls require preventive maintenance or periodic replacement to provide employee protection. It is the direct responsibility of the supervisor to insure engineering controls operate properly.

Engineering controls such as biological safety cabinets, safety centrifuges and mechanical pipetting devices are to be decontaminated immediately (or as soon as feasible) when overtly contaminated, or after a spill of blood or other potentially infectious materials. Engineering controls shall also be decontaminated at the end of the work shift. Decontamination should be performed with a 10% household chlorine bleach solution or an EPA registered tuberculocidal disinfectant.

Engineering controls can be organized into four categories: (1) controls that reduce needlestick opportunities; (2) controls that contain spills; (3) controls that contain splashes and aerosols; and (4) controls that decontaminate.
7.1 Controls That Reduce Needlestick Opportunities

Needlesticks can occur in situations where needles must be manipulated or dissembled. There are devices or systems that reduce the need to use needles or decrease the danger of accidental needlesticks. These or similar devices should be utilized whenever feasible.

“Sharps with Engineered Sharps Injury Protection” include non-needle sharps or needle devices containing built-in safety features that are used for collecting fluids, administering medications or other fluids, or any procedure involving the risk of sharps injury. This description covers a broad array of devices, including:

- syringes with a sliding sheath that shields the attached needle after use;
- needles that retract into a syringe after use;
- shielded or retracting catheters;
- intravenous (IV) medication delivery systems that use a catheter port with a needle housed in a protective covering.

“Needleless Systems” are devices that provide an alternative to needles for various procedures to reduce the risk of injury from contaminated sharps. Examples include: IV medication systems which administer medication or fluids through a catheter port using non-needle connections, and jet injection systems that deliver liquid medication beneath the skin or through a muscle.

Safe sharps devices, such as needleless systems and needles that include safety features must be strongly evaluated by the employer to prevent or minimize exposure potential. The employer must also evaluate whether these devices could prevent future incidents as part of its responsibility under the law in evaluating exposures that occur in the workplace. The employer must solicit input from non-managerial employees regarding the identification, evaluation and selection of effective engineering controls, including safer medical devices. The employer must document each evaluation and continue to pursue engineering controls that are designed to prevent occupational exposure. Finally, where a new engineering control is issued, suitable training on its use must be provided to the employees and documented. See Appendix D for the Sharps Safety Device Evaluation Record and Form.

Where engineering controls will reduce employee exposure either by directly removing, eliminating or isolating the hazard, they must be used. Contact the Yale Environmental Health and Safety for information on needleless systems or needle protected devices or evaluation of safety devices.

Needlesticks occur in situations where needles must be manipulated or dissembled. Each of the following devices or systems reduces the need to use needles or decreases the chance of accidental needlesticks. These or similar devices should be utilized whenever feasible. Additional information and a list of different types of safety devices is available at http://www.healthsystem.virginia.edu/pub/epinet/new/safetydevice.html

7.1.1 Catheter Safety Systems

Products like the Criticon Protective I.V. Catheter Safety System use a needle to pierce the patient's skin. A needle guard begins to cover the needle as the catheter is threaded into the vein. The needle is drawn completely into the needle guard and locked into place before removal from the catheter hub.

Considerations and Limitations of Catheter Safety Systems:
1. A needle must be used when the patient's skin must be pierced.
2. Needle guard and needle units must be discarded into a needle disposal sharps container.

7.1.2 I.V. Access Safety Systems

Products such as the Baxter Interlink Access System utilize blunt plastic cannulas instead of needles to access I.V. tubing, catheters, syringes and multiple and single dose vials.

Considerations and Limitations of I.V. Access Safety Systems:
1. A needle must be used when the patient's skin must be pierced.
2. All needles and syringes must be discarded into a needle disposal sharps container.

7.1.3 Needle Protection Devices

Devices such as the Needle-Pro Needle Protection Device are puncture-resistant sheaths that snap over a used needle. The sheath is attached to the syringe at the base of the needle. After the needle is used, the sheath is snapped over the needle.

Considerations and Limitations of Needle Protection Devices:
1. The sheath protrudes from the needle/syringe unit and may interfere with the procedure being performed.
2. All needles and all syringes must be discarded into a needle disposal sharps container.

7.1.4 Retracting Needle Devices

Products such as the Retractable Technologies, Inc. Vanish Point syringe device has a retractable needle that is retracted into the syringe barrel. After injecting, continue depressing plunger to activate automated retraction mechanism while needle is still in patient. Needle retraction occurs directly from the patient into syringe barrel.

Considerations and Limitations of Needle Protection Devices:
1. A needle must be used when the patient's skin must be pierced.
2. All needles and all syringes must be discarded into a needle disposal sharps container.

7.1.5 Air Bubble Removal Devices

Products like the Filter-Pro Air Bubble Removal Device may be used in place of a needle when air bubbles are expelled from a syringe. Attach the device to the syringe. Hold the device end up, tap the syringe to move the bubbles to the top, push the plunger and expel the air from the sample. Stop pushing the plunger when all the air is expelled from the syringe and the sample wets the filter at the end of the device. The filter should self seal when wet. Detach the device when you are ready to process the sample.

Considerations and Limitations of Air Bubble Removal Devices:
1. If a needle is required to fill the syringe, it must be mechanically removed from the syringe. Never remove a needle by hand.
2. Remove the air bubble removal device in a biological safety cabinet or behind a shield. The syringe contents may be under pressure.
3. All needles, syringes and air bubble removal devices must be discarded into a needle disposal sharps container.

7.1.6 Syringe Needle Shields

Products like the Safety-Lok Syringe enable the user to cover the needle with a shield before disposal. Before use, the syringe is sheathed in a built-in shield. After use, the shield may be slipped forward over the needle and locked into place.

Considerations and Limitations of Syringe Needle Shields:
1. Users may need to adjust to the increased width of the syringe.
2. All needles and syringes must be discarded into a needle disposal sharps container.

7.1.7 Add On Safety Devices

The Point-Lok device by Sims Protex is a freestanding needle protection device that locks onto and contains a single contaminated needle. The Point-Lok device is leak resistant and therefore minimizes exposure to fluids. It provides needlestick prevention from point of use to point of disposal. It is an alternative for needle protection
when an integral engineered sharps safety device is not available. This point-of-use device is small but accommodates a wide range of needle and stylet sizes from 16 to 30 gauge.

Considerations and Limitations of device:

1. All needles and syringes must be discarded into a needle disposal sharps container.

7.1.8 Quick-release scalpel blade handles

Bard-Parker SafetyLock Surgical Blade System offers ease of use, convenience, and a protective design to help minimize accidental blade injury, especially when disarming. Simple downward pressure on the SafetyLock handle lever releases the used blade safely and efficiently without the need for a hemostat.

Considerations and Limitations of Quick-Release Scalpel Blade Handles:

1. Scalpel blade must be discarded into a needle disposal sharps container.

7.1.9 Disposable scalpels with safety features

Single use disposable scalpels with blades that retract or sheath the blade.

Considerations and Limitations of Disposal Scalpel with Safety Features:

1. Most of these devices do not lock with the blade retracted.
2. Scalpel blade must be discarded into a needle disposal sharps container.

7.1.10 Alternative Skin Closure Device

Alternative skin closure devices such as skin adhesives increase patient comfort and reduce physical pain, as well as the anxiety associated with needles used for anesthetic administration. The application and setting of skin adhesive is faster than wound closure with sutures. No need for suture removal. Skin adhesive sloughs off the skin during the wound healing period (usually five to ten days). Although physicians may want to check a healed wound after this time has elapsed, a follow-up visit for suture or staple removal is not required.

Considerations and Limitations of Alternative Skin Closure Device Features:

1. Should not be used over areas of increased tension such as the knee, elbow, or finger knuckles, unless these joints are immobilized.

7.1.10 Sharps Disposal Containers

Sharps disposal containers are used to contain and discard used and unused sharps waste. Containers for sharps disposal must be easily accessible and located as close as possible to the immediate area where sharps are used or found. Sharps containers must be puncture-resistant, closeable and leak-proof on the sides and bottom. Proper use of sharps containers eliminates the need to recap, bend, break or manipulate sharps waste by hand. A contaminated sharp is any object contaminated with human blood, blood products, or other potentially infectious material that is also capable of puncturing the skin.

Waste items that can puncture or tear plastic bags are considered sharps. Examples of sharps and additional information regarding sharps waste disposal are provided in the Yale University Biological Safety Manual. Contact EHS Environmental Affairs at 203-432-6545 if you have waste handling questions. If you dispose of your sharps waste through the hospital system contact Yale New Haven Hospital regarding sharps waste disposal.

Considerations and Limitations of Sharps Disposal Containers:

1. The container must be closed before removal or replacement to prevent spillage or protrusion of contents during handling, storage or transport.
2. The container must be kept in an upright position to prevent spillage or protrusion of contents.
3. The container must be replaced before it becomes overfilled to prevent protrusion of contents and Needlesticks.
7.1.11 Sharps Reprocessing Containers

All reusable sharps (such as surgical needles and scalpels) must be placed in puncture-resistant containers that are leak-proof on the sides and bottom. Reusable sharps containers must be red in color or labeled with the biohazard symbol with the word "Biohazard" in the legend. Reusable sharps must be placed in reprocessing containers immediately or as soon as feasible after use. Contaminated reusable sharps must be stored or processed in a way that does not require an employee to reach into the container by hand, risking the possibility of an injury or needlestick.

7.1.12 Tong, Forceps and Hemostat

Mechanical devices for handling or collecting contaminated sharp objects, such as needles and syringes, scalpels, and broken glass. Never manipulate sharps directly by hand. Pick up contaminated broken glass with tongs, scoops, or dust pan and brush. Use forceps or a hemostat to change scalpel blades or use scalpel blades with safety features. If needles must be removed from the syringe, use forceps, a hemostat, or a safe sharps device designed for removing scalpel blades.

7.2 Controls That Contain Spills

7.2.1 Plastic Backed Towels, Bench Coats or Diapers and Spill Trays

Plastic backed towels, bench coats or diapers and spill trays used in conjunction with the biohazard warning label symbol provide a flexible, clear definition of work areas where potentially infectious materials are in use. They also absorb and/or contain potentially infectious materials in the event of a spill and facilitate clean up when work is completed. Because spill trays are reusable, the volume of medical waste generated each day may be reduced. Long term use of spill trays may be more cost effective than the plastic backed towels or diapers.

Considerations and Limitations of Plastic Backed Towels, Bench Coats or Diapers and Spill Trays:

1. Towels, coats or diapers must be removed, decontaminated, discarded and replaced when visibly soiled or at the end of the work shift.
2. Notebooks, pens and other common use items must not be placed in the defined biohazard work area.
3. Infectious materials spilled on spill trays may splatter or aerosolize.

7.2.2 Specimen Containers and Specimen Transport Bags

Human blood or other potentially infectious materials must be placed in containers, which prevent leakage during collection, handling, processing, storage, transport or shipping. These containers include test tubes, freezer vials, etc. All specimen containers must be labeled with a biohazard symbol and the term "biohazard" when the specimens leave the facility, or when the specimen containers are not readily identifiable as containing specimens. Specimen transport bags may be used if placed in a rigid puncture proof container, that is red or labeled with a biohazard symbol and the term “biohazard”, when specimen containers are packaged and sent to another facility. Be certain to select specimen bags with the biohazard symbol and the word "Biohazard" in the legend. Some bags have outside pockets to help keep accompanying paperwork from being contaminated.

Considerations and Limitations of Specimen Containers and Specimen Transport Bags

1. Examine all containers for cracks, chips or other flaws before use. Containers with cracks, chips or flaws may break or leak. Do not use defective containers - discard them through the medical waste stream.
2. Primary containers that become contaminated on the outside or can be punctured by the specimen must be placed within a secondary container. Secondary containers are subject to all labeling requirements.
3. Handle all specimen transport bags with surgical gloves. Assume the specimen container or the paperwork in outer pocket may be contaminated.
7.3 Controls That Contain Splashes and Aerosols

Controls that contain splashes and aerosols include vacuum line trap and filter systems, splash guards, biological safety cabinets (BSCs), mechanical pipetting devices, safety centrifuges, safety blenders and safety sonicators. The following types of controls should be incorporated into protocols whenever possible.

7.3.1 Vacuum Line Chemical Traps and Filters

Vacuum line chemical traps and filters prevent suction of human blood and other potentially infectious materials into the vacuum lines. The trap systems also prevent vacuum lines from clogging with non-infectious material. A vacuum filter is located between the overflow flask and the vacuum line.

![Diagram of vacuum line with labels: A. Collection Flask, B. Overflow Flask, C. Filter, D. Vacuum line](image)

Contact Yale Environmental Health and Safety for information regarding vacuum line filters. Vacuum line filters are available in the stock rooms.

Considerations and Limitations of Vacuum Line Chemical Traps and Filters:

1. The collection flasks should be monitored and emptied or replaced before they are filled. All connections or seals shall be tight to assure the vacuum is adequate.

2. Add full strength chemical disinfectant to collection flasks and allow the aspirated fluids to dilute disinfectant to appropriate concentration. (For example: Start with 100 ml household chlorine bleach, aspirate 900 ml fluids and discard.). Change bleach solutions in the collection flask periodically as the concentration of chlorine will dissipate upon exposure to light and air. Periodically flush dilute bleach solutions through sections of plastic tubing for decontamination.

3. Vacuum line filters shall be examined and replaced if clogged or if liquid makes contact with the filter. Used filters shall be discarded in the medical waste stream.

4. If the collection and overflow flask is located on the floor outside the biological safety cabinet, ensure that it is protected from physical damage by placing the flasks in a protective container. A cardboard box or Styrofoam container is suitable for this purpose.

7.3.2 Splash Guards

Splash guards are clear plastic shields that prevent potentially infectious material from splashing onto laboratory workers when working on the open bench.

Considerations and Limitations of Splash Guards:

1. Splash guards protect against splashes - not aerosols.

2. Decontaminate the inside portion of the shield at the end of the experiment.
7.3.3 Biological Safety Cabinets

Biological Safety Cabinets (BSCs) offer personal, product and environmental protection. A BSC isolates biohazards from the worker by confining the contaminant within the cabinet and removing the contaminants through High Efficiency Particulate Air (HEPA) filters. The cabinet's intake and exhaust air is filtered through a HEPA filter before flowing into or out of the BSC work area. Any aerosols generated within the cabinet work area are contained within the BSC. See Appendix E for a discussion of the proper use of the class II BSC.

Considerations and Limitations of Biological Safety Cabinets:

1. Many BSCs are equipped with germicidal ultra-violet (UV) lamps. The germicidal effect of the UV lamp is affected by time of exposure, distance, presence of dust or debris and the UV lamp intensity. Therefore UV lamp intensity must be monitored. Even though UV lamps maintain a blue visible glow throughout their lifetime, it does not mean the lamps still have a germicidal effect. Routine surface decontamination of the BSC is more effective then the use of UV lamps.

2. Biological Safety Cabinets (BSCs) must be certified after installation and before use, after being relocated and on an annual basis. Consult the Clean Air Device Program Guide for information regarding BSC certification, use and repair.

3. BSCs must be professionally decontaminated before the unit is relocated, stored or if service to the interior of the unit is required. Consult the Clean Air Device Program Guide for decontamination service information.

7.3.4 Mechanical Pipetting Devices

Mechanical pipetting devices must be used in place of mouth pipetting or mouth aspiration. Mechanical pipetting devices prevent: (1) oral contamination by aspiration of infectious fluids or aerosols; (2) transfer of infectious material from fingers to the mouth by the proximal end of the pipette; (3) contamination of the work environment; and (4) possible injuries from sharp or broken pipettes.

Considerations and Limitations of Mechanical Pipetting Devices:

1. Even though mechanical pipetting devices used with biohazards are decontaminated after use, they should be labeled with the biohazard symbol. When using mechanical pipetting devices, be careful not to create aerosols while dispensing liquids.

2. To prevent internal contamination of pipetting device from aerosols generated during pipetting operations, use cotton plugged or filtered pipettes.

7.3.5 Safety Centrifuges

Safety centrifuges such as centrifuges with automatic locking mechanisms or solid lids prevent the centrifuge lid from being opened while the rotor is still in motion, thereby preventing the release of aerosols. The locking device is released after the centrifuge head has stopped revolving. Centrifuges without automatic locking mechanisms or solid lids shall be replaced by those with automatic locking mechanisms and solid lids as soon as possible. Use safety buckets or gasketed carriers for low to moderate centrifugation.

Considerations and Limitations of Safety Centrifuges:

1. Ensure that o-rings are in place on centrifuge rotors and other locations where they are required for proper use of equipment.

2. Small table top centrifuges may be operated within a biological safety cabinet to protect workers from any aerosols generated. Place the centrifuge to the rear of the biological safety cabinet and do not perform any work in the biological safety cabinet while the centrifuge is in operation.

7.3.6 Centrifuge Safety Buckets or Carriers with Covers

Safety buckets or carriers with covers are designed to retain the contents of the centrifuge tube in the event of breakage or leakage.
Considerations and Limitations of Safety Cups or Buckets:

1. A centrifuge tube that leaks or breaks is likely to be under pressure. Routinely open safety cups inside a BSC to contain the release of any potential aerosols.

7.3.7 Safety Blenders and Safety Sonicators

Safety Blenders and Safety Sonicators are designed to contain aerosols during operation.

Considerations and Limitations of Safety Blenders and Safety Sonicators:

Safety Blenders and Sonicators must be opened inside a BSC to prevent the release of aerosols since contents may be under pressure. Wait at least 5 minutes before opening safety blenders and sonicators to allow aerosols to settle.

7.4 Controls That Decontaminate

Physical decontamination controls use low temperature, incineration, high temperature (either dry or moist heat), osmotic pressure, sonic and ultrasonic waves, ultraviolet light, x-rays and gamma rays to achieve their effect. Decontamination renders an item safe for further handling.

Moist heat has greater penetrating power than dry heat. One of the most effective physical decontamination controls is steam sterilization (autoclaving) which generates moisture and high temperature (pressurized steam) within a sealed chamber. Autoclaves can sterilize all items that are heat stable (not damaged by steam or high temperature). In gravity autoclaves, cycles of 250°F (121°C), 15 to 18 lbs. pressure for one hour may be required for decontamination. In the newer vacuum autoclaves, decontamination may require 270°F (132°C), 27 to 30 lbs. pressure for 45 minutes. Use a biological indicator to verify your autoclave technique. Contact the Office of Environmental Health and Safety for more information on the Biological Indicator test kit. Personal protective equipment (PPE) such as rubberized aprons, full face shields and heat and liquid resistant gloves must be worn when operating autoclaves.

Reusable glass and other labware can be decontaminated in hot air ovens. Higher temperatures and longer time periods are required to achieve decontamination. Hot air decontamination is achieved at 160° to 180°C in 1-4 hours. A minimum of 2 hours is required to destroy spores. The effectiveness of hot air ovens is increased if the oven has a circulating fan.

When autoclaves or hot air ovens cannot be used, an alternative method such as chemical decontamination may be employed. Items must be soaked in a tuberculocidal disinfectant or a 10% bleach solution for at least 30 minutes. Heavily soiled items must be cleaned first.

Considerations and Limitations of Autoclaves and Hot Air Ovens:

1. Whatever the temperature and time requirement for decontamination, the contents of each load must be positioned so that steam penetrates into, or heated air flows freely among all items to be decontaminated. Tightly sealed or stoppered materials may not be effectively decontaminated and may become dangerously pressurized causing injury when removed.

2. A routine autoclave maintenance program is recommended. Regular chemical "tape" monitoring of temperature and periodic biological monitoring should be performed to evaluate the effectiveness of the autoclave. Place biological indicators at locations inside the load, the area slowest to heat up, throughout the autoclave are the best indication of sterilization. Autoclaves should be tested periodically. Contact the Office of Environmental Health and Safety for assistance in testing your autoclave.

3. Items containing chemicals (such as phenol-chloroform) should not be placed in an autoclave or a hot air oven (remove chemicals first).

SECTION 8.0 WORK PRACTICE CONTROLS
Work practice controls reduce the likelihood of employee exposure to infectious agents by altering the manner in which a task is performed. The protection provided by work practice controls is based upon employee behavior and attitude.

Proper work practice controls ensure that engineering controls and personal protective equipment are used effectively. Proper work practice controls protect others from exposure to pathogens in the work area or facility, reduce possible cross contamination and improve the quality of the work performed. Routine use of safe work practices also provides a margin of safety for unrecognized hazards.

REMEMBER: Safety is a shared responsibility. Your attitude and work practices are critical for your own health and safety, and for the welfare of those around you.

A work practice essential to reducing employee exposure to bloodborne pathogens is the practice of Universal Precautions. According to the concept of Universal Precautions, all human blood and certain fluids are treated as if known to be infectious for HIV, HBV and other bloodborne pathogens.

8.1 General Work Practices

Organize and plan work procedures with your safety and the safety of others in mind. Keep an uncluttered work space. Always make sure all necessary safety materials and exposure control equipment are available and in good working order. Keep EPA registered tuberculocidal disinfectant or 10% household chlorine bleach and paper towels nearby in case of a spill. Decontaminate all work surfaces with 1:10 dilution of household chlorine bleach or manufacturer’s recommended dilution of an EPA registered tuberculocidal disinfectant upon completion of work and after any spill before leaving your work area.

Know the location of the eyewash and know how to use it. Test the eyewash once a week to flush the system.

Always remove laboratory coats, gowns, smocks, gloves, shoe covers and all other personal protective equipment and wash hands before leaving clinical or laboratory areas for general access areas such as lunchrooms, libraries and administrative offices.

Avoid touching your face, skin, or handle clean surfaces, materials or equipment while wearing gloves.

Use mechanical pipetting devices – never pipette by mouth.

Beards or mustaches may be undesirable in work places with potential airborne contamination. Facial hair retains particulate contamination more persistently than clean shaven skin. Clean shaven faces enhance the fit of facial masks and are required when respirators are used.

Eating, drinking, smoking, applying cosmetics and lip balm and handling contact lenses are prohibited in potentially contaminated work sites. Hand creams and lotions are permitted because they are not considered to be cosmetics. Use non-petroleum based hand creams only. Petroleum based hand creams can compromise the integrity of some brands of gloves.

Food and drink must never be stored in refrigerators, freezers, cabinets or bench tops where blood or other potentially infectious materials may be present.

8.2 Handwashing

Handwashing removes microorganisms that may have contaminated hands during manipulation of specimens, equipment and supplies or while treating patients or contacting environmental surfaces. Each clinical or laboratory area must have readily accessible handwashing facilities. Wash hands with soap and running water immediately, or as soon as feasible, after removing gloves and other personal protective equipment. Wash hands when leaving the clinical or laboratory area for general access areas such as lunchrooms, libraries, and administrative offices. Wash
hands and change gloves between patients. Wash any other skin areas with soap and water, and flush mucous membranes with water immediately, or as soon as feasible, following contact with blood or other potentially infectious materials.

Employees working in field situations must use antiseptic hand cleansers in conjunction with clean cloths or paper towels, or antiseptic towelettes (antiseptic towelettes may not be appropriate for use on all mucous membrane areas). Hands must still be washed with soap and running water as soon as feasible even when antiseptic hand cleansers have been used.

8.3 Handling Disposable Needles and Syringes

Hypodermic needle and syringe units must only be used for parenteral injection and aspiration of fluids from humans, laboratory animals and diaphragm bottles.

Use extreme caution when handling needles and syringes. Use needle-locking syringes or disposable syringe-needle units (i.e. the needle is integral to the syringe) whenever possible. Avoid autoinoculation and aerosol generation during use and disposal.

Contaminated needles must not be sheared, bent, recapped or removed unless the supervisor can demonstrate to the Biological Safety Committee that no alternative is feasible or that such action is required by a specific medical procedure. If the appeal is approved by the Biological Safety Committee, then such recapping or removal of a needle must be accomplished through the use of a mechanical device or a one handed technique. The one hand scoop method with the hand holding the sharp is used to scoop up the cap from a flat surface. (Keep one hand behind your back).

All needles and syringes shall be discarded promptly in a needle disposal sharps container after use. (Consult the Biological Safety Manual for more information regarding needle disposal.)

8.4 Reusable Sharps

Contaminated reusable sharps must be placed in special containers as soon as possible after use and stored or processed in a way that does not require employees to reach into the reprocessing container by hand and risk a needlestick or other injury.

Unlike disposable sharps that are immediately discarded within a leak-proof puncture-resistant container after use, procedures involving reusable sharps (such as large bore needles, razor blades, scalpels and fine tip forceps) may present additional opportunities for needlestick or other injuries. Employees may face potential exposure to contaminated reusable sharps during collection, transport, decontamination, or while removing or changing blades.

According to the OSHA, “reusable sharps must be immediately placed in a leakproof puncture-resistant (sides and bottom) container that is labeled with a biohazard symbol or colored red.” Containers must be maintained and used in a manner that prevents an employee from manually handling contaminated sharps.

The following precautions will help minimize your risk of percutaneous injuries from contaminated sharps.

- Don’t change scalpel blades by hand. Use a forceps, hemostat or safety device and keep your hands away from the blade.
- Place a leakproof puncture-resistant tray that contains a suitable disinfectant (such as 10% household bleach) in the immediate work area. Label the collection tray with the biohazard symbol label. Reusable sharps may also be autoclaved.
- Place reusable sharps in the collection tray immediately following use. Lay sharps in the same direction within the collection tray.
- Allow a sufficient contact time for disinfection (at least 20 minutes). If items are covered with debris, wipe clean with a small bristle brush, keeping your hands away from the blade. Decontaminate the brush after use,
After decontamination, remove reusable sharps from the container with tongs or forceps. If bleach is used as a disinfectant, rinse items with ethanol or water to remove any corrosive residues.

Substitute plastic for glass wherever feasible.

8.5 Footwear

Wear close-toed shoes at all times. Sandals or open-toed shoes do not provide adequate foot protection and are inappropriate in clinical, laboratory or animal care areas. A dedicated pair of work shoes may reduce the amount and type of contamination introduced into the workplace by street shoes. This practice can also minimize the possibility of bringing microbial contamination from the workplace into the home. Steel-toed shoes are recommended while working in animal care areas.

8.6 Splash and Aerosol Control

All procedures involving blood or other potentially infectious materials must be performed in a manner that minimizes splashing, spraying, spattering and generation of droplets. This precaution decreases the chances of direct personal exposure and reduces the contamination of bench tops, instruments or other surfaces in the work area. Liquid cultures of infectious material, sealed ampoules and vacutainers are best opened in a biological safety cabinet (BSC). If a BSC is not available, use a splash guard.

Avoid mixing biohazardous materials by drawing and expulsion through pipettes. When delivering pipette contents into a container, allow the contents to run down the container wall or deliver the contents as close as possible to the fluid or agar level. Avoid dropping pipette contents from a height. Mix covered solutions by swirling, inverting or vortexing.

8.7 Housekeeping for Clinical and Laboratory Workers - Surface/Equipment Decontamination

Clinical and laboratory workers are responsible for certain housekeeping activities. Work surfaces are to be decontaminated immediately (or as soon as feasible) when overtly contaminated, or after a spill of blood or other potentially infectious materials. Work surfaces must also be cleaned at the end of the work shift. Remove and replace bench covers, pads or plastic back toweling when overtly contaminated and at the end of the experiment and at least once daily.

Equipment that may become contaminated with blood or other potentially infectious materials must be decontaminated: (1) when visibly contaminated; (2) at the end of the work shift; and (3) prior to servicing or shipping. Heavily soiled equipment that is also contaminated must be prewashed before being decontaminated. Most disinfectants or sterilants cannot effectively penetrate organic material present on heavily soiled equipment. Allow at least 15 minutes of contact time when using disinfectants to decontaminate work surfaces, equipment or spills. See section 6.3 for labeling procedures after equipment has been decontaminated.

OSHA requires EPA registered tuberculocidal disinfectants or 10% household (chlorine) bleach be used to disinfect surfaces contaminated with human blood or other potentially infectious materials. Household (chlorine) bleach sold commercially has a concentration of 5.25 w/v (52,500 ppm) available chlorine. A 1:10 dilution (10% solution) results in a 0.5 w/v (5,250 ppm chlorine) which inactivates bloodborne pathogens. The list of EPA registered tuberculocidal disinfectants is available on the web at [http://www.epa.gov/oppad001/list_b_tuberculocide.pdf](http://www.epa.gov/oppad001/list_b_tuberculocide.pdf)

Ethanol or isopropanol (70%) are effective disinfectants, but are not accepted by OSHA as being tuberculocidal. These alcohols are effective cleansers, and may be used in conjunction with a tuberculocidal disinfectant.

Quaternary ammonium compounds are not believed to be effective against HBV— even though they are considered to be tuberculocidal. Products registered by EPA as HIV effective are not necessarily tuberculocidal, and are not necessarily effective against agents such as HBV, which is more resistant to inactivation than HIV.
Contaminated broken glassware must not be picked up directly by hand. Use mechanical devices (brush and dust pan, tongs or forceps) to facilitate clean up. Decontaminate the mechanical devices with a tuberculocidal disinfectant. Vacuum cleaners are not appropriate for clean up of broken contaminated glass.

Waterbaths and waterbath sonicators used for inactivating, incubating or testing of infectious substances should contain a disinfectant or other microstatic agent to minimize bacterial, fungal or algae growth. Change the water periodically or whenever growth is observed.

Freezers and refrigerators shall be checked periodically. Promptly remove any broken vials, ampoules or tubes containing human material or other potentially infectious materials and decontaminate the inside of the freezer or refrigerator.

General trash receptacles must be inspected daily to insure that regulated sharps are not inadvertently discarded in the general waste stream. Improperly discarded sharps can result in puncture wounds and cuts to custodians and other support staff.

Custodians are responsible for activities such as sweeping or mopping floors, removing general trash, and cleaning general environmental surfaces such as floors and walls. Custodians are instructed not to touch any laboratory or clinical equipment, materials, supplies or special wastes unless instructed by their supervisor to do so. Custodians are instructed not to remove general trash if they find medical waste in the trash receptacle or to pick up medical waste from the floor. Custodians are not responsible for cleaning and decontaminating laboratory spills.

SECTION 9.0 PERSONAL PROTECTIVE EQUIPMENT (PPE)

Engineering and work practice controls provide the first level of protection against exposure to bloodborne pathogens. Personal protective equipment (PPE) is used to provide additional protection when the potential for an occupational exposure remains after engineering and work practice controls have been instituted.

Barrier precautions is another term for the use of PPE. PPE is used to prevent blood or other potentially infectious materials from making direct contact with an employee's clothing or body. The type and amount of PPE required depends upon the task to be performed and the type of anticipated exposure. The types of PPE utilized to prevent exposure to potentially infectious materials include disposable (single use) gloves, rubber utility gloves, protective body clothing (gowns, coats, jumpsuits, aprons), face and eye protection (face shield or surgical mask and protective eyewear), emergency ventilation devices, surgical caps, hoods or head covers, and shoe protection.

Supervisors must discuss with their employees the type and proper use and limitations of PPE needed to perform job tasks safely.

9.1 General Guidelines for Personal Protective Equipment

Yale must provide all PPE at no charge to an employee. Yale is also responsible for cleaning, repairing, disposing of and replacing PPE. The supervisor must be aware of his/her responsibilities regarding PPE maintenance and replacement.

The PPE must be easily accessible and of proper size and must not permit blood or other potentially infectious materials to pass through or to reach the employee's outer or inner clothing (including uniforms), skin, eyes, mouth, or other mucous membranes. Garments penetrated by blood or other potentially infectious materials must be removed immediately or as soon as feasible.

Hypoallergenic gloves, latex free gloves, glove liners, powderless gloves or different glove brands must be provided to employees who exhibit allergic reactions to the gloves normally provided. It is important to note that latex gloves have proved effective in preventing transmission of many infectious diseases to health care workers, but for some
workers, exposures to latex may result in skin rashes; hives; flushing; itching; nasal, eye, or sinus symptoms; asthma; and (rarely) shock. Reports of such allergic reactions to latex have increase in recent years – especially among health care workers.

All PPE must be removed prior to leaving the work area for common areas such as cafeterias, offices, etc. PPE must be placed in an appropriately designated area or container, for storage, washing, decontamination or disposal. Don’t touch clean surfaces, such as phones, door knobs or computers, with gloved hands.

Supervisors are responsible for taking necessary measures to insure their employees wear the PPE outlined for specific tasks.

### 9.2 Considerations and Limitations of Personal Protective Equipment

#### 9.2.1 Gloves

1. Gloves must be inspected prior to use for holes or tears. Glove quality can vary with age, manufacturer and elastomeric materials used in fabrication.
2. Gloves must be replaced as soon as feasible if contaminated, torn, punctured or the integrity of the glove barrier is compromised.
3. Gloves do not protect against injuries from needles or other sharp objects.
4. Contaminated gloves must be discarded in the medical waste stream. Uncontaminated gloves may be discarded in the general trash.
5. Rubber utility gloves may be more desirable than disposable gloves when performing certain procedures such as cleaning. Rubber utility gloves may be washed and reused as long as the integrity of the glove is not compromised. Take care not to contaminate the inside of the gloves. Avoid grasping the outside of a contaminated glove with bare hands.
6. Always wash hands thoroughly with soap and water after glove removal.

#### 9.2.2 Gowns, Lab Coats, Aprons and Jumpsuit

1. Gowns, lab coats and jumpsuits protect the wearer's clothing and skin from contamination. As with all PPE, the type of clothing needed will depend on the task being performed and the degree of exposure anticipated.
2. Long sleeved protective clothing with snug fitting cuffs are preferred over open or short sleeves. Snug fitting cuffs prevent splashes, splatters and aerosols from making contact with exposed skin on the lower arms. Longer single use gloves can be pulled over snug fitting cuffs to seal out any infectious materials.
3. Plastic, vinyl or rubber aprons are usually worn over other protective body clothing when extra protection is desired. Aprons are generally used for protection against liquid spills, splashes or soiling of blood or other potentially infectious materials. Plastic, vinyl or rubber aprons may also be used to provide protection from steam and hot water in locations such as animal handling facilities, autoclave rooms and laboratory glass washing rooms.
4. Protective clothing must be removed as soon as feasible if contaminated or penetrated by blood or other potentially infectious materials and autoclaved before being discarded or laundered.

#### 9.2.3 Face and Mucous Membrane Protection

1. Face and mucous membrane protection must be worn whenever there is potential for the generation of splashes, spray, splatter or droplets of blood or other potentially infectious material in the eyes, nose, mouth or other facial areas.
2. Eye protection may prevent damage to the eye in addition to preventing exposure to bloodborne pathogens. Certain disinfectants and other chemicals can damage the eye or cause blindness if splashed in the eye.
Note Contact lenses provide no protection in the laboratory and should not be substituted for required eye protection. The use of contact lenses may increase the risk of eye damage because infectious agents and other microbes may become trapped between the contact lens and the cornea. The liquid between the contact lens and cornea is an excellent growth medium for microorganisms. If contact lenses must be worn, barrier eye protection shall be worn.

3. Surgical masks are generally protective against droplets, splashes and sprays. Masks must cover both the nose and the mouth. Masks must fit the face closely, so the air passes through the mask before being inhaled. Some surgical masks are available with attached eye shields. Moisture from expired air may eventually saturate the mask, making breathing difficult. Change the mask once it has been compromised. Note: surgical masks do not protect the worker from aerosol exposure. Use a respirator to prevent inhalation of infectious aerosols.

Proper selection of respirators is very important and should not be made without input from the Office of Environmental Health and Safety (EHS). New regulations concerning respirators require initial and annual training and fit-testing, and well as medical surveillance of all respirator wearers. Please make sure that the Office of Environmental Health and Safety is notified whenever the use of a respirator is being considered. The Respirator Administrator in EHS can assist in evaluating the procedure, selecting the proper respirator, and provide the required training and fit testing. The Employee Health Office must also be notified so that medical surveillance and clearance can be issued prior to wearing the respirator. A copy of Yale University’s Respiratory Protection Program is available at http://ehs.yale.edu/.

9.2.4 Head Covers

1. Head covers are worn when gross contamination or splashing on the head is reasonably anticipated. These situations may arise when performing autopsies, orthopedic surgery or working in animal facilities.

9.2.5 Shoe Covers

1. Shoe covers or boots shall be worn when gross contamination is reasonably anticipated. Circumstances where shoe protection may be necessary include animal rooms, surgery and autopsy rooms, etc. Shoe covers are required to prevent contamination migration and direct and indirect transmission.

9.2.6 Emergency Ventilation Devices

1. Emergency ventilation devices such as mouthpieces and resuscitation bags protect personnel while performing artificial resuscitation. Emergency ventilation devices must be readily available for use in areas where the need to perform artificial resuscitation is anticipated.

9.3 Decontamination of Personal Protective Equipment

Disposable gloves must not be washed or decontaminated for re-use. Disinfecting agents (including soap and water) often cause deterioration of glove material (e.g., latex). Washing with surfactants can result in “wicking” or enhanced penetration of liquids through the gloves. Wicking can transport potentially infectious materials to the skin inside the glove.

Rubber utility gloves may be decontaminated and discarded in the general waste stream. Contaminated utility gloves shall be discarded in the medical waste stream. Contaminated laboratory clothing (such as lab coats) should be placed in a biohazard bag and autoclaved if extensively contaminated or “spot treated” if contamination is isolated before being discarded or laundered. Never take used PPE home to launder. Reusable face and mucous membrane protection (face shields, goggles, safety glasses, surgical masks) must be decontaminated with 10% household chlorine bleach or an EPA registered tuberculocidal disinfectant after use and when visibly soiled.

SECTION 10.0 HEPATITIS B VACCINE INFORMATION
Hepatitis B vaccination provides the most effective protection from Hepatitis B virus. "Occupationally exposed" employees are strongly encouraged to receive the Hepatitis B vaccine.

The Hepatitis B Vaccine Notification Form (Appendix G) must be completed for each occupationally exposed employee. The Employer is not required to offer the vaccination to those employees who have already received the vaccine, or to those whose antibody status has shown the employee to be immune, or to those for whom the vaccine is contraindicated for medical reasons.

The Hepatitis B vaccine is offered free of charge by the Department of Employee Health. Call Employee Health (203-432-0071) to arrange to receive the vaccine. Pre-exposure vaccination is preferred. If an "occupationally exposed" employee initially declines the vaccine, he/she can choose to be vaccinated at a later time. The vaccine is also offered to unvaccinated employees who experience an "exposure incident". Employees who have already received the vaccine should have a blood test to demonstrate immunity to the virus.

10.1  Hepatitis B Vaccine

Hepatitis B vaccine is 96-99% effective in preventing HBV infection. Booster doses of hepatitis B vaccine are not necessary. If a routine booster dose of Hepatitis B vaccine is recommended by the US Public Health Service at a future date, the employer must make the booster dose available to its employees. The vaccine also protects against Hepatitis D viral (HDV) infection, as HDV requires the co-existence of HBV infection for viral replication.

The vaccine is a series of three injections given intramuscularly. The second dose is given 1 month after the first. The third dose is given 5 months after the second. Remember: All three shots must be given for the vaccination to be complete. Employees must be tested for antibody to Hepatitis B surface antigen 1 to 2 months after the completion of the series. Employees who do not respond to the primary vaccination series must be revaccinated with a second three-dose vaccine series and retested. Non-responders must be medically evaluated.

The first Hepatitis B vaccine (licensed in 1982) was made from inactivated human sera from people with chronic HBV infections. This vaccine is no longer available in the United States.

The newer synthetic vaccines (first licensed in 1987) are much safer. Recombinant DNA technology enabled the Hepatitis B surface antigen (HBsAg) gene to be inserted into common baker's yeast cell DNA. These altered yeast cells produce HBV protein markers, but no complete virus particles. It is virtually impossible to become infected with bloodborne pathogens from the vaccine. No whole or live virus particles or human sera are used in the vaccine preparation.

10.2  Medical Considerations

The only medical reason for not receiving the vaccine is a history of allergy to yeast or any of the vaccine components. Employees with a history of chronic illness or immunosuppression should consult the Employee Health physician or their personal physician before receiving the vaccine. Pregnant women or nursing mothers should not receive the vaccine until it has been discussed with the physician. Vaccine administration should be delayed if possible in persons with a current febrile illness or other active infections.

The most commonly reported side effects are local redness and soreness at the injection site, low grade fever, headache or dizziness in 1-10% of those vaccinated.

Further questions regarding the vaccine should be directed to the Department of Employee Health (203-432-0071).
A bloodborne pathogen emergency is an unplanned release of human blood or "other potentially infectious materials". This release may be the result of: (1) a spill; (2) an aerosol release; (3) an injured co-worker; or (4) an unanticipated encounter with human blood or "other potentially infectious materials".

If an "exposure incident" occurs in conjunction with an emergency, follow the exposure incident procedures outlined in section 12.0 in addition to the emergency action procedures described here.

11.1 Decontamination procedures for a spill or other release

Only "occupationally exposed" employees can perform decontamination procedures. Protective clothing and equipment appropriate for the situation must be worn. Household chlorine bleach or an appropriate EPA registered tuberculocidal disinfectant shall be used to clean all spills of human blood and "other potentially infectious materials". Make up a fresh 1:10 dilution of household chlorine bleach solution. Mix up a fresh dilution of an appropriate EPA registered tuberculocidal disinfectant following the manufacturer's instructions.

11.2 Emergency Actions to Take - Spills of Blood or Other Potentially Infectious Materials

Stop work immediately. Presume the spilled material is contaminated with bloodborne pathogens. Inform others in the immediate area that a spill has occurred. Employees who are not considered "occupationally exposed" must leave the area and not attempt to clean up the spill. These individuals must notify their supervisors and obtain assistance from employees who are considered "occupationally exposed". If the spill occurred in a public access area (hallways, waiting rooms, restrooms, etc.), or if you do not know how to proceed, notify Yale Environmental Health and Safety (203-785-3555).

If you will be cleaning up the spill, first contain it to prevent it from spreading to uncontaminated areas. Place paper towels or other absorbent materials over the spill. Pour enough disinfectant into the spill puddle to double its size (if possible). Allow the disinfectant to remain in contact with the spilled material for at least 15 minutes. Carefully mop up the liquid, or soak it up with paper towels or disposable pads. Pick up any glass (or other sharps) with tongs. Discard the towels or pads in a red bag. Remove all spilled materials and decontaminate the area again with an appropriate disinfectant. If a mop is used, soak the mop in fresh disinfectant for 20 minutes before rinsing for reuse. Wash reusable gloves with the disinfectant. Wash hands thoroughly with soap and water afterward. See Appendix F, Universal Precautions Spill Response Guide for further information on spill response to BL2 and blood spills.

11.3 Emergency Actions to Take - Aerosol Generation

Stop work immediately. Presume the aerosolized material is contaminated with bloodborne pathogens. Inform all others in the area that an aerosol may have been generated. All persons shall evacuate the room immediately for at least 30 minutes. Notify the Yale Environmental Health and Safety at 203-785-3555. Label the area off-limits for at least 30 minutes. Decontaminate all exposed environmental surfaces after 30 minutes have passed and before releasing the room for normal use.

11.4 Emergency Actions to Take - Injured Co-worker

Assess the situation. If the injured worker can provide his or her own first aid, assist the person only by supplying bandages and dressings. The injured person will clean and decontaminate any contaminated work surfaces, if possible. If further assistance is needed, persons who are not considered "occupationally exposed" shall notify their supervisors and obtain assistance from workers who are considered "occupationally exposed".
Workers who assist injured co-workers must follow Universal Precautions while performing Good Samaritan acts. Employees must use their best judgment if the injury is life threatening. Employees who incur an exposure incident during a Good Samaritan act will notify their supervisor and seek treatment from Acute Care at 55 Lock Street.

11.5 Emergency Actions to Take - Unanticipated Encounters

A situation may arise where an unanticipated occupational exposure to human blood or "other potentially infectious materials" may occur. In all likelihood, employees experiencing an unanticipated encounter will not have previously been considered "occupationally exposed". These employees must attempt to avoid contact with human blood and "other infectious materials". They must notify their supervisors and Yale Environmental Health and Safety at 203-785-3555. If the employees at the scene are considered "occupationally exposed" they shall follow the instructions for spills or aerosols as appropriate.

SECTION 12.0 EXPOSURE INCIDENT RESPONSIBILITIES/PROCEDURES

An "exposure incident" is specific contact (eye, mouth, other mucous membrane, non-intact skin, or parenteral) with blood or "other potentially infectious materials" that results from the performance of an employee's duties.

12.1 Employee's Responsibilities

An employee who sustains a known or potential "exposure incident" must wash the area immediately with soap and water. When handwashing sinks are not available, antiseptic hand cleanser or towelettes may be used for skin only. The exposed area must be washed with soap and running water as soon as possible. The employee must report the incident to his/her supervisor, and seek medical assistance at the Yale Health Center’s Acute Care immediately.

12.2 Supervisor's Responsibilities

The supervisor must complete a Department Head's Report of Injury form and a Health Service Report form, documenting the route of exposure and the circumstances under which the incident occurred. The injured employee will be sent with the forms to Acute Care at 55 Lock Street for treatment and counseling. It is essential that the employee gets to medical assistance immediately, especially for high risk exposures, employees can be sent without forms. However, the forms must be sent within 48 hours.

12.3 University Health Services' Responsibilities

Yale Health will provide the post-exposure evaluation and follow-up at no cost to employees who experience "exposure incidents".

12.4 Post-Exposure Evaluation and Follow-Up

All employees who have an "exposure incident" will be offered a confidential post-exposure medical evaluation and follow-up through the Department of Employee Health. The post-exposure medical evaluation and follow-up includes the following:

1. A review/evaluation of the route of exposure and the circumstances under which the incident occurred.
2. An attempt to identify the source individual, if possible, and his/her HIV and HBV infection status.
3. The employee will be offered the option of having blood drawn for baseline blood collection (storage) or for HIV and HBV serological status testing.
4. The employee will be offered post exposure prophylaxis when medically indicated.
5. The employee will be given appropriate treatment and counseling concerning precautions to take during the
period after the exposure incident. The employee will also be given information on what potential illnesses to
be alert for and to report any similar experiences to appropriate personnel.

The University must provide the employee with a copy of the evaluating health care professional's written opinion
within 15 working days of the completion of the original evaluation. The opinion for post-exposure evaluation and
follow-up will indicate: (1) that the employee has been informed of the results of the evaluation; and (2) that the
employee has been told about any medical conditions resulting from exposure to blood or "other potentially
infectious materials" that require further evaluation or treatment. All other findings or diagnoses will remain
confidential and will not be included in the written report. All laboratory tests are conducted at no cost to the
employee. Contact the Department of Employee Health (203-432-0071) if you have post-exposure evaluation or
follow-up questions.
REFERENCES

Abram Benenson, editor, Control of Communicable Diseases, 1995, APHA.


Centers for Disease Control, "Update on Adult Immunization" MMWR Vol. 40, No. RR-12, November 15, 1991, pp. 31-33.


Centers for Disease Control, "Recommendations for Prevention of HIV Transmission in Health-Care Settings", MMWR August 21, 1987, 36(2S), pp. 2S-18S.


APPENDIX A: OSHA BLOODBORNE PATHOGEN STANDARD

The Standard

General Industry

Part 1910 of title 29 of the Code of Federal Regulations is amended as follows:

PART 1910-[AMENDED]

Subpart Z-Amended
1. The general authority citation for subpart Z of 29 CFR part 1910 continues to read as follows and a new citation for §1910.1030 is added:

Authority: Secs. 6 and 8. Occupational Safety and Health Act. 29 U.S.C. 655, 657, Secretary of Labor's Orders Nos. 12-71 (36 FR 8784), 8-76 (41 FR 25059), or 9-83 (48 FR 35736), as applicable and 29 CFR part 1911.

Section 1910.1030 also issued under 29 U.S.C. 653.

2. Section 1910.1030 is added to read as follows:

1910.1030(a) Scope and Application. This section applies to all occupational exposure to blood or other potentially infectious materials as defined by paragraph (b) of this section.

1910.1030(b) Definitions. For purposes of this section, the following shall apply:

Assistant Secretary means the Assistant Secretary of Labor for Occupational Safety and Health, or designated representative.

Blood means human blood, human blood components, and products made from human blood.

Bloodborne Pathogens means pathogenic microorganisms that are present in human blood and can cause disease in humans. These pathogens include, but are not limited to, hepatitis B virus (HBV) and human immunodeficiency virus (HIV).

Clinical Laboratory means a workplace where diagnostic or other screening procedures are performed on blood or other potentially infectious materials.

Contaminated means the presence or the reasonably anticipated presence of blood or other potentially infectious materials on an item or surface.

Contaminated Laundry means laundry which has been soiled with blood or other potentially infectious materials or may contain sharps.

Contaminated Sharps means any contaminated object that can penetrate the skin including, but not limited to, needles, scalpels, broken glass, broken capillary tubes, and exposed ends of dental wires.

Decontamination means the use of physical or chemical means to remove, inactivate, or destroy bloodborne pathogens on a surface or item to the point where they are no longer capable of transmitting infectious particles and the surface or item is rendered safe for handling, use, or disposal.

Director means the Director of the National Institute for Occupational Safety and Health, U.S. Department of Health and Human Services, or designated representative.

Engineering Controls means controls (e.g., sharps disposal containers, self-sheathing needles, safer medical devices, such as sharps with engineered sharps injury protections and needleless systems) that isolate or remove the bloodborne pathogens hazard from the workplace.

Exposure Incident means a specific eye, mouth, other mucous membrane, non-intact skin, or parenteral contact with blood or other potentially infectious materials that results from the performance of an employee's duties.

Handwashing Facilities means a facility providing an adequate supply of running potable water, soap and single use towels or hot air drying machines.

Licensed Healthcare Professional is a person whose legally permitted scope of practice allows him or her to independently perform the activities required by paragraph (f) Hepatitis B Vaccination and Post-exposure Evaluation and Follow-up.

HBV means hepatitis B virus.

HIV means human immunodeficiency virus.

Needleless systems means a device that does not use needles for:

(1) The collection of bodily fluids or withdrawal of body fluids after initial venous or arterial access is established; (2) The administration of medication or fluids; or (3) Any other procedure involving the potential for occupational exposure to bloodborne pathogens due to percutaneous injuries from contaminated sharps.

Occupational Exposure means reasonably anticipated skin, eye, mucous membrane, or parenteral contact with blood or other potentially infectious materials that may result from the performance of an employee’s duties.

Other Potentially Infectious Materials means (1) The following human body fluids: semen, vaginal secretions, cerebrospinal fluid, synovial fluid, pleural fluid, pericardial fluid, peritoneal fluid, amniotic fluid, saliva in dental procedures, any body fluid that is visibly contaminated with blood, and all body fluids in situations where it is difficult
Examples include, but are not limited to, hospital and clinic pathological and microbiological wastes containing blood materials during handling; contaminated sharps; and infectious materials and are capable of releasing these items that are caked with dried blood or other potentially infectious materials in a liquid or semi-liquid state if compressed; that would release blood or other potentially infectious other potentially infectious materials; contaminated items

Regulated Waste means liquid or semi-liquid blood or other potentially infectious materials; contaminated items that would release blood or other potentially infectious materials in a liquid or semi-liquid state if compressed; items that are caked with dried blood or other potentially infectious materials and are capable of releasing these materials during handling; contaminated sharps; and pathological and microbiological wastes containing blood or other potentially infectious materials.

Personal Protective Equipment is specialized clothing or equipment worn by an employee for protection against a hazard. General work clothes (e.g., uniforms, pants, shirts or blouses) not intended to function as protection against a hazard are not considered to be personal protective equipment.

Production Facility means a facility engaged in industrial-scale, large-volume or high concentration production of HIV or HBV.

Regulated Waste means liquid or semi-liquid blood or other potentially infectious materials; contaminated items that would release blood or other potentially infectious materials in a liquid or semi-liquid state if compressed; items that are caked with dried blood or other potentially infectious materials and are capable of releasing these materials during handling; contaminated sharps; and pathological and microbiological wastes containing blood or other potentially infectious materials.

Research Laboratory means a laboratory producing or using research-laboratory-scale amounts of HIV or HBV. Research laboratories may produce high concentrations of HIV or HBV but not in the volume found in production facilities.

Sharps with engineered sharps injury protections means a nonneedle sharp or a needle device used for withdrawing body fluids, accessing a vein or artery, or administering medications or other fluids, with a built-in safety feature or mechanism that effectively reduces the risk of an exposure incident.

Source Individual means any individual, living or dead, whose blood or other potentially infectious materials may be a source of occupational exposure to the employee. Examples include, but are not limited to, hospital and clinic patients; clients in institutions for the developmentally disabled; trauma victims; clients of drug and alcohol treatment facilities; residents of hospices and nursing homes; human remains; and individuals who donate or sell blood or blood components.

Sterilize means the use of a physical or chemical procedure to destroy all microbial life including highly resistant bacterial endospores.

Universal Precautions is an approach to infection control. According to the concept of Universal Precautions, all human blood and certain human body fluids are treated as if known to be infectious for HIV, HBV, and other bloodborne pathogens.

Work Practice Controls means controls that reduce the likelihood of exposure by altering the manner in which a task is performed (e.g., prohibiting recapping of needles by a two-handed technique).

1910.1030(c)
Exposure Control --

1910.1030(c)(1)
Exposure Control Plan.

1910.1030(c)(1)(i)
Each employer having an employee(s) with occupational exposure as defined by paragraph (b) of this section shall establish a written Exposure Control Plan designed to eliminate or minimize employee exposure.

1910.1030(c)(1)(ii)
The Exposure Control Plan shall contain at least the following elements:

1910.1030(c)(1)(ii)(A)
The exposure determination required by paragraph (c)(2),

1910.1030(c)(1)(ii)(B)
The schedule and method of implementation for paragraphs (d) Methods of Compliance, (e) HIV and HBV Research Laboratories and Production Facilities, (f) Hepatitis B Vaccination and Post-Exposure Evaluation and Follow-up, (g) Communication of Hazards to Employees, and (h) Recordkeeping, of this standard, and

1910.1030(c)(1)(ii)(C)
The procedure for the evaluation of circumstances surrounding exposure incidents as required by paragraph (f)(3)(i) of this standard.

1910.1030(c)(1)(iii)
Each employer shall ensure that a copy of the Exposure Control Plan is accessible to employees in accordance with 29 CFR 1910.1020(e).

1910.1030(c)(1)(iv)
The Exposure Control Plan shall be reviewed and updated at least annually and whenever necessary to reflect new or modified tasks and procedures which affect occupational exposure and to reflect new or revised employee positions with occupational exposure. The review and update of such plans shall also:

1910.1030(c)(1)(iv)(A)
Reflect changes in technology that eliminate or reduce exposure to bloodborne pathogens; and

1910.1030(c)(1)(iv)(B)
Document annually consideration and implementation of appropriate commercially available and effective safer medical devices designed to eliminate or minimize occupational exposure.

1910.1030(c)(1)(v)
An employer, who is required to establish an Exposure Control Plan shall solicit input from non-managerial

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For Clinical and Laboratory Personnel
employees responsible for direct patient care who are potentially exposed to injuries from contaminated sharps in the identification, evaluation, and selection of effective engineering and work practice controls and shall document the solicitation in the Exposure Control Plan.

1910.1030(c)(1)(vi)
The Exposure Control Plan shall be made available to the Assistant Secretary and the Director upon request for examination and copying.

1910.1030(c)(2)
Exposure Determination.

1910.1030(c)(2)(i)
Each employer who has an employee(s) with occupational exposure as defined by paragraph (b) of this section shall prepare an exposure determination. This exposure determination shall contain the following:

1910.1030(c)(2)(i)(A)
A list of all job classifications in which all employees in those job classifications have occupational exposure;

1910.1030(c)(2)(i)(B)
A list of job classifications in which some employees have occupational exposure, and

1910.1030(c)(2)(i)(C)
A list of all tasks and procedures or groups of closely related task and procedures in which occupational exposure occurs and that are performed by employees in job classifications listed in accordance with the provisions of paragraph (c)(2)(i)(B) of this standard.

1910.1030(c)(2)(ii)
This exposure determination shall be made without regard to the use of personal protective equipment.

1910.1030(d)
Methods of Compliance --

1910.1030(d)(1)
General. Universal precautions shall be observed to prevent contact with blood or other potentially infectious materials. Under circumstances in which differentiation between body fluid types is difficult or impossible, all body fluids shall be considered potentially infectious materials.

1910.1030(d)(2)
Engineering and Work Practice Controls.

1910.1030(d)(2)(i)
Engineering and work practice controls shall be used to eliminate or minimize employee exposure. Where occupational exposure remains after institution of these controls, personal protective equipment shall also be used.

1910.1030(d)(2)(ii)
Engineering controls shall be examined and maintained or replaced on a regular schedule to ensure their effectiveness.

1910.1030(d)(2)(iii)
Employers shall provide handwashing facilities which are readily accessible to employees.

1910.1030(d)(2)(iv)
When provision of handwashing facilities is not feasible, the employer shall provide either an appropriate antiseptic hand cleanser in conjunction with clean cloth/paper towels or antiseptic towelettes. When antiseptic hand cleansers or towelettes are used, hands shall be washed with soap and running water as soon as feasible.

1910.1030(d)(2)(v)
Employers shall ensure that employees wash their hands immediately or as soon as feasible after removal of gloves or other personal protective equipment.

1910.1030(d)(2)(vi)
Employers shall ensure that employees wash hands and any other skin with soap and water, or flush mucous membranes with water immediately or as soon as feasible following contact of such body areas with blood or other potentially infectious materials.

1910.1030(d)(2)(vii)
Contaminated needles and other contaminated sharps shall not be bent, recapped, or removed except as noted in paragraphs (d)(2)(vii)(A) and (d)(2)(vii)(B) below. Shearing or breaking of contaminated needles is prohibited.

1910.1030(d)(2)(vii)(A)
Contaminated needles and other contaminated sharps shall not be bent, recapped or removed unless the employer can demonstrate that no alternative is feasible or that such action is required by a specific medical or dental procedure.

1910.1030(d)(2)(vii)(B)
Such bending, recapping or needle removal must be accomplished through the use of a mechanical device or a one-handed technique.

1910.1030(d)(2)(viii)
Immediately or as soon as possible after use, contaminated reusable sharps shall be placed in appropriate containers until properly reprocessed. These containers shall be:

1910.1030(d)(2)(viii)(A)
Puncture resistant;

1910.1030(d)(2)(viii)(B)
Labeled or color-coded in accordance with this standard;

1910.1030(d)(2)(viii)(C)
Leakproof on the sides and bottom; and

1910.1030(d)(2)(viii)(D)
In accordance with the requirements set forth in paragraph (d)(4)(ii)(E) for reusable sharps.

1910.1030(d)(2)(ix)
Eating, drinking, smoking, applying cosmetics or lip balm, and handling contact lenses are prohibited in work areas where there is a reasonable likelihood of occupational exposure.

1910.1030(d)(2)(x)
Food and drink shall not be kept in refrigerators, freezers, shelves, cabinets or on countertops or benchtops where blood or other potentially infectious materials are present.
1910.1030(d)(2)(xi)
All procedures involving blood or other potentially infectious materials shall be performed in such a manner as to minimize splashing, spraying, spattering, and generation of droplets of these substances.

1910.1030(d)(2)(xii)
Mouth pipetting/suctioning of blood or other potentially infectious materials is prohibited.

1910.1030(d)(2)(xiii)
Specimens of blood or other potentially infectious materials shall be placed in a container which prevents leakage during collection, handling, processing, storage, transport, or shipping.

1910.1030(d)(2)(xiv)(A)
The container for storage, transport, or shipping shall be labeled or color-coded according to paragraph (g)(1)(i) and closed prior to being stored, transported, or shipped. When a facility utilizes Universal Precautions in the handling of all specimens, the labeling/color-coding of specimens is not necessary provided containers are recognizable as containing specimens. This exemption only applies while such specimens/containers remain within the facility. Labeling or color-coding in accordance with paragraph (g)(1)(i) is required when such specimens/containers leave the facility.

1910.1030(d)(2)(xiv)(B)
If outside contamination of the primary container occurs, the primary container shall be placed within a second container which prevents leakage during handling, processing, storage, transport, or shipping and is labeled or color-coded according to the requirements of this standard.

1910.1030(d)(2)(xiv)(C)
If the specimen could puncture the primary container, the primary container shall be placed within a secondary container which is puncture-resistant in addition to the above characteristics.

1910.1030(d)(2)(xv)
Equipment which may become contaminated with blood or other potentially infectious materials shall be examined prior to servicing or shipping and shall be decontaminated as necessary, unless the employer can demonstrate that decontamination of such equipment or portions of such equipment is not feasible.

1910.1030(d)(2)(xv)(A)
A readily observable label in accordance with paragraph (g)(1)(i)(H) shall be attached to the equipment stating which portions remain contaminated.

1910.1030(d)(2)(xv)(B)
The employer shall ensure that this information is conveyed to all affected employees, the servicing representative, and/or the manufacturer, as appropriate, prior to handling, servicing, or shipping so that appropriate precautions will be taken.

1910.1030(d)(3)
Personal Protective Equipment --

1910.1030(d)(3)(i)
Provision. When there is occupational exposure, the employer shall provide, at no cost to the employee, appropriate personal protective equipment such as, but not limited to, gloves, gowns, laboratory coats, face shields or masks and eye protection, and mouthpieces, resuscitation bags, pocket masks, or other ventilation devices. Personal protective equipment will be considered "appropriate" only if it does not permit blood or other potentially infectious materials to pass through to or reach the employee's work clothes, street clothes, undergarments, skin, eyes, mouth, or other mucous membranes under normal conditions of use and for the duration of time which the protective equipment will be used.

1910.1030(d)(3)(ii)
Use. The employer shall ensure that the employee uses appropriate personal protective equipment unless the employer shows that the employee temporarily and briefly declined to use personal protective equipment when, under rare and extraordinary circumstances, it was the employee's professional judgment that in the specific instance its use would have prevented the delivery of health care or public safety services or would have posed an increased hazard to the safety of the worker or co-worker. When the employee makes this judgement, the circumstances shall be investigated and documented in order to determine whether changes can be instituted to prevent such occurrences in the future.

1910.1030(d)(3)(iii)
Accessibility. The employer shall ensure that appropriate personal protective equipment in the appropriate sizes is readily accessible at the worksite or is issued to employees. Hypoallergenic gloves, glove liners, powderless gloves, or other similar alternatives shall be readily accessible to those employees who are allergic to the gloves normally provided.

1910.1030(d)(3)(iv)
Cleaning, Laundering, and Disposal. The employer shall clean, launder, and dispose of personal protective equipment required by paragraphs (d) and (e) of this standard, at no cost to the employee.

1910.1030(d)(3)(v)
Repair and Replacement. The employer shall repair or replace personal protective equipment as needed to maintain its effectiveness, at no cost to the employee.

1910.1030(d)(3)(vi)
If a garment(s) is penetrated by blood or other potentially infectious materials, the garment(s) shall be removed immediately or as soon as feasible.

1910.1030(d)(3)(vii)
All personal protective equipment shall be removed prior to leaving the work area.

1910.1030(d)(3)(viii)
When personal protective equipment is removed it shall be placed in an appropriately designated area or container for storage, washing, decontamination or disposal.
Gloves. Gloves shall be worn when it can be reasonably anticipated that the employee may have hand contact with blood, other potentially infectious materials, mucous membranes, and non-intact skin; when performing vascular access procedures except as specified in paragraph (d)(3)(ix)(D); and when handling or touching contaminated items or surfaces.

Disposable (single use) gloves such as surgical or examination gloves, shall be replaced as soon as practical when contaminated or as soon as feasible if they are torn, punctured, or when their ability to function as a barrier is compromised.

Disposable (single use) gloves shall not be washed or decontaminated for re-use.

Utility gloves may be decontaminated for re-use if the integrity of the glove is not compromised. However, they must be discarded if they are cracked, peeling, torn, punctured, or exhibit other signs of deterioration or when their ability to function as a barrier is compromised.

If an employer in a volunteer blood donation center judges that routine gloving for all phlebotomies is not necessary then the employer shall:

Periodically reevaluate this policy;

Make gloves available to all employees who wish to use them for phlebotomy;

Not discourage the use of gloves for phlebotomy; and

Require that gloves be used for phlebotomy in the following circumstances:

When the employee has cuts, scratches, or other breaks in his or her skin;

When the employee judges that hand contamination with blood may occur, for example, when performing phlebotomy on an uncooperative source individual; and

When the employee is receiving training in phlebotomy.

Masks, Eye Protection, and Face Shields. Masks in combination with eye protection devices, such as goggles or glasses with solid side shields, or chin-length face shields, shall be worn whenever splashes, spray, spatter, or droplets of blood or other potentially infectious materials may be generated and eye, nose, or mouth contamination can be reasonably anticipated.

Gowns, Aprons, and Other Protective Body Clothing. Appropriate protective clothing such as, but not limited to, gowns, aprons, lab coats, clinic jackets, or similar outer garments shall be worn in occupational exposure situations. The type and characteristics will depend upon the task and degree of exposure anticipated.

Surgical caps or hoods and/or shoe covers or boots shall be worn in instances when gross contamination can reasonably be anticipated (e.g., autopsies, orthopaedic surgery).

Housekeeping –

General. Employers shall ensure that the worksite is maintained in a clean and sanitary condition. The employer shall determine and implement an appropriate written schedule for cleaning and method of decontamination based upon the location within the facility, type of surface to be cleaned, type of soil present, and tasks or procedures being performed in the area.

All equipment and environmental and working surfaces shall be cleaned and decontaminated after contact with blood or other potentially infectious materials.

Contaminated work surfaces shall be decontaminated with an appropriate disinfectant after completion of procedures; immediately or as soon as feasible when surfaces are overtly contaminated or after any spill of blood or other potentially infectious materials; and at the end of the work shift if the surface may have become contaminated since the last cleaning.

Protective coverings, such as plastic wrap, aluminum foil, or imperviously-backed absorbent paper used to cover equipment and environmental surfaces, shall be removed and replaced as soon as feasible when they become overtly contaminated or at the end of the workshift if they may have become contaminated during the shift.

All bins, pails, cans, and similar receptacles intended for reuse which have a reasonable likelihood for becoming contaminated with blood or other potentially infectious materials shall be inspected and decontaminated on a regularly scheduled basis and cleaned and decontaminated immediately or as soon as feasible upon visible contamination.

Broken glassware which may be contaminated shall not be picked up directly with the hands. It shall be cleaned up using mechanical means, such as a brush and dust pan, tongs, or forceps.
1910.1030(d)(4)(ii)(E) Reusable sharps that are contaminated with blood or other potentially infectious materials shall not be stored or processed in a manner that requires employees to reach by hand into the containers where these sharps have been placed.

1910.1030(d)(4)(iii) Regulated Waste --


1910.1030(d)(4)(iii)(A)(1) Contaminated sharps shall be discarded immediately or as soon as feasible in containers that are:


1910.1030(d)(4)(iii)(A)(1)(iii) Leakproof on sides and bottom; and

1910.1030(d)(4)(iii)(A)(1)(iv) Labeled or color-coded in accordance with paragraph (g)(1)(i) of this standard.

1910.1030(d)(4)(iii)(A)(2) During use, containers for contaminated sharps shall be:

1910.1030(d)(4)(iii)(A)(2)(i) Easily accessible to personnel and located as close as is feasible to the immediate area where sharps are used or can be reasonably anticipated to be found (e.g., laundries);

1910.1030(d)(4)(iii)(A)(2)(ii) Maintained upright throughout use; and


1910.1030(d)(4)(iii)(A)(3) When moving containers of contaminated sharps from the area of use, the containers shall be:

1910.1030(d)(4)(iii)(A)(3)(i) Closed immediately prior to removal or replacement to prevent spillage or protrusion of contents during handling, storage, transport, or shipping;

1910.1030(d)(4)(iii)(A)(3)(ii) Placed in a secondary container if leakage is possible. The second container shall be:


1910.1030(d)(4)(iii)(A)(3)(ii)(B) Constructed to contain all contents and prevent leakage during handling, storage, transport, or shipping; and

1910.1030(d)(4)(iii)(A)(3)(ii)(C) Labeled or color-coded according to paragraph (g)(1)(i) of this standard.

1910.1030(d)(4)(iii)(A)(4) Reusable containers shall not be opened, emptied, or cleaned manually or in any other manner which would expose employees to the risk of percutaneous injury.

1910.1030(d)(4)(iii)(B) Other Regulated Waste Containment --

1910.1030(d)(4)(iii)(B)(1) Regulated waste shall be placed in containers which are:


1910.1030(d)(4)(iii)(B)(1)(ii) Constructed to contain all contents and prevent leakage of fluids during handling, storage, transport or shipping;

1910.1030(d)(4)(iii)(B)(1)(iii) Labeled or color-coded in accordance with paragraph (g)(1)(i) this standard; and

1910.1030(d)(4)(iii)(B)(1)(iv) Closed prior to removal to prevent spillage or protrusion of contents during handling, storage, transport, or shipping.

1910.1030(d)(4)(iii)(B)(2) If outside contamination of the regulated waste container occurs, it shall be placed in a second container. The second container shall be:


1910.1030(d)(4)(iii)(B)(2)(ii) Constructed to contain all contents and prevent leakage of fluids during handling, storage, transport or shipping;

1910.1030(d)(4)(iii)(B)(2)(iii) Labeled or color-coded in accordance with paragraph (g)(1)(i) of this standard; and

1910.1030(d)(4)(iii)(B)(2)(iv) Closed prior to removal to prevent spillage or protrusion of contents during handling, storage, transport, or shipping.

1910.1030(d)(4)(iii)(C) Disposal of all regulated waste shall be in accordance with applicable regulations of the United States, States and Territories, and political subdivisions of States and Territories.


1910.1030(d)(4)(iv)(A) Contaminated laundry shall be handled as little as possible with a minimum of agitation.

1910.1030(d)(4)(iv)(A)(1) Contaminated laundry shall be bagged or containerized at the location where it was used and shall not be sorted or rinsed in the location of use.

1910.1030(d)(4)(iv)(A)(2) Contaminated laundry shall be placed and transported in bags or containers labeled or color-coded in accordance
with paragraph (g)(1)(i) of this standard. When a facility utilizes Universal Precautions in the handling of all soiled laundry, alternative labeling or color-coding is sufficient if it permits all employees to recognize the containers as requiring compliance with Universal Precautions.

1910.1030(d)(4)(iv)(A)(3) Whenever contaminated laundry is wet and presents a reasonable likelihood of soak-through of or leakage from the bag or container, the laundry shall be placed and transported in bags or containers which prevent soak-through and/or leakage of fluids to the exterior.

1910.1030(d)(4)(iv)(B) The employer shall ensure that employees who have contact with contaminated laundry wear protective gloves and other appropriate personal protective equipment.

1910.1030(d)(4)(iv)(C) When a facility ships contaminated laundry off-site to a second facility which does not utilize Universal Precautions in the handling of all laundry, the facility generating the contaminated laundry must place such laundry in bags or containers which are labeled or color-coded in accordance with paragraph (g)(1)(i).

1910.1030(e) HIV and HBV Research Laboratories and Production Facilities.

1910.1030(e)(1) This paragraph applies to research laboratories and production facilities engaged in the culture, production, concentration, experimentation, and manipulation of HIV and HBV. It does not apply to clinical or diagnostic laboratories engaged solely in the analysis of blood, tissues, or organs. These requirements apply in addition to the other requirements of the standard.

1910.1030(e)(2) Research laboratories and production facilities shall meet the following criteria:

1910.1030(e)(2)(i) Standard Microbiological Practices. All regulated waste shall either be incinerated or decontaminated by a method such as autoclaving known to effectively destroy bloodborne pathogens.


1910.1030(e)(2)(ii)(A) Laboratory doors shall be kept closed when work involving HIV or HBV is in progress.

1910.1030(e)(2)(ii)(B) Contaminated materials that are to be decontaminated at a site away from the work area shall be placed in a durable, leakproof, labeled or color-coded container that is closed before being removed from the work area.

1910.1030(e)(2)(ii)(C) Access to the work area shall be limited to authorized persons. Written policies and procedures shall be established whereby only persons who have been advised of the potential biohazard, who meet any specific entry requirements, and who comply with all entry and exit procedures shall be allowed to enter the work areas and animal rooms.

1910.1030(e)(2)(ii)(D) When other potentially infectious materials or infected animals are present in the work area or containment module, a hazard warning sign incorporating the universal biohazard symbol shall be posted on all access doors. The hazard warning sign shall comply with paragraph (g)(1)(ii) of this standard.

1910.1030(e)(2)(ii)(E) All activities involving other potentially infectious materials shall be conducted in biological safety cabinets or other physical-containment devices within the containment module. No work with these other potentially infectious materials shall be conducted on the open bench.

1910.1030(e)(2)(ii)(F) Laboratory coats, gowns, smocks, uniforms, or other appropriate protective clothing shall be used in the work area and animal rooms. Protective clothing shall not be worn outside of the work area and shall be decontaminated before being laundered.

1910.1030(e)(2)(ii)(G) Special care shall be taken to avoid skin contact with other potentially infectious materials. Gloves shall be worn when handling infected animals and when making hand contact with other potentially infectious materials is unavoidable.

1910.1030(e)(2)(ii)(H) Before disposal all waste from work areas and from animal rooms shall either be incinerated or decontaminated by a method such as autoclaving known to effectively destroy bloodborne pathogens.

1910.1030(e)(2)(ii)(I) Vacuum lines shall be protected with liquid disinfectant traps and high-efficiency particulate air (HEPA) filters or filters of equivalent or superior efficiency and which are checked routinely and maintained or replaced as necessary.

1910.1030(e)(2)(ii)(J) Hypodermic 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., the needle is integral to the syringe) shall be used for the injection or aspiration of other potentially infectious materials. Extreme caution shall be used when handling needles and syringes. A needle shall not be bent, sheared, replaced in the sheath or guard, or removed from the syringe following use. The needle and syringe shall be promptly placed in a puncture-resistant container and autoclaved or decontaminated before reuse or disposal.

1910.1030(e)(2)(ii)(K) All spills shall be immediately contained and cleaned up by appropriate professional staff or others properly trained and
equipped to work with potentially concentrated infectious materials.

1910.1030(e)(2)(ii)(L)
A spill or accident that results in an exposure incident shall be immediately reported to the laboratory director or other responsible person.

1910.1030(e)(2)(ii)(M)
A biosafety manual shall be prepared or adopted and periodically reviewed and updated at least annually or more often if necessary. Personnel shall be advised of potential hazards, shall be required to read instructions on practices and procedures, and shall be required to follow them.

1910.1030(e)(2)(iii)
Containment Equipment.

1910.1030(e)(2)(iii)(A)
Certified biological safety cabinets (Class I, II, or III) or other appropriate combinations of personal protection or physical containment devices, such as special protective clothing, respirators, centrifuge safety cups, sealed centrifuge rotors, and containment caging for animals, shall be used for all activities with other potentially infectious materials that pose a threat of exposure to droplets, splashes, spills, or aerosols.

1910.1030(e)(2)(iii)(B)
Biological safety cabinets shall be certified when installed, whenever they are moved and at least annually.

1910.1030(e)(3)
HIV and HBV research laboratories shall meet the following criteria:

1910.1030(e)(3)(i)
Each laboratory shall contain a facility for hand washing and an eye wash facility which is readily available within the work area.

1910.1030(e)(3)(ii)
An autoclave for decontamination of regulated waste shall be available.

1910.1030(e)(4)
HIV and HBV production facilities shall meet the following criteria:

1910.1030(e)(4)(i)
The work areas shall be separated from areas that are open to unrestricted traffic flow within the building. Passage through two sets of doors shall be the basic requirement for entry into the work area from access corridors or other contiguous areas. Physical separation of the high-containment work area from access corridors or other areas or activities may also be provided by a double-doored clothes-change room (showers may be included), airlock, or other access facility that requires passing through two sets of doors before entering the work area.

1910.1030(e)(4)(ii)
The surfaces of doors, walls, floors and ceilings in the work area shall be water resistant so that they can be easily cleaned. Penetrations in these surfaces shall be sealed or capable of being sealed to facilitate decontamination.

1910.1030(e)(4)(iii)
Each work area shall contain a sink for washing hands and a readily available eye wash facility. The sink shall be foot, elbow, or automatically operated and shall be located near the exit door of the work area.

1910.1030(e)(4)(iv)
Access doors to the work area or containment module shall be self-closing.

1910.1030(e)(4)(v)
An autoclave for decontamination of regulated waste shall be available within or as near as possible to the work area.

1910.1030(e)(4)(vi)
A ducted exhaust-air ventilation system shall be provided. This system shall create directional airflow that draws air into the work area through the entry area. The exhaust air shall not be recirculated to any other area of the building, shall be discharged to the outside, and shall be dispersed away from occupied areas and air intakes. The proper direction of the airflow shall be verified (i.e., into the work area).

1910.1030(e)(5)
Training Requirements. Additional training requirements for employees in HIV and HBV research laboratories and HIV and HBV production facilities are specified in paragraph (g)(2)(ix).

1910.1030(f)
Hepatitis B Vaccination and Post-exposure Evaluation and Follow-up --

1910.1030(f)(1)
General.

1910.1030(f)(1)(i)
The employer shall make available the hepatitis B vaccine and vaccination series to all employees who have occupational exposure, and post-exposure evaluation and follow-up to all employees who have had an exposure incident.

1910.1030(f)(1)(ii)
The employer shall ensure that all medical evaluations and procedures including the hepatitis B vaccine and vaccination series and post-exposure evaluation and follow-up to all employees who have had an exposure incident.

1910.1030(f)(1)(ii)(A)
Made available at no cost to the employee;

1910.1030(f)(1)(ii)(B)
Made available to the employee at a reasonable time and place;

1910.1030(f)(1)(ii)(C)
Performed by or under the supervision of a licensed physician or by or under the supervision of another licensed healthcare professional; and
Provided according to recommendations of the U.S. Public Health Service current at the time these evaluations and procedures take place, except as specified by this paragraph (f).

1910.1030(f)(1)(iii)
The employer shall ensure that all laboratory tests are conducted by an accredited laboratory at no cost to the employee.

1910.1030(f)(2)
Hepatitis B Vaccination.

1910.1030(f)(2)(i)
Hepatitis B vaccination shall be made available after the employee has received the training required in paragraph (g)(2)(vii)(l) and within 10 working days of initial assignment to all employees who have occupational exposure unless the employee has previously received the complete hepatitis B vaccination series, antibody testing has revealed that the employee is immune, or the vaccine is contraindicated for medical reasons.

1910.1030(f)(2)(ii)
The employer shall not make participation in a prescreening program a prerequisite for receiving hepatitis B vaccination.

1910.1030(f)(2)(iii)
If the employee initially declines hepatitis B vaccination but at a later date while still covered under the standard decides to accept the vaccination, the employer shall make available hepatitis B vaccination at that time.

1910.1030(f)(2)(iv)
The employer shall assure that employees who decline to accept hepatitis B vaccination offered by the employer sign the statement in Appendix A.

1910.1030(f)(2)(v)
If a routine booster dose(s) of hepatitis B vaccine is recommended by the U.S. Public Health Service at a future date, such booster dose(s) shall be made available in accordance with section (f)(1)(ii).

1910.1030(f)(3)
Post-exposure Evaluation and Follow-up. Following a report of an exposure incident, the employer shall make immediately available to the exposed employee a confidential medical evaluation and follow-up, including at least the following elements:

1910.1030(f)(3)(i)
Documentation of the route(s) of exposure, and the circumstances under which the exposure incident occurred;

1910.1030(f)(3)(ii)
Identification and documentation of the source individual, unless the employer can establish that identification is infeasible or prohibited by state or local law;

1910.1030(f)(3)(ii)(A)
The source individual's blood shall be tested as soon as feasible and after consent is obtained in order to determine HBV and HIV infectivity. If consent is not obtained, the employer shall establish that legally required consent cannot be obtained. When the source individual's consent is not required by law, the source individual's blood, if available, shall be tested and the results documented.

1910.1030(f)(3)(ii)(B)
When the source individual is already known to be infected with HBV or HIV, testing for the source individual's known HBV or HIV status need not be repeated.

1910.1030(f)(3)(ii)(C)
Results of the source individual's testing shall be made available to the exposed employee, and the employee shall be informed of applicable laws and regulations concerning disclosure of the identity and infectious status of the source individual.

1910.1030(f)(3)(iii)
Collection and testing of blood for HBV and HIV serological status;

1910.1030(f)(3)(iii)(A)
The exposed employee's blood shall be collected as soon as feasible and tested after consent is obtained.

1910.1030(f)(3)(iii)(B)
If the employee consents to baseline blood collection, but does not give consent at that time for HIV serologic testing, the sample shall be preserved for at least 90 days. If, within 90 days of the exposure incident, the employee elects to have the baseline sample tested, such testing shall be done as soon as feasible.

1910.1030(f)(3)(iv)
Post-exposure prophylaxis, when medically indicated, as recommended by the U.S. Public Health Service;

1910.1030(f)(3)(v)
Counseling; and

1910.1030(f)(3)(vi)
Evaluation of reported illnesses.

1910.1030(f)(4)
Information Provided to the Healthcare Professional.

1910.1030(f)(4)(i)
The employer shall ensure that the healthcare professional responsible for the employee's Hepatitis B vaccination is provided a copy of this regulation.

1910.1030(f)(4)(ii)
The employer shall ensure that the healthcare professional evaluating an employee after an exposure incident is provided the following information:

1910.1030(f)(4)(ii)(A)
A copy of this regulation;

1910.1030(f)(4)(ii)(B)
A description of the exposed employee's duties as they relate to the exposure incident;

1910.1030(f)(4)(ii)(C)
Documentation of the route(s) of exposure and circumstances under which exposure occurred;
1910.1030(f)(4)(ii)(D)
Results of the source individual's blood testing, if available; and

1910.1030(f)(4)(ii)(E)
All medical records relevant to the appropriate treatment of the employee including vaccination status which are the employer’s responsibility to maintain.

1910.1030(f)(5)
Healthcare Professional’s Written Opinion. The employer shall obtain and provide the employee with a copy of the evaluating healthcare professional’s written opinion within 15 days of the completion of the evaluation.

1910.1030(f)(5)(i)
The healthcare professional’s written opinion for Hepatitis B vaccination shall be limited to whether Hepatitis B vaccination is indicated for an employee, and if the employee has received such vaccination.

1910.1030(f)(5)(ii)
The healthcare professional’s written opinion for post-exposure evaluation and follow-up shall be limited to the following information:

1910.1030(f)(5)(ii)(A)
That the employee has been informed of the results of the evaluation; and

1910.1030(f)(5)(ii)(B)
That the employee has been told about any medical conditions resulting from exposure to blood or other potentially infectious materials which require further evaluation or treatment.

1910.1030(f)(5)(iii)
All other findings or diagnoses shall remain confidential and shall not be included in the written report.

1910.1030(f)(6)
Medical Recordkeeping. Medical records required by this standard shall be maintained in accordance with paragraph (h)(1) of this section.

1910.1030(g)
Communication of Hazards to Employees —

1910.1030(g)(1)
Labels and Signs —

1910.1030(g)(1)(i)
Labels.

1910.1030(g)(1)(i)(A)
Warning labels shall be affixed to containers of regulated waste, refrigerators and freezers containing blood or other potentially infectious material; and other containers used to store, transport or ship blood or other potentially infectious materials, except as provided in paragraph (g)(1)(i)(E), (F) and (G).

1910.1030(g)(1)(i)(B)
Labels required by this section shall include the following legend:

1910.1030(g)(1)(i)(C)
These labels shall be fluorescent orange or orange-red or predominantly so, with lettering and symbols in a contrasting color.

1910.1030(g)(1)(i)(D)
Labels shall be affixed as close as feasible to the container by string, wire, adhesive, or other method that prevents their loss or unintentional removal.

1910.1030(g)(1)(i)(E)
Red bags or red containers may be substituted for labels.

1910.1030(g)(1)(i)(F)
Containers of blood, blood components, or blood products that are labeled as to their contents and have been released for transfusion or other clinical use are exempted from the labeling requirements of paragraph (g).

1910.1030(g)(1)(i)(G)
Individual containers of blood or other potentially infectious materials that are placed in a labeled container during storage, transport, shipment or disposal are exempted from the labeling requirement.

1910.1030(g)(1)(i)(H)
Labels required for contaminated equipment shall be in accordance with this paragraph and shall also state which portions of the equipment remain contaminated.

1910.1030(g)(1)(i)(I)
Regulated waste that has been decontaminated need not be labeled or color-coded.

1910.1030(g)(1)(ii)
Signs.

1910.1030(g)(1)(ii)(A)
The employer shall post signs at the entrance to work areas specified in paragraph (e), HIV and HBV Research Laboratory and Production Facilities, which shall bear the following legend:
1910.1030(g)(1)(ii)(B)
These signs shall be fluorescent orange-red or predominantly so, with lettering and symbols in a contrasting color.

1910.1030(g)(2)
Information and Training.

1910.1030(g)(2)(i)
Employers shall ensure that all employees with occupational exposure participate in a training program which must be provided at no cost to the employee and during working hours.

1910.1030(g)(2)(ii)
Training shall be provided as follows:
1910.1030(g)(2)(ii)(A)
At the time of initial assignment to tasks where occupational exposure may take place;
1910.1030(g)(2)(ii)(B)
Within 90 days after the effective date of the standard; and
1910.1030(g)(2)(ii)(C)
At least annually thereafter.

1910.1030(g)(2)(iii)
For employees who have received training on bloodborne pathogens in the year preceding the effective date of the standard, only training with respect to the provisions of the standard which were not included need be provided.

1910.1030(g)(2)(iv)
Annual training for all employees shall be provided within one year of their previous training.

1910.1030(g)(2)(v)
Employers shall provide additional training when changes such as modification of tasks or procedures or institution of new tasks or procedures affect the employee’s occupational exposure. The additional training may be limited to addressing the new exposures created.

1910.1030(g)(2)(vi)
Material appropriate in content and vocabulary to educational level, literacy, and language of employees shall be used.

1910.1030(g)(2)(vii)
The training program shall contain at a minimum the following elements:
1910.1030(g)(2)(vii)(A)
An accessible copy of the regulatory text of this standard and an explanation of its contents;
1910.1030(g)(2)(vii)(B)
A general explanation of the epidemiology and symptoms of bloodborne diseases;
Additional Initial Training for Employees in HIV and HBV Laboratories and Production Facilities. Employees in HIV or HBV research laboratories and HIV or HBV production facilities shall receive the following initial training in addition to the above training requirements.

1910.1030(g)(2)(ix)(A)
The employer shall assure that employees demonstrate proficiency in standard microbiological practices and techniques and in the practices and operations specific to the facility before being allowed to work with HIV or HBV.

1910.1030(g)(2)(ix)(B)
The employer shall assure that employees have prior experience in the handling of human pathogens or tissue cultures before working with HIV or HBV.

1910.1030(g)(2)(ix)(C)
The employer shall provide a training program to employees who have no prior experience in handling human pathogens. Initial work activities shall not include the handling of infectious agents. A progression of work activities shall be assigned as techniques are learned and proficiency is developed. The employer shall assure that employees participate in work activities involving infectious agents only after proficiency has been demonstrated.

1910.1030(h) Recordkeeping --

1910.1030(h)(1) Medical Records.

1910.1030(h)(1)(i)
The employer shall establish and maintain an accurate record for each employee with occupational exposure, in accordance with 29 CFR 1910.1020.

1910.1030(h)(1)(ii)
This record shall include:

1910.1030(h)(1)(ii)(A)
The name and social security number of the employee;

1910.1030(h)(1)(ii)(B)
A copy of the employee's hepatitis B vaccination status including the dates of all the hepatitis B vaccinations and any medical records relative to the employee's ability to receive vaccination as required by paragraph (f)(2);

1910.1030(h)(1)(ii)(C)
A copy of all results of examinations, medical testing, and follow-up procedures as required by paragraph (f)(3);

1910.1030(h)(1)(ii)(D)
The employer's copy of the healthcare professional's written opinion as required by paragraph (f)(5); and

1910.1030(h)(1)(ii)(E)
A copy of the information provided to the healthcare professional as required by paragraphs (f)(4)(ii)(B)(C) and (D).

1910.1030(h)(1)(iii)
Confidentiality. The employer shall ensure that employee medical records required by paragraph (h)(1) are:

1910.1030(h)(1)(iii)(A)
Kept confidential; and

1910.1030(h)(1)(iii)(B)
Not disclosed or reported without the employee's express written consent to any person within or outside the workplace except as required by this section or as may be required by law.

1910.1030(h)(1)(iv)
The employer shall maintain the records required by paragraph (h) for at least the duration of employment plus 30 years in accordance with 29 CFR 1910.1020.

1910.1030(h)(2) Training Records.

1910.1030(h)(2)(i)
Training records shall include the following information:

1910.1030(h)(2)(i)(A)
The dates of the training sessions;

1910.1030(h)(2)(i)(B)
The contents or a summary of the training sessions;

1910.1030(h)(2)(i)(C)
The names and qualifications of persons conducting the training; and

1910.1030(h)(2)(i)(D)
The names and job titles of all persons attending the training sessions.

1910.1030(h)(2)(ii)
Training records shall be maintained for 3 years from the date on which the training occurred.

1910.1030(h)(3) Availability.

1910.1030(h)(3)(i)
The employer shall ensure that all records required to be maintained by this section shall be made available upon request to the Assistant Secretary and the Director for examination and copying.

1910.1030(h)(3)(ii)
Employee training records required by this paragraph shall be provided upon request for examination and copying to employees, to employee representatives, to the Director, and to the Assistant Secretary.

1910.1030(h)(3)(iii)
Employee medical records required by this paragraph shall be provided upon request for examination and copying to the subject employee, to anyone having written consent of the subject employee, to the Director, and to the Assistant Secretary in accordance with 29 CFR 1910.1020.

1910.1030(h)(4) Transfer of Records.

1910.1030(h)(4)(i)

The employer shall comply with the requirements involving transfer of records set forth in 29 CFR 1910.1020(h).

1910.1030(h)(4)(ii)
If the employer ceases to do business and there is no successor employer to receive and retain the records for the prescribed period, the employer shall notify the Director, at least three months prior to their disposal and transmit them to the Director, if required by the Director to do so, within that three month period.

1910.1030(h)(5)
Sharps injury log.

1910.1030(h)(5)(i)
The employer shall establish and maintain a sharps injury log for the recording of percutaneous injuries from contaminated sharps. The information in the sharps injury log shall be recorded and maintained in such manner as to protect the confidentiality of the injured employee. The sharps injury log shall contain, at a minimum:

1910.1030(h)(5)(i)(A) The type and brand of device involved in the incident,
1910.1030(h)(5)(i)(B) The department or work area where the exposure incident occurred, and
1910.1030(h)(5)(i)(C) An explanation of how the incident occurred.

1910.1030(h)(5)(ii)
The requirement to establish and maintain a sharps injury log shall apply to any employer who is required to maintain a log of occupational injuries and illnesses under 29 CFR 1904.

1910.1030(h)(5)(iii)
The sharps injury log shall be maintained for the period required by 29 CFR 1904.6.

1910.1030(i)
Dates --

1910.1030(i)(1) Effective Date. The standard shall become effective on March 6, 1992.

1910.1030(i)(2) The Exposure Control Plan required by paragraph (c) of this section shall be completed on or before May 5, 1992.

1910.1030(i)(3) Paragraph (g)(2) Information and Training and (h) Recordkeeping shall take effect on or before June 4, 1992.


Appendix A to Section 1910.1030 -Hepatitis B Vaccine Declination (Mandatory)

I understand that due to my occupational exposure to blood or other potentially infectious materials I may be at risk of acquiring hepatitis B virus (HBV) infection. I have been given the opportunity to be vaccinated with hepatitis B vaccine, at no charge to myself. However, I decline hepatitis B vaccination at this time. I understand that by declining this vaccine, I continue to be at risk of acquiring hepatitis B, a serious disease. If in the future I continue to have occupational exposure to blood or other potentially infectious materials and I want to be vaccinated with hepatitis B vaccine, I can receive the vaccination series at no charge to me.

APPENDIX B: SYMPTOMS, ROUTES OF ENTRY AND TRANSMISSION OF COMMON BLOODBORNE PATHOGENS

<table>
<thead>
<tr>
<th>AGENT</th>
<th>DISEASE</th>
<th>INCUBATION PERIOD (initial infection)</th>
<th>PRINCIPAL METHOD OF SPREAD (TRANSMISSION)</th>
<th>SYMPTOMS TO WATCH FOR</th>
<th>IMMUNITY</th>
<th>CARRIER STATE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plasmodium Species</td>
<td>malaria</td>
<td>10-35 days</td>
<td>Arthropod bite, transfusion</td>
<td>weakness, sweating, headache, fever, chills, muscle pain</td>
<td>gradual</td>
<td>no</td>
</tr>
<tr>
<td>Treponema pallidum</td>
<td>acquired syphilis</td>
<td>9-90 days</td>
<td>Sexual</td>
<td>sore in genital area</td>
<td>no</td>
<td>no</td>
</tr>
<tr>
<td>HIV</td>
<td>AIDS</td>
<td>2 mos-10 yrs</td>
<td>Sexual, broken skin/mucous membrane, Transfusion</td>
<td>flu-like illness</td>
<td>no</td>
<td>yes</td>
</tr>
<tr>
<td>HBV</td>
<td>viral bloodborne hepatitis</td>
<td>40-180 days</td>
<td>Sexual, broken skin/mucous membrane, transfusion</td>
<td>fatigue, vomiting, fever</td>
<td>yes</td>
<td>yes</td>
</tr>
<tr>
<td>HDV</td>
<td>Same as HBV</td>
<td>2-10 weeks</td>
<td>Same as HBV</td>
<td>same as HBV</td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td>HCV</td>
<td>Same as HBV</td>
<td>2 wks-6 mos</td>
<td>Transfusion</td>
<td>same as HBV</td>
<td>?</td>
<td>yes</td>
</tr>
<tr>
<td>Leptospira interrogans</td>
<td>leptospirosis</td>
<td>4-20 days</td>
<td>Broken skin/mucous membrane, inhalation</td>
<td>impaired senses, headache, fever, chills, muscle pain, rash</td>
<td>no</td>
<td>no</td>
</tr>
<tr>
<td>Borrelia species</td>
<td>relapsing fever</td>
<td>2-15 days</td>
<td>Arthropod bite</td>
<td>sudden fever, severe headache, chills, nerve and muscle pain</td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td>Borrelia burgdorferi</td>
<td>Lyme disease</td>
<td>2-30 days</td>
<td>Arthropod bite</td>
<td>spread red rash, swollen glands, stiff neck</td>
<td>no</td>
<td>no</td>
</tr>
</tbody>
</table>

HBV is the only bloodborne pathogen that has an approved vaccine.
<table>
<thead>
<tr>
<th>AGENT</th>
<th>DISEASE</th>
<th>INCUBATION PERIOD (initial infection)</th>
<th>PRINCIPAL METHOD OF SPREAD (TRANSMISSION)</th>
<th>SYMPTOMS TO WATCH FOR</th>
<th>IMMUNITY</th>
<th>CARRIER STATE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brucella species</td>
<td>brucellosis</td>
<td>5-60 days</td>
<td>Broken skin/mucous membrane, inhalation, ingestion</td>
<td>sudden fever night sweats, chills, severe headache, body aches</td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td>Francisella tularensis</td>
<td>tularemia</td>
<td>1-10 days</td>
<td>Arthropod bite, ingestion, inhalation</td>
<td>flu-like illness</td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td>Babesia species</td>
<td>babesiosis</td>
<td>1 wk to 12 mos</td>
<td>Insect bite, transfusion</td>
<td>same as malaria</td>
<td>no</td>
<td>no</td>
</tr>
<tr>
<td>Streptobacillus moniliformis</td>
<td>streptobacillary rat-bite fever</td>
<td>3-20 days</td>
<td>Rat bite, ingestion</td>
<td>rash, headache, chills, fever, arthritic symptoms</td>
<td>no?</td>
<td>no</td>
</tr>
<tr>
<td>Spirillum minus</td>
<td>spirillary (rat-bite) fever</td>
<td>1-3 weeks</td>
<td>Arthropod bite</td>
<td>same as streptobacillus</td>
<td>no?</td>
<td>no</td>
</tr>
<tr>
<td>Creutzfeldt-Jakob virus</td>
<td>Creutzfeldt-Jakob disease</td>
<td>15 mos-20 yrs</td>
<td>Broken skin/mucous membrane</td>
<td>confusion, madness</td>
<td>?</td>
<td>?</td>
</tr>
<tr>
<td>Marburg virus</td>
<td>hemorrhagic fever</td>
<td>3 days-2 weeks</td>
<td>Broken skin/mucous membrane</td>
<td>sudden fever, weakness, severe leg pain, rash, stomach pain</td>
<td>?</td>
<td>no</td>
</tr>
<tr>
<td>Junin virus Machupo virus</td>
<td>hemorrhagic fever</td>
<td>7-16 days</td>
<td>Scratched or broken skin/mucous membrane, inhalation of dried infected rodent feces and urine</td>
<td>same as Marburg/Ebola and sweats, sore throat</td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td>Crimean-Congo Virus</td>
<td>hemorrhagic fever</td>
<td>3-12 days</td>
<td>Arthropod bite, also infected rodents and domesticated animals, broken skin/mucous membrane</td>
<td>same as Marburg/Ebola</td>
<td>some</td>
<td>no</td>
</tr>
<tr>
<td>HTLV-I</td>
<td>leukemia/lymphoma (T-cell origin)</td>
<td>Unknown</td>
<td>Transfusion</td>
<td>-</td>
<td>no</td>
<td>yes</td>
</tr>
<tr>
<td>Several arboviruses</td>
<td>Colorado tick fever</td>
<td>3-12 days</td>
<td>Arthropod bite, transfusion</td>
<td>fever, headache, stiff neck, confusion</td>
<td>yes</td>
<td>no</td>
</tr>
</tbody>
</table>
APPENDIX C: SAMPLE OF EQUIPMENT DECONTAMINATION TAG

BIOSAFETY NOTICE

This equipment’s exterior and interior surfaces were decontaminated, and are free of Biological Hazards. This notice does not apply to radiation or chemical hazards (if any).

This equipment is released for: (circle one)

- Service/Repair
- Relocation
- Discard

Decontamination performed by: ____________________________________________

Chemical or disinfectant used: ____________________________________________

Date of decontamination: ________________________________________________

Location of equipment: __________________________________________________

Lab telephone number: __________________________________________________

Note: The following areas ____________________________________________ of this equipment remain contaminated and a biohazard warning label has been attached near the contaminated area.

Additional forms are available through the Office of Environmental Health and Safety.

Yale University
Office of Environmental Health and Safety 203-785-3550
APPENDIX D: Sharps Safety Device Evaluation Record and Form

<table>
<thead>
<tr>
<th>Sharps Safety Device Evaluation Record</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Evaluation performed due to:</strong></td>
</tr>
<tr>
<td>- Follow-up to an injury/exposure involving a contaminated sharp</td>
</tr>
<tr>
<td>- Proactive review of sharps use with human material, other potentially infectious materials (e.g. human or animal pathogens)</td>
</tr>
<tr>
<td><strong>Evaluation Date:</strong> ________________</td>
</tr>
<tr>
<td><strong>Principal Investigator:</strong> ________________</td>
</tr>
<tr>
<td><strong>Department:</strong> ___________________</td>
</tr>
<tr>
<td><strong>Building/Room#:</strong> ___________________</td>
</tr>
<tr>
<td><strong>Contact Employee:</strong> ________________</td>
</tr>
<tr>
<td><strong>Phone:</strong> ___________________</td>
</tr>
<tr>
<td><strong>Fax:</strong> ___________________</td>
</tr>
<tr>
<td><strong>Email:</strong> ___________________</td>
</tr>
<tr>
<td><strong>Procedure involving a contaminated sharp:</strong></td>
</tr>
<tr>
<td><strong>Type/Brand of sharp currently in use:</strong></td>
</tr>
<tr>
<td><strong>Recommendation:</strong></td>
</tr>
<tr>
<td>- Elimination of sharp from procedure</td>
</tr>
<tr>
<td>- Substitution with a safe sharps device</td>
</tr>
<tr>
<td>- Use of engineering controls</td>
</tr>
<tr>
<td>- Implementation of safe work practices</td>
</tr>
<tr>
<td>- Personal Protective Equipment</td>
</tr>
<tr>
<td>- No recommendation needed at this time, effective safety device(s) currently in use.</td>
</tr>
<tr>
<td><strong>Device(s) in use:</strong></td>
</tr>
<tr>
<td><strong>Results of training and evaluation of new device:</strong></td>
</tr>
<tr>
<td>- <strong>Type/Brand of sharp(s) evaluated:</strong></td>
</tr>
<tr>
<td>- <strong>List employees involved in formal evaluation of safe sharps device(s):</strong> ________________</td>
</tr>
<tr>
<td>- <strong>Training date for work with new safe sharps device(s):</strong> ________________</td>
</tr>
<tr>
<td>- <strong>Device(s) formally in use following evaluation (selection/use date):</strong> ________________</td>
</tr>
</tbody>
</table>

Complete and return all forms to: Office of Environmental Health and Safety  
135 College St., First Floor  
Fax# 203-785-7588
SAFETY DEVICE EVALUATION FORM

Device: ____________________________ Number of times used: ____________

Product Name/Supplier: ______________________________________________________

Applications: __________________________________________________________________

Reviewer: ____________________________ Department: ____________________________ Date: ______

Circle the most appropriate answer for each question. Not applicable (N/A) may be used if the question does not apply to this particular product.

SAFETY SYRINGES

During Use:
1. The safety feature can be activated using a one-handed technique 1 2 3 4 5 N/A
2. The safety feature does not obstruct vision of the tip of the sharp 1 2 3 4 5 N/A
3. Use of this product requires you to use the safety feature 1 2 3 4 5 N/A
4. This product does not require more time to use than a non-safety device 1 2 3 4 5 N/A
5. The safety feature works well with a wide variety of hand sizes 1 2 3 4 5 N/A
6. The device is easy to handle while wearing gloves 1 2 3 4 5 N/A
7. This device does not interfere with uses that do not require a needle 1 2 3 4 5 N/A
8. This device offers a good view of any aspirated fluid 1 2 3 4 5 N/A
9. This device will work with all required syringe and needle sizes 1 2 3 4 5 N/A
10. This device provides a better alternative to traditional recapping 1 2 3 4 5 N/A

After Use:
11. There is a clear and unmistakable change (audible or visible) that occurs when the safety feature is activated 1 2 3 4 5 N/A
12. The safety feature operates reliably 1 2 3 4 5 N/A
13. The exposed sharp is permanently blunted or covered after use and prior to disposal 1 2 3 4 5 N/A
14. This device is no more difficult to process after use than non-safety devices 1 2 3 4 5 N/A

Training:
15. The user does not need extensive training for correct operation 1 2 3 4 5 N/A
16. The design of the device suggests proper use 1 2 3 4 5 N/A
17. It is not easy to skip a crucial step in proper use of the device 1 2 3 4 5 N/A

I.V. ACCESS DEVICES

1. The safety feature can be activated using a one-handed technique 1 2 3 4 5 N/A
2. The safety feature does not interfere with normal use of this product 1 2 3 4 5 N/A
3. Use of this product requires you to use the safety feature 1 2 3 4 5 N/A
4. This product does not require more time to use than a non-safety device 1 2 3 4 5 N/A
5. The safety feature works well with a wide variety of hand sizes 1 2 3 4 5 N/A
6. The device allows for rapid visualization of flashback in the catheter or chamber 1 2 3 4 5 N/A
7. Use of this product increases the number of sticks to the patient 1 2 3 4 5 N/A
8. The product stops the flow of blood after the needle is removed from the catheter (or after the butterfly is inserted) and just prior to line connections or hep-lock capping 1 2 3 4 5 N/A
9. A clear and unmistakable change (either audible or visible) occurs when the safety feature is activated 1 2 3 4 5 N/A
10. The safety feature operates reliably 1 2 3 4 5 N/A
11. The exposed sharp is blunted or covered after use and prior to disposal 1 2 3 4 5 N/A
12. The product does not need extensive training to be operated correctly 1 2 3 4 5 N/A
### SAFETY FEATURE EVALUATION FORM (continue)

#### I.V. CONNECTORS

<p>| | | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
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<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Use of this connector eliminates the need for exposed needles in connections</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. The safety feature <strong>does not</strong> interfere with normal use of this product</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Use of this product requires you to use the safety feature</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. This product <strong>does not</strong> require more time to use than a non-safety device</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. The safety feature works well with a wide variety of hand sizes</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. The safety feature allows you to collect blood directly into a vacuum tube, eliminating the need for needles</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. The connector can be secured (locked) to Y-sites, hep-locks, and central lines</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. A clear and unmistakable change (either audible or visible) occurs when the safety feature is activated</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9. The safety feature operates reliably</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10. The exposed sharp is blunted or covered after use and prior to disposal</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. The product does not need extensive training to be operated correctly</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### VACUUM TUBE BLOOD COLLECTION SYSTEMS

<p>| | | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The safety feature can be activated using a one-handed technique</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. The safety feature <strong>does not</strong> interfere with normal use of this product</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Use of this product requires you to use the safety feature</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. This product does not require more time to use than a non-safety device</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. The safety feature works well with a wide variety of hand sizes</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. The safety feature works with a butterfly</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. A clear and unmistakable change (either audible or visible) occurs when the safety feature is activated</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. The safety feature operates reliably</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9. The exposed sharp is blunted or covered after use and prior to disposal</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10. The inner vacuum tube needle (rubber sleeved needle) does not present a danger of exposure</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. The product does not need extensive training to be operated correctly</td>
<td>1 2 3 4 5 N/A</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

### Patient Safety and Comfort

<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
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</thead>
<tbody>
<tr>
<td>Does the device minimize the risk of infection to the patient (e.g., through cross-contamination)?</td>
<td>Yes</td>
<td>No</td>
<td></td>
</tr>
<tr>
<td>Can the device be used without causing more patient discomfort than a conventional device?</td>
<td>Yes</td>
<td>No</td>
<td></td>
</tr>
</tbody>
</table>

*For IV devices*: Does the device attach comfortably (i.e., without causing patient discomfort at the catheter … port or IV tubing)

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<tbody>
<tr>
<td>Yes</td>
<td>No</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Are there other questions that you feel should be asked regarding the safety/utility of this product?

Would you recommend using this device? Yes No

**Comments** (e.g., describe problems, list incompatibilities)
Sharps Safety Device Evaluation Form

Device: ____________________________ Number of times used: ________________

Product Name/Supplier: ____________________________

Applications: ____________________________

Reviewer: ____________________________ Department: ____________________________ Date: ________________

**Circle** the most appropriate answer for each question. Not applicable (N/A) may be used if the question does not apply to this particular product.

<table>
<thead>
<tr>
<th>HEALTHCARE WORKER SAFETY:</th>
<th>agree...........disagree</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The device prevent needlesticks during use (i.e., before disposal)</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>After use, the safety mechanism remain activated through disposal of the device</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>2. The device provide protection one of the following ways: Either intrinsically or</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>automatically (N/A° if a specific action by the user is required to activate the safety</td>
<td></td>
</tr>
<tr>
<td>mechanism.)</td>
<td></td>
</tr>
<tr>
<td>3. If “N/A,” the mechanism activated in one of the following ways: either by one-handed</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>technique or by a two-handed technique accomplished as part of the usual procedure</td>
<td></td>
</tr>
<tr>
<td>4. During the use of device, the user’s hands remain behind the needle until activation</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>of the safety mechanism is complete</td>
<td></td>
</tr>
<tr>
<td>5. The safety mechanism is reliable when activated properly</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>6. The device minimize the risk of user exposure to the patient’s blood</td>
<td>1 2 3 4 5 N/A</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PATIENT SAFETY AND COMFORT:</th>
<th>1 2 3 4 5 N/A</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The device minimize the risk of infection to the patient (e.g., through cross-contamination)</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>2. The device can be used without causing more patient discomfort than a conventional device</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>3. <em>For IV devices</em>: the device is attach comfortably (i.e., without causing patient discomfort) at the catheter port or IV tubing</td>
<td>1 2 3 4 5 N/A</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>EASE OF USE AND TRAINING:</th>
<th>1 2 3 4 5 N/A</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The device operation is obvious. That is, can the device be used properly without</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>extensive training</td>
<td></td>
</tr>
<tr>
<td>2. The device can be used by a left-handed person as easily as by a right handed person</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>3. The technique required for using the device is the same as that for using a conventional device</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>4. It is easy to identify the type and size of the product from the packaging</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>5. <em>For intravenous (IV) catheters and blood collection needle sets</em>: The device can provide a visible blood flashback during initial insertion</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>6. Ease of using device</td>
<td>1 2 3 4 5 N/A</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>COMPATIBILITY:</th>
<th>1 2 3 4 5 N/A</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. The device is compatible with devices (e.g., blood collection tubes) from a variety of suppliers</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>2. <em>For IV devices</em>:</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>A. The device is compatible with intralipid solutions</td>
<td></td>
</tr>
<tr>
<td>B. The device is attach securely at the catheter port</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>C. The device is attach securely or lock at a Y-site (e.g., for piggybacking)</td>
<td>1 2 3 4 5 N/A</td>
</tr>
<tr>
<td>3. The device is easy to dispose of in sharps containers of all sizes (if required)</td>
<td>1 2 3 4 5 N/A</td>
</tr>
</tbody>
</table>

Would you recommend using this device? Yes No

COMMENTS (e.g., describe problems, list incompatibilities)
APPENDIX E: PROPER USE OF BIOLOGICAL SAFETY CABINETS

- A properly balanced and properly used Biological Safety Cabinet (BSC) will do an excellent job of controlling airborne contaminants only if appropriate contamination control procedures and aseptic techniques are also employed.

- Position the BSC away from doorways, high traffic areas, room ventilation systems, air conditioners, and low ceilings. Common room air currents can disrupt the protective air barrier of a BSC. The minimum distance from the top of the BSC to the ceiling is ten inches; this will allow for proper airflow and repairs when needed.

- All BSCs shall be professionally certified at the time of installation and annually thereafter. If a BSC is to be moved, it shall be professionally formaldehyde decontaminated before moving, and recertified before work commences. Contact the Office of Environmental Health and Safety at 737-2125 for assistance with professional certification and decontamination. All service, repairs, and certifications must be performed by the vendor contracted by Yale University.

- Keep the insides and tops of BSCs free of unnecessary equipment or supplies. Clutter inside and on top of the BSC may effect proper air flow or damage the exhaust HEPA filter.

- Some BSCs are equipped with ultraviolet (UV) lights. If good procedures are followed, UV lights are not needed. All UV lights shall be turned off whenever the laboratory is occupied.

- Infectious agents assigned to Biosafety Level 4 shall not be used on campus or in this type of cabinet.

- Avoid using toxic, explosive, flammable, or radioactive substances unless a safety professional has approved them for work in your BSC.

- To begin the BSC operation, turn on the fluorescent lights, confirm the air intake and exhaust grills are clear, and turn on the blower. If a drain valve is present, make certain it is closed.

- Wash hands and arms with germicidal soap before and after work in the BSC. Operators shall wear long sleeved gowns with tight fitting cuffs, and gloves. This measure protects the operator's hands and arms from contamination, and minimizes the shedding of skin flora into the work area.

- Disinfect interior surfaces of the work area using freshly prepared 10% household chlorine bleach or an EPA registered tuberculocidal disinfectant. Use a contact time appropriate for the agent in use. If bleach is used in the BSC, follow up with a wipe down of surfaces with 70% ethanol to remove any corrosive residues from work surface.

- Everything needed for the complete procedure shall be placed in the BSC before starting work. Nothing shall pass in or out through the air barrier until the procedure is completed. Place a pan containing an appropriate disinfectant into the BSC for discarding contaminated materials. Avoid overloading the work area, and thereby compromising the efficacy of the BSC.

- Work supplies are best arranged to segregate clean from dirty materials.

- Set the view screen at the proper height.

- Wait five minutes after all materials have been placed in the BSC before beginning work. This will enable the BSC to purge airborne contaminants from the work area.

- Work as far to the back of the BSC workspace as possible.

- Always use mechanical pipetting aids.

- Avoid using open flames inside BSCs. If a flame must be present, use a burner with a pilot light and place it to the rear of the workspace. Flames disrupt the unidirectional airflow and contribute to the heat load inside the BSC. Flames have shortened the lifetime of HEPA filters, burned holes through HEPA filters and have caused explosions in BSCs.

- Do not work in a BSC while a warning light or alarm is signaling.
After completion of work, enclose or cover all equipment and materials. Wipe down items such as flasks and bottles with an appropriate disinfectant prior to removal from the BSC. Allow the BSC to run for five minutes to purge airborne contaminants from the work area.

Decontaminate interior surfaces with freshly prepared 10% household chlorine bleach or an EPA registered tuberculocidal disinfectant after removal of all materials, cultures and apparatus. Use a contact time appropriate for the agent in use. If bleach is used in the BSC, follow up with a wipe down of surfaces with 70% ethanol to remove any corrosive residues from work surface.

Periodically decontaminate under work grills and work surfaces if these parts are removable.

When the blower is shut off, the air barrier is destroyed. Within seconds, the inside of the cabinet becomes contaminated with microorganisms from the laboratory. For this reason, some manufacturers recommend that BSCs be left operating continuously (24 hours a day).

ACCIDENTS OR SPILLS: In the event of a spill, all surfaces and items shall be surface decontaminated before being removed from the BSC. If the spill results in puddles, flood the area with an appropriate disinfectant for a sufficient time to achieve a complete kill. If a drain system is involved, consult the BSC manufacturer's specific instructions regarding decontamination. After a spill is decontaminated, the area shall be thoroughly cleaned and dried. Residual materials can support the growth and multiplication of microorganisms, and can jeopardize the product protection normally provided by BSCs.

Revised 6/05
APPENDIX F: UNIVERSAL PRECAUTIONS SPILL RESPONSE GUIDE

Prepare and maintain a spill response kit. Basic equipment is some concentrated disinfectant (chlorine bleach), a package of paper towels, household rubber gloves, full face protection, biohazard bags, and dust pan/brush, forceps to pick up broken glass. The contents of the kit can be kept in a small sharps container or plastic container.

**Biosafety Level 2 (BL2) Spill**

1. Avoid inhaling airborne material, while quickly leaving the room. Notify others to leave. Close door, and post with a warning sign.
2. Remove contaminated clothing, turn exposed areas inward, and place in a biohazard bag.
3. Wash all exposed skin with disinfectant.
4. Inform Supervisor, and, if assistance is needed, consult OHS Biosafety (203-785-3550).

**Clean-up of BL2 Spill:**

1. Allow aerosols to disperse for at least 30 minutes before reentering the laboratory. Assemble clean-up materials (disinfectant, paper towels, biohazard bags, and forceps).
2. Put on protective clothing (lab coat or tyvek, face protection, utility gloves, and booties if necessary). Depending on the nature of the spill, it may be advisable to wear a HEPA filtered respirator instead of a surgical mask.
3. Cover the area with disinfectant-soaked towels, and then carefully pour disinfectant around the spill. Avoid enlarging the contaminated area. Use more concentrated disinfectant as it is diluted by the spill. Allow at least a 20 minute contact time.
4. Handle any sharps objects with forceps and discard in a sharps container. Wipe surrounding areas (where the spill may have splashed) with disinfectant.
5. Soak up the disinfectant and spill, and place the materials into a biohazard bag.
6. Spray the area with 10% household bleach solution and allow to air-dry (or wipe down with disinfectant-soaked towels after a 10 minute contact time). Place all contaminated paper towels and any contaminated protective clothing into a biohazard bag and autoclave.
7. Wash hands and exposed skin areas with disinfectant or antiseptic soap and water.

**Blood Spills** (For blood or other material with a high organic content and low concentration of infectious microorganisms)

1. Wear gloves, eye protection, and a lab coat (or tyvek).
2. Absorb blood with paper towels or disinfectant-soaked paper towels and place in a biohazard bag. Collect any sharp objects with forceps or other mechanical device and place in a sharps container.
3. Using a detergent solution, clean the spill site of all visible blood.
4. Spray the spill site with 10% household bleach and allow to air-dry for 15 minutes.
5. After the 15 minute contact time, wipe the area down with disinfectant-soaked paper towels. Discard all disposable materials used to decontaminate the spill and any contaminated personal protective equipment into a biohazard bag. Decontaminate any reusable items with disinfectant.
6. Wash your hands.
APPENDIX G: HEPATITIS B VACCINE NOTIFICATION FORM

Yale University

HEPATITIS B VACCINE NOTIFICATION FORM

I understand that due to my occupational exposure to blood or other potentially infectious materials I may be at risk of acquiring hepatitis B virus (HBV) infection.

I have been given the opportunity to be vaccinated with hepatitis B vaccine, at no charge to myself.

☐ However, I have declined Hepatitis B vaccination at this time. I understand that by declining this vaccine, I continue to be at risk of acquiring Hepatitis B, a serious disease. If in the future I continue to have occupational exposure to blood or other potentially infectious materials and I want to be vaccinated with hepatitis B vaccine, I can receive the vaccination series at no charge to me.

☐ Yes, I wish to be vaccinated against Hepatitis B if recommended by the Department of Employee Health.

☐ I have already received the Hepatitis B vaccine.

________________________________________________________________________
Name (Please Print)  Signature

________________________________________________________________________
Department  Net ID

________________________________________________________________________
Campus address  Date

Telephone (8:30 a.m. - 5:00 p.m.)

Please return completed copy to: Environmental Health & Safety
Bloodborne Pathogen Program
135 College Street, Suite 100
APPENDIX H: BLOODBORNE PATHOGEN QUESTIONNAIRE

Yale University

CLINICAL AND LABORATORY BLOODBORNE PATHOGEN QUESTIONNAIRE

Name: ____________________________ Date: ________________
(Please Print)

Department: ________________________ Net ID: ________________

This Questionnaire is designed to survey employee awareness about Bloodborne Pathogen workplace precautions. It will also be a record of your training.

T=true  F=false
T  F   1. Universal Precautions is an approach to infection control that eliminates or minimizes an employee’s exposure to infectious agents that may be present in human blood, tissue and certain body fluids.

T  F   2. Employees with cuts or dermatitis on their hands, are not required to wear gloves when performing a procedure that may cause exposure to blood and other potential infectious materials.

T  F   3. Hands are washed immediately or as soon as feasible after gloves are removed.

T  F   4. HBV and HIV are transmitted by mucous membrane or parenteral exposure to human blood and body fluids.

T  F   5. A Biological Safety Cabinet is an Engineering Control that offers personal, product, and environmental protection.

T  F   6. Do not recap, bend, break, or manipulate sharps waste by hand.

T  F   7. Face and eye protection are worn when blood or other potentially infectious materials may splash or splatter in the eyes, nose, mouth or other facial areas.

T  F   8. Hepatitis B vaccine is offered to all occupationally exposed employees and protects against Hepatitis B virus, as well as against Hepatitis D virus

T  F   9. Where engineering controls will reduce employee exposure either by removing, eliminating, or isolating the hazard, they must be used.

T  F  10. Employers must solicit input from non-managerial employees who would use a sharp with engineered sharp injury protection device when identifying, evaluating, and selecting a safer sharps device

T  F  12. Use 10% household bleach or an EPA registered tuberculocidal disinfectant to decontaminate work surfaces.
11. Work surfaces are decontaminated:
   a) immediately or soon as feasible when overtly contaminate
   b) after a spill of blood or other potentially infectious materials
   c) at the end of the work shift
   d) all of the above

12. Disposable (single use) gloves are worn when:
   a) contact with blood, mucous membranes, and non-intact skin is anticipated
   b) performing vascular access procedures
   c) handling or touching contaminated surfaces or items
   d) cleaning up blood spills
   e) all of the above

13. Remove Personal Protective Equipment (PPE) and wash hands:
   a) when leaving the clinic or laboratory area for a common area (e.g. lunchroom)
   b) after performing vascular access procedures between patients
   c) before touching phones, computers, door knobs, or elevator buttons
   d) all of the above

14. An “occupationally exposed” employee is defined as:
   a) an employee who has a needlestick from a used needle
   b) an employee who has blood splashed in his/her face
   c) an employee who reasonably anticipates skin, eye, mucous membrane or parenteral contact with human blood or other potentially infectious materials
   d) all of the above

15. If you are stuck by a needle or other sharp item you should:
   a) wash the exposed area, remove gloves, contact supervisor, report to Acute Care
   b) remove gloves, report to Acute Care, wash the exposed area, contact supervisor
   c) remove gloves, wash the exposed area, contact supervisor, report to Acute Care
   d) contact supervisor, report to Acute Care, remove gloves, wash exposed area