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National Action Week for the Bone and Joint Decade — October 12–20, 2006

National Action Week for the Bone and Joint Decade, a global, multidisciplinary initiative promoting the care of persons with bone and joint disorders, is being observed October 12–20. This initiative focuses on improving quality of life and advancing the understanding and treatment of musculoskeletal conditions through research, prevention, and education. CDC, the National Institutes of Health, the World Health Organization, and the United Nations are among the governmental and nongovernmental organizations supporting this initiative. In 2002, the United States officially proclaimed the years 2002–2011 as the National Bone and Joint Decade.

Bone and joint disorders are the leading causes of disability in the United States (1) and impose substantial burdens on the health-care system and society (2). The United States Bone and Joint Decade organization is committed to raising awareness of the growing burden of musculoskeletal conditions and promoting their prevention, advancing research, and improving diagnosis and treatment. This week, two education programs will be inaugurated by the U.S. group: Straighten Up America and Joint Health and Arthritis. Additional information regarding the United States Bone and Joint Decade is available at http://www.usbjd.org/rd/?naw. Information about activities worldwide is available at http://www.boneandjointdecade.org.

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Prevalence of Doctor-Diagnosed Arthritis and Arthritis-Attributable Activity Limitation — United States, 2003–2005

Arthritis is highly prevalent among U.S. adults, the leading cause of disability (1), and associated with substantial activity limitation, work disability, reduced quality of life, and high health-care costs (2-4). As the population ages, arthritis is expected to affect an estimated 67 million adults in the United States by 2030 (5). This report updates estimates of the national prevalence of doctor-diagnosed arthritis and arthritisattributable activity limitation in the adult U.S. population, using data from the National Health Interview Survey (NHIS) for 2003-2005. The findings indicated that an estimated 21.6% of the adult U.S. population (46.4 million persons) had doctor-diagnosed arthritis, and 8.3% (17.4 million) had arthritis-attributable activity limitations. Public and private health agencies should promote measures to increase the availability of evidence-based arthritis prevention and management interventions.

INSIDE



Recommended Adult Immunization Schedule — United States, October 2006–September 2007

- 1093 Update on Vaccine-Derived Polioviruses
- 1097 West Nile Virus Activity United States, January 1-October 10, 2006
- 1098 Botulism Associated with Commercial Carrot Juice Georgia and Florida, September 2006
- 1100 Notice to Readers
- 1101 QuickStats

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NHIS is an annual, household-based survey of a representative sample of the U.S. civilian, noninstitutionalized population, using in-person interviews. This study used the sample adult core component of the NHIS survey, which collects information on adults aged ≥18 years residing in selected households. In 2003, 2004, and 2005, the sample sizes were 30,852, 31,326, and 31,428, respectively, for the adult core component, and the final response rates were 74.2%, 72.5%, and 69.0%, respectively. Respondents were defined as having doctor-diagnosed arthritis if they answered "yes" to the question, "Have you ever been told by a doctor or other health professional that you have some form of arthritis, rheumatoid arthritis, gout, lupus, or fibromyalgia?" Those who answered "yes" were asked, "Are you limited in any way in any of your usual activities because of arthritis or joint symptoms?" Persons responding "yes" to both questions were defined as having an arthritis-attributable activity limitation.

For this study, prevalence estimates are presented overall and by sex, age group, race/ethnicity, education level, body mass index (BMI)* category, and physical activity level. Physical activity level of respondents was determined from six questions that asked about frequency and duration of participation in leisure-time activities of moderate and vigorous intensity; those reporting no participation in such activities were classified as inactive, and all others as active. Estimates were calculated by using combined data from 2003-2005 and applying an annual average weighting; 95% confidence intervals (CIs) were calculated using sample design factors and statistical software to account for the multistage probability sample. To facilitate comparisons between demographic subgroups, estimates were age adjusted to the standard 2000 U.S. population (6). All differences noted in this report are statistically significant (p<0.05) with nonoverlapping 95% CIs.

In unadjusted analyses for 2003–2005 (Table), the prevalence of doctor-diagnosed arthritis among adults was estimated at 21.6%, or 46.4 million persons. Prevalence was higher among women (25.4%) compared with men (17.6%), older age groups (50% for persons aged ≥65 years and 29.3% for persons aged 45–64 years) compared with younger age groups (7.9% for persons aged 18–44 years), and non-Hispanic whites (24.3%) compared with non-Hispanic blacks (19.2%) and Hispanics (11.4%). Prevalence also was higher among those who were obese (31.6%) or overweight (21.7%) compared with those who were normal weight or underweight (16.3%)

^{*}BMI was calculated using self-reported weight and height as follows: weight (kg) / height (m²). Categories were defined as follows: underweight/normal weight, ≤24.9; overweight, 25.0–29.9; and obese, ≥30.0.

TABLE. Unadjusted and age-adjusted* estimates of the prevalence of doctor-diagnosed arthritis and arthritis-attributable activity limitations[†] among adults aged ≥18 years, by selected characteristics — National Health Interview Survey, United States, 2003–2005

	Adult population prevalence									Proport	tion with	la.
	Do	ctor-diagn					ble activity	/ limitation	activ	arunnus-a ity limitati		-
		(46.4 millio					lion persor			doctor-dia		•
	Una	djusted	Age	adjusted	Una	djusted	Age	adjusted	Una	djusted	Age a	djusted
Characteristic	% (95% CI§)	%	(95% CI)	%	(95% CI)	%	(95% CI)	%	(95% CI)	%	(95% CI)
Sex												
Men	17.6	(<u>+</u> 0.5)	18.1	(<u>+</u> 0.5)	6.4	(<u>+</u> 0.3)	6.6	(<u>+</u> 0.4)	36.6	(<u>+</u> 1.4)	35.2	(<u>+</u> 1.6)
Women	25.4	(<u>+</u> 0.6)	24.4	(± 0.5)	10.0	(<u>+</u> 0.3)	9.7	(<u>+</u> 0.3)	40.1	(<u>+</u> 1.0)	37.5	(<u>+</u> 1.3)
Age (yrs)												
18–44	7.9	(<u>+</u> 0.3)		_	2.7	(<u>+</u> 0.2)		_	33.9	(<u>+</u> 1.8)		_
45–64	29.3	(<u>+</u> 0.7)		_	11.3	(<u>+</u> 0.5)		_	38.6	(<u>+</u> 1.4)		_
≥65	50.0	(<u>+</u> 0.9)		_	20.5	(<u>+</u> 0.7)		_	41.5	(<u>+</u> 1.3)		_
Race/Ethnicity												
White, non-Hispanic	24.3	(<u>+</u> 0.5)	22.6	(<u>+</u> 0.4)	8.9	(<u>+</u> 0.3)	8.4	(<u>+</u> 0.3)	37.4	(<u>+</u> 0.9)	35.0	(<u>+</u> 1.2)
Black, non-Hispanic	19.2	(<u>+</u> 0.9)	21.4	(<u>+</u> 0.9)	8.8	(<u>+</u> 0.6)	10.0	(<u>+</u> 0.6)	45.7	(<u>+</u> 2.4)	43.4	(<u>+</u> 3.1)
Hispanic	11.4	(<u>+</u> 0.6)	16.5	(±0.8)	5.0	(<u>+</u> 0.4)	7.3	(<u>+</u> 0.6)	43.8	(<u>+</u> 2.7)	42.3	(<u>+</u> 3.2)
Other non-Hispanic	14.7	(<u>+</u> 1.3)	17.3	(<u>+</u> 0.5)	5.7	(±0.8)	7.0	(<u>+</u> 1.0)	41.5	(<u>+</u> 4.6)	40.0	(<u>+</u> 5.3)
Education												
Did not graduate from high school		(<u>+</u> 1.0)	23.2	\— <i>'</i>	13.6	·— /	11.7	(<u>+</u> 0.6)	50.6	(<u>+</u> 1.6)	49.3	(<u>+</u> 3.0)
High school graduate or more	20.8	(<u>+</u> 0.4)	21.2	(<u>+</u> 0.4)	7.4	(<u>+</u> 0.2)	7.7	(<u>+</u> 0.2)	36.1	(<u>+</u> 0.9)	34.5	(<u>+</u> 1.1)
Body mass index (BMI [¶])												
Underweight/Normal weight	16.3	(<u>+</u> 0.5)	17.4	(<u>+</u> 0.5)	5.5	(± 0.3)	5.9	(<u>+</u> 0.3)	34.3	(<u>+</u> 1.4)	32.4	(<u>+</u> 1.8)
Overweight	21.7	(<u>+</u> 0.6)	20.5	(<u>+</u> 0.5)	7.5	(<u>+</u> 0.3)	7.1	(<u>+</u> 0.3)	35.0	(<u>+</u> 1.3)	33.4	(<u>+</u> 1.8)
Obese	31.6	(<u>+</u> 0.8)	29.3	(<u>+</u> 0.7)	14.4	(<u>+</u> 0.6)	13.8	(<u>+</u> 0.6)	46.4	(<u>+</u> 1.6)	43.2	(<u>+</u> 1.9)
Physical activity level												
Inactive	25.0	(<u>+</u> 0.6)	22.3	(<u>+</u> 0.5)	13.2	(± 0.5)	11.7	(<u>+</u> 0.4)	52.6	(±1.3)	49.8	(<u>+</u> 2.1)
Active	19.5	(<u>+</u> 0.5)	20.8	(<u>+</u> 0.5)	6.1	(<u>+</u> 0.3)	6.6	(<u>+</u> 0.3)	31.3	(<u>+</u> 1.0)	29.9	(<u>+</u> 1.2)
Total	21.6	(<u>+</u> 0.4)	21.5	(<u>+</u> 0.4)	8.3	(<u>+</u> 0.2)	8.3	(<u>+</u> 0.2)	38.8	(<u>+</u> 0.8)	36.6	(<u>+</u> 1.0)

and among those who were physically inactive (25.0%) compared with those who were physically active (19.5%). After adjustment for age, all of these differences (except among age groups) were slightly attenuated but remained significant, with the exception of differences between non-Hispanics whites (22.6%) and non-Hispanic blacks (21.4%).

Unadjusted analyses for arthritis-attributable activity limitation among adults indicated an estimated overall prevalence of 8.3%, or 17.4 million persons, with differences among groups that were similar to those for doctor-diagnosed arthritis prevalence. The exception was a similar prevalence for non-Hispanic blacks (8.8%) and non-Hispanic whites (8.9%). Age-adjusted analyses identified differences among groups that were similar to the unadjusted figures, except that prevalence among non-Hispanic blacks (10.0%) significantly exceeded that for non-Hispanic whites (8.4%).

In unadjusted analyses of all adults reporting arthritis, 38.8% reported arthritis-attributable activity limitation (Table). Proportions were significantly higher among women (40.1%) compared with men (36.6%) and among non-Hispanic blacks (45.7%) and Hispanics (43.8%) compared with non-Hispanic whites (37.4%). Persons with arthritis and activity limitations also were more likely to have less than a high school education (50.6% versus 36.1%) or to be obese (46.4% versus 34.3%) underweight/normal weight) or physically inactive (52.6%) versus 31.3%). Age-adjusted analyses eliminated the significant difference between men and women, but did not otherwise change the results.

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^{*}Adjusted to the projected 2000 population aged ≥18 years by three age groups: 18–44 years, 45–64 years, and ≥65 years.

† Doctor-diagnosed arthritis was defined as those answering "yes" to the question, "Have you ever been told by a doctor or other health professional that you have some form of arthritis, rheumatoid arthritis, gout, lupus, or fibromyalgia?" Those who answered "yes" were asked, "Are you limited in any way in any of your usual activities because of arthritis or joint symptoms?" Persons responding "yes" to both questions were defined as having an arthritis attributable s activity limitation.

Confidence interval

[¶]BMI = weight (kg) / height (m²). Underweight/normal weight, ≤24.9; overweight, 25.0–29.9; and obese, ≥30.0.

Editorial Note: The findings in this report indicate that 21.6% (46.4 million) of U.S. adults reported doctor-diagnosed arthritis, and 8.3% (17.4 million) reported arthritisattributable activity limitation during 2003–2005. This represents an increase from 2002, when an estimated 20.8% (42.7 million) reported doctor-diagnosed arthritis and 7.8% (16.0 million) reported arthritis-attributable activity limitation (2). The increase in both the prevalence of doctor-diagnosed arthritis and arthritis-attributable activity limitation is consistent with future projections, largely based on the aging of the population (5). However, in 2003, the NHIS transitioned to a weighting structure based on the 2000 U.S. Census population; therefore, interpretation of this increased prevalence should be made with caution. Additional years of data are needed to determine whether these growth trends will be lasting.

Disparities exist with regard to arthritis and activity limitations. Women, older adults, persons with little education, or those who are obese, overweight, or physically inactive are more likely affected. In unadjusted analyses, doctor-diagnosed arthritis was less prevalent among non-Hispanic blacks and Hispanics than among non-Hisipanic whites; however, both groups reported greater proportions of persons with arthritisattributable activity limitation.

In contrast to previous estimates of arthritis prevalence based on 1 year of data, prevalences for a 3-year period were used to reduce the year-to-year fluctuation that can result from smaller sample sizes from a single year. This approach might provide more reliable estimates, especially for smaller groups such as certain racial/ethnic populations and older adults.

The findings in this report are subject to at least three limitations. First, doctor-diagnosed arthritis was self-reported and not confirmed by a health-care professional, although selfreport of arthritis has been determined valid for surveillance purposes (7). Second, the cross-sectional study design does permit determining the temporal sequence of arthritis onset and selected characteristics (e.g., obesity or physical inactivity). However, other studies have identified excess body weight as a risk factor for incident osteoarthritis, the most common type of arthritis, and physical activity has been determined to prevent or delay onset of functional limitation and disability among adults with osteoarthritis (8). Finally, certain factors that might contribute to differences in arthritis prevalence (e.g., history of joint injury or comorbid conditions such as cardiovascular disease, diabetes, or depression) were not analyzed.

Population-based national surveillance of arthritis prevalence and associated effects such as arthritis-attributable activity limitation are important to identify groups at greatest risk, target interventions, and measure progress toward achieving national health objectives (9). Currently, the CDC Arthritis Program is focusing on expanding the availability of evidence-based physical-activity and self-management interventions proven to reduce pain and improve function among adults with arthritis. Such interventions include those related to safe physical activity for persons with arthritis (e.g., Arthritis Foundation's Exercise Program, Arthritis Foundation's Aquatics Program, and EnhanceFitness) and self-management education (e.g., Arthritis Foundation's Self-Help Course and the Chronic Disease Self-Management Program). In addition, the CDC Arthritis Program is working with 35 state health department programs and various local chapters of the Arthritis Foundation to disseminate a health communications campaign designed to promote greater physical activity among adults with arthritis. The campaign, "Physical Activity. The Arthritis Pain Reliever," was developed to target an audience of low-income men and women aged ≥45 years with arthritis. A similar campaign targeted to Spanish-speaking adults, "Buenos Dias Artritis," is being developed and tested.† Further research is needed to investigate possible underlying reasons for the differences among groups in arthritis prevalence and activity limitation and to develop more targeted solutions to improve the quality of life for all adults with arthritis, particularly among those most affected.

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[†]Additional information on arthritis programs is available at http://www.cdc.gov/

Update on Vaccine-Derived Polioviruses

In 1988, the World Health Assembly resolved to eradicate polio worldwide. The Global Polio Eradication Initiative (PEI) of the World Health Organization (WHO) has led to a decline in global polio incidence, from an estimated 350,000 cases in 1988 to fewer than 2,000 reported cases in 2005, and polio remains endemic to only four countries (Afghanistan, India, Nigeria, and Pakistan) (1). However, two additional obstacles to global eradication involve vaccine-derived polioviruses (VDPVs). Polio outbreaks continue to be associated with circulating vaccine-derived polioviruses (cVDPVs) in areas with low oral poliovirus vaccine (OPV) coverage. In addition, long-term excretion of neurovirulent immunodeficiency-associated vaccine-derived polioviruses (iVDPVs) can lead to poliovirus spread to contacts. Overcoming these obstacles is challenging. High rates of OPV coverage will prevent all poliovirus spread, including spread of VDPVs, but will not prevent establishment of prolonged VDPV infections in certain persons with B-cell immunodeficiencies (i.e., having defects in antibody production). Inevitable gaps in vaccination coverage will give rise to cVDPVs as long as OPV use continues. This report updates a previous report on VDPVs and describes the potential implications of VDPVs in the final stages of global polio eradication (2). The findings underscore the critical need to strengthen strategies to prevent emergence of VDPVs and to stop all OPV use once wild polioviruses (WPVs) are eradicated (2–5).

Biologic Properties of VDPVs

The critical biologic properties of VDPVs are their capacity to cause paralytic polio in humans and their potential or demonstrated capacity for sustained circulation. VDPVs have lost key attenuating mutations and resemble WPVs biologically (2). All known cVDPVs (except those from China) (Table 1), but no iVDPVs, are recombinants with nonstructural protein sequences derived from species C enteroviruses, a property associated with poliovirus circulation (2). Most VDPVs are antigenic variants of the Sabin strains, but antigenic evolution appears to be faster in iVDPVs than in cVDPVs. Unlike cVDPV isolates, iVDPV isolates commonly contain mixed VDPV populations. These biologic distinctions (and the differing conditions favoring iVDPV and cVDPV emergence) have helped in recognition of the likely origins of many ambiguous VDPVs (aVDPVs) (2).

Categories of VDPVs

VDPVs differ from the majority of vaccine-related isolates by having genetic properties consistent with prolonged replication or transmission. Because poliovirus genomes evolve at a rate of approximately 1% per year, vaccine-related isolates

TABLE 1. Outbreaks of circulating vaccine-derived polioviruses (cVDPVs) — worldwide, 1988–2006

Location	Years	Reported no. of polio cases	Serotype	No. of clinical isolates (% VP1 divergence from Sabin strain)	Recombination with species C enteroviruses*	Estimated duration of circulation	Routine vaccination coverage with 3 doses of oral polio vaccine (OPV)
Egypt [†]	1988-1993	30	2	30 (4.0-7.0)	Yes	10.0 yrs	Reported high
Haiti [§]	2000-2001	8	1	8 (1.9-2.6)	Yes	2.5 yrs	<30% nationwide
Dominican Republic§	2000-2001	13	1	13 (1.9-2.6)	Yes	>0.5 yr	<30% around most cases
Philippines	2001	3	1	4 (3.1–3.5)	Yes	2.5 yrs	OPV shortage previous 2 yrs
Madagascar [¶] **	2002	4	2	6 (2.5-3.0)	Yes	2.5 yrs	<50% nationwide
China ^{††} §§	2004	2	1	4 (1.0-1.2)	No	1.0 yr	<50% around cases
Madagascar ^{§§}	2005	3	2	3 (1.1–1.8)	Yes	1.0 yr	<50% nationwide
Indonesia ^{§§} ¶¶	2005	46	1	46 (1.1–3.0)	Yes	2.0 yrs	<40% in Madura
Cambodia ^{§§}	2005–2006	3	3	3 (1.9–2.4)	Yes	>1.0 yr	<50% around cases

^{*} All cVDPV isolates except those from China were vaccine/nonvaccine recombinants.

[†] Inferred retrospectively from sequence studies of stored isolates. Not investigated in the field.

[§] Common outbreak. In 2000, cVDPV spread from Haiti to the Dominican Republic.

[¶] In 2001, an unrelated type 2 ambiguous VDPV isolate (1% VP1 divergence) was obtained from a patient with acute flaccid paralysis in a separate community in Madagascar.

^{**} VDPVs were isolated from four polio patients (March–April 2002) and from two healthy children (from a stool survey of 316 healthy children conducted in June 2002 in the outbreak area).

^{††} Localized outbreak in Guizhou province.

^{§§} New cases reported since publication of previous report (2).

[🅅] Localized outbreak on Madura Island off coast of Java.

that differ from the corresponding OPV strain by more than 1% of nucleotide positions (usually determined by sequencing the genomic region encoding the major viral surface protein, VP1) are estimated to have replicated for at least 1 year after administration of an OPV dose, substantially longer than the normal period of vaccine virus replication of 4-6 weeks. Poliovirus isolates are divided into three categories, identified by the extent of VP1 nucleotide sequence divergence from the corresponding Sabin OPV strain: 1) OPV-like viruses (<1% divergent), 2) VDPVs (1%-15% divergent), and 3) WPVs (>15% divergent) (2). VDPVs are further divided into 1) iVDPVs isolated from persons with primary immunodeficiencies who have prolonged VDPV infections after exposure to OPV, 2) cVDPVs that emerge in communities with inadequate OPV coverage, and 3) aVDPVs, which are clinical isolates from persons with no known immunodeficiency and environmental isolates whose ultimate source has not been identified (2).

iVDPVs

A small proportion of immunodeficient persons exposed to OPV have excreted iVDPV over prolonged periods (>6 months). WHO maintains an iVDPV registry; since the introduction of OPV in 1961-1962, only 30 persons excreting iVDPVs have been identified. Persons with primary B-cell immunodeficiencies, but not persons with T-cell immunodeficiencies (e.g., from human immunodeficiency virus infection), are at risk for iVDPV infections (6). Approximately 70% of iVDPV infections have spontaneously ceased within 3 years of exposure to OPV, or the patients have died from complications of their immunodeficiency. Five persons excreted virus for 3-8 years, and in three persons, the duration of excretion exceeded 9 years (Table 2). Eighteen (60%) documented iVDPV infections were associated with type 2 poliovirus infection, eight (27%) with type 1, one (3%) with type 3, and three (9%) with mixed infections (Table 2, Figure). The first reports of iVDPVs came from high-income countries (e.g., the United States, countries of Western Europe, and Japan) but recent reports of iVDPVs include middleincome countries (Table 2). No iVDPVs have been reported from low-income countries, where survival rates for persons with B-cell immunodeficiencies are low (7). Exposure usually is from receipt of OPV, but three of the known iVDPV infections occurred in unimmunized persons (Table 2). Strategies for resolving iVDPV infections are needed, both because of the risk for paralytic disease to infected persons and the risk for transmission to the wider community. No antiviral drug that has been shown to resolve iVDPV infections is currently available. However, new antiviral drugs broadly effective against VDPVs are under development (8).

cVDPVs

VDPVs do not circulate when high vaccination coverage leads to high population immunity. However, low vaccination coverage increases the proportion of nonimmune persons in a population; this increases the potential for VDPVs to circulate. Under circumstances of low vaccination coverage, cVDPVs have produced several localized polio outbreaks. Eight independent outbreaks (i.e., two or more polio cases) in eight countries have been associated with cVDPVs (Table 1, Figure). The largest documented outbreak (46 polio cases) occurred on the Indonesian island of Madura. Genetic studies on stored isolates suggest that a type 2 cVDPV circulated endemically in Egypt for 10 years (approximately from 1983 to 1993) and probably caused more polio cases than were reported (2). Outbreaks of cVDPVs have been associated with all three poliovirus serotypes. Two independent type 2 cVDPV outbreaks occurred in Madagascar in 2002 and 2005 (2), possibly signaling a higher potential for the emergence of type 2 cVDPVs.

aVDPVs

aVDPVs are VDPV isolates that cannot be clearly assigned to either of the other two well-defined categories. They have been isolated from paralyzed persons with no evidence of additional paralyzed VDPV-infected persons among household or community contacts. Highly divergent (>12% VP1 nucleotide divergence) aVDPVs also have been isolated from sewage in Estonia, Israel, and Slovakia. The sewage isolates have similar genetic and antigenic properties as iVDPVs, but measures to identify the infected persons have been unsuccessful. In 1966, aVDPVs were found in Belarus after local suspension of OPV use; in 1999, they were found in Russia among children in orphanages (2). A growing number of aVDPVs having VP1 sequence divergence slightly above 1% have been found by the Global Polio Laboratory Network.

Limited person-to-person transmission for certain aVDPVs has occurred. In 2005, a type 3 aVDPV was isolated from one polio patient and seven nonparalyzed contacts in Madagascar. Similarly, a type 1 VDPV was isolated from one patient and seven contacts in Romania in 2002, a type 2 VDPV was isolated from one patient and two contacts in Laos in 2004 (2), a type 1 VDPV was isolated from an unimmunized severe combined immunodeficiency (SCID) patient and four community members in rural Minnesota in 2005 (9), and a type 1 VDPV was isolated from one patient and six contacts in Myanmar in 2006. Other aVDPVs with genetic properties resembling those of cVDPVs were found in Peru in 1983, in Pakistan in 2000, and in Nigeria in 2002 and 2006 (2).

TABLE 2. Selected characteristics of persons excreting immunodeficiency-associated vaccine-derived polioviruses (iVDPVs) — worldwide, 1962–2006

	Year				Maximum % VP1 divergence from Sabin	Estimated interval between last oral polio vaccine (OPV) dose and detection of iVDPV		Estimated duration of virus excretion
Location	detected			Serotype		infection (yrs)*	Outcome	(yrs)
United Kingdom	1962	Hypogammaglobulinemia	No	1	Unknown	0†	Died	2.7
United Kingdom	1962	Hypogammaglobulinemia	No	3	2.3	0†	Died	1.8
Japan	1977	X-linked agammaglobulinemia	Yes	2	Unknown	1.5	Died	3.4
United States	1980	Agammaglobulinemia	Yes	2	Unknown	1	Died	1§
United States	1981	Common variable immunodeficiency	Yes	1	10.0	7.1	Died	7.6
United States	1986	X-linked agammaglobulinemia	Yes	2	2.0	0.4	Survived	0.4
United States [¶]	1986	Common variable immunodeficiency	Yes	1	5.4	4.7	Survived	9.6
	1992	_	_	2	11.8	_		_
United Kingdom	1987	Common variable immunodeficiency	No	2	4.1	4	Survived	3.6
United States	1989	Agammaglobulinemia	Yes	1	1.1	0.3	NA**	Unknown
Germany	1990	Common variable immunodeficiency	Yes	1	8.3	4	Survived	9.5
United States	1990	Severe combined immunodeficiency	Yes	2	1.9	0.5	Died	0.8
United States	1991	Common variable immunodeficiency	Yes	2	1.4	0.4	Survived	0.6
Iran	1995	Antibody deficiency	Yes	2	2.2	Unimmunized	Died	1.5
United Kingdom	1995	Common variable immunodeficiency	No	2	12.9	15.7	Survived	20
United States	1995	Severe combined immunodeficiency	Yes	2	2.1	0.3	Died	3.7
Argentina	1998	X-linked agammaglobulinemia	Yes	1	2.8	Unimmunized	Survived	2
Germany	2000	Antibody deficiency	Yes	1	3.5	NA	Survived	1.5
Taiwan	2001	Common variable immunodeficiency	Yes	1	3.5	1.6	Survived	3
United Kingdom	2002	Common variable immunodeficiency	No	2	3.3	NA	Survived	3.3
United Kingdom	2002	Immunodeficiency-centromeric						
		instability-facial abnormalities syndron		2	2.5	NA	Survived	2.5
Kazakhstan	2002	Hypogammaglobulinemia	Yes	2	2.3	NA	Died	2
Kuwait	2002	Major histocompatibility complex class	I					
		molecule deficiency	No	2	2.0	0.9	Died	0.4
Peru	2003	Agammaglobulinemia	Yes	2	1.2	0.6	Survived	<1.0
Thailand	2003	Hypogammaglobulinemia	Yes	2	2.2	0.3	NA	< 0.5
China ^{††}	2005	X-linked agammaglobulinemia	Yes	2+3	2.7	0.6	NA	NA
Iran ^{††}	2005	Common variable immunodeficiency	Yes	2	1.4	0.7	Died	0.7
Morocco ^{††}	2005	Severe combined immunodeficiency	Yes	2	2.5	1	Died	1
Syria ^{††}	2005	Hypogammaglobulinemia	Yes	2	1.3	<0.1	Survived	<0.1
United States††	2005	Severe combined immunodeficiency	No	1	2.3	Unimmunized	Survived	<0.5 ^{§§}
Tunisia ^{††}	2006	Severe combined immunodeficiency	No	2	2.0	NA	Survived	0.1

^{*} Several estimates are approximate because of no follow-up sampling, long sampling intervals, or uncertain date of associated OPV exposure. Because criteria for estimates varied in different studies, certain estimates were rounded off to the nearest integer.

Risk Factors for VDPV Emergence

The key factors favoring cVDPV emergence and spread are the same as for WPV circulation: low OPV coverage, poor sanitation, high population densities, and (usually) tropical conditions. In all but the remaining polio-endemic areas, immunity to polio is no longer acquired from natural infection; immunization is the only current means to prevent the spread of emerging VDPVs or imported WPVs (3).

Although OPV is not recommended for immunodeficient patients, it is often inadvertently administered because certain primary immunodeficiencies (e.g., common variable immunodeficiency [CVID]) develop later in life. Certain persons with CVID who excrete iVDPVs had onset of polio several years after the implicated OPV dose was administered, and three have demonstrated no signs of paralysis. Survival of patients with primary immunodeficiencies can be extended

[†] Immunodeficient children were administered OPV, and virus excretion was monitored.

[§] Neural isolate obtained at autopsy, approximately 4.3 years after last OPV dose.

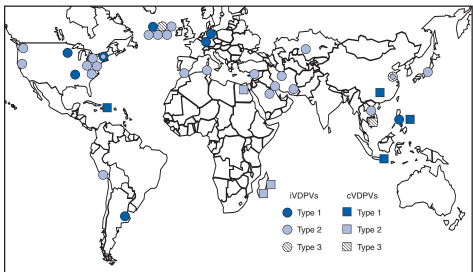
Two different iVDPVs were isolated from the same patient (a type 1 in 1986 and a type 2 in 1992).

^{**} Information not available.

^{††} New cases reported since publication of previous report (2).

^{§§} Excretion stopped after bone marrow transplant.

FIGURE. Locations of persons excreting immunodeficiency-associated vaccinederived polioviruses (iVDPVs), 1962–2006, and polio outbreaks associated with circulating vaccine-derived polioviruses (cVDPVs), 1988–2006



in upper- and middle-income countries by intravenous immunoglobulin therapy; however, for patients in low-income countries, such therapy often is too expensive and difficult to obtain (7).

Global VDPV Surveillance

Since the cVDPV outbreak in Haiti and the Dominican Republic in 2000-2001 (Figure, Table 1), all polioviruses isolated in the WHO Global Poliovirus Laboratory Network from patients with acute flaccid paralysis have been characterized by one molecular method, to identify polioviruses by their genetic properties (usually using the polymerase chain reaction), and one antigenic method, to detect antigenic differences from the OPV strains (using either an enzyme-linked immunosorbent assay [ELISA] or panels of specific neutralizing monoclonal antibodies) (10). Isolates found to be genetically related to an OPV strain but with antigenic differences are possible VDPVs. VP1 sequencing is routinely performed on all possible VDPV and WPV isolates. Approximately 12,000 isolates from all WHO regions have been routinely screened for VDPVs since 2001 (10). Temporal or geographic clustering of vaccine-related isolates of the same serotype has prompted the detection and investigation of cVDPV outbreaks in eight countries (Table 1).

Reported by: WHO Global Poliovirus Laboratory Network. Immunization, Vaccines and Biologicals Dept, WHO, Geneva, Switzerland. Div of Viral Diseases and Global Immunization Div, National Center for Immunization and Respiratory Diseases (proposed), CDC.

Editorial Note: VDPVs will continue to emerge as long as OPV is used. Intensified surveillance has indicated that cVDPVs can emerge repeatedly under conditions of low OPV coverage (e.g., Madagascar). VDPVs also can be found in developed countries with no paralytic cases (e.g., Estonia, Israel, and Slovakia) and can circulate in isolated pockets of unimmunized persons in countries with overall high rates of vaccination coverage (e.g., China and the United States). Although iVDPVs can emerge in middle-income developing countries, cVDPVs have not been found in some areas of high biologic risk, such as in northern India, presumably because of the current high rates of OPV coverage.

Occurrences of VDPVs, including cVDPV-related outbreaks, are rare events, and all recent outbreaks of cVDPVs have been rapidly interrupted using OPV campaigns. The recent increase in the detection of VDPVs is probably primarily attributable to intensified surveillance and improved laboratory methods. Enhanced surveillance for VDPVs has allowed for better understanding of the risks associated with the different types of VDPVs. Areas with continued use of OPV but lacking optimal coverage (e.g., Indonesia in 2005) are at increased risk for cVDPV emergence. The importance of detecting aVDPVs with limited VP1 divergence is not clear; the presence of aVDPVs in certain settings might not have any public health consequences, whereas aVDPVs found elsewhere might signal conditions favoring the emergence of a cVDPV.

Under certain circumstances, OPV viruses regain both neurovirulence and the capacity to circulate and cause outbreaks and therefore are of concern to the PEI. After global eradication of WPVs, the continued use of OPV would continually generate cVDPVs and could eventually pose a challenge to the goal of stopping all poliovirus infections in the human population. The increasing risk of cVDPV emergence in countries with widening immunity gaps and the ongoing risks for vaccine-associated paralytic polio and iVDPVs have prompted an evaluation of the feasibility of orderly cessation of OPV use as soon as possible in the posteradication era (4) while population immunity and surveillance sensitivity are still high (6). Continued development and implementation of a comprehensive strategy to minimize the risks for VDPV emergence in the posteradication era presents a challenge to the PEI and to the public health and scientific communities.

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 June 2005. MMWR 2005;54:958

 –61.

West Nile Virus Activity — United States, January 1–October 10, 2006

This report summarizes West Nile virus (WNV) surveillance data reported to CDC through ArboNET as of 3 a.m. Mountain Daylight Time, October 10, 2006. A total of 41 states and the District of Columbia had reported 3,135 cases of human WNV illness to CDC (Table, Figure). A total of 1,717 (55%) cases for which such data were available occurred in males; median age of patients was 50 years (range: 3 months–99 years). Dates of illness onset ranged from January 6 to September 25; a total of 97 cases were fatal.

A total of 260 presumptive West Nile viremic blood donors (PVDs) have been reported to ArboNET during 2006. Of these, 40 were reported from Nebraska; 27 from Texas; 24 from Utah; 21 from Colorado; 15 from California; 14 from Louisiana; 11 each from North Dakota and South Dakota; 10 each from Iowa and Wisconsin; nine each from Arizona, Mississippi, and Oklahoma; eight from Kansas; six from Idaho; five each from Minnesota and Virginia; four each from Kentucky and Missouri; three each from Illinois, Montana, and Nevada; two from Michigan; and one each from Arkansas, Maryland, New York, Ohio, Oregon, Pennsylvania, and

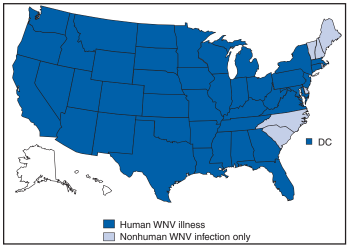
TABLE. Number of human cases of West Nile virus (WNV) illness, by state — United States, 2006*

State N	Neuroinvasive disease [†]	West Nile fever§	Other clinical/ unspecified ¹	Total reported to CDC**	Deaths
Alabama	4	0	1	5	0
Arizona	15	14	16	45	3
Arkansas	21	5	0	26	0
California	65	164	13	242	3
Colorado	54	219	0	273	3
Connecticut	6	2	0	8	1
District of Colu	umbia 0	1	0	1	0
Florida	3	0	0	3	0
Georgia	2	4	1	7	1
Idaho	94	542	6	642	10
Illinois	111	55	23	189	9
Indiana	11	5	12	28	0
Iowa	17	12	0	29	0
Kansas	14	10	0	24	3
Kentucky	5	1	0	6	1
Louisiana	66	49	0	115	0
Maryland	2	1	1	4	0
Massachusett	s 2	1	0	3	0
Michigan	29	2	6	37	3
Minnesota	29	34	Ö	63	3
Mississippi	72	79	0	151	6
Missouri	41	9	1	51	2
Montana	10	19	1	30	0
Nebraska	33	123	0	156	1
Nevada	34	73	14	121	i
New Jersey	2	2	1	5	0
New Mexico	1	2	0	3	0
New York	7	3	1	11	2
North Dakota	20	115	0	135	1
Ohio	27	7	0	34	3
Oklahoma	21	12	1	34	5
Oregon	4	42	8	54 54	0
Pennsylvania	7	1	0	8	2
South Dakota	37	71	0	108	3
Tennessee	7	1	0	8	ა 1
Texas	, 175	81	0	256	23
Utah	175 48	88	0	236 136	23 4
	48 0		2	2	
Virginia	-	0		_	0
Washington	0 1	2	0	2 1	0 0
West Virginia	=	0	0	=	-
Wisconsin	10 14	8	0	18 61	1 2
Wyoming		36	11	61	_
Total	1,121	1,895	119	3,135	97

- * As of October 10, 2006.
- [†] Cases with neurologic manifestations (i.e., West Nile meningitis, West Nile encephalitis, and West Nile myelitis).
- § Cases with no evidence of neuroinvasion.
- ¶ Illnesses for which sufficient clinical information was not provided.
- ** Total number of human cases of WNV illness reported to ArboNET by state and local health departments.

Wyoming. Of the 260 PVDs, three persons (median age: 73 years [range: 26–74 years]) subsequently had neuroinvasive illness, one person aged 41 years had other illness, and 54 persons (median age: 46 years [range: 17–70 years]) had West Nile fever.

FIGURE. Areas reporting West Nile virus (WNV) activity — United States, 2006*



^{*} As of October 10, 2006.

In addition, 2,746 dead corvids and 636 other dead birds with WNV infection have been reported in 41 states and New York City during 2006. WNV infections have been reported in horses in 34 states, in one squirrel in Kansas, and in two unidentified animal species in North Carolina and Wyoming. WNV seroconversions have been reported in 682 sentinel chicken flocks in 12 states (Arizona, Arkansas, California, Florida, Iowa, Montana, Nevada, North Carolina, North Dakota, Pennsylvania, Utah, and Virginia). A total of 10,157 WNV-positive mosquito pools have been reported from 38 states, the District of Columbia, and New York City.

Additional information about national WNV activity is available from CDC at http://www.cdc.gov/ncidod/dvbid/westnile/index.htm and at http://westnilemaps.usgs.gov.

Botulism Associated with Commercial Carrot Juice — Georgia and Florida, September 2006

On October 6, this report was posted as an MMWR Dispatch on the MMWR website (http://www.cdc.gov/mmwr).

On September 8, 2006, the Georgia Division of Public Health (GDPH) and CDC were notified of three suspected cases of foodborne botulism in Washington County, Georgia. On September 25, the Florida Department of Health and CDC were notified of an additional suspected case in Tampa, Florida. This report describes the joint investigation and con-

trol measures undertaken by state and local health departments, CDC, and the Food and Drug Administration (FDA).

On September 8, the three patients from Washington County, Georgia, went to a local hospital with cranial nerve palsies and progressive descending flaccid paralysis resulting in respiratory failure; the patients had shared meals on September 7. On the evening of September 8, physicians suspected foodborne botulism, notified the state health department, and collected clinical specimens for testing at CDC. On the same evening, CDC provided clinical consultation and dispatched botulinum antitoxin, which was administered to each of the patients the following morning. After receiving antitoxin, the patients had no progression of neurologic symptoms, but they remain hospitalized and on ventilators.

On September 9, the Washington County Health Department, Richmond County Health Department, and GDPH launched an investigation. The three patients had consumed several food items during their two meals together on September 7, including juice from a single 1-liter bottle of Bolthouse Farms carrot juice. The bottle had a "best if used by" date of September 18, 2006. Clinical specimens and left-over food and juice were collected and sent to CDC for testing. On September 13, botulinum toxin type A was identified in the serum and stool of all three patients. On September 15, leftover carrot juice recovered from the home of one of the patients also tested positive for botulinum toxin type A.

During September 8-15, FDA, the Georgia Department of Agriculture, the Georgia Hospital Association, and public health officials in all 50 states were notified of the outbreak and the implicated product as information became available. After these notifications, no additional cases of botulism in Georgia were reported to the state and local health departments or to CDC. During this time, FDA launched an investigation of the Bolthouse Farms, Inc., manufacturing plant in Bakersfield, California. FDA and CDC tested other bottles of the implicated brand of carrot juice, including bottles from different lots, and all were negative for botulinum toxin. Because botulinum toxin was found only in the bottle of carrot juice consumed by the three patients, a lapse in refrigeration of the carrot-juice bottle during transport or storage was suspected, which would have allowed for growth of Clostridium botulinum and subsequent production of botulinum toxin. Based on the CDC test results, on September 17, FDA issued a consumer advisory on the importance of keeping carrot juice refrigerated. However, information obtained from patient interviews regarding storage and transport of the carrot juice did not confirm mishandling by the patients.

On September 25, officials at the Florida Department of Health, the Hillsborough County Health Department, and CDC were notified that a patient had been hospitalized in Tampa, Florida, on September 16, with respiratory failure and descending paralysis. On September 28, botulinum toxin type A was identified in the patient's serum. Circulating toxin persisted more than 10 days after illness onset in this completely paralyzed patient, indicating ingestion of a massive toxin dose. Accordingly, the patient was treated with antitoxin, which prevents binding of circulating botulinum toxin to nerve endings. The patient remains hospitalized, paralyzed, and on a ventilator. The Hillsborough County Health Department collected an open, 450-milliliter bottle of Bolthouse Farms carrot juice, which had been found by a family member in the hotel room where the patient had been staying during the month before being hospitalized. The hotel room had no refrigerator. The bottle, which had a "best if used by" date of September 19, 2006, had a different lot number than the bottle associated with the Georgia cases. On September 29, botulinum toxin was identified in carrot juice from the bottle found in the patient's hotel room; the toxin was subsequently identified as botulinum toxin type A. The Hillsborough County Health Department and CDC notified FDA, public health officials in all 50 states, and infection-control practitioners in Hillsborough County about the botulism case and implicated product. The manufacturer provided FDA with bottles of carrot juice from the same lot as the bottle found in the patient's room. FDA tested juice from all of these bottles, and it was negative for botulinum toxin.

C. botulinum spores are found in the environment and can be present naturally in carrot juice and other foods that have not undergone the retort canning process, which involves high temperatures and high pressure. Anaerobic conditions, low acidity (pH>4.6), low salt and sugar concentrations, and temperatures >39°F (>4°C) promote germination of C. botulinum spores and botulinum toxin production. Carrot juice has low acidity, with a natural pH of approximately 6.0; therefore, in the absence of another inhibitor, refrigeration at temperatures <40°F (<4°C) is necessary to prevent germination of C. botulinum spores and production of botulinum toxin. Inhibiting C. botulinum growth in other ways, such as through acidification, can retard its growth in juice that is not properly refrigerated.

Acidification has been used as a solution to previous foodborne botulism outbreaks. In 1985, 36 patients in the United States and Canada were identified with botulism after eating at a restaurant in Vancouver, British Columbia. A casecontrol study implicated commercially produced, chopped garlic in soybean oil stored at room temperature as the source of the outbreak (1). In 1989, a second outbreak of botulism associated with chopped garlic in oil occurred when three

patients in New York were identified with botulism after consuming a meal containing unrefrigerated, commercially produced, chopped garlic in virgin olive oil (2). After these outbreaks, FDA rules were altered to require that garlic-in-oil products contain an acidifying agent such as phosphoric or citric acid.

The carrot juice consumed by these four patients was manufactured by Bolthouse Farms, Inc., and distributed in all 50 states, Mexico, Canada, and Hong Kong with the labels "Bolthouse Farms 100% Carrot Juice," "Earthbound Farm Organic Carrot Juice," and "President's Choice Organics 100% Pure Carrot Juice." Investigations of these cases by state and local health departments and investigations of the manufacturer by FDA are ongoing. On September 29, GDPH and the Georgia Department of Agriculture recommended that Georgia residents not purchase or consume Bolthouse Farms carrot juice. The same day, the FDA warned consumers not to drink Bolthouse Farms carrot juice with "best if used by" dates of November 11, 2006 or earlier (i.e., all bottles produced before the date the warning was issued), and Bolthouse Farms issued a voluntary recall of these products. Additional information regarding the recall is available from the Bolthouse Farms website at http://www.bolthouse.com/ bolthouserecallFAQ.pdf or from FDA (telephone, 888-723-3366).

Suspected botulism cases should be reported immediately to local or state public health officials, who then should call the 24-hour CDC Emergency Operations Center at 770-488-7100; the center will immediately connect them with an on-call botulism specialist. Health-care providers and public health officials are encouraged to inquire specifically about consumption of carrot juice as part of the food history of suspect botulism cases. Additional information on botulism is available at http://www.cdc.gov/ncidod/dbmd/diseaseinfo/botulism_g.htm.

Reported by: C Shuler, DVM, C Drenzek, DVM, S Lance, DVM, PhD, G Gonzalez, MD, J Miller, MSPH, M Tobin-D'Angelo, MD, J Gabel, DVM, C Burnett, MPH, Georgia Div of Public Health. D Atrubin, MPH, Florida Dept of Health. J Sobel, MD, P Juliao, PhD, S Maslanka, PhD, Div of Foodborne, Bacterial, and Mycotic Diseases, National Center for Zoonotic, Vector-Borne, and Enteric Diseases; P Wiersma, MD, A Sheth, MD, EIS officers, CDC.

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Notice to Readers

Availability of Provisional AIDS and HIV/AIDS Data in MMWR Table IV and Pediatric HIV Surveillance Data in MMWR Table I

CDC is upgrading the national HIV/AIDS surveillance data management system. Because of this transition, CDC will not update AIDS or HIV/AIDS surveillance data for display in quarterly *MMWR* Table IV for the last two quarters of 2006. In addition, CDC will not provide monthly updates of HIV infection data for persons aged <13 years in *MMWR* Table I for the remainder of this year. Explanatory footnotes will be included with Tables I and IV during the period when no updates are available.

Erratum: Vol. 55, No. 35

In the MMWR report, "Update: Delayed-Onset Pseudomonas fluorescens Bloodstream Infections After Exposure to Contaminated Heparin Flush — Michigan and South Dakota, 2005–2006," the last line of the first column on page 961 should read, "The patients all had indwelling central venous catheters and received treatment during October 2004–February 2005 at clinics known to have used the contaminated flush."

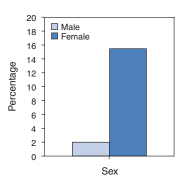
Erratum: Vol. 55, No. 39

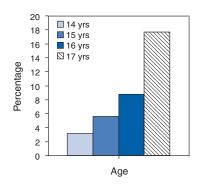
In the MMWR report, "Childhood Influenza Vaccination Coverage — United States, 2004–05 Influenza Season," on page 1061, an error occurred in the fifth sentence of the first paragraph. The sentence should read, "Others recommended to receive influenza vaccination include children aged 5–18 years who have certain high-risk medical conditions, are on chronic aspirin therapy, or who are household contacts of persons at high risk for influenza complications (1)."

QuickStats

FROM THE NATIONAL CENTER FOR HEALTH STATISTICS

Percentage of Teens Aged 14–17 Years Who Used Indoor Tanning Devices
During the Preceding 12 Months, by Sex and Age —
United States, 2005*





*Data are based on household interviews of a sample of the civilian, noninstitutionalized population.

The World Health Organization recommends that no person aged <18 years use a tanning bed because of the associated increased risk for skin cancer. In addition, CDC recommends that school programs to prevent skin cancer advise students to avoid using sunlamps and tanning beds. Nonetheless, in 2005, 8.7% of teens aged 14–17 years used indoor tanning devices. Girls aged 14–17 years were seven times more likely to use these devices than boys in the same age group. The use of indoor tanning devices increased with age from 14 to 17 years.

SOURCES: National Health Interview Survey, 2005. Available at http://www.cdc.gov/nchs/nhis.htm.

World Health Organization. The World Health Organization recommends that no person under 18 should use a sunbed. Available at http://www.who.int/mediacentre/news/notes/2005/np07/en/index.html.

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TABLE I. Provisional cases of infrequently reported notifiable diseases (<1,000 cases reported during the preceding year) — United States, week ending October 7, 2006 (40th Week)*

	Current	Cum	5-year weekly	Total o	cases ren	orted for	rpreviou	s vears	
Disease	week	2006	average [†]	2005	2004	2003	2002	2001	States reporting cases during current week (No.
Anthrax		1	0				2	23	
Botulism:		-	-				_		
foodborne	1	8	0	19	16	20	28	39	FL (1)
infant	1	64	2	90	87	76	69	97	UT (1)
other (wound & unspecified)		42	1	33	30	33	21	19	J. (.)
Brucellosis	_	78	2	122	114	104	125	136	
Chancroid	_	23	0	17	30	54	67	38	
Cholera	_	6	0	8	5	2	2	3	
Cyclosporiasis§		91	1	734	171	75	156	147	
Diphtheria		91		734	17.1	1	1	2	
		_	_			,		2	
Domestic arboviral diseases ^{§,¶} :		20	5	80	112	108	164	128	
California serogroup	_	33			–				
eastern equine	_	6	0	21	6	14	10	9	
Powassan	_	1	_	1	1	_	1	N	
St. Louis	_	3	1	13	12	41	28	79	
western equine	_	_	_	_	_	_	_	_	
Ehrlichiosis§:									
human granulocytic	3	282	8	790	537	362	511	261	NY (1), PA (1), AL (1)
human monocytic	4	281	8	522	338	321	216	142	NY (1), MO (2), TN (1)
human (other & unspecified)	1	130	1	122	59	44	23	6	MO (1)
Haemophilus influenzae,**									
invasive disease (age <5 yrs):									
serotype b	_	8	0	9	19	32	34	_	
nonserotype b	2	67	2	135	135	117	144	_	CT (1), OK (1)
unknown serotype	1	152	2	217	177	227	153	_	ID (1)
Hansen disease§	3	55	1	88	105	95	96	79	NH (1), FL (2)
Hantavirus pulmonary syndrome§	1	25	0	29	24	26	19	8	AZ (1)
Hemolytic uremic syndrome, postdiarrheal§	7	183	5	221	200	178	216	202	OH (1), MN (2), NC (2), AL (1), UT (1)
Hepatitis C viral, acute	11	594	32	771	713	1,102	1,835	3,976	OH (2), MI (3), MO (1), MD (1), OK (1), TX (1),
Tropanio o viiai, aoato		001	OL.		7 10	1,102	1,000	0,070	WA (1), OR (1)
HIV infection, pediatric (age <13 yrs) ^{§,††}	_	52	4	380	436	504	420	543	(.),(.)
Influenza-associated pediatric mortality ^{§,§§,¶¶}	_	40	0	45	_	N	N	N	
Listeriosis	7	498	19	892	753	696	665	613	NY (2), PA (1), OH (1), FL (1), AL (1), WA (1)
Measles	***	43	0	66	37	56	44	116	(1), (2), (1), (1), (1), (1), (1), (1), (1)
Meningococcal disease,††† invasive:		40	O	00	07	50	77	110	
A, C, Y, & W-135	1	170	3	297				_	WA (1)
serogroup B	'	108	2	157					WA (1)
		14	0	27					
other serogroup	9		5		050		070		NV (1) DA (1) MO (1) NO (6)
Mumps		5,791		314	258	231	270	266	NY (1), PA (1), MO (1), NC (6)
Plague	1	12	0	8	3	1	2	2	UT (1)
Poliomyelitis, paralytic	_		0	1	_	_	_	_	
Psittacosis§		17	0	19	12	12	18	25	110 (0) TH (1) TV (1)
Q fever [§]	4	120	1	139	70	71	61	26	MO (2), TN (1), TX (1)
Rabies, human	_	1	0	2	7	2	3	1	
Rubella	1	8	0	11	10	7	18	23	NY (1)
Rubella, congenital syndrome	_	1	_	1	_	1	1	3	
SARS-CoV ^{§,§§}	_	_	_	_	_	8	N	N	
Smallpox§	_	_	_	_	_	_	_	_	
Streptococcal toxic-shock syndrome§	_	78	1	129	132	161	118	77	
Streptococcus pneumoniae,§									
invasive disease (age <5 yrs)	15	787	11	1,257	1,162	845	513	498	NY (3), MI (2), MN (2), NE (2), OK (1), TX (2), CO (3)
Syphilis, congenital (age <1 yr)	5	205	8	361	353	413	412	441	NY (2), IL (1), NC (1), AZ (1)
Tetanus	_	17	Ō	27	34	20	25	37	
Toxic-shock syndrome (other than streptococca	al)§ —	72	2	96	95	133	109	127	
Trichinellosis	_	11	0	19	5	6	14	22	
Tularemia§	_	68	3	154	134	129	90	129	
Typhoid fever	_	220	9	324	322	356	321	368	
Vancomycin-intermediate Staphylococcus aure	us§ —	2	_	2	522	330 N	N	308 N	
Vancomycin-intermediate <i>Staphylococcus aureus</i> §	us ³ —	_	0	3	1	N	N	N	
	_	_					1		
Yellow fever	_	_	_	_	_	_	- 1	_	

N: Not notifiable. Cum: Cumulative year-to-date counts. —: No reported cases.

Incidence data for reporting year 2006 is provisional, whereas data for 2001, 2002, 2003, 2004, and 2005 are finalized.

Calculated by summing the incidence counts for the current week, the two weeks preceding the current week, and the two weeks following the current week, for a total of 5 preceding years. Additional information is available at http://www.cdc.gov/epo/dphsi/phs/files/5yearweeklyaverage.pdf. Not notifiable in all states.

Not notifiable in all states.

Includes both neuroinvasive and non-neuroinvasive. Updated weekly from reports to the Division of Vector-Borne Infectious Diseases, National Center for Zoonotic, Vector-Borne, and Enteric Diseases (proposed) (ArboNET Surveillance).

Data for *H. influenzae* (all ages, all serotypes) are available in Table II.

Updated monthly from reports to the Division of HIV/AIDS Prevention, National Center for HIV/AIDS, Viral Hepatitis, STD, and TB Prevention (proposed). Implementation of HIV reporting influences the number of cases reported. Pediatric HIV data will not be updated monthly for the remainder of this year due to upgrading of the national HIV/AIDS surveillance data management system. Data for HIV/AIDS are available in Table IV quarterly.

Updated weekly from reports to the Influenza Division, National Center for Immunization and Respiratory Diseases (proposed).

Cumulative totals for 2005 and 2006 do not include reports from states where influenza-associated pediatric mortality is not a notifiable condition.

No measles cases were reported for the current week.

Data for meningococcal disease (all serogroups and unknown serogroups) are available in Table II.



Recommended Adult Immunization Schedule — United States, October 2006–September 2007

Weekly

October 13, 2006 / Vol. 55 / No. 40

The Advisory Committee on Immunization Practices (ACIP) annually reviews the recommended Adult Immunization Schedule to ensure that the schedule reflects current recommendations for the licensed vaccines. In June 2006, ACIP approved the Adult Immunization Schedule for October 2006–September 2007. This schedule has also been approved by the American Academy of Family Physicians and the American College of Obstetricians and Gynecologists.

Changes in the Schedule for October 2006–September 2007

The 2006–2007 schedule differs from the previous schedule as follows:

- The broken red line has been deleted on the age-based schedule (Figure 1). Vaccination of persons with specific risk factors is now shown only with purple bars.
- Human papillomavirus (HPV) vaccine has been added to the age-based schedule, with a yellow bar indicating that the vaccine is recommended for women ≤26 years.
- Tetanus, diphtheria, and acellular pertussis (Tdap) vaccine has been added to the age-based schedule, with a hatched yellow bar indicating that Tdap is a one-time, 1-dose recommendation for persons ≤64 years.
- The purple bar for varicella vaccine has been shortened in anticipation of the recommendation for the use of zoster vaccine in persons aged ≥60 years.
- A new column has been added to the medical/other indications schedule (Figure 2) to clarify indications for hepatitis A and B vaccines. The indications "chronic liver disease" and "recipients of clotting factor concentrates" have been removed from the previous schedule's third and fifth columns, respectively, and combined into a new column. The column has a yellow bar for hepatitis A and B vaccines, clarifying that these vaccines are recommended for all persons with these medical indications.
- HPV vaccine has been added to the medical/other indications schedule, with a yellow bar to indicate the vaccine

The Recommended Adult Immunization Schedule has been approved by the Advisory Committee on Immunization Practices, the American College of Obstetricians and Gynecologists, and the American Academy of Family Physicians. The standard MMWR footnote format has been modified for publication of this schedule.

Suggested citation: Centers for Disease Control and Prevention. Recommended Adult Immunization Schedule—United States, October 2006–September 2007. MMWR 2006;55:Q1–Q4.

- is recommended for women aged ≤26 years with all indications except pregnancy.
- Tdap was added to the medical/other indications schedule, with a hatched yellow bar to indicate that Tdap is a one-time, 1-dose recommendation for all indications except pregnancy.
- The tetanus and diphtheria footnote (#1) has been reworded to reflect ACIP recommendations for use of Tdap.
- A footnote (#2) has been added to reflect ACIP recommendations for HPV vaccination for all women aged ≤26 years.
- The measles, mumps, and rubella (MMR) footnote (#3) has been reworded to reflect ACIP recommendations to administer a second dose of mumps vaccine to adults in certain age groups and with certain risk factors.
- The varicella footnote (#4) has been reworded in accordance with ACIP recommendations for administering a routine second dose for all adults without evidence of immunity. The footnote also has been revised to reflect the new definition of immunity to varicella.
- The influenza footnote (#5) has been revised to reflect recent ACIP recommendations to vaccinate close contacts of children aged 0–59 months rather than 0–23 months (1).
- The hepatitis B footnote (#9) has been revised to reflect recommendations to vaccinate any adult seeking protection from hepatitis B virus infection and vaccinate adults in specific settings (e.g., sexually transmitted disease clinics) (2).

The Adult Immunization Schedule is available in English and Spanish at http://www.cdc.gov/nip/recs/adult-schedule.htm. General information about adult vaccinations, including recommendations concerning vaccination of person with HIV and other immunosuppressive conditions, is available from state and local health departments and at http://www.cdc.gov/nip. Vaccine information statements are available at http://www.cdc.gov/nip/publications/vis. ACIP statements for each recommended vaccine and provisional vaccine recommendations can be viewed, downloaded, and printed at http://www.cdc.gov/nip/publications/acip-list.htm. Instructions for reporting adverse events to the Vaccine Adverse Event Reporting System are available at http://www.vaers.hhs.gov or by telephone, 800-822-7967.

References

- CDC. Prevention and control of influenza: recommendations of the Advisory Committee on Immunization Practices (ACIP). MMWR 2006;55(No. RR-10).
- 2. CDC. A comprehensive immunization strategy to eliminate transmission of hepatitis B virus infection in the United States: recommendations of the Advisory Committee on Immunization Practices (ACIP). Part II: immunization of adults. MMWR. In press 2006.

FIGURE 1. Recommended adult immunization schedule, by vaccine and age group — United States, October 2006–September 2007

		Age group (yrs)	
Vaccine	19–49	50–64	≥65
Tetanus, diphtheria, pertussis (Td/Tdap) ¹ *	Substitute 1 do	1-dose Td booster every 10 yrs	3
Human papillomavirus (HPV) ^{2*}	3 doses (females)		
Measles, mumps, rubella (MMR) ³ *	1 or 2 doses	1	dose
Varicella ⁴ *	2 doses (0, 4–8 wks)	2 doses (0, 4–8 wks)	
Influenza ⁵ *	1 dose annually	1 dose	e annually
Pneumococcal (polysaccharide) ^{6,7}	1–2 (doses	1 dose
Hepatitis A ⁸ *	2	doses (0, 6–12 mos, or 0, 6–18 m	os)
Hepatitis B ⁹ *		3 doses (0, 1–2, 4–6 mos)	
Meningococcal ¹⁰		1 or more doses	

^{*} Covered by the Vaccine Injury Compensation Program.

NOTE: These recommendations must be read along with the footnotes, which can be found on pages Q2-Q4 of this schedule.

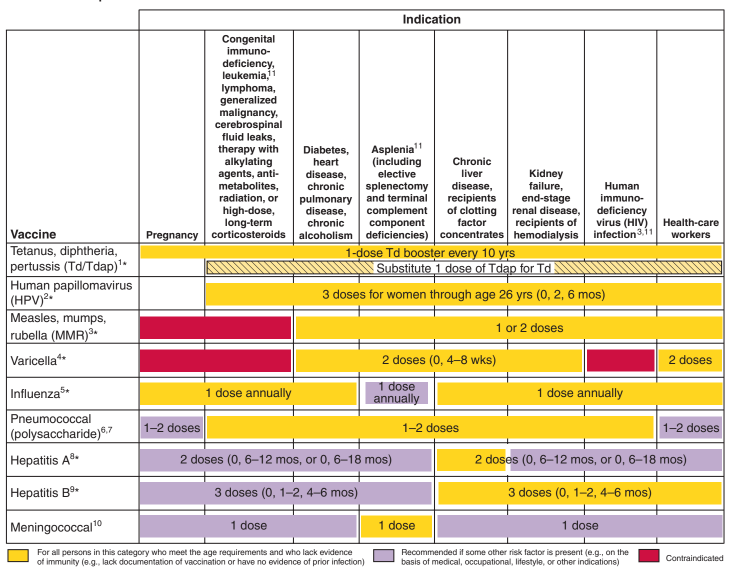
Approved by the Advisory Committee on Immunization Practices, the American College of Obstetricians and Gynecologists, and the American Academy of Family Physicians

- 1. Tetanus, diphtheria, and acellular pertussis (Td/Tdap) vaccination. Adults with uncertain histories of a complete primary vaccination series with diphtheria and tetanus toxoid-containing vaccines should begin or complete a primary vaccination series. A primary series for adults is 3 doses; administer the first 2 doses at least 4 weeks apart and the third dose 6-12 months after the second. Administer a booster dose to adults who have completed a primary series and if the last vaccination was received >10 years previously. Tdap or tetanus and diphtheria (Td) vaccine may be used; Tdap should replace a single dose of Td for adults aged <65 years who have not previously received a dose of Tdap (either in the primary series, as a booster, or for wound management). Only one of two Tdap products (Adacel [sanofi pasteur, Swiftwater, Pennsylvania]) is licensed for use in adults. If the person is pregnant and received the last Td vaccination ≥10 years previously, administer Td during the second or third trimester; if the person received the last Td vaccination in <10 years, administer Tdap during the immediate postpartum period. A one-time administration of 1-dose of Tdap with an interval as short as 2 years from a previous Td vaccination is recommended for postpartum women, close contacts of infants aged <12 months, and all health-care workers with direct patient contact. In certain situations, Td can be deferred during pregnancy and Tdap substituted in the immediate postpartum period, or Tdap can be given instead of Td to a pregnant woman after an informed discussion with the woman (see http://www.cdc.gov/nip/ publications/acip-list.htm). Consult the ACIP statement for recommendations for administering Td as prophylaxis in wound management (http://www.cdc.gov/mmwr/preview/mmwrhtml/ 00041645.htm).
- 2. Human papillomavirus (HPV) vaccination. HPV vaccination is recommended for all women aged ≤26 years who have not completed the vaccine series. Ideally, vaccine should be administered before potential exposure to HPV through sexual activity; however, women

who are sexually active should still be vaccinated. Sexually active women who have not been infected with any of the HPV vaccine types receive the full benefit of the vaccination. Vaccination is less beneficial for women who have already been infected with one or more of the four HPV vaccine types. A complete series consists of 3 doses. The second dose should be administered 2 months after the first dose; the third dose should be administered 6 months after the first dose. Vaccination is not recommended during pregnancy. If a woman is found to be pregnant after initiating the vaccination series, the remainder of the 3-dose regimen should be delayed until after completion of the pregnancy.

3. Measles, mumps, rubella (MMR) vaccination. Measles component: adults born before 1957 can be considered immune to measles. Adults born during or after 1957 should receive >1 dose of MMR unless they have a medical contraindication, documentation of ≥1 dose, history of measles based on health-care provider diagnosis, or laboratory evidence of immunity. A second dose of MMR is recommended for adults who 1) have been recently exposed to measles or in an outbreak setting; 2) have been previously vaccinated with killed measles vaccine; 3) have been vaccinated with an unknown type of measles vaccine during 1963-1967; 4) are students in postsecondary educational institutions; 5) work in a health-care facility; or 6) plan to travel internationally. Withhold MMR or other measles-containing vaccines from HIV-infected persons with severe immunosuppression. Mumps component: adults born before 1957 can generally be considered immune to mumps. Adults born during or after 1957 should receive 1 dose of MMR unless they have a medical contraindication, history of mumps based on health-care provider diagnosis, or laboratory evidence of immunity. A second dose of MMR is recommended for adults who 1) are in an age group that is affected during a mumps outbreak; 2) are students in postsecondary educational institutions; 3) work in a healthcare facility; or 4) plan to travel internationally. For unvaccinated healthcare workers born before 1957 who do not have other evidence of

FIGURE 2. Recommended adult immunization schedule, by vaccine and medical and other indications — United States, October 2006–September 2007



^{*} Covered by the Vaccine Injury Compensation Program.

NOTE: These recommendations must be read along with the footnotes, which can be found on pages Q2-Q4 of this schedule.

mumps immunity, consider giving 1 dose on a routine basis and strongly consider giving a second dose during an outbreak. *Rubella component*: administer 1 dose of MMR vaccine to women whose rubella vaccination history is unreliable or who lack laboratory evidence of immunity. For women of childbearing age, regardless of birth year, routinely determine rubella immunity and counsel women regarding congenital rubella syndrome. Do not vaccinate women who are pregnant or who might become pregnant within 4 weeks of receiving vaccine. Women who do not have evidence of immunity should receive MMR vaccine upon completion or termination of pregnancy and before discharge from the health-care facility.

4. Varicella vaccination. All adults without evidence of immunity to varicella should receive 2 doses of varicella vaccine. Special consideration should be given to those who 1) have close contact with persons at high risk for severe disease (e.g., health-care workers and family contacts of immunocompromised persons) or 2) are at high risk for exposure or transmission (e.g., teachers of young children; child care 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; and international travelers). Evidence of immunity to varicella in adults includes any of the following: 1) documentation of 2 doses of varicella vaccine at least 4 weeks apart; 2) U.S.-born before 1980 (although for health-care workers and pregnant women, birth before 1980 should not be considered evidence of immunity); 3) history of varicella based on diagnosis or verification of varicella by a health-care provider (for a patient reporting a history of or presenting with an atypical case, a mild case, or both, health-care providers should seek either an epidemiologic link with a typical varicella case or evidence of laboratory confirmation, if it was performed at the time of acute disease); 4) history of herpes zoster based on health-care provider diagnosis; or 5) laboratory evidence of immunity or laboratory confirmation of disease. Do not vaccinate women who are pregnant or might become pregnant within 4 weeks of receiving the vaccine. Assess pregnant women for evidence of varicella immunity. Women who do not have evidence of immunity should receive dose 1 of varicella vaccine upon completion or termination of pregnancy and before discharge from the health-care facility. Dose 2 should be administered 4-8 weeks after dose 1.

- 5. Influenza vaccination. Medical indications: chronic disorders of the cardiovascular or pulmonary systems, including asthma; chronic metabolic diseases, including diabetes mellitus, renal dysfunction, hemoglobinopathies, or immunosuppression (including immunosuppression caused by medications or HIV); any condition that compromises respiratory function or the handling of respiratory secretions or that can increase the risk of aspiration (e.g., cognitive dysfunction, spinal cord injury, or seizure disorder or other neuromuscular disorder); and pregnancy during the influenza season. No data exist on the risk for severe or complicated influenza disease among persons with asplenia; however, influenza is a risk factor for secondary bacterial infections that can cause severe disease among persons with asplenia. Occupational indications: health-care workers and employees of long-term-care and assisted living facilities. Other indications: residents of nursing homes and other long-term-care and assisted living facilities; persons likely to transmit influenza to persons at high risk (i.e., in-home household contacts and caregivers of children aged 0-59 months, or persons of all ages with high-risk conditions); and anyone who would like to be vaccinated. Healthy, nonpregnant persons aged 5-49 years without high-risk medical conditions who are not contacts of severely immunocompromised persons in special care units can receive either intranasally administered influenza vaccine (FluMist[®]) or inactivated vaccine. Other persons should receive the inactivated vaccine.
- 6. Pneumococcal polysaccharide vaccination. Medical indications: chronic disorders of the pulmonary system (excluding asthma); cardiovascular diseases; diabetes mellitus; chronic liver diseases, including liver disease as a result of alcohol abuse (e.g., cirrhosis); chronic renal failure or nephrotic syndrome; functional or anatomic asplenia (e.g., sickle cell disease or splenectomy [if elective splenectomy is planned, vaccinate at least 2 weeks before surgery]); immunosuppressive conditions (e.g., congenital immunodeficiency, HIV infection [vaccinate as close to diagnosis as possible when CD4 cell counts are highest], leukemia, lymphoma, multiple myeloma, Hodgkin disease, generalized malignancy, or organ or bone marrow transplantation); chemotherapy with alkylating agents, antimetabolites, or high-dose, long-term corticosteroids; and cochlear implants. Other indications: Alaska Natives and certain American Indian populations and residents of nursing homes or other long-term—care facilities.
- **7. Revaccination with pneumococcal polysaccharide vaccine.** One-time revaccination after 5 years for persons with chronic renal failure or nephrotic syndrome; functional or anatomic asplenia (e.g., sickle cell disease or splenectomy); immunosuppressive conditions (e.g., congenital immunodeficiency, HIV infection, leukemia, lymphoma, multiple myeloma, Hodgkin disease, generalized malignancy, or organ or bone marrow transplantation); or chemotherapy with alkylating agents, antimetabolites, or high-dose, long-term corticosteroids. For persons aged ≥65 years, one-time revaccination if they were vaccinated ≥5 years previously and were aged <65 years at the time of primary vaccination.
- 8. Hepatitis A vaccination. Medical indications: persons with chronic liver disease and persons who receive clotting factor concentrates. Behavioral indications: men who have sex with men and persons who use illegal drugs. Occupational indications: persons working with hepatitis A virus (HAV)—infected primates or with HAV in a research laboratory setting. Other indications: persons traveling to or working in countries that have high or intermediate endemicity of hepatitis A (a

- list of countries is available at http://www.cdc.gov/travel/diseases.htm) and any person who would like to obtain immunity. Current vaccines should be administered in a 2-dose schedule at either 0 and 6–12 months, or 0 and 6–18 months. If the combined hepatitis A and hepatitis B vaccine is used, administer 3 doses at 0, 1, and 6 months.
- 9. Hepatitis B vaccination. Medical indications: persons with endstage renal disease, including patients receiving hemodialysis; persons seeking evaluation or treatment for a sexually transmitted disease (STD); persons with HIV infection; persons with chronic liver disease; and persons who receive clotting factor concentrates. Occupational indications: health-care workers and public-safety workers who are exposed to blood or other potentially infectious body fluids. Behavioral indications: sexually active persons who are not in a long-term, mutually monogamous relationship (i.e., persons with >1 sex partner during the previous 6 months); current or recent injection-drug users; and men who have sex with men. Other indications: household contacts and sex partners of persons with chronic hepatitis B virus (HBV) infection; clients and staff members of institutions for persons with developmental disabilities; all clients of STD clinics; international travelers to countries with high or intermediate prevalence of chronic HBV infection (a list of countries is available at http://www.cdc.gov/travel/diseases.htm); and any adult seeking protection from HBV infection. Settings where hepatitis B vaccination is recommended for all adults: STD treatment facilities; HIV testing and treatment facilities; facilities providing drug-abuse treatment and prevention services; health-care settings providing services for injection-drug users or men who have sex with men; correctional facilities; end-stage renal disease programs and facilities for chronic hemodialysis patients; and institutions and nonresidential daycare facilities for persons with developmental disabilities. Special formulation indications: for adult patients receiving hemodialysis and other immunocompromised adults, 1 dose of 40 µg/mL (Recombivax HB) or 2 doses of 20 μg/mL (Engerix-B[®]).
- 10. Meningococcal vaccination. Medical indications: adults with anatomic or functional asplenia, or terminal complement component deficiencies. Other indications: first-year college students living in dormitories; microbiologists who are routinely exposed to isolates of Neisseria meningitidis; military recruits; and persons who travel to or live in countries in which meningococcal disease is hyperendemic or epidemic (e.g., the "meningitis belt" of sub-Saharan Africa during the dry season [December-June]), particularly if their contact with local populations will be prolonged. Vaccination is required by the government of Saudi Arabia for all travelers to Mecca during the annual Hajj. Meningococcal conjugate vaccine is preferred for adults with any of the preceding indications who are aged ≤55 years, although meningococcal polysaccharide vaccine (MPSV4) is an acceptable alternative. Revaccination after 5 years might be indicated for adults previously vaccinated with MPSV4 who remain at high risk for infection (e.g., persons residing in areas in which disease is epidemic)
- 11. Selected conditions for which *Haemophilus influenzae* type b (Hib) vaccine may be used. Hib conjugate vaccines are licensed for children aged 6 weeks–71 months. No efficacy data are available on which to base a recommendation concerning use of Hib vaccine for older children and adults with the chronic conditions associated with an increased risk for Hib disease. However, studies suggest good immunogenicity in patients who have sickle cell disease, leukemia, or HIV infection or who have had splenectomies; administering vaccine to these patients is not contraindicated.

This schedule indicates the recommended age groups and medical indications for routine administration of currently licensed vaccines for persons aged \geq 19 years, as of October 1, 2006. Licensed combination vaccines may be used whenever any components of the combination are indicated and when the vaccines's other components are not contraindicated. For detailed recommendations on all vaccines, including those used primarily for travelers or that are issued during the year, consult the manufacturers' package inserts and the complete statements from the Advisory Committee on Immunization Practices (http://www.cdc.gov/nip/publications/acip-list.htm).

Report all clinically significant postvaccination reactions to the Vaccine Adverse Event Reporting System (VAERS). Reporting forms and instructions on filing a VAERS report are available at http://www.vaers.hhs.gov or by telephone, 800-822-7967.

Information on how to file a Vaccine Injury Compensation Program claim is available at http://www.hrsa.gov/vaccinecompensation or by telephone, 800-338-2382. To file a claim for vaccine injury, contact the U.S. Court of Federal Claims, 717 Madison Place, N.W., Washington, D.C. 20005; telephone, 202-357-6400.

Additional information about the vaccines in this schedule and contraindications for vaccination is also available at http://www.cdc.gov/nip or from the CDC-INFO Contact Center at 800-CDC-INFO (800-232-4636) in English and Spanish, 24 hours a day, 7 days a week.

TABLE II. Provisional cases of selected notifiable diseases, United States, weeks ending October 7, 2006, and October 8, 2005 (40th Week)*

(40th Week)*	Chlamydia [†]						Cossis	lioidomy	cosic		Cryptosporidiosis				
	-	Pre	vious	ııa [,]				ious	cosis				iosporic	iiosis	
Danielius and	Current	52 v	veeks	Cum	Cum	Current	52 w	eeks	Cum	Cum	Current	52 w	reeks	Cum	Cum
Reporting area United States	10,833	Med 18,961	Max 35,170	2006 723,079	2005 736,812	week 10	Med 149	Max 1,643	2006 6,216	2005 3,222	week 111	Med 69	Max 594	2006 3,594	2005 5,738
New England	1,068	623	1,550	24,919	24,817	_	0	1,043	0,210	3,222	8	4	29	235	285
Connecticut	400	167	1,214	7,196	7,275	N	0	0	N	N	_	0	26	26	64
Maine§ Massachusetts	51 543	43 284	74 442	1,725 11,392	1,714 11,086	N —	0	0	N —	N	_	0 1	4 14	32 88	25 129
New Hampshire Rhode Island	51	37 60	65 100	1,511 2,244	1,424 2,572	_	0	0	_	_	4	1 0	4 6	36 11	30 7
Vermont§	23	19	43	851	746	N	0	0	N	N	4	0	5	42	30
Mid. Atlantic New Jersey	1,475 132	2,380 375	3,696 497	91,650 14,080	90,744 14,850	N	0	0	_ N	_ N	3	11 0	444 3	403 9	2,286 52
New York (Úpstate)	444	499	1,727	18,506	17,969	N	Ō	Ō	N	N	2	3	441	130	1,900
New York City Pennsylvania	439 460	731 739	1,570 1,074	29,148 29,916	29,441 28,484	N N	0	0	N N	N N	1	1 5	9 13	50 214	118 216
E.N. Central	1,836	3,123	12,578	121,131	123,466	_	1	3	37	9	18	16	109	878	1,364
Illinois Indiana	647 320	964 393	1,691 510	39,135 15,042	38,714 15,526	N	0	0	N	N		2 1	8 18	72 68	141 59
Michigan	684	645	9,888	26,632	20,545	_	0	3	33	9	2	2	7	102	89
Ohio Wisconsin	41 144	686 399	1,433 531	25,115 15,207	33,067 15,614	N	0 0	1 0	4 N	N	10 1	5 5	76 52	285 351	650 425
W.N. Central	488	1,152	1,456	44,315	45,432	_	0	12	1	4	14	11	73	645	516
Iowa Kansas	_	154 154	225 269	5,730 5,443	5,484 5,665	N N	0	0 0	N N	N N	_	1 1	28 7	151 58	113 32
Minnesota Missouri	332	230 441	346 612	8,222 17,497	9,493 17,469	_	0	12 1		3 1	11 1	2 2	22 18	155 145	103 221
Nebraska [§]	71	95	176	4,109	3,960	N	0	1	N	N	2	1	16	72	19
North Dakota South Dakota	21 64	33 51	58 116	1,273 2,041	1,230 2,131	N N	0 0	0 0	N N	N N	_	0 1	4 7	8 56	1 27
S. Atlantic	3,117	3,454	4,927	138,603	137,469	_	0	1	3	1	46	14	63	753	546
Delaware District of Columbia	63 23	68 52	92 103	2,714 1,829	2,572 2,957	N	0 0	0 0	N —	N —	_	0 0	3 3	11 12	3 9
Florida Georgia	840 1	942 635	1,154 2,142	37,497 22,464	33,379 24,134	N	0	0	N	N	30 4	6 3	32 11	356 154	243 113
Maryland§	307	331 562	486	13,456	14,176		0	1	3	1 N	8	0	3	15 79	26 69
North Carolina South Carolina§	1,017 349	306	1,772 1,306	25,920 13,895	24,902 14,846	N	0	Ō	N N	N	4	1	11 13	81	18
Virginia§ West Virginia	495 22	423 57	840 226	18,390 2,438	18,458 2,045	N N	0	0	N N	N N	_	1 0	6 3	38 7	53 12
E.S. Central	552	1,419	1,947	55,669	53,492		0	0			11	3	20	140	173
Alabama [§] Kentucky	4	391 160	756 402	15,314 6,427	11,982 6,858	N N	0	0	N N	N N	10 1	1 1	7 19	61 31	21 117
Mississippi Tennessee§	 548	382 495	802 599	14,283 19,645	16,599 18,053	N	0	0	N	N	_	0 1	3 5	14 34	2 33
W.S. Central	615	2,151	3,605	82,185	84,961	_	0	1	1	_	4	4	26	194	185
Arkansas Louisiana	181 225	158 265	333 761	6,270 11,278	6,692 12,966	_	0	0 1	_ 1	N	1	0	2 7	18 41	4 72
Oklahoma	209	228	2,159	9,209	8,735	N	0	Ô	N	N	3	1	4	32	36
Texas [§] Mountain	1,300	1,392 1,027	1,774 1,839	55,428 38,608	56,568 48,496	N 10	0 114	0 452	N 4,330	N 2,100	— 5	2 2	20 38	103 280	73 111
Arizona	881	354	642	14,119	16,579	10	111	448	4,256	2,020	_	0	2	19	9
Colorado Idaho [§]	62 191	156 50	482 159	4,512 2,236	11,670 1,998	N N	0	0 0	N N	N N	2 1	1 0	7 5	58 25	38 13
Montana Nevada§	 166	42 77	195 432	1,825 3,732	1,775 5,642	<u>N</u>	0	0 4	N 21	N 48	_	0	26 1	104 4	16 11
New Mexico [§]	_	172	339	7,422	6,504	_	0	3	13	16	_	0	4	16	10
Utah Wyoming	_	94 27	170 55	3,731 1,031	3,457 871	_	1 0	3 2	38 2	13 3		0 0	3 11	16 38	11 3
Pacific	382	3,315	5,079	125,999	127,935	_	43	1,179	1,844	1,108	2	2	52	66	272
Alaska California	57 —	85 2,570	152 4,231	3,247 98,604	3,263 99,331	_	0 43	0 1,179	1,844	1,108	_	0 0	1 14	4	3 156
Hawaii Oregon [§]	_	103 174	135 315	3,948 6,624	4,264 6,836	N N	0	0	N N	N N		0	1 6	4 58	1 60
Washington	325	350	604	13,576	14,241	N	0	0	N	N	_	Ó	38	_	52
American Samoa C.N.M.I.	U	0	46 0	U	U U	U U	0	0	U	U U	U U	0	0	U U	U
Guam	_	18	37	_	629	_	0	0	_	_	_	0	0	_	_
Puerto Rico U.S. Virgin Islands	_	75 5	161 16	2,945 178	3,187 196	N —	0	0 0	N —	N —	N —	0 0	0 0	N —	N —
C N M I : Commonwe	alth of Nort						-	-				-			

C.N.M.I.: Commonwealth of Northern Mariana Islands.
U: Unavailable. —: No reported cases. N: Not notifiable. Cum: Cumulative year-to-one in the common state of the co Cum: Cumulative year-to-date counts. Med: Median. Max: Maximum.

TABLE II. (*Continued*) Provisional cases of selected notifiable diseases, United States, weeks ending October 7, 2006, and October 8, 2005 (40th Week)*

	Giardiasis Previous							onorrhe	a		Hae	All age	es, all se	<i>zae</i> , inva rotypes	sive
	Current		rious eeks	Cum	Cum	Current		/ious /eeks	Cum	Cum	Current		vious veeks	Cum	Cum
Reporting area	week	Med	Max	2006	2005	week	Med	Max	2006	2005	week	Med	Max	2006	2005
United States	297	322	1,029	12,510	14,626	3,548	6,499	,	250,416	- ,	14	38	142	1,550	1,777
New England Connecticut	23 8	24 0	75 37	967 222	1,331 280	145 82	107 41	288 241	4,211 1,698	4,476 1,908	1 1	2	19 9	127 38	136 40
Maine [†] Massachusetts	6	2 10	13 25	133 357	168 602	2 58	2 46	6 86	98 1,855	110 1,936	_	0 1	4 7	17 52	8 68
New Hampshire Rhode Island	1	0 0	9 25	24 92	50 86	1	4 8	9 19	149 360	127 350	_	0 0	2 7	7 4	7 7
Vermont [†]	8	3	10	139	145	2	1	4	51	45	_	0	2	9	6
Mid. Atlantic New Jersey	39 —	61 9	254 13	2,239 297	2,646 351	448 88	636 102	1,014 143	24,324 3,767	26,068 4,399		7 1	30 4	299 45	333 67
New York (Upstate) New York City	27 2	24 10	227 31	911 438	909 708	107 98	123 175	455 357	4,787 7,198	5,240 7,873	2	2 1	27 4	103 36	97 61
Pennsylvania	10	15	29	593	678	155	215	394	8,572	8,556	_	3	8	115	108
E.N. Central Illinois	28 —	49 9	86 21	1,846 317	2,612 615	654 207	1,285 377	7,047 709	49,286 14,969	50,104 15,252	2	5 1	14 6	219 47	308 104
Indiana Michigan	N 3	0 14	0 22	N 502	N 631	137 262	161 252	237 5,880	6,713 11,097	6,259 8,512	2	1 0	11 3	66 18	54 19
Ohio Wisconsin	25	16 10	32 40	624 403	605 761	15 33	329 134	648 172	11,390 5,117	15,637 4,444	_	2	6 4	65 23	94 37
W.N. Central	19	29	260	1,423	1,614	173	364	436	14,024	14,447	2	2	15	112	88
Iowa Kansas	1	5 3	15 11	223 148	215 158	_	33 44	46 124	1,199 1,519	1,219 2,010	_	0	1 3	1 14	9
Minnesota Missouri	 15	2 10	238 32	477 420	667 365	 155	62 190	105 251	2,113 7,753	2,660 7,309	_ 2	0	9	56 30	37 29
Nebraska†	3	2	8	86	103	11	23	56	1,062	899	_	0	2	7	12
North Dakota South Dakota	_	0 1	7 7	12 57	12 94	1 6	3 6	7 15	87 291	79 271	_	0	3 0	4	1
S. Atlantic Delaware	80	49 1	95 4	1,920 33	2,118 44	1,187 27	1,491 27	2,334 44	61,369 1,132	60,107 663	1	10 0	26 1	409 1	420
District of Columbia		1	5	52	41	24	34	61	1,238	1,633	_	0	1	4	7
Florida Georgia	40 29	18 10	39 44	821 411	743 565	381 4	437 300	553 1,014	17,853 10,865	15,359 11,261	1	3 2	9 12	133 80	103 90
Maryland [†] North Carolina	8 N	4 0	11 0	158 N	161 N	102 332	128 284	186 766	5,022 13,093	5,324 11,903	_	1 0	5 9	53 46	58 68
South Carolina† Virginia†		1 8	7 50	69 359	86 446	151 148	132 130	704 288	6,262 5,161	6,798 6,628	_	1 1	3 8	27 49	28 43
West Virginia	_	0	5	17	32	18	17	42	743	538	_	0	4	16	23
E.S. Central Alabama†	20 13	8 4	40 29	377 204	329 149	205	564 183	863 310	22,544 7,110	21,249 6,880	_	2 0	7 5	79 20	94 17
Kentucky Mississippi	N	0	0	N	<u>N</u>	7	55 141	132 435	2,301 5,607	2,353 5.401	_	0	1 1	4	11
Tennessee [†]	7	4	12	173	180	198	187	237	7,526	6,615	_	1	4	52	66
W.S. Central Arkansas	11 6	5 2	31 6	209 92	248 66	291 97	879 79	1,430 142	35,567 3,241	34,532 3,499	3	1 0	15 2	54 7	95 7
Louisiana Oklahoma	<u> </u>	0 2	3 24	18 99	51 131	123 71	161 82	354 764	6,889 3,440	7,183 3,482		0 1	2 14	5 40	32 51
Texas [†]	N	0	0	N	N		541	836	21,997	20,368	_	Ö	2	2	5
Mountain Arizona	47 —	30 3	56 36	1,230 116	1,147 108	362 198	217 90	552 201	8,715 3,541	10,447 3,768	2	4 1	8 7	158 73	183 92
Colorado Idaho†	21 7	9 3	33 11	439 134	400 111	84 15	41 2	90 10	1,595 132	2,459 84	1 1	1 0	4 1	42 4	37 4
Montana Nevada [†]	<u>,</u>	2	11 6	79 41	58 83	65	2 24	20 194	145 1,230	121		0	0 1		
New Mexico†	_	1	6	47	68	_	31	64	1,348	2,215 1,211	_	0	4	21	21
Utah Wyoming	18 1	7 1	19 4	344 30	299 20	_	17 2	25 6	631 93	530 59	_	0	4 1	15 3	8 7
Pacific	30	59	202	2,299	2,581	83	807	963	30,376	31,379	1	2	15	93	120
Alaska California	3	1 43	15 105	75 1,606	86 1,833	11	11 659	23 830	451 24,950	447 26,144	_	0	9	9 21	26 50
Hawaii Oregon [†]	7	1 8	3 14	37 308	52 344	<u>1</u>	18 28	29 58	707 1,016	788 1,174		0 1	1 6	14 47	8 36
Washington	20	6	90	273	266	71	75	142	3,252	2,826	_	0	4	2	_
American Samoa C.N.M.I.	U U	0	0	U U	U	U U	0	2	U	U	U U	0	0	U U	U U
Guam Puerto Rico	4	0 2	0 12	<u> </u>	11 211	_	1 5	15 16	188	71 286	_	1 0	2 1	_ 1	7 3
U.S. Virgin Islands	_	0	0	_	_	_	0	5	30	45	_	0	0	_	_

Med: Median.

Max: Maximum.

Cum: Cumulative year-to-date counts.

C.N.M.I.: Commonwealth of Northern Mariana Islands.
U: Unavailable. —: No reported cases. N: Not notifiable. Cum: Cumulative year-to* Incidence data for reporting year 2006 is provisional.

† Contains data reported through the National Electronic Disease Surveillance System (NEDSS).

TABLE II. (*Continued*) Provisional cases of selected notifiable diseases, United States, weeks ending October 7, 2006, and October 8, 2005 (40th Week)*

(40th Week)*				Нер	atitis (viral	, acute), by	type							_	
		Drov	A rious				Previ	В					gionello vious	sis	
Departing area	Current		eeks	Cum 2006	Cum 2005	Current		eks	Cum 2006	Cum 2005	Current	52 v	/eeks	Cum 2006	Cum
Reporting area United States	week 17	69	Max 245	2,463	3,245	week 45	84	Max 597	3,064	4,035	week 37	Med 44	127	1,636	2005 1,574
New England	_	3	20	144	371	_	1	9	49	119	9	2	12	99	106
Connecticut	_	1	2	34	43	_	0	3	_	39	9	0	8	38	22
Maine† Massachusetts	_	0 1	2 13	6 51	3 234	_	0 0	2 5	15 14	12 39	_	0 1	2 6	7 27	5 49
New Hampshire Rhode Island	_	0 0	16 4	36 9	75 10	_	0	2 4	11 8	24 1	_	0	1 10	1 20	8 16
Vermont†	=	0	2	8	6	=	0	1	1	4	_	0	3	6	6
Mid. Atlantic	4	7	18	270	525	3	8	55	317	527	16	13	42	553	537
New Jersey New York (Upstate)	4	2 1	7 14	61 67	109 81	_	2 1	8 43	80 49	195 45	11	1 5	10 29	61 230	91 137
New York City Pennsylvania	_	2 1	12 5	92 50	252 83		2	4 9	60 128	110 177	 5	1 4	9 17	46 216	83 226
E.N. Central	2	7	12	224	284	2	8	24	311	438	2	8	25	335	326
Illinois	_	1	4	50	103	_	2	7	57	124	_	1	4	21	46
Indiana Michigan	1	0 2	5 8	23 78	14 88	1	0 3	17 7	42 105	33 143	_	0 2	3 7	24 86	21 89
Ohio Wisconsin	1	1 1	4 5	45 28	41 38	1	2	10 4	101 6	104 34	2	4 0	19 5	171 33	141 29
W.N. Central	1	2	30	103	70	_	4	22	125	213	_	1	15	52	63
Iowa	_	0	2	8	18	_	0	3	14	21	_	0	3	10	4
Kansas Minnesota	_	0 0	5 29	24 9	13 3	_	0 0	2 13	8 17	24 29	_	0 0	2 11	3 11	2 16
Missouri Nebraska [†]	1	1 0	3 3	39 15	28 8	_	2	7 1	74 11	111 22	_	0	3 2	18 6	24 3
North Dakota	_	0	2	_	_	_	0	0	_	_	_	0	1	_	2
South Dakota	_	0	3	8		_	0	1	1	6	_	0	6	4	12
S. Atlantic Delaware	6	11 0	29 2	419 10	572 5	12	23 1	66 4	889 35	1,081 25	6	9	19 2	322 8	304 13
District of Columbia Florida	<u> </u>	0 4	2 13	6 165	3 228	 8	0 8	2 19	5 322	10 370		0 3	5 9	16 130	9 84
Georgia	_	1	7	50	108	1	3	7	124	166	_	0	4	15	27
Maryland† North Carolina	1	1 0	6 20	53 67	58 70	_	3 0	10 23	128 123	121 128	4	1 0	6 5	65 29	87 24
South Carolina†	_	0	2	17	34	3	2	7	63	121	_	0	1	2	11
Virginia† West Virginia	1	1 0	11 3	46 5	63 3	_	1 0	18 18	43 46	113 27	_	1 0	7 3	49 8	35 14
E.S. Central	2	2	8	97	216	4	6	15	249	284	1	1	9	67	62
Alabama† Kentucky	_	0 0	3 5	13 29	40 22	_ 1	1 1	8 5	78 57	67 54	1	0 0	2 4	9 23	12 21
Mississippi Tennessee [†]	2	0	1 5	5 50	17 137		0 2	2	11 103	44 119	_	0	1 7	1 34	3
W.S. Central	2	4	5 77	138	376	3 17	14	315	578	478	_	1	32	43	26 38
Arkansas	_	0	9	35	16	1	1	4	37	53	_	0	3	3	5
Louisiana Oklahoma		0 0	4 2	15 6	56 4	12	0 0	3 17	25 43	62 37	_	0	2	4 1	1 7
Texas [†]	_	2	73	82	300	4	12	295	473	326	_	0	26	35	25
Mountain Arizona	_	5 2	18 16	192 108	253 135	_	4 1	39 23	126 33	429 276	3	2 1	7 4	95 32	79 19
Colorado	_	1	4	33	34	_	1	5	29	44	1	0	2	21	17
Idaho† Montana	_	0 0	2	9 9	20 7	_	0	2 7	10	12 3	1	0 0	3 1	11 5	3 5
Nevada† New Mexico†	_	0	2	7 12	19 19	_	0	4 3	15 15	42 18	_	0	2 1	4 4	17 3
Utah	_	0	2	11	18	_	0	5	24	32	1	0	1	18	11
Wyoming	_	0	1	3	1	_	0	1	_	2	_	0	0	_	4
Pacific Alaska	_	19 0	163 0	876 —	578 4	7	9 0	61 1	420 5	466 7	_	1 0	9 1	70 —	59 —
California Hawaii	_	15 0	162 2	793 9	477 21	_	7 0	41 1	317 5	312 6	_	1 0	9	70 —	57 2
Oregon [†]	_	0	5	37	38	1	1	5	54	84	N	0	0	N	N
Washington	_	1	13	37	38	6	0	18	39	57	_	0	0	_	_
American Samoa C.N.M.I.	U U	0 0	0 0	U	1 U	U U	0	0	U U	U	U U	0	0	U U	U
Guam Puerto Rico	_	0 0	0 5		2 58		0	0		18 38		0	0	- 1	_
U.S. Virgin Islands	_	0	0	23 —	58	_	0	0	24 —	38	_	0	0	_	_
<u> </u>															

Cum: Cumulative year-to-date counts. Med: Median. Max: Maximum.

C.N.M.I.: Commonwealth of Northern Mariana Islands.
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† Contains data reported through the National Electronic Disease Surveillance System (NEDSS).

TABLE II. (*Continued*) Provisional cases of selected notifiable diseases, United States, weeks ending October 7, 2006, and October 8, 2005 (40th Week)*

(40th Week)*												
		Due	Lyme dis	ease		_		Duan	Malaria	l		
	Current		vious reeks	Cum	Cum	c	urrent	Prev 52 w		Cum	Cum	
Reporting area	week	Med	Max	2006	2005		week	Med	Max	2006	2005	
United States	101	252	2,153	13,234	17,704		13	24	125	916	1,096	
New England	29	37	780	2,217	3,094		_	1	11	44	63	
Connecticut Maine [†]	11 16	13 1	753 34	1,519 164	491 213		_	0 0	5 1	11 4	16 5	
Massachusetts		1	35	33	2,129		_	0	3	19	34	
New Hampshire	2	6	59	422	188		_	0	3	9	5	
Rhode Island Vermont [†]	_	0 1	5 14	1 78	32 41		_	0 0	8 1	_ 1	2 1	
Mid. Atlantic	62	153	1,176	7,681	10,314		2	4	13	163	297	
New Jersey	_	22	168	1,656	3,111		_	1	3	28	69	
New York (Upstate) New York City	51 —	75 1	1,150 18	3,244 124	3,084 349		2	1 2	11 8	36 64	39 160	
Pennsylvania	11	40	222	2,657	3,770		_	1	3	35	29	
E.N. Central	1	10	134	1,161	1,605		2	2	7	101	117	
Illinois	_	0	2	_	119		_	1	4	42	66	
Indiana Michigan	_ 1	0 1	3 6	16 41	26 47		_	0 0	3 2	9 16	4 19	
Ohio	_	1	6	38	49		2	0	3	27	18	
Wisconsin	_	10	129	1,066	1,364		_	0	3	7	10	
W.N. Central	_	7	168	497	646		_	0	32	33	43	
Iowa Kansas	_	1 0	8 2	77 4	86 3		_	0 0	1 2	1 6	8 5	
Minnesota	_	5	167	398	539		_	0	30	14	11	
Missouri Nobraskat	_	0	3	9	13		_	0	1 1	6	16	
Nebraska† North Dakota	_	0	1 3	8 —	3		_	0	1	4 1	3	
South Dakota	_	0	1	1	2		_	0	1	1	_	
S. Atlantic	8	32	108	1,416	1,841		6	6	15	258	234	
Delaware District of Columbia		8 0	28 7	404 41	567 8		_	0 0	1 2	5 3	3 8	
Florida	3	1	5	35	34		3	1	6	51	40	
Georgia	_	0	1	3	5		_	1	6	66	43	
Maryland† North Carolina	2	14 0	65 4	676 24	984 42		3	1 0	5 8	57 24	86 24	
South Carolina [†]	_	0	2	10	19		_	0	2	8	7	
Virginia† West Virginia	1	3 0	25 44	214 9	172 10		_	1 0	9 2	42 2	22 1	
E.S. Central		0	3	21	31		_	0	3	19	23	
Alabama†	_	0	1	6	2		_	0	2	8	4	
Kentucky	_	0	2	7	5		_	0	2	3	8	
Mississippi Tennessee [†]	_	0	0 2	 8	 24		_	0 0	1 2	3 5	 11	
W.S. Central	1	0	3	15	69		_	2	31	55	105	
Arkansas		0	1	_	4		_	0	1	2	5	
Louisiana Oklahoma	_	0	0	_	3		_	0	1	4 7	3 9	
Okianoma Texas†	1	0	0 3	15	62		_	0 1	2 29	7 42	9 88	
Mountain	_	0	3	22	20		1	1	9	53	44	
Arizona	_	0	2	4	7		_	0	9	17	10	
Colorado Idaho†	_	0 0	1 2	5 4			_	0 0	2 1	11 1	21 —	
Montana	_	0	0	_	_		_	0	1	2	_	
Nevada†	_	0	1	1	3		_	0	1	1	3	
New Mexico [†] Utah	_	0	1 1	1 6	3 2		1	0 0	1 2	4 17	3 5	
Wyoming	_	Ö	i	1	3		_	Ö	0		2	
Pacific	_	4	17	204	84		2	5	13	190	170	
Alaska California	_	0	1	100	4		_	0	4	23	5	
California Hawaii	N	4 0	16 0	190 N	54 N		_	4 0	10 2	127 4	125 15	
Oregon [†]	_	Ō	2	9	18		_	0	1	9	10	
Washington	_	0	3	3	8		2	0	5	27	15	
American Samoa C.N.M.I.	U U	0	0	U	U U		U U	0 0	0 0	U U	U U	
Guam	_	0	0	_	_		_	0	0	_	_	
Puerto Rico	N	0	0	N	N		_	0	1	_	3	
U.S. Virgin Islands		0	0	_	_		_	0	0		_	

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U: Unavailable. —: No reported cases. N: Not notifiable. Cum: Cumulative year-to-common to the common state of the Cum: Cumulative year-to-date counts. Med: Median. Max: Maximum.

TABLE II. (*Continued*) Provisional cases of selected notifiable diseases, United States, weeks ending October 7, 2006, and October 8, 2005 (40th Week)*

(40th Week)*				Menir	ngococcal	disease, inv	asive								
			All serog	roups				<u> </u>	ınknown				Pertus	ssis	
Reporting area	Current week	Prev 52 w Med	ious eeks Max	Cum 2006	Cum 2005	Current week	Previo		Cum 2006	Cum 2005	Current week		vious veeks Max	Cum 2006	Cum 2005
United States	5	20	85	840	962	4	13	58	548	587	129	265	2,877	9,945	17,567
New England Connecticut Maine [†]	_	1 0 0	3 2 1	35 9 4	61 12 2	_	0 0 0	2 2 1	25 2 3	22 1 2	1 	29 1 1	83 5 11	959 35 63	1,057 52 40
Massachusetts New Hampshire Rhode Island Vermont [†]		0 0 0 0	2 2 1 1	15 5 — 2	28 12 2 5		0 0 0	2 2 0 0	15 5 —	5 12 — 2	_ _ _ 1	19 2 0 1	43 36 17 14	594 129 45 93	805 54 29 77
Mid. Atlantic New Jersey New York (Upstate)	1 	3 0 1	14 2 7	121 11 31	118 27 31	1 _ _	2 0 0	11 2 5	90 11 4	90 27 11	50 — 34	34 3 14	137 13 123	1,412 156 644	1,052 143 402
New York City Pennsylvania	<u>1</u>	0 1	6 5	42 37	18 42	<u>1</u>	0	6 5	42 33	18 34	16	1 11	8 26	64 548	86 421
E.N. Central Illinois Indiana Michigan	1 - -	3 0 0 0	11 4 5 3	96 18 19 19	120 27 18 24	1 — —	1 0 0 0	6 4 1 3	65 18 6 8	99 27 8 15	23 — 5 4	39 7 4 7	133 35 75 24	1,406 230 189 389	2,986 685 252 246
Ohio Wisconsin	1	1 0	5 2	37 3	32 19	1	1 0	4 2	30 3	30 19	14	14 4	30 41	459 139	907 896
W.N. Central lowa	_	1 0	4 2	45 13	64 15	_	0	3 1	15 5	28 1	16 —	28 6	552 63	955 212	2,908 733
Kansas Minnesota Missouri	_	0 0 0	1 2 2	1 11 13	9 11 22	_	0 0 0	1 1 1	1 3 2	9 4 11	14 1	7 0 7	28 485 42	226 160 241	334 966 361
Nebraska [†] North Dakota	_	0	2 1	5 1	4	_	0	1	3 1	3	1	2	9 25	73 26	232 112
South Dakota S. Atlantic	_	0	1 14	1 149	3 182	2	0 2	0 7	61	— 77	9	0 20	46	17 758	170 1,133
Delaware District of Columbia Florida Georgia	2	0 0 1 0	1 1 6 2	4 1 59 12	4 5 69 14	 2	0 0 0	1 1 5 2	4 1 21 12	4 4 27 14	2 3	0 0 4 0	1 3 9 3	3 6 172 15	15 7 165 41
Maryland [†] North Carolina South Carolina [†]		0 0 0	2 11 2	11 24 18	19 28 13	=	0 0	1 3 2	2 7 8	3 6 8	1 1 2	3 0 3	9 22 22	97 155 129	162 98 322
Virginia† West Virginia	_	0	4 2	15 5	24 6	_	0	3	6	9 2	_	2	27 9	155 26	284 39
E.S. Central Alabama [†] Kentucky	_	1 0 0	4 1 2	31 5 7	47 5 16	_	1 0 0	4 1 2	25 4 7	36 3 16	1	7 1 1	16 7 5	261 54 53	435 72 130
Mississippi Tennessee [†]	_	0	1 2	3 16	5 21	_	0	1 2	3 11	5 12	1	1 2	4 10	35 119	48 185
W.S. Central Arkansas Louisiana	_	1 0 0	23 3 2	52 9 6	93 12 28	_	0 0 0	6 2 1	23 6 3	23 3 5	_	16 1 0	360 21 3	520 47 11	1,842 245 44
Oklahoma Texas [†]		0	4 16	8 29	14 39	=	0	0 4	- 14	2 13	Ξ	0 14	124 215	18 444	1,552
Mountain Arizona Colorado	_	1 0 0	5 3 2	57 16 19	80 31 17	_	0 0 0	4 3 1	27 16 2	21 10	22 2 5	61 9 20	230 177 40	2,099 402 650	3,260 819 1,040
Idaho† Montana	=	0	2 1	3 4	4	=	0	2 1	2 2	3		2 2	8 9	74 96	174 549
Nevada† New Mexico† Utah	_	0 0 0	1 1 1	3 3 5	12 5 11		0 0 0	0 1 0	1	2 4 2	 13	0 2 15	9 6 39	39 60 716	43 152 439
Wyoming Pacific	1	0 5	2 29	4 254	— 197	_	0 5	2 25	4 217	— 191	7	1 42	8 1,334	62 1,575	44 2,894
Alaska California Hawaii Oregon [†]		0 3 0 1	1 14 1 7	2 156 7 60	3 128 11 36		0 3 0 1	1 14 1 4	2 156 7 41	3 128 6 36	_ _ _	2 27 2 2	15 1,136 4 8	61 1,099 65 94	110 1,359 142 598
Washington	1	0	25	29	19	_	0	11	11	18	7	7	195	256	685
American Samoa C.N.M.I. Guam	U U —	0 0 0	0 0 0	=	<u>_</u> 1	U U —	0 0 0	0 0 0	U U —	U U 1	U U	0 0 0	0 0 0	U U	U U 2
Puerto Rico U.S. Virgin Islands	_	0	1 0	_	6	_	0	1 0		<u>6</u>	_	0	1 0	<u>1</u>	5

Cum: Cumulative year-to-date counts. Med: Median. Max: Maximum.

C.N.M.I.: Commonwealth of Northern Mariana Islands.
U: Unavailable. —: No reported cases. N: Not notifiable. Cum: Cumulative year-to* Incidence data for reporting year 2006 is provisional.

† Contains data reported through the National Electronic Disease Surveillance System (NEDSS).

TABLE II. (*Continued*) Provisional cases of selected notifiable diseases, United States, weeks ending October 7, 2006, and October 8, 2005 (40th Week)*

(40th Week)*	Rabies, animal							Salmonellosis							
			, .	mal		Roc			tted fever				almonello vious	osis	
	Current	Prev 52 w		Cum	Cum	Current	Prev 52 w		Cum	Cum	Current		weeks	Cum	Cum
Reporting area	week	Med	Max	2006	2005	week	Med	Max	2006	2005	week	Med	Max	2006	2005
United States	30	109	173	4,654	4,807	3	37	246	1,599	1,349	640	809	2,291	30,471	33,347
New England Connecticut	7 4	11 3	26 14	529 160	574 156	_	0	2	2	7	6	30 0	383 375	1,533 375	1,776 392
Maine [†]	_	2	7 17	84 178	50	N	0	0	N 1	N 5	_	2 18	10 53	91 782	139 941
Massachusetts New Hampshire	1	0	5	39	288 12	_	0	1	1	1	4	2	24	160	144
Rhode Island Vermont [†]		0 1	4 4	20 48	20 48	_	0	2 0	_	1	_	0 1	17 5	73 52	81 79
Mid. Atlantic	1	23	59	1,103	779	1	1	6	56	79	59	86	272	3,551	4,069
New Jersey New York (Upstate)	N	0 11	0 22	N 416	N 435	_	0	2 2	7 4	25 1	 37	14 22	43 233	630 966	805 965
New York City	1	0	5	20	23	_	Ō	4	13	6	2	18	36	694	955
Pennsylvania E.N. Central	_ 1	14 1	42 18	667 141	321 163	1	1	3 6	32 32	47 37	20 72	29 99	67 172	1,261 3,926	1,344 4,597
Illinois	1	0	7	44	47	_	0	1	3	11	_	25	45	854	1,527
Indiana Michigan	_	0	2 5	11 41	11 35	_	0	1 1	5 2	 5	35 3	15 17	67 32	711 749	479 749
Ohio Wisconsin	N	0	9	45 N	70 N	_	0	4 1	21 1	19 2	34	23 15	56 27	983 629	1,058 784
W.N. Central	4	5	20	251	281	1	2	15	190	141	35	42	107	1,994	2,040
Iowa Kansas		0 1	7 5	52 61			0	1	4 2	5 5		7 6	21 16	335 259	339 296
Minnesota	_	1	6	36	61	_	0	2	4	2	22	10	60	552	443
Missouri Nebraska†	4	1 0	8 0	65 —	65 —	1	2 0	11 5	159 21	117 7	12 1	14 4	36 9	587 142	634 170
North Dakota South Dakota	_	0	7	16 21	28 57	_	0	1	_	 5	_	0 3	46 7	19 100	28 130
S. Atlantic	_	36	118	1,562	1,719	_	16	94	894	674	321	207	450	8,192	9,223
Delaware District of Columbia	_	0	0	_	_	_	0	3	18 1	7		2	9	117 50	105 45
Florida	_	0	99	136	201	_	0	3	15	13	139	95	228	3,449	3,593
Georgia Marvland [†]	_	2 7	9 13	100 254	214 308	_	0 1	3 6	28 60	84 60	52 18	27 12	100 29	1,243 540	1,464 645
North Carolina South Carolina [†]	_	9 3	22 10	397 129	391 176	_	15 0	87 6	663 23	356 61	85 21	32 19	130 51	1,231 720	1,219 1,113
Virginia [†]	_	10	27	458	383	_	2	13	83	86	4	20	57	751	905
West Virginia E.S. Central	 8	1 4	13 16	88 197	46 128	_	0 5	2 26	3 273	5 247	30	2 54	19 149	91 2,197	134 2,329
Alabama [†]	8	1	7	69	67	_	1	8	78	64	5	14	71	729	550
Kentucky Mississippi	_	0 0	5 2	23 4	16 5	_	0 0	1 1	1 2	3 13	7	8 12	22 47	350 541	393 729
Tennessee [†]	_	2	9	101	40	_	3	20	192	167	18	14	31	577	657
W.S. Central Arkansas	1 1	14 0	34 4	549 26	744 31	1	1 0	161 10	101 46	137 98	62 44	83 14	922 45	2,904 703	3,226 566
Louisiana Oklahoma	_	0	0 9	<u></u>	67	_	0	1 154	1 35	6 7	2 16	12 7	35 48	400 382	734 326
Texas [†]	_	12	29	471	646	1	0	3	19	26	-	48	839	1,419	1,600
Mountain	8	3	12	156	236	_	0	6	44	25	19	50	84	1,864	1,836
Arizona Colorado	6	2 0	10 1	119 —	152 16	_	Ö	6 1	8 2	12 4	2 6	15 12	67 30	587 515	499 472
Idaho [†] Montana	_	0	12 2	2 13	 15	_	0	3 2	13 2	3 1	3	3 3	9 16	137 107	114 69
Nevada [†] New Mexico [†]	_	0	1 2	1 7	14 9	_	0	0 2		_ 3	_	1	17 12	72 179	146 212
Utah	1	0	1	9	14	_	0	2	6	_	8	5	15	230	255
Wyoming	1	0	2	5	16	_	0	1	6	2	_	1	5	37	69
Pacific Alaska	_	4 0	10 4	166 14	183 1	_	0 0	1 0	7 —	_	36 1	110 1	426 7	4,310 62	4,251 45
California Hawaii	_	3 0	10 0	135	176 —	_	0	1 0	5	_	_	88 5	292 10	3,369 181	3,217 237
Oregon [†]	 U	0	4	17 U	6 U	N	0	1	2 N	2 N	1 34	7 7	16 124	324 374	331 421
Washington American Samoa	U	0	0	U	U	U	0	0	N U	U	34 U	0	124	374 U	7
C.N.M.I.	ŭ	0	0	Ü	ŭ	Ü	0	0	U	U	Ü	0	0	Ü	U
Guam Puerto Rico	_	0 1	0 6	66	55	N	0	0	N	N	11	1 6	3 35	189	30 506
U.S. Virgin Islands	_	0	0	_	_	_	0	0	_	_	_	0	0	_	_

C.N.M.I.: Commonwealth of Northern Mariana Islands.
U: Unavailable. —: No reported cases. N: No U: Unavailable. —: No reported cases. N: Not notifiable. Cum: Cumulative year-to
* Incidence data for reporting year 2006 is provisional.

† Contains data reported through the National Electronic Disease Surveillance System (NEDSS). Cum: Cumulative year-to-date counts. Med: Median. Max: Maximum.

TABLE II. (Continued) Provisional cases of selected notifiable diseases, United States, weeks ending October 7, 2006, and October 8, 2005 (40th Week)*

(40th Week)*	Shiga toxin-producing E. coli (STEC)†					Shigellosis					Streptococcal disease, invasive, group A				
	Current	Prev 52 w		Cum	Cum	Current	Prev 52 w		Cum	Cum	Current		ious eeks	Cum	Cum
Reporting area	week	Med	Max	2006	2005	week	Med	Max	2006	2005	week	Med	Max	2006	2005
United States	54	56	297	2,264	2,419	220	242	1,013	8,895	11,307	42	87	283	3,828	3,635
New England Connecticut	1	3	59 58	217 58	184 50	_	4	57 51	205 51	259 47	2 U	4	15 3	177 U	235 82
Maine§ Massachusetts	_	0 1	8 9	30 82	28 71	_	0 3	2 11	3 128	13 157	_	0 2	2 6	15 101	12 107
New Hampshire Rhode Island	1	0	3 2	22 8	14 5	_	0	4 6	7 11	12 14	2	0	9	44 5	16 9
Vermont§	_	0	2	2	16	_	Ö	2	5	16	_	Ö	2	12	9
Mid. Atlantic New Jersey	3	5 0	107 3	158 3	286 62	5 —	14 4	72 26	589 206	1,046 267	5 —	15 3	43 8	699 122	732 151
New York (Upstate) New York City	_	0	103 4	12 27	110 13	4 1	4 4	60 12	188 128	219 347	4	4	32 9	251 74	209 143
Pennsylvania	_	0	5	5	101		2	5	67	213	1	6	13	252	229
E.N. Central Illinois	11	11 1	52 7	503 59	503 120	5	20 7	38 16	703 229	890 305	3	14 4	43 11	666 144	757 252
Indiana	4	1	8 7	68	50	3	2	18 10	113	121 192	_	2	11	92	86 179
Michigan Ohio	7	1 3	18	69 150	78 126	2	3	11	119 130	82	3	3 4	12 19	183 205	161
Wisconsin W.N. Central	 10	2 8	39 35	157 335	129 397	 21	3 34	9 77	112 1,225	190 1,220	10	1 5	4 57	42 285	79 225
Iowa	_	2	8	108	82		2	10	77	68	N	0	0	N	N
Kansas Minnesota	7	0 3	3 27	186	39 120	17	3 2	20 10	103 122	167 70	9	1	5 52	46 136	35 86
Missouri Nebraska [§]	5 —	2 1	13 8	143 55	82 42	4	13 2	69 14	574 102	783 83	1	1 0	5 4	61 25	57 18
North Dakota South Dakota	_	0	15 5	 29	6 26	_	0 4	18 21	63 184	4 45	_	0	5 3	9 8	9 20
S. Atlantic	3	7 0	39 2	350	322	118	54 0	122	2,150	1,682 10	12	22	43	927	725
Delaware District of Columbia	_	0	1	7 2	8 —	1	0	2	8 14	9		0	2	10 13	5 8
Florida Georgia	1 —	2 1	29 6	75 69	75 44	73 43	26 17	66 41	1,064 714	818 435	9	6 5	16 11	234 176	188 153
Maryland [§] North Carolina	2 7	1 1	8 10	69 90	66 44	_	2 1	10 21	94 125	68 149	1	4 0	12 26	169 138	143 104
South Carolina§ Virginia§	_	0	2	6	9 74	1	1	9	69 60	84 108	_	1 2	6 11	51 110	30 72
West Virginia	_	0	2	7	2	_	Ö	2	2	1	_	0	6	26	22
E.S. Central Alabama [§]	5 1	3	15 5	171 29	135 26	32 17	13 3	31 17	530 176	1,017 194		3 0	11 0	161 N	141 N
Kentucky	3	1 0	8	69	51 8	3	4 1	12	171	257	<u></u>	0	5 0	33	28
Mississippi Tennessee [§]	_	0	1 4	24	50	12	3	8 10	61 122	75 491	_	3	9	128	113
W.S. Central Arkansas	8 6	1 0	52 3	37 19	84 11	6 1	34 1	596 7	1,153 81	2,813 51	2	7 0	58 5	304 24	254 15
Louisiana	_	0	1	_	18	_	1	25	90	119	_	Ö	1	5	5
Oklahoma Texas [§]	2	0 1	8 44	18 64	22 33	<u>5</u>	3 29	286 308	100 882	524 2,119	2	2 4	14 43	81 194	93 141
Mountain Arizona	1	5 1	16 8	233 76	239 23	20 4	22 11	60 30	889 468	641 340	6	11 6	78 57	523 277	486 206
Colorado	_	1	8	87	62	10	3	18	180	108	5	3	8	112	149
Idaho [§] Montana		1 0	7 1	56 —	32 14	_	0	4 6	15 12	10 5	_	0	2	8 —	3
Nevada [§] New Mexico [§]	_	0 0	3 1	10 4	18 22	_	0 2	8 10	30 114	44 96	_	0 1	2 7	63	8 68
Utah Wyoming	5 —	1 0	13 3	103 17	61 7	5 1	1 0	6 3	62 8	33 5	1	1 0	7 1	60 3	49 3
Pacific	12	7	55	260	269	13	40	148	1,451	1,739	2	2	9	86	80
Alaska California	_	0 4	1 18	 161	9 103	_	0 32	2 104	9 1,189	11 1,488	_	0 0	0 0	_	_
Hawaii Oregon [§]		0 2	2 47	12 99	10 72	_ 1	1 2	4 31	33 110	28 112	2 N	2	9	86 N	80 N
Washington	12	1	32	87	75	12	2	43	110	100	N	0	0	N	N
American Samoa C.N.M.I.	U U	0	0	U	U U	U U	0	0	U	7 U	U U	0	0	U U	U U
Guam Puerto Rico	_	0	0	_	2	<u>_</u>	0	3 2	12	16 5		0	0	 N	N
U.S. Virgin Islands	_	0	0	_	_	<u>.</u>	0	0		_		ő	0		

C.N.M.I.: Commonwealth of Northern Mariana Islands. U: Unavailable. —: No reported cases. N: No N: Not notifiable. Cum: Cumulative year-to-date counts. Med: Median. Max: Maximum.

[†] Incidence data for reporting year 2006 is provisional.
† Includes *E. coli* O157:H7; Shiga toxin positive, serogroup non-0157; and Shiga toxin positive, not serogrouped. Contains data reported through the National Electronic Disease Surveillance System (NEDSS).

TABLE II. (*Continued*) Provisional cases of selected notifiable diseases, United States, weeks ending October 7, 2006, and October 8, 2005 (40th Week)*

(40th Week)*	Strepto	Syphilis, primary and secondary					Varicella (chickenpox)								
	Drug resistant, all ages Previous					Previous				Previous					
Reporting area	Current week	Med Med	eeks Max	Cum 2006	Cum 2005	Current week	52 we Med	eks Max	Cum 2006	Cum 2005	Current week	Med Med	veeks Max	Cum 2006	Cum 2005
United States	84	50	334	1,976	2,040	79	172	334	6,711	6,463	442	802	3,204	31,532	21,295
New England		1	24	30	176	2	4	17	157	159	16	39	144	1,147	4,043
Connecticut Maine†	U —	0 0	7 2	U 8	74 N	1	0 0	11 2	34 7	34 1	U —	0 5	58 20	U 151	1,196 240
Massachusetts New Hampshire	_	0	6 0		76 —	1	2	6 2	97 10	98 13	 10	0 7	54 47	94 376	1,816 231
Rhode Island	_	0	11	10	17	_	0	6	7	12	_	0	0	_	_
Vermont [†]	_	0	2	12	9	_	0	1	2	1	6	12	50	526	560
Mid. Atlantic New Jersey	1 N	3 0	15 0	125 N	169 N	13	21 3	35 7	855 128	801 111	84	103 0	183 0	3,659	3,605
New York (Úpstate)	_	1	10	44	65	5	2	14	117	62	_	0	0	_	_
New York City Pennsylvania	U 1	0 2	0 9	U 81	U 104	4 4	10 5	23 9	410 200	481 147	— 84	0 103	0 183	3,659	3,605
E.N. Central	7	11	41	442	510	8	17	38	678	700	121	237	587	11,255	4,425
Illinois Indiana	_ 1	0 2	3 21	15 117	26 160	1 2	8 1	23 4	313 68	393 51	_	2	7 475	68 475	78 251
Michigan	_	0	4	17	33	1	2	19	90	63	33	102	174	3,266	2,637
Ohio Wisconsin	6 N	6 0	32 0	293 N	291 N	<u>4</u>	4 1	8 4	159 48	167 26	88 —	93 12	420 52	6,816 630	1,118 341
W.N. Central	62	1	191	96	34	1	5	10	194	195	7	23	84	1,108	350
Iowa Kansas	N N	0 0	0	N N	N N	_	0 0	2 2	11 16	8 15	N	0	0 8	N 20	N
Minnesota	60	0	191	60	_	_	0	3	21	57	_	0	0	_	_
Missouri Nebraska†	2	1 0	3 0	35 —	27 2	_	3 0	8 1	130 3	110 4	7	19 0	82 0	1,006	242
North Dakota South Dakota	_	0	1 1	_ 1	2	_ 1	0	1 3	1 12	_ 1	_	0 1	25 12	44 38	20 88
S. Atlantic	13	26	53	1,035	835	25	42	186	1,619	1,584	22	90	860	3,328	1,646
Delaware	_	0	2	´ —	1	_	0	2	16	9	_	1	5	52	25
District of Columbia Florida	1 10	0 14	3 36	23 574	13 460	5 10	2 15	9 29	101 575	86 537	2	0 0	5 0	30	28 —
Georgia Maryland†	2	7 0	29 0	343	265	1	7 5	147 19	277 229	332 247	_	0	0	_	_
North Carolina	N	0	0	N	N	5	5	17	229	205	_	0	0	_	_
South Carolina† Virginia†	N	0 0	0 0	N	N	1	1 3	7 12	54 133	58 107	10	15 30	53 812	809 1,287	439 350
West Virginia	_	1	14	95	96	1	0	1	5	3	10	26	70	1,150	804
E.S. Central Alabama [†]	N	3 0	13 0	152 N	143 N	7	13 4	25 19	547 238	356 115	_	1 1	70 70	91 90	95 95
Kentucky	_	0	5	29	26	1	1	8	56	36	N	0	0	N	N
Mississippi Tennessee [†]	_	0 3	0 13	123	1 116	6	0 5	6 13	47 206	39 166	N	0	1 0	1 N	N
W.S. Central	_	0	4	17	99	12	27	43	1,153	953	134	183	1,757	8,838	5,089
Arkansas Louisiana	_	0	3 4	12 5	12 87	1 10	1 4	5 17	60 190	40 199	_	7 0	110 8	590 44	110
Oklahoma	N	0	0	N	N	1	1	6	57	29	_	0	0	_	_
Texas [†]	N	0 1	0	N 79	N 74	 10	21 7	36	846	685	134	170	1,647	8,204	4,979
Mountain Arizona	1 N	0	0	N	/4 N	7	3	25 16	322 144	332 132	58 —	54 0	138 0	2,106 —	2,042
Colorado Idaho†	N N	0 0	0	N N	N N	_	1 0	3 1	32 2	36 20	35	33 0	76 0	1,152	1,408
Montana	_	0	1	_	_	_	0	1	1	5	_	0	2	2	_
Nevada† New Mexico†	_	0 0	3 1	4 1	29 —	3	1 1	12 5	83 52	91 40	_	0 3	2 34	4 307	2 172
Utah Wyoming		0 1	8 4	34 40	23 22	_	0	1 0	8	8	23	10 0	55 8	608 33	409 51
Pacific		0	0	_	_	1	33	49	1,186	1,383	_	0	0	_	_
Alaska		0	0	_	_	_	0	4	9	6	_	0	0	_	_
California Hawaii	N —	0 0	0 0	N —	N	_	28 0	39 2	1,007 15	1,235 9	N	0 0	0 0	N	N
Oregon [†] Washington	N N	0	0	N N	N N	_ 1	0 3	6 10	13 142	26 107	N N	0	0	N N	N N
American Samoa	_	0	0	_	_	U	0	0	U	U	U	0	0	U	U
C.N.M.I. Guam	_	0	0	_	_	Ü	0	0	Ü	U 3	U	0	0 12	U	U 389
Puerto Rico	N	0	0	N	N	_	2	10	86	164	3	8	47	284	554
U.S. Virgin Islands		0	0			_	0	0			_	0	0		

Cum: Cumulative year-to-date counts.

Med: Median.

Max: Maximum.

C.N.M.I.: Commonwealth of Northern Mariana Islands.
U: Unavailable. —: No reported cases. N: Not notifiable. Cum: Cumulative year-to-t* Incidence data for reporting year 2006 is provisional.

* Contains data reported through the National Electronic Disease Surveillance System (NEDSS).

TABLE II. (Continued) Provisional cases of selected notifiable diseases, United States, weeks ending October 7, 2006, and October 8, 2005 (40th Week)*

(40th Week)*					West Nile	virus disease†						
		1	Neuroinva	sive	West Mile	il us disease		No	n-neuroin	vasive		
		Prev	ious					Prev	rious			
Reporting area	Current week	52 w Med	eeks Max	Cum 2006	Cum 2005		rrent eek	52 w Med	eeks Max	Cum 2006	Cum 2005	
United States	_	1	161	1,119	1,246		_	1	344	1,895	1,653	
New England	_	0	3	8	9	-	_	0	2	3	4	
Connecticut Maine [§]	_	0	2 0	6	4		_	0 0	1 0	2	2	
Massachusetts	_	0	1	2	4			0	1	1	2	
New Hampshire	_	0	0	_	_		_	0	0	_	_	
Rhode Island Vermont§	_	0	0 0	_	1		_	0 0	0 0	_	_	
Mid. Atlantic	_	0	6	16	47		_	0	3	6	22	
New Jersey	_	0	2	2	3	-	_	0	1	2	3	
New York (Upstate) New York City	_	0 0	1 4	7	19 11		_	0 0	1 2	3	5 3	
Pennsylvania	_	0	2	7	14		_	0	1	1	11	
E.N. Central	_	0	35	188	255	-	_	0	18	77	153	
Illinois	_	0	21	111	136		_	0	16	55	114	
Indiana Michigan	_	0 0	4 7	11 29	10 54		_	0 0	2 1	5 2	11 8	
Ohio	_	0	11	27	45		_	0	3	7	14	
Wisconsin	_	0	2	10	10	-	_	0	2	8	6	
W.N. Central	_	0	31	191	161		_	0	73	374	461	
Iowa Kansas	_	0	2 3	17 14	13 13	-	_	0 0	4 3	12 10	23 N	
Minnesota	_	0	6	29	18	-	_	0	7	34	27	
Missouri Nabraaka [§]	_	0	12	41	17		_	0 0	3 24	9	13	
Nebraska [§] North Dakota	_	0	7 5	33 20	53 12		_	0	27	123 115	131 74	
South Dakota	_	Ö	7	37	35		_	Ö	22	71	193	
S. Atlantic	_	0	3	8	29	-	_	0	3	6	26	
Delaware District of Columbia	_	0	0 0	_	1 3			0 0	1 1	_ 1	_ 1	
Florida	_	0	2	3	8			0	0		11	
Georgia	_	0	1	2	7	-	_	0	2	4	10	
Maryland§ North Carolina	_	0	1 0	2	4 2		_	0 0	1 0	1	1 2	
South Carolina§	_	0	1	_	4			0	0	_	_	
Virginia [§]	_	0	0	_	_		_	0	0	_	1	
West Virginia	_	0	1	1	_		N	0	0	N	N	
E.S. Central Alabama§	_	0	12 1	86 4	62 6		_	0 0	15 2	81 —	36 2	
Kentucky	_	Ö	i	3	4			0	1	1	_	
Mississippi	_	0	9	72	38		_	0	15	79	31	
Tennessee§	_	0	3	7	14		_	0	1	1	3	
W.S. Central Arkansas	_	1 0	55 4	283 21	249 12			0 0	25 2	147 5	145 15	
Louisiana	_	0	14	66	106	-	_	0	8	49	53	
Oklahoma Texas [§]	_	0	6 35	21 175	16 115	-	_	0 0	3 14	12 81	11 66	
Mountain	_	0	59	270	134	-		0	196	993	228	
Arizona	_	0	59 4	15	44	-	_	0	196	993 14	49	
Colorado	_	0	10	54	20		_	0	43	219	85	
Idaho§ Montana	_	0	29 3	94 10	3 8		_	0 0	128 7	542 19	10 17	
Nevada [§]	=	0	9	34	14			0	13	73	17	
New Mexico§	_	0	1	1	19		_	0	1	2	13	
Utah Wyoming	_	0	8 7	48 14	21 5			0 0	17 7	88 36	31 6	
Pacific	_	0	15	69	300		_	0	45	208	578	
Alaska	_	0	0	_	_	-	_	0	0	_	_	
California	_	0	15 0	65	299		_	0	33	164	572	
Hawaii Oregon [§]	_	0	2	4	1			0 0	0 12	42	6	
Washington	_	Ö	0			-	_	Ö	2	2	_	
American Samoa	U	0	0	U	U		U	0	0	U	U	
C.N.M.I. Guam	U —	0	0	U	U		U	0 0	0	U	U	
Puerto Rico	_	0	0	_	_	-		0	0	_	_	
U.S. Virgin Islands	_	Ö	Ō	_	_	-	_	Ö	Ō	_	_	

C.N.M.I.: Commonwealth of Northern Mariana Islands. U: Unavailable. —: No reported cases. N: No N: Not notifiable. Cum: Cumulative year-to-date counts. Med: Median. Max: Maximum.

Updated weekly from reports to the Division of Vector-Borne Infectious Diseases, National Center for Zoonotic, Vector-Borne, and Enteric Diseases (proposed) (ArboNET) Surveillance).

Scontains data reported through the National Electronic Disease Surveillance System (NEDSS).

TABLE III. Deaths in 122 U.S. cities.* week ending October 7, 2006 (40th Week)

TABLE III. Deaths in 122 U.S. cities,* week ending October 7, 2006 (40th Week) All causes, by age (years) All causes, by age (years)															
		All c	auses, b	y age (ye	ears)					All o	causes, b	y age (ye	ears)		
Reporting Area	All Ages	<u>≥</u> 65	45-64	25-44	1-24	<1	P&I [†] Total	Reporting Area	All Ages	<u>≥</u> 65	45-64	25-44	1-24	<1	P&I [†] Total
New England	516	358	115	25	14	4	41	S. Atlantic	988	612	245	75	31	25	65
Boston, MA	124	79	27	7	8	3	9	Atlanta, GA	63	40	18	3	2	_	2
Bridgeport, CT	35 7	21 5	12 2	2	_	_	1	Baltimore, MD	116 119	64	32	10	7 7	3 4	12 14
Cambridge, MA Fall River, MA	27	22	4	_	1	_	1	Charlotte, NC Jacksonville, FL	114	73 64	23 30	12 11	6	3	6
Hartford, CT	48	36	11	_	1	_	6	Miami, FL	97	69	21	4	_	3	11
Lowell, MA	25	18	5	1	1	_	i i	Norfolk, VA	51	29	11	6	2	3	2
Lynn, MA	13	10	3	_	_	_	2	Richmond, VA	42	25	14	3	_	_	3
New Bedford, MA	19	14	.5	_	_	_	_	Savannah, GA	50	31	13	5	_	1	2
New Haven, CT	44	29	11	3	_	1	6	St. Petersburg, FL	47	32	8	3	1	3	5
Providence, RI Somerville, MA	60 5	43 3	12 2	4	1	_	7	Tampa, FL Washington, D.C.	172 100	110 62	42 30	14 3	4 2	2	5 1
Springfield, MA	30	21	5	4	_	_	_	Wilmington, DE	17	13	3	1	_	_	2
Waterbury, CT	34	24	6	3	1	_	4	"							
Worcester, MA	45	33	10	1	1	_	4	E.S. Central Birmingham, AL	874 173	542 106	235 47	62 13	18 2	17 5	63 14
Mid. Atlantic	1,995	1,362	436	129	36	31	108	Chattanooga, TN	58	39	11	5	1	2	3
Albany, NY	40	26	10	1	3	_	4	Knoxville, TN	101	70	24	6	_	1	5
Allentown, PA	25	17	6	1	_	1	1	Lexington, KY	94	65	22	3	2	2	6
Buffalo, NY	62	40	17	3	2	_	6	Memphis, TN	137	82	33	14	3	5	13
Camden, NJ	19	8	7	3	_	1	_	Mobile, AL	91	59	18	11	3	_	6
Elizabeth, NJ Erie, PA	10 38	8 31	2 5	1	_	1	3	Montgomery, AL Nashville, TN	65 155	37 84	24 56	3 7	1 6	_	7 9
Jersey City, NJ	19	12	7		_		2	· ·							
New York City, NY	1,024	705	219	69	16	14	41	W.S. Central	1,239	775	307	93	29	35	47
Newark, NJ	49	19	15	8	2	5	1	Austin, TX Baton Rouge, LA	77 64	46 35	22 21	6 6	2 2	1	1 1
Paterson, NJ	26	15	8	2	_	1	1	Corpus Christi, TX	48	35	10	3	_	_	2
Philadelphia, PA	321	201	89	21	8	2	19	Dallas, TX	181	103	43	14	3	18	9
Pittsburgh, PA [§] Reading, PA	24 25	18 18	4 3	1 2	_ 1	1 1	_	El Paso, TX	74	53	13	5	2	1	3
Rochester, NY	119	85	25	6	i	2	10	Fort Worth, TX	94	63	21	3	2	5	2
Schenectady, NY	21	16	1	4		_	2	Houston, TX	344	199	99	29	11	6	11
Scranton, PA	25	23	2	_	_	_	2	Little Rock, AR New Orleans, LA ¹	71 U	41 U	20 U	4 U	3 U	3 U	U
Syracuse, NY	85	71	7	3	2	2	8	San Antonio, TX	141	90	38	11	1	1	8
Trenton, NJ	21	14	6	1	_	_	_	Shreveport, LA	55	41	9	3	2		8
Utica, NY Yonkers, NY	14 28	12 23	1 2	1 2	1	_	2 4	Tulsa, OK	90	69	11	9	1	_	2
								Mountain	966	611	233	68	27	27	60
E.N. Central Akron, OH	1,967 48	1,314 28	436 13	134 2	33 1	50 4	136 4	Albuquerque, NM	133	86	38	7	2	_	7
Canton, OH	25	19	6	_			3	Boise, ID	63	41	12	5	2	3	4
Chicago, IL	330	199	72	34	13	12	25	Colorado Springs, CO		40	6	2	_	3	2
Cincinnati, OH	101	64	27	5	1	4	10	Denver, CO Las Vegas, NV	81 244	48 164	27 57	3 16	4	1	4 17
Cleveland, OH	232	169	47	13	_	3	7	Ogden, UT	24	16	4	3	1	_	1
Columbus, OH	196	144	33	11	2	6	22	Phoenix, AZ	189	92	54	26	10	7	14
Dayton, OH Detroit, MI	119 152	84 75	30 53	3 19	1 2	1 3	8 11	Pueblo, CO	25	20	4	1	_	_	2
Evansville, IN	49	38	7	3	1	_	2	Salt Like City, UT	156	104	31	5	6	10	9
Fort Wayne, IN	36	20	12	3	1	_	1	Tucson, AZ	U	U	U	U	U	U	U
Gary, IN	11	4	4	2	1	_	_	Pacific	1,602	1,096	338	93	42	33	117
Grand Rapids, MI	45	29	11	2	_	3	5	Berkeley, CA	14	11	2	_	_	1	3
Indianapolis, IN Lansing, MI	187 44	124 34	47 9	10 1	2	4	8 2	Fresno, CA Glendale, CA	152 8	97 6	35 2	11	4	5	5 3
Milwaukee, WI	95	71	13	6	2	3	3	Honolulu, HI	66	52	8	3	_	1	5
Peoria, IL	36	24	9	1	1	1	3	Long Beach, CA	52	40	6	3	2	1	8
Rockford, IL	54	41	8	2	2	1	4	Los Angeles, CA	211	120	55	19	11	6	16
South Bend, IN	52	31	11	.5	1	4	3	Pasadena, CA	27	18	8	1	_	_	1
Toledo, OH	104	75	16	10	2	1	11	Portland, OR	130	90	29	7	3	1	5
Youngstown, OH	51	41	8	2	_	_	4	Sacramento, CA San Diego, CA	236 170	165 118	51 34	9 13	6 2	5 3	15 15
W.N. Central	566	362	127	32	19	26	34	San Francisco, CA	104	79	21	3	_	1	17
Des Moines, IA	53	36	16	1	_	_	3	San Jose, CA	161	115	28	11	3	4	11
Duluth, MN Kansas City, KS	24 26	19 18	5 5	1	_	_	2 1	Santa Cruz, CA	25	17	4	1	2	1	1
Kansas City, MO	107	70	24	5	2	6	6	Seattle, WA	89	54	21	7	5	2	5
Lincoln, NE	34	21	10	3	_	_	2	Spokane, WA	65	47	13	2	1	2	4
Minneapolis, MN	53	34	7	7	3	2	4	Tacoma, WA	92	67	21	3	1	_	3
Omaha, NE	72	51	11	4	2	4	5	Total	10,713**	7,032	2,472	711	249	248	671
St. Louis, MO	78 50	35 41	24	7	5	7	4								
St. Paul, MN Wichita, KS	59 60	41 37	12 13	1 3	3 2	2 5	3 4								
vviolina, NO			10				4	I							

U: Unavailable. —:No reported cases.

* Mortality data in this table are voluntarily reported from 122 cities in the United States, most of which have populations of ≥100,000. A death is reported by the place of its occurrence and by the week that the death certificate was filed. Fetal deaths are not included.

† Pneumonia and influenza.

§ Because of changes in reporting methods in this Pennsylvania city, these numbers are partial counts for the current week. Complete counts will be available in 4 to 6 weeks.

¶ Because of Hurricane Katrina, weekly reporting of deaths has been temporarily disrupted.

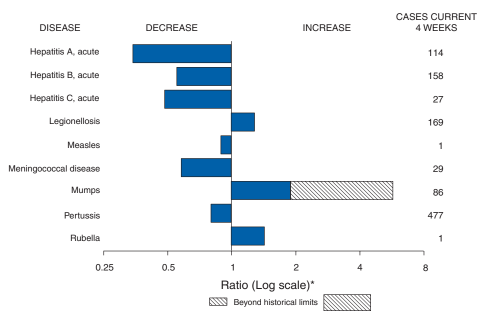
** Total includes unknown ages.

TABLE IV. Provisional cases of selected notifiable diseases,* United States, guarter ending September 30, 2006 (39th Week)

States, quarter end	ing Septemb		06 (39th V		
		Tu Pre	3		
	Current		arters	Cum	Cum
Reporting area	quarter	Min	Max	2006	2005
United States	3,090	2,384	3,589	8,100	9,661
New England Connecticut Maine Massachusetts New Hampshire Rhode Island Vermont	90 22 3 53 5 7	60 11 2 31 0 0	167 35 4 113 5 12	224 52 8 142 9 10 3	278 60 13 161 4 35 5
Mid. Atlantic New Jersey New York (Upstate) New York City Pennsylvania	505 137 61 250 57	472 101 50 232 57	605 137 110 269 102	1,501 370 172 719 240	1,491 361 192 715 223
E.N. Central Illinois Indiana Michigan Ohio Wisconsin	284 120 39 44 68 13	219 91 24 30 54 13	371 158 40 93 83 19	815 369 91 129 180 46	948 453 106 153 177 59
W.N. Central lowa Kansas Minnesota Missouri Nebraska North Dakota South Dakota	131 5 28 62 30 6 —	95 4 15 34 15 1 0	142 19 31 62 38 6 4 5	331 18 80 150 68 10 —	348 36 46 143 70 40 2 11
S. Atlantic Delaware District of Columbia Florida Georgia Maryland North Carolina South Carolina Virginia West Virginia	608 9 18 247 64 48 100 33 83 6	369 3 10 177 8 26 57 6 49 4	839 9 18 354 131 79 126 47 131 7	1,549 18 48 646 203 153 231 47 188 15	2,041 19 42 740 418 212 203 164 224 19
E.S. Central Alabama Kentucky Mississippi Tennessee	147 44 21 23 59	147 44 12 23 59	211 52 43 36 84	473 145 59 82 187	531 168 81 67 215
W.S. Central Arkansas Louisiana Oklahoma Texas	410 26 — 27 357	136 19 0 23 76	462 37 0 55 388	1,008 82 — 105 821	1,338 79 — 102 1,157
Mountain Arizona Colorado Idaho Montana Nevada New Mexico Utah Wyoming	139 90 3 — — 21 13 11	65 25 0 0 0 9 7 4	193 115 34 7 2 24 13 11	296 176 17 — 45 31 25 2	402 166 67 16 8 88 32 25
Pacific Alaska California Hawaii Oregon Washington	776 12 652 43 — 69	528 11 446 18 0 45	776 19 652 43 28 77	1,903 42 1,578 92 — 191	2,284 46 1,891 87 75 185
American Samoa C.N.M.I. Guam Puerto Rico U.S. Virgin Islands	U — 62 —	0 0 0 0	2 0 8 62 0	U — 79 —	U U 55 76 —

C.N.M.I.: Commonwealth of Northern Mariana Islands.
U: Unavailable. —: No reported cases. N: Not notifiable.
Cum: Cumulative year-to-date counts. Min: Minimum. Max: Maximum.
* AIDS and HIV/AIDS data are not updated for this quarter because of upgrading of the national HIV/AIDS surveillance data management system.

FIGURE I. Selected notifiable disease reports, United States, comparison of provisional 4-week totals October 7, 2006, with historical data



^{*} Ratio of current 4-week total to mean of 15 4-week totals (from previous, comparable, and subsequent 4-week periods for the past 5 years). The point where the hatched area begins is based on the mean and two standard deviations of these 4-week totals.

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