CIRCULAR

The Hospital Infection Control Committee (HICC) of Lok Nayak Hospital is in the process to update the hospital infection control manual.

The HIC committee has revised the manual that has to be uploaded on Lok Nayak hospital and Maulana Azad Medical college (MAMC) website for 15 days to obtain the feedback and comments of various stake holders to the infection prevention and control programme. The comments and feedback can be submitted to the HIC committee at hicclnh@gmail.com in prescribed format detailed below:

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Chapter No./Name</th>
<th>Section No. / Paragraph/Line</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
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</tbody>
</table>

Comments can be send to HICC latest by June 30 2016.

Dr. Shagufta Vij
Secretary, HICC
Lok Nayak Hospital

Copy to:
1. Personnel Branch, Medical Director, LN Hospital
2. Guard File.
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<tr>
<th>S.No</th>
<th>Date Of Amendment</th>
<th>Amendment made</th>
<th>Reason of Amendment made</th>
<th>Sign of HICC secretary</th>
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<tbody>
<tr>
<td>1.</td>
<td></td>
<td>To frame antibiotic policy for the hospital has been added in the “purpose”</td>
<td>As a step towards optimizing antimicrobial therapy</td>
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<tr>
<td>2.</td>
<td></td>
<td>“Objectives” of the Infection control program has been revised</td>
<td>To improve the scope of the program as per scope of services provided by the hospital</td>
<td></td>
</tr>
<tr>
<td>3.</td>
<td></td>
<td>Terms of reference of HICC has been added</td>
<td>To define the scope of activities carried out by HICC</td>
<td></td>
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<td>4.</td>
<td></td>
<td>List of committee members revised</td>
<td>For an effective implementation and monitoring of Hospital infection control practices</td>
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<tr>
<td>5.</td>
<td></td>
<td>Incharges of all high risk areas included as committee members</td>
<td>To strengthen the implementation of infection control processes in high risk areas</td>
<td></td>
</tr>
<tr>
<td>6.</td>
<td></td>
<td>Nursing sister incharges have been included as committee members</td>
<td>For an effective implementation and monitoring of Hospital infection control practices</td>
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<tr>
<td>7.</td>
<td></td>
<td>List of notifiable/reportable diseases has been added</td>
<td>For better regulatory compliance</td>
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<tr>
<td>8.</td>
<td></td>
<td>List of alert organisms and alert conditions has been added</td>
<td>To implement effective infection control practices for such organisms and conditions</td>
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<td>9.</td>
<td></td>
<td>Surveillance activities for high risk areas have been updated</td>
<td>To facilitate effective implementation and monitoring.</td>
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<td>10.</td>
<td></td>
<td>Policy of monitoring of efficacy of disinfectants added</td>
<td>To monitor efficacy of disinfection by glutaraldehyde</td>
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<td>11.</td>
<td></td>
<td>Elaborated the surveillance activities.</td>
<td>Further detailing of surveillance activities has been done to facilitate effective implementation and monitoring.</td>
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<td>12.</td>
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<td>CSSD process has been re-defined. Recall policy for CSSD has been added</td>
<td>To make the process more clear</td>
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<td>13.</td>
<td></td>
<td>Infection control procedures and practices has been elaborated</td>
<td>To make the process more clear</td>
<td></td>
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<tr>
<td>14.</td>
<td></td>
<td>Respiratory hygiene and cough etiquettes has been added</td>
<td>To implement effective infection control practices</td>
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<td>15.</td>
<td>Hand hygiene policy elaborated with the inclusion of surveillance of hand hygiene compliance</td>
<td>To make the process more clear and to facilitate effective implementation and monitoring.</td>
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<td>16.</td>
<td>Safe injection practices and safe drug administration has been added</td>
<td>To aware HCW about the safe injection practices.</td>
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<td>17.</td>
<td>Management of sharps disposal, injuries and PEP has been elaborated</td>
<td>To make the process more clear</td>
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<td>18.</td>
<td>Isolation policy and list of diseases which need isolation has been included</td>
<td>To facilitate effective infection control practices to contain transmission of infection within hospital</td>
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<td>19.</td>
<td>Steam sterilization process has been defined and various levels of monitoring of such processes.</td>
<td>To meet sterilization standards as per national and international guidelines/requirements has been added.</td>
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<td>20.</td>
<td>ETO process and monitoring</td>
<td>For improved employee safety and to meet sterilization standards as per national and international guidelines/requirements have been added.</td>
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<td>21.</td>
<td>Method of instrument cleaning and endoscopes reprocessing has been added</td>
<td>To make the sterilization process more effective</td>
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<td>22.</td>
<td>Added new list of disinfectants and their role in several departments</td>
<td>To implement and promote rational and appropriate use of disinfectants</td>
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<td>23.</td>
<td>Chapter on care of systems and indwelling devices has been added</td>
<td>To promote infection prevention practices and to reduce device associated hospital acquired infection</td>
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<td>24.</td>
<td>Added chapter on special care units</td>
<td>To implement good infection control practices in high risk areas</td>
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<tr>
<td>25.</td>
<td>Visitors policy in emergency services</td>
<td>To minimize risk of HCAI among patients, staff and visitors</td>
<td></td>
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<tr>
<td>26.</td>
<td>Chapter on Food safety has been added</td>
<td>To make the process more clear</td>
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<tr>
<td>27.</td>
<td>Elaboration of methods for laundry and linen management</td>
<td>To ensure linen handling at all levels in safe manner</td>
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<td>Description</td>
<td>Purpose/Reason</td>
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<td>28.</td>
<td>Multidose vial policy has been added</td>
<td>To promote safe injection practices</td>
<td></td>
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<tr>
<td>29.</td>
<td>Antimicrobial Stewardship Programme is included</td>
<td>For rational use of antibiotics</td>
<td></td>
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<tr>
<td>30.</td>
<td>BMW policy has been included</td>
<td>As per revised management rules March 2016</td>
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</tr>
<tr>
<td>31.</td>
<td>Case definitions used for diagnosis of HACIs</td>
<td>To make HCW aware about the definitions</td>
<td></td>
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</tr>
<tr>
<td>32.</td>
<td>List of appendices added: HIC indicators, Housekeeping checklist, Daily round checklist, List of disinfectants, Dialysis checklist, NSI form</td>
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</table>
1. Introduction

1.1 Infections which arise in healthcare are termed Healthcare associated infection' (HAI). HAIs are those infections that were neither present nor incubating at the time the patient was admitted to health care facility. The majority of HAI become evident 48 hours or more following admission. However, it may not become clinically evident until after discharge.

1.2 Lok Nayak hospital recognizes the control of hospital associated infections (HAI) as an essential part of patient care. LNH is committed to fulfilling its responsibility by ensuring that proper safeguards are instituted to identify and prevent HAI. All aspects of hospital functions are included in this activity.

1.3 Infection Control includes the prevention and management of HAIs through the application of research based knowledge to practices that include: standard precautions, decontamination, waste management, surveillance and audit.

1.4 The overall aim of this document is to provide evidence based information in the prevention and control of infection at LNH. To fulfill this aim hospital infection control committee has been formed that looks after the infection control needs of the hospital. It is relevant to all staff including doctors, nurses, other clinical professionals and managers working at LNH to help to fulfill their legal and professional obligations with regard to both communicable diseases and infection control.

1.5 This document will be reviewed and updated by the HICC, LNH yearly.
2. Organization of Infection Prevention and Control Program

Lok Nayak Hospital recognizes the prevention and control of hospital associated infections (HAI) as an important issue and is committed to fulfilling its responsibility by ensuring that proper safeguards are instituted to identify and prevent HAI. All aspects of hospital function are included in this activity.

2.1 Purpose:

2.1.1 To establish standards in prevention, control measures and minimize HAIs in patients, staff and visitors.
2.1.2 To define policies and procedures for implementing and monitoring of HAIs at LNH.
2.1.3 To establish antibiotic stewardship program with at least yearly updation of evidence based antibiotic policy with monitoring of its adherence by the prescribing authorities and monitoring antibiotic utilisation in various areas of the hospital.

2.2 Components of the LNH Infection Prevention and Control Program

2.2.1 Establishing and regular updating of hospital infection control manual
2.2.2 Minimizing HAIs through continuous monitoring of healthcare associated infection
2.2.3 Surveillance

2.2.3.1 Laboratory based surveillance of HAIs
2.2.3.2 Ward based surveillance of HAIs
2.2.3.3 Surveillance and regular feedback of device associated infection
2.2.3.4 Surveillance and regular feedback of surgical site infection

2.2.4 Improvement of hand hygiene compliance
2.2.5 Investigation and control of outbreaks
2.2.6 Monitoring of emergence of antimicrobial resistance
2.2.7 To recommend antibiotic policy for the hospital based on local antibiograms and evidence based published national/international guidelines.
2.2.8 Identify areas if irrational use of antibiotics and curb irrational use of antibiotics in hospital areas
2.2.9 Identification of high risk areas and establish steps to mitigate risk of HAIs to patients, staff and visitors
2.2.10 Establish sterilization and disinfection protocols and establish mechanisms to monitor the same.
2.2.11 Monitoring of staff health to prevent, staff to patient and patient to staff spread of infection.
2.2.12 Monitoring and promotion of bio-medical waste management as per government regulations
2.2.13 Training of staff in prevention and control of HAI.

2.3 Objectives and Terms of Reference

2.3.1 Objectives of the program

2.3.1.1 To minimize healthcare associated infections among patients, staff and visitors
2.3.1.2 To establish antimicrobial stewardship program and promote rational use of antimicrobials
2.3.1.3 To provide education and training to healthcare workers, patients and visitors regarding policies and procedures to minimise healthcare associated infections

2.3.2 Terms of reference

2.3.2.1 To develop a documented healthcare associated infections prevention and control program
2.3.2.2 To identify and reduce risks of healthcare associated infections among patient, staff and visitors and implement risk mitigation strategies for the same
2.3.2.3 To meet and monitor all statutory requirement related to healthcare associated infections asked by various government authorities from time to time
2.3.2.4 To perform surveillance activities to capture and monitor infection prevention and control data
2.3.2.5 To take action to prevent and control healthcare associated infections in patient, visitors and healthcare workers
2.3.2.6 To ensure adequate and appropriate resources for prevention and control of healthcare associated infections
2.3.2.7 To identify and take appropriate action to control outbreaks of infection in the hospital
2.3.2.8 To document policies and procedures and sterilization activities and ensure its implementation and monitoring
2.3.2.9 To ensure appropriate and safe handling of Biomedical waste management in hospital premises
2.3.2.10 To plan, support and implement regular training of healthcare regarding infection control and prevention

2.4 Constitution of Hospital Infection Control Committee (HICC)

2.4.1 Members

<table>
<thead>
<tr>
<th>SNO</th>
<th>Designation/Departments</th>
<th>NAME</th>
<th>Committee Organisation</th>
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<tbody>
<tr>
<td>1</td>
<td>Medical Director</td>
<td>DR. D.K. TEMPE</td>
<td>Chairperson</td>
</tr>
<tr>
<td>2</td>
<td>Prof. &amp; HOD Microbiology (MAMC)</td>
<td>DR. C.P. BAVEJA</td>
<td>Vice Chairperson</td>
</tr>
<tr>
<td>3</td>
<td>Asst. Prof. Microbiology</td>
<td>DR. VIKAS MANCHANDA</td>
<td>Infection Control Officer</td>
</tr>
<tr>
<td>4</td>
<td>Addl. M.S NABH</td>
<td>DR. S.D SHARMA</td>
<td>Member</td>
</tr>
<tr>
<td>5</td>
<td>Prof &amp; HOD Neonatology</td>
<td>DR. S.RAMJI</td>
<td>Member</td>
</tr>
<tr>
<td>6</td>
<td>Prof &amp; HOD Surgery</td>
<td>DR. S.K.TUDDU</td>
<td>Member</td>
</tr>
<tr>
<td>7</td>
<td>Prof &amp; HOD GynaObs</td>
<td>DR. SUDHA PRASAD</td>
<td>Member</td>
</tr>
<tr>
<td>8</td>
<td>Prof &amp; HOD Anesthesia</td>
<td>DR. U.C VERMA</td>
<td>Member</td>
</tr>
<tr>
<td>9</td>
<td>Prof &amp; HOD Orthopedics</td>
<td>DR. A.K DHAL</td>
<td>Member</td>
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<tr>
<td>10</td>
<td>Prof &amp; HOD ENT</td>
<td>DR. J.C PASSEY</td>
<td>Member</td>
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<tr>
<td>11</td>
<td>C.C.M.O Causality</td>
<td>DR. RITU SAXENA</td>
<td>Member</td>
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<td>12</td>
<td>M.O I/C CSSD &amp; OT Complex</td>
<td>DR. PAWANINDER Lal</td>
<td>Member</td>
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<tr>
<td>13</td>
<td>I/C ICU</td>
<td>DR. ANIL MISHRA</td>
<td>Member</td>
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<tr>
<td>14</td>
<td>I/C CCU</td>
<td>DR. M.K. DAGA</td>
<td>Member</td>
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<tr>
<td>15</td>
<td>I/C RCU</td>
<td>DR. D.P BATHURIA</td>
<td>Member</td>
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<td>16</td>
<td>I/C PICU</td>
<td>DR. U.JHAMBA</td>
<td>Member</td>
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<td>17</td>
<td>I/C Nursery Referral</td>
<td>DR. N.B. MATHUR</td>
<td>Member</td>
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<td>18</td>
<td>I/C Dialysis Unit</td>
<td>DR. RAJIV KOLI</td>
<td>Member</td>
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<tr>
<td>19</td>
<td>Incharge Nursing Services (D.N.S)</td>
<td>MRS. SANDAL VATS</td>
<td>Member</td>
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<td>D.N.S Emergency Block</td>
<td>MRS. BIMLESH KAIN</td>
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<td>D.N.S Gyna Block</td>
<td>MRS. VEENA SINGH</td>
<td>Member</td>
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<td>22</td>
<td>D.N.S Ortho / OT Block</td>
<td>MRS. RAMPYARI MEHRA</td>
<td>Member</td>
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<td>D.N.S Surgery /OPD Block</td>
<td>MRS. HIRAMANI LUGUN</td>
<td>Member</td>
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<td>24</td>
<td>D.N.S Paeds Block</td>
<td>MRS. PUSPA MINZ</td>
<td>Member</td>
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<td>MRS. CECILIA KUJUR</td>
<td>Member</td>
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<td>26</td>
<td>A.N.S (HICC)</td>
<td>MRS. R.E.LEPCHA</td>
<td>Member</td>
</tr>
<tr>
<td>27</td>
<td>Sister Incharge ICU</td>
<td>MRS. RUPA SINGH</td>
<td>Member</td>
</tr>
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<td>28</td>
<td>Sister Incharge RCU/CCU</td>
<td>MRS. KUNJU MOL</td>
<td>Member</td>
</tr>
<tr>
<td>29</td>
<td>Sister Incharge NICU /Ward 16</td>
<td>MRS. TRIPTA GIRDHAR</td>
<td>Member</td>
</tr>
<tr>
<td>30</td>
<td>Sister Incharge PICU</td>
<td>MRS MARRY BABU</td>
<td>Member</td>
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<tr>
<td>31</td>
<td>Sister Incharge Nursery</td>
<td>MRS. KOHLE KHANNE</td>
<td>Member</td>
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<td>32</td>
<td>Sister Incharge Dialysis</td>
<td>MRS. POONAM</td>
<td>Member</td>
</tr>
<tr>
<td>33</td>
<td>Infection Control Nurses</td>
<td>ALL ICNS</td>
<td>Members</td>
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<tr>
<td>34</td>
<td>S.R Microbiology L.N.H</td>
<td>DR. SHILPI GUPTA</td>
<td>Infection Control SR</td>
</tr>
<tr>
<td>35</td>
<td>Occupational Epidemiologist</td>
<td>DR. SHAGUFTA VIJ</td>
<td>Secretary</td>
</tr>
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Co –Opted Members

<table>
<thead>
<tr>
<th>SNO</th>
<th>Designation/Departments</th>
<th>NAME</th>
<th>Committee Organisation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Addl. M.S Procurement</td>
<td>DR ASHOK KUMAR MITTAL</td>
<td>Member</td>
</tr>
<tr>
<td>2</td>
<td>Addl. M.S Out Sourced Services</td>
<td>DR. SATISH KR.SHARMA</td>
<td>Member</td>
</tr>
<tr>
<td>3</td>
<td>Addl. M.S Sanitation &amp; Bmw</td>
<td>DR. S.S GAMBHIR</td>
<td>Member</td>
</tr>
<tr>
<td>4</td>
<td>Prof &amp; HOD Pediatric Medicine</td>
<td>DR. SANGEETA YADAV</td>
<td>Member</td>
</tr>
<tr>
<td>5</td>
<td>Prof &amp; HOD Medicine</td>
<td>DR. N.GUPTA</td>
<td>Member</td>
</tr>
<tr>
<td>6</td>
<td>Hod Blood Bank/B.T.O</td>
<td>DR. SUNITA MEENA</td>
<td>Member</td>
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<td>7</td>
<td>M.O I/C PWD</td>
<td>DR. DHEERAJ KUMAR</td>
<td>Member</td>
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<td>8</td>
<td>M.O I/C Laundry /Linen Stores / Gen Store</td>
<td>DR. VINITA JAISWAL</td>
<td>Member</td>
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<td>9</td>
<td>D M S OPD</td>
<td>DR. VIKAS RAMPAL</td>
<td>Member</td>
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<td>10</td>
<td>M.O I/C Stationary Store/Kitchen</td>
<td>DR. SANGEETA BHASIN</td>
<td>Member</td>
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<td>11</td>
<td>M.O I/C (MRD) Nodal Officer Vector Borne Disease</td>
<td>DR. SUDHA RANI</td>
<td>Member</td>
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</tbody>
</table>

2.4.2 Meetings of HICC
2.4.2.1 The infection control team meets at least quarterly or more frequently as necessary. Documentation of meetings and recommendations are kept by the secretary.

2.4.2.2 Minimum Quorum required: Chairperson, Infection Control Team [ICO and ICNs (atleast 5)] and 50% of other members

2.4.2.3 The ICN (Infection Control Nurse) conduct rounds and report the findings to the ICO/Senior Resident infection control on daily basis. Registers are maintained by ICNs.

2.4.4 Hospital Infection control team (ICT)

The infection control team consists of:
1. Microbiologist (Infection control officer)
2. Senior Resident (Infection Control)
3. Infection Control Nurses

2.4.5 Responsibilities of the Infection Control Team

2.4.5.1 Advise staff on all aspects of infection control and maintain a safe environment for patients and staff.

2.4.5.2 Advise management of at risk patients.

2.4.5.3 Carry out targeted surveillance of healthcare associated infections and act upon data obtained e.g. investigates clusters of infection above expected levels.

2.4.5.4 Recommend antibiotic policy for different areas of the hospital.

2.4.5.5 Provide a manual of policies and procedures for aseptic, isolation and antiseptic techniques.

2.4.5.6 Investigate outbreaks of infection and take corrective measures.

2.4.5.7 Provide relevant information on infection problems to management.

2.4.5.8 Assist in training of all new employees as to the importance of infection control and the relevant policies and procedures.

2.4.5.9 Have written procedures for maintenance of cleanliness.

2.4.5.10 Surveillance of infection, data analyses, and implementation of corrective steps. This is based on reviews of lab reports, reports from nursing in charge etc.

2.4.5.11 Surveillance of Biomedical Waste management

2.4.5.12 Supervision of isolation procedures

2.4.5.13 Monitors employee health programme.

2.4.5.14 Addresses all requirements of infection control and employee health as specified by Central laws, State laws and NABH.

2.4.6 Infection Control Nurse (ICN)

The duties of the ICN are primarily associated with ensuring the practice of infection control measures by healthcare workers. Thus the ICN is the link between the HICC and the wards/ICUs etc. in identifying problems and implementing solutions.

2.4.6.1 Duties of infection Control Nurse includes:

2.4.6.1.1 The ICN conducts Infection control rounds daily and maintains the registers.

2.4.6.1.2 The ICN is involved in education of practices minimising healthcare associated infections and hand hygiene among Healthcare workers.

2.4.6.1.3 Maintains registers and data of Sharps/Needle stick injuries and Post–exposure prophylaxis.

2.4.6.1.4 Initiates and ensure proper immunization for Hepatitis B Virus Immunoglobulin and HBsAg vaccine, in consultation with microbiologist (Member HICC) in case of suspected exposure to any hospital worker.

2.4.6.1.5 Ensures that all positive culture cases are been tracked and for each positive culture from inpatient unit a hospital infection information sheet or surgical site infection Sheet is filled and keep record for each positive culture case. All probable cases of healthcare associated infections and anomalous/irrational use of antibiotics must be discussed in HICC meetings.

2.4.6.1.6 Track the indicators of infection control and present the data to the HICC meetings on regular basis.

2.4.6.1.7 Conducts special tasks given to him/her as per components and objectives of the hospital infection prevention and control.

Selection of ICNs

ICNs can be selected through following process:-

1. Any staff nurse can volunteer to enrol for ICN provisionally
2. Nominated nursing staff can be enrolled for ICN provisionally
3. All staff nurses have to undergo written exam provided by surveillance and infection control division. Only staff nurses scoring more than 90% of marks shall be enrolled as
ICN. Till the time provisional ICNs qualify the exam, they can work under supervision of qualified ICN. Provisional ICN shall not work independently or take independent rounds.

2.4.7 Infection Control Officer (ICO):
The microbiologist serves as Infection Control Officer.

2.4.7.1 Duties of Infection Control Officer:
2.4.7.1.1 The ICO supervises the surveillance of healthcare associated infections as well as preventive and control programs.
2.4.7.1.2 Co-ordinate with the chairperson and HICC in planning infection control programme and measures.
2.4.7.1.3 Keeps a track of any developing outbreaks.
2.4.7.1.4 Participate, guides in research activities related to infection control practices and publish them.
2.4.7.1.5 Developing guidelines for appropriate collection, transport & handling of specimens.
2.4.7.1.6 Ensuring laboratory practices meet appropriate standards.
2.4.7.1.7 Ensuring safe laboratory practices to prevent infection in staff.
2.4.7.1.8 Performing antimicrobial susceptibly testing following internationally recognized method & providing summary reports of prevalence of resistance.
2.4.7.1.9 Monitoring sterilization, disinfection & the environment where necessary.

2.5 Review and Revision of Infection Control Manual
2.5.1 Written policies and procedures shall be reviewed at least in a year. Signature of chairperson HICC, Secretary HICC, ANS and Infection Control Nurses shall be affixed on controlled copies. There shall be atleast five controlled copies that shall be distributed to the following: MS, ICO, ICN, NS and Hospital Library. All department shall have atleast one copy of the manual. Digital version shall be available through hospital website to all.
3. SURVEILLANCE AND REPORTING OF HOSPITAL ACQUIRED INFECTIONS.

3.1 Statutory Notifications
Infectious diseases, which are listed in section 3.1.2 whether confirmed or suspected, must be notified by the attending doctor to the Consultant for Communicable Disease Control (CCDC) who is MO/Ic MRD.

3.1.1 Prompt notification and reporting of disease is essential. The objectives of notification are:
   a. Regulatory obligation by Govt. of NCT of Delhi
   b. To collect accurate and complete epidemiological information on the disease.
   c. To ensure prompt and appropriate control measures to prevent the spread of infection.
Any doctor who considers that a patient is suffering from a notifiable/reportable disease/ has a statutory duty to notify the Nodal officers (Medicine, Paediatrics and Dermatology).
   - Nodal Officers should provide weekly data to ICN.
   - ICN should monitor Infection control practices in wards for these diseases and should provide feedback to ICO, CCDC and nodal officer.

3.1.2 Notifiable/Reportable Diseases
- Measles
- Cholera
- Smallpox
- Plague
- Diphtheria
- Dengue hemorrhagic fever/ Dengue
- Acute flaccid paralysis
- Swine flu
- Malaria
- HIV/AIDS

In case of an epidemic:
- Acute gastroenteritis
- Viral hepatitis
- Meningococcemia

3.2 Healthcare associated Infection Surveillance
Surveillance of health care associated infections means recording and counting of infections arising in the hospital. It is done so that we know the extent of any problems that exist.

3.2.1 Aims
The main objectives of surveillance of hospital acquired infections are:

Objective of Surveillance
- Establish endemic baseline rate.
- Reducing infection rates in the hospital.
- Identifying and containing the outbreaks.
- Evaluating and monitoring infection control measures.
- Monitoring antimicrobial susceptibility patterns

Surveillance is part of the routine infection control programme. It helps to identify risks of infection and reinforces the need for good practices. Preventing outbreaks depends on prompt recognition of one or more infections with alert organisms and instituting special control measures to reduce the risk of spread of the organism. Collection of accurate data allows comparison with other units and measurement of response to changes in practice. All patients which are diagnosed with HAI are followed up till separation (discharge, death, LAMA, Abscond) for monitoring of ALOS, outcome of HAI. All bed side X-Rays of IPD are monitored on daily basis to detect hospital acquired pneumonia. Efforts will be made to contact all patient undergoing surgery at LNH. Telephonic follow up till the 90 days of surgery (if implant placed up to one year) are done to detect possible SSI.

Policy describes following key points
1. Passive methods of surveillance
   a. Methods
   b. Action plan
c. Response statement
2. Active methods of surveillance

3.2.2 Passive Surveillance
Passive surveillance shall be done laboratory based ward surveillance in conjunction with “Alert organism/Alert condition” surveillance. The system is managed by the Infection Control Team and details are reported back to the Infection Control Committee.

3.2.2.1 Laboratory-Based Ward Liaison Surveillance (Alert Organisms).
All positive microbiology reports from inpatient will be screened and may result in a case review, a search for other carriers or infected patients and ward visits by the Infection Control Nurse. Approximately 70% of infections and alert organisms can be detected in this way. A patient may be placed in source isolation if considered to be a source of infection to other patients.

3.2.2.2 Ward Based Surveillance (Alert Conditions)
Alert conditions are medical syndromes such as Acinetobacter bacteraemia or Pseudomonas pneumonia which immediately suspected healthcare associated infection. It is the responsibility of the ward staff to notify the infection control team if they suspect an infection which may be a risk to others. Appropriate specimens must be taken and sent promptly, properly labelled, to the laboratory. Source isolation precautions must be instituted immediately that infection is suspected.

3.2.2.3 Action Plan
When organism/s is/are detected by the laboratory based surveillance or ward based surveillance, microbiologist and the treating clinician will discuss the possibility of healthcare associated infections and action will be recorded in Hospital acquired infection assessment form. Every effort will be made to evaluate critically each and every positive culture report from the in-patient units including critical care areas. The record will be maintained by ICN and the data will be presented at least once a month at HICC meeting to review the case critically for possible HAI infections and the feedback will be provided to the concerned unit head.

3.2.2.4 Response
Appropriate measures will be taken in case of suspected outbreak or sudden increase in rates of suspected healthcare associated infections. Control measures to prevent spread of infection and decrease the incidence of healthcare associated infections may be suggested in feedback report to the concerned units. The report will be prepared at least biannually and will be submitted to the unit heads. In case urgent intervention is required the response may be communicated more frequently.

 Clinicians must tell the Infection Control Team about any Alert Condition/s.

List of ALERT ORGANISMS (suggested list but NOT limited to)

**BACTERIA**
1. Methicillin-resistant *Staphylococcus aureus*
2. Other resistant *Staphylococcus aureus*
3. Penicillin-resistant *Streptococcus pneumoniae*
4. *Haemophilus influenzae*
6. Glycopeptide-resistant enterococci
7. *Neisseria meningitidis*.
8. Pan-resistant Gram negative bacilli
9. *Mycobacterium tuberculosis*
10. Any unusual bacteria

**VIRUSES**
1. Hepatitis B
2. Hepatitis C
3. HIV
4. Rotavirus
5. Small round structured virus (Norovirus)
6. Respiratory syncytial virus
7. Varicella zoster
8. Influenza virus
9. Parvovirus
10. Measles
11. Novel H1N1
12. Dengue

Examples of ALERT CONDITIONS
1. Post-surgical sepsis
2. Exanthema (acute rash illness)
3. Chicken pox or shingles
4. Mumps, measles, rubella, parvovirus
5. Whooping cough
6. Poliomyelitis
7. Diphtheria
8. Meningococcal Meningitis
9. Hepatitis B and Hepatitis C Viral Infection
10. Pyrexia of unknown origin
11. Typhoid and paratyphoid fevers
12. Viral haemorrhagic fever
13. Swine flu

3.2.3 TARGETED SURVEILLANCE
Detailed targeted surveillance in specific areas is performed. An example would be surgical site infection (SSI) surveillance. Results are feedback to HICC.

3.2.4 Active Surveillance
2.3.4.1 Active surveillance of HAI
ICN collects positive culture reports from the microbiology. The ICN in consultation with ICO may proceed for investigation of HAI.

2.3.4.2 Active surveillance of High Risk Areas and other areas of significance
High risk areas of the hospital are identified includes:

High risk areas of hospital include:
- Intensive care units (NICU, PICU, CCU/RCU, MSICU)
- Operation theatres (OT I, OT II, OT III, Urology OT, Burns and Plastic OT, Gynae OT, PPOT, Septic OT
  - HDU
  - Dialysis unit
  - Kitchen
  - CSSD
  - Blood bank
  - Drinking water facilities

I. Operation Theatres
As per guidelines for Environmental Infection Control in health care facilities recommended by the Centers for Disease Control (CDC) and the Healthcare Infection Control Practices Advisory Committee (HICPAC), 2003, Microbial Sampling of Air and inanimate surfaces (i.e. Environmental Sampling including surface swabs) is not recommended.
The air quality testing shall be done only under following conditions:-
- To support an investigation of an outbreak of disease or infections.
- For the purpose of research.
- After any major construction periods to qualitatively detect breaks in environmental infection control measures.
Surprise air checks can be undertaken to monitor general OT discipline at least once in a month.
Fogging of OTs will be done on the basis of these reports and/or clinical out of procedures carried out in the operating areas.
Records are kept with nursing incharge OT and the results must be produced in HICC meetings biannually or more frequently. In case of unacceptable results decision on corrective measures are taken by HICC.

Monitoring of disinfectants (Glutaraldehyde 2%)
The efficacy of the Glutaraldehyde shall be tested by surprise check at least once in a month and records are to be kept with ICN. The data shall be presented in HICC meeting at least once in 3 months.

II. Intensive care units
Surveillance samples to be taken when there is suspected outbreak or same isolate irrespective of their antibiotic sensitivity are isolated from 3 or more patients.

Surveillance samples include but not limited to:

- **Clinical Material**
  - Central line tips
  - ET tube secretions
  - Urine samples from catheterized patients
  - Nasogastric tube materials

- **Environmental Sampling**
  - Water samples from humidifiers
  - Sampling of drugs prepared for patients
  - Ventilators
  - Walls
  - Floors
  - Suction tubing
  - Disinfectants on dressing trolleys

Surveillance clinical samples are sent per patient on basis of clinical data or microbiological reports. Any positive sample will be analyzed critically to detect healthcare associated infections. The data will be maintained by ICN and presented in subsequent HICC meeting.

Colonization swabs (nasal and rectal) will be collected from time to time to monitor anti-microbial resistance and multi drug resistant organism.

**III. Transfusion services unit**

The blood samples from bags must be sent for culture periodically. Blood component FFP and platelets shall be screened for contamination, as and when required. The record will be maintained by blood bank officer and chairman/Secretary HICC must be updated about the data atleast once in a month and presented in HICC meetings.

**IV. Food handlers**

Screening of food handlers is done biannually. Samples include nasal swabs, urine and stool samples. Records to be maintained by the dietician and ICN.

**V. Drinking Water**

Bacteriological surveillance is to be done at least once in 2 months in the microbiology laboratory for live bacterial contamination and once in six months in an accredited laboratory. Responsibility of sending the samples and records maintenance is of Dietary department.

**VI. CSSD**

Cleaning protocols of CSSD:

Environmental surveillance is done monthly basis to check the Air quality of the sterile zone.
- Floor is mopped daily with soap and water.
- Fogging of sterile storage room may be done based on air surveillance reports or as per needs.
- Trolleys, shelves and tables are wiped with disinfectant every day.

**Structure:**

The different Standard Operating Procedures in the CSSD are followed. CSSD has been divided into 3 zones. There should not be cris-crossing of processes within CSSD. The three zones are:

1. **Protective zone**
2. **Clean zone**
3. **Sterile zone**

**1. Protective Zone includes:**
   i) Receiving Window
   ii) Cleaning Area
   iii) Decontamination Area
   iv) Drying Area
   v) Assembling and Packaging Area

**2. Clean Zone includes:**
   i) Autoclaving Area

**3. Sterile Zone includes:**
i) Sterile storage room
ii) Issuing window.

1. Protective Zone

i. Receiving Area: Items are brought to CSSD from respective wards, ICU’s, O.T.’s & casualty by nursing orderly. The CSSD assistant receives them & checks the status of items.

ii. Cleaning Area: In this area all instruments are primarily cleaned and rinsed with plain water to remove visible particles.

iii. Decontamination Area: In this area soiled instruments including heat sensitive items like oxytubings, nebulisation chamber, airway etc. and other supplies are decontaminated with the help of glutaraldehyde 2%, enzyme solution(s) 1%.

iv. Drying Area: In this area all cleaned items are dried with the help of drying cabinet at a temperature of 45° C for 45 minutes.

v. Assembling & Packaging Area:
Here all the instruments are assembled and packed for sterilization after cleaning & drying. Labels and autoclave indicator tapes are pasted on all the packs.

vi. Packing Area: In this section Various types of dressings like gauge pieces; cotton pads and bandages etc. are also prepared in this area.

2. Clean Zone

i. Autoclaving Area: In this area sterilization process is carried out by autoclaves. Before that autoclave indicators are pasted on the packs. Then technician places the packs in the autoclave machine and starts the machine as per cycle of appropriate temperature and pressure recommended by the manufacturer for 30 minutes.

3. Sterile Zone

i. Sterile storage Area: In this area sterile items are placed in racks after completion of autoclave before that adequacy of sterilization is confirmed by indicators.

ii. Issuing Window: All the sterile instruments and other supplies are distributed to concerned departments at a separate window after entry of all the items in the appropriate issuing register.

Quality Indicators ( Before use & after use).

Monitoring protocol of Autoclave:

1. Temperature, Pressure and time of each cycle is recorded is followed according to manufacturer’s recommendations. Records should be maintained

2. Various quality indicators are used to check the efficacy of sterilization:

a) Exposure control: Autoclave indicators tape is pasted on all packs to be kept in autoclave.

b) Load Control: Biological indicators are used once a week (Monday) in all autoclave machines in first load and with every load which contain any implant. This indicator gives us rapid results, i.e. positive result in one hour and negative result in 3 hours. If result is positive means sterilization is not adequate that whole load is recalled & re-autoclaved.

c) Pack control: Class 5 chemical indicator - It is used in every pack.

d) Equipment control: Bowie-dick test pack – It is used once daily in each machine.

3. Air cultures are taken once in a month

4. Wet pack is not accepted as sterile. These are repacked and resterilized (even if the indicators show the appropriate changes.

5. There are different trolleys for carrying sterile and unsterile instruments White & Red respectively.

6. No person is allowed to enter in sterile room without Personal Protective Equipments (PPE) (i.e. Cap, mask, gown, & slippers etc.)

7. All sterile items must be used within 72 hours after 72 hours items should send to CSSD for re-autoclaving

Recall policy:

Actions to be taken if any monitoring indicators fail:

a) Recall the item immediately with the help of load number

b) CSSD supervisor are informed immediately.

c) CSSD personnel should try and discover the cause of the failure and arrange for corrective action.

d) The item are reprocessed and then supplied after confirmation of sterility.

Record keeping:
a) Entry of all the items made in CSSD receipt register including date, time, type of instruments in the pack, name of department, procedure used for, case infected not, name and signature of person receiving the items.
b) Inventory of sterile packs is checked so that they are not distributed directly to the user department.
c) Record of all the indicators tests and culture report is kept.
d) Result of load control, equipment control and glutaraldehyde solution monitoring results are submit to the HIC department on monthly basis.
e) Recall event should be documented and record should be maintained in a register.

3.2.5 Special Studies
Special studies will be conducted as needed. These may include:
The investigation of clusters of infections above expected levels.
   a. The investigation of single cases of unusual or epidemiologically significant HA infections.
   b. Prevalence and incidence studies, collection of routine or special data as needed and sampling of personnel or the environment as needed.

3.2.5 Surveillance of Hand Hygiene Compliance
3.2.5.1 Direct observations can be made by any of the infection control team members. This can usually be accomplished well through regular observations, especially at odd hours.
3.2.5.2 Data for all categories of staff should be gathered including faculty, residents, nursing, ward boys and other health care workers involved in direct patient care.
3.2.5.3 This should be followed by awareness drives and educational activities.
3.2.5.4 Provision of accessible alcoholic rubs should preferably be made at each bedside.
3.2.5.5 Data generated should be presented in HICC meeting regularly.
4. INFECTION CONTROL PROCEDURES AND PRACTICES

Since it is impossible to identify some infectious patients (especially those infected with HIV, Hepatitis B or C) a system of standard precautions MUST be adopted in all health care work.

4.1 According to HICPAC and the CDC, “Standard Precautions combine the major features of Universal Precautions and Body Substance Isolation and are based on the principle that all blood, body fluids, secretions, excretions except sweat, non-intact skin, and mucous membranes may contain transmissible infectious agents.” Standard Precautions are a group of infection prevention practices that apply to all patients and residents, regardless of suspected or confirmed infection status, in any setting in which healthcare is delivered and include:

1) Hand hygiene
2) Use of personal protective equipment (e.g., gloves, gowns, facemasks), depending on the anticipated exposure
3) Respiratory hygiene and cough etiquette
4) Management of spillage
5) Safe injection practices

4.1.1 Hand Hygiene

4.1.1.1 Purpose
Hand washing is THE SINGLE most important measure in reducing the spread of infection. Hands are the principle route of cross infection. The level of hand hygiene will be determined by the activity or area of practice.

4.1.1.2 Scope
All procedures that require hand hygiene should be done through appropriate hand hygiene.

4.1.1.3 Responsibilities
All hospital staff including Nurses, Doctors, O.T. Technicians, Lab Technicians, Nursing orderlies, food handlers and housekeeping staff.

4.1.1.4 When to wash hands
This is determined by actions – those completed and those about to be performed. A list is given below.

4.1.1.5 Routine washing (Social Hand Wash)

- Before preparing, eating, drinking or handling food.
- Before and after smoking.
- After visiting the toilet.
- Before starting work (remove jewellery, e.g. rings) and after leaving an occupational area.
- Before and after physical contact with each client in clinical situations, eg bathing, assisting to move, toileting.
- After handling contaminated items such as dressings, bedpans, urinals, urine drainage bags and nappies.
- Before putting on gloves and after removing them.
- Before and after removing any protective clothing.
- After blowing your nose, covering a sneeze.
- Whenever hands become visibly soiled.
- When hands are visibly soiled,
- Before starting work,
- Handling food and following patient contact.
4.1.1.6 The “My 5 Moments for Hand Hygiene” approach

![Diagram of hand hygiene moments]

1. Before touching a patient
2. Before clean/ aseptic procedure
3. After body fluid exposure risk
4. After touching a patient
5. After touching patient surroundings

4.1.1.7 Sequence of events
- Wet hands under running water.
- Dispense one dose of soap into cupped hand.
- Hand wash for 40-60 seconds vigorously and thoroughly by following six step techniques, without adding more water. (See Figure 1)
- Rinse hands thoroughly under running water.
- Dry hands with single use brown paper.

4.1.1.8 Hand disinfection - Aseptic/hygiene hand wash
Hand disinfection with alcohol based hand rub (eg sterilium, 70% alcohol) preferably with chlorhexidine and alcohol are practice at least in following condition:
- Whenever touching any patient esp. in inpatient units and critical care areas.
- After handling any potentially infectious object
- Before putting on gloves and after removing them.
- Prior to invasive procedures
- Visibly clean hands
- In high dependency areas and after attending patients in isolation or with known transmissible condition.

Broken skin, cuts and abrasions in any area of exposed skin, particularly the hands and forearms, are covered with a waterproof dressing. Wear gloves if hands are extensively affected. Wrist watches/bracelets are removed.
Alcohol is an effective decontamination agent but should only be used on visibly clean hands. It is also a valuable agent for use, but should only be used 2-3 times consecutively before a hand wash as build up can occur.
- Dispense the required amount of solution onto the hands.
- Ensure solution covers all hand surfaces.
- Rub vigorously, using hand washing technique, until dry.
It is recommended that everyone involved in providing healthcare in the community must be trained in hand decontamination, the use of protective clothing and safe disposal of sharps, and this includes patients and healthcare personnel.

4.1.1.9 Hand Care
- Keep nails clean and short.
- Remove rings with stones or ridges.
- Do not wear artificial or gel nails or nail polish.
- When washing hands, wrist watches are removed.
- Sleeves are rolled up to the elbow.
- Nailbrushes should not be used for routine hand washing as they damage the skin and encourage shedding of cells.
- Nailbrushes, where used, must be single use disposable or single use autoclaveable.

Gloves are worn before:
- Before inserting a central intravascular catheter.
- Before inserting indwelling urinary catheters, peripheral vascular catheters, or other invasive devices that do not require a surgical procedure.
- For cleaning up any spillage of body fluids.

The physical action of washing and rinsing hands under such circumstances is recommended because alcohols, chlorhexidine, iodophors, and other antiseptic agents have poor activity against spores.

4.1.1.10 Hand-hygiene Technique
When decontaminating hands with an alcohol-based hand rub
- Apply product to palm of one hand and rub hands together,
- Cover all surfaces of hands and fingers by six step technique, until hands are dry.
- Follow the manufacturer’s recommendations regarding the volume of product to use.

When washing hands with soap and water
- Wet hands first with water
- Apply an amount of product recommended by the manufacturer to hands
- Rub hands together vigorously for atleast 40-60 seconds
- Cover all surfaces of the hands and fingers by following six step technique
- Rinse hands with water and,
- Dry thoroughly with a disposable towel/ Paper
- Use sterile paper towel to turn off the faucet or elbow taps if available.

Surgical hand antisepsis
- Remove rings, watches, and bracelets before beginning the surgical hand scrub
- Finger nails are kept short and well maintained.
- Hands and forearms must be free of open lesions and breaks in skin integrity.
- Wear complete operating room attire including mask, cap, and goggles if required.
- Keep clothing away from sink and splashes
- Turn on water.
- Keep arms level well away from body and hands up above elbows for duration of scrub.
- Wet hands and forearms
- Apply antiseptic hand wash solutions
- Lather hands and forearms for at least one minute from fingertips to three inches above elbows starting with hands to forearm, forearm to elbow.
- Wash hands thoroughly, using the following six steps to facilitate eradication of all bacteria and 10 seconds/step.

1. Palm to palm
2. Palm over dorsum
3. Palm to palm, fingers interlaced
4. Back to fingers to opposing palms
5. Rotate thumbs in palm
6. Rotate fingers in palm
7. Rinse
Steps of hand washing

How to Handwash?

WASH HANDS WHEN VISIBLE SOILED! OTHERWISE, USE HANDRUB

Duration of the entire procedure: 40-60 seconds

1. Apply enough soap to cover all hand surfaces.
2. Rub hands palm to palm.
3. Lather hands and forearms for at least two minutes in the same manner.
4. Recommended scrub time is between 2-6 minutes, longer times are not necessary.
5. Rinse hands and forearms under running water.
6. Keep hands higher than the elbow at all times.
7. Thoroughly dry hands and forearms with a sterile paper towel keeping hands raised.
8. Proceed to OT keeping hands above the elbow and out from scrub clothes. Allow hands and forearms to dry thoroughly before donning sterile gloves.
9. Between short cases only, hands may be disinfected by using 2 or more applications of alcohol handrub.
10. Before applying the alcohol solution, prewash hands & forearms with a non-antiseptic soap & dry hands & forearms completely.

All clinical areas including consultation chambers, each floor & critical care areas should have:
- Hand washing facilities appropriate to the area.
- Clear unobstructed access to the hand washing sink
- Hand washing sinks for that purpose only and clear of inappropriate items.
- Liquid soap and alcohol hand rubs available at every sink.
- Hand washing posters are placed by each sink.
4.1.1.11 Hand Hygiene Audit
- To ensure that the hand washing protocols are followed in the LNH Hospital.
- A monthly report is generated and analyzed and corrective actions taken by training.
- The audits are done in the prescribed format.

4.1.1.12 Patient Hand Hygiene
Hand hygiene for patients must be encouraged as it is equally as important in the prevention and control of infection. Staff must ensure that patients are afforded an opportunity to hand wash prior to meals, after having used a bedpan/urinal or toilet or when hands are otherwise soiled.

4.1.1.13 Quality Assurance
- Completion of mandatory training on Hand Hygiene by all Healthcare Doctors, paramedical, housekeeping and Nurses.
- Monitor and record adherence to hand hygiene.
- Provide feedback to healthcare workers about their performance.

4.1.2 PERSONAL PROTECTIVE EQUIPMENT (PPE)
In determining the type of personal protective equipment to use for a given procedure, HCWs should consider the following factors:
- Probability of exposure to blood and body substances;
- Amount likely to be encountered;
- Type of body substance involved; and
- Probable route of transmission of infectious agents
Full protective wear, including double gloves, protective eye/face-shields, protective footwear and impermeable gowns or aprons, is recommended for operating room or mortuary procedures.

4.1.2.1 Risk assessment
The risk assessment should take account of various factors that include:
- Nature of the task to be undertaken.
- Risk of contamination to either patient or user.
- Barrier efficacy of gloves, both surgical and examination gloves can fail.
- Sterile or non-sterile gloves required.
- Patient/user sensitization.

A. Gloves
The use of disposable gloves is part of the Standard Precautions concept, which offers consistent guidelines for infection control programmes. As part of personal protective equipment, gloves prevent contact with blood, body fluids, and mucous membranes. They also protect the patient from contamination by the micro-organisms from the wearer’s hands; gloves are single use items and are changed after each procedure to further minimize the risk of infection.

Gloves are worn when dealing with:
Any blood or other body fluids, such as synovial fluid, peritoneal fluid, amniotic fluid, pleural fluid.
- Any wound or broken skin.
- Handling chemicals or disinfectants, which could cause skin irritation
As a general rule, if the risk is to the patient then ‘Sterile’ gloves are required. If the risk is to the user then ‘Non-sterile’ gloves will probably be sufficient. When handling chemical disinfectants you may need to wear industrial or domestic gloves.

B. Gowns and aprons
The purpose of wearing gowns and aprons is to protect susceptible patients from infection and protect the wearer from contamination as well as maintaining the uniform or clothes worn under the apron in a clean and dry state.
Gowns and aprons should not be worn outside the area they are intended to be used. Remove your gowns/aprons when moving out of area they are intended to be used.

C. Face Protection
Protective eye or face wear are considered where risk of blood or other bodily fluids splashing into eyes is a possibility, including the preparation of some cytotoxic chemotherapy and during the physical decontamination or cleaning of instruments.

D. Masks
There is no clear guidance available for the efficacy of masks in the prevention of airborne infections. However, they may offer protection against potential splashing of the mouth and face during certain procedures such as minor operations, physical decontamination or cleaning instruments with a brush. The type of mask best suited to a particular situation depends on the body substances likely to be encountered and the nature of the activity.

There are two main types of masks used in health care:
• **Surgical masks** — fluid-repellent paper filter masks worn during surgical and dental procedures
• **Particulate filter personal respiratory protection devices (P2 respiratory protection devices)** — close fitting masks capable of filtering 0.3-µm particles and worn when attending patients with active pulmonary tuberculosis

Mask must:
- be fitted and worn according to the manufacturer’s instructions;
- Not be touched by hand while being worn;
- cover both mouth and nose while worn;
- be removed as soon as practicable after they become moist or visibly soiled;
- be removed by touching the strings and loops only; and not be worn loosely around the neck, but be removed and discarded as soon as practicable after use.

E. Shoe cover
Shoe cover must be worn before entering to the ICU, Operating Theatre, Dialysis, CSSD and HDU.

F. Protective foot wear:
Protective foot wear should be used when handling biomedical waste as unnoticed cuts and wounds are quite common in the legs. Footwear is also essential to protect legs from ‘sharps’ injury.

G. Head cap:
Head cap covers the hairs of the health care provider in order to prevent the contamination of the sterile high risk areas.

4.1.3 RESPIRATORY HYGIENE AND COUGH ETIQUETTE
4.1.3.1 The strategy is targeted at the patients and accompanying family members and friends with undiagnosed transmissible respiratory infections, and applies to any persons with signs of illness including cough, congestion, rhinorrhea, or increased production of respiratory secretions when entering a health care facilities.

4.1.3.2 The elements of respiratory hygiene/ cough etiquette include
1. Education of healthcare facility staff, patients and visitors
2. Source control measures (Covering the mouth/nose with tissue or a cloth when coughing or sneezing)
3. Hand hygiene after contact with respiratory secretions
4. Spatial separation, ideally >3 feet, of persons with respiratory infections in common waiting areas when possible
5. Masks should be provided to the coughing patients to contain dispersion of respiratory secretions into the air from infected patients
6. Healthcare personnel are advised to observe Droplet precautions (i.e. wear a mask) and hand hygiene when examining and caring for patients with signs and symptoms of a respiratory infections.
7. Healthcare personnel who have a respiratory infection are advised to avoid direct patient contact, especially with high risk patients. If this is not possible then a mask should be worn before providing patient care.

4.1.4 MANAGEMENT OF SPILLAGE
It is vital that any spillage must be attended to as soon as possible. Assessment of hazards and associated risks to health must be undertaken to ensure the health and safety of employees, patients and other visitors to the primary health care premises.

4.1.4.1 Responsibilities
Department Heads are responsible for the development and implementation of a policy that deals with spillages, and should exposure occur, they are also required to ensure that any risks to staff, patients and visitors are minimized. All staff has the responsibility for ensuring that they adhere to any policies and procedures to minimize the hazards resulting from any spillage. All staff involved in the clinical care of patients or the safe handling of waste are aware of how to deal safely with any spillage should it occur.

“ALL SPILLS LARGE (>30ml) OR SMALL(<30ml) MUST BE REPORTED TO HOSPITAL INFECTION CONTROL NURSE (ICN) IMMEDIATELY”

4.1.4.2 Body Fluid Spillage
Body fluid spills are divided into two categories, those which are visibly contaminated with blood and those which are not.

Blood Spillage or other Body Fluid visibly contaminated with Blood.
- Spillages of blood are dealt with as soon as possible.
- Splashes of blood (or any body fluid) on the skin are washed off immediately with soap and water.
- If there is broken glass do not touch even with gloved hands- use a paper or plastic scoop and dispose in the sharps box.

4.1.4.3 Management blood spillage.
1. Make the people aware about spill
2. Cordon off the area.
3. Identify the spill kit.
4. Wear PPE.
5. Put soaking paper (brown paper, newspaper and tissue paper) over the spill.
6. Make fresh bleaching solution by using 0.75gm of bleaching powder in 100ml water) which equivalent to 0.5 to 1% strength.
7. Pour this prepared solution over the recovered spill.
8. Leave for contact time ideally 20 minutes but if the area where the spill is occurred is a very busy area then minimum 2-5 minutes.
9. After contact time put another paper covering the soaked paper and then remove the soaked paper and put it in the RED bag.
10. Discard this red bag in main red bin the unit.
11. Clean the area with soap and water.
12. Remove the PPE & discard it in the red bag.
13. Do the hand washing.

4.1.4.5 Large Spill
In case of large spill Inform to HIC dept. or ICN
Immediate action has to be taken with the help of large spill kit available at the concern department. The recommended practice is:

Procedure to manage large spill
1. Cordon off the area and make the people aware about the spill.
2. Put on the PPE.
3. If there is any sharp material present along with the spill, then first remove it with the help of plastic scoop or with x-ray film.
4. Put large size gauze pad over the spill to soak large amount of spill and discard the pads in red bag.
5. Put soaking paper over the rest amount of spill.
6. Make fresh bleaching solution by using 7.5gm bleaching powder in 1 Ltr. of water
7. Put this bleaching solution over the spill and wait for contact time (20 min)
8. Take another paper and with the help of this paper, remove the paper which is already put on the spill.
9. Discard all the papers in red bag.
10. Wash the area with soap and water.
11. Remove the PPE and discard in red bag.
12. Do hand washing.
13. Fill the incident reporting form and send it to the HIC department.
14. It is the responsibility of person who had done the spill to manage it. For anonymous spills nursing staff posted in the area shall be responsible to get it managed. Ultimate responsibility of implementation of the policy lies with Nursing Incharge of the area where spill has occurred.
4.1.4.6 Role of ICN in the large spill management.
- To ensure proper spill management
- Ensure incident reporting form is filled with proper details.
- Root cause analysis of incident and ensure that preventive action is taken.

4.1.4.7 Urine Spills visibly contaminated with Blood
Chlorine releasing agents are **not** be used for urine spillages even if it contains visible blood. If a chlorine releasing agent is used with urine the resulting fumes are considered a hazard. The recommended practice is:
- Wearing non-sterile, non-powdered latex gloves and plastic apron.
- Soak up with paper.
- Use detergent and water on area after soaking up the spill.
- A chlorine-releasing agent may now be used on the area if necessary.
- Discard gloves, waste materials and apron in a Reg bag.
- Wash hands thoroughly

4.1.4.8 Spillages of Body Fluids not visibly contaminated with Blood
These spillages will include faeces, vomit, urine and sputum.
- Always wear protective clothing, i.e. plastic disposable apron, disposable powder-free, non-sterile latex or similar.
- Use paper towels to soak up the spill.
- If there is broken glass do not use hands even if gloved - use a paper or plastic scoop and dispose in the sharps box.
- Discard paper towels and any other waste from the spillage into clinical waste bags.
- Clean the contaminated area with water and detergent.
- Discard gloves and apron into a red bag
- Wash hands.

4.1.4.9 Mercury Spillages
As per the Delhi Govt. policy of mercury free hospital, every attempt has been made to make hospital mercury free. Mercury containing equipments are replaced and no mercury containing equipments are purchased by the Hospital.
5. SHARPS MANAGEMENT, SHARP INJURIES AND POST EXPOSURE PROPHYLAXIS

5.1 Sharp Management
5.1.1 Introduction
Safe handling and disposal of sharps is a vital component of the Standard Precautions approach to reduce the risk of transmission of blood borne virus.
Good practice involves:
- Correct assembly of the sharps container with proper size opening.
- Labelling of the container upon assembly as “SHARP CONTAINER” with Biohazard symbol and department name.
- Sharps container should not be more than two thirds full.
- Sharps containers are properly sealed before sending it for final segregation.
- Being aware of the first aid treatment following a needle-stick injury.
- Being aware of the follow up treatment after a used needle-stick injury.

5.1.2 Disposal of Sharps
- An adequate number of sharps containers, are located and conveniently placed in clinical areas.
- Ensure that the sharps containers have been assembled correctly.
- Make sure the department’s name is identified on the sharps bin.
- It is the responsibility of the person using the sharp to dispose of it safely.
- Sharps (needles, scalpel blades, razor blades and glass ampoules etc) are placed directly into a container.
- Whenever possible, take a sharps bin to the point of use.
- Needle must not to be recapped, bent or broken.
- If it is necessary to disassemble a needle and syringe, such as before transferring blood from a syringe to a pathological specimen bottle, the needles are placed in the sharps container before transferring the blood.
- Sharps containers are sealed closed when two-thirds to three-quarters full.
- Sharps containers when carried are held away from the body.
- Use needle safety devices where there are clear indications that they will provide a safer system of working.
- Needle collection tray in needle destroyer must be emptied in the morning by the coming nursing staff or more frequently if required. It should never be overfilled.
- Stray sharps should not be present.

5.2 Sharp injuries
This part is designed as guidance for all Health Care Workers in handling needle-stick injuries and exposure to blood and body fluids. An exposure that might place HCW at risk for HBV, HCV, or HIV infection is defined as:
- **Sharp Injury** - a percutaneous injury (e.g., a needle stick injury (NSI) or cut with a sharp object
- **Blood and body fluid exposure (BBF)** - Contact of mucous membrane or non-intact skin (e.g., exposed skin that is chapped, abraded or affected with dermatitis) – Contact with blood, tissue, or other body fluids that are potentially infectious
- **Contamination, from an Infected Known or Highly Suspected Person to another**
  - Recipient the Risks are:
    - Hepatitis B virus 1:3
    - Hepatitis C virus 1:30
    - Human Immunodeficiency Virus 1:300

It has been estimated that the risk of acquiring HIV through mucous membrane exposure splashed with contaminated body fluids is much less (probably 1 per 1000 injuries) 0.1%.

5.2.1 Main Risks From Needle-Stick Injury And Blood Contamination
The main concern is the transmission of bloodborne viruses, i.e.
- **HEPATITIS B** (HBV)
- **HEPATITIS C** (HCV)
- **HUMAN IMMUNODEFICIENCY VIRUS** (HIV)

5.2.2 Body Fluids Likely To Be Infectious
There is more experience of occupational exposure in the health care situation and in these circumstances the **highest risk** of transmission is from exposure to **liquid blood**. The risk is lower for other body fluids or body tissues from an infected patient. **Those, which represent a lower risk, are:**

- Cerebrospinal Fluid.
- Peritoneal Fluid, Pleural Fluid, Pericardial Fluid, Synovial Fluid, Amniotic Fluid
- Semen.
- Vaginal Secretions.
- Breast Milk.
- Any other body fluid containing visible blood, eg saliva.
- Bleeding gums in association with bites.
- Unfixed tissues and organs, ie those which have not been preserved in formalin.

### 5.2.3 Risks from Injuries

The risk of transmission is higher (particularly for HIV) when there is:

- A deep injury, i.e. when the injury is deeper than a superficial scratch drawing blood.
- Visible blood on the device that caused the injury (including teeth).
- Injury with a needle that had come from the source patient’s artery or vein.
- Terminal HIV related illness in the source patient.

**When does NSI Occur?**

- Recapping needles (Most important)
- Performing activities involving needles and sharps in a hurry
- Handling and passing needles or sharp after use
- Failing to dispose of used needles properly in puncture-resistant sharps containers
- Poor healthcare waste management practices
- Ignoring Universal Work Precautions

**Infections transmitted by NSI / BBF**

- **Blastomycosis**
- **Hepatitis B**
- **Malaria**
- **S. aureus**
- **Brucellosis**
- **Hepatitis C**
- **Mycobacteriosis**
- **S. pyogenes**
- **Cryptococcosis**
- **Herpes**
- **Mycoplasmosis**
- **Syphilis**
- **Diphtheria**
- **HIV**
- **Scrub typhus**
- **Toxoplasmosis**
- **Ebola fever**
- **Leptospirosis**
- **Tuberculosis**
- **Gonorrhoea**
- **Rocky mountain fever**

### 5.3 Management of the exposed site

#### 5.3.1 First Aid

**For skin** – if the skin is broken after a needle stick or sharp instrument:
- Immediately wash the wound & surrounding skin with water & soap and rinse.
- Do not scrub.
- Do not use antiseptics or skin scrub (bleach, chlorine, alcohol, betadine )

**After a splash of blood or body fluid:**
- To unbroken skin:
- Wash the area immediately
- Do not use antiseptics.

**For the eye:**
- Irrigate the exposed eye immediately with water or normal saline.
- Sit in a chair, tilt head back and ask a colleague to gently pour water or normal saline over the eye.
- If wearing contact lenses, leave them in place while irrigating, as they form a barrier over the eye and will help to protect it. Once the eyes cleaned, remove the contact lens and clean them in normal manner. This will make them to wear again.
- Do not use soap or disinfectant on the eye.

**For Mouth:**
- Spit fluid out immediately
- Rinse the mouth thoroughly, using water or saline and split again. Repeat this process several times.
- Do not use soap or disinfectant in the mouth.

Consult the designated physician of the institution for management of the exposure immediately.
### 5.3.2 SUMMARY OF DO’S & DONT’S

<table>
<thead>
<tr>
<th><strong>DO</strong></th>
<th><strong>DONT</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Remove gloves, if appropriate</td>
<td>• Do not panic</td>
</tr>
<tr>
<td>Wash the exposed site thoroughly with running water</td>
<td>• Do not put pricked finger in mouth</td>
</tr>
<tr>
<td>Irrigate with water or saline if eyes or mouth have been exposed</td>
<td>• Do not squeeze wound to bleed it</td>
</tr>
<tr>
<td>Wash the skin with soap and water</td>
<td>• Do not use bleach, chlorine, alcohol, betadine, iodine or any antiseptic or detergent</td>
</tr>
</tbody>
</table>

**Note:** Do consult the designated physician immediately as per institutional guidelines for management of the occupational exposure. Report all needle stick injuries to unit head / casualty medical officer. Fill the requisite proforma and send blood sample to microbiology laboratory for testing of HIV / HBsAg / HCV after pre-test counseling and consent of both patient and health care worker.

### 5.3.3 Establish eligibility for PEP

The HIV sero-conversion rate after an AEB (accidental exposure to blood) for percutaneous exposure is 0.3%. The risk of infection transmission is proportional to the amount of HIV transmitted (=amount of the contaminated fluid and the viral load)

Healthcare worker must inform ICN of the injury in designated form. After routine duty hours CMO on duty should be informed in designated form. The designate person shall assess the risk of HIV and HCV transmission following an AEB. This evaluation **must be made rapidly**, so as to start any treatment as soon as possible after the accident (ideally within 2 hours but certainly within 72 hours). This assessment must be made thoroughly (because not every AEB requires prophylactic treatment).

The first dose of PEP should be administered within the first 72 hours of exposure and the risk evaluated as soon as possible. If the risk is insignificant, PEP could be discontinued, if already commenced.

**PEP must be initiated as soon as possible, preferably within 2 hours**

Two main factors determine the risk of infection: the nature of exposure and the status of the source patient. Availability of PEP needs to be ensured at emergency Department for round-the-clock availability of PEP. The utilization data should be prepared on monthly basis as per NACO/DSACS guidelines.

**Post Exposure Prophylaxis**

- Post exposure prophylaxis is available for HIV in the form of antiretroviral drugs which are prescribed on the basis of NACO guidelines. HBV vaccine is available in routine hours and anti HBV immunoglobulin will be made available to the exposed worker as soon as possible after consulting with Microbiologist. For HBV PEP following criteria will act as guideline:

**Post-HIV exposure management / prophylaxis (PEP)**

- It is necessary to determine the status of the exposure and the HIV status of the exposure source before starting post-exposure prophylaxis(PEP)

**Immediate measures:**

- wash with soap and water
- No added advantage with antiseptic/bleach

**Next step:**

- Prompt reporting in accident/incident reporting forms
- Post-exposure treatment is begin as soon as possible
- Preferably within two hours
- Not recommended after seventy -two hours
- Late PEP? May be yes
• Is PEP needed for all types of exposures? No

1. Post exposure Prophylaxis:
The decision to start PEP is made on the basis of degree of exposure to HIV and the HIV status of the source from whom the exposure/infection has occurred. PEP is started, as early as possible, after an exposure. Incase PEP is initiated after 72 hours of exposure is of limited use and hence is not recommended. In case of anticipated delay of serology reports one dose of PEP may be given.

2. Determination of the Exposure Code (EC)
• Exposure code can be defined as per the flow chart given below

2. Status Code (SC)
Determined as per flow chart below.
3. Determine Post-Exposure Prophylaxis (PEP) Recommendation

<table>
<thead>
<tr>
<th>EC</th>
<th>HIV SC</th>
<th>PEP</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>Consider basic</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>Recommend basic regimen</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>Recommend expanded regimen</td>
</tr>
<tr>
<td>3</td>
<td>1 or 2</td>
<td>Recommend expanded regimen</td>
</tr>
<tr>
<td>1,2,3</td>
<td>Unknown</td>
<td>If exposure setting suggests risks of HIV Exposure, consider basic regimen</td>
</tr>
</tbody>
</table>

**Basic regimen (Three Drug Regimen):**
- Tenofovir 300 mg + Lamivudine 300 mg + Efavirenz 600 mg once daily for 28 days. **Expanded regimen: (Three drug regimen)**
- Basic regimen (+ Indinavir – 800 mg/thrice a day, or any other protease Inhibitor.)

4. Testing and Counseling
   The health care provider are tested for HIV as per the following schedule) to monitor seroconversion.
   - Base-line HIV test - at time of exposure
   - Repeat HIV test - at **six weeks** following exposure
   - 2nd repeat HIV test - at **twelve weeks** following exposure
   - 3rd repeat HIV test - at **6 months** following exposure

   - On all four occasions, HCW must be provided with a pre-test and post-test counseling. HIV testing are carried out on three ERS (Elisa/ Rapid/ Simple) test kits or antigen preparations as per NACO guidelines.
   - The HCW are advised to refrain from donating blood, semen or organs/tissues and abstain from sexual intercourse.
   - In case sexual intercourse is undertaken a latex condom be used consistently. In addition, women HCW should not breast-feed their infants.

5. Duration of PEP:
   - PEP is started, as early as possible, after an exposure. It has been seen that PEP started after 72 hours of exposure is of no use and hence is not recommended.
   - The optimal course of PEP is not unknown, but 4 weeks of drug therapy appears to provide protection against HIV.
   - If the HIV test is found to be positive at anytime within 12 weeks, the HCW are referred to a physician for treatment.

   - In case, exposed worker refuses PEP or refuses to get the laboratory testing done for monitoring of PEP, the same is documented on PEP refusal form.
5.3.4 Assessing the nature of exposure and risk of transmission

Three categories of exposure can be described based on the amount of blood/fluid involved and the entry port. These categories are intended to help in assessing the severity of the exposure but may not cover all possibilities.

### 5.3.4.1 Categories of exposure

<table>
<thead>
<tr>
<th>Category</th>
<th>Definition &amp; example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mild exposure</td>
<td>Mucous membrane/non-intact skin with small volumes&lt;br&gt;Or contact with the eyes or mucous membranes, subcutaneous injections following small-bore needles</td>
</tr>
<tr>
<td>Moderate exposure</td>
<td>Mucous membrane/non intact skin with large volumes or percutaneous&lt;br&gt;Superficial exposure with solid needle&lt;br&gt;Eg: a cut or needle stick injury penetrating gloves</td>
</tr>
<tr>
<td>Severe exposure</td>
<td>Percutaneous with large volume. Eg:&lt;br&gt;An accident with a high caliber needle (&gt;18G) visibly contaminated with blood;&lt;br&gt;A deep wound (haemorrhagic wound and/or very painful);&lt;br&gt;Transmission of a significant volume of blood;&lt;br&gt;An accident with material that has previously been used intravenously or intra-arterially.</td>
</tr>
</tbody>
</table>

The wearing of gloves during any of these accidents constitutes a protective factor. **Note:** In case of an AEB with material such as discarded sharps/needles, contaminated for over 48 hours, the risk of infection becomes negligible for HIV, but still remains significant for HBV. HBV survives longer than HIV outside the body.

5.3.5 Assessing the HIV status of the source of exposure

PEP needs to be started as soon as possible after the exposure and within 72 hours. In animal studies, initiating PEP within 12, 24 or 36 hours of exposure was more effective than initiating PEP 48 hours or 72 hours following exposure. PEP is not effective when given more than 72 hours after exposure. A baseline rapid HIV testing should be done before starting PEP.

Initiating of PEP where indicated should not be delayed while waiting for the results of HIV testing of the source of exposure. Informed consent should be taken before testing of the source as per national HIV testing guidelines.

### 5.3.5.1 Categories of situations depending on results of the source

<table>
<thead>
<tr>
<th>Source HIV status</th>
<th>Definition of risk in source</th>
</tr>
</thead>
<tbody>
<tr>
<td>HIV negative</td>
<td>Source is not HIV infected but consider HBV &amp; HCV</td>
</tr>
<tr>
<td>Low risk</td>
<td>HIV positive &amp; clinically asymptomatic</td>
</tr>
<tr>
<td>High risk</td>
<td>HIV positive &amp; clinically symptomatic</td>
</tr>
<tr>
<td>Unknown</td>
<td>Status of the patient is unknown &amp; neither the patient nor his/her blood is available for testing. The risk assessment will be based only upon the exposure</td>
</tr>
</tbody>
</table>

HIV infection is not detected during the primary infection period by routine use HIV tests. During the window period which lasts for approximately 6 weeks, the antibody level is still too low for detection, but infected persons can still have a high viral load. This implies that a positive HIV test result can help in taking the decision to start the PEP, but a negative test result does not exclude HIV infection. In districts or some population groups with a high HIV prevalence, a higher proportion of HIV infected individuals are found in the window period. In these situations, a negative result has even less value for decision making on PEP.

5.3.5.2 Assessment of the exposed individual
The exposed individual should have confidential counseling & assessment by an experienced physician. The exposed individual should be assessed for pre-existing HIV infection, intended for people who are HIV negative at the time of their potential exposure to HIV. Exposed individuals who are known or discovered to be HIV positive should not receive PEP. They should be offered counseling & information on prevention of transmission of & referred to clinical & laboratory assessment to determine eligibility for antiretroviral therapy (ART). Besides the medical assessment, counseling exposed HCP is essential to allay fear & start PEP (if required) at the earliest.

5.3.5.3 Counseling for PEP

Exposed persons should receive appropriate information about what PEP is about & the risk & benefits of PEP in order to provide informed consent. It should be clear that PEP is not mandatory.

<table>
<thead>
<tr>
<th>Key information to provide informed consent to the client after occupational exposure</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>The risk of acquiring HIV infection from the specific exposure</strong></td>
</tr>
<tr>
<td>• Ask client for understanding of HIV transmission risk after exposure</td>
</tr>
<tr>
<td>• The risk of getting HIV infection from a person known to be HIV positive is estimated to be</td>
</tr>
<tr>
<td>• Sharps injury :3 in 1000 exposures (0.3%)</td>
</tr>
<tr>
<td>• Mucous membrane splash: 1 in 1000 exposures(0.1%)</td>
</tr>
<tr>
<td>• The risk in increased with large exposure eg: needle stick from hollow bore needles with visible blood, from artery/vein &amp; from source patients with high viral load</td>
</tr>
<tr>
<td><strong>What is known about PEP efficacy</strong></td>
</tr>
<tr>
<td>• Ask clients understanding of PEP</td>
</tr>
<tr>
<td>• PEP is provided to prevent potential transmission of the HIV virus</td>
</tr>
<tr>
<td>• PEP is not 100%effective &amp; should be given within 72 hours</td>
</tr>
<tr>
<td>• Balance risk &amp; benefits of PEP: PEP may prevent HIV transmission, versus possible risk of side effects</td>
</tr>
<tr>
<td>• Information about clients risk of HIV infection based upon a risk assessment</td>
</tr>
<tr>
<td>• The importance of being tested &amp; receiving appropriate posttest counseling</td>
</tr>
<tr>
<td>• That PEP medicines will be discontinued if their initial HIV test is positive</td>
</tr>
<tr>
<td>• Clients possibility of prior HIV infection should be assessed.</td>
</tr>
<tr>
<td>• Counsel for HIV testing &amp; follow-up psychosocial support- where possible rapid testing should be used based on national testing guidelines</td>
</tr>
<tr>
<td>• Inform if the baseline test is positive, then the PEP will be discontinued</td>
</tr>
<tr>
<td>• Arrange referral to ART centre for assessment if found HIV positive</td>
</tr>
<tr>
<td>• Importance of adhering to medication once started</td>
</tr>
<tr>
<td>• Duration of the course of medicine (4weeks)</td>
</tr>
<tr>
<td>• Discuss dosing of the PEP medicine eg: pill should be taken twice a day for 28 days</td>
</tr>
<tr>
<td>• Depending on the nature &amp; risk of exposure, 2drugs or 3 drugs may be used</td>
</tr>
<tr>
<td>• Side effects may be important with use of 3 drugs.</td>
</tr>
<tr>
<td>• Expert opinion/consultation by phone or referral may be needed with a HIV specialist if 3rd drug is used.</td>
</tr>
<tr>
<td>• Arrange for special leave from work (2 weeks initially).</td>
</tr>
<tr>
<td>• Common side effects that may be experienced</td>
</tr>
<tr>
<td>• Discuss possible side effects of the PEP medicines eg: nausea, fatigue, headache</td>
</tr>
<tr>
<td>• Side effects often improve over time. It is often</td>
</tr>
</tbody>
</table>
minor & do not need specialized supervision.
- Symptomatic relief can also be given by using other drugs.

That they can stop at any time but will not get the benefit of PEP – if the source is HIV positive
- Animal studies suggest that taking less than 4 weeks of PEP does not work.
- If client decides to stop at any time, he needs to contact the physician before stopping the medication.
- Arrange for follow up visit & decide further course of action.

Prevention during the PEP period
- After any AEB, the exposed person should not have unprotected intercourse until it is confirmed, 3 months after the exposure, that he is not HIV infected.
- It is also advised to avoid pregnancy.
- Use of condoms is essential.

If client is pregnant – she can still take PEP during pregnancy
- The PEP drugs used are safe for pregnancy.
- If the client gets HIV during the pregnancy due to the exposure, the baby will have some risk of becoming HIV infected.

Safety of PEP if the client is breastfeeding
- The PEP drugs used are safe during breastfeeding.
- May consider stopping breast feeding if PEP is indicated.

Educate client on the possible signs & symptoms of early HIV seroconversion
- Signs & symptoms of early HIV seroconversion: fever, rash, oral ulcer, pharangitis, malaise, fatigue, joint pains, weight loss, mayalgia, headache (similar to flu like symptoms)

Risk of acquiring Hepatitis B & C from a specific exposure & availability of prophylaxis for this
- Risk of Hepatitis B is 9-30% from a needle stick exposure – client can be given vaccinations.
- Risk of Hepatitis C is 1-10% after a needle stick exposure – there is no vaccination for this.

HIV RNA testing by Reverse transcriptase polymerase chain reaction (RT-PCR) during PEP has a very poor positive predictive value & should be strongly discouraged.

Pregnancy testing should also be available, but its unavailability should not prevent the provision of PEP.

Other laboratory testing such as haemoglobin estimation should be available, especially when AZT is used in areas where anaemia is common.

Testing of other blood borne diseases such as syphilis, malaria & kala azar may also be useful, depending on the nature of risk, symptoms of the source patient, local prevalence & laboratory capacity.

Follow up of an Exposed Person

Whether or not PEP prophylaxis has been started, follow up is indicated to monitor for possible infections & provide psychological support.

Clinical follow up

In addition, in the weeks following an AEB, the exposed person must be monitored for the eventual appearance of signs indicating an HIV seroconversion: acute fever, generalized lymphadenopathy, cutaneous eruptions, pharangitis, non-specific flu symptoms & ulcers of the mouth & genital area. These symptoms appear in 50-70% of individuals with an primary infection & almost always within 3-6 weeks after exposure. When a primary infection is suspected, referral to an ART centre or for expert opinion should be arranged rapidly.

An exposed person should be advised to use precautions (eg- avoid blood/ tissue donations, breastfeeding, unprotected sexual relations or pregnancy. Condom use is essential.
Adherence and side effect counseling should be provided & reinforced at every follow-up visit. Psychological support & mental health counseling is often required.

**Laboratory follow up**

**Follow up HIV testing:** exposed persons should have post PEP HIV tests. Testing at the completion of PEP may give an initial indication of seroconversion outcome if the available antibody test is very sensitive. However, testing at 4-6 weeks may not be enough as use of PEP may prolong the time of seroconversion; & there is not enough time to diagnose all persons who seroconvert. Therefore testing at 3 months & 6 months is recommended. Very few cases of seroconversion after 6 months has been reported. Hence, no further testing is recommended if the HIV test at 6 months is negative.

<table>
<thead>
<tr>
<th>Recommended follow up laboratory tests</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Timing</strong></td>
</tr>
<tr>
<td>Weeks 2 &amp; 4</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Week 6</td>
</tr>
<tr>
<td>Month 3</td>
</tr>
<tr>
<td>Transaminases</td>
</tr>
<tr>
<td>Month 6</td>
</tr>
<tr>
<td>Transaminases</td>
</tr>
</tbody>
</table>

**Hepatitis B**

All health staff should be vaccinated against hepatitis B. the vaccination for Hepatitis B consists of 3 doses: initial, 1 month & 6 months. Sero conversion after completing the full course is 99%.

If the exposed person is unvaccinated or unclear vaccination status give complete hepatitis B vaccine series.

**Guidelines for Post exposure prophylaxis** of persons with nonoccupational exposures to blood or body fluids that contain blood by exposure type and vaccination status

<table>
<thead>
<tr>
<th>Exposure</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HBsAg Positive source</strong></td>
<td>Unvaccinated person</td>
</tr>
<tr>
<td>Percutaneous ( e.g. bite or needle stick ) or mucosal exposure to HBsAg positive blood or body fluids</td>
<td>Administer hepatitis B vaccine series and hepatitis B immune globulin (HBIG).</td>
</tr>
<tr>
<td>Sex or needle sharing contact of an HBsAg positive person</td>
<td>Administer hepatitis B vaccine series and HBIG</td>
</tr>
<tr>
<td>Victim of sexual assault/abuse by a perpetrator who is HBsAg positive</td>
<td>Administer hepatitis B vaccine series and HBIG</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Source with unknown HBsAg status</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Victim of sexual assault/abuse by a perpetrator with unknown HBsAg status</td>
<td>Administer hepatitis B vaccine series.</td>
</tr>
<tr>
<td>Percutaneous ( e.g. bite or needle stick ) or mucosal exposure to potentially infectious blood or body fluids form a source with unknown HBsAg status</td>
<td>Administer hepatitis B vaccine series.</td>
</tr>
<tr>
<td>Sex or needle sharing contact of person with unknown HBsAg status.</td>
<td>Administer hepatitis B vaccine series.</td>
</tr>
</tbody>
</table>
When indicated immunoprophylaxis should be initiated as soon as possible, preferably within 24 hours. Studies are limited on the maximum interval after exposure during which postexposure prophylaxis is effective, but the interval is unlikely to exceed 7 days for percutaneous exposures or 14 days for sexual exposures. The hepatitis B vaccine series should be completed.

These guidelines apply to nonoccupational exposures. Guidelines for management of occupational exposure have been published separately and also can be used for management of nonoccupational exposure, if feasible.

A person who is in the process of being vaccinated but who has not completed the vaccine series should complete the series and receive treatment as indicated.

A person who has written documentation of a complete hepatitis B vaccine series and who did not receive post vaccination testing.

**Determination of HBIG (Immunglobulin)**

For percutaneous, needlestick, ocular, or mucous-membrane exposure to blood known to contain HBsAg and for human bites from HBsAg carriers that penetrate the skin, a single dose of HBIG (0.06 ml/kg or 5.0 ml for adults) should be given as soon as possible after exposure and within 24 hours if possible. HB vaccine 1 ml (20 ug) should be given IM at a separate site as soon as possible, but within 7 days of exposure, with the second and third doses given after one month and 6 months, respectively. If HBIG is unavailable, immunoglobulin may be given in an equivalent dosage (0.06 ml/kg or 5.0 ml for adults). If an individual has received at least two doses of HB vaccine before an accidental exposure, no treatment is necessary if serologic tests show adequate levels (> 10MIU/DL) of anti-HBs. For persons who choose not to receive HB vaccine, the previously recommended two doses HBIG regimen may be used.

**HBV PROPHYLAXIS FOR REPORTED EXPOSURE INCIDENTS**

<table>
<thead>
<tr>
<th>HBV status of person exposed</th>
<th>Significant exposure</th>
<th>Non-significant exposure</th>
</tr>
</thead>
<tbody>
<tr>
<td>HBsAg positive source</td>
<td>Accelerated course of HB vaccine* HBIG x 1</td>
<td>Initiate course of HB vaccine</td>
</tr>
<tr>
<td>Unknown source</td>
<td>HBIG x 1</td>
<td>HBIG x 1</td>
</tr>
<tr>
<td>HBsAg negative source</td>
<td>HBIG x 1</td>
<td>No HBIG</td>
</tr>
<tr>
<td>Continued risk</td>
<td>HBIG x 1</td>
<td>No HBIG</td>
</tr>
<tr>
<td>No further risk</td>
<td>HBIG x 1</td>
<td>No HBIG</td>
</tr>
</tbody>
</table>

**Hepatitis C**

There is presently no prophylaxis available against hepatitis C. Post exposure management for HCV is based on early identification of chronic HCV disease & referral to a specialist for management. In the absence of PEP for HCV, recommendations for post exposure management are intended to achieve early identification of chronic disease and, if present, referral for evaluation of treatment options. However, a theoretical argument is that intervention with antivirals when HCV RNA first becomes detectable might prevent the development of chronic infection. Data from studies conducted outside the United States suggest that a short course of interferon started early in the course of acute hepatitis C is associated with a higher rate of resolved infection than that achieved when therapy is begun after chronic hepatitis C has been well established. These studies used various treatment
regimens and included persons with acute disease whose peak ALT levels were 500–1,000 IU/L at the time therapy was initiated (2.6–4 months after exposure). No studies have evaluated the treatment of acute infection in persons with no evidence of liver disease (i.e., HCV RNA-positive <6 months duration with normal ALT levels); among patients with chronic HCV infection, the efficacy of antivirals has been demonstrated only among patients who also had evidence of chronic liver disease (i.e., abnormal ALT levels). In addition, treatment started early in the course of chronic HCV infection (i.e., 6 months after onset of infection) might be as effective as treatment started during acute infection. Because 15%–25% of patients with acute HCV infection spontaneously resolve their infection, treatment of these patients during the acute phase could expose them unnecessarily to the discomfort and side effects of antiviral therapy. Data upon which to base a recommendation for therapy of acute infection are insufficient because a) No data exist regarding the effect of treating patients with acute infection who have no evidence of disease,
b) Treatment started early in the course of chronic infection might be just as effective and would eliminate the need to treat persons who will spontaneously resolve their infection, and
c) The appropriate regimen is unknown

Pregnancy and PEP:
• Based on limited information, anti-retroviral therapy taken during 2nd and 3rd trimester of pregnancy has not caused serious side effects in mothers or infants. There is very little information on the safety in the 1st trimester. If the HCW is pregnant at the time of exposure to HIV, the designated authority/physician must be consulted about the use of the drugs for PEP.

Side-effects of these drugs:
• Most of the drugs used for PEP have usually been tolerated well except for nausea, vomiting, tiredness, or headache.

Follow-Up Of HCW With Sharps Injury Or BBF For HBV & HCV Seroconversion:
• SGOT and SGPT test - at six weeks following exposure and at twelve weeks following exposure
• In case above mentioned parameters are found deranged then HCW should be screened for seroconversion. If found positive, HCW should be referred to Hepatologist.

References:
• NACO PEP Guidelines
• CDC Guidelines for the Management of Occupational Exposures to HBV, HCV, and HIV and Recommendations for Post exposure Prophylaxis
• MMWR. DEC 8, 2006/55(RR16)
6. ISOLATION POLICY AND TRANSMISSION BASED PRECAUTIONS

6.1 Isolation Policy
CRITERIA FOR ISOLATION AND PROCEDURES
6.1.1 Aim
- To prevent the transmission of pathogenic microorganisms within the hospital.
- To recognize the importance of all body fluids, secretions and excretions in the transmission of healthcare associated pathogens.
- To practice adequate precautions for infections transmitted by airborne Droplet & contact

6.1.2 Measures for Reduction of Transmission
6.1.2.1 Hand Washing
- Frequent hand washing is the most important measure.
6.1.2.2 Patient care Handwash
- Wash hands after touching blood, body fluids, secretions, excretions and contaminated items, whether gloves are worn or not.
- Wash hands immediately after gloves are removed.
- Wash hands between tasks and procedures on the same patient to prevent cross contamination of different body sites.
- Use a plain soap for routine hand washing.
- Use antiseptic soap or an alcohol based disinfectant followed by thorough hand washing for accidental skin contamination. Antimicrobial hand washing products are used for hand washing before personnel care for newborns and when otherwise indicated during their care, between patients in high-risk units, and before personnel take care of severely immunocompromised patients.
6.1.2.3 Surgical Hand Wash
- Procedural hand hygiene includes a full surgical scrub using running water and 4% chlorhexidine scrub solution from the fingertips to the elbow. The scrub is performed for a minimum of 2 to 3 minutes.
6.1.2.4 Gloves
- Clean, unsterile gloves may be worn as a protective barrier during procedures. Sterile gloves are worn when sterile procedures are undertaken.
6.1.2.5 Personal Protective Equipment: (PPE)
- Gowns: A clean, no sterile, gown is worn to prevent contamination of clothing and skin of personnel from exposure to blood and body fluids. When gowns are worn to attend to a patient requiring barrier nursing, they are removed before leaving the patients environment and hand washing is done.
- Masks and goggles: This equipment is worn to provide barrier protection. Mask should cover both the nose and the mouth.

6.2 Patient Isolation
Patients are isolated when suffering from highly transmissible diseases e.g. chicken pox. These patients are provided with isolation through designated isolation areas (e.g. isolation room in swine flu ward – when no patient of swine flu is admitted or in a single room at private wards).

6.2.1 Barrier Nursing
Barrier nursing: The aim is to erect a barrier to the passage of infectious pathogenic organisms between the contagious patient and other patients and staff in the hospital, and hence to the outside world. Preferably, all contagious patients are isolated in separate rooms, but when such patients must be nursed in a ward with others, screens are placed around the bed or beds they occupy. Cohort nursing may be practiced as re-infection with the same organism is unlikely. The nurses, attending consultants as also any visitors must wear gowns, masks, and sometimes rubber gloves and they observe strict rules that minimize the risk of passing on infectious agents. Surgical standards of cleanliness in hand washing are observed after they have been attending the patient.
- Bedding is carefully moved in order to minimize the transmission of airborne particles, such as dust or droplets that could carry contagious material.
Barrier nursing must be continued until subsequent cultures give a negative report. Infected with epidemiologically important microorganisms such as MRSA, Pan-resistant gram-negative bacteria are kept in their patient care unit with alert of zero tolerance barrier nursing.

6.2.2 Cleaning of Equipment and articles
Contaminated disposable articles are bagged appropriately in leak proof bags and disposed. Critical reusable medical equipment is disinfected or sterilized after use. Non-critical equipment is cleaned, disinfected after use.

6.2.3 Laundry
Soiled linen are handled as little as possible and with minimum agitation to prevent gross microbial contamination of the air and of persons handling the linen. All soiled linen are bagged in red bag with proper labels and put into small carts at the location where it was used than transferred into the big carts; it should not be sorted or pre-rinsed in patient-care areas and transported to the laundry from the pre-defined corridors.

6.2.4 Eating Utensils
Routine cleaning with detergent and hot water is sufficient.

6.2.5 Terminal Cleaning
Terminal cleaning of patient unit should be done with appropriate disinfectant solution.
- Bed should be cleaned properly including Bed frames, side rails and mattress initially with soap and water followed by disinfectant solution.
- Other equipments like I/V stands, bed lockers etc should be cleaned with soap and water followed by disinfectant solution.
- All metal items should me clean with bacillocid (0.5%) and non-metal items can be clean with superoxide water.
- If any electrical items like infusion pumps etc. used should be clean with spirit twice.
- All the used items like oxygen mask, O2 tubes, suction jars and tubing’s should be send to CSSD for HLD.
- Ventilator is used for the patient then whole ventilator tubing’s should be send to CSSD for autoclaving after primary cleaning. Ventilator surface also should be disinfected.
- Ventilator switching from one patient to another is strongly discouraged.
- All wall tiles and floor should clean with soap and water.
- swine flu ward – when no patient of swine flu are admitted or in a single room at private wards).

6.2.6 Isolation policy for certain groups of organism
1. MRSA: When MRSA is isolated in the lab the microbiologist will inform the sister-in-charge/duty doctor/head of unit. Patient is isolated and barrier nursed. Hand washing is strictly adhered to by all concerned. Linen is changed on a daily basis. Dirty linen is carefully packed in red bag with proper label and sent to laundry.

2. Multi-resistant bacteria e.g. Imipenem resistant Acinetobacter, multi-resistant *Pseudomonas aeruginosa*: The aim is to curtail the spread of such bacteria. Hence patient is to be placed on strict barrier nursing precautions irrespective of whether the organism is a coloniser or the cause of infection.

3. Pulmonary tuberculosis: Masks are used during the care of all patients with sputum positive pulmonary tuberculosis.

Note: Isolation precautions are to be followed until all previous culture sites are negative.

3. HIV/HBsAg/ HCV infected patients: Universal precautions.

6.2.7 List of diseases which need isolation precautions

<table>
<thead>
<tr>
<th>Condition</th>
<th>PPE REQUIRED</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chicken-Pox (Varicella)</td>
<td>Gloves, plastic apron for contact.</td>
<td>Preferably Single room. <strong>Staffs who have not had Chicken-Pox should not nurse these patients.</strong></td>
</tr>
</tbody>
</table>
### german measles (rubella)
Gloves/apron for direct contact
Single room. Check on any non-immune pregnant staff.

### hepatitis
**type a infection**
Gloves
None

**type b serum**
Gloves
Single room if bleeding.

**type c serum**
Gloves
Single room if bleeding.

### HIV/AIDS
Gloves/apron in contact with body fluids
Single room if bleeding or has an opportunistic infection.

### Herpes Zoster
Gloves/apron for direct contact
Single room. Staffs who have not had Chicken-Pox /vaccinated should not nurse these patients.

### impetigo
Gloves for direct contact.
Single room.

### Measles (including encephalitis)
Gloves/apron in direct contact.
Isolation room.

### Infection with Multi – resistant organisms, including MRSA ,VRE
Gloves/Apron for direct contact
Strict hand washing is essential. Isolation room/cohort nursing

### scabies
Gloves for contact until treated. 24 hours after treatment not infectious.
All staff in contact need treatment also other patients.

### Tuberculosis
**Pulmonary**
Gloves/apron for direct contact. Masks to be worn by staff for 2 weeks after patient starting treatment.
Masks must be worn in open cases of tuberculosis. Transfer to infectious disease hospital

---

### 6.3 Transmission Based Precautions
Besides standard precautions, specific transmission based precautions are observed according to the mode of transmission of the various conditions to protect health care workers and other patients from cross infections.

**Table: Transmission based Precautions**

<table>
<thead>
<tr>
<th>Precautions</th>
<th>Mode of Transmission is Contact (category I)</th>
<th>Mode of Transmission is Droplet (categoryII)</th>
<th>Mode of Transmission is Airborne (categoryIII)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mask</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Gown</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Gloves</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Patient Transport</td>
<td>Receiving department to be informed of precautions</td>
<td>1. Mask the patient</td>
<td>3. Mask the patient</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Receiving department to be informed of precautions</td>
<td>4. Inform the receiving department of precautions</td>
</tr>
<tr>
<td>Environment Cleaning</td>
<td>5. Dedicate or change solutions and equipment after use</td>
<td>Routine</td>
<td>Routine</td>
</tr>
<tr>
<td></td>
<td>6. Change privacy curtain when isolation is discontinued or patient is discharged</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patient Care Equipment (Special Handling)</td>
<td>7. Yes, dedicated equipments</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>
Table 2 Reference Table of Standard and Transmission based precautions. Precautions for Various Diseases and Conditions

<table>
<thead>
<tr>
<th>Diseases / Condition</th>
<th>Precaution Category</th>
<th>Infective Material</th>
<th>Duration for Precautions</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abscess, draining, major</td>
<td>Contact, drainage</td>
<td>Drainage</td>
<td>Until drainage contained</td>
<td>Major = drainage not contained by dressing</td>
</tr>
<tr>
<td>Acid Fast Bacillus Positive</td>
<td>See Tuberculosis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acquired immunodeficiency syndrome (AIDS)</td>
<td>Standard, blood and bloody body fluids</td>
<td>AIDS is specified communicable disease.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Actinomycosis</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amebiasis (Dysentery) Adult</td>
<td>Standard, faeces</td>
<td></td>
<td></td>
<td>Consider Contact precautions for adults with poor hygiene and/or who contaminate the environment.</td>
</tr>
<tr>
<td>Pediatric</td>
<td>Contact, faeces</td>
<td></td>
<td>Until formed or normal stools × 24 hours</td>
<td></td>
</tr>
<tr>
<td>Arthropod borne viral encephalitis (Jap B)</td>
<td>Standard, blood and bloody body fluids</td>
<td>Arthropod borne viral fever is a specified communicable disease.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arthropod borne viral fevers (Dengue)</td>
<td>Standard, blood and bloody body fluids</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aspergillosis</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bronchiolitis Adult</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pediatric</td>
<td>Contact, respiratory secretions</td>
<td>Duration of symptoms</td>
<td>Various etiologic agents, such as respiratory syncytial virus, parainfluenza viruses, adenoviruses have been associated with this condition</td>
<td></td>
</tr>
<tr>
<td>Candidiasis</td>
<td>All forms, including mucocutaneous (moniliasis, thrush)</td>
<td>Standard</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cellulitis (Uncontrolled drainage)</td>
<td>Contact, drainage</td>
<td>Drainage</td>
<td>Until drainage contained</td>
<td></td>
</tr>
<tr>
<td>Chancroid (Soft chancre)</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chickenpox (Varicella) Caused by Varicella zoster virus.</td>
<td>Airborne AND Contact Respiratory Secretion and Lesions</td>
<td>Until all lesions are crusted</td>
<td>Negative pressure room is required. Neonates born to mothers with active</td>
<td></td>
</tr>
</tbody>
</table>
Varicella should be placed on Airborne and Contact isolation at birth. Exposed susceptible patients should be placed on Airborne and contact isolation beginning 10 days after first exposure and continuing until 21 days after last exposure (up to 28 days if VZIG given). First exposure is defined as day one. Consult attending physician to assess need for VZIG.

<table>
<thead>
<tr>
<th>Disease</th>
<th>Isolation Type</th>
<th>Exposed Isolation</th>
<th>Period of Communicability</th>
</tr>
</thead>
<tbody>
<tr>
<td>Varicella</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cholera</td>
<td>Contact</td>
<td>Faeces</td>
<td>Until formed or normal stools × 24 hours</td>
</tr>
<tr>
<td>Clostridium difficile diarrhea</td>
<td>Contact</td>
<td>Faeces</td>
<td>Until formed normal stools or no stools × 48 hours</td>
</tr>
<tr>
<td>Clostridium perfringens (Gas gangrene)</td>
<td>Standard</td>
<td>Faeces</td>
<td></td>
</tr>
<tr>
<td>Congenital rubella</td>
<td>Contact and Droplet</td>
<td>Respiratory secretions and urine</td>
<td>During any admission for the 1st year after birth unless nasopharyngeal and urine cultures after 3 months of age are negative for rubella virus</td>
</tr>
<tr>
<td>Conjunctivitis (Pink eye Acute bacterial Chlamydia, Gonococcal Acute Viral)</td>
<td>Standard</td>
<td>Eye Discharge</td>
<td>Duration of symptoms</td>
</tr>
<tr>
<td>Croup Pediatric</td>
<td>Contact</td>
<td>Respiratory secretions</td>
<td>Duration of symptoms</td>
</tr>
<tr>
<td>Cryptosporidiosis</td>
<td>Standard</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cryptosporidiosis Adult</td>
<td>Standard</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Susceptible persons should stay out of room.

Viral Agents such as parainfluenza viruses and influenza A virus have been associated with this condition.

Consider contact precautions for adults with poor hygiene and/or who contaminate environment.
<table>
<thead>
<tr>
<th>Illness</th>
<th>Contact</th>
<th>Faeces</th>
<th>Until formed or normal stools 24 hours</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cytomegalovirus infection</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Decubitus ulcer major</td>
<td>Contact</td>
<td>Drainage</td>
<td>Until drainage contained</td>
<td>Major = drainage not contained by dressing.</td>
</tr>
<tr>
<td>Dengue</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diarrhea, acute</td>
<td>Contact</td>
<td>Faeces</td>
<td>Until formed or normal stools × 24 hours</td>
<td></td>
</tr>
<tr>
<td>Diarrhea, acute</td>
<td>Contact</td>
<td>Faeces</td>
<td>Until formed or normal stools × 24 hours</td>
<td></td>
</tr>
<tr>
<td>Diphtheria (Corynebacterium diphtheriae Cutaneous)</td>
<td>Contact</td>
<td>Lesion secretions.</td>
<td>Until 2 cultures from skin lesions taken at least 24 hours apart after cessation of antimicrobial therapy are negative</td>
<td></td>
</tr>
<tr>
<td>Diphtheria Pharyngeal</td>
<td>Droplet</td>
<td>Respiratory secretions</td>
<td>Until 2 cultures from both nose and throat taken at least 24 hours apart after cessation of antimicrobial therapy are negative for corynebacterium diphtheriae</td>
<td></td>
</tr>
<tr>
<td>Epiglottitis</td>
<td>Droplet</td>
<td>Respiratory secretions</td>
<td>For 24 hours after start of effective therapy.</td>
<td></td>
</tr>
<tr>
<td>Haemophilus influenza Type B</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Epstein-Barr virus infection (including infectious mononucleosis)</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Food Poisoning (Botulism, clostridium perfringens or Staphylococcus)</td>
<td>Contact</td>
<td>Faeces</td>
<td>Until formed or normal stools × 24 hours.</td>
<td></td>
</tr>
<tr>
<td>Furunculosis, staphylococcal (Pediatric)</td>
<td>Contact</td>
<td>Drainage</td>
<td>Until drainage stops</td>
<td></td>
</tr>
<tr>
<td>Gonorrhea</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Guillain-Barre syndrome</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Helicobacter pylori</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hepatitis Viral</td>
<td>Standard</td>
<td></td>
<td></td>
<td>For Hepatitis A &amp; E consider contact precautions for adults with poor hygiene and/or who contaminate the environment.</td>
</tr>
<tr>
<td>Hepatitis A, Hepatitis E Adult</td>
<td>Standard</td>
<td></td>
<td></td>
<td>For 7 days after onset of symptoms</td>
</tr>
<tr>
<td>Disease</td>
<td>Standard</td>
<td>Blood and bloody fluids</td>
<td>Duration of symptoms.</td>
<td></td>
</tr>
<tr>
<td>------------------------------------------------------------------------</td>
<td>-----------</td>
<td>-------------------------</td>
<td>-----------------------</td>
<td></td>
</tr>
<tr>
<td>Hepatitis B (HBsAg + )</td>
<td>Standard</td>
<td>Standard</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hepatitis C and other specified non A, non B</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Herpes simplex (Herpes virus hominis) Encephalitis</td>
<td>Contact</td>
<td>Lesion, secretions, possibly all body secretions and excretions.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neonatal</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mucocutaneous, disseminated or primary severe</td>
<td>Contact</td>
<td>Lesion secretions</td>
<td>Duration of symptoms.</td>
<td></td>
</tr>
<tr>
<td>Mucocutaneous, recurrent skin, oral or genital</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Herpes Zoster Caused by Varicella zoster virus (shingles)</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Localized in normal patient.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Localized in immunocompromised patient, and/or disseminated in any patient.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Airborne</td>
<td>Lesion secretions and possibly respiratory secretions.</td>
<td>For 72 hours after start of effective antiviral therapy or if untreated until all lesions are crusted.</td>
<td></td>
</tr>
</tbody>
</table>

Hepatitis B and C are specified communicable disease. For staff issues for all types of Hepatitis.

Precautions are indicated for infants delivered either vaginally or by caesarean section (if membranes have been ruptured more than 4-6 hrs) to women with active genital herpes simplex infections, until neonatal HSV infection has been ruled out.

For localized lesions, try to contain with dressings. Roommates should not be susceptible to chickenpox.

Negative pressure isolation room required.

Exposed susceptible patients should be placed on Airborne and contact isolation beginning 10 days after first exposure and continuing until
<table>
<thead>
<tr>
<th>Disease</th>
<th>Transmissibility</th>
<th>Contact</th>
<th>Duration</th>
<th>Isolation</th>
<th>Precautions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human immunodeficiency virus (HIV)</td>
<td>Standard</td>
<td>Blood &amp; Bloody body fluids</td>
<td>For 7 days after onset of symptoms.</td>
<td></td>
<td>If private room is unavailable, consider cohorting patients with influenza.</td>
</tr>
<tr>
<td>Influenza</td>
<td>Droplet</td>
<td>Nasopharyngeal secretions.</td>
<td>For 4 days after start of rash, except in immunocompromised patients for whom precautions should be maintained for duration of illness.</td>
<td></td>
<td>Negative pressure room is required. Exposed susceptible patients should be placed on Airborne isolation beginning 5 days after first exposure through 21 days after last exposure.</td>
</tr>
<tr>
<td>Leprosy (Hansen's disease)</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leptospirosis</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Malaria</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Measles (Rubella)</td>
<td>Airborne</td>
<td>Respiratory secretions.</td>
<td>For 24 hours after start of effective therapy</td>
<td></td>
<td>Bacterial Meningitis is a specified communicable disease.</td>
</tr>
<tr>
<td>Meningitis Unknown etiology</td>
<td>Droplet</td>
<td>Possibly Respiratory Secretions</td>
<td>Until etiology Known</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neisseria meningitidis (meningococcal) known or suspected</td>
<td>Droplet</td>
<td>Respiratory secretions</td>
<td>For 24 hours after start of effective therapy</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haemophilus influenzae Type b known or suspected</td>
<td>Droplet</td>
<td>Respiratory secretions</td>
<td>For 24 hours after start of effective therapy</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other Bacterial, Fungal</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aseptic (Viral or non (bacterial)</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Meningococcemia (meningococcal sepsis)</td>
<td>Droplet</td>
<td>Respiratory secretions</td>
<td>For 24 hours after start of effective therapy</td>
<td></td>
<td>Meningococcemia is a specified communicable disease.</td>
</tr>
<tr>
<td>Methicillin Resistant Staphylococcus aureus (MRSA)</td>
<td>Contact</td>
<td>Any body fluid or site</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mucormycosis</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| Mumps (infectious parotitis) | Droplet | Respiratory secretions | For 9 days after onset of swelling | | Exposed susceptible patients should be
<table>
<thead>
<tr>
<th>Disease Description</th>
<th>Isolation Type</th>
<th>Isolation Period</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mycobacterium (non-tuberculosis, atypical, non TB complex) Pulmonary</td>
<td>Standard</td>
<td>Duration of symptoms</td>
<td>Placed on Droplet isolation beginning 12 days after first contact through 26 days after last exposure.</td>
</tr>
<tr>
<td>Mycoplasma pneumonia (Primary atypical pneumonia)</td>
<td>Droplet</td>
<td>Respiratory secretions</td>
<td>Duration of symptoms</td>
</tr>
<tr>
<td>Nocardiosis</td>
<td>Standard</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pertussis (Whooping cough)</td>
<td>Droplet</td>
<td>Respiratory secretions</td>
<td>For 5 days after start of effective therapy or 3 weeks after onset of paroxysms if not treated</td>
</tr>
<tr>
<td>Pharyngitis</td>
<td>Standard</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plague (Yersinia pestis) Bubonic</td>
<td>Standard</td>
<td></td>
<td>Bubonic plague is a specified communicable disease.</td>
</tr>
<tr>
<td>Pneumonic</td>
<td>Droplet</td>
<td>Respiratory</td>
<td>For 3 days after start of effective therapy</td>
</tr>
<tr>
<td>Pneumococcal Infections, Invasive</td>
<td>Standard</td>
<td></td>
<td>Invasive (cultured from sterile site) pneumococcal infections are a specified communicable disease.</td>
</tr>
<tr>
<td>Pneumonia</td>
<td>Standard</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haemophilus influenza Type b Adult</td>
<td>Standard</td>
<td></td>
<td>For 24 hours after start of effective therapy</td>
</tr>
<tr>
<td>Pediatric</td>
<td>Droplet</td>
<td>Respiratory secretions</td>
<td></td>
</tr>
<tr>
<td>Neisseria meningitidis (meningococcal) known or suspected</td>
<td>Droplet</td>
<td>Respiratory secretions</td>
<td>For 24 hours after start of effective therapy</td>
</tr>
<tr>
<td>Mycoplasma (Primary atypical pneumonia) known or suspected</td>
<td>Droplet</td>
<td>Respiratory secretions</td>
<td>Duration of symptoms</td>
</tr>
<tr>
<td>Pneumocystis carinii</td>
<td>Standard</td>
<td></td>
<td>Ensure roommate not immunocompromised</td>
</tr>
<tr>
<td>Staphylococcus aureus</td>
<td>Standard</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Streptococcus. Group A</td>
<td>Standard</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

DRAFT
<table>
<thead>
<tr>
<th>Adult</th>
<th>Pediatrics</th>
<th>Standard</th>
<th>Respiratory</th>
<th>start of effective</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other bacterial</td>
<td>Standard</td>
<td>Droplet</td>
<td>secretions.</td>
<td>therapy</td>
</tr>
<tr>
<td>including gram negative and etiology unknown.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fungal</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Viral</td>
<td>Standard</td>
<td>Respiratory</td>
<td>secretions.</td>
<td></td>
</tr>
<tr>
<td>Adult</td>
<td></td>
<td>Duration of</td>
<td>symptoms</td>
<td></td>
</tr>
<tr>
<td>Pediatrics</td>
<td>Droplet</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Poliomyelitis</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pseudo membranous colitis</td>
<td>Contact</td>
<td>Faeces</td>
<td>Until Clostridium difficile ruled out.</td>
<td></td>
</tr>
<tr>
<td>Rabies</td>
<td>Standard</td>
<td></td>
<td></td>
<td>Rabies is a specified communicable disease.</td>
</tr>
<tr>
<td>Rheumatic fever</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ritter's disease (Staphylococcal scalded skin syndrome)</td>
<td>Standard</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adult</td>
<td></td>
<td></td>
<td></td>
<td>Consider contact precautions for adults with poor hygiene and/or who contaminate the environment.</td>
</tr>
<tr>
<td>Pediatric</td>
<td>Contact</td>
<td>Faeces</td>
<td>Until formed or normal stools x 24 hrs.</td>
<td></td>
</tr>
<tr>
<td>Rubella (German measles)</td>
<td>Droplet</td>
<td>Respiratory Secretions</td>
<td>Until 7 days after onset of rash</td>
<td>Exposed susceptible patients should be placed on Droplet isolation beginning 12 days after first contact through 26 days after last exposure.</td>
</tr>
<tr>
<td>Salmonellae Including Typhoid fever or Salmonella typhi (case/carrier)</td>
<td>Standard</td>
<td></td>
<td></td>
<td>Consider contact precautions with poor hygiene and/or who contaminate the environment.</td>
</tr>
<tr>
<td>Adult</td>
<td></td>
<td></td>
<td></td>
<td>Typhoid fever is a specified communicable disease.</td>
</tr>
<tr>
<td>Pediatric</td>
<td>Contact</td>
<td>Faeces</td>
<td>Until formed or normal stools x 24 hours</td>
<td></td>
</tr>
<tr>
<td>Shigellosis</td>
<td>Standard</td>
<td></td>
<td></td>
<td>Consider Contact precautions for adults with poor hygiene and/or who contaminate the environment.</td>
</tr>
<tr>
<td>Condition</td>
<td>Contact Type</td>
<td>Faeces Type</td>
<td>Duration After Start of Effective Therapy</td>
<td>Isolation Requirement</td>
</tr>
<tr>
<td>--------------------------------------------------------------------------</td>
<td>--------------</td>
<td>-------------</td>
<td>-------------------------------------------</td>
<td>-----------------------</td>
</tr>
<tr>
<td>Pediatric Contact Faeces Until formed or normal stools × 24 hours</td>
<td>Standard</td>
<td>Drainage</td>
<td>For 24 hours after start of effective therapy.</td>
<td>Major = drainage not contained by dressing.</td>
</tr>
<tr>
<td>Streptococcal infection (Group A Streptococcus)</td>
<td>Standard</td>
<td>Drainage</td>
<td>For 24 hours after start of effective therapy.</td>
<td></td>
</tr>
<tr>
<td>Skin, wound or major burn Contact Drainage For 24 hours after start of effective therapy. Major = drainage not contained by dressing.</td>
<td>Standard</td>
<td>Drainage</td>
<td>For 24 hours after start of effective therapy.</td>
<td></td>
</tr>
<tr>
<td>Necrotizing fascitis, myositis or other soft tissue necrosis.</td>
<td>Contact</td>
<td>Drainage</td>
<td>For 24 hours after start of effective therapy.</td>
<td></td>
</tr>
<tr>
<td>Pneumonia Adult Standard Droplet Respiratory secretions For 24 hours after start of effective therapy.</td>
<td>Standard</td>
<td>Respiratory secretions</td>
<td>For 24 hours after start of effective therapy.</td>
<td></td>
</tr>
<tr>
<td>Pediatric Droplet Respiratory secretions For 24 hours after start of effective therapy.</td>
<td>Standard</td>
<td>Respiratory secretions</td>
<td>For 24 hours after start of effective therapy.</td>
<td></td>
</tr>
<tr>
<td>Scarlet fever Pediatric Droplet Respiratory secretions For 24 hours after start of effective therapy.</td>
<td>Standard</td>
<td>Respiratory secretions</td>
<td>For 24 hours after start of effective therapy.</td>
<td></td>
</tr>
<tr>
<td>Toxic Shock Syndrome (TSS) Standard</td>
<td>Standard</td>
<td>Respiratory secretions</td>
<td>Prior to discontinuing isolation</td>
<td></td>
</tr>
<tr>
<td>Syphilis Standard</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tetanus Standard</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Toxoplasmosis Standard</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trachoma Standard</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tuberculosis (Mycobacterium tuberculosis. M. africanum M. bovis)</td>
<td>Airborne</td>
<td>Respiratory secretions</td>
<td>Prior to discontinuing isolation</td>
<td>Negative pressure isolation room is required.</td>
</tr>
<tr>
<td>Confirmed or suspected pulmonary, laryngeal, or military.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Skin-test (mantoux), positive with no evidence of current pulmonary disease.</td>
<td>Standard</td>
<td>Respiratory secretions</td>
<td>Prior to discontinuing isolation</td>
<td></td>
</tr>
<tr>
<td>Extra pulmonary, meningitis, and drainage lesion (including scrofula).</td>
<td>Standard</td>
<td>Respiratory secretions</td>
<td>Prior to discontinuing isolation</td>
<td>Assess for pulmonary disease.</td>
</tr>
<tr>
<td>UTI Including pyelonephritis with or without urinary catheter</td>
<td>Standard</td>
<td>Respiratory secretions</td>
<td>Prior to discontinuing isolation</td>
<td></td>
</tr>
</tbody>
</table>
7. STERILISATION, DISINFECTION AND DECONTAMINATION

7.1 STERILISATION
Definition
Sterilization is defined as a process where all microbes are removed from a defined object, inclusive of bacterial endospores.

7.1.1 Methods:
- **Heat Sterilization:**
  - **Moist Heat:** Exposure to saturated steam at 121°C for 15-20 min OR 134°C for 4 min in any autoclave.
  - **Dry Heat:** Exposure to dry heat at 160°C for 120 min.
- **Chemical Sterilization:** (for heat sensitive items)
  - Ethylene oxide
- **Low temperature Sterilization**
  - Plasma sterilizer using Per acetic acid or hydrogen peroxide.

7.1.2 Packing & Loading
For effective sterilization, selection of packaging material plays important role apart from sterilization parameters. The following are keys in selecting a suitable packaging material.
1. The packaging material must be permeable to sterilizing agent.
2. The packaging material must be impermeable to bacteria and other contaminants.
3. The packaging material must resist tears and punctures.
4. It should facilitate aseptic presentation of packaged content.

**Proper loading of material inside sterilizer is very critical for efficient sterilization. Relative humidity in the processing area should be at least 35%.**
- When loading sterilizer there should be space between item to facilitate circulation and penetration of sterilant.
- There should be no contact between items and chamber wall.
- In mixed load linen should be kept on top racks and metal on bottom
- Peel pouches should be kept on the edge facing same direction
- Textile should be kept on the edge
- Instrument sets should be placed flat

7.1.3 Monitoring:
- Mechanical, chemical and biological monitors can be used to evaluate the effectiveness of the sterilization process.
- Each load is monitored with mechanical (time, temperature, pressure) and chemical (internal and external) indicators.
- Biological indicators (spores) should be used weekly to monitor the effectiveness of sterilization.
- Chemical indicators as strips should be used with every batch.

7.1.4 ETO monitoring
- Use to sterilize items that are moisture or heat sensitive.
- Essential parameters of ETO sterilization includes:
  - Temperature – Should be 40-55°C
  - Exposure time – 16 hours

**AN1087 Dosimeters** are placed with every run. They change color from yellow to blue when exposed to Ethylene oxide. They integrate the effects of time, temperature and the concentration of Ethylene oxide in contact with the crystals in the capillary tube.
For a load to be considered sterile, the color change from yellow to blue must extend past the triangular mark on the label. No laboratory testing is required. The information is available immediately at the end of a sterilization cycle.

**Biological Indicators – Done weekly**
Each AN1080 Biological and Chemical Sterilizer Control pouch is a complete sterility control. Steritest eliminates the possibility of a false positive by including both a spore strip and an ampoule of sterile culture broth sealed in a transparent, gas permeable, waterproof, plastic pouch.

Place the unopened Steritest with the items to be sterilized. At the end of the cycle, remove the Steritest and look at the Dosimeter. A color change from yellow to blue that extends to the triangular mark on the Dosimeter label indicates that a dose of Ethylene oxide sufficient for sterilization has been delivered. Without opening the Steritest pouch, manipulate the ampoule of culture broth inside of its break shield so that the neck of the ampoule is broken.

Gently shake the broth down to cover the spore disk. Incubate the Steritest at 37.5°C for 72 hours. A change in the color of the broth from blue to orange indicates growth of bacteria and therefore an unsterile load.

7.2 Disinfection
7.2.1 Disinfection is a process where most microbes are removed from defined object or surface, expect bacterial spores.

High level disinfection is that which kills all microganism and high number of bacterial spores.

7.2.2 Classification of Disinfectants
(a) High Level Disinfectants:
- They destroy all microorganisms including vegetative bacteria, most bacterial spores, fungi, viruses including enteroviruses and mycobacterium tuberculosis except some bacterial spores. Ex.: 2% Glutaraldehyde, Ethylene Oxide, 1%Sodium Hypochlorite (10,000ppm of chlorine)
- Used for semi critical instruments and equipments (those that are in contact with intact mucous membrane without penetration)
- For gastrointestinal endoscopes, endotracheal tubes, anesthesia breathing circuits, respiratory therapy equipments.

(b) Intermediate Level Disinfectants:
- They destroy vegetative bacteria, Mycobacterium tuberculosis, most viruses e.g. enteroviruses and fungi but not bacterial spores. Ex.: Isopropyl alcohol (70%), ethyl alcohol, sodium hypochlorite (0.1%), Chlorhexidine, hydrogen peroxide, phenolic solutions.

(c) Low Level Disinfectents:
- They destroy most vegetative bacteria, fungi and enveloped virus e.g. HIV but will not kill bacterial spores, Mycobacteria and non enveloped viruses like enterovirus. Ex: Quaternary ammonium compounds like benzylkonium chloride, some soaps.

7.2.4 Guidelines for Selection of Disinfectants:
There is no ideal disinfectant. Each application requires careful view of following:
- Type and number of organisms.
- Type and amount of organic matter
- Contact time
- Type of surface (Rough / Corrugated)
- Type of water (hard / soft)
- Manufacturers data on efficacy
- Safety and environmental aspects (chlorine is not free from toxicity)
- Cost, shelf life and convenience of use
- Residual activity

7.2.5 Two Approaches for Selection of Disinfectants:
1. Accept the manufacturers data
2. Validate yourself

7.2.6 Guidelines for Use of Disinfectants

<table>
<thead>
<tr>
<th>Name of Disinfectant</th>
<th>Method of Dilution</th>
<th>Contact Time</th>
<th>In Use Span/ Use</th>
</tr>
</thead>
</table>
1. **Aldehyde Solutions:**
   - a. **Glutaraldehyde (2%)**
     - Add activator powder / liquid to the liquid in 5 liter jar and use undiluted
     - Disinfection: 20-30 mins
     - Sterilization: 10 hours
     - 14 days used for heat sensitive instruments e.g. Endoscopes
   - b. **OPA (orthophthalyl aldehyde)**
     - Same as above
     - Long acting (28 days)
   - c. **Glutaraldehyde + Formaldehyde + Benzyl chloride**
     - water 1 part : 49 parts (20 ml + 980 ml)
     - Disinfection: 15 min
     - Sterilization: 5 hours , 30 min
     - Used as surface disinfectant or 2% solution in operation theaters and 0.5% in wards, dressing room. Can be used in a low pressure sprayer.
     - 14 days used for heat sensitive instruments e.g. Endoscopes

2. **(Glutaraldehyde + formaldehyde)**
   - 20 ml H₂O₂ + 80 ml normal saline = 6% H₂O₂ (use freshly prepared)
   - Disinfection: 15 min
   - Sterilization: 5 hours, 30 min
   - 14 days (used for instrument sterilization)

3. **6% Hydrogen Peroxide (Available as 30% stabilized solution)**
   - 20 ml H₂O₂ + 80 ml normal saline = 6% H₂O₂ (use freshly prepared)
   - 6-8 minutes
   - Use immediately after preparation for surgical dressings.

4. **1% Sodium Hypochlorite**
   - Ex.: Polar Bleach 5%
   - Polar Bleach 10%
   - 5%: 80 ml water + 20 ml bleach to make it 1% solution.
   - 10%: 90 ml water + 10 ml bleach
   - 20-30 minutes
   - 8 hours
   - Used for blood spills and laboratory decontamination

5. **Calcium hypochlorite**
   - Ex.: Bleaching powder (70% available chlorine)
   - 1.4 gms / liter of water for visibly contaminated articles
   - 20-30 min.
   - 24 hours
   - Disinfection of toilets, bathrooms and may be used if liquid bleach not available

6. **Formaldehyde (40%)**
   - Ex.: Formalin
   - Ready to use
   - 30 minutes
   - Then open the area after 6 hours
   - No longer recommended for fumigation.

7. **Formaldehyde (40%)**
   - Ex.: Formalin

8. **70% Alcohol**
   - Ready to use
   - 2-5 minutes
   - 24 hours used for surface disinfection

9. **Chlorhexidine (2%) w/v**
   - 4% Chlorhexidine w/v
   - Ready to use
   - 2-3 minutes
   - 2%: Upto 6-8 hours for disinfection of hands
   - 4%: Used before a procedure.

10. **Povidine Iodine 10%**
    - Ready to use
    - Allowed to dry
    - For skin preparation before surgery

11. **1% Triclosan**
    - Ready to use
    - Antiseptic soap or bathing liquid
    - For MRSA (Methicillin resistant Staphylococcus aureus)

12. **(2 propanol - 1 propanol, macetronium ethyl sulfate)**
    - Ready to use
    - 30 seconds
    - Hand rub

13. **(Stabilized H₂O₂ 11% w/v with 0.01% w/v diluted silver nitrate solution)**
    - 10 % w/v solution
    - 60 minutes
    - Surface disinfection
    - For fumigation

14. **(Glutaraldehyde + formaldehyde)**
    - 20 ml H₂O₂ + 80 ml normal saline = 6% H₂O₂ (use freshly prepared)
    - Disinfection: 15 min
    - Sterilization: 5 hours, 30 min
    - 14 days (used for instrument sterilization)

### 7.2.6 General Guidelines For Disinfection
7.2.6.1 Critical instruments /equipments - (that are those penetrating skin or mucous membrane or enter sterile tissue or vascular system) should undergo sterilisation before and after use. e.g. surgical instruments and implants.

7.2.6.2 Semi-critical instruments /equipments - (that are those in contact with intact mucous membrane without penetration or skin that is not intact) should undergo high level. e.g laryngoscopes, Anaesthesia equipment.

7.2.6.3 Non-critical instruments /equipments - (that are those in contact with intact skin and no contact with mucous membrane) requires only intermediate or low level disinfection before and after use.e.g. ECG electrodes

<table>
<thead>
<tr>
<th>Classification</th>
<th>Item Use</th>
<th>Goal</th>
<th>Appropriate Process</th>
</tr>
</thead>
<tbody>
<tr>
<td>Critical item</td>
<td>Items entering sterile tissue, the body cavity, the vascular system and non-intact mucous membranes E.g surgical instruments</td>
<td>Objects will be sterile (free of all microorganisms including bacterial spores)</td>
<td>Sterilization (or use of single use sterile products) (steam sterilization)</td>
</tr>
<tr>
<td>Semi-Critical items</td>
<td>Items that make contact, directly or indirectly, with intact mucous membranes or non-intact skin. E.g. endoscopes, anaesthetic equipments, Respiratory therapy Equipment Endocavitary probes Tonometer Diaphragm</td>
<td>Objects will be free of all microorganisms, with the exception of high numbers of bacterial spores</td>
<td>High level disinfection · Thermal disinfection · Chemical disinfection (glutaraldehyde, OPA) It is always preferable to sterilize semi-critical items whenever they are compatible with available sterilization processes.</td>
</tr>
<tr>
<td>Non-Critical items</td>
<td>Objects that come into contact with intact skin but not mucous membranes E.g crutches, BP cuffs, Tabletops Bed pans, bed rail, bedside table, ECG leads etc</td>
<td>Objects will be clean</td>
<td>Low level disinfection · Cleaning (manual or mechanical)</td>
</tr>
</tbody>
</table>

7.2.7 Instrument cleaning process

STEP 1 - Decontamination

- Decontaminate instruments and other items by placing them in a plastic container of 0.5% Hypochloride solution/Bleaching Solution. Let them soak for 10 minutes. A container of this solution should be kept in every operating theatre and procedure room, so that used items can be place directly into the bucket.
- Users should put instruments and other items into the solution as soon as they are finished using each item. Open or unlock jointed instruments, such as haemostats and scissors. Disassemble those instruments with sliding or multiple parts.
- After 10 minutes, remove the items from the Hypochlorite solution/Bleaching Solution and either rinse with water or clean immediately. Do not leave items in the solution for more than 10 minutes, since excessive soaking in the solution can damage instruments and other items. Always wear gloves when removing instruments and other items from a chlorine solution. Dried out instruments then can be taken for further processing.

STEP 1 has to be performed at User area. All other steps to be performed at CSSD.

STEP 2- Primary Cleaning

- Cleaning is the removal of foreign material (e.g., soil, and organic material) from objects and is normally accomplished using water with detergents or enzymatic products.
Thorough cleaning is required before high-level disinfection and sterilization because inorganic and organic materials that remain on the surfaces of instruments interfere with the effectiveness of these processes.

If soiled materials dry or bake onto the instruments, the removal process becomes more difficult and the disinfection or sterilization process less effective or ineffective.

Surgical instruments should be pre-soaked or rinsed to prevent drying of blood and to soften or remove blood from the instruments.

### 7.2.8 Steps of Cleaning

Always wear utility gloves, a mask, and protective eyewear when cleaning instruments and other items. Avoid using steel wool or abrasive cleansers. These products can scratch or pit metal or stainless steel, resulting in grooves that can become a nesting place for microorganisms. This also increases the potential for corrosion of the instruments and other items.

**Step 1**
**Decontamination**

**Step 2**
Using a soft brush or old toothbrush, detergent, and water, scrub instruments and other items vigorously to completely remove all blood, other body fluids, tissue, and other foreign matter. Hold items under the surface of the water while scrubbing and cleaning to avoid splashing. Disassemble instruments and other items with multiple parts, and be sure to brush in the grooves, teeth, and joints of items, where organic material can collect and stick.

**Step 3**
Rinse items thoroughly with clean running water to remove all detergent. Any detergent left on the items can reduce the effectiveness of further chemical processing.

**Step 4**
Allow items to air-dry (or dry them with a clean towel).

Note: Instruments that will be further processed with chemical solutions must dry completely to avoid diluting the chemicals; items that will be high-level disinfected by boiling do not need to be dried first.

### 7.2.9 Endoscopes - cleaning and disinfection

1. **Mechanical cleaning:** This is the most important step. Flush the air/water channel for 10-15 seconds to eject any blood or mucus. Aspirate detergent through the biopsy/suction channel to remove gross debris. Use a cleaning brush suitable for the instrument and channel size to brush through the suction channel.

2. **Disinfection:** The endoscope and all internal channels are soaked in 2% Glutaraldehyde for 20 minutes.

3. **Rinsing:** Following disinfection, rinse the instrument internally and externally to remove all traces of disinfectant.

4. **Drying:** Dry the endoscope externally. Flush air through each channel.

5. Store: store the endoscope in a way that prevents recontamination and promotes drying (e.g., hung vertically).
6. Monitoring: Monitoring of disinfection procedure of endoscope is done on regular basics (through round sheet) and disinfectant is checked on regular basic.

7.3 Decontamination

This encompasses cleaning, disinfecting and sterilizing of equipment/device:

7.3.1 Decontamination Procedure for Equipment

Pre-cleaning of any item / medical device is an essential step prior to disinfection

<table>
<thead>
<tr>
<th>Article</th>
<th>Standard Procedure</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ambubag</td>
<td>Should be cleaned with detergent and water, dried and sterilized.</td>
<td></td>
</tr>
<tr>
<td>Applanator (Tonometer Prisms)</td>
<td>Immersion in 0.05% hypochlorite (500 parts per million available chlorine) for 10 minutes.</td>
<td>A fresh solution should be prepared at the start of each clinic.</td>
</tr>
<tr>
<td>Arterial catheters</td>
<td>Sterile, single use only, must be discarded after use.</td>
<td></td>
</tr>
<tr>
<td>Baby equipment feeding bottles &amp; teats</td>
<td>1. Disposable – single use. 2. Re-usable – should be returned to CSSD or washed in hot detergent and water, rinsed and immersed in Milton fluid, freshly made up from tablets according to manufacturer's instructions.</td>
<td>Should be soaked for a minimum of 1 hour.</td>
</tr>
<tr>
<td>Baby weighing scales</td>
<td>A fresh liner should be used for each baby. Clean tray as necessary with detergent and water.</td>
<td>If contaminated should be wiped with hypochlorite 1000ppm after washing.</td>
</tr>
<tr>
<td>Baby bath</td>
<td>Should be cleaned after each use with detergent and water</td>
<td></td>
</tr>
<tr>
<td>Beds and couches Frame</td>
<td>Should be cleaned with detergent and water between patients and as required</td>
<td>If contaminated with body fluids, see spillage policy. If used in isolation room after cleaning, should be wiped with a disinfectant</td>
</tr>
<tr>
<td>Mattresses and pillows</td>
<td>Should be cleaned with detergent and water between patients and as required</td>
<td>If contaminated with body fluids, the blood spills policy should be implemented. Should not be used if cover is damaged. Contaminated pillows must be discarded. Torn mattress covers must be replaced before mattress is re-used.</td>
</tr>
<tr>
<td>Bedpans and urinals</td>
<td>Should be cleaned and disinfected with 0.5% sodium hypochlorite or hot water. It must be ensured that the item is dry before re-use.</td>
<td></td>
</tr>
<tr>
<td>Breast pumps</td>
<td>Should be washed with detergent and water, immersed in sodium hypochlorite, freshly made up from tablets according to</td>
<td></td>
</tr>
<tr>
<td>Item</td>
<td>Instructions</td>
<td>Manufacturer's Instructions</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>------------------------------------------------------------------------------</td>
<td>-----------------------------------------------</td>
</tr>
</tbody>
</table>
| Brushes Nail Toilet         | 1. Disposable – single use.  
2. Re-usable-to be returned to CSSD after each use.  
Should be rinsed well in flush water and stored dry. | Should not be left on sink after use.          |
| Carpets                    | Vacuum daily                                                                 | Should be shampooed or steam cleaned in isolation rooms as part of terminal cleans. |
| Commodes                   | Seat and arms should be cleaned with detergent and water, and dried.         | If soiled or used in isolation, should be wiped with sodium hypochlorite 2% and dried, after cleaning |
| Cradles                    | Should be cleaned with detergent and water and dried.                        |                                               |
| Crockery and cutlery       | Should be heat disinfected in dishwasher.  
If washed in sink with water and detergent |                                               |
| Curtains                   | Should be changed as part of a rolling program by domestic services.        | Should be changed as part of terminal clean.  |
| Denture pots               | 1. To be cleaned by patients themselves with detergent and water  
2. Disposable with lid-single use. |                                               |
| Drainage bottles           | 1. Disposable – single use  
2. reusable- rinse and return to CSSD                                      |                                               |
<p>| Drip Stands                | Should be cleaned with detergent and water and dried.                        | After use in isolation, should be wiped with sodium hypochlorite 2% and dried after cleaning. |
| Ear Pieces for auroscope   | Should be cleaned with detergent and water and dried.                        | To be returned to CSSD after use in isolation. |
| Earphones                  | Should be cleaned with detergent and water and dried.                        | Foam should be replaced after use in isolation. |
| Leads and monitors         | Should be dismantled to smallest components and cleaned with detergent and water and dried. |                                               |
| Eye protection             | Should be cleaned with detergent and water and dried.                        | For blood splashes blood spillage policy should be followed. |
| Floors                     | Should be vacuumed daily. A damp mop with detergent and water should be used. | For blood splashes blood spillage policy should be followed. |
| Flower vases               | Should be cleaned with detergent and water and dried.                        |                                               |
| Furniture                  | Should be damp dusted with detergent and water.                              |                                               |
| Humidifiers                | Should be cleaned and sterilized at low temperature.                         |                                               |
| Incubators                 | Should be cleaned with detergent and water and switch on to dry.             | Terminal sterilization with ethylene oxide gas may be required after some infections. |</p>
<table>
<thead>
<tr>
<th>Equipment</th>
<th>Cleaning Instructions</th>
<th>Sterilization Instructions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intravenous monitoring pumps</td>
<td>Should be cleaned with detergent and water and dried.</td>
<td>After use in isolation wipe with sodium hypochlorite 2% and dry, after cleaning</td>
</tr>
<tr>
<td>(and feed pumps)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Instruments</td>
<td>After single use to be returned to CSSD</td>
<td></td>
</tr>
<tr>
<td>Linen</td>
<td>Should be soaked in hot water, returned to laundry</td>
<td></td>
</tr>
<tr>
<td>Mops</td>
<td>Disposable use for one day. Re-usable to be laundered in washing machine.</td>
<td>Mops must not be stored wet or cleaned in disinfectant solutions.</td>
</tr>
<tr>
<td>Peak flow</td>
<td>Disposable – single patient use.</td>
<td></td>
</tr>
<tr>
<td>Nebulizers</td>
<td>Cleaning and low temperature sterilization.</td>
<td></td>
</tr>
<tr>
<td>Pressure relieving devices</td>
<td>Should be clean with detergent and water and dried.</td>
<td></td>
</tr>
<tr>
<td>Proctoscopes</td>
<td>Disposable - single use, re-usables to be rinsed and returned to CSSD.</td>
<td></td>
</tr>
<tr>
<td>Raised toilet seats</td>
<td>Should be cleaned after each use with detergent.</td>
<td></td>
</tr>
<tr>
<td>Razors</td>
<td>Safety – single use disposable Electric – patients own. Detachable head and clean with 70% isopropyl alcohol swab.</td>
<td></td>
</tr>
<tr>
<td>Shaving brush</td>
<td>Should not be used, unless supplied by the patients for their own use.</td>
<td></td>
</tr>
<tr>
<td>Skin disinfection</td>
<td>Showers are preferred to bath or bed baths.</td>
<td></td>
</tr>
<tr>
<td>Soap dispensers</td>
<td>Should be cleaned weekly with detergent and water and dried.</td>
<td></td>
</tr>
<tr>
<td>Sphygmo-manometer cuffs</td>
<td>After use in isolation, should be laundered in washing machine.</td>
<td></td>
</tr>
<tr>
<td>Spillages</td>
<td>Should be cleaned with detergent</td>
<td></td>
</tr>
<tr>
<td>Sputum pots</td>
<td>Disposable with close fitting lid. Should be discarded into clinical waste for incineration.</td>
<td></td>
</tr>
<tr>
<td>Stethoscopes</td>
<td>Should be cleaned with detergent and water and dried. Should be wiped with 70% alcohol.</td>
<td></td>
</tr>
<tr>
<td>Suction bottles</td>
<td>Disposal liners. Must be sealed when 75% full and placed in yellow plastic bag. Re-usable, should be cleaned with sodium hypochlorite and dried. Must be changed daily and in between each patient. To be stored dry when not in use.</td>
<td></td>
</tr>
<tr>
<td>Telephones</td>
<td>To be wiped with 70% alcohol.</td>
<td></td>
</tr>
<tr>
<td>Thermometers</td>
<td>To be covered with disposable sleeve before use and stored dry in individual holder. In</td>
<td></td>
</tr>
</tbody>
</table>
between patients, should be cleaned and wiped with 70% isopropyl alcohol (swab). If disposable sleeve not used in between patients, should be washed in general purpose detergent and tepid water then wiped with 70% alcohol (swab). To be stored in individual holder inverted.

<table>
<thead>
<tr>
<th>Toilet seats</th>
<th>To be cleaned at least twice daily with detergent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Toys</td>
<td>Toys should be cleaned with detergent and water and dried. For isolated patients, toys that cannot be decontaminated to be avoided. Heavily contaminated toys may have to be destroyed.</td>
</tr>
<tr>
<td>Trolley (Dressing)</td>
<td>To be cleaned daily with detergent and water. After each use should be wiped with 70% isopropyl alcohol.</td>
</tr>
<tr>
<td>Urine measuring jugs</td>
<td>To be heat disinfected after each use in bed pan washer.</td>
</tr>
<tr>
<td>Ventilators</td>
<td>To be sent to respiratory therapy unit.</td>
</tr>
<tr>
<td>Vomit bowls</td>
<td>Contents must be emptied into sluice then rinsed and washed and disinfected with hot water and detergent.</td>
</tr>
<tr>
<td>Walls</td>
<td>Should be cleaned with detergent and water as part of planned preventive maintenance program.</td>
</tr>
<tr>
<td>Wash bowls</td>
<td>Patients must have own dedicated bowl. After each patient's use, should be cleaned with detergent.</td>
</tr>
<tr>
<td>Wheel chairs</td>
<td>Patient's own – should be cleaned with detergent and water as necessary. Hospital – clean between patients with detergent and water</td>
</tr>
</tbody>
</table>
8. CARE OF SYSTEMS AND INDWELLING DEVICES

8.1 General Guidelines
To be followed for all procedures:
1. Hand washing is mandatory before, after and in-between procedures and patients.
2. Each health care worker are familiar with the personal protection (Universal precautions) required for each procedure. These precautions are strictly adhered to.
3. Follow proper waste segregation & disposal after each procedure.

8.2 Vascular Care
8.2.1 Hand washing
Wash hands before every attempted intravascular catheter insertion. Antimicrobial handwashing soaps are desirable, and are preferred before attempted insertions of central intravenous catheters, catheters requiring cut downs, and arterial catheters.

8.2.2 Preparation of skin
Povidone-iodine (PVP) or 70% alcohol may be used for cleaning the skin. Insertion sites are scrubbed with a generous amount of antiseptic. Beginning at the centre of the insertion site, use a circular motion and move outward. Antiseptics should have a contact time of at least 30 seconds prior to catheter insertion.

Antiseptics should not be wiped off with alcohol prior to catheter insertion.

8.2.3 Applying dressings
Sterile dressings are applied to cover catheter insertion sites. Unsterile adhesive tape should not be placed in direct contact with the catheter-skin interface.

8.2.4 Inspecting catheter insertion sites
Intravascular catheters are inspected daily and whenever patients have unexplained fever or complaints of pain, tenderness, or drainage at the site for evidence of catheter related complications. Inspect for signs of infection (redness, swelling, drainage, tenderness) or phlebitis and also palpate gently through intact dressings.

8.2.5 Manipulation of intravascular catheter systems
Strict aseptic techniques are maintained when manipulating intravascular catheter systems. Examples of such manipulations include the following:
- Placing a heparin lock
- Starting and stopping an infusion
- Changing an intravascular catheter site dressing
- Changing an intravascular administration set

8.2.6 Flushing IV lines
Solutions used for flushing IV lines should not contain glucose which can support the growth of microorganisms. Do not reuse syringes used for flushing. One syringe is used for flushing only one IV line once.

8.2.7 Peripheral IV sites (short term catheters)
- **Dressing changes**
  - Peripheral IV site dressings should not usually require routine changes, since peripheral IV catheters are removed within 72 hours.
- **Replacement of Peripheral IV Catheters**
  - Peripheral IV catheters are removed 72 hours after insertion, provided no IV-related complications, requiring catheter removal are encountered earlier. A new peripheral IV catheter, if required, may be inserted at a new site.

8.2.8 Central intravascular catheters (long term catheters)
- **Dressing changes**
  - Central IV catheter dressings are changed every 72 hours.
- **Replacement of central IV catheters**
  - Central IV catheters do not require routine removal and reinsertion. The catheter can be kept for a maximum of 3 months, provided there is no sign of catheter related infection or other complications.

8.2.9 Catheter related Infection
At the time of catheter removal, the site is examined for the presence of swelling, erythema, lymphangitis, increased tenderness and palpable venous thrombosis. Any antimicrobial ointment or blood present on the skin around the catheter is first removed with alcohol. The catheter is withdrawn with sterile forceps, the externalized portion being kept directed upward and away from the skin surface.
(If infection is suspected, after removal, the wound is milked in an attempt to express purulence. For 5.7 cm catheters, the entire length, beginning several millimeters inside the former skin surface catheter interface, is aseptically cut and sent for culture. With longer catheter, (20.3 cm and 60.9 cm in length), two 5-7 cm segments are cultured a proximal one beginning several millimeters inside the former skin catheter interface and the tip. Catheter segments are transported to the laboratory in a sterile container.)

Three way with extension is used only when multiple simultaneous infusates or Central Venous Pressure monitoring are required.

8.3 Respiratory Care

In addition to the general guidelines that are to be adhered to, the following should also be noted with regard to respiratory care:

Mouth flora influences development of healthcare associated pneumonia in ventilated patients. Frequent chlorhexidine mouthwashes minimise the chances of pneumonia.

32.12.3.1 Ventilator Care

- Sterile water is to be used in nebulizers and humidifiers. This are replaced once or twice a day.
- Pneumatic circuits (masks, Y connection and tubes) are to be changed every 24-48 hours. Condensate in tubing should not be drained into the humidifier or airway as they contain large numbers of pathogenic organisms. This are drained only into water traps. Use disposable circuits if cost permits.
- Use heat and moisture exchanging filter (HMEF) at Y connection for all patients if feasible and cost permits. Heat and moisture exchanging filter (HMEF) is to be changed every 24-48 hours. It should not be removed from circuit except at the time of changing.
- Oxygen masks, venture devices and nebulizer chambers are cleaned carefully and then send to CSSD for HLD.
- Humidifier domes are periodically send to CSSD. Ambu bags are cleaned thoroughly and periodically send to CSSD for HLD.
- Microbiological surveillance of respiratory therapy equipment is practised in our hospital.

8.3.2 Tracheostomy Care / Endotracheal Tube

- Careful attention to post-operative wound care is mandatory.
- The patient should receive aerosol therapy to prevent dessication of the tracheal and bronchial mucosa or the formation of crusts. The skin around the tracheostomy tube is cleaned with betadine (Povidone-iodine 5%) every four hours or more frequently, if necessary.
- In case of metal tracheostomy tubes, the inner cannula is cleaned every four hours and more often if necessary to prevent the formation of crusts. The inner cannula is cleaned with water, immersed in hydrogen peroxide for 15 minutes and then rinsed with fresh & sterile normal saline. The plastic tracheostomy tubes are removed, another plastic tube is inserted, and the tube is cleaned, with hydrogen peroxide, and rinsed well before reuse.
- The tracheostomy tape securing the tube are changed every 24 hours. This tape must be tied securely at all times.
- The first complete tube change are performed no earlier than 4-5 days to allow time for the tract to be formed. Subsequent changes are done weekly or as necessary.
- Clean technique is used to change the tracheostomy tube unless there is a medical indication for sterile technique.
- The obturator are at the bedside (preferably taped to the head of the bed) to be used if the tracheostomy tube accidently is dislodged or is removed for any reason.

8.3.3 Suctioning of endotracheal / tracheostomy tube

Employees are instructed and supervised by trained personnel in proper technique before performing this procedure on their own. Assess the patient using auscultation and vital signs prior to suctioning.

8.3.3.1 Sterile Suctioning

1. Wash your hands.
2. Use a catheter with a blunt tip.
3. The wall suction are set no higher than 120 mm Hg for adults and between 60 and 80 mm Hg for children.
4. Attach the suction catheter to the suction tubing; do not touch the catheter with bare hands (leave it in its protective covering).
5. Put on sterile gloves. The wearing of a mask is also strongly recommended.
6. However, if saline does need to be instilled, ½ cc of sterile saline is put into the tracheostomy tube on inspiration only.
7. If on a respirator, pre-oxygenate the patient by connecting the resuscitation bag to the artificial airway and ventilating the patient with three or four deep breaths. A mechanical ventilator on 100% oxygen may also be used by depressing the manual ventilation button three or four times.
8. Insert the catheter gently through the inner cannula until resistance is met. Do not apply suction during insertion.
9. Withdraw the catheter approximately 1 cm and institute suctioning.
10. Carefully withdraw the catheter, rotating it gently between the thumb and forefinger applying intermittent suctioning.
11. Continuous suctioning for longer than 10 seconds may create an unacceptable level of hypoxia.
12. The patient are given time to rest between suctioning episodes. If possible, this time are from two to three minutes. If the patient is receiving oxygen or ventilatory support, reapply the oxygen or ventilator for at least two minutes before re-suctioning.
13. Observe for unfavourable reactions such as increased heart rate, hypoxia, arrhythmia, hypotension, cardiac arrest, etc.
14. If oral suctioning is necessary, it are done after the tracheostomy is suctioned.
15. When suctioning is completed, clear the catheter and tubing of mucous and debris with sterile water or saline.
16. Discard the catheter, water container, and gloves appropriately.
17. Wash hands.
18. The tubing and suction canister are changed every 24 hours. The canister are labeled with the date and time when they are changed. If debris adheres to the side of the tubing or the canister, either or both are changed. The tubing are secured between suctioning periods so that it will not fall to the bed, floor, etc.

8.4 Urinary Catheter
8.4.1 Urethral catheterization
Personnel
Only persons who know the correct technique of aseptic insertion and maintenance of catheters should handle catheters.
Catheter Use
Urinary catheters are inserted only when necessary and left in place only as long as medically necessary and are changed after 7 days.
Hand hygiene
Hand hygiene is performed immediately before and after any manipulation of the catheter site or apparatus.
Catheter Insertion
Catheters are inserted using aseptic technique and sterile equipment. Use an appropriate antiseptic solution for periurethral cleaning. As small a catheter as possible, consistent with good drainage, are used to minimize urethral trauma. Indwelling catheters are properly secured after insertion to prevent movement and urethral traction.
Anchoring the catheter
Strapping of the catheter is done to the lower anterior abdominal wall in male patients. This is to prevent direct transmission of the weight of the bag on the catheter, so that pulling and inadvertent dislodgment of the catheter does not occur. This also helps to prevent stricture of the penile urethra if the patient is on a catheter for a long duration.

8.5 Wound Care
Surgical wounds
- Surgical wounds after an elective surgery are inspected on the third post-operative day, or earlier if wound infection is suspected.
- All personnel doing dressings should wash their hands before the procedure. Ideally, a two member technique is followed. One to open the wound, and one to do the dressing.
- If two health care workers are not available, then, take off the dressing, wash hands again before applying a new dressing.
- A clean, dry wound may be left open without any dressing after inspection.
- If there is any evidence of wound infection, or purulent discharge, then dressings are done daily, using povidone-iodine to clean the wound and applying dry absorbent dressings.
8.5.1 Collection of wound swabs
The superficial wound site:
1. Wound site should be gently washed with sterile saline. This process helps removing the colonizers from the wound.
2. Sampling should be done from the margin and floor of the wound to maximize the microbiological yield.
3. Paired wound swab (one for gram staining and another for culture) should be sent.

8.5.2 In deep wound site with pus discharge/ooze
1. Wound site should be gently washed with sterile saline. This process helps removing the colonizers from the wound.
2. The pus/discharge should be actively expressed and collected on the cotton swab.
3. Paired wound swab (one for gram staining and another for culture) should be sent.

All specimens should be sent immediately with the laboratory request clearly mentioning of wound site, body side (in case of limbs/face), diagnosis and surgical procedure (if any) undertaken.
9. SPECIAL CARE UNITS

9.1 Intensive Care Units
9.1.1 Design of the Unit
- Space around and between beds are adequate for placement and easy access to equipment and to patients (6-8 feet)
- A single, closed cubicle is used only for patients needing isolation; e.g. open tuberculosis, anthrax, enteric fever, cholera, MRSA colonization or infection with other multi-drug resistant organisms.
- Good housekeeping practices are followed. This includes regular cleaning of all areas, maintenance, linen and curtain changes etc. Clean floor at least four times a day.

9.1.2 Procedures to be followed by health care personnel
- Hand washing: Importance of this need not be over-emphasized in the ICU setting. Five moments of hand hygiene must be complied with hand hygiene actions. Appropriate steps must be performed while doing hand hygiene.
- Standard Precautions: as appropriate, are followed by all staff while handling patients or samples. Wear plastic aprons and gloves for all procedures. Remove and discard them immediately after each patient. Use gloves for all patient contact. Wear masks while examining patients with 'uncertain' diagnosis.

9.1.3 Instruments
Although disposable items are ideal, reusable items are often used, for reducing the cost. Separate thermometers are used for each patient or must be disinfected before reuse in other patients. Separate AMBU bag and mask are used for each patient. These must be reused after proper disinfection procedures in CSSD. Trolleys are to be adequately loaded and are used for bedside procedures.

9.1.4 Microbiological monitoring
Environmental surveillance will be done as per guidelines for high risk areas mentioned in chapter 3. Passive surveillance will be used to detect healthcare associated outbreaks.

9.1.5 Visitors policy
Minimum Visitors are allowed inside intensive units for control of infections.

9.2 Dialysis Unit
The purpose of this policy is to optimize the treatment and minimize the risk of the transmission of infections from patient to patient and between patients and employees.

To prevent cross infection following disinfection and equipment maintenance should be done as per provisions in Schedules.

9.2.1 Haemodialysis machines:
- Priming of kit (Haemodialyser and Arteriovenous tubing) should be done thoroughly with Normal saline without coming in contact with the floor surface and priming bucket surface area.
- Kit has to be kept in recirculation mode by connecting Hansen connectors to dialyzer and giving 2000 IU inj. Heparin.
- Machine should be disinfected with 4% sodium hypochlorite/citric acid on daily basis.
- Bleaching of machines should be done with 5% chlorine once a month
- Conductivity of the haemodialysis machine shall be monitored by lab method on a weekly basis
- Dialysate sterility should be checked on a monthly basis
- Calibration of machines should be undertaken on a quarterly basis

9.2.2 RO Unit
RO maintenance should be done on weekly basis by regeneration of softener and giving backwashes. Disinfection of RO unit including loop lines and storage tanks should be done using 1% sodium hypochlorite solution on a monthly basis.

The following tests on the RO unit output water should be undertaken:
- Conductivity: Daily
- Hardness test: Once/week
- Chloramine test: Once/week
- Culture: Once/month
- Endotoxin Assay: Once/month
A detailed examination of RO water should be undertaken on quarterly basis as per AAMI guidelines.

9.2.3 Reprocessor machine:
- Reprocessing machine should be sanitized with sodium hypochlorite on a weekly basis
- Ends of dialyzer connectors should be dipped in disinfectant solution after every process
- Fibre Bundle Volume and number of times Haemodialyser was being used should be recorded
- Haemodialyser kits should be stored in separate boxes for multiple uses

9.2.4 Blood lines and multidose vials should not be re-used

9.2.5 Staff members shall be vaccinated properly and proper care needs to be taken reardin isolation to prevent cross infection

9.2.6 Log of disinfection activities should be maintained for verification.

9.2.1 Disinfection Schedule for Hemodialysis
- Disinfection of HD machine with Hemoclean.
- Hot disinfection of HD machine with calfree: After every dialysis.
- Front cleaning of HD machine with Hemoclean:
- Disinfection/ washing of R.O. inlet filter of H.D. Machine with Hemoclean:
- Disinfection of R.O tank with hemoclean: 1st week of every month.
- Charging of R.O system: as per the recommendations.
- Culture of dialysate & R.O water: 1st week every month.
- Washing Biocarbonate container: After every dialysis.
- Carbolization of Hemodialysis room: Daily.
- Changing glutaradehyde container : Every 14 Days.
- Washing of H.D. Room : 1st week every month.
- Fumigation of H.D. Room with (Hydrogen peroxide+ Silver nitrate) e.g. Ecoshield: 1st week every month.

9.2.2 Catheter Infection on Treatment

a) Localized Exit Site Infection:
Erythema or crust but no purulent discharge, it can be treated with local applicator of antibiotics.

b) Septicemia Infection:
Fever with chills at the initiation of the dialysis. Two set of blood samples with culture, with atleast one drawn percutaneous site and other through the catheter are obtained in the case of CRBSI (Catheter related Blood Stream Infection). Prophylactic antibiotic (Ceftazidime and Amikacin) are started and also take a sample for Blood culture. Antibiotics will be discontinued if the blood culture has no growth and antibiotic regimen adjusted only when bacterial sensitivity is available. Antibiotics are continued in uncomplicated case of catheter related bacteria.

9.2.3 Specimen Collection and Handling
- Extreme caution must be employed when drawing blood for laboratory testing. Gloves and face shields will be worn while drawing specimens.
- Blood spills will be cleaned immediately with solution of bleach. During cleaning, gloves will be worn.
- Any specimen collected from a patient on Isolation is labelled according to Infection Control policy.
- Bacterial monitoring of water for preparing dialysis fluids and dialysate fluid are collected and immediately sent to Microbiology department on a monthly basis.
- Specimens are clearly labelled and should include the following information: initials of person collecting specimen, date, time, specimen source (i.e., dialysate fluid or dialysis water), and the machine from which the source was collected.

9.2.4 Environment
The environment shall be thoroughly cleaned between each treatment and as necessary for spills of blood and body fluids.
Terminal cleaning procedures must be used between the patients.
10. HOUSE KEEPING

10.1 House Keeping In Wards

10.1.1 Patient Care Environment Cleaning
A patient admitted to the hospital can develop infection due to bacteria that survive in the environment. Therefore, it is important to clean the environment thoroughly on a regular basis. This will reduce the bacterial load and make the environment unsuitable for growth of micro-organisms.

1. The floor is to be cleaned at least twice times in 24 hours. Detergent and copious amounts of water are used during cleaning.
2. The walls are to be washed with a brush, using detergent and water once a week.
3. High dusting is to be done with a wet mop.
4. Fans and lights are cleaned with soap and water once a month.
5. All work surfaces are to be disinfected by wiping with appropriate disinficant and then cleaned with detergent and water twice a day.
6. Cupboards, shelves, beds, lockers, IV stands, stools and other fixtures are to be cleaned with detergent and water once a week.
7. Curtains are to be changed once a month or whenever soiled. These curtains are to be sent for regular laundering. In certain areas, eg. Transplant units and ICUs, more frequent changes are required.
8. Patient's cot is to be cleaned every week with detergent and water. 1% hypochlorite to be used when soiled with blood or body fluids. In the isolation ward, cleaning is done daily.
9. Store rooms are to be mopped once a day and high dusted once a week.
10. The floor of bathrooms is to be cleaned with a broom and detergent once a day and then disinfected.
11. Toilets are cleaned with a brush using a detergent twice a day (in the morning and evening). Disinfection and stain removal solution may be used.
12. Wash basins are to be cleaned every morning.
13. Regular AC maintenance is required. The AC section should draw up a protocol for this.

10.1.2 Patient linen
- Bed linen is to be changed daily and whenever soiled with blood or body fluids.
- Patient's gown is to be changed every day and whenever soiled with blood or body fluids.
- Dry dirty linen is to be sent to the laundry for regular wash.
- Linen soiled with blood or body fluids, and all linen used by patients diagnosed to have HIV, HBV, HCV and MRSA, are send in red bag to the laundry.

10.1.3 Miscellaneous items
- Kidney basins, basins, bed pans, urinals, etc to be cleaned with detergent and water and disinfected with 0.5% hypochlorite solution.

10.2 House Keeping In The Operation Theatre
Theatre complex are absolutely clean at all items. Dust should not accumulate at any region in the theatre.
Soap solution is recommended for cleaning floors and other surfaces. Operating rooms are cleaned daily and the entire theatre complex is cleaned thoroughly once a week.

10.2.1 Environment

STEPS to be followed for maintenance of the housekeeping in O.T.

10.2.1.1 Before the start of the 1st case
Wipe all equipment, furniture, room lights, suction points, OT table, surgical light reflectors, other light fittings, slabs etc with soap or disinfecant solution(2% Bacillocid). This is completed at least one hour before the start of surgery.

10.2.1.2 Between two surgeries
- Spill- Clean spills with a 0.5% bleaching solution.
- Wipe OT table, surgical light reflectors, slabs etc are disinfected with sterisol 1% or with available disinfecant solution.
- Instrument tables (trolley Mayo stands & other flat surfaces. Wipe all flat surfaces that have come in immediate contact with a patient or body fluids with a disinfecant cleaning solution.
- Waste- Collect and remove all waste from the operating room in closed leak proof containers.
- Sharps containers. Close and remove containers from the operating room when they are three quarters full.

10.2.1.3 After the last case

The same procedure as mentioned above is followed and in addition the following are carried out.
- Wipe over heads light, cabinets, waste receptacles, equipments, furniture with ecoshield.
- Wash floor and wet mop with liquid soap and then remove water and wet mop with Bacilloflor solution.
- Clean the storage shelves scrub & clean sluice room.

10.2.1.4 Linen & gloves

Gather all soiled linen and towels in the receptacles provided. Take them to the service corridor (behind the theatre) and place them in trolleys to be taken for sorting. The dirty linen is then sent to the laundry. Use gloves while handling dirty linen.

10.2.1.5 Instruments

Used instruments are cleaned immediately by the scrub nurse and the Nursing Orderly. Reusable sharps are decontaminated in Lysol / hypochlorite and then washed in the room adjacent to the respective OR by scrubbing with a brush, liquid soap and vim. They are then sent for sterilization in the CSSD. After septic cases the instruments are sent in the instrument tray for autoclaving. Once disinfected, they are taken back to the same instrument cleaning area for a manual wash described earlier. They are then packed and re-autoclaved before use.

10.2.1.6 Weekly cleaning procedure
- Remove all portable equipment.
- Damp wipe lights and other fixtures with detergent.
- Clean doors, hinges, facings, glass inserts and rinse with a cloth moistened with detergent.
- Wipe down walls with clean cloth mop with detergent.
- Scrub floor using detergent and water or Bacilloflor.
- Stainless steel surfaces – clean with detergent, rinse & clean with warm water.
- Replace portable equipment: Clean wheel castors by rolling across towelling saturated with detergent.
- Wash (clean) and dry all furniture and equipment (OT table, suction holders, foot & sitting stools, Mayo stands, IV poles, basin stands, X-ray view boxes, hamper stands, all tables in the room, holes to oxygen tank, kick buckets and holder, and wall cupboards)
- After washing floors, allow disinfectant solution to remain on the floor for 5 minutes to ensure destruction of bacteria (Bacilloflor)

10.2.1.7 Maintenance and Repairs
- Machinery and equipment are checked, cleaned and repaired routinely
- Urgent repairs are carried out at the end of the days list
- Air conditioners and suction points are checked, cleaned and repaired on a weekly basis.

Preventive maintenance on all theatre equipment to be carried out weekly and major work to be done at least once every year.

10.3 Mopping Schedule for Various Departments

10.3.1 ICU, Dialysis, HDU
Floors should be mopped in each shift with detergent and water.

10.3.2 OPD, LABS and Wards
Floor should be mopped at least thrice daily with soap and water.

10.3.3 CSSD
Floor should be mopped twice daily with soap and water

10.3.4 OT
Floors should be mopped with Bacillocid after each surgery

10.4 Bio-Medical Waste collection schedule
- Segregated BMW is Collected twice daily from each departments except laboratories. In laboratories BMW is collected three times in a day.
11. INVESTIGATION OF AN OUTBREAK

The occurrence of two or more similar cases relating to place and time is identified as a cluster or an outbreak and needs investigation to discover the route of transmission of infection, and possible sources of infection in order to apply measures to prevent further spread. If the cases occur in steadily increasing numbers and are separated by an interval approximating the incubation period, the spread of the disease is probably due to person to person spread. On the other hand if a large number of cases occur following a shared exposure e.g an operation, it is termed a common source outbreak, implying a common source for the occurrence of the disease.

11.1 Epidemiological Methods
The investigation of an outbreak may require expert epidemiological advice on procedures. Formulation of a hypothesis regarding source and spread is made before undertaking microbiological investigations in order that the most appropriate specimens are collected.

32.15.1.1 Steps to be taken for investigation of an outbreak

**Step 1**
- Recognition of the outbreak. Is there an increase in the number of cases of a particular infection or a rise in prevalence of an organism? Such findings indicate a possible outbreak.
- Preliminary investigation must be begun by developing a case definition, identifying the site, pathogen and affected population. Define the outbreak in time person and place.
- Determination of the magnitude of the problem and if immediate control measures are required. If so general control measures such as isolation or cohorting of infected cases; strict hand washing and asepsis are immediately applied.
- Verification of the diagnosis. Each case are reviewed to meet the definition.
- Confirmation that an outbreak exists by comparing the present rate of occurrence with the endemic rate are made.

**Step 2**
- The appropriate departments and personnel and the hospital administration are notified and involved.

**Step 3**
- Additional cases must be searched for by examining the clinical and microbiological records.
- Line listings for every case, patient details, place and time of occurrence and infection details are developed.
- An epidemic curve based on place and time of occurrence are developed, the date analyzed, the common features of the cases e.g age, sex, exposure to various risk factors, underlying diseases etc. are identified.
- A hypothesis based on literature search and the features common to the cases; are formulated to arrive at a hypothesis about suspected causes of the outbreak.
- Microbiological investigations depending upon the suspected epidemiology of the causative organism are carried out. This will include (a) microbial culture of cases, carriers and environments (b) epidemiological typing of the isolates to identify clonal relatedness.
- The hypothesis is tested by reviewing additional cases in a case control study, cohort study, and microbiological study.

**Step 4**
- Specific control measures are implemented as soon as the cause of outbreak of identified.
- Monitoring for further cases and effectiveness of control measures are done.
- A report are prepared for presentation to the HICC, departments involved in the outbreak and administration.

11.2 Immediate Control Measures
Control measures are initiated during the process of investigation. An intensive review of infection control measures is made and general control measures initiated at once. General measures include:
- Strict hand washing ;
- Intensification of environmental cleaning and hygiene.
- Adherence to aseptic protocols, and
- Strengthening of disinfection and sterilization.

11.3 Microbiological Study
- Microbiological study is planned depending upon the known epidemiology of the infection problem. The study is carried out to identify possible sources and routes of transmission. The
investigation may include cultures from other body sites of the patient, other patients, staff and environment. Careful selection of specimens to be cultured is essential to obtain meaningful data.

11.4 Specific Control Measures
✓ Specific control measures are instituted on the basis of nature of agent and characteristics of the high-risk group and the possible sources. These measures may include:
  ➢ Identification and elimination of the contaminated product;
  ➢ Modification of nursing procedures;
  ➢ Identification and treatment of carriers, and
  ➢ Rectification of lapse in technique or procedure

11.5 Evaluation Of Efficacy Of Control Measures
➢ The efficacy of control measures are evaluated by a continued followed-up of cases after the outbreak clinically as well as microbiologically. Control measures are effective if cases cease to occur or return to the endemic level.
  The outbreak should be documented.
12. VISITORS POLICY

12.1 Introduction
Although instructing and preparing visitors for patients in isolation is time consuming and often frustrating, their presence is valuable to the emotional well-being of the patient.

- The ward sisters and the doctors concerned shall have the responsibility of informing the patients' relatives of the measures to be taken and the importance of restriction of visitors. This is done at admission of the patient.
- The patient and the relatives must be given health education about the cause, spread and prevention of the infection, in detail. The need for isolation and restriction of visitors are discussed with them.
- Hand washing after all contact with the patient will have to be stressed.
- No more than two adult visitors are allowed 'at a time' during the hospital visiting hours and the length of stay are governed by the needs of the patient.
- Children below 12 years are not allowed into the isolation areas. The policy of our hospital is to allow one female attendant to stay in the ward with the patient. The attendants are individually trained to avoid infection.
- Before entering the room, visitors must enquire at the nurses’ station for instructions and for gown and mask if indicated. Visitor’s footwear, bags etc., are left outside the room. Only articles that can be discarded, disinfected or sterilized are taken into the room.
- Visitors are not allowed to sit on the patient's bed.
- Visitors should wash their hands well with soap and water before entering and when leaving the room.
- Active immunization of attendants and other follow up steps, where applicable must be conducted by the physician in-charge.

12.2 Emergency Service
Standard precautions are to be strictly adhered and all patients are to be treated as potentially infected with blood-borne pathogens. Importance of this cannot be over emphasized in this area.

1. Wash hands with soap and water before and after patient contact.
2. Wear gloves preferably for all patient contact. It is a must for all invasive procedures, however minor. Examination gloves are placed in the shelves in all patient care areas.
3. Wear masks for all situations where a splash is expected, and where infection that spreads through the respiratory route is possible diagnosis.
4. Wear plastic aprons, in addition to a mask if splash to the body area is expected.
5. Use disposal needles and discard them into the sharps container which is placed in all patient care areas. Dispose IV cannula, stylettes, scalpel blades and razor blades into the sharps containers immediately after use.
6. Attendants and Sweepers are to wear gloves while handling lab samples and performing sanitation work.

12.3 Additional Precautions for Patients known to harbor Blood Borne Pathogens
• Use plastic aprons during procedures where body fluids may be split.
• Disinfect all items following discharge, transfer or death of the patient (as per hospital protocol refer to the chapter on housekeeping). Mattress, pillow and mackintosh are to be disinfected with 1% sodium hypochlorite solution and dried in sunlight.

12.3.1 Infectious Diseases
Refer to the chapter on Isolation Policies

12.3.2 Wound and Skin Infections
• Hands are to be washed before and after handling the patient.
• Wear gloves while handling infected wounds.
• Cover the wounds (as far as possible) before transferring the patient
• Dispose waste as per hospital guidelines
13. FOOD SAFETY

13.1 Background
Experience has shown that outbreaks of food poisoning in hospitals are notable not only because of the public interest that is generated, but because they are clinically serious and can result in the deaths of patients.

13.2 Aim
The aim is to ensure that food is provided to patients and staff in a safe and hygienic manner.

13.3 Principles of food safety
Within the hospital, any worker who handles food, or whose actions could affect its safety, must includes workers who clean articles or equipment that come into contact with food. Food and personal hygiene regulations are enforced by dietician of LNH who will make periodic visits to assess compliance.
The Infection Control Team also performs an audit. The manager of an area that contains a kitchen (Dietician, LNH) is the person deemed to be responsible for all acts of omission and commission in Kitchen area. The manager must:
- Make sure that food is supplied in a hygienic way;
- Identify food safety hazards;
- Know which steps in the processes are critical for food safety;
- Ensure that safety controls are in place, maintained and reviewed.

13.4 Basic Requirements
As a minimum kitchen should:
- Be clean and maintained in good repair;
- Be designed and constructed to permit good hygiene practices;
- Have an adequate supply of drinking water;
- Be protected against pests;
- Contain facilities for the disposal of kitchen waste;
- Have adequate hand washing facilities;
- Be provided with adequate drainage.

Food trolleys must be:
- Be adequate clean and maintained in good repair;
- Are be reserved for food only;
- Allow for separation of different products;
- Are cleaned between loads.

Staff must maintain a high degree of personal cleanliness and their practice must also be clean and hygienic. Food handlers must wear a clean uniform and protective over-clothes such as a plastic apron.

Food handlers must:
- Routinely wash their hands when handling food;
- Report any illness such as infected wounds, skin infections, diarrhoea or vomiting to their manager and occupational health immediately. If such illness is reported they must be excluded from food handling areas. Such action is the responsibility of the dietician of LNH, his or her manager.

- It is the responsibility of staff to ensure that the equipment and facilities are clean and fit for use.

13.5 Refrigerator and Freezer Use
The use of refrigerators/freezers must be carefully controlled by the dietician responsible. Controls to ensure food safety include:
- The removal of outer packaging when possible;
- The immediate storage of chilled foods after delivery checks are completed;
- Food will be stored at temperatures below 8°C refrigerator;
- Food will be packaged, wrapped or covered as protection;
- Food must be labelled with:
  - the name of the product;
  - date before which it must be used;
  - date of refrigeration;
- Food is stored within the shelf life.

13.6 Microwave Ovens
Microwave Ovens are not to be used for the heating or re-heating of patient’s food.
When used for the processing of food belonging to staff, the following applies:
- Only containers approved for use by the manufacturer are to be used.
- A core temperature of 75oC must be achieved.

13.7 Training
Food handlers must be trained in food hygiene matters to a level appropriate to their job.

13.8 Standard of Food
Guidelines to ensure that food served to patients, visitors and employees is processed in a manner that avoids contamination:-
- All food is prepared and served into covered containers and set into trays in the main kitchen and then sent to wards. This activity is supervised by trained personnel.
  - Cold storage temperatures are maintained appropriately and scrupulously.
  - Hot and cold food is transported in such a manner that appropriate temperatures will be maintained during transportation.
  - Food returned to the kitchen is discarded into black bags. Mouths of bags are tied before disposal.
  - Housekeeping is done according to the set procedures of the department.
  - The arrangement of work stations in the kitchen are such that there is no contamination of cooked food from raw food. There are no interchange of personnel working on raw food and those on cooked food.
  - Personnel handling and serving the food are trained to observe universal precautions to protect themselves.
  - Personnel are also trained to protect food consumers from body substances of handling personnel.

Training should include the following aspects.
- Hand washing should cover exposed portions of arms and hands with special attention to fingernails and areas between fingers.
- Clothing is free from obvious dirt and food spills.
- Food should not be consumed in preparation or serving areas.
- Utensils are used to handle food.
- Clean gloves may be used.

Pest control of entire facility and thorough cleaning with disinfectants should be done at defined intervals to ensure pest free food operations and safe environment.

13.9 Screening of Kitchen Workers
- Kitchen Workers must be screened for Nasal MRSA carriage, and stool parasite examination.
- Surveillance is conducted biannually for detection of carriage of *Salmonella* and MRSA. Stool samples and nasal swabs are submitted to the microbiology laboratory. Surveillance is also done after worker re-joins duty after period of leave more than two weeks. Records are maintained by in charge of the department

13.9.1 Food borne diseases
- **Bacterial disease**
  Typhoid and paratyphoid fever, Salmonellosis, Staphylococcal intoxication, *Clostridium perfringens* *B.cereus* food poisoning, *E.coli* diarrhoea, Streptococcal infection, Shigellosis, Brucella
- **Viral diseases**
  Viral hepatitis, gastroenteritis

- **Parasites**
  Taeniasis, Hydatidosis, Trichinosis, Ascariasis, amoebiasis, Oxyuriasis

13.10 Cleaning procedures for food service department facilities

<table>
<thead>
<tr>
<th>S.No.</th>
<th>Equipment/ Work area</th>
<th>Cleaning Procedure &amp; Frequency</th>
<th>Responsibility</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PRODUCTION AREA</strong></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>1.</td>
<td>Cold storage</td>
<td>Daily mop/ Sweeping</td>
<td>Kitchen mates</td>
</tr>
<tr>
<td>2.</td>
<td>Wet grinder</td>
<td>After every use thorough cleaning with water</td>
<td>Cooks/ Mates</td>
</tr>
<tr>
<td>3.</td>
<td>Knife</td>
<td>After every use. Cleaning with cold water</td>
<td>Cooks/ Mates</td>
</tr>
<tr>
<td>4.</td>
<td>Work Table Sinks</td>
<td>Twice a day cleaning with soap/water. And regular mopping with water after every use.</td>
<td>Cooks / Kitchen mates</td>
</tr>
<tr>
<td><strong>PANTRY</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.</td>
<td>Work Tables</td>
<td>Twice a day cleaning with soap/ water. And regular mopping with water after every use.</td>
<td>Kitchen mates</td>
</tr>
<tr>
<td><strong>HOT KITCHEN</strong></td>
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<td></td>
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</tr>
<tr>
<td>1.</td>
<td>Tiling pans</td>
<td>After every use washed with water</td>
<td>Cooks/ Kitchen Mates.</td>
</tr>
<tr>
<td>2.</td>
<td>Sinks / Work Tables</td>
<td>As required</td>
<td>As required</td>
</tr>
<tr>
<td><strong>DISHWASH AREA</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.</td>
<td>Tables &amp; Sinks</td>
<td>Twice a day with water and soap</td>
<td>Food Attendants.</td>
</tr>
</tbody>
</table>
14. LAUNDRY AND LINEN MANAGEMENT

14.1 Introduction
The purpose of this policy is the prevention of infection or injury in patients and health care staff involved in the use, handling or laundering of hospital linen.

14.2 Categories of Linen
14.2.1 Dirty Linen (Used linen, but not visibly soiled with blood or blood tinged body secretions)
Used linen, which may be slightly contaminated with excreta, blood and body fluids are not classed as infected.

14.2.2 Soiled Linen (Known, or potentially, infected/infested linen)
All linen which is:
- Grossly contaminated with excreta, blood or body fluids,
- Or contaminated linen from a patient who is known, or clinically suspected, to be infectious. For example salmonella, hepatitis A, B or C, open pulmonary tuberculosis, HIV.

14.3 Specific Items
14.3.1 Mattress overlays
These must be protected by waterproof covers, which are cleaned with soap and water between patients. Alcohol wipes MUST NOT be used to clean these items as alcohol damages the cover which may allow fluid to pass through to the mattress foam, the life of the mattress and its ability to protect patients form cross infection is then reduced. If the cover is damaged or punctured, and the article itself is contaminated it must be condemned and disposed of as clinical waste. Replacement covers can be purchased and may be used providing the mattress itself is not soiled stained or has a smell.

14.3.2 Staff uniforms
Must be sent to the laundry contained in the appropriate bags and labelled with the name of the individual, ward and hospital to ensure it is returned. After washing, uniforms are protected from contamination with dust during storage.

14.4 Handling And Storage Of Used Linen In Ward/Department
- Used linen must be handled with care to prevent environmental contamination with excretion or secretions, skin scales or bacteria. Linen must be bagged at the bedside, never shaken or allowed to touch the floor.
- No extraneous items must be placed in the laundry bags, especially sharp objects. This may contribute to a health & safety risk for the laundry workers.
- All linen bags must be placed in the correct colour bag, securely tied, labelled as appropriate and stored in a room or area designated for the purpose, which is safe and separate from patient areas.
- Gloves may also be required if linen is wet. Hands must be washed after handling soiled or infected linen.
- Linen are held away from the body to prevent contamination of clothing.

14.5 Transporting Used Linen from Ward / Department to Pick-Up Point
- Laundry bags must be securely tied.
- The pick-up point must be dry and secure and separate from the clean linen area.
- The frequency of collection will depend on the volume of laundry.
- Linen handlers must have heavy-duty rubber gloves available. Guidance on hand washing technique and frequency must be given.

14.6 Transporting Used Linen from the Pick-Up Point to the Laundry
- Frequency of collection will be dependent on the volume of laundry and the predefined schedule.
- Laundry is responsible for cleaning and disinfection of the Trolley in order to prevent contamination of clean linen:
  i. After any spillage
  ii. After transportation of dirty laundry, if it is to be used for clean laundry next
  iii. at least weekly
- There must be no contact between clean and soiled linen at any time. So, clean and dirty/soiled linen are transported separately from separate corridors, clean linen are transported in white trolleys while dirty linen are transported in a red trolley, if the linen is soiled it are first tied in a red bag.

14.7 Return of Clean Linen TO THE USER
Contamination of clean linen must be prevented by:
- Storage in a clean, dry area or cage
- Transport in a white trolley which is cleaned and disinfected prior to loading with clean linen. Linen that is (or thought to be) contaminated must be returned to the laundry for re-processing.

14.8 Infection Control Issues In The Laundry
- No person shall be permitted to work in or about the processing or handling of any article to be supplied to the hospital while suffering from an infection or skin disease. All contractors' staff must report such conditions to the contractor.
- Personal protective clothing will be available and worn when handling linen.
- All such clothing must be removed and changed each time the person leaves the department.
  a) Heavy duty rubber gloves  b) Apron
- Disposable items must not be re-used. Reusable gloves must be cleaned and dried at least daily.
- A hand hygiene facility complete with soap and paper towels, must be available close to the working areas.
- Staff must be aware of the possibility of extraneous items and sharps containers must be available.
- Staff must be aware of actions to take in the event of a sharps injury.
- Systems and machinery will be designed and operated so as to reduce the risk of re-infection of linen during the course of the laundering process and, to prevent articles being re-infected after laundering and prior to re-issue to the hospital.

14.9 Spillage Of Contaminated Linen
Wearing gloves, replace the linen in an appropriate bag. Clean the surface as per spill management policy and wash the surface with detergent and water and dry. Wash hands thoroughly after removing gloves.

14.10 APPENDIX - Thermal disinfection times and temperatures and environmental issues in the laundry
14.10.1 Disinfection of used (soiled and fouled) linen
- A sluice cycle is incorporated into washing machines for the removal of organic matter from fouled linen.
- Put 200gm of bleaching powder (25Ltr water) in one sluice cycle to disinfect soiled linen.
- Wash loads will have a mixing time of 8 minutes added to the temperature holding times.
- The wash temperatures will be maintained:
14.10.2 Disinfection of suspected (or known) infected linen
- The temperatures described previously will adequately disinfect linen.
- This linen must not be processed in a batch continuous washing machine, but are processed in a washer extractor.
14.10.3 Disinfection of heat-labile linen (Blanket)
- If soiled, than first dip the blanket in Bleaching powder (0.5%) for 20 minutes, than sluicing will be done to wash off any organic material stick to it.
- Linen in this category must be laundered in a machine at 40°C and dried at 60°C using tumble dryers.
- Bleaching powder (0.5%) may be used in the penultimate rinse.

14.11 General measures to prevent infection
- All surfaces will be kept free from dust, debris and pests. There will be a system for regular cleaning of the environment including high level surfaces.
- All washing machines will be kept clean and free from algae.
- All washing machines are fitted with accurate heat sensors that are correctly positioned. These must be tested at predefined interval and calibrated. Records must be kept of this and of regular monitoring of wash temperatures.
15. Staff Health Programme

15.1 Health Evaluation at Placement

❖ A medical checkup is performed at placement according to protocol laid down by Govt of NCT of Delhi. After induction to the hospital services immunisation and health appraisal is conducted by preventive health clinic medical officer in conjunction with HICC nominated member. Data is maintained by Preventive Health Clinic Medical Officer and monthly presented in HICC meeting.
❖ The data is collected in prescribed form.
❖ Vaccination for Hepatitis B is provided to all staff members who are not vaccinated or those vaccinated but do not have protective anti-HBs levels. These schedules are completed by the staff member within three months of start of employment. All staff are encouraged to get their Anti-HBs titers done to ensure their safety after vaccination.
❖ Vaccination for Salmonellosis is mandatory for kitchen staff and must be vaccinated within three months of their employment.
❖ Vaccination Varicella, Meningococcal Disease etc. will be carried out in staff exposed during the outbreak or as and when required as decided by HICC time to time.

15.2 Employee Health Programme

❖ Employee health education: Periodic education programs are conducted for paramedical staff by the ICN. All employees MUST attend the program within month of their induction to the hospital and then at least twice a year. The attendance record is kept by ICN. All employee are instructed to adhere to universal precautions, nursing barrier/isolation policies, hand washing protocols and waste management.
❖ All infections including contagious and other diagnosed communicable diseases e.g. hepatitis, mumps, rubella, measles, chicken pox, diarrhea, productive cough more than three weeks, rashes etc., MUST to be reported by staff to their immediate supervisor and thereby to ICN at which time appropriate action to protect the patients/staff in the hospital will be taken. Work restrictions may be imposed in situations which call for such action.
❖ All staff is informed that they should report exposure to potentially infectious body fluid to their immediate supervisor who in turn informs the ICN or secretary HICC in absence of ICN. Action is taken after assessment of risk at each situation (refer PEP guidelines). It is MANDATORY to report all such kind of exposures on prescribed form. Work restrictions may be imposed in situations which call for such action.
❖ Personnel shall adhere to policies and practices to minimize the potential spread of diseases and/or infection. Personnel shall adhere to existing employee health requirements.
16. MULTIDOSE VIAL POLICY
MULTIDOSE VIALS, BOTTLES, DROPPERS, UNIT DOSE AMPULES/SYRINGES

16.1 General
To establish a uniform policy on shelf life and handling of all multidose vials, bottles, droppers, unit dose ampoules/syringes, and single dose medications in use in the LNH.

16.2 Specific
16.2.1 Multi-Dose Vials
a) Multiple dose/multi-dose medication vials must be handled in accordance with the manufacturer's instructions to include:
   1. Place the expiration date on the opened vial. The expiration date is 28 days after the vial is opened or the manufacturer's recommended expiration date (whichever comes first) and discard at time of expiration.
   2. Refrigerate multi-dose vials after they are opened if recommended by the manufacturer.
   3. Cleanse the access diaphragm of multi-dose vials with 70% alcohol (such as alcohol swabs) before inserting a device into the vial.
   4. Use a sterile device to access a multi-dose vial and avoid touch contamination of the device before penetrating the access diaphragm.
   5. Discard the multi-dose vial be if user suspects vial sterility has been compromised.
   6. Vials of saline or water may be used as multi-dose only if they contain a preservative.
   7. Visual inspection of the vial should be accomplished each time medication is withdrawn to determine that the stopper is intact and that no unusual particulate matter is in the vial.
   Check the vial for
      a. Turbidity
      b. Discoloration
      c. Integrity of rubber stopper seal.
   8. Avoid opening more than one multidose vial of the same medication at the same time.
   9. Refrigeration of opened multidose vials is product specific (i.e., insulin, heparin, etc., will be refrigerated).
      ➢ Routine refrigeration of opened multidose vials is not recommended.
   10. Read the label.
      All components on labels include:
         1. Name of Drug:
         2. Drug dilution/concentration:
         3. Date and Time of Opening:
         4. Date and Time of Expiry (28 days of opening date or expiry by manufacturer whichever is earlier)
         5. Signature of the staff labelling the vial.

16.2.2 Single Use vials
1. Single use parenteral drugs do not contain preservatives and should be immediately discarded by the original user after the dose is withdrawn. Those vials containing medications that have limited storage capability should be dated and initialed and disposed of in accordance with the manufacturer's recommended instructions.
2. Single dose containers are preferred over multidose containers. If this is not possible, the smallest multidose container available should be used. This lessens the risk of contamination/cross contamination.
3. A sterile needleless device or blunt syringe must be used to withdraw the required amount of medication from single dose vials.
4. Unit dose glass ampoules/syringes are specifically designed for single dose only. Any unused portions of medications must be discarded immediately and not left on the unit for any period of time.

16.2.3 Precautions For Maintaining Drug Integrity
All drugs will be clearly labeled. The identification of a drug shall not be assumed if unlabeled. When drugs are unlabeled or labels are defaced, these drugs will not be used.
Instructions on drugs should be read carefully to determine the temperature range at which the drugs are to be kept.
Some medications are clearly labeled: DO NOT REFRIGERATE. Thus the refrigerator should not be used arbitrarily as a storage place for drugs.
All drugs will be checked prior to use and monthly for expiration to ensure that outdated drugs are not used. If drugs have only lot number but no expiration date, the lot number may be checked by pharmacy for expiration date.
A multidose vial labeled to expire in a given month will expire on the last day of that month.

16.2.4 Multidose Ophthalmic Drops
Multidose Ophthalmic Drops for inpatients may be ordered from unit dose and used only for that patient.
Other multidose ophthalmic drops may be used for more than one patient provided the dropper surface is not contaminated by touching any part of the eye, eyeball, face, or eyelid.

16.2.5 Ampoules
All ampoules formulations should be discarded immediately after use.

16.3 Responsibilities
Nursing Staff
• Must follow the requirements of the policy.
• It is the responsibility of each person using a multiple-dose vial to determine its safety for future use based on any suspected or known compromise to the solution’s sterility.

Unit incharges of all departments
• Must ensure employee compliance with the policy.

Surveillance and infection control Division
• Will bring these policies to the HICC
17. Biomedical Waste Management

Waste management policy at LNH has been implemented in accordance with the rules of Delhi State Pollution Control Board and Biomedical Waste Management and Handling (Second amendment) Rules, 2000.

17.1 Environmental Protection Act, 1986
The Government of India (GOI) enacted the Environmental Protection Act, 1986, (EPA) under Article 253 of the Constitution. The purpose of this Act is to serve as an “umbrella” legislation designed to provide a framework for central government coordination for the activities of various established central and state authorities. As this is an “umbrella” and all-encompassing legislation, this is relevant to the health sector activities as well. There are rules / notifications that have been brought out under this Act, which are directly relevant to the health sector.

Under the Environmental Protection Act, the Bio-Medical Waste Management Rules were introduced. These Rules are directly relevant to the health sector. The salient features of these Rules are as follows:

- Bio-medical waste means waste that is generated during the diagnosis, treatment or immunization of human beings or animals or in research activities pertaining thereto or in the production or testing of biological.
- It is the duty of every occupier of an institution generating bio-medical waste which includes a hospital, nursing home, clinic, dispensary, veterinary institution, animal house, pathological laboratory, blood bank by whatever name called to take all steps to ensure that such waste is handled without any adverse effect to human health and the environment.
- Bio-Medical waste shall not be mixed with other wastes.
- Bio-Medical waste shall be segregated into containers/bags at the point of generation in accordance with Schedule II of these Rules prior to its storage, transportation, treatment and disposal. The containers shall be labelled according to Schedule III of these Rules.
- Bio-Medical waste shall be treated and disposed of in accordance with Schedule I of these Rules, which gives the categories of waste and methods for treatment and disposal. The Rules also require compliance with the standards prescribed in Schedule V, which gives standards for different treatment technologies.

17.2.1 Objectives
1. To prevent infection by maintaining good hygiene and sanitation.
2. To protect the patient, patient attendants and all health care personnel from avoidable exposure to infection.
3. To prevent environmental pollution.
4. To manage waste in a clean, healthy, economical and safe manner.
5. To minimize waste

17.2.2 Steps in Waste Management
- Segregation
- Transport to site of temporary storage.
- Final disposal

17.2.2.1 Segregation
Segregation is done at source. A colour code is followed and appropriately coded waste bags are placed in bins in all patient care areas. Labelling of all the bags with predesigned labels with information including hospital name, patient care unit name and date is done before usage of bags. A liaison nurse has been designated to carry our surprise rounds in every unit of hospital and check for proper segregation and educating new staff then and there. Regular employee education programs are held at LNH for constant sensitization of healthcare workers.

17.2.2.2 Transportation
- Waste from various patient care areas is removed twice a day or more frequently if necessary. All bags are tied at the mouth to avoid spillage during transport. The bags are then transported in larger moving carts carried by the house keeping department. The bags are transported to the central waste receiving terminal.
- Avoid the transport of too many bags at one time and contact of the bag with the body of personnel
- Mixing of segregated wastes should NEVER be done. It the duty of unit in charge to ensure that the bag from their unit is properly transported without mixing.
17.2.2.3 Final disposal
No final disposal of waste is undertaken within the hospital premises. This is undertaken by an outsourcing agency recognized by Govt of NCT of Delhi.

Annual report of waste generated is maintained by MOI/C Bio-medical waste management.

17.2.3 Disposal of Contaminated Needles and Syringes
Contaminated needles are burnt in needle destroyer and the trays are emptied in sharps container when use of needle destroyer is possible. Contaminated needle are disposed of by placing them uncapped into a puncture resistant container. Containers are closed and are handed over to the medical waste disposal contractors.

17.3 Waste Management Committee
Waste Management committee is responsible for making Hospital specific action plan for hospital waste management and its supervision, monitoring and implementation.

17.3.1 Terms of Reference
• To seek a commitment from Management to comply with all relevant Legislation (Delhi State Pollution Control Board and Biomedical Waste Management Handling Rules)
• To conduct a waste audit and prepare a comprehensive report of current waste generation, segregation, handling, storage and disposal practices and costs
• To provide appropriate Personal Protective equipment and offer staff vaccinations
• To develop spill management strategies for all waste categories
• To implement an ongoing waste management training program which caters for all staff
• To promote waste management principles throughout hospital (signs, posters, notice boards, bulletins, etc)
• To improve waste segregation
• To liaise with the corporation authorities and private waste contractors with regard to the transport and disposal of waste external to the hospital.
• To conduct a Waste Management Audit annually and review the Waste Management Plan
• To conduct on-going audits of waste

17.3.2 Membership of the Group
• Microbiologist/Clinician – Consultant for Bio-waste management (CBM)
• Liaison Nurse to report to CBM and Infection Control Nurse
• Housekeeping in-charge

The Committee is represented at the Hospital Risk Management and Safety Committee, where progress reports are made at each meeting. Minutes of the meeting are maintained.

17.3.3 Meetings
The Group will meet quarterly as a part of HICC meeting or more frequently if necessary

17.3.4 Record keeping on biomedical waste
Annual report—Every occupier/operator shall submit an annual report to the prescribed authority in below mention form by 31st January every year. The prescribed authority will send this information in a compiled form to the central pollution control board by 31st March every year.

SCHEDULE I
Biomedical wastes categories and their segregation, collection, treatment, processing and disposal options
Category Type of Waste Type of Bag or Container to be used Treatment and Disposal options

<table>
<thead>
<tr>
<th>Category</th>
<th>Type of Waste</th>
<th>Type of Bag or Container to be used</th>
<th>Treatment and Disposal options</th>
</tr>
</thead>
<tbody>
<tr>
<td>YELLOW</td>
<td>(a) Human Anatomical Waste:</td>
<td></td>
<td>Yellow coloured non-chlorinated plastic bags Incineration or Plasma Pyrolysis or deep burial*</td>
</tr>
<tr>
<td></td>
<td>Human tissues, organs, body parts and fetus below the viability period (as per the Medical Termination of Pregnancy)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**Hospital Infection and Prevention and Control Manual**

Lok Nayak Hospital

April 2016

<table>
<thead>
<tr>
<th>(b) Animal Anatomical Waste</th>
<th>(c) Soiled Waste</th>
<th>(d) Expired or Discarded Medicines</th>
<th>(e) Chemical Waste</th>
<th>(f) Chemical Liquid Waste</th>
</tr>
</thead>
<tbody>
<tr>
<td>Experimental animal carcases, body parts, organs, tissues, including the waste generated from animals used in experiments or testing in veterinary hospitals or colleges or animal houses.</td>
<td>Items contaminated with blood, body fluids like dressings, plaster casts, cotton swabs and waste to be sent for energy recovery.</td>
<td>Pharmaceutical waste like antibiotics, cytotoxic drugs including all items contaminated with cytotoxic drugs along with glass or plastic ampoules, vials etc.</td>
<td>Chemicals used in production of biological and used or discarded disinfectants.</td>
<td>Liquid waste generated due to use of chemicals in production of biological and used or discarded disinfectants, Silver X-ray film developing liquid, discarded Formalin,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Yellow coloured non-chlorinated plastic bags or containers</td>
<td></td>
<td>Separate collection system leading to effluent treatment system</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Expired cytotoxic drugs and items contaminated with cytotoxic drugs to be returned back to the manufacturer or supplier for incineration at temperature &gt;1200°C or to common biomedical waste treatment facility or hazardous waste treatment, storage and disposal facility for incineration at &gt;1200°C Or Encapsulation or Plasma Pyrolysis at &gt;1200°C. All other discarded medicines shall be either sent back to manufacturer or disposed by incineration.</td>
<td>Disposed of by incineration or Plasma Pyrolysis or Encapsulation in hazardous waste treatment, storage and disposal facility.</td>
<td>After resource recovery, the chemical liquid waste shall be pre-treated before mixing with other wastewater. The combined discharge shall conform to the discharge norms given in Schedule-III.</td>
</tr>
</tbody>
</table>
infected secretions, aspirated body fluids, liquid from laboratories and floor washings, cleaning, house-keeping and disinfecting activities etc.

(g) Discarded linen, mattresses, beddings contaminated with blood or body fluid.

Non-chlorinated yellow plastic bags or suitable packing material
Non-chlorinated chemical disinfection followed by incineration or Plasma Pyrolysis or for energy recovery.
In absence of above facilities, shredding or mutilation or combination of sterilization and shredding. Treated waste to be sent for energy recovery or incineration or Plasma Pyrolysis.

(h) Microbiology, Biotechnology and other clinical laboratory waste:
Blood bags, Laboratory cultures, stocks or specimens of microorganisms, live or attenuated vaccines, human and animal cell cultures used in research, industrial laboratories, production of biological, residual toxins, dishes and devices used for cultures.

Autoclave safe plastic bags or containers
Pre-treat to sterilize with nonchlorinated chemicals on-site as per National AIDS Control Organisation or World Health Organisation guidelines thereafter for Incineration.

RED Contaminated Waste (Recyclable)
(a) Wastes generated from disposable items such as tubing, bottles, intravenous tubes and sets, catheters, urine bags, syringes (without needles and fixed needle syringes) and vaccutainers with their needles cut) and gloves.

Red coloured non-chlorinated plastic bags or containers
Autoclaving or micro-waving/hydroclaving followed by shredding or mutilation or combination of sterilization and shredding. Treated waste to be sent to registered or authorized recyclers or for energy recovery or plastics to diesel or fuel oil or for road making, whichever is possible. Plastic waste should not be sent to landfill sites.

WHITE (TRANSLUCENT) Waste sharps including Metals:
Needles, syringes with fixed needles, needles from needle tip cutter or burner, scalpels, blades,
Puncture proof, Leak proof, tamper proof containers
Autoclaving or Dry Heat Sterilization followed by shredding or mutilation or encapsulation in metal container or cement concrete; combination of...
or any other contaminated sharp object that may cause puncture and cuts. This includes both used, discarded and contaminated metal sharps

<table>
<thead>
<tr>
<th>BLUE</th>
<th>(a) Glassware: Broken or discarded and contaminated glass including medicine vials and ampoules except those contaminated with cytotoxic wastes.</th>
<th>Cardboard boxes with blue colored marking</th>
<th>Disinfection (by soaking the washed glass waste after cleaning with detergent and Sodium Hypochlorite treatment) or through autoclaving or microwaving or hydroclaving and then sent for recycling.</th>
</tr>
</thead>
<tbody>
<tr>
<td>(b) Metallic Body Implants</td>
<td>Cardboard boxes with blue colored marking</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Form - IV**

*(See rule 13)*

**ANNUAL REPORT**

[To be submitted to the prescribed authority on or before 30th June every year for the period from January to December of the preceding year, by the occupier of health care facility (HCF) or common bio-medical waste treatment facility (CBWTF)]

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Particulars</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Particulars of the Occupier</td>
</tr>
<tr>
<td></td>
<td>(i) Name of the authorised person (occupier or operator of facility)</td>
</tr>
<tr>
<td></td>
<td>(ii) Name of HCF or CBMWTF</td>
</tr>
<tr>
<td></td>
<td>(iii) Address for Correspondence</td>
</tr>
<tr>
<td></td>
<td>(iv) Address of Facility</td>
</tr>
<tr>
<td></td>
<td>(v) Tel. No, Fax. No</td>
</tr>
<tr>
<td></td>
<td>(vi) E-mail ID</td>
</tr>
<tr>
<td></td>
<td>(vii) URL of Website</td>
</tr>
<tr>
<td></td>
<td>(viii) GPS coordinates of HCF or CBMWTF</td>
</tr>
<tr>
<td></td>
<td>(ix) Ownership of HCF or CBMWTF</td>
</tr>
<tr>
<td></td>
<td>(x) Status of Authorisation under the Bio-Medical Waste (Management and Handling) Rules</td>
</tr>
<tr>
<td></td>
<td>(xi). Status of Consents under Water Act and Air Act</td>
</tr>
<tr>
<td>2.</td>
<td>Type of Health Care Facility:</td>
</tr>
<tr>
<td></td>
<td>(i) Bedded Hospital</td>
</tr>
<tr>
<td></td>
<td>(ii) Non-bedded hospital (Clinic or Blood Bank or Clinical Laboratory or Research Institute or Veterinary Hospital or any other)</td>
</tr>
<tr>
<td></td>
<td>(iii) License number and its date of expiry</td>
</tr>
<tr>
<td>3.</td>
<td>Details of CBMWTF:</td>
</tr>
<tr>
<td></td>
<td>(i) Number healthcare facilities covered by</td>
</tr>
<tr>
<td>CBMWTF</td>
<td></td>
</tr>
<tr>
<td>--------</td>
<td>---</td>
</tr>
<tr>
<td>(ii) No of beds covered by CBMWTF</td>
<td>:</td>
</tr>
<tr>
<td>(iii) Installed treatment and disposal capacity of CBMWTF:</td>
<td>: _______ Kg per day</td>
</tr>
<tr>
<td>(iv) Quantity of biomedical waste treated or disposed by CBMWTF</td>
<td>: _______ Kg/day</td>
</tr>
</tbody>
</table>

4. Quantity of waste generated or disposed in Kg per annum (on monthly average basis)
   : Yellow Category : 
   : Red Category : 
   : White: 
   : Blue Category : 
   : General Solid waste: 

5. Details of the Storage, treatment, transportation, processing and Disposal Facility
   : (i) Details of the on-site storage Facility : Size : 
   : Capacity : 
   : Provision of on-site storage : (cold storage or any other provision) |
   : (ii) Quantity of recyclable wastes sold to authorized recyclers after treatment in Kg per annum. : Red Category (like plastic, glass etc.) |
   : (iv) No of vehicles used for collection and transportation of biomedical waste |
   : (v) Details of incineration ash and ETP sludge generated and disposed during the treatment of wastes in Kg per annum 
   : Quantity generated 
   : Where disposed |
   : (vi) Name of the Common Bio-Medical Waste Treatment Facility Operator through which wastes are disposed of : |
   : (vii) List of member HCF not handed over biomedical waste: |

6. Do you have bio-medical waste management committee? If yes, attach minutes of the meetings held during the reporting period 

7. Details of the accident occurred during the year
   : (i) Number of Accidents occurred 
   : (ii) Number of the persons affected 
   : (iii) Remedial Action taken (Please attach details if any) 
   : (iv) Any Fatality occurred, details. |

8. Details of the accident occurred during the year
   : (i) Number of Accidents occurred 
   : (ii) Number of the persons affected 
   : (iii) Remedial Action taken (Please attach details if any) 
   : (iv) Any Fatality occurred, details. |

9. Are you meeting the standards of air Pollution from the incinerator? How many times in last year could not met the standards?

10. Liquid waste generated and treatment methods in place. How many times you have not met the standards in a year? 

11. Is the disinfection method or sterilization
meeting the log 4 standards? How many times you have not met the standards in a year?

12. Any other relevant information : (Air Pollution Control Devices attached with the Incinerator)

Certified that the above report is for the period from
..........................................................................................................................
..........................................................................................................................
..........................................................................................................................
..........................................................................................................................
..........................................................................................................................

Name and Signature of the Head of the Institution

Date:
Place

FORM – I

ACCIDENT REPORTING

1. Date and time of accident :

2. Type of Accident :

3. Sequence of events leading to accident :

4. Has the Authority been informed immediately :

5. The type of waste involved in accident :

6. Assessment of the effects of the accidents on human health and the environment:

7. Emergency measures taken :

8. Steps taken to alleviate the effects of accidents :

9. Steps taken to prevent the recurrence of such an accident :

10. Does you facility has an Emergency Control policy? If yes give details:

    Date : ...................... Signature ....................... 

    Place: ...................... Designation .....................
**SCHEDULE IV**
[See rule 8(3) and (5)]

**Part A**

**LABEL FOR BIO-MEDICAL WASTE CONTAINERS or BAGS**

**CYTOTOXIC HAZARD SYMBOL**

**HANDLE WITH CARE**

**Part B**

**LABEL FOR TRANSPORTING BIO-MEDICAL WASTE BAGS OR CONTAINERS**

Day ............

Month ............

Year ............

Date of generation ....................

Waste category Number ........

Waste quantity ........

Sender's Name and Address Receiver's Name and Address:

Phone Number ........

Fax Number ........

Contact Person ........

In case of emergency please contact:

Name and Address:

Phone No.

Note : Label shall be non-washable and prominently visible.
WARD SISTER’S BMW RECORD SHEET

WARD NO. __________________________ DEPARTMENT OF __________________________

<table>
<thead>
<tr>
<th>DATE &amp; TIME</th>
<th>WEIGHT OF BAGS</th>
<th>NAME &amp; SIGN OF NURSING STAFF</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>RED</td>
<td>YELLOW</td>
</tr>
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</tbody>
</table>

IT IS MANDATORY TO DEPOSIT THIS DOCUMENT TO THE DEPARTMENT OF SANITATION, HK & BMWM AT THE END OF EVERY MONTH

SIGN OF NURSING STAFF
18. Antimicrobial Stewardship Programme

18.1 The past 30 years have brought multidrug-resistant pneumococcal, gonococci, and Salmonella spp. and extremely drug-resistant tuberculosis to patients in the community. Vancomycin-resistant enterococci and vancomycin-resistant S. aureus have also emerged. Extremely drug-resistant gram-negative bacteria, such as carbapenemase-producing Klebsiella pneumoniae and other carbapenem-resistant Enterobacteriaceae spp., extended-spectrum beta-lactamase-producing Enterobacteriaceae, P. aeruginosa, and Acinetobacter baumannii have spread widely among patients in healthcare settings; in some cases these pathogens have been panresistant, that is, resistant to all available antibiotics.

Unfortunately, during the last decade there has also been a dramatic drop in the development and approval of new antibacterial agents. The antimicrobial armamentarium has been depleted and our ability to treat infectious diseases has been severely compromised. Resistant infections not only result in increased morbidity and mortality but also dramatically increase healthcare costs. It is ironic that in the twenty-first century we are encountering bacterial infections for which we have no treatment. A multifaceted approach is necessary to prevent, detect, and control the emergence of antimicrobial-resistant organisms. This includes ensuring the availability of adequate and appropriate therapeutic agents, the existence of diagnostic capacity to rapidly and reliably detect specific pathogens and their antimicrobial susceptibilities, and the promotion of robust infection prevention, control, and antimicrobial stewardship programs. This document focuses on issues relating to antimicrobial stewardship. Other issues important to the emergence, transmission, and management of antimicrobial resistance are addressed else.

18.2 Definition
Antimicrobial stewardship refers to coordinated interventions designed to improve and measure the appropriate use of antimicrobial agents by promoting the selection of the optimal antimicrobial drug regimen including dosing, duration of therapy, and route of administration.

18.3 Objectives
The major objectives of antimicrobial stewardship are to achieve best clinical outcomes related to antimicrobial use while minimizing toxicity and other adverse events, thereby limiting the selective pressure on bacterial populations that drives the emergence of antimicrobial-resistant strains. Antimicrobial stewardship may also reduce excessive costs attributable to suboptimal antimicrobial use.

A Multidisciplinary inter professional antimicrobial stewardship committee [Drug and therapeutics committee (DTC)] should be in place which is physician directed or supervised.

There should be Antibiotic Management Team (AMT).

Team members include:

a) A clinical microbiologist.
b) An infection control nurse

18.4 Antibiotic policy
Antibiotic policy is to be prepared by the antimicrobial stewardship team in consultation with microbiology departments and physicians and surgeons from various departments. The policy is reviewed and updated annually.

18.5 Antimicrobial Stewardship Program
18.5.1 Antimicrobial Stewardship Program included Monitoring of following activities at LNH:
1. Rational use of antibiotic are been monitored – On daily basis for seven indicator antibiotics (Vancomycin, Meropenem, Ofl oxacin, Ciprofloxacin, Cefoperazone + Sulbactam, Colistin and Levofloxacin, Daptomycin, Tigecycline, Ceftaroline and any other antibiotic outside hospital formulary) by ICNs on daily rounds and details recorded on preformatted template. Other antibiotics are also checked for rational combinations and doses. Treating doctors are asked to explain the reasons for initiating these antibiotics in writing. These patients are the discussed for rationality with Clinical Microbiologists. Irrational antibiotic therapy, if identified is communicated to treating physician or surgeon for immediate discontinuation/modification. Irrational combination of antibiotics or doses is also monitored.
2. Pre – surgical prophylaxis and and post operative antibiotic therapy are also monitored on daily basis. In case of irrationality it is been informed to the concerned department and necessary actions are taken.
3. Defined Daily Dose (DDD) for antibiotics are monitoring for the usage pattern.
4. No. of doses administered are also monitored per thousand patient days.
5. The data analysis is done and discussed during periodic HICC meetings.
6. Adherence to antibiotic policy is also discussed in the HICC meeting.
7. Prescription audits of in patients and outpatients are conducted periodically.
8. Antibiotic costing and savings thus done is being monitored.
Appendix 1. HIC Indicators

Various indicators used for hospital associated infections include:
2. Catheter related blood stream infections (CRBSI)
3. Surgical site infections (SSI)
4. Catheter associated urinary tract infections (CAUTI)
5. Ventilator associated pneumonia (VAP)
6. Hospital acquired blood stream infections (HA BSI)
7. Device utilisation rates for central line catheters, Foley’s catheter and ventilators.
8. Antibiotic usage and resistance monitoring (AUR)

1. To calculate Hospital acquired infections in various units:
   Data to be calculated include:
   \[ \text{No. of patients with healthcare associated infections in particular unit} \times 1000 \]
   \[ \text{No. of patient days in that particular unit} \]

2. To calculate CRBSI, data to be collected include:
   \[ \text{No. of patients developed CRBSI} \times 1000 \]
   \[ \text{Total no. of catheter days} \]

3. To calculate SSI in surgical unit, data to be collected include:
   \[ \text{No. of patients with SSI in surgical department} \times 1000 \]
   \[ \text{No. of patient undergoing surgery in the department} \]

4. To calculate CAUTI, data to be collected include:
   \[ \text{No. of patients developed CAUTI} \times 1000 \]
   \[ \text{Total no. of urinary catheter days} \]

5. To calculate VAP, data to be collected include:
   \[ \text{No. of patients developed VAP} \times 1000 \]
   \[ \text{Total no. of Ventilator days} \]

6. To Calculate Hospital acquired BSI
   \[ \text{No. of patients developed BSI (HAI)} \times 1000 \]
   \[ \text{Total No.of patient days} \]

7. To calculate Device (Ventilator, central line, Foley’s Catheter) Utilization Rate:
   \[ \text{No. of Device days} \]
   \[ \text{No. of Patient days} \]

Device-days are the total number of days of exposure to the device (ventilator, urinary catheter or central line) by all of the patients during the selected time period.
Patient-days are the total number of days that patients are in a particular unit during the specified time period.

Calculation of device associated infection rate:

Device- associated Infection Rate= \[ \frac{\text{No. of device –associated infections for a specific site}}{\text{Number of device days}} \times 1000 \]

This is done for 3 devices namely.
1. Central line- Sample from CVP tip
2. Ventilator -Sample from endotracheal tube secretions
3. Foley’s Catheter - Urine sample

Data is collected in a prescribed format.

8. Antibiotic Utilization rate: Antibiotic used (gms)
Defined drug Dose (gms)

**Calculation of Hand Hygiene Compliance:**

\[
\text{Compliance (\%) = \frac{\text{Actions}}{\text{Opportunities}} \times 100}
\]

**Case definitions used for diagnosis of HCAIs**

Case definitions as described by National Healthcare Safety Network (NHSN), CDC are being used. The summary diagrams of the common HCAIs are summarized below:

Healthcare associated infection (HAI) is acquired in a hospital by a patient, that is, it was not present or incubating at the time of admission. This also includes infection acquired in the hospital but appearing after discharge. These infections can occur from inadvertent exposure to pathogenic bacteria’s, viruses, fungi or spores.

**BLOOD STREAM INFECTIONS**

<table>
<thead>
<tr>
<th>LABORATORY CONFIRMED BLOOD STREAM INFECTION</th>
<th>CENTRAL LINE-ASSOCIATED BLOOD STREAM INFECTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>LCBI -1</strong></td>
<td><strong>LCBI -2</strong></td>
</tr>
<tr>
<td>Patient has a recognized pathogen cultured from one or more blood cultures AND organism cultured from blood is not related to an infection at another site</td>
<td>Patient has at least one of the following signs or symptoms: fever (&gt;38.0°C), chills, or hypotension AND organism cultured from blood is not related to an infection at another site AND the same common commensal (i.e., diphtheroids [Corynebacterium spp. not C. diphtheriae], Bacillus spp. [not B. anthracis], Propionibacterium spp., coagulase-negative staphylococci [including S. epidermidis], viridans group streptococci, Aerococcus spp., and Micrococcus spp.) is cultured from two or more blood cultures drawn on separate occasions.</td>
</tr>
</tbody>
</table>
### SURGICAL SITE INFECTION

Must meet the following criteria:

<table>
<thead>
<tr>
<th>Superficial SSI</th>
<th>Deep SSI</th>
<th>Organ/Space SSI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infection occurs within 30 days after any NHSN operative procedure (where day 1 = the procedure date), including those coded as ‘OTH’* AND involves only skin and subcutaneous tissue of the incision AND patient has at least one of the following: a. purulent drainage from the superficial incision. b. organisms isolated from an aseptically-obtained culture from the superficial incision or subcutaneous tissue. c. superficial incision that is deliberately opened by a surgeon, attending physician** or other designee and is culture positive or not cultured AND patient has at least one of the following signs or symptoms: pain or tenderness; localized swelling; erythema; or heat. A culture negative finding does not meet this criterion. d. diagnosis of a superficial incisional SSI by the surgeon or attending physician** or other designee.</td>
<td>Infection occurs within 30 or 90 days after the NHSN operative procedure (where day 1 = the procedure date) according to the list in Table 3 AND involves deep soft tissues of the incision (e.g., fascial and muscle layers) AND patient has at least one of the following: a. purulent drainage from the deep incision. b. a deep incision that spontaneously dehisces, or is deliberately opened or aspirated by a surgeon, attending physician** or other designee and is culture positive or not cultured AND patient has at least one of the following signs or symptoms: fever (&gt;38°C); localized pain or tenderness. A culture negative finding does not meet this criterion. c. an abscess or other evidence of infection involving the deep incision that is detected on gross anatomical or histopathologic exam, or imaging test. ** The term attending physician for the purposes of application of the NHSN SSI criteria may be interpreted to mean the surgeon(s), infectious disease, other physician on the case, emergency physician or physician’s designee (nurse practitioner or physician’s assistant).</td>
<td>Infection occurs within 30 or 90 days after the NHSN operative procedure (where day 1 = the procedure date) according to the list in Table 3 AND infection involves any part of the body deeper than the fascial/muscle layers, that is opened or manipulated during the operative procedure AND patient has at least one of the following: a. purulent drainage from a drain that is placed into the organ/space (e.g., closed suction drainage system, open drain, T-tube drain, CT guided drainage) b. organisms isolated from an aseptically-obtained culture of fluid or tissue in the organ/space c. an abscess or other evidence of infection involving the organ/space that is detected on gross anatomical or histopathologic exam, or imaging test AND meets at least one criterion for a specific organ/space infection site listed in Table 4. These criteria are in the Surveillance Definitions for Specific Types of Infections chapter.</td>
</tr>
</tbody>
</table>
## URINARY TRACT INFECTION

### Symptomatic UTI (SUTI)
Must meet at least **one** of the following criteria

<table>
<thead>
<tr>
<th>SUTI 1a</th>
<th>SUTI 1b</th>
<th>SUTI 2</th>
<th>ABUTI</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Catheter-associated Urinary Tract Infection (CAUTI)</strong></td>
<td><strong>Non-Catheter-associated Urinary Tract Infection (Non-CAUTI)</strong></td>
<td><strong>CAUTI or Non-CAUTI in patients 1 year of age or less</strong></td>
<td><strong>Asymptomatic bacteremic UTI (ABUTI)</strong></td>
</tr>
</tbody>
</table>

**Patient must meet 1, 2, and 3 below:**
1. Patient had an indwelling urinary catheter that had been in place for \(>2\) days on the date of event (day of device placement = Day 1) AND was either:
   - Still present on the date of event†, OR
   - Removed the day before the date of event‡

   _OR_
   - Patient did not have a urinary catheter in place on the date of event nor the day before the date of event

2. Patient has at least **one** of the following signs or symptoms:
   - fever (\(>38.0^\circ\text{C}\))
   - suprapubic tenderness*
   - costovertebral angle pain or tenderness*
   - urinary urgency*
   - urinary frequency*
   - dysuria*

3. Patient has a urine culture with no more than two species of organisms, at least one of which is a bacteria of \(\geq105\) CFU/ml

**Patient must meet 1, 2, and 3 below:**
1. One of the following is true:
   - Patient has/had an indwelling urinary catheter but it has/had not been in place \(>2\) calendar days on the date of event†

**Patient must meet 1, 2, and 3 below:**
1. Patient has at least **one** of the following signs or symptoms:
   - fever (\(>38.0^\circ\text{C}\)) in a patient that is \(\leq65\) years of age
   - suprapubic tenderness*
   - costovertebral angle pain or tenderness*
   - urinary frequency*
   - dysuria*

**Patient must meet 1, 2, and 3 below:**
1. Patient is \(\leq1\) year of age (with or without an indwelling urinary catheter)

2. Patient has a urine culture with no more than two species of organisms, at least one of which is a bacteria of \(\geq105\) CFU/ml

**Patient must meet 1, 2, and 3 below:**
1. Patient has/had an indwelling urinary catheter but it has/had not been in place \(>2\) calendar days on the date of event†

2. Patient has at least **one** of the following signs or symptoms:
   - hypothermia (\(<36.0^\circ\text{C}\))
   - apnea*
   - bradycardia*
   - lethargy*
   - vomiting*
   - suprapubic tenderness*

3. Patient has a urine culture with no more than two species of organisms, at least one of which is a bacteria of \(\geq105\) CFU/ml

**Patient must meet 1, 2, and 3 below:**
1. Patient with or without an indwelling urinary catheter has no signs or symptoms of SUTI 1 or 2 according to age

2. Patient has a urine culture with no more than two species of organisms, at least one of which is a bacteria of \(\geq105\) CFU/ml (see Comment section below)

3. Patient has a positive blood culture with at least **one** matching bacteriato the urine culture, or meets LCBI criterion 2 (without fever) and matching common commensal(s) in the urine.
VENTILATOR ASSOCIATED PNEUMONIA (VAP)

**Imaging**

- Patient with underlying diseases:
  - New or progressive and persistent infiltrate
  - Consolidation
  - Cavitation
  - Pneumatoceles, in ≤1 y.o.

- Patient without underlying diseases:
  - New or progressive and persistent infiltrate
  - Consolidation
  - Cavitation
  - Pneumatoceles, in ≤1 y.o.

**Signs and Symptoms**

- Infants ≤1 y.o.
  - Worsening gas exchange (e.g., O₂ desats [e.g., pulse oximetry <94%, fē O₂ req. or ↑ ventilation demand])
  - Temperature instability
  - Leukopenia (<4,000 WBC/mm³) or leukocytosis (≥15,000 WBC/mm³) and left shift (≥10% band forms)
  - New onset of purulent sputum or change in character of sputum, or ↑ respiratory secretions, or ↑ suctioning requirements
  - Arness, tachypnea, nasal flaring with retractions of chest wall or grunting
  - Wheezing, rales, or rhonchi
  - Cough
  - Bradycardia (<100 beats/min) or tachycardia (>170 beats/min)

- Children >1 or ≤12 y.o.
  - At least three of the following:
    - Fever (>38.0°C/100.4°F) or hypothermia (<36.0°C/96.8°F)
    - Leukopenia (<4,000 WBC/mm³) or leukocytosis (≥15,000 WBC/mm³)
    - New onset of purulent sputum or change in character of sputum, or ↑ respiratory secretions, or ↑ suctioning requirements
    - New onset of worsening cough, or dyspnea, anemia, or tachypnea
    - Rales or bronchial breath sounds
    - Worsening gas exchange (e.g., O₂ desats [e.g., pulse oximetry <94%, fē O₂ req. or ↑ ventilation demand])

**VENTILATOR ASSOCIATED PNEUMONIA**
# APPENDIX 2. List of disinfectants currently available in the hospital and their use

<table>
<thead>
<tr>
<th>S.no</th>
<th>Department</th>
<th>(70% ethyl alcohol or Isopropyl alcohol)</th>
<th>0.5%-1% Sodium Hypochlorite</th>
<th>2% Glutaraldehyde (5% Glutaraldehyde + 11.2% chemically bound formaldehyde + 5% benzalkonium chloride) Currently available formulation-Sanillocid</th>
<th>Bleaching powder (70% available chlorine)</th>
<th>(0.5% chlorhexidine + 70% ethyl alcohol) Currently available formulation- Nanzilon</th>
<th>5% Povidone iodine</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>O.T.</td>
<td>For skin disinfection, Trolley tops, cautery leads</td>
<td>Disinfection of infected plastics (syringes, cannula caps)</td>
<td>Disinfection of sharp instruments or heat labile instruments (scissors, laryngoscope) (0.5%) Tables, trolleys, tiles, floor cleaning, surgery tables.</td>
<td>Spill management</td>
<td>Hand hygiene</td>
<td>Preoperative Skin preparation</td>
</tr>
<tr>
<td>2.</td>
<td>ICU’s</td>
<td>For skin disinfection, Trolley tops, monitors leads, BP cuff</td>
<td>Surface disinfection (Bed frames, trolleys, tiles). Disinfection of infected plastics (syringes, cannula caps), for cleaning of patient’s furniture and fittings. Disinfection of suction jars &amp; tubings, laryngoscope, O2 Humidifiers</td>
<td>Terminal cleaning of furniture of patients on weekly basis (Bed frames, trolleys, tiles).</td>
<td>Spill management</td>
<td>Hand hygiene</td>
<td></td>
</tr>
<tr>
<td>3.</td>
<td>Laboratory</td>
<td>Surface cleaning( tables, Biosafety cabinets, work stations)</td>
<td>Disinfection of used syringes, slides, cover slips and culture loops etc.</td>
<td></td>
<td>NR</td>
<td>NR</td>
<td>Hand hygiene</td>
</tr>
<tr>
<td>4.</td>
<td>Wards</td>
<td>For skin disinfection,&amp; Surface cleaning (Trolley tops etc)</td>
<td>Disinfection of infected plastics (syringes, cannula caps, patient patient furniture and fixtures.)</td>
<td>Disinfection Heat labile and other instruments( scissors etc) Terminal cleaning of furniture of patients on weekly basis (Bed frames, trolleys, tiles).</td>
<td></td>
<td></td>
<td>Hand hygiene</td>
</tr>
<tr>
<td>5.</td>
<td>Dressing room</td>
<td>For skin disinfection, Trolley tops etc</td>
<td>Disinfection of infected plastics (syringes etc)</td>
<td>Disinfection of instruments</td>
<td>NR</td>
<td>NR</td>
<td>Hand hygiene</td>
</tr>
</tbody>
</table>

NR – Not required
APPENDIX 3. Housekeeping Check List for OTs

Before start of OT daily cleaning of parts surrounding

<table>
<thead>
<tr>
<th>Date</th>
<th>OT Table</th>
<th>OT Light</th>
<th>Boyle’s App/Anaesthesia trolley</th>
<th>IV Stand</th>
<th>Cautery Machine &amp; Cautery, Paddle</th>
<th>Instrument trolley (Especially trolley top)</th>
<th>Door Handle</th>
<th>Suction Machine</th>
<th>Hand Washing Area/ Scrubbing Area</th>
<th>AC Point checking</th>
<th>Floor Cleaning</th>
<th>Prepare bleach solution</th>
</tr>
</thead>
</table>

During Surgery

<table>
<thead>
<tr>
<th>Date</th>
<th>Any Spillage</th>
<th>Management of Spill</th>
</tr>
</thead>
</table>

DRAFT
# In between surgery

<table>
<thead>
<tr>
<th>Date</th>
<th>OT table</th>
<th>Patients surroundings</th>
<th>Cleaning of suction tubing and jar</th>
</tr>
</thead>
</table>

# At the end of day

<table>
<thead>
<tr>
<th>Date</th>
<th>OT Table</th>
<th>OT Light</th>
<th>Boyle’s App</th>
<th>IV Stand</th>
<th>Cautery</th>
<th>Machine &amp; Cautery, Paddle</th>
<th>Instrument trolley/ Specially trolley</th>
<th>Door Handle</th>
<th>Cleaning of suction jar followed by sterilization</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Date</th>
<th>Cleaning done by 1.NO(M)</th>
<th>2.HK(M)</th>
<th>1. NO(E)</th>
<th>2. HK(E)</th>
<th>Supervised by</th>
<th>Sister incharge</th>
</tr>
</thead>
</table>

# Weekly cleaning

<table>
<thead>
<tr>
<th>Date</th>
<th>Check all suction and ac points working</th>
<th>Remove all portable items.</th>
<th>Remove dust from inaccessible area with wet mop</th>
<th>Thorough cleaning of surfaces by three bucket system</th>
<th>Wash the OT floor with soap &amp; water</th>
<th>Clean AC filters/ AC ducts</th>
<th>Clean doors, walls, windows</th>
<th>Seal all crevices, holes before fumigation</th>
<th>Replace all portable items back after cleaning</th>
<th>All the AC point sealed</th>
<th>Complete fumigation process as per protocol</th>
</tr>
</thead>
</table>
APPENDIX 4. Daily round format

<table>
<thead>
<tr>
<th>Name of Unit:</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Checklist/ Date</td>
<td></td>
</tr>
<tr>
<td>BMW</td>
<td></td>
</tr>
<tr>
<td>Segregation</td>
<td></td>
</tr>
<tr>
<td>Sharps disposal</td>
<td></td>
</tr>
<tr>
<td>Patient bundle care</td>
<td></td>
</tr>
<tr>
<td>Intra vascular device</td>
<td></td>
</tr>
<tr>
<td>Urinary bag</td>
<td></td>
</tr>
<tr>
<td>Ventilator</td>
<td></td>
</tr>
<tr>
<td>Bed sore</td>
<td></td>
</tr>
<tr>
<td>Dressing Trolley</td>
<td></td>
</tr>
<tr>
<td>Crash cart</td>
<td></td>
</tr>
<tr>
<td>Disinfection &amp; sterilization</td>
<td></td>
</tr>
<tr>
<td>Suction apparatus</td>
<td></td>
</tr>
<tr>
<td>Reusable items</td>
<td></td>
</tr>
<tr>
<td>Others</td>
<td></td>
</tr>
<tr>
<td>Blood spillage Policy</td>
<td></td>
</tr>
<tr>
<td>Cleanliness</td>
<td></td>
</tr>
<tr>
<td>Personnel protec Equip</td>
<td></td>
</tr>
<tr>
<td>Usage</td>
<td></td>
</tr>
<tr>
<td>Availability</td>
<td></td>
</tr>
<tr>
<td>Hand Washing</td>
<td></td>
</tr>
<tr>
<td>Surveillance reports</td>
<td></td>
</tr>
<tr>
<td>Last report</td>
<td></td>
</tr>
<tr>
<td>Date sent</td>
<td></td>
</tr>
<tr>
<td>Who took sample</td>
<td></td>
</tr>
<tr>
<td>Sites for sampling</td>
<td></td>
</tr>
<tr>
<td>Before/ after cleaning</td>
<td></td>
</tr>
<tr>
<td>Remarks</td>
<td></td>
</tr>
<tr>
<td>Name of Sister incharge</td>
<td></td>
</tr>
<tr>
<td>Signature of Sister incharge</td>
<td></td>
</tr>
<tr>
<td>REMARKS</td>
<td></td>
</tr>
</tbody>
</table>

A: Appropriate;  IA: Inappropriate
## Appendix 5. PROFORMA FOR OCCUPATIONAL EXPOSURE TO BLOOD, BODY FLUIDS AND SHARP INJURIES

### Categories of Exposure:

**Mild:** Mucous membrane/non-intact skin with small volumes e.g: a superficial wound (erosion of the epidermis) with a low calibre needle, or contact with eyes mucous membrane, subcutaneous injection following small-bore needle.

**Moderate:** Mucous membrane /non-intact skin with large volumes OR percutaneous superficial exposure with solid needle e.g: a cut or needle stick injury penetrating the gloves.

**Severe:** Percutaneous with large volume e.g.
(a) - an accident with a high calibre needle (>18 G) visibly contaminated with blood.
(b) - a deep wound.
(c) - transmission of a significant volume of blood.

### Categories of Source:

- HIV Negative
- Low risk
- High risk
- Unknown

### Practice of Standard Precautions:

<table>
<thead>
<tr>
<th>Category</th>
<th>Yes/No</th>
</tr>
</thead>
<tbody>
<tr>
<td>First Aid measures:</td>
<td>(Wash / Bleed / Antiseptic / TT)</td>
</tr>
<tr>
<td>Action taken in Casualty</td>
<td>Yes/No</td>
</tr>
<tr>
<td>Hepatitis B Vaccination</td>
<td>Yes/No</td>
</tr>
<tr>
<td>HBIG</td>
<td>Yes/No</td>
</tr>
<tr>
<td>Anti HBSAg Titre</td>
<td>Yes/No</td>
</tr>
<tr>
<td>If yes; Level of antibody</td>
<td>Responder/Non-responder</td>
</tr>
<tr>
<td>PEP advised/taken</td>
<td>Yes/No</td>
</tr>
<tr>
<td>Consent/Signature:</td>
<td>Consent/ Signature:</td>
</tr>
<tr>
<td>Contact no.:</td>
<td>Address:</td>
</tr>
</tbody>
</table>

---

**Signature and Stamp of Unit INCHARGE/CMO**
<table>
<thead>
<tr>
<th>HEALTH CARE WORKER</th>
<th>SOURCE PATIENT</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>TEST</strong></td>
<td><strong>TEST</strong></td>
</tr>
<tr>
<td>HIV</td>
<td>HIV</td>
</tr>
<tr>
<td>HBsAg</td>
<td>HBsAg</td>
</tr>
<tr>
<td>HCV</td>
<td>HCV</td>
</tr>
</tbody>
</table>

* Above tests done by Rapid testing methods only
** All Reactive / Positive results must be correlated clinically and confirmed by ELISA

<table>
<thead>
<tr>
<th>TECHNICIAN:</th>
<th>MICROBIOLOGIST</th>
</tr>
</thead>
<tbody>
<tr>
<td>DATE / TIME:</td>
<td></td>
</tr>
</tbody>
</table>
## Appendix 6. CHECKLIST FOR INFECTION CONTROL ROUND IN DIALYSIS UNIT

<table>
<thead>
<tr>
<th>Action Expected</th>
<th>Expected Frequency</th>
<th>Last 2 dates when complied</th>
<th>Overall compliance (Yes/No/Partial)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>1. HAEMODIALYSIS MACHINE</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AV tubing completely immersed in disinfectant after use</td>
<td>After every use</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Disinfection of Haemodialysis machine with 4% Hypo</td>
<td>Once in a day</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Disinfection of Haemodialysis machine surface area with 1% Hypo</td>
<td>Once in a day</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bleaching of machines with 5% chlorine</td>
<td>Once in a week</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Conductivity test of RO water</td>
<td>Once in a month</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dialysate sterility</td>
<td>Once in a month</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calibration of machine</td>
<td>Quarterly</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Conductivity test</td>
<td>Once in a day</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RO maintenance by backwashes and regeneration of softener</td>
<td>Once in a week</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hardness test</td>
<td>Once in a week</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chloramine test</td>
<td>Once in a week</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Disinfection of RO unit including Loop lines and Storage tanks</td>
<td>Once in a month</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Culture of RO unit output water</td>
<td>Once in a month</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Endotoxin assay of RO water</td>
<td>Once in a month</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Detailed examination of RO water under AAMI guidelines</td>
<td>Quarterly</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>REPROCESSOR MACHINE</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ends of dialyzer connectors dipped in disinfectant</td>
<td>After every use</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of times haemodialyser used</td>
<td>Expected frequency:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Disinfection of reprocessor machine with 1% hypo</td>
<td>Once in a week</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Date of round:
Appendix 7: EOGas Sterilizer Operation Reference Chart

**Warm up**

Make sure sterilizer is connected to power. Sterilizer should come up to temperature in about 15 min (50°C).

**PREPARATION OF ITEMS FOR STERILIZATION**

1. Disassemble and wash with detergent and water.
2. Air-dry; do not oven-dry.
3. Prehumidify items that cannot be washed.
4. Wrap in paper, cloth, or EO permeable film.

**LOAD STERILIZATION BAG**

1. Place wrapped items in medium (#5), or large (#6) sterilization bag.
2. Mark Dosimeter card with time/date of sterilization and place in the core of the load. Include biological control if appropriate.
3. Place fresh Humidichip in sterilization bag.
4. Match EOGas cartridge size (#5, #6) with number on sterilization bag. Remove trigger safety and place EOGas cartridge in sterilization bag.
5. Heat-seal sterilization bag. DO NOT ACTIVATE EOGas CARTRIDGE.
6. Place sterilization bag in sterilizer cabinet.
7. Humidify sterilization bag contents for 2 hours at 50°C.
8. Remove sterilization bag from sterilizer and humidify for an additional 2 hours at room temperature before sterilization.

**BIOLOGICAL CONTROLS**

Challenge the EOGas procedure on the schedule recommended by your governing body or whenever changing packaging materials or techniques. Use a Steritest or other appropriate biological control. Position control in core of load.

**START STERILIZATION CYCLE**

1. Press Load button on sterilizer control panel.
2. When purge cycle is complete, the door is unlocked and the cabinet can be opened.
3. Activate EOGas cartridge within sterilization bag by depressing trigger button.
4. Immediately place sterilization bag into sterilizer cabinet and close door.

1. Leave sterilization bag undisturbed in the sterilizer for 16 hours.
2. Sterilization and aeration proceed simultaneously during the 16-hour cycle.
3. After 16 hours, remove the sterilization bag from the cabinet. Check the Dosimeter; make sure the blue line has progressed beyond the triangular mark.
4. Remove the sterile items from the sterilization bag. Discard the sterilization bag and used cartridge.

STERILIZATION & AERATION

AERATION
The 16-hour EOGas cycle at 50°C includes an aeration period adequate for most EO absorbing materials. Large gas absorbent items, especially implants, or devices that will contact blood or living tissue, require additional, post sterilization aeration.