Chickenpox (Varicella)

Disease Plan

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Last updated: July 1, 2016, by Jeffrey Eason.

Questions about this disease plan?

Contact the Utah Department of Health Bureau of Epidemiology: 801-538-6191.
WHY IS CHICKENPOX IMPORTANT TO PUBLIC HEALTH?

Chickenpox is an acute, highly contagious infectious disease. It is caused by the varicella-zoster virus (VZV) and causes a blister-like rash, itching, fatigue, and fever. VZV spreads in the air through coughing or sneezing. It can also be spread by touching or breathing in the virus particles that come from chickenpox lesions. VZV can be serious, especially in babies, adults, and people with weakened immune systems, but is generally a mild disease in healthy individuals. It spreads easily from infected people to others who have never had chickenpox or received the chickenpox vaccine. Morbidity and mortality associated with VZV has decreased dramatically since the introduction of the varicella vaccine in 1995. Chickenpox occurs worldwide and occurs throughout the year in temperate regions, but the incidence typically peaks in the months of March through May. Continued surveillance is needed to evaluate the effectiveness of immunization policy in controlling disease and detecting and controlling outbreaks. Chickenpox surveillance data is needed to document and monitor the impact of the Chickenpox vaccination program on VZV incidence, morbidity, and mortality, evaluate the effectiveness of prevention strategies, and to evaluate vaccine effectiveness under conditions of routine use.

DISEASE AND EPIDEMIOLOGY

Clinical Description
VZV causes two distinctly different clinical diseases, chickenpox and shingles. Below clinical descriptions are provided for chickenpox, including breakthrough VZV and shingles.

1) Chickenpox
Primary infection with VZV is known as chickenpox. It often starts with a mild prodrome of low-grade fever and malaise. After a day or two of prodrome or as the first evidence of illness, the generalized, pruritic (itchy) rash begins along with constitutional symptoms (fever, anorexia, listlessness).

The rash initially appears on the face and trunk, and then spreads to the arms and legs. Lesions may also occur on mucous membranes of the oropharynx, respiratory tract, genital area, conjunctiva, and the cornea. Lesions develop into superficial, delicate vesicles that break, leaving open sores that dry and crust over into brown scabs. Chickenpox sores appear in crops, with lesions (typically 250–500 lesions) in several stages of development (maculopapules, vesicles, scabs) present at the same time. Serious complications can occur, most commonly in infants, adults, and immunocompromised people. Complications include
secondary bacterial infections of skin lesions, pneumonia, cerebellar ataxia, encephalitis, and hemorrhagic conditions; rarely (in about 1 in 40,000 cases), these complications may result in death.

The acute disease lasts from 3-5 days, with resolution of the lesions taking 1-2 weeks.

2) Breakthrough varicella or mild varicella-like syndrome (MVLS)
Breakthrough varicella or MVLS refers to infection after exposure to wild-type virus in those who have been previously vaccinated. It can be defined as varicella infection that develops more than 42 days after vaccination. Studies have shown that breakthrough varicella or MVLS occurs in less than 1-3% of vaccinated children.

Most cases of breakthrough varicella/MVLS in vaccinated children are mild. Patients usually have fewer than 50 lesions and they usually do not present with a fever. Cases with breakthrough varicella/MVLS may have a maculopapular rash rather than a vesicular rash. Often, the lesions in breakthrough varicella/MVLS do not scab and crust over. However, it is estimated that as many as 25% of children who develop breakthrough varicella/MVLS may still present with symptoms similar to those in unvaccinated cases.

3) Shingles
Herpes zoster or shingles occurs when varicella-zoster virus reactivates from its latent state in a posterior dorsal root ganglion. Herpes zoster is common among older age groups or those who are immunosuppressed. About 20% of the population will have an episode of herpes zoster. This disease presents with vesicular eruptions, typically unilateral and in the dermatome (area of skin) served by a nerve branch, most often on the trunk. Shingles can be quite painful. Disease resolution takes several weeks to a month.

Causative Agent
Chickenpox is caused by the varicella-zoster virus, a member of the Herpesvirus group.

Differential Diagnosis
The differential diagnosis includes smallpox, herpes simplex virus (HSV), coxsackie viruses, and impetigo. Smallpox lesions start on the extremities, whereas chickenpox lesions start on the trunk. Smallpox lesions are all in the same stage of development (crop) versus chickenpox lesions that are in different stages of development. The last case of wild smallpox occurred in Somalia in 1977.

Laboratory Identification
Chickenpox is generally diagnosed clinically. Laboratory testing may be appropriate if the presentation is unusual. The clinical diagnosis of VZV in the U.S. has become increasingly challenging as a growing proportion of cases occur in vaccinated people, in whom disease is mild and rash is atypical. Although not routinely recommended, laboratory diagnosis is likely to become increasingly useful. For laboratory confirmation, fluids from skin lesions are the preferred specimen.
Lab test methods include:

**Serology**
This is the lowest cost alternative. Both IgG and IgM testing can be performed.
- IgG testing requires that an acute and convalescent serum sample be drawn, 10-14 days apart. Acute and convalescent specimens must be labeled as such; paired sera testing is preferred. Transport serum at 2-8°C. IgG testing has generally low sensitivity to reliably detect vaccine-induced immunity.
  - NOTE: A single IgG result cannot be used to confirm a VZV case.
- Because results early in the disease may be equivocal, if initial IgM testing is negative, a second sample should be collected 10-14 days after the first. Transport serum at 2-8°C.
  - NOTE: IgM testing is less sensitive than PCR testing on skin lesions. Tests may not be reliable and false negative IgM results are not uncommon. A positive IgM result, although suggestive of a primary infection, does not exclude re-infection or reactivation of latent VZV. Nevertheless, a positive result from a person with a rash is usually interpreted as laboratory confirmation of VZV.

**PCR**
Polymerase chain reaction (PCR) testing is the method of choice for laboratory diagnosis, but it may not be available or costly. Vesicular fluid and scabs are suitable specimens. Biopsy tissue and CSF can also be used as specimens for diagnosis. Vesicular fluid should be collected using a Dacron swab with a plastic shaft (wood, cotton, and calcium alginate may interfere with the test). Each sample should be placed in a snap cap tube or other suitable container. DO NOT place the specimen in transport medium. Samples for PCR may be transported at ambient temperature or at 2-8°C. PCR samples are not suitable for other testing methods. The sensitivity of the test is improved by submitting five specimens from different lesions.

**DFA**
This is a lower cost alternative to PCR, but is somewhat less sensitive. DFA is also more rapid and more sensitive than culture. Vesicular fluid, tissue, or skin scrapings are all acceptable specimens. The specimen should be immediately placed in viral transport media and be transported at 2-8°C.

**Culture**
Because the VZV is difficult to culture, culturing is not routinely recommended. Additionally, results may take up to a week and are not readily available. Viral culture is only recommended for cases of severe disease, particularly in the immunocompromised. It can distinguish VZV from HSV. Vesicular fluid, tissue, or skin scrapings are all acceptable specimens. The specimen should be immediately placed in viral transport media and be transported at 2-8°C.

**Tzanck Smear**
Tzanck smears of lesion specimens are inexpensive and can be performed at bedside, although they do not distinguish between VZV and HSV infections. Vesicle scraping and swab of lesion base (must include cells) are acceptable specimens. This test is less sensitive and accurate than the DFA. It is known to most dermatologists as a rapid test for diagnosing herpes virus...
infections. The Tzanck smear should not be used to replace more standard diagnostic techniques, but rather as an adjunct to the physical examination.

**Utah Public Health Laboratory (UPHL):** UPHL performs a Herpes PCR Panel (HSV and VZV) by PCR. If PCR testing is to be requested, UPHL should be contacted before specimen submission.

**Treatment**
Several antivirals are active against VZV, including acyclovir, valacyclovir, famciclovir, and foscarnet. Valacyclovir and famciclovir are approved for use only in adults. Acyclovir is available in an oral tablet or in intravenous form. These drugs may be beneficial if given within 24 hours of rash onset. The decision to initiate antiviral therapy depends upon the patient’s age, comorbid conditions, and clinical presentation. Although VZV is usually a self-limiting disease if complications develop, they can be life-threatening. Studies suggest that antivirals may reduce severity of disease and the number of days in which new lesions appear. Antiviral drugs have not been shown to decrease transmission of chickenpox, reduce the duration of absence from school. Antivirals are not recommended for routine treatment of otherwise healthy infants and children.

Antivirals may be considered for:
- Persons older than 13 years
- Persons with chronic cutaneous or pulmonary disorders
- Persons receiving long-term salicylate therapy
- Children receiving short, intermittent or aerosolized courses of corticosteroids

Immunocompromised children and adults with viral-mediated complications should receive intravenous administration of antivirals. For immunocompetent adults with uncomplicated VZV infection, oral antiviral therapy in addition to supportive care is recommended. Antiviral therapy can reduce the severity of symptoms and may decrease the risk of complications. Antiviral therapy is administered regardless of the individual’s vaccination status. Oral valacyclovir is the recommended choice for these individuals because it can be given in fewer doses compared with acyclovir, and has equivalent activity. For immunocompromised individuals who present with VZV, antiviral treatment, in addition to supportive care, is also recommended. For most patients, initial therapy with intravenous acyclovir is suggested rather than oral agents.

Treatment should be administered for 7-10 days. Treatment for newborn infants is different than that of adults. Acyclovir reduces the risk of mortality in severe VZV. Newborns with severe infection should be treated with acyclovir (30 mg/kg per day in three divided doses IV) for 10 days. To be effective, antiviral treatment must be started as soon as possible after the onset of symptoms because most viral replication has stopped by 72 hours after appearance of the rash. Fever control rarely is needed in VZV because temperature elevations usually are mild. The use of aspirin should be avoided because of the increased risk of developing Reye syndrome. Use of acetaminophen may prolong the illness. If temperature elevation persists, the infant should be evaluated for secondary complications. Breastfeeding is encouraged in newborns exposed to, or infected with, VZV because antibodies in breast milk may be protective.
Case Fatality
For normal childhood chickenpox, mortality is less than 2 per 100,000 cases/year. The risk in adults is 15 times higher. Neonates (usually 5-10 days) and patients with leukemia are susceptible to severe and possibly fatal chickenpox infection, with case fatality rates of 30% and 5-10%, respectively.

Reservoir
Humans are the only known hosts of VZV.

Transmission
Chickenpox is primarily spread by droplet or airborne spread of respiratory secretions or vesicle fluid, but may also spread by direct contact with an infected person or contact with an object that has touched a blister’s fluid (shirts, pants, etc.). Secondary attack rates among susceptible household contacts can be as high as 90%. Chickenpox is extremely contagious. Immunologically susceptible children/adults can acquire chickenpox from direct contact with herpes zoster lesion fluid.

An immunized person with breakthrough chickenpox disease is about two thirds less contagious than an unimmunized case of chickenpox.

High-risk groups for exposure and transmission of VZV include:
- Teachers of young children
- Childcare employees
- Residents and staff members of institutional settings, including correctional institutions
- College students
- Military personnel
- Adolescents and adults living in households with children
- Non-pregnant women of childbearing age
- International travelers

A person with shingles can spread the VZV to others that have never had chickenpox or received the chickenpox vaccine. With shingles, the virus is spread through contact with fluid from the rash blisters, and not through sneezing, coughing or causal contact. Shingles is less contagious than chickenpox, and the risk of a person with shingles spreading the virus is low if the rash is covered.

Susceptibility
Anyone can contract chickenpox, although it is most common in children under 15 years. Recovery from chickenpox usually results in lifetime immunity. Children who are immunocompromised (especially with leukemia) have a more serious illness that can take longer to resolve.
Incubation Period
The incubation period is usually 14-16 days, but can range from 10-21 days. The incubation period may be prolonged for as long as 28 days after receipt of Varicella-zoster immune globulin (VariZIG) or intravenous immune globulin (IVIG) and can be shortened in immunocompromised individuals. VZV can develop 2-16 days after birth in infants born to mothers with active VZV around the time of delivery.

Period of Communicability
A patient is contagious for roughly 7-10 days starting 1-2 days (sometimes 5) before rash onset, and continuing until scabs have formed on all lesions (usually 5 days after rash onset). Vaccinated persons with VZV may develop lesions that do not crust (macules and papules only). Isolation guidance for these persons is to exclude until no new lesions appear within a 24-hour period. Patients with altered immunity may be contagious for a longer period of time.

Epidemiology
Chickenpox can occur sporadically year-round, however, outbreaks occur most frequently in winter and early spring. In the U.S., incidence is highest between March and May, and lowest between September and November. Chickenpox affects males and females equally, and people of all races are susceptible.

Before the availability of a vaccine, chickenpox mainly affected children, with at least 90% of the population acquiring chickenpox by the age of 15 years. The highest age-specific incidence was among children 1-4 years of age, followed by children 5-9 years of age. Since the introduction of the vaccine, the incidence of chickenpox and disease-related hospitalizations has decreased by 70-80%. Cases have declined across all age groups, but mostly in children 1-4 and 5-9 years of age.

Figure 1. VZV case: Utah Cases (Confirmed and Probable)
Since 2009, 44% of all chickenpox cases reported to the Utah Department of Health are in children 5-9 years of age, followed by 30% in children 10-14 years of age. In recent years, the number of cases of chickenpox reported in Utah has declined

✓ PUBLIC HEALTH CONTROL MEASURES

Public Health Responsibility

- Prevent illness in high-risk individuals through disease investigation and administration of vaccine.
- Promote vaccination to reduce disease burden in the community.
- Provide education to the general public (regarding disease transmission) and to clinicians (regarding disease diagnosis, reporting, and prevention).
- Monitor disease trends.
- Monitor the impact of vaccination on incidence, morbidity, and mortality.

Prevention

The most effective control is widespread active immunization. Each year, more than 3.5 million cases of VZV, 9,000 hospitalizations, and 100 deaths are prevented by VZV vaccination in the United States.

Chemoprophylaxis

Persons without evidence of immunity to VZV, and who do not have a contraindication to vaccination, should be vaccinated. Studies among children have shown that vaccine administered within three days of exposure to rash is most effective in preventing disease (≥90%); however, vaccine administered within 5 days of exposure to rash is about 70% effective in preventing disease and 100% effective in modifying disease. In a VZV outbreak setting, ongoing exposures are likely and may continue for weeks and even months. Thus, to limit disease transmission during an outbreak and to provide protection against subsequent exposures, ACIP recommends that all persons without evidence of immunity to VZV be offered vaccine, even if more than 5 days have passed since first exposure to the disease.

School Settings

One dose of the VZV vaccine has been successful for outbreak prevention and control in school settings. No data is available regarding a potential benefit of administering a second dose to 1-dose vaccine recipients after exposure. However, administration of the second dose is recommended for exposed people to bring them up-to-date on vaccination. If a person is unvaccinated and receives one dose of the vaccine, that person may immediately return to school after vaccination. If a person has already had one dose of the vaccine previously, a second dose is recommended for outbreak control. Children who are vaccinated with a first or second dose during an outbreak may immediately return to school after vaccination. For outbreaks among preschool-aged children in particular, a second dose of VZV vaccine is recommended to provide optimal protection for children 1-4 years of age.
VariZIG was approved by the FDA in December of 2012 for use in the U.S. for post exposure prophylaxis of VZV for persons at high risk for severe disease who lack evidence of immunity to VZV, and for whom VZV vaccine is contraindicated. VariZIG can prevent chickenpox if given within 10 days of exposure. (Note: FDA approved the 10 day time frame from four days in May 2011. VariZIG should be administered as soon as possible following VZV virus exposure, ideally within 96 hours for greatest effectiveness).

It is recommended for use in high-risk persons including:

- Immunocompromised patients without evidence of immunity to VZV.
- Newborn infants whose mothers have signs and symptoms of VZV around the time of delivery (i.e., five days before, to two days after).
- Hospitalized premature infants born at ≥28 weeks of gestation whose mothers do not have evidence of immunity to VZV.
- Hospitalized premature infants born at < 28 weeks of gestation or who weigh ≤1,000 g at birth, regardless of their mother’s evidence of immunity to VZV.
- Pregnant women without evidence of immunity.

**VariZIG Administration**

VariZIG is supplied in 125-IU vials and should be administered intramuscularly as directed by the manufacturer. The recommended dose is 125 IU/10 kg of body weight, up to a maximum of 625 IU (five vials). The minimum dose is 62.5 IU (0.5 vial) for patients weighing ≤2.0 kg and 125 IU (one vial) for patients weighing 2.1–10.0 kg. If VariZIG is not available, IVIG can be considered (also within 10 days of exposure). Additionally, in the absence of both VariZIG and IVIG, some experts recommend prophylaxis with acyclovir (80 mg/kg/day, administered 4 times per day for 7 days; maximum dose, 800 mg, 4 times per day), beginning 7–10 days after exposure for people without evidence of immunity, and with contraindications for VZV vaccination.

VZV vaccine should be administered ≥5 months after VariZIG administration. Because VZV zoster immune globulin might prolong the incubation period by ≥1 week, any patient who receives VariZIG should be observed closely for signs and symptoms of VZV for 28 days after exposure. Antiviral therapy should be instituted immediately if signs or symptoms of VZV occur. Most common adverse reactions following VariZIG administration were pain at injection site (2%) and headache (2%). Contraindications for VariZIG administration include a history of anaphylactic or severe systemic reactions to human immune globulins, and IgA-deficient patients with antibodies against IgA and a history of hypersensitivity.

**Vaccine**

Vaccination is the primary method of prevention. A live, attenuated vaccine has been available since 1995. It is recommended that the following groups be vaccinated:
• All children <13 years of age should routinely be given two doses of VZV-containing vaccine. The first dose should be given at 12-15 months of age, and the second at 4-6 years of age. The second dose can be given at an earlier age provided it has been at least three months since the first dose. However, if the second dose is given at least 28 days after the first, the second dose does not need to be repeated. In 2014, the percentage of children in Utah aged 19-35 months that received the VZV vaccine was 86.3%.
• A second dose catch-up VZV vaccination is recommended for children and adolescents who previously had received one dose. Catch-up second dose can be administered at any interval longer than three months after the first dose.
• All other people ≥13 years of age without evidence of immunity should be vaccinated with two doses of VZV vaccine separated by 4-8 weeks.

For the 2014-2015 school year in Utah, students entering kindergarten or seventh grade are required to have two doses of the chickenpox vaccine. A child one year of age and older attending an early childhood program must have received one dose of the chickenpox vaccine prior to entry. For more information regarding the vaccination requirements in Utah for 2014/2015, see the “Utah Immunization Guidebook” available at http://www.immunize-utah.org/pdf/Immunization_Guidebook.pdf.

A single dose of the VZV vaccine is 80% effective against infection of any severity, and more than 95% effective against severe disease. Breakthrough infection (chickenpox in a vaccinated individual) can occur, but is usually milder with fewer lesions. Two doses are more effective than one dose, having a mean effectiveness of approximately 93% against VZV of any severity. Children with breakthrough chickenpox usually do not have a fever, tend to be out of school for a shorter amount of time than children with natural infections, and have lesions that generally do not blister and scab.

Ninety-seven percent of healthy children under age 13 years will develop detectable levels of antibody after one vaccination; however, only 80% of persons 13 years and older will respond to a single dose. Two doses administered 4-8 weeks apart for persons 13 years and older produces detectable levels of antibodies in 99% of vaccines.

A vaccine for shingles was licensed in 2006 and recommended for persons 60 years of age and older. The vaccine reduces the risk of shingles by 51% and the risk of post-herpetic neuralgia (a painful complication of shingles) by 67%. Vaccine efficacy lasts for at least three years, although the ultimate length of efficacy is unknown.

The Advisory Committee on Immunization Practices (ACIP) recommends vaccination for all children, adolescents, and adults who do not have evidence of immunity. Vaccination is especially important for:
• Healthcare personnel
• Household contacts of immunocompromised persons
• Persons living or working in environments where transmission is likely (teachers, childcare workers, residents and staff of institutional settings)
• Persons in environments where transmission has been reported
• Non-pregnant women of childbearing age
• Adolescents and adults in households with children
• International travelers

Because increasing use of VZV vaccine will lead to an expected decrease in circulating wild-
type VZV, unimmunized children may acquire infection at an older age, when they are more
susceptible to severe infection. Mathematical models predict that if immunization rates in
children are >90%, a greater proportion of VZV infections will occur at older ages; however,
overall disease burden will decline. This emphasizes the need for high levels of immunization
among children to prevent late-onset disease.

Contraindications to the VZV vaccination include:
• Immunocompromised persons
• History of anaphylactic reactions to any component of the vaccine
• Pregnancy
• Acute severe illness
• History of congenital immune disorders in first degree relatives, unless the immune
  competence of potential vaccine recipient has been demonstrated

Isolation and Quarantine Requirements

Isolation: Non-hospitalized patients should be voluntarily isolated in their house until all of their
lesions have scabbed over.

For children with suspected breakthrough VZV consisting of only maculopapular lesions and
occurring greater than 42 days after receiving a vaccine dose, isolation from school or childcare
is recommended until 24 hours have elapsed since the appearance of any new lesions. As
lesions in these children may not form crusts, parents do not have to wait for crusting to occur or
for the lesion to completely resolve before the child can resume physical and social contact with
others.

Hospital: Airborne and contact isolation is required for all chickenpox cases until all lesions are
crusted. Only immune individuals should care for the patient. VariZIG should be considered for
susceptible high-risk contacts of the sick person.

Quarantine: Only in the school setting should contacts of cases be routinely quarantined.
Recommendations follow in the School/childcare outbreak control recommendations section.

CASE INVESTIGATION

Reporting
Chickenpox should be reported within three working days of identification to public health in
Utah. Herpes zoster is not a reportable condition. Report any illness to public health authorities
that meets any of the following criteria:
1. Any person with a generalized, maculopapulovesicular rash characteristic of VZV.
2. Any person with a positive laboratory test confirming VZV infection. Acceptable tests are: isolation of VZV from a clinical specimen; positive direct fluorescent antibody (DFA) for VZV DNA from a clinical specimen; a positive polymerase chain reaction (PCR) for VZV DNA on a clinical specimen; or a significant rise (i.e., at least a 4-fold rise) in serum VZV immunoglobulin G (IgG) antibody level on paired sera by any standard serologic assay.
3. A person whose healthcare record (i.e., hospital discharge or outpatient record) contains a diagnosis of VZV.
4. A person whose death certificate lists VZV as the cause of death or a contributing cause of death.

Other recommended reporting procedures
- All cases of VZV should be reported.
- Reporting should be on-going and routine.
- Frequency of reporting should follow the state health department’s routine schedule.

Figure 2. Chickenpox reporting table

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Reporting</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Clinical Evidence</strong></td>
<td></td>
</tr>
<tr>
<td>Generalized maculopapulovesicular rash</td>
<td>S</td>
</tr>
<tr>
<td>Healthcare record contains a diagnosis of varicella</td>
<td>S</td>
</tr>
<tr>
<td>Death certificate lists varicella as the cause of death or a contributing cause of death</td>
<td>S</td>
</tr>
<tr>
<td><strong>Laboratory Evidence</strong></td>
<td></td>
</tr>
<tr>
<td>Isolation of varicella virus from a clinical specimen</td>
<td>S</td>
</tr>
<tr>
<td>Positive direct fluorescent antibody (DFA) for varicella from a clinical specimen</td>
<td>S</td>
</tr>
<tr>
<td>Positive polymerase chain reaction (PCR) for varicella-specific nucleic acid on a clinical specimen</td>
<td>S</td>
</tr>
<tr>
<td>A significant rise in serum varicella immunoglobulin G (IgG) antibody level by any standard serologic assay</td>
<td>S</td>
</tr>
</tbody>
</table>

Notes: S = This criterion alone is sufficient to identify a case for reporting.

Disease-specific data elements to be included in the initial report are:
- Clinical factors
  - Number of lesions (by category: 500) and actual number of lesions if <50
  - Fever
  - Complications
  - Hospitalization
- Epidemiologic factors
  - Predisposing conditions
  - Previous VZV history
  - Transmission setting
  - Association with an outbreak
• Immunization history
  o Doses of VZV-containing vaccine received
  o Date(s) of dose(s) of VZV-containing vaccine received

Case Definition
Varicella (Chickenpox) (CSTE, 2009)

Clinical Case Definition
An illness with acute onset of diffuse (generalized) maculopapulovesicular rash characteristic of VZV, without other apparent cause. In vaccinated persons who develop VZV more than 42 days after vaccination (i.e., breakthrough disease), the rash may by atypical in appearance (maculopapular with few or no vesicles).

Laboratory Criteria
• Isolation of VZV from a clinical specimen, OR
• Direct fluorescent antibody (DFA), OR
• Polymerase chain reaction (PCR), OR
• Significant rise (at least a 4 fold rise) in serum VZV IgG antibody level.

Case Classification
Probable:
• An acute illness with:
  o Diffuse (generalized) maculopapulovesicular rash; AND
  o Lack of laboratory confirmation; AND
  o Lack of epidemiologic linkage to another probable or confirmed case.

Confirmed:
• An acute illness with diffuse (generalized) maculopapulovesicular rash, AND
  o Epidemiologic linkage to another probable or confirmed case, OR
  o Laboratory confirmation by any of the following:
    ▪ Isolation of VZV virus from a clinical specimen, OR
    ▪ VZV antigen detected by direct fluorescent antibody test, OR
    ▪ VZV-specific nucleic acid detected by polymerase chain reaction (PCR), OR
    ▪ Significant rise in serum anti-VZV immunoglobulin G (IgG) antibody level by any standard serologic assay.

Note: Two probable cases that are epidemiologically linked would be considered confirmed, even in the absence of laboratory confirmation.

In vaccinated persons who develop VZV more than 42 days after vaccination (breakthrough disease), the disease is almost always mild with fewer than 50 skin lesions and shorter duration of illness. The rash may also be atypical in appearance (maculopapular with few or no vesicles).

Laboratory confirmation of cases of VZV is not routinely recommended; laboratory confirmation is recommended for fatal cases and in other special circumstances.
### Figure 3. Classification table

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Confirmed</th>
<th>Probable</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Clinical Evidence</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acute illness with diffuse (generalized) maculopapulovesicular rash</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td><strong>Laboratory Evidence</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Isolation of varicella virus from a clinical specimen</td>
<td></td>
<td>O</td>
</tr>
<tr>
<td>Detection of varicella antigen by direct fluorescent antibody test;</td>
<td></td>
<td>O</td>
</tr>
<tr>
<td>Detection of varicella-specific nucleic acid by polymerase chain reaction (PCR)</td>
<td></td>
<td>O</td>
</tr>
<tr>
<td>Significant rise in serum anti-varicella immunoglobulin G (IgG) antibody level by any standard serologic assay</td>
<td></td>
<td>O</td>
</tr>
<tr>
<td><strong>Epidemiologic Evidence</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Contact of a confirmed or probable case of varicella infection</td>
<td></td>
<td>O</td>
</tr>
<tr>
<td>Member of a risk group as defined by the public health authorities during an outbreak</td>
<td></td>
<td>O</td>
</tr>
</tbody>
</table>

**Notes:**
- S = This criterion alone is sufficient to classify a case.
- N = All ―“N” criteria in the same column are necessary to classify a case.
- O = At least one of these ―“O” (Optional) criteria in each category (i.e., clinical evidence and laboratory evidence) in the same column—in conjunction with all ―“N” criteria in the same column—is required to classify a case.

### Case Investigation Process

The majority of chickenpox case investigations will not require any case management or contact tracing. In the event that a case is identified in a person at high-risk for complications, or in a setting where persons at high-risk for complications may have been exposed, the local health department may choose to more thoroughly investigate.

### Outbreaks

Schools and childcare facilities are the most common sites for chickenpox outbreaks. A chickenpox outbreak in a school setting is defined as:

- Five or more cases within a 2-month period in the same school facility.

An outbreak is considered over when no new cases occur within two months from the date the last case was considered infectious.

**Note:** Because childcare facilities, after school programs, preschools, Head Start programs, charter schools, and private schools can differ significantly from traditional public school settings, the local health department may make adjustments to the outbreak definition for settings where it is deemed necessary.
A Confirmed Outbreak

Once an outbreak is confirmed, parents should be notified of their child’s possible exposure to chickenpox. They should be educated on the signs and symptoms of the disease, the availability of a second dose of vaccine, told to keep children home if they suspect they are ill, and to inform the school if their child has chickenpox. Teachers and other school employees should also be educated on the signs and symptoms of chickenpox. Children with signs and symptoms should be sent home and encouraged to consult a medical professional. Pregnant teachers and employees should be identified, and their immune status determined. Those without an appropriate immune history should be encouraged to consult with a medical professional. Laboratory diagnosis is not necessary (or even required) to make a chickenpox diagnosis.

Exclusions of vaccine-exempt students should occur once at least five cases have occurred within a 2-month period among students in the same kindergarten class or grade level. Exclusions should only be made to students within the same kindergarten class and grade level. Vaccine-exempt students should remain out of school for days 10-21 after the last known case is considered non-infectious. Days 10-21 are the critical days for vaccine-exempt students to be out of school, however, local health departments may choose to exclude students for the full 21 days after exposure to simplify the exclusion policy. Vaccination within three days, and possibly five days, of exposure has been shown to prevent or reduce the severity of disease. However, since an adequate immune response can take up to two weeks to develop, disease may still result, even if vaccination occurs soon after exposure. Parents should still be cautioned to watch for signs of illness even if their child receives the vaccine within five days of exposure, and should keep the child home if they suspect their child is ill. If parents of vaccine-exempt students choose to vaccinate their children, the exclusionary period does not apply, and students may return to school as soon as they are vaccinated.

Since the 2006-2007 school year in Utah, all students in kindergarten through 4th grade as well as 7th graders are required to be vaccinated for chickenpox. Exclusions do not apply to students without a vaccination requirement, although any sick child, regardless of vaccination requirement, should be excluded until all lesions have scabbed.

Current CDC recommendations for dealing with chickenpox outbreaks can be found here: http://www.cdc.gov/chickenpox/outbreaks/manual.html.

Under Utah Administrative Code, students who have vaccine exemptions can be excluded from school as stated below:


1) A local or state health department representative may exclude a student who has claimed an exemption or who is conditionally enrolled from school attendance if there is good cause to believe that the student has a vaccine preventable disease and:
   a) has been exposed to a vaccine-preventable disease; or
b) will be exposed to a vaccine-preventable disease as a result of school attendance.

2) An excluded student may not attend school until the local health officer is satisfied that a student is no longer at risk of contracting or transmitting a vaccine-preventable disease.

**A Prolonged Outbreak**

Because persons can be infectious before symptoms appear, exclusions can have a limited value on preventing transmission. Outbreaks in the school setting have been documented to persist for up to six months. Chickenpox vaccination has been effective in controlling chickenpox outbreaks. ACIP now recommends that a second dose of vaccine be offered to exposed persons with a history of one dose of chickenpox vaccine, providing that the appropriate time interval has passed since the first dose (three months for people 12 months to 12 years of age, and at least four weeks for people ≥13 years of age).

*Optional*: A second dose of vaccine should be recommended to all students if cases are still being diagnosed 60 days after the initial rash onset of the first identified case, or once 20 cases have been identified. Efforts should be focused on students with active cases in their kindergarten class or grade level, or siblings of active cases.

The local health department may evaluate individual cases and outbreaks and make any necessary adjustments to exemptions.

**Identifying Case Contacts**

Close contacts are people who have the following contact with the case patient during the infectious period (7-10 days starting 1-2 days [sometimes 5] before rash onset until scabs have formed on all lesions [usually 5 days after rash onset]).

- Household and immediate family members (those who spend many hours together or sleep under the same roof)
- Those who have direct contact with respiratory secretions
- Those who have direct or indirect contact with blister fluid
- Healthcare workers with face-to-face contact with a patient
- Those who share confined space during the communicable period. Such contacts may include:
  - Core groups of close friends, social contacts, boyfriends, girlfriends
  - Students within the same kindergarten class or grade level
  - Contacts at church activities and employment
  - Participants in extracurricular activities (such as fieldtrips)
  - Children attending after-school care or a playgroup

**Healthcare Personnel**

Nosocomial outbreaks continue to be reported with source patients including those who did not have evidence of clinical disease at the time of transmission. Even if patients with VZV can be promptly placed in negative pressure rooms, as recommended by the CDC, susceptible healthcare providers remain at risk of acquiring disease that is transmitted before clinical disease is apparent. An estimated 14-40% of healthcare providers are susceptible to VZV.
While 97-99% of those who have a positive history of VZV are seropositive, variable percentages of those with an unclear or no history of VZV are seropositive. All healthcare workers (HCWs) should be immune to chickenpox, either through natural infection or vaccination. HCWs without a reliable history should have laboratory documentation of seroconversion. If a susceptible HCW is exposed to VZV, they should be furloughed from work for days 10-21 after exposure. Because receipt of VariZIG can prolong the incubation period by one week, susceptible employees receiving VariZIG should be furloughed for days 10-28 after exposure. Because of the possibility of breakthrough disease, several options for monitoring vaccinated HCWs that are potentially exposed to VZV exist. Serological testing for immunity immediately after exposure is recommended. Those with detectable antibodies are unlikely to become infected. Persons without detectable antibodies may be retested in 5-6 days. If detectable antibodies are present, these persons are unlikely to become ill. If a HCW is tested twice, and both times antibodies are not detected, the HCW may be furloughed for days 10-28 after exposure, or monitored daily for symptoms of chickenpox.

**Case Contact Management**

**School/childcare outbreak control recommendations**

*One case identified*

Persons with symptoms should be excluded from school/childcare until all lesions have scabbed over, even if no laboratory confirmation is performed, or an outbreak is not recognized.

*Optional:* Parents of vaccine-exempt students should be notified of their child’s possible exposure once **one** case of chickenpox occurs in the school.

*Optional:* Parents of students in the same kindergarten class or grade level of a chickenpox case should be educated on the signs and symptoms of the disease, the availability of a second dose of vaccine, told to keep children home if they suspect they are ill, and to inform the school if their child has chickenpox.

**Treatment**

Determine if case is at high-risk for complications; this includes:

- Persons older than 13 years
- Persons with a chronic cutaneous or pulmonary disorders
- Persons receiving long-term salicylate therapy
- Children receiving short, intermittent or aerosolized courses of corticosteroids
- Immunocompromised children and adults with viral-mediated complications should receive intravenous administration of VariZIG or IGIV

Consider antiviral treatment if case is at high-risk for complications.

**Isolation**

Children with chickenpox should be excluded from school or child-care settings until all lesions have crusted over. Voluntary isolation from work and other settings where close contact may transmit the disease is desirable. Educate patient to refrain from contact until all lesions have
crusted over. Follow the isolation and quarantine requirements (listed above) for healthcare workers or residents in a hospital or long term care facility.

Education
Provide an educational fact sheet to the patient (or parent of the patient). If the patient works at or attends a childcare or school, work with the school administration to send notification letters to other students/parents/teachers as necessary.

Asymptomatic contacts
- Exclude vaccine-exempt children without history of disease from school/childcare or contact with children when necessary.
- Recommend vaccine to susceptible persons.
- Recommend medical consultation for exposed susceptible persons at high-risk of severe disease (VariZIG may be warranted).
- Provide educational materials informing of exposure and recommending vaccination.

Symptomatic contacts
- Exclude from school/childcare settings until infectious period has passed.
- Consider vaccination of susceptible persons to lessen disease severity.
- Consider treatment of active cases with antiviral therapy (adolescents, adults, and high-risk only).


✓ REFERENCES


Centers for Disease Control and Prevention. *Immunization of Health-Care Workers: Recommendations of the Advisory Committee on Immunization Practices (ACIP) and the Hospital Infection Control Practices Advisory Committee (HICPAC)*. MMWR 1997; 46(RR-18): 1-42.

http://www.cdc.gov/mmwr/preview/mmwrhtml/mm6228a4.htm.


✔ VERSION CONTROL

Update. Aug 2015: General update to document format and revisions of thematic order.

Update. Feb 2016: Added the importance of public health section and additional information to the clinical description. Update to testing (serology, culture, PCR, DFA). Added Tzanck stain form of testing. Additional information to incubation period and period of communicability. Update to Utah epidemiologic statistics. Added antiviral treatment information for adults and newborns. Update to chemoprophylaxis and vaccine information. Update to transmission section. Added narrative of reporting requirements. Updated UT-NEDSS Minimum/Required Fields.
**✓ UT-NEDSS Minimum/Required Fields by Tab**

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