

MORBIDITY AND MORTALITY

WEEKLY REPORT

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Update: Investigation of Bioterrorism-Related Anthrax and Interim Guidelines for Exposure Management and Antimicrobial Therapy, October 2001

Since October 3, 2001, CDC and state and local public health authorities have been investigating cases of bioterrorism-related anthrax. This report updates previous findings, provides new information on case investigations in two additional areas, presents the susceptibility patterns of *Bacillus anthracis* isolates, and provides interim recommendations for managing potential threats and exposures and for treating anthrax.

As of October 24, investigations in the District of Columbia (DC), Florida, New Jersey, New York City (NYC), Maryland, Pennsylvania, and Virginia have identified 15 (11 confirmed and four suspected) cases of anthrax according to the CDC surveillance case definition (1). Seven of the 15 cases were inhalational anthrax and eight were cutaneous. Of the seven inhalational cases, five occurred in postal workers in New Jersey and DC, and one in a person who sorted and distributed mail at a media company in Florida. Two letters mailed to two different recipients in NYC and one letter mailed to a recipient in DC are known to have contained *B. anthracis* spores. Six cases were identified in employees of media companies; one was a 7-month-old infant who visited a media company; and eight cases are consistent with exposures along the postal route of letters known to be contaminated with *B. anthracis* spores in New Jersey and DC. Using molecular typing, analysis of *B. anthracis* isolates from cases in Florida, NYC, and DC indicated that the isolates are indistinguishable (2). Epidemiologic investigations and surveillance in other locations are continuing; no additional cases have been identified.

Florida

As of October 24, investigations in Florida have identified two confirmed cases of inhalational anthrax in persons who worked at the same media company; no additional cases of disease have been identified since the last report (1). A pleural biopsy for the second confirmed patient was positive for *B. anthracis* by immunohistochemical (IHC) staining. In addition, a >4-fold increase in levels of serum antibody (lgG) to the protective antigen (PA) component of the anthrax toxin using enzyme-linked immunosorbant assay (ELISA) was demonstrated.

Environmental sampling of the work site revealed *B. anthracis* contamination and implicated one or more mailed letters or packages as the likely source of exposure. Several environmental specimens from regional and local postal centers that provided mail services to the work site were culture-positive for *B. anthracis*. Thirty postal workers

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had no evidence of *B. anthracis* exposure by nasal swab testing. No cases of disease have been identified among postal workers. On the basis of the positive environmental swabs, focused clean-up procedures continue at regional and local postal centers. The Environmental Protection Agency (EPA), in consultation with health officials, is conducting decontamination of the work site.

Approximately 1,100 persons were started on antimicrobial prophylaxis for suspected *B. anthracis* exposure; 555 worked either full- or part-time in the affected building. The majority of other persons reported spending at least 1 hour in the affected building since August 1. Additional follow-up for compliance with prophylaxis recommendations and monitoring adverse events associated with long-term antimicrobial prophylaxis is ongoing.

New York

Investigations in NYC have identified five (three confirmed and two suspected) cutaneous anthrax cases; three cases (one confirmed and two suspected) have been identified since the last report (1). These five cases were associated with four media companies (A–D). The two previously reported cases were related to work sites A and B, and the three additional cases were related to work sites C, D, and A, respectively. No cases among postal workers have been identified.

On October 1, a 27-year-old woman who regularly handled mail at work site C sought medical care at a local hospital for two lesions on the left cheek, which developed surrounding erythema and edema and local adenopathy. A biopsy obtained on October 16 was positive by IHC staining for the cell wall antigen of *B. anthracis* and serologic testing was weakly reactive. No suspicious letter was identified from her work site.

Two suspected cases of cutaneous anthrax also have been detected. The first suspected case, a 29-year-old woman with onset of illness on September 22, frequently handled mail at work site D. At her work site, an unopened letter postmarked September 18, which contained powder contaminated with *B. anthracis* was found on October 19. The second suspected case, a 23-year-old woman with onset of illness on September 28, handled a suspicious letter postmarked September 18 from work site A. All three patients were treated with ciprofloxacin and have shown clinical improvement. A total of three persons were confirmed by nasal swabs to have been exposed to *B. anthracis*, presumably acquired during handling and processing of specimens during the investigation of the first confirmed case (1).

In work site A, potentially exposed persons were identified and prescribed antimicrobial prophylaxis. An environmental investigation of work site A was conducted subsequently; environmental samples taken from work site A were culture-positive for *B. anthracis*. Of 1,360 persons who were tested by nasal swabs from work site A, all were confirmed negative. Nasal swabs were obtained from 1,202 persons from work sites B, C, and D; 1,183 tested negative and 19 are pending final results. Environmental samples taken from work site A were positive. Testing of environmental specimens from work sites B, C, and D is ongoing.

Prophylaxis was recommended for potentially exposed persons at work site A. Antimicrobial prophylaxis was initiated for nine persons who had recent contact with the sealed letter containing *B. anthracis* in work site D.

New Jersey

To date, investigations in New Jersey and Pennsylvania have identified four (two confirmed and two suspected) anthrax cases. Cutaneous disease has been diagnosed in three patients and one has illness suspected to be inhalational anthrax, but laboratory tests to confirm the diagnosis are pending. All four of these patients worked at one of two postal facilities in New Jersey. Although no specific contaminated letter was identified, contaminated letters destined for both NYC and DC passed through at least one of these postal facilities in New Jersey.

On October 1, a 45-year-old female mail carrier sought medical care at a local hospital for a 4-day history of worsening skin lesions on her right forearm. A biopsy was obtained and arrived at CDC on October 17 and later that night was found positive by IHC. In addition, tissue was positive for *B. anthracis* by polymerase chain reaction (PCR), and serologic testing was reactive. The patient's condition improved on antimicrobial therapy.

On October 16, a 35-year-old male mail processor, with a history of a chronic, bullouslike skin condition, was taken to a local hospital complaining of a 2-day history of a large pustular lesion on his neck. He returned 1 day later with increasing ulceration of the skin lesion associated with fatigue, chills, and a swollen throat; he was afebile but had vesicles and bullae around the pustular lesion. Biopsy was positive by IHC, and serologic testing was reactive to *B. anthracis*. The patient's lesions responded to antimicrobial therapy.

Two suspected cases also have been detected. The first case occurred in a 39-yearold male machine mechanic who was taken to a local hospital on September 26 for two bullous, vesicular lesions with surrounding erythema, edema, and induration on the right forearm, which progressed to black eschars. The patient was treated for cellulitis with ceftriaxone followed by amoxicillin/clavulanate. The patient was reported to CDC on October 17 and serologic testing at CDC was reactive to *B. anthracis*. No biopsy was obtained. The patient's condition improved.

On October 14, the second suspected case occurred in a 56-year-old female postal worker who sought medical care for fever, diarrhea, and vomiting at a local hospital. On October 19, the patient was admitted to the hospital with chills, dry cough, and pleuritic chest pain. A chest radiograph showed a small right infiltrate and bilateral effusions, but no evidence of a widened mediastinum. The next day, her respiratory status and pleural effusions worsened. A chest computerized tomography (CT) showed an enlarged mediastinal and cervical lymph nodes without parenchymal disease. The pleural fluid was positive for *B. anthracis* by PCR. Bilateral pleural effusions have complicated the patient's hospital course and she continues to require supplemental oxygen.

On October 20, the postal facility was closed; the New Jersey Department of Health and Senior Services recommended that postal workers at both postal facilities initiate antimicrobial prophylaxis pending further epidemiologic and environmental investigation. Both facilities have been closed pending results of further environmental evaluation. Environmental sampling is being conducted at both postal facilities. In one facility, 13 of 23 samples from high-risk areas were preliminarily culture-positive for *B. anthracis.* Clean-up efforts are ongoing. Results of cultures from samples taken in the second facility and results from approximately 600 nasal swab cultures obtained from postal employees are pending.

District of Columbia

To date, investigations in DC, Maryland, and Virginia have identified four confirmed anthrax cases. All patients had inhalational illness and all worked at a single postal facility in DC.

On October 15, a staff member in the office of a U.S. Senator noted a small burst of dust released while opening a tightly sealed letter. The U.S. Capitol Police and Federal Bureau of Investigation (FBI) were notified and the area was vacated and secured immediately; ventilation systems for the Senator's offices were deactivated within 45 minutes of recognizing the threat. The letter and surrounding carpet were removed and sent for testing. On October 16, the letter tested positive for *B. anthracis* by PCR, and an epidemiologic investigation was initiated by the health officials from the Office of Attending Physician, U.S. Capitol; DC Department of Health (DCDOH); Infectious Disease Service, National Naval Medical Center; and CDC.

Based on the initial investigation, the area of exposure was determined to consist of two floors in the southeast quadrant of the building where the U.S. Senator's office is located. Approximately 340 staff and visitors potentially were exposed. Beginning October 15, nasal swab testing was performed on these persons and approximately 5,000 additional persons who referred themselves for testing. Twenty-eight persons had evidence of exposure by nasal swab testing; 13 were in the immediate office space where the letter was opened, nine were in adjacent areas, and six were first responders. Antimicrobial prophylaxis was administered to persons from the area of exposure and firstresponders to the incident. Environmental specimens were collected at the affected building and other buildings in the U.S. Capitol complex. To date, environmental specimens are positive from the area of exposure as well as two mail rooms in the U.S. Capitol complex; one of the mail rooms did not process the contaminated letter. None of the mail room personnel and none of the postal workers at the post office serving the mail rooms had positive nasal swabs. These mail handlers were all offered prophylactic antibiotics. Initially, a single positive environmental sample for the post office serving these mail rooms was positive. Subsequent samples from this post office and the mail distribution center serving this post office were positive.

On October 19, enhanced regional surveillance activities (a collaborative effort between DCDOH, Maryland Department of Health and Mental Hygiene, and the Virginia Department of Health) identified a case of pulmonary illness in a postal worker. The postal worker, a 56-year-old man, sought medical care at a Virginia hospital for fever, chills, chest heaviness, malaise, and minimally productive cough of 3 days' duration. Initial evaluation in the emergency department (ED) revealed a widened mediastinum on a chest radiograph; a subsequent CT scan revealed mediastinal lymphadenopathy and small, bilateral pleural effusions. The patient was hospitalized for suspected inhalational anthrax and was treated with broad spectrum antimicrobial agents, including ciprofloxacin. Blood cultures grew gram-positive rods within 15 hours of collection, later confirmed to be *B. anthracis* at the Virginia State Health Laboratory and CDC on October 21. The patient is clinically stable and remains hospitalized.

On October 20, a second postal worker, also a 56-year-old man, who worked at the same distribution center, was admitted to the hospital with a 3-day history of progressively worsening headache and night sweats. He had no fever, stiff neck, or other symptoms or signs consistent with meningitis. He had a mild sore throat and occasional dry cough. Because the patient was linked epidemiologically to the index case of inhalational

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anthrax, a chest radiograph and chest CT scan were performed that revealed mediastinal lymphadenopathy and a right middle lobe infiltrate. Antimicrobial therapy was initiated. Blood cultures grew *B. anthracis* within 18 hours. The patient is clinically stable and remains hospitalized.

On October 21, a third postal worker, a 55-year-old man, who worked at the same distribution center was admitted to the hospital with suspected inhalational anthrax. The patient had initially sought medical care at a physician's office on October 18 for 2 days of progressive fatigue, myalgias, and fever. The patient had a temperature of 102 F (38.9 C) and normal white blood cell count and was sent home. The patient returned to the ED on October 21 with persistent symptoms, including chills, vague chest tightness, and temperature of 102 F (38.9 C). Chest radiograph revealed right middle and lower lobe alveolar infiltrates and right hilar and peritracheal soft tissue fullness. Evaluation revealed hypoxia, leukocytosis, and hemoconcentration. Antimicrobial therapy was initiated, and the patient was mechanically ventilated. The patient's condition deteriorated, and he died on October 21. Blood cultures obtained on admission to the hospital grew grampositive bacilli, which were confirmed later as *B. anthracis* at CDC.

On October 22, a fourth postal worker, a 47-year-old man, who worked at the same distribution center was admitted to the hospital with suspected inhalational anthrax. The patient had initially presented to the ED on October 21 with complaints of 5 days of progressive fatigue, nausea, vomiting, and diarrhea, and syncope. The patient was afebrile and had orthostatic hypotension. A chest radiograph was obtained and reported to be normal. The patient received intravenous fluids and was discharged. He returned to the ED 26 hours later following another syncopal episode and persistent gastrointestinal complaints. The patient was afebrile, hypotensive, diaphoretic, and in respiratory distress. A second chest radiograph and a chest CT revealed mediastinal lymphadenopathy and bilateral pleural effusions. Subsequent review of the first chest radiograph revealed an ill-defined area of increased density in the right subhilar region. Laboratory evaluation revealed leukocytosis and hemoconcentration. Antimicrobial therapy was initiated, and the patient was mechanically ventilated. Peripheral blood smear demonstrated grampositive bacilli; blood cultures grew gram-positive bacilli within 18 hours and were confirmed as *B. anthracis* at CDC. The patient died on October 22.

On October 20, CDC and DCDOH initiated an investigation of the postal facility where the four patients were employed. Although no specific exposure event was identified, the contaminated tightly sealed letter that was mailed to the Senator's office was processed at this facility on October 12 before entering the Capitol mail distribution system. The postal facility was closed on October 21, and antimicrobial prophylaxis was recommended to employees working in proximity to the same mail sorting area of the first patient. In addition, visitors to nonpublic operations areas of this facility also were offered antimicrobial prophylaxis.

On October 22, because of concern about the potential for unrecognized aerosol exposures among postal workers, antimicrobial therapy was recommended for all workers and visitors to nonpublic areas in this postal facility. Subsequently, this recommendation has been extended to all postal workers in the DC area directly served by this postal facility pending results of ongoing epidemiologic and environmental investigation.

The first patient also worked at a second postal facility. On October 21, this facility also was closed. Antimicrobial prophylaxis also was recommended for workers at this facility pending further epidemiologic and environmental testing.

Susceptibility Testing of B. anthracis Isolates

Antimicrobial susceptibility patterns were determined for 11 *B. anthracis* isolates associated with intentional exposures in Florida, NYC, and DC. Susceptibility breakpoints for interpreting minimum inhibitory concentration (MIC) results for *B. anthracis* have not been determined by the National Committee for Clinical Laboratory Standards (NCCLS); thus, breakpoints for staphylococci were used (*3*). All *B. anthracis* isolates were susceptible to ciprofloxacin (MIC \leq 0.06 µg/mL), doxycycline (MIC \leq 0.03 µg/mL), chloramphenicol (MIC=4 µg/mL), clindamycin (MIC \leq 0.5 µg/mL), tetracycline (MIC=0.06 µg/mL), rifampin (MIC \leq 0.5 µg/mL), and vancomycin (MIC=1–2 µg/mL). Limited testing of imipenem suggests that these organisms are also susceptible to this agent (MIC \leq 0.12 µg/mL) and are likely susceptible to meropenem. Susceptibility of the isolates was considered intermediate to erythromycin (MIC=1 µg/mL) and borderline susceptible to azithromycin (MIC=2 µg/mL); clarithromycin was considered susceptible (MIC=0.25 µg/mL).

B. anthracis isolates were susceptible to penicillin (MIC range: $\leq 0.06 \text{ ug/mL}-0.12 \text{ µg/mL}$) and amoxicillin (MIC $\leq 0.06 \text{ µg/mL}$); ceftriaxone (MIC=16) was considered intermediate. NCCLS has not defined either a *B. anthracis* or staphylococcal interpretive breakpoint for ceftriaxone results; thus, breakpoints for gram-negative organisms were used to interpret ceftriaxone results. These ceftriaxone MICs and additional laboratory data at CDC indicate the presence in *B. anthracis* isolates of a cephalosporinase, an enzyme that inhibits the antibacterial activity of cephalosporins such as ceftriaxone. Additional studies were performed with some of the *B. anthracis* isolates to identify other betalactamases, the general class of enzymes that inactivate penicillins, cephalosporins, and related drugs. These preliminary studies indicate the presence of a class B cephalosporinase and suggest that a penicillinase also may be present. These enzymes often are present in naturally occurring *B. anthracis* isolates.

This information is current as of October 24, 2001, 9 p.m. eastern daylight time. Intensive surveillance activities and environmental and case investigations are in progress to identify and treat all U.S. Postal Service workers and others at potential risk for anthrax. Surveillance also is being conducted to monitor adverse events associated with antimicrobial prophylaxis for anthrax. CDC and FBI are collaborating to accelerate all aspects of the investigation surrounding these events.

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Editorial Note: Bioterrorism attacks using *B. anthracis* spores sent through the mail have resulted in 15 anthrax cases and three deaths. The initial anthrax cases occurred among persons with known or suspected contact with opened letters contaminated with *B. anthracis* spores. Later, investigations identified four confirmed cases and one suspected case among postal workers who had no known contact with contaminated opened letters. This suggests that sealed envelopes contaminated with *B. anthracis*

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passing through the postal system may be the source of exposure. The number of contaminated envelopes passing through the postal system is not known. In addition, automated sorting could damage envelopes and release spores into postal environments; other circumstances that could contribute to the contamination of postal facility environments may be identified.

Because these cases are the result of intentional exposures, FBI and other law enforcement authorities are investigating these events as criminal acts and are working to identify and eliminate the source of these exposures. Until that occurs, the possibility of further exposure to *B. anthracis* and subsequent clinical illness exists. Clinicians and laboratorians should be vigilant for symptoms or laboratory findings that indicate *B. anthracis* infection, particularly among mail handlers. Information to guide health-care providers and laboratorians is available at http://www.bt.cdc.gov>.

Managing Threats

Letters containing *B. anthracis* spores have been sent to persons in NYC and DC. Prompt identification of a threat and institution of appropriate measures may prevent inhalational anthrax. To prevent exposure to *B. anthracis* and subsequent infection, suspicious letters or packages should be recognized and appropriate protective steps taken.

Characteristics of suspicious packages and letters include inappropriate or unusual labeling, strange return address or no return address, postmarks from a city or state different from the return address, excessive packaging material, and others. If a package appears suspicious, it should not be opened. The package should be handled as little as possible. The room should be vacated and secured promptly and appropriate security or law enforcement agencies promptly notified (Box 1).

Box 1. Handling of Suspicious Packages or Envelopes

- Do not shake or empty the contents of a suspicious package or envelope.
- Do not carry the package or envelope, show it to others, or allow others to examine it.
- Put the package or envelope on a stable surface; do not sniff, touch, taste, or look closely at it or any contents that may have spilled.
- Alert others in the area about the suspicious package or envelope. Leave the area, close any doors, and take actions to prevent others from entering the area. If possible, shut off the ventilation system.
- Wash hands with soap and water to prevent spreading potentially infectious material to face or skin. Seek additional instructions for exposed or potentially exposed persons.
- If at work, notify a supervisor, a security officer, or a law enforcement official. If at home, contact the local law enforcement agency.
- If possible, create a list of persons who were in the room or area when this suspicious letter or package was recognized and a list of persons who also may have handled this package or letter. Give the list to both the local public health authorities and law enforcement officials.

Managing Exposures

Identification of a patient with anthrax or a confirmed exposure to *B. anthracis* should prompt an epidemiologic investigation. The highest priority is to identify at-risk persons and initiate appropriate interventions to protect them. The exposure circumstances are the most important factors that direct decisions about prophylaxis. Persons with an exposure or contact with an item or environment known, or suspected to be contaminated with *B. anthracis*—regardless of laboratory tests results—should be offered antimicrobial prophylaxis. Exposure or contact, not laboratory test results, is the basis for initiating such treatment. Culture of nasal swabs is used to detect anthrax spores. Nasal swabs can occasionally document exposure, but cannot rule out exposure to *B. anthracis*. As an adjunct to epidemiologic evaluations, nasal swabs may provide clues to help assess the exposure circumstances. In addition, rapid evaluation of contaminated powder, including particle size and characteristics, may prove useful in assessing the risk for inhalational anthrax.

CDC is working with U.S. Postal Service employees and managers on several strategies to address the risk for anthrax among workers involved in mail handling. These strategies include personal protective equipment for workers handling mail and engineering controls in mail facilities. Clinicians and laboratorians should be vigilant for symptoms or laboratory findings that indicate possible anthrax infection, particularly among workers involved in mail sorting and distribution. Information to guide health-care providers and laboratories is available at <http://www.bt.cdc.gov>(1).

Antimicrobial Treatment

A high index of clinical suspicion and rapid administration of effective antimicrobial therapy is essential for prompt diagnosis and effective treatment of anthrax. Limited clinical experience is available and no controlled trials in humans have been performed to validate current treatment recommendations for inhalational anthrax. Based on studies in nonhuman primates and other animal and in vitro data, ciprofloxacin or doxycycline should be used for initial intravenous therapy until antimicrobial susceptibility results are known (Table 1). Because of the mortality associated with inhalational anthrax, two or more antimicrobial agents predicted to be effective are recommended; however, controlled studies to support a multiple drug approach are not available. Other agents with in vitro activity suggested for use in conjunction with ciprofloxacin or doxycycline include rifampin, vancomycin, imipenem, chloramphenicol, penicillin and ampicillin, clindamycin, and clarithromycin; but other than for penicillin, limited or no data exist regarding the use of these agents in the treatment of inhalational *B. anthracis* infection. Cephalosorins and trimethoprim-sulfamethoxazole should not be used for therapy. Regimens being used to treat patients described in this report include ciprofloxacin, rifampin, and vancomycin; and ciprofloxacin, rifampin, and clindamycin.

Penicillin is labelled for use to treat inhalational anthrax. However, preliminary data indicate the presence of constitutive and inducible beta-lactamases in the *B. anthracis* isolates from Florida, NYC, and DC. Thus, treatment of systemic *B. anthracis* infection using a penicillin alone (i.e., penicillin G and ampicillin) is not recommended. The *B. anthracis* genome sequence shows that this organism encodes two beta-lactamases: a penicillinase and a cephalosporinase. Data in the literature also show that some beta-lactamase negative *B. anthracis* strains for which the penicillin MICs are 0.06 µg/mL increase to 64 µg/mL and become beta-lactamase positive when exposed to semisynthetic penicillins (4). The frequency of this induction event is unknown. Although

Category	Initial therapy (intravenous) ^{s,1}	Duration
Adults	Ciprofloxacin 400 mg every 12 hrs* or Doxycycline 100 mg every 12 hrs ^{+†} and One or two additional antimicrobials [¶]	IV treatment initially**. Switch to oral antimicrobial therapy when clinically appropriate: Ciprofloxacin 500 mg po BID or Doxycycline 100 mg po BID Continue for 60 days (IV and po combined) ^{§§}
Children	Ciprofloxacin 10–15 mg/kg every 12hrs ¶**** or Doxycycline: ^{†††,††} >8 yrs and >45 kg: 100 mg every 12 hrs >8 yrs and ≤45 kg: 2.2 mg/kg every 12 hrs ≤8 yrs: 2.2 mg/kg every 12 hrs and One or two additional antimicrobials [¶]	IV treatment initially**. Switch to oral antimicrobial therapy when clinically appropriate: Ciprofloxacin 10–15 mg/kg po every 12 hrs*** or Doxycycline: ⁺⁺⁺ >8 yrs and >45 kg: 100 mg po BID >8 yrs and ≤45 kg: 2.2 mg/kg po BID ≤8 yrs: 2.2 mg/kg po BID
Pregnant women⁵⁵⁵	Same for nonpregnant adults (the high death rate from the infection outweighs the risk posed by the antimicrobial agent)	IV treatment initially. Switch to oral antimicrobial therapy when clinically appropriate. [†] Oral therapy regimens same for nonpregnant adults
Immunocompromised persons	Same for nonimmunocompromised persons and children	Same for nonimmunocompromised persons and children
 For gastrointestina Ciprofloxacin or d anthrax. Steroids may be co experience with ba Other agents with imipenem, clindam in <i>Bacillus anthraci</i> specialist is advise Initial therapy may ciprofloxacin or do If meningitis is sus tion. Because of the po continued for 60 da If intravenous ciprof absorbed from the concentrations are 	In and oropharyngeal anthrax, use regimens recomm loxycycline should be considered an essential para posidered as an adjunct therapy for patients with ser- acterial meningitis of other etiologies. In <i>in vitro</i> activity include rifampin, vancomycin, j pycin, and clarithromycin. Because of concerns of co- is, penicillin and ampicillin should not be used alone ed. y be altered based on clinical course of the patient paycycline) may be adequate as the patient improve pected, doxycycline may be less optimal because of tential persistence of spores after an aerosol exper- ays. of loxacin is not available, oral ciprofloxacin may be gastrointestinal tract with no substantial loss by fi attained 1–2 hours after oral dosing but may not be	nenged for innalational anthrax. art of first-line therapy for inhalational vere edema and for meningitis based or penicillin, ampicillin, chloramphenicol, institutive and inducible beta-lactamases . Consultation with an infectious disease ; one or two antimicrobial agents (e.g. es. of poor central nervous system penetra- posure, antimicrobial therapy should be acceptable because it is rapidly and well rst-pass metabolism. Maximum serum achieved if vomiting or ileus are present

TABLE 1. Inhalational anthrax treatment protocol^{*,†} for cases associated with this bioterrorism attack

*** In children, ciprofloxacin dosage should not exceed 1 g/day.
 *** The American Academy of Pediatrics recommends treatment of young children with tetracyclines for serious infections (e.g., Rocky Mountain spotted fever).
 *** Although tetracyclines are not recommended during pregnancy, their use may be indicated for life-threatening

^{§§§} Although tetracyclines are not recommended during pregnancy, their use may be indicated for life-threatening illness. Adverse effects on developing teeth and bones are dose related; therefore, doxycycline might be used for a short time (7–14 days) before 6 months of gestation.

amoxicillin/clavulanic acid is more active than amoxicillin alone against beta-lactamase, producing strains *in vitro*, the combination may not be clinically effective for inhalational anthrax where large numbers of organisms are likely to be present.

Toxin-mediated morbidity is a major complication of systemic anthrax. Corticosteroids have been suggested as adjunct therapy for inhalational anthrax associated with extensive edema, respiratory compromise, and meningitis (5).

For cutaneous anthrax, ciprofloxacin and doxycycline also are first-line therapy (Table 2). As for inhalational disease, intravenous therapy with a multidrug regimen is recommended for cutaneous anthrax with signs of systemic involvement, for extensive edema, or for lesions on the head and neck (Table 2). In cutaneous anthrax, antimicrobial treatment may render lesions culture negative in 24 hours, although progression to eschar formation still occurs (5). Some experts recommend that corticosteroids be considered for extensive edema or swelling of the head and neck region associated with cutaneous anthrax. Cutaneous anthrax is typically treated for 7–10 days; however, in this bioterrorism attack, the risk for simultaneous aerosol exposure appears to be high. Although infection may produce an effective immune response, a potential for reactivation of latent infection may exist. Therefore, persons with cutaneous anthrax associated with this attack should be treated for 60 days.

Category	Initial therapy (oral) [†]	Duration
Adults*	Ciprofloxacin 500 mg BID or Doxycycline 100 mg BID	60 days⁵
Children*	Ciprofloxacin 10–15 mg/kg every 12 hrs (not to exceed 1 g/day) [†] or Doxycycline: [¶] >8 yrs and >45 kg: 100 mg every 12 hrs >8 yrs and <u><</u> 45 kg: 2.2 mg/kg every 12 hrs <u><</u> 8 yrs: 2.2 mg/kg every 12 hrs	60 days⁵
Pregnant women*,**	Ciprofloxacin 500 mg BID or Doxycycline 100 mg BID	60 days⁵
Immunocompromised persons*	Same for nonimmunocompromised persons and children	60 days⁵

TABLE 2. Cutaneous anthrax treatment protocol* for cases associated with this bioterrorism attack

 Cutaneous anthrax with signs of systemic involvement, extensive edema, or lesions on the head or neck require intravenous therapy, and a multidrug approach is recommended. Table 1.
 [†] Ciprofloxacin or doxycycline should be considered first-line therapy. Amoxicillin 500 mg po TID for adults or 80

[†] Ciprofloxacin or doxycycline should be considered first-line therapy. Amoxicillin 500 mg po TID for adults or 80 mg/kg/day divided every 8 hours for children is an option for completion of therapy after clinical improvement. Oral amoxicillin dose is based on the need to achieve appropriate minimum inhibitory concentration levels.

[§] Previous guidelines have suggested treating cutaneous anthrax for 7–10 days, but 60 days is recommended in the setting of this attack, given the likelihood of exposure to aerosolized *B. anthracis* (6).

¹ The American Academy of Pediatrics recommends treatment of young children with tetracyclines for serious infections (e.g., Rocky Mountain spotted fever).

** Although tetracyclines or ciprofloxacin are not recommended during pregnancy, their use may be indicated for life-threatening illness. Adverse effects on developing teeth and bones are dose related; therefore, doxycycline might be used for a short time (7–14 days) before 6 months of gestation.

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Prophylaxis for inhalational anthrax exposure has been addressed in a previous report (1) and indicates the use of either ciprofloxacin or doxycycline as first line agents. High-dose penicillin (e.g., amoxicillin or penicillin VK) may be an option for antimicrobial prophylaxis when ciprofloxacin or doxycycline are contraindicated. The likelihood of beta-lactamase induction events that would increase the penicillin MIC is lower when only small numbers of vegetative cells are present, such as during antimicrobial prophylaxis.

All medications may have undesirable side effects and allergic reactions may result from any medication. Clinicians prescribing these medications should be aware of their side effects and consult an infectious disease specialist as needed. Patients should be urged to inform their health-care provider of any adverse event.

This is the first bioterrorism-related anthrax attack in the United States, and the public health ramifications of this attack continue to evolve. Additional updates and recommendations will be published in *MMWR*.

References

- CDC. Update: investigation of anthrax associated with intentional exposure and interim public health guidelines, October 2001. MMWR 2001;50:889–97.
- Keim P, Price LB, Klevytska AM, et al. Multiple-locus variable-number tandem repeat analysis reveals genetic relationships with *Bacillus anthracis*. J Baceriol 2000;182:2928–36.
- National Committee for Clinical Laboratory Standards. Performance standards for antimicrobial susceptibility testing. Wayne, Pennsylvania: National Committee for Clinical Laboratory Standards, 2001; 11th informational supplement M100-S11.
- Lightfoot NF, Scott RJ, Turnbull PC. Antimicrobial susceptibility of *Bacillus anthracis*. Salisbury Med Bull 1990;68:95S-98S.
- 5. Dixon TC, Meselson M, Guillemin J, Hanna PC. Anthrax. N Engl J Med 1999;341:815–26.
- Inglesby TV, Henderson DA, Bartlett JG, et al. Anthrax as a biological weapon: medical and public health management. JAMA 1999;281:1735–45.

Methicillin-Resistant *Staphylococcus aureus* Skin or Soft Tissue Infections in a State Prison — Mississippi, 2000

On October 25, 2000, the Mississippi State Department of Health (MSDH) notified CDC that, since November 1999, 31 inmates had acquired methicillin-resistant *Staphylococcus aureus* (MRSA) skin or soft tissue infections at a state prison. During November 1998–October 1999, no MRSA infections had been reported at the prison, which houses approximately 1,200 female and 1,800 male inmates. This report summarizes the case investigation and the nasal culture prevalence survey conducted by MSDH and CDC during November 2000. Findings indicate that MRSA infections were transmitted person-to-person within the prison, and that the number of asymptomatic carriers was unexpectedly high for a nonhealth-care setting. Correctional facilities can reduce the increasing prevalence of MRSA disease by identifying and appropriately treating infected persons and by instituting prevention measures.

A case of MRSA infection was defined as a skin or soft tissue lesion occurring in a state prison inmate with symptoms (e.g., pus, pain, warmth, or tenderness) and with MRSA cultured from the site of infection during November 1999–November 2000. Cases were identified by interviews with physicians and inmates and a review of the prison's medical, laboratory, and pharmacy records. Fifty-nine inmates had an illness that met

Methicillin-Resistant Staphylococcus aureus - Continued

the case definition (Figure 1); 46 (78%) were women, and the median age was 33 years (range: 19–70 years). The median length of incarceration was 397 days (range: 3–3,717 days).

Records of 45 (76%) infected inmates were reviewed. Three (7%) had been hospitalized during the year preceding infection. Twenty-six (58%) had infections on the legs and seven (16%) on the arms. Fifteen (33%) were diagnosed with furuncles, 12 (27%) with skin abscesses, and 11 (24%) with open wounds; 21 (47%) had cellulitis, and two (4%) had systemic infections requiring hospitalization. Infections resolved after a median of 3 weeks (range: 1–36 weeks). Systemic antimicrobials were used to treat 44 (98%) infected inmates, 35 (78%) received topical antimicrobials, six (13%) required incision

FIGURE 1. Number of cases of methicillin-resistant *Staphylococcus aureus** (MRSA) in a state prison, by month and year of onset — Mississippi, November 1999–November 2000[†]



Month and Year of Onset

* Defined as a skin or soft tissue lesion occurring in a state prison inmate with symptoms (e.g., pus, pain, warmth, or tenderness) and with MRSA cultured from the site of infection during November 1999–November 2000.

† n=59.

Methicillin-Resistant Staphylococcus aureus - Continued

and drainage, and wound dressing was prescribed for 21 (47%). Nineteen (90%) of the 21 infected inmates with wound dressings changed their dressings themselves. During interviews, 15 (33%) infected inmates reported helping or being helped by other inmates with wound care or dressing changes. Twenty-six (58%) reported lancing their own boils or other inmates' boils with fingernails or tweezers; 40 (89%) shared personal items (e.g., linen, pillows, clothing, and tweezers) that potentially were contaminated by wound drainage.

To assess the extent of MRSA carriage among the inmates, swab specimens of both anterior nares were collected from all female and a one third systematic sample of male inmates. Of 1,757 inmates sampled, 86 (4.9%) were MRSA carriers. More women (73 of 1,241 [5.9%]) were carriers than men (13 of 516 [2.5%]) (p=0.003), and inmates who had been incarcerated for >60 days were more likely to be carriers (84 of 1,565 [5.4%]) than those who had served less time (one of 142 [0.7%]) (p=0.01).

Of the 59 infection-associated isolates, 41 (69%) were tested and genotyped at CDC. All 41 isolates were confirmed as MRSA and 40 (98%) were susceptible to gentamicin, rifampin, trimethoprim-sulfamethoxazole, clindamycin, vancomycin, and chloramphenicol; three (7%) were resistant to levofloxacin. Pulsed-field gel electrophoresis of isolates revealed that three MRSA strains predominated: genotype A (24 [59%]), genotype B (seven [17%]), and genotype C (four [10%]).

During December 2000, CDC and MSDH provided the Mississippi State Department of Corrections and the prison with control measures such as optimizing antimicrobial treatment of infected inmates, reinforcing infection control practices (e.g., implementing Standard Precautions [1] at prison clinics, educating inmates in personal hygiene and wound care), using antibacterial soap, and establishing an MRSA skin infection surveillance system.

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Editorial Note: *S. aureus* is an important and common pathogen in humans. It is found in the nose or on the skin of many healthy, asymptomatic persons (i.e., carriers) and can cause infections with clinical manifestations ranging from pustules to sepsis and death. Most transmission occurs through the contaminated hands of a person infected with or carrying *S. aureus*. MRSA infections frequently are encountered in health-care settings (*2*). Since the 1960s, treatment of these infections has become more difficult because *S. aureus* has progressively acquired resistance to previously effective antimicrobial agents (*2*). In 1999, 2,538 (53.5%) of 4,744 intensive care unit patients with hospital-acquired *S. aureus*-associated infection had MRSA (*3*). Less information is available on long-term–care facilities, where prevalence of MRSA carriage may range from zero to 33% of the residents (*4*).

Risk factors for infection with MRSA in health-care settings include prolonged hospital stay, exposure to multiple or prolonged broad-spectrum antimicrobial therapy, stay in an intensive care or burn unit, proximity to patients colonized or infected with MRSA, use of invasive devices, surgical procedures, underlying illnesses, and MRSA nasal carriage (5).

Although community-onset MRSA infections have been reported recently (6), little is known about their epidemiology or prevalence of carriage. Community outbreaks have occurred among injection-drug users; aboriginals in Canada, New Zealand, and Australia; Native Americans/Alaska Natives in the United States; and players of close-contact

Methicillin-Resistant Staphylococcus aureus — *Continued*

sports (6). Reported most commonly have been uncomplicated skin infections; however, community-acquired MRSA infections can be severe. Four deaths from community-acquired MRSA in children were reported in Minnesota and North Dakota in 1999 (7).

Disease transmission can occur easily among inmates at correctional facilities. In 1999, approximately two million persons were incarcerated in the United States (8). Skin or soft tissue infections are recognized problems in these facilities (9). MRSA disease in prisons can be controlled or prevented using several approaches. First, severe skin disease or treatment failures of presumed *S. aureus* skin infection should be evaluated with appropriate cultures or other diagnostic tests. Efforts to monitor the etiology of skin disease should be linked to these data to determine whether MRSA is a problem in the facility. MRSA outbreaks can be reported to CDC (telephone [800] 893-0485) through state departments of corrections and state health departments. Second, optimal treatment of MRSA disease should be based on the infecting organism's antimicrobial susceptibility result and, when available, input by infectious disease expertise. Third, close contact among inmates may place them at increased risk for transmission of skincolonizing or skin-infecting organisms. To prevent skin disease, all inmates should practice good personal hygiene, including daily showers. Inmates should avoid touching wounds or drainage of others and should have access to sinks and plain soap (in this setting, the usefulness of antibacterial soap is unknown). Hands should be washed with soap as soon as possible after touching wounds or dressings. Personnel that provide wound care should follow Standard Precautions (1).

References

- 1. Garner JS. Guideline for isolation precautions in hospitals. Hospital Infection Control Practices Advisory Committee. Infect Control Hosp Epidemiol 1996;17:53–80.
- 2. Lowy FD. Staphylococcus aureus infections. N Engl J Med 1998;339:520-32.
- CDC. Semiannual report: aggregated data from the National Nosocomial Infections Surveillance system. Available at http://www.cdc.gov/ncidod/hip/SURVEILL/NNIS.HTM. Accessed September 2001.
- 4. Strausbaugh LJ, Jacobson C, Sewell DL, Potter S, Ward TT. Methicillin-resistant Staphylococcus aureus in extended-care facilities: experiences in a Veterans' Affairs nursing home and a review of the literature. Infect Control Hosp Epidemiol 1991;12:36–45.
- Herwaldt LA. Control of methicillin-resistant Staphylococcus aureus in the hospital setting. Am J Med 1999;106:11S-18S,48S-52S.
- 6. Cookson BD. Methicillin-resistant *Staphylococcus aureus* in the community: new battlefronts, or are the battles lost? Infect Control Hosp Epidemiol 2000;21:398–403.
- 7. CDC. Four pediatric deaths from community-acquired methicillin-resistant *Staphylococcus aureus*—Minnesota and North Dakota, 1997–1999. MMWR 1999;48:707–10.
- 8. Beck AJ. Bureau of Justice Statistics Bulletin, prisoners in 1999. Washington, DC: US Department of Justice, Office of Justice Programs, Bureau of Justice Statistics, 2000:16.
- 9. Duncan WC, Dodge BG, Knox JM. Prevention of superficial pyogenic skin infections. Arch Dermatol 1969;99:465-8.

Shigella sonnei Outbreak Among Men Who Have Sex with Men — San Francisco, California, 2000–2001

Shigella sonnei causes approximately 10,000 cases of gastroenteritis each year in the United States (1). These infections occur predominately among young children and usually are associated with poor hygienic conditions in child-care settings. Outbreaks of

Shigella sonnei Outbreak - Continued

shigellosis among men who have sex with men (MSM) have occurred because of direct or indirect oral-anal contact (2,3) but usually are caused by *Shigella flexneri* (4). This report describes an investigation of *S. sonnei* cases that occurred among MSM in San Francisco during May–December 2000. Following efforts to heighten awareness, the number of cases has declined, but new cases continue to occur at low levels in this risk group (Figure 1). The increased incidence of sexually transmitted *Shigella* and other sexually transmitted diseases (STDs) in MSM require renewed and innovative prevention efforts.

During June–December 2000, 230 cases of culture-confirmed* *S. sonnei* infection were reported to the San Francisco Department of Public Health; an average of 21 cases (range: 13–29 cases) occurred during the same period from 1996 to 1999. Based on data obtained from 230 reported cases, the median age was 39 years (range: 16–77 years) and 211 (92%) patients were males. Of 199 males for whom information was available, 141 (71%) were non-Hispanic whites, 159 (80%) were residents of predominantly gay neighborhoods, and 121 (61%) were self-reported MSM. Sexual behavior was unknown for 62 (31%) patients, and 16 (8%) were self-reported heterosexuals. On the basis of denominator data obtained from the annual San Francisco HIV/AIDS epidemiology report, the rate of *S. sonnei* infection among MSM was 259 per 100,000 population. The rate among all other groups, including women and heterosexual men, was 16 (*5*).

FIGURE 1. Number of adult *Shigella sonnei* infections, by month, year, and sex — San Francisco, California, January 2000–September 2001



Month and Year

^{*}Defined as culture-confirmed *S. sonnei* infection in residents of San Francisco County aged \geq 15 years.

Shigella sonnei Outbreak — Continued

Among persons aged \geq 18 years with *S. sonnei* and symptom onset during May– December 2000, 106 were selected randomly for telephone interview; 35 (33%) could not be contacted and four (4%) refused to participate. Of the 67 (63%) who agreed to participate, 64 (96%) were male. Among the 64 male respondents, 62 (97%) were MSM, 42 (66%) were college graduates, and 29 (46%) had an annual income >\$45,000. Of the respondents, 49 (78%) had health insurance coverage, 45 (70%) thought they became ill from a sexual partner, and 35 (55%) reported concurrent infection with human immunodeficiency virus (HIV).

The median duration of symptoms for male respondents was 7 days (range: 2–90 days); 62 (97%) reported diarrhea, 50 (78%) abdominal cramps, 49 (77%) fever, 47 (73%) weight loss, and 20 (31%) blood in stool.

In the week before illness, 50 (78%) of the 64 males reported being sexually active, including 34 (53%) who had multiple sex partners; 32 (50%) answered "yes" to, "The week before your illness did you put your tongue in a partner's anus?" Forty-seven (73%) answered "yes" to, "The week before your illness did you have a penis in your mouth?"

Of the 14 patients who reported sexual activity during the week of or the week following illness, three (21%) answered "yes" to, "During [or after] your illness did you have a tongue in your anus?" All 14 persons who were sexually active during and after illness reported diarrhea (duration: 3–23 days) for which they were prescribed antibiotics.

Local response to the outbreak included a press release, development of an Internet web site, and a media campaign with newspaper and Internet articles for the gay community. Approximately 2,000 notices were mailed to community agencies and providers, 10 presentations were conducted for community agencies, and 4,000 health alerts were distributed through a mass mailing to 40 acquired immunodeficiency syndrome-related agencies and their clients, several large gay and lesbian fairs, bars, sex clubs, and the city STD clinic.

Free *Shigella* screening was offered for 1 month at the city STD clinic. Of 119 patients screened, five reported having diarrhea at presentation to the STD clinic. Two of the five had *S. sonnei* isolated from their rectal swab samples; no *Shigella* species were isolated from the 114 remaining clients.

A convenience sample of *S. sonnei* from outbreak-related patients and controls (women and children with *S. sonnei* infection in the outbreak period and region) was subtyped by pulsed-field gel electrophoresis (PFGE). Of 26 outbreak-related isolates, 23 (88%) shared one of two closely related patterns, and only one (12%) of eight isolates from controls had a similar PFGE patterns.

Of 20 randomly selected isolates from outbreak-related patients, 19 were resistant to trimethoprim-sulfamethoxazole, tetracycline, ampicillin, sulfisoxazole, and streptomycin. All isolates were susceptible to ciprofloxacin, nalidixic acid, and ceftriaxone.

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Editorial Note: This report indicates that *S. sonnei* can cause large community outbreaks through sexual transmission among MSM. The strains circulating among MSM were different from those circulating in the rest of the community, indicating unique

Shigella sonnei Outbreak — Continued

transmission. The recent increases in STDs and enteric infections in MSM follow a 10-year decline (4). The rate of *S. sonnei* remained low in MSM until the summer of 2000 in San Francisco. These trends paralleled changes in sexual behavior that increased the risk for HIV and other STDs (6).

Approximately half of the patients in this report were infected with HIV compared with an estimated prevalence of 20% among MSM in San Francisco (7), suggesting that MSM with HIV infection are more likely to participate in sexual behaviors that place them at risk for shigellosis. Standard HIV management includes stool bacteria cultures of persons with diarrhea. However, HIV-infected persons with shigellosis might have more severe illness (8) leading to more frequent diagnosis and reporting.

The findings in this report are subject to at least two limitations. First, approximately a third of the selected cases could not be contacted, and those who were might have had difficulty accurately recalling events that occurred up to 6 months preceding the interview. Second, the magnitude of this outbreak probably was underestimated because reporting shigellosis in California is required of physicians but not of laboratories, and many cases probably were undiagnosed and unreported.

Because most patients in this outbreak were sexually active with multiple partners, the potential for ongoing transmission is high. In San Francisco and other communities with high rates of shigellosis in adult men, clinicians should obtain stool cultures and sexual orientation data from men with diarrhea and report suspected cases of shigellosis to the health department. Appropriate antimicrobial therapy will decrease the duration, transmission, and severity of symptoms and should be prescribed based on the severity of illness or the need to protect close contacts. Patients in certain occupations (i.e., foodhandlers, child-care providers, and health-care workers) and children who attend child care often are required to have a negative stool culture documented following treatment. The incubation period of shigellosis is 1–4 days, and *shigellae* are shed in stool from several days to several weeks after illness. Persons who receive appropriate antimicrobial therapy will be culture negative at 72 hours (9).

Patients with shigellosis should be counseled to abstain from sexual behavior that is likely to transmit infection for at least 3 days after starting an appropriate course of antimicrobial therapy (9). Because antimicrobial resistance is common, in cases in which antimicrobial susceptibility data are not available, patients should be counseled on abstaining from high-risk sexual behavior until at least one negative posttreatment stool culture is obtained. Patients also should be counseled on methods to avoid or reduce the risk for sexual transmission of enteric infections such as *Shigella* and hepatitis A, should be educated to avoid sexual practices that might result in fecal-oral transmission, and should be advised to wash with soap and water the perianal/perineal area, other body parts, and sex toys before and after sexual activity.

References

- 1. CDC. *Shigella* surveillance: annual tabulation summary, 1999. Atlanta, Georgia: US Department of Health and Human Services, CDC, 2000.
- Bader M, Pedersen AHB, Williams R, Spearman J, Anderson H. Venereal transmission of shigellosis in Seattle-King County. STD 1977;4:89–91.
- Vugia DJ, Shallow S, Samuel MC, Aragon T, Reingold AL, Bradford WZ. Risk factors for shigellosis in San Francisco adults. 38th annual meeting of the Infectious Diseases Society of America, New Orleans, Louisiana, 2000, 507.
- Department of Public Health, City and County of San Francisco. Shigellosis in San Francisco, 1977–1985. San Francisco Epidemiologic Bulletin 1986;2:1–3.

Shigella sonnei Outbreak — Continued

- Department of Public Health, City and County of San Francisco. 2000 HIV/AIDS Epidemiology Annual Report. San Francisco Department of Public Health, HIV Seroepidemiology and AIDS Surveillance Section, 2000, 1–58. Available at http://www.dph.sf.ca.us/PHP/AIDSSurvUnit.htm. Accessed October 2001.
- CDC. Increases in unsafe sex and rectal gonorrhea among men who have sex with men— San Francisco, California, 1994–1997. MMWR 1999;48:45–8.
- 7. Catania JA, Osmond D, Stall RD, et al. The continuing HIV epidemic among men who have sex with men. Am J Public Health 2001;91:907–14.
- 8. Baer JT, Vugia DJ, Reingold AL, Aragon T, Angula FJ, Bradford WZ. HIV infection as a risk factor for shigellosis. Emerg Infect Dis 1999;5:820–3.
- Lolekha S, Vibulbandhitkit S, Poonyarit P. Response to antimicrobial therapy shigellosis in Thailand. Reviews of Infectious Diseases 1991 March-April;13 Suppl 4:S342-6.

Weekly Update: West Nile Virus Activity — United States, October 17–23, 2001

The following report summarizes West Nile virus (WNV) surveillance data reported to CDC through ArboNET and verified by states and other jurisdictions as of October 23, 2001.

During the week of October 17–23, six human cases of WNV encephalitis or meningitis were reported in Pennsylvania (three), New Jersey (two), and Florida (one). During the same period, WNV infections were reported in 101 crows, 45 other birds, and 26 horses. A total of 31 WNV-positive mosquito pools were reported in five states (Connecticut, Florida, Georgia, New York, and Ohio).

During 2001, 37 human cases of WNV encephalitis or meningitis have been reported in Florida (10), Maryland (six), New York (six), New Jersey (six), Connecticut (five), Pennsylvania (three), and Georgia (one); one death occurred in Georgia. Among these 37 cases, 20 (54%) were in males, the median age was 69 years (range: 36–81 years), and dates of illness onset ranged from July 13 to October 7. A total of 3,796 crows and 1,394 other birds with WNV infection were reported from 25 states and the District of Columbia (Figure 1); 151 WNV infections in other animals (all horses) were reported from 11 states (Alabama, Connecticut, Florida, Georgia, Kentucky, Louisiana, Massachusetts, Mississippi, New York, Pennsylvania, and Virginia); and 725 WNV-positive mosquito pools were reported from 14 states (Connecticut, Florida, Georgia, Illinois, Kentucky, Maryland, Massachusetts, Michigan, New Hampshire, New Jersey, New York, Ohio, Pennsylvania, and Rhode Island).

Additional information about WNV activity is available at <http://www.cdc.gov/ncidod/ dvbid/westnile/index.htm> and <http://cindi.usgs.gov/hazard/event/west_nile/ west_nile.html>. Update: West Nile Virus - Continued





* As of October 23, 2001.

[†] Mississippi reported WNV infection in a horse but no birds.

Notice to Readers

National Lead Poisoning Prevention Week — October 21–27, 2001

October 21–27 is National Lead Poisoning Prevention Week (NLPPW), and this year's theme is "Treat Yourself to Lead-Safe Living: Harvest the Rewards." Childhood lead poisoning is considered the most preventable environmental disease of young children, but approximately one million children have elevated blood lead levels. One of the national health objectives for 2010 is to eliminate childhood lead poisoning in the United States (objective 8-11) (1). The goal of NLPPW is 1) to raise awareness about this serious health issue and the importance of screening at-risk children at aged 1–2 years and children aged 3–5 years who have not been screened previously and 2) to urge persons to take precautions to minimize exposure to lead.

In commemoration of NLPPW, events such as state proclamations, free screenings, lead-awareness community events, and educational campaigns will be conducted nationwide. CDC, the Environmental Protection Agency, and the U.S. Department of Housing and Urban Development are collaborating to coordinate activities and offer assistance to campaigns at the local level. Additional information about NLPPW activities is available from state or local health departments.

Additional information about preventing childhood lead poisoning is available at http: //cdc.gov/nceh/lead or from the National Lead Information Center, telephone (800) 424-LEAD ([800] 424-5323).

Notices to Readers — Continued

Reference

1. US Department of Health and Human Services. Healthy people 2010 (conference ed, 2 vols). Washington, DC: US Department of Health and Human Services, 2000.

Notice to Readers

Availability of Final Recommendations on Reducing the Risk for Transmission of Enteric Pathogens at Petting Zoos, Open Farms, Animal Exhibits, and Other Venues

Final Recommendations on "Reducing the Risk for Transmission of Enteric Pathogens at Petting Zoos, Open Farms, Animal Exhibits, and Other Venues Where the Public Has Contact With Farm Animals" are available on the Internet. Draft recommendations were published in *MMWR* on April 20, 2001. Readers were invited to submit comments and suggestions before July 1. Twenty-six submissions were received and reviewed. The final recommendations are posted under "Outbreak Reports and Publications" at http:// www.cdc.gov/ncidod/dbmd/outbreak/recomm_farm_animal.htm.



FIGURE I. Selected notifiable disease reports, United States, comparison of provisional 4-week totals ending October 20, 2001, with historical data

- * No rubella cases were reported for the current 4-week period yielding a ratio for week 42 of zero (0).
- [†] 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.

		Cum. 2001		Cum. 2001
Anthrax		8	Poliomyelitis, paralytic	-
Brucellosis [†]		73	Psittacosis [†]	17
Cholera		3	Qfever [†]	18
Cyclosporiasis	[†]	121	Rabies, human	1
Diphtheria		2	Rocky Mountain spotted fever (RMSF)	443
Ehrlichiosis:	human granulocytic (HGE)†	168	Rubella, congenital syndrome	-
	human monocytic (HME) [†]	70	Streptococcal disease, invasive, group A	2,927
Encephalitis:	California serogroup viral [†]	72	Streptococcal toxic-shock syndrome [†]	47
·	eastern equine ^Ť	7	Syphilis, congenital [¶]	166
	St. Louis [†]	1	Tetanus	22
	western equine [†]	-	Toxic-shock syndrome	96
Hansen diseas	se (leprosy) [†]	70	Trichinosis	21
Hantavirus pu	Imonary syndrome [†]	7	Tularemia [†]	90
Hemolytic ure	mic syndrome, postdiarrheal [†]	119	Typhoid fever	212
HIV infection,	pediatric ^{t§}	153	Yellow fever	-
Plague	-	2		

TABLE I. Summary of provisional cases of selected notifiable diseases, United States, cumulative, week ending October 20, 2001 (42nd Week)*

-: No reported cases. *Incidence data for reporting year 2001 are provisional and cumulative (year-to-date).

[†] Not notifiable in all states.

⁵ Updated monthly from reports to the Division of HIV/AIDS Prevention — Surveillance and Epidemiology, National Center for HIV,

STD, and TB Prevention (NCHSTP). Last update September 25, 2001. ¹Updated from reports to the Division of STD Prevention, NCHSTP.

		20					Escherichia coli O157:H7 [†]			7 [†]
	All Cum.	Cum.	Chlan Cum.	nydia ^s Cum.	Cryptos Cum.	Cum.	Cum.	Cum.	Cum.	LIS Cum.
	2001	2000	2001	2000	2001	2000	2001	2000	2001	2000
NITED STATES NEW ENGLAND Maine N.H. Vt. Vt. Mass. R.I. Conn.	29,580 1,129 36 31 13 602 78 369	29,975 1,586 27 27 29 998 75 430	561,232 18,403 877 1,093 491 7,747 2,379 5,816	560,703 18,613 1,174 885 428 7,925 2,158 6,043	2,346 99 16 10 30 39 4	2,493 124 18 21 26 32 3 24	2,385 210 25 31 13 109 12 20	3,843 334 25 31 31 151 18 78	1,936 204 26 24 8 105 10 31	3,139 349 27 32 33 160 16 81
MID. ATLANTIC Upstate N.Y. N.Y. City N.J. Pa.	6,710 731 3,385 1,389 1,205	6,678 662 3,609 1,295 1,112	61,857 10,951 23,969 8,694 18,243	52,600 1,801 21,493 8,800 20,506	218 86 73 7 52	321 100 152 15 54	176 136 9 31 N	385 253 21 111 N	165 121 10 34	269 59 15 111 84
E.N. CENTRAL Ohio Ind. III. Mich. Wis.	2,238 430 264 992 413 139	2,865 430 282 1,568 437 148	85,919 17,985 11,797 21,767 24,186 10,184	95,892 25,207 10,704 26,993 19,788 13,200	896 150 70 1 154 521	846 233 56 107 82 368	614 154 71 132 79 178	938 230 107 175 128 298	450 137 39 128 69 77	668 203 80 143 102 140
W.N. CENTRAL Minn. Iowa Mo. N. Dak. S. Dak. S. Dak. Nebr. Kans.	637 108 71 312 2 22 52 70	680 129 69 318 2 7 53 102	28,973 5,646 3,797 10,595 750 1,414 2,175 4,596	31,835 6,549 4,320 10,822 714 1,480 3,039 4,911	351 137 72 35 12 6 88 1	261 55 70 27 9 15 76 9	389 151 74 42 17 37 51 17	551 138 167 96 15 53 57 25	394 186 59 68 30 40 - 11	529 169 137 86 19 57 45 16
S. ATLANTIC Del. Md. D.C. Va. W. Va. N.C. S.C. Ga. Fla.	9,497 203 1,506 644 723 61 726 577 1,031 4,026	8,257 156 1,056 569 556 46 505 639 991 3,739	107,104 2,041 9,096 2,372 14,718 1,874 16,445 9,110 22,775 28,673	106,106 2,328 11,513 2,584 12,569 1,745 18,185 7,831 22,427 26,924	268 6 32 10 22 24 - 103 69	398 5 9 13 16 3 21 - 147 184	192 4 23 - 47 10 41 9 26 32	317 2 30 1 61 14 77 21 35 76	120 6 1 36 8 28 11 15 15	255 1 55 11 65 16 36 70
E.S. CENTRAL Ky. Tenn. Ala. Miss.	1,423 278 456 347 342	1,529 159 635 417 318	39,174 7,205 11,713 10,731 9,525	41,189 6,394 11,893 12,800 10,102	39 4 12 13 10	44 5 11 15 13	115 57 35 16 7	118 39 48 8 23	95 46 36 6 7	97 31 47 9 10
W.S. CENTRAL Ark. La. Okla. Tex.	3,141 159 665 186 2,131	3,006 149 493 259 2,105	83,888 5,903 14,077 8,325 55,583	84,931 5,414 14,928 7,501 57,088	32 6 7 12 7	143 10 10 16 107	82 11 4 25 42	211 54 13 17 127	86 - 25 24 37	261 37 44 15 165
MOUNTAIN Mont. Idaho Wyo. Colo. N. Mex. Ariz. Utah Nev.	1,073 14 17 3 231 103 437 90 178	1,105 11 19 7 259 116 348 108 237	32,592 1,542 1,492 660 6,963 4,738 11,575 1,572 4,110	31,184 1,104 1,468 656 8,763 4,045 10,220 1,722 3,206	178 28 20 6 34 21 7 58 4	146 10 19 5 60 14 10 24 4	236 16 54 81 13 22 30 15	373 30 60 17 143 19 44 47 13	120 - 1 53 9 22 34 1	271 - 34 9 103 16 34 65 10
PACIFIC Wash. Oreg. Calif. Alaska Hawaii	3,732 395 154 3,112 16 55	4,269 379 113 3,669 15 93	103,322 10,941 5,853 81,303 2,150 3,075	98,353 10,410 5,322 77,714 2,014 2,893	265 43 43 175 1 3	210 U 16 194 -	371 103 61 186 4 17	616 195 125 255 27 14	302 62 57 176 1 6	440 191 107 128 3 11
Guam P.R. V.I. Amer. Samoa C.N.M.I.	10 934 2 -	13 1,023 27 -	1,930 53 U 103	413 U U U	- - U	- - U U	N 1 - U	N 6 - U U		

 TABLE II. Provisional cases of selected notifiable diseases, United States, weeks ending October 20, 2001, and October 21, 2000 (42nd Week)*

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

 * Incidence data for reporting year 2001 are provisional and cumulative (year-to-date). Incidence data for reporting year 2000 are finalized and cumulative (year-to-date).

 * Individual cases can be reported through both the National Electronic Telecommunications System for Surveillance (NETSS) and the Public Health Laboratory Information System (PHLIS).

 * Chamydia refers to genital infections caused by *C. trachomatis*.

 * Updated monthly from reports to the Division of HIV/AIDS Prevention — Surveillance and Epidemiology, National Center for HIV, STD, and TB Prevention. Last updated September 25, 2001.

	Gonorrhea		Hepati Non-A, I	tis C; Non-B	Legione	llosis	Listeriosis	Lyme Listeriosis Diseas		
Reporting Area	Cum. 2001	Cum. 2000	Cum. 2001	Cum.	Cum. 2001	Cum.	Cum. 2001	Cum. 2001	Cum. 2000	
UNITED STATES	258,213	287,298	2,643	2,587	802	884	372	10,228	13,915	
NEW ENGLAND Maine N.H. Vt. Mass. R.I. Conn.	5,298 94 152 53 2,445 662 1,892	5,258 76 89 54 2,180 519 2,340	14 - 6 8 - -	24 2 4 13 5	54 8 10 5 13 9 9	50 2 5 16 8 17	33 1 4 2 18 1 7	3,227 113 14 653 436 2,011	4,336 60 31 1,081 414 2,750	
MID. ATLANTIC Upstate N.Y. N.Y. City N.J. Pa.	30,926 6,778 10,061 5,321 8,766	31,134 5,710 9,325 5,954 10,145	1,313 51 - 1,214 48	581 33 - 511 37	159 54 16 7 82	241 72 39 20 110	57 25 8 10 14	5,182 2,871 2 927 1,382	7,352 3,173 167 2,323 1,689	
E.N. CENTRAL Ohio Ind. III. Mich. Wis.	46,030 10,140 5,132 13,370 13,787 3,601	57,559 15,392 5,046 16,964 14,469 5,688	148 8 1 13 126 -	195 10 19 166	216 103 19 - 63 31	230 93 30 28 42 37	50 13 8 1 21 7	505 100 20 - 1 384	739 55 22 33 23 606	
W.N. CENTRAL Minn. Iowa Mo. N. Dak. S. Dak. Nobr	12,410 1,846 997 6,584 33 228 710	14,428 2,566 1,014 7,100 59 251 1,211	566 9 545 - -	468 5 1 451 -	45 9 7 19 1 3	53 7 13 23 2	15 - 2 8 - -	335 279 29 22 -	280 187 29 45 1 -	
Kans.	2,012	2,227	9	4	5 1	4	4	2	15	
S. ATLANTIC Del. Md. D.C. Va. W. Va. N.C. S.C. Ga. Fla.	66,427 1,212 4,938 2,187 8,638 536 13,705 6,143 12,541 16,527	75,220 1,392 7,944 2,091 8,368 529 14,870 6,980 14,473 18,573	94 - - - 9 18 6 - 46	92 2 12 3 14 14 2 3 39	168 11 31 7 20 N 7 10 9 73	161 8 57 4 31 N 13 4 6 38	61 - 11 5 4 5 11 14	731 49 468 10 110 11 35 5 - 43	980 167 574 5 130 26 42 7 - 29	
E.S. CENTRAL Ky. Tenn. Ala. Miss.	25,342 2,873 7,894 8,308 6,267	29,697 2,839 9,500 9,894 7,464	169 8 57 4 100	385 31 80 9 265	49 11 24 12 2	30 17 9 3 1	19 5 8 6	51 22 20 8 1	47 11 28 5 3	
W.S. CENTRAL Ark. La. Okla. Tex.	41,069 3,593 9,667 3,789 24,020	44,851 3,174 11,053 3,306 27,318	171 4 83 3 81	623 8 369 8 238	5 - 2 3 -	21 - 7 2 12	17 1 - 2 14	79 - 1 - 78	74 5 7 62	
MOUNTAIN Mont. Idaho Wyo. Colo. N. Mex. Ariz. Utah Nev.	8,177 86 65 2,373 799 3,150 119 1,524	8,528 38 69 40 2,614 885 3,456 172 1,254	58 1 2 6 18 11 9 3 8	63 4 3 12 13 17 - 12	46 - 3 1 13 2 18 5 4	33 1 5 11 1 7 8 -	30 - 1 7 7 6 2 6	11 - 6 1 - - 1 2	10 - 2 3 - - 2 3	
PACIFIC Wash. Oreg. Calif. Alaska Hawaii	22,534 2,426 923 18,365 344 476	20,623 1,835 756 17,377 280 375	110 19 12 79 -	156 28 25 101 - 2	60 8 N 48 - 4	65 15 N 49 - 1	90 7 8 69 - 6	107 8 7 90 2 N	97 7 9 79 2 N	
Guam P.R. V.I. Amer. Samoa C.N.M.I.	461 6 U 10	44 412 - U U	- 1 - U	3 1 - U U	2 - U	1 - U U	- - - -	N U U	N - U U	

TABLE II. (Cont'd) Provisional cases of selected notifiable diseases, United States, weeks ending October 20, 2001, and October 21, 2000 (42nd Week)*

N: Not notifiable. U: Unavailable. - : No reported cases. * Incidence data for reporting year 2001 are provisional and cumulative (year-to-date). Incidence data for reporting year 2000 are finalized and cumulative (year-to-date).

					Salmonellosis [†]						
	Ma	aria	Rabie	s, Animal	NE	TSS	PH	ILIS			
Reporting Area	Cum. 2001	Cum. 2000	Cum. 2001	Cum. 2000	Cum. 2001	Cum. 2000	Cum. 2001	Cum. 2000			
UNITED STATES	928	1,198	5,487	5,813	29,419	31,840	24,208	27,066			
NEW ENGLAND Maine N.H. Vt. Mass. R.I. Conn.	63 4 2 1 26 7 23	64 6 1 2 30 8 17	611 58 20 56 220 56 201	678 109 19 52 224 49 225	2,030 158 151 69 1,132 113 407	1,877 108 119 99 1,086 117 348	1,946 137 137 63 1,043 150 416	1,905 88 124 95 1,080 131 387			
MID. ATLANTIC Upstate N.Y. N.Y. City N.J. Pa.	238 57 122 25 34	320 61 183 42 34	1,025 664 24 163 174	1,068 682 11 162 213	3,436 1,014 827 651 944	4,155 1,009 1,024 994 1,128	3,212 1,043 1,091 657 421	4,439 1,097 1,107 862 1,373			
E.N. CENTRAL Ohio Ind. III. Mich. Wis.	92 22 16 1 35 18	121 16 5 59 28 13	119 42 3 24 44 6	145 48 - 22 64 11	3,967 1,129 448 1,026 678 686	4,406 1,189 523 1,306 745 643	3,627 1,061 399 1,049 689 429	2,972 1,206 524 101 806 335			
W.N. CENTRAL Minn. Iowa Mo. N. Dak. S. Dak. Nebr. Kans.	30 6 11 - 2 5	47 13 2 15 2 1 8 6	289 42 71 38 33 25 4 76	477 73 69 49 106 85 2 93	1,828 487 291 515 53 139 125 218	1,999 454 305 593 48 83 192 324	2,023 609 277 763 73 111 - 190	2,184 587 295 741 68 93 132 268			
S. ATLANTIC Del. Md. D.C. Va. W. Va. N.C. S.C. Ga. Fla.	235 2 100 13 44 1 13 6 12 44	269 4 89 15 47 3 31 2 19 59	1,874 30 279 121 496 99 294 166	1,989 42 349 - 473 101 480 142 268 134	7,120 79 685 72 1,132 113 1,055 681 1,236 2,067	6,510 101 667 52 829 136 910 628 1,165 2,022	4,873 87 750 U 747 121 905 595 1,210 458	5,006 114 589 U 795 125 955 478 1,482 468			
E.S. CENTRAL Ky. Tenn. Ala. Miss.	31 12 11 6 2	42 17 11 13 1	184 29 96 57 2	175 19 90 65 1	2,135 319 526 597 693	1,974 321 514 550 589	1,600 192 663 474 271	1,534 223 687 512 112			
W.S. CENTRAL Ark. La. Okla. Tex.	11 3 4 3 1	67 3 11 8 45	876 20 57 799	763 20 3 51 689	3,163 754 313 397 1,699	4,097 605 717 326 2,449	2,068 92 566 292 1,118	2,494 492 596 251 1,155			
MOUNTAIN Mont. Idaho Wyo. Colo. N. Mex. Ariz. Utah Nev.	46 2 3 19 3 8 3 8 3 8	41 1 20 - 7 5 5	216 31 28 20 - 14 108 14 14 1	241 60 9 50 - 19 85 10 8	1,778 60 116 50 495 243 512 179 123	2,288 79 103 55 609 199 585 420 238	1,451 4 43 484 205 517 175 23	2,144 95 47 591 182 630 419 180			
PACIFIC Wash. Oreg. Calif. Alaska Hawaii	182 9 10 153 1 9	227 24 35 158 10	293 3 253 37	277 7 244 26	3,962 429 202 2,989 34 308	4,534 474 253 3,555 52 200	3,408 491 2,71 2,335 28 283	4,388 566 311 3,272 33 206			
Guam P.R. V.I. Amer. Samoa C.N.M.I.	3 - U -	2 5 - U U	73 - U	65 - U U	455 - U 11	22 556 - U U	U U U U U	U U U U			

TABLE II. (Cont'd) Provisional cases of selected notifiable diseases, United States, weeks ending October 20, 2001, and October 21, 2000 (42nd Week)*

N: Not notifiable. U: Unavailable. -: No reported cases. * Incidence data for reporting year 2001 are provisional and cumulative (year-to-date). Incidence data for reporting year 2000 are finalized and cumulative (year-to-date).

¹ Individual cases can be reported through both the National Electronic Telecommunications System for Surveillance (NETSS) and the Public Health Laboratory Information System (PHLIS).

	NET	Shige SS	llosis⁺ P	HLIS	Sy (Primary 8	philis Secondary)	Tuberculosis		
Demostin a Anna	Cum.	Cum.	Cum.	Cum.	Cum.	Cum.	Cum.	Cum.	
UNITED STATES	2001 14.325	2000 18.128	<u>2001</u> 6.689	10.403	<u>2001</u> 4.639	2000 4.937	9.543	2000 11.364	
NEW ENGLAND Maine N.H. Vt. Mass. R.I. Conn.	218 6 7 171 17 11	344 10 6 4 244 24 56	239 2 3 5 164 23 42	331 11 8 223 26 63	49 - 1 2 27 9 10	70 1 49 4 15	334 8 13 4 193 29 87	337 16 16 4 194 27 80	
MID. ATLANTIC Upstate N.Y. N.Y. City N.J. Pa.	1,076 424 286 185 181	2,188 618 859 466 245	669 101 319 184 65	1,408 184 587 400 237	420 24 220 115 61	230 9 96 59 66	1,783 280 869 396 238	1,809 241 974 430 164	
E.N. CENTRAL Ohio Ind. III. Mich. Wis.	3,578 2,491 184 395 262 246	3,557 311 1,335 1,036 587 288	1,585 1,047 34 288 192 24	1,038 253 139 61 536 49	768 68 135 229 316 20	1,001 64 293 348 254 42	1,040 191 83 500 203 63	1,140 235 112 534 187 72	
W.N. CENTRAL Minn. Iowa Mo. N. Dak. S. Dak. Nebr. Kans.	1,486 360 335 276 20 372 63 60	2,021 668 437 592 16 7 106 195	1,102 384 276 174 27 206 35	1,730 747 304 418 49 4 98 110	74 26 4 21 - 5 18	58 15 10 26 - - 2 5	363 176 34 109 3 12 29	411 128 33 151 2 14 19 64	
S. ATLANTIC Del. Md. D.C. Va. W. Va. N.C. S.C. Ga. Fla.	1,986 14 128 50 286 8 290 223 250 737	2,446 20 172 67 375 4 298 111 209 1,190	639 10 78 U 124 8 143 112 130 34	1,004 20 97 U 313 3 238 81 157 95	1,628 9 191 43 90 3 374 197 299 422	1,639 8 248 34 111 3 405 185 314 331	1,919 15 176 51 194 26 274 153 365 665	2,269 14 199 24 215 24 276 223 500 794	
E.S. CENTRAL Ky. Tenn. Ala. Miss.	1,297 599 83 184 431	925 380 305 63 177	480 236 85 130 29	481 85 341 49 6	511 39 263 97 112	730 67 437 104 122	652 90 237 220 105	767 98 295 251 123	
W.S. CENTRAL Ark. La. Okla. Tex.	1,899 485 121 56 1,237	2,827 170 236 99 2,322	1,098 155 137 17 789	896 50 145 38 663	582 28 134 58 362	680 84 182 100 314	750 123 115 512	1,666 157 146 126 1,237	
MOUNTAIN Mont. Idaho Wyo. Colo. N. Mex. Ariz. Utah Nev.	793 4 33 204 109 326 49 65	966 7 43 5 209 123 397 70 112	564 - 1 213 72 224 46 8	711 25 3 172 96 276 73 66	197 - 1 35 18 126 8 8	192 - 1 8 15 161 1 5	379 6 8 3 90 24 166 30 52	417 14 7 2 69 36 165 41 83	
PACIFIC Wash. Oreg. Calif. Alaska Hawaii	1,992 171 72 1,686 6 57	2,854 395 151 2,270 7 31	313 167 91 - 6 49	2,804 364 98 2,310 3 29	410 41 13 346 - 10	337 53 11 272 - 1	2,323 193 84 1,887 40 119	2,548 195 81 2,074 88 110	
Guam P.R. V.I. Amer. Samoa C.N.M.I.	- 8 - U 4	34 29 Ū U	U U U U	U U U U	- 172 - U 4	3 127 - U U	- 76 U 23	47 119 - U U	

TABLE II. (Cont'd) Provisional cases of selected notifiable diseases, United States, weeks ending October 20, 2001, and October 21, 2000 (42nd Week)*

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 * Individual cases can be reported through both the National Electronic Telecommunications System for Surveillance (NETSS) and the Public Health Laboratory Information System (PHLIS).

	H. influ	uenzae,	Hepatitis (Viral), By Type				Measles (Rubeola)					
	Inva	asive	A		В	-	Indige	nous	Impo	orted⁺	Tota	l
Reporting Area	Cum. 2001 [§]	Cum. 2000	Cum. 2001	Cum. 2000	Cum. 2001	Cum. 2000	2001	Cum. 2001	2001	Cum. 2001	Cum. 2001	Cum. 2000
UNITED STATES	1,037	1,008	8,271	10,682	5,226	5,673	-	49	-	42	91	71
NEW ENGLAND	73	83	510	319	82	93	-	4	-	1	5	6
Naine N.H.	2	12	10	17 18	5 14	5 15	-	-	-	-	-	- 3
Vt. Mass	3 35	7 36	12 222	8 120	4	6 13	-	1	-	- 1	1	3
R.I.	3	4	46	22	25	18	-	- 1	-	-	-	-
Conn.	26	23	204	134	31	30	-	1	-	-	1	-
Upstate N.Y.	58	79	207	207	114	949 107	-	4	-	4	5	10
N.Y. City	37 38	50 35	227 159	425 241	338 169	461 149	-	2	-	1 1	3 1	10
Pa.	20	24	180	377	217	232	-	1	-	5	6	1
E.N. CENTRAL	140	151	918 100	1,384	724	593	-	-	-	10	10	7
Ind.	43	40 26	92	87	42	93 41	-	-	-	3 4	3	-
III. Mich	10 8	51 9	304 278	603 399	124 469	104 320	-	-	-	3	3	3
Wis.	22	20	54	72		35	-	-	-	-	-	-
W.N. CENTRAL	54	61	342	587	166	244	-	4	-	-	4	1
lowa	-	-	30	61	20	27	-	-	-	-	-	-
Mo. N. Dak.	13 7	19 2	91 3	238 3	88 1	121 2	-	2	-	-	2	-
S. Dak.	- 1	1	2	1	1	1 27		-		-	-	-
Kans.	1	4	152	93	16	22	-	-	-	-	-	-
S. ATLANTIC	304	231	1,940	1,184	1,158	1,004	-	4	-	1	5	3
Del. Md.	- 73	- 70	227	13 173	118	13	-	2	-	- 1	- 3	-
D.C. Va	- 25		43 110	23 129	11 145	27 136	-	- 1	-	-	- 1	- 2
W. Va.	14	8	18	52	20	11	-	-	-	-	-	-
N.C. S.C.	42 5	20 7	173 65	121 69	1/3	205 13	-	-	-	-	-	-
Ga.	72 73	55 36	752	223 381	305 360	162 330	-	1	-	-	1	- 1
E S CENTRAL	63	39	323	347	361	376	_	2		_	2	-
Ky.	2	12	114	44	41	64	-	2	-	-	2	-
Ala.	33 26	9	68	46	73	48	-	-	-	-	-	-
Miss.	2	2	16	136	54	88	-	-	-	-	-	-
W.S. CENTRAL Ark.	37	61 2	1,136 61	2,009 121	536 80	944 85	-	1	-	-	1	-
La.	3	16	56	72	39	132	-	-	-	-	-	-
Tex.	- 34	2	914	1,598	347	597	-	1	-	-	1	-
MOUNTAIN	122	100	630	738	419	425	-	1	-	1	2	12
Mont. Idaho	- 1	1 4	10 53	7 22	3 10	6 6	-	-	-	- 1	- 1	-
Wyo.	-	1	7	4	2	3	-	-	-	-	-	-
N. Mex.	20	25 20	78 31	62	91 124	76 117	-	-	-	-	-	- 2
Ariz.	54	35 10	342	369	128	156	, i	1	ū	-	1	- 3
Nev.	10	4	49	56	38	41	-	-	-	-	-	7
PACIFIC	91	94	1,699	2,864	942	1,045	-	29	-	18	47	21
Oreg.	3 17	5 28	67	148	86	87 93	-	4	-	2 -	4	- 3
Calif. Alaska	43	33	1,495 14	2,451 11	715 9	844 10	-	10	-	11	21	14 1
Hawaii	22	22	3	13	16	11	-	2	-	5	7	3
Guam P B	- 1	1	- 01	1 217	-	9	U	-	U	-	-	- 2
V.I.	-	4 	-	-	-	200	Ŭ	-	Ŭ	-		-
Amer. Samoa C.N.M.I.	U -	U	U -	U U	U 28	U	U	U -	U	U -	U -	U

TABLE III. Provisional cases of selected notifiable diseases preventable
by vaccination, United States, weeks ending October 20, 2001,
and October 21, 2000 (42nd Week)*

N: Not notifiable. U: Unavailable. -: No reported cases. * Incidence data for reporting year 2001 are provisional and cumulative (year-to-date). Incidence data for reporting year 2000 are finalized and cumulative (year-to-date). * For imported measles, cases include only those resulting from importation from other countries. * Of 219 cases among children aged <5 years, serotype was reported for 113, and of those, 20 were type b.

	Mening Dis	gococcal ease	Mumps				Pertussis		Rubella		
Reporting Area	Cum. 2001	Cum. 2000	2001	Cum. 2001	Cum. 2000	2001	Cum. 2001	Cum. 2000	2001	Cum. 2001	Cum. 2000
UNITED STATES	1,737	1,802	5	179	273	91	3,784	5,495	-	20	146
NEW ENGLAND Maine N.H. Vt. Mass. R.I. Conn.	95 4 12 5 49 4 21	112 8 11 3 64 9 17			4 - - 1 1 2		333 21 26 27 237 5 17	1,399 41 102 205 995 16 40			12 - 2 - 8 1 1
MID. ATLANTIC Upstate N.Y. N.Y. City N.J. Pa.	181 51 33 43 54	205 63 38 39 65	- - -	19 3 9 3 4	22 9 6 3 4	3 3 - -	253 127 38 18 70	558 275 73 30 180	- - - -	5 1 3 1	9 1 8 - -
E.N. CENTRAL Ohio Ind. III. Mich. Wis.	231 75 36 22 56 42	322 77 36 76 95 38	1 - 1 - -	17 1 2 11 3	20 7 1 6 5 1	29 - 7 3 19 -	569 257 74 62 115 61	620 263 86 90 76 105		3 - 1 2 -	1 - - 1 -
W.N. CENTRAL Minn. Iowa Mo. N. Dak. S. Dak. Nebr. Kans.	124 18 25 44 6 5 12 14	126 18 28 60 2 5 6 7	- - - - - U	7 3 - - 1 3	17 - 7 4 1 - 2 3	35 35 - - - U -	241 105 19 86 4 4 4 4 19	460 282 46 65 6 4 21 36	- - - - U	3 - 1 - - - 1	1 - - - - 1
S. ATLANTIC Del. Md. D.C. Va. W. Va. N.C. S.C. Ga. Fla	326 4 37 - 35 12 60 31 40 107	254 1 26 - 37 12 34 20 42 82	1 - - 1 - - -	34 - 6 - 5 5 7 5	39 - 9 - 5 10 2 4	12 - - 5 - 7	204 31 1 36 2 63 31 14 26	399 8 104 3 90 1 77 26 35 55		6 1 - - - 2 - 3	94 1 - - 64 27 - 2
E.S. CENTRAL Ky. Tenn. Ala. Miss.	117 19 55 30 13	119 25 48 33 13	- - -	6 1 1 - 4	5 1 2 2	- - -	124 31 55 34 4	99 50 29 17 3	- - - -	- - - -	6 1 1 4
W.S. CENTRAL Ark. La. Okla. Tex.	193 18 58 26 91	188 11 42 25 110	- - - -	10 1 2 7	29 1 5 23	7 1 - 6	374 25 2 11 336	316 33 19 21 243		1 - - 1	8 1 1 - 6
MOUNTAIN Mont. Idaho Wyo. Colo. N. Mex. Ariz. Utah Nev.	83 4 7 5 29 12 13 7 6	76 4 7 26 7 22 7 3	- - - - - U	11 1 1 2 1 3	18 - 1 - 1 4 5 6	3 - - 3 - U -	1,138 31 168 1 227 129 498 71 13	643 35 57 4 373 81 63 18 12	- - - - - U	1 - - 1 - - - -	2 - - 1 - 1 - -
PACIFIC Wash. Oreg. Calif. Alaska Hawaii	387 58 34 281 2 12	400 46 55 283 8 8 8	3 - N 3 -	75 1 N 37 1 36	119 9 N 82 8 20	2 2 - - -	548 132 44 334 7 31	1,001 337 101 507 19 37	- - - -	1 - - - 1	13 7 6 -
Guam P.R. V.I. Amer. Samoa C. N.M.I.	- 4 - U	- 9 - U		- - - U	14 - - U U		2 - U	3 7 - U		- - - U	1 - - U

TABLE III. (Cont'd) Provisional cases of selected notifiable diseases preventable by vaccination, United States, weeks ending October 20, 2001, and October 21, 2000 (42nd Week)*

N: Not notifiable.

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		All Cau	ises, By	Age (Ye	ears)		P&I⁺	P&I [†]		All Causes, By Age (Years)					P&I⁺
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	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 Waterbury, Conn.	416 U 35 14 28 59 55 18 59 58 44 59 5 6 36	311 U 27 11 27 38 21 13 12 36 40 25 25	63 U 5 3 1 16 2 3 3 2 8 3 7 3	29 U 3 - 32 1 1 1 8 - 32	9 U - 2 - 1 - 2 2 - 1 - 1	4 U - - 3 1 -	¥∪32323127 · · 31	S. ATLANTIC Atlanta, Ga. Baltimore, Md. Charlotte, N.C. Jacksonville, Fla. Morfolk, Va. Richmond, Va. Savannah, Ga. St. Petersburg, F Tampa, Fla. Washington, D.C Wilmington, D.C	1,240 144 164 88 167 148 76 55 55 55 55 55 55 55 55 55 55 55 55 55	818 84 99 57 107 97 53 38 40 43 122 65 13	272 41 37 17 41 35 13 11 13 10 31 23	88 13 16 8 16 10 4 2 1 1 9 8 -	354942233 - 53 - 53 - 1	26 2 2 2 1 4 3 4 1 - 3 4 -	78 5 16 10 9 14 1 8 - 6 9 -
Worcester, Mass. MID. ATLANTIC Albany, N.Y. Allentown, Pa. Buffalo, N.Y. Camden, N.J. Elizabeth, N.J. Erie, Pa.S	61 2,560 45 15 96 24 18 29	48 1,547 36 13 75 17 10 22	7 513 7 1 15 5 6 5 7	5 415 1 4 1 2 1	1 51 - - 1 - 1	- 33 1 - 2 - -	16 122 5 9 4 -	E.S. CENTRAL Birmingham, Ala Chattanooga, Te Knoxville, Tenn. Lexington, Ky. Memphis, Tenn. Mobile, Ala. Montgomery, Al Nashville, Tenn.	825 a. 186 nn. 56 75 66 149 72 a. 50 171	553 132 42 46 40 95 53 30 115	165 28 11 18 14 39 11 14 30	68 20 5 6 8 4 3 21	21 4 2 1 4 3 3 2	14 2 - 5 3 1 - 3	48 9 4 7 4 8 2 2 12
New York City, N.J. New York City, 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. Yonkers, N.Y.	42 42 42 42 42 42 42 42 42 42	819 U 14 176 21 105 21 26 82 29 15 U	7 340 55 55 9 2 27 3 6 11 7 2 U	361 U 26 1 1 6 2 2 4 1 1 U	31 U 10 - 1 1 2 2 U	16 U 2 5 - 4 - 3 - U	51U2195173583 U	W.S. CENTRAL Austin, Tex. Baton Rouge, La Corpus Christi, T Dallas, Tex. El Paso, Tex. Ft. Worth, Tex. Houston, Tex. Little Rock, Ark. New Orleans, La. San Antonio, Te: Shreveport, La. Tulsa, Okla.	1,369 74 83 67 206 69 113 358 U U x. 196 78 125	895 52 48 47 125 48 73 220 U U 136 56 90	305 11 20 17 55 18 25 81 U U 38 17 23	117 7 11 19 2 10 41 U 16 3 8	30 2 3 1 5 1 3 9 U 5 1 -	22 2 2 2 2 7 U U 1 1 4	94 6 - 4 15 3 7 25 U U 11 12 11
E.N. CENTRAL Akron, Ohio Canton, Ohio Chicago, III. Cincinnati, Ohio Cleveland, Ohio Columbus, Ohio Dayton, Ohio Detroit, Mich. Evansville, Ind. Fort Wayne, Ind. Gary, Ind. Grand Rapids, Mid Indianapolis, Ind. Lansing, Mich. Milwaukee, Wis. Peoria, III. Rockford, III. South Bend, Ind. Toledo, Ohio Youngstown, Ohio	1,662 55 37 U 42 138 173 138 43 43 66 188 43 66 188 89 90 61 67 54 54 115 0 76	1,156 38 28 0 27 85 121 107 107 30 51 7 45 125 43 79 39 45 36 89 45	$\begin{array}{c} 324\\ 3213\\ 6\\ U\\ 1\\ 32\\ 36\\ 16\\ 5\\ 7\\ 6\\ 7\\ 28\\ 12\\ 16\\ 15\\ 16\\ 15\\ 1\\ 18\\ 18\\ 18\\ \end{array}$	115 2 U 2 13 12 7 23 5 5 2 6 14 3 5 1 4 4 3 2	33 	34 2 1 U 1 5 - 2 4 1 2 - 5 3 - 1 1 2 3 1	124 3 6 U 4 7 6 8 4 4 8 4 4 2 9 4 7 8 5 3 2	MOUNTAIN Albuquerque, N Boise, Idaho Colo. Springs, C Denver, Colo. Las Vegas, Nev. Ogden, Utah Phoenix, Ariz. Pueblo, Colo. Salt Lake City, U Tucson, Ariz. PACIFIC Berkeley, Calif. Fresno, Calif. Glendale, Calif. Honolulu, Hawa Long Beach, Cali Los Angeles, Cal Pasadena, Calif. Portland, Oreg. Sacramento, Cal	1,043 .M. 120 35 olo. 50 103 246 31 177 tah 111 139 1,607 19 148 100 if. 68 if. 310 25 129 if. 221	689 86 21 32 67 154 23 75 108 23 75 100 1,138 98 64 64 50 227 16 8 85 227	207 21 8 8 22 58 4 35 7 18 26 28 1 3 26 3 8 9 5 4 32 6 4 2 28 4 35 7 18 2 28 4 35 7 18 2 2 8 2 2 8 2 2 58 4 35 7 1 8 2 2 58 4 35 7 1 8 2 2 58 4 35 7 1 8 2 2 58 4 35 7 1 8 2 2 58 4 35 7 1 8 2 2 58 4 35 7 1 8 2 2 58 4 35 7 1 8 2 2 58 4 35 7 1 8 2 2 58 4 35 7 1 8 2 2 5 7 1 8 2 2 5 7 1 8 2 2 5 7 1 8 2 2 5 7 1 8 2 2 5 7 1 8 2 2 5 7 1 8 2 2 5 7 1 8 2 2 5 7 1 8 2 2 5 7 1 8 2 2 5 7 18 2 2 2 5 7 2 5 2 5 7 2 5 7 2 5 7 2 5 2 5	87 7 2 3 9 25 4 19 - 11 7 114 - 11 - 6 8 18 21 13	34 6 2 2 2 2 4 - 12 1 3 2 38 - 8 1 4 - 6 6	26 2535-3-44 3635-115444	65 3 1 3 3 17 7 13 1 6 1 108 3 6 2 6 7 14 2 6 22
W.N. CENTRAL Des Moines, Iowa Duluth, Minn. Kansas City, Kans. Kansas City, Mo. Lincoln, Nebr. Minneapolis, Min Omaha, Nebr. St. Louis, Mo. St. Paul, Minn. Wichita, Kans.	734 36 30 96 41 n. 205 76 109 83 37	503 26 22 12 51 29 148 62 57 64 32	147 8 6 27 8 35 8 33 12 4	52 2 3 12 3 10 3 11 5 1	11 - - 2 - 3 1 4 1 -	17 - 4 1 5 2 4 1 -	51 5 2 4 4 8 9 2 3 4 3 4	San Diego, Calif. San Francisco, C San Jose, Calif. Santa Cruz, Calif Seattle, Wash. Spokane, Wash. Tacoma, Wash. TOTAL	. 136 alif. U 176 5. 31 104 48 99 11,456 [¶]	90 U 124 24 66 40 76 7,610	26 U 36 5 22 5 13 2,277	13 U 10 2 11 1 8 1,085	3 U 4 - 3 1 2 262	4 U 2 1 - 212	9 U 14 3 4 7 733

TABLE IV. Deaths in 122 U.S. cities,* week ending October 20, 2001 (42nd Week)

U: Unavailable. * Mortality data

U: Unavailable. -:No reported cases. * Mortality data in this table are reported voluntarily 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. [¶] Total includes unknown ages.

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