In order to contain the spread of SARS (severe acute respiratory syndrome), the Ontario Ministry of Health and Long-Term Care directs all hospitals to undertake the following procedures effective immediately:

Preamble

The stringent control measures put in place by the Provincial Operations Committee in March 2003 have had a dramatic impact on the epidemiology of SARS in Toronto. The initial outbreak was characterized by nosocomial transmission. Heightened infection control practices, with modification of clinical activity as well as active surveillance, have significantly decreased hospital-based transmission to both patients and staff.

While nosocomial infections, in particular infections involving healthcare workers, have decreased, they have not disappeared. Many of these infections appear to have occurred during procedures in intensive care units in which staff were exposed to a very high burden of respiratory secretions (high frequency oscillation, difficult intubations, bronchoscopy, non-invasive ventilation, aerosolized therapy etc.).

These directives are based on the current available epidemiology and may be updated as further information becomes available.

High-risk procedures should be avoided whenever possible. However, in situations where such procedures are unavoidable (e.g., intubation of a SARS patient in respiratory distress), staff should follow the additional level of staff precautions outlined in this document for these high-risk procedures.
All staff working in SARS units or with SARS patients must follow the Directives for Acute Care Hospitals regarding infection control measures for SARS units (version 03-05(R) dated April 24, 2003).

Principles:

- High-risk procedures performed on SARS patients expose staff to a high viral burden and must be avoided as much as possible.
- High-risk procedures should be performed:
  - in a private room with negative pressure
  - by the most experienced staff
  - with minimum numbers of staff
  - with strict adherence to SARS precautions and hand disinfection
- ICUs must have access to an infection control consultant to assist with the review of practices.

SPECIFIC DIRECTIVES FOR:

A) ALL PATIENTS DURING THE SARS OUTBREAK (for all patients in all critical care areas in the province)

Protection/Equipment

- For emergency endotracheal intubation of non-SARS patients, ensure that each patient unit has:
  - a manual resuscitation bag with bacterial/viral filter
  - in-line suction catheters
  - intubation equipment
  - full protective apparel for the individual performing the intubation, and for the respiratory therapist. This includes gowns, gloves, flip down hoods or full-face shields with N95 masks or equivalent. Masks must be fit tested according to existing guidelines.
- Protective apparel must be removed carefully at the end of the procedure to reduce the risk of contamination and re-aerosolization of droplets.

Procedures

- Nebulized therapies should be avoided. Ventolin or atrovent can be delivered using the metered dose inhaler and aerochamber.
- Oxygen should be delivered DRY avoiding nebulized humidity. Maximum flow rate for nasal prongs should be 6 litres per minute.
- If a patient requires up to 50% oxygen by mask use a venti-mask. If a patient requires more than 50% oxygen then the respiratory therapist (RT) should be notified. The nebulizer system shall be emptied of the water from the prefilled
water bottle. The water bottle should remain DRY. The RT will monitor the patient and wean to nasal prongs as soon as the patient can tolerate.

- Patients should receive frequent mouth care.
- Patients with tracheostomies should be provided with humidity.
- Patients who require oxygen greater than 50% should be referred to RT for set up and ongoing monitoring.
- High frequency oscillation and non-invasive ventilation (CPAP/BiPAP) should be avoided. If ventilation is essential for the patient, the patient should be screened in consultation with infectious diseases/infection control staff, to ensure that a diagnosis of SARS has been ruled out. The procedure should be performed in a private room.

B) SARS PATIENTS IN CRITICAL CARE AREAS

1. Bronchoscopy should be avoided if possible in patients known or suspected to have SARS.

2. Equipment:

   - In the ICUs and SARS units and other high risk areas identified by the hospital, include with the arrest cart (crash cart):
     - manual resuscitation bag with bacterial/viral filter
     - in-line suction catheters
     - personal protection system (PPS) – an apparatus consisting of head, face and neck protection with or without enclosed body protection

3. Intubation and bronchoscopy:

   Personal Protection:
     - Those performing the intubation should wear a full head, face and neck protection. This may consist of positive airway pressure respirator (PAPR) or another type of PPS (see appendix “Parameters to Guide the Selection of Personal Protective Systems”).
     - The system chosen should allow for safe performance of the procedure and not fog when in use.
     - Staff must be trained in the use of the specific type of PPS chosen.

   Use of the Positive Airway Pressure Respirators and Personal Protection Systems:
     - An N95 mask or equivalent should be worn underneath the respirator and be left in place once the respirator hood is removed until staff has left the room.
     - Staff using this equipment must receive proper instruction on the application and removal to avoid contamination.
     - A practice session should be carried out prior to use and written instructions should be given to staff. Staff training sessions must be documented. The
hospital Infection Control Practitioner must review the written procedure/instructions.

- Ensure that all disposable components of the equipment are carefully removed at the end of the procedure and reusable items are thoroughly cleaned using hospital disinfectant or disinfectant wipes.
- The application and removal of PAPR/PPS equipment requires the assistance of another person and should not be done alone.

**Personnel:**

- The procedure should be performed by the most experienced staff members available. The number of persons in the room should be kept to a maximum of 2-3 persons (note: hospitals may wish to consider a SARS intubation team).

**Procedure:**

- The procedure should be done in a negative pressure room. If none are available, it must be done in a private room with the door closed.
- After hand-washing and prior to entering the room, the code team will apply the personal protective equipment as per directive 03-05 (April 23, 2003) and manufacturer’s instructions.
- Staff performing the intubation will apply the personal protection system (PPS).
- The intubation should be done while the patient is sedated and paralysed if medical condition permits.
- The ventilator and in-line suction device should be in the patient room to reduce time needed for bag ventilation and disconnecting bag from the endotracheal tube suctioning.
- Remove protective equipment following directive 03-05 (April 23, 2003) and manufacturer’s instructions after intubation.
- Minimize re-entry to the room by staff for approximately 2 hours post procedure.

**Cleaning:**

- Excess medications should be discarded at the end of the procedure.
- Immediate clean up of room and equipment should be done slowly and in such a way as to reduce the re-release of aerosols.
- Potentially contaminated surfaces in the room should be wiped with a hospital-approved disinfectant.

**4. Management of SARS patients with mechanical ventilation:**

**Note:** Infectious respiratory secretions from SARS patients will contaminate respiratory equipment and be expelled into the surrounding environment.

**Procedure:**

- Ventilators
- A hydrophobic submicron filter must be placed between the endotracheal tube and the ventilator circuit tubing.
- A second bacterial/viral filter must be placed in the expiratory circuit of the ventilator.
- Filters should be changed when fluid build-up impedes ventilation (at least every 24 hrs).
- Disposal of filters should be considered a high-risk exposure and staff must protect themselves using full personal protective equipment following the maximal precautions policy.
- Filters and respiratory circuits for known SARS cases should be single use and disposed of after use.
- Filters are to be bagged, sealed, and then placed in a biohazardous bag for disposal.
- Heated wire circuits should be used on both the inspiratory and expiratory sides of the circuit.
- A water trap/filter combination should be placed at the end of the expiratory circuit.

- Manual Resuscitation Bags:
  - A hydrophobic submicron filter must be placed between the endotracheal tube and the bag.
  - Filters should be changed when fluid buildup impedes ventilation (at least every 24 hours).
  - Disposal of filters should be considered a high-risk exposure and staff must protect themselves following maximum precautions using full SARS protective equipment.
  - Filters are to be bagged, sealed, and then placed in a biohazardous bag for disposal.
  - Equipment used for manual bagging should be disposed of after use, not cleaned.

C) PATIENTS WITH RESPIRATORY SYMPTOMS OR UNEXPLAINED FEVER AND UNKNOWN SARS RISK

1. Treat as SARS until another diagnosis has been confirmed. Follow all policies as described above until that time.
D) PATIENTS WITH NO RESPIRATORY SYMPTOMS, OR WITH RESPIRATORY SYMPTOMS/FEVER DUE TO A KNOWN CAUSE OTHER THAN SARS:

1. Treat as per current hospital policy regarding non-SARS patients.

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APPENDIX

Parameters to Guide the Selection of Personal Protective Systems

1. Provides barrier precautions for droplets/splashing and completely covers all of the face and head, or can be easily combined with other protective apparel to provide full coverage.

2. Provides filtration at <0.3 micron with 95% filter efficiency. For hooded devices, the circulating air within the hood should not affect the wearing of a N95 mask nor impede its effectiveness.

3. Hooded devices should be able to create positive pressure.

4. Consideration should be given to the extent of CO₂ build up within hood or respirator. Current guidelines recommend that CO₂ should not exceed 5000 ppm as a time weighted average (TWA).

5. Where applicable, the system should be able to provide full fit testing, similar to N95 masks.

6. The equipment should be able to be assembled with little chance of error, and disassembled easily.

7. Ability to clean the surface of the equipment with hospital grade disinfectants. Single use of high-risk components is preferred, or the product is easily disassembled and cleaned and tolerates high-level disinfection or sterilization.

8. The filter for any system should be easy to remove and dispose.

9. Demonstrable ease of donning with minimal amount of time.

10. Ability to remove equipment with minimal contamination of the wearer and the equipment.

11. The device provides a good field of vision, and a clear view with no distortion in order to perform procedures such as intubation and bronchoscopy.

12. The device should not interfere with communications between team members and allow for clinical assessments of patients such as auscultation.

13. Wearer is able to easily perform procedures based on ergonomic factors.
14. The system or device should be comfortable to wear for at least a continuous 2-hour period.

15. The equipment should be easily worn and managed by staff of varying sizes.

16. The equipment or device should allow for the wearer to remain cool and comfortable.

Examples of Personal Protective Systems:
- 3M PAPR Hood
- Stryker T4 System
- Full face respirators

A listing of powered air-purifying respirators can be found by doing the following two steps:
http://www2.cdc.gov/drds/cel/cel/_form.asp
Highlight “HEPA (PAPR only)” in the section with the heading “For Protection Against:”
Click on “View Results” at the bottom of the page on the screen.