# Prevention and Control of Influenza with Vaccines: Recommendations of the Advisory Committee on Immunization Practices (ACIP) — United States, 2012–13 Influenza Season

In 2010, the Advisory Committee on Immunization Practices (ACIP) first recommended annual influenza vaccination for all persons aged  $\geq 6$  months in the United States (1). Annual influenza vaccination of all persons aged ≥6 months continues to be recommended. This document 1) describes influenza vaccine virus strains included in the U.S. seasonal influenza vaccine for 2012-13; 2) provides guidance for the use of influenza vaccines during the 2012-13 season, including an updated vaccination schedule for children aged 6 months through 8 years and a description of available vaccine products and indications; 3) discusses febrile seizures associated with administration of influenza and 13-valent pneumococcal conjugate (PCV-13) vaccines; 4) provides vaccination recommendations for persons with a history of egg allergy; and 5) discusses the development of quadrivalent influenza vaccines for use in future influenza seasons. Information regarding issues related to influenza vaccination that are not addressed in this update is available in CDC's Prevention and Control of Influenza with Vaccines: Recommendations of the Advisory Committee on Immunization Practices (ACIP), 2010 and associated updates (1,2).

Methodology for the formulation of the ACIP annual vaccine recommendations has been described previously (1). The ACIP

Recommendations for routine use of vaccines in children and adolescents are issued by CDC and are harmonized to the greatest extent possible with recommendations made by the American Academy of Pediatrics, the American Academy of Family Physicians (AAFP), and the American College of Obstetrics and Gynecology (ACOG). CDC recommendations for routine use of vaccines in adults are harmonized to the greatest extent possible with recommendations made by AAFP, ACOG, and the American College of Physicians. The Advisory Committee on Immunization Practices (ACIP) is chartered as a federal advisory committee to provide expert external advice and guidance to the Director of CDC on use of vaccines in the civilian population of the United States. ACIP members are named by the Secretary of the U.S. Department of Health and Human Services. ACIP recommendations become CDC policy once approved by the Director of CDC, on the date published by MMWR.

Influenza Work Group meets every 2–4 weeks throughout the year. Work Group membership includes several voting members of ACIP and representatives of ACIP Liaison Organizations. Meetings are held by teleconference and include discussion of influenza-related issues, such as influenza surveillance, vaccine effectiveness and safety, coverage in groups recommended for vaccination, program feasibility, cost-effectiveness, and anticipated vaccine supply. Presentations are requested from invited experts, and published and unpublished data are discussed. CDC's Influenza Division provides data on influenza surveillance, antiviral resistance, and vaccine effectiveness. CDC's Immunization Safety Office provides information on vaccine safety, and CDC's Immunization Services Division provides information on vaccine distribution and coverage.

#### Vaccine Strains for the 2012-13 Influenza Season

U.S. influenza vaccines for 2012–13 will contain A/California/7/2009 (H1N1)-like, A/Victoria/361/2011 (H3N2)-like, and B/Wisconsin/1/2010-like (Yamagata lineage) antigens. The influenza A(H3N2) and B antigens differ from the respective 2010–11 and 2011–12 seasonal vaccine antigens (3). The influenza A(H1N1) vaccine virus strain is derived from an influenza A(H1N1)pdm09 (2009[H1N1]) virus and was included in the 2009(H1N1) monovalent pandemic vaccine as well as the 2010–11 and 2011–12 seasonal vaccines.

#### Recommendations for Vaccination

Routine annual influenza vaccination is recommended for all persons aged ≥6 months. To permit time for production of protective antibody levels (4,5), vaccination optimally should occur before onset of influenza activity in the community. Therefore, vaccination providers should offer vaccination as soon as vaccine is available. Vaccination should be offered throughout the influenza season (i.e., as long as influenza viruses are circulating in the community).

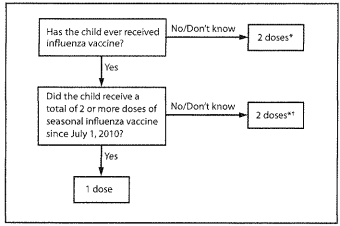
#### Vaccine Dose Considerations for Children Aged 6 Months Through 8 Years

Children aged 6 months through 8 years require 2 doses of influenza vaccine (administered a minimum of 4 weeks apart) during their first season of vaccination to optimize immune response. In a study of children aged 5 through 8 years receiving trivalent inactivated influenza vaccine (TIV) for the first time,

the proportion of children with protective antibody responses was significantly higher after 2 doses compared with a single dose (6). Several studies have indicated that the time interval between two initial doses (from 4 weeks up to 1 year) of the same antigen might not be critical (7-9). However, because of the antigenic novelty of the 2009(H1N1) pandemic virus, which is anticipated to continue circulating during 2012–13, exposure history to this antigen also must be considered. Children who last received seasonal (trivalent) influenza vaccine before the 2010–11 season but did not receive a vaccine containing 2009(H1N1) antigen (either seasonal vaccine since July 2010 or monovalent 2009[H1N1] vaccine) will not have received this antigen. These children are recommended to receive 2 doses this season, even if 2 doses of seasonal influenza vaccine were received before the 2010-11 season. This is illustrated in two approaches for determining the number of doses required for children aged 6 months through 8 years, both of which are acceptable (Figure 1).

- 1. The first approach takes into consideration only doses of seasonal influenza vaccine received since July 1, 2010. This recommendation is harmonized with that of the American Academy of Pediatrics (10). This approach has the advantage of simplicity, particularly in settings in which ascertaining vaccination history before the 2010–11 season is difficult. Using this approach, children aged 6 months through 8 years need only 1 dose of vaccine in 2012–13 if they received a total of 2 or more doses of seasonal vaccine since July 1, 2010. Children who did not receive a total of 2 or more doses of seasonal vaccine since July 1, 2010, require 2 doses in 2012–13.
- 2. In settings where adequate vaccination history from before the 2010–11 season is available, the second approach may be used. By this approach, if a child aged 6 months through 8 years is known to have received at least 2 seasonal influenza vaccines during any previous season, and at least 1 dose of a 2009(H1N1)-containing vaccine (i.e., either 2010–11 or 2011–12 seasonal vaccine or the monovalent 2009[H1N1] vaccine), then the child needs only 1 dose for 2012–13. Using this approach, children aged 6 months through 8 years need only 1 dose of vaccine in 2012–13 if they have received any of the following:
  - 2 or more doses of seasonal influenza vaccine since July 1, 2010; or
  - 2 or more doses of seasonal influenza vaccine before July 1, 2010, and 1 or more doses of monovalent 2009(H1N1) vaccine; or
  - 1 or more doses of seasonal influenza vaccine before July 1, 2010, and 1 or more doses of seasonal influenza vaccine since July 1, 2010.

FIGURE 1. Influenza vaccine dosing algorithm for aged children 6 months through 8 years — Advisory Committee on Immunization Practices, United States, 2012–13 influenza season



\* Doses should be administered at least 4 weeks apart.

<sup>†</sup> For simplicity, this algorithm takes into consideration only doses of seasonal influenza vaccine received since July 1, 2010. As an alternative approach in settings where vaccination history from before July 1, 2010, is available, if a child aged 6 months through 8 years is known to have received at least 2 seasonal influenza vaccines during any previous season, and at least 1 dose of a 2009(H1N1)-containing vaccine (i.e., either 2010–11 or 2011–12 seasonal vaccine or the monovalent 2009[H1N1] vaccine), then the child needs only 1 dose for 2012–13. Using this approach, children aged 6 months through 8 years need only 1 dose of vaccine in 2012–13 if they have received any of the following: 1) 2 or more doses of seasonal influenza vaccine since July 1, 2010; 2) 2 or more doses of seasonal influenza vaccine; or 3) 1 or more doses of seasonal influenza vaccine; or 3) 1 or more doses of seasonal influenza vaccine before July 1, 2010, and 1 or more doses of seasonal influenza vaccine since July 1, 2010. Children for whom one of these conditions is not met require 2 doses in 2012–2013.

Children for whom one of these conditions is not met require 2 doses in 2012–13.

#### **Available Vaccine Products and Indications**

Multiple influenza vaccines (with the same antigenic composition) are expected to be available during the 2012–13 season (Table). Current package inserts should be consulted for updated information and description of additional components of various vaccine formulations, indications, contraindications, and precautions.

TIV preparations, with the exception of Fluzone Intradermal (Sanofi Pasteur), should be administered intramuscularly. For adults and older children, the deltoid is the preferred site. Infants and younger children should be vaccinated in the anterolateral thigh. Specific guidance regarding site and needle length for intramuscular administration can be found in ACIP's General Recommendations on Immunization (11). For intramuscular TIV preparations, children aged 6 through 35 months receive 0.25 mL per dose; persons aged ≥36 months receive 0.5 mL per dose (Table). Fluzone Intradermal is administered intradermally

TABLE. Influenza vaccine information, by age group — United States, 2012-13 influenza season\*

Vaccine	Trade name	Manufacturer	Presentation	Mercury content (μg Hg per 0.5 mL dose)	t Ovalbumin content (μg per 0.5mL dose)†	Age group	No. of doses	Route
TIV	Fluzone	Sanofi Pasteur	0.25 mL prefilled syringe 0.5 mL prefilled syringe 0.5 mL vial 5.0 mL multidose vial	0.0 0.0 0.0 25.0	\$ \$ \$	6–35 mos ≥36 mos ≥36 mos ≥6 mos	1 or 25 1 or 25 1 or 25 1 or 25	IM** IM** IM**
TIV	Agriflu	Novartis Vaccines	0.5 mL prefilled syringe	0	<0.4	≥18 yrs	1	IM**
TIV	Fluvirin	Novartis Vaccines	0.5 mL prefilled syringe 5.0 mL multidose vial	≤1 25.0	≤1 ≤1	≥4 yrs	1 or 25	IM**
TIV	Fluarix	GlaxoSmithKline	0.5 mL prefilled syringe	0	≤0.05	≥3 yrs	1 or 25	IM**
TIV	FluLaval	ID Biomedical Corporation of Quebec (distributed by GlaxoSmithKline)	5.0 mL multidose vial	<25.0	≤0.3	≥18 yrs	1	IM**
TIV	Afluria	CSL Biotherapies	0.5 mL prefilled syringe	0.0	≤1	≥9 yrs††	1	IM**
		(distributed by Merck)	5.0 mL multidose vial	24.5	≤1			
TIV high-dose <sup>§§</sup>	Fluzone High-Dose	Sanofi Pasteur	0.5 mL prefilled syringe	0.0	\$	≥65 yrs	1	IM**
TIV intradermal <sup>15</sup>	Fluzone Intradermal	Sanofi Pasteur	0.1 mL prefilled microinjection system	0.0 (per 0.1 mL)	······\$	1864 yrs	1	ID
LAIV	FluMist***	Medimmune	0.2 mL prefilled intranasal sprayer	0.0 (per 0.2 mL)	<0.24 (per 0.2mL) <sup>†††</sup>	2–49 yrs <sup>§§§</sup>	1 or 2¶	IN

Abbreviations: TIV = trivalent inactivated vaccine; LAIV = live-attenuated influenza vaccine; IM = intramuscular; ID = intradermal; IN = intranasal.

via a single-dose, prefilled microinjection syringe. The preferred site for administration is over the deltoid muscle.

Age indications for the various TIV products differ. All TIV preparations contain the same quantity of hemagglutinin (15  $\mu$ g per vaccine virus strain per 0.5 mL dose; 45  $\mu$ g total), except Fluzone Intradermal and Fluzone High-Dose (Sanofi Pasteur). Fluzone Intradermal is indicated for persons aged 18 through 64 years and contains 9  $\mu$ g of hemagglutinin per vaccine virus strain (27  $\mu$ g total) in a 0.1 mL dose. Fluzone

High-Dose is indicated for persons aged  $\geq 65$  years and contains 60  $\mu$ g of hemagglutinin per vaccine virus strain (180  $\mu$ g total) in a 0.5 mL dose. Within specified age indications, ACIP expresses no preference for any given TIV formulation over another.

The intranasally administered live-attenuated influenza vaccine (LAIV), FluMist (MedImmune), is indicated for healthy, nonpregnant persons aged 2 through 49 years. No preference is indicated for LAIV versus TIV in this age group

<sup>\*</sup>Vaccination providers should consult Food and Drug Administration-approved prescribing information for 2012–13 influenza vaccines for the most updated information, including indications, contraindications, and precautions.

<sup>†</sup> Data on maximum ovalbumin content is supplied in package inserts of certain vaccines. Persons with a history of mild allergy to egg (specifically, those who experience only hives) should receive TIV with additional precautions (Figure 2).

<sup>§</sup> Information is not included in package insert but is available upon request from the manufacturer, Sanofi Pasteur, by contacting 1-800-822-2463 or mis.emails@sanofipasteur.com.

Figure 1 describes two approaches for determining the number of doses needed for children aged 6 months through 8 years.

<sup>\*\*</sup> For adults and older children, the recommended site of vaccination is the deltoid muscle. The preferred site for infants and young children is the anterolateral aspect of the thigh.

<sup>††</sup> Age indication per package insert is ≥5 years; however, the Advisory Committee on Immunization Practices recommends that Affuria not be used in children aged 6 months through 8 years because of increased risk for febrile reactions noted in this age group with CSL's 2010 Southern Hemisphere TiV. If no other age-appropriate, licensed inactivated seasonal influenza vaccine is available for a child aged 5 through 8 years who has a medical condition that increases the child's risk for influenza complications, Afluria can be used; however, vaccination providers should discuss with the parents or caregivers the benefits and risks of influenza vaccination with Afluria before administering this vaccine. Afluria may be used in persons aged ≥9 years.

<sup>\$\$</sup> A 0.5-mL dose contains 60  $\mu g$  of each vaccine antigen (180  $\mu g$  total).

<sup>55</sup> A 0.1-mL dose contains 9  $\mu$ g of each vaccine antigen (27  $\mu$ g total).

<sup>\*\*\*</sup> A new quadrivalent formulation of FluMist was approved by the Food and Drug Administration in February 2012. It is anticipated that this formulation will replace the currently available seasonal trivalent LAIV formulation for the 2013–14 season. FluMist is shipped refrigerated and stored in the refrigerator at 35°F–46°F (2°C–8°C) after arrival in the vaccination clinic. The dose is 0.2 mL divided equally between each nostril. Health-care providers should consult the medical record, when available, to identify children aged 2 through 4 years with asthma or recurrent wheezing that might indicate asthma. In addition, to identify children who might be at greater risk for asthma and possibly at increased risk for wheezing after receiving LAIV, parents or caregivers of children aged 2 through 4 years should be asked, "In the past 12 months, has a health-care provider ever told you that your child had wheezing or asthma?" Children whose parents or caregivers answer "yes" to this question and children who have asthma or who had a wheezing episode noted in the medical record within the past 12 months should not receive FluMist.

<sup>†††</sup> Insufficient data available for use of LAIV in egg-allergic persons.

<sup>\$55</sup> Flumist is indicated for healthy, nonpregnant persons aged 2 through 49 years. Persons who care for severely immunosuppressed persons who require a protective environment should not receive FluMist given the theoretical risk for transmission of the live-attenuated vaccine virus.

(1). Persons with a history of egg allergy should receive TIV rather than LAIV. Persons who care for severely immunosuppressed persons who require a protective environment should not receive LAIV given the theoretical risk for transmission of the live-attenuated vaccine virus.

#### Febrile Seizures Associated with TIV and PCV13

Febrile seizures are common in young children. At least one febrile seizure is experienced by 2%-5% of children, and nearly all children who have a febrile seizure recover quickly and are healthy afterwards (12). Before the 2010-11 influenza season, an increased risk for febrile seizures after TIV administration had not been observed in the United States (13, 14). During the 2010-11 influenza season, CDC and the Food and Drug Administration (FDA) conducted enhanced monitoring for febrile seizures after influenza vaccination because of reports of an increased risk for fever and febrile seizures in young children in Australia associated with a 2010 Southern Hemisphere vaccine produced by CSL Biotherapies (up to nine febrile seizures per 1,000 doses) (15). Because of the findings in Australia, ACIP does not recommend the U.S.-licensed CSL Biotherapies' TIV, Afluria, for children aged <9 years (2,16) (Table).

Surveillance for U.S.-licensed influenza vaccines during the 2010–11 season subsequently detected safety signals for febrile seizures in young children after TIV administration (17,18). Further assessment determined that the increased risk was in children aged 6 months through 4 years on the day of vaccination to the day after (the 0–1 day risk window). The risk was higher when children received concomitant PCV13 (i.e., when the two vaccines are administered at the same health-care visit) and peaked at approximately age 16 months (18). No increased risk was observed in children aged ≥5 years after TIV or in children of any age after LAIV. The magnitude of the increased risk for febrile seizures in young children in the United States (<1 per 1,000 children vaccinated) was substantially lower than the risk observed in Australia in 2010 (15).

After evaluating the data on febrile seizures from the 2010–11 influenza season and taking into consideration benefits and risks of vaccination, no policy change was recommended for use of TIV or PCV13 for the 2011–12 season (16,19,20). Surveillance data on febrile seizures in young children after administration of influenza vaccine for the 2011–12 influenza season (same vaccine formulation as 2010–11) were consistent with those from the 2010–11 influenza season (CDC, unpublished data, 2012). No changes in the use of TIV or PCV13 are recommended for the 2012–13 influenza season. As stated previously, ACIP does not

recommend the U.S.-licensed CSL Biotherapies' TIV, Afluria, for children aged <9 years (2,16) (Table).

## Influenza Vaccination of Persons with a History of Egg Allergy

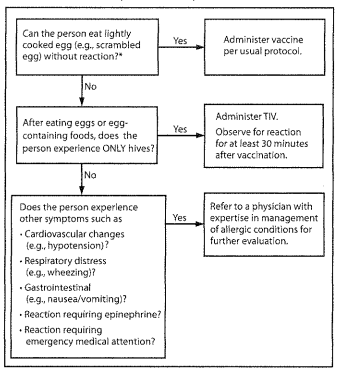
Severe allergic and anaphylactic reactions can occur in response to a number of influenza vaccine components, but such reactions are rare. All currently available influenza vaccines are prepared by means of inoculation of virus into chicken eggs. The use of influenza vaccines for persons with a history of egg allergy has been reviewed recently by ACIP (16). For the 2011–12 influenza season, ACIP recommended that persons with egg allergy who report only hives after egg exposure should receive TIV, with several additional safety measures, as described in this document. Recent examination of VAERS data indicated no disproportionate reporting of allergy or anaphylaxis after influenza vaccination during the 2011–12 season (21). For the 2012–13 influenza season, ACIP recommends the following:

- Persons with a history of egg allergy who have experienced only hives after exposure to egg should receive influenza vaccine, with the following additional safety measures (Figure 2):
  - a) Because studies published to date involved use of TIV, TIV rather than LAIV should be used (22);
  - b) Vaccine should be administered by a health-care provider who is familiar with the potential manifestations of egg allergy; and
  - c) Vaccine recipients should be observed for at least 30 minutes for signs of a reaction after administration of each vaccine dose (22).

Other measures, such as dividing and administering the vaccine by a two-step approach and skin testing with vaccine, are not necessary (22).

2. Persons who report having had reactions to egg involving such symptoms as angioedema, respiratory distress, lightheadedness, or recurrent emesis; or who required epinephrine or another emergency medical intervention, particularly those that occurred immediately or within a short time (minutes to hours) after egg exposure, are more likely to have a serious systemic or anaphylactic reaction upon reexposure to egg proteins. Before receipt of vaccine, such persons should be referred to a physician with expertise in the management of allergic conditions for further risk assessment (Figure 2).

FIGURE 2. Recommendations regarding influenza vaccination for persons who report allergy to eggs — Advisory Committee on Immunization Practices, United States, 2012–13 influenza season



Abbreviation: TIV = trivalent inactivated vaccine.

- \* Persons with egg allergy might tolerate egg in baked products (e.g., bread or cake). Tolerance to egg-containing foods does not exclude the possibility of egg allergy.
- 3. All vaccines should be administered in settings in which personnel and equipment for rapid recognition and treatment of anaphylaxis are available. ACIP recommends that all vaccination providers should be familiar with the office emergency plan (11).
- 4. Some persons who report allergy to egg might not be eggallergic. Those who are able to eat lightly cooked egg (e.g., scrambled egg) without reaction are unlikely to be allergic. Egg-allergic persons might tolerate egg in baked products (e.g., bread or cake). Tolerance to egg-containing foods does not exclude the possibility of egg allergy (23). Egg allergy can be confirmed by a consistent medical history of adverse reactions to eggs and egg-containing foods, plus skin and/ or blood testing for immunoglobulin E antibodies to egg proteins.
- 5. A previous severe allergic reaction to influenza vaccine, regardless of the component suspected to be responsible for the reaction, is a contraindication to future receipt of the vaccine.

#### **Quadrivalent Influenza Vaccines**

All currently available influenza vaccines are trivalent and contain A(H1N1), A(H3N2), and B viral antigens. There are two antigenically distinct lineages of influenza B viruses referred to as Victoria and Yamagata lineages (24). Immunization against B virus strains of one lineage provides limited cross-protection against strains in the other lineage (25). Because of this and the difficulty of predicting which B virus lineage will predominate during a given season, inclusion of a second influenza B vaccine virus strain in seasonal influenza vaccines has been proposed. A recent analysis indicates that the impact of such a quadrivalent vaccine could result in a modest reduction in influenza-associated outcomes, depending upon adequate vaccine supply, coverage, effectiveness, and incidence of influenza associated with the two B lineages (26).

In February 2012, FDA approved a new seasonal quadrivalent LAIV, FluMist Quadrivalent (MedImmune). This vaccine currently is not anticipated to be available until the 2013–14 influenza season, at which time it is expected to replace the currently available seasonal trivalent FluMist formulation (Table). Inactivated quadrivalent influenza vaccines currently are in development. These vaccines will be addressed in the ACIP influenza statement as they are approved and become available commercially.

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# Evaluation of Rapid Influenza Diagnostic Tests for Influenza A (H3N2)v Virus and Updated Case Count — United States, 2012

On August 10, 2012, this report was posted as an MMWR Early Release on the MMWR website (http://www.cdc.gov/mmwr). Previous reports have described cases of influenza A (H3N2) variant (H3N2v) virus\* infection with the influenza A (H1N1)pdm09 M gene detected in the United States during July 2011–July 2012 (1–3). This report provides 1) an update on the number of reported cases of H3N2v infections from July 12 to August 9, 2012, in the United States, 2) an updated results interpretation for the CDC Flu Real-Time Reverse Transcription Polymerase Chain Reaction (rRT-PCR) Dx Panel for A(H3N2)v for public health laboratories, and 3) an evaluation of rapid influenza diagnostic tests for the detection of H3N2v viruses.

From July 12 to August 9, a total of 153 cases of H3N2v infections were reported in Indiana (120 cases), Ohio (31), Hawaii (one), and Illinois (one). Of the 138 reported cases for which demographic information was available, 128 (93%) occurred in persons aged <18 years, and 10 (7%) occurred in adults. The median age of patients was 7 years. Two persons were hospitalized as a result of their illness; no deaths occurred. The patient in Hawaii was exposed to swine on the job, and no additional cases were found in Hawaii. The 152 patients reported from Illinois, Indiana, and Ohio resided in 27 counties; all reported direct or indirect exposure to swine, the majority at agricultural fairs.

H3N2v viruses can be detected by qualified U.S. public health laboratories using the CDC Flu rRT-PCR Dx Panel. Initially, if specimens tested positive for influenza A, H3, and pandemic influenza A markers and negative for H1 and pandemic H1 markers, they were reported as inconclusive until confirmed as influenza A (H3N2)v at the CDC laboratory (1). On August 7, CDC updated the results interpretation of the CDC Flu rRT-PCR Dx Panel for H3N2v for public health laboratories. Specimens with these findings may now be reported as "presumptive positive for influenza A (H3N2)v virus" and, for the ongoing investigations, cases with presumptive-positive test results at the state or local public health laboratory will now be classified as confirmed, as are those cases confirmed at CDC.

The CDC Flu rRT-PCR Dx Panel is available in public health laboratories but is not a point-of-care test available to clinicians. Rapid influenza diagnostic tests (RIDTs) frequently are used for the diagnosis of influenza infection in clinical settings, and the recent outbreaks of H3N2v virus (2,3) have highlighted the need to evaluate commercially available, widely used RIDTs for their ability to detect H3N2v viruses. As an initial assessment, CDC conducted an evaluation of seven FDA-cleared RIDTs with seven H3N2v viruses (Table 1). Five 10-fold dilutions in physiological saline of each virus grown in Madin-Darby Canine Kidney (MDCK) cells were tested with all of the RIDTs in duplicate. Tests with BinaxNOW, Directigen, FluAlert, QuickVue, and Sofia were performed according to the procedures in the kit inserts for nasal washes or aspirates. Xpect tests were performed according to their procedure for nasal washes and swab specimens transported in liquid media. For the Veritor test, 100 μL of diluted specimen was added directly to the reagent tube. Positive and negative controls contained in each RIDT were run before testing the viruses in the study to verify performance of each assay lot, with the exception of FluAlert, which does not provide controls.

Only four of seven RIDTs in this study (Directigen, Sofia, Veritor, and Xpect) detected all influenza A (H3N2)v viruses (Table 2). BinaxNOW detected five of seven, and QuickVue detected three of seven. FluAlert detected only one of seven.

#### Reported by

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#### **Editorial Note**

The H3N2v viruses identified since July 12, 2012, are similar to the 13 H3N2v viruses identified during July 2011–April 2012 (1); all sequenced viruses had the M gene from the influenza A (H1N1)pdm09 virus. As of August 9, all H3N2v patients for whom contact information was available reported contact with swine or attended an agricultural fair where swine were

<sup>\*</sup>Influenza viruses that circulate in swine are called swine influenza viruses when isolated from swine, but are called variant viruses when isolated from humans. A variant virus (human isolate) might or might not have the M gene from the influenza A (H1N1)pdm09 virus, along with other genetic changes. Seasonal influenza A (H3N2) viruses that circulate worldwide in the human population have significant antigenic and genetic differences from influenza A (H3N2) viruses circulating in swine. Additional information is available at http://www.who.int/influenza/gisrs\_laboratory/terminology\_ah3n2v/en/index.html.

TABLE 1. Evaluation of seven FDA-cleared RIDTs for the ability to detect H3N2v viral antigens — CDC, United States, 2012

RIDT (manufacturer)	Abbreviated name	Approved specimens*	Analyzer for interpretation
BinaxNOW Influenza A&B (Alere)	BinaxNOW	NP swab Nasal wash/aspirate/swab	No
Directigen EZ Flu A+B (Becton-Dickinson)	Directigen	NP wash/aspirate/swab Throat swab	No
SAS FluAlert A&B (SA Scientific)	FluAlert	Nasal wash/aspirate	No
QuickVue Influenza A+B Test (Quidel)	QuickVue	NP swab Nasal wash/aspirate/swab	No
Sofia Influenza A+B (Quidel)	Sofia	NP aspirate/swab/wash Nasal wash	Required
BD Veritor System for Rapid Detection of Flu A+B (Becton Dickinson)	Veritor	NP swab/nasal swab	Required
Xpect Flu A&B (Remel)	Xpect	Nasal wash/swab Throat swab	No

**Abbreviations:** FDA = Food and Drug Administration; RIDTs = rapid influenza diagnostic tests; NP = nasopharyngeal.

present. During 2011, evidence of limited human-to-human transmission of H3N2v was observed in some cases, and human-to-human transmission might occur in the current outbreak. Enhanced surveillance for influenza H3N2v virus infection is indicated, especially in regions and states with confirmed H3N2v cases. The initial goal of enhanced surveillance is to detect the source and geographic spread of these viruses, but once cases are detected, particular emphasis should be placed on detection of ongoing transmission within the community through investigation of close contacts of patients with confirmed cases. In addition, surveillance in hospitals will be important to determine whether severe illnesses are occurring as a result of H3N2v infections.

The predominance of children among persons with confirmed H3N2v infections is consistent with serologic studies that found children less likely to have cross-protective antibodies than adults (4). However, confirmation of cases in adults highlights the fact

that persons of any age can be infected. Persons who are at increased risk for influenza complications (e.g., those with underlying chronic medical conditions, or who are pregnant, or aged <5 or ≥65 years, or who have weakened immune systems [5]) should avoid exposure to pigs and swine barns this summer, particularly if ill swine have been identified. Persons with increased risk for complications who develop influenza-like illness should see their health-care provider promptly to determine whether treatment with antiviral medications is warranted. Clinicians should consider antiviral treatment with oral oseltamivir or inhaled zanamivir in patients with suspected or confirmed H3N2v infection. Antiviral treatment is most effective when started as soon as possible after influenza illness onset (5).

The sensitivity of RIDTs to detect seasonal influenza viruses compared with virus isolation or rRT-PCR varies among commercial kits but has been shown to be low in some reports

(6–9). In this evaluation of seven RIDTs, the ability to detect H3N2v virus varied substantially among the tests. This evaluation emphasizes the fact that a negative RIDT result should not be considered as conclusive evidence of lack of infection with influenza A (H3N2)v. More data are needed on the clinical performance of all RIDTs in detecting H3N2v virus in various respiratory specimens. Results from RIDTs, both positive and negative, always should be interpreted in the broader context of the circulating influenza strains present in the area, level of clinical suspicion, severity of illness, and risk for complications in a patient with suspected infection. Clinicians should minimize the occurrence of false RIDT results by strictly following the manufacturer's instructions, collecting specimens soon after onset of influenza-like illness (ideally within the first 72 hours), and confirming RIDT results by sending a specimen to a public health laboratory (10). Additional CDC guidance on interpretation of RIDTs for testing of patients

TABLE 2. Number of 10-fold virus dilutions (maximum = five) detected by seven FDA-cleared RIDTs, by H3N2v strain designation — CDC, United States, 2012

						RIDT			
Subtype	Strain designation	TCID <sub>50</sub> /mL	BinaxNOW	Directigen	FluAlert	QuickVue	Sofia	Veritor	Xpect
H3N2v	A/Kansas/13/2009	10 <sup>4,5</sup>	1	4	U	U	2	4	4
H3N2v	A/Pennsylvania/14/2010	104.5	2	4	U	2	2	4	3
H3N2v	A/Minnesota/11/2010	10 <sup>4.5</sup>	U	3	U	U	3	3	2
H3N2v	A/Indiana/08/2011	10 <sup>6.0</sup>	1	3	U	U	2	3	2
H3N2v	A/Indiana/10/2011	10 <sup>4.0</sup>	U	3	U	U	2	4	2
H3N2v	A/West Virginia/06/2011	10 <sup>6.0</sup>	2	3	U	2	4	4	2
H3N2v	A/lowa/07/2011	10 <sup>4.5</sup>	2	4	1	1	3	4	3

Abbreviations: FDA = Food and Drug Administration; TCID<sub>50</sub>/mL = infectious titer of stock virus; RIDT = rapid influenza diagnostic test; U = undetected at any concentration tested.

<sup>\*</sup> Approved respiratory specimens according to manufacturer's package insert. Test performance has only been demonstrated for these specimen types.

#### What is known on this topic?

From July 2011 to April 2012, 13 cases of influenza A (H3N2)v virus infection in humans were reported. In July 2012, four new cases were reported from Indiana, all in persons who had contact with swine at a county fair.

What is added by this report?

From July 12 to August 9, 2012, a total of 153 cases of H3N2v virus infection were reported in four states (Hawaii, Illinois, Indiana, and Ohio); all patients for whom such information was available reported direct or indirect contact with swine. Testing of the sensitivity of rapid influenza diagnostic tests (RIDTs) for detection of influenza A (H3N2)v virus produced mixed results regarding the detection capabilities of the individual tests.

What are the implications for public health?

With the substantial increase in the number of cases of H3N2v virus infection during July–August, enhanced surveillance for detection of these cases is indicated. Health-care workers also should note that the sensitivity of RIDTs to detect H3N2v virus infection varies, and a negative RIDT should not be considered evidence of lack of infection with influenza A (H3N2)v.

with suspected H3N2v infection is available at http://www.cdc.gov/flu/swineflu/h3n2v-testing.htm.

Specimens from patients with influenza-like illness in whom H3N2v is suspected should be sent to public health laboratories for additional diagnostic testing. Public health laboratories are requested to continue to contact the CDC Influenza Division immediately when they identify these viruses to coordinate transfer of the specimen to CDC for additional testing.

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#### Notes from the Field

# Lymphocytic Choriomeningitis Virus Infections in Employees of a Rodent Breeding Facility — Indiana, May–June 2012

In late April 2012, an infectious disease physician contacted CDC regarding a patient with aseptic meningitis who worked at a rodent breeding facility in Indiana. Lymphocytic choriomeningitis virus (LCMV) infection was suspected, and LCMV-specific antibody was detected in blood and cerebrospinal fluid from the patient, confirming the diagnosis. LCMV is an arenavirus carried by the common house mouse. Persons become infected through close contact with infected rodents, through infected organ transplantation, or from mother to fetus. In immunocompetent adults, symptoms can range from mild febrile illness to meningeal symptoms (e.g., headache, stiff neck, or sensitivity to light). Congenitally infected infants can have a range of severe birth defects including hydrocephalus, chorioretinitis, blindness, and mental retardation (1). Infections in organ recipients, who are immunosuppressed, can have a case-fatality rate approaching 90% (2).

CDC notified the Indiana State Department of Health of a potential outbreak of LCMV infection at the rodent breeding facility and subsequently notified county health officials and the Indiana Board of Animal Health. A serosurvey was performed; 52 current and former employees of the facility consented to serum testing. Of the 52 tested, 13 (25%) demonstrated recent LCMV infection as evidenced by the prescence of immunoglobulin M (IgM) and IgG by enzyme-linked immunosorbent assay (ELISA). Nine employees who showed laboratory evidence of recent exposure reported experiencing a clinical illness consistent with LCMV; symptoms ranged from severe influenza-like illness to meningeal symptoms that required hospitalization. Of the persons experiencing illness, 89% were male; ages ranged from 20 to 48 years. No employees, including those not tested, were known to be pregnant at the time of the serosurvey. All employees who experienced clinical illness have since recovered. Three additional employees had evidence of a previous LCMV infection, with detectable anti-LCMV IgG and no IgM.

The rodent facility bred and raised mice and rats for sale as live and frozen feeder animals for reptiles or birds of prey. The facility housed approximately 155,000 adult mice and 14,000 adult rats. A representative sample of healthy-appearing adult rodents was tested for evidence of LCMV infection by ELISA and polymerase chain reaction. Of 1,421 mice tested,

296 (20.8%) had detectable anti-LCMV IgG, and 10 (0.7%) had detectable LCMV RNA. Of 399 rats tested, none were positive by ELISA or polymerase chain reaction. All living mice at the facility were euthanized. All rodents remaining in cold storage at the time of diagnosis also were disposed of in accordance with local environmental regulations. The buildings and equipment housing the mice were cleaned and disinfected. Used litter and contaminated feed were disposed of in accordance with local environmental regulations. Live mice distributed from the facility before the LCMV diagnosis currently are being followed to the point of purchase through an ongoing investigation.

Any persons with direct or indirect contact with these animals should be made aware of the public health risk and should seek medical evaluation if they have had any recent illness. Pregnant women or immunocompromised persons should be cautioned to avoid contact with rodents in general. Wild mice in the United States have a prevalence of LCMV estimated at 3.9%–13.4% (3). Any additional rodent populations that have come into direct contact with potentially infected mice should be depopulated.

Employers of rodent breeding facilities of all kinds should make their employees aware that working with rodents can expose them to LCMV and should educate workers regarding risks for exposure, including potential health effects. Employers also should work with their local health departments to develop guidance material on disease prevention and provide the recommended personal protective equipment for employees. Routine serologic testing of rodents can be used to detect and control LCMV infections. Evidence of LCMV infection in rodents should be dealt with promptly to prevent human illness from occurring. Purchasers of frozen rodents used to feed another pet should be reminded to always wear plastic gloves when handling the rodents and to wash their hands afterward.

#### Reported by

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#### Announcement

#### **Updated Online NCHHSTP Atlas**

CDC's National Center for HIV/AIDS, Viral Hepatitis, STD, and TB Prevention (NCHHSTP) recently launched an update to the NCHHSTP Atlas, an interactive, online mapping tool and platform for accessing data collected by the center. With this update, the atlas allows users to observe disease trends and patterns of not only human immunodeficiency virus infection, acquired immunodeficiency syndrome, and some sexually transmitted infections (i.e., chlamydia, gonorrhea, and primary and secondary syphilis), but also of acute viral hepatitis A, B, and C, and tuberculosis. The atlas also allows users to create detailed reports, maps, and other graphics of these surveillance data.

The interactive atlas is a valuable tool to help public health professionals, researchers, community leaders, health-care providers, and others view overlapping disease trends, set research priorities, and plan prevention and care services. The NCHHSTP Atlas is available at http://www.cdc.gov/nchhstp/atlas.

#### Notices to Readers

### Scanned 1952–1982 Issues of *MMWR* Available Online

Issues from the first 30 years of MMWR are now available to the public online as one of the collections in "CDC Stacks," an institutional repository. As with other documents in CDC Stacks, the MMWR issues are in portable document format (PDF), and the text can be searched electronically.

In addition to the first 30 years of MMWR, CDC Stacks contains documents spanning the history of the agency, including CDC Open Access, Influenza Surveillance Reports, and CDC Guidelines and Recommendations. CDC Stacks allows users to browse journal articles by public health subjects and explore collections of documents on relevant topics. New documents are added each week. CDC Stacks is available at http://stacks.cdc.gov. The MMWR collection is available at http://stacks.cdc.gov/mmwr.

### Final 2011 Reports of Nationally Notifiable Infectious Diseases

The tables listed in this report on pages 625–637 summarize finalized data, as of June 30, 2012, from the National Notifiable Diseases Surveillance System (NNDSS) for 2011. These data will be published in more detail in the Summary of Notifiable Diseases — United States, 2011 (1). Because no cases were reported in the United States during 2011, the following diseases do not appear in these early release tables: diphtheria; eastern equine encephalitis virus disease, nonneuroinvasive; poliomyelitis, paralytic; poliovirus infection, nonparalytic; rubella, congenital syndrome; severe acute respiratory syndrome-associated coronavirus disease; smallpox; vancomycin-resistant Staphylococcus aureus; western equine encephalitis virus disease, neuroinvasive and nonneuroinvasive; yellow fever; and viral hemorrhagic fevers.

Policies for reporting NNDSS data to CDC can vary by disease or reporting jurisdiction depending on case status classification (i.e., confirmed, probable, or suspected). The publication criteria used for the 2011 finalized tables are listed in the "Print Criteria" column of the NNDSS event code list, available at http://wwwn.cdc.gov/nndss/document/nndss\_event\_code\_list\_july\_2011\_28\_final.pdf. The NNDSS website is updated annually to include the latest national surveillance case definitions approved by the Council of State and Territorial Epidemiologists (CSTE) for enumerating data on nationally notifiable infectious diseases.

Population estimates are from the National Center for Health Statistics postcensal estimates of the resident population of the United States for July 1, 2010–July 1, 2011, by year, county, single-year of age (0 to ≥85 years), bridged-race, (white, black or African American, American Indian or Alaska Native, Asian or Pacific Islander), Hispanic origin (not Hispanic or Latino, Hispanic or Latino), and sex (vintage 2010), prepared under a collaborative arrangement with the U.S. Census Bureau. Population estimates for states are available at http://www.cdc.gov/nchs/nvss/bridged\_race/data\_documentation.htm#vintage2010 as of May 31, 2012. Population estimates for territories are 2010 estimates from the U.S. Census Bureau (2).

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TABLE 2. Reported cases of notifiable diseases,\* by geographic division and area — United States, 2011

						Arl	ooviral diseases	i <sup>†</sup>			
	Total resident			serogroup irus	Eastern equine encephalitis virus	Powas	san virus		encephalitis irus	West	Nile virus
Area	population (in thousands)	Anthrax	Neuro- invasive	Nonneuro- invasive	Neuro- invasive	Neuro- invasive	Nonneuro- invasive	Neuro- invasive	Nonneuro- invasive	Neuro- invasive	Nonneuro- invasive
United States	309,049	1	120	17	4	12	4	4	2	486	226
New England	14,474				1					15	2
Connecticut	3,527		******	remer	*****	Term	*****			8	1
Maine Massachusetts	- 1,313 6,631				1		******		-	5	_ 1
New Hampshire	1,324						-				
Rhode Island	1,057					*****				1	
Vermont	622		****		1	1				1 35	
Mid. Atlantic New Jersey	40,943 8,733		****	******		1		-		33 2	22 5
New York (Upstate)	11,146	-			1					19	14
New York City	8,431	-	******							9	2
Pennsylvania	12,633					1				5	1
E.N. Central Illinois	46,521 12,944		51 1	12	1	2	2			73 22	28
Indiana	6,445		2							7	12 2
Michigan	9,931	*****	1				*****	*****	******	32	2
Ohio	11,532		44	6						10	11
Wisconsin	5,669		3	6	1	2	2			2	1
W.N. Central Iowa	20,451 3,023		1		1	9	2		1	31 5	29 4
Kansas	2,841									3 4	4
Minnesota	5,290		1			9	2	rmen	*****	1	1
Missouri	6,012		****	*****	1	****		******	1	6	4
Nebraska North Dakota	1,811 654		****							14 1	15 3
South Dakota	820										2
S. Atlantic	59,659	1	52	5					1	67	27
Delaware	891						*****			1	-
District of Columbia	611	north A	****	******	*****	****				10	5
Florida Georgia	18,678 9,908	1 —	1 2						_	20 14	4 8
Maryland	5,737								1	10	9
North Carolina	9,459		26					****	*****	2	
South Carolina Virginia	4,597 7,952		1	1			******			8	1
West Virginia	1,826		22	4			<del></del>			2	
E.S. Central	18,367		15			-	-	1		56	24
Alabama	4,730		1					1		5	
Kentucky	4,339		1		*****		-			4	1
Mississippi Tennessee	2,960 6,338		1 12	******	****					31 16	21 2
W.S. Central	36,376						MATERIAL PROPERTY.	3	*****	28	17
Arkansas	2,910				******			3	****	1	
Louisiana	4,529									6	4
Oklahoma Texas	3,724 25,213									1 20	7
Mountain	22,380		1					******	*****	71	35
Arizona	6,677		1	*****			notes.		******	49	20
Colorado	5,095									2	5
Idaho	1,560					****			MANAGE	1	2
Montana Nevada	980 2,655								<del></del>	1 12	4
New Mexico	2,034		*****	******	*****					4	
Utah	2,831		*****	-	****		*****	***************************************		1	2
Wyoming	548		_	MANA	****		******	****	- Angelogo	1	2
Pacific Alaska	49,878 709						*****			110	48
California	709 37,267				_					110	48
Hawali	1,300										_
Oregon	3,856	****		•		*****	****	******			
Washington	6,746										
Territories	EE										
American Samoa C.N.M.I.	55 54										
Guam	159						_				
Puerto Rico	3,722	******	*****	*******	***************************************	******	-				
U.S. Virgin Islands	106	*****				*****					

N: Not reportable U: Unavailable —: No reported cases C.N.M.I.: Commonwealth of Northern Mariana Islands.

N: Not reportable U: Unavailable —: No reported cases C.N.M.I.: Commonwealth of Northern Mariana Islands.

No cases of diphtheria; eastern equine encephalitis virus disease, nonneuroinvasive; poliomyelitis, paralytic; poliovirus infection, nonparalytic; rubella, congenital syndrome; severe acute respiratory syndrome-associated coronavirus disease; smailpox; vancomycin-resistant Staphylococcus aureus; western equine encephalitis virus disease, neuroinvasive and nonneuroinvasive; yellow fever; and viral hemorrhagic fevers were reported in the United States during 2011. Data on hepatitis B virus, perinatal infection, chronic hepatitis B and hepatitis C virus infection (past or present) are not included because they are undergoing data quality review. Data on human immunodeficiency virus (HIV) infections are not included because HIV infection reporting has been implemented on different dates and using different methods than for acquired immunodeficiency syndrome (AIDS) case reporting.

<sup>†</sup> Totals reported to the Division of Vector-Borne Infectious Diseases (DVBD), National Center for Emerging and Zoonotic Infectious Diseases (NCZVED) (ArboNET Surveillance), as of April 17, 2012.

TABLE 2. (Continued) Reported cases of notifiable diseases,\* by geographic division and area — United States, 2011

	-		Bot	ulism		_		Chlamydia trachomati
Area	Babesiosis	Total	Foodborne	Infant	Other <sup>†</sup>	Brucellosis	Chancroid <sup>§</sup>	trachomati infection <sup>§</sup>
Jnited States	1,128	153	24	97	32	79	8	1,412,791
lew England	378					1	2	48,146
Connecticut	74							13,649
Maine	9						amn	3,094
Massachusetts	208		***	****	****	1	2	22,764
New Hampshire	13	****	****					3,010
Rhode Island	73					_		4,146
Vermont	1							1,483
hid. Atlantic	584	29	2	27		7		181,856
New Jersey	166	11		11		1		26,209
New York (Upstate)	361	2	1	1	-hours			37,494
New York City	57	4	1	3		3		65,269
Pennsylvania	N	12		12		3		52,884
.N. Central	80	3	2	1		10	1	219,580
Illinois						8		64,939
Indiana		1	1			***		27,801
Michigan		*****	*****			1	1	49,568
Ohio	N	2	1	1		1		52,653
Wisconsin	80							24,619
/.N. Central	74	2		1	1	1		78,726
lowa	/ <del>4</del>			1	<u>'</u>	i		10,705
Kansas	N	1	******	1			THE STATE OF THE S	10,598
Minnesota	73	1		,	1			16,902
Missouri	Ň					-	*****	27,887
Nebraska	19							6,780
North Dakota	1							2,445
South Dakota	N				shapet	77777		3,409
		0						
. Atlantic Delaware	5 1	9 2	1	8		13	2	293,101
District of Columbia			-	2			*****	4,508 6,585
Florida		*****			****	6		76,033
Georgia	***	1	1	***		5		54,403
	4	2		2		1		27,212
Maryland North Carolina	N N	2				<u> </u>		54,819
South Carolina				2		1	2	28,932
Virginia	N	2		2				36,314
West Virginia		<u></u>	****	<u>~</u>				4,295
		7		7				
.S. Central Alabama	2 1	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	<del></del>	,		4		98,576
Kentucky	N,	2		2	-	1		29,626 16,629
Mississippi	14	2		2	****	1		21,216
Tennessee	1	3		3		2		31,105
V.S. Central		6	1	4	1	15	1	187,144
Arkansas			-	North Control of the		3		16,052
Louisiana								31,614
Oklahoma	N	]	1			1		14,596
Texas	N	5		4	1	11	1	124,882
lountain	****	26	10	15	1	10	1	90,226
Arizona	-	5	2	3	_	3	1	29,251
Colorado		4		3	1		munn	21,811
ldaho	N	2		2		2	*****	4,699
Montana		_						3,406
Nevada	N	1		1	-			10,507
New Mexico	***	2		2		2	******	11,374
Utah	N	12	8	4		3		7,086
Wyoming	navara.	*****			*****	*****	******	2,092
acific	5	71	8	34	29	18	1	215,436
Alaska	Lancate Control of the Control of th	6	6					5,739
California	4	58	1	30	27	15	1	166,773
Hawaii				<del></del>		1		6,001
Oregon	1	2	1	1		1		13,643
Washington	*****	5		3	2	1		23,280
erritories								
American Samoa	.gama	*****	****		******		-	*****
C.N.M.L				****				
Guam								1,071
Puerto Rico	N				N	*****		5,634
					• •			21001

N: Not reportable U: Unavailable —: No reported cases C.N.M.I.: Commonwealth of Northern Mariana Islands.

<sup>\*</sup> No cases of diphtheria; eastern equine encephalitis virus disease, nonneuroinvasive; poliomyelitis, paralytic; poliovirus infection, nonparalytic; rubella, congenital syndrome; severe acute respiratory syndrome-associated coronavirus disease; smallpox; vancomycin-resistant Staphylococcus aureus; western equine encephalitis virus disease, neuroinvasive and nonneuroinvasive; yellow fever; and viral hemorrhagic fevers were reported in the United States during 2011. Data on hepatitis B virus, perinatal infection, chronic hepatitis B and hepatitis C virus infection (past or present) are not included because they are undergoing data quality review. Data on human immunodeficiency virus (HIV) infections are not included because HIV infection reporting has been implemented on different dates and using different methods than for acquired immunodeficiency syndrome (AIDS) case reporting.

<sup>†</sup> Includes cases reported as wound and unspecified botulism.

<sup>&</sup>lt;sup>5</sup> Totals reported to the Division of STD Prevention, NCHHSTP, as of June 7, 2012.

TABLE 2. (Continued) Reported cases of notifiable diseases,\* by geographic division and area — United States, 2011

				Cryptosporidiosis			Dengue	virus infection <sup>†</sup>
Area	Cholera	Coccidioidomycosis	Total	Confirmed	Probable	Cyclosporiasis	Dengue fever	Dengue hemorrhagic fever
United States	40	22,634	9,250	6,130	3,120	151	251	3
lew England	4	2	418	358	60	12	4	
Connecticut		N	71	71	*****	10	1	
Maine		N	51	19	32	Ŋ		
Massachusetts New Hampshire	4	1	168 68	168 40	28	2		
Rhode Island		1	12	12			****	
Vermont	****	N	48	48		N	3	
Aid. Atlantic	14	6	904	824	80	38	69	
New Jersey	1	N	56	55	1	. 8		
New York (Upstate)	2	N	234	226	8	11	8	
New York City Pennsylvania	10 1	N 6	86 528	86 457	71	19 N	45 16	
•	2	56	2,676	1,476	1,200	7	21	2
.N. Central Illinois	1	N	213	31	182		6	2
Indiana		N	261	79	182	*****	2	Ar-
Michigan	1	36	358	325	33	7	6	
Ohio		20	1,106	303	803		2	
Wisconsin			738	738		_	5	-
V.N. Central	1	130	1,563	714	849	3	13	uname.
Iowa Kansas	1	N N	364 42	61 42	303	1	5 1	
Minnesota		104	309	309	******		6	****
Missouri		18	495	156	339	1		
Nebraska		8	175	124	51	1		
North Dakota		N	32	1	31	N	1	******
South Dakota		N	146	21	125			
. Atlantic	13	5	1,239	791	448	69	92	1
Delaware District of Columbia		MAAAA	7 N	7		1 N	2	*****
Florida	11	· N	437	203	234	58	 66	****
Georgia	1	N	307	307		6	6	
Maryland		5	70	66	4	1	6	
North Carolina		N	115	69	46	1	4	
South Carolina	1	N N	132 140	66 54	66 86	2	1 7	<del></del> 1
Virginia West Virginia		N N	31	54 19	12	<u>~</u>		
.S. Central	2		457	301	156	2	11	
Alabama		N	138	16	122	N	4	*****
Kentucky	2	N	177	160	17	N	4	****
Mississippi		N	50	50		N	******	
Tennessee		N	92	75	17	2	3	
V.S. Central	1	3	712	579	133	15	10	
Arkansas		N 3	32	32				
Louisiana Oklahoma		3 N	87 89	87 2	<del></del> 87	1	3	
Texas	1	Ň	504	458	46	14	7	******
lountain	1	16,712	641	552	89	1	6	
Arizona		16,467	46	42	4	*****	2	
Colorado		N	147	126	21	******	-	******
Idaho		N	111	79	32	N		
Montana		5	77 17	77 3	14	N N		
Nevada New Mexico	1	104 75	134	134	[4	11	1 2	M14444
Utah		58	63	62	1		1	NAME OF THE PARTY
Wyoming		3	46	29	17			
acific	2	5,720	640	535	105	4	25	
Alaska	1	N	12	12		N		<del></del>
California	1	5,697	332	332			5	****
Hawaii		N 12	1 207	1 179	30	*****	11	<del></del>
Oregon Washington		10 13	207 88	179	28 77	4	9	
		10				.,1		
erritories American Samoa		N	N			N		
C.N.M.I.		N		****				
Guam	*******	*****			*****	****	*****	
Puerto Rico	1	N	N	_		N	1,507	34
U,S. Virgin Islands		****	•					

N: Not reportable U: Unavailable —: No reported cases C.N.M.I.: Commonwealth of Northern Mariana Islands.

<sup>\*</sup> No cases of diphtheria; eastern equine encephalitis virus disease, nonneuroinvasive; poliomyelitis, paralytic; poliovirus infection, nonparalytic; rubella, congenital syndrome; severe acute  $respiratory syndrome-associated coronavirus disease; small pox; van comycin-resistant {\it Staphylococcus aureus}; western equine encephalitis virus disease, neuroinvasive and nonneuroinvasive; {\it Staphylococcus aureus}; {\it Continuous aureus}; {\it Conti$ yellow fever; and viral hemorrhagic fevers were reported in the United States during 2011. Data on hepatitis B virus, perinatal infection, chronic hepatitis C virus infection (past or present) are not included because they are undergoing data quality review. Data on human immunodeficiency virus (HIV) infections are not included because HIV infection reporting has been implemented on different dates and using different methods than for acquired immunodeficiency syndrome (AIDS) case reporting.

† Total number of reported laboratory-positive dengue cases including all confirmed cases [by anti-dengue virus (DENV) molecular diagnostic methods or seroconversion of anti-DENV [gM] and all probable cases (by a single, positive anti-DENV [gM]). Totals reported to the Division of Vector-Borne Diseases (DVBD), National Center for Emerging and Zoonotic Infectious Diseases

<sup>(</sup>NCEZID) (ArboNET Surveillance), as of April 17, 2012.

TABLE 2. (Continued) Reported cases of notifiable diseases,\* by geographic division and area — United States, 2011

		Ehrlichiosis/	Anaplasmosis			
Area	Anaplasma phagocytophilum	Ehrlichia chaffeensis	Ehrlichia ewingii	Undetermined	Giardiasis	Gonorrhea <sup>†</sup>
United States	. 2,575	850	13	148	16,747	321,849
New England	461	4		2	1,594	5,612
Connecticut	152	<del></del>	*****		233	2,449
Maine	26	1	Wester	-	171	272
Massachusetts New Hampshire	172 31	1	****	1	758 130	2,353
Rhode Island	72	2	******	1	130 79	130 360
Vermont	8				223	48
Mid. Atlantic	482	108		25	3,293	41,824
New Jersey	126	60		7	437	7,348
New York (Upstate)	314	41	******	11	1,144	6,240
New York City	36	4		MANAGEMENT	917	14,466
Pennsylvania	6	3		7	795	13,770
E.N. Central	710	42	****	58	2,657	58,022
Illinois	11	25		***	407	17,037
Indiana			****	18	324	6,569
Michigan Ohio	9	4	****	5	550	12,901
Wisconsin	690	6 7	*****	1 34	799 577	16,726 4,789
	808	178	6	25		
W.N. Central Iowa	N N	176 N	N N	N N	1,769 271	16,420
Kansas	6	18		1	139	1,920 2,209
Minnesota	770	7	1	10	672	2,284
Missouri	25	151	5	13	344	7,802
Nebraska	1	1	******	1	179	1,352
North Dakota	3	<del></del>			54	251
South Dakota	3	1			110	602
S. Atlantic	72	272	6	16	2,756	79,089
Delaware	1	15	2		34	827
District of Columbia Florida	N 11	N 15	N	N	56	2,569
Georgia	11	23	1	3	1,255 651	19,689 16,428
Maryland	7	33	2	<del></del>	291	6,458
North Carolina	21	83		1	N	17,454
South Carolina		2		1	117	8,350
Virginia	21	100	1	9	290	6,518
West Virginia	****	1	- Marine	2	62	796
E.S. Central	15	78	1	14	171	27,134
Alabama	4	5		MAAAAA	171	9,132
Kentucky Mississippi	<del>-</del>	16 3		******	N	4,521
Tennessee	10	54	1	14	N N	5,814 7,667
	20	167		1	349	
W.S. Central Arkansas	8	53		₹ *********	123	49,001 4,687
Louisiana	1	<del></del>	****	1	226	9,169
Oklahoma	9	110		·		4,215
Texas	2	4			N	30,930
Mountain	1		Time.	5	1,326	11,336
Arizona			****	4	133	4,564
Colorado	N	N	N	N	445	2,363
Idaho	N	N	N	N	178	162
Montana	N	N	N	N	86	85
Nevada New Mexico	N.	N	N.	N.	79	2,000
Utah	14	14	N 	N 1	108 256	1,839 277
Wyoming	1		MARKE		41	46
Pacific	6	1	Patricia	2	2,832	33,411
Alaska	Ň	N	N	N	101	984
California				2	1,728	27,516
Hawaii	N	` N	N	Ñ	38	685
Oregon	6	*****	***		436	1,489
Washington		1	Market .		529	2,737
Territories						
American Samoa	N	N	N	N		
C.N.M.I.					*******	
Guam Duarta Disa	N	N	N	N	0.4	96
Puerto Rico U.S. Virgin Islands	N	N	N	N	84	341 139

N: Not reportable U: Unavailable —: No reported cases C.N.M.i.: Commonwealth of Northern Mariana Islands.

<sup>\*</sup> No cases of diphtheria; eastern equine encephalitis virus disease, nonneuroinvasive; poliomyelitis, paralytic; poliovirus infection, nonparalytic; rubella, congenital syndrome; severe acute respiratory syndrome-associated coronavirus disease; smallpox; vancomycin-resistant Staphylococcus aureus; western equine encephalitis virus disease, neuro invasive and nonneuro invasive; yellow fever; and viral hemorrhagic fevers were reported in the United States during 2011. Data on hepatitis B virus, perinatal infection, chronic hepatitis B and hepatitis C virus infection (past or present) are not included because they are undergoing data quality review. Data on human immunodeficiency virus (HIV) infections are not included because HIV infection reporting has been implemented on different dates and using different methods than for acquired immunodeficiency syndrome (AIDS) case reporting.

† Totals reported to the Division of STD Prevention, NCHHSTP, as of June 7, 2012.

TABLE 2. (Continued) Reported cases of notifiable diseases,\* by geographic division and area — United States, 2011

	•	Haemophilus influ	enza, invasive disease	3		Hantavirus	Hemolytic uremic
Area	All ages, serotypes	Serotype b	Age <5 years Nonserotype b	Unknown serotype	Hansen disease (leprosy)	pulmonary syndrome	syndrome, postdiarrheal
United States	3,539	14	145	226	82	23	290
New England	252		9	6	3	2.5	12
Connecticut	65			4		N	2
Maine	26		1		N		ž
Massachusetts	121		7		2		5
New Hampshire	17		1	1			****
Rhode Island	16				1		2
Vermont	7	MALANA		1	N		1
Mid. Atlantic	771	****	13	45	4	1	21
New Jersey New York (Upstate)	123 195		8	9 1	 N	1	4 13
New York City	187	*****		15	4	-	4
Pennsylvania	266	*****	5	20			N
E.N. Central	645	3	30	28	3		36
Illinois	188	-	6	8		man	7
Indiana	117	1	9		1	*****	
Michigan	72			14			9
Ohio	173	2	15		2		5
Wisconsin	95 *			6	*****	*****	15
W.N. Central	224	2	4	23	2	2	49
lowa	3					1	13
Kansas Minnesota	23			3			4
Missouri	71 80	. 1	3	13			12
Nebraska	30	1	1	4	4		. 20
North Dakota	16			3	N		
South Dakota	1	www				1	Author
S. Atlantic	783	2	25	46	14		24
Delaware	6	Manage					
District of Columbia	ī				· ·	N	N
Florida	232			23	11	*****	4
Georgia	140		10	10		****	7
Maryland	95 0.5	1	7	1	2		2
North Carolina South Carolina	85 79		2	8 3	uphraphite Saparithy	********	5 3
Virginia	108	1	5		1		3
West Virginia	37		1	1	Ň		
E.S. Central	225	3	14	7	1	****	25
Alabama	57	1	5	·		N	9
Kentucky	41		ì	4	****		Ñ
Mississippi	19	1	1		1	N	1
Tennessee	108	1	7	3		4	15
W.S. Central	163		9	13	19		41
Arkansas	35		5		2		12
Louisiana	53		•	13	1		
Oklahoma Texas	73 2		4 N	A.I	N 16	-	7
				N	16	4.6	22
Mountain	294	3	31	16	2	16	25
Arizona Colorado	95 67	1	13 5	2	****	3 3	5 6
Idaho	21		2	1	******	3	3
Montana	3		<u></u>			2	1
Nevada	17	****		3	1	2	2
New Mexico	47	*****	2	10		5	2
Utah	42	2	9	*****	1		5
Wyoming	2				*****		1
Pacific	182	1	10	42	34	4	57
Alaska	26	-		11		N	N
California	44			27	14	*****	42
Hawaii	32	MARIA	······	4	20		1
Oregon Washington	72 8	1	3 7	***************************************	N	2 2	14
		r	,		14		<del>-</del>
Territories						ki	k.1
American Samoa C.N.M.I.	<del>_</del>					N 	N 
Guam		Marine	was a			N	
Puerto Rico				-		14	N
U.S. Virgin Islands	N					N	N

N: Not reportable U: Unavailable —: No reported cases C.N.M.J.: Commonwealth of Northern Mariana Islands.

\*No cases of diphtheria; eastern equine encephalitis virus disease, nonneuroinvasive; poliomyelitis, paralytic; poliovirus infection, nonparalytic; rubella, congenital syndrome; severe acute respiratory syndrome-associated coronavirus disease; smallpox; vancomycin-resistant Staphylococcus aureus; western equine encephalitis virus disease, neuroinvasive and nonneuroinvasive; yellow fever; and viral hemorrhagic fevers were reported in the United States during 2011. Data on hepatitis B virus, perinatal infection, chronic hepatitis B and hepatitis C virus infection (past or present) are not included because they are undergoing data quality review. Data on human immunodeficiency virus (HIV) infections are not included because HIV infection reporting has been implemented on different dates and using different methods than for acquired immunodeficiency syndrome (AIDS) case reporting.

TABLE 2. (Continued) Reported cases of notifiable diseases,\* by geographic division and area — United States, 2011

	Н	epatitis, viral, acute		HIV	Influenza-associated		
Area	A	В	С	diagnoses†	pediatric mortality <sup>§</sup>	Legionellosis	Listeriosis
Inited States	1,398	2,903	1,229	35,266	118	4,202	870
ew England	77	97	88	1,003	4	406	61
Connecticut	18	19	47	305	1	81	18
Maine	6	8	12	46	1	18	4
Massachusetts	39	67	23	523	1	240	32
New Hampshire		3	N	40		26	4
Rhode Island	8	U	ñ	88		29	3
Vermont	6		6	1	1	12	
id. Atlantic	252	291	140	5,628	15	1,353	158
New Jersey	79	73	53	812	4	235	33
New York (Upstate)	47	54	44	1,301	2	400 216	48 30
New York City	66	80 84	8 35	2,246 1,269	3 6	502	47
Pennsylvania	60						
N. Central	214	353	143	3,641	19	864	116
llinois	73	85	6	1,351	7	151	34
ndiana	24	70 91	84	434 610	2	71 187	11 29
Michigan Ohio	70 39	90	32 6	987	6 1	386	29
Visconsin	8	17	15	259	3	69	13
			35	1,085	9	122	62
.N. Centrai	59	124				11	62 S
owa	8	15 15	8	116 126		14	14
Kansas Viinnesota	4 27	20	17	283	3	29	6
Missouri	13	60	8	481	1	55	21
Vebraska	5	12	2	46		8	9
North Dakota				12	1	3	6
South Dakota	2	2		21	4	2	1
Atlantic	222	775	284	10,925	22	640	111
Delaware	2	13	U	99		24	
District of Columbia				495	*****	N	N
Florida	87	213	64	4,890	2	185	38
Georgia	27	142	53	1,431	4	55	9
Maryland	26	62	35	851		143	19
North Carolina	31	109	60	1,439	10	83	21
South Carolina	11	39	1	771	<del></del>	25	6
Virginia	30	84	25	857	5	93	15
West Virginia	8	113	46	92	1	32	3
S. Central	48	519	248	2,191	2	180	22
Alabama	8	119	23	592		29	9
Kentucky	10	151	142	233	2	53	4
Mississippi	7	57	U	552	****	14	4
Tennessee	23	192	83	814		84	5
'.S. Central	157	423	97	4,967	16	165	79
Arkansas	3	. 57		199	-	14	6
_oulsiana	5	62	7	1,281	1	25	7
Oklahoma	11	100	53 37	262 3,225	4	15 111	15 51
l exas	138	204			11.		
ountain	129	88	85	1,410	12	147	98
Arizona	77	14	U	494	4	46	8
Iolorado	21	23	28	362	3	41 9	51
daho	6 3	2	12 9	16 17		1	5 3
Montana Nevada	3 5	29	10	320	3	16	5 5
vevada Vew Mexico	7	10	14	111	1	12	15
Jtah	8	10	10	76	1	18	5
Vyoming	ž		2	. 14	-	4	6
cific	240	233	109	4,416	19	325	163
CITIC Naska	4	3		25			103
Talifornia	186	157	48	3,679	16	261	123
Hawaii	8	6		50	1	5	12
Oregon	11	32	20	213	i	22	9
Washington	31	35	41	449	1	37	19
rritories				······································			
erritories American Samoa	<b>1475</b>	*****				N	N
.N.M.L						14	
Buam	43	120	70	1	MAAAAA		
uerto Rico	21	28	Ň	436		9	
J.S. Virgin Islands		5		22		1	

N: Not reportable U: Unavailable —: No reported cases C.N.M.I.: Commonwealth of Northern Mariana Islands.

\* No cases of diphtheria; eastern equine encephalitis virus disease, nonneuroinvasive; poliomyelitis, paralytic; poliovirus infection, nonparalytic; rubella, congenital syndrome; severe acute respiratory syndrome-associated coronavirus disease; smallpox; vancomycin-resistant Staphylococcus aureus; western equine encephalitis virus disease, neuroinvasive and nonneuroinvasive; yellow fever; and viral hemorrhagic fevers were reported in the United States during 2011. Data on hepatitis B virus, perinatal infection, chronic hepatitis B and hepatitis C virus infection (past or present) are not included because they are undergoing data quality review. Data on human immunodeficiency virus (HIV) infections are not included because HIV infection reporting has been implemented on different dates and using different methods than for acquired immunodeficiency syndrome (AIDS) case reporting.

<sup>†</sup> Data on HIV diagnoses include persons with a diagnosis of HIV infection regardless of stage of disease (i.e., AIDS status) at diagnosis. Total number of HIV diagnoses case counts was reported to the Division of HIV/AIDS Prevention, National Center for HIV/AIDS, Viral Hepatitis, STD, and T8 Prevention (NCHHSTP) through December 31, 2011.

<sup>§</sup> Totals reported to the Division of Influenza, National Center for Immunization and Respiratory Diseases (NCIRD), as of December 31, 2011.

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TABLE 2. (Continued) Reported cases of notifiable diseases,\* by geographic division and area — United States, 2011

		Lyme disease				Measles	· · · · · · · · · · · · · · · · · · ·
Area	Total	Confirmed	Probable	Malaria	Total	Indigenous	Imported <sup>†</sup>
United States	33,097	24,364	8,733	1,724	222	142	80
New England	8,602	6,080	2,522	109	28	18	10
Connecticut	3,039	2,004	1,035	20	1	******	1
Maine	1,006	801	205	6			
Massachusetts	2,476	1,801	675	68	24	17	7 1
New Hampshire Rhode Island	1,299 159	887 111	412 48	3 6	1	*****	1
Vermont	623	476	147	6	1	1	
Mid. Atlantic	14,114	11,255	2,859	438	49	35	14
New Jersey	4,262	3,398	864	97	4	3	1
New York (Upstate)	3,759	2,678	1,081	53	7	4	3
New York City	731	440	291	227	25	16	9
Pennsylvania	5,362	4,739	623	61	13	12	1
E.N. Central	4,094	2,808	1,286	174	21	15	6
Illinois	194	194		66	3	1	2
Indiana	94	81	13	14	14	13	1
Michigan Ohio	104 53	89 36	15 17	34 41		1	1
Wisconsin	3,649	2,408	1,241	19	2	*****	2
	2,291	1,304	987	109	34	30	4
W.N. Central lowa	100	72	28	22	1		1
Kansas	17	11	6	10	6	6	
Minnesota	2,124	1,185	939	46	26	23	3
Missouri	8	5	3	21			
Nebraska	11	7	4	8	****	-	
North Dakota	27	22	5		1	1	
South Dakota	4	2	2	2			
S. Atlantic	3,637	2,720	917	478	20	7	13
Delaware District of Columbia	873 N	767	106	7 18	1 N	1	
Florida	115	78	37	99	8	3	5
Georgia	32	32		91			
Maryland	1,351	938	413	128	2	*****	2
North Carolina	88	18	70	49	2		2
South Carolina	37	24	13	7	7		
Virginia West Virginia	1,023 118	756 107	267 11	78 1		3	4
~	69	20	49	41	4	1	3
E.S. Central Alabama	24	9	15	9	*	******	J
Kentucky	3	3	1.3	10	1		1
Mississippi	5	3	2	1			
Tennessee	37	5	32	21	3	1	2
W.S. Central	78	31	47	121	6	5	1
Arkansas	<del></del>			7			-
Louisiana	2	1	1	2	*****	****	
Oklahoma	2	2		10			a
Texas	74	28	46	102	6	5 **-	1
Mountain	52	32	20	67	22	15	7
Arizona Colorado	15	8	7	21 24	2	******	2
Idaho	4	3	1	2			
Montana	11	9	2	2	*****		****
Nevada	5	3	2	8	1	******	1
New Mexico	6	2	4	5	4	1	3
Utah	9	6	3 1	5	15	14	1
Wyoming	2	114		187	38	16	22
Pacific	160	9	46 2	187		16	22
Alaska California	11 92	79	13	129	31	12	19
Hawaii	92 N	N	N	7	J1	12	
Oregon	38	9	29	22	3	2	1
Washington	19	17	2	24	4	2	2
Territories	·····						
American Samoa	N			1			
C.N.M.I.		*******	*******	Moreok	-	********	******
Guam	**	******	-		*****		
Puerto Rico	N N	*****		1	*****	AAAAAA.	
U,S. Virgin Islands	N						

N: Not reportable U: Unavailable —: No reported cases C.N.M.I.: Commonwealth of Northern Mariana Islands.

\* No cases of diphtheria; eastern equine encephalitis virus disease, nonneuroinvasive; poliomyelitis, paralytic; poliovirus infection, nonparalytic; rubella, congenital syndrome; severe acute respiratory syndrome-associated coronavirus disease; smallpox; vancomycin-resistant Staphylococcus aureus; western equine encephalitis virus disease, neuroinvasive and nonneuroinvasive; yellow fever; and viral hemorrhagic fevers were reported in the United States during 2011. Data on hepatitis B virus, perinatal infection, chronic hepatitis B and hepatitis C virus infection. (past or present) are not included because they are undergoing data quality review. Data on human immunodeficiency virus (HIV) infections are not included because HIV infection reporting has been implemented on different dates and using different methods than for acquired immunodeficiency syndrome (AIDS) case reporting.

<sup>†</sup> Imported cases include only those directly related to importation from other countries.

TABLE 2. (Continued) Reported cases of notifiable diseases,\* by geographic division and area --- United States, 2011

TABLE 2. (Continued) Re		***************************************	Meningococcal dise				
Area	All serogroups	Serogroup A, C, Y, and W-135	Serogroup B	Serogroup other	Serogroup unknown	Mumps	Novel Influenza A virus infections
United States	759	257	159	20	323	404	14
New England	29	18	7	2	2	12	2
Connecticut	3	2	****		ī		
Maine	.5	3	2	*****	-	2	2
Massachusetts	14 1	8 1	3	2	1	4	
New Hampshire Rhode Island	1	:	1	•	*******	5	****
Vermont	5	4	1			í	
Mid. Atlantic	92	20	5	1	66	55	3
New Jersey	13			*******	13	13	<del></del>
New York (Upstate)	23	18	4	1	******	10	
New York City	31		******		31	29	
Pennsylvania	25	2	1		22	3	3
E.N. Central	115	59	44	6	6	110	3
Illinois	35	19	12	1	3	78	
Indiana	25	12	12	1	1	3	2
Michigan Ohio	12 24	4 13	6 7	1 2	2	9 16	*****
Wisconsin	19	11	7	1	<u>~</u>	4	1
W.N. Central	63	15	15	3	30	35	4
lowa	14	6	6	1	1	8	3
Kansas	5				s s	4	
Minnesota	15	6	8	1		2	1
Missouri	15	****	and the same of th	www	15	11	- mariners
Nebraska	11	3	1	1	6	6	*****
North Dakota			*****			4	
South Dakota	3				3	<del>-</del>	
5. Atlantic	135	42	23	4	66	46	2
Delaware District of Columbia	1		****	******	1	2	
Florida	51				51	11	****
Georgia	14	10	1	2	1	5	
Maryland	15	10	4	1		2	<del></del>
North Carolina	15	10	4	******	1	9	<del></del>
South Carolina	9	5	4			3	
Virginia	18	3	8		7	13	
West Virginia	11	4	2	1	4	1	
E.S. Central	31	13	10	2	6	6	
Alabama Kentucky	11 8	4 3	5 1	1	2 3	2	
Mississippi	3	1	1	1	<del></del>	3	
Tennessee	9	5	à	·	1	1	-
W.S. Central	70	25	20	1	24	76	
Arkansas	12	5	5		2	4	
Louisiana	16	****		NAMES OF THE PERSON OF THE PER	16		
Oklahoma	12	7	4	1		4	
Texas	30	13	11	******	6	68	
Mountain	55	32	17		6	11	
Arizona	16	7	5		4	***	
Colorado	9 7	5	4 1			7	*****
Idaho Montana	4	6	4	******		2	*****
Nevada	5	3	1		1		
New Mexico	3	2			i	1	
Utah	11	9	2	****		****	
Wyoming		****		******		1	Mandan
Pacific	169	33	18	1	117	53	****
Alaska	2	-			2	1	
California	110	1	*****		110	43	
Hawaii	4 31	1 22	6	1	2 3	3 4	
Oregon Washington	22	10	12		3	2	name a
Territorles American Samoa							
C.N.M.I.	-			*******		NAME OF TAXABLE PARTY.	MARKE
Guam				******		3	MARINA.
Puerto Rico				****		4	
U.S. Virgin Islands		Account.	*******				

N: No reportable U: Unavailable —: No reported cases C.N.M.L: Commonwealth of Northern Mariana Islands.

\* No cases of diphtheria; eastern equine encephalitis virus disease, nonneuroinvasive; poliomyelitis, paralytic; poliovirus infection, nonparalytic; rubella, congenital syndrome; severe acute respiratory syndrome-associated coronavirus disease; smallpox; vancomycin-resistant Staphylococcus aureus; western equine encephalitis virus disease, neuroinvasive and nonneuroinvasive; yellow fever; and viral hemorrhagic fevers were reported in the United States during 2011. Data on hepatitis B virus, perinatal infection, chronic hepatitis B and hepatitis C virus infection (past or present) are not included because they are undergoing data quality review. Data on human immunodeficiency virus (HIV) infections are not included because HIV infection reporting has been implemented on different dates and using different methods than for acquired immunodeficiency syndrome (AIDS) case reporting.

TABLE 2. (Continued) Reported cases of notifiable diseases,\* by geographic division and area --- United States, 2011

					Q fever		Rai	abies
Area	Pertussis	Plague	Psittacosis	Total	Acute	Chronic	Animal	Human
Jnited States	18,719	3	2	134	110	24	4,357	6
ew England	870			2	1	1	344	2
Connecticut	68		N			<u>,</u>	195	-
Maine	205			2	1	1	66	_
Massachusetts	271						-	2
New Hampshire	170				N	N	25	-
		*****	******	*****				
Rhode Island	62						27	
Vermont	94	*****	-		N	Ŋ	31	
fid. Atlantic	2,305		1	14	11	3	835	2
New Jersey	312			6	6			1
New York (Upstate)	928			5	2	3	370	ī
New York City	323	*****		1	1		13	~~~
Pennsylvania	742		1	2	2		452	
.N. Central	4,526		1	20	16	4	195	
Illinois	1,509			4	4		51	
Indiana	367			1	1		28	Monte
Michigan	691		ī	10	8	2	65	
Ohio	767			10	1	Z.	51	******
Wisconsin	1,192			4	2	2	N	
V.N. Central	1,636	-		5	3	2	197	
lowa	232			more	N	N	25	*****
Kansas	145	_	****				31	
Minnesota	658			1	1	****	56	
Missouri	438			1	·	1	29	
Nebraska	56			2	1	1	33	*****
North Dakota	70	*****		********			23	*******
South Dakota	37			1	1			
		aurana	mam.					
Atlantic	1,506			18	15	3	1,147	1
Delaware	29					****	Market .	Auma
District of Columbia	9				N	N	was.	
Florida	312			3	3		120	
Georgia	179			2	2		****	
Maryland	123			2	2	****	305	
North Carolina	198			5	5		505	
South Carolina				2	1			
	156		nme			1	N	1
Virginia	399			3	1	2	618	
West Virginia	101	*****	Referen	1	1	-	104	
.S. Central	481			2	-	2	162	
Alabama	143			1		1	83	
Kentucky	179			1		1	16	
Mississippi	49						-	
Tennessee	110	****	-	******	*****	****	63	-
/.S. Central	1,140			27	24	3	1,144	******
	80				5		60	
Arkansas				5				
Louisiana	31						6	
Oklahoma	68	-		3	3		60	vew.
Texas	961	****	N	19	16	3	1,018	*******
iountain	2,574	2	anneau.	21	18	3	75	unique.
Arizona	867			2	1	1	N	
Colorado	416			3	2	1		
Idaho	192						6	
Montana	134	*****		15	14	1	Ň	
								· ·
Nevada	34			****	****		17	*****
New Mexico	273	2				Property Control of the Control of t	19	******
Utah	645		•			******	7	*****
Wyoming	13	_	****	1	1		26	*****
cific	3,681	1	MARTINE .	25	22	3	258	1
Alaska	27	****	-			*****	14	
California	2,319			16	16		216	1
Hawaii	59				<del></del>			
	314			1		1	17	
Oregon		1						
Washington	962	buret		8	6	2	11	
erritories								
			N		NI.	K)	N	<b>k</b> 1
American Samoa	LABORANI		N		N	N	N	N
C.N.M.I.		****		*****	****		*******	***************************************
Guam	7					N		
Puerto Rico	8		N				47	
J.S. Virgin Islands								

N: Not reportable U: Unavailable —: No reported cases C.N.M.I.; Commonwealth of Northern Mariana Islands.

<sup>\*</sup> No cases of diphtheria; eastern equine encephalitis virus disease, nonneuroinvasive; poliomyelitis, paralytic; poliovirus infection, nonparalytic; rubella, congenital syndrome; severe acute respiratory syndrome-associated coronavirus disease; smallpox; vancomycin-resistant Staphylococcus aureus; western equine encephalitis virus disease, neuroinvasive and nonneuroinvasive; yellow fever; and viral hemorrhagic fevers were reported in the United States during 2011. Data on hepatitis B virus, perinatal infection, chronic hepatitis B and hepatitis C virus infection (past or present) are not included because they are undergoing data quality review. Data on human immunodeficiency virus (HIV) infections are not included because HIV infection reporting has been implemented on different dates and using different methods than for acquired immunodeficiency syndrome (AIDS) case reporting.