Section 2 Respiratory and Direct Contact



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This section provides a general overview of the communicable diseases transmitted through respiratory and direct contact. The information in this introduction provides both general considerations and key concepts. Specific procedures and information are included within each disease chapter.

Objectives

- 1. Individuals infected with organisms that are transmitted through respiratory (droplet or aerosolization) and direct contact will be identified, investigated and managed in a timely manner.
- 2. The complications secondary to infection will be minimized in individuals through the timely identification, investigation and implementation of control measures.
- 3. Outbreaks will be contained through the timely identification of the source and contacts and through the implementation of prevention and control measures.
- 4. Health authorities will work to deliver immunization programs according to the provincially funded recommended immunization schedule. Immunization coverage rates are a useful indicator to evaluate programs and plan service delivery to provide the best protection to the population.
- 5. Information will be managed in a confidential manner and will be shared in accordance with Appendix B Interjurisdictional Communication, *The Public Health Act*, 1994 and *The Health Information Protection Act* and their respective regulations.
- 6. Information that is required for notification purposes will be extracted from the electronic case management system.

Background

As a group, acute respiratory diseases are one of the leading causes of death from any infectious disease (Heymann, 2008). It has been demonstrated that Canada's ability to fight an outbreak, such as Severe Acute Respiratory Syndrome (SARS), is more closely tied to specific strengths of the public health system than to the health system that provides individual health services. Key aspects of the public health system include the capacity to detect, prevent, understand, and manage outbreaks of significant infectious diseases.



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An effective response to an outbreak of severe respiratory illness requires cooperation and collaboration among all jurisdictions – regional, First Nations, provincial, national, and international partners to ensure the timely mitigation of risk and prevention of further illness.

Reporting Requirements

See <u>Reporting Requirements in General Information - Section 1</u> of the manual for guidelines. Refer to Appendix A – Reporting and Follow-up Timelines.

Methods of Control

Primary Prevention

Many of the organisms that cause respiratory diseases are spread via respiratory droplets generated by coughing and sneezing. These organisms are also spread from person to person when they are in close contact with one another or through touching something with organisms on it and then touching their mouth or nose. "In shelters and other homeless service programs large numbers of people may live together and regularly move in and out. People often share sleeping and bathroom facilities. This means people may have contact with others who have an infection" (Toronto Public Health, 2006).

Coughing and sneezing can also generate small airborne particles that can be inhaled causing infection in the recipient. In general, the following measures are the best way to avoid contact with respiratory droplets or secretions.

Hand Hygiene

"Proper handwashing with soap and water is an important barrier to many infectious diseases and promotes better health and well-being" and "handwashing is one of the most practical and effective ways of preventing the spread of disease" (World Health Organization as quoted by College of Registered Nurses of Manitoba, 2010).



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This is especially important after touching surfaces or objects that might be contaminated with respiratory droplets, or after touching persons who are ill with respiratory symptoms. Alcohol hand gels are an adequate substitute when soap and clean water are not readily available and your hands are not visibly soiled. Alcohol gels are not effective if hands are soiled with protein material. Refer to Attachment – Handwashing.

Personal Protective Measures

Avoiding crowds and practicing respiratory hygiene and cough etiquette can help reduce the spread of respiratory illnesses:

- cover the nose and mouth when coughing or sneezing;
- tissues should be used to contain secretions and should be properly disposed of at the earliest opportunity;
- practice hand hygiene after handling items that may be contaminated with respiratory secretions.

Avoiding sharing of personal items (eating/drinking utensils, towels, toothbrushes, etc.) may reduce the risk of transmission of bacteria and viruses. Bacteria and viruses that cause respiratory illness may survive on hard non-porous surfaces and be transmitted to others, via hand contamination and self-inoculation. These surfaces should be cleaned and sanitized on a regular basis, especially when people are sick.

Immunization

A number of communicable diseases transmitted by respiratory contact and direct contact are preventable through vaccination. "Vaccination programs are considered to be the most cost-beneficial health intervention and one of the few that systematically demonstrate far more benefits than costs" (Health Canada, 2002, p. 2). Immunization history should be obtained from and reviewed with clients. Every effort should be made to update a client's immunizations as per the recommendations in the Saskatchewan Immunization Manual ¹



¹ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

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Secondary Prevention

Asymptomatic carriage can serve as a source of infection to others. When a case occurs in a setting with susceptible individuals (i.e., long term care facilities), active screening of individuals may be an appropriate action so prevention and control measures can be implemented to interrupt the chain of infection.

The specific level of intervention and contact tracing varies according to the disease and should be individualized based on the guidelines in the specific agent.

General Guidelines for Investigation of Diseases Transmitted through Respiratory and Direct Contact

These guidelines aim to assist in the collection of information and define control measures for organisms that are transmitted through respiratory and direct contact. Refer to the <u>General Information - Roles of Stakeholders</u> section of the manual. Appendix C - Major Legislation identifies the significant Acts and Regulations that are applicable to Communicable Disease Control in Saskatchewan. The following questions/guidelines² can assist you to determine the approach you will need to follow to prevent and control the disease.

- 1. What is the source of the disease? Can it be identified? Communication with the case is important to determine the **risk factors**, **exposures** and **potential exposures of others** to the disease.
- 2. Who else may have been exposed to the disease? When determining the possible source and possible contacts exposed, the incubation period, mode of transmission and period of communicability are important considerations. Key considerations include:
 - Recent exposure to someone else who is sick with similar symptoms.
 - Travel history.
 - Attendance in childcare, school, daycare, healthcare settings.
- 3. Is an outbreak present?



² These questions were adapted from http://www.health.gov.nl.ca/health/publications/diseasecontrol/dcresp.pdf

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- 4. What is the potential impact of the disease for the individual? Their household/family? Their community?
- 5. Are there people who are more likely to develop symptoms or serious manifestations of the infection?
- 6. Is there a population who are more likely to be susceptible to the infection?
- 7. What interventions are available to prevent the transmission of the infection? Refer to disease specific measures and implement necessary activities.
- 8. Is there a high risk for transmission to others (e.g., highly communicable agent, common vehicle for transmission such as food or water, etc.)? Determine if this individual is in a situation where there is a high risk of transmission of the organism (childcare, health care worker, environmental conditions conducive to transmission, etc.). Who else may have been exposed to the disease? Conduct contact tracing to:
 - Determine if the contact is in a high-risk group.
 - Inform contacts of any prophylaxis and/or exclusion measures:
 - information that should be gathered from the contacts relates to their level of risk, the need for testing, the potential benefit of prophylaxis (as detailed in the disease sections specifically) and immunization history.
 - interventions such as **exclusion/isolation/quarantine** may be appropriate depending on the nature of the disease and the status of the contacts that have been identified.
- 9. Educate case and contacts regarding:
 - The nature of the disease including the incubation period, period of communicability, mode of transmission, etc.
 - Self-care measures.
 - Personal protective measures, which should always include hand washing, not sharing personal items (eating and drinking implements, towels, lip balms, etc.).
 - Disease control measures they must follow (treatment, exclusion, etc.).
 - Publicly funded treatment and chemoprophylaxis is indicated for certain diseases such as meningococcal disease, tuberculosis and *Haemophilus influenzae* type b invasive disease. Publicly funded control measures may be used in long-term care facilities in the event of an influenza outbreak.



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10. Obtain an immunization history from case and appropriate contacts.

- Immunizations should be offered to cases and contacts that are not up-to-date or who are eligible for vaccines as per the Saskatchewan Immunization Manual³ Chapter 5: Immunization Schedules and Chapter 7: Immunization of Special Populations.
- Depending on the organism and other circumstances, it may be prudent to offer immunization for the disease for both the case and the contact(s).
 Refer to disease section for details.
- 11. Document case management and follow-up information on the electronic case management and surveillance system.
- 12. Communication with other stakeholders (physicians, acute and long term care, schools, daycares, etc.) is vital for a coordinated and efficient response to a single case or an outbreak of communicable disease. Maintaining **confidentiality** according to the corresponding legislation is important.

Special Considerations

Certain individuals and certain environments may be considered higher-risk. The following sections outline some circumstances that may need to be considered when doing your investigation.

Immunocompromised/Immunosuppression

The status of an individual's immune system may have an impact on the individual's response to the disease. When there are circumstances that have an impact on an individual's immune system, additional interventions may be required.

Elderly and Infants

Elderly people and infants may be more susceptible to some communicable diseases. Some organisms are also more virulent in these individuals.



³ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

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Immigrants/Refugees

Depending on the diseases required to be screened for under immigration regulations and adherence with treatment regimes prior to emigrating, the immigrant or refugee may carry with them diseases specific to their country of origin. Additionally, these individuals may be more susceptible to certain diseases as immunization programs in their country of origin may be different from Saskatchewan or Canadian standards.

Individuals with Suboptimal Personal Hygiene Practices

Individuals with poor practices of personal hygiene (i.e., mentally or physically handicapped) may serve as a vehicle of transmission due to the lack of self-care measures that are useful in interrupting the chain of infection.

Child Care Centres

Young children have limited ability to implement the individual measures to reduce the risk of spread of diseases. This provides an increased opportunity for transmission. This also necessitates early identification and diligent infection control practices. Refer to the Saskatchewan Ministry of Health Infection Control Manual for Child Care Facilities. This serves as an excellent resource for daycare settings to assist in minimizing the risk and spread of communicable diseases.

Health Care Facilities and Institutional Settings

Health care facilities present as a high-risk environment for two reasons:

- 1. Typically, the clients/patients within the facility are there because either they have a medical condition that puts them at greater risk for contracting an infection or they are already infected and experiencing complications of a communicable disease.
- 2. Health Care Workers serve as a vehicle for transmission of a communicable disease to a high-risk individual.

To avoid this, familiarity with and adherence to Infection Control Guidelines and Practices is of paramount importance.

Travel



⁴ http://www.saskatchewan.ca/live/births-deaths-marriages-and-divorces/starting-a-family/early-learning-and-child-care/child-care.

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Travel to destinations where one can be exposed to communicable diseases that are not common or endemic in Saskatchewan or Canada.

Environments Where Individuals are in Close Proximity to Others

This may be related to crowded living conditions such as multi-family homes and homeless shelters. It may also be related to environments where people are in close proximity to groups of people such as in schools, airport/bus terminals, public transportation vehicles, etc.

Public Health Agency of Canada (2007) states "Statistics Canada uses the measure of persons per room (PPR) to assess crowding in houses. PPR is calculated by dividing the number of persons living in a dwelling by the number of rooms." Rooms refers to all rooms within a dwelling excluding bathrooms, halls, vestibules and rooms used solely for business purposes. This statistic is not sensitive to the size of the house or the rooms, or to the composition of the household (age of occupants, etc.). For diseases transmitted through the respiratory route it is found that the higher the number of persons per room, the greater the risk for transmission within the household. Greater than one person per room puts the occupants at greater risk for these illnesses.



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Notification Timeline:

From Lab/Practitioner to Public Health: Immediate.
From Public Health to Ministry of Health: Immediate.

Public Health Follow-up Timeline: Initiate within 24-48 hrs.

Information

Case Definition (Public Health Agency of Canada, 2008)

Confirmed Case	Clinical illness* or systemic manifestations compatible with diphtheria								
	in a person with an upper respiratory tract infection or infection at								
	another site (e.g., wound, cutaneous) PLUS at least one of the								
	following:								
	• Laboratory confirmation of infection:								
	 isolation of Corynebacterium diphtheriae with confirmation of toxin from an appropriate clinical specimen,¹ including the exudative membrane 								
	OR								
	 isolation of other toxigenic Corynebacterium species 								
	(<i>C. ulcerans</i> or <i>C. pseudotuberculosis</i>) from an appropriate clinical specimen, including the exudative membrane								
	OR								
	 histopathologic diagnosis of diphtheria. 								
	OR								
	Epidemiologic link (contact within two weeks prior to onset of								
	symptoms) to a laboratory-confirmed case.								
Probable Case	Clinical illness* in the absence of laboratory confirmation or epidemiologic link to a laboratory-confirmed case.								
Suspected Case	Upper respiratory tract infection (nasopharyngitis, laryngitis or								
Suspected Case	tonsillitis) with or without a nasal, tonsillar, pharyngeal and/or								
	laryngeal membrane.								
*	raryingear memorane.								

Clinical illness is characterized as an upper respiratory tract infection (nasopharyngitis, laryngitis or tonsillitis) with or without an adherent nasal, tonsillar, pharyngeal and/or laryngeal membrane, plus at least one of the following:

- gradually increasing stridor;
- cardiac (myocarditis) and/or neurologic involvement (motor and/or sensory palsies) one to six weeks after onset;
- death, with no known cause.



Refer to Specimen Collection and Transport for details on appropriate clinical specimens.

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Causative Agent

Corynebacterium diphtheriae (C. diphtheriae), a gram positive bacillus. Diphtheria is caused by toxigenic strains of the bacteria C. diphtheriae of gravis, mitis or intermedius biotypes.

Symptoms

The various clinical forms of diphtheria are caused by an exotoxin produced by toxigenic strains of the bacteria; all toxigenic strains produce an identical toxin. Toxin production occurs following infection of a *C. diphtheriae* strain by a corynebacteriophage containing the tox gene.

Non-toxigenic strains can also produce a mild, localized disease resembling that caused by toxigenic strains.

- Infections that are not apparent tend to outnumber clinical cases, and both toxigenic and non-toxigenic strains of *C. diphtheriae* may be harboured in the nasopharynx, skin, and other sites of asymptomatic carriers.
- Pharyngeal diphtheria is a febrile illness beginning with a low-grade fever, a sore throat, and a yellow-white discharge over the tonsils, uvula, and throat. This discharge becomes grey, patchy, and membranous and may involve the larynx, where it can present an airway obstruction, particularly in infants and young children. There may be marked edema of the neck (classic bull neck appearance).
- <u>Nasal diphtheria</u> is often a mild form of the disease and is characterized by one-sided nasal secretions.
- Diphtheria may also present as a cutaneous, vaginal, or conjunctival infection.
- <u>Cutaneous or mucous membrane diphtheria</u> is usually found in warmer climates or among the homeless and will present as a shallow ulcer coated with a pseudomembrane.

Complications

- Affects distant tissues and organs after 2 to 6 weeks, in particular cranial and peripheral motor and sensory palsies, and myocarditis.
- A case-fatality rate of 5% to 10% is reported for non-cutaneous diphtheria, with the highest rates among the very young and the elderly (Manitoba Health, 2001).



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Incubation Period

Usually 2-5 days, sometimes as long as 10 days.

Reservoir/Source

Humans are the only reservoir - harboured in the nasopharynx, skin, and other sites.

Mode of Transmission

- Direct transmission of toxigenic strains or indirect transmission by transfer of the bacteriophage from a person infected with a toxigenic strain to a non-toxigenic strain in a carrier.
- Contact with nasopharyngeal secretions of a case or carrier.
- Rarely, contact with articles soiled with discharges from infected skin lesions.
- Raw milk has also served as a vehicle for transmission.

Period of Communicability

- Variable, until virulent bacilli have disappeared from discharges and lesions. For example:
 - effective antibiotic therapy promptly ends shedding within 4 days;
 - without treatment, infectivity usually last 2 weeks or less (seldom more than 4 weeks):
 - the rare chronic carrier may shed organisms for 6 months or more.

Specimen Collection and Transport

The diagnostic specimen is a throat swab in reduced charcoal transport medium.

Material for culture should be obtained by collecting throat swabs and placing them in Amies transport medium. Swabs should be taken from the inflamed areas of the throat and nasopharynx in symptomatic patients. Swabs should be taken for culture before antibiotic therapy is initiated. Confirmatory diagnosis requires isolation and identification of the organism, and toxigenicity testing, and may take several days.

If cutaneous diphtheria is suspected, swabs should be collected from the base of the lesion. Specimens should be transported as soon as possible.

Do not wait for laboratory results before initiating treatment.



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Methods of Control/Role of Investigator

Prevention and Education

Refer to the <u>Respiratory and Direct Contact Introduction and General Considerations</u> section of the manual that highlights topics for client education that should be considered and as well as provides information on high-risk groups and activities. As diphtheria is a vaccine-preventable illness, attention should be placed on immunization.

Immunization

Immunize infants, children and adults according to the recommended schedule. Refer to Saskatchewan Immunization Manual.¹

Education

- Education should be provided regarding respiratory etiquette and measures to prevent transmission.
- Educate the public about the disease and the need for active immunization. Immunization information fact sheets can be used to guide discussion.

Management

See <u>Attachment – Recommendations for the Management of Diphtheria Cases and Contacts Algorithm.</u>

I. Case

Collaborate with the primary care provider to determine respective roles and responsibilities (e.g., contact tracing, education, and follow-up).

History

- Determine case status including a review of the immunization history. Do not wait for laboratory results before initiating treatment.
- Obtain travel history or history of immigration within the past week.
- Identify contacts (refer to Contact Definition).
- Refer to <u>Attachment Diphtheria Case Investigation Worksheet</u> to guide followup.



¹ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

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Treatment/Supportive Therapy

- **Diphtheria Antitoxin** For pharyngeal diphtheria, early administration of diphtheria antitoxin is recommended to neutralize the circulating diphtheria toxin. It should be given in the early stages if diphtheria is suspected. "The site and size of the diphtheria membrane, the degree of toxic effects, and the duration of illness are guides for estimating the dose of antitoxin" (American Academy of Pediatrics, 2009, p. 281). Dosage should be coordinated by the clinician, infectious disease (ID) specialist and Medical Health Officer (MHO). Diphtheria antitoxin can be obtained from Population Health Branch, Saskatchewan Ministry Health. See Appendix D Publicly Funded Medications for Chemoprophylaxis/Treatment.
- Antimicrobial therapy is not a substitute for antitoxin treatment.
- Likewise, antitoxin treatment is not a substitute for antibiotic therapy (Health Canada, 1998).
- Antimicrobial Therapy Treatment choices are governed by the most recent guidelines. The public health practitioner should direct any questions regarding the current treatment protocols to the physician or MHO. Refer to Heymann, other texts and clinical treatment guidelines for specific treatment details. See Appendix H Sources for Clinical Treatment Guidelines. Heymann (2008) indicates the following:
 - Procaine penicillin G IM or parenteral erythromycin is recommended until the patient can swallow comfortably, at which point the treatment may be given orally. A total of 14 days treatment is recommended.
 - Supportive treatment, in hospital or home is advised under strict isolation involving routine contact precautions for cutaneous and droplet precautions in instances of pharyngeal until 2 consecutive throat cultures are negative for diphtheria bacilli. These cultures should be taken not less than 24 hours apart and not less than 24 hours after the completion of a 14-day course of antibiotics
- For cutaneous diphtheria, the skin lesions should be cleaned with soap and water, and a course of oral antibiotics should be given for a 10-day period. Antitoxin may be of some use in cutaneous disease, because of toxic sequelae (American Academy of Pediatrics, 2009).

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Immunization

Immunization against diphtheria should begin during convalescence because there is no guarantee that immunity to diphtheria is conferred by natural infection.

Exclusion

Exclude and isolate <u>all</u> cases from work, school, daycare, and other public environments using precautions appropriate to the site of infection until two cultures (nasal and pharyngeal) taken 24 hours apart and at least 24 hours after completion of a 14-day course of appropriate antibiotics, are negative.

Referrals

To the appropriate specialist(s) including an ID specialist.

II. Contacts/Contact Investigation

Refer to <u>Attachment – Diphtheria Contact Investigation Worksheet</u> to guide follow-up.

Contact Definition

- Close contacts are defined as:
 - household members;
 - friends, relatives, and caretakers who regularly visit the home;
 - kissing and/or sexual contacts;
 - those who share the same room at school or work;
 - healthcare staff exposed to oropharyngeal secretions of the infected person (staff who have taken appropriate isolation precautions need not be considered contacts).

Follow up of contacts involves:

Education

• All contacts (or their parents if children are contacts) should be provided with information on the disease, risk factors, prevention and necessary follow-up tests, treatments, and exclusion requirements.



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Testing/Prophylaxis

- Collect appropriate screening and case-finding specimens (see <u>Specimen Collection</u>). Samples for culturing should be taken from nasal and pharyngeal swabs **before** antibiotic treatment is started (Health Canada, 1998).
 - A single intramuscular dose of benzethine penicillin G or a 7 to 10 day course of oral erythromycin is recommended for all close contacts exposed to diphtheria regardless of their immunization status (Heymann, 2008).
 - If carrier status is determined, refer to <u>Carrier Management</u>.
 - Follow-up surveillance should continue for 7 days. "All close contacts should be kept under daily surveillance for 7 days from the date of last contact with the case and assessed clinically for signs and symptoms of diphtheria" (Health Canada, 1998).

Immunization

- Assess the immunization status of all contacts.
- Previously immunized contacts who have not received a booster dose within 10 years should receive a booster dose of diphtheria toxoid.
- Under-immunized contacts should have a primary series initiated.

Exclusion

- Exclude under-immunized contacts from school, daycare, health care, and food handling until 2 cultures taken 24 hours apart and at least 24 hours after completion of a 14 day course of appropriate antibiotics, are negative.
- Exclude adult contacts from the workplace until bacteriologic examination proves them not to be carriers for those occupations that involve handling food (especially milk) or close association with under-immunized adults or children (Heymann, 2008).
- Keep all close contacts under active daily surveillance for signs and symptoms for 7 days. Refer to <u>Attachment Diphtheria Contact Investigation Worksheet</u>. Exclude anyone who becomes symptomatic or whose cultures return positive (Heymann, 2008).



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III. Carrier Management Carrier Definition

One who harbours, and may disseminate, the bacterium without discernable clinical disease.

Testing

- Follow-up pharyngeal cultures should be obtained from contacts proven to be carriers at a minimum of 2 weeks after completion of therapy.
- If cultures are positive, an additional 10-day course of erythromycin should be given.

Treating

• For carriers, a single intramuscular dose of benzathine penicillin G (600,000 units for persons < 6 years of age, and 1.2 million units for persons ≥ 6 years of age), or 7 to 10 day course of oral erythromycin (40 mg/kg/day to a maximum of 1 g/day for children, and 1 g/day) divided in 4 doses for adults has been recommended (Heymann, 2008).

Immunization

• Ensure appropriate immunizations are up-to-date.

Exclusion

- Standard and droplet precautions should be observed for hospitalized carriers with cutaneous diphtheria until 2 negative cultures are obtained from lesions at least 2 weeks after completion of antibiotics.
- Carriers should be excluded from food handling and working with children who
 are under-immunized until 2 negative cultures have been obtained after
 completion of antibiotics.
- Carriers should pay strict attention to personal hygiene, particularly:
 - respiratory etiquette;
 - hand hygiene;
 - keeping infected wounds covered.



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IV. Environment

Child Care Centre Control Measures

Although an outbreak of diphtheria would be rare, if it occurs the following should be implemented:

- Provide information (not personal information) to the parents of the children in the daycare or school. See Attachment Diphtheria Template Letter to Parents.
- An immunization information sheet containing diphtheria can be provided and used as a guide.

Children who have not completed the primary series of immunization against diphtheria should begin, or finish the series. These children should be referred to their local public health office.

Institutional Control Measures

- Consultation between Public Health/MHO and infection control staff is important.
- Strict isolation of cases in hospital until two consecutive negative cultures are obtained from throat and nasopharyngeal swabs are obtained at least 24 hours apart and at least 2 weeks after completion of antibiotic therapy. If cultures are difficult to obtain, isolation should be not be discontinued until 14 days after the beginning of antibiotic therapy.
 - Droplet precautions should be used for pharyngeal diphtheria.
 - Contact precautions should be used for cutaneous diphtheria.

Epidemic Measures

Immunize the largest possible proportion of the population group involved especially infants and preschool children.

If unimmunized adults are affected, immunize the groups most affected and individuals at high risk of exposure to cases. Provide a second dose of vaccine one month later to ensure two doses are received.

Travellers to countries where epidemics occur should have their diphtheria status reviewed and updated when necessary (American Academy of Pediatrics, 2009).



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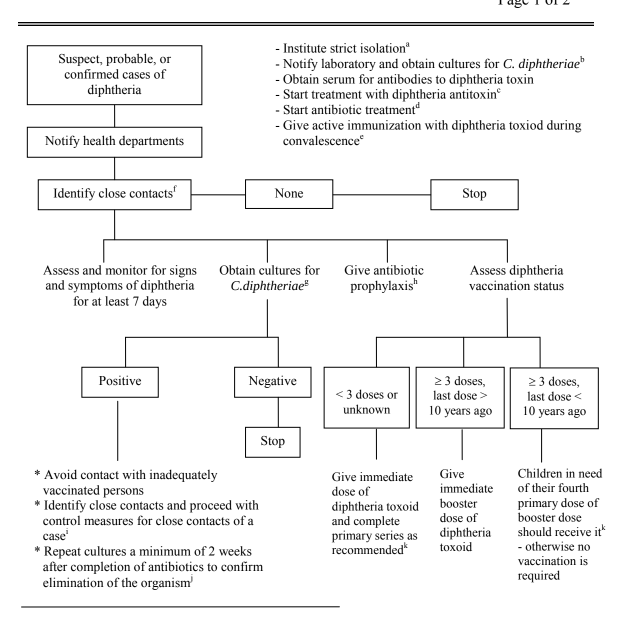
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Diphtheria

Attachment – Recommendations for the Management of Diphtheria Cases and Contacts Algorithm

Reviewed: October, 2010 Section: 2-30 Page 1 of 2



Adapted from CDC Diphtheria Worksheet which was based on Farizo et al. (24), Clinical Infectious Diseases 1993, 16:59-68.

Diphtheria

Attachment – Recommendations for the Management of Diphtheria Cases and Contacts Algorithm

Reviewed: October, 2010 Section: 2-30 Page 2 of 2

- a. Strict isolation with contact and droplet precautions for all potentially infectious cases, as well as a private room and the use of masks, gowns, and gloves for all persons entering the room. Maintain isolation until elimination of the organism is demonstrated by negative cultures of two samples obtained at least 24 hours apart after completion of antimicrobial therapy.
- b. Both nasal and pharyngeal swabs should be obtained for culture.
- c. The recommended dosage and route of administration of antitoxin depends on the extent and duration of disease. Refer to Guidelines for the Control of Diphtheria in Canada at http://www.phac-aspc.gc.ca/publicat/ccdr-rmtc/98vol24/24s3/24s3e_e.html for detailed dosage recommendations for equine diphtheria antitoxin.
- d. Antibiotic therapy is not a substitute for antitoxin treatment. **Refer to Guidelines for the Control of Diphtheria in Canada for detailed antibiotic dosage recommendations for cases.** Eliminations of *C. diphtheriae* should be confirmed by two negative cultures of throat and nasopharyngeal swabs taken at least 24 hours apart, a minimum of 2 weeks after antibiotics are completed. Persistent carriage of the organism should be treated with an additional 10-day oral course of erythromycin with follow-up cultures.
- e. Vaccination is required because clinical diphtheria does not necessarily confer immunity.
- f. Close contacts include household members and other persons with a history of direct contact with a case (e.g., caretakers, relatives, or friends who regularly visit the home) as well as health-care personnel exposed to oral or respiratory secretions of a case.
- g. Both nasal and pharyngeal swabs should be obtained for culture. Swabs should also be taken from any wounds or skin lesions.
- h. Antibiotic therapy is not a substitute for antitoxin treatment. Refer to Guidelines for the Control of Diphtheria in Canada for detailed antibiotic dosage recommendations for contacts and carriers.
- i. Control measures for contacts of a case should be given a higher priority than control measures for contacts of a carrier.
- j. Eliminations of *C. diphtheriae* should be confirmed by two negative cultures of throat and nasopharyngeal swabs taken at least 24 hours apart, a minimum of 2 weeks after antibiotics are completed. Persistent carriage of the organism should be treated with an additional 10-day oral course of erythromycin with follow-up cultures.
- k. Refer to the Saskatchewan Immunization Manual at http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx or NACI for recommendations for schedule of vaccination.



Diphtheria Attachment – Diphtheria Case Investigation Worksheet

Reviewed: October, 2010 Section: 2-30 Page 1 of 3

Please see the following pages for the Diphtheria Case Investigation Worksheet.



Diphtheria Case Investigation Worksheet

Shaded areas are mandatory for reporting to Saskatchewan Ministry of Health [Indicates field in iPHIS] Please use YYYY/MM/DD for all dates

	Date Reported		Name (Last,	Name (Last, First)							HSN			
	Birth Date	Age	🔲 1	Male Female Jnknown	Pregnar Yes No Unk		Ethnicity Arab/West Asia Asian Black Inuit Latin-American Métis		Sou Wh		orth American Indian uth Asian hite iknown her:			
	Address (Street and No.) City Province		e	Post	tal Code	Pho	Phone							
If residential facility or daycare please indicate name:														
	Date Symptom Onset	Date Diag		Date Hospital	lized		His	story of immur	nization aş	ation against diphtheria				
MATION	or lab diagnosis)		Childhood prin series? Yes No Unknown	nary	If < 18 years old, number of doses?	adult' Y N	es	Date of last dose or Unknown						
PATIENT INFORMATION	Description of Clinical Picture)]]]	Dutcome Recovered, no Recovered, res Unknown Died – Date:	sidual	effects	□ P	Diphtheria as cause of death: Primary Contributing Incidental				
	Symptoms Fever			<u>:</u>	Signs				Complications					
				nx e	☐ Mid ☐ To c ☐ Belo ☐ Stridor ☐ Wheezing ☐ Palatal wea ☐ Tachycardi ☐ EKG abnor	embra teral side on t side mandi way to laviel ow cla kness a maliti	only only bular only o clavicle e vicle	Airway obstruction Date of onset: Intubation/traech required Myocarditis Date of onset: (Poly)neuritis Date of onset: Other Describe:						
ATORY	Specimen culture diphtheria? Yes or No Unknown	e for	If Yes, date sp	ecimen obt	tained:	Culture result? Positive Negative Unknown		ime of lab perf lture:	Forming	☐ Mi ☐ Gra	ure positive, biotype? tis avis ermedious Ifanti			
LABORATORY	If culture positive Positive Negative Unknown Not done	oxigenicity tes	ting?	Type of specimen? (check all that apply) Clinical swab Piece of membrane C. diphtheriae isolate				PCR result? Positive Negative Unknown Not done						

(please turn over)

	Treated with Antibiotics	s? Yes No	Unknown								
ANTIBIOTICS	As an Outpatient? If yes, Date Initiated:		Number of Days of Therapy:	Antibiotic Therapy i Hospital? Yes	n If yes, Dat	e Initiated:	Name of Antibiotic:	Number of Days of Therapy:			
AN	Were Antibiotics given in the 24 Hours before Culture? ☐ Yes ☐ No ☐ Unknown										
NFO	To access Diphtheria A completed and returned	access Diphtheria Antitoxin, Special Access Program Form A* must be mpleted and returned to Saskatchewan Ministry of Health. Amount of DAT admini									
XINI	Date Requested:						units				
ANTITOXIN INFO	Date Administered:										
	Country of Residence Canada Other	anada or \square Unknown									
	History of International Travel?	Country(s) Visited:			Dates						
	(2 weeks Prior to Onset)										
E)	Yes No Unknown										
EXPOSURE	History of	Province(s) Visited:			Dates						
EXPO	Interprovincial Travel?				To:		From:				
	(2 weeks Prior to Onset)										
	☐ Yes ☐ No ☐ Unknown				To:		From: _				
	Known Exposure to Dip Carrier? Yes No Unknown	ohtheria Case or	rnational Travelers?	Travelers? Known Exposure to Immigrants? Yes No Unknown							
& REPORTING	Has this Suspected Case Yes No Unknown	e been reported to the	Saskatchewan N	Ministry of	Health? If Yes, D	ate Reporte	d:				
	Person Informed:		Pho	one:		Fax:					
ATION	Reporting Physician:			Pho	one:		Fax:				
CONFIRMATION	Final Diagnosis	Final Diag	nosis Confirmed?	Case Status or Class onfirmed obable ot a case	sification:						
*htt	p://www.hc-sc.gc.ca	/dhp-mps/acces/dr	rugs-drogues/	/index-en	g.php						
Sign	nature:		Tit	tle:			Date:				

Diphtheria Attachment – Diphtheria Contact Investigation Worksheet

Reviewed: October, 2010 Section: 2-30 Page 1 of 2

Please see the following pages for the Diphtheria Contact Investigation Worksheet.



Diphtheria Contact Investigation Worksheet

*Close Contact = household members; friends; relatives and caretakers who regularly visit the home; kissing and/or sexual contacts; those who share the same room at school or work; health-care staff exposed to oropharyngeal secretions of the infected person (staff who have taken appropriate isolation precautions need not be considered contacts).

Close contacts that develop signs/ symptoms should be followed as a case - refer to Diphtheria Case Investigation Worksheet.

	CON	NTACT	INFO	RMATIC	ΟN								
Name	Age Relation to case												
Contact Phone #													
Contact I none ii	Activ	e Surve	illance	for S/S	Da	ıy 1	Day 2	Day 3	3 1	Day 4	Day 5	Day 6	Day 7
					Da	ıy ı	Day 2	Day .	,	Day 4	Day 3	Day 0	Day 1
	Indicate Yes					C	D	-14	D:	4	N4:	D-46	C14
Vaccinated? \square Yes \square No \square Unknown If vaccinated # of doses: $\square \le 3$ \square Unknown	Culture taken	Yes	No	Unknov	vn	Cu	Culture Results		Positive		Negative	Date of Culture	
Time since last dose: $\square < 10 \text{ yrs } \square > 10 \text{ yrs}$	Nasopharyngeal Oropharyngeal												
	1												
Antibiotic Prophylaxis: Yes No	Medication:					D.	1-4 4						
Name			A	ge		Re	elation to c	ase					
Contact Phone #													
	Activ	ve Surv	eillanc	e for S/S	Da	y 1	Day 2	Day	3	Day 4	Day 5	Day 6	Day 7
	Indicate Yes	or No i	f S/S i	s present									
Vaccinated? Yes No Unknown	Culture taken	Yes	No	Unknov	vn	Cu	lture Resu	ılts	Posi	tive	Negative	Date of	Culture
If vaccinated # of doses: $\square \le 3$ \square Unknown	Nasopharyngeal												
Time since last dose: $\square < 10 \text{ yrs } \square > 10 \text{ yrs}$	Oropharyngeal												
Antibiotic Prophylaxis: Yes No	Medication:	,		•						•			
Name			A	ge		Re	elation to c	ease					
Contact Phone #													
Contact I none #	Activ	Survo	illance	for S/S	Da	y 1	Day 2	Day 3	3 1	Day 4	Day 5	Day 6	Day 7
					Da	ıy ı	Day 2	Day .	,	Day 4	Day 3	Day 0	Day 1
	Indicate Yes		1			C 1		14	n :		NY 4°	D (C	C II
Vaccinated? \square Yes \square No \square Unknown If vaccinated # of doses: $\square \le 3$ \square Unknown	Culture taken	Yes	No	Unknov	vn	Cui	ture Resu	its	Positive Ne		Negative	Date of Culture	
Time since last dose: $\square < 10 \text{ yrs } \square > 10 \text{ yrs}$	Nasopharyngeal												
	Oropharyngeal												
Antibiotic Prophylaxis: Yes No	Medication:												
Name			A	ge		Re	elation to c	case					
Contact Phone #													
	Activ	ve Surv	eillanc	e for S/S	Da	y 1	Day 2	Day	3	Day 4	Day 5	Day 6	Day 7
	Indicate Yes	or No i	f S/S is	s present									
Vaccinated? Yes No Unknown	Culture taken	Yes	No	Unknov	vn	Cu	lture Resu	ılts	Posi	tive	Negative	Date of	Culture
If vaccinated # of doses: $\square \le 3 \square$ Unknown	Nasopharyngeal												
Time since last dose: $\square < 10 \text{ yrs } \square > 10 \text{ yrs}$	Oropharyngeal												
Antibiotic Prophylaxis: Yes No	Medication:												
Name			A	ge		Re	elation to c	ease					
Contact Phone #													
	Active	e Surve	illance	for S/S	Da	ıy 1	Day 2	Day 3	3]	Day 4	Day 5	Day 6	Day 7
	Indicate Yes	or No i	f S/S i	s present					1				
Vaccinated? ☐ Yes ☐ No ☐ Unknown	Culture taken	Yes	No	Unknov	vn	Cul	ture Resu	lts	Posi	tive	Negative	Date of	Culture
If vaccinated $\#$ of doses: $\square \le 3 \square$ Unknown	Nasopharyngeal												
Time since last dose: $\square < 10 \text{ yrs } \square > 10 \text{ yrs}$	Oropharyngeal												
Antibiotic Prophylaxis: Yes No	Medication:		•	•				1				•	

Diphtheria Attachment – Diphtheria Template Letter to Parents

Reviewed: October, 2010	Section: 2-30 Page 1 of 1
Date	
Dear Parent,	
There has been a case of diphtheria diagnosed in the daycare/school attends. Diphtheria is a rare disease which may cause fever, sore to white discharge over the back of the throat. An information sheet included with this letter.	hroat, and a yellow-
Public health will be reviewing immunization records for all the climmunizations to any child who requires further immunization.	nildren and providing
All children who have been in contact with diphtheria should have nasal swab collected and then should be started on preventive med family doctor to have swabs taken and antibiotics started.	
If the lab tests indicate that your child is infected with diphtheria y providing advice about further treatment and testing.	our physician will be
If you have any questions or concerns contact the local Public Heaphysician, or the HealthLine at 1-877-800-0002.	lth office, your family
Sincerely,	
Medical Health Officer	
Phone:	



Notification Timeline:

From Lab/Practitioner to Public Health: Immediate.
From Public Health to Ministry of Health: Immediate.
Public Health Follow-up Timeline: Immediate.

Public Health Purpose for Notification of invasive Group A Streptococcal (iGAS) Disease (adapted from Health Protection Surveillance Center

• To measure the burden of iGAS, identify populations at increased risk and provide a basis for epidemiologic studies;

- To ensure early detection of clusters/outbreaks of iGAS so effective control measures can be implemented;
- To prevent mortality and serious morbidity from iGAS through contact tracing and initiation of chemoprophylaxis;
- To monitor trends in iGAS;
- To monitor the effectiveness of prevention and control measures;
- To inform health care planning; to support ongoing research into sources, transmission, risk factors, pathogenesis and control of iGAS; and
- To inform the public and medical community about iGAS.

Surveillance Case Definition¹ (Public Health Agency of Canada, 2008)

Laboratory confirmation of infection with or without clinical evidence of invasive disease:* isolation of group A streptococcus (Streptococcus pyogenes) from a normally sterile site (blood, cerebrospinal fluid (CSF), pleural fluid, pericardial fluid, peritoneal fluid, deep tissue specimen taken during surgery [e.g., muscle collected during debridement for necrotizing fasciitis], bone or joint fluid excluding the middle ear and superficial wound aspirates [e.g., skin and soft tissue abscesses^]).

¹ Surveillance case definitions ensure uniform reporting to allow comparability of surveillance data. The definition is not intended to be used for clinical or laboratory diagnosis or management of cases.



Probable case	Clinical evidence of invasive disease* in the absence of another identified
	etiology and with non-confirmatory laboratory evidence of infection:
	 isolation of group A streptococcus from a non-sterile site
	OR
	positive group A streptococcus antigen detection.

- *Clinical evidence of invasive disease may be manifested as one or more of several conditions.

 These include:
 - a) Streptococcal Toxic Shock Syndrome (STSS), which is characterized by hypotension (systolic blood pressure ≤ 90 mmHg in adults or < 5th percentile for age for children) and at least two of the following signs:
 - i. Renal impairment (creatinine level $\geq 177 \mu mol/L$ for adults).
 - ii. Coagulopathy (platelet count ≤ 100,000/mm3 or disseminated intravascular coagulation).
 - iii. Liver function abnormality (SGOT [AST], SGPT [ALT], or total bilirubin $\geq 2x$ upper limit of normal).
 - iv. Adult respiratory distress syndrome (ARDS).
 - v. Generalized erythematous macular rash that may desquamate.
 - b) Soft-tissue necrosis, including necrotizing fasciitis, myositis or gangrene.
 - c) Meningitis.

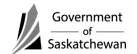
^ Wounds are not considered a sterile site with the exception of isolation of group A streptococcus (GAS) and the presence of necrotizing fasciitis and/or STSS.

Pneumonia with isolation of GAS from a sterile site, or from a bronchoalveolar lavage (BAL) when no other cause has been identified, should be regarded as a form of invasive disease for the purposes of public health management; however, as BAL does not provide a sterile site specimen, the latter would not meet the national case definition and would not be nationally notifiable.

Epidemiology and Occurrence

iGAS in Canada²

Figure 1 shows the number of cases and incidence rates of iGAS reported to the Canadian Notifiable Disease Surveillance System from 2000 to 2016. The graph shows the steady increase in the incidence rate of iGAS; doubling between 2004 (2.7/100 000) and 2016 (6.0/100,000).



² National Epidemiologic Summary as of February 28, 2018

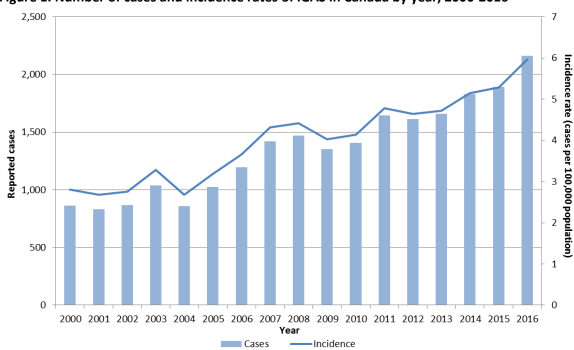


Figure 1. Number of cases and incidence rates of iGAS in Canada by year, 2000-2016

iGAS in Saskatchewan³

Figure 2 shows the number of cases and incidence rates of iGAS in Saskatchewan between 2004-2017. The upsurge in 2008, which was seen across the most westerly provinces and was related to Indigenous people, was not sustained over the following years.

Starting in 2013, a gradual increase in the iGAS trend was noted in Saskatchewan with a doubling of cases by 2017. No definitive reason has been established for this upward trend other than it reflects an upward trend reported in other Canadian provinces.



³ Saskatchewan Ministry of Health (2018)

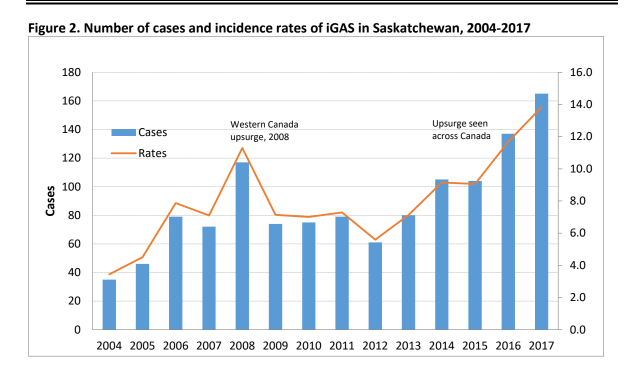
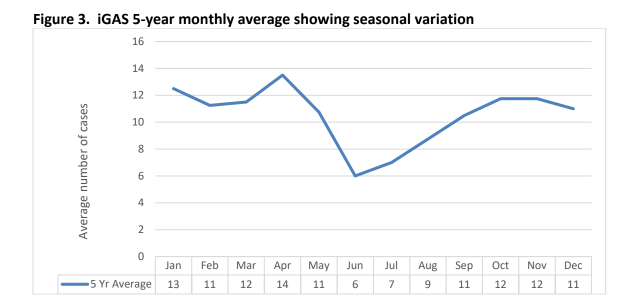


Figure 3 demonstrates the seasonal impacts and a lower incidence in the summer months when there is less crowding of individuals.



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Additional Background Information

Causative Agent

Group A streptococcus – Streptococcus pyogenes, a gram-positive coccus.

Symptoms

- Early signs and symptoms of necrotizing fasciitis include: fever, severe pain, redness and swelling at the site of wound.
- Symptoms of STSS may include pain (abrupt in onset and severe), pneumonia, acute myocardial infarction, or pericarditis, fever, chills, myalgia, nausea, vomiting, diarrhea, confusion, clinical signs of soft tissue infection (localized swelling and erythema).
- Clinical evidence for STSS is outlined in the case definition, above.
- Refer to clinical textbooks for symptoms of other clinical presentations related to GAS (meningitis, etc.).

Reservoir/Source

Humans.

Incubation Period

The incubation period of iGAS infection has not been determined (Public Health Agency of Canada, 2006).

Period of Communicability

The specified period of infectivity of the index case is:

• 7 days prior to the onset of illness, until 24 hours after the start of treatment.

Mode of Transmission

- Large respiratory droplets.
- Direct person to person contact with patient and or carrier.



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Risk Groups/Risk Factors

GAS infection can occur in anyone but risk of iGAS is significantly associated with the following:

- chronic conditions (HIV infection, cancer, heart disease, diabetes, lung disease);
- alcohol abuse;
- injection drug use;
- varicella;
- crowded living conditions;
- suboptimal hygiene practices;
- immunosuppressive therapy;
- elderly (65 years and older);
- systemic steroid use;
- Aboriginal persons.

Specimen Collection and Transport

To confirm the diagnosis of GAS, specimens should be cultured from:

- a sterile site (e.g., blood, CSF, joint fluid) or;
- an aspirate from a non-sterile site, in individuals with clinical signs of hypotension and/or invasive disease such as necrotizing fasciitis.

All GAS isolates from iGAS disease are to be sent to the Roy Romanow Provincial Laboratory (RRPL) for typing and screening for toxin genes. Characterization of the organism (emm type, whole genome sequencing) becomes important for monitoring virulence or identifying transmission patterns.

Public Health Investigation

I. Case

Refer to <u>Attachment – Invasive Group A Streptococcal Disease Data Collection</u> Worksheet to assist.

<u>History</u>

- Presentation of illness and for severity of disease.
- Health conditions that may render the individual more susceptible to invasive disease (see Risk Factors).
- Contact details refer to Attachment Contact Follow-up Form in the <u>Respiratory</u> and Direct Contact Introduction and General Considerations section.



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Public Health Interventions

Assessment

Assess for contacts as per Table 1.

Communication

• When clients are hospitalized, communication with hospital staff and or infection control staff is important.

Education

 All cases should be provided disease information as well as information on prevention and control measures including period of communicability.

Exclusion and Isolation

- Individuals are communicable until at least 24 hours after antibiotics are started.
- Strict enforcement of standard infection control practices including contact and droplet precautions. Refer to local Infection Control Manuals.

Immunization

- There is no immunization for GAS.
- If the case has any risk factors, they may be eligible for other immunizations. If not up-to-date, offer vaccines as appropriate.

Referrals

- Inform clients that supportive services (physiotherapy, occupational therapy, Home Care) are available if necessary. Refer client to primary caregiver for referrals.
- Consultation with the Medical Health Officer (MHO) may be required to determine if chemoprophylaxis is to be offered to contacts.

Treatment/Supportive Therapy

- For patient management, the client's physician is to consult an infectious disease specialist.
- Antibiotic treatment is required.
- Client may need to be hospitalized.
- In the case of necrotizing fasciitis, surgical intervention may be required.



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II. Contacts/Contact Investigation Contact Definition/Categorization

Table 1. Definition of Close Contacts

- Household contacts of a case who has spent at least 4 hours/day on average in the previous 7 days or 20 hours/week with the case.
- Non-household persons who share the same bed with the case or had sexual relations with the case.
- Persons who have had direct mucous membrane contact with the oral or nasal secretions of a case (e.g., mouth-to-mouth resuscitation, open mouth kissing) or direct contact with an open skin lesion of the case.
- Injection drug users who have shared needles with the case.

Source: Guidelines for the Prevention and Control of Invasive Group A Streptococcal Disease, 2006.

In order to be considered a close contact, there must have been exposure to the case during the period of communicability (see above). School classmates (kindergarten and older), work colleagues, as well as social or sports contacts of a case are not usually considered close contacts, unless they fit into one of the above categories.

Public Health Interventions

Assessment

- Assess for symptoms.
- Assess for risk factors.

Education

All close contacts (irrespective of whether prophylaxis is given of confirmed cases of severe disease should be alerted to signs and symptoms of iGAS disease, and be advised to seek medical attention immediately should they develop febrile illness or any other clinical manifestations of GAS infection within 30 days of diagnosis in the index case.

Chemoprophylaxis

- Chemoprophylaxis is used to prevent disease in colonized individuals and in those who
 have recently been exposed, thereby decreasing transmission of a strain known to
 cause severe infection.
- **NOTE:** Chemoprophylaxis should only be offered to close contacts of a confirmed severe case (cases of STSS, soft-tissue necrosis including necrotizing fasciitis, myositis, or gangrene, meningitis, GAS pneumonia or other life-threatening conditions) or a confirmed case resulting in death (Public Health Agency of Canada, 2006).
- Chemoprophylaxis is not routinely recommended for contacts of cases that are not severe (i.e., bacteremia or septic arthritis). These cases often have milder disease



than those with invasive disease. Their contacts are also likely to have milder disease as well since there is consistency in type and severity of disease with particular strains of GAS.

- Refer to contact definition for listing of those who require prophylaxis. A close contact will be given prophylaxis if they were in contact with the case during the period of communicability (noted above).
- Even though the incubation period is not known, most subsequent cases occur within 7 days after last contact with an infectious case (Public Health Agency of Canada, 2006). Close contacts should ideally be given antibiotics within 24 hours of case identification; however it is still advisable for up to 7 days. The benefits of starting prophylaxis should be discussed with the MHO if it is beyond 7 days of last contact with the infectious case.
- Refer to <u>Attachment Recommended Chemoprophylaxis Regimens for Close Contacts.</u> **Testing**
- Not routinely done Refer to <u>Attachment Investigation and Control Approaches for Long Term Care Facilities</u> for the screening procedures for instances in long term care⁴ (LTC) facilities.

Exclusion

No need to exclude contacts from day care, school or work.

III. Environment

Table 2. Impetus f	Table 2. Impetus for Action for Organization-based Outbreaks or Clusters						
Long-term care facility	 An incidence rate of culture-confirmed GAS infections > 1 per 100 residents per month, OR At least 2 cases of culture-confirmed infection in one month in facilities with less than 200 residents, OR An incidence rate of suspected GAS infections of > 4 per 100 residents per month. 						
Child care centre	 One severe case of iGAS disease in a child attending a child care centre. 						
Hospital	 One or more linked invasive or non-invasive GAS cases in either patients or staff occurring within one month of an iGAS case (see Annex 3 – National Guidelines, Oct 2006). 						

Source: Adapted from Guidelines for the Prevention and Control of Invasive Group A Streptococcal Disease, 2006.

⁴ Adapted from Guidelines for the Prevention and Control of Invasive Group A Streptococcal Disease, October 2006.



Section 2 - 40 – Group A Streptococcal Disease - invasive (iGAS)
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Child Care Centre Control Measures

- Although outbreaks of iGAS disease occurring among children attending a child care centre are rare, when a case occurs the following needs to occur:
 - strict enforcement of standard infection control practices refer to the Saskatchewan Ministry of Health Infection Control Manual for Child Care Facilities;⁵
 - details of the Child Care Centre (size, attendees, etc.) should be collected.

Institutional Control Measures

- Residents of LTC facilities are at increased risk of morbidity and mortality due to iGAS disease because of their older age and higher prevalence of underlying conditions.
- Strict enforcement of standard infection control practices including contact and
 droplet precautions are required. Refer to Local Infection Control Manual. In LTC
 facility outbreaks, the implicated strain is usually widespread within the facility and
 limited provision of chemoprophylaxis to close contacts is not the optimal approach.
 Refer to Attachment Investigation and Control Approaches for Long Term Care Facilities for detailed information regarding investigation and control approaches that may be useful.

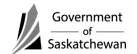
IV. Epidemic Measures

- Determine source and manner of spread.
- Investigate promptly the extent of the exposure.
- If there is exposure of groups like schools, LTCs, daycare centres, it may be necessary to administer preventative antimicrobial therapy to the whole group.
- Consider extensive consultation with various specialties including: infectious disease, laboratory medicine, Saskatchewan Ministry of Health, others as appropriate.

Prevention Measures

Refer to the <u>Respiratory and Direct Contact Introduction and General Considerations</u> section of the manual that highlights topics for client education that should be considered as well as provides information on high-risk individuals and environments.

⁵ http://www.saskatchewan.ca/live/births-deaths-marriages-and-divorces/starting-a-family/early-learning-and-child-care/child-care.



Section 2 - 40 – Group A Streptococcal Disease - invasive (iGAS)
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Education

- Good hygiene, especially hand washing is important to prevent the spread of bacteria.
- Provide information sheet, Attachment Invasive Group A Streptococcal Disease.
- Non-severe cases will be dealt with on a case-by-case basis in consultation with the MHO.



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Revisions

Date	Change
September 2018	Updated to align with Panorama configuration
	• Incorporated the purpose for notification of cases to public health
	Provided clarification in the case definition on the limited
	applicability of specimens from wounds.
	Incorporated an Epidemiology and Occurrence section to the
	chapter.
	Rearranged and updated the style into the new format of the
	Manual.
	References reaffirmed or updated as necessary.



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References

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Streptococcal Invasive Disease (group A) Data Collection Worksheet



Please complete all sections.

Panorama QA complete:	□No	Panorama Client ID: Panorama Investigation ID:				
Initials: A) CLIENT INFORMATION			I HN -> SUBIF		J	RSONAL INFORMATION
Last Name:		First Name: and Middle Name:	LINE - JODGE		Name (Goes by	
			1		,	,.
DOB: YYYY / MM / DD	Age:	Health Card Province:			Communication phone, text):	n Method: (specify -
Phone #: Primary Home:		Health Card Number (PHN):			dress: \square Work	□ Personal
☐ Mobile contact: ☐ Workplace:				Lillan Auc	11C33. — VVOIN	- Felsonai
Place of Employment/School:		Gender: □ Male	□ Female		Other	□ Unknown
Alternate Contact:		Address Type: ☐ No fixed ☐ Postal Address Mailing (Postal address):	☐ Primary Hor	ne □Temp	oorary □Legal	Land Description
Alt. Contact phone:		Street Address or FN Communit	ty (Primary Hon	ne):		
		Address at time of infection (if r	not the same):			
B) INVESTIGATION INFORMATION	SUBJI	ECT SUMMARY->RESPIRATORY &	DIRECT CONTA	ACT ENCOUP	NTER GROUP->(CREATE INVESTIGATION
Disease Summary Classification: CASE	Date	Classification: CONTACT	Date	,	LAB TEST INFORMATION: Date specimen collected:	
□ Confirmed	YYYY / MM / DD	□ Contact	YYYY / MM	/ DD	YYYY / MM /	
□ Does Not Meet Case Definition	YYYY / MM / DD	□ Not a Contact	YYYY / MM	/ DD	Specimen type	2:
☐ Person Under Investigation	YYYY / MM / DD	☐ Person Under Investigation	YYYY / MM	/ DD	□ Blood □ CSF	
□ Probable	YYYY / MM / DD				□ Other	
Disposition:		<u>1</u>				
FOLLOW UP: ☐ In progress	YYYY / MM / DD	□ Complete		YYYY / N	MM / DD	
☐ Incomplete - Declined	YYYY / MM / DD	□ Not required		YYYY / N		
☐ Incomplete – Lost contact	YYYY / MM / DD	☐ Referred – Ou	at of province	YYYY / N	/IM / DD	
☐ Incomplete – Unable to locate	YYYY / MM / DD	(specify where)				
REPORTING NOTIFICATION		Location:				
Name of Attending Physician or Nur	se:					
Physician/Nurse Phone number:		Date Received	d (Public Health	ı): YYYY /	' MM / DD	
Type of Reporting Source: ☐ Heal	th Care Facility	ab Report	ioner □Phy	/sician [□ Other	

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Streptococcal Invasive Gisease (group A) Data Collection Worksheet

Please complete all sections.

 Panorama Client ID:
Panorama Investigation ID:

C١	SIGNS & SYMPTOMS	(Rold text - nart o	f case definition
·	SIGNS & STIVIP I CIVIS	IDUIU LEXL – DUI L O	ı cuse uenninili

LHN-> INVESTIGATION->SIGNS & SYMPTOMS

Description	No	Yes – Date of onset	Description	No	Yes - Date of onset
Acute respiratory distress syndrome (ARDS) - CXR/CT*		YYYY / MM / DD	Muscle inflammation (myositis)		YYYY / MM / DD
Arthritis - septic		YYYY / MM / DD	Necrosis - skin and tissue		YYYY / MM / DD
Cardiac - myocardial infarction		YYYY / MM / DD	Necrotizing fasciitis		YYYY / MM / DD
Cellulitis		YYYY / MM / DD	Confusion		YYYY / MM / DD
Chills		YYYY / MM / DD	Pain - severe		YYYY / MM / DD
Fever		YYYY / MM / DD	Cardiac - pericarditis		YYYY / MM / DD
Gangrene		YYYY / MM / DD	Pharyngitis (sore throat)		YYYY / MM / DD
Hypotension*		YYYY / MM / DD	Pneumonia		YYYY / MM / DD
Infection - soft tissue		YYYY / MM / DD	Rash - erythematous macular *		YYYY / MM / DD
Infection - wound		YYYY / MM / DD	Renal impairment * (refer to CDC Manual for parameters)		YYYY / MM / DD
Lab - liver function abnormality* (refer to CDC Manual for parameters)		YYYY / MM / DD	Sepsis (e.g. bacteremia, septicemia, etc.)		YYYY / MM / DD
Lab - platelet count low* (refer to CDC Manual for parameters)		YYYY / MM / DD	Skin - pain and swelling		YYYY / MM / DD
Meningitis		YYYY / MM / DD	Streptococcal toxic shock syndrome (STSS) Includes 2 or more of the S/S with an *		YYYY / MM / DD

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_		пν	L	JD	ΑI	IUI	ıA	Nυ	LU	IVI	IVΙL	JIVI	LA	DIL	_I I Y

LHN-> INVESTIGATION->INCUBATION & COMMUNICABILITY

Communicability for Case (period for transmission):

Earliest Possible Communicability Date: YYYY / MM / DD

Latest Possible Communicability Date: YYYY / MM / DD

Communicability Calculation Details:

E) RISK FACTORS (RF followed by + impact the Immunization Forecaster)

LHN-> SUBJECT->RISK FACTORS

DESCRIPTION	YES	N – No NA – not asked U - Unknown	DESCRIPTION	YES	N – No NA – not asked U - Unknown
Chronic Medical Condition - Cardiac Disease +			Medical Risk Factor - Varicella	YYYY / MM / DD	
Chronic Medical Condition - Diabetes Mellitus +			Medical Treatment - Surgery/surgical wound	YYYY / MM / DD	
Chronic Medical Condition - Liver disease +			Setting - Crowded living conditions (>1 person per room excluding bathrooms)		
Chronic Medical Condition - Lung disease +			Special Population – Homeless +		
Chronic Medical Condition - Renal disease +			Special Population - Lives in a communal setting		
Contact to a known case (Add'l Info)	YYYY / MM / DD		Special Population - LTC Facility +		

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Streptococcal Invasive Gisease (group A) Data Collection Worksheet

Please complete all sections.

 Panorama Client ID:	
Panorama Investigation ID:	

YYYY / MM / DD

DESCRIPTION		YES	N – No NA – not asked U - Unknown	DESCRIPTION	YES	N – No NA – not asked U - Unknown		
Immunocompromise	d - HIV +			Special Population - Self-reported Indigenous identity				
Immunocompromise Related to underlying treatment				Substance Use - Alcohol				
Medical Risk Factor -	Postpartum			Substance Use - Injection drug use (including steroids) +				
Medical Risk Factor History of injury (Add	'l Info)	YYYY / MM / DD		Travel - Outside of Canada (Add'l Info)	YYYY / MM / DD			
Medical Risk Factor - or dermatological con		YYYY / MM / DD		Travel -Outside of Saskatchewan, but within Canada (Add'l Info)	YYYY / MM / DD			
TREATMENT					/IEDICATIONS->MEDI	CATIONS SUMMA		
Medication (Panoran Prescribed by:				Started on: YYYY / MM / DD				
i) INTERVENTIONS				INVESTIGATION->TREATMENT & INTE	ERVENTIONS->INTER	/ENTION SLIMM/		
Intervention Type an	d Sub Type:			INVESTIGATION-ZIREATIVIENT & INTE	ERVENTIONS-ZINTERV	VENTION SOIVINA		
Assessment: ☐ Assessed for conta Investigator name	cts	Y	YYY / MM / DD	Education/counselling: Inves ☐ Prevention/Control measures ☐ Disease information provided		/ / MM / DD / / MM / DD		
Communication: Phone call attempt Phone call attempt	ted (evening)	Y	YYY / MM / DD YYY / MM / DD	Immunization: ☐ Eligible Immunization(s) recommended YYYY / MM / DD Investigator name				
☐ Home visit attemp ☐ Letter sent ☐ Text message sent		Y	YYY / MM / DD YYY / MM / DD YYY / MM / DD	Isolation: ☐ Facility isolation YYYY / MM / DD Investigator name	☐ Home isolation	on YYYY/MM/ DD		
☐ Other communicat☐ Letter (See Docum Investigator name			YYY / MM / DD YYY / MM / DD	Referral Consult with MHO Investigator name	YYYY / MM /	DD		
General: Investigator ☐ Disease-Info/Prev-		Y	YYY/ MM / DD		/ MM / DD			
Disease-Info/Prev-C	Cont/Assess'd for C Intervention	Comments	YYY/ MM / DD	☐ Document Management	Next follow-up	Initials		
	subtype				Date YYYY / MM / D			
YYYY / MM / DD					YYYY / MM / D			
YYYY / MM / DD					YYYY / MM / D			
YYYY / MM / DD					YYYY / MM / E			
YYYY / MM / DD					YYYY / MM / D			
YYYY / MM / DD					YYYY / MM / D	DD D		
YYYY / MM / DD					YYYY / MM / D)D		
YYYY / MM / DD					YYYY / MM / D)D		
YYYY / MM / DD					YYYY / MM / D	DD		
YYYY / MM / DD					YYYY / MM / D)D		

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YYYY / MM / DD

Streptococcal Invasive Gisease (group A) Data Collection Worksheet

Please complete all sections. Panorama Client ID: _ Panorama Investigation ID: H) OUTCOMES (optional except for severe influenza, LHN-> INVESTIGATION-> OUTCOMES

ransmission	Exposure Name	Setting type	Date/Time	# of contacts
Event ID system-generated an be documented selow		(Select the most appropriate setting for the TE; if >1 select multiple settings will be entered into Panorama)	- 11.7	
		☐ Childcare worker/attendee ☐ Health care setting ☐ Household ☐ Sexual exposure ☐ Type of community contact ☐ Congregate/communal living setting		
		☐ Childcare worker/attendee ☐ Health care setting ☐ Household ☐ Sexual exposure ☐ Type of community contact ☐ Congregate/communal living setting		
		☐ Childcare worker/attendee ☐ Health care setting ☐ Household ☐ Sexual exposure ☐ Type of community contact ☐ Congregate/communal living setting		
		☐ Childcare worker/attendee ☐ Health care setting ☐ Household ☐ Sexual exposure ☐ Type of community contact ☐ Congregate/communal living setting		
	iGAS Contacts – Inv	☐ Multiple Settings	YYYY / MM / DD	
	ID#		to YYYY / MM / DD	

September 1, 2018 Page 4 of 4 Attachment – Recommended Chemoprophylaxis Regimens for Close Contacts

Drug	Dosage	Comments
First line - First generation cephalosporins: cephalexin, cephadroxil, cephradine	Children and adults: 25 to 50 mg/kg/day, to a maximum of 1 g/day, in	Recommended drug for pregnant and lactating women.
	2 to 4 divided doses x 10 days	Should be used with caution in patients with allergy to penicillin.
		Use of cephalosporins with nephrotoxic drugs (e.g. aminoglycosides, vancomycin) may increase the risk of cephalosporin- induced nephrotoxicity.
Second line - Erythromycin	Children: 5 to 7.5 mg/kg every 6 hours or 10 to 15 mg/kg every 12 hours (base) x 10 days (to a maximum of the adult dose)	Erythromycin estolate is contraindicated in persons with pre-existing liver disease or dysfunction and during pregnancy.
	Adults: 500 mg every 12 hours (base) x 10 days	Sensitivity testing is recommended in areas where macrolide resistance is unknown or known to be $\geq 10\%$.
Second line - Clarithromycin	Children: 15 mg/kg/day in divided doses every 12	Contraindicated in pregnancy.
	hours (to a maximum of the adult dose) Adults: 250 mg po bid x 10 days	Sensitivity testing is recommended in areas where macrolide resistance is unknown or known to be ≥10%.
Second line - Clindamycin	Children: 8 to 16 mg/kg/day divided into 3 or 4 equal doses x 10 days (to a maximum of the adult dose) Adults: 150 mg every 6 hours x 10 days	Alternative for persons who are unable to tolerate beta-lactam antibiotics.

Source: Guidelines for the Prevention and Control of Invasive Group A Streptococcal Disease, 2006

All prophylactic regimes are administered orally and taken for 10 days.

Note: All persons who receive chemoprophylaxis should watch for signs and symptoms of invasive GAS disease, for 30 days after the diagnosis of invasive disease in the index patient.



Attachment – Investigation and Control Approach for Long Term Care (LTC) Facilities

Page 1 of 2

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Background

Residents of LTC facilities are at increased risk of morbidity and mortality due to iGAS disease because of their older age and higher prevalence of underlying conditions. When a culture-confirmed case of iGAS disease occurs in a LTC facility, there is a 38% likelihood that a second positive blood culture-confirmed case of the same strain will be detected in the facility within six weeks. A number of outbreaks of iGAS infections have been documented in LTC facilities. Infection is often spread through person-to-person contact, with clustering of cases by room or care unit in some instances. Staff may be a source of or conduit of infection either through poor infection control practices or asymptomatic carriage. However, hospital staff who are carriers are more likely to be the source of infection in outbreaks in acute care facilities, whereas outbreaks in LTC facilities are more often patient-propagated. In LTC facility outbreaks, the implicated strain is usually widespread within the facility and limited provision of chemoprophylaxis to close contacts is not the optimal approach.

Procedure

In addition to strict enforcement of standard infection control practices, the following approach may be useful in the investigation and control of iGAS disease in LTC facilities:

- 1. When a confirmed case of iGAS disease occurs in a LTC facility such as a nursing home, the facility should:
 - a. Report the case to the local Medical Health Officer (MHO) or designate.
 - b. Review the facility's nosocomial infection reports, for the previous 4 to 6 weeks, for culture-confirmed cases of GAS disease and cases of skin and soft tissue infections (e.g., pharyngitis and cellulitis). An excess of GAS infection and clinically compatible illness, or LTC facility outbreak, is defined in Table 2 Impetus for Action for Organization-based Outbreaks or Clusters.
 - c. Assess the potential for a source of infection from outside the facility (e.g., regular visits from children who have recently been ill).



- 2. If an excess of GAS infection is identified, the following actions should be considered:
 - a. Consult the local Infection Control Practitioner/MHO/CD Epidemiologist team, as to the most practical approach. This could comprise the concentric-circles approach, (i.e., begin screening the closest contacts and extending the investigation from there).
 - b. Anyone colonized with GAS should receive chemoprophylaxis.
 - c. Non-patient care staff¹ should be asked about possible recent GAS infections. Those with a positive history should be screened for GAS and those persons positive should be treated with antibiotics.
 - Treatment choices are governed by the most recent guidelines.
 The public health practitioner should direct any questions regarding the current treatment protocols to the physician or MHO. Refer to Heymann and other texts and clinical treatment guidelines for specific treatment details. See Appendix H Sources for Clinical Treatment Guidelines.
 - d. All GAS isolates should have further typing. This should be coordinated through the MHO and the Saskatchewan Disease Control Lab (SDCL). Culturing for a test of cure is recommended for individuals found to have the outbreak-related strain. Culturing for a test of cure is not necessary for individuals infected with a non-outbreak-related strain of GAS.
 - e. Re-screen all GAS positive residents and staff including their throat and skin lesion(s) 14 days after the treatment has been started. If this screen is positive, the individual should be retreated with antibiotics and rescreened in 14 days. If still colonized, discontinue treatment unless the facility has an ongoing problem with GAS infection.
 - f. Active surveillance for GAS infection should be initiated and continued for 1 to 2 months as determined by the local outbreak team.
 - g. Appropriate specimens should be taken for culture to rule out GAS when suspected infections are detected by active surveillance.
- 3. If no excess is identified, especially if there is evidence of an outside source of infection for the index case, then active surveillance alone for 2 to 4 weeks to ensure the absence of additional cases is warranted.



¹ This includes maintenance and housekeeping staff for example.

Notification Timeline:

From Lab/Practitioner to Public Health: Immediate.

From Public Health to Ministry of Health: Within 72 hours. Public Health Follow-up Timeline: Initiate within 48 hrs.

Public Health Purpose for Notification of *Haemophilus Influenzae* **Disease** (adapted from British Columbia Center for Disease Control [2017])

- To minimize mortality and serious morbidity from *Haemophilus Influenzae* B;
- To rapidly identify close contacts of the case and to provide recommendations for appropriate preventive measures for close contacts so as to prevent further spread of infection and disease;
- To provide information about the disease, its transmission, and methods of prevention;
- To identify clusters or outbreaks of infection and to initiate appropriate prevention and control measures;
- To track epidemiology trends of meningococcal disease in Saskatchewan including risk factors and distribution;
- To monitor the effectiveness of prevention and control measures;
- To plan expansion or introduction of future immunization programs;
- To provide timely clinical care including diagnosis and treatment using current, evidence-based guidelines;
- To make timely and evidence informed actions on outbreaks; and
- To inform the public and medical community about *Haemophilus Influenza* disease.

Information

Table 1. Differences between *Haemophilus Influenzae* Invasive B (Hib) and Non-Hib Typeable Strains

	Hib	Non-Hib Typeable Strains a,c,d,e,f
Reportable	Yes	Yes
Public Health Follow- Up	Yes	No
Invasive Disease	More common	Less common



Communicability	Not considered communicable after 24-48 hours of effective antimicrobial therapy	Unknown
Hospitalized Patients	Routine and droplet precautions until 24 hours after initiation of antimicrobial therapy	Not defined
Treatment	Third generation cephalosporin or chloramphenicol in combination with ampicillin	No defined regimen. Ceftriaxone and cefotaxime have been used successfully
Management of Contacts	Recommended	Not recommended
Prevention	Vaccine	No vaccine

Source: Manitoba Health Communicable Disease Management Protocol, 2007.

Surveillance Case Definitions¹ (Public Health Agency of Canada, May 2008)

Table 2. Haemophilus Influenzae B Invasive Disease

Confirmed Case	Clinical evidence ¹ of invasive disease with laboratory confirmation of
	infection:
	• isolation of <i>H. influenzae</i> (serotype b) (Hib) from a normally sterile
	site^
	OR
	• isolation of <i>H. influenzae</i> (serotype b) from the epiglottis in a
	person with epiglottitis.
Probable Case	Clinical evidence of invasive disease with laboratory evidence of
	infection:
	• demonstration of <i>H. influenzae</i> type b antigen in cerebrospinal
	fluid
	OR
	• demonstration of <i>H. influenzae</i> DNA in a normally sterile site
	OR
	 buccal cellulitis or epiglottitis in a child < 5 years of age with no
	other causative organisms isolated.
¹Clinical illness assoc	ciated with invasive disease due to <i>H. influenzae</i> includes meningitis,

bacteraemia, epiglottitis, pneumonia, pericarditis, septic arthritis and empyema.

¹ Surveillance case definitions ensure uniform reporting to allow comparability of surveillance data. The definition is not intended to be used for clinical or laboratory diagnosis or management of cases.



[^]Includes: blood, cerebrospinal, joint, pleural, pericardial, or peritoneal fluid.

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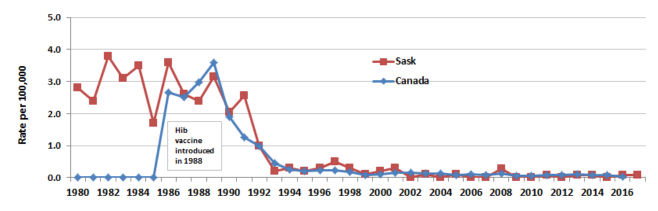
Confirmed Case	Clinical evidence ¹ of invasive disease with laboratory confirmation of infection:
	 isolation of <i>H. influenzae</i> (serotype a,c,d,e,f, undifferentiated and non-typeable isolates) from a normally sterile site OR
	• isolation of <i>H. influenzae</i> (serotype a,c,d,e,f, undifferentiated and non-typeable isolates) from the epiglottis in a person with epiglottitis.

¹Clinical illness associated with invasive disease due to *H. influenzae* includes meningitis, bacteraemia, epiglottitis, pneumonia, pericarditis, septic arthritis and empyema.

Epidemiology and Occurrence

Saskatchewan introduced Hib vaccine in 1988 resulting in a dramatic decline in reported cases. The highest number of cases were reported in 1984 and 1986 (36 cases), mainly infants. The average number of reported cases in the pre-vaccine era was 21 cases per year; the yearly average over the decades 1998 - 2017 was about one case per year (19 cases). These are largely unimmunized children. Between 1979 and 1993 there were 96 deaths and there have been no reported deaths caused by invasive Hib since then.

Figure 1. H.Influenza Rates Saskatchewan and Canada before and after Immunization Program



Additional Background Information

Causative Agent

Haemophilus influenzae, a gram-negative coccobacilli. Encapsulated strains are classified into sub-types. Serotype b (Hib) is the most pathogenic.

Symptoms

Illnesses often caused by *H. influenzae* type b include meningitis, epiglottitis, pneumonia, and bacteremia.

Symptoms include fever, lethargy, drowsiness, rapid or difficult breathing, sore throat, stiff neck and bulging fontanelles in infants. Most cases are in children 2 months to 4 years of age (Heymann, 2015; American Academy of Pediatrics, 2015).

Reservoir/Source

Upper respiratory tract of humans.

Incubation Period

Unknown, probably variable, and possibly as short as 2-4 days.²

Period of Communicability

As long as organisms are present, asymptomatic carriage may occur indefinitely in up to 2-5% of children. Communicability ends within 24-48 hours after the beginning of antibiotic therapy.

Mode of Transmission

Person-to-person from direct contact or droplet contact of oral or nasal secretions, e.g., saliva, nasal mucus, or respiratory secretions.

Specimen Collection and Transport

If invasive disease, blood cultures and CSF specimens should be submitted as per local lab specimen collection and transport guidelines.

² Most "secondary" cases in families usually occur within 2 weeks and in childcare settings within 60 days. However, this may be transmission from an asymptomatic carrier rather than the index case.



Public Health Investigation

I. Case

Refer to <u>Attachment – Haemophilus Influenzae</u> Type B (invasive) Data Collection Worksheet to assist.

History

- Onset of illness, presentation and treatment (with what and when) to determine incubation period and period of communicability which helps to identify the possible source and contacts to be followed.
- Review immunization history of the case.
- Determine case status including a review of the immunization history. Do not wait for specific typing results before initiating public health follow-up.
- Identify contacts (refer to contact definition).
- Determine if case has underlying medical conditions or falls into a risk category.

Public Health Interventions

Assessment

- Assess for contacts Aggressive contact tracing, identification, and appropriate management, is the foundation to the prevention of secondary cases. Refer to Contact Definition.
- Obtain names, addresses, and phone numbers of all possible contacts. This information may need to be obtained from someone close to the case.

Communication

 Letters can be sent to classrooms and other group settings where individual contact tracing is not required to inform them of the exposure, symptom monitoring and when to seek medical attention (see Sample letter)

Education

 All cases should be provided disease information as well as information on prevention and control measures including period of communicability and avoiding contact with vulnerable individuals.

Immunization

 Ensure the client's entire immunization status is up-to-date once they have recovered.³

³ Life-long immunity is expected following natural infection with Hib in individuals who were older than 24 months at the time of the disease. Take the opportunity to update any other immunizations that the client is eligible for.



- Grabenstein (2011) indicates that children with invasive Hib disease, when
 younger than 24 months, may not develop adequate anticapsular antibodies and
 remain at risk for a subsequent episode of the disease. After recovery from this
 illness episode, these children should be re-vaccinated with Hib vaccine
 according to age at presentation as if they have not been previously immunized.
 Children who were older than 24 months of age at time of disease do not need
 to be immunized as they should develop a protective immune response.
- Refer to Saskatchewan Immunization Manual Chapter 5⁴

Isolation

 Respiratory isolation for 24 hours following initiation of appropriate antibiotic treatment

Referrals

- When clients are hospitalized, communication with hospital staff and or infection control staff is important.
- Inform clients that supportive services (physiotherapy, occupational therapy, Home Care) are available if necessary. Refer client to primary caregiver for referrals.

Treatment/Supportive Therapy

The public health practitioner should direct any questions regarding the current treatment protocols to the physician or Medical Health Officer (MHO). The following serves as a reference for the public health investigator:

- Antibiotic treatment is require. For patient management the client's physician should consult an infectious disease specialist.
- In addition to therapeutic antibiotics, the case should receive chemoprophlaxis with rifampin before hospital discharge unless the infection was treated with an antibiotic that is effective in eliminating Hib colonization (American Academy of Pediatric, 2009 and Heymann, 2008). Refer to Attachment Rifampin Chemoprophylaxis Dosage Guide for Haemophilus influenzae Type b for information on dosing.

⁴https://www.ehealthsask.ca/services/Manuals/Documents/Ch.%205%20Immunization%20Schedules%20Aug%202018.pdf



II. Contacts/Contact Investigation

Contact Definition (American Academy of Pediatrics, 2009)

- Contacts are defined as:
 - a person residing with the case of invasive Hib disease
 - non-residents who have spent 4 or more hours per day with the index case for at least 5 of the 7 days preceding the day of hospital admission of the case.
- Complete the Attachment Contact Follow-up Form in the <u>Respiratory and</u> <u>Direct Contact Introduction and General Considerations</u> for all identified contacts.
- Consult with the MHO immediately to determine whether rifampin chemoprophylaxis and/or Hib immunization is necessary.

Public Health Interventions

Assessment

Assess for symptoms.

Chemoprophylaxis

When indicated, prophylaxis should be initiated as soon as possible given that most secondary cases in households occur during the first week after hospitalization of the index case. As some secondary cases occur later, initiation of prophylaxis seven days or more after hospitalization of the index case may still be of some benefit (American Academy of Pediatrics, 2009). Discuss with the MHO.

Recommended for:

- 1. All household contacts, regardless of age, in the following circumstances:
 - household with at least 1 contact younger than 4 years of age who is unimmunized or incompletely immunized⁵ for age;
 - household with a child younger than 12 months of age if the child has not received the primary series of three doses;
 - household with an immunocompromised child, regardless of that child's Hib immunization status (i.e., even if fully immunized).

⁵ Complete immunization is determined by the age at when they received their first dose, their current age and the number of doses received to date. Please refer to the Saskatchewan Immunization Manual for further details: http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx



- **2. Preschool/daycare contacts (including staff)**, regardless of age, when **2** or more cases of invasive Hib disease have occurred within 60 days among attendees.
 - If the index case attends preschool or day care, and the decision is to provide rifampin to all contacts, inform all parents of the situation. Together with the facility operator, plan and provide parent education about invasive Hib disease. It is especially important to discuss contraindications and side effects of rifampin.
- **3. The case**, if younger than 2 years of age or is a member of a household with a susceptible contact, and who had been treated with a regimen other than cefotaxime sodium or ceftriaxone sodium; chemoprophylaxis usually is provided just before discharge from hospital (American Academy of Pediatrics, 2009).

Chemoprophylaxis MAY be considered in the following situations at the discretion of the Medical Health Officer:

 Health care workers who have administered mouth-to-mouth resuscitation to the case (British Columbia Centre for Disease Control, 2005).

Testing

 Contacts of an index case should **not** be swabbed for culture of Hib prior to initiating rifampin chemoprophylaxis since the result has no bearing on the decision to administer rifampin.

Immunization

- Post-exposure Hib immunization is not known to decrease the risk of transmission. Rather, the situation presents an opportunity for completion of Hib immunization of contacts.
- Offer immunization to contacts less than 60 months of age who are unimmunized or not completely immunized⁶ for age and to individuals older than 5 years of age who have chronic conditions associated with increased risk of invasive Hib disease. Refer to Saskatchewan Immunization Manual⁵ – Chapter 5: Immunization Schedules and Chapter 7: Immunization of Special Populations).

Exclusion

 Any individual who is eligible to receive prophylaxis should be isolated at home until 24 hours after prophylaxis has been initiated.

⁶ Complete immunization is determined by the age at when they received their first dose, their current age and the number of doses received to date. Please refer to the Saskatchewan Immunization Manual for further details: http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspxl.



- New attendees to a daycare should not be permitted until rifampin prophylaxis has been completed.
- Persons entering a setting (new staff and attendees) where rifampin is being given should also receive it and be excluded as above.

III. Environment

Child Care Centre/Schools Control Measures

Ensure each parent receives information about Hib disease. See <u>Attachment – Sample Fact Sheet on Haemophilus Influenzae Type B Disease</u>.

Management of the centre. Three situations may occur:

- **1.** If **one case** of invasive Hib disease occurs in a centre with one child under 24 months attending:
 - Notify and educate staff and parents of contacts of the case to be alert for anyone with fever, sore throat, headache, stiff neck, drowsiness, rapid or difficult breathing, excessive irritability, or symptoms at the site of infection.
 Seek prompt evaluation by a physician for any ill child.
 - The centre director must notify public health if any additional children become ill.
 - The advisability of rifampin prophylaxis in exposed childcare groups with unimmunized or incompletely immunized children is controversial. Discuss this with the MHO.
 - Notify parents of other childcare centre attendees of the occurrence of a case of Hib disease (see the appropriate <u>Attachment Sample Letter about</u> <u>Haemophilus Influenzae</u> Type B Invasive Disease <u>Prophylaxis Recommended</u> or <u>Sample Letter about <u>Haemophilus Influenzae</u> Type B Invasive Disease <u>Prophylaxis NOT Recommended</u>).
 </u>
 - Assess immunization status of children.
 - Recommend age-appropriate Hib immunization for all incompletely immunized or unimmunized children.
- **2.** If **one case** of invasive Hib disease occurs in a centre and all children in the centre are at least 24 months of age, regardless of immunization status:
 - Educate parents and staff to be alert for anyone with fever, sore throat, headache, stiff neck, drowsiness, rapid or difficult breathing, excessive irritability, or symptoms at the site of infection. Seek prompt evaluation by a physician for any ill child.



- Notify all parents of the occurrence of a case of Hib disease (see <u>Attachment Sample Letter about Haemophilus Influenzae Type B Invasive Disease Prophylaxis NOT Recommended</u>).
- The centre director must notify public health if any additional children become ill.
- Do not recommend prophylaxis or vaccine.
- **3.** If **two or more cases** of invasive Hib disease occur within 60 days and incompletely immunized children attend the centre, carry out the same procedures as for one case but prophylaxis for all attendees and staff is recommended.
 - Notify all parents of the occurrence of a case of Hib disease (see <u>Attachment Sample Letter about Haemophilus Influenzae Type B Invasive Disease Prophylaxis Recommended</u>).
 - Absent attendees should be contacted to determine if they are contacts/cases.
 - All new attendees entering a setting where rifampin has been used within two months must be age-appropriately immunized (Government of Manitoba, 2007).
 - See <u>Chemoprophylaxis</u> section.

IV. Epidemic Measures

Not applicable

Prevention and Education

Refer to the Respiratory and Direct Contact Introduction and General Considerations section of the manual that highlights topics for client education that should be considered as well as provides information on high-risk groups and activities.

Immunization

- Immunize infants, children and adults according to the recommended schedule.
 Refer to Saskatchewan Immunization Manual.⁷
- Complete immunization is determined by the age at when they received their first dose, their current age and the number of doses received to date. Please refer to the Saskatchewan Immunization Manual² for further details.



⁷ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx .

Respiratory and Direct Contact Section 2-50 - Haemophilus Influenzae Page **11** of **14** 2018 09 01

Education

- Educate the public about the disease and the need for active immunization.
- Immunization information fact sheets can be used to guide discussion.
- Education should be provided regarding respiratory etiquette and measures to prevent transmission.



Revisions

Date	Change
September 2018	 Updated to align with Panorama configuration; Clarified the purpose for notification of cases to public health; Incorporated an Epidemiology and Occurrence section into the chapter; Incorporated Haemophilus Influenzae Infection (invasive) Data Collection Worksheet; Rearranged and updated the style into the new format of the Manual.
	 Implemented boxes to draw attention to treatment and chemoprophylaxis information.



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Respiratory and Direct Contact Section 2-50 - Haemophilus Influenzae Page **14** of **14** 2018 09 01

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Saskatchewan Haemophilus inflenzae infection (invasive) Data Collection Worksheet Please complete all sections.

Panorama QA complete: Yes Initials:	□No				Panorama Client ID: ma Investigation ID:		
A) CLIENT INFORMATION			IHN -> SUBJEC		AILS -> PERSONAL INFORMATIO		
Last Name:		First Name: and Middle Name:		Alternate Nan			
DOB: YYYY / MM / DD	Age:	Health Card Province:			nmunication Method: (specify -		
Phone #: Primary Home: Mobile contact: Workplace:	☐ Mobile contact:		Health Card Number (Phin).		i.e. home phone, text): Email Address: □Work □Personal		
Place of Employment/School:		Gender: □ Male	□ Female	□Othe	er 🗆 Unknown		
Alternate Contact:		Address Type: ☐ No fixed ☐ Postal Address Mailing (Postal address): Street Address or FN Communi			ry □ Legal Land Description		
		Address at time of investigation					
B) INVESTIGATION INFORMATION	LHN-> SUBJEC	T SUMMARY-> RESPIRATORY & D	DIRECT CONTAC	T ENCOUNTER G	GROUP-> CREATE INVESTIGATION		
Disease Summary Classification:	Date	Classification: CONTACT	Date		B TEST INFORMATION: te specimen collected:		
□ Confirmed	YYYY / MM / DD	□ Contact	YYYY / MM	/ DD YYY	YY / MM / DD		
□ Does Not Meet Case	YYYY / MM / DD	□ Not a Contact	YYYY / MM	/ DD Spe	ecimen type:		
☐ Person Under Investigation	YYYY / MM / DD	□ Person Under Investigation	YYYY / MM	/ DD	□ Blood □ Urine		
□ Probable	YYYY / MM / DD	1			□ Stool		
Disposition:		<u>4</u>)		•			
FOLLOW UP: ☐ In progress	YYYY / MM / DD	☐ Complete		YYYY / MM	/ חח		
☐ Incomplete - Declined	YYYY / MM / DD	□ Not required	ı	YYYY / MM			
☐ Incomplete – Lost contact	YYYY / MM / DD	☐ Referred – Ou		YYYY / MM			
☐ Incomplete – Unable to locate	YYYY / MM / DD	(specify where)					
REPORTING NOTIFICATION		Location:					
Name of Attending Physician or Nu	irse:						
Physician/Nurse Phone number:		Date Receive	d (Public Health	n): YYYY / MN	M / DD		
Type of Reporting Source: ☐ Hea	————————ા alth Care Facility □ ૄા	_ab Report □ Nurse Practiti	ioner □Phy	/sician □ Ot	her		

September 1, 2018 Page 1 of 4

Haemophilus inflenzae infection (invasive) Data Collection Worksheet

Please complete all sections.

 Panorama Client ID:
Panorama Investigation ID:

	No Yes – Date of o	acot D-	scription	Nia	Yes - Date of onset
Description	No Yes – Date of o	nset De	scription	No	
Arthritis - septic	YYYY / MM /	DD Let	hargy (fatigue, drowsiness, weakness, etc)		YYYY / MM / DD
Bulging fontanelle	YYYY / MM /	DD Me	ningitis		
Cardiac - pericarditis	YYYY / MM /	DD Ne	ck stiffness (nuchal rigidity)		YYYY / MM / DD
Cellulitis	YYYY / MM /	DD Coi	nfusion		YYYY / MM / DD
Dyspnea (shortness of breath)	YYYY / MM /	DD Pn	eumonia		YYYY / MM / DD
Epiglottitis	YYYY / MM /	DD Res	spiratory compromise		YYYY / MM / DD
Fever	YYYY / MM /	DD Sep	osis (e.g. bactremia, septicemia, etc.)		YYYY / MM / DD
nfection - empyema	YYYY / MM /	DD			
Other s/s	I I				
INCUBATION AND COMMUNICABILIT	Υ		LHN-> INVESTIGATION	l->INCU	BATION & COMMUNICABILIT
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ommunicability for Case (period for tra arliest Possible Communicability Date communicability Calculation Details: RISK FACTORS DESCRIPTION Ontact - Daycare Ontact to a known case (Add'l Info) pecial population — Attends Childcare pecial population — Attends school ravel - Outside of Canada (Add'l Info) ravel - Outside of Saskatchewan, but vithin Canada (Add'l Info)	Yes Start Date YYYY / MM / DD TE YYYY / MM / DD AE YYYY / MM / DD TE	NA, U	Add'I Info	L	HN-> SUBJECT->RISK FACTOR
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Communicability for Case (period for trace arliest Possible Communicability Date Communicability Date Communicability Calculation Details: RISK FACTORS DESCRIPTION Contact - Daycare Contact to a known case (Add'l Info) Special population – Attends Childcare Special population – Attends school Travel - Outside of Canada (Add'l Info) Travel - Outside of Saskatchewan, but within Canada (Add'l Info) IMMUNIZATION HISTORY INTERPRET Interpretation Date: YYYY / Interpretation of Disease Immunity:	Yes Start Date YYYY / MM / DD TE OTE TO T	ed (for age)	Add'I Info LHN -> INVESTIGATION-> IMMUNIZATION F	HISTOR	HN-> SUBJECT->RISK FACTOR
Communicability for Case (period for trace arliest Possible Communicability Date Communicability Date Communicability Calculation Details: RISK FACTORS	Yes Start Date YYYY / MM / DD TE YYYY / MM / DD AE YYYY / MM / DD TE YYYY / MM / DD	ed (for age)	LHN -> INVESTIGATION-> IMMUNIZATION F	HISTOR	HN-> SUBJECT->RISK FACTOR

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Haemophilus inflenzae infection (invasive) Data Collection Worksheet

Please complete **all** sections

		Panorama Client ID:Panorama Investigation ID:
G) TREATMENT		INVESTIGATION-> MEDICATIONS->MEDICATIONS SUMMARY
Medication (Panorama = Other Meds) :		
Prescribed by:	Started on:	YYYY / MMM / DD

Prescribed by:				_ Started on: YYYY / MMM / DD		
H) INTERVENTIONS				INVESTIGATION->TREATMENT & INTERV	ENTIONS->INTERVENTION	ON SUMMARY
Intervention Type a	nd Sub Type:			THE CONTROL OF THE PROPERTY OF THE PERSON		J. (J. () () () () () () () () () (
Assessment:	7.			Isolation:		
☐ Assessed for cont	tacts		YYYY / MM / DD	☐ Facility isolation YYYY / MM / DD ☐ Home isolation YYYY / MM / DD	Investigator name Investigator name	
Investigator name						
Communication: ☐ Other communication:	ation (See Investigator I	Notes)	YYYY / MM / DD	Testing:	2000/ / 200	. / 55
Investigator name Letter (See Document)			YYYY / MM / DD	☐ Laboratory testing recommended Investigator name	YYYY / MN	1 / DD
Investigator name	ment Management)		TTTT / IVIIVI / DD	Treatment:		
□ Letter			YYYY / MM / DD	☐ Treatment not recommended	YYYY / MIV	1 / DD
Investigator name				Investigator name		
General: Investigate			1000//2424/55	Other Investigation Findings:		
☐ Disease-Info/Prev☐ Disease-Info/Prev☐	v-Control v-Cont/Assess'd for Con	tacts	YYYY/ MM / DD YYYY/ MM / DD	☐ Investigator Notes ☐ See Doc	ument Management	
	ng: Investigator name		,	Referral:		
☐ Prevention/Conti	-		YYYY / MM / DD	_	y Care Provider	
☐ Disease informat	ion provided		YYYY / MM / DD	☐ Infectious Disease Specialist		
	ations recommended mmunization recomme mmunization given	nded	YYYY / MM / DD YYYY / MM / DD YYYY / MM / DD			
Date	Intervention subtype	Comme	nts		Next follow-up Date	Initials
YYYY / MM / DD					YYYY / MM / DD	
YYYY / MM / DD					YYYY / MM / DD	
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YYYY / MM / DD					YYYY / MM / DD	
YYYY / MM / DD					YYYY / MM / DD	

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Haemophilus inflenzae infection (invasive) Data Collection Worksheet

Please complete **all** sections

				a Client ID: tigation ID:
OUTCOMES (op	ntional except for severe inf	luenza,	LHN-> INVE	STIGATION-> OUTCOM
☐ Not yet recover☐ Recovered☐ Fatal	red/recovering YYYY / MM YYYY / MM YYYY / MM	1 / DD □ Intubation /ventilation YYYY / MM / [DD □ Unknown	
Cause of Death: (if	Fatal was selected)			
Transmission Transmission Event ID	Exposure Name	Setting type (Consider the following settings for TE; if >1 select	Date/Time	# of contacts
		"multiple settings" in Panorama) □ Congregate/Communal living □ Health Care setting □ Type of community contact □ Household Exposure □ Public facilities (e.g daycare)		
		□ Congregate/Communal living □ Health Care setting □ Type of community contact □ Household Exposure □ Public facilities(e.g daycare)		
		□ Congregate/Communal living □ Health Care setting □ Type of community contact □ Household Exposure □ Public facilities (e.g daycare)		
	Hib Contacts – Inv ID#	☐ Multiple Settings	YYYY / MM / DD to YYYY / MM / DD	

September 1, 2018 Page 4 of 4

1. What is Haemophilus influenzae type b disease?

• *Haemophilus influenzae* type b (Hib) is a bacterial infection that can cause deadly brain infections and other serious infections like meningitis (swelling of the covering of the brain and spinal cord), blood infections, pneumonia (lung infection), and infections of other parts of the body.

2. How is Hib disease spread?

 Hib is carried in the nose and throat of children and adults who may be healthy or have mild symptoms. It is then spread to another person through contact with discharges or droplets from the nose or mouth through activities such as kissing, sharing food, utensils, or glasses, or coughing directly on someone.

3. What will decrease the risk of getting Hib disease?

- The best way to prevent infection with Hib is through immunization.
- Hib vaccine is recommended for all children starting at 2 months of age. Children need a total of 4 shots, given at 2, 4, 6, and 18 months. This vaccine is included in routine immunization provided by Saskatchewan Ministry of Health. Make sure your child's immunizations are up to date.

4. What are the signs and symptoms of Hib disease?

- Hib infections are sometimes difficult to recognize. In general, any infection that seems more serious than usual should be brought to a doctor's attention.
 Symptoms to look for are:
 - fever:
 - drowsiness:
 - stiff neck;
 - rapid or difficult breathing;
 - loss of appetite;
 - skin or joints that are red, tender, or swollen.
- If you child develops any of these symptoms, see a doctor immediately for treatment.

5. How is Hib disease diagnosed?

• Lab tests look for the bacteria from various sites (blood, cerebrospinal fluid, etc.) from individuals who are ill.

6. How is Hib disease treated?



Attachment – Sample Fact Sheet on Haemophilus Influenzae Type B Disease
Page 2 of 2
2011 11 01

• Hib is treated with antibiotics. Treatment with antibiotics should be started immediately to reduce serious complications.

7. Who should receive preventive treatment?

Medications to prevent getting or spreading Hib may be needed for those who live
in the same house with a child who had Hib disease, and for children and
employees in childcare settings. Check with your local public health office for
advice.

8. What are the long-term complications of Hib disease?

- Hib can infect the throat and then can spread causing meningitis, pneumonia, or ear, skin, joint, or blood infections.
- If Hib meningitis occurs, death occurs in one out of 20 children and permanent brain damage in 10-30 % of the survivors.

9. Who is at risk of getting Hib disease?

• Since the vaccine was introduced, the disease usually occurs in children who are too young to be immunized or children who have not finished their immunizations.

For more information contact: Your local public heath office, OR your physician or nurse practitioner, OR the HealthLine at 1-877-800-0002.

References:

American Academy of Pediatrics, 2009. Control of Communicable Disease Manual, Heymann (2008).



Attachment – Sample Letter about *Haemophius Influenzae* Type B Invasive Disease – Prophylaxis NOT Recommended
Page **1** of **1**2011 11 01

Date
Dear Parent/Guardian:
This letter is to let you know that your child had contact with a child who has been diagnosed with an infection caused by <i>Haemophilus influenzae</i> type b (Hib). Hib is a bacteria ("germ") that causes serious infections. More information about Hib is included in the attached Fact Sheet.
Hib infections are sometimes difficult to recognize. In general, any infection that seems more serious than usual should be brought to a doctor's attention. Symptoms to look for: • drowsiness; • stiff neck; • rapid or difficult breathing; • extreme irritability; • skin or joints that are red, tender, or swollen.
Notify Public Health at if your child becomes ill with any of the symptoms listed above.
The risk of your child getting this illness is low and Public Health is NOT recommending that your child receive any medicine. Further you should watch your child for fever, excessive sleepiness, trouble breathing, stiff neck, sore throat, or joint or skin infection. Call your doctor immediately if your child becomes sick.
Your child may have received immunizations for Hib as an infant. You should however make sure your child's immunizations are up to date. This will help protect your child. If you have other children under 5 years of age that have not been completely immunized for Hib, they should receive the vaccine.
If either you or your physician require(s) further information, please callYours sincerely,
Medical Health Officer



Attachment – Sample Letter about *Haemophilus Influenzae* Type B Invasive Disease
- Prophylaxis Recommended
Page **1** of **1**2011 11 01

Date
Dear Parent/Guardian:
This letter is to let you know that your child had contact with a child who has been diagnosed with an infection caused by <i>Haemophilus influenzae</i> type b (Hib). Hib is a bacteria ("germ") that causes serious infections. More information about Hib is included in the attached Fact Sheet.
Hib infections are sometimes difficult to recognize. In general, any infection that seems more serious than usual should be brought to a doctor's attention. Symptoms to look for: • drowsiness; • stiff neck; • rapid or difficult breathing; • extreme irritability; • skin or joints that are red, tender, or swollen.
Notify Public Health at if your child becomes ill with any of the symptoms listed above.
Because your child was at the daycare with an infected child, he or she is considered a "close contact." Public Health recommends that all close contacts be given medication to prevent further spread of the disease. Please contact us as soon as possible. The most common medication recommended to prevent infection is called rifampin.
Your child may have received immunizations for Hib as an infant. You should however make sure your child's immunizations are up to date. This will help protect your child, but he or she still needs to take medication and should be watched carefully for signs and symptoms. If you have other children under 5 years of age that have not been completely immunized for Hib, they should receive the vaccine.
If you have any questions please call
Sincerely,
Medical Health Officer



Dosage Guide based on the noted weight in kg below. Calculate dose based on exact weight. Maximum dose 600 mg once every 24 hrs x 4 days (doses).										
Weight in kg Dosage by age	sed on ex	6	thi. Max	amum d 8	9	mg once 10	15	20	25	30 Max or adult dose
<1 mo of age 10 mg/kg (25 mg/ml suspension)	2.0 ml	2.4 ml	2.8 ml	3.2 ml	3.6 ml	4.0 ml				
>1 mo of age 20mg/kg (25 mg/ml suspension)	4.0 ml	4.8 ml	5.6 ml	6.4 ml	7.2 ml	8.0 ml	12.0 ml	16 ml	20 ml	24 ml

Recommendations

- 1. Use the appropriate weight-specific dose noted in the first column in the chart above for infants and children.
- 2. Rifampin Pediatric Suspension can be prepared as follows:
 - Add contents of 3 300mg caps or 6 150 mg caps of Ripampin to 36 mls of simple syrup to yield a 25 mg/ml suspension.
 - SHAKE WELL.
- 3. Store prepared suspension and simple syrup at room temperature because of their tendency to crystallize if refrigerated.
- 4. Discard prepared suspension after treatment course is completed. Preparation expires after 28 days.
- 5. As much as possible, use only one preparation form per client (i.e., capsule(s) only, or suspension only).
- 6. Give client a Rifampin information sheet. See <u>Appendix F Patient Information Sheets Rifampin</u>

Note:

- Rifampin is contraindicated in pregnancy. Discuss Ceftriaxone dose with MHO.
- If necessary, discuss alternative treatments with MHO for non-pregnant adults.



Notification Timeline:

From Lab/Practitioner to Public Health¹:

Severe or Novel: Within 24 hours.

Non-severe or non-novel: Within 48 hours.

From Public Health to Ministry of Health:

Individual case reporting of severe or novel: Within 24 hours (see Attachment –

Severe Influenza in Panorama)

Individual case reporting of non-severe or non-novel: Not required (see

Attachment - Non-severe Influenza in Panorama)

Outbreaks: Initial report within 24 hours.

Updates as necessary.

Final report within 30 days of completing the investigation.

Public Health Follow-up Timeline:

Severe or Novel: Within 24 hours.

Non-severe or non-novel: No follow-up required.

Public Health Purpose for Notification of Influenza

- Timely detection of severe morbidity and mortality caused by common strains of the influenza virus or its variants. This may include the exacerbation of underlying medical conditions resulting in the need for intensive medical care.
- To provide an early warning mechanism in order that available control measures may be implemented at the appropriate time to minimize transmission.
- To track epidemiology trends of severe influenza in Saskatchewan including risk factors and distribution; and
- To inform the public and medical community about influenza.

¹ Local public health is encouraged to collaborate with their partners in ERs and hospitals to ensure all roles and responsibilities are well understood and agreed upon, specifically the timely reporting to public health upon admitting a client with severe influenza. The Severe Influenza Notification Form should be given to ERs and hospitals along with the fax number where to send completed forms.



2018 11 01

Surveillance Case Definitions² (Public Health Agency of Canada, May 2008)

Confirmed Case

Clinical illness^a with laboratory confirmation of infection:

isolation of influenza virus from an appropriate clinical specimen

OR

 demonstration of influenza virus antigen in an appropriate clinical specimen

OR

 significant rise (e.g., 4 fold or greater) in influenza IgG titre between acute and convalescent sera

OR

detection of influenza virus RNA.

^aClinical illness defined as influenza-like illness (ILI) is characterized as abrupt onset of respiratory illness with fever and cough and with one or more of the following:

- sore throat;
- arthralgia;
- myalgia;
- prostration that could be due to influenza virus.

In children under 5, gastrointestinal symptoms may also be present. In patients under 5, or 65 and older, fever may not be prominent.

Note: Illness associated with novel influenza viruses may present with other symptoms.

Other definitions

Severe Influenza b (Saskatchewan Ministry of Health, adapted from Public

Health Agency of

Canada)

A person requiring intensive medical care with:

I. Respiratory symptoms

 Fever (over 38 degrees Celsius)^c AND new onset of or exacerbation of chronic cough or breathing difficulty

AND

II. Evidence of severe illness progression

- Either radiographic evidence of infiltrates consistent with pneumonia OR acute respiratory distress syndrome (ARDS)
 OR
- Severe ILI, which may also include complications, such as encephalitis or other severe and life threatening complications or exacerbation of existing medical conditions

AND

requiring mechanical ventilation

² Surveillance case definitions ensure uniform reporting to allow comparability of surveillance data. The definition is not intended to be used for clinical or laboratory diagnosis or management of cases.



	AND III. Diagnostic criteria Results of laboratory investigations are positive for influenza A or B virus
Severe influenza case - deceased	 I. A person meeting the definition of severe influenza case resulting in death. OR
	 II. Autopsy performed with findings consistent with severe influenza Autopsy findings consistent with the pathology of ARDS AND III. Diagnostic criteria Results of laboratory investigations are positive for influenza virus

^b The indicator of severe influenza is requiring intensive medical care which means mechanical ventilation defined as artificial ventilation where mechanical means is used to assist or replace spontaneous breathing. This includes mechanisms such as continuous positive airway pressure (CPAP), ventilators, and respirators.

Novel Influenza A (Saskatchewan Ministry of Health, adapted from US Centers for Disease Prevention and Control, 2014)

Confirmed	 A case of human infection with a novel influenza A virus confirmed by National Microbiology Laboratory or using methods agreed upon as noted in Laboratory Criteria d.
Probable	 A case meeting the clinical criteria ^e and epidemiologically linked ^c to a confirmed case, but for which no confirmatory laboratory testing for influenza virus infection has been performed or test results are inconclusive for a novel influenza A virus infection.
Suspect	 A case meeting the clinical criteria, pending laboratory confirmation. Any case of human infection with an influenza A virus that is different from currently circulating human influenza H1 and H3 viruses is classified as a suspected case until the confirmation process is complete.
Epi-linked ^f	 The patient has had contact with one or more persons who either have or had the disease, AND Transmission of the agent by the usual modes of transmission is
d	plausible.

^d A human case of infection with an influenza A virus subtype that is different from currently circulating human influenza H1 and H3 viruses. Novel subtypes include, but are not limited to, H2, H5, H7, and H9 subtypes. Influenza H1 and H3 subtypes originating from a non-human species or from genetic reassortment between animal and human viruses are also novel subtypes. Novel subtypes will be detected with methods available for detection of currently circulating human influenza viruses at state



^C Age should be taken into consideration in the clinical assessment.

public health laboratories (e.g., real-time reverse transcriptase polymerase chain reaction [RT-PCR]). Confirmation that an influenza A virus represents a novel virus will be performed by NML. Once a novel virus has been identified by NML, confirmation may be made by public health laboratories following NML-approved protocols for that specific virus, or by laboratories using an authorized test specific for detection of that novel influenza virus.

^e An illness compatible with influenza virus infection (fever >38 degrees Celsius, with cough and/or sore throat).

^f A case may be considered epidemiologically linked to a laboratory-confirmed case if at least one case in the chain of transmission is laboratory confirmed. Laboratory testing for the purposes of case classification should use methods verified by NML. Currently, only viral isolation, RT-PCR, gene sequencing, or a 4-fold rise in strain-specific serum antibody titers are considered confirmatory.

Notifying the local Medical Health Officer of a Severe Novel Influenza case

Emergency Rooms (ERs) and hospitals must update the local medical health officers <u>immediately</u> when a patient with severe influenza is suspected or determined to be infected with a novel strain.

Epidemiology and Occurrence

The occurrence and epidemiology of seasonal influenza varies by year. Generally, it occurs in the winter months between October and March. It has a more severe manifestation in those with Risk Factors. Refer to the Weekly Influenza Activity in Saskatchewan Report for current information.

Table 1. Severe Influenza and Related Deaths per Influenza Season

Influenza season (year)	Severe influenza cases	Deaths	Total influenza
2013-14	56	16	1590
2014-15	26	9	1732
2015-16	54	17	3192
2016-17	26	4	2153
2017-18	51	16	3446



Additional Background Information

Causative Agent

Three strains of human influenza virus exist: they are type A, B, and C. Influenza types A and B are associated with seasonal epidemics. Emergence of completely new subtypes (antigenic shift) occurs at irregular intervals and occurs only with type A viruses. They are responsible for pandemics and result from the unpredictable recombination of human, swine, or avian (usually duck) antigens. The relatively minor antigenic changes (i.e., antigenic drift) of A and B viruses, that are responsible for frequent epidemics and regional outbreaks, occur constantly.

Symptoms

Acute upper respiratory tract infection (URTI) characterized by abrupt onset of fever and chills; headache; malaise; myalgia; prostration; sore throat and cough (Taubenberger, 2008). Abdominal pain, nausea, and vomiting may also be present. Refer to Case
Definition and ILI for details.

Reservoir/Source

Primarily humans. Birds and mammalian reservoirs such as swine are likely sources of new human subtypes thought to emerge through genetic re-assortment.

Incubation Period

Usually 1-3 days.

Period of Communicability

Contagious from 24 hours before the onset of symptoms to 3-5 days after peak symptoms appear.

Mode of Transmission

- Breathing droplets that have been sneezed or coughed into the air by someone with influenza, or having the droplets land on the surface of your eye.
- Shaking hands with an infected person or touching a contaminated surface, and then touching your own eyes, nose or mouth.

Risk Factors

Risk factors are associated with individual susceptibility and settings that create opportunities for acquisition or transmission to others. This includes:

- Individuals with the following medical conditions:
 - Cardiac Disease;



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- o Diabetes mellitus;
- Lung disease including asthma;
- o Cancer;
- o Renal disease;
- Immunocompromised related to underlying disease or treatment;
- Transplant candidates or recipients;
- Neurological conditions that impede the clearance of respiratory secretions
- Individuals that are morbidly obese;
- Pregnant women;
- Children under the age of 5;
- Adults 65 years of age and older;
- Children in childcare;
- Individuals in long term care facilities, homeless shelters or crowded living conditions or communal settings;
- Individuals that use alcohol, tobacco or other drugs; and
- Indigenous individuals.

Specimen Collection and Transport

The recommended specimens for diagnosis of influenza are nasopharyngeal specimens collected on a flocked swab or a vigorous throat swab taken within the first 48 hours of infection. Refer to Roy Romanow Provincial Laboratory (RRPL) Compendium of Tests at https://rrpl-testviewer.ehealthsask.ca/. The specimen should reach the lab in 24 hours.

Each specimen is tested by three methods:

- 1. PCR for influenza viruses;
- 2. the rapid DFA (direct fluorescent antibody) microscopy;
- 3. virus isolation in tissue culture cells.

All specimens are tested by PCR within 24 hours of receipt. If PCR is negative, DFA and culture results will be available within 5 days of receipt of specimen.



Treatment/Supportive Therapy

Treatment for clinical management is at the discretion of the primary care provider. The following serves as a reference for the public health investigator:

- Supportive care for symptoms is all that is indicated for most cases of influenza.
- An appropriate antiviral may be effective in reducing the duration of the illness when initiated by the attending physician within 48 hrs of the onset of signs and symptoms.
- Refer to Association of Medical Microbiology and Infectious Disease Canada (AMMI) guidelines on the use of antivirals (http://www.ammi.ca/guidelines/).
- Antibiotic therapy is not indicated unless bacterial complications arise.
- Because of the association with Reye's syndrome, salicylates (e.g., Aspirin) should be avoided in children with influenza.

Public Health Investigation

I. Case

History

Refer to <u>Attachment – Influenza Data Collection Worksheet</u> to assist.

- Investigation of influenza cases when influenza is not circulating broadly is recommended to determine if influenza has been imported from another area (i.e. travel-related).
- During influenza season, investigations are limited to cases with severe presentations or those infected with a novel strain.
 - o Review if immunized for the current influenza season.

Public Health Interventions

No follow-up required for non-novel and non-severe cases during seasonal influenza.

Assessment

In the case of novel influenza – assess for contacts.

Communication

- Communication with health care providers is important to determine the clinical presentation of severe cases.
- Letters can be sent to group settings where cases of novel influenza have appeared to inform them of the exposure, symptom monitoring and when to seek medical attention.



Education

 All recovered severe cases should be provided information on influenza immunization programs for future reference.

Exclusion and Isolation

- See Epidemic Measures.
- For additional information on infection prevention and control measures for individuals in health care facilities refer to Regional Infection Control Manual.
- Health Care Workers (HCWs) refer to Regional Management of Employees and Other Health Care Workers during Influenza Outbreaks in Health Care Facilities.

Immunization

Offer relevant immunizations if eligible.

II. Contacts/Contact Investigation

Contact tracing is not required except in the case of novel influenza. See Epidemic Measures.

III. Environment

Child Care Centres/Institutional Control Measures

- Child care centres refer to the Saskatchewan Ministry of Health Infection Control Manual for Child Care Facilities.³
- Health care facilities refer to regional infection control manual.

IV. Epidemic Measures

- Child care centre (CCC) control measures:
 - o Educate as per <u>Prevention Measures</u>.
 - Children with influenza or influenza-like illness should not attend until the child has been without fever (without the use of fever reducing medications) for 24 hours (Centers for Disease Control, July 2009).
 - For CCC with children under 5 years, review immunization records of those under
 5 years of age and offer influenza if eligible.
- Institutional control measures:
 - Educate as per Prevention Measures.
 - Persons in the community with influenza or influenza-like illness should not visit until 5 days after onset of symptoms. Exceptional circumstances should be discussed with facility manager and MHO.

³ http://publications.gov.sk.ca/documents/11/96181-infection-control-manual-child-care-centres.pdf.



- Every effort should be made to control influenza outbreaks within institutions to optimize the protection of the patients, staff and the community. The use of antivirals has been used to control outbreaks in Special Care Homes. Refer to Use of Oseltamivir for the Management of Influenza Outbreaks in Special Care Homes.
- Infection control measures included in the region's infection control manual should be reviewed with staff.
- Ensure cases are reported to local public health.
- Refer to the <u>Outbreaks</u> section of the manual for additional details about managing an outbreak in a Special Care Home.

NOTE: The MHO is the only designated Public Health Official legislated to declare and/or end an outbreak.

V. Pandemic Measures

See local, provincial, national pandemic plans.

Prevention Measures

Immunization

- Refer to the National Committee on Immunization Statement on Influenza Vaccination for the current season at http://www.phac-aspc.gc.ca/naci-ccni/indexeng.php.
- Eligible persons should be immunized annually because of declining immunity and change in virus variants.
- Refer to Saskatchewan Ministry of Health's Seasonal Influenza Program for recommendations on risk groups, dosages and schedules.
- Adults do not benefit from multiple doses in the same year; re-immunization may be considered in outbreak situations or for high-risk travellers; discuss with the Medical Health Officer (MHO).
- Educate the public about the disease: transmission, symptoms, and preventive measures especially hand hygiene and cough etiquette.
- Encourage immunization of health care workers. Lower mortality in long-term care facilities has been demonstrated in institutions where health care workers are immunized than in those where they were not.
- Administration of influenza vaccine to international travellers should be considered
 refer to Saskatchewan International Travel Manual.



Respiratory and Direct Contact
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Education

• Education should be provided regarding respiratory etiquette and measures to prevent transmission.

Surveillance

The province-wide influenza surveillance system is a component of the national FluWatch program in Canada, a program that is ongoing year-round. The national program is part of international surveillance by World Health Organization (WHO). The Saskatchewan program comprises several components including year-round laboratory surveillance, sentinel physicians submitting specimens on respiratory patients during influenza season, active community sentinel site surveillance, emergency room (ER) influenza-like-illness surveillance, HealthLine calls and outbreaks during the influenza season.

- Laboratory surveillance is conducted through epidemiological analyses (number of cases, week of specimen collection, age category, health area, etiological type) associated with positive influenza specimens processed by RRPL and Royal University Hospital.
- Sentinel physicians are a network of local physicians across the province who submit specimens to the Virology Section of RRPL from patients presenting with influenzalike symptoms. They also report weekly on the number of patients with influenzalike symptoms among all the patients seen on a chosen day in their practice.
- Active community sentinel site surveillance supplements the other components of
 influenza surveillance. Public Health in all health areas conduct active community
 surveillance and report weekly to the, Ministry of Health at cdc@health.gov.sk.ca on the
 Attachment Community Influenza-Like Illness Weekly Surveillance Form. Sentinel
 institutions such as schools, long-term care facilities and workplaces provide weekly
 reports of absenteeism/illness related to influenza-like illness. See Attachment Details
 of Influenza Surveillance Programs.
- Emergency Room (ER) monitoring of visitors to ERs with influenza-like symptoms-is an
 early alert system of increased influenza-like activity in the community. See Attachment
 Details of Influenza Surveillance Programs.
- *HealthLine* inquiries from callers with influenza-like symptoms is an early alert system of increased influenza-like activity in communities across the province.
- Outbreaks from confirmed influenza are analyzed from the Outbreak Notifications
 reported to Population Health Branch, Ministry of Health. These are typically institutional
 outbreaks in facilities where vulnerable residents live. These counts represent the
 effectiveness of the seasonal influenza vaccine among residents and
 staff and infection control measures in the facility. Outbreak notification are



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emailed to the Ministry of Health at cdc@health.gov.sk.ca at all phases of the outbreak investigation- initial notification to public health, updates and final summary report. See Attachment - Outbreak Notifications.

 Deaths of individuals receiving intensive medical care are counted for reporting and surveillance purposes as an indicator of severe influenza. It is considered the outcome of a severe influenza event.



Respiratory and Direct Contact Section 2 – 60 – Influenza Page 12 of 14 2018 11 01

Revisions

Date	Change
November 2018	 Updated to incorporate severe and novel case definitions Incorporated the purpose for notification of cases to public health Updated to align with Panorama configuration Incorporated an Epidemiology and Occurrence as a placeholder Rearranged and updated the style into the new format of the Manual. Incorporated details of Influenza Surveillance Program within and added an attachment with further details.



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Severe Influenza Notification Form



YYYY / MMM / DD

) PERSON REPORTING – HEALTH CARE	PROVII	DER INFORMATION					
Hospital Name and Unit:			Patient informatio	Patient information sticker or addressograph			
Location:							
Attending Physician or Nurse:							
Phone number:							
ADDITIONAL CUENT INFORMATION			a cattala A				
ADDITIONAL CLIENT INFORMATION	(not inc						
Last Name:		First Name: and	Middle Name:	HSN:	DOB:		
Place of Employment/School:		Comments:					
Allowed a Control							
Alternate Contact:							
Relationship:							
phone:							
DISEASE EVENT HISTORY Presentation: □ Severe Date of Influenza Immunization: YYYY SIGNS & SYMPTOMS			lts pending? eceive vaccine or □ U	Inknown			
Description	No	Yes – Date of onset	Description	No	Yes - Date of onset		
Acute onset of symptoms		YYYY / MMM / DD	Muscle inflammation (myo	sitis)	YYYY / MMM / DD		
Acute respiratory distress syndrome ARDS)		YYYY / MMM / DD	Myalgia (muscle pain)		YYYY / MMM / DD		
Arthralgia		YYYY / MMM / DD	Nasal congestion		YYYY / MMM / DD		
Bronchiolitis		YYYY / MMM / DD	Neurologic - delerium		YYYY / MMM / DD		
Cardiac - myocarditis		YYYY / MMM / DD	Otitis media		YYYY / MMM / DD		
Chills		YYYY / MMM / DD	Pain - abdominal		YYYY / MMM / DD		

Pharyngitis (sore throat)

Respiratory compromise

Respiratory failure - requiring mechanical

Pneumonia - CXR/CT

Prostration

ventilation

Seizures

Sinusitis

Reye's syndrome

YYYY / MMM / DD

Coryza or rhinitis

Encephalitis

Headache

Malaise

Fever

Croup (laryngotracheobronchitis)

Dyspnea (shortness of breath)

Gastrointestinal symptoms



Severe Influenza Notification Form



E) RISK FACTORS	T			
DESCRIPTION	Yes Start date if applicable	N, NA, U	Add'l Info	
Access to healthcare services > 4 hours by road				
Chronic Medical Condition - Cardiac Disease				
Chronic Medical Condition - Diabetes Mellitus				
Chronic Medical Condition - Lung Disease+				
Chronic Medical Condition - Malignancies/Cancer				
Chronic Medical Condition - Morbid Obesity				
Chronic Medical Condition - Neurological conditions that impede the clearance of respiratory/oral secretions				
Chronic Medical Condition - Other (add'l info)				
Chronic Medical Condition - Renal Disease				
Contact to a known case (add'l info)	YYYY / MM/DD			
Exposure - Second hand smoke				
Immunocompromised - Related to underlying disease or treatment				
Immunocompromised - Transplant Candidate or Recipient - Solid Organ/Tissue				
Setting - Crowded living conditions (>1 person per				
room excluding bathrooms) Special Population - Attends childcare				
Special Population - Homeless				
Special Population - Lives in a communal setting				
Special Population - LTC Facility				
Special Population - Pregnancy				
Special Population - Self-reported Indigenous identity				
Substance Use - Alcohol				
Substance Use - Injection drug use (including steroids)+				
Substance Use - Tobacco				
Travel - Outside of Canada (Add'l Info)	YYYY / MM/DD			
Travel - Outside of Saskatchewan, but within Canada (Add'l Info)	YYYY / MM/DD			
F) OUTCOMES (For hospitalization and ICU, please in	iclude admission date: for in	ntubation/ve	ntilation, please use date in	itiated.)
□ Not yet recovered/recovering YYYY / MM / DD	☐ ICU/intensive medi		·	talization YYYY / MM / DD
☐ Intubation /ventilation	□ Unknown		YY / MM / DD	tanzation in a party population
□ Fatal YYYY / MM / DD	□ Other			
How was Influenza Related to Cause of Death: (if Fata	l was selected)			
Initial Report				Date initial report completed:
completed by:				YYYY / MM / DD
<u>'</u>				ı
PLEASE FAX TO THE PUBLIC HEALTH OFFICE	:			

THANK YOU.

CONFIDENTIAL FAX #:



Please complete the following sections:



Panorama QA complete: ☐ Yes ☐ No Initials:

Severe - intensive medical care - Sections D, F, G, and I;

Novel - Sections D, E, F, H, I, J, K and L;

Panorama Client ID:	
Panorama Investigation ID:	

A) CLIENT INFORMATION			LHN -> SUBJE	CT -> CLIENT	DETAILS -> PERSONAL INFORMATION	
Last Name:		First Name: and Middle Name:		Alternate N	lame (Goes by):	
DOB: YYYY / MM / DD Age: Phone #: Primary Home:		Health Card Province: Health Card Number (PHN):		Preferred Communication Method: (specify i.e. home phone, text): Email Address: □Work □Personal		
Place of Employment/School:		Gender: □ Male	□ Female	<u> □</u> 01	ther 🗖 Unknown	
Alternate Contact: Relationship: Alt. Contact phone:		Address Type: No fixed Postal Address Primary Home Temporary Legal Land Description Mailing (Postal address): Street Address or FN Community (Primary Home): Address at time of infection if not the same:				
B) INVESTIGATION INFORMATION	LHN ->SUBJE	CT SUMMARY-> RESPIRATORY &	DIRECT CONTA	CT ENCOUNT	ER GROUP-> CREATE INVESTIGATION	
Disease Summary Classification: CASE	Date	Classification: CONTACT	Date		LAB TEST INFORMATION: Date specimen collected:	
□ Confirmed	YYYY / MM / DD	□ Contact	YYYY / MM	/ DD	YYYY / MM / DD	
☐ Does Not Meet Case Definition	YYYY / MM / DD	□ Not a Contact	YYYY / MM / DD		Specimen type:	
☐ Person Under Investigation	YYYY / MM / DD	□ Person Under Investigation	YYYY / MM / DD		□ Nasopharyngeal □ Swab	
□ Probable	YYYY / MM / DD					
Disposition: FOLLOW UP: ☐ In progress ☐ Incomplete - Declined ☐ Incomplete - Lost contact ☐ Incomplete - Unable to locate	YYYY / MM / DD YYYY / MM / DD YYYY / MM / DD YYYY / MM / DD	☐ Complete ☐ Not required ☐ Referred – Ou (specify where)		YYYY / MI YYYY / MI YYYY / MI	M / DD	
REPORTING NOTIFICATION Name of Attending Physician or Nu	rse:	Location:				
Physician/Nurse Phone number:		Date Received	d (Public Health	n): YYYY /	MM / DD	
Type of Reporting Source: ☐ Hea	llth Care Facility □L	ab Report	oner Phy	ysician 🗆	Other	
C) DISEASE EVENT HISTORY		LHN-> INVESTIGATI	ON->DISEASE S	SUMMARY (U	PDATE)->DISEASE EVENT HISTORY	
_	Severe - intensive medic Novel	Complete section			Cases;	

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 \square Other

Please complete all sections.

Panorama Client ID:	
Panorama Investigation ID:	

SIGNS & SYMPTOMS	INVESTIGATION->SIGNS & SYMPTON
	SIGNS & SYMPTOMS

Description	No	Yes – Date of onset	Description	No	Yes - Date of onset
Acute onset of symptoms		YYYY / MMM / DD	Muscle inflammation (myositis)		YYYY / MMM / DD
Acute respiratory distress syndrome (ARDS)		YYYY / MMM / DD	Myalgia (muscle pain)		YYYY / MMM / DD
Arthralgia		YYYY / MMM / DD	Nasal congestion		YYYY / MMM / DD
Bronchiolitis		YYYY / MMM / DD	Neurologic - delerium		YYYY / MMM / DD
Cardiac - myocarditis		YYYY / MMM / DD	Otitis media		YYYY / MMM / DD
Chills		YYYY / MMM / DD	Pain - abdominal		YYYY / MMM / DD
Coryza or rhinitis		YYYY / MMM / DD	Pharyngitis (sore throat)		YYYY / MMM / DD
Cough		YYYY / MMM / DD	Pneumonia - CXR/CT		YYYY / MMM / DD
Croup (laryngotracheobronchitis)		YYYY / MMM / DD	Prostration		YYYY / MMM / DD
Dyspnea (shortness of breath)		YYYY / MMM / DD	Respiratory compromise		YYYY / MMM / DD
Encephalitis		YYYY / MMM / DD	Respiratory failure - requiring mechanical ventilation		YYYY / MMM / DD
Fever		YYYY / MMM / DD	Reye's syndrome		YYYY / MMM / DD
Gastrointestinal symptoms		YYYY / MMM / DD	Seizures		YYYY / MMM / DD
Headache		YYYY / MMM / DD	Sinusitis		YYYY / MMM / DD
Malaise		YYYY / MMM / DD			YYYY / MMM / DD

Maiaise	YYYY / MININI / DD			TTTT / IVIIVIIVI / DD
E) INCUBATION AND COMMUNICABILITY	FOR NOVEL INFLUENZA ONLY	LHN-> INVESTIGATION	I->INCl	JBATION & COMMUNICABILIT
Incubation for Case (period for acquisitio	n):			
Earliest Possible Exposure Date: YYYY /	MM / DD	Latest Possible Exposure Date:	YYYY	/ MM / DD
Exposure Calculation details:				
Communicability for Case (period for tran	•			
Earliest Possible Communicability Date:	YYYY / MM / DD	Latest Possible Communicability	Date:	YYYY / MM / DD
Communicability Calculation Details:				

F) RISK FACTORS FOR NOVEL AND SEVERE INFLI	JENZA ONLY			LHN-> SUBJECT->RISK FACTOR
DESCRIPTION	Start date Yes	N, NA, U	Add'l Info	
Access to healthcare services > 4 hours by road				
Chronic Medical Condition - Cardiac Disease+				
Chronic Medical Condition - Diabetes Mellitus+				
Chronic Medical Condition - Lung Disease+				
Chronic Medical Condition - Malignancies/Cancer+				
Chronic Medical Condition - Morbid Obesity				
Chronic Medical Condition - Neurological				
conditions that impede the clearance of				
respiratory/oral secretions+				
Chronic Medical Condition - Other (add'l info)				
Chronic Medical Condition - Renal Disease+				
Contact to a known case (add'l info)	YYYY / MM/DD			
Exposure - Second hand smoke				
Immunocompromised - Related to underlying				
disease or treatment				
Immunocompromised - Transplant Candidate or				
Recipient - Solid Organ/Tissue+				
Setting - Crowded living conditions (>1 person per				
room excluding bathrooms)				

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Please complete all sections.

Panorama Client ID:	
Panorama Investigation ID:	

DESCRIPTION		St Ye	art date es	N, NA, U	Add'l Info		
Special Population -	Attends childcare		<u></u>				
Special Population -	Homeless+						
Special Population -	Lives in a communal se	tting					
Special Population -	LTC Facility+						
Special Population -	Pregnancy						
	Self-reported Indigenou	ıs					
identity Substance Use - Alco	ohol						
	ction drug use (includin	g					
steroids)+ Substance Use - Tob	acco						
Travel - Outside of C		YY	YY / MM/DD				
Travel - Outside of S Canada (Add'l Info)	askatchewan, but withir	n AE	YY / MM/DD				
G) IMMUNIZATION	HISTORY INTERPRETAT	IONI SLIMMAR	·v	I HINI >	INVESTIGATION-> IMMUNIZATION	HISTORY INTERDRETAT	ION SHIMMARY
Interpretation Date			v.	LIIIV->	INVESTIGATION-> INIVIONIZATION	III310KI INTEKEKETAT	TON SOMMAN
Interpretation of Dis	sease Immunity:	Disease Case	- Fully immunize	ed (for age)	☐ Disease Case - Partially	immunized	
☐ Disease Case – Ur	nimmunized \Box	Disease Case	- Unclear immur	nization his	tory Valid doses received:		
Reason:							
	history by investigator						
H) INTERVENTION Intervention Type a	and Sub Type:		l	LHN -> INVI	ESTIGATION->TREATMENT & INTERV	/ENTIONS->INTERVENT	ION SUMMARY
Assessment:	mu sub Type.			Isolat			
☐ Assessed for con Investigator name	tacts	Y	/YY / MM / DD		cility isolation Investigator name ome isolation Investigator name	YYYY / N	
Communication:					r Investigation Findings:		
☐ Other communic Investigator name	ation (see Investigator N	Notes) Y	/YY / MM / DD		vestigator Notes e document management	YYYY / N YYYY / N	-
	ment Management)	YY	/YY / MM / DD		e document management	1111 / IV	TIVI / DD
Investigator name General: Investigat	or name			Ouar	antine:		
☐ Disease-Info/Pre		Y	/YY/ MM / DD		uarantine	YYYY / N	1M / DD
l _	v-Cont/Assess'd for Cont		/YY/ MM / DD	Inves	tigator name		
Education/counsell				Testi	_		
☐ Prevention/Cont☐ Disease informat			/YY / MM / DD /YY / MM / DD		b testing recommended YYYY / tigator name	MM / DD	
Exclusion: Investiga			TTT / IVIIVI / DD	Refer	0		
□ Work YYYY / I		ool YYYY / N		□ Cc	nsultation with MHO Investiga	ator name YYYY / N	1M / DD
□ School YYYY / I	MM / DD □ Daycar	e YYYY / N	/IIVI / DD			ator name YYYY / N	-
Immunization:	Investigator name			□ In	fection prevention and Control Inves	tigator name YYYY / N	1M / DD
☐ Eligible Immuniza	ation recommended		/YY / MM / DD				
•	mmunization recomme		/YY / MM / DD				
☐ Disease-specific i	Intervention	Comments	/YY / MM / DD			Next follow-up	Initials
	subtype	Comments				Date Date	Anticuts
YYYY / MM / DD						YYYY / MM / DD	
YYYY / MM / DD				-		YYYY / MM / DD	
YYYY / MM / DD			- 			YYYY / MM / DD	

November 1, 2018 Page 3 of 4

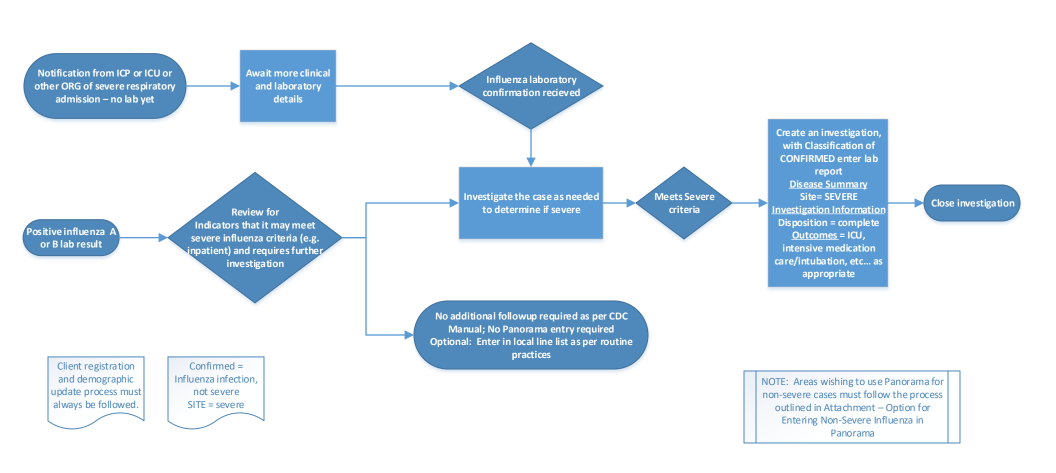
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☐ ER Visit	YYYY / MIV				
☐ Fatal	YYYY / MM	1 / DD Cause of Death: (if Fatal was selected)			
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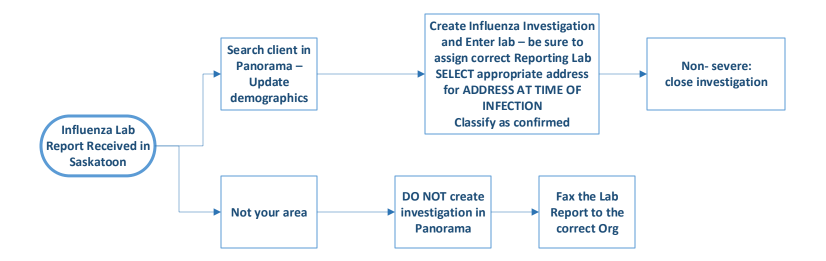
November 1, 2018 Page 4 of 4

Attachment - Severe Influenza in Panorama 2018-19 Influenza Season – DRAFT Following Discussion IOM Key User Group Oct 24, 2018



January 29, 2019 Page 1

Attachment – Non-Severe Influenza in Panorama



Confirmed = Influenza infection, not severe SITE = severe

Attachment – Community Influenza-like Illness Weekly Surveillance Form

Page 1 of 2

2018 12 01

Please see the following pages for the Community Influenza-like Illness Weekly Surveillance Form.





Name of former RHA:	Surveillance date:
	(dd/mm/yy)
Reported by:	Ph:
ACTIVITY LEVEL THIS WEEK (check one):	
0. Community sentinel sites report no influenza-like	ke activity at all.
 Sporadically occurring influenza-like illness but n 	no lab confirmations.
 Sporadically occurring influenza-like illness wher confirmed influenza case but no outbreaks deter 	-
 At least one lab confirmed influenza in your heal outbreak(s) in schools, worksites or a laboratory a residential institution(s). Re-emergent school of there has been an intervening 8 weeks of Level 0 	y confirmed influenza outbreak(s) in outbreaks are considered NEW if
The following influenza/influenza-like outbreaks had the surveillance week. List only those outbreaks meeting th	
surveillance week. List only those outbreaks meeting th	
surveillance week. List only those outbreaks meeting th	his criterion.
surveillance week. List only those outbreaks meeting th	his criterion.
surveillance week. List only those outbreaks meeting th	his criterion.
surveillance week. List only those outbreaks meeting th	his criterion.

Outbreak reporting for influenza follows the same protocol as for all other outbreaks. This table does not constitute an outbreak notification. An outbreak notification form must still be submitted on each outbreak from a communicable disease.

PLEASE EMAIL THIS COMPLETED FORM TO THE MINISTRY OF HEALTH AT cdc@health.gov.sk.ca BY 4 PM EACH THURSDAY. THANK YOU.

ILI is the acute onset of respiratory illness with fever and cough and with one or more of the following: sore throat, arthralgia, myalgia, or prostration that could be due to influenza virus. In children under 5, gastrointestinal symptoms may also be present. In patients under 5 or 65 and older, fever may not be prominent.

Reviewed each influenza season

Influenza
Section 2-60
Attachment – Details of Influenza Surveillance Programs
Page 1 of 4
2018 11 01

Protocol for Establishing Surveillance of Influenza like Illness (ILI) in Community Settings

Involving community partners in influenza-like illness surveillance is needed to determine the arrival and ongoing activity of the influenza virus in the community. ILI surveillance provides the health area with a window of opportunity for alerting long term and acute care facilities that the influenza virus is in their community. Once influenza is identified in the community there is usually a two-week period before the influenza virus spreads to the institutional settings.

Steps in setting a Community Influenza like Illness Surveillance Program

- 1. Decide which community partners to include in your community surveillance program. Facilities such as schools, nursing homes, workplaces and daycares are often the best choices for sentinel sites as influenza-like illness commonly occurs among crowded populations in enclosed places. A variety of sentinel sites ensures ongoing surveillance during school breaks.
- 2. Contact community partners asking for their participation in the influenza-like illness surveillance program. Contact is typically made in early October prior to the beginning of the influenza season. Phone contact should be made one week later to determine if community partners are interested in participating in the influenza-like illness surveillance program.
- 3. Receive weekly reports or emails from your community partners in the annual influenza-like illness surveillance program. The community surveillance season usually begins late October to mid-November, depending on the season, and runs until approximately the end of April. Community partners are requested to fax their "Community Influenza-Like illness Surveillance Reports" to the public health CD/Immunization Coordinator by noon on Wednesday each week. Community partners should be given other options regarding the reporting mechanism (i.e., by email or phone). Their participation involves monitoring the number or proportion of individuals with influenza-like illness that are absent from their facility (on a daily basis) over the past week (Thursday through Wednesday). Client confidentiality is maintained at all times as the names of the individuals are not required.
- 4. Record the data on the Attachment Community Influenza-Like-Illness Weekly Surveillance Form.
- 5. Email the completed <u>Community Influenza-Like-Illness Weekly Surveillance Form</u> to the Ministry of Health at <u>cdc@health.gov.sk.ca</u> <u>no later than 4:00 every Thursday</u>. The email Subject line should read "Community ILI surveillance <former *RHA name*> week ending *<Saturday's date>*".



Emergency Room Influenza-Like Illness Monitoring

• ER monitoring is done for a twenty-four hour period on at least one weekday (the exact time period will vary with the ER schedule). The ER should report to public health services in the health areas on Wednesdays and health areas will report to the Ministry of Health by 4:00 pm on Thursday each week. This may increase to include one weekend day if there is indication that ILI activity is increasing and laboratory-confirmations support the need to do so. FNIHB and NITHA will continue to report to the local health region in which the ER or health clinic is located.

ER surveillance data should be interpreted in conjunction with other influenza surveillance data to establish a clear epidemiological profile of influenza-like illness activity in the community. Other pieces of information to consider are:

- congruence with ILI activity reported by community sentinel sites;
- congruence with the number of respiratory calls to HealthLine;
- congruence with rate of ILI patients seen by sentinel physicians;
- number and distribution of lab-confirmed cases reported to the MHO;
- number and location of respiratory outbreaks in institutions (residential and academic) and workplaces.

Aberrations in trends away from the baseline rates should be investigated further. A rise may indicate an increase in ILI activity in the community or a decrease could indicate an artifactual change in data capture mechanisms. ERs should be reminded of the ILI definition and that symptom onset is abrupt (within 4-6 hours of feeling well).

The count of ILI patients is captured in each of four broad age categories, preschool (~0-4 years), school age (~5-19 years), working age group (~20-64 years), seniors (~65 years plus) as a proportion of total ER admissions in each of those age categories. The age group in which to place a patient will be determined in part by the age groups used by the ER's administrative database. The categories are approximate but provide a general profile of the age groups most affected by ILI.

<u>Instructions for ERs to complete the electronic Excel reporting template:</u>

- By correctly entering data to the template, sums and rates by age group will be calculated automatically. Be careful not to delete data from the shaded or coloured fields to avoid deleting the formulas.
- For the chosen 24-hour surveillance period, the ER will tally the number of patients seen in each of the four broad categories (the broad age groupings may be defined slightly differently across ERs). The number of patients to the ER will be entered on the spreadsheet (right hand boxes).
- The ER will tally the number of patients with influenza-like symptoms in each of the age categories and enter the information on the spreadsheet (left hand boxes).
- Some ERs will wish to capture data on more than one 24-hour period per week. Data for each surveillance 24-hour period will be recorded in a separate row.
- The ER will send the report to local public health services by Thursday morning, 8:00 am.



Instructions for transposing the data from an ER tally sheet to the reporting spreadsheet:

- Public Health services should enter the data from each ER into the Excel reporting format by age group both the number of ILI patients seen and total number of patients seen in the corresponding age group during a 24 hour period on a designated surveillance weekday.
- Data from the row line titled "ILI Sum" on the ER report should be entered in the appropriate cells on lines 9 through 15 of the HR report, one row line for each ER submitting a surveillance report to local public health. Do not enter daily counts for ERs undertaking surveillance on multiple days but rather use the summed data for the reporting week (i.e., "ILI sum" row).
- If more than one ER in your health authority submits data electronically, it will be automatically consolidated when entered into the Excel surveillance tool, resulting in an overall weekly ILI rate for your jurisdiction (i.e., "ILI sum" row).

·			Influenza	a-like IIIn	in Emergency Rooms									
<health region=""></health>			Patient	s with IL	I		Total patients seen for all reasons including ILI						ng ILI	
	Pre school	School age	Working age	Seniors		Total ILI		Pre school	School age	Working age	Seniors		Total patients	
	Approx	Approx	Approx	Approx	Age	all age		Approx 0-4 vr	Approx 5-19 yr	Approx 20-64 vr	Approx 65 +	Age unknown	all age groups	
ILI Su	1	0	0	0	0	0	L	0	0	0	0	0	0	
Rate/1000 patients	•	0.0	0.0	0.0	0.0	0.0	_							
Percent I Emergency Date in		0.0 Approx	0.0 Approx	0.0 Approx	0.0 Age	0.0		Approx	Approx	Apprx	Approx	Age	Total	
room collecte	d 0-4	5-19	20-64	65 +	unknown	Total ILI		0-4	5-19	20-64	65 +	unknown	patients	
						0 0 0			<u>-</u>	The tota rows wil row. Complet	l be dis	played	in the I	LI Sum

NOTE – all shaded cells include formulas that will automatically calculate the totals, rates and percentages.

The completed <u>ER Influenza-Like-Illness Weekly Surveillance Form</u> should be emailed to the Ministry of Health at <u>cdc@health.gov.sk.ca</u> <u>no later than 4:00 pm every Thursday</u>. The email Subject line should read "ER ILI surveillance – <former *RHA name*> week ending <*Saturday's date*>"



Influenza Section 2-60 Attachment – Details of Influenza Surveillance Programs Page 4 of 4 2018 11 01

Revisions

Date	Change
November 2018	New; replaces the annual Influenza Surveillance package.

Attachment – Influenza-like Illness Surveillance in Emergency Rooms
Page **1** of **3**2010 11 01

Please see the following pages for the Influenza-like Illness Surveillance in Emergency Rooms. The excel format includes formulas. Please use the excel document to assist in data submission. The excel document can be located at

https://www.ehealthsask.ca/services/Manuals/Documents/Sec 2-60 Attachment - Influenza ER ILI Data Collection and Submission Tool 2018 11 15.xls



Attachment - Influenza-like Illness Surveillance in Emergency Rooms

Page 2 of 3

2010 11 01

<former Regi</former 				Patient	s with IL	I		Total patients seen for all reasons including IL						
		Pre school	School age	Working age	Seniors		Total ILI		Pre school	School age	Working age	Seniors		Total patients
		Approx 0-4 yr	Approx 5-19 yr	Approx 20-64 yr	Approx 65 +	Age unknown	all age groups		Approx 0-4 yr	Approx 5-19 yr	Approx 20-64 yr	Approx 65 +	Age unknown	all age groups
	ILI Sum	0	0	0	0	0	0		0	0	0	0	0	0
Rate/1000) patients	0.0	0.0	0.0	0.0	0.0	0.0							
	Percent ILI	0.0	0.0	0.0	0.0	0.0	0.0							
Emergency room	Date info collected	Approx 0-4	Approx 5-19	Approx 20-64	Approx 65 +	Age unknown	Total ILI		Approx 0-4	Approx 5-19	Apprx 20-64	Approx 65 +	Age unknown	Total patients
							0							0
							0							0

Influenza-like Illness Surveillance in Emergency Rooms

Instructions for ERs to complete the electronic Excel reporting template:

•For the chosen 24-hour surveillance period(s), the ER will tally the total number of patients seen in each of the four broad age categories. The number of patients seen will be entered on the spreadsheet (right hand boxes)

•The ER will tally the number of patients with ILI symptoms in each of the age categories and enter the information on the spreadsheet (left hand boxes).

•By correctly entering data to the template, sums and rates by age group are calculated automatically in the coloured cells. Be careful not to delete any data in the colored cells to avoid deleting the formulas from these cells.

- Some ERs will wish to capture data on more than one 24-hour period per week. Data for each surveillance 24-hour period will be recorded in a separate row.
- The ER will send the report to local public health services by Thursday morning, 8:00 am.

ILI will manifest as: Acute* onset of respiratory illness with fever and cough and with one or more of the following - sore throat, arthralgia, myalgia, or prostration which could be due to influenza virus.

In children under 5, gastrointestinal symptoms may also be present. In patients under 5 years or 65 and older, fever may not be prominent. The acute (rapid) onset onset of symptoms differentiates ILI patients from those with other viral respiratory illnesses circulating in the community.

*rapid, within 4-6 hours of feeling well

Reviewed October 2018

*This is a replica of the excel document.

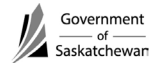


This is an example of an emergency room weekly report to Public Health.

<st joseph's<="" th=""><th>S></th><th></th><th></th><th>Patient</th><th>s with IL</th><th>I</th><th></th><th>Total pa</th><th>atients se</th><th>en for al</th><th>l reason:</th><th>s includi</th><th>ng ILI</th></st>	S>			Patient	s with IL	I		Total pa	atients se	en for al	l reason:	s includi	ng ILI
		Pre school	School age	Working age	Seniors		Total ILI	Pre school	School age	Working age	Seniors		Total patients
		Approx 0-4 yr	Approx 5-19	Approx 20-64	Approx 65 +	Age unknown	all age groups	Approx 0-4	Approx 5-19	Approx 20-64	Approx 65 +	Age unknown	all age groups
	ILI Sum	6	9	17	10	0	42	18	22	80	26	0	146
Rate/1000	patients	333.3	409.1	212.5	384.6	0.0	287.7						
	Percent ILI	33.3	40.9	21.2	38.5	0.0	28.8						
Emergency room	Date info collected	Approx 0-4	Approx 5-19	Approx 20-64	Approx 65 +	Age unknown	Total ILI	Approx 0-4	Approx 5-19	Apprx 20-64	Approx 65 +	Age unknown	Total patients
	<monday date=""></monday>	2	8	12	6		28	10	12	45	15		82
	<tuesday date></tuesday 	4	1	5	4		14	8	10	35	11		64

This is an example of a former health region weekly report to the Ministry of Health.

<health reg<="" th=""><th>ion></th><th></th><th></th><th>Patients</th><th>s with ILI</th><th></th><th></th><th>Total pa</th><th>itients se</th><th>en for al</th><th>l reasons</th><th>s includii</th><th>ng ILI</th></health>	ion>			Patients	s with ILI			Total pa	itients se	en for al	l reasons	s includii	ng ILI
		Pre school	School age	Working age	Seniors		Total ILI	Pre school	School age	Working age	Seniors		Total patients
		Approx 0-4 yr	Approx 5-19	Approx 20-64	Approx 65 +	Age unknown	all age groups	Approx 0-4	Approx 5-19	Approx 20-64	Approx 65 +	Age unknown	all age groups
	ILI Sum	9	22	39	14	0	84	26	49	147	42	0	264
Rate/1000	patients	346.2	449.0	265.3	333.3	0.0	318.2						
Pro	portion ILI	34.6	44.9	26.5	33.3	0.0	31.8						
Emergency room	Date info collected	Approx 0-4	Approx 5-19	Approx 20-64	Approx 65 +	Age unknown	Total ILI	Approx 0-4	Approx 5-19	Apprx 20-64	Approx 65 +	Age unknown	Total patients
St.Joseph's	Jan 25	3	14	20	8	0	45	15	26	102	25	0	168
St. Paul's	Jan 24	2	6	13	2		23	5	10	24	7		46
St. Peter's	Jan 26	4	2	6	4		16	6	13	21	10		50
<name></name>							0						0
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Legionellosis

Date Reviewed: February, 2011 Section: 2-70

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Notification Timeline:

From Lab/Practitioner to Public Health: Within 48 hours. From Public Health to Ministry of Health: Within 3 days.

Immediate if outbreak is suspected or if single nosocomial or occupational case.

Public Health Follow-up Timeline: Initiate within 24 to 48 hours.

Information

Case Definition (Public Health Agency of Canada, May 2008)

Table 1. National Case Definition for Legionellosis			
Confirmed Case	Clinical illness* with laboratory confirmation of infection:		
	• isolation of <i>Legionella</i> species or detection of the antigen from respiratory secretions, lung tissue, pleural fluid or other normally sterile fluids		
	OR		
	• a significant (e.g., fourfold or greater) rise in <i>Legionella</i> species IgG titre between acute and convalescent sera OR		
	• IgG titre > 1:128 against <i>Legionella</i> species		
	OR		
	• demonstration of <i>L. pneumophila</i> antigen in urine		
Probable Case	Clinical illness* with demonstration of <i>Legionella</i> species DNA.		
*Legionellosis comprises two distinct illnesses: Legionnaires' disease, characterized			

^{*}Legionellosis comprises two distinct illnesses: Legionnaires' disease, characterized by fever, myalgia, cough and pneumonia, and Pontiac fever, a milder illness without pneumonia.

Causative Agent

Some species of *Legionella*, a genus of Gram-negative bacilli. Over 35 species have been described, but most cases of legionellosis are caused by *L. pneumophila* serogroup 1.

Symptoms

Legionellosis is an acute bacterial infection and there are two manifestations recognized: Legionnaire's disease and Pontiac fever.



Legionellosis

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Initial Symptoms for both Manifestations	Pontiac Fever ¹	Legionnaire's Disease
 Fever and chills. Temperatures can reach 39°C-40.5°C. Myalgia. Anorexia. Malaise. Headache. Nonproductive cough, abdominal pain and diarrhea may also be present. 	 No pneumonia or multi system involvement. Patients generally recover in two to five days without treatment. 	 Chest x-ray is usually consistent with pneumonia. May progress to multisystem failure with confusion, disorientation, increasing respiratory distress and disseminated legionellosis. Death may occur especially in persons with pre existing medical conditions or a depressed immune system.

Incubation Period

- Legionnaire's disease 2-10 days usually 5-6 days.
- Pontiac fever 5-66 hours, usually 24-48 hours.

Reservoir/Source

The bacterium is ubiquitous in nature and is primarily aquatic. Hot water systems (i.e., showers), air conditioning cooling towers, evaporative condensers, humidifiers, whirlpool spas, respiratory therapy devices and decorative fountains have all been implicated in causing disease and outbreaks.



¹ Believed to be caused by a reaction to inhaled antigen rather than bacterial invasion. Pontiac fever has only been recognized during outbreaks.

Legionellosis

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The bacteria have been isolated from water found in the previously mentioned areas as well as from water in taps, hot tubs, and from creeks, ponds and the soil of their banks. It has been known to survive for months in tap and distilled water. The organism can survive for years in water at 2°C to 8°C and is resistant to usual levels of chlorination (Mandell, 2000).

Mode of Transmission

It is most commonly associated with water-droplet transmission from cooling towers. *Legionella* are transmitted directly from the environment to humans with the most common source thought to be aerosolization of water containing *L. pneumophila*.

Risk Groups/Risk Factors

Illness occurs most frequently with increasing age (most cases are at least 50 years of age), especially in persons who smoke and in those:

- with diabetes mellitus:
- with chronic lung disease;
- that require intubation;
- with renal diseases or;
- with malignancy and;
- who are immunocompromised especially solid organ transplant recipients.

The disease is rare in those under 10 years of age; however, nosocomial infection in neonates has been reported. Several outbreaks have occurred among hospitalized patients. Unrecognized infections are common (Alberta Health and Wellness, 2007).

Period of Communicability

Person to person transmission of these bacteria has not been documented.

Specimen Collection and Transport

- Urine for L. pneumophila serogroup 1 antigen in a sterile specimen container.
- Bronchoalveolar lavage (BAL).
- Urine and BAL should be refrigerated during transport.
- Blood for serology in a plain tube (red top).



Legionellosis

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Methods of Control/Role of Investigator

Prevention and Education

- All devices and equipment that contain, produce, or distribute water or water aerosols must be properly maintained. This may involve draining systems that are not in use, mechanically cleaning systems to remove scale and sediment, and using biocides to limit the growth of *Legionella* (Heymann, 2008).
- Environmental sampling and routine surveillance for this organism is not recommended due to the ubiquitous nature of the organism, the multiplicity of potential sources in the environment, likely recolonization of environmental sources, and the frequency of environmental bacteria in the absence of clinical disease.
- Bacteria can normally be found in the environment with the absence of clinical illness (Alberta Health and Wellness, 2007).

Management

I. Case

History

Source of infection

Inquire about:

- possible exposures to air conditioners, humidifiers, etc., where they work or live;
- presence of other people with similar symptoms to determine if a common source exposure is present.

With the identification of a single laboratory-confirmed case in a health care facility initiate an investigation. This is especially important in facilities serving highly susceptible, immunocompromised patients.

For outbreaks in any other facility, search for:

- common exposures amongst cases;
- common possible environmental source(s) of infection.

Treatment/Supportive Therapy

Cases with Pontiac fever generally recover spontaneously in two to five days without treatment.



Legionellosis

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Antibiotics:

Treatment choices are governed by the most recent guidelines. The public health practitioner should direct any questions regarding the current treatment protocols to the physician or Medical Health Officer. See Appendix H - Sources for Clinical Treatment Guidelines.

Heymann (2008, p. 339) indicates the following:

• "The recommended treatment for Legionnaire's disease is either a respiratory fluoroquinolone, such as levofloxacin, or a newer marcrolide (azithromycin). Observational studies suggest that levofloxacin may be more effective than macrolides, especially in severe cases. Rifampicin has been used as an adjunct in patients failing standard therapy, but data to support this approach are lacking. Penicillin, the cephalosporins and the aminoglycosides are ineffective."

Exclusion

None.

Immunization

Not applicable.

Referrals

Infection control should be notified if the case occurs in a health care facility (acute or long-term).

II. Contacts/Contact Investigation

- Inquire about additional cases in household or school/business setting to determine if a common environmental source exists.
- Quarantine and immunization of contacts are not applicable.

III. Environment

Environmental prevention and control measures (e.g., cleaning, maintenance, decontamination, superheating, superchlorination, etc.) should be referred to the Public Health Inspector Manager/Environmental Health Officer and will likely require engineering expertise.



Legionellosis

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Infection Control Measures

Routine/Standard precautions are recommended.

Epidemic Measures

- In epidemic situation, investigation of common exposures and possible environmental sources of infection is required.
- Decontamination of implicated sources may be necessary and expert advice may be required.
- Culturing from environmental sources should only be considered once cases have been confirmed.



Legionellosis

Date Reviewed: February, 2011 Section: 2-70 Page 7 of 8

References

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Legionellosis Data Collection Worksheet

PANORA

Panorama QA complete:	□Yes	□No
Initials:		

Please complete all sections. Panorama Client ID: Panorama Investigation ID:

A) CLIENT INFORMATION				LHN -> SUBJ	ECT -> CLIE	NT DETAILS ->	PERSONAL INFORMATION
Last Name:		First Name: and M	1iddle Name:		Alternate	Name (Goes l	by):
DOB: YYYY / MM / DD Phone #: Primary Home:	Age:	Health Card Proving Health Card Numb			home pho	one, text):	ion Method: (specify - i.e.
□ workplace.							
Place of Employment/School:		Gender: □ Mal	e	□ Female		Other	□ Unknown
Alternate Contact: Relationship: Alt. Contact phone:		Address Type: No fixed Postal ad Postal ad Street Address or Address at time of	dress): FN Community	y (Primary Hom		oorary □ Leg	gal Land Description
B) INVESTIGATION INFORMATION		LHN->	SUBJECT SUN	IMARY-> ENTE	RIC ENCOL	INTER GROUP	->CREATE INVESTIGATION
Disease Summary Classification: CASE	Date	Classification: CONTACT		Date	:		FORMATION: nen collected:
□ Confirmed	YYYY / MM / DD	□ Contact		YYYY / MM	/ DD	YYYY / MM	/ DD
□ Does Not Meet Case Definition	YYYY / MM / DD	□ Not a Contact		YYYY / MM	/ DD	Specimen ty	
☐ Person Under Investigation	YYYY / MM / DD	☐ Person Under Ir	nvestigation	YYYY / MM	/ DD	□ Blood	2
□Probable	YYYY / MM / DD					□ Respi	iratory Secretions
Disposition: FOLLOW UP: ☐ In progress ☐ Incomplete - Declined ☐ Incomplete — Lost contact ☐ Incomplete — Unable to locate	YYYY / MM / DD YYYY / MM / DD YYYY / MM / DD YYYY / MM / DD		Complete Not required Referred – Out ecify where)	t of province	YYYY / N	MM / DD MM / DD MM / DD	
REPORTING NOTIFICATION			Location:				
Name of Attending Physician or Nu	irse:						
Physician/Nurse Phone number:			Date Received	(Public Health): YYYY ,	/ MM / DD	
Type of Reporting Source: ☐ Hea	alth Care Facility □L	ab Report	Nurse Practitic	oner □Phy	sician	□ Other	

DISEASE EVENT HISTORY LHN->INVESTIGATION->DISEASE SUMMARY (UPDATE)->DISEASE EVENT HISTORY Site Description: \square Legionnaires' disease $\hfill\Box$ Pontiac fever \square Other

Legionellosis Data Collection Worksheet

Please complete **all** sections

D) SIGNS & SYMPTOMS (Bold text = part of case definition)

Investigator name

Panorama Client ID:	
Panorama Investigation ID:	

LHN-> INVESTIGATION->SIGNS & SYMPTOMS

Description	NO	res – i	Date of onset	Description	on	NO	Yes - Date of onset
Loss of appetite (anorexia)		YYYY / I	MMM / DD	Headache			YYYY / MMM / DD
Chills		YYYY / I	MMM / DD	Malaise			YYYY / MMM / DD
Confusion				Myalgia (muscle pain)		
Cough		YYYY / I	MMM / DD	Pain - abd	lominal		YYYY / MMM / DD
Diarrhea		YYYY / I	MMM / DD	Pneumon	ia		YYYY / MMM / DD
Fever		YYYY / I	MMM / DD	Respirato	ry distress	+	YYYY / MMM / DD
E) INCUBATION AND COMMUNICABILITY					LHN-> INVESTIGATION-	->INCUB/	ATION & COMMUNICABILITY
Incubation for Case(period for acquisition):					Latert Bassilla Farrage Bata	000/ / 5	40.4 / DD
Earliest Possible Exposure Date: YYYY / MN	VI / L	טט			Latest Possible Exposure Date:	YYY / IV	לוט / טט
Exposure Calculation details:							
,							
F) RISK FACTORS N—No, NA–Not asked,	11_11	Inknow	n			111	N-> SUBJECT->RISK FACTORS
•	0-0	IIKIIOW	Yes		Add'l Info	LITT	N-> 30BJECT->RISK FACTORS
DESCRIPTION			Start date	N, NA, U			
Chronic Medical Condition - Malignancies/Co	ancer	+					
Immunocompromised - Related to underlyin	ng dise	ease					
or treatment Immunocompromised - Transplant Candidat	e or						
Recipient - Solid Organ/Tissue+							
Travel - Outside of within Canada (Add'l Info)		YYYY / MM/DD AE				
Travel - Outside of Saskatchewan, but within	Cana	ida	YYYY / MM/DD				
(add'l info) Water - Aerosol - Air conditioning unit			AE YYYY / MM/DD				
Water - Aerosol - Other (add'l info)			YYYY / MM/DD				
			YYYY / MM/DD				
Water - Aerosol - Room/central humidifier							
Water - Aerosol - Shower head			YYYY / MM/DD				
G) TREATMENT					LUNE SINVECTION SINVERTION SINVERTION	ICATIONI	C - BAEDICATIONIC CLIBABA A DV
G) TREATMENT					LHN-> INVESTIGATION-> MEDI	CATIONS	5->IVIEDICATIONS SUIVIIVIARY
Medication (Panorama = Other Meds) :							
Prescribed by:				Sta	rted on: YYYY / MMM / DD		
Tresensed by:				5ta	rted on.		
H) INTERVENTION			ı	.HN-> INVE	STIGATION->TREATMENT & INTERVEN	NTIONS->	INTERVENTION SUMMARY
Intervention Type and Sub Type:							
Assessment: Investigator name				Immi	unization: Investigator name		
☐ Assessed for contacts		YY	YY / MM / DD		gible immunizations recommended		YYYY / MM / DD
(individuals exposed to same source) Communication:				Defe	l		
☐ Other communication (See Investigator N	otes)	YY	YY / MM / DD	Refe	ection Prevention and Control		YYYY / MM / DD
Investigator name	/			Inves	tigator name		
☐ Letter (See Document Management) Investigator name		YY	YY / MM / DD		nsultation with MHO		YYYY / MM / DD
General: Investigator name					tigator name r Investigation Findings:		
☐ Disease-Info/Prev-Control		YY	YY/ MM / DD		vestigator Notes		
☐ Disease-Info/Prev-Cont/Assess'd for Conta	acts		YY/ MM / DD		cument Management Notes		
Education/counselling:							
☐ Prevention/Control measures		YY	YY / MM / DD				
☐ Disease information provided		YY	YY / MM / DD				

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Legionellosis Data Collection Worksheet

Please complete **all** sections

Panorama Client ID:	
Panorama Investigation ID:	

Date	Intervention subtype	Comments	Next follow-up Date	Initials
YYYY / MM / DD			YYYY / MM / DD	
YYYY / MM / DD			YYYY / MM / DD	
YYYY / MM / DD			YYYY / MM / DD	
YYYY / MM / DD			YYYY / MM / DD	
YYYY / MM / DD			YYYY / MM / DD	
YYYY / MM / DD			YYYY / MM / DD	
I) OUTCOMES (option	onal except for severe i	nfluenza)	LHN-> INVESTIGATIO	DN-> OUTCOMES
·	//recovering YYYY / N	•	italization YYYY / MN	-
	YYYY / N		YYYY / MIV	I / DD
□ Fatal	YYYY / N	M / DD Unknown		
Cause of Death: (if Fa	atal was selected)			
J) EXPOSURES Acquisition Event Acquisition Event ID:		LHN-> INVESTIGATION-> EXPOSURE SU	MMARY-> ACQUISITIO	ON QUICK ENTRY
Acquisition Start YYY	Y / MM / DD to	Acquisition End: YYYY / MM / DD		
Location Name:				
Setting Type	_	_		
□ Travel	☐ Exposure or consur	nption of potentially contaminated food or water	ly source	
Initial Report completed by:			Date initial report of	•
			TITT / IVIIVIIVI / DD	,

April 30, 2019 Page 3 of 3

Leprosy (Hansen's Disease)

Date Reviewed: February, 2011 Section: 2-80 Page 1 of 5

Notification Timeline:

From Lab/Practitioner to Public Health: Within 48 hours. From Public Health to Ministry of Health: Within 2 weeks. Public Health Follow-up Timeline: Initiate within 72 hours.

Information

Case Definition (Public Health Agency of Canada, May 2008)

Table 1. National	Surveillance Case Definition for Leprosy (Hansen's Disease)
Confirmed Case	Clinical evidence of illness (see symptoms) with laboratory
	confirmation:
	positive acid fast stain with typical morphology for
	Mycobacterium leprae
	OR
	histopathological report from skin or nerve biopsy compatible
	with leprosy
Probable Case	Clinical illness (see symptoms) in a person who is
	epidemiologically linked to a confirmed case

Causative Agent

Mycobacterium leprae.

Symptoms (Public Health Agency of Canada, May 2008)

<u>Tuberculoid or paucibacillary disease</u>: one or a few well-demarcated, hypopigmented and anesthetic skin lesions, frequently with active, spreading edges and a clearing centre; peripheral nerve swelling or thickening may also occur.

<u>Lepromatous or multibacillary disease</u>: erythematous papules and nodules or an infiltration of the face, hands and feet with lesions in a bilateral and symmetrical distribution that progress to thickening of the skin and loss of normal hair distribution, particularly on the face (madarosis).

<u>Borderline (dimorphous)</u>: skin lesions characteristic of both the tuberculoid and lepromatous forms.

<u>Indeterminate:</u> early lesions, usually hypopigmented macules, without developed tuberculoid or lepromatous features.



Leprosy (Hansen's Disease)

Date Reviewed: February, 2011 Section: 2-80 Page 2 of 5

Incubation Period

9 months to 20 years. Tuberculoid is an average of 4 years, and 8 years for the lepromatous form. Rarely seen in children under 3 years.

Reservoir/Source

Humans are the reservoir of proven significance however it has been shown that the armadillo, mangabey monkey and chimpanzee can be infected.

Mode of Transmission

Transmission is person to person with nasal secretions, normally containing the highest bacterial load, often causing infection when spread to the skin or respiratory tract of another. Close contact is necessary for transmission. Untreated multibacillary leprosy (high levels of bacillus) has been proven to be the major source of human transmission.

Risk Groups/Risk Factors

- Leprosy is a disease of poverty.
- Approximately 95% of people are genetically immune to infection with *M. leprae*.
- HIV clients are not at increased risk of becoming infected.

Period of Communicability

Clinical and laboratory evidence suggest that infectiousness is lost in most instances within a day of beginning treatment with multidrug therapy (Heymann, 2008).

Specimen Collection and Transport

For specimen collection instructions, consult with Saskatchewan Disease Control Laboratory (SDCL) Medical Director at (306) 787-8636.

Methods of Control/Role of Investigator

Prevention and Education

Refer to the <u>Respiratory and Direct Contact Introduction and General Considerations</u> section of the manual that highlights topics for client education that should be considered as well as provides information on high-risk groups and activities.



Leprosy (Hansen's Disease)

Date Reviewed: February, 2011 Section: 2-80 Page 3 of 5

- The best preventive measure is early diagnosis and treatment of cases.
- Health education should stress the importance of effective multi-drug therapy, the non-infectivity of persons under continuous treatment and the importance of completing therapy.

Management

I. Case

History

- No public health interventions are required; communicability is low, particularly after initiation of treatment.
- Persons with leprosy require medical follow-up from an infectious diseases specialist.
- Manage infectious persons with routine infection control precautions. Handwashing is the most effective measure to prevent transmission when caring for patients.
- Hospitalization is reserved only for managing reactions, surgical correction of deformities and the treatment of ulcers resulting from the anesthesia of the extremities.

<u>Treatment/Supportive Therapy</u>

- Consultation with an infectious disease specialist, internist, dermatologist or pediatrician is recommended. See Appendix H - Sources for Clinical Treatment Guidelines.
 - Multi-drug chemotherapy is necessary for all patients. There is widespread prevalence of dapsone resistance, and the emerging resistance to rifampin.

Exclusion:

No restrictions in employment or attendance at school are indicated for persons whose disease is regarded as non-infectious.

II. Contacts/Contact Investigation

Household and other close contacts should be examined initially, and then annually for at least 5 years. Consult specialist.



Leprosy (Hansen's Disease)

Date Reviewed: February, 2011 Section: 2-80 Page 4 of 5

- Manage infectious persons with routine infection control precautions.
 Handwashing is the most effective measure to prevent transmission when caring for patients.
- Chemoprophylaxis is not recommended.

III. Environment

Isolation of cases and quarantine of individuals is not necessary and often leads to stigmatization. No restrictions for employment or school are indicated.

Epidemic Measures

Not applicable.



Leprosy (Hansen's Disease)

Date Reviewed: February, 2011 Section: 2-80 Page 5 of 5

References

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Leprosy Data Collection Worksheet



Please complete all sections.

Panorama QA complete: ☐ Yes Initials:	□No			Pan	Panorama Client ID:
A) CLIENT INFORMATION			LHN -> SUBJE	CT -> CLIEN	T DETAILS -> PERSONAL INFORMATION
Last Name:		First Name: and Middle	Name:	Alternate	Name (Goes by):
DOB: YYYY / MM / DD Phone #: Primary Home: Mobile contact:	Age:	Health Card Province: _ Health Card Number (P		i.e. home	Communication Method: (specify - phone, text): dress: Work Personal
☐ Workplace:					
Place of Employment/School:		Gender: Male	□ Female		Other
Alternate Contact:		Address Type: ☐ No fixed ☐ Postal A Mailing (Postal address		me □Tem _l	porary □ Legal Land Description
Relationship:		Street Address or FN Co	ommunity (Primary Hor	ne):	
Alt. Contact phone:		Address at time of infec	ction if not the same:		
B) IMMIGRATION INFORMATION		SUBJECT -> CLIE	NT DETAILS -> PERSON	AL INFORM	ATION->IMMIGRATION INFORMATION
Country Born in: Country Emigrated from:		Arrival Date: YYYY / 1	MMM / DD O	R Arri	val Year
C) INVESTIGATION INFORMATION		LHN-> SUBJECT SUI	MMARY-> ZOONOTIC 8	& VECTORBO	ORNE GROUP->CREATE INVESTIGATION
Disease Summary Classification: CASE	Date	Classification: CONTACT	Date	?	LAB TEST INFORMATION: Date specimen collected:
□ Confirmed	YYYY / MM / DD	□ Contact	YYYY / MM	/ DD	YYYY / MM / DD
□ Does Not Meet Case	YYYY / MM / DD	□ Not a Contact	YYYY / MM	/ DD	
☐ Person Under Investigation	YYYY / MM / DD	☐ Person Under Investigation	YYYY / MM	/ DD	
□ Probable	YYYY / MM / DD				
Disposition: FOLLOW UP: ☐ In progress ☐ Incomplete - Declined ☐ Incomplete — Lost contact ☐ Incomplete — Unable to locate	YYYY / MM / DD YYYY / MM / DD YYYY / MM / DD YYYY / MM / DD	☐ Comp ☐ Not ri ☐ Refer (specify	equired red – Out of province	YYYY / I	MM / DD MM / DD MM / DD
REPORTING NOTIFICATION Name of Attending Physician or Nu	ırse:	Locat	ion:		
Physician/Nurse Phone number:		Date	Received (Public Health	n): YYYY	/ MM / DD
Type of Reporting Source: ☐ Hea	alth Care Facility 🗆 🗅 La	ab Report □ Nurse	e Practitioner	ysician	□ Other

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Leprosy Data Collection Worksheet

Please complete **all** sections

 Panorama Client ID:
anorama Investigation ID:

	. □Tuł	ວerculoid	□Borderline	, 🗆	Other 🗆 Unknown				
) SIGNS & SYMPTOMS						-> INVEST	TIGATION->	SIGNS 8	& SYMPT
Description	No		te of onset	Des	cription	No	Yes - Date		
Alopecia (loss of normal hair distribution)			MMM / DD	Rash	h - papules - erythematous		YYYY / N	_	
Bleeding - nose (epistaxis)			MMM / DD	Skin	- infiltrative disorders		YYYY / N		
Iritis (inflammation of the iris)	\top _	YYYY / N	MMM / DD		- lesions - hypopigmented and esthetic (painless)		YYYY / N	/IMM /	DD
Keratitis (inflammation of the cornea)	T	YYYY / N	MMM / DD	Skin	nodules		YYYY / N	/MM	DD
Neurologic - peripheral nerve - swelling or thickening (neuritis)				Skin	- thickening				
Neuropathy	T_{-}	YYYY / N	MMM / DD	Skin	- nodules - erythematous	<u> </u>	YYYY / N	/MM	DD
Rash - macules - hypopigmented	\top	YYYY / N	MMM / DD	†	11000102 2. j	\dashv	YYYY / N	/MM	DD
.) RISK FACTORS (during risk period) DESCRIPTION	YES		N – No		DESCRIPTION	YES	LHN-> SUB.	JECT->R	
DESCRIPTION	125		NA – not aske U - Unknown	ed	DESCRIP HON	120		NA – n	not asked known
Contact - Visitor from an endemic country	,	/ MM / DD	U - Ufikilowii	,	Travel - Outside of Canada (Add'l Info)	AE	/ MM / DD	U - on	knowii
Contact to a known case (Add'l Info)	YYYY /	/ MM / DD			Travel - Outside of Saskatchewan, but within Canada (Add'l Info)	YYYY ,	/ MM / DD		
Special Population - From or residence in an endemic country (Add'l Info)	YYYY /	/ MM / DD							
) MEDICATIONS					INVESTIGATION-> M	1EDICATIO	ONS->MEDI	CATION	IS SUMN
Medication (Panorama = Other Meds)	:								
Prescribed by:					Started on: YYYY / MMM / DD				
) INTERVENTIONS)NS->INTER\	VENTIO	N SUMI
) INTERVENTIONS Intervention Type and Sub Type:					Started on: YYYY / MMM / DD)NS->INTER\	VENTIO	N SUMI
) INTERVENTIONS		YY	YY / MM / [LHN	Started on: YYYY / MMM / DD	E RVENTIO		Y / MM	
Interventions Intervention Type and Sub Type: Assessment: Assessed for contacts Investigator name Communication: Other communication (See Investig	ator Note		YY / MM / [LHN	Started on: YYYY / MMM / DD N-> INVESTIGATION->TREATMENT & INTE Education/counseling: Investigator na Prevention/Control measures Disease information provided	ame YYY'	YYYYY Y / MM /	Y / MM	1 / DD
Interventions Intervention Type and Sub Type: Assessment: Assessed for contacts Investigator name Communication: Other communication (See Investig Investigator name Letter (See Document Management Investigator name		es) YY		LHN DD	Started on: YYYY / MMM / DD N-> INVESTIGATION->TREATMENT & INTE Education/counseling: Investigator na	ame YYY'	YYYYY Y / MM /	Y / MM DD	1 / DD
Intervention Type and Sub Type: Assessment: Assessed for contacts Investigator name Communication: Other communication (See Investig Investigator name Letter (See Document Management Investigator name General: Investigator name		es) YY'	'YY / MM / [LHIN DD DD	Started on: YYYY / MMM / DD N-> INVESTIGATION->TREATMENT & INTE Education/counseling: Investigator na	ame YYY' iigator na	YYYYYY / MM / Imme	Y / MIM DD Y / MIM	1 / DD
Intervention Type and Sub Type: Assessment: Assessed for contacts Investigator name Communication: Other communication (See Investig Investigator name Letter (See Document Management Investigator name General: Investigator name Disease-Info/Prev-Control	t)	es) YYY	YY / MM / [YY / MM / [LHIN DD DD	Started on: YYYY / MMM / DD N-> INVESTIGATION->TREATMENT & INTE Education/counseling: Investigator na	ame YYY' iigator na	YYYYY Y / MM /	Y / MIM DD Y / MIM	1 / DD
Intervention Type and Sub Type: Assessment:	r Contact	es) YYY	'YY / MM / [LHIN DD DD	Started on: YYYY / MMM / DD N-> INVESTIGATION->TREATMENT & INTE Education/counseling: Investigator na	ame YYYY iigator na d	YYYYYY / MM / Imme	y / MM DD y / MM	1 / DD
Intervention Type and Sub Type: Assessment: Assessed for contacts Investigator name Communication: Other communication (See Investig Investigator name Letter (See Document Management Investigator name General: Investigator name Disease-Info/Prev-Control Disease-Info/Prev-Cont/Assess'd for	r Contact	es) YYY YYY tts YYY	YY / MM / [YY / MM / [LHIN DD DD	Started on: YYYY / MMM / DD N-> INVESTIGATION->TREATMENT & INTE Education/counseling: Investigator na	ame YYYY iigator na d	YYYYY Y / MM / Ime YYYYY Document M	y / MM DD y / MM	1 / DD

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YYYY / MM / DD

Leprosy Data Collection Worksheet

Please complete **all** sections

Panorama Client ID:	
Panorama Investigation ID:	

YYYY / MM / DD								
YYYY / MM / DD	_							
YYYY / MM / DD	_							
YYYY / MM / DD								
YYYY / MM / DD								
YYYY / MM / DD								
YYYY / MM / DD								
YYYY / MM / DD								
1000/ / BABA / DD								
YYYY / MM / DD								
YYYY / MM / DD								
D) OUTCOMES (opti			□ ICII/intensive medi	ral care VVVV / MM / DD	Пнос		/ESTIGATION-	
D) OUTCOMES (opti	ional except for severe /recovering YYYY / N	MM / DD		cal care YYYY / MM / DD		pitalization	/ESTIGATION- YYYY / MM YYYY / MM	/ DD
D) OUTCOMES (opti	/recovering YYYY / N	MM / DD MM / DD	☐ Intubation /ventilati		□ Unkr	pitalization	YYYY / MM	/ DD
D) OUTCOMES (option of the property of the pro	/recovering YYYY / N	MM / DD MM / DD MM / DD	☐ Intubation /ventilati	on YYYY / MM / DD	□ Unkr	pitalization	YYYY / MM	/ DD

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Notification Timeline:

From Public Health to Ministry of Health: Immediate.

From Public Health to Ministry of Health: Immediate.

Public Health Follow-up Timeline: Immediate.

Public Health Purpose for Notification of Measles

- To prevent transmission of measles from imported cases and further local transmission;
- To prevent mortality and serious morbidity from measles through contact tracing;
- To track epidemiology trends of measles in Saskatchewan including risk factors and distribution;
- To monitor the effectiveness of prevention and control measures;
- To make timely and evidence informed actions on outbreaks; and
- To inform the public and medical community about measles.

Surveillance Case Definition¹ (Public Health Agency of Canada, 2013)

Commined
(Public
Health
Agency of
Canada,
2013)

Confirmed Case Laboratory confirmation of infection in the absence of recent immunization ^a with measles-containing vaccine:

- isolation of measles virus from an appropriate clinical specimen ^b
- detection of measles virus ribonucleic acid (RNA) (e.g. PCR) ^c
- seroconversion or a significant (e.g., fourfold or greater) rise in measles immunoglobulin G (IgG) titre by any standard serologic assay between acute and convalescent sera

OF

 positive serologic test for measles immunoglobulin M (IgM) antibody using a recommended assay^d in a person who is either epidemiologically linked to a laboratory-confirmed case or has recently travelled to an area of known measles activity.

OR

Clinical illness in a person with an epidemiologic link to a laboratory-confirmed case.

¹ Surveillance case definitions ensure uniform reporting to allow comparability of surveillance data. The definition is not intended to be used for clinical or laboratory diagnosis or management of cases.



Probable Case	Clinical illness
(Public Health	in the absence of appropriate laboratory tests
Agency of	OR
Canada, 2013)	in the absence of an epidemiologic link to a laboratory-confirmed case
,	OR
	• in a person who has recently travelled to an area of known measles activity.
Clinical Case	Clinical illness is characterized by all of the following features:
(Public Health	• fever of 38.3° C or greater;
Agency of	cough, coryza or conjunctivitis;
Canada, 2013)	generalized maculopapular rash for at least 3 days.

^a The most frequent reaction to measles-mumps-rubella (MMR) immunization is malaise and fever (with or without rash) occurring 6-23 days after immunization. However, this should be determined for each case, as these reactions and the timeframe can vary (Pubic Health Agency of Canada, 2015).

Most acute measles cases develop IgM after 3 days post rash onset. Therefore, a suspected measles case in which serum collected \leq 3 days after rash onset initially tests IgM negative should have a second serum specimen collected > 3 days after onset for retesting for IgM.

Further strain characterization is indicated for epidemiologic, public health and control purposes.

Refer to <u>Specimen Collection and Transport</u> for further details about interpretation of lab results in the context of past immunization for measles.

Epidemiology and Occurrence

Measles became reportable in Canada in 1924 (PHAC). Prior to the development of a vaccine (1924 to 1958), an average of 45,000 cases were reported annually. Through the use of vaccines, Canada eliminated measles in 1998 (PHAC, 2013), however sporadic cases and outbreaks continue as a result of importations (PHAC, 2018). To achieve herd immunity, the recommended 2-dose immunization coverage rate is ≥ 95% (PHAC, 2014). While Canada's overall coverage is high, pockets of susceptible individuals and communities remain so the risk of domestic transmission following an importation of measles remains a reality.

Saskatchewan UNDER CONTRUCTION



^b See Specimen Collection and Transport

^c Confirmation of genotype is required in recently vaccinated individuals (within the past 6-45 days) to determine if illness is related to wild virus or vaccine-related.

^d IgM serology has the potential for false-positive findings. If the clinical presentation is inconsistent with a diagnosis of measles or in the absence of recent travel/exposure history, IgM results must be confirmed by the other listed confirmatory methods.

Table 1. Evolution of the Measles Immunization Program in Saskatchewan

1966	Measles vaccine introduced for ages 1-3 (Lirugen - live, further attenuated)		
1970	Measles vaccine extended to ages 1-7 (ATTENUVAX - live, further		
	attenuated)		
1970	Rubella vaccine for grade 7 girls. (MERUVAX and MERUVAX-II); Cendevax		
	(rubella vaccine) used ~1970-72		
1970-1975	Rubella vaccine for grade 1 students (MERUVAX)		
1971	Rubella vaccine available to physicians for susceptible women at premarital		
	exams		
1979	MMR vaccine for age 1 year		
1981 - 1982	Review of measles immunization for children ages 1-14, followed by		
	program to raise coverage to > 98%		
Fall 1991 to	Mass MMR immunization for teen-aged boys in high schools and post-		
1992	secondary institutions		
Fall 1996	Second dose Measles & Rubella (MR) added to 18 months.		
	Catch-up program included school entry, Grade 6 and 8.		
Spring 1997	MR immunization of Grades 9-12		
2001	MMR used exclusively for all 1st and 2nd doses; MR discontinued by Berna		
2003 - 2004	2 dose mumps catch-up in Grade 6		
2007 - 2013	2-dose mumps catch-up for eligible Grade 12 students		
2008 - 2013	2-dose mumps catch-up for eligible Grade 8 students		
2011 - 2013	2nd dose provided to eligible Grade 6 students		
May 2013	Adult born since Jan. 1, 1970 eligible for 2 MMR doses		

Saskatchewan Immunization Manual (2018)

The Roy Romanow Provincial Laboratory conducted a review of measles immunity in February 2014 to inform risk populations. Based on this review, approximately 93% of samples tested from individuals born prior to 1965 indicated immunity for measles, while approximately 83% of samples from persons born between 1965 and 1980 indicated immunity to measles. Factors to consider include age, history of exposure to measles, and availability of medical records.

Additional Background Information

Causative Agent

Measles virus, an RNA virus, a member of the family paramyxovirus, genus *Morbillivirus*.



Symptoms

Measles is an acute, highly communicable disease with a prodrome that lasts two to four days (range one to seven days). The prodrome is characterized by fever followed by conjunctivitis, coryza, or cough.

- Koplik spots on the buccal mucosa are considered pathognomic for measles. They occur one to two days before the rash.
- A characteristic red maculo-papular rash appears on the third to seventh day beginning behind the ears and on the face. The rash gradually spreads downwards to the trunk and then the extremities. The skin lesions are usually discrete but may become confluent.
- Fever often rises as the rash appears. The rash may last four to seven days and often fades in the same sequence as it appears.
- Symptoms are more severe in infants and they are more likely to experience complications.
- Immunocompromised individuals experience more severe disease and may have a prolonged course. These individuals may not develop the characteristic rash.
- Other symptoms of measles include anorexia, diarrhea (especially in infants), and generalized lymphadenopathy.
- Individuals who have been previously exposed to measles antigen (e.g., previously vaccinated), may have a modified clinical presentation (Centers for Disease Control and Prevention, 2018).

Complications (Heymann, 2015)

- Diarrhea (8%), otitis media (7%), pneumonia (6%), seizures (0.7%), encephalitis (0.1%), and laryngotracheobronchitis (croup).
- Very rarely, sub-acute sclerosing panencephalitis (SSPE) develops 7-10 years after infection as a late sequelae (Centers for Disease Control and Prevention, 2018).
- The case-fatality rate can be as high as 10-30% in developing countries (typically 3-5%); it is approximately 0.1-0.2% in Canada.

Reservoir

Humans.

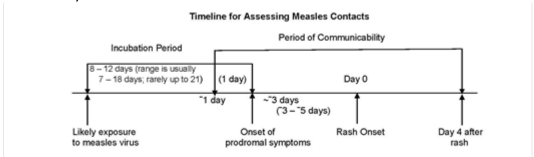
Incubation Period

- About 10 days (range 7 to 18 days) from exposure to onset of fever.
- Usually 14 days until rash appears (range 9 to 21 days).



Period of Communicability

- Measles is highly communicable, with greater than 90% secondary attack rates among susceptible persons.
- Cases are infectious from one day before onset of prodrome, usually about four days before onset of the rash, and continue until four days after rash onset (Heyman, 2015).
- Maximum communicability occurs from onset of prodrome through the first three to four days of rash.



(Adapted from BCCDC, 2014)

Mode of Transmission

- Large respiratory droplets.
- Airborne transmission via droplet nuclei has been documented.
- Direct person-to-person contact with the nasal or throat secretions of the infected person.
- Indirect contact with articles freshly soiled with the respiratory secretions.

Risk Factors

Risk factors are associated with individual susceptibility and settings that create opportunities for acquisition or transmission to others.

- Non-immune individuals.
- Immunocompromised individuals.
- Infants.
- Children in childcare settings.
- Child care workers.
- Health care workers (HCWs).
- Students at post-secondary institutions.
- Travellers.
- Military personnel.



• Infection during pregnancy is associated with an increased frequency of spontaneous abortion, premature labor and preterm birth and low birth weight.

Specimen Collection and Transport

Specimen collection needs to be done in coordination and consultation with public health and infection control to ensure it is completed in a that will reduce further transmission (e.g. in home collection by immune employee or arrangements with the lab for end of day collection).

When sending specimens for measles testing, laboratory requisitions should be clearly marked "suspect case of measles" to facilitate rapid testing. When an outbreak number has been assigned, the outbreak number should be included on the requisition and the transport tote should be marked that outbreak specimens are included.

Molecular isolation/detection² of the virus is preferred to confirm the diagnosis of suspected measles cases because of the complications in interpreting positive IgM serology in the absence of an epidemiological link to a confirmed case and the contradictory serological results in previously immunized individuals (Public Health Agency of Canada, 2013). There is no single laboratory test capable of confirming with confidence 100% of true measles cases. Therefore, to confirm the diagnosis the following specimens should be submitted to Roy Romanow Provincial Laboratory (RRPL):

- Urine, throat and nasopharyngeal secretions for isolation of measles virus (i.e. PCR):
 - Collect nasopharyngeal swab or aspirate, or a throat swab³ as soon as possible after the onset of the rash (within four to seven⁴ days). Place in viral transport medium.
 - Collect approximately 50 ml of urine within seven days after the onset of rash.
- Serum sample for measles IgM and IgG (acute and convalescent):
 - o IgM response begins with onset of rash and will persist for one to two months.
 - IgG response begins about one week after the onset of rash and will persist for a lifetime.
 - Convalescent sera should be drawn 10 to 30 days after the initial serology to assess the rise in IgG titre (seroconversion).

⁴ Measles virus may be still detected after seven days from the onset of rash, but with rapidly decreasing sensitivity.



² Isolation permits measles virus genotyping which provides confirmation of epidemiologic data showing measles virus transmission routes and the differentiation of wild-type from vaccine strain measles virus in cases where vaccine may be implicated in serious illness.

³ Nasopharyngeal and throat swabs must be collected in physicians' office.

Negative results do not definitively rule out measles because both methods are affected by timing of specimen collection and quality of handling.

Treatment/Supportive Therapy

- There is no specific treatment available for measles.
- Supportive therapy as indicated.
- Vitamin A supplementation of children with measles has been associated with decreased morbidity and mortality rates. The World Health Organization (WHO) currently recommends vitamin A for all children with acute measles, regardless of their country of residence. Vitamin A for measles is administered once daily for two days at the following doses:
 - o 200 000 IU for children 12 months of age or older;
 - o 100 000 IU for children six through 11 months of age;
 - 50 000 IU for infants younger than six months of age (American Academy of Pediatrics, 2015).

Public Health Investigation

I. Single Case/Household Cluster

 All reports of probable and laboratory-confirmed measles cases should be investigated immediately. Refer to <u>Attachment – Measles Data Collection Worksheet</u> to assist.

History

- Determine measles immunization history including number of doses, date(s) administered,⁵ and type of vaccine.
- Determine if there is an opportunity for <u>acquisition</u> through:
 - In the 7-21 days before the onset of rash, there was a history of travel or contact with a person who had recent travel.
 - o contact with a confirmed or probable case of measles.
- Health conditions that may render the individual more susceptible to infection or alter the period of communicability (e.g. immunocompromised).

ii. they are epidemiologically linked to a laboratory-confirmed case(Centers for Disease Control and Prevention, 2013).



⁵ Fever and rash may occur in the 6-23 days following administration of a measles-containing vaccine. The inability to rely on positive IgM serology when vaccine was administered in the 6-45 days prior to onset of rash can create challenges in confirming a diagnosis of measles. In this case, specimens for viral isolation should also be obtained (see Specimen Collection and Transport section above); if wild type measles virus is isolated, the case can be confirmed. When strain typing to confirm wild type virus is not available in these situations, the case should only be confirmed if:

i. they meet the clinical case definition, and

- Identify opportunities for <u>transmission</u> events and contacts exposed during the infectious period, which includes four days prior to and four days after the rash appears:
 - household;
 - daycare/school;
 - workplaces;
 - health care facilities⁶ (including physicians' offices and waiting rooms).
- Identify locations, dates, times and details of any event the case has attended during the infectious period. This includes gatherings of all sizes in both public and private forums such as:
 - social or religious functions;
 - sports activities;
 - shopping excursions;
 - concerts;
 - conferences and meetings.
- Identify routes, dates, times and details of public transportation (flights, buses, taxis, etc.).
 - Obtain details about the public transportation involved (e.g., company of carrier, seating information, depots/terminals/gates involved, etc.).

Public Health Interventions

Assessment

Assess for contacts paying particular attention to vulnerable contacts as per Table 3.

Communication

 Letters can be sent to other group settings where individual contact tracing is not required (i.e. in the same workplace, but do not share the same work schedule or location of work) to inform them of the exposure, symptom monitoring and when to seek medical attention (see Sample letter).

Education

 All cases should be provided disease information as well as information on prevention and control measures including period of communicability, to self-isolate at home (no visitors).

Exclusion and Isolation

Exclusion and isolation of cases should be implemented as outlined in Table 2.

⁶ In acute care settings, Infection Control and Occupational/ Employee Health should also be involved.



Table 2. Exclusion Requirement for Confirmed, Probable, Clinical Cases and Persons Under Investigation for Measles

Context	Exclusion Requirement	Timeframe
Community Settings.	Self-isolation at home.	
	Exclude from daycare, schools, and workplaces.	Immediately and up to and including four days after
Avoid exposing non- household contacts (i.e. no outside visitors) onset of rasl		onset of rash.
Hospitalized Settings ⁷ 1. Immunocompetent patients.	Airborne precautions.	Immediately and up to and including four days after onset of rash (Public Health Agency of Canada, 2013).
2. Immuncompromised patients.	Airborne precautions.	Immediately and up to and including four days after onset of rash, or for the duration of illness because viral excretion is expected to be prolonged ⁸ (Public Health Agency of Canada, 2013). Consult with Medical Microbiologist in charge of Infection Control and/or ID Specialist for an individual assessment

Immunization

 Review the immunization history. Consider potential issues with vaccines administered for cases who are fully immunized. Immunization of case is not indicated, but may be provided to offer protection against other vaccine antigens (e.g. mumps or rubella) if eligible.

Referrals

Not applicable.

⁸ An immunocompromised person may shed virus for several weeks after the acute illness (CDC, 2015)



⁷ Refer to <u>Health Care Facility Control Measures</u> for further details and additional measures to be taken with cases.

II. Contacts/Contact Investigation

Identification of contacts and contact investigation should proceed immediately and should be re-evaluated once laboratory results are available. Contact Investigation Worksheet should be used to support investigation.

Contacts should be prioritized based on individual and public health risk including:

- high risk contacts;
- employees in health care settings (direct and indirect patient care staff);
- other susceptible contacts;
- public exposures.

Table 3. Contact Definitions (Adapted from Public Health Agency of Canada, 2013)

A. Contact

A contact is defined as any individual who has:

- spent any length of time in a room or enclosed space with a measles case during that case's infectious period (i.e., from one day before onset of prodrome, usually about four days before onset of the rash, and continue until four days after rash onset); or
- spent time in the same room as in infectious case of measles or in a room that the case vacated in the previous two hours. 9

Individualized (person-by-person) contact investigation should include:

- 1. household contacts;
- 2. in a daycare/educational facility all employees, volunteers, students, bus drivers, members of a sports team or club;
- 3. in a workplace individuals who share the same schedule or office location as the case;
- 4. in a health care facility individuals who shared the same room, waiting room or exam room and did not use appropriate protection (i.e., N95 respirator). NOTE: This is not limited to patients in these settings but includes anyone attending appointments with the patient.

B. High Risk Contacts

- Infants <1 year of age.
- Pregnant women.
- Immunocompromised individuals.

C. Susceptible Contacts

Employees in health care and daycare settings are considered susceptible if they have:

- NO laboratory evidence of immunity, AND
- NO documented evidence of two doses of measles-containing vaccine (given at the appropriate interval as outlined in the Saskatchewan Immunization Manual for vaccine type [MMR or MMRV]).

Non-health care/daycare workers 10, may be susceptible if they have:

¹⁰ Generally, individuals born before 1970 are considered immune. During outbreak situations and in consultation with the Medical Health Officer, this date may be expanded to 1965 based on the review of the RRPL data in February 2014



⁹ This would include doctors' offices, emergency departments, waiting rooms, classrooms, laboratories, locker rooms, etc. **There is no minimum duration of time for which the case must be present in the room.**

- NO laboratory evidence of immunity, AND
- NO documented evidence of two doses of measles-containing vaccine (given at the appropriate interval as outlined in the Saskatchewan Immunization Manual for vaccine type [MMR or MMRV]), AND
- NO history of measles disease¹¹.

See <u>Attachment – Immunoprophylaxis and Exclusion Considerations for Contacts</u> for further assessment and management

Public Health Interventions

Assessment

Individualized (person-by-person) contact investigations (Table 3) include assessment of immunization records.

Assessment varies by setting:

Individuals in Health Care Settings Who Are Contacts

- Coordination between Public Health, Occupational/Employee Health services as well as Infection Prevention and Control for the facility involved is required so there can be a systematic approach to:
 - Review immunization records and immune status for all employees (both direct and indirect patient care staff), support exclusion requirements as necessary and monitor for suspicious cases within their facility. See Figure 4, <u>Attachment – Immunoprophylaxis and Exclusion Considerations for Contacts</u>.
 - Review immunization records and immune status for all other individuals exposed (both patients and visitors), implementation of exclusion requirements as necessary and active surveillance for secondary cases. See Attachment-Immunoprophylaxis and Exclusion Considerations for Contacts.

Individuals in Child Care Centers Who Are Contacts

- Vaccination history should be reviewed for all employees, attendees and volunteers in daycare settings and appropriate action taken as per <u>Attachment – Immunoprophylaxis</u> and Exclusion Considerations for Contacts.
- Parents may also be considered as potential contacts based on their child's risk of becoming infected.

which indicated thatapproximately 93% of samples tested from individuals born prior to 1965 indicated immunity for measles, while approximately 83% of samples from persons born between 1965 and 1980 indicated immunity to measles. Factors to consider include age, history of exposure to measles, and availability of medical records.

11 Clinical judgement is required to determine if documentation is necessary.



Individuals Exposed in Public Venues

Gatherings apply to events of any size in both public and private fora. They can include (but are not limited to) social or religious functions, sports activities, shopping excursions, concerts, conferences and meetings as well as public transit.

Communication

- Person-by-person individualized investigation of contacts identified in Table 3 should include direct notification where possible.
- Identifiable contacts should, at a minimum, be provided with a letter that includes all details as outlined in education.
- When exposures involve public settings where individuals cannot be identified, news, social media as well as public websites should be used to communicate the exposure setting to the public.
 - Details to be provided in the messaging include dates and times (including two hours after the infected individual vacated the venue). Attachment –
 <u>Information for People who May Have Been Exposed to Measles in a Public Facility</u> should be used in the messaging or, at a minimum, be made available so exposed individuals have relevant information about measles and what to do if they develop symptoms.

Education

Close contacts of confirmed cases should be educated about measles and the signs and symptoms of measles. They should also be advised:

- that measles is communicable to others 4 days before the onset of the rash and until 4 days after the rash appears. They should be advised to use selfisolation (work, school, travel and other activities) as per exclusion;
- o to limit new or further exposure to other individuals; and
- to call ahead to their health care provider's office if signs and symptoms appear so arrangements can be made to see the patient in a way that reduces the chance of exposing other individuals to measles.

Refer to <u>Attachment – Template Letter to Measles Contacts</u>. Refer to <u>Attachment – Infection Prevention and Control Measures in Physicians' Offices</u> and <u>Attachment – Infection Prevention and Control Measures for Patients Suspected or Known to be Infected with Measles</u> for infection prevention and control measures in these settings.

Exclusion

 Exclusion of susceptible contacts that meet the criteria in Table 3 (C) is outlined in Figures 1–6, Attachment – Immunoprophylaxis and Exclusion Considerations for Contacts.



- Exclusion may be applied in all circumstances where the contact may be exposing other individuals (this includes work or school settings, organized groups and activities and public places including public transit).
- Consideration should be given to the number of susceptible individuals in that setting; the presence of high risk individuals (e.g. susceptible infants, or immunocompromised individuals); and the reliability of the contact to adhere to public health direction regarding early recognition and self-isolation.
- When exclusion is recommended, it should apply:
 - o From five days after first exposure and up to 21 days after last exposure; or
 - o Until serological confirmation of immunity is provided.
- If the contact develops symptoms compatible with measles, exclusion criteria for cases should be applied.
- When Ig has been provided, extend the exclusion period to 28 days after the last exposure.

Immunoprophylaxis

- There are limited data on the effectiveness of measles vaccine or immune globulin (Ig) for the prevention of measles. The use of either of these products may provide some protection or alter the clinical course of disease when provided within the timeframes outlined in Table 1, Attachment Immunoprophylaxis and Exclusion Considerations for Contacts, (Centers for Disease Control and Prevention, 2018).
- Post-exposure vaccination is preferable to the use of Ig whenever feasible to prevent secondary cases. In addition, contact follow-up provides an opportunity to improve vaccination coverage in general.
- Figures 1-6 in <u>Attachment Immunoprophylaxis and Exclusion Considerations for Contacts</u> outline the appropriate immunoprophylaxis recommendations based on the age and setting of contacts based on their immunization history.

Testing

- Routine screening for immune status of susceptible contacts is not recommended.
 Figures 4–5, Attachment – Immunoprophylaxis and Exclusion Considerations for
 Contacts outline the testing for contacts who are employees in health care settings or patients in hospital settings.
- Under certain circumstances it would be beneficial to evaluate immunity of individuals involved through immunization history or immunity serology. Figures 3-5 should be referenced if the MHO determines testing is recommended for other contacts.
- No laboratory testing for measles required if asymptomatic.
- Confirmatory testing is recommended for contacts that develop symptoms.



III. Environment

Child Care Centre/Schools Control Measures

Strict enforcement of infection control measures – refer to Saskatchewan Ministry of Health Infection Control Manual for Child Care Facilities. ¹²

Recommendations for the facility and attendees/employees must be individualized based on the characteristics and operations of the daycare/school and the susceptibility of the population in attendance.

- The school or child care centre must report immediately to public health any person suspected of having or diagnosed with measles.
- Contact tracing must be completed. Information about staff and attendees, must be
 obtained as soon as possible so immunization records can be reviewed to determine
 their susceptibility and their need for post-exposure immunoprophylaxis (see
 Attachment Immunoprophylaxis and Exclusion Considerations for Contacts). Provide
 Attachment Template Letter to Schools or Group Exposed to a Measles Case.
- Inform parents of the need for unimmunized/under immunized children to be immunized immediately.
- Contacts should be excluded as outlined in <u>Figures 1-3 Attachment Immunoprophylaxis and Exclusion Considerations for Contacts</u>.
- Individuals who attend the daycare but were not present during the exposure period
 (i.e. are not considered contacts) should not return to daycare until their
 immunizations have been brought up to date for age. However, the risks and benefits
 of returning to daycare need to be considered and exclusion may be indicated until
 transmission within the facility can be ruled out.
- Active surveillance of absent contacts should be conducted on a daily basis to determine if reason for absenteeism is related to measles. This allows public health to implement additional measures in a timely manner.
- Case finding for the source, concurrent and secondary cases should be targeted to one
 incubation period before (i.e. 21 days) the current case and for 21 days after the onset
 of rash of the last case in the setting.
- Evaluate parents and siblings of attendees to detect cases and identify susceptible individuals. Those who are susceptible should be immunized as per the Saskatchewan Immunization Manual.¹³



¹² http://www.saskatchewan.ca/live/births-deaths-marriages-and-divorces/starting-a-family/early-learning-and-child-care/child-care.

¹³ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx

Health Care Facilities Control Measures

Health care workers (HCWs)¹⁴ have an increased risk of exposure to measles and should have proof of immunity or adequate protection upon employment. See Chapter 7, Section 3.2 (Health Care Workers) of the Saskatchewan Immunization Manual¹⁵ and other relevant Saskatchewan Ministry of Health policies/memos.

- All individuals suspected of having or diagnosed with measles must be reported immediately to the local public health office, infection control and occupational/employee health.
- Strict enforcement of infection prevention and control measures. See <u>Attachment Infection Prevention and Control Precautions for Patients Suspected or Known to be Infected with Measles</u> and to the Authority's Infection Control Manual for additional details.
 - Airborne precautions in addition to Routine/Standard precautions should be taken immediately from the time measles diagnosis is being considered up to an including four days after onset of rash (Public Health Agency of Canada, 2013).
 - Immunocompromised patients should be isolated for the duration of their illness (Public Health Agency of Canada, 2013)
- Provide measles-containing vaccine to susceptible contacts (or Ig to high risk susceptible contacts) according to <u>Figure 4–5</u>, <u>Attachment – Immunoprophylaxis and</u> <u>Exclusion Considerations for Contacts</u>.
- Employees in health care settings who are contacts should be managed as per <u>Figure 4</u>, Attachment Immunoprophylaxis and Exclusion Considerations for Contacts.
- Patients in health care settings who are contacts should be managed as per <u>Figure 5</u>,
 <u>Attachment Immunoprophylaxis and Exclusion Considerations for Contacts</u>.
- Public Health should ensure that:
 - all susceptible contacts (Table 3), have been immunized as soon as possible;
 - no further cases of related illness have been detected (over the subsequent 21 day period).
 - If a person acquired measles while in hospital, a case finding for the source investigation should be conducted in partnership with public health and infection control.

¹⁵ http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter7.pdf.



¹⁴ Health care workers should be considered as ALL employees in health care settings. This includes direct and indirect patient care staff.

Outpatient Departments (including Lab and Radiology)/Physicians' Offices

Physicians' offices have been identified as the setting for transmission of secondary cases of measles. Strict application of infection prevention and control measures are required to reduce further transmission.

When measles is circulating in the community, contacts should be instructed to call HealthLine so the MHO/public health can provide direction for seeking medical attention in a way that reduces the risk of further transmission. In addition to staff using personal protective equipment, the following practical measures can be used ¹⁶:

- arrange to see patients with clinical signs of measles at the end of the day;
- provide signage and procedural masks at the entrance instructing patients to don a mask before entering the facility;
- immediately take patients to a separate examination room and only allow staff who are considered immune to measles (documented serology or have been appropriately immunized) to interact with the patient;
- ensure the exam room used by the patient is not used by other patients for two hours after the patient leaves the facility (regardless if the room is cleaned by an immune employee before the 2 hour period lapses).
- Susceptible staff should be immunized as soon as possible.

IV. Epidemic Measures

- Immediate reporting (within 24 hours) of probable and clinical cases or persons under investigation for measles.
- Determine source and manner of spread.
- Determine extent of exposure and transmission.
- If there is exposure of groups like schools, health care facilities, daycare centres, etc., it
 may be necessary to implement a coordinated immunization program for all
 unimmunized and incompletely immunized individuals to limit spread. The decision for
 this will be made in consultation with the Medical Health Officer and Saskatchewan
 Ministry of Health.
 - If vaccine supply is limited, priority should be given to young children (>6 months) for whom the risk is greatest.

¹⁶ See Attachment – Infection Prevention and Control Measures in Physicians' Offices.



- In institutional settings all individuals without adequate protection should be immunized (Heymann, 2015).
- In community-wide outbreaks, alternative measures such as broad immunization catch up programs may be considered and the date of presumed immunity expanded from 1970 to 1965.

Prevention Measures

Refer to the <u>Respiratory and Direct Contact Introduction and General Considerations</u> section of the manual that highlights topics for client education that should be considered as well as provides information on high-risk groups and activities.

Immunization

- Routine immunization of children with two doses of a measles containing vaccine in accordance with the recommended schedule in the Saskatchewan Immunization Manual.¹⁷ One dose of measles-containing vaccine given after the first birthday is 95% effective in preventing measles. Most cases of vaccine failure following one dose occur in individuals who had an inadequate immune response to the vaccine and are not related to waning immunity (American Academy of Pediatrics, 2015).
- Those born in 1970 or later who have not had two doses of measles vaccine or have not had natural measles infection should be vaccinated for measles as per the Saskatchewan Immunization Manual¹⁸
- Individuals who are travelling abroad should have a pre-travel consultation and be offered MMR is appropriate.

Education

- Education should be provided regarding respiratory etiquette and measures to prevent transmission.
- Educate the public about the disease and the need for active immunization for measles. Immunization information fact sheets¹⁹ can be used to guide discussion.



¹⁷ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx see Chapter 5, Appendix 5.2

¹⁸ This differs from the CDC year of presumed natural immunity of prior to 1957.

¹⁹ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

Revisions

Date	Change
May 2019	Updated notification timeline from Lab/Practitioner to public
,	health and from Public Health to the Ministry of Health.
	Updated Public Health purpose to include prevention of local
	transmission.
	 Included reference to PCR in case definition.
	Risk Factors - Added Child Care Worker;
	Specimen Collection - Added footnote regarding
	nasopharyngeal/throat swabs collected in physicians' office;
	Exclusion of Cases - Updated exclusion criteria to remove caveat
	regarding other susceptible individuals not yet exposed.
	Exclusion in Table - Added context on the prolonged duration of
	illness for immunocompromised individuals and to consult
	Medical Microbiologist or ID Specialist.
	• Corrected reference in Contact Exclusion to Table 3(C) rather than
	3(A).
	Updated Public Health Interventions:
	 Clarified that coordination is required with
	Employee/Occupational Health and Infection Control is
	required for exposures in Health Care and Daycare Settings.
	 Included discretion of MHO for serological testing of contacts
	 Provided more explicit information for contacts who develop
	symptoms to seek advise from public health via HealthLine
	before seeking medical attention
	 Added caveat that 1970 is generally considered cut off year
	for presumed immunity, but included information about
	RRPL information to use 1965 during outbreak situations.
	Prevention Measures
	 Updated footnote with reference to 1965 in Prevention
	Measures to explain the Saskatchewan context for this date.
	 Updated the year for eligibility for measles vaccine in
	Prevention Measures to align with the Saskatchewan
	Immunization Manual.
	 Added reference to recommending a pre-travel consultation
	for international travel considerations.
	Updated references as applicable.



May 2018	Updated to align with Panorama configuration
	Clarified the purpose for notification of cases to public health
	Incorporated an Epidemiology and Occurrence as a placeholder
	and included Saskatchewan Immunization program history from
	Sask Immunization Manual to provide context.
	Rearranged and updated the style into the new format of the
	Manual.
	References reaffirmed or updated as necessary.

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Please complete all sections.

Panorama QA complete: ☐ Ye Initials:	s □No			Panorama Client ID: Panorama Investigation ID:		
A) CLIENT INFORMATION				LHN -> SUBJE	CT -> CLIENT DETAILS -> PE	ERSONAL INFORMATION
Last Name:		First Name:	and Middle Na	nme:	Alternate Name (Goes by	/):
DOB: YYYY / MM / DD Phone #: Primary Home: Mobile contact: Workplace:	Age:		Province: Number (PHN		Preferred Communicatio i.e. home phone, text): Email Address: □Work	,
Place of Employment/School:		Gender: [□ Male	□ Female	□Other	□ Unknown
Alternate Contact: Relationship: Alt. Contact phone:		Mailing (Pos	□ Postal Add stal address): ess or FN Comi	ress	me □Temporary □Lega	al Land Description
L	ON SL	JBJECT SUMMARY	Y-> RESPIRATO	ORY &DIRECT CONTA	ACT ENCOUNTER GROUP->	CREATE INVESTIGATION
Disease Summary Classification: CASE:	Date	Classification: CONTACT:		Date	LAB TEST INFOR	MATION:
□ Confirmed	YYYY / MM / DD	□ Contact		YYYY / MM / DD	Date specimen co	ollected:
□ Does Not Meet Case	YYYY / MM / DD	□ Not a Contact	t	YYYY / MM / DD	YYYY / MM / L	DD.
☐ Person Under Investigation	YYYY / MM / DD	☐ Person Under Investigation		YYYY / MM / DD	□ Blood □ Ur	
□ Probable	YYYY / MM / DD				□ Nasopharyng	eal
□ Clinical	YYYY / MM / DD					
Disposition:		1				
FOLLOW UP:						
☐ In progress	YYYY	/ MM / DD	☐ Complet	e	YY	YY / MM / DD
☐ Incomplete - Declined	YYYY	/ MM / DD	□ Not requ		YY	YY / MM / DD
☐ Incomplete – Lost contact	YYYY	/ MM / DD	☐ Referred	I – Out of province	YY	YY / MM / DD
☐ Incomplete – Unable to locat	e YYYY	/ MM / DD	(Specify	where)	YY	YY / MM / DD
REPORTING NOTIFICATION Name of Attending Physician or	Nurse:		Location:			
Provider's Phone number:			Date Received (Public Health): YYYY / MM / DD			
Type of Reporting Source: ☐ F	Health Care Facility	□ Lab Report	□ Nurse P	ractitioner \square Ph	ysician Dother	

September 1, 2018 Page 1 of 4

Please complete all sections

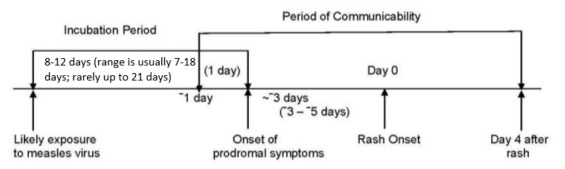
Panorama Client ID:	
Panorama Investigation ID:	

C)	SIGNS & SYMPTOMS	(Bold text = part	of case definition

LUNIS	INIVECTIO A	TION SCIENC	& SYMPTOMS
LHIV->	INVESTIGA	110N->2IGN2	& SYIVIP LUIVIS

Description	No	Yes – Date of onset	Description	No	Yes - Date of onset
Conjunctiva - inflammation (conjunctivitis)		YYYY / MMM / DD	Koplik spots		YYYY / MMM / DD
Coryza or rhinitis		YYYY / MMM / DD	Lymphadenopathy - generalized		YYYY / MMM / DD
Cough		YYYY / MMM / DD	Pain – photophobia (light sensitivity)		YYYY / MMM / DD
Fever		YYYY / MMM / DD	Rash – maculopapular (3 days)		YYYY / MMM / DD
Other s/s					

Timeline for Assessing Measles Contacts



LHN-> INVESTIGATION->INCUBATION & COMMUNICABILITY

Incubation for Case (period for acquisition): Earliest Possible Exposure Date: YYYY / MM / DD	Latest Possible Exposure Date: YYYY / MM / DD
Exposure Calculation details:	
Communicability for Case (period for transmission):	
Earliest Possible Communicability Date: YYYY / MM / DD	Latest Possible Communicability Date: YYYY / MM / DD
Communicability Calculation Details:	

E) RISK FACTORS (RF followed by + impact the Immunization Forecaster)

LHN-> SUBJECT->RISK FACTORS

DESCRIPTION	State Date Yes	N, NA, U	Add'l Info
Contact - At risk population (international travellers or immigrants)	YYYY / MM/DD		
Contact – Persons with similar symptoms	YYYY / MM/DD		
Contact to a known case (Add'l Info)	YYYY / MM/DD		
Immunocompromised - Related to underlying disease or treatment	YYYY / MM/DD		
Occupation - Health Care Worker - IOM Risk Factor	YYYY / MM/DD TE		
Special Population - Attends childcare	YYYY / MM/DD TE		
Special Population - Attends school	YYYY / MM/DD TE		
Special Population - Lives in a communal setting	YYYY / MM/DD TE		
Special Population - Post secondary education institution	YYYY / MM/DD TE		
Travel - Outside of Canada (Add'l Info)	YYYY / MM/DD AE/TE		
Travel - Outside of Saskatchewan, but within Canada (specify)_	YYYY / MM/DD AE/TE		
Other risk factor (Add'l Info)	YYYY / MM/DD		

September 1, 2018 Page 2 of 4

Please complete **all** sections

Panorama Client ID:	
Panorama Investigation ID:	

F) IMMUNIZATION	HISTORY INTERPRE	TATION SUMMARY	LHN -> INVESTIGATION-> IMMUNIZ	ZATION HISTORY INTERPR	ETATION SUMMARY
Interpretation Date	YYYY /	/ MM / DD			
Interpretation of Di	sease Immunity:	☐ IOM - Fully immunized (for age)	☐ IOM - Partially	immunized	
□ IOM – Unimmuni	zed	☐ IOM - Unclear immunization history			
Reason:					
☐ Previous disease		\square Previous responder	r/Previous history of immunity	☐ Date Of Birth	
☐ IOM - Interpretat	ion of history by inv	vestigator			
G) INTERVENTIONS	od Cook Towns		INVESTIGATION->TREATMENT &	INTERVENTIONS->INTER\	/ENTION SUMMAR
Intervention Type ar Assessment:	ia Sub Type:		Immunization: Investigator	name	
☐ Assessed for conta	acts	YYYY / MM / DD	☐ Eligible Immunization recomme		/ MM / DD
Investigator name			☐ Disease-specific immunization		/ MM / DD
			☐ Disease-specific immunization	given YYYY	/ MM / DD
Communication:	1		Isolation:	1000	/ 2424 / DD
☐ Other communica Investigator name	ition (see investigat	or Notes) YYYY / MM / DD	☐ Facility isolation Investigator name	YYYY	/ MM / DD
Letter (See Docum	nent Management)	YYYY / MM / DD	☐ Home isolation	YYYY	/ MM / DD
Investigator name	2	,, 55	Investigator name		, ,
General: Investigato	r name		Other Investigation Findings:		
☐ Disease-Info/Prev	-Control	YYYY/ MM / DD	☐ Investigator Notes		/ MM / DD
☐ Disease-Info/Prev-	Cont/Assess'd for C	Contacts YYYY/ MM / DD	☐ Document Management	YYYY	/ MM / DD
Education/counselling			Quarantine:		
☐ Prevention/Contro	ol measures	YYYY / MM / DD	Quarantine	YYYY	/ MM / DD
Investigator name Disease information	on provided	YYYY / MM / DD	Investigator name		
Investigator name	on provided	TTTT / IVIIVI / DD			
Exclusion: Investigat	or name		Testing:		
□ Work YYYY / N		☐ Preschool YYYY / MM / DD	<u> </u>	YYYY / MM / DD	
☐ School YYYY / N		□ Daycare YYYY / MM / DD	Investigator name		
Date	Intervention subtype	Comments		Next follow-up Date	Initials
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD		+		YYYY / MM / DD	
TITT / WINT / DD				1111 / 101101 / 00	
YYYY / MM / DD				YYYY / MM / DD	
				,	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
TITT / IVIIVI / DD				TITT / IVIIVI / DD	
YYYY / MM / DD				YYYY / MM / DD	

September 1, 2018 Page 3 of 4

Please complete **all** sections

 Panorama Client ID:
Panorama Investigation ID:

OUTCOMES (op	tional except fo	or severe influer	za,		LI	-IN-> INVES	TIGATION-> OUTCO
Not yet recovere Recovered Fatal		YYYY / MM / YYYY / MM / YYYY / MM /	DD Intubation /ventilation	YYYY / MM / DD	☐ Unknow		YY / MM / DD YY / MM / DD
ause of Death: (if	Fatal was selec	ted)					
EXPOSURES Acquisition Ever			INVESTIGATION-> EXF	OSURE SUMMARY	-> ACQUISITIO	N EVENT SUI	VIMARY > QUICK EN
posure Name:							
cquisition Start	YYYY / MM	/ DD to Acqu	isition End: YYYY / MM / DD				
ocation Name:							
etting Type	_		_	_		_	
Travel Transmission I		are setting	☐ Public facilities	□ Recrea	tional facilities		Most likely source
i ransmission i	Events		LHN -> INVESTIGATION-> EXPOS	URE SUMMARY -> 1	FRANSMISSION	EVENT SUN	/IMARY -> QUICK EN
ransmission vent ID	Exposure N	Name	Setting type (Consider the following settings for TE; "multiple settings" in Panorama)	f >1 select	Date/Time		# of contacts
			□ Congregate/Communal living □ Hea	Ith Care setting	YYYY / MI	/ DD	
			☐ Type of community contact ☐ Ho	usehold Exposure	to		
			☐ Public facilities		YYYY / MI	/ DD	
			☐ Congregate/Communal living ☐ Hea	lth Care setting	YYYY / MI	/ / DD	
			☐ Type of community contact ☐ Ho	usehold Exposure	to		
			□ Public facilities		YYYY / MI	/ DD	
			☐ Congregate/Communal living ☐ Hea	Ith Care setting	YYYY / MI	/ / DD	
			☐ Type of community contact ☐ Ho	usehold Exposure	to	,	
			Public facilities□	•	YYYY / MI	/ DD	
			□ Congregate/Communal living □ Hea	Ith Care setting	YYYY / MI	1 / DD	
				usehold Exposure	to	vi / DD	
			□ Public facilities□	aseriola Exposure	YYYY / MI	/ / DD	
			□ Congregate/Communal living □ Hea	Ith Care setting		. /	
				_	YYYY / MI	Л / DD	
				usehold Exposure	to YYYY / MI	л / DD	
			Public facilities				_
			□ Congregate/Communal living □ Hea —	_	YYYY / MI	M / DD	
			☐ Type of community contact ☐ Ho	usehold Exposure	to	4 / 55	
			□ Public facilities□		YYYY / MI	W / DD	
			☐ Multiple Settings		YYYY / MI	/ DD	
	Measles – Inv	ID#			to YYYY / M	M / DD	
TOTAL NUMBER	OF CONTACTS	<u>L</u>			,	,	
LHN	-> INVESTIGAT	TION-> EXPOSUI	RE SUMMARY -> TRANSMISSION EVENT	SUMMARY -> TE H	YPERLINK -> UI	NKNOWN/A	NONYMOUS CONTA
nonymous contac	cts: (te	otal number of i	ndividuals [including groups that 1:1 foll	ow-up is not requir	ed or is not fea	sible])	
	1						
itial Report Impleted by:						Date initial	report completed:

September 1, 2018 Page 4 of 4

Measles
Section 2-90
Attachment – Letter Template to a Measles Case
Page 1 of 2
April 2014

Please see the following pages for the Letter Template to a Measles Case.



<DATE>

<mr./ms. Name of Case>
<address>
<city sk postal code>

Re: Temporary Exclusion from Work and Public for <INDIVIDUAL> until <DATE>

Dear <MR./MS. NAME OF CASE>

As we have discussed, you are <SUSPECTED/CONFIRMED> to have measles disease. Because this is a very contagious disease, all precautions need to be taken to prevent possible spread of infection to others. You are considered to be contagious from <DATE> to <DATE>.

Your assistance is important to prevent spreading this disease to individuals who have not been immunized or who have not had the disease previously. This means that you are required to remain in your home (not to be out in public or at school/work) until <DATE>. This also means that during this time, there cannot be visitors in the home. Should you require medical attention, it is important to call ahead to your health care provider so they can plan to see you in a way that reduces the chance of exposing other individuals to measles.

Thank you for your cooperation in identifying individuals and locations where people may have been exposed to measles and for your cooperation during this period. We appreciate that you are doing your best to prevent further spread of infection. Please feel free to call < PHONE NUMBER> as needed.

The Medical Health Officer has authority under *The Public Health Act, 1994* of Saskatchewan to enforce compliance with this requirement.

Sincerely,

<NAME OF PUBLIC HEALTH DESIGNATE>
<TITLE>

cc: Medical Health Officer

2019 05 01

Table 1. Vaccination or Immune Globulin (Ig) for Susceptible Contacts – See <u>Table 3</u> (Person-by-person contact investigation)

If measles vaccine is given within 72 hours of exposure, it may provide some protection.

Do not delay providing vaccine to contacts that are not up-to-date, even if >72 hours have lapsed in order to provide protection from future exposures. Immune globulin is available in two products:

- IMIg (intramuscular immune globulin)
- IVIg (intravenous immune globulin)

Denviction	Time since Exposure to Measles		
Population	≤ 72 hours	73 hours – 6 days	
Susceptible infants 0-6 months of age;	IMIg (0.5 mL/kg)		
Susceptible immunocompetent infants 6-12 months of age;	MMR vaccine	IMIg (0.5 mL/kg)	
Susceptible immunocompetent persons 12 months of age or older	MMR vaccine series		
susceptible pregnant women	IVIg (400 mg/kg) OR IMIg (0.5 mL/kg) to maximum of 15 mL (limited protection if 30 kg or more);		
immunocompromised individuals 6 months of age or older;*	IVIg (400 mg/kg) OR IMIg 0.5 mL/kg to maximum of 15 mL (limited protection if 30 kg or more);		
Individuals with confirmed measles immunity	N/A		

^{*} Ig is recommended for immunocompromised individuals for whom measles vaccination is contraindicated and past measles vaccination is no longer considered to be effective as outlined in the Saskatchewan Immunization Manual, Chapter 7¹. Maximum doses and sites are outlined in SIM, Chapter 8²

Source: Canada Communicable Disease Report, 2018 (Tuvis)

The following figures outline when vaccine or immune globulin should be provided to different populations and when exclusion/self-isolation should be implemented.

When exclusion is recommended, it should apply:

- From five days after first exposure and up to 21 days after last exposure; or
- Until serological confirmation of immunity is provided.

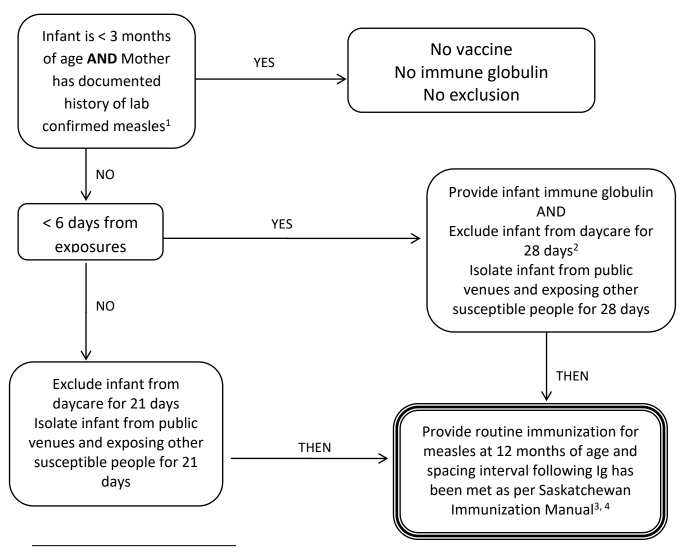
If a contact develops symptoms compatible with measles, exclusion criteria for cases should be applied. When Ig has been provided, extend the exclusion period to 28 days after the last exposure.

² https://www.ehealthsask.ca/services/Manuals/Documents/Ch.%208%20Administration%20of%20Bio%20Prods.%20Oct%202018.pdf



¹ http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter7 and

Figure 1. Infants < 6 months of age



¹ Maternal antibodies from vaccination wane more quickly than antibodies from natural infection. Considering the vulnerable population, immune globulin is recommended for infants < 3 months if mother's immunity is vaccine-induced.

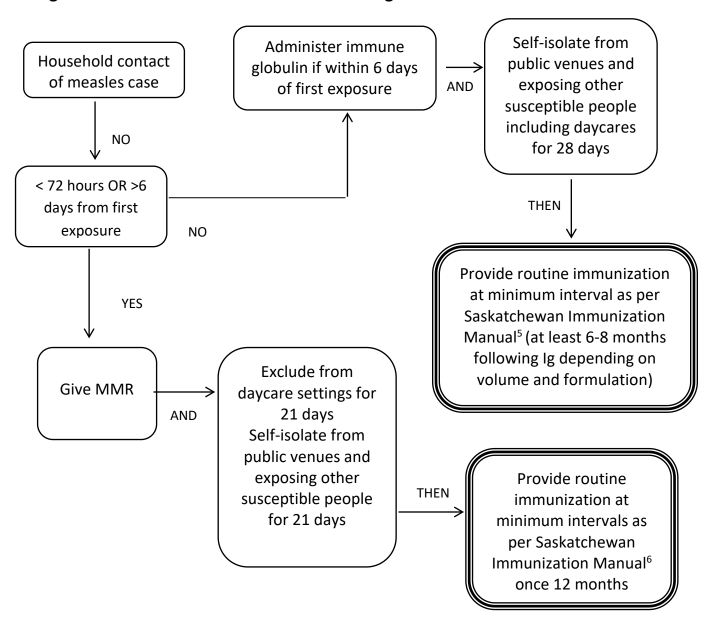


² Immune globulin may not prevent measles, and may cause a longer incubation period up to 28 days

³ http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter5

⁴ If risk of measles is ongoing and Ig was not given, MMR may be given at 6 months of age.

Figure 2. Infants 6 month to <12 months of age⁵



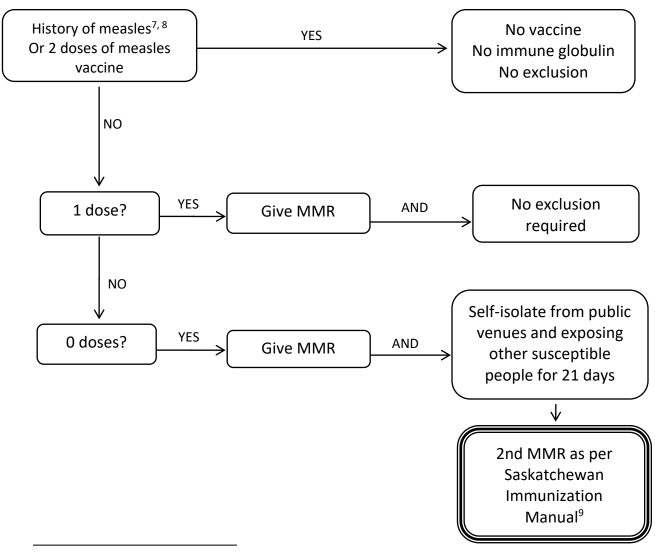
⁵ No previous measles-containing vaccine previously provided for travel or past measles exposure.

⁶ http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter5

Attachment – Immunoprophylaxis and Exclusion Considerations for Contacts
Page 4 of 8

2019 05 01

Figure 3. Immunocompetent Children and Adults (Non-Health Care Settings)



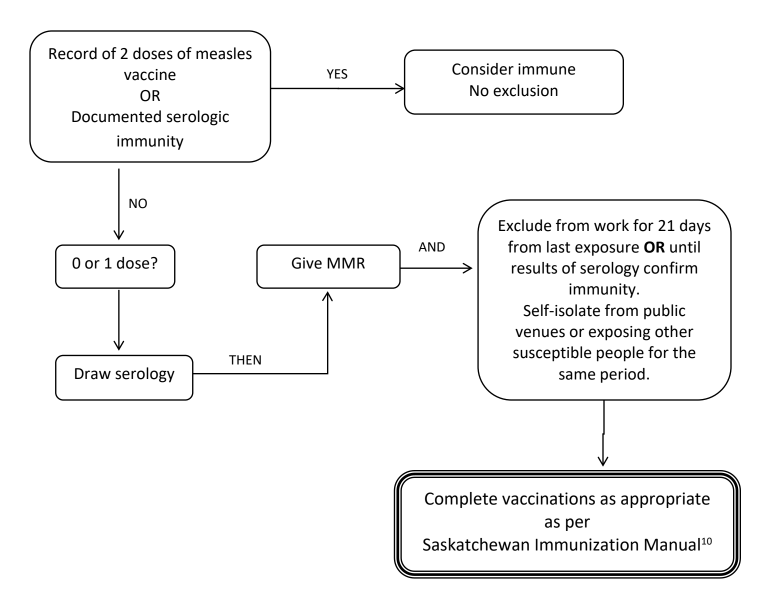
⁷ Generally, people born prior to 1970 are considered to have natural immunity. During outbreak situation, this date may be expanded to 1965 based on a review of RRPL (February 2014): approximately 93% of samples tested from individuals born prior to 1965 indicated immunity for measles, while approximately 83% of samples from persons born between 1965 and 1980 indicated immunity to measles. Factors to consider include age, history of exposure to measles, and availability of medical records.



⁸ Clinical judgement is required to determine if serology is necessary. If born in Canada in 1970 or later, previous rubella immunity serves as a proxy for measles immunity based on past measles-rubella vaccine.

⁹ http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter5 and http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter7

Figure 4. Health Care Settings - All Employees

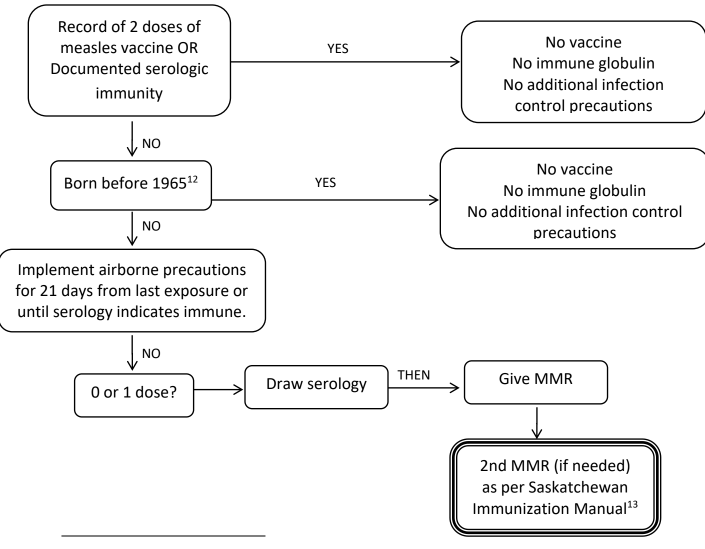


¹⁰ http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter7

Attachment – Immunoprophylaxis and Exclusion Considerations for Contacts

Page 6 of 8 2019 05 01

Figure 5. Health Care Settings - Patients¹¹



¹¹ If immunocompromised, consult with MHO and attending physician.

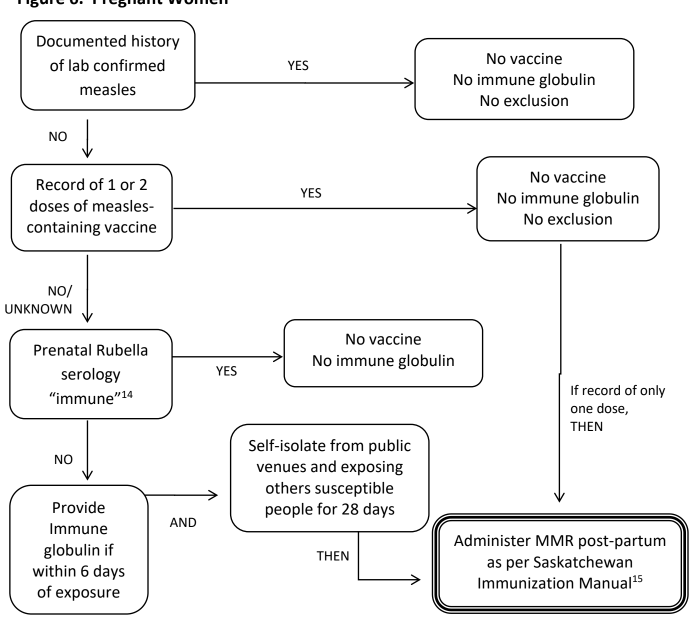


¹² Generally, people born prior to 1970 are considered to have natural immunity. During outbreak situation, this date may be expanded to 1965 based on a review of RRPL (February 2014): approximately 93% of samples tested from individuals born prior to 1965 indicated immunity for measles, while approximately 83% of samples from persons born between 1965 and 1980 indicated immunity to measles. Factors to consider include age, history of exposure to measles, and availability of medical records.

¹³ http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter5 and http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter7

2019 05 01





¹⁴ For women born in Canada after 1970, rubella immunity is a proxy for immunization with measles/rubella vaccine. This may not be try for foreign born women.



¹⁵ http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter5 and http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter7

Revisions

Date	Change
April 2019	Figure 1 – added footnote #4 regarding use of MMR in ongoing exposure situations; added caveat about use of MMR as per spacing guidelines following Ig. Figure 2 – simplified flowchart; updated spacing interval following Ig from 5 months to 6-8 months; corrected footnote 5 to refer to measles exposure (not disease) Figure 3 – added footnote that rubella immunity may serve as proxy for measles immunity; adjusted to refer to year of birth 1965 for immunity during outbreaks Figure 4 – simplified flowchart for 0 or 1 doses of vaccine Figure 5 – simplified flowchart for 0 or 1 doses of vaccine Figure 6 – added footnote that rubella immunity may serve as a proxy for measles immunity
September 2018	Updated the dosage and formulation recommendations for immunoglobulin to align with the September 2018 National Advisory Committee on Immunization recommendations.



Measles
Section 2-90
Attachment – Letter Template to a Measles Contact
Page 1 of 2
April 2014

Please see the following pages for the Letter Template to a Measles Contact.



<DATE>

<MR./MS. NAME OF CONTACT REQUIRING EXCLUSION>
<ADDRESS>
<CITY SK POSTAL CODE>

Re: Temporary Exclusion from Work and Public for <INDIVIDUAL> until <DATE>

Dear < MR./MS. NAME OF CONTACT REQUIRING EXCLUSION>

As we have discussed, you have been exposed to measles, a highly contagious disease. Because this is a very contagious disease, and you have not been immunized previously, you are at increased risk of developing infection. Until it is determined that you have not been infected, all precautions need to be taken to prevent possible spread of infection to others. You are considered to be contagious from <DATE> to <DATE>.

Your assistance is very important to prevent spreading this disease to individuals who have not been immunized or who have not had the disease previously. Measles is contagious from 4 - 5 days before a person develops a rash until 4 days after the rash appears. This means that you are required to remain in your home (not to be out in public or at school/work) and should not have visitors to your home from <DATE> (5 days after 1st exposure) to <DATE> (21 days after last exposure). It is during this time that you may develop infection.

If you develop symptoms during this time, it is important to call ahead to your health care provider's office so arrangements can be made for you to be seen a way that reduces the chance of exposing other individuals to measles. Early symptoms include:

- high fever;
- cough;
- runny nose;
- red eyes.

A rash then develops after a day or 2 and usually starts on the face then spreads over the rest of the body.

If you do develop symptoms, we will be in touch with you to gather a list of individuals that you have been in contact with so we can offer immunization and education as necessary.

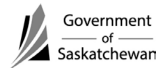
Thank you for your cooperation during this period. We appreciate your assistance in preventing the possible spread of infection. Please feel free to call <PHONE NUMBER> as needed.

Sincerely,

<NAME OF PUBLIC HEALTH DESIGNATE>

cc: Medical Health Officer

Please see the following pages for the Letter Template to a School or Group Exposed to a Measles Case.



<DATE>

<NAME SCHOOL/SPORTS GROUP/ETC.>
<ADDRESS>
<CITY SK POSTAL CODE>

Re: Possible Exposure to Measles between <DATE> to <DATE>

Dear < NAME SCHOOL/SPORTS GROUP/ETC.>

We are investigating a person with red measles (Rubeola) who, while infectious, may have exposed others during <SCHOOL/SPORTS GROUP ACTIVITY/ETC.> Measles is a highly contagious disease spread through the air (by coughing, sneezing, talking). Public health is actively obtaining immunization records for individuals who have been exposed and may be contacting you to ask for your assistance in obtaining this information. Individuals who have had two doses of measles-containing vaccine (commonly provided as measles, mumps rubella [MMR] vaccine) or who have had a lab-confirmed infection in the past are considered immune and not at risk for infection.

Individuals who have not been immunized are at risk of developing infection and may be asked to stay out of <SCHOOL/SPORTS GROUP ACTIVITY/ETC.> until immunization has been provided.

Individuals who have been exposed may develop symptoms as early as 7 or as late as 21 days after the exposure. Early symptoms usually include:

- high fever;
- cough;
- runny nose;
- red eyes.

A rash then develops after a day or 2 and usually starts on the face then spreads over the rest of the body.

If you develop symptoms during this time, it is important to call ahead to your health care provider's office and inform them of your exposure so arrangements can be made for you to be assessed in a way that reduces the chance of exposing other individuals to measles.

Measles is contagious before early symptoms develop, which is 4 - 5 days before a person develops a rash, and remains contagious until 4 days after the rash appears. If you have symptoms and are waiting for laboratory confirmation, it is very important that you avoid contact with others who are not immune until the 5th day after the rash appears (you are no longer considered contagious after that time). If you require further medical attention, it is important call ahead to your health care provider as mentioned above.

We appreciate your assistance in preventing the possible spread of infection. Please feel free to call NUMBER as needed.

Sincerely,

<NAME OF PUBLIC HEALTH DESIGNATE>

cc: Medical Health Officer

Attachment – Information for People Who May Have Been Exposed to Measles in a Public Facility Page 1 of 1 April 2014

Public Health is investigating persons with measles who, while infectious, visited one or more public venues.

Measles is a highly contagious disease. Individuals who have had two doses of measles-containing vaccine (commonly provided as MMR or MMR-V) are considered immune and not at risk for infection. Individuals born before 1965 are likely to have been exposed to measles in childhood, and are considered to have a natural immunity to it.

Individuals born in 1965 or later and who have not been immunized are at risk of developing infection. Symptoms may develop as early as 7 or as late as 21 days after the exposure. Early symptoms usually include:

- high fever;
- cough;
- runny nose;
- red eyes.

A rash then develops after a day or 2 and usually starts on the face then spreads over the rest of the body.

If you develop symptoms compatible with measles in 7 to 21 days after being exposed, it is important to call ahead to your health care provider's office and inform them of your exposure so arrangements can be made for you to be assessed in a way that reduces the chance of exposing other individuals to measles.

Measles is contagious before symptoms develop, which is 4 to 5 days before a person develops a rash, and remains contagious until 4 days after the rash appears. If you have symptoms and are waiting for laboratory confirmation, it is very important that you avoid contact with others who are not immune until the 5th day after the rash appeared (you are no longer considered contagious after that time). If you require further medical attention, it is important to call ahead to your health care provider as mentioned above.

We appreciate your assistance in preventing the possible spread of infection. Please call your local public health office or the HealthLine at 811 as needed.



Measles
Section 2-90
Attachment – Infection Prevention and Control Measures in
Physicians' Offices Page 1 of 2
April 2014

Please see the following pages for the Infection Prevention and Control Measures in Physicians' Offices.



Infection Prevention and Control Measures in Physicians' Offices

- For each patient encounter, screen the patient to determine whether the patient has any signs/symptoms of measles. Screen the patient:
 - at time of booking;
 - upon arrival in the waiting room; or
 - in exam room.

NOTE: Symptoms of measles include prodromal fever, conjunctivitis, coryza, cough and small spots with white or bluish white centers on an erythematous base on the buccal mucosa (Koplik spots). Three to five days after the start of the symptoms a red, blotchy (maculopapular) rash appears on the face and then progresses down the body.

- Book anyone with symptoms at the end of the day if clinical status allows; ensure other patients are not in the office.
- Use airborne precautions as measles virus remains suspended in the air; health care providers (HCP) entering the patient's room must wear a fit-tested N95 respirator unless the HCP has documented immunity to measles.
- Where possible immune staff should provide care to patients suspected to have measles.
- Post signage at the entrance (Attachment Measles Alert Poster) instructing patients
 with signs and symptoms of measles or other respiratory symptoms to put on a surgical
 mask before entering the clinic.
- Provide surgical face masks for symptomatic patients as close to the entry of the clinical
 office as possible, with instructions on how to put on and take off mask. Consider having
 client call when in the parking lot (outside) and having immune staff go out to provide
 patient with a surgical mask. Instruct patient to wear the mask at all times while in the
 clinical office. Pediatric masks should be available on an individual patient basis if
 needed.
- Quickly triage the patient out of the common waiting areas and move the patient to an examining room. If possible, the patient should enter and exit through a separate entrance and go directly in and out of the examination room. Close door to examination room. The room should be a single room with a solid door (closed).
- Place a "DO NOT ENTER" sign for staff on the closed door.
- Keep the door closed to allow sufficient time (**two hours**) for the air to change in the room and be free of droplet nuclei before using the room for a non-immune patient.
- A bathroom used by a suspect case, to collect urine for measles virus, should also be off
 limits to non-immune patients for two hours. The bathroom door should be closed and
 have a "DO NOT ENTER" sign for staff/patients placed on the closed door. This is another
 reason to have patients seen at the end of the day.
- Routine cleaning for the room/equipment (once free of droplet nuclei) is sufficient.
- Ensure/encourage appropriate hand hygiene by staff and patient.

These recommendations align with Ontario's Infection Prevention and Control for Clinical Office Practice (June 2013) at:

http://www.publichealthontario.ca/en/eRepository/IPAC Clinical Office Practice 2013.pdf

Measles

Section 2-90

Attachment – Infection Prevention and Control Measures for Patients Suspected or Known to be Infected with Measles

Page 1 of 2

April 2019

Please see the following pages for the Non-Saskatchewan Measles Alert Poster.



Measles Alert

Do you have a **FEVER** and **RASH**, with **COUGH** or **RUNNY NOSE** or **RED EYES**?

Have you been **EXPOSED TO SOMEONE** with suspected or confirmed measles?



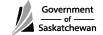
PLEASE: Put on a mask.

Clean your hands with alcohol hand rub.

Report to the nurse or front desk immediately.

Measles is very contagious.

Help prevent the spread of measles.



Measles

Section 2-90

Attachment – Infection Prevention and Control Measures for Patients Suspected or Known to be Infected with Measles

Page 1 of 2

April 2014

Please see the following pages for the Measles Alert Poster.



Measles Alert

Do you have a **FEVER** and **RASH**, with **COUGH** or **RUNNY NOSE** or **RED EYES**?

Have you been **EXPOSED TO SOMEONE** with suspected or confirmed measles?



PLEASE: Put on a mask.
Clean your hands with alcohol hand rub.
Report to the nurse or front desk immediately.

Measles has been confirmed in Saskatchewan.

Measles is very contagious.

Help prevent the spread of measles.



Measles

Section 2-90

Attachment – Infection Prevention and Control Measures for Patients Suspected or Known to be Infected with Measles

Page 1 of 2

April 2014

Please see the following pages for the Infection Prevention and Control Measures for Patients Suspected or Known to be Infected with Measles.



Infection Prevention and Control Precautions for patients suspected or known to be infected with Measles (Rubeola)

Always follow Routine Practices including a Point of Care Risk Assessment

Triage

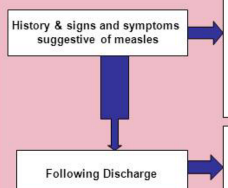
Display posters at patient/client entrances with instructions for anyone with measles symptoms to identify themselves immediately to staff

If patient presents with symptoms of measles1

- ➤ Triage suspected patients as expeditiously as possible into an airborne infection isolation room/AIIR (negative pressure room) to avoid exposure to contacts in waiting rooms.
 - o If a negative pressure room is unavailable, the patient should be managed in a private room with the door closed.
- > Place appropriate signage (Airborne Precautions) outside of the room in a noticeable location (if applicable)

Assessment

Where possible, only immune² staff should provide care to patients suspected or confirmed to have measles (Staff should check with local Employee Health and Wellness/Occupational Health and Safety nurse for their immune status)



- ➤ A fitted **N**-95 respirator should be worn by all non-immune Healthcare Workers (HCWs) who enter the room.
- ➤ Patient should be restricted to their room, but instructed or assisted with respiratory and hand hygiene, if transport out of room is required
- > Patients should be provided with a surgical mask to wear when they are outside of the exam room
- > Postpone elective procedures that generate droplets
- > DO NOT use the room occupied by a patient with suspected or confirmed case of the measles for 2 hours after they have been discharged
- > Leave the door closed and leave alert signage posted until 2 hours have elapsed and a terminal clean has been performed

Communication

Inform:

- ·Patient of the requirements for isolation precautions/visitor restrictions
- •IMMEDIATE notification of all <u>suspect and confirmed</u> cases to your local Public Health Office (306-) or the MHO on call (306-).

The Public Health Office will provide additional guidance for case management.

- •Your Infection Control Professional
- •Your local Housekeeping department after 2 hours have elapsed following discharge, to perform a terminal clean of the exam room as per regional policies and procedures.
- If exposure of a Healthcare Worker (HCW) is suspected, contact local Employee Health and Wellness/ Occupational Health and Safety office.
- Symptoms include Fever 38.3° C or higher, cough, runny nose or red eyes, red blotchy rash appearing three to seven days after fever starts, beginning behind the ears and on the face and spreading down to the body and then to the arms and legs
- Immune Serological evidence of measles IgG antibodies; or Documentation of 2 doses of measles-containing vaccines for all HCWs.

Notification Timeline:

From Lab/Practitioner to Public Health: Immediate.

From Public Health to Saskatchewan Health: Within 72 hours. Public Health Follow-up Timeline: Initiate within 24-48 hours.

Public Health Purpose for Notification of Meningococcal Disease (adapted from British Columbia Center for Disease Control [2017])

- To minimize mortality and serious morbidity from meningococcal disease;
- To rapidly identify close contacts of the case and to provide recommendations for appropriate preventive measures for close contacts so as to prevent further spread of infection and disease;
- To provide information about the disease, its transmission, and methods of prevention;
- To identify clusters or outbreaks of infection and to initiate appropriate prevention and control measures;
- To track epidemiology trends of meningococcal disease in Saskatchewan including risk factors and distribution;
- To monitor the effectiveness of prevention and control measures;
- To plan expansion or introduction of future immunization programs;
- To provide timely clinical care including diagnosis and treatment using current, evidence-based guidelines
- To make timely and evidence informed actions on outbreaks; and
- To inform the public and medical community about meningococcal disease.

Surveillance Case Definition¹ (Public Health Agency of Canada, May 2008)

Confirmed Case	Clinical evidence of invasive disease with laboratory confirmation of infection:
	 isolation of Neisseria meningitidis from a normally sterile site (blood, cerebrospinal fluid (CSF), joint, pleural or pericardial fluid)

¹ Surveillance case definitions ensure uniform reporting to allow comparability of surveillance data. The definition is not intended to be used for clinical or laboratory diagnosis or management of cases.



	OR	
	 demonstration of N. meningitidis DNA by an appropriately validated nucleic acid test (NAT)² from a normally sterile site. 	
Probable Case	Clinical evidence ¹ of invasive disease with purpura fulminans or petechiae, with no other apparent cause and with non-	
	confirmatory laboratory evidence:	
	 detection of N. meningitidis antigen in the CSF. 	
¹ Clinical illness associated with invasive meningococcal disease usually manifests itself as meningitis		
and/or septicaemia, although other manifestations may be observed (e.g., orbital cellulitis, septic arthritis). Invasive disease may progress rapidly to petechiae or purpura fulminans, shock and death.		

Both confirmed and probable cases of <u>invasive</u> meningococcal disease (IMD) are notifiable to the provincial and national level.

²Each jurisdiction will have a validation process for the NAT that they have in place.

At this time, conjunctivitis and pneumonia cases due to *N. meningitidis* are not notifiable nor reported to the Ministry of Health or the Public Health Agency of Canada, however case definitions are as follows:

- A <u>conjunctivitis case</u> requires isolation of *N. meningitidis* from the eye or the conjunctival sac in association with purulent conjunctivitis.
- A <u>pneumonia case</u> is one with a Gram strain (if done) showing Gram-negative diplococci and a polymorphonuclear cell response from sputum or respiratory aspirate, isolation with heavy growth of *N. meningitides*, <u>and</u> clinical or radiological evidence of pneumonia.
- Patients with N. meningitidis conjunctivitis or pneumonia should be treated by the clinician with appropriate systemic antibiotics (Public Health Agency of Canada, 2005).

Epidemiology and Occurrence

Under development

Additional Background Information

Causative Agent

Neisseria meningitidis, the meningococcus, is a Gram-negative, aerobic diplococcus. *Neisseria* are divided into Serogroups including A, B, C, W-135, X and Y (Heymann, 2015).



Symptoms

- Sudden onset of fever, intense headache, nausea and often vomiting, stiff neck, and photophobia.
- Petechial rash with pink macules or, very rarely, vesicles (Heymann, 2015).
- Delirium and coma often appear.
- Occasional fulminating cases exhibit sudden prostration, ecchymoses, and shock.

Complications

10-20% of survivors suffer long-term sequelae:

- Neurological deficits
- Hearing loss
- Limb loss

The case fatality rate is 8-15% (Heymann, 2015)

Reservoir/Source

Humans are the only reservoir. Asymptomatic colonization in the upper respiratory tract occurs in up to 5-10% of people. Less than 1% of colonized individuals develop disease (Heymann, 2015).

Incubation Period

2 to 10 days, commonly 3 to 4 days (Heymann, 2015).

Period of Communicability

As long as 7 days before the onset of symptoms until meningococci are no longer present in discharges from the nose and mouth, usually within 24 hours of the beginning of appropriate antibiotic treatment. Up to 5-10% of people can be asymptomatic carriers; communicability is difficult to determine in carriers (Heymann, 2015).

Mode of Transmission

- Person-to-person by direct contact with respiratory droplets from the nose and throat of an infected person. Can be carried for distances < 1 meter by droplets generated by coughing or sneezing (Public Health Agency of Canada, 2005).
- Fomite transmission is insignificant (Heymann, 2015).



 The likelihood/risk of person-to-person transmission of meningococcal disease is related to the type of contact and length of the contact with the confirmed case (Public Health Agency of Canada, 2005).

Specimen Collection and Transport

- Cultures of blood and CSF are indicated in all patients with suspected invasive disease.
- Cultures of petechial (purpuric lesions) scrapings, synovial fluid, pleural fluid and pericardial fluid are positive in some patients.
- In accordance with the Saskatchewan Disease Control Regulations, section 21.1, <u>all</u> clinical isolates **must** be forwarded to the Roy Romanow Provincial Laboratory (RRPL) for serotyping. Ideally this should be done when the initial gram stain or positive preliminary culture results are available if not ordered sooner by the Medical Health Officer (MHO). Further characterization may be done by National Microbiology Lab.
- **Note:** Since *N. meningitidis* can be part of the normal nasopharyngeal flora, isolation of this organism from the throat is not helpful in determining the cause of the disease.

Refer to the RRPL Compendium of Tests for details on specimen collection and transportation – available online at https://rrpl-testviewer.ehealthsask.ca/.

Risk Factors/Risk Groups

Susceptibility to the clinical disease is low and decreases with age. Persons deficient in certain complement components are especially prone to recurrent disease.

Increased risk of IMD is associated with the following:

- functional or anatomic asplenia;
- underlying immune deficiencies (properdin deficiency, deficiency of terminal complement components or factor D deficiency);
- candidates and recipients of solid organ transplant;
- recipients of hematopoietic stem cell transplant;
- infants, adolescents, and young adults;
- crowded housing/living conditions;
- low socioeconomic status;



- active or passive exposure to tobacco smoke and concurrent upper respiratory tract infections (U.S. Centers for Disease Control and Prevention, 2015);
- young people living in an institutional setting such as military recruits or enlisted personnel and university students in a dormitory setting or at residential camps;
- living in the African meningitis belt (area from Senegal to Ethiopia).

Public Health Investigation

I. Case

Refer to <u>Attachment – Meningococcal Disease (invasive) Data Collection Worksheet</u> to assist.

History

- Determine if case has been laboratory confirmed and if molecular serotyping has been completed.
- Onset of illness, presentation and treatment (with what and when) to
 determine incubation period and period of communicability which helps to
 identify the possible source and contacts to be followed.
- Travel history may be of significance in contact tracing.
- Determine if case has underlying medical conditions or falls into a risk category.
- Try to determine acquisition exposures as well as transmission exposures (e.g. student residence, sporting events, childcare or occupational settings etc.).
- Liaison with school authorities when a case is a student.
- Review immunization history of the case.

Public Health Interventions

Assessment

- Assess for contacts Aggressive contact tracing, identification, and appropriate management, is the foundation to the prevention of secondary cases. Refer to Table 2 - <u>Contact Definition</u>.
- Obtain names, addresses, and phone numbers of all possible contacts. This information may need to be obtained from someone close to the case.

Communication

• Letters can be sent to classrooms and other group settings where individual contact tracing is not required (i.e. involving school age and adults where there are no vulnerable contacts) to inform them of the exposure, symptom monitoring and when to seek medical attention (see Sample letter).



Education

 All cases should be provided disease information as well as information on prevention and control measures including period of communicability and avoiding contact with vulnerable individuals.

Exclusion

• Individuals with the disease are generally in hospital so are not attending activities. They are isolated until 24 hours after initiation of an appropriate antibiotic (e.g., ceftriaxone, etc.). Otherwise cases or contacts generally do not need to be excluded from any activities.

Immunization

 Case follow-up should be used as an opportunity to recommend immunizations they are eligible for as per the Saskatchewan Immunization Manual. The case should be assessed for underlying risk factors and should be offered vaccine as outlined in the Saskatchewan Immunization Manual, Appendix 7.1.²

Referrals

- When clients are hospitalized, communication with hospital staff and or infection control staff is important.
- Refer client to primary caregiver for referrals (physiotherapy, occupational therapy, Home Care) are available if necessary.

Treatment

The public health practitioner should direct any questions regarding the current treatment protocols to the physician or MHO. The following serves as a reference for the public health investigator:

- Antibiotic treatment is required and should be started as soon as presumptive diagnosis is made. For patient management the client's physician should consult an infectious disease specialist.
- In addition to therapeutic antibiotics, the case should receive chemoprophlaxis before
 hospital discharge unless the infection was treated with an antibiotic that is effective in
 nasopharyngeal eradication of N. meningitidis (Public Health Agency of Canada, 2005).



² http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter7

II. Contacts/Contact Investigation Contact Definition

Table 2: Definition of Close Contacts (Public Health Agency of Canada, 2005)

- Household contacts of a case.
- Persons who share sleeping arrangements with the case (e.g. shared bedrooms or dorm rooms in residences).
- Persons with intimate contact with the case.
- Persons who have direct contamination of their nose or mouth with the oral/nasal secretions of a case (e.g., kissing on the mouth, shared cigarettes, shared drinking bottles) in the 7 days before onset of illness.
- Health care workers (HCWs) who have had intensive (e.g., intubating, resuscitating or closely examining the oropharynx), unprotected contact (without using droplet precautions) with infected patients in the 7 days before onset of illness and completion of the first 24 hours of treatment.³
- Children and staff in childcare and nursery school facilities during the 7 days before onset of illness.
- Airline passengers sitting immediately on either side of the case (but not across the aisle) when the total time spent aboard the aircraft was at least 8 hours during the 7 days before onset of illness.

Chemoprophylaxis is not recommended for close contacts of patients with evidence of *N. meningitidis* only in non-sterile sites such as an oropharyngeal swab, endotracheal secretions, or conjunctival swab. (US Centers for Disease Prevention and Control, 2018).

Prior to the introduction of the routine use of chemoprophylaxis for household contacts, they were 500 to 1,200 times at greater risk of IMD than the general population. There is also an increased risk in child care settings, although the risk is lower than in household settings. Risk is not increased in social contacts, therefore the individual relationship to the case must be considered as outlined in Table 2 for school, transportation, social, and workplace contacts.

³ HCWs are rarely at risk even when caring for infected patients and chemoprophylaxis is rarely warranted except when they meet the definition of a close contact (Heymann, 2015).



Public Health Interventions

Assessment

Assess for symptoms.

Communication

Individual follow-up of contacts in in larger daycares, classrooms, schools, teams, workplaces, etc., is generally <u>not recommended</u>. These individuals should be informed by letter from public health, advising them to see their physician if they develop symptoms. These persons, if they become symptomatic, should not be assumed to have pertussis but should be assessed, tested and treated appropriately.

Chemoprophylaxis

The purpose of chemoprophylaxis is to eradicate nasopharyngeal colonization by *N. meningitidis* and thus prevent disease in contacts and further transmission to susceptible persons (Public Health Agency of Canada, 2005).

- Antimicrobial chemoprophylaxis should be administered as soon as possible, ideally less than 24 hours, after identification of the index patient but is still recommended for up to 10 days after the last contact with the index case while they were infectious (Public Health Agency of Canada, 2005; U.S. Centers for Disease Control and Prevention, 2011)⁴.
- Chemoprophylaxis should be offered for close contacts (as defined in Table 2).
- It should also be considered for close contacts of a case that is strongly suspected to be IMD, if laboratory confirmation cannot be obtained within 24 hours.
- Chemoprophylaxis is <u>not</u> routinely recommended for HCWs including emergency personnel^{5.} Only health care personnel who were managing an airway⁶ or exposed to respiratory secretions of a patient with meningococcal disease (US Centers for Disease Prevention and Control, 2018).

⁵ HCWs are rarely at risk even when caring for infected patients and chemoprophylaxis is rarely warranted except when they meet the definition of a close contact (Heymann, 2015). 6 intubating, resuscitating or closely examining the oropharynx



⁴ Chemoprophylaxis is unlikely to be of benefit if given > 10 days after the most recent exposure to an infectious case (Public Health Agency of Canada, 2005).

- For residents of an institutional living or residential camp setting, only contacts that share a room with the case need prophylaxis. If there are other persons who meet the contact definition, they should also receive prophylaxis.
- Refer to <u>Attachment Meningococcal Chemoprophylaxis Guidelines</u> for details.

Education

- Close contacts of confirmed cases should be educated about meningococcal disease and the signs and symptoms of meningococcal disease (meningitis and meningococcemia).
- They should be advised to seek immediate medical attention if they develop febrile illness or any other signs (see Symptoms).
- They should also be advised about the modes of transmission, period of communicability, and measures that they can take to reduce the risk of acquiring the disease.
- Reinforce proper hand washing and personal protective measures as per <u>Respiratory and Direct Contact Introduction and General Considerations</u> regarding diseases transmitted via respiratory and direct contact.
- Exposed household contacts and daycare contacts should be observed and advised to seek prompt medical attention if they develop a febrile illness.
- Meningococcal Disease (Neisseria meningitidis) information sheet can be provided.
- Advise individuals of the increased risk from overcrowding in living quarters and workplaces, such as schools, camps, and ships.

Exclusion

Due to the low secondary attack rate and the short duration of chemoprophylaxis, contacts do not need to be excluded from day care, school, or work.

<u>Immunoprophylaxis</u>

 Close contacts of individuals with meningococcal infections have an increased risk of developing IMD; this risk is greatest for household contacts. The increased risk of meningococcal disease for household contacts persists for up to one year after disease in the index case and beyond any protection from antibiotic chemoprophylaxis. In general, this prolonged risk is not seen among other contacts that do not have ongoing exposure (Public Health Agency of Canada, 2005).



When the serogroup is vaccine-preventable (i.e., serogroup A, B, C, W-135 or Y), the following individuals should be considered for immunoprophylaxis in addition to chemoprophylaxis:

- household contacts of a case;
- persons who share sleeping arrangements with the case;
- persons who have direct exposure of their nose or mouth with oral/nasal secretions of a case (e.g., kissing on the mouth, shared cigarettes, shared drinking bottles); and
- children and staff in child care and nursery school facilities.

The following individuals are close contacts who **do not** require immunoprophylaxis (they should <u>only receive</u> chemoprophylaxis) as they do not have ongoing exposure:

- HCWs who have managed the airway⁷ of a meningococcal case.
- Airline passengers sitting immediately on either side of the case (but not across the aisle) when the total time spent aboard the aircraft was at least 8 hours.

Vaccination history should be reviewed for eligible close contacts (date and type of previous meningococcal vaccine). When indicated, immunization should be carried out as soon as possible. For those who were previously immunized, revaccination should be provided based on the following criteria:

- Individuals with underlying medical risk factors (as per Saskatchewan Immunization Manual, Appendix 7.18) should be revaccinated if it has been **more than four weeks** since a previous meningococcal vaccine was received (Public Health Agency of Canada, 2015).
- Individuals who were immunized at less than 1 year of age should be revaccinated if it has been **more than 4 weeks** since a previous meningococcal vaccine was received (Public Health Agency of Canada, 2015).
- Individuals who were immunized after their first birthday and individuals without underlying medical risk factors should be revaccinated if they have not been vaccinated with a meningococcal vaccine in the past year (Public Health Agency of Canada, 2015).

⁸ http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter7.



⁷ intubating, resuscitating or closely examining the oropharynx

To the extent possible, Saskatchewan follows the recommendations in the 2015 Canadian Immunization Guide for post-exposure vaccination of close contacts for vaccine preventable meningococcal serogroups⁹.

Special Considerations for Immunoprophylaxis

Serogroup B:

Recommendations for post-exposure use of meningococcal B vaccine are not included in the current version of the evergreen Canadian Immunization Guide (as of April 2015). Refer to Chapter 10 of the Saskatchewan Immunization Manual for the multicomponent meningococcal B vaccine (4CMenB) schedule and complete the series that they are eligible for based on their age.

Serogroup C:

Individuals 11 years of age and older who are contacts to serogroup C can receive either Men-C-C or Men-C-ACYW-135. Saskatchewan parameters for which vaccine to provide are outlined in Age and Older.

Testing

Testing of asymptomatic contacts is of no value and is not recommended.

III. Environment

Child Care Centre/Schools Control Measures

Ensure each parent receives the information sheet about <u>Meningococcal Disease</u> (*Neisseria meningitidis*).

⁹ https://www.canada.ca/en/public-health/services/publications/healthy-living/canadian-immunization-guide-part-4-active-vaccines/page-13-meningococcal-vaccine.html



Management of the centre/school:

- Notify and educate staff and parents of contacts of the case to be alert for anyone with sudden onset of fever, intense headache, nausea and often vomiting, stiff neck and photophobia. Seek prompt evaluation by a physician for any ill child.
- The centre director must notify public health if any additional children become ill.
- All children and care providers should be evaluated as to whether they meet criteria for contact definition for meningococcal disease and be given chemoprophylaxis as appropriate.
- Notify parents of all childcare centre attendees of the occurrence of a case of meningococcal disease (see the appropriate Attachment – Sample Letter about Invasive Meningococcal Disease – Prophylaxis Recommended or Sample Letter about Invasive Meningococcal Disease – Prophylaxis NOT Recommended) and a Meningococcal Disease (Neisseria meningitidis) information sheet.
- Assess immunization status of children and staff and immunize as per Immunoprophylaxis section.

Special Considerations for Funeral Homes

Follow routine infection control practices when handling cadavers.

Traditionally, cadavers with meningococcal disease have been considered a possible source of infection. The risk is likely very low if the deceased person had been treated with an effective antibiotic for at least 24 hours before death.

In instances when the deceased had not been treated with an effective antibiotic before death, it is prudent for those who have occupational contact with a cadaver to follow routine infection control practices with additional droplet and contact precautions (Public Health Agency of Canada, 2005).

IV. Epidemic Measures

<u>Outbreaks</u>

An outbreak is defined as increased transmission of *N. meningitidis* in a population, manifested by an increase in cases of the same serogroup.



Outbreaks can be subdivided into organization-based or community-based outbreaks using the criteria shown in Table 3.

Table 3: Types of Outbreak

Organization- based	Increased transmission of <i>N. meningitidis</i> in an organization or institution with two or more cases of the same serogroup occurring within a 4-week interval. This includes restricted populations, such as schools, day cares, sports groups, or social groups, as well as nursing homes or long-term care facilities.
Community- based	Increased transmission of <i>N. meningitidis</i> in a community, with three or more confirmed cases of the same serogroup occurring within a three-month interval AND an age-specific incidence OR specific community population incidence of approximately 10/100,000, where there is an absence of an epidemiologic link between cases. This is not an absolute threshold and should be considered in the context of other factors.

Regardless of the type of outbreak, contact tracing, identification of close contacts, and provision of chemoprophylaxis to close contacts need to be conducted as described for sporadic cases.

When evidence suggests that an outbreak is occurring with increased transmission of *N. meningitidis* involving a vaccine-preventable serogroup in a delineated population, vaccination of persons at high risk should be considered. The specific epidemiology of the outbreak needs to be ascertained to define the group at risk. Decisions regarding the use of vaccine in communities with a higher than expected rate of disease should be made in consultation with the chief MHO.

When an outbreak occurs:

- 1. Communication strategy should be in place to provide timely information to the public. This would include:
 - why some people are being immunized and not others;
 - why some people are being given rifampin and not others;
 - not sharing of drinking equipment, cigarettes, etc. especially at sports and high school events;
 - low risk to people entering outbreak area.



- A communication strategy aimed at the health care community should also be developed. This includes notification of local hospital emergency departments, labs, infection control departments, and physicians/nurse practitioners.
- 3. An outbreak advisory committee comprising ministry and local public health representatives, clinicians, and medical laboratory personnel should be established. Keep other jurisdictions informed about the outbreak and related control strategies.
- 4. A communication strategy is prepared before a decision is made to undertake an outbreak immunization program.

Prevention and Education

Refer to the <u>Respiratory and Direct Contact Introduction and General Considerations</u> section of the manual that highlights topics for client education that should be considered as well as provides information on high-risk groups and activities.

Education

- Educate the public about the disease and the need for active immunization. Immunization information fact sheets can be used to guide discussion.
- Education should be provided regarding respiratory etiquette and measures to prevent transmission.

Immunization

- Immunize infants, children, and adults according to the recommended ageappropriate schedules. Refer to Saskatchewan Immunization Manual.¹⁰
- Provide the appropriate vaccine to travelers at risk (refer to local travel health consultant for details). The risk to travelers planning to have prolonged contact with the local population in areas experiencing endemic/epidemic meningococcal A or C diseases may be reduced by immunization.
- Consider vaccination/revaccination for individuals at risk in outbreak situations if one of the vaccine preventable serogroups has been confirmed.



¹⁰ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

Revisions

Date	Change
September 2018	 Updated to align with Panorama configuration; Incorporated Meningococcal Disease Data Collection Worksheet; Clarified the purpose for notification of cases to public health; Incorporated an Epidemiology and Occurrence placeholder into the chapter; Rearranged and updated the style into the new format of the Manual. Implemented boxes to draw attention to treatment, chemo and Immunoprophylaxis information. Removed reference to treatment of conjunctivitis and chemoprophylaxis for contacts to cases with conjunctivitis based on more recent references. Updated chemoprophylaxis recommendations for HCWs based on more recent references. References reaffirmed or updated as necessary.



References

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Meningococcal Disease (invasive) Data Collection Worksheet

Panorama QA complete: ☐ Yes Initials:	□No	Please complete a	all sections.	Panorama Client ID: Panorama Investigation ID:			
A) CLIENT INFORMATION			LHN -> SUBJEC	T -> CLIENT DETAILS -> PERSONAL INFORMATION			
Last Name: First Na			liddle Name:	Alternate Name (Goes by):			
DOB: YYYY / MM / DD Phone #: Primary Home:	Age:	Health Card Proving Health Card Number		Preferred Communication Method: (specify - i.e. home phone, text): Email Address: □Work □Personal			
Place of Employment/School:		Gender: Mal	e □ Female	□Other □ Unknown			
Alternate Contact: Relationship: Alt. Contact phone:		□ No fixed □ Pos Mailing (Postal ad Street Address or	Address Type: No fixed Postal Address Primary Home Temporary Legal Land Description Mailing (Postal address): Street Address or FN Community (Primary Home): Address at time of infection if not the same:				
B) INVESTIGATION INFORMATION	N LHN ->SUBJ	ECT SUMMARY-> RESP	IRATORY & DIRECT CONTAC	T ENCOUNTER GROUP-> CREATE INVESTIGATION			
Disease Summary Classification: CASE:	Date	Classification: CONTACT:	Date	LAB TEST INFORMATION:			
□ Confirmed	YYYY / MMM / DD	□ Contact	YYYY / MMM / D	Date specimen collected:			
□ Does Not Meet Case	YYYY / MMM / DD	□ Not a Contact	YYYY / MMM / D				
☐ Person Under Investigation	YYYY / MMM / DD	□ Person Under Investigation	YYYY / MMM / D	□ CSF			
□ Probable	YYYY / MMM / DD			☐ Joint fluid ☐ Pericardial fluid			
Disposition: FOLLOW UP: ☐ In progress ☐ Incomplete - Declined ☐ Incomplete — Lost contact ☐ Incomplete — Unable to loca	YYYY / MM / DD YYYY / MM / DD YYYY / MM / DD		Complete Not required Referred – Out of province pecify where)	YYYY / MM / DD YYYY / MM / DD YYYY / MM / DD			
REPORTING NOTIFICATION Name of Attending Physician or	Loc	cation:					
Provider's Phone number:			te Received (Public Health):	YYYY / MMM / DD			
Type of Reporting Source: ☐ Health Care Facility ☐ Lab Report ☐ Other			Nurse Practitioner Ph	ysician			
C) DISEASE EVENT HISTORY		LUNI	> INVESTIGATION SPICEAGE	S SIMMARY (LIDDATE) SDISEASE EVENT LISTARY			
Site / Presentation:	☐ Meningitis	□ Sepsis	Unkn	SUMMARY (UPDATE)->DISEASE EVENT HISTORY OWN			

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Meningococcal Disease (invasive) Data Collection Worksheet

Please complete all sections.

Panorama Client ID:
Panorama Investigation ID:

D) SIGNS & SYMPTOMS (Bold text = part of case definition)	LHN-> INVESTIGATION-> SIGNS & SYMPTOMS

Description	No	Yes – Date of onset	Description	No	Yes - Date of onset
Arthritis - septic		YYYY / MMM / DD	Neurologic - delerium		YYYY / MMM / DD
Bruising - ecchymoses		YYYY / MMM / DD	Pain - photophobia (sensitivity to light)		YYYY / MMM / DD
Cellulitis - orbital		YYYY / MMM / DD	Prostration		YYYY / MMM / DD
Coma		YYYY / MMM / DD	Purpura fulminans (coagulation of small blood vessels)		YYYY / MMM / DD
Fever		YYYY / MMM / DD	Rash - maculopapular		YYYY / MMM / DD
Headache		YYYY / MMM / DD	Rash - petechial		YYYY / MMM / DD
Meningitis		YYYY / MMM / DD	Sepsis (e.g. bacteremia, septicemia, etc.)		YYYY / MMM / DD
Nausea		YYYY / MMM / DD	Shock		YYYY / MMM / DD
Neck stiffness (nuchal rigidity)		YYYY / MMM / DD			YYYY / MMM / DD

E) INCUBATION AND COMMUNICABILITY	LHN-> INVESTIGATION->INCUBATION & COMMUNICABILITY			
Incubation for Case (period for acquisition): Earliest Possible Exposure Date: YYYY / MM / DD	Latest Possible Exposure Date: YYYY / MM / DD			
Exposure Calculation details:				
Communicability for Case (period for transmission): Earliest Possible Communicability Date: YYYY / MM / DD	Latest Possible Communicability Date: YYYY / MM / DD			
Communicability Calculation Details:				

RISK FACTORS (RF followed by + impact the Immunization Forecaster) LHN-> SUBJECT->RISK FAC				
DESCRIPTION	Yes Start Date	N, NA, U	Add'l Info	
Chronic Medical Condition - Cochlear Implant +				
Chronic Medical Condition Congenital or Acquired, or Functional Asplenia +				
Contact At risk population (international travellers or immigrants) (i.e. risk areas)				
Contact - IMD Case: serogroup A, Y, or W-135 +	YYYY / MM/DD			
Contact - IMD Case: serogroup B +	YYYY / MM/DD			
Contact - IMD Case: serogroup C +	YYYY / MM/DD			
Contact to a known case (Add'l Info)	YYYY / MM/DD			
Immunocompromised – Acquired Complement Deficiency +				
Immunocompromised – Congenital immunodeficiency +				
Immunocompromised - Related to disease or treatment (Add'l Info)				
Immunocompromised - Transplant Candidate or Recipient - Solid Organ/Tissue +				
Occupation - Health care worker - IOM Risk Factor	TE			
Occupation - Child care worker	TE			
Behaviour - Sharing personal items (cigarettes, water bottles, etc)	TE			
Setting - Crowded living conditions (>1 person per room excluding bathrooms)	TE			
Special Population – Attends childcare	TE			
Special Population - Attends school	TE			
Special Population - Lives in a communal setting	TE			

September 1, 2018 Page 2 of 4

Meningococcal Disease (invasive) Data Collection Worksheet

Please complete all sections.

 Panorama Client ID:
Panorama Investigation ID:

DESCRIPTION			Yes Start Date	N, NA, U	Add'l Info			
Special Population - institution	Post secondary education	on	TE					
Travel: Outside of Ca	anada (Add'l Info)		YYYY / MM/DD AE					
Travel Outside of Sas (Add'l Info)	skatchewan, but within (Canada	YYYY / MM/DD AE					
Other risk factor (Ad	ld'l Info)							
G) COMPLICATIONS			ı	II.	•	LHN-	> INVESTIGATION->CO	MPLICATIONS
Description			es Pate of onset		Description		Yes Date of onset	
Disseminated intrav	ascular coagulation (DIC	:)	YYYY / MMM	/ DD	Gangrene		YYYY / MMN	1 / DD
Other complications								
H) IMMUNIZATION	HISTORY INTERPRETATION	ON SUMI	MARY	LHN	-> INVESTIGATION-> IMN	IUNIZATION H	ISTORY INTERPRETATION	ON SUMMARY
Interpretation Date	e: YYYY / MI	M / DD	serotype:					
Interpretation of Di	isease Immunity:	IOM - F	ully immunized (f	or age)	□ IOM - P	artially immun	ized	
☐ IOM – Unimmun			nclear immuniza		Valid doses receive	ed: Do	ses needed:	
Reason:								
□ Previous disease			☐ Previous re	sponder/P	revious history of immunit	V	□ Date Of Birth	
□ IOM – Interpreta	ition of history by investi	igator		-,,	, , , , , , , , , , , , , , , , , , , ,	,		
I) TREATMENT		0			I HN-> INVESTIG	ATION-> MEDI	CATIONS->MEDICATIO	NS SIIMMARV
THEATMENT					LINE / NEV LOTTO	ATTOM > IVIEDI	CATIONS PINEDICATIO	NO SOMMAN
Medication (Panora	ma = Other Meds) :							
Prescribed by:					Started on: YYYY / N	MMM / DD		
J) INTERVENTIONS				II	IVESTIGATION->TREATME	NT & INTERVE	NTIONS->INTERVENTION	ON SUMMARY
Intervention Type a	nd Sub Type:			,				
Assessment: Assessed for con	Investigator name stacts		YYYY / MM /	DD C	nmunization: Inves Eligible Immunization re Disease-specific immuni Disease-specific immuni	zation recomm	nended YYYY / N	MM / DD
Communication:					nmunoprophylaxis	_		
	ation (see Investigator N	lotes)	YYYY / MM /	DD E	Immunoprophylaxis (Co	ntacts only)		
Investigator name Letter (See Docur	ment Management)		YYYY / MM /	DD				
Investigator name	= :		, , , , , , , , ,					
General: Investigato					olation:			
☐ Disease-Info/Prev			YYYY/ MM / DI	Ĭ F	_	tigator name tigator name	YYYY / N YYYY / N	
☐ Disease-Info/Prev	-Cont/Assess'd for Conta	acts	YYYY/ MM / DI	D	- Home isolation hives	tigator name	1111 / 10	IIVI / DD
Education/counselli	ing.			Т	esting:			
☐ Prevention/Contr	-		YYYY / MM /	טט	Lab testing recommende	ed YYYY / I	MM / DD	
☐ Disease informati	ion provided		YYYY / MM /	DD	nvestigator name			
Investigator name	atornamo			D	oforral:			
Exclusion: Investigator name □ Daycare YYYY / MM / DD □ Preschool YYYY / MM / DD			_ ا	Referral: □ Consultation with MHO □ Primary Care Provider				
☐ School YYYY	/ MM / DD 🗆 Wo		YYYY / MM /		- consultation with with		ary care i rovider	
Other Investigation Investigator note	-	□ Docur	ment Manageme	nt				
Date	Intervention subtype	Comme	ents	•			Next follow-up Date	Initials
YYYY / MM / DD							YYYY / MM / DD	
YYYY / MM / DD							YYYY / MM / DD	
YYYY / MM / DD							YYYY / MM / DD	

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Meningococcal Disease (invasive) Data Collection Worksheet Please complete all sections. Panorama Client ID: Panorama Investigation ID: ___ YYYY / MM / DD K) OUTCOMES (optional except for severe influenza) LHN-> INVESTIGATION-> OUTCOMES □ Not yet recovered/recovering YYYY / MM / DD ☐ ICU/intensive medical care YYYY / MM / DD ☐ Hospitalization YYYY / MM / DD ☐ Recovered YYYY / MM / DD ☐ Intubation /ventilation YYYY / MM / DD □ Unknown YYYY / MM / DD ☐ Fatal YYYY / MM / DD Cause of Death: (if Fatal was selected) _ L) Acquisition Event LHN-> INVESTIGATION-> EXPOSURE SUMMARY-> ACQUISITION EVENT SUMMARY -> QUICK ENTRY Acquisition Event ID:_ Exposure Name: Acquisition Start YYYY / MM / DD to Acquisition End: YYYY / MM / DD Location Name: ___ **Setting Type** ☐ Travel ☐ Health care setting Public facilities ☐ Recreational facilities ☐ Most likely source M) Transmission Events LHN -> INVESTIGATION-> EXPOSURE SUMMARY -> TRANSMISSION EVENT SUMMARY -> QUICK ENTRY Transmission Exposure Name Setting type Date/Time # of contacts

Transmission	Exposure Name	Setting type	Date/Time	# of contacts
Event ID		(Consider the following settings for TE; if >1 select		
		"multiple settings" in Panorama)		
		☐ Congregate/Communal living ☐ Health Care setting	YYYY / MM / DD	
		☐ Type of community contact ☐ Household Exposure	to	
		☐ Public facilities (daycare, school, etc)	YYYY / MM / DD	
		☐Congregate/Communal living ☐ Health Care setting	YYYY / MM / DD	
		☐ Type of community contact ☐ Household Exposure	to	
		☐ Public facilities (daycare, school, etc)	YYYY / MM / DD	
		☐ Congregate/Communal living ☐ Health Care setting	YYYY / MM / DD	
		☐ Type of community contact ☐ Household Exposure	to	
		☐ Public facilities (daycare, school, etc)☐	YYYY / MM / DD	
		☐Congregate/Communal living ☐ Health Care setting	YYYY / MM / DD	
		☐ Type of community contact ☐ Household Exposure	to	
		☐ Public facilities (daycare, school, etc)☐	YYYY / MM / DD	
		☐ Multiple Settings	YYYY / MM / DD	
	Meningococcal Contacts –		to	
	Inv ID#		YYYY / MM / DD	

N) TOTAL NUMBER OF CONTACTS

LHIN -> IINVESTIGA	ATION-> EXPOSURE SUMMARY -> TRANSMISSION EVENT SUMMARY -> TE HYPERLINK -> UNKNOWN/ANONYMOUS CONTACTS
A	(Andrew or beautified in the last transport of the state
Anonymous contacts:	_ (total number of individuals [including groups that 1:1 follow-up is not required or is not feasible])

ı		I
	Initial Report	Date initial report completed:
	completed by:	YYYY / MMM / DD

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Meningococcal Disease - invasive Attachment - Meningococcal Chemoprophylaxis Guidelines

Date Reviewed: May, 2015 Section: 2-100 Page 1 of 2

Chemoprophylaxis* for Close Contacts of Individuals with Meningococcal Infection					
Drug***	Dosage**	Comments			
Rifampin	Adults: • 600 mg orally every 12 hours for 4 doses	Should not be used in pregnancy - Ceftriaxone is a safer alternative.			
	Children ≥ 1 month of age (up to 60 kg): 10 mg/kg (maximum 600 mg) orally every 12 hours for 4 doses	Urine and tears may be stained red. Advise against wearing of soft contact lenses as they can also be stained. Can reduce effectiveness of oral contraceptives. Advise use of alternative/ additional contraceptive measures.			
	Infants < 1 month of age: 5 mg/kg per dose orally every 12 hours for 4 doses	Refer to <u>Rifampin Chemoprophylaxis Dosage</u> <u>Guide for <i>Neisseria meningitidis</i></u> for information on dosing.			
Ceftriaxone	Adults and adolescents ≥ 12 years: ■ 250 mg IM x 1 dose	Recommended drug for pregnant women. Alternative for persons who cannot tolerate oral medication.			
	Children < 12 years: 125 mg IM x 1 dose	Dilute in 1% lidocaine to reduce pain at injection site.			
Ciprofloxacin	Adults ≥ 18 years of age: 500 mg PO x 1 dose	Contraindicated during pregnancy and lactation. Only approved for persons > 18 years of age. Not recommended for prepubertal children			

^{*}Chemoprophylaxis should be offered to all persons having close contact with an IMD case during the infectious period (the 7 days before onset of symptoms in the case to 24 hours after onset of effective treatment) regardless of their immunization status. Chemoprophylaxis is unlikely to be of benefit if given > 10 days after the most recent exposure to an infectious case. If antibiotics such as penicillin, which do not reliably eliminate nasopharyngeal carriage, have been used for treatment, the index case should also receive antibiotics that clear nasal carriage before

discharge.

(Source: Public Health Agency of Canada, 2005)



^{**}PO, orally; IM, intramuscularly.

^{***} See Appendix F - Patient Information Sheets for medication fact sheets.

Rifampin Chemoprophylaxis Dosage Guide for Neisseria meningitidis

	Dosage Guide based on the noted weight in kg below. Calculate dose based on exact weight. Maximum dose 600 mg.															
Weight in kg Dosage by age	5	6	7	8	9	10	15	20	25	30	35	40	45	50	55	60 Max or adult dose
<1 mo of age 5 mg/kg/dose (25 mg/ml suspension) 1 Dose PO q 12 h x 4 doses	1.0 ml	1.2 ml	1.4 ml	1.6 ml	1.8 ml	2.0 ml	3.0 ml	4.0 ml								
>1 mo of age 10 mg/kg/dose (max dose 600 mg) 1 Dose PO q 12 h x 4 doses	2.0 ml	2.4 ml	2.8 ml	3.2 ml	3.6 ml	4.0 ml	6.0 ml	8.0 ml	10.0 ml	12.0 ml	14.0 ml	16.0 ml	18.0 ml	20.0 ml	22.0 ml	24.0 ml

Recommendations

- 1. Use the appropriate weight-specific dose noted in the first column in the chart above for infants and children.
- 2. Rifampin Pediatric Suspension should be prepared by a pharmacist as follows:
 - Add contents of 3 (300mg) caps or 6 (150 mg caps) of Rifampin to 36 mls of simple syrup to yield a 25 mg/ml suspension.
 - SHAKE WELL.
- 3. Store prepared suspension and simple syrup at room temperature because of their tendency to crystallize if refrigerated.
- 4. Discard prepared suspension after treatment course is completed. Preparation expires after 28 days.
- 5. As much as possible, use only one preparation form per client (i.e., capsule(s) only, or suspension only).
- 6. Give client a <u>Rifampin</u> information sheet.

Note:

- Rifampin is contraindicated in pregnancy. Discuss Ceftriaxone dose with MHO.
- If necessary, discuss alternative treatments with MHO for non-pregnant adults.

Meningococcal Disease - invasive

Attachment – Immunoprophylaxis Guidelines for Serogroup C Contacts Who Are 11 Years of Age and Older

Date Reviewed: May, 2015 Section: 2-100

Page 1 of 2

Individuals 11 years of age and older who are contacts to serogroup C can receive either Men-C-C or Men-C-ACYW-135. Saskatchewan parameters for which vaccine to provide are outlined as follows:

Contact Group	Vaccine	Recommendation	
Individuals 11 years and older with underlying risk factors (as	Men-C-ACYW-135	Provide to individuals who: • have not received a previous dose of Men-C-ACYW-135 as part of their routine immunization • OR • are due for a Men-C-ACYW-135 booster dose as per high-risk immunization schedule.	
per SIM Appendix 7.1 ¹)	Men-C-C	Provide to high-risk individuals who: • have had a dose of Men-C-ACYW-135 more than 4 weeks ago BUT • are not yet due for their routine Men-C-ACYW-135 booster. 1	
Grade 6 students (regardless of age)	Men-C-ACYW-135	Provide to individuals who: • have not received a dose of meningococcal C-containing vaccine in the past year AND • are eligible for Men-C-ACYW-135 as part of the routine school immunization program.	
uge)	Men-C-C	If Men-C-C is provided at the time of exposure, Men-C-ACYW-135 should be provided a minimum of 4 weeks after Men-C-C to complet the routine immunization Grade 6 program.	



¹ http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter7

Meningococcal Disease - invasive Attachment –Immunoprophylaxis Guidelines for Serogroup C Contacts Who Are 11 Years of Age and Older

Date Reviewed: May, 2015 Section: 2-100

Page 2 of 2

Contact Group	Vaccine	Recommendation
Individuals born Jan 1, 2000 or later (up to age	Men-C-ACYW-135	Provide to individuals who: • have not received a dose of meningococcal C-containing vaccine in the past year AND • have not received a single dose of Men-C-ACYW-135 as part of the routine school immunization program.
22) ²	Men-C-C	Provide to individuals who: • have received one dose of Men-C-ACYW- 135 AND • it has been more than 1 year since their last meningococcal C-containing vaccine.
Individuals 11 years and older with no risk factors and not eligible for the Grade 6 program	Men-C-C	Provide to individuals who have not received a dose of meningococcal C-containing vaccine in the past year.



² http://www.ehealthsask.ca/services/manuals/Documents/sim-chapter5

Notification Timeline:

From Lab/Practitioner to Public Health: Within 48 hours.
From Public Health to Ministry of Health: Within 2 weeks.
Public Health Follow-up Timeline: Initiate within 72 hrs.

Public Health Purpose for Notification of Mumps (Adapted from Massachusetts, 2017)

- To prevent mortality and serious morbidity from mumps through rapid contact tracing;
- To prevent transmission of mumps from imported cases;
- To track epidemiology trends of mumps in Saskatchewan including risk factors and distribution;
- To monitor the effectiveness of prevention and control measures including the immunization program;
- To inform decisions about future immunization programs
- To make timely and evidence informed actions on outbreaks; and
- To inform the public and medical community about mumps.

Information

Surveillance Case Definition¹ (Public Health Agency of Canada, 2008)

Confirmed Case	Clinical illness ^a and laboratory confirmation of infection in the		
	absence of recent immunization ^b with mumps-containing		
	vaccine:		
	isolation of mumps virus from an appropriate clinical		
	specimen		
	OR		
	 detection of mumps virus ribonucleic acid (RNA) 		
	OR		
	• seroconversion or a significant rise (e.g., fourfold or greater)		
	in mumps immunoglobulin G (IgG) titre by any standard		
	serologic assay between acute and convalescent sera		

¹ Surveillance case definitions ensure uniform reporting to allow comparability of surveillance data. The definition is not intended to be used for clinical or laboratory diagnosis or management of cases.



	 OR positive serologic test for mumps immunoglobulin M (IgM) antibody ^c in a person who is either epidemiologically linked to a laboratory-confirmed case or has recently travelled to an area of known mumps activity. OR Clinical illness in a person with an epidemiologic link to a laboratory-confirmed case.
Probable Case	Clinical illness ^a • in the absence of appropriate laboratory tests
	 OR in the absence of an epidemiologic link to a laboratory-confirmed case.

^a Clinical illness is characterized by acute onset of unilateral or bilateral tender, self-limited swelling of the parotid or other salivary gland, lasting > 2 days, and without other apparent cause.

A laboratory-confirmed case may not exhibit clinical illness, as up to 30% of cases are asymptomatic.

^c IgM serology has the potential for false-positive findings. *If the clinical presentation is inconsistent with a diagnosis of mumps or in the absence of recent travel/exposure history, IgM results must be confirmed by the other listed confirmatory methods.

Further strain characterization is indicated for epidemiologic, public health and control purposes.

Although the case definition indicates that a positive serologic test for mumps IgM is a confirmed case, the challenge with relying on the IgM serology alone is that other etiologic agents (e.g., infection with parainfluenza virus, Epstein-Barr virus (EBV), or *Mycoplasma pneumoniae*) cross react and result in a false positive IgM for mumps. The positive predictive value of mumps IgM is low when the incidence of mumps is low in the community and most results will be false positives. If you have any questions regarding the interpretation of lab results, please call the to Roy Romanow Provincial Laboratory (RRPL) or the local Medical Health Officer (MHO).

To confirm diagnosis of mumps, the following must be taken into consideration:

- lab information;
- clinical presentation;
- case history.



^b The most frequent reaction to measles-mumps-rubella (MMR) immunization is malaise and fever (with or without rash) occurring 7-12 days after immunization. Parotitis has occasionally occurred after immunization. However, this should be determined for each case, as these reactions and the time frame can vary (*Canadian Immunization Guide*, 7th edition).

Epidemiology and Occurrence

Canada

With the introduction of vaccine in 1969 in Canada, the number of reported mumps cases nationally decreased by more than 99% from an average of 34,000 cases per year in the early 1950s to fewer than 400 cases in the early 1990 and an annual average of 79 cases in the period 2000–2006. From 1996 to 2006, only five outbreaks primarily involving pre-school or school-aged children, adolescents, and young adults were reported.

Over time, the age distribution of mumps cases has changed. In Canada, the proportion of reported cases aged 20 years and older increased from 14% in 1988–1990 to 64% in 2003–2005 while the proportion of cases aged 1–9 decreased from 49% to 17% during the same period (Public Health Agency of Canada, 2009).

In Saskatchewan, between 2000-2016, a total of 32 cases of mumps were reported with zero cases reported in 6 of these years.

In 2017, 77 cases were largely related to three outbreaks: one involved a mine worksite; another involved a sports team which likely contracted the virus during interprovincial sports events. Mumps was introduced into a remote community from a neighbouring province.

Table 1. Evolution of the Mumps Immunization Program in Saskatchewan

1979	MMR vaccine for age 1 year			
Fall 1991 to	Mass MMR immunization for teen-aged boys in high schools and post-			
1992	secondary institutions			
2001	MMR used exclusively for all 1st and 2nd doses; MR discontinued by Berna			
2003 - 2004	2 dose mumps catch-up in Grade 6			
2007 - 2013	2-dose mumps catch-up for eligible Grade 12 students			
2008 - 2013	2-dose mumps catch-up for eligible Grade 8 students			
2011 - 2013	2nd dose provided to eligible Grade 6 students			
May 2013	Adult born since Jan. 1, 1970 eligible for 2 MMR doses			

Saskatchewan Immunization Manual (2018)



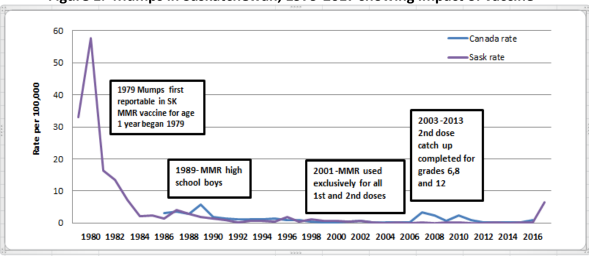


Figure 1. Mumps in Saskatchewan, 1979-2017 showing impact of vaccine

Additional Background Information

Causative Agent

Mumps virus, a member of the family *Paramyxoviridae*, genus *Rubulavirus*.

Symptoms

Prodromal symptoms are non-specific and include myalgia, headache, malaise and low-grade fever.

This acute viral illness is characterized by fever, swelling and tenderness of one or more salivary glands, usually the parotid and sometimes the sublingual or submaxillary glands. Children often experience respiratory symptoms. Up to 30% of infected cases can be asymptomatic.

- Orchitis can occur in as many as 20-30% of postpubertal males.
- Asceptic meningitis occurs in up to 10% of cases and rarely, encephalitis may occur as a complication (Heymann, 2015).
- Other rare complications may include arthritis, mastitis, glomerulonephritis, myocarditis, endocardial fibroelastosis, thrombocytopenia, cerebellar ataxia, transverse myelitis, ascending polyradiculitis, pancreatitis, oophoritis, hearing impairment etc. (American Academy of Pediatrics, 2018).
- During the first trimester of pregnancy, mumps is associated with an increased rate of spontaneous abortion (Heymann, 2015).



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Reservoir/Source

Humans are the only known natural hosts.

Incubation Period

Range from 12-25 days (usually16-18 days) (American Academy of Pediatrics, 2018).

Period of Communicability

Can be isolated for up to 7 days before the onset of symptoms and for as long as 9 days after the onset of the illness. The period of maximum infectiousness is between 2 days before to 5 days after the onset of illness (Heymann, 2015)

Mode of Transmission

Droplet spread or direct contact with the respiratory secretions of an infected person; airborne transmission also occurs.

Specimen Collection and Transport

For the diagnosis of mumps the recommended specimens to be collected are:

serum sample

AND

- a swab from around opening of Stenson's duct
 OR
- o a urine sample.

The buccal swab and urine sample will be tested by polymerase chain reaction (PCR). The respiratory pathogens currently in circulation can interfere with the serologic diagnosis (cross-reactions leading to false positive IgM).

- It is recommended that a buccal swab be obtained at the same time as the serological sample.
- Samples should be collected when the patient first presents with symptoms; these have the best chance of having a positive result by PCR if mumps infection is present. Among symptomatic persons who have received a dose of MMR, the virus may be cleared rapidly.



Public Health Investigation

I. Case

Control measures must be implemented immediately for all confirmed, probable or clinical cases. Awaiting lab confirmation must not delay the initiation of control measures. Refer to Attachment - Mumps Data Collection Worksheet to assist.

History

- Determine mumps immunization history including number of doses, date(s) administered, and type of vaccine.
- If the case has been fully immunized against mumps, further details of immunizations are required (lot numbers, where the vaccines were received, etc.).
- Determine if there is an opportunity for <u>acquisition</u> through:
 - o contact with a confirmed or probable case of mumps.
 - history of travel (seven to 21 days before onset of rash), or contact (seven to
 21 days before onset of rash) with a person who had recent travel.
- Health conditions that may render the individual more susceptible to infection or alter the period of communicability (e.g. immunocompromised).
- Identify contacts (refer to contact definition) by inquiring about opportunities for <u>transmission</u> events during the infectious period, which includes seven days prior to and five days after the parotitis appears:
 - o household;
 - daycare/school;
 - o workplaces;
 - o health care facilities² (including physicians' offices and waiting rooms).

Public Health Interventions

Assessment

Assess for contacts paying particular attention to susceptible contacts as per <u>Table 3</u>.

Communication

• Letters can be sent to classrooms and other group settings where individual contact tracing is not required (i.e. involving school age and adults where there are no vulnerable contacts) to inform them of the exposure, symptom monitoring and when to seek medical attention (see Sample letter).

² In acute care settings, Infection Control and Occupational/ Employee Health should also be involved.



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Education

 All cases should be provided disease information as well as information on prevention and control measures including period of communicability and avoiding contact with susceptible individuals.

Exclusion and Isolation

Exclusion and isolation of cases should be implemented as outlined in Table 2.

Table 2. Exclusion Requirements for Cases

Who	Exclusion Requirements	Timeframe
Cases (including confirmed,	Exclude from childcare, school,	For at least 5 days from parotitis
clinical and suspect).3	post-secondary institutions, and	onset. This should be extended to
	workplaces.	9 days if the case remains
		symptomatic.
	Avoid contact with susceptible	
	people.	
Health Care Workers	Cases should be excluded from	For at least 5 days from parotitis
(HCWs) who are cases	work.	onset. This should be extended to
(including confirmed,		9 days if the case remains
clinical and suspect). ³		symptomatic.
Note – Advise case to	Cases who work with vulnerable	For 9 days from parotitis onset.
immediately notify	patients (i.e.,	
Occupational Health and/or	immunocompromised).	
Infection Control for the		
facility in which they work.		
Cases in the hospital or	The case should be on droplet	For at least 5 days from parotitis
other health care facility.	precautions.	onset. This should be extended to
		9 days if the case remains
		symptomatic.

Immunization

Ensure the client's entire immunization status is up-to-date once they have recovered.4

Treatment

- Supportive therapy as there is no specific treatment for mumps.
- Individuals with severe central nervous system involvement may require hospitalization.

⁴ Life-long immunity is expected following natural infection with mumps.



³ The exclusion of epidemiologically-linked contacts with symptoms can be discontinued before five days if laboratory results rule out a diagnosis of mumps.

II. Contacts/Contact Investigation

Table 3. Contact Definition (Public Health Agency of Canada, 2009)

Definition of Close Contact

Contacts of confirmed cases are defined as any of the following during the infectious period (approximately 7 days before to 5 days after symptom onset):

- household contacts of a case;
- persons who sleep in the same room as the case;
- direct contact with the oral/nasal secretions of a case (e.g., face-to-face contact
 where droplet contact may occur, sharing cigarettes/drinking
 glasses/food/cosmetics (lip gloss), kissing on the mouth, children and staff in
 child care and nursery school facilities, etc.);
- children and staff in child care and school facilities;
- HCWs who have unprotected face-to-face interaction (within 1 metre) to an infectious mumps case in the facility.

Definition of Susceptible Contacts

- Those born in 1970 or later who have not received two doses of mumpscontaining vaccine (at least four weeks apart) after their first birthday AND
 - who have not had laboratory confirmed mumps OR
 - o who do not have documented immunity due to mumps illness.

Serological screening to identify susceptible contacts is impractical and unnecessary, since there are no additional risks of immunizing those already immune.

In Canada, it is assumed that people who were born before 1970 are generally considered immune due to natural immunity.

Public Health Interventions

Assessment

Assess for signs and symptoms and immunization history.

Communication

• Identifiable contacts should, at a minimum, be provided with a letter that includes all details as outlined in education.

Education

- All contacts of confirmed cases should be educated about mumps including the signs and symptoms, period of communicability and measures to prevent transmission of respiratory viruses – handwashing, not sharing water bottles, etc.
- The risk of exposure should also be communicated to all students and parents and other contacts.



• Individuals should be advised to visit one's health-care provider should any symptoms develop.

Exclusion and Immunization

- Exclusion of susceptible contacts that meet the criteria in <u>Table 3</u> is outlined in <u>Table 4</u>.
- If the contact develops symptoms compatible with mumps, exclusion criteria for cases should be applied.

Table 4. Exclusion and Immunization Requirements for Contacts

Non-HCW Contacts who	Required Immunizations	Exclusion	Timeframe	
are:		Requirements		
Immune.	None.	None.	None.	
Susceptible (in school,	As per Saskatchewan	None.	None.	
childcare or workplace	Immunization Manual. ⁵			
setting).				

The following additional requirements apply to Health Care Workers who are Contacts

- Advise the health-care worker to contact Occupational Health and/or Infection Control for the facility in which they work.
- Public Health will notify Occupational Health and/or Infection Control that contacts to mumps have been identified in their facility. Personal details of the contacts will not be disclosed.
- Provide information on mumps disease and its symptoms.
- Assess immunization status.

Table 5. Exclusion and Immunization Requirements for Contacts who are Health Care Workers

History of Immunization	Required	Exclusion Requirements
	Immunizations	
Documented 2 doses of	None.	None.
mumps-containing		
vaccine.		
Documented 1 dose of	Provide second dose	Return to work immediately.
mumps-containing	of mumps-containing	
vaccine.	vaccine.	



⁵ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

History of Immunization	Required Immunizations	Exclusion Requirements
Undocumented immunization history.	1. Draw blood for mumps IgG serology. 2. Provide a dose of mumps-containing vaccine (after serology taken).	While waiting for serology results, exclude case from work for period of communicability, which starts on day 10 after exposure where exposure is day 1: a. If IgG positive, then consider immune and can return to work; consider a second dose of MMR for adequate measles and rubella protection. b. If IgG negative, then consider susceptible, provide a second dose of mumps-containing vaccine 28 days after the first and exclude from work on day
		10 after first exposure until day 26 after last exposure.

Testing

Attempt to confirm diagnosis in any contacts that develop symptoms consistent with mumps.⁶

Prophylaxis/Immunization

Although immunization with live virus mumps-containing vaccine has not been demonstrated to be effective in preventing infection after exposure, the following still applies:

Immunization of <u>susceptible contacts</u> with mumps-containing vaccine, recognizing
that immunization after exposure may not prevent disease if the individual is already
infected. See <u>Table 4</u>. <u>Exclusion and Immunization Requirements for Contacts</u>.
 Serological screening to identify susceptible contacts is impractical and unnecessary,
since there is no risk to those already immune.

⁶ This recommendation is applicable when sporadic cases are occurring. Recommendations for testing during an outbreak should be discussed with the MHO.



III. Environment

Child Care Centre/Schools Control Measures

Strict enforcement of infection control measures. Refer to the Saskatchewan Ministry of Health Infection Control Manual for Child Care Facilities.⁷

Health Facilities Control Measures

Strict enforcement of infection control measures. Refer to your Regional Infection Control Manual.

Cases should be on isolation and in a private room for at least 5 days from parotitis onset. This should be extended to 9 days if the case remains symptomatic. Refer to <u>Table 2</u> (Exclusion Requirements for Cases) and <u>Table 5</u> (Exclusion and Immunization Requirements for Contacts who are Health Care Workers).

IV. Epidemic Measures

The resources required for contact tracing and the management of contacts may put significant demands on public health and laboratory capacity.

- Logistics for providing immunization to susceptible contacts, including prioritization of vaccine supply, should be carefully considered.
- Serological screening to identify susceptible individuals is impractical and unnecessary, since there is no risk to those already immune.

When determining means to control outbreaks, exclusion of susceptible students from affected schools, thought to be at risk of transmission, should be considered. Excluded students can be readmitted following immunization. Immunization is not known to prevent mumps in those already exposed, but will protect against future exposures if the individual has had time to mount an immune response. Those who continue to be unimmunized due to medical, religious, or other reasons should be excluded until at least 26 days after the onset of parotitis in the last person with mumps in the affected school.

In the event of an outbreak, promote awareness in the community affected by the outbreak and among healthcare personnel:

- share information about the settings within which transmission is occurring;
- transmission patterns among fully vaccinated populations;

⁷ http://www.saskatchewan.ca/live/births-deaths-marriages-and-divorces/starting-a-family/early-learning-and-child-care/child-care



Respiratory and Direct Contact

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remind healthcare personnel to not rule out mumps on the assumption that
individuals have evidence of mumps immunity because outbreaks have occurred in
highly vaccinated populations in high transmission settings, including school settings
(e.g., elementary school, middle school, high school, and college students) (Centers
for Disease Prevention and Control, 2018).

Prevention Measures

Refer to the <u>Respiratory and Direct Contact Introduction and General Considerations</u> section of the manual that highlights topics for client education that should be considered as well as provides information on high-risk groups and activities.

Immunization

Immunize infants, children, and adults according to the recommended schedule. Refer to the Saskatchewan Immunization Manual.⁸

Education

- Education should be provided regarding respiratory etiquette and measures to prevent transmission of mumps.
- Educate the public about the disease and the need for active immunization with a mumps-containing vaccine. Immunization information fact sheets can be used to guide discussion.



⁸ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

Respiratory and Direct Contact

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Revisions

Date	Change
September 2018	 Updated to align with Panorama configuration Clarified the purpose for notification of cases to public health Incorporated an Epidemiology and Occurrence section with Canadian information and included Saskatchewan Immunization program history from Sask Immunization Manual to provide context. Updated period of communicability to remove outer limit of 14 days following parotitis. Rearranged and updated the style into the new format of the Manual. Added information into Epidemic section regarding transmission among fully vaccinated individuals. References reaffirmed or updated as necessary.



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Panorama QA complete: ☐Yes	□No	Please complete all sections. Panorama	Client ID:
Initials:		Panorama Invest	gation ID:

A) CLIENT INFORMATION			LHN -> SUBJE	CT -> CLIENT	DETAILS -> PERSONAL INFORMATION	
Last Name:		First Name: and Middle Name:		Alternate I	Name (Goes by):	
		Health Card Province: Health Card Number (PHN): 		Preferred Communication Method: (specify - i.e. home phone, text): Email Address: Work Personal		
Place of Employment/School:		Gender: □ Male	□ Female		Other	
Alternate Contact: Relationship: Alt. Contact phone:		Mailing (Postal address): Street Address or FN Communit Address at time of infection if n	ty (Primary Hon	ne):	porary □ Legal Land Description	
Disease Summary Classification:	Date	Classification:	Date		LAB TEST INFORMATION:	
□ Confirmed	YYYY / MM / DD	☐ Contact	YYYY / MM	/ DD	Date specimen collected: YYYY / MM / DD	
□ Does Not Meet Case	YYYY / MM / DD	□ Not a Contact	YYYY / MM	/ DD	Specimen type:	
☐ Person Under Investigation	YYYY / MM / DD	☐ Person Under Investigation	YYYY / MM / DD		□ Blood	
□ Probable	YYYY / MM / DD				□ Urine □ Stool	
Disposition: FOLLOW UP: In progress YYYY / MM / DD Complete YYYY / MM / DD Incomplete - Declined YYYY / MM / DD Referred - Out of province YYYY / MM / DD Incomplete - Unable to locate YYYY / MM / DD (specify where) REPORTING NOTIFICATION Name of Attending Physician or Nurse: Date Received (Public Health): YYYY / MM / DD					MM / DD MM / DD	
Type of Reporting Source: ☐ Hea	alth Care Facility	ab Report	oner □Phy	rsician [□ Other	

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Please complete all sections.

Panorama Client ID:	
Panorama Investigation ID:	

C) SIGNS & SYMPTOMS (Bold text = par			T		1	IGATION->SIGNS & SYMPTOMS
Description	No	Yes – Date of onset	Description		No	Yes - Date of onset
Abortion - spontaneous (miscarriage)		YYYY / MM / DD	Lab - platel	et count low		YYYY / MM / DD
Coryza or rhinitis		YYYY / MM / DD	Lethargy (fa	atigue, drowsiness, weakness, etc)		YYYY / MM / DD
Cough		YYYY / MM / DD	Meningitis -	- aseptic		YYYY / MM / DD
Encephalitis		YYYY / MM / DD	Orchitis (inf	lamed testicle)		YYYY / MM / DD
Hearing loss		YYYY / MM / DD	Pain - saliva	ary glands		YYYY / MM / DD
Infection - upper respiratory tract		YYYY / MM / DD	Parotid gla	nd - inflammation (parotitis)		YYYY / MM / DD
Other S/S						
D) INCUBATION AND COMMUNICABILIT	Υ			LHN-> INVESTIGATIO	N->INC	UBATION & COMMUNICABILITY
Incubation for Case (period for acquisiti Earliest Possible Exposure Date: YYYY / Exposure Calculation details: Communicability for Case (period for tra	MM /	on):		Latest Possible Exposure Date:		
Earliest Possible Communicability Date	: YYY	Y / MM / DD		Latest Possible Communicability	y Date:	YYYY / MM / DD
Communicability Calculation Details:						
E) RISK FACTORS						LHN-> SUBJECT->RISK FACTORS
DESCRIPTION		Start date Yes	N, NA, U	Add'l Info		
Contact - At risk population (international or immigrants)	al travell					
Contact to a known case (Add'l Info)		YYYY / MM/DD				
Immunocompromised - Related to under disease or treatment	rlying					
Occupation - Health Care Worker - IOM	Risk Fact	or TE				
Risk Behaviour - Sharing personal items water bottles)	(cigarett	es, TE				
Special Population - Attends childcare		TE				
Special Population - Attends school		TE				
Special Population - Lives in a communa	l setting	TE				
Special Population - Post secondary eduinstitution	cation	TE				
Special Population - Pregnancy						
Travel - Outside of Canada (Add'l Info)		YYYY / MM/DD AE				
Travel - Outside of Saskatchewan, but w Canada (Add'l Info)	ithin	YYYY / MM/DD AE				
F) IMMUNIZATION HISTORY INTERPRET	ATION S	SUMMARY	LHN ->	INVESTIGATION-> IMMUNIZATION	HISTO	RY INTERPRETATION SUMMARY
Interpretation Date: YYYY /	MM /	DD				
Interpretation of Disease Immunity:	□ Dise	ease Case - Fully immu	nized (for age)	Disease Case - Partiall	y immu	nized
☐ Disease Case – Unimmunized	□ Dise	ease Case - Unclear imi	munization his	tory Valid doses received:	D	oses needed:
Reason: ☐ Previous disease ☐ Interpretation of history by investigat	or	☐ Previous resp	oonder/Previo	us history of immunity	□ Dat	e Of Birth

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Please complete all sections.

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Panorama Investigation ID:	

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			Exclusion: Investigator name		
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			□ School YYYY / MM / DD	☐ Daycare YYYY	/ MM / DD
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n ti ie c C e e i e i e i e i e i e i e i e i e i	ion (see Investigator Nent Management) name Control Cont/Assess'd for Cont g: Investigato Investigato Intervention subtype recovering YYYY / Nert YY	ion (see Investigator Notes) ent Management) name Control cont/Assess'd for Contacts g: Investigator name I measures n provided Intervention Comme	ion (see Investigator Notes) ion (s	Disease-specific immunization re Disease-specific immunization re Disease-specific immunization gin Solation: Facility solation YYYY / MM / DD Home isolation YYYY / MM / DD Disease-specific immunization gin Solation: Facility isolation YYYY / MM / DD Home isolation YYYY / MM / DD Disease-specific immunization gin Facility isolation YYYY / MM / DD Disease-specific immunization gin Teating: Quarantine: Quarantine: Quarantine: Quarantine: Quarantine: Quarantine: Quarantine: Disease-specific immunization YYYY / MM / DD Disease-specific immunization YYYY / MM	lagement

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Please complete all sections.

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Panorama Investigation ID:	

Exposure Name:				
acquisition Start Y	YYYY / MM / DD to Acqu	uisition End: YYYY / MM / DD		
ocation Name:				
Setting Type				-
□ Travel	Health care setting	2 Public facilities 2 Recrea	ational facilities	☐ Most likely source
Transmission	Events	LHN -> INVESTIGATION-> EXPOSURE SUMMARY ->	> TRANSMISSIO	N EVENT SUMMARY -> QUICK EN
Transmission Event ID	Exposure Name	Setting type (Consider the following settings for TE; if >1 select "multiple settings" in Panorama)	Date/Tim	e # of contacts
		□Congregate/Communal living □Health Care setting	YYYY / M	IM / DD
		☐ Type of community contact ☐ Household Exposure	to	
		☐ Public facilities	YYYY / M	IM / DD
	-	☐ Congregate/Communal living ☐ Health Care setting		IM / DD
		☐ Type of community contact ☐ Household Exposure	to	
		☐ Public facilities	YYYY / N	1M / DD
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		☐ Type of community contact ☐ Household Exposure	to	
		□ Public facilities	YYYY / N	1M / DD
		□Congregate/Communal living □Health Care setting		IM / DD
		☐ Type of community contact ☐ Household Exposure	to	
		☐ Public facilities☐	YYYY / N	1M / DD
		□ Congregate/Communal living □ Health Care setting	,	IM / DD
		☐ Type of community contact ☐ Household Exposure	to	
		☐ Public facilities☐	YYYY / N	1M / DD
		□ Congregate/Communal living □ Health Care setting	YYYY / M	IM / DD
		☐ Type of community contact ☐ Household Exposure	to	
		☐ Public facilities☐	YYYY / N	1M / DD
		☐ Multiple Settings	YYYY / M	IM / DD
	Mumps Contacts – Inv ID#		to	
			YYYY / N	1M / DD
) TOTAL NUMBER		LIDE CLINARARDY > TRANSPARICCIONI EVENT CLINARARDY > TE	TANDEDI ININ 2 I	AND STRANGER (AND IN AND IN AN
LITTE		URE SUMMARY -> TRANSMISSION EVENT SUMMARY -> TE		•
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Initial Report				Date initial report completed:
completed by:				YYYY / MM / DD

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Respiratory and Direct Contact

Neonatal Group B Streptococcus

Date Reviewed: August, 2011 Section: 2-120 Page 1 of 6

Notification Timeline:

From Lab/Practitioner to Public Health: Immediate.

From Public Health to Ministry of Health: Within 2 weeks.

Public Health Follow-up Timeline: Within 72 hours.

Information

Case Definition (Public Health Agency of Canada, May 2008)

Confirmed Case	Clinical illness ¹ in an infant less than 1 month of age with laboratory confirmation of infection:
	• isolation of group B <i>Streptococcus</i> (<i>Streptococcus agalactiae</i>) from a normally sterile site (such as blood or cerebrospinal fluid) OR
	• demonstration of group B <i>Streptococcus</i> DNA in a normally sterile site.
Probable Case	Clinical illness ¹ in an infant less than 1 month of age with laboratory
	confirmation of infection:
	• detection of group B <i>Streptococcus</i> antigen in a normally sterile site.

There are two forms of clinical illness; <u>early onset</u> disease (1-7 days), characterized by sepsis, respiratory distress, apnea, shock, pneumonia, and meningitis; and <u>late onset</u> (7 days to 1 month), characterized by sepsis and meningitis.

Even though the case definition is for infants < 1 month, follow-up of infants between 1 to 3 months may be considered.

Causative Agent

Streptococcus agalactiae, group B Streptococcus (GBS).

Symptoms

There are 2 distinct forms:

<u>Early-onset disease</u> – lethargy, poor feeding, jaundice, fever, grunting respirations and other signs of respiratory distress, pallor and hypotension. Respiratory distress is usually present at or within a few hours after birth. Diagnosed as sepsis, pneumonia and less frequently meningitis, osteomyelitis or septic arthritis. It is acquired in utero or during delivery; low-birth weight, premature infants are more susceptible.



Respiratory and Direct Contact

Neonatal Group B Streptococcus

Date Reviewed: August, 2011 Section: 2-120 Page 2 of 6

• <u>Late-onset disease</u> – lethargy, poor feeding, irritability and fever. Diagnosed as sepsis and meningitis and, less frequently, bone and joint infections.

Incubation Period

- Early-onset 1 to 7 days.
- Late-onset 7 days to 1 month.

Reservoir/Source

Humans. Heymann (2008) says about 10-30% of pregnant women harbour group B streptococci in the genital tract, and about 1-2% of their offspring may develop symptomatic infection.

Mode of Transmission

- Early-onset is acquired in utero or during delivery.
- Late-onset is acquired through person-to-person contact and occurs in full-term infants.
- Nosocomial transmission may occur if appropriate infection prevention and control measures are not taken.

Risk Factors/Risk Group

The American Academy of Pediatrics (2009) indicates that the risk for GBS is increased in the following:

- maternal age younger than 20 years;
- previous baby with GBS disease;
- urinary tract infection due to GBS during the pregnancy;
- GBS carriage late in pregnancy;
- maternal temperature of 38 degrees Celsius or higher during labour;
- rupture of membranes 18 hours or more before delivery;
- preterm infants born at less than 37 weeks gestation.

Period of Communicability

The administration of intravenous antibiotics (generally penicillin) to women colonized with group B streptococci at the onset and throughout labour interrupts transmission to newborn infants, decreasing infection and mortality. (This is consistent with Society of Obstetricians and Gynaecologists of Canada Guidelines, Jan 2007.)



Neonatal Group B Streptococcus

Date Reviewed: August, 2011 Section: 2-120 Page 3 of 6

Specimen Collection and Transport

- Take a vaginal and rectal swab for culture at 35-37 weeks gestation. Cultures
 collected earlier do not accurately predict whether a woman will have GBS at
 delivery.
- For diagnosis in a neonate, culture of sterile fluid (blood or CSF) is required.

Methods of Control/Role of Investigator

Prevention and Education

There are limited effective primary prevention strategies for the early onset form of this disease. Refer to the <u>Respiratory and Direct Contact Introduction and General</u>
<u>Considerations</u> section of the manual that highlights topics for client education that should be considered as well as provides information on high-risk groups and activities.

Prevention of the late onset form of this disease is best accommodated via handwashing.

Studies that looked at screening versus risk-based approach found that risk of early-onset disease was significantly lower among the infants of screened women compared to those in the risk-based approach. As such, pregnant women are to be tested late in pregnancy (35-37 weeks) to determine whether or not they are positive for GBS, so they can be treated during labour.

Intrapartum therapy of women with positive screenings and certain other risk factors has been found to be the most effective in preventing neonatal GBS disease (Dobson & Money, 2004).

Immunization

Immunization strategies have been researched for many years, but currently, there is no vaccine for group B *Streptococcus*.

Education

- Prenatal education of high risk mothers about screening and intrapartum treatment.
- Physicians should be aware of the need for testing of pregnant women and appropriate treatment of the women who screen positive.



Neonatal Group B Streptococcus

Date Reviewed: August, 2011 Section: 2-120 Page 4 of 6

Management

I. Case

History

See Risk Factors/Risk Groups above.

Immunization

Not applicable.

Treatment/Supportive Therapy

- Treatment choices are governed by the most recent guidelines. The public health practitioner should direct any questions regarding the current treatment protocols to the physician or Medical Health Officer. See Appendix H Sources for Clinical Treatment Guidelines.
- See <u>Attachment Recommendations for Prevention and Management of Neonatal</u> Group B *Streptococcus*.

Exclusion

Not applicable.

Referrals

15-30% of survivors of group B streptococcal meningitis have permanent neurologic sequelae (hearing/vision loss or learning disabilities). Referral by physician to appropriate disciplines.

II. Contacts/Contact Investigation Contact Definition

No contact tracing is required.

Testing

Test only if symptomatic.

Prophylaxis/Immunization

Not applicable.



Neonatal Group B Streptococcus

Date Reviewed: August, 2011 Section: 2-120 Page 5 of 6

Exclusion

Not applicable.

III. Environment

Child Care Centres/Institutional Control Measures

Neonatal nurseries – hand hygiene is the best way to prevent the spread to other infants (American Academy of Pediatrics, 2009).

Epidemic Measures

- Contact precautions and cohorting of ill and colonized infants is recommended during an outbreak.
- Epidemiologic evaluation of late-onset cases in a special care nursery may be required to determine a common source and prevent spread to others.



Neonatal Group B Streptococcus

Date Reviewed: August, 2011 Section: 2-120 Page 6 of 6

References

American Academy of Pediatrics. (2009). *Red book: 2009 Report of the Committee on Infectious Diseases* (28th ed.). Elk Grove Village, IL: Author.

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Neonatal Group B Streptococcus

Attachment – Recommendations for Prevention and Management of Neonatal Group B Streptococcus

Date Reviewed: August, 2011 Section: 2-120 Page 1 of 1

The following are recommendations for pregnant women (Society of Obstetricians and Gynaecologists of Canada [SOGC], 2004):

- 1. Offer all women screening for group B *streptococcus* (GBS) disease at 35 to 37 weeks' gestation with culture done from one swab first to the vagina then to the rectal area.
- 2. Treat the following women intrapartum at time of labour or rupture of membranes with IV antibiotics:
 - all women positive by GBS culture screening done at 35 to 37 weeks;
 - any women with an infant previously infected with GBS;
 - any women with documented GBS bacteriuria (regardless of level of colony-forming units per mL) in this pregnancy.
- 3. Treat women at less than 37 weeks' gestation with IV antibiotics unless there has been a negative GBS vaginal/rectal swab culture within 5 weeks.
- 4. Treat women with intrapartum fever with IV antibiotics (i.e., chorioamnionitis must be treated, but broader spectrum antibiotics would be advised).
- 5. If a woman is GBS-positive by culture screening or by history of bacteriuria, with prelabour rupture of membranes at term, treat with GBS antibiotic prophylaxis and initiate induction of labour with IV oxytocin.
- 6. If GBS culture result is unknown and the woman has ruptured membranes at term for greater than 18 hours, treat with GBS antibiotic prophylaxis.

Neonatal Management (SOGC, 2004)

- 1. Infants delivered by women who have received intrapartum antibiotics at least 4 hours before delivery, do not need a septic workup. These infants should be observed in hospital for the first 24 hours for signs of infection, but do not need additional therapy or investigations.
- 2. Infants who appear well despite their mothers being GBS colonized and not receiving adequate antibiotics (< 4 hours) should be observed for 48 hours and evaluated or treated if signs of sepsis develop.
- 3. Infants of mothers with chorioamnionitis should undergo a diagnostic evaluation for sepsis and be treated with antibiotics. (Sepsis workup includes a complete blood-cell count and differential, blood culture, and chest radiograph, including a lumbar puncture if feasible.)



Notification Timeline:

From Lab/Practitioner to Public Health: Immediate.

From Public Health to Ministry of Health: Within 2 weeks.

Public Health Follow-up Timeline: Immediate.

Public Health Purpose for Notification of Pertussis (adapted from British Columbia Center for Disease Control [2017])

- To minimize mortality and serious morbidity from pertussis in young children through contact tracing;
- To track epidemiology trends of pertussis in Saskatchewan including risk factors and distribution;
- To identify locations where increased transmission of pertussis may be occurring in order to inform other interventions;
- To monitor the effectiveness of prevention and control measures;
- To plan expansion or introduction of future immunization programs;
- To provide timely clinical care including diagnosis and treatment using current, evidence-based guidelines;
- To make timely and evidence informed actions on outbreaks; and
- To inform the public and medical community about pertussis.

Surveillance Case Definition¹ (Public Health Agency of Canada, May 2008)

Surveillance Cas	e Definition (Public Health Agency of Canada, May 2008)
Confirmed Case	Laboratory confirmation of infection:
	• isolation of <i>Bordetella pertussis</i> (e.g. from a culture) from an appropriate clinical specimen
	OR
	• detection of <i>B. pertussis</i> DNA (e.g NAAT or PCR) from an appropriate
	clinical specimen AND one or more of the following:
	 cough lasting 2 weeks or longer
	 paroxysmal cough of any duration
	cough with inspiratory "whoop"
	 cough ending in vomiting or gagging, or associated with apnea.
	OR

¹ Surveillance case definitions ensure uniform reporting to allow comparability of surveillance data. The definition is not intended to be used for clinical or laboratory diagnosis or management of cases.



	Epidemiologic link to a laboratory-confirmed case AND one or more of
	the following for which there is no other known cause:
	 paroxysmal cough of any duration
	cough with inspiratory "whoop"
	 cough ending in vomiting or gagging, or associated with apnea.
Probable Case	Cough lasting 2 weeks or longer in the absence of appropriate
	laboratory tests and not epidemiologically linked to a laboratory-
	confirmed case AND one or more of the following, with no other known
	cause:
	paroxysmal cough of any duration
	cough with inspiratory "whoop"
	• cough ending in vomiting or gagging, or associated with apnea.
Suspect Case	One or more of the following, with no other known cause:
	paroxysmal cough of any duration
	cough with inspiratory "whoop"
	• cough ending in vomiting or gagging, or associated with apnea.
Public health follo	w-up of probable and suspect cases should be considered based on the
epidemiology of p	pertussis in the community and the involvement of vulnerable populations.

Epidemiology and Occurrence

Pertussis is a cyclical disease which peaks at 4 to 5 year intervals (see Figure 1). Infants are the most vulnerable and are often infected by older siblings, parents or caregivers. Figure 2 shows the rates of pertussis in infants relative to children 1-19 years of age.

- An adolescent pertussis vaccine (Tdap) was introduced to students in Grade 8 in 2003. This widened the gap in the rate of illness in these age groups; the gap was narrowed following the implementation of a Tdap program for all adults in 2010, especially parents and caregivers of infants, in an effort to reduce the risk to these vulnerable infants.
- In October 2017, it was recommended that all pregnant women be offered Tdap in the third trimester irrespective of prior Tdap receipt.
- The waning of immunity conferred by pertussis vaccine in infancy was reflected in an increase of incidence in 2015 to 2017, mainly among the 10-14 year old cohort.



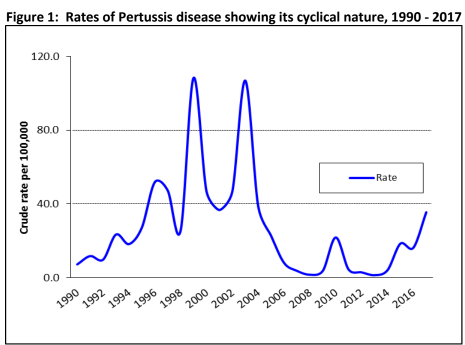


Figure 2: Pertussis Rates in Infants versus Children Aged 1-19 by year, 2002-2017 Cocooning Grade 8 TDaP 400.0 introduced. Infant Tdap offered booster to all caregivers offered introduced late Tdap, late 2010. pregnant 2003-2004 women at 300.0 27 weeks or later, Oct Rate per 100,000 2017 200.0 100.0 Infants 1-19 yrs 0.0

2010

2011

2012

2013

2014

2017

2003

2004

2005

2006

2007

2008

2009

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Additional Background Information

Causative Agent

Bordetella pertussis.

Symptoms

<u>Catarrhal Stage</u>: starts with mild respiratory symptoms of cough, rhinorrhea and possible fever.

<u>Paroxysmal Stage</u>: paroxysms of cough characterized by inspiratory whoop and vomiting after cough.

Convalescent Stage: gradual recovery with cough lasting 1-2 months or longer.

Infants less than 6 months can have an atypical presentation with short catarrhal stage, gagging, gasping or apnea as prominent early manifestations, absence of whoop and prolonged convalescence.

Complications among infants include pneumonia, seizures, encephalopathy and death. Complications in adolescents and adults include syncope, sleep disturbance, incontinence, rib fracture and pneumonia.

Reservoir/Source

Humans.

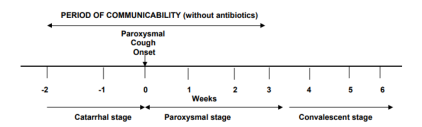
Incubation Period

6-20 days (average 9-10 days).

Period of Communicability

- Highly communicable in the early catarrhal stage and the beginning of the paroxysmal stage (first 2 weeks).
- Communicability decreases after the catarrhal and paroxysmal stages and becomes negligible 3 weeks after onset of symptoms.
- Case is no longer contagious after completing 5 days of treatment.





Mode of Transmission

Person-to-person by direct contact with discharges from respiratory secretions via aerosolized droplet.

Specimen Collection and Transport

Nasopharyngeal swab in Regan Lowe transport medium. See the Saskatchewan Disease Control Laboratory Compendium for further details at https://rrpl-testviewer.ehealthsask.ca/

Public Health Investigation

I. Case

Refer to Attachment – Pertussis Data Collection Worksheet to assist.

History

- Key elements to inquire about include:
 - Immunization history of case.
 - Onset of illness and treatment (with what and when) to determine incubation period and period of communicability which helps to identify the possible source and contacts to be followed.
 - Travel history may be of significance in contact tracing.
 - Underlying medical conditions and severity of illness (e.g. if hospitalization was required).
 - o Current health status of household contacts (are contacts symptomatic?).
 - Identify contacts (refer to <u>Table 2 Definitions of Contacts</u>) paying particular attention to vulnerable contacts (infants and women in the third trimester).
 - Occupational considerations exist for health care settings see <u>Special</u>
 Considerations for Cases and Contacts in the Health Care Setting



Public Health Interventions

Assessment

 Assess for contacts paying particular attention to vulnerable contacts as per Table 2.

Communication

 Letters can be sent to classrooms and other group settings where individual contact tracing is not required (i.e. involving school age and adults where there are no vulnerable contacts) to inform them of the exposure, symptom monitoring and when to seek medical attention (see Sample letter).

Education

 All cases should be provided disease information as well as information on prevention and control measures including period of communicability and avoiding contact with vulnerable individuals.

Exclusion

- There is limited evidence supporting the use of exclusion; by the time a
 person is diagnosed with pertussis, they have likely exposed most of their
 contacts. Therefore exclusion is no longer recommended in most
 situations; however the consensus was to use exclusion if there are
 vulnerable individuals involved (see Table 2 Definitions of Contacts).
 - Cases should be excluded from school or daycare/preschool where there
 are vulnerable persons, for 5 days after they start the medication, or 21
 days from onset of cough if untreated. If there are no vulnerable persons
 in the school or day care, the case can return to school or
 daycare/preschool as soon as he/she feels well enough to do so.
 - 2. Adult cases who have close contact with vulnerable persons at work should be excluded from work for 5 days after they start the medications, or 21 days from onset of cough if untreated. If there are no vulnerable persons in the workplace, the case can return to work as soon as he/she feels well enough to do so.
- When exclusion is recommended, it should continue for 5 days after they start the appropriate medication, or 21 days from onset of cough if untreated or until test results come back negative for pertussis.
- Exclusion is not recommended in most other situations as there is limited
 evidence to support it since a person who has been diagnosed with pertussis
 may have likely exposed most of their contacts. Please refer to Special Considerations for Cases and Contacts in the Health Care Setting below for
 additional recommendations.



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Immunization

 Case follow-up should be used as an opportunity to recommend immunizations they are eligible for as per the Saskatchewan Immunization Manual. Infants and children who have recovered from pertussis should complete their pertussis immunization series, as natural infection does not confer life-long immunity (American Academy of Pediatrics, 2015).

Treatment

Treatment recommendations have been summarized in <u>Attachment –</u>
 Pertussis Treatment and Chemoprophylaxis Guidelines.

Who Should be Treated

Treatment is recommended for all individuals that are laboratory confirmed, clinically diagnosed and epidemiologically linked to another case, or probable cases (clinically diagnosed) during an outbreak.

- 1. **All cases** laboratory confirmed **OR** clinically diagnosed and epidemiologically linked to another case **OR** clinically diagnosed during an outbreak.
- 2. **All symptomatic household contacts** the assumption is that these symptomatic people will also have pertussis. *Sometimes symptomatic household contacts may be reluctant to take antibiotics without a confirmed diagnosis. If there are no vulnerable persons in the household, it is acceptable to wait for results of testing.*
- 3. All other community contacts who are symptomatic should **not** be assumed to have pertussis unless clinical symptoms are very predictive, but should be assessed, tested and treated appropriately.



II. Contacts/Contact Investigation

Table 2 Definitions of Cor	Table 2. Definitions of Contacts						
Close Contact	 Individuals that have shared respiratory secretions (e.g., kissing) or shared the same confined air space for more than an hour, or have had face-to-face exposure for more than 5 minutes. 						
Vulnerable Contact	 Children less than 1 year of age, because they have a higher rate of mortality from pertussis infection. Pregnant women in the third trimester, because if infectious at the time of birth they may pass the infection to their newborn. 						
Household Contact	 Household contact is living in the same household as the case including family² day care setting. 						
Occupational Contact	 Contact of Health Care Workers (HCW's) oral or nasal mucosa with infected secretions from the pertussis case. OR Sharing the same confined air space (within 2 metres) for more than an hour with the pertussis case, without implementing droplet precautions. OR Having had face-to-face exposure for more than 						
	5 minutes with a pertussis case without implementing droplet precautions.						

Public Health Interventions

Assessment

- Assess for symptoms.
- Assess for vulnerable individuals in their household. Recommend chemoprophylaxis as appropriate.

Communication

 Individual follow-up of contacts in in larger daycares, classrooms, schools, teams, workplaces, etc., is generally <u>not recommended</u>. These individuals should be informed by letter from public health, advising them to see their physician if they develop symptoms. These persons, if they become symptomatic, should not be assumed to have pertussis but should be assessed, tested and treated appropriately.

² Family day care refers to day cares that are run out of an individual's home to a limited number of children (*The Child Care Act, 2003*).



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Education

 All contacts should be provided disease information on symptom monitoring, prevention and control measures including avoiding contact with vulnerable individuals.

Exclusion

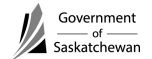
- Symptomatic family daycare contacts should be excluded from daycare
 where there are vulnerable persons, until they have completed 5 days of
 appropriate antibiotic or until test results come back negative for pertussis.
 In other words, if there are no vulnerable persons in the family day care, the
 symptomatic day care contact can return to day care as soon as he or she
 feels well enough to do so.
- **Symptomatic contacts** (non-household, non family-daycare) who have been assessed and tested but are not being treated until the test results are back, do not need to be excluded. They should be asked to **avoid close contact with vulnerable persons** until their diagnosis is established.

Immunization

- Immunization status of exposed individuals should be reviewed. Priority should be given to infants, children, and pregnant women in their third trimester.
- Accelerated immunization with the first dose at 6 weeks of age and the second and third doses using minimum intervals may be indicated in case of an outbreak in a defined community. See Saskatchewan Immunization Manual³ and discuss with Medical Health Officer.
- Immunizations should be completed for those whose schedules are incomplete.

Testing

 Non-immediate household and non-family day care contacts who are symptomatic should be assessed, tested and treated as appropriate.



³ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

Chemoprophylaxis

Chemoprophylaxis efficacy is related to early implementation and is **unlikely to be of benefit after 21 days** has elapsed since the first contact with a case. **Prophylaxis is generally not recommended for contacts in larger daycares, classrooms, schools, teams, workplaces, etc.** Contacts will be informed, usually by letter from public health, and advised to see their physician/nurse practitioner if they develop symptoms. The letter will inform these contacts that if they become symptomatic they should be assessed, tested and treated appropriately.

- See Attachment Pertussis Treatment and Chemoprophylaxis Guidelines.
- Chemoprophylaxis should be offered to the following contacts:
 - 1. **All symptomatic immediate household contacts** persons in a family day care setting are considered immediate household contacts. The assumption is that these symptomatic people will also have pertussis.
 - 2. **Symptomatic vulnerable persons** who have had "close contact" with a case should be started on antibiotics until their diagnosis is established.
 - 3. **Asymptomatic immediate household contacts**, including family-daycare attendees, where there is a vulnerable person in the household. The vulnerable person being ill does not eliminate the need for chemoprophylaxis of household contacts.
 - 4. Outside of the immediate household or family day care, offer prophylaxis only to asymptomatic vulnerable persons who have had "close contact" with a case.
 - 5. Non immediate-household and non family-daycare contacts who are symptomatic should not be assumed to have pertussis unless clinical symptoms are very predictive, but should be assessed, tested and treated appropriately.
- Chemoprophylaxis efficacy is related to early implementation and is unlikely to be of benefit after 21 days has elapsed since the first contact with a case.
- Chemoprophylaxis for all people in larger daycares, classrooms, schools, teams, workplaces, etc., is generally <u>not recommended</u>. These individuals should be informed by letter from public health, advising them to see their physician if they develop symptoms. These persons, if they become symptomatic, should not be assumed to have pertussis but should be assessed, tested and treated appropriately.



<u>Special Consideration for Cases and Contacts in the Health Care Setting</u> (Ontario Hospital Association, 2015)

Collaboration with Occupational Health/Employee Health is important in appropriate management of health care workers (HCWs). HCWs who do not provide direct patient care, such as housekeeping staff, may be managed as in the community setting. HCWs who do not provide direct patient care, such as housekeeping staff, may be managed as in the community setting. Community contacts who are health care workers should be managed as outlined below.

Prevention is always the primary goal and HCWs should protect themselves and their patients by being vaccinated as per the *Saskatchewan Immunization Manual*⁴ – Chapter 7: Immunization of Special Populations, Section 3.2 Health Care Workers. Status of vaccination with Tdap (tetanus, diphtheria, and acellular pertussis vaccine) should be evaluated for all HCW contacts.

The most effective control of transmission of pertussis in hospital settings includes isolation of the suspected or known case and use of droplet precautions. In addition, the following outlines appropriate management:

Management of Health Care Workers

- 1. HCWs who are considered **vulnerable contacts**⁵ should be offered chemoprophylaxis.
- 2. HCWs who are **confirmed cases** of pertussis:
 - Should be referred for appropriate antibiotic treatment.
 - Should be excluded from work until after 5 days of treatment or for 21 days from onset of cough if untreated.

[•] who may expose these vulnerable patient populations (e.g. hospitalized infants or pregnant women).



⁴ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

⁵ HCW vulnerable or high risk contacts include:

[•] pregnant women in their third trimester,

[•] household contact of infants under 12 months of age or a woman who is in her third trimester of pregnancy; OR

- 3. HCWs who are **symptomatic contacts** to pertussis case:
 - Should be referred for clinical management, which should include laboratory investigation (nasopharyngeal swab) and appropriate antibiotic treatment.
 - Should be excluded from work until after 5 days of treatment or for 21 days from onset of cough if untreated, or until swab comes back negative for pertussis. A surgical mask is not sufficient for protection of patients and other staff.
- 4. HCWs who are **asymptomatic contacts** to pertussis case:
 - Should be given chemoprophylaxis with an appropriate antibiotic if they are vulnerable or work or live with a vulnerable contact(s) (American Academy of Pediatrics, 2015).
 - Should be advised of early symptoms of pertussis and be put under surveillance by their employee health nurse.
 - Report development of symptoms to Occupational Health and Safety/Employee Health Department for an individual assessment.
 - Those with no history of an adult dose of Tdap vaccine should be given vaccine.
 - Exclusion of asymptomatic contacts is not indicated.

III. Environment

Child Care Centre/Schools Control Measures

Strict enforcement of infection control measures. Refer to the *Infection Control Manual for Child Care Facilities*. Notification of parents of children in either of these settings where a case has occurred is important. This can be accomplished via a letter sent through the school or daycare.

Chemoprophylaxis for all people in larger daycares, classrooms, schools, teams, workplaces, etc., is generally <u>not recommended</u>. They should be informed by letter from public health, and advised to see their physician if they develop symptoms. Review immunization histories of childcare attendees.

Health Facilities Control Measures

Strict enforcement of infection control measures. Refer to the Health Authority Infection Control Manual. Refer to <u>Special Considerations for Cases and Contacts in the Health Care Setting</u> for additional information.

⁶ http://publications.gov.sk.ca/documents/13/105320-infection-control-manual-child-care-centres.pdf



IV. Epidemic Measures

- Enhanced surveillance including details about immunization history of case and household contacts.
- Accelerated immunization with the first dose at 6 weeks of age and the second and third doses at 4 week intervals may be indicated at a community level.
- Immunizations should be completed for those whose schedule is incomplete.
- Additional measures may be instituted by the medical health officer to help contain the outbreak.
- As of October 2017, an enhanced outbreak measure is to provide pregnant women at 27 weeks gestation or later, irrespective of prior Tdap receipt, an additional dose of Tdap to offer protection to their newborn until they are eligible to be vaccinated.

Prevention Measures

Refer to the <u>Respiratory and Direct Contact Introduction and General Considerations</u> section of the manual that highlights topics for client education that should be considered as well as provides information on high-risk groups and activities.

Immunization

Immunize infants, children, pregnant women and adults according to the recommendations in the *Saskatchewan Immunization Manual*.

Education

- Education should be provided regarding respiratory etiquette and measures to prevent transmission of pertussis by practising good hand hygiene and not sharing drinking glasses or utensils.
- Educate the public about the disease and the need for active immunization. Immunization information fact sheets can be used to guide discussion.



Revisions

Date	Change
November 2018	Clarified which HCW require chemoprophylaxis.
September 2018	Updated to align with Panorama configuration.
	Updated Epidemiology and Occurrence section with 2017 data.
	Incorporated incubation and communicability graphic.
	Updated Special Considerations for Cases and Contacts in the
	Health Care Setting based on Ontario Hospital Association 2017
	updates.
	Updated purpose for notification based on BCCDC objectives of
	surveillance (2017).
September 2017	Clarified the purpose for notification of cases to public health.
	Incorporated an Epidemiology and Occurrence section to the
	chapter indicating timeframes of when changes were made to
	pertussis immunization program.
	Incorporated reference regarding when public health
	management should be considered for probable and suspect
	cases.
	Incorporated reference to outbreak measure of enhanced
	immunization of pregnant women in 3 rd trimester.
	Incorporated clarification on the use of chemoprophylaxis for
	health care workers.
	Rearranged and updated the style into the new format of the
	Manual.
	References reaffirmed or updated as necessary.



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Pertussis Data Collection Worksheet Panorama QA complete: ☐ Yes □No Panorama Client ID: Please complete all sections. Panorama Investigation ID: __ Initials: A) CLIENT INFORMATION LHN -> SUBJECT -> CLIENT DETAILS -> PERSONAL INFORMATION Last Name: First Name: and Middle Name: Alternate Name (Goes by): Preferred Communication Method: (specify - i.e. DOB: YYYY / MM / DD Age: ____ Health Card Province: ___ home phone, text): Health Card Number (PHN): Phone #: □ Primary Home: Email Address: □Work □Personal ☐ Mobile contact: ☐ Workplace: Place of Employment/School: Gender: ☐ Male ☐ Female Other □ Unknown Address Type: □ No fixed □ Postal Address □ Primary Home □ Temporary □ Legal Land Description Alternate Contact: _____ Mailing (Postal address): Relationship: Alt. Contact phone: ___ Street Address or FN Community (Primary Home): Address at time of infection if not the same: LHN-> SUBJECT SUMMARY-> RESPIRATORY & DIRECT CONTACT ENCOUNTER GROUP->CREATE INVESTIGATION B) INVESTIGATION INFORMATION Disease Summary Classification: Classification: LAB TEST INFORMATION: Date **CONTACT** Date CASE Date specimen collected: ☐ Confirmed YYYY / MM / DD □ Contact YYYY / MM / DD YYYY / MM / DD ☐ Does Not Meet Case YYYY / MM / DD ☐ Not a Contact YYYY / MM / DD Specimen type: Person Under Investigation YYYY / MM / DD ☐ Person Under Investigation YYYY / MM / DD □ Nasopharyngeal □ Throat ☐ Probable YYYY / MM / DD YYYY / MM / DD ☐ Suspect Disposition: **FOLLOW UP:** ☐ In progress YYYY / MM / DD ☐ Complete YYYY / MM / DD ☐ Not required YYYY / MM / DD ☐ Incomplete - Declined YYYY / MM / DD ☐ Incomplete – Lost contact YYYY / MM / DD ☐ Referred – Out of province YYYY / MM / DD ☐ Incomplete – Unable to locate YYYY / MM / DD (specify where) REPORTING NOTIFICATION Location: Name of Attending Physician or Nurse: Physician/Nurse Phone number: Date Received (Public Health): YYYY / MM / DD \square Lab Report ☐ Nurse Practitioner ☐ Physician □ Other___

September 1, 2018 Page 1 of 4

Pertussis Data Collection Worksheet

Please complete all sections.

Panorama Client ID:	
Panorama Investigation ID:	

C) SIGNS & SYMPTOMS (Bold text = part of case definition)

- 1	HN->	INV	FSTIG	ATION-	>SIGNS	R,	SYMPTOMS

Description	No	Yes - Date of onset	Description	No	Yes - Date of onset
Apnea		YYYY / MM / DD	Cough – paroxysmal		YYYY / MM / DD
Coryza or rhinitis		YYYY / MM / DD	Cough – with whoop		YYYY / MM / DD
Cough		YYYY / MM / DD	Cough > 2 weeks		YYYY / MM / DD
Cough – with apnea		YYYY / MM / DD	Gagging - infant		YYYY / MM / DD
Cough – with vomiting		YYYY / MM / DD	Gasping - infant		YYYY / MM / DD

Paroxysmal Cough Onset -2 -1 0 1 2 3 4 5 6 Weeks Catarrhal stage Paroxysmal stage Convalescent stage

D) INCUBATION AND COMMUNICABILITY

LHN-> INVESTIGATION->INCUBATION & COMMUNICABILITY

Incubation for Case (period for acquisition):

Earliest Possible Exposure Date: YYYY / MM / DD

Latest Possible Exposure Date: YYYY / MM / DD

 ${\it Exposure \ Calculation \ details:}$

Communicability for Case (period for transmission):

Earliest Possible Communicability Date: YYYY / MM / DD Latest Possible Communicability Date: YYYY / MM / DD

Communicability Calculation Details:

F) RISK FACTORS (RE followed by + impact the Immunization Forecaster)

LHN-> SUBJECT->RISK FACTORS

DESCRIPTION	Yes	N –No NA – not asked U - unknown	DESCRIPTION	Yes	N –No NA – not asked U - unknown
Special Population - Pregnancy	YYYY / MM / DD		Setting - Crowded living conditions (>1 person per room excluding bathrooms)		
Contact - Persons with similar symptoms	YYYY / MM / DD		Special Population - Lives in a communal setting		
Contact to a known case (Add'l Info)	YYYY / MM / DD		Travel - Outside of Canada (Add'l Info)	AE/TE YYYY / MM / DD	
Immunocompromised - Related to underlying disease or treatment			Travel - Outside of Saskatchewan, but within Canada (Add'l Info)	AE/TE YYYY / MM / DD	
Maternal Tdap not received between 27 weeks and 2 weeks prior to delivery (For infant cases <1 year)	YYYY / MM / DD				

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Pertussis Data Collection Worksheet

Please complete all sections.

Panorama Client ID:	
Panorama Investigation ID:	

F) IMMUNIZATIO	ON HISTORY INTERI	PRETATION SUM	MARY LHI	-> INVESTIGATION-> IMMUNIZ	ATION HISTORY INTERPRET	TATION SUMMARY
Interpretation Da	ate: YY	YY / MM / DD				
Interpretation of	Disease Immunity	: 🗆 IOM - Fu	ally immunized (for age)	☐ IOM - Partially i	immunized	
□ IOM – Unimm			nclear immunization history	Valid doses received:	Doses needed:	
Reason:	□ IOM - Inter	pretation of histo	ory by investigator			
G) TREATMENT				LHN -> INVESTIGATION	I-> MEDICATIONS->MEDICA	ATIONS SUMMARY
Medication (Pana	orama = Other Med	ds) :				-
Prescribed by:				Started on: YYYY / MM / D	D	
H) INTERVENTIO	N		LHN -> I	NVESTIGATION->TREATMENT &	INTERVENTIONS->INTERVE	ENTION SUMMARY
Intervention Type	e and Sub Type:					
Investigator name	ant or < 1 year of ag e	ge)	YYYY / MM / DD	Immunization: ☐ Eligible immunizations record ☐ Disease-specific immunization ☐ Disease-specific immunization ☐ Investigator name	on recommended YYYY /	/ MM / DD / MM / DD / MM / DD
Other Investigation	_	D		111100000000000000000000000000000000000		
☐ Investigator No Communication: ☐ Other communication of the communica	nication (see Invest	ee Document Ma	YYYY / MM / DD	Referral: Other (specify)	YYYY	/ MM / DD
	cument Manageme	ent)	YYYY / MM / DD	Investigator name		
General: Investig ☐ Disease-Info/P ☐ Disease-Info/P		for Contacts	YYYY/ MM / DD YYYY/ MM / DD	Testing: ☐ Laboratory testing recommendation investigator name	ended YYYY /	/ MM / DD
Education/couns Prevention/Co Disease inform		name	YYYY / MM / DD YYYY / MM / DD	Treatment: ☐ Treatment not recommende Investigator name	ed YYYY /	/ MM / DD
Exclusion: Invest Daycare YY School YY		□ Preschool □ Work	YYYY / MM / DD YYYY / MM / DD			
Date	Intervention subtype	Comments			Next follow-up Date	Initials
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	

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Pertussis Data Collection Worksheet

Please complete all sections.

Panorama Client ID:	
Panorama Investigation ID:	

I) OUTCOMES (op	tional except for severe influ	enza,	L	.HN-> INVEST	IGATION-> OUTCOMES
☐ Not yet recoverd☐ Recovered☐ Fatal	ed/recovering YYYY / MM YYYY / MM YYYY / MM	/ DD □ Intubation /ventilation YYYY / MM / DI	□ Unkno		/Y / MM / DD /Y / MM / DD
	Fatal was selected)				
Transmission Transmission Event ID	Exposure Name	Setting type	Date/Tim		# of contacts
		□ Congregate/Communal living □ Health Care setting □ Type of community contact □ Household Exposure			
		□ Congregate/Communal living □ Health Care setting □ Type of community contact □ Household Exposure			
		□ Congregate/Communal living □ Health Care setting □ Type of community contact □ Household Exposure			
	Pertussis Contacts – Inv	☐ Multiple Settings	to	,	
	-> INVESTIGATION-> EXPOSU	JRE SUMMARY -> TRANSMISSION EVENT SUMMARY -> TE H		NKNOWN/AI	NONYMOUS CONTACTS
Anonymous contac	cts: (total number o	f individuals [including groups that do not require 1:1 follow-	-up])		
Initial Report completed by:				Date initial	report completed:

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Drug ¹	Dosage	Comments
Azithromycin	Infants <6 months: 10 mg/kg/day orally for 5 days.	Preferred antibiotic for infants under 1 month of age.
	Children (>= 6 months to 50 kg): 10 mg/kg/day (to a maximum of 500 mg)orally on the first day followed by 5mg/kg/day (to a maximum of 250 mg) once a day for the next 4 days (5 days total).	Azithromycin is likely safe in pregnancy. No teratogenicity in humans or animals (Rx Files, 2013).
	Adults (50 kg and over): 500 mg orally on the first day followed by 250 mg daily for the next 4 days (5 days total).	
Clarithromycin	Children (up to 33 kg): 15 mg/kg/day provided in a divided dose bid for 7 days (not to exceed maximum of adult dose). Adults (33 kg and over): 250-500 mg po bid for 7 days	Clarithromycin should not be used in pregnancy except where no alternative therapy is appropriate (eCPS, 2015)
Erythromycin	Children (up to 25 kg): Erythromycin estolate: 40 mg/kg/day (to maximum of 1 g per day) provided in a divided dose tid for 7 days. The estolate is a liquid preparation, only used for children or people with difficulty swallowing. Adults: Erythromycin 250 mg qid for 7 days (to maximum of 1 g per day). Some experts recommend 2 g daily in divided doses, for example: a) The Anti-infective Guidelines for Community Acquired Infections: 2001, recommends 1-2 g po daily in divided doses. b) b) The Sanford Guide to Antimicrobial Therapy, 2002, recommends 500 mg qid po.	When prescribing erythromycin prophylactically for neonates one should consider that there have been reports of infantile hypertrophic pyloric stenosis (IHPS) associated with its use as pertussis prophylaxis for newborns. The risk of IHPS after treatment with azithromycin and clarithromycin is unknown. Erythromycin estolate is contraindicated in individuals with existing liver disease or dysfunction, and in pregnancy (CPS, 2010).

Prescribers of macrolide antibiotics for infants <2 months of age on should monitor for signs and symptoms of pyloric stenosis.

For those who are allergic to macrolides, the following may be used although its efficacy is not proven:

- 1. Children: trimethoprim 8mg/kg/day-sulfamethoxazole 40mg/kg/day for 10 days.
- 2. Adults: 2 tabs bid or 1 double strength (DS) tab bid.

¹ Refer to the product monograph and/or the current version of the CPS before prescribing medications.



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References

Jensen, B., Regier, L. D., (Ed.) (2013). *Rx files, Drug Comparison Charts* (9th ed.). Saskatoon, SK: Saskatoon Health Region.

Canadian Pharmacists Association. (2015). Online Compendium of pharmaceuticals and specialties (eCPS): The Canadian drug reference for health professionals. Ottawa, Canada: Author.

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Section 2 - 150 – Pneumococcal Disease - invasive Page **1** of **8** 2018 09 01

Notification Timeline:

From Lab/Practitioner to Public Health: Within 48 hours. From Public Health to Ministry of Health: Within 2 weeks. Public Health Follow-up Timeline: Initiate within 72 hrs.

Public Health Purpose for Notification of Pneumococcal Disease - invasive (adapted from British Columbia Center for Disease Control [2017])

- To track epidemiology trends of invasive pneumococcal disease (IPD) in Saskatchewan including characteristics, risk factors and distribution;
- To monitor the effectiveness of prevention and control measures;
- To plan expansion or introduction of future immunization programs;
- To make timely and evidence informed actions on outbreaks; and
- To inform the public and medical community about IPD.

Surveillance Case Definition¹ (Public Health Agency of Canada, May 2008)

Trable reality of canada, way 2000)				
Confirmed Case	Clinical evidence of invasive disease ¹ with laboratory			
	confirmation of infection:			
	• isolation of Streptococcus pneumoniae from a normally			
	sterile site (excluding the middle ear and pleural cavity)			
	OR			
	• demonstration of <i>S. pneumoniae</i> DNA from a normally sterile			
	site (excluding the middle ear and pleural cavity)			
Probable Case	Clinical evidence of invasive disease ¹ with no other apparent			
	cause and with nonconfirmatory laboratory evidence:			
	• demonstration of <i>S. pneumoniae</i> antigen from a normally			
	sterile site (excluding the middle ear and pleural cavity)			
¹ Clinical illness asso	ociated with invasive disease manifests itself mainly as pneumonia			
with bacteremia, ba	acteremia without a known site of infection, and meningitis.			
Pneumonia without	bacteremia is not notifiable.			

¹ Surveillance case definitions ensure uniform reporting to allow comparability of surveillance data. The definition is not intended to be used for clinical or laboratory diagnosis or management of cases.



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Epidemiology and Occurrence

Under Development

Additional Background Information

Causative Agent

Streptococcus pneumoniae is a gram-positive coccus that replicates in chains. It has a capsule made up of polysaccharides, which lead to the differentiation of over 90 sero-types.

Reservoir/Source

Humans - can be colonized in the upper respiratory tract but not develop infection or disease in the host.

- When the bacterium migrates in the respiratory tract and is not cleared effectively because of cillia impairment or mechanical obstruction, it can replicate and cause disease.
- When bacteremia occurs it can be spread to a variety of sites where replication leads to disease outcomes.

Pathophysiology

Invasive pneumococcal disease (IPD) can present as meningitis, endocarditis, septic arthritis, and peritonitis.

- Meningitis
 - > Streptococcus pneumoniae is the most common etiological agent of bacterial meningitis in adults. It may arise from direct extension of infection from the middle ear, sinuses, or from bacterial seeding to the choroid plexus in the brain following bacteremia.
 - Local extension to the meninges via the sinuses or dura mater defects or the pleura via the lungs can also lead to invasive disease development.
- Peritonitis in adults, endocarditis, pericarditis and septic arthritis can occur spontaneously or secondarily to a prosthesis or underlying rheumatoid illness.
- Osteomyelitis in adults tends to involve the vertebrae.
- Unusual pneumococcal infections may suggest underlying immunodeficiencies of some cause.



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Streptococcus pneumoniae can colonize the upper respiratory tract and adhere to the cells lining the nasopharanx. Impairment of ciliary action plays an important role in the development of infection in the respiratory tract.

The organism causes disease through its ability to escape phagocytosis because of its capsular structure. It is therefore able to replicate in tissues and fluids and create an intense inflammatory response causing the various familiar clinical pictures to appear. The organism does not produce any clinically significant toxins.

Symptoms

Common symptoms of IPD (e.g., infections of the meninges, joints, etc.) are:

- fever;
- malaise;
- associated symptoms of severe systemic infection symptoms vary depending on the site of infection (see Pathophysiology section above).

In non-invasive disease, direct spread in the respiratory tract can lead to the development of disease entities such as otitis media, sinusitis, and pneumonia.

Incubation Period

The incubation period is dependent on a number of factors including site of infection, bacterial load and underlying conditions that support the development of infection. In invasive disease the clinical picture usually starts developing within a few hours of infection ocurring and is a reflection of the intense inflamatory response to the organism.

- Meningitis unknown; probably short, 1-4 days.
- Pneumonia not well determined; may be as short as 1-3 days.

Period of Communicability

- Unknown.
- May be as long as the bacterium is present in the respiratory tract.
- May be prolonged especially in immunocompromised hosts.
- Probably less than 24-48 hours after effective antimicrobial therapy has begun.

Mode of Transmission

- Contact with respiratory secretions or direct oral contact.
- Person to person via droplet spread is thought to be the most prevalent form of transmission but infrequently leads to illness.



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Risk Groups/Risk Factors (Fauci, et al., 2007)

Settings with increased risk of exposure:

- daycare centres;
- military training camps;
- prisons;
- homeless shelters;
- air pollution;
- over-crowded living conditions;
- poor socioeconomic status.

Host factors:

- respiratory infection, inflammation (viral respiratory illness such as influenza);
- chronic obstructive pulmonary disease (COPD);
- immunosuppression due to illness or therapy;
- asplenia;
- age (infancy or elderly);
- alcoholism;
- allergies;
- cigarette smoking;
- malnutrition;
- chronic disease (including HIV, liver/kidney disease, diabetes, etc.);
- fatigue, stress and/or exposure to cold.

Specimen Collection and Transport

Specimen type is dependent on the relevant clinical disease. Material can be obtained from the infectious focus, blood or CSF. Blood cultures should be done in all cases of suspected invasive disease. Recovery of pneumococci from an upper respiratory tract culture is not indicative of the etiologic diagnosis of pneumococcal disease in the respiratory tract.

Where appropriate, material obtained can be gram stained and subsequently cultured using standard microbiological techniques. All isolates from a normally sterile site should be tested for antibiotic sensitivity as results from this will assist in case management and antibiotic therapy.

Isolates of *S. pneumoniae* from IPD cases should be referred to Roy Romanow Provincial Laboratory (RRPL) for serotyping.



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Public Health Investigation

I. Case

History

Refer to <u>Attachment – Pneumococcal Disease (invasive) Data Collection Worksheet</u> to assist.

Key elements to inquire about include:

- Presentation of illness.
- Medical history including underlying medical conditions that may predispose the individual to invasive disease (see risk factors/risk groups).
- Settings with increased risk of exposure (see risk factors/risk groups).
- Immunization history of case.

Public Health Interventions

Education

 All cases should be provided disease information as well as information on prevention and control measures including period of communicability and avoiding contact with vulnerable individuals.

Immunization

- Immunization to be offered if incomplete.
- If case meets eligibility criteria, immunizations should be started as per Saskatchewan Immunization Manual².

Isolation

- Clients are no longer communicable once on effective antibiotic therapy for 24-48 hours.
- Clients may return to work or school/daycare settings when they have clinically recovered and are able to resume normal activities.

Referrals

Specialist care and long-term follow up may be indicated in certain circumstances.

Treatment/Supportive Therapy

Treatment for clinical management is under the direction of the primary care provider. The public health practitioner should direct any questions regarding the current treatment protocols to the physician or Medical Health Officer (MHO).

II. Contacts/Contact Investigation

No contact tracing is required.



²https://www.ehealthsask.ca/services/Manuals/Pages/SIM.aspx

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III. Environment

Child Care Centres/Institutional Control Measures

• Standard precautions for hospitalized patients (refer to local infection control manual). No specific measures.

IV. Epidemic Measures

- No specific measures.
- Immunization may be indicated for use in outbreaks.
- Outbreaks should be reported immediately to Saskatchewan Ministry of Health.

Prevention Measures

Refer to the <u>Respiratory and Direct Contact Introduction and General Considerations</u> section of the manual that highlights topics for client education that should be considered as well as provides information on high-risk groups and activities.

Immunization

- Routine immunization of all children with the Pneu-C (conjugate pneumococcal vaccine) as per Saskatchewan Immunization Manual.³
- The reader is referred to both the Saskatchewan Immunization Manual,¹ the latest version of the Canadian Immunization Guide and the latest guidelines/memos indicating provincial policies for further information.

Prophylactic Antibiotic Therapy

• Individuals with certain risk conditions may be placed on long-term prophylactic antibiotic therapy by their physician.



³ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

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Revisions

Date	Change
September 2018	Clarified the purpose for notification of cases to public health.
	 Incorporated an Epidemiology and Occurrence section as a placeholder.
	Rearranged and updated the style into the new format of the
	Manual.



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References

- American Academy of Pediatrics. (2015). *Red book: 2015 Report of the Committee on Infectious Diseases* (30th ed.). Elk Grove Village, IL: Author.
- Fauci, A. S., Braunwald, E., Kasper, D., Hause, S. L., Longo, D. L., Jameson, J. L., et al. (2007). *Harrison's principles of internal medicine* (17th ed.). Whitby, ON: The McGraw-Hill Companies.
- Heymann, D. L. (Ed.). (2015). *Control of communicable diseases manual* (20th ed.). Washington, DC: American Public Health Association.
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- Public Health Agency of Canada. (2006). *Canadian immunization guide* (7th ed.). Ottawa, Canada: Public Works and Government Services Canada.
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Pneumococcal Disease (invasive) Data Collection Worksheet



Panorama QA complete: ☐Yes Initials:	□No	Please complete all sections.		Par	Panorama Client ID: Panorama Investigation ID:		
A) CLIENT INFORMATION			LHN -> SUBJE	CT -> CLIEN	T DETAILS -> PEF	RSONAL INFORMATIO	
Last Name:		First Name: and Middle Name:	Alternate Name (Goes by):				
DOB: YYYY / MM / DD Phone #: Primary Home:	Age:	Health Card Province: Health Card Number (PHN):	Preferred Communication Method: (specify - i.e. home phone, text): Email Address: □Work □Personal				
Place of Employment/School:		Gender: □ Male	□ Female		Other	□ Unknown	
Alternate Contact: Relationship: Alt. Contact phone:		Address Type: □ No fixed □ Postal Address Mailing (Postal address): Street Address or FN Communi Address at time of infection if r	ity (Primary Hon		porary □Legal	Land Description	
B) INVESTIGATION INFORMATION Disease Summary Classification: CASE	SUBJE Date	CT SUMMARY-> RESPIRATORY &	DIRECT CONTA	CT ENCOU	LAB TEST INFO Date specimen	RMATION:	
□ Confirmed	YYYY / MM / DD	☐ Person Under Investigation	YYYY / MM	/ DD	YYYY / MM / [DD	
□ Does Not Meet Case	YYYY / MM / DD	□ Probable	YYYY / MM	/ DD	Specimen type: Blood Other	: □ CSF	
Disposition: FOLLOW UP: ☐ In progress ☐ Incomplete - Declined ☐ Incomplete - Lost contact ☐ Incomplete - Unable to locate REPORTING NOTIFICATION Name of Attending Physician or Nu Physician/Nurse Phone number:	YYYY / MM / DD	☐ Complete ☐ Not required ☐ Referred – Or (specify where) Location:		YYYY / N	MM / DD MM / DD		
Priysician/Nurse Phone number:		Date Receive	ea (Public Health	ij: YYYY /	ואוואו / טט		
Type of Reporting Source: Hea	lth Care Facility □L	ab Report □ Nurse Practit	ioner □Phy	rsician	□ Other		

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Pneumococcal Disease (invasive) Data Collection Worksheet

Please complete all sections.

				Pa		norama Client ID: a Investigation ID:
DISEASE EVENT HISTORY			IN	/ESTIGATION->DISEASE SUMMAR	Y (UPE	DATE)->DISEASE EVENT HISTOR
Site / Presentation:	sis	☐ Meningitis		☐ Pneumonia with bacteremia		□ Other
) SIGNS & SYMPTOMS (Bold text = par	t of cas	se definition)		LHN->	INVEST	FIGATION->SIGNS & SYMPTON
Description	No	Yes – Date of onset	Description		No	Yes - Date of onset
Arthritis - septic		YYYY / MM / DD	Malaise			YYYY / MMM / DD
Cardiac - endocarditis		YYYY / MM / DD	Meningitis			YYYY / MMM / DD
Cardiac - pericarditis		YYYY / MM / DD	Peritonitis			YYYY / MMM / DD
Fever		YYYY / MM / DD	Pneumonia			YYYY / MMM / DD
Osteomyelitis			Sepsis (e.g. ba	actremia, septicemia, etc.)		
E) RISK FACTORS (RF followed by + imp	act the	Immunization Forecas	ter)		ı	LHN-> SUBJECT->RISK FACTOR
DESCRIPTION		Yes Start date	N, NA, U	Add'l Info		
Chronic Medical Condition - Cardiac Disc	ease+	Start date				
Chronic Medical Condition - Diabetes M	lellitus-	+				
Chronic Medical Condition - Liver Disease	se+					
Chronic Medical Condition - Lung Diseas	se+					
Chronic Medical Condition - Other (Add	'l Info))					
Contact to a known case (Add'l Info)		YYYY / MM/DD				
Exposure - Second hand smoke						
Immunocompromised - Related to unde disease or treatment	erlying					
Special Population - Attends childcare						
Special Population – Homeless +						
Special Population - Lives in a communa	l settin	g				
Substance Use - Alcohol						
Substance Use - Tobacco						
:) IMMUNIZATION HISTORY INTERPRE	FATION	SUMMARY	LHN -> IN	VESTIGATION-> IMMUNIZATION	HISTO	RY INTERPRETATION SUMMAR
Interpretation Date: YYYY /	MM /	DD				
Interpretation of Disease Immunity:		M - Fully immunized (fo	0 ,	☐ IOM - Partially immuni		
☐ IOM – Unimmunized ☐ ION Reason:	1 - Uncl	ear immunization histor	y Valid do	ses received: Doses need	ed:	
☐ IOM – Interpretation of history by inv	estigat	or				
G) TREATMENT				LHN -> INVESTIGATION-> MED	ICATIO	DNS-> MEDICATIONS SUMMAR
Medication (Panorama = Other Meds) :						
Prescribed by:			C+	ed on: YYYY / MM / DD		

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Pneumococcal Disease (invasive) Data Collection Worksheet

Please complete all sections.

Panorama Client ID: ___

			Par	norama investigation ib	:
H) INTERVENTION		LHN	-> INVESTIGATION->TREATMENT & INTERV	'ENTIONS->INTERVENT	ION SUMMAR
Intervention Type a					
General: Investigato	or name		Immunization:		
☐ Disease-Info/Prev	v-Control	YYYY/ MM / DD	☐ Eligible Immunization recommended	ed YYYY / MM / DD	
			Disease-specific immunization recomm		
Education/counselli			☐ Disease-specific immunization given	YYYY / N	IM / DD
□ Prevention/Contr		YYYY / MM / DD	Investigator name		
☐ Disease informati	ion provided	YYYY / MM / DD			
Other Investigation	Findings:		Isolation:		
☐ Investigator Note	es -	☐ See Document Management	Facility isolation YYYY / MM / DD	Investigator name	
	·	Τ	☐ Home isolation YYYY / MM / DD	Investigator name	I _I .
Date	Intervention subtype	Comments		Next follow-up Date	Initials
YYYY / MM / DD	subtype			YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
YYYY / MM / DD				YYYY / MM / DD	
OUTCOMES (option	onal except for severe	influenza,		LHN-> INVESTIGATION	N-> OUTCOM
□ Not yet recovered	d/recovering YYYY / I	MM / DD	nedical care YYYY / MM / DD	oitalization YYYY / MN	4 / DD
□ Recovered		_	•	nown YYYY / MN	
□ Fatal	*	•	YYYY / MM / DD	10WII 1111 / 1VII	11 / 00
	atal was selected)		1111 / 10101 / 55		
cause of Beatin (ii re	atai was selectea)				
Initial Report				Date initial report c	omnleted:
completed by:				VVVV / MMA / DD	•

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Date Reviewed: August, 2011 Section: 2-160
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Notification Timeline:

From Lab/Practitioner to Public Health: Within 48 hours (or immediate if an outbreak is suspected).

From Public Health to Ministry of Health: Within 72 hours (or immediate if an outbreak is suspected).

Public Health Follow-up Timeline: Initiate within 24-48 hrs.

Information

Case Definition (Public Health Agency of Canada, May 2008)

Case Definition (Fublic Health Agency of Canada, Way 2008)			
Confirmed Case	Laboratory confirmation of infection in the absence of recent immunization ¹ with rubella containing vaccine:		
	• isolation of rubella virus from an appropriate clinical specimen OR		
	• detection of rubella virus RNA OR		
	 seroconversion or a significant (e.g., fourfold or greater) rise in rubella IgG titre by any standard serologic assay between acute and convalescent sera OR 		
	• positive serologic test for rubella IgM antibody using a recommended assay* in a person with an epidemiologic link to a laboratory-confirmed case or who has recently travelled to an area of known rubella activity.		
	OR		
	Clinical illness ² in a person with an epidemiologic link to a laboratory-confirmed case.		
Probable Case	Clinical illness ²		
	• in the absence of appropriate laboratory tests OR		
	• in the absence of an epidemiologic link to a laboratory-confirmed case OR		
	in a person who has recently travelled to an area of known rubella activity.		



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The most frequent reaction to measles-mumps-rubella (MMR) immunization is malaise and fever (with or without rash) occurring 7-12 days after immunization. However, this should be determined for each case, as these reactions and time frames can vary (*Canadian Immunization Guide*, 2006).

² Clinical illness is characterized by fever and rash, and at least one of the following:

- arthralgia/arthritis
- lymphadenopathy
- conjunctivitis

*IgM serology has the potential for false-positive findings. If the clinical presentation is inconsistent with a diagnosis of rubella or in the absence of recent travel/exposure history, IgM results must be confirmed by the other listed confirmatory methods. Rubella avidity serology is recommended for IgM positive results in pregnant women. Most acute rubella cases develop IgM after 5 days post rash onset. Therefore, a suspected rubella case in which serum collected < 5 days after rash onset initially tests IgM negative should have a second serum collected > 5 days after onset for retesting for IgM. Further strain characterization is indicated for epidemiologic, public health and control purposes.

Causative Agent

Rubella virus, an RNA virus of the genus Rubivirus.

Symptoms

Adults may experience a 1 to 5 day prodrome of mild fever, malaise, headache, and conjunctiva. Characteristic postauricular and suboccipital lymphadenopathy is followed by a diffuse maculopapular rash 5 to 10 days later. Children usually have few or no symptoms.

Complications (American Academy of Pediatrics, 2009)

- Encephalitis.
- Thrombocytopenia.
- Maternal rubella during pregnancy can result in miscarriage, fetal death or a variety of congenital anomalies. Refer to <u>Congenital Rubella Syndrome/Infection</u> in the Respiratory and Direct Contact section of the manual.

Incubation Period

Usually 16-18 days, but ranges from 14-23 days, (American Academy of Pediatrics, 2009).

Reservoir/Source

Humans.



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Mode of Transmission

Spread by direct or droplet contact with nasopharyngeal secretions of an infected individual. In congenital rubella syndrome, the virus is transmitted to the fetus during pregnancy in 25% of cases of women who were exposed to rubella during their first trimester of pregnancy.

Period of Communicability

Approximately 1 week before to 4-5 days after onset of the rash.

Specimen Collection and Transport

To facilitate rapid testing, laboratory requisitions should be clearly marked "suspect case of rubella" when sending specimens for rubella testing.

To confirm the diagnosis the following specimens should be submitted to Saskatchewan Disease Control Laboratory (SDCL):

- Submit 5 mL serum samples for rubella IgM and IgG (acute and convalescent).
 - IgM response begins with onset of rash and will persist for 1 to 2 months. Only a small proportion of cases will have IgM present in serum samples collected on the day the rash appears. The proportion with IgM rises rapidly until the great majority of cases have IgM by day 5 post-onset of rash.
 - IgG response begins about 1 week after the onset of symptoms and will persist for a lifetime.
 - Convalescent sera should be drawn 10 to 20 days after the initial serology to assess the rise in IgG titre (seroconversion). This interval may be shorter if maternal rubella is being investigated.
 - Rubella specific IgM serology is the standard test for routine diagnosis of rubella but demonstration of a significant increase in the rubella specific IgG titre is a reliable alternative serologic method for diagnosis.
- Nasopharyngeal secretions, for isolation of rubella virus. Collect nasopharyngeal swab or a throat swab, and place in virus transport medium, within 4 days after the onset of symptoms. Refer to the SDCL Compendium of Tests at http://sdcl-testviewer.ehealthsask.ca/ for specimen collection instructions.
- Refrigerate specimens immediately and ship on ice to SDCL. Specimen must be received within 24 hr of collection.



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Methods of Control/Role of Investigator

Prevention and Education

Refer to the <u>Respiratory and Direct Contact Introduction and General Considerations</u> section of the manual that highlights topics for client education that should be considered as well as provides information on high-risk groups and activities.

Immunization

- Immunize infants, children and adults according to the recommended schedule. Refer to Saskatchewan Immunization Manual.¹
- Because of the implication of congenital rubella syndrome, special attention to immune status should be paid to women in their preconception, prenatal and postnatal period. If necessary, immunizations should be offered in accordance with the Saskatchewan Immunization Manual.¹
- Postpartum women who are non-immune should be given rubella-containing vaccine before discharge from hospital. Refer to Saskatchewan Immunization Manual¹ for details.

Education

- Education should be provided regarding respiratory etiquette and measures to prevent transmission of rubella.
- Educate the public about the disease and the need for active immunization with a rubella-containing vaccine. Immunization information fact sheets can be used to guide discussion.

Management

The primary goal of rubella control is to prevent defects in the infants of women who acquire the disease while pregnant. Educate all individuals who are considered contacts. Provide information about rubella to all individuals who may have been exposed to the virus, especially women who may be pregnant or of reproductive age.



¹ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

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Information about the signs and symptoms of the disease and the importance of isolation from other possible contacts, including health care workers, daycares and schools and especially other pregnant women is essential.

I. Case

History

- Determine case status and immunization history including a review of the number and dates of rubella-containing vaccine.
- Determine the source of infection. Discuss social events, visitors from out of province, travel out of province and any contact with others who have been ill or with infants who may have congenital rubella syndrome.
- Discuss in detail the dates, names and places where the individual may have been
 in contact with others during the period of communicability and record contact
 details on the Attachment Contact Follow-up Form in the Respiratory and Direct
 Contact Introduction and General Considerations section of the manual.

Immunization

Investigate immunization history, record date and place.

Treatment/Supportive Therapy

None. Supportive care in the home if symptoms of fever and headache indicate encephalitis, the case should seek medical attention.

Exclusion

Exclude cases from school, daycare, and work for 7 days following the onset of rash (Health Canada 1999, American Academy of Pediatrics 2009).

Referrals

In case of infection with wild rubella virus early in pregnancy, referral to family physician for appropriate counselling should be provided.

II. Contacts/Contact Investigation Contact Definition/Categorization

• Anyone who is likely to have been exposed to the nose or throat secretions of a person with rubella during their infectious period.



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• Investigate all household and close contacts, with special emphasis on exposure to pregnant women, and determine susceptibility. See Definition of Susceptible Contacts. The following settings should be considered:

- work, school, childcare centres;
- social events;
- medical or clinical facilities may be considered as well.
- Individuals are considered immune if they:
 - were born in Canada prior to 1970;
 - were born in Canada in 1970 or later and have documented evidence of immunization with live rubella-containing vaccine after their first birthday;
 - were born outside Canada and have documented evidence of immunization with live rubella-containing vaccine after their first birthday,
 - have laboratory-documented evidence of rubella or laboratory evidence of immunity.

Definition of Susceptible Contacts

- Infants less than one year of age.
- Immunocompromised individuals.
- Persons born in Canada in 1970 or later and people born outside of Canada who do not have:
 - documented evidence of vaccination with one dose of live rubella-containing vaccine received after their first birthday

OR

laboratory evidence of immunity

OR

• a history of laboratory-confirmed rubella.

Prophylaxis/Testing/Immunization

 All pregnant women who have been exposed to the virus should have a blood test for rubella antibody if not already documented. Immune globulin may be suggested for those who are non-immune in consultation with the infectious disease specialist and gynaecologist. The value of this approach has not been established.



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Immunize all susceptible contacts with the exception of pregnant or immunosuppressed individuals. All individuals who have been exposed to the virus and who have no medical contraindications to the rubella vaccine should be given rubella-containing vaccine immediately.² Post pubertal females should be advised not to get pregnant for 1 month after receiving rubella-containing vaccine.

• Follow up all contacts within one week to confirm that they have been immunized and/or that they have or have not developed symptoms.

Exclusion

Exclude all suspected cases from school, daycare or work. If possible do not send them home on public transportation or on the school bus.

III. Environment

Child Care Centres/Institutional Control Measures

- Investigate immune status of health care/daycare workers and immunize all who are non-immune, except in the case of pregnancy or immunosuppression.
- Health care workers who are susceptible must not work with patients suspected or confirmed to have rubella. These workers can become infected and may also become a source for transmission (Health Canada, 2002).
- Inform parents of children in daycare centres of the need for susceptible children 12 months of age or older to be immunized immediately.
- Cases in a hospital or institution should be managed under strict contact and droplet isolation precautions.

Epidemic Measures

- Ensure prompt reporting of all confirmed and suspected cases. The medical community and general public should be made aware of rubella epidemics in order to identify and protect any pregnant women who may be susceptible.
- Active surveillance for infants with congenital rubella syndrome (CRS) should be carried out until 9 months after the last reported case of rubella.



² Although live-virus rubella vaccine given after exposure has not been demonstrated to prevent illness, vaccine theoretically could prevent illness if administered within 3 days of exposure. If this exposure does not result in illness, immunization will provide protection in the future (American Academy of Pediatrics, p. 582, 2009).

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• There is a special concern when rubella cases are identified in unimmunized or underimmunized communities and additional control measures may be implemented.

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Congenital Rubella Syndrome/Infection (CRS/CRI)

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Notification Timeline:

From Lab/Practitioner to Public Health: Within 48 hours. From Public Health to Ministry of Health: Within 72 hours. Public Health Follow-up Timeline: Initiate within 72 hrs.

Information

Case Definition (Public Health Agency of Canada May 2008)

Table 1. Nationa	al Case Definition for Congenital Rubella Syndrome (CRS)		
Confirmed Case	Live birth: two clinically compatible manifestations (any combination		
	from <u>Table 3</u> , Columns A and B) with laboratory confirmation of		
	infection:		
	• isolation of rubella virus from an appropriate clinical specimen OR		
	detection of rubella virus RNA		
	OR		
	positive serologic test for IgM antibody in the absence of recent immunization with rubella-containing vaccine		
	OR		
	rubella IgG persisting for longer than would be expected		
	(approximately six months after birth) from passive transfer of		
	maternal antibody, or in the absence of recent immunization.		
	Still birth: two clinically compatible manifestations with isolation of		
	rubella virus from an appropriate clinical specimen.		
Probable Case	In the absence of appropriate laboratory tests, a case that has at least:		
	• any two clinically compatible manifestations listed in <u>Table 3</u> ,		
	Column A		
	OR		
	• one manifestation listed in <u>Table 3</u> , Column A, plus one listed in		
	<u>Table 3,</u> Column B.		
Not a Case	rubella antibody titre absent in the infant		
	OR		
	rubella antibody titre absent in the mother		
	OR		
	• rubella antibody titre declining in the infant consistent with the normal decline after birth of passively transferred maternal antibody.		



Congenital Rubella Syndrome/Infection (CRS/CRI)

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Case Definition (Public Health Agency of Canada, May 2008)

Table 2. National Case Definition for Congenital Rubella Infection (CRI)			
Confirmed Case	Laboratory confirmation of infection but with no clinically compatible		
	manifestations:		
	• isolation of rubella virus from an appropriate clinical specimen		
	OR		
	detection of rubella virus RNA		
	OR		
	• positive serologic test for rubella IgM antibody in the absence of		
	recent immunization with rubella-containing vaccine		
	OR		
	• rubella IgG persisting for longer than would be expected		
	(approximately six months after birth) from passive transfer of		
	maternal antibody, or in the absence of recent immunization.		

Table 3. Congenital Rubella Syndrome: Clinically Compatible Manifestations			
(Public Health Agency of Canada, May 2008)			
Column A	Column B		
1. Cataracts or congenital glaucoma	1. Purpura.		
(either one or both count as one).	2. Hepatosplenomegaly.		
2. Congenital heart defect.	3. Microcephaly.		
3. Sensorineural hearing loss.	4. Micro ophthalmia.		
4. Pigmentary retinopathy.	5. Mental retardation.		
	6. Meningoencephalitis.		
	7. Radiolucent bone disease.		
	8. Developmental or late onset conditions		
	such as diabetes and progressive		
	panencephalitis and any other conditions		
	possibly caused by rubella virus.		

Causative Agent

Rubella virus, an RNA virus of the genus Rubivirus.



Congenital Rubella Syndrome/Infection (CRS/CRI)

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Symptoms

In addition to the manifestations identified in <u>Table 3</u>, the following may also be seen (American Academy of Pediatrics, 2009):

- growth retardation;
- interstitial pneumonitis;
- thrombocytopenia;
- dermal erythropoesis ("blueberry muffin" lesions).

Moderate to severe cases of CRS are usually recognizable at birth. Mild cases that involve slight cardiac involvement or deafness may not be detected for months or even years. A frequent late manifestation of CRS is insulin-dependent diabetes mellitus (Heymann, 2008).

Fetal infections during the 1st trimester are at the greatest risk of intrauterine death, spontaneous abortion and congenital malformations of major organ systems. Infection in the first 20 weeks of gestation is most often associated with CRS and birth defects. Infections after the first 20 weeks of gestation are most often associated with CRI (Alberta Health & Wellness, 2005).

Incubation Period

Not applicable.

Reservoir/Source

Humans.

Mode of Transmission

- From an infected mother to her developing fetus.
- The occurrence of congenital defects is up to 85% if infection associated with maternal rash occurs during the first 12 weeks of gestation, 54% during 13-16 weeks, and 25% during the end of the second trimester (American Academy of Pediatrics, 2009).

Period of Communicability

Infants with CRS/CRI can shed virus in their pharyngeal secretions and urine for up to a year or more.



Congenital Rubella Syndrome/Infection (CRS/CRI)

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Specimen Collection and Transport

Laboratory confirmation of CRS/CRI is done by:

- detection of IgM in cord blood or serum of the infant
- detection of persistent rubella IgG in the infant (beyond approximately 6 months at which time maternally acquired antibodies usually wane)
 OR
- detection of rubella virus in samples (e.g., respiratory specimens collected during the first few months of life) (Alberta Health & Wellness, 2005).

Contact Saskatchewan Disease Control Laboratory (SDCL) Virology Section for additional information about specimen collection.

Methods of Control/Role of Investigator

Prevention and Education

Refer to the <u>Respiratory and Direct Contact Introduction and General Considerations</u> section of the manual that highlights topics for client education that should be considered as well as provides information on high-risk groups and activities.

Immunization

- Immunize infants, children and adults according to the recommended schedule. Refer to Saskatchewan Immunization Manual.¹
- Special attention must be paid to the immune status of women in their preconception, prenatal and postnatal period. If necessary, immunizations should be offered in accordance with the Saskatchewan Immunization Manual.¹
- Postpartum women who are non-immune should be given rubella-containing vaccine before discharge from the hospital. Refer to the Saskatchewan Immunization Manual.¹



¹ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

Congenital Rubella Syndrome/Infection (CRS/CRI)

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Education

• Educate the public about the disease and the need for active immunization with a rubella-containing vaccine. Immunization information fact sheets can be used to guide discussion.

Management

I. Case

History

Confirm the diagnosis.

Treatment/Supportive Therapy

There is no specific treatment for CRS.

Exclusion

- The infant should be isolated after birth. Routine practices, as well as droplet and contact precautions should be strictly enforced.
- Health care workers who are susceptible must not work with patients suspected or confirmed to have rubella. These workers can become infected and subsequently become a source for transmission (Health Canada, 2002).
- Once discharged from hospital, only persons that are immune to rubella should have contact with and care for the infected newborn.
- Children with CRS/CRI should be presumed infectious at least through to age one year, unless nasopharyngeal and urine cultures are negative for virus after three months of age. The Medical Health Officer (MHO) should determine a schedule of nasopharyngeal swabs and urine cultures for the first year of life in consultation with the physician and SDCL.
- Viral isolation is not always successful and repeated attempts at viral isolation testing may be necessary – the pediatrician may consult with MHO who is to consult with SDCL for guidance in this regard.

Referrals

- The family physician may make referrals to specialists for infants with CRS/CRI, as appropriate (ophthalmologists, audiologists, heart specialists, etc.).
- The infant should continue to be monitored for clinical manifestations by their physician.



Congenital Rubella Syndrome/Infection (CRS/CRI)

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II. Contacts/Contact Investigation

Susceptible (non-immune) persons should avoid contact with the infant until they are immunized. This is particularly relevant for non-immune pregnant women and children less than 12 months of age.

III. Environment

Child Care Centres/Institutional Control Measures

- Contact and droplet isolation precautions should be implemented in hospitals to infants with CRS/CRI who are under 12 months, unless urine and pharyngeal virus cultures are negative for rubella virus after 3 months of age.
- Investigate immune status of health care/daycare workers and immunize all who are non-immune, except in the case of pregnancy or immunosuppression.



Congenital Rubella Syndrome/Infection (CRS/CRI)

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Severe Acute Respiratory Infection (SARI)

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Notification Timeline

From Lab/Practitioner to Public Health: Immediate.

From Public Health to Ministry of Health: Upon notification from lab or

physician.

Public Health Follow-up Timeline: Within 24-48 hours.

Information

Case Definition (adapted from Public Health Agency of Canada, 2013)

To confirm the diagnosis of a case of SARI, the case must meet criteria in each of the categories listed below for hospitalized cases (A) or for cases who are deceased (B):

- 1. Respiratory symptoms.
- 2. Severity.
- 3. Unknown diagnosis.
- 4. Epidemiological exposure, as detailed in the specific case definitions below.

SARI Case (A)

A person admitted to hospital with the following:

- 1. Respiratory symptoms, i.e.:
 - Fever¹ of over 38 degrees Celsius **AND** new onset of (or exacerbation of chronic) cough or breathing difficulty.

AND

- 2. Evidence of severe illness progression, i.e.:
 - Either radiographic evidence of infiltrates consistent with pneumonia, or a diagnosis of acute respiratory distress syndrome (ARDS) or severe influenza-like illness (ILI),² which may also include complications such as encephalitis, myocarditis or other severe and life threatening complications.

AND

3. Either admission to the ICU/other area of the hospital where critically ill patients are cared for OR mechanical ventilation.

AND

¹ As per the ILI definition, fever may not be prominent in patients under 5 years or 65 years and older as well as in immunosuppressed individuals. Failure to take temperature should not rule out a history of self-reported fever. Clinical judgment should always prevail with regard to these groups.

² **Severe ILI:** In addition to the symptoms of ILI noted below, severe ILI may also include complications such as encephalitis, myocarditis or other severe and life threatening complications.



Severe Acute Respiratory Infection (SARI)

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4. No alternate diagnosis within the first 72 hours³ of hospitalization, i.e.:

• Results of preliminary clinical and/or laboratory investigations, within the first 72 hours of hospitalization, cannot ascertain a diagnosis that reasonably explains the illness.

AND

5. One or more of the following exposures/conditions, i.e.:

- Residence, recent travel (within ≤ 14 days of illness onset) to a country where human cases of novel influenza virus or other emerging/re-emerging pathogens have been detected or are known to be circulating in animals⁴.
- Close contact⁵ with an ill person who has been to an affected area/site within the 14 days prior to onset of symptoms.
- Exposure to settings in which there had been mass die offs or illness in domestic poultry or swine in the previous six weeks.
- Occupational exposure involving **direct** health care, laboratory or animal exposure, i.e.:
 - Health care exposure involving health care workers who work in an environment where patients with SARI are being cared for, particularly patients requiring intensive care.

OR

• **Laboratory exposure** in a person who works directly with Laboratory biological specimens.

OR

- Animal exposure in a person employed as one of the following:
 - Poultry/swine farm worker;
 - Poultry/swine processing plant worker;
 - Poultry/swine culler (catching, bagging, transporting or disposing of dead birds/swine);

⁵ Close contact is defined as: Anyone who provided care for the patient, including a health care worker or family member, or who had other similarly close physical contact; Anyone who stayed at the same place (e.g. lived with, visited) as a probable or confirmed case while the case was ill.



³ It is suggested that laboratory investigation, including laboratory testing for influenza and other respiratory pathogens should be started as soon as possible upon presentation (i.e., do not wait 72 hours to initiate testing) and it requires immediate infection control and public health action. Refer to Attachment - Severe Acute Respiratory Illness (SARI) Screening Tool and discuss with the Medical Health Officer and Infection Control.

⁴ Refer to the World Health Organization Human Animal Interface for the most recent information http://www.who.int/influenza/human animal interface/en/)

Severe Acute Respiratory Infection (SARI)

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- Worker in live animal market;
- Dealer or trader of pet birds, pigs or other potentially affected animals;
- Chef working with live or recently killed domestic poultry, swine or other potentially affected animals;
- Veterinarian worker;
- Public health inspector/regulator.

OR

SARI Case (B)

A deceased person with the following:

- 1. A history of respiratory symptoms, i.e.:
 - History of unexplained acute respiratory illness (including fever and new onset of (or exacerbation of chronic) cough or breathing difficulty) resulting in death.

AND

- 2. Autopsy performed with findings consistent with SARI, i.e.:
 - Autopsy findings consistent with the pathology of ARDS without an identifiable cause.

AND

3. No alternate diagnosis that reasonably explains the illness.

AND

4. One or more of exposures/conditions, as listed in (A).

SARI Case Exclusion Criteria

A person should not be reported as a case of SARI if an alternate diagnosis can reasonably explain their illness.

Health Care Facility Surveillance for SARI

It is recommended that regions/jurisdictions use the <u>Attachment – Severe Acute</u> <u>Respiratory Illness (SARI) Screening Tool</u> in their acute and integrated health care facilities to ensure the early recognition of potential SARI cases and the prompt notification of Infection Control and Medical Health Officers (MHOs). This will ensure that sporadic cases of SARI are reported and assessed using this case definition.



Severe Acute Respiratory Infection (SARI)

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Causative Agent

Varies; includes several emerging respiratory pathogens including but not limited to influenza A (H5N1), other novel influenza virus, SARS-CoV (coronavirus), etc.

Symptoms

• Fever (> 38 degrees Celsius).

- New onset of (or exacerbation of chronic) cough or breathing difficulty.
- Radiographic evidence of infiltrates consistent with pneumonia or acute respiratory distress syndrome (ARDS) or Severe ILI.

Incubation Period

Varies depending on the organism; for example:

- SARS-CoV is 3 to 10 days.
- Avian influenza ranges from 2-8 days and as long as 17 days.

Reservoir/Source

Varies depending on the organism; for example:

- SARS-CoV is unknown.
- Avian influenza primarily birds, but can affect humans and pigs as well.

Mode of Transmission

- Direct contact with respiratory secretions or body fluids of a confirmed, suspect of probable case or direct contact with suspected animals implicated in transmission.
- Airborne via aerosol-generating medical procedures.⁶
- SARS-CoV person to person by close contact. Primarily through droplets and fomites
- Avian influenza refer to Vector-Borne and Zoonotic Diseases Avian Influenza section of the manual. (The virus is transmitted through close contact with dead or sick birds. There is limited human-to-human transmission occurring at this time.)
- MERS-CoV contact with camels or their milk or urine; person to person by close contact.

⁶ **Aerosol Generating Medical Procedure**: A medical or surgical procedure that involves manipulation or stimulation of a patient's airway in a manner that may stimulate coughing and/or promote the generation of aerosols.



Severe Acute Respiratory Infection (SARI)

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Period of Communicability

- Varies depending on the specific organism suspected or identified.
- Not completely understood for SARS-CoV initial studies suggest that transmission does not occur before onset of clinical symptoms and maximum period of communicability is less than 21 days.
- Difficult to determine when there is no evidence of direct human-to-human transmission (avian influenza).

Specimen Collection and Transport

Appropriate testing for routine respiratory pathogens should be reinforced.

The following are suggested laboratory diagnostic tests that should be considered in the **initial** laboratory work-up of patients presenting with symptoms of SARI. Relevant medical history, as well as clinical signs and symptoms will dictate appropriate ongoing testing for each patient, (The Public Health Agency of Canada, 2013).

Specimens should be sent on a STAT basis. Refer to the Saskatchewan Disease Control Laboratory (SDCL) Compendium of Tests⁷, Time or Temperature Sensitive, STAT and Outbreak Samples Policy for details on submitting STAT samples. The MHO may be able to assist in expediting testing.

The initial specimens must be clearly marked "SARI Screen".

- Blood culture.
- Sputum for C&S.
- Nasopharyngeal swab in viral transport for:
 - influenza PCR;
 - respiratory virus culture;
 - direct antigen testing.
- CBC and differential.
- Liver function tests.
- Stool for viral studies (only if the patient has diarrhea).
- Arrange for other testing as recommended by MHO and/or Infectious Disease (ID) Specialist.



⁷ http://sdcl-testviewer.ehealthsask.ca/

Severe Acute Respiratory Infection (SARI)

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Methods of Control/Role of Investigator

Infection control procedures are paramount. Contact, droplet and airborne precautions must be implemented as necessary for patients in health care facilities and should be done in consultation with Infection Control and MHO. Refer to Infection Prevention and Control Measures and Initial Management of Persons who May Be Infected with a Novel Respiratory Virus.

Prevention and Education

Refer to the <u>Respiratory and Direct Contact Introduction and General Considerations</u> section of the manual that highlights topics for client education and provides information on high-risk groups and activities.

Refer to Vector-Borne and Zoonotic Diseases Avian Influenza section of the manual for additional prevention measures if poultry is involved as a host or source of infection.

SARI alerts should trigger MHOs to inform clinicians about the SARI screening tool and reinforce the "Think, Tell and Test" message.

- <u>THINK</u> about the possibility of an emerging respiratory infection, e.g., novel respiratory virus and how the spread can be prevented (implementation of appropriate infection control measures).
- TELL the local MHO and local infection control and consult with ID Specialist.
- <u>TEST</u> for pathogens only after appropriate consultation with the MHO and ID Specialist and based on clinical and epidemiologic symptoms.

Refer to Specimen Collection and Transport above, Attachment – Severe Acute
Respiratory Illness (SARI) Screening Tool or Laboratory Testing for Persons Who May
Be Infected with a Novel Respiratory Virus.

- Educate cases and contacts on the appropriate infection control measures that must be taken to reduce the spread.
- Provide education and instructions for staff who have cared for the case before appropriate precautions were implemented (i.e., had unprotected close contact with the case). This should include specific advice on how to self-monitor for fever and symptoms of respiratory illness for 14 days.



Severe Acute Respiratory Infection (SARI)

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Management

I. Case

Contact, droplet and airborne precautions must be implemented as necessary for all clients being investigated for SARI.

History

- Complete the <u>Attachment Severe Acute Respiratory Illness (SARI) Screening Tool</u> and Attachment Emerging Respiratory Pathogens and Severe Acute Respiratory Infection (SARI) Case Report.
- If person-to-person spread is typical for the suspected organism, identify those who may have been exposed to this case and follow-up as per Contact Investigation below.
- If the case was severely ill with the respiratory illness during air travel (i.e., on return to Canada), then the MHO should contact Health Canada's Centre for Emergency Preparedness and Response (CEPR), to request passenger contact information (e.g., airplane manifest). Follow-up of passengers may be considered if the case meets the SARI case definition and there is an identified concern of SARI globally and travel exposure occurred during the incubation period (within 14 days prior to the onset of illness), or the case is found to have another illness with significant public health implications.

Immunization

• Review immunization history specifically for Pneu-P-23 (pneumococcal 23 polysaccharide vaccine) and Influenza. If high-risk, offer as appropriate.

Treatment/Supportive Therapy

• Consult with ID Specialist.

Exclusion

- The period of exclusion will be based on the specific organism.
- While laboratory results are pending, appropriate infection control measures should be implemented including exclusion where appropriate.



Severe Acute Respiratory Infection (SARI)

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Referrals

- All SARI cases should be managed in consultation with the ID specialist and MHO.
- If no organism is identified, consultation with colleagues to determine further action is recommended.

II. Contacts/Contact Investigation

<u>Close Contact</u> means having cared for, lived with, or had face-to-face (within 1 metre) contact with, or has had direct contact with respiratory secretions and/or body fluids of a person with SARI (Public Health Agency of Canada, 2003).

- Household contacts, intimate contacts and health care providers should be the initial priority.
- Follow-up of the other close contacts should occur if the contacts can be reached within 14 days of their last contact with an infectious case. 8

The extent of investigation for remote contacts is dependent on the extent of illness in the close contacts and specific organism and will be directed by the MHO. See Attachment – Sample Severe Acute Respiratory Infection Contact Management Form.

Testing

• Consult with MHO for recommendations.

Prophylaxis/Immunization

• Review immunization history for contacts. The opportunity should be taken to catch up on immunizations for which the contact meets the eligibility criteria.

⁸ This recommendation takes into account the need to prioritize limited public health resources. It is acknowledged that some cases may be symptomatic and missed if no attempt is made to reach potentially ill contacts identified beyond the 14-day time frame. Therefore this should be considered a reasonable approach to contact management and should not preclude any jurisdiction from undertaking a more complete contact investigation.



Severe Acute Respiratory Infection (SARI)

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Education

- Public health should ensure that contacts receive education/instructions regarding
 infection control measures, self-monitoring, and who to contact if they become ill
 with respiratory symptoms. This should include informing the contact that if they
 develop symptoms (i.e., fever, cough or difficulty breathing), they should do the
 following:
 - Phone their personal physician so that decisions regarding the need for a clinical assessment can be individualized.
 - Health care providers should be asked to check in with their respective occupational health departments prior to returning to work.
 - Hospital/home isolation⁹ may be recommended until symptoms have resolved/returned to baseline.

Exclusion

- If the close contact is **symptomatic** (i.e., has fever, cough or difficulty breathing), manage as a case.
- No exclusion recommended if the close contact is **asymptomatic** (i.e., is afebrile and has no respiratory symptoms that are different from their baseline status):
 - Self-monitor for fever and new respiratory symptoms for 14 days following last contact with the case.

III. Environment

Child Care Centres/Institutional Control Measures

- Facilities should promptly initiate contact, droplet and airborne precautions (in addition to Routine/Standard Precautions) and consult their local infection control policies. Infection Control and the MHO should be consulted on all SARI cases.
- Patients with suspected SARI should be moved to a designated isolation room ASAP (or negative pressure room if available).

⁹ The symptomatic contact should be isolated in their home unless hospitalization is clinically indicated. These individuals would be instructed to stay home from work/school/other activities, wash their hands frequently and avoid direct face to face contact with others for the duration of their illness. The extent of the isolation requirements should be based on the severity of illness in the case, the composition of the household (e.g., presence of immunocompromised individuals) and any available evidence regarding communicability and ease of transmission.



Severe Acute Respiratory Infection (SARI)

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Refer to provincial and national guidelines and discuss with the local MHO or Infection Control Practitioner for Infection Control guidance. Initial precautions may be more conservative and include airborne as well as contact and droplet precautions.

Epidemic Measures

If SARI cases are identified in a health care facility, it is important to heighten surveillance to assist in early identification and implementation of control measures and further outbreak control measures as required.

PHAC may be in a position to provide direction. Saskatchewan Ministry of Health will participate in communication messages and provide direction. Specific measures include:

- Use media to clearly inform the general public about the disease, risk of transmission/infection, signs and symptoms, and how to avoid contact with cases.
- Provide HealthLine with updated information to address concerns from the public.
- Ensure that health care workers are well informed of infection control measures and have appropriate facilities for triage.
- Promote the location of the triage facilities to the public.



Severe Acute Respiratory Infection (SARI)

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Attachment – Severe Acute Respiratory Illness (SARI) Screening Tool Page 1 of 2 2020 01 20

Please see the following pages for the Severe Acute Respiratory Illness (SARI) Screening Tool.

SEVERE ACUTE RESPIRATORY ILLNESS (SARI)* SCREENING TOOL

Place surgical mask on all patients presenting with severe acute respiratory symptoms (unless the

patient's clinical condition will be compromised by wearing the mask).

PHYSICIANS to complete

Date/Time

For all persons with severe acute respiratory illness* presenting to the Emergency Department or admitted to Hospital.

*SARI may be caused by respiratory pathogens of known or unknown origin including novel respiratory viruses (Avian Influenza H7N9, H5N1, Novel Coronaviruses e.g. MERS CoV, Wuhan, etc.)

Addressograph/Patient Name:

		Ensure that it remains in place during any transportation of the patient for medical investigations/examinations, including Chest X-ray			
СОМР	COMPLETE THE FOLLOWING SCREENING QUESTIONS - Indicating Yes or No for each of the criteria				
PATIE	NT pre	sents with SARI-defining features:			
Yes	No	Fever (over >38° C), and			
Yes	No	Cough or breathing difficulty, and			
Yes	No	Radiographic evidence of infiltrates consistent with pneumonia or Respiratory Distress Syndrome			
NOT	E: If an	swered "NO" to any of the above, there is no need to proceed with this screening tool.			
IN THE	14 DAY	S BEFORE THE ONSET OF SYMPTOMS, WERE ANY OF THE FOLLOWING PRESENT:			
Yes	No	1.a) Close contact with a suspect or probable case of SARI [Close contact means having cared for, lived with, or had face to face (within 2 meter or having had direct contact with respiratory secretions and/or body fluids of a perso			
Yes	No	1.b) Travel to a country where there is a Public Health Agency of Canada public health notice of respiratory illness in effect: http://www.phac-aspc.gc.ca/phn-asp/index-eng.php			
Yes	No	1.c) Recent exposure/close contact to a potential source of a SARI which may include reports of illness or die offs in domestic poultry flocks or illness in other animal vectors such as camels or swine.			
Yes	No	2. Current illness is inconsistent with other known cause.			
-	If you answered "NO" to questions 1 (a, b & c) and 2 The patient has not had any exposures of concern, and does have another explanation for their symptoms If you answered "YES" to questions Initiate Contact & Droplet Precautions (in addition to Routine Practices)				
·	1 (a, b or c) or 2 with negative pressure (AIIR). If not available, place in a private room with the door closed.				
◆ Ev	1. THINK infection control • Everyone entering the room should observe hand hygiene, airborne and contact precautions (N95 pone of pone pone). Not pone				
2. TELL your Medical Health Officer (Regional contact ##) or if after hours, the MHO on call. ### The MHO will call Roy Romanow Provincial Laboratory (RRPL) to expedite STAT testing (306-798-1234).			Done	Not Done	
3. TEL	3. TELL Infection Control (Monday to Friday) – insert Regional contact ##			Not Done	

Nasopharyngeal and oropharyngeal swab in viral transport media

4. **CONSULT** an Infectious Disease Specialist – insert Regional contact ##

5. TEST - Collect specimens and clearly mark specimens "URGENT: for SARI Screen"

- CXR
- CBC and differential
- Endotracheal secretions, Broncoalveolar lavage (BAL)

Collect the specimens when clinically indicated

- Serum for Mycoplasma pneumoniae and Chlamydia pneumoniae serology.
- If patient has diarrhea, send stool for viral studies.
- Arrange other testing as recommended by MHO and/or ID specialist (document on this form).
- Local lab to contact RRPL and confirm details related to delivery/arrival for the STAT specimens.

Not

Not

Done

Done

Done

Done

Liver function tests

Blood culture

Sputum C & S

Notification Timeline:

From Lab/Practitioner to Public Health: Within 48 hours.

From Public Health to Ministry of Health: Immediate for known outbreaks. Individual cases are not reportable to the Ministry.

Public Health Follow-up Timeline: Less than 48 hours for prenatal and neonatal cases and contacts.

Information

Table 1: Case Definition (Public Health Agency of Canada, 2008)

Confirmed case	Clinical evidence of illness ¹ and laboratory confirmation of infection:		
	 isolation or direct antigen detection of varicella-zoster virus (VZV) from an appropriate clinical specimen OR detection of VZV DNA 		
	OR		
	 seroconversion or a significant rise (e.g., fourfold or greater) by any standard serologic assay in varicella-zoster IgG titre between acute and convalescent sera OR 		
	 positive serologic test for varicella-zoster IgM antibody OR 		
	 clinical evidence of illness¹ in a person with an epidemiologic link to a laboratory-confirmed case of chickenpox or VZV infection. 		
Probable Case	Clinical evidence of illness ¹ in the absence of laboratory confirmation or epidemiologic link to a laboratory confirmed		
¹ Clinical illness is share	ctarized by a rash with rapid evolution of macules to papules vesicles and		

¹Clinical illness is characterized by a rash with rapid evolution of macules to papules, vesicles, and crusts; all stages are simultaneously present; lesions are superficial and may appear in crops.



^{*}Refer to Specimen Collection and Transport for details on appropriate clinical specimens.

Causative Agent

Human herpesvirus3 (alpha); member of VZV (Heymann, 2015).

Symptoms

Varicella may or may not begin with a prodromal period. The prodromal period, when present, is characterized by fever, malaise and upper respiratory tract infection followed by the characteristic lesions. The lesions appear in successive crops over the first 2-5 days of the rash and tend to develop on the trunk and face, with progression to the extremities. They progress rapidly from macules to papules, vesicles and crusts, all stages are simultaneously present; lesions are superficial, distribution is centrifugal. Ulcerated lesions may also be present on mucous membranes including the oropharynx, upper respiratory tract, conjunctiva and rectal and vaginal mucosa. In adults, these symptoms may be more severe (Mandell, Bennett & Dolin, 2000).

Complications

Varicella is generally considered a mild infection; however, 5-10% of otherwise healthy children may develop complications that may be fatal. Complications may include pneumonia, secondary bacterial infections, soft tissue infections, bacteraemia, septicemia, septic arthritis, necrotizing fasciitis, toxic shock-like syndrome, thrombocytopenia, cerebellar ataxia, encephalitis and hepatitis (American Academy of Pediatrics, 2015; Heymann, 2015).

Primary varicella is a more severe disease in adults, with a case fatality rate 10 to 30 times higher than in children. Moreover, in both adults and children, the majority who die of varicella have no identifiable risk factor for severe disease (Health Canada, 1999).

Neonates who develop varicella at 5-10 days are at increased risk for severe generalized varicella. The case-fatality rate for neonates whose mother developed varicella five days before delivery to within two days following delivery and who did not receive Varicella- Zoster Immune Globulin (Varlg) or antiviral therapy can reach 30% (Heymann, 2015).

Incubation Period

Usually 14-16 days but it can be as early as 10 days or as late as 21 days (Heymann, 2015).



Reservoir/Source

Humans.

Mode of Transmission

- Direct or indirect contact of oral or nasal mucous membranes with respiratory secretions or vesicular fluid.
- Inhalation of airborne virus.
- Indirect transmission may occur through contact with respiratory secretions or discharge from lesions on freshly soiled linens or towels.
- Transmission of vaccine virus is rare (Public Health Agency of Canada, 2006).
- Transmission can occur from direct contact with fluids from localized shingles lesions but is rare if the lesions are covered. Disseminated zoster can be transmitted by airborne route. (Household transmission rates have been noted to be approximately 15% [Stankus, Dlugopolski & Packer, 2000]).
- In utero infection through transplacental passage during maternal infection.

Risk Groups/Risk Factors

- Neonates born to non-immune mothers.
- Newborns of mothers who develop varicella between five days prior to delivery and 48 hours after the delivery.
- Infants.
- Adolescents (American Academy of Pediatrics, 2015).
- Individuals with chronic cutaneous/pulmonary disorder (American Academy of Pediatrics, 2015).
- Pregnant women who have never had varicella vaccine, varicella disease or shingles.
- Immunocompromised individuals.
- Cancer patients, especially lymphoid tissue, with or without steroid therapy.

Period of Communicability

- From one to two days before onset of rash and continuing until all lesions are crusted, approximately five days (Heymann, 2015; American Academy of Pediatrics, 2015).
- In immuno-competent individuals most virus replication has stopped by 72 hours after onset of the rash. The time may be longer in immunocompromised individuals (Mandell et al., 2000).



Specimen Collection and Transport

- Swabs from the base of a freshly de-roofed lesion for culture and direct fluorescent antibody (DFA) or polymerase chain reaction (PCR).
- Cerebrospinal fluid (CSF) for culture or PCR.
- Blood for serology.

Methods of Control/Role of Investigator

Prevention and Education

Refer to the <u>Respiratory and Direct Contact Introduction and General Considerations</u> section of the manual that highlights topics for client education that should be considered as well as provides information on high-risk groups and activities.

Immunization

Immunize infants, children, and adults according to the recommended schedules in the Saskatchewan Immunization Manual.¹

Education

- Education should be provided regarding respiratory etiquette, hand hygiene and other measures to prevent transmission.
- Educate the public about the disease and the need for active immunization. Immunization information fact sheets can be used to guide discussion.

Management

I. Case

History

- Assess risk factors and exposure history. The source of infection could be a case of varicella or herpes zoster (rarely unless disseminated).
- Identify contacts (refer to contact definition).

Immunization

Assess immunization history.



¹ http://www.ehealthsask.<u>ca/services/manuals/Pages/SIM.aspx</u>.

Education

- Practicing good hand hygiene.
- Not sharing personal items such as drinking glasses, eating utensils, or towels.
- Respiratory etiquette.
- Cases should avoid contact with high risk individuals who have not yet been exposed.

Treatment/Supportive Therapy

- Supportive therapy as indicated.
- Treatment with antivirals has a limited window of opportunity to affect the
 outcome of varicella-zoster infection. Acyclovir therapy initiated within 24 hours
 after onset of the rash is effective in accelerating skin lesion healing and can be
 used for generally healthy population (at increased risk of moderate to severe
 varicella) as soon as possible after rash onset (Public Health Agency of Canada,
 2006).

Exclusion

- Cases should not be cared for by susceptible persons.
- Children with chickenpox may remain in school/daycare as long as they are feeling well enough to take part in normal activities (Canadian Pediatric Society, 2016).
 - Exclusion for five days after the appearance of the rash should still be considered when the child has severe illness or is going into a new setting where the classmates have not already been exposed.
- In health care facilities, the appropriate infection control measures should be implemented because of the risk of serious varicella in susceptible immunocompromised individuals. Refer to <u>Health Facility Control Measures</u>.
- Air travel is not recommended until lesions are crusted.
- Swimming in public pools is not recommended until lesions have healed and crusts are no longer present (Alberta Health and Wellness, 2008).

Referrals

Not applicable.



II. Contacts/Contact Investigation

<u>Identify susceptible contacts</u> with <u>significant exposure</u> (see Contact Definition).

Table 2: Contact Definition

Contact	Anyone who shared the same airspace with a case	
Contact	·	
	during the infectious period (48 hours before to five days	
	after onset of rash).	
Significant Exposure ²	<u>Varicella</u>	
(Public Health Agency of	 Continuous household contact (living in the same 	
Canada, 2016 and 2013) dwelling) with a person with varicella.		
	Close contact with an infectious person, such as	
	close indoor contact (e.g., in the same room) or	
	face-to-face contact ³ .	
	 Being in the same hospital room for >1 hour, or >15 	
	minutes of face-to-face contact, with a patient with	
	varicella.	
	 Touching the lesions of a person with active 	
	varicella.	
	Zoster	
		
	Touching a zoster rash, exposed lesion or vesicle	
	fluid or articles freshly soiled by discharges from	
	vesicles;	
	 Contact with an individual who has disseminated zoster; 	
	Contact with articles freshly soiled by mucous	
	membrane secretions of a person with	
	disseminated zoster; or	
	·	
	Exposure to an immunocompromised person with	
	localized zoster anywhere on the body because	
	their viral shedding may be greater.	

² Verbal history of infection is not acceptable following a significant exposure to varicella in individuals at <u>high risk for varicella complications</u> and cannot be accepted as evidence of immunity

³ Experts differ in their opinion about the duration of contact; some suggest five minutes and others up to one hour, but do agree that it does not include transitory contact (Centers for Disease Control and Prevention, 2016)



Susceptible Contacts

- Newborns of mothers who develop varicella between five days prior to delivery and 48 hours (two days) after delivery.
- Hematopoietic stem cell transplant (HSCT)
 recipients regardless of pre-transplant varicella
 immune status or history of varicella disease or
 vaccination.
- Immunocompromised individuals.
- Hospitalized patients, especially premature infants.
 - Preterm infants >/= 28 weeks gestation whose mother lacks a reliable history of chickenpox or serologic immunity (American Academy of Pediatrics, 2009).
 - ➤ Preterm infants < 28 weeks gestation or birth weight of 1,000 g or less, regardless of the maternal history of chickenpox or serostatus (American Academy of Pediatrics, 2009).
- Pregnant women who do not have documentation of immunity to varicella (routine prenatal screening includes varicella immunity).
- Healthy individuals who (Public Health Agency of Canada, 2015):
 - Do not report having a health care provider diagnosed or self-diagnosed history of varicella or zoster prior to implementation of a one dose varicella program⁴
 - Do not have documented evidence of immunization with two doses of varicella containing vaccine, or
 - Do not have previous laboratory evidence of immunity⁵ to varicella.

⁵ Laboratory testing should be conducted only once in a lifetime. If a person has been found to be seropositive, it is not necessary to test again.



⁴ One-dose varicella program was implemented in Saskatchewan on January 1, 2005

Education

- Close contacts of confirmed cases should be educated about varicella and its signs and symptoms.
- They should also be advised that varicella is communicable to others long before the rash appears.
- Adult contacts (including pregnant women), and any individual with immunocompromising conditions, should be advised to see a physician if early signs and symptoms appear.
- Household contacts of confirmed and probable cases should avoid contact with susceptible/high risk groups/individuals during the incubation period.

History

- Assess risk factors.
- History of vaccination.
- History of varicella disease and/or shingles.

Preventive Measures

Immunize individuals as per the Saskatchewan Immunization Manual⁶.

Prophylaxis Immunization

Although varicella vaccine has been shown to be effective in preventing or reducing the severity of the disease if given to susceptible individuals within 72 hours and no longer than five days after exposure, Saskatchewan Ministry of Health, at this time, does not routinely provide publicly funded immunization for contacts of chickenpox. The exception is children who fall into the target group who have not yet been immunized, and who do not have contraindications to immunization.

Immune Globulin Prophylaxis

Susceptible individuals at higher risk for severe disease (see list below), should be evaluated immediately for administration of Varlg. The National Advisory Committee for Immunization (NACI) (2016) recommends:

 For optimum benefit, Varlg should be administered as soon as possible (ideally within 96 hours) following <u>first</u> exposure.



⁶ http://www.ehealthsask.<u>ca/services/manuals/Pages/SIM.aspx</u>

- In instances of prolonged exposures, where the exact timing of transmission may be unknown, it may be used within 96 hours of the most recent exposure.
- If more than 96 hours but less than 10 days have elapsed since the last exposure, the susceptible high-risk individuals' clinician may determine that Varlg would be useful to attenuate (rather than prevent) disease. The benefit of administering Varlg after 96 hours is uncertain.

Dosage: 125 units/10 kg of body weight, to a maximum of 625 units IM. Refer to Appendix D – Publicly Funded Medications for Chemoprophylaxis/Treatment for information on how to access Varlg from Canadian Blood Services.

NACI recommends Varig for the following susceptible <u>high-risk groups</u> after exposure to VZV (Public Health Agency of Canada, 2016):

- 1. Susceptible pregnant women.
- 2. Newborn infants of mothers who have varicella that began during the five days before to 48 hours after delivery.
- 3. Selected neonates in neonatal or pediatric intensive care units for the management of significant varicella exposure in consultation with the infectious diseases/infection control specialist.
- 4. Susceptible immunocompromised individuals, including (including those with HIV with CD4 cell count < 200 × 106/L or CD4 percentage < 15%) and HSCT recipients regardless of pre-transplant varicella immune status or history of varicella disease or vaccination.

Testing

Adolescents and adults who have a negative or uncertain past history of varicella and no documentation of vaccination should have serologic tests to establish susceptibility, since as many as 70 to 95% of such individuals have immunity to varicella. However, delays in obtaining test results should not delay appropriate post-exposure varicella management (Public Health Agency of Canada, 2006).

Chemoprophylaxis

Clinicians may want to consult with specialists to determine if and when acyclovir should be used for specific contacts in circumstances where the timeframe for VarIg has elapsed.

Acyclovir is generally not recommended for immunocompetent contacts.



Treatment

Antiviral drugs such as acyclovir appear useful in preventing or modifying varicella in exposed individuals if given within a week of exposure.

Exclusion

Susceptible caregivers, including healthcare workers (HCWs) exposed to chickenpox should be excluded from contact with high-risk patients from 8-21 days after exposure. Extend to 28 days if Varlg was given as it may prolong the incubation period if it is unable to fully protect against infection in the susceptible person who received it (Health Canada, 2002).

III. Environment

Prevent the spread of infection by using a household cleaner to wash any articles soiled with fluid from chickenpox blisters. Keep the infected person away from others who have not had chickenpox.

Health Facilities Control Measures

- HCWs should have proof of immunity or previous immunization assessed upon employment. Refer to the Saskatchewan Immunization Manual⁷ – Chapter 7: Immunization of Special Populations, Section 3.2 Health Care Workers and other relevant Saskatchewan Ministry of Health policies/memos.
- A suspected or confirmed case of varicella occurring within a facility must be reported immediately to the local public health office and to infection control.
- Strict enforcement of infection control practices (routine practices as well as contact and airborne precautions) should be taken for a minimum of five days and until all lesions are crusted (Health Canada, 2002 and Health Canada, 1999).
- Immunocompromised cases should be isolated with contact and airborne precautions for the duration of their illness which can be up to a week (American Academy of Pediatrics, 2015).
- Provide varicella vaccine or Varlg to susceptible contacts as described in contact management.



⁷ http://www.ehealthsask.ca/services/manuals/Pages/SIM.aspx.

- Susceptible contacts who are HCWs should be excluded from working with highrisk susceptible patients during the potential period of communicability (from
 eight days, after first exposure to 21 days from last exposure to an infectious
 client) or to day 28 for those who received immune globulin as it may prolong
 the incubation period (Public Health Agency of Canada, 2006).
- Health care facilities may, after consultation with the Medical Health Officer (MHO), provide HCWs immunization and other follow up. HCWs must be instructed to call public health if they develop any signs or symptoms suggestive of varicella.
- HCWs who are symptomatic should be excluded from work until all lesions are dry and crusted and no new lesions are forming.
- Occupational Health (OH) should not exclude HCWs with a localized, postimmunization varicella-like rash that can be covered with an occlusive dressing.
- OH should exclude HCWs with a postimmunization varicella-like rash if the rash cannot be covered and if the HCWs are involved in the care of high-risk patients, (e.g., immunocompromised and newborn patients) for the duration of the rash.
- OH should inform Infection Control as soon as possible of a suspected or confirmed case.

Epidemic Measures

- Follow as per case and contact management.
- The use of varicella vaccine may be considered in the management of outbreaks in consultation with Saskatchewan Ministry of Health.



Respiratory and Direct Contact Section 2 - 210 – Varicella (Chickenpox) Page **12** of **15** 2017 05 04

Revisions

Date	Change
March 2016	Updated recommendations on use of VarIg based on NACI
	Statement 2015.
March 2017	Updated definition of susceptible individuals based on NACI
	Statement (2015) and included contact to zoster under significant
	exposure definition as per PHAC (2015).
	References reaffirmed or updated as necessary.



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Varicella Data Collection Worksheet Panorama QA complete: ☐ Yes □No Panorama Client ID: Please complete all sections. Panorama Investigation ID: __ Initials: A) CLIENT INFORMATION LHN -> SUBJECT -> CLIENT DETAILS -> PERSONAL INFORMATION Last Name: First Name: and Middle Name: Alternate Name (Goes by): Preferred Communication Method: (specify - i.e. DOB: YYYY / MM / DD Age: ____ Health Card Province: ___ home phone, text): Health Card Number (PHN): Phone #: □ Primary Home: Email Address: □Work □Personal ☐ Mobile contact: ☐ Workplace: Place of Employment/School: Gender: ☐ Male □ Other ☐ Female □ Unknown Address Type: □ No fixed □ Postal Address □ Primary Home □ Temporary □ Legal Land Description Alternate Contact: _____ Mailing (Postal address): Relationship: Alt. Contact phone: ___ Street Address or FN Community (Primary Home): Address at time of infection if not the same: LHN-> SUBJECT SUMMARY-> RESPIRATORY & DIRECT CONTACT ENCOUNTER GROUP->CREATE INVESTIGATION **B)** INVESTIGATION INFORMATION Disease Summary Classification: Classification: LAB TEST INFORMATION: Date **CONTACT** Date CASE Date specimen collected: ☐ Confirmed YYYY / MM / DD □ Contact YYYY / MM / DD YYYY / MM / DD ☐ Does Not Meet Case Definition YYYY / MM / DD ☐ Not a Contact YYYY / MM / DD YYYY / MM / DD Person Under Investigation YYYY / MM / DD ☐ Person Under Investigation Probable YYYY / MM / DD YYYY / MM / DD ☐ Suspect Disposition: **FOLLOW UP:** ☐ In progress YYYY / MM / DD □ Complete YYYY / MM / DD \square Incomplete - Declined YYYY / MM / DD □ Not required YYYY / MM / DD ☐ Incomplete – Lost contact YYYY / MM / DD ☐ Referred – Out of province YYYY / MM / DD ☐ Incomplete – Unable to locate YYYY / MM / DD (specify where) REPORTING NOTIFICATION Location: Name of Attending Physician or Nurse: Date Received (Public Health): YYYY / MM / DD Physician/Nurse Phone number: Type of Reporting Source: Health Care Facility □ Lab Report ☐ Nurse Practitioner ☐ Physician □ Other_ INVESTIGATION->DISEASE SUMMARY (UPDATE)->DISEASE EVENT HISTORY C) DISEASE EVENT HISTORY

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☐ Reactivation

Site / Presentation: ☐ Severe ☐ Neonatal ☐ Case with high risk contacts

☐ Acute

Staging:

Varicella Data Collection Worksheet

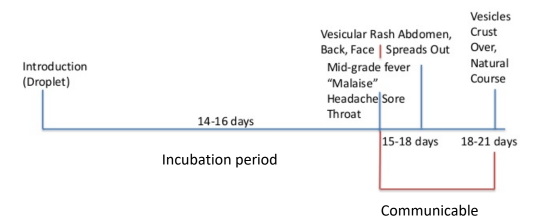
Please complete all sections.

Panorama Client ID:	
Panorama Investigation ID:	

D) SIGNS & SYMPTOMS (Bold text = part of case definition)

LHN-> INVESTIGATION->SIGNS & SYMPTOMS

Description	Yes	Date of	Description	Yes	Date of
	Date of onset	recovery		Date of onset	recovery
Fever		YYYY / MMM / DD	Rash - crusted lesions or scabs		YYYY / MMM / DD
Lesion - less than 50 lesions (Mild)		YYYY / MMM / DD	Rash - herpes zoster (shingles)		YYYY / MMM / DD
Lesion - 50 to 249 lesions (Mild - moderate)		YYYY / MMM / DD	Rash - itchy		YYYY / MMM / DD
Lesion - 250 to 499 lesions (Moderate)		YYYY / MMM / DD	Rash - macules, papules, and vesicles		YYYY / MMM / DD
Lesion - 500 or more lesions (Severe)		YYYY / MMM / DD	Rash - painful		YYYY / MMM / DD
Lesions - conjunctiva		YYYY / MMM / DD	Rash - ulcerated lesions		YYYY / MMM / DD
Lesions - mucous membrane - ulcerated		YYYY / MMM / DD	Rash - unilateral red painful blisters		YYYY / MMM / DD
Malaise		YYYY / MMM / DD	Infection - upper respiratory tract		YYYY / MMM / DD



E) INCUBATION AND COMMUNICABILITY

LHN-> INVESTIGATION->INCUBATION & COMMUNICABILITY

z, mees, men, and commenter state.			
Incubation for Case (period for acquisition):			
Earliest Possible Exposure Date: YYYY / MM / DD	Latest Possible Exposure Date: YYYY / MM / DD		
Exposure Calculation details:			
Exposure culculation details.			
Communicability for Case (period for transmission):			
Earliest Possible Communicability Date: YYYY / MM / DD	Latest Possible Communicability Date: YYYY / MM / DD		
Communicability Calculation Details:			

F) RISK FACTORS (RF followed by + impact the Immunization Forecaster)

LHN-> SUBJECT->RISK FACTORS

DESCRIPTION	YES	N – No	DESCRIPTION	YES	N – No
		NA – not asked			NA – not asked
		U - Unknown			U - Unknown
Contact to a known case (Add'l Info)	YYYY / MM / DD		Special Population - Pregnancy	YYYY / MM / DD	
	AE				
Immunocompromised - Related to			Travel - Outside of Canada (specify)		
underlying disease or treatment			, ,		
Occupation - Health Care Worker -	TE		Travel - Outside of Saskatchewan, but		
IOM Risk Factor			within Canada (specify)		
Special Population - Infant born to an	YYYY / MM / DD				
infected mother					

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Varicella Data Collection Worksheet

Please complete all sections.

Panorama Client ID:	
anorama Investigation ID:	

G) IMMUNIZATIO	ON HISTORY INTERF	PRETATION SUM	MARY LHIN	-> INVESTIGATION-> IMMUNIZA	ATION HISTORY INTERPRET	TATION SUMMARY
Interpretation Da	ate: YYY	Y / MM / DD				
Interpretation of	Disease Immunity:	: 🗆 IOM - Ft	ully immunized (for age)	☐ IOM - Partially in	mmunized	
□ IOM – Unimm	unized	□ IOM - U	nclear immunization history	Valid doses received:	Doses needed:	
Reason:	☐ IOM - Interp	pretation of histo	ory by investigator			
H) TREATMENT				LHN -> INVESTIGATION	-> MEDICATIONS->MEDIC/	ATIONS SUMMARY
Medication (Pana	orama = Other Med	ls) :				_
Prescribed by:				Started on: YYYY / MM / DI		
i) intervention	N		LHN -> I	NVESTIGATION->TREATMENT & I	NTERVENTIONS->INTERVE	ENTION SUMMARY
Intervention Type	e and Sub Type:					
Assessment: ☐ Assessed for conception of the c	ant or < 1 year of ag	ge)	YYYY / MM / DD	Immunization: ☐ Eligible immunizations recon ☐ Disease-specific immunizatio ☐ Disease-specific immunizatio	on recommended YYYY /	/ MM / DD / MM / DD / MM / DD
Other Investigati	_			Investigator name		
☐ Investigator No Communication:	otes	ee Document Ma	nnagement			
_	nication (see Invest	igator Notes)	YYYY / MM / DD	Referral: ☐ Other (specify)	VVVV	/ MM / DD
Investigator name ☐ Letter (See Do Investigator name	cument Manageme	ent)	YYYY / MM / DD	Investigator name		, IVIIVI , DD
General: Investig ☐ Disease-Info/P	ator name		YYYY/ MM / DD	Testing: ☐ Laboratory testing recomme	ndod VVVV	/ MM / DD
· ·	rev-Cont/Assess'd f	or Contacts	YYYY/ MM / DD	Investigator name	inded 1111 /	/ IVIIVI / DD
Education/couns Prevention/Co Disease inform		name	YYYY / MM / DD YYYY / MM / DD			
Exclusion: Invest ☐ Daycare YY ☐ School YY	0	□ Preschool □ Work	YYYY / MM / DD YYYY / MM / DD			
Date	Intervention subtype	Comments			Next follow-up Date	Initials
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	
YYYY/MM/DD					YYYY/MM/DD	

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Varicella Data Collection Worksheet

Please complete all sections.

Panorama Client ID:	
Panorama Investigation ID:	

J) OUTCOMES (op	LI	.HN-> INVESTIGATION-> OUTCOMES			
☐ Recovered ☐ Fatal	ed/recovering YYYY / MM YYYY / MM YYYY / MM Fatal was selected)	/ DD □ Intubation /ventilation YYYY / MM / D / DD □ Other	□ Unkno	talization YYYY / MM /	
K) Transmission Transmission Event ID	Events Exposure Name	LHN -> INVESTIGATION-> EXPOSURE SUMMARY -> Setting type	Date/Time		
		☐ Congregate/Communal living ☐ Health Care setting ☐ Household Exposure			
		☐ Congregate/Communal living ☐ Health Care setting ☐ Household Exposure			
		☐ Congregate/Communal living ☐ Health Care setting ☐ Household Exposure			
	varicella Contacts – Inv ID#	☐ Multiple Settings	to		
•	-> INVESTIGATION-> EXPOS	URE SUMMARY -> TRANSMISSION EVENT SUMMARY -> TE H of individuals [including groups that do not require 1:1 follow		NKNOWN/ANONYMOUS Date initial report com	

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