



MORBIDITY AND MORTALITY WEEKLY REPORT

- 901 H. influenzae Type b Disease U.S.
- 906 Self-Perceived Excellent and Very Good Health Among Blacks — Kansas, 1995
- 911 Mass Vaccination with Oral Poliovirus Vaccine Asia and Europe, 1996
- 914 Births and Deaths U.S., 1995
- 919 Adult Blood Lead Epidemiology and Surveillance — United States, 1996
- 921 Impact of HIV Protease Inhibitors on the Treatment of HIV-Infected Tuberculosis Patients with Rifampin
- 926 AIDS Rates

Progress Toward Elimination of Haemophilus influenzae Type b Disease Among Infants and Children — United States, 1987–1995

Before effective vaccines were available, *Haemophilus influenzae* type b (Hib) was the most common cause of bacterial meningitis among children in the United States, and an estimated one of 200 children aged <5 years developed invasive Hib disease (1–4). From December 1987—when Hib conjugate vaccines were introduced—through 1994, the incidence of invasive Hib disease declined 95% among children aged <5 years (4,5). Eliminating invasive Hib disease among children aged <5 years by 1996 is a goal of the Childhood Immunization Initiative (CII) (6). This report summarizes data about trends in invasive *H. influenzae* (Hi) disease during 1987–1995 from three separate surveillance systems (CDC's National Notifiable Diseases Surveillance System [NNDSS]; the National Bacterial Meningitis and Bacteremia Reporting System [NBMBRS]; and an active, multistate, laboratory-based surveillance system). The findings underscore the need for age-appropriate vaccination of infants and for complete investigation and reporting of cases of invasive Hi disease (2).

National Surveillance

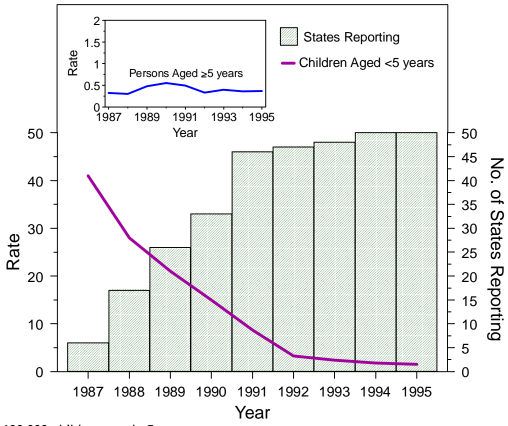
State health agencies report weekly provisional notifiable diseases data to NNDSS through the National Electronic Telecommunications System for Surveillance (NETSS) (7,8). Because the primary purpose of NNDSS is timely national surveillance, the information transmitted includes only basic demographic data about persons with invasive Hi disease. NETSS permits electronic transmission of critical supplemental information (e.g., the type of clinical illness, serotype causing disease, Hib vaccination status, and clinical outcome) for cases of Hi disease; these data were reported consistently by approximately 25 states. NBMBRS collects information about invasive bacterial diseases in the United States and includes detailed information about each case identical to the supplemental information transmitted through NETSS. Approximately 11 states participated consistently in reporting through NBMBRS.

During June–July 1996, all states were contacted to obtain additional supplemental information about Hi disease among children aged <5 years who had onset of disease in 1995. Three states confirmed that no Hi cases had been reported in children aged <5 years while the remaining 47 states, the District of Columbia, and New York City submitted additional supplemental data.

During 1987–1995, the incidence of invasive Hi disease among children aged <5 years declined 96% (from 41 cases per 100,000 children to 1.6) (Figure 1) (5). However, the incidence of Hi disease among persons aged ≥5 years remained stable during this period. Of the 317 cases of invasive Hi disease among children aged <5 years reported in 1995, serotype data were available for 201 (63%) cases. In 1995, Hib accounted for 86 (43%) isolates for which serotype was known; of these, 47 (55%) cases occurred among non-Hispanic whites, 13 (15%) among non-Hispanic blacks, five (6%) among American Indians/Alaskan Natives, two (2%) among Asians/Pacific Islanders, and 12 (14%) among Hispanics; race/ethnicity data were missing for seven (8%).

Of the 74 (86%) Hib case-patients identified in 1995 for whom information about age and vaccination status were available, 33 (45%) were unvaccinated and 41 (55%) had received at least one Hib-containing vaccine; all but three children had their first vaccination at age <7 months (Table 1). Twenty-two children were age-appropriately vaccinated before disease onset, including three children who began the series late. Among 18 children who had completed a primary series before onset of Hib disease, each had completed the two- or three-dose series with one of the three licensed vac-

FIGURE 1. Incidence rate* of invasive *Haemophilus influenzae* (Hi) disease among children aged <5 years, incidence rate[†] of invasive Hi disease among persons aged ≥5 years, and number of states reporting Hi surveillance data — United States, National Notifiable Diseases Surveillance System, 1987–1995[§]



^{*}Per 100,000 children aged <5 years.

[†]Per 100,000 persons aged ≥5 years.

[§]Because of the low number of states reporting surveillance data during 1987–1990, rates for those years were race-adjusted using the 1990 U.S. population.

TABLE 1. Number of children aged <5 years with invasive *Haemophilus influenzae* type b (Hib) disease, by age group and number of Hib vaccine doses received — United States, 1995*

Age group		No.	vaccine de		Vaccination		
(mos.)	0	1	2	3	4	status unknown	Total
0- 1	7	-	-	_	_	2	9
2- 3	9	5	_	_	_	3	17
4- 5	6	9	2	_	_	2	19
6–11	8	3	3	4	_	3	21
12–60	3	2	1	7	5	2	20
Total	33	19	6	11	5	12	86

^{*}Reported through the National Notifiable Diseases Surveillance System, the National Bacterial Meningitis and Bacteremia Reporting System, and states reporting additional supplementary information.

cines (i.e., vaccines had not been used interchangeably). Vaccine manufacturer lot numbers were available for 12 of the 18 children; among these children, no single lot was used to complete the series.

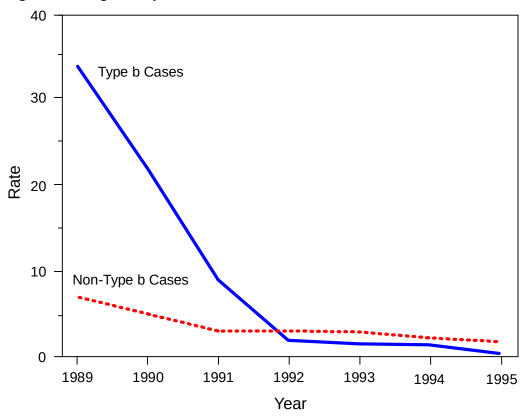
Laboratory-Based Surveillance

The active laboratory-based system coordinated by CDC includes surveillance projects which, during 1989–1994, maintained continuous surveillance of 10.5 million persons in four areas (three counties in the San Francisco Bay area, eight counties in metropolitan Atlanta, four counties in Tennessee, and the state of Oklahoma) (5). During 1995, the population under surveillance was 12.8 million persons in four areas (three counties in the San Francisco Bay area, eight counties in metropolitan Atlanta, five counties in Tennessee, and the state of Maryland). Information routinely obtained about all cases of invasive Hi disease included serotype, clinical syndrome and outcome, vaccination status, and demographic information. Because blacks were overrepresented in the surveillance population, rates were race-adjusted to the 1990 U.S. population.

During 1989–1995, the race-adjusted incidence of invasive Hib disease among children aged <5 years decreased substantially compared with the decrease in incidence of non-type b Hi disease among children (Figure 2). Among children aged <5 years, during 1989–1995 the incidence of invasive Hib cases declined 99% (from 34 cases per 100,000 in 1989 to 0.4 cases in 1995). During 1995, Hib accounted for 18% of all the Hi isolates serotyped from children aged <5 years. Information about vaccination status was available for one of the four children aged <5 years with invasive Hib disease. This infant had received one dose of vaccine, although he was aged 4.5 months at disease onset and was eligible to have received two doses. One child was too young to be vaccinated, and the other two children for whom vaccination information was not available were aged 6 and 9 months.

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FIGURE 2. Race-adjusted incidence rate* for invasive *Haemophilus influenzae* type b and non-type b disease detected through active laboratory-based surveillance[†] among children aged <5 years — United States, 1989–1995



*Per 100,000 children aged <5 years.

Editorial Note: The findings in this report document the continued decline in the incidence of invasive Hib disease among children aged <5 years in the United States—a trend associated with the increased use of Hib conjugate vaccines. These findings also affirm two of the primary barriers to eliminating invasive Hib disease among children: disease among undervaccinated children and among children too young to complete the primary series of Hib vaccination.

Identification of the serotype of cases of invasive Hi disease is essential for evaluating changes in the epidemiology of Hib disease. In the United States, the percentage of invasive Hi disease cases among children with serotype information has increased substantially, from 41% of 340 cases in 1994 to 63% of 317 cases in 1995 (5). These data suggest that the proportion of Hi cases caused by Hib has been decreasing.

Invasive Hib disease now occurs primarily among undervaccinated children and among infants too young to have completed the primary series of vaccinations. However, among children aged 19–35 months, Hib vaccination coverage with three or

[†]During 1989–1994, the surveillance area population was 10.5 million in four areas (three counties in the San Francisco Bay area, eight counties in metropolitan Atlanta, four counties in Tennessee, and the state of Oklahoma). In 1995, the surveillance area population was 12.8 million persons in four areas (three counties in the San Francisco Bay area, eight counties in metropolitan Atlanta, five counties in Tennessee, and the state of Maryland).

more doses of Hib vaccine increased substantially from the second quarter of 1994 (76%) to the second quarter of 1995 (92%) (9). Although widespread vaccination with conjugate vaccine has reduced Hib colonization rates among young children, circulation of the organism continues (1,4). Population groups with low levels of vaccination probably contribute to the ongoing occurrence of disease and regional differences in disease incidence. Consequently, all health-care providers who counsel parents about childhood vaccination should emphasize the importance of protecting infants against Hib disease and ensure that the vaccine is administered in a timely manner, especially to children of low socioeconomic status who may be more likely to be undervaccinated or unvaccinated (2,10). The small proportion of Hib cases reported through national surveillance among children who had completed a primary Hib vaccine series with vaccines from different manufacturers and with different lot numbers suggests that vaccine failure occurs infrequently.

The findings in this report indicate that the incidence of invasive Hib disease among preschool-aged children has continued to decline, consistent with the goal to eliminate invasive Hib disease among children aged <5 years. However, to attain this goal, timely and complete vaccination coverage is needed among all preschool-aged children (2). Because conjugate vaccines reduce Hib carriage and interrupt transmission of the organism, complete coverage among preschool-aged children will help to eliminate disease among infants who are too young to be completely vaccinated (1,2,4).

To monitor progress toward meeting the goal of elimination of invasive Hib disease among children aged <5 years and to evaluate changes in the epidemiology of invasive Hi disease, national surveillance for Hi should be strengthened. To optimize surveillance efforts, case reporting should include four elements. First, because Hib vaccines protect against Hi serotype b organisms only, serotyping should be performed for all cases of invasive Hi disease—state health departments are encouraged to identify laboratories in which serotyping is conducted for all Hi isolates or to send Hi isolates to CDC for serotyping. In addition, assessments of children in older age groups (5-14 years) will be needed to document persistence of protection by Hib conjugate vaccines. Second, to improve characterization of groups at risk for undervaccination and Hib disease, vaccination status of all children with invasive Hib disease should be assessed. Third, to ensure continued high levels of vaccine effectiveness and to enable systematic evaluation of factors associated with vaccine failure in persons with Hib disease, the date, vaccine manufacturer, and vaccine lot number should be included in the case report. Fourth, important indicators of the severity of Hi infections should be reported, including the type of clinical syndrome, specimen source (e.g., cerebrospinal fluid, blood, or joint fluid), and clinical outcome.

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Factors Associated with Self-Perceived Excellent and Very Good Health Among Blacks — Kansas, 1995

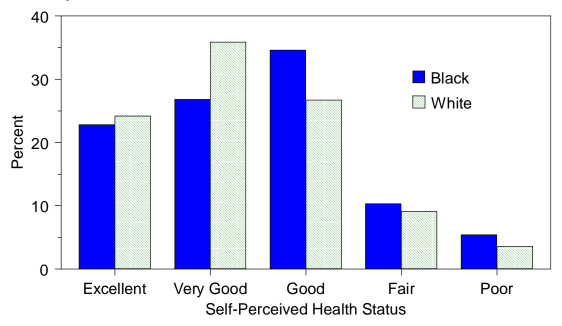
Self-perceived health is related to functional status, morbidity, and mortality and is an important measure in determining health status and health-related quality of life (HRQOL) scales (1). In 1994, the group of health professionals who established the health status indicators for *Healthy People 2000* (2) recommended that states examine the indicators for major demographic subgroups (e.g., racial/ethnic groups). The Kansas Department of Health and Environment (KDHE) analyzed data from the 1995 Behavioral Risk Factor Surveillance System (BRFSS) supplemental survey of blacks in Kansas to determine the relation between self-perceived excellent and very good health (EVGH) and physical functioning, mental functioning, role limitations, access to care, and health behaviors among blacks—the largest racial/ethnic minority group in that state. This report summarizes the findings of the analysis, which indicate that several factors related to demographics, physical functioning, and health behaviors were associated with EVGH.

BRFSS is a population-based, random-digit-dialed telephone survey of the noninstitutionalized U.S. adult population aged ≥18 years. In 1995, BRFSS was conducted in 50 states and provides population-based estimates of the prevalence of health behaviors, health-care access, and selected chronic conditions. Since 1993, BRFSS has included the option of a health status module (3-5). In Kansas, a supplemental survey to the statewide BRFSS was conducted in 1995 using all telephone prefixes having an estimated ≥10% of households self-identified as black race based on 1990 census data. Self-identified blacks aged ≥18 years (n=518; response rate=83%) participated in a structured interview that included the question, "Would you say in general your health is . . . excellent, very good, good, fair, or poor?" In addition, respondents were asked several questions about their health status, access to care, health behaviors, and demographics. All analyses were unweighted. Odds ratios (ORs) and 95% confidence intervals (CIs) were used to measure the association between categorical variables and EVGH. A chi-square test for trend was conducted to determine whether income was associated with EVGH. Factors bivariately associated with EVGH were analyzed by sex-specific logistic regression models.

Overall, the crude percentage of EVGH reported by blacks surveyed was 49% (54% of men and 47% of women); in comparison, 60% of whites surveyed in the 1995 statewide BRFSS reported EVGH (Figure 1). Annual household income <\$25,000 (OR=0.3, 95% Cl=0.2–0.4) and reporting hypertension were negatively associated with EVGH

FIGURE 1. Percentage of respondents reporting self-perceived health status, by race*

— Kansas, 1995



^{*}Data for whites were obtained from the 1995 statewide Behavioral Risk Factor Surveillance System (BRFSS) survey. Data for blacks were obtained from the 1995 supplemental survey for BRFSS.

among both black men and black women (OR=0.3, 95% Cl=0.2–0.4). The sex-specific percentage of EVGH varied by other demographic and descriptive characteristics (Table 1). Among women, factors positively associated with EVGH by bivariate analysis were working for wages (OR=2.0, 95% Cl=1.2–3.3), being aged <45 years (OR=1.9, 95% Cl=1.1–3.2), and reporting no days of poor physical health during the previous 30 days (OR=3.6, 95% Cl=2.0–6.4). Of the 10 factors negatively associated with EVGH among women by bivariate analysis, the association was strongest for having diabetes (OR=0.08, 95% Cl=0.01–0.3), having any limitations (OR=0.2, 95% Cl=0.05–0.4), and being unable to work (OR=0.2, 95% Cl=0.02–0.7) (Table 1). Among men, factors positively associated with EVGH by bivariate analysis included working for wages (OR=2.2, 95% Cl=1.2–4.2) and age <45 years (OR=2.7, 95% Cl=1.5–5.2). Of the eight factors negatively associated with EVGH among men by bivariate analysis, the association was strongest for being unable to work (OR=0.1, 95% Cl=0.0–0.8), having any limitations (OR=0.2, 95% Cl=0.05–0.4), and needing daily medications (OR=0.2, 95% Cl=0.1–0.5) (Table 1).

Because of interaction between sex, insurance status, and income with EVGH, sex-specific logistic regression models were constructed to adjust for factors independently associated with EVGH. Multivariate analysis indicated that, among women, factors negatively associated with EVGH included diabetes (OR=0.2, 95% Cl=0.04–0.9), any limitations (OR=0.3, 95% Cl=0.01–0.6), annual household income <\$25,000 (OR=0.3, 95% Cl=0.2–0.5), hypertension (OR=0.4, 95% Cl=0.2–0.8), and having smoked at least 100 cigarettes (OR=0.3, 95% Cl=0.2–0.5). Among men, those with health insurance and an annual household income ≥\$25,000 were 17 times more likely to report EVGH than those with no health insurance and an annual household income <\$25,000.

TABLE 1. Unweighted percentage of black survey respondents who reported excellent or very good health (EVGH), by sex and selected characteristics — supplemental Behavioral Risk Factor Surveillance System survey, Kansas, 1995*

		Md (n=2			Wor (n=3	
Characteristic	Sample size	(%)	(95% CI†)	Sample size	(%)	(95% CI)
	3126	(/0)	(35 /6 CI*)	3126	(/0)	(95 /6 CI)
Age group (yrs)§		()			/·	
18–24	37	(64.9)	(47.5%–79.8%)	59	(55.9)	(42.4%–68.8%)
25–44	92	(62.0)	(51.2%–71.9%)	154	(50.0)	(41.8%–58.2%)
45–64 65, 74	60	(43.3)	(30.6%–56.8%)	63	(41.3)	(29.0%–54.4%)
65–74	11	(18.2)	(2.3%–51.8%)	23	(21.7)	(7.5%–43.7%)
≥75	5	(20.0)	(0.5%–71.6%)	10	(40.0)	(12.2%–73.8%)
Education level§						
Less than high school	27	(25.9)	(11.1%–46.3%)	30	(36.7)	(19.9%–56.1%)
High school graduate	68	(50.0)	(37.6%–62.4%)	123	(37.4)	(28.8%-46.6%)
Some college	70	(62.9)	(50.5%–74.1%)	103	(54.4)	(44.3%–64.2%)
College graduate	40	(62.5)	(45.8%–77.3%)	53	(60.4)	(46.0%–73.5%)
Annual household income¶						
<\$14,999	15	(33.3)	(11.8%-61.6%)	35	(22.9)	(10.4%-40.1%)
\$15,000-\$24,999	63	(34.9)	(23.3%-48.0%)	119	(37.0)	(28.3%-46.3%)
\$25,000-\$49,999	83	(68.7)	(57.6%–78.4%)	93	(66.7)	(56.1%–76.1%)
≥\$50,000	28	(67.9)	(47.6%–84.1%)	34	(52.9)	(35.1%–70.2%)
Employment status§						
Employed	154	(60.4)	(52.2%-68.2%)	199	(54.3)	(47.1%–61.3%)
Unemployed	12	(41.7)	(15.2%–72.3%)	22	(36.4)	(17.2%–59.3%)
Homemaker	0	_	_	16	(43.8)	(19.8%–70.1%)
Student	7	(71.4)	(29.0%-96.3%)	20	(50.0)	(27.2%–72.8%)
Retired	23	(26.1)	(10.2%-48.4%)	36	(27.8)	(14.2%-45.2%)
Unable to work	9	(11.1)	(0.3%-48.2%)	16	(12.5)	(1.6%–38.3%)
Marital status§						
Married	78	(52.6)	(40.9%–64.0%)	99	(53.5)	(43.2%-63.6%)
Divorced	38	(57.9)	(40.8%–73.7%)	70	(43.6)	(32.4%–56.7%)
Widowed	9	(44.4)	(13.7%–78.8%)	31	(29.0)	(14.2%-48.0%)
Separated	10	(30.0)	(6.7%-65.2%)	18	(38.9)	(17.3%–64.3%)
Never married	60	(58.3)	(44.9%–70.9%)	85	(45.9)	(35.0%–57.0%)
Unmarried couple	10	(50.0)	(18.7%–81.3%)	6	(50.0)	(11.8%–88.2%)
Have health insurance§						
Yes	175	(53.7)	(46.0%–61.3%)	260	(45.8)	(39.6%-52.0%)
No	30	(53.3)	(34.3%–71.7%)	49	(53.1)	(38.3%–67.5%)
Needed to see a doctor but didn't because of cost					•	,
Yes	25	(48.0)	(27.8%–68.7%)	49	(28.6)	(16.6%-43.3%)
No	180	(54.4)	(46.9%–61.9%)	260	(50.4)	(44.1%–56.6%)

^{*} n=518.

[†]Confidence interval.

[§]Excludes two men and two women for whom this characteristic was missing.

[¶]Excludes 18 men and 30 women for whom either EVGH or income data were missing.

^{**} Excludes three men and one woman for whom either EVGH or smoking data were missing.

TABLE 1. Unweighted percentage of black survey respondents who reported excellent or very good health (EVGH), by sex and selected characteristics — supplemental Behavioral Risk Factor Surveillance System survey, Kansas, 1995*

		Mo (n=2			Wor (n=3	
Characteristic	Sample size	(%)	(95% CI†)	Sample size	(%)	(95% CI)
Have diabetes§						
Yes	7	(14.3)	(0.4%-57.9%)	24	(4.2)	(0.1%-21.1%)
No	198	(55.1)	(47.8%–62.1%)	285	(50.5)	(44.6%–56.5%)
Have hypertension§						
Yes	63	(31.7)	(20.6%-44.7%)	112	(28.6)	(20.4%-37.9%)
No	142	(63.4)	(54.9%-71.3%)	197	(57.4)	(50.1%-64.4%)
Have high blood cholesterol level§						
Yes	35	(42.9)	(26.3%-60.6%)	51	(31.4)	(19.1%-45.9%)
No	170	(55.9)	(48.1%–63.5%)	258	(50.0)	(43.7%–56.3%)
Take daily medications						
Yes	47	(27.7)	(15.6%-42.6%)	84	(25.0)	(16.2%-35.6%)
No	158	(61.4)	(53.3%–69.0%)	225	(55.1)	(48.4%–61.7%)
Have any activity limitation§						
Yes	47	(31.9)	(19.1%–47.1%)	78	(25.6)	(16.4%-36.8%)
Duration ≤1 yr	22	(45.4)	(22.4%–67.8%)	34	(38.2)	(22.2%-56.4%)
2–4 yrs	10	(40.0)	(12.2%–73.8%)	20	(15.0)	(3.2%–37.9%)
≥ <i>5 yrs</i>	15	(6.7)	(0.2%–31.9%)	24	(16.7)	(4.7%–37.4%)
No	158	(60.1)	(52.0%–67.8%)	231	(54.1)	(47.5%–60.7%)
Have smoked ≥100 cigarettes during lifetime**						
Yes	111	(50.5)	(40.8%-60.1%)	188	(36.1)	(29.3%-43.5%)
No	93	(58.1)	(47.4%–68.2%)	122	(54.3)	(44.8%–63.2%)

^{*} n=518.

Multivariate analysis indicated that factors negatively associated with EVGH among men included the duration of activity limitations in years (OR=0.8, 95% Cl=0.7–1.0) and hypertension (OR=0.4, 95% Cl=0.2–0.9).

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[†]Confidence interval.

[§]Excludes two men and two women for whom this characteristic was missing.

[¶]Excludes 18 men and 30 women for whom either EVGH or income data were missing.

^{**}Excludes three men and one woman for whom either EVGH or smoking data were missing.

racial/ethnic minority groups. In addition to this assessment of blacks, KDHE has implemented a similar supplemental BRFSS survey of Hispanics (the second largest and most rapidly increasing racial/ethnic minority group in the state).

The finding that self-perceived EVGH was lower among blacks than among whites is consistent with mortality patterns in Kansas, which, when compared with those of whites, indicate the average age at death for black men and black women is 12.5 years younger and 13.4 years younger, respectively. The positive association between EVGH and an annual household income ≥\$25,000 for both black men and black women underscores previous studies indicating the impact of economics on self-perceived health status (6). In addition, the findings suggest that efforts to improve the health status of blacks and, therefore, self-perceived health and HRQOL, should be directed toward preventing activity limitations and hypertension among men and toward preventing diabetes, smoking initiation, and activity limitations among women.

The findings in this report are subject to at least five limitations. First, temporal variations in the relation between variables associated with EVGH have not been characterized, and the potential for increasing self-perceived EVGH by altering these factors is unknown. Second, many of the biologically and psychologically plausible determinants of EVGH (e.g., chronic conditions such as heart disease, cancer, and arthritis) were not associated with EVGH, probably reflecting the small sample size or misclassification. Third, the findings of the study may not be generalizable to all blacks in the state because black households without telephones and those in areas of the state with <10% of the households self-identified as black were not eligible for survey selection. Fourth, identification of self-perceived health status may vary by cultural groups, and comparisons of that status across groups may reflect cultural differences rather than health status. Finally, the factors studied were chosen based on hypotheses, some limited literature, and available data and were not identified by the respondents as factors important to them and their health status.

The relation between the duration of activity limitations and EVGH among black men in Kansas suggests the need to better characterize the types and causes of activity limitations among this group to enable development of appropriate interventions. The findings also suggest that, for some women, strategies for improving self-perceived health might include programs for diabetes self-management, education of health-care providers, prevention of smoking initiation, and promotion of general health. KDHE, in collaboration with an advisory group to the BRFSS supplemental survey of blacks, is implementing interventions to improve HRQOL among black residents of Kansas. Examples of such efforts include extended diabetes outreach and education to black women and the involvement of churches to reach both men and women with hypertension. KDHE is planning future BRFSS surveys to assess the impact of these programs.

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Update: Mass Vaccination with Oral Poliovirus Vaccine — Asia and Europe, 1996

In 1995, a group of 18 geographically contiguous countries in Europe, Central and South Asia, and the Middle East cooperated in conducting coordinated National Immunization Days (NIDs)* with oral poliovirus vaccine (OPV) (1). The World Health Organization (WHO) had designated this effort "Operation MECACAR" (MEditerranean, CAucasus, and Central Asian Republics)[†]. Operation MECACAR was repeated in 1996 with the addition of the Russian Federation (Figure 1). This report describes OPV coverage achieved during the mass vaccination campaign in 1996 and summarizes the impact of the NIDs on poliomyelitis incidence during 1995 and the first 6 months of 1996.

To maximize the geographic area covered and the number of children targeted simultaneously for mass vaccination campaigns with OPV, adjoining countries in Europe (Armenia, Azerbaijan, Bulgaria [1995 only], Georgia, Russia [starting in 1996], and Turkey), Central Asia (Kazakstan, Kyrgyzstan, Tajikistan, Turkmenistan, and Uzbekistan), South Asia (Afghanistan, Iran, and Pakistan), and the Middle East (Iraq, Jordan, Lebanon, Syria, and West Bank and Gaza) conducted synchronized NIDs during 1995 and 1996 and are planning to conduct NIDs in 1997. Based on the desirability of scheduling mass vaccination campaigns during the low polio incidence season, all rounds of NIDs were scheduled during March–May. The only exception was Pakistan, which conducted NIDs in December 1995 and January 1996 to synchronize efforts with India (2) and China (3).

A total of 62 million children were targeted to receive two doses each of OPV, including 16 million children aged <4 years in Europe and Central Asia (the European Region of WHO [EURO]) and 46 million children aged <5 years in South Asia and the Middle East (the Eastern Mediterranean Region of WHO [EMRO]). Reported coverage was 92%–99% for each round in EURO and 84%–100% in EMRO (Table 1).

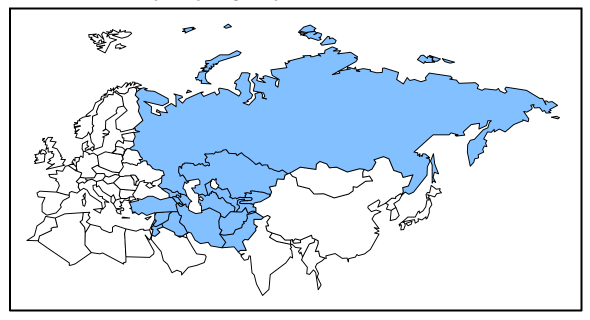
The mass vaccination efforts were followed by declines among participating countries in the incidence of polio reported to EURO and EMRO. In EURO, 50 cases of polio were reported in 1995, compared with 201 cases in 1994. For the first 6 months of 1996, a total of 13 cases of polio were reported from participating countries in EURO (all from Russia and Turkey). In EMRO, 647 cases of polio were reported from participating countries in 1995, compared with 691 cases in 1994. For the first 6 months of

^{*}Mass campaigns over a short period (days to weeks) in which two doses of oral poliovirus vaccine are administered to all children aged <5 years, regardless of prior vaccination history, with an interval of 4–6 weeks between doses.

[†]Operation MECACAR is supported by a coalition of organizations that include WHO, United Nations Children's Fund (UNICEF), other bilateral and multilateral organizations, and Rotary International.

Oral Polio Vaccine — Continued

FIGURE 1. Countries participating in Operation MECACAR — 1996



1996, a total of 60 cases of polio were reported from these countries in South Asia and the Middle East (Table 1).

Reported by: Regional Office for Europe, Copenhagen; Regional Office for Eastern Mediterranean Region, Alexandria; Global Programme for Vaccines and Immunization, World Health Organization, Geneva, Switzerland. Respiratory and Enterovirus Br, Div of Viral and Rickettsial Diseases, National Center for Infectious Diseases; Polio Eradication Activity, National Immunization Program, CDC.

Editorial Note: In 1988, the World Health Assembly, the governing body of WHO, resolved to eradicate polio from the world by the year 2000 (4); efforts to achieve this goal have progressed rapidly in many regions. For example, in 1994, the Western Hemisphere was certified as free of wild poliovirus (5), and substantial progress has been achieved in the rest of the world (6). These successful efforts have been associated with the implementation of recommended strategies for polio eradication, which include 1) achieving and maintaining high routine vaccination coverage with OPV; 2) instituting sensitive surveillance systems for polio; 3) introducing NIDs; and 4) conducting other supplemental vaccination activities (e.g., mopping-up operations). In 1995, approximately 300 million children aged <5 years (an estimated one half of the world's children in this age group) received OPV administered during NIDs (6). By the end of 1996, all polio-endemic countries of Europe and Asia will have conducted NIDs, and 29 countries in the African Region of WHO will implement NIDs (n=26) or Sub-National Immunization Days (SNIDs) (n=three) (6).

The collaboration among the 18 countries in EURO and EMRO for polio eradication has included countries characterized by different political systems and economies and racial/ethnic and religious diversity. In 1996, as a result of a large polio outbreak in Chechnya in 1995 (7) and suboptimal routine vaccination coverage in some states, Russia joined the coalition of countries comprising Operation MECACAR. Bulgaria conducted successful NIDs in 1995 and increased coverage especially among Gypsy populations, but did not participate in Operation MECACAR in 1996. In Albania, NIDs

Oral Polio Vaccine — Continued

TABLE 1. Percentage of oral poliovirus vaccine (OPV) coverage achieved during National Immunization Days* in 1996, and reported poliomyelitis cases during 1994–June 1996, by World Health Organization (WHO) region and countries participating in Operation MECACAR

	19	96						
WHO region/	% Co	verage	No. reported polio cases					
Country	Round 1	Round 2	1994	1995	1996 [†]			
EURO§								
Armenia	99%	98%	5	3	0			
Azerbaijan	97%	98%	16	5	0			
Georgia	92%	95%	0	0	0			
Kazakhstan	97%	99%	4	1	0			
Kyrgyzstan	98%	99%	0	0	0			
Russian Federation	99%	99%	5¶	154¶	3			
Tajikistan	95%	99%	26	0	0			
Turkey	93%	96%	27	32	10			
Turkmenistan	99%	99%	6	8	0			
Uzbekistan	98%	98%	117	1	0			
Total			201	50	13			
EMRO**								
Afghanistan	87%	84%	NR ^{††}	NR	NR			
Iran	99%	100%	93	101	15 ^{§§}			
Iraq	98%	98%	63	34	3			
Jordan ^{¶¶}	>100%	>100%	4	0	0			
Lebanon	99%	95%	2	0	0			
Pakistan ^{¶¶}	>100%	>100%>	527	508	42			
Syria ^{¶¶}	>100%	>100%>	2	4	0			
West Bank								
and Gaza¶	>100%	>100%>	NR	0	0			
Total			691	647	60			

^{*}Mass campaigns over a short period (days to weeks) in which two doses of OPV are administered to all children aged <5 years, regardless of prior vaccination history, with an interval of 4–6 weeks between doses.

were conducted during April and May 1996. In the fall of 1996, NIDs will be conducted in Moldova and Ukraine, while Romania and Serbia and Montenegro will conduct SNIDs.

The primary purpose of mass vaccination campaigns with OPV is to interrupt the transmission of wild poliovirus. These mass vaccination efforts have interrupted wild poliovirus transmission in many areas of Europe, as reflected by the rapid decrease in cases in EURO following the efforts conducted in 1995 and 1996; of the 50 countries in EURO, only four (Albania [8], Russia, Turkey, and Ukraine) have reported polio cases

[†]January–June.

[§]Comprising countries in Europe and Central Asia.

Russia joined Operation MECACAR in 1996; cases reported in 1994 and 1995 are excluded from the regional total.

^{**}Comprising countries in South Asia and the Middle East.

^{††}Not reported.

^{§§} January–May.

[¶]Coverage levels above 100% result from vaccinating children outside the target age group (numerator problem) and uncertainties about the exact target age group (denominator problem).

Oral Polio Vaccine — Continued

in 1996. Despite some dramatic successes in EMRO (Iran, Jordan, Lebanon, and West Bank and Gaza), other countries (Afghanistan, Iraq, Pakistan, and Syria) reported more limited progress in 1995. For example, Pakistan continued to report more than half (64%) of the regional total number of polio cases in EMRO, and Syria detected four polio cases associated with isolation of wild poliovirus. However, preliminary data for January–June 1996 suggest a substantial decline in the number of reported cases in both EURO and EMRO countries (Table 1).

The technical basis for achieving worldwide polio eradication already exists; however, insufficient political will, inadequate funding, and other barriers will need to be addressed to ensure that polio will be eradicated by the year 2000.

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Preliminary Data on Births and Deaths — United States, 1995

Timely and accurate health data are essential to public health surveillance efforts for monitoring trends in vital events, diseases, injuries, and disabilities. More timely release of accurate vital statistics has been identified as a priority by health agencies at the federal, state, and local levels and in academia and the private sector. In response to the need for faster release of high-quality data, CDC's National Center for Health Statistics (NCHS), in collaboration with state vital statistics offices, has initiated a new system to speed the transmission of vital statistics from states to CDC. This initiative has resulted in the availability of near-final natality and mortality data approximately 1 year before final data usually are released. This report presents the selected findings of an analysis of preliminary birth and death records for 1995. The number of births in the United States declined for the fifth consecutive year to an estimated 3,900,089 in 1995, 1% fewer than the final 1994 total of 3,952,767 (Table 1) (1). In addition, the estimated number of deaths in 1995 totaled 2,312,180 (Table 2), 1% more than the previous record high of 2,278,994 in 1994 (3).

Detailed natality statistics, which previously have not been available before release of the final data, are based on 90% of the births registered in 1995. Mortality statistics are based on up to 90% of deaths occurring in 1995; the only mechanism previously used for releasing provisional detailed mortality statistics before the release of final data was the Current Mortality Sample, based on a 10% sample of deaths. The death

TABLE 1. Total number of births, percentage of births with selected demographic and health characteristics, and birth rates by maternal age, by race* and ethnicity of mother — United States, final 1994 data and preliminary 1995 data

	W	nite	Bla	ack	Hisp	anic [†]	To	tal [§]
Births	1994	1995	1994	1995	1994	1995	1994	1995
Number	3,121,004	3,105,315	636,391	598,558	665,026	671,849	3,952,767	3,900,089
				Perce	ntage			
Births to mothers aged								
<20 years	11.3%	11.5%	23.2%	23.2%	17.8%	18.0%	13.1%	13.2%
Births to unmarried								
mothers	25.4%	25.3%	70.4%	69.5%	43.1%	40.8%	32.6%	32.0%
Low birthweight¶	6.1%	6.2%	13.2%	13.0%	6.2%	6.3%	7.3%	7.3%
Births delivered								
by cesarean	21.2%	20.8%	21.8%	21.8%	20.5%	20.1%	21.2%	20.8%
Prenatal care beginning								
during first trimester	82.8%	83.5%	68.3%	70.3%	68.9%	70.4%	80.2%	81.2%
Maternal age				Rate	es **			
10-14 years	0.8	0.8	4.6	4.2	2.7	2.7	1.4	1.3
15–19 years	51.1	50.3	104.5	95.5	107.7	106.2	58.9	56.9
20–24 years	106.2	106.6	146.0	136.5	188.2	186.9	111.1	110.0
25–29 years	115.5	115.2	104.0	97.7	153.2	151.8	113.9	112.4
30–34 years	83.2	84.7	65.8	63.4	95.4	94.2	81.5	82.5
35–39 years	33.7	34.3	28.9	28.3	44.3	43.9	33.7	34.1
40–44 years	6.2	6.3	5.9	5.9	10.7	10.5	6.4	6.5
45–49 years	0.3	0.3	0.3	0.2	0.6	0.5	0.3	0.3
15–44 years ^{††}	64.9	64.5	76.9	71.7	105.6	103.7	66.7	65.6

^{*}The full preliminary report provides birth rates for American Indians/Alaskan Natives and Asians/Pacific Islanders (3); percentages for these races by characteristics will be available in the final report.

† Persons of Hispanic origin may be of any race.

§ Includes races other than white and black.

¶ Birthweight of <2500 grams (<5 lb 8 oz).

**Per 1000 population in specified group.

†† Rates computed by relating total births, regardless of age of mother, to women aged 15–44 years.

TABLE 2. Preliminary number of deaths and death rates for the 15 leading causes of death for all ages, races, and sexes, by rank — United States, 1995

Rank*	Causes of death (ICD-9 [†] code)	No.	Crude death rate [§]	Age-adjusted death rate¶	% Change 1994 to 1995**
1	Diseases of heart (390–398, 402, 404–429)	738,781	281.2	138.2	- 1.6
2	Malignant neoplasms, including neoplasms of lymphatic and				
_	hematopoietic tissues (140–208)	537,969	204.7	129.8	- 1.3
3	Cerebrovascular diseases (430–438)	158,061	60.2	26.7	8.0
4	Chronic obstructive pulmonary diseases and allied conditions				
	(490–496)	104,756	39.9	21.2	1.0
5	Accidents and adverse effects				
	(E800-E949) ^{††}	89,703	34.1	29.2	- 3.6
6	Pneumonia and influenza (480–487)	83,528	31.8	13.0	0
7	Diabetes mellitus (250)	59,085	22.5	13.2	2.3
8	Human immunodeficiency virus	•			
	infection (042–044)§§	42,506	16.2	15.4	0
9	Suicide (E950–E959)	30,893	11.8	11.0	- 1.8
10	Chronic liver disease and cirrhosis	,			
	(571)	24,848	9.5	7.5	- 5.1
11	Nephritis, nephrotic syndrome, and	,		-	-
	nephrosis (580–589)	23,845	9.1	4.4	2.3
12	Homicide and legal intervention	•			
	(E960–E978)	21,577	8.2	8.8	-14.6
13	Septicemia (038)	21,123	8.0	4.1	2.5
14	Alzheimer's disease (331)	20,415	7.8	2.7	8.0
15	Atherosclerosis (440)	16,781	6.4	2.3	0
	All Causes	2,312,180	880.0	503.7	- 0.7

^{*}Based on number of deaths.

[†] International Classification of Diseases, Ninth Revision.

§ Per 100,000 population.

¶ Per 100,000 U.S. standard million population.

** Percentage change between 1995 estimated age-adjusted death rates and 1994 final age-adjusted death rates.

†† When a death occurs under "accidental" circumstances, the preferred term within the public health community is "unintentional". injury."

These codes are not part of ICD-9, but were introduced by CDC's National Center for Health Statistics for classifying and coding human immunodeficiency virus infection (2).

certificate data are processed in two parts: estimates of demographic characteristics are based on approximately 90% of 1995 deaths and medical (cause-of-death) estimates are based on approximately 80% of deaths. CDC receives independent monthly counts of birth and death records from state vital statistics offices. To produce the estimates in this report, the records from the preliminary files were weighted to these independent counts of births, infant deaths, and total deaths registered during 1995. Differences between the preliminary estimates and the final mortality data are likely to be greatest for causes for which reporting of deaths was delayed (e.g., when the case was referred to a coroner or medical examiner for investigation).

Births

In 1995, the preliminary birth rate for teenagers (56.9 births per 1000 females aged 15-19 years) declined 3% from 1994 and sustained a decline since 1991. Rates declined up to 3% for white, American Indian/Alaska Native, Asian/Pacific Islander, and Hispanic teenagers, and 9% for black teenagers. From 1994 to 1995, declines also occurred in the number and proportion of births to unmarried mothers—the number declined 3% to an estimated 1,248,028, and the proportion from 32.6% to 32.0%. For the first time since 1976, the birth rate for unmarried women aged 15-44 years declined—from 46.9 per 1000 in 1994 to 44.9 in 1995, a 4% decline. In addition, 1995 was the first year since 1940 (when national statistics first became available) during which concurrent declines occurred in the number, rate, and proportion of births to unmarried women. Compared with 1994, the 1995 proportions of births to unmarried women for whites and blacks declined by approximately 1%, and for Hispanics, by 5%. In 1995, the incidence of low birthweight (birthweight of <2500 grams [<5 lb 8 oz]) was unchanged from 1994 (7.3%); the proportion of births by cesarean delivery (20.8%) declined for the sixth consecutive year, and the proportion of mothers beginning prenatal care during the first trimester (81.2%) increased for the sixth consecutive year.

Deaths

In 1995, although the preliminary crude death rate (880.0 deaths per 100,000 population) increased slightly from 1994 (875.4), the age-adjusted death rate* was a record low (503.7). The overall estimated life expectancy in 1995 (75.8 years) increased slightly from 1994 (75.7) and equaled the record high for 1992. Record highs were reached for black females (74.0 years), black males (65.4), and white males (73.4). The life expectancy for white females was 79.6 years, unchanged from the previous year and slightly below the record high (79.8) in 1992.

From 1994 to 1995, the age-adjusted death rate for the two leading causes of death continued to decline—heart disease mortality by 1.6% and cancer mortality by 1.3%. Preliminary age-adjusted death rates also declined for homicide by 14.6%, chronic liver disease and cirrhosis by 5.1%, and mortality attributed to accidents[†] (including motor-vehicle and other injuries) by 3.6%. For the first time since human immunodeficiency virus (HIV) infection was included in U.S. death statistics, the age-adjusted

^{*}Age-adjusted death rates adjust for differing age distributions of population groups and are more effective for comparing relative risks for mortality among groups and over time. They should be used as relative indexes rather than as actual measures of risk. The age-adjusted rates were computed using the U.S. standard million population.

[†]When a death occurs under "accidental" circumstances, the preferred term within the public health community is "unintentional injury."

rates for HIV-related deaths—the leading cause of death among persons aged 25–44 years—did not increase. Age-adjusted death rates declined for suicides, injuries from firearms, drug and alcohol-induced causes, and workplace-related injuries. Age-adjusted death rates increased from 1994 to 1995 for diabetes by 2.3% and for Alzheimer's disease by 8.0%.

From 1994 to 1995, declines occurred in overall infant mortality, neonatal mortality (age <28 days), and postneonatal mortality (age 28 days through 11 months) for both white and black infants. The preliminary infant mortality rate in 1995 (7.5 deaths per 1000 live births) was lower than in 1994 (8.0).

Reported by: Div of Vital Statistics, National Center for Health Statistics, CDC.

Editorial Note: The data from the new system for releasing vital statistics provides an earlier indication of potential shifts in trends and has important ramifications for planning public health program policies and strategies. Preliminary data about live births and deaths for 1995 are available almost a year in advance of the release of the final data. CDC plans to improve the timeliness by releasing statistics based on these preliminary files twice a year (4). Each release will be based on data for a 12-month period beginning in either January or July. Statistics for January 1995–December 1995 were released on October 4, 1996. The next release, scheduled for April 1997, will cover July 1995–June 1996.

In previous years, full-year provisional mortality estimates released by NCHS have been based on a 10% sample of death certificates. However, the new data system combines expedited electronic transmittal of data from the states with more rapid data processing within NCHS to make available preliminary data for the full year based on 80%–90% of records. Although the preliminary data are based on substantial samples of births and deaths, statistics based on the final data will differ from the preliminary in some cases. In particular, the final 1995 infant mortality rate probably will exceed the rate based on preliminary data but will remain less than the 1994 rate (8.0). In addition, estimates based on final data for certain causes of death (e.g., homicide and unintentional injury) may be higher than estimates based on preliminary data.

Changes in methodologic procedures also can contribute to year-to-year differences. For example, approximately half of the 1995 decline in unmarried childbearing is the result of changes in reporting procedures in California, which particularly affected births to Hispanic women. Beginning in 1995, reporting procedures in California more accurately ascertain the marital status of Hispanic mothers than in 1994 and prior years. However, even when these reporting changes were taken into account, the 1995 decline is significant. Changes in diagnostic practices also may have accounted for the increase in the death rate for Alzheimer's disease. Finally, the earlier release of more accurate natality and mortality data should assist efforts to prolong and improve the quality of life and to prevent disease, injury, and premature death.

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Adult Blood Lead Epidemiology and Surveillance — United States, Second Quarter, 1996

CDC's National Institute for Occupational Safety and Health Adult Blood Lead Epidemiology and Surveillance program (ABLES) monitors laboratory-reported elevated blood lead levels (BLLs) among adults in the United States. This report presents ABLES data through the second quarter of 1996 and compares these data with the second quarter of 1995.

During April 1–June 30, 1996, the 6305 reports of BLLs \geq 25 µg/dL represented a 7% decrease from the 6782 reports for the second quarter of 1995 (1), which now include previously unpublished data for Minnesota and an estimate for Ohio. For the first 2 quarters of 1996, the number of reports of BLLs \geq 25 µg/dL decreased by 9% compared with the number reported for the first 2 quarters of 1995 (1), which also include previously unpublished data for Minnesota and an estimate for Ohio (Table 1). The cumulative number of reports in 1996 decreased at each reporting level compared with data for 1995. This overall trend of decreasing reports is consistent with the first quarter report for 1996 (2).

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Adult Blood Lead Epidemiology and Surveillance — Continued

TABLE 1. Number of reports of elevated blood lead levels (BLLs) among adults, number of adults with elevated BLLs, and percentage change in number of reports — 25 states,* second quarter, 1996

Reported BLL (μg/dL)	Second qu	arter, 1996	Cumulative	Cumulative	% Change
	No. reports [†]	No. persons§	reports, 1995 [¶]	reports, 1996	1995–1996
25–39	5,024	3,508	10,527	9,978	- 5%
40-49	959	674	2,697	2,111	-22%
50-59	224	159	554	431	-22%
≥60	98	55	236	200	-15%
Total	6,305	4,396	14,014	12,720	- 9%

^{*}Alabama, Arizona, California, Connecticut, Illinois, Iowa, Maine, Maryland, Massachusetts, Michigan, Minnesota, New Hampshire, New Jersey, New York, North Carolina, Ohio, Oklahoma, Oregon, Pennsylvania, South Carolina, Texas, Utah, Vermont, Washington, and Wisconsin.

Editorial Note: The findings in this report suggest that exposure to lead may be decreasing; however, variation in national quarterly reporting totals may result from 1) changes in the number of participating states; 2) changes in staffing and funding in state-based surveillance programs; and 3) interstate differences in worker BLL testing by lead-using industries. ABLES data also may be underreported when compared to estimates of the number of adults exposed to lead derived from the Third National Health and Nutrition Examination Survey (3).

The findings in this report document the continuing hazard of work-related lead exposures as an occupational health problem in the United States. ABLES enhances surveillance for this preventable condition by expanding the number of participating states, reducing variability in reporting, and distinguishing between new and recurring elevated BLLs in adults.

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[†]Data for Alabama were missing; first quarter 1995 data were used as an estimate.

[§]Individual reports for persons are categorized according to the highest reported BLL for the person during the given quarter. Pennsylvania provides the number of reports but no information on persons. The data on persons from Pennsylvania included in this table are estimates based on the proportions from the other 24 states combined and the number of reports received from Pennsylvania. Data for Alabama were missing; first quarter 1995 data were used as an estimate.

Data for Minnesota and Ohio are included for the first time in addition to previously published 1995 totals (1). For Minnesota, first and second quarter data for 1995 were used; for Ohio, first and second quarter data for 1996 were used as an estimate.

Notice to Readers

Clinical Update: Impact of HIV Protease Inhibitors on the Treatment of HIV-Infected Tuberculosis Patients with Rifampin

In 1995 and 1996, the Food and Drug Administration (FDA) approved three products in the new protease inhibitor class of drugs—saquinavir (InviraseTM), ritonavir (NorvirTM), and indinavir (CrixivanTM). Another drug in this class of agents, nelfinavir (ViraceptTM) (Agouron Pharmaceuticals), is expected to be available soon from the manufacturer through an expanded-access program. All four drugs, which inhibit HIV protease and thus interfere with viral maturation and replication, are the most potent antiretroviral agents available to treat patients with HIV disease (1). However, these protease inhibitors interact with rifamycin derivatives, such as rifampin and rifabutin, which are used to treat and prevent the mycobacterial infections commonly observed in HIV-infected patients. Rifamycins accelerate the metabolism of protease inhibitors (through induction of hepatic P450 cytochrome oxidases), resulting in subtherapeutic levels of the protease inhibitors. In addition, protease inhibitors retard the metabolism of rifamycins, resulting in increased serum levels of rifamycins and the likelihood of increased drug toxicity (2). This report describes approaches for managing patients who are candidates for or who are undergoing protease inhibitor therapy when tuberculosis (TB) is diagnosed and presents interim recommendations for managing these patients until additional data are available and formal guidelines are issued.

BACKGROUND

Rifampin is an essential component of the currently recommended regimen for treating TB (3). This regimen is efficacious in treating HIV-infected patients with TB and consists of isoniazid and rifampin for a minimum of 6 months, plus pyrazinamide and either ethambutol or streptomycin for the first 2 months (4,5). Therefore, the pharmacokinetic interactions between protease inhibitors and rifampin are important for health-care workers involved in TB control and the care of patients co-infected with TB and HIV because clinicians may decrease or restrict the use of rifampin in the treatment of patients who are candidates for therapy with both protease inhibitors and rifampin. Prompt initiation of appropriate drug therapy for patients with HIV infection who acquire TB is critical because TB may be rapidly fatal (6). Drug therapy is a critical personal health measure for curing TB and minimizes the impact of this disease on the progression of HIV infection; in addition, drug therapy is a major public health measure for interrupting the transmission of *Mycobacterium tuberculosis* to persons in the community.

Currently, the manufacturers' product labeling for protease inhibitors contraindicates, does not recommend, or discourages the concurrent administration of rifampin and protease inhibitors. Because of the common association of TB and HIV infection, an increasing number of patients probably will be considered candidates for rifampin and protease inhibitors. The management of these patients is complex, requires an individualized approach, and should be undertaken only by or in consultation with an expert. In addition, all HIV-infected patients at risk for TB infection should be carefully evaluated and administered isoniazid for preventive treatment if indicated, regardless of their status for being prescribed protease inhibitor therapy.

MANAGEMENT OPTIONS

TB Management for Patients for Whom Protease Inhibitor Therapy Is Being Considered but Has Not Yet Been Initiated

For HIV-infected patients diagnosed with drug-susceptible TB and for whom protease inhibitor therapy is being considered but has not been initiated, the suggested management strategy is to complete TB treatment with a regimen containing rifampin before starting therapy with a protease inhibitor. The duration of this anti-TB regimen is at least 6 months, and therapy should be administered following current guidelines published by the American Thoracic Society and CDC (3), including the recommendation to carefully assess clinical and bacteriologic response in patients co-infected with HIV and to prolong treatment if response is slow or suboptimal. Antiretroviral agents other than protease inhibitors may be used concurrently with this regimen. Directly observed therapy (DOT) is routinely recommended for the treatment of TB to ensure adherence with the recommended regimen and is available through local health department TB-control programs. Among patients who adhere to therapy, four-drug regimens are expected to be effective even in those infected with strains of M. tuberculosis resistant to isoniazid or streptomycin alone. However, the management of patients with drug-resistant TB should be evaluated on a case-by-case basis and individualized based on the results of drug-susceptibility studies.

TB Management Options for Patients Undergoing Protease Inhibitor Therapy

There are three options for managing HIV-infected patients with TB who are undergoing protease inhibitor therapy when TB is diagnosed. One option is to discontinue therapy with the protease inhibitor while undergoing a TB treatment regimen that includes rifampin. However, because of the potential that interruptions in the administration of the prescribed protease inhibitor can induce HIV resistance to the protease inhibitor and possibly to other drugs within the protease inhibitor class (1) and because discontinuation of protease inhibitor therapy may be associated with a detrimental effect on the patient's clinical status, some clinicians may be reluctant to discontinue protease inhibitor therapy for the duration of TB treatment. In such cases, two additional options may be considered. Because the risks and benefits of all these options are unknown, clinicians should individualize management decisions on a case-by-case basis to provide optimal patient care.

Option I. This option involves discontinuing therapy with the protease inhibitor and completing a short (minimum 6 months) course of TB treatment with a regimen containing rifampin. This anti-TB regimen should be administered following current guidelines published by the American Thoracic Society and CDC (3), and the duration of therapy should be prolonged in patients with slow or suboptimal responses. Protease inhibitor therapy may be resumed when treatment with rifampin is discontinued. Antiretroviral agents other than protease inhibitors may be used concurrently with rifampin. Although the risks associated with a complete discontinuation of protease inhibitor therapy while undergoing TB treatment are unclear, they may be serious; however, the risks and complications associated with TB treatment regimens that do not include rifampin are known. Potential consequences include prolonged duration of therapy to at least 18–24 months, increased likelihood of treatment failure and mortality (7,8), slower conversion of sputum culture to negative with patients remaining infectious for longer periods of time, and the adverse effect of TB disease on the

progression of HIV infection (9,10). Therefore, nonrifampin-containing regimens are not recommended for the treatment of rifampin-susceptible TB.

Option II. To minimize the interruption of protease inhibitor therapy, one option is to use a four-drug TB treatment regimen that includes rifampin (i.e., daily isoniazid, pyrazinamide, rifampin, and ethambutol or streptomycin) for a minimum of 2 months and until bacteriologic response is achieved (i.e., sputum conversion to culture-negative status), and the results from susceptibility testing are available. After bacteriologic response and drug susceptibility have been documented (usually 3 months), treatment may be modified to a 16-month continuation-phase regimen consisting of isoniazid (15 mg/kg) and ethambutol (50 mg/kg) two times per week. This regimen allows the reintroduction of protease inhibitor therapy. Some experts consulted for this report recommended adding a third drug, such as streptomycin, during this continuation phase if the infecting organism is not resistant to the drug. Option II cannot be recommended for patients with proven isoniazid-resistant TB.

Option III. The other management option is to continue protease inhibitor therapy with indinavir (800 mg every 8 hours) and administer a four-drug, 9-month TBtreatment regimen containing daily rifabutin (150 mg/day) instead of rifampin. When this option is used for TB management, clinicians should conduct careful monitoring, possibly including measuring serum concentrations of rifabutin—a service available only in specialized centers in the United States. This alternative TB therapy is recommended based on the pharmacokinetic characteristics of rifabutin and limited data from clinical trials. Rifabutin is a rifamycin derivative with comparable anti-TB activity in vitro but with less hepatic P450 cytochromic enzyme-inducing effect than rifampin (11,12). An international multicenter study indicated that a 6-month regimen containing rifabutin, at the daily dose of either 150 mg or 300 mg, was as effective for treating TB as a similar regimen containing rifampin (13). In a small clinical trial, a rifabutincontaining regimen was effective in treating TB in patients co-infected with HIV (14). In addition, limited data from pharmacokinetic studies suggest that the combination of rifabutin at 150 mg/day and indinavir resulted in acceptable levels of both drugs (15). Option III cannot be recommended for patients undergoing therapy with ritonavir or saquinavir. For these patients, the decision to change the prescribed protease inhibitor to indinavir and to prescribe rifabutin for TB therapy should be made in consultation with an expert in the use of protease inhibitors to manage HIV infection. In the United States, rifabutin is approved by FDA for the prevention of disease caused by M. avium complex (MAC) but not for the treatment or prevention of TB.

ADDITIONAL RECOMMENDATIONS

Neither option II nor option III have been studied in large clinical trials of HIV-infected patients or patients undergoing protease inhibitor therapy during TB treatment. For these reasons, if either of these options are selected for managing patients with TB, CDC recommends the following interim guidelines until additional data are available and formal guidelines are issued: 1) on initiation of therapy, perform frequent bacteriologic evaluations to document sputum conversion to culture-negative status, and after culture conversion, to detect any possible treatment failures, 2) extend the duration of therapy to at least 18 months for option II or 9 months for option III, 3) use only indinavir with option III, 4) carefully monitor for drug toxicity, 5) use DOT throughout, and 6) reevaluate periodically during the first 2 years after comple-

tion of therapy (including an assessment of bacteriologic status at six months) and instruct patients to promptly report symptoms compatible with relapse of TB disease. The management of HIV-infected patients diagnosed with drug-resistant TB or diagnosed clinically with TB but without culture and susceptibility-testing results should be evaluated on a case-by-case basis and performed in consultation with a TB expert.

CONCLUSIONS

In the future, concurrent use of protease inhibitors with rifampin might be possible by modifying the doses of both to compensate for the drug interaction. For example, based on limited data submitted to FDA during the new drug application review for ritonavir, a slight increase in the dosage of ritonavir and a reduction by half in the dosage of rifampin may have resulted in satisfactory levels of both drugs. However, this option cannot be recommended until data from larger, more detailed studies are available and will require careful monitoring of the serum levels of rifampin.

Interactions between protease inhibitors and the rifamycins also have complicated prophylaxis and treatment for disseminated MAC disease. Rifabutin is one of the drugs recommended for MAC prophylaxis (16). According to the manufacturer of indinavir, rifabutin at half the dose (150 mg) can be used for MAC prophylaxis simultaneously with indinavir. Other options for MAC prophylaxis are clarithromycin and azithromycin (17,18), two macrolide antibiotics approved by FDA for this purpose and for which interactions with protease inhibitors are expected to be less of an issue. In November 1996, a joint working group of the Public Health Service and Infectious Disease Society of America will update recommendations for MAC prophylaxis.

To reduce the likelihood of drug interactions while providing optimal anti-TB care for HIV-infected persons, health-care workers involved in the care of patients with TB and health-care workers involved in HIV clinical care are encouraged to coordinate efforts and thus ensure the best possible outcome for these patients. CDC's Research and Evaluation Branch, Division of Tuberculosis Elimination, National Center for HIV, STD, and TB Prevention, (telephone [404) 639-8123]) requests the inclusion of clinical information in the comments section of TB surveillance reports from private practitioners or health department staff who manage HIV-infected patients undergoing protease inhibitor therapy when TB is diagnosed.

Reported by: Center for Drug Evaluation and Research, Food and Drug Administration. Div of Tuberculosis Elimination, and Div of HIV/AIDS Prevention, National Center for HIV, STD, and TB Prevention, CDC.

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Erratum: Vol. 45, No. 40

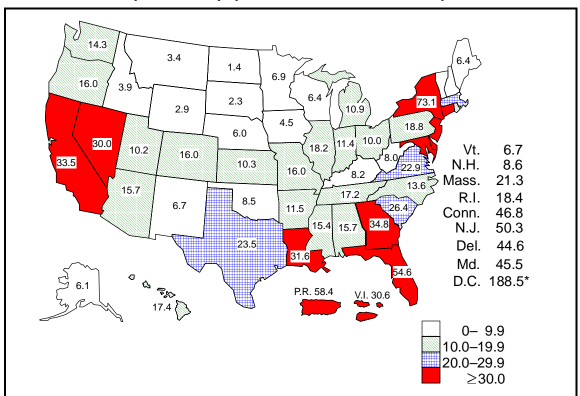
In the box, "National Adult Immunization Awareness Week—October 20–26, 1996," on page 853, the World Wide Web site given for the National Coalition for Adult Immunization is incorrect. The correct site is http://www.medscape.com/ncai.

AIDS Rates

The following map provides information about the reported number of acquired immunodeficiency syndrome (AIDS) cases per 100,000 population, by state of residence from July 1995 through June 1996. The accompanying table lists the metropolitan areas with the 50 highest annual rates of AIDS per 100,000 population.

More detailed information about AIDS cases is provided in the *HIV/AIDS Surveil-lance Report*, single copies of which are available from the CDC National AIDS Clearinghouse, P.O. Box 6003, Rockville, MD 20849-6003; telephone (800) 458-5231 or (301) 217-0023. Internet users can view an electronic copy of the report by accessing CDC's World Wide Web home page (http://www.cdc.gov), then selecting "Publications & Products."

AIDS annual rates per 100,000 population — United States, July 1995-June 1996



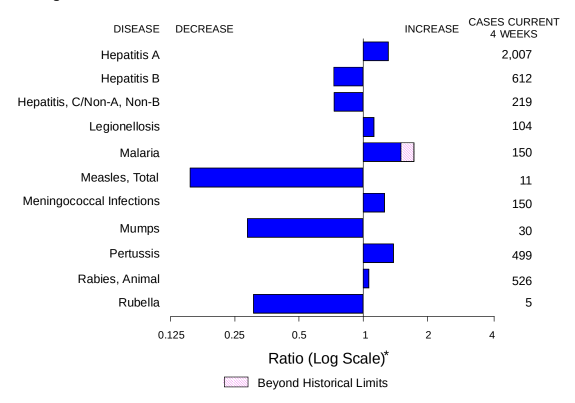
^{*}This rate represents only persons residing within the geographic boundaries of the District and differs from the rate for the larger Washington, D.C., metropolitan area (see table).

Metropolitan areas* with the 50 highest AIDS annual rates per 100,000 population — United States, July 1995–June 1996

Metropolitan area		Metropolitan area	
of residence	Rate	of residence	Rate
New York, N.Y.	132.0	Dallas, Tex.	34.1
Miami, Fla.	117.3	Las Vegas, Nev.	33.0
Jersey City, N.J.	115.2	Middlesex, N.J.	30.8
San Francisco, Calif.	109.8	Oakland, Calif.	30.4
Fort Lauderdale, Fla.	86.7	Richmond, Va.	30.1
West Palm Beach, Fla.	86.4	Rochester, N.Y.	29.8
Newark, N.J.	79.1	Austin, Tex.	29.6
San Juan, Puerto Rico	67.0	Springfield, Mass.	28.5
Baltimore, Md.	61.5	Memphis, Tenn.	28.3
New Orleans, La.	55.8	Monmouth-Ocean, N.J.	27.4
New Haven, Conn.	53.2	Nashville, Tenn.	26.9
Atlanta, Ga.	51.6	San Antonio, Tex.	25.6
Wilmington, Del.	50.2	Charleston, S.C.	25.3
Hartford, Conn.	46.2	Seattle, Wash.	24.7
Washington, D.C.	45.8	Denver, Colo.	24.4
San Diego, Calif.	43.4	Albany-Schenectady, N.Y.	24.4
Los Angeles, Calif.	42.9	Nassau-Suffolk, N.Y.	24.1
Orlando, Fla.	42.6	Birmingham, Ala.	24.0
Jacksonville, Fla.	39.6	Chicago, III.	23.4
Baton Rouge, La.	39.2	Sarasota, Fla.	23.4
Bergen-Passaic, N.J.	37.5	Indianapolis, Ind.	22.8
Houston, Tex.	35.9	Riverside-San Bernardino, Calif.	22.5
Norfolk, Va.	35.3	Bakersfield, Calif.	22.2
Tampa-Saint Petersburg, Fla.	34.6	Mobile, Ala.	21.1
Philadelphia, Pa.	34.3	Kansas City, Mo.	20.7
•		Portland, Ore.	20.7

^{*}Includes only metropolitan areas with a population ≥500,000. Metropolitan areas are named for a central city or county, may include several cities and counties, and may cross state boundaries.

FIGURE I. Selected notifiable disease reports, comparison of provisional 4-week totals ending October 19, 1996, with historical data — United States



^{*}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.

TABLE I. Summary — provisional cases of selected notifiable diseases, United States, cumulative, week ending October 19, 1996 (42nd Week)

	Cum. 1996		Cum. 1996
Anthrax Brucellosis Cholera Congenital rubella syndrome Cryptosporidiosis* Diphtheria Encephalitis: California* eastern equine* St. Louis* western equine* Hansen Disease Hantavirus pulmonary syndrome*†	64 3 1 1,758 1 84 2 - 88 15	HIV infection, pediatric*§ Plague Poliomyelitis, paralytic¶ Psittacosis Rabies, human Rocky Mountain spotted fever (RMSF) Streptococcal toxic-shock syndrome* Syphilis, congenital** Tetanus Toxic-shock syndrome Trichinosis Typhoid fever	216 2 35 1 591 13 225 23 107 16 291

^{-:} no reported cases

^{-:} no reported cases

*Not notifiable in all states.

† Updated weekly from reports to the Division of Viral and Rickettsial Diseases, National Center for Infectious Diseases (NCID).

§ Updated monthly to the Division of HIV/AIDS Prevention, National Center for HIV, STD, and TB Prevention (NCHSTP), last update September 24, 1996.

¶ Three suspected cases of polio with onset in 1996 has been reported to date.

**Updated quarterly from reports to the Division of STD Prevention, NCHSTP.

TABLE II. Provisional cases of selected notifiable diseases, United States, weeks ending October 19, 1996, and October 21, 1995 (42nd Week)

	AIE	os*	Chlamydia	Esche coli O NETSS [†]	richia 157:H7 PHLIS [§]	Gono	rrhea		atitis A,NB	Legion	ellosis
Reporting Area	Cum. 1996	Cum. 1995	Cum. 1996	Cum. 1996	Cum. 1996	Cum. 1996	Cum. 1995	Cum. 1996	Cum. 1995	Cum. 1996	Cum. 1995
UNITED STATES	51,611	57,393	305,921	2,201	1,191	237,839	318,083	2,690	3,231	739	958
NEW ENGLAND	2,065	2,824	13,447	302	75	5,648	6,182	96	104	57	30
Maine N.H.	32 66	82 77	707 397	21 37	36	49 80	73 93	8	12	2 3	5 2
Vt.	18	28	U	30	29	42	49	32	11	4	-
Mass. R.I.	997 129	1,235 187	5,649 1,552	138 15	10	1,808 416	2,189 427	50 6	74 7	26 22	19 4
Conn.	823	1,215	5,142	61	-	3,253	3,351	-	-	N	Ň
MID. ATLANTIC	14,243	15,909	34,507	193	42	26,439	35,139	253	367	184	163
Upstate N.Y. N.Y. City	1,855 7,855	1,832 8,405	N 15,878	132 12	15 -	5,381 8,618	7,521 14,166	200 1	187 1	60 7	43 5
N.J.	2,905	3,770	4,161	49	5	3,971	3,327	-	143	12	24
Pa.	1,628	1,902	14,468	N	22	8,469	10,125	52	36	105	91
E.N. CENTRAL Ohio	4,076 871	4,229 877	65,707 14,523	512 149	342 84	45,630 10,460	64,056 19,974	372 32	270 12	195 84	283 127
Ind.	498	423	8,368	72	48	5,428	7,521	8	4	39	69
III. Mich.	1,808 685	1,727 916	19,613 16,107	202 89	84 68	14,408 11,893	16,514 14,660	58 274	73 181	9 46	29 27
Wis.	214	286	7,096	N	58	3,441	5,387	-	-	17	31
W.N. CENTRAL	1,221	1,286	22,204	512	324	9,961	16,230	102	69	40	68
Minn. Iowa	226 72	302 91	2,702 3,486	231 107	212 81	U 914	2,430 1,295	2 44	4 13	5 9	6 19
Mo.	626	559	9,654	60	-	6,553	9,200	33	18	9	14
N. Dak. S. Dak.	10 10	4 17	2 784	15 21	15 -	115	26 179	_	5 1	2	3 3
Nebr.	83	84	2,049	49	4	783	953	7	15	12	16
Kans.	194	229	3,527	29	12	1,596	2,147	16	13	3	7
S. ATLANTIC Del.	13,079 232	14,724 265	44,113 1,148	119 1	60 2	77,226 1,181	88,536 1,828	212 1	202	121 11	152 2
Md.	1,961	2,226	5,608	N	8	11,762	10,718	i	7	26	24
D.C. Va.	1,001 896	872 1,151	N 9,323	- N	29	3,497 7,168	3,809 8,693	13	- 17	8 17	4 21
W. Va.	88	84	1	N	2	442	543	9	43	1	4
N.C. S.C.	677 667	837 814	-	37 9	12 7	15,166 8,757	19,681 9,852	43 25	47 19	9 5	31 30
Ga.	1,867	1,791	9,554	30	-	14,852	16,682	U	15	3	14
Fla.	5,690	6,684	18,479	30	-	14,401	16,730	120	54	41	22
E.S. CENTRAL Ky.	1,749 309	1,817 220	24,264 5,350	57 12	52 8	25,993 3,384	33,044 3,857	461 27	822 28	39 4	50 10
Tenn.	647	725	10,559	24	41	9,405	11,207	337	792	19	24
Ala. Miss.	470 323	520 352	6,759 U	10 11	3	10,817 2,387	13,578 4,402	5 92	2 U	3 13	6 10
W.S. CENTRAL	5,138	5,070	32,328	62	12	24,548	44,919	381	273	18	20
Ark.	207	223	-	13	3	2,653	4,609	13	6	2	6
La. Okla.	1,177 189	849 207	6,101 6,113	6 10	4 1	6,547 3,973	8,938 4,778	175 69	151 40	1 5	3 4
Tex.	3,565	3,791	20,114	33	4	11,375	26,594	124	76	10	7
MOUNTAIN Mont.	1,533 33	1,819 20	13,303	178 23	91	5,611 25	7,663 59	468 14	398 13	37 1	99 4
Idaho	32	38	1,236	30	13	23 87	115	93	45	-	2
Wyo.	5 406	13 571	466	11	9 36	32 1 077	46	146 49	166	3 7	12 35
Colo. N. Mex.	139	571 148	3,192	63 10	-	1,077 722	2,323 870	49 64	60 42	2	35 4
Ariz.	461	550	5,344	N	22	2,786	2,995	62	41	16	9
Utah Nev.	144 313	113 366	1,248 1,817	26 15	- 11	242 640	209 1,046	22 18	11 20	3 5	14 19
PACIFIC	8,506	9,715	56,048	266	193	16,783	22,314	345	726	48	93
Wash.	538	779	7,459	85	72	1,640	2,218	48	186	6	20
Oreg. Calif.	359 7,440	374 8,295	U 42,322	66 111	37 74	494 14,011	628 18,437	6 113	34 444	1 36	68
Alaska	28	62	964	4	2	349	558	3	1	1	-
Hawaii Guam	141 4	205	1,075 168	N N	8	289 31	473 89	175 1	61 6	4 2	5 1
P.R.	1,792	1,951	N	16	Ū	298	470	82	189	-	-
V.I. Amer. Samoa	17	30	N	N N	U U	-	26	-	-	-	-
C.N.M.I.	1	-	N	N	Ü	11	51	-	5	-	-

U: Unavailable

^{-:} no reported cases

C.N.M.I.: Commonwealth of Northern Mariana Islands

^{*}Updated monthly to the Division of HIV/AIDS Prevention, National Center for HIV, STD, and TB Prevention, last update September 24, 1996.

†National Electronic Telecommunications System for Surveillance.

§Public Health Laboratory Information System.

TABLE II. (Cont'd.) Provisional cases of selected notifiable diseases, United States, weeks ending October 19, 1996, and October 21, 1995 (42nd Week)

		me ease	Mal	aria	Mening Dise		Syp	hilis Secondary)	Tubero	ulosis	Rabies.	Animal
Reporting Area	Cum. 1996	Cum. 1995	Cum. 1996	Cum. 1995	Cum. 1996	Cum. 1995	Cum. 1996	Cum. 1995	Cum. 1996	Cum. 1995	Cum. 1996	Cum. 1995
UNITED STATES	11,266	9,154	1,191	1,079	2,582	2,449	8,817	13,325	15,003	16,989	5,496	6,417
NEW ENGLAND	3,518	1,777	47 7	42 6	113 12	116 9	148	296	341 21	402 11	600	1,285
Maine N.H.	43 42	24 22	2	1	5	20	1	2 1	11	16	89 51	46 129
Vt. Mass.	15 292	8 119	4 18	1 14	4 44	9 39	- 67	- 51	1 171	2 225	123 94	151 378
R.I.	430	295	6	4 16	13 35	5 34	3 77	3 239	27 110	40 108	33	282 299
Conn. MID. ATLANTIC	2,696 6,729	1,309 5,998	330	292	232	303	347	681	2,692	3,489	210 1,168	1,642
Upstate N.Y.	3,463	3,053	72	54	70	83	62	72	335	416	874	969
N.Y. City N.J.	252 1,393	371 1,552	172 59	161 58	32 55	44 70	106 77	302 136	1,315 602	1,953 623	109	289
Pa.	1,621	1,022	27	19	75	106	102	171	440	497	185	384
E.N. CENTRAL Ohio	67 41	390 25	110 13	140 11	345 129	347 97	1,271 470	2,290 729	1,633 240	1,597 217	85 11	92 12
Ind. III.	23 3	16 16	13 35	16 71	54 92	49 89	173 347	267 884	143 851	145 831	7 23	14 15
Mich.	-	5	36	21	38	65	142	238	309	330	31	37
Wis. W.N. CENTRAL	U 137	328 159	13 43	21 24	32 208	47 155	139 289	172 626	90 385	74 469	13 445	14 313
Minn.	59	80	19	4	25	26	51	37	88	114	25	25
Iowa Mo.	20 22	12 43	3 9	3 8	41 88	29 57	16 189	39 513	53 160	52 179	207 17	110 28
N. Dak. S. Dak.	1	-	1	1 2	3 9	1 6	-	-	6 17	3 20	56 105	24 82
Nebr.	5	5	3	3	19	14	11	11	13	20	5	5
Kans. S. ATLANTIC	30 566	19 563	8 250	3 212	23 527	22 410	22 3,075	26 3,339	48 2,865	81 2,953	30 2,288	39 1,795
Del.	78	38	3	1	2	6	34	14	20	49	61	81
Md. D.C.	331 3	374 3	69 7	55 16	66 10	36 6	533 115	388 92	242 110	320 87	518 9	347 11
Va. W. Va.	44 11	48 22	39 5	50 4	49 11	56 8	328 3	497 9	234 50	202 59	502 82	366 101
N.C. S.C.	62 6	49 16	25 11	15 1	66 49	68 52	887 314	926 485	407 281	357 263	594 74	398 109
Ga.	1	10	26	27	122	79	545	621	521	576	246	238
Fla. E.S. CENTRAL	30 57	3 63	65 26	43 23	152 184	99 172	316 2,004	307	1,000 1,034	1,040	202 173	144 244
Ky.	15	13	3	3	25	39	122	2,748 150	186	1,175 258	36	25
Tenn. Ala.	19 6	28 7	13 3	9 8	50 62	67 35	654 457	718 530	312 346	352 338	66 68	83 127
Miss.	17	15	7	3	47	31	771	1,350	190	227	3	9
W.S. CENTRAL Ark.	99 23	96 7	38	48 2	293 33	289 29	1,181 124	2,679 412	1,778 158	2,441 195	322 21	534 41
La. Okla.	2 20	7 40	6	5 1	50 32	43 32	429 151	830 155	59 137	228 321	13 27	24 28
Tex.	54	42	32	40	178	185	477	1,282	1,424	1,697	261	441
MOUNTAIN Mont.	7	12	52 7	53 3	149 5	177 2	112	178 4	503 14	533 10	132 20	160 41
ldaho	1 2	3	7	1	22	10 8	4 2	1	7	12	-	3 24
Wyo. Colo.	-	-	22	23	32	44	23	96	72	59	26 41	9
N. Mex. Ariz.	1 -	1 1	2 6	5 10	23 37	32 51	1 67	6 36	67 197	66 257	6 30	6 51
Utah Nev.	1 2	1 6	4	6 5	15 12	15 15	2 13	4 31	39 101	37 88	4 5	15 11
PACIFIC	86	96	295	245	531	480	390	488	3,772	3,930	283	352
Wash. Oreg.	14 13	10 15	20 18	21 15	84 93	77 87	6 11	12 19	211 134	225 106	6 1	14 2
Calif.	58	71	246	196	341	302	372	455	3,229	3,383	268	329 7
Alaska Hawaii	1	-	3 8	3 10	8 5	10 4	1	2	51 147	62 154	8 -	-
Guam P.R.	-	-	-	1 1	1 4	2 23	3 108	8 237	35 63	87 162	- 41	- 36
V.I.	-	-	-	2	-	-	-	-	-	-	4 I -	-
Amer. Samoa C.N.M.I.	-	-	-	1	-	-	1	9	-	4 31	-	-

U: Unavailable

-: no reported cases

TABLE III. Provisional cases of selected notifiable diseases preventable by vaccination, United States, weeks ending October 19, 1996, and October 21, 1995 (42nd Week)

	H. influ	ienzae,		Hepatitis (vi	al), by type			Measles	-	
	inva			A Cum	Cum		Ind	igenous	lm	ported [†]
Reporting Area	Cum. 1996*	Cum. 1995	Cum. 1996	Cum. 1995	Cum. 1996	Cum. 1995	1996	Cum. 1996	1996	Cum. 1996
UNITED STATES	830	913	22,387	24,143	7,784	7,982	4	404	2	46
NEW ENGLAND	24	36	319	251	159	185	-	11	-	4
Maine N.H.	9	3 9	16 14	26 11	2 14	7 18	-	-	-	-
Vt. Mass.	1 12	2 11	7 165	5 105	10 55	5 71	-	1 9	-	1 3
R.I. Conn.	2	5 6	17 100	28 76	9 69	8 76	-	1	-	-
MID. ATLANTIC	150	133	1,519	1,476	1,200	1,134	_	23	-	5
Upstate N.Y. N.Y. City	44 31	36 32	363 485	373 695	280 491	309 342	-	9	-	3
N.J.	48	18	278	223	205	308	-	3	-	-
Pa. E.N. CENTRAL	27 137	47 155	393 1,856	185 2,686	224 803	175 909	-	11 5	-	2 7
Ohio	80	77	636	1,512	105	91	-	2	-	3
Ind. III.	12 32	19 40	256 460	152 549	128 210	185 240	-	2	-	1
Mich. Wis.	7 6	17 2	356 148	305 168	305 55	330 63	-	- 1	-	3
W.N. CENTRAL	40	69	2,014	1,596	364	524	-	20	_	2
Minn. Iowa	25 5	38 3	108 305	163 70	51 62	49 41	-	16	-	2
Mo.	7	21	954	1,123	179	359	-	3	-	-
N. Dak. S. Dak.	1	1	110 41	22 54	2 5	4 2	-	-	-	-
Nebr. Kans.	1 1	3 3	190 306	41 123	36 29	28 41	-	1	-	-
S. ATLANTIC	158	177	1,159	942	1,215	1,031	-	5	-	9
Del. Md.	2 51	- 58	15 200	9 179	7 246	7 209	-	1 -	-	2
D.C. Va.	6 9	- 26	35 140	22 173	29 117	19 93	-	1	-	3
W. Va.	7	7	13	21	21	45	-	-	-	-
N.C. S.C.	22 4	25 2	139 44	89 40	277 74	224 43	-	3	-	1 -
Ga. Fla.	37 20	54 5	150 423	52 357	32 412	62 329	-	-	-	2 1
E.S. CENTRAL	25	10	1,048	1,678	668	692	-	2	-	-
Ky. Tenn.	4 12	4	38 687	41 1,400	52 381	60 547	-	2	-	-
Ala. Miss.	8 1	5 1	148 175	72 165	57 178	85 U	-	-	-	-
W.S. CENTRAL	34	57	4,760	3,576	1,048	1,111	_	26	_	2
Ark. La.	- 4	6 1	417 157	469 111	65 120	53 163	-	-	-	-
Okla.	27	21	1,980	928	59	139	-	-	-	-
Tex. MOUNTAIN	3 86	29 99	2,206 3,581	2,068 3,363	804 914	756 686	-	26 152	-	2 5
Mont.	-	-	98	124	12	19	-	1	-	-
Wyo.	1 35	3 6	29	273 97	76 36	80 24	-	1	-	-
Colo. N. Mex.	13 10	15 12	383 316	434 696	117 318	106 258	-	4 16	-	3
Ariz. Utah	11 8	25 10	1,425 813	915 596	210 81	97 53	-	8 117	-	2
Nev.	8	28	326	228	64	49	-	5	-	-
PACIFIC Wash.	176 4	177 9	6,131 471	8,575 715	1,413 83	1,710 156	4	160 51	2	12
Oreg.	22	23	702	2,265	87	99	-	4	-	-
Calif. Alaska	146 2	140 1	4,861 36	5,410 41	1,217 14	1,431 11	-	36 63	-	5 -
Hawaii	2	4	61	144	12	13	4	6	2	7
Guam P.R.	1	3	2 99	7 84	310	4 503	U -	7	U -	-
V.I. Amer. Samoa	-	-	-	8 6	-	14	U	-	U U	-
C.N.M.I.	10	11	1	24	5	22	ŭ	-	ŭ	-

U: Unavailable

^{-:} no reported cases

 $^{^{*}}$ Of 197 cases among children aged <5 years, serotype was reported for 45 and of those, 14 were type b.

[†]For imported measles, cases include only those resulting from importation from other countries.

TABLE III. (Cont'd.) Provisional cases of selected notifiable diseases preventable by vaccination, United States, weeks ending October 19, 1996, and October 21, 1995 (42nd Week)

	Measles (Rub		l	JCI 2 1,	1990 (-	TETTO	VVCCR,						
	То	Total		Mumps			Pertussis			Rubella			
Reporting Area	Cum. 1996	Cum. 1995	1996	Cum. 1996	Cum. 1995	1996	Cum. 1996	Cum. 1995	1996	Cum. 1996	Cum. 1995		
UNITED STATES	450	281	5	515	690	122	4,213	3,519	4	207	109		
NEW ENGLAND	15	9	-	2	11	24	875	486	-	27	46		
Maine N.H.	-	-	-	-	4 1	-	20 90	39 43	-	-	1		
Vt. Mass.	2 12	2	-	2	2	4 20	98 610	67 307	-	2 21	- 7		
R.I.	-	5	-	-	1	-	30	4	-	-	-		
Conn. MID. ATLANTIC	1 28	2 12	-	- 74	3 101	- 17	27 384	26 289	-	4 11	38 13		
Upstate N.Y.	-	1	-	22	24	11	221	135	-	4	3		
N.Y. City N.J.	12 3	5 6	-	16 2	15 17	-	29 16	44 17	-	4 2	8 2		
Pa.	13	-	-	34	45	6	118	93	-	1	-		
E.N. CENTRAL Ohio	12 5	15 2	1	88 39	126 42	7 1	429 193	442 122	-	3	3		
Ind.	-	-	1	8	8	3	55	42	-	-	-		
III. Mich.	3 3	2 5	-	19 21	35 41	3	141 35	89 62	-	1 2	3		
Wis.	1	6	-	1	-	-	5	127	-	-	-		
W.N. CENTRAL Minn.	22 18	2	1	15 5	40 4	10 8	318 251	235 120	-	-	-		
lowa	-	-	-	1	9	1	17	10	-	-	-		
Mo. N. Dak.	3	1 -	1 -	6 2	22 1	-	33 1	55 8	-	-	-		
S. Dak. Nebr.	-	-	-	-	- 4	- 1	4 8	11 10	-	-	-		
Kans.	1	1	-	1	-	-	4	21	-	-	-		
S. ATLANTIC	14	14	3	90	99	26	500	300	1	93	9		
Del. Md.	1 2	1	-	25	30	1 2	13 173	10 39	-	-	1		
D.C. Va.	1 3	-	-	1 12	- 21	-	2 71	6 19	-	2 2	-		
W. Va.	-	-	-	-	-	-	2	-	-	-	-		
N.C. S.C.	4 -	-	1 1	20 6	16 10	21 1	100 37	110 23	1 -	78 1	1 -		
Ga. Fla.	2 1	2 11	- 1	3 23	6 16	- 1	17 85	19 74	-	10	- 7		
E.S. CENTRAL	2	-	-	21	11		85	265	_	2	1		
Ky.	2	-	-	3	4	-	39 19	22 206	-	-	1		
Tenn. Ala.	-	-	-	3	4	-	18	35	-	2	-		
Miss.	-	-	-	15	3	-	9	2	N	N	N		
W.S. CENTRAL Ark.	28	31 2	-	28 2	47 7	5 2	101 12	260 34	-	3	7 -		
La. Okla.	-	18	-	13	12	1 2	9 10	18 31	-	1 -	-		
Tex.	28	11	-	13	28	-	70	177	-	2	7		
MOUNTAIN	157	68	-	21	30 1	4	353	524	-	7	4		
Mont. Idaho	1	-	-	-	3	-	28 102	3 99	-	3	-		
Wyo. Colo.	1 7	26	-	3	2	-	5 91	1 85	-	2	-		
N. Mex.	16	31	Ν	N	N	4	54	99	-	-	-		
Ariz. Utah	8 119	10 -	-	1 2	2 11	-	27 19	153 27	-	1 -	3 1		
Nev.	5	1	-	15	11	-	27	57	-	1	-		
PACIFIC Wash.	172 51	130 19	-	176 19	225 12	29 28	1,168 531	718 250	3	61 2	26 1		
Oreg. Calif.	4 41	1 108	-	128	192	1	33 573	44 375	- 3	1 55	20		
Alaska	63	-	-	2	12	-	4	1	-	-	-		
Hawaii	13	2	-	27	9	-	27	48	-	3	5		
Guam P.R.	- 7	3	U -	5 1	4 2	U -	1 1	2 1	U	-	1 -		
V.I. Amer. Samoa	-	-	U U	-	3	U U	-	-	U U	-	-		
C.N.M.I.	-	-	Ü	-	1	Ü	-	-	Ü	-	-		

U: Unavailable

-: no reported cases

TABLE IV. Deaths in 121 U.S. cities,* week ending October 19, 1996 (42nd Week)

	-	All Causes, By Age (Years)						P&I [†]	All Causes, By Age (Years)						
Reporting Area	All Ages	>65	45-64	25-44	1-24	<1	Total	Reporting Area	All Ages	>65	45-64	25-44	1-24	<1	P&l [†] Total
NEW ENGLAND Boston, Mass. Bridgeport, Conn. Cambridge, Mass. Fall River, Mass. Hartford, Conn. Lowell, Mass. Lynn, Mass. New Bedford, Mas. New Haven, Conn. Providence, R.I. Somerville, Mass. Springfield, Mass.		406 75 34 17 22 40 24 13 27 26 33 7	28 7 3 2 12 5 3 10 10 2	37 13 2 1 - 4 1 2 2 3 3	8 1 - 1 2 - - 3 - 1	12 2 2 - 1 - - 4 2	34 4 1 1 3 1 2 1 3	S. ATLANTIC Atlanta, Ga. Baltimore, Md. Charlotte, N.C. Jacksonville, Fla. Miami, Fla. Norfolk, Va. Richmond, Va. Savannah, Ga. St. Petersburg, Fla. Tampa, Fla. Washington, D.C. Wilmington, Del.	1,020 80 243 U 111 109 39 68 36 49 148 114 23	648 40 150 U 77 63 26 41 23 36 108 64 20	209 25 42 U 21 29 8 18 9 6 25 23 3	112 10 39 U 10 12 3 5 2 4 15	25 5 5 U 3 1	24 5 U 3 2 1 4 2 3	51 13 13 10 1 1 3 5 3 4 16 2
Waterbury, Conn. Worcester, Mass. MID. ATLANTIC Albany, N.Y. Allentown, Pa. Buffalo, N.Y. Camden, N.J. Elizabeth, N.J. Erie, Pa.§ Jersey City, N.J. New York City, N.Y. Newark, N.J. Paterson, N.J. Philadelphia, Pa. Pittsburgh, Pa.§ Reading, Pa. Rochester, N.Y. Schenectady, N.Y. Scranton, Pa.§ Syracuse, N.Y. Trenton, N.J. Utica, N.Y.	19 57 2,382 60 23 82 38 82 54 41 1,219 67 31 300 84 7 128 24 26 85 388 23	13 43 1,631 19 60 23 14 41 41 28 839 27 53 5 94 41 15 20 61 18 18	5 11 461 9 47 17 8 8 9 239 17 18 24 47 15 15 14	1 2 193 2 - 3 3 4 3 3 106 4 25 5 - 5 1 1 6 4 2	44 1 1 3 20 2 2 9 2 -	1 52 3 - 11 15 4 1 9 6 - - - 4 1	2 7 124 6 4 3 6 2 37 10 1 8 3 2 1 7 1 8 2 1	E.S. CENTRAL Birmingham, Ala. Chattanooga, Tenn. Knoxville, Tenn. Lexington, Ky. Memphis, Tenn. Mobile, Ala. Montgomery, Ala. Nashville, Tenn. W.S. CENTRAL Austin, Tex. Baton Rouge, La. Corpus Christi, Tex. Dallas, Tex. El Paso, Tex. Ft. Worth, Tex. Houston, Tex. Little Rock, Ark. New Orleans, La. San Antonio, Tex. Shreveport, La. Tulsa, Okla.	676 109 75 74 59 146 60 24 129 1,355 53 40	446 78 48 50 42 88 43 16 81 871 29 26 38 92 61 75 239 37 53 133 26 62	137 222 17 15 13 32 9 5 24 264 11 8 10 34 13 82 12 18 38 11	53 7 5 6 3 12 6 1 13 121 6 4 3 7 31 4 11 20 6 2	15 - 3 1 4 1 - 6 59 1 1 2 3 3 5 9 3 4 6 8 4	24 11 5 10 11 22 5 39 6 11 3 4 3 3 9 2 3 4 1	38 72 65 91 11 7 78 32 46 55 36 8 10 22
Yonkers, N.Y. E.N. CENTRAL Akron, Ohio Canton, Ohio Chicago, III. Cincinnati, Ohio Cleveland, Ohio Cleveland, Ohio Dayton, Ohio Detroit, Mich. Evansville, Ind. Fort Wayne, Ind. Gary, Ind. Grand Rapids, Micl Indianapolis, Ind. Madison, Wis. Milwaukee, Wis. Peoria, III. Rockford, III. South Bend, Ind. Toledo, Ohio Youngstown, Ohio W.N. CENTRAL Des Moines, Iowa Duluth, Minn. Kansas City, Kans. Kansas City, Kans. Kansas City, Mo. Lincoln, Nebr. Minneapolis, Minn. Omaha, Nebr. St. Louis, Mo. St. Paul, Minn. Wichita, Kans.	183 61 100 44 56 47 103 64 880 110 26 33 124 34	27 1,375 22 22 29 68 109 126 84 129 36 46 U 44 127 45 68 33 48 31 72 56 606 80 00 18 20 76 26 160 57 82 29 58	9 10 65 20 32 32 32 17 34 5 8 U 7 35 9 9 6 10 7 5 147 24 6 32 16 18 5	171 33 500 76 18 82 22 3 10 12 4 5 1 1 1 3 10 2 5 7 4 1 3 10 17 3 17 4 7 4 7 4 7 7 4 7 7 7 7 7 7 7 7 7 7	49 - 2 18 1 - 6 17 2 2 U 1 3 - 2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	544 	3 110 · 2761 157821U · 125546261 5712182176262	MOUNTAIN Albuquerque, N.M. Colo. Springs, Colo Denver, Colo. Las Vegas, Nev. Ogden, Utah Phoenix, Ariz. Pueblo, Colo. Salt Lake City, Utah Tucson, Ariz. PACIFIC Berkeley, Calif. Fresno, Calif. Glendale, Calif. Honolulu, Hawaii Long Beach, Calif. Los Angeles, Calif. Pasadena, Calif. Portland, Oreg. Sacramento, Calif. San Diego, Calif. San Diego, Calif. San Jose, Calif. Santa Cruz, Calif. Seattle, Wash. Spokane, Wash. Tacoma, Wash. Tacoma, Wash.	113 131 188 129 26 104 143 1,234 15 5 5 9 82 45 327 25 99 0 134	545 67 38 72 80 15 74 20 71 108 858 12 41 25 222 16 69 99 71 11 68 14 10 36 53 7,386	151 21 12 21 28 1 26 2 13 27 212 3 U 2 14 13 49 5 19 U 21 22 16 1 3 6 8 2,039	83 7 2 16 14 2 2 11 3 14 4 4 5 33 1 8 U 11 14 4 4 5 33 1 1 4 4 9 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	21 5 6 2 1 5 2 36 U 1 2 2 2 2 2 2 2 3 3 3 3 2 2 2 2 2 2 2 2	17 2 1 4 2 2 - 5 1 2 2 2 4 - U 1 2 5 3 3 1 1 1 1 7 7 3 3 2771	60 2 2 12 12 6 1 9 14 81 1 U 2 7 4 18 3 4 U 11 15 2 2 4 4 6 3 3

U: Unavailable -: no reported cases

*Mortality data in this table are voluntarily reported from 121 cities in the United States, most of which have populations of 100,000 or more. 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 these 3 Pennsylvania cities, these numbers are partial counts for the current week. Complete counts will be available in 4 to 6 weeks.

Total includes unknown ages.

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