Prevention and Control of Infections with Hepatitis Viruses in Correctional Settings
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Prevention and Control of Infections with Hepatitis Viruses in Correctional Settings

Prepared by
Cindy Weinbaum, M.D.
Rob Lyerla, Ph.D.
Harold S. Margolis, M.D.
Division of Viral Hepatitis
National Center for Infectious Diseases

Summary

This report consolidates previous recommendations and adds new ones for the prevention and control of infections with hepatitis viruses in correctional settings. These recommendations provide guidelines for juvenile and adult correctional systems regarding 1) identification and investigation of acute viral hepatitis; 2) preexposure and postexposure immunization for hepatitis A and hepatitis B; 3) prevention of hepatitis C virus infection and its consequences; 4) health education; and 5) release planning. Implementation of these recommendations can reduce transmission of infections with hepatitis viruses among adults at risk in both correctional facilities and the outside community. These recommendations were developed after consultation with other federal agencies and specialists in the fields of corrections, correctional health care, and public health at a meeting in Atlanta, March 5–7, 2001. This report can serve as a resource for correctional health, corrections, and public health professionals involved in planning and implementing health-care programs for incarcerated persons.

Introduction

Persons incarcerated in correctional systems comprise approximately 0.7% of the U.S. population and have a disproportionately greater burden of infectious diseases, including infections with hepatitis viruses and other infections of public health importance (e.g., human immunodeficiency virus [HIV], sexually transmitted disease [STD], and tuberculosis [TB] infections) (1). In 2000, >8 million inmates of prisons and jails were released and returned to the community (A. Beck, Bureau of Justice Statistics, personal communication, 2002). Recent estimates indicate 12%–39% of all Americans with chronic hepatitis B virus (HBV) or hepatitis C virus (HCV) infections were released from correctional custody during the previous year (1) (Table 1). The significance of including incarcerated populations in community-based disease prevention and control strategies is now recognized by public health, correctional health and corrections professionals (2,3). Improved access to medical care and prevention services for incarcerated populations can benefit communities by reducing disease transmission and medical costs (4–8). Inmates who participate in health-related programs while incarcerated have lower recidivism rates and are more likely to maintain health-conscious behaviors (4). Finally, because incarcerated persons have a high frequency of infection with hepatitis viruses, community prevention and control of these infections and their disease consequences require inclusion of this population (9–11). However, implementation of preventive health programs for incarcerated persons has substantial challenges.

Correctional staff are included in groups at potential risk for occupationally acquired infections with bloodborne pathogens. Therefore, recommendations are also reviewed for prevention and control of occupationally acquired infections with hepatitis viruses among correctional workers.

Definitions

Adolescent: Person aged ≥10 and <19 years.
Adult: Person aged ≥19 years.
Anti-HAV: Total antibody to hepatitis A virus (HAV) detected in serum of persons with acute or resolved HAV infection; indicates a protective immune response to infection, vaccination, and passively acquired antibody.
Anti-HBc: Antibody to hepatitis B core antigen; positive test indicates past or current infection with HBV.
Anti-HBs: Antibody to hepatitis B surface antigen; indicates immunity to HBV infection, either from HBV infection or immunization.
Anti-HCV: Antibody to HCV; positive test indicates past or current infection with HCV.
Arrestee: Person placed under arrest by law enforcement who has not been formally charged with a crime.

The material in this report originated in the National Center for Infectious Diseases, James M. Hughes, M.D., Director, and the Division of Viral Hepatitis, Harold S. Margolis, M.D., Director.
TABLE 1. Estimated chronic infections with hepatitis viruses among inmates and releasees — United States, 1997

<table>
<thead>
<tr>
<th>Chronic infection</th>
<th>Number of jail and prison inmates with condition*</th>
<th>Number among noninmate population with condition</th>
<th>Number among total U.S. population with condition</th>
<th>Number of releasees with condition and as percentage of U.S. population†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hepatitis B virus</td>
<td>34,000 (2%)§</td>
<td>1 million—1.25 million (0.5%)§</td>
<td>1.036 million—1.29 million</td>
<td>155,000 (12%—15%)</td>
</tr>
<tr>
<td>Hepatitis C virus</td>
<td>255,000 (15%)**</td>
<td>2.7 million (1.3%)††</td>
<td>2.97 million</td>
<td>1.3 million (39%)</td>
</tr>
</tbody>
</table>


* Based on 1.7 million inmates in prisons and jails, 1997 (15).
† Based on estimated 7.75 million unduplicated released inmates (2); A. Beck, Ph.D. Bureau of Justice Statistics, personal communication, 2002.
§ (31,83,84,85,86,88,89,90,92,94).
¶ Data from CDC, National Center for Health Statistics, National Health and Nutrition Examination Survey (NHANES III), adjusted to include persons of Asian origin (76).
** (88,121,122); L. Wang, Ph.D., New York State Department of Health, personal communication, 2001; D. Lau, M.D., University of Texas Medical Branch—Galveston, personal communication, 2001.
†† Based on data from NHANES III (107).

Body fluids, potentially infectious: Semen, vaginal secretions, cerebrospinal fluid, synovial fluid, pleural fluid, pericardial fluid, peritoneal fluid, and amniotic fluid. Potentially infectious body fluids include any body fluid visibly contaminated with blood, and all body fluids in situations where identifying blood contamination is difficult or impossible.

Detainee: Person arrested and legally charged with a crime who is held in a correctional facility before trial.

HAV: Hepatitis A virus, the infectious agent that causes HAV infection and hepatitis A.

HBIG: Hepatitis B immune globulin; sterile preparation of high-titer antibodies (immunoglobulins) to hepatitis B surface antigen obtained from pooled human plasma of immunized persons and which provides protection against HBV infection.

HBeAg: Hepatitis B e antigen; positive test correlates with HBV replication and infectivity.

HBsAg: Hepatitis B surface antigen; positive test indicates an active HBV infection.

HBV: Hepatitis B virus, the infectious agent that causes HBV infection, hepatitis B, and chronic liver disease.

HBV DNA: Deoxyribonucleic acid from HBV; positive test indicates active infection.

HCC: Hepatocellular carcinoma; a primary liver cancer caused by chronic HBV or HCV infection that is usually fatal.

HCV: Hepatitis C virus, the infectious agent that causes HCV infection, hepatitis C, and chronic liver disease.

HCV RNA: Ribonucleic acid from HCV; positive test indicates active infection.

HDV: Hepatitis D virus, a viroid (incomplete virus) that requires an active (acute or chronic) HBV infection to replicate and cause delta hepatitis virus infection, delta hepatitis, and chronic liver disease.

IDUs: Injection-drug users; persons who have ever used needles to inject illicit drugs.

IgM anti-HAV: Immunoglobulin M antibody to HAV; positive test indicates acute HAV infection.

IgM anti-HBe: Immunoglobulin M antibody to hepatitis B core antigen; positive test indicates acute HBV infection.

Immune globulin (IG): Sterile preparation of antibodies (immunoglobulins) made from pooled human plasma that contains anti-HAV and provides protection against hepatitis A.

Infant: Person aged ≤1 year.

Inmate: Incarcerated person.

Jail: Locally operated correctional facility that confines persons pending arraignment, awaiting trial and sentencing, or, after sentencing, to serve their sentence (generally >1 year).

Juvenile: Person aged <19 years, in custody of the legal system.

Prison: Adult correctional facility under the jurisdiction of state or federal authorities that confines persons with a sentence of >1 year.

Seroconversion: The change of a serologic test from negative to positive.

Sero protection: Level of antibodies necessary to protect against infection.

Correctional Populations

Juveniles

In 1997, approximately 12% of persons aged 16 years reported at least one arrest (12). In 1999, a reported 108,965 juvenile offenders were held in residential placement facilities (13). In 1994, the average length of stay in public facilities for juvenile releases was 2 weeks for those detained and 5 months for those committed; the stay in private facilities (primarily a committed population) averaged 3.5 months (12). Of arrested juveniles not incarcerated, the majority are diverted to alternative programs (teen courts, restorative justice) where they remain under supervision of the juvenile justice system.
Approximately 74% of incarcerated juvenile offenders are held in public facilities, the rest in facilities operated by private contractors (14). Adult jails hold >7,600 juveniles and approximately 3,100 are held in adult prisons (15). Females account for 27% of juveniles arrested and 13% of those in residential placement (14,16). Of juveniles arrested in 1999, approximately 72% were white, 25% black, and 3% of other races. However, a disproportionate number of racial and ethnic minorities were detained in residential placement (40% black and 18% Hispanic).

Adults

At the end of 2001, adult jail and prison populations totaled 1.96 million — a 71% increase from 1990 (13). Prior incarceration as juveniles was reported by 9% of adults in federal prisons and 20% in state prisons (17). According to 2000 data, racial/ethnic minorities were overrepresented, with 46% black, 36% white, 16% Hispanic, and 2% other races. Approximately 6.6% of adult inmates are female, a 111% increase since 1990; of incoming women to state prisons, 5% are pregnant (18). Among adult U.S. residents, approximately 1 in every 112 men and 1 in every 1,724 women were sentenced to state or federal prisons in 2001 (13).

The estimated 12.6 million admissions and 12.6 million releases from local jails, and 625,000 admissions and 606,000 releases from prisons represent annual turnover rates in these facilities of 1300% and 40% respectively (1,15) (A. Beck, Ph.D., Bureau of Justice Statistics, personal communication, 2002).

Staff

In 2000, >457,000 custody and security officers worked in the U.S. correctional system, including both public and private sectors (19). These officers comprise approximately two-thirds of all correctional staff, which also includes professional, technical, educational, clerical, maintenance, food service, and administrative workers (20,21).

Health Care in the Correctional System

Upon incarceration, all adults and the majority of juveniles lose access to the usual public and private health-care and disease-prevention services. Their health care becomes the sole responsibility of either the correctional system (federal, tribal, state, or local), or less frequently, the public health system (22). For the majority of persons, entry into the correctional system provides the opportunity to access health care. In one series, approximately 78% of newly incarcerated females had abnormal Papanicolaou smears, and >50% had vaginal infections or STDs (23). However, the rapid turnover of the incarcerated population, especially in the jail setting, and the suboptimal funding of correctional health and prevention services, often limits the correctional system in providing both curative and preventive care.

Infectious diseases — including acquired immune deficiency syndrome (AIDS), STDs, TB, and viral hepatitis — are more prevalent among correctional inmates than the general population. In 1997, an estimated 46,000–76,000 prison and jail inmates had serologic evidence of syphilis; 8,900 had AIDS (4% of the U.S. AIDS burden); and 1,400 had active TB (4% of the annual U.S. TB burden) (1).

Among incarcerated persons, shared risk factors (e.g., injection-drug use) can result in populations coinfected with HBV, HCV, or HIV. Coinfections can make treatment of chronic viral hepatitis, AIDS, and TB more difficult because of the need to use multiple drugs, which increases the chance of hepatotoxicity and other adverse events. In addition, both TB chemoprophylaxis and HIV postexposure prophylaxis can be complicated by the presence of chronic liver disease (24,25).

Risk Factors for Viral Hepatitis Transmission among Incarcerated Persons

Drug Use

During 1990–1999, the rate of arrest for substance abuse violations among persons aged 10–17 years increased by 132% (12,26). Injection-drug use is reported by 3.3%–6% of incarcerated juveniles (A. Thomas, M.D., Oregon Health Division; and R. Bair, M.D., Bexar County Juvenile Detention Center, San Antonio, Texas, personal communications, 2001). Among juvenile detainees, 53% of males and 38% of females tested positive for marijuana use at the time of arrest, ≤17% tested positive for cocaine, and ≤18% were positive for methamphetamine (27).

Arrested adults also have a high prevalence of illicit drug use. In 2000, 21% of state prisoners and 59% of federal prisoners were incarcerated for drug offenses (13). In 1997 inmate surveys, 83% of state prisoners and 73% of federal prisoners reported past drug use, and 57% of state prisoners and 45% of federal prisoners reported using drugs in the month before their offense (28). Among jail inmates, drug use in the month before incarceration was reported by 55%, and injection-drug use was reported by 18% (29). However, urine testing at entry has indicated drug use might be substantially
underreported by jail inmates (30). Injection-drug use during incarceration has been reported by 3%–28% of adult inmates (31–34). While some correctional systems offer substance-abuse treatment and education programs, the demand generally exceeds the capacity of existing programs (20). There appear to be no comprehensive risk-reduction programs available within correctional facilities.

**Sexual Behavior**

All states have laws prohibiting sex between adult residents of correctional systems (35). Despite these laws, 2%–30% of inmates have sex while incarcerated (31,36–38). Outbreaks of syphilis and hepatitis B among inmates reflect sexual activity in correctional facilities (31,33,39,40). Although two state prison systems and five city/county correctional systems make condoms available to adult inmates and detainees for use in their facilities (Vermont, Mississippi, New York City, Philadelphia, San Francisco, Washington D.C., Los Angeles), no juvenile correctional systems are known to provide condoms (E. Dunlap, National Juvenile Detention Association, personal communication, 2001).

**Percutaneous Exposures of Uncertain Risk**

Percutaneous exposures have the potential to transfer infectious blood and transmit bloodborne pathogens. Tattoos and other percutaneous exposures (e.g., bites and abrasions) are common in correctional facilities and have the potential to expose residents and correctional staff to blood and body fluids (34,41,42). Case-control studies indicate tattooing is not a risk factor for acquiring acute hepatitis B or hepatitis C (43,44). However, results from seroprevalence studies of noninstitutionalized populations have been variable, and studies of highly select groups might not be relatable to other populations (45). One study of a limited number of injection-drug users suggested an increased risk for both HBV and HCV infection among those tattooed while in prison (46), but limited studies of both adult and juvenile inmate populations have not confirmed this finding (33) (R. Bair, M.D., Bexar County Juvenile Detention Center, San Antonio, Texas, personal communication, 2001).

**Occupational Exposures**

Correctional employees have reported injuries from human bites, needles, and other sharp instruments, as well as skin and mucous membrane exposures to blood and body fluids (41,42). Occupational transmission of HBV infection among hospital-based workers has been linked to percutaneous and mucous membrane exposures, and HBV infection has been primarily associated with percutaneous exposure. Transmissions of HBV and HCV infections have not been associated with intact skin exposures (10,47). Limited data from correctional workers have indicated 21% reported blood contact with intact skin, and 7% reported a percutaneous exposure (including needle stick, cut with a contaminated object, or bite) or mucous membrane exposure (48).

**Epidemiology and Outcome of Infection with Hepatitis Viruses**

**Hepatitis A Virus Infection**

HAV infection is acquired by the fecal-oral route, produces a self-limited disease that does not result in chronic infection or long-term liver disease, and usually produces symptoms of acute viral hepatitis among adolescents and adults after an average incubation period of 28 days (range: 15–50 days). Signs and symptoms usually last <2 months, although 10%–15% of symptomatic persons have prolonged or relapsing disease lasting ≤6 months (49). Peak infectivity occurs during the 2-week period before the onset of jaundice or elevation of liver enzymes, when the concentration of virus in stool is highest (11). Persons with chronic liver disease who acquire hepatitis A are at increased risk for fulminant hepatitis (50).

**Epidemiology of HAV Infection**

In the United States, the majority of cases of hepatitis A occur through person-to-person transmission during communitywide outbreaks (11,51). Viral transmission can occur through close personal contact (e.g., household contact, sexual contact, drug use, or children playing), and contaminated food or water (e.g., infected food-handlers, or raw shellfish). Viremia occurs during HAV infection, and transmission has occurred from parenteral blood exposure (blood transfusion, injection-drug use) on occasion (56). The most frequently reported source of infection (12%–26%) is household or sexual contact with a person with HAV infection; however, 45%–50% of patients have no identified source for their infection (51,52). Historically, the highest rates of disease have occurred in 11 western U.S. states and selected counties, and accounted for approximately 50% of cases during 1987–1997 (11,52).

HAV infection is common among injection-drug users. Injection drug use has been reported by 5%–19% of hepatitis A patients. In certain communities, hepatitis A outbreaks involving users of injected and noninjected methamphetamine have accounted for approximately 30% of reported cases (11,51,53,54). Cross-sectional serologic surveys demonstrate...
that users of illicit drugs have a higher prevalence of infection than the general U.S. population (11,55). Although there is potential for parenteral transmission during HAV infection, the majority of transmissions among users of illicit drugs is believed to occur through fecal contamination of drug paraphernalia and subsequent percutaneous inoculation, as well as from close personal contact with household or other contacts (56,57).

Hepatitis A outbreaks among men who have sex with men (MSM) are frequently reported, and cyclic outbreaks occur in urban areas of the United States (58,59). HAV-infected MSM report more frequent oral-anal contact, longer duration of sexual activity, and a larger number of sex partners than persons without serologic evidence of infection (60–63).

HAV Infection in Correctional Settings

No hepatitis A outbreaks have been reported from correctional settings, although a substantial proportion of incarcerated persons have risk factors for infection (e.g., drug use or MSM). The prevalence of prior HAV infection among incarcerated persons is estimated at 22%–39%, which is similar to age-adjusted prevalence rates in the general U.S. population (11) (C. Shapiro, M.D., CDC, personal communication, 2002; T. Lincoln, M.D., Hampden County Correctional Center, Ludlow, Massachusetts; and D. Lau, M.D., University of Texas Medical Branch—Galveston; personal communications, 2001). Employment in a correctional setting has not been identified as a risk factor for HAV infection.

Hepatitis B Virus Infection

HBV is a bloodborne pathogen, transmitted by percutaneous or permucosal (e.g., sexual) exposure to infectious blood or body fluids (e.g., semen, or saliva). HBV circulates in high titers in the blood and lower titers in other body fluids (e.g., semen, vaginal fluid, or saliva), and is approximately 100 times more infectious than HIV and 10 times more infectious than HCV (47).

Acute hepatitis B develops in approximately 30%–50% of adults at the time of the initial infection and is characterized by anorexia, nausea, vomiting, and often jaundice. The risk of progression to chronic infection varies with age, being highest among young children and infants (30%–90% respectively) and lowest among adolescents and adults (2%–6%) (64).

The majority of persons with chronic HBV infection are asymptomatic and one third have no evidence of liver disease, despite high levels of viral replication in hepatocytes (65). The remainder have chronic hepatitis (mild, moderate, or severe) that can lead to cirrhosis and HCC. Persons with chronic HBV infection have a 15%–25% lifetime risk of death from chronic liver disease or HCC (66–70). Rates of progression to cirrhosis and HCC vary according to age at acquisition of chronic infection, HBeAg status; coinfection with hepatitis delta virus (HDV), HIV, HCV, and alcohol abuse (69,71–75). HBV-related liver disease and HCC cause approximately 3,000 deaths in the United States annually (S. Goldstein, M.D., CDC, unpublished data, 2002).

Epidemiology of HBV Infection

An estimated 5% of the civilian, noninstitutionalized U.S. population has serologic evidence of past or present HBV infection, and 0.4%–0.5% have chronic infection and serve as the primary source of infection for others (9,76). Overall prevalence of HBV infection differs among racial/ethnic populations and is highest among persons who have immigrated from areas with a high endemicity of HBV infection (e.g., Asia, Pacific Islands, Africa, and the Middle East) (77). Prevalence of infection among blacks is four times the prevalence among whites (11.9% compared with 2.6%) (76).

The incidence of reported cases of acute hepatitis B declined by 76% during 1987–1998 (8). Nonetheless, an estimated 78,000 persons were infected with HBV in 2001 (G. Armstrong, M.D., CDC, unpublished data, 2002). Disease incidence is highest among blacks, followed by Hispanics and whites, and highest among persons aged 25–39 years (5,52). The age of newly infected persons has increased from a median of 27 years during 1982–1988 to 32 years during 1994–1998, probably as a result of vaccination of adolescents and young adults, and changes in high-risk behaviors in certain populations (8). Before national prevention programs that began in 1990, perinatal and early childhood transmission accounted for 30% of chronic HBV infections (78).

Sex is the predominant mode of HBV transmission among adults and adolescents, accounting for more than half of all newly acquired infections (8). Among reported cases of acute hepatitis B, approximately 40% reported heterosexual exposure to an infected partner or to multiple partners, and 15% were MSM. In addition, 14% of persons with acute hepatitis B reported injection-drug use. Thirty-three percent of persons with acute hepatitis B cannot identify a risk factor for infection, although approximately 50% of those persons have a past history of known risk factors for infection (8).

HBV Infection in Correctional Settings

Juveniles. The majority of juvenile offenders have behaviors that place them at risk for HBV infection (e.g., injection-drug use or unprotected sex with multiple partners). The prevalence of past HBV infection among noninstitutionalized high-risk juveniles (e.g., homeless, drug-using, or HIV-positive) ranges from 3.6%–19% (79–81) (B.M. Beech, Ph.D,
University of Memphis, Tennessee, 2002), compared with the <3% prevalence of infection among adolescents in the general population (76,82). Among incarcerated juveniles, prevalence of past HBV infection ranges from 0% to 6% (79,82) (A. Thomas, M.D., Oregon Health Division; and R. Bair, M.D., Bexar County Juvenile Detention Center, San Antonio, Texas, personal communications, 2001). HBV transmission has not been observed in juvenile correctional settings.

Adults. The prevalence of serologic markers for current or past HBV infection among prison inmates is 13%–47%, and varies by region. Prevalence is higher among women (37%–47%) than men (13%–32%) (31,83–88) (T. Lincoln, M.D., Hampden County Correctional Center, Ludlow, Massachusetts, 2001). Chronic HBV infection is diagnosed in 1.0%–3.7% of prison inmates, 2–6 times the national prevalence estimate of 0.5% (31,83,86,88–94), and comparable to rates of chronic infection among injection-drug users (5%–10%) (95–98), and among MSM (1.5%–6%) (99) (D. MacKellar, CDC, personal communication, 2002).

Upon release, susceptible inmates are often at increased risk for infection because they resume high-risk behaviors. A study of recidivist women reported an HBV seroconversion rate of 12.2/100 person-years between incarcerations (100), compared to an estimated incidence of 0.03 per 100 person-years for the U.S. population (G. Armstrong, CDC, personal communication, 2002).

The majority of HBV infections among incarcerated persons are acquired in the community. However, infection is also transmitted within correctional settings and incidence rates have ranged from 0.82% to 3.8%/year (31,34,84). Following identification of a single case of acute hepatitis B in a state prison, serologic testing identified acute HBV infection in 1.2% of the population (33,34). Highest rate of acute infection (8%) was found in the dormitory of the index case and was associated with sex with another inmate. No other risk factors were associated with infection. Acute infections were also identified in other prison dormitories, and chronic HBV infection was found in 1% of the inmate population. Serologic testing of susceptible inmates one year later found an additional 3.8% had become newly infected with HBV.

Among cases of acute hepatitis B reported to CDC’s Sentinel Counties Study of Viral Hepatitis, 5.6% have a history of incarceration during the disease incubation period (8). HBV transmission in the prison setting can occur through sexual activity, injection-drug use, and percutaneous exposures that are not apparent, as it does in households where persons with chronic HBV infection reside (101,102).

Data are lacking regarding the prevalence of HBV infection among short- and long-term residents of jails. However, the demographic and risk factor profiles of jail and prison inmates are similar, and the burden of HBV infection and risk of transmission might be expected to be similar, especially among long-term jail residents (13,15,28,29).

Correctional Staff. The overall prevalence of HBV infection was 12.6% in the only study performed among correctional workers, a rate not significantly different from that of the general population after adjusting for age and race (48). Percutaneous and mucous membrane exposures to blood were relatively infrequent and the most frequently reported exposure was blood on the skin, which was not associated with HBV infection.

Hepatitis C Virus Infection

HCV, a bloodborne pathogen, is most efficiently transmitted by direct percutaneous exposure to infectious blood. Of persons newly infected with HCV, only 20%–30% have symptoms of acute hepatitis (10,103,104). Chronic infection develops among 75%–85% of persons infected as older adults (aged >45 years) and among 50%–60% of persons infected as juveniles or young adults (105).

The majority of persons with chronic HCV infection are asymptomatic, and approximately 30% have no evidence of liver disease. Among chronically infected persons, biochemical evidence of chronic liver disease develops among 70% of those infected as adults, but (on the basis of limited data) in only 10% of those infected as juveniles (105). The risk for progression to cirrhosis also varies by age at infection, from 10%–20% among persons infected as older adults to <5% among persons infected as juveniles or younger adults. In addition to age, clinical progression is also accelerated by alcohol intake, chronic coinfection with HBV, and male sex (105). Coinfection with HIV increases HCV viral loads, the rate of progression to fibrosis and cirrhosis, and liver-related mortality (106). Hepatocellular carcinoma develops among 1%–5% of persons with chronic hepatitis C.

Epidemiology of HCV Infection

An estimated 3.9 million persons (1.8%) in the civilian, noninstitutionalized U.S. population have been infected with HCV, of whom approximately 2.7 million (1.3%) are chronically infected. In 1990, approximately two-thirds of persons infected with HCV were aged 30–49 years (107). Blacks had a higher prevalence of HCV infection than whites (3.2% compared with 1.5%), and among black males 40–49 years old, prevalence was 9.8% (107).

The highest prevalence of HCV infection (70%–90%) is reported among those persons with substantial or repeated direct percutaneous exposures to blood (e.g., injection-drug users, persons with hemophilia treated with clotting factor
concentrates that did not undergo viral inactivation, and recipients of transfusions from HCV-positive donors). Moderate infection prevalence (10%) has been reported among long-term hemodialysis patients, and lower prevalence is found among persons with high-risk sexual practices (5%) and healthcare workers (1%–2%) (10). HCV is not transmitted efficiently through occupational exposure. The risk of acquiring HCV infection from a contaminated needle stick is <2%, and transmission rarely has been documented from mucous membrane or nonintact skin exposures (47).

The highest incidence of acute hepatitis C is among persons aged 20–39 years (108,109). Blacks and whites have a similar incidence of acute disease, and incidence rates are higher among males than females. Although the incidence of acute hepatitis C has declined by >80% since 1989, primarily as a result of a decrease in cases among IDUs, the major risk factor for HCV infection remains injection-drug use, which accounts for 60% of newly reported cases (10,110,111). No association has been determined between newly acquired HCV infection and military service, medical, surgical, or dental procedures, tattooing, acupuncture, ear piercing, or foreign travel (43,44). If transmission from such exposures does occur, the frequency has been too low to detect.

Although the number of cases of acute hepatitis C among IDUs has declined dramatically since 1989, both the incidence and prevalence of HCV infection remain high among this group (98,112,113). Among IDUs, HCV is transmitted through the transfer of infected blood by sharing syringes, needles, or drug preparation equipment contaminated with the blood of an infected person (114–116). HCV infection is acquired more rapidly after the initiation of injection-drug use than either HBV or HIV infection, and the rate of HCV infection among juvenile injection-drug users is 4 times higher than the rate of HIV infection. In 1980s studies, approximately 80% of newly initiated injection-drug users were infected with HCV ≤2 years (98,117,118). This rapid acquisition of HCV infection is probably caused by the high prevalence of chronic HCV infection among IDUs, which results in a greater likelihood of exposure to an HCV-infected person through sharing of needles, syringes, and other drug paraphernalia. More recent studies suggest the rate of HCV acquisition has slowed so that approximately one third of IDUs are infected ≤2 years, however, incidence remains high at 10%–15%/year (112,116,119,120).

HCV Infection in Correctional Settings

Juveniles. The prevalence of HCV antibody among detained or incarcerated juveniles is estimated to be 2%–3.5%. A history of injection-drug use is the predominant risk behavior, and regardless of reported risk behaviors, the prevalence is higher among females than among males (3%–7% versus 2%–3%) (A. Thomas, M.D., Oregon Health Division; and R. Bair, M.D., Bexar County Juvenile Detention Center, San Antonio, Texas, personal communications, 2001). The extent to which HCV infection is transmitted in juvenile correctional institutions is not known.

Adults. Among prison inmates, 16%–41% have serologic evidence of HCV infection; approximately 12%–35% have chronic HCV infection; and rates vary by geographic region (88,107,121,122) (L. Wang, Ph.D., New York State Department of Health; D. Lau, M.D., University of Texas Medical Branch—Galveston, personal communication, 2001). HCV infection is primarily associated with a history of injection-drug use. In a Wisconsin study of 1,148 inmates, among the 310 (27%) with a history of injection-drug use and serologic evidence of HBV infection or biochemical evidence of liver disease, 91% were found to be anti-HCV–positive (J. Pfister, M.S., Wisconsin State Laboratory of Hygiene, personal communication, 2001). Among HCV-positive entering jail inmates in Massachusetts, 85% reported needle-sharing, prior drug use, or a history of hepatitis (T. Lincoln, M.D., Hampden County Correctional Center, Ludlow, Massachusetts, personal communication, 2001).

The risk of HCV acquisition during incarceration is not well-established. The only published study to examine the incidence of HCV infection among prison inmates reported a rate of 1.1 infections/100 person-years of incarceration among males (121).

Correctional Staff. No published studies have reported the prevalence of HCV infection among correctional staff. In one unpublished study performed among correctional health-care workers, the prevalence of HCV infection was 2% (R. Gershon, Dr.P.H., Columbia University, New York, personal communication, 2002), which is not higher than that in the general population. This finding is similar to that of studies among other occupational groups, including hospital-based health-care workers, surgeons, and public safety workers (10,123).

Prevention and Control of Viral Hepatitis

Primary prevention of infection with hepatitis viruses can be achieved either through immunization (HAV, HBV) or through behavioral interventions to reduce the risk factors for infection (HCV). In addition, identification of persons with chronic HBV and HCV infection provides the opportunity to initiate activities (e.g., counseling, treatment, or vaccination) that can prevent further disease transmission and reduce the progression of chronic liver disease. This section summarizes
current information and practices to prevent infection with hepatitis viruses, including immunization, antiviral treatment, and risk-reduction counseling.

**Prevention of HAV Infection**

**Strategy to Prevent HAV Infection**

**Preexposure Immunization.** Vaccination is the most effective means to prevent HAV infection and reduce disease incidence. In the United States, preexposure vaccination is recommended for persons at highest risk for infection and persons in whom infection would result in adverse consequences (Box 1). In addition, routine vaccination is recommended for persons aged 2–19 years living in states and communities that have had the highest rates of disease (11) because the conditions that contribute to communitywide disease transmission continue to exist.

**Postexposure Prophylaxis.** Passive immunization with immune globulin (IG) is >85% effective in preventing hepatitis A after exposure of an unvaccinated person to an infected person if administered <2 weeks after exposure (11). Anti-HAV testing is not recommended because it would delay IG administration and is likely not to be cost-effective. Although limited data indicate hepatitis A vaccine might provide protection when administered soon after exposure, this has not been evaluated in controlled clinical trials, and use of hepatitis A vaccine alone is not recommended for postexposure prophylaxis. However, persons who receive IG postexposure prophylaxis, and for whom hepatitis A vaccine is also recommended, require vaccination (11).

**Detection and Management of Acute HAV Infection**

The diagnosis of hepatitis A is based on a positive serologic test for IgM anti-HAV in a person with clinical signs or symptoms of acute viral hepatitis. Serologic confirmation of HAV infection is required because hepatitis A cannot be distinguished from the other forms of viral hepatitis on the basis of clinical presentation alone (Box 2). Although management of clinical illness is supportive, progression to acute liver failure can occur (especially in persons with chronic liver disease), and 10%–15% of patients have relapsing illness.

**Contact Tracing.** Cases of acute hepatitis A are reported to the appropriate public health authorities, and a contact investigation is initiated by correctional officials to identify persons who would benefit from postexposure prophylaxis. Cellmates, sexual contacts, and persons having ongoing close personal contact with the index case are administered IG (Box 3) (11).

**Current Practices: Prevention of HAV in Correctional Settings**

Nationally, the extent to which juvenile correctional systems vaccinate against hepatitis A is unknown. A recent assessment found that in six of the 17 states where routine childhood vaccination is recommended, vaccination was also being conducted in juvenile detention facilities (CDC, unpublished data). A limited number of adult correctional systems routinely offer hepatitis A vaccination to all persons at risk for infection, whereas others offer vaccination only to inmates infected with HCV.

**Prevention of HBV Infection**

**Strategy to Prevent HBV Infection**

Prevention of acute and chronic HBV infection and elimination of HBV transmission in all age groups is most effectively achieved through hepatitis B vaccination (9). The national strategy to eliminate HBV transmission has four components: 1) prevention of perinatal HBV infection through maternal screening and postexposure prophylaxis of newborns of HBsAg-positive mothers; 2) hepatitis B vaccination of all infants to prevent infection in childhood and at later ages; 3) vaccination of all adolescents not previously vaccinated to

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**BOX 1. Groups for whom hepatitis A vaccination is recommended**

<table>
<thead>
<tr>
<th>Persons at increased risk for infection</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Travelers to countries with high endemicity for hepatitis A virus infection</td>
</tr>
<tr>
<td>• Men who have sex with men</td>
</tr>
<tr>
<td>• Users of injection and noninjection illegal drugs</td>
</tr>
<tr>
<td>• Persons who receive blood product replacement therapy for clotting factor disorders</td>
</tr>
<tr>
<td>• Children and adolescents living in states with historically elevated rates of hepatitis A*</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Persons at increased risk for adverse consequences of hepatitis A</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Persons with chronic liver disease of any etiology</td>
</tr>
</tbody>
</table>

* Routine vaccination recommended: Alaska, Arizona, California, Idaho, Nevada, New Mexico, Oklahoma, Oregon, South Dakota, Utah, Washington; routine vaccination should be considered: Arkansas, Colorado, Missouri, Montana, Texas, Wyoming.
### BOX 2. Diagnostic testing for infection with hepatitis viruses*

<table>
<thead>
<tr>
<th>Hepatitis Type</th>
<th>Test(s) Required</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute hepatitis A</td>
<td>- Immunoglobulin M antibody to hepatitis A virus (IgM anti-HAV) positive</td>
</tr>
<tr>
<td>Acute hepatitis B</td>
<td>- IgM antibody to hepatitis B core antigen (IgM anti-HBc) positive and - Hepatitis B surface antigen (HBsAg) positive</td>
</tr>
<tr>
<td>Acute hepatitis C</td>
<td>- Serum alanine aminotransferase (ALT) levels &gt;7 times the upper limit of normal and - Antibody to hepatitis C virus (anti-HCV) positive (repeat reactive) by screening immunoassay, and confirmed by a more specific assay (e.g. recombinant immunoblot assay [RIBA®] for anti-HCV or nucleic acid testing for HCV RNA) or - Anti-HCV–positive (repeat reactive) by screening immunoassay and a signal-to-cutoff ratio predictive of a true positive or determined for the particular assay (e.g., ≥3.8 for screening enzyme immunoassay [EIA])</td>
</tr>
</tbody>
</table>

* See Table 2 for interpretations of other markers of HBV infections.

### BOX 3. Contact investigation and postexposure prophylaxis after identification of a case of hepatitis A

- Contact investigation should be coordinated with local and state health departments. If the index patient is a food handler, public health officials should be directly involved in the investigation to evaluate the risk for transmission and the need for postexposure prophylaxis.
- The following persons, if not previously vaccinated, should be considered candidates for postexposure prophylaxis if exposed to an index patient with hepatitis A during the two weeks prior to the onset of symptoms. A single dose of immune globulin (IG) (0.02 mL/kg intramuscular) should be administered as soon as possible (but not >2 weeks after the last exposure) to — cellmates or dormitory mates — sex contacts — other close contacts based on epidemiologic investigation — other food handlers if the index patient was a food handler
- IG is not routinely indicated when an index case occurs in a school, work setting, or temporary housing unit.
- When a person with hepatitis A is admitted to a hospital, standard and contact precautions are indicated. Staff members are at low risk of infection and postexposure prophylaxis is not indicated.

* See Table 2 for interpretations of other markers of HBV infections.

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Prevent infection in this age group and at later ages; and 4) vaccination of adults and adolescents in groups at increased risk for infection (Box 4) (9,124).

Hepatitis B vaccination has been included in routine health-care visits for adolescents, however, this has not occurred for adults at risk of infection (9,125). Although the majority of persons aged <19 years not covered by private insurance are covered under the Vaccines for Children Program,* similar coverage does not exist for adults, and cost reimbursement is a substantial barrier to vaccination of adults (126).

Older juveniles and adults at risk for HBV infection are usually not vaccinated during health-care visits. Approximately 56% of persons with hepatitis B have either been treated for an STD (36%) or incarcerated (29%), factors for which routine hepatitis B vaccination is recommended (8,127).

* The Vaccines for Children (VFC) Program was established in 1994 for federal purchase of vaccine to be administered by a qualified health provider to juveniles aged <19 years who are American Indian or Alaska Native, uninsured or underinsured, or on Medicaid. VFC supports purchase of hepatitis A and hepatitis B vaccines, and HBIG. The VFC website is available at http://www.cdc.gov/nip/vfc/Default.htm.
Identification of persons with chronic HBV infection through diagnostic testing can reduce risks for chronic liver disease and further transmission of infection. Appropriate medical management and antiviral therapy of persons with chronic HBV infection can reduce the risk of cirrhosis and HCC. Additional morbidity from other hepatic insults can be reduced through hepatitis A vaccination, alcohol-reduction counseling, and risk-reduction education. The high rate of HBV infection in sex and close contacts, including cellmates, can be prevented through vaccination.

Prevention of Perinatal HBV Infection. Perinatal HBV infections can be prevented through routine testing to identify pregnant women who test positive for HBsAg and through timely postexposure immunization (prophylaxis) of their infants (78,128,129). Independent of maternal HBsAg status, hepatitis B vaccination is recommended for all infants beginning soon after birth and before their release from the hospital (130). Initiating hepatitis B vaccination soon after birth serves as a safety net to prevent HBV infection in infants whose mothers were not tested (131).

Adolescent Vaccination. Universal vaccination of infants against hepatitis B was first recommended in the United States in 1991 (9) and catch-up vaccination of all adolescents was recommended in 1995 to achieve elimination of HBV transmission in a more timely manner (132–135). Hepatitis B vaccination is now required by 33 states for entry to middle school or seventh grade. Three states have laws that require vaccination for college entry, and certain colleges require hepatitis B vaccination for matriculation (136) (S. Ainsworth, American College Health Association, personal communication, 2002).

Juvenile correctional vaccination programs have been established to prevent infections among detained persons at high risk for infection who might not be reached by school programs or entry requirements. Implementation of programs in juvenile correctional settings has been complicated by population mobility and need for parental consent in certain jurisdictions. However, recidivism can result in inmates being offered second and third vaccine doses (137) (G. Shostak, M.P.H., Massachusetts Department of Youth Services, personal communication, 2001).

Adult Vaccination. Routine vaccination of infants, young children and adolescents is expected to eventually eliminate transmission of HBV among adults in the United States. However, decades will pass before vaccinated children become protected adults, and vaccination of adults at increased risk for infection continues to be essential to reducing the high incidence of disease in this age group.

Vaccination coverage among adults at occupational risk for HBV infection has successfully reduced infection incidence by >90% (138). This was achieved by requiring employers to provide education and hepatitis B vaccination at no cost to employees (139). However, early efforts to vaccinate other adults (the majority of whom receive prevention services in the public sector) had limited success, primarily because of a lack of sustained programs and coverage for vaccine cost. More recently, demonstration projects and programs funded by state and local health departments to deliver hepatitis B vaccine in correctional facilities and STD and substance-abuse–treatment centers have demonstrated that immunization is feasible and high vaccination coverage can be achieved (140,141).

Previously, a major barrier to vaccination of adults at high risk was the practice of only offering vaccine to persons likely to complete the series. Although administration of the

Box 4. Groups recommended for preexposure hepatitis B vaccination

**Universal**
- All infants
- All children and adolescents not previously vaccinated

**Based on risk**
- Inmates of long-term correctional facilities
- Injection-drug users
- Sexually active men who have sex with men
- Men and women with >1 partner in the previous 6 months, a history of a sexually transmitted disease (STD), or treatment in an STD clinic
- Household contacts (including cellmates) and sex partners of persons with chronic HBV infection
- Persons in occupational groups with exposure to blood or body fluids
- Hemodialysis patients
- Recipients of clotting factor concentrates
- Long-term international travelers
- Clients and staff of institutions for the developmentally disabled

complete vaccine series should be the goal of any immunization program, high first-dose and modest second-dose vaccination coverage rates have been achieved when vaccine is offered to all persons in settings that serve populations at high risk (140). Protective levels of antibody develop after 1 dose of hepatitis B vaccine among 30%–50% and after 2 doses of vaccine among 75% of healthy young adults (142–144).

The transient nature of adult populations in correctional facilities often prevents completion of the full hepatitis B vaccine series. Ensuring follow-up with subsequent doses requires that an immunization record is included in the medical record of all inmates, is transferred among correctional facilities, and is provided to the inmate as part of release planning.

Testing for HBV Infection

Pregnant Women. HBsAg testing is recommended for all pregnant women early during each pregnancy or as soon as the pregnancy is recognized, irrespective of hepatitis B vaccination history or previous test results (9,145–147). In addition, women with risk factors for HBV infection during their pregnancy (e.g., intercurrent STDs, multiple sex partners, sex partners and household contacts of HBsAg positive persons, or clinically apparent hepatitis) need retesting for HBsAg late in pregnancy because of the high risk for HBV infection (9,147). Women determined to have chronic infection need evaluation for chronic liver disease, and close contacts (e.g., sex, household, prison cell, or dormitory) require vaccination because of their high risk for infection (9).

Prevaccination Testing. Proof of previous hepatitis B vaccination through use of an immunization registry, medical records, or vaccination card can be used to determine whether to exclude inmates from vaccination. When inmate vaccination status is unknown, testing for immunity to HBV infection can save vaccine costs among populations with high rates of infection or vaccination coverage (Box 5). However, vaccination of a person immune to HBV infection because of prior vaccination or infection does not increase the risk for adverse events. Testing is not indicated before vaccination of adolescents or younger children because of the low prevalence of HBV infection in these age groups (9,148).

As hepatitis B vaccination coverage increases among adolescents, a higher proportion of adults will be immune to HBV infection. Correctional systems should be aware of state hepatitis B vaccination requirements for middle school entry, which typically achieves high vaccination coverage rates. If adequate immunization records are not routinely available for incoming inmates, periodic serologic surveys are necessary to determine the prevalence of immunity to HBV infection among the incoming inmate population and guide policies for prevaccination testing.

BOX 5. Method to determine cost-effectiveness of prevaccination screening for hepatitis B vaccination*

<table>
<thead>
<tr>
<th>Equation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>( T = P_1 \times [P_2 + P_2(P_3)] \times v )</td>
<td>where</td>
</tr>
<tr>
<td>( T = ) cost of serologic test for anti-HBc or anti-HBs;</td>
<td></td>
</tr>
<tr>
<td>( P_1 = ) prevalence of past infection/immunization;</td>
<td></td>
</tr>
<tr>
<td>( P_2 = ) percentage of recipients of first dose who actually receive a second dose;</td>
<td></td>
</tr>
<tr>
<td>( P_3 = ) percentage of recipients of doses 1 and 2 who receive dose 3;</td>
<td></td>
</tr>
<tr>
<td>([P_2 + P_2(P_3)] = ) average number of doses for a person starting the series; and</td>
<td></td>
</tr>
<tr>
<td>( v = ) cost per dose of vaccine, including administrative costs.</td>
<td></td>
</tr>
</tbody>
</table>

* Using the formula for hepatitis A vaccination assumes no vaccination is done at the time of the blood draw. For hepatitis A vaccination, \( T = P_1 \times v \).

Among populations with a high prevalence of immunity as a result of vaccination, testing for chronic HBV infection is not warranted. However, among populations with a high prevalence of HBV infection, testing to identify inmates with chronic HBV infection is necessary to initiate appropriate medical follow-up and immunization of close contacts.

Postvaccination Testing. Testing to determine antibody response to vaccination is not necessary for healthy juveniles and adults (Appendix). For immunocompromised persons (e.g., hemodialysis patients, or HIV-infected) and persons with continued known exposure to HBV infection (e.g., infants born to HBsAg-positive mothers, sex partners of HBsAg-positive persons, health-care workers) testing is needed to verify response to vaccination and the need for possible revaccination, or to identify HBV infection (9,149,150)

Prevention of HBV Infection After Exposure

Immunization (active, passive, passive-active) within a relatively short period of time after exposure to HBV can effectively prevent acute and chronic infection. Initiation of the hepatitis B vaccine series within 12–24 hours of exposure has been demonstrated 70%–90% effective in preventing HBV infection (131,151). The combination of vaccine and HBIG achieves a similar level of efficacy (Box 6) (128,129). Among persons known to have been nonresponders to vaccination, 1 dose of HBIG is 70%–90% effective in preventing hepatitis B
TABLE 2. Interpretation of hepatitis B virus serologic testing

<table>
<thead>
<tr>
<th>Interpretation</th>
<th>HBsAg*</th>
<th>Total anti-HBc†</th>
<th>IgM§ anti-HBc</th>
<th>Anti-HBs¶</th>
</tr>
</thead>
<tbody>
<tr>
<td>Susceptible, never infected</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Acute infection, early incubation period**</td>
<td>+</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Acute infection</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>Acute resolving infection</td>
<td>–</td>
<td>+</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>Past infection, recovered and immune</td>
<td>–</td>
<td>+</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>Chronic infection</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>False positive (i.e., susceptible), past infection, or low-level chronic infection</td>
<td>–</td>
<td>+</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>Immune from vaccination if antibody concentration ≥10 milli international units per milliliter (mIU/mL)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>+</td>
</tr>
</tbody>
</table>

* Hepatitis B surface antigen.
† Antibody to hepatitis B core antigen.
§ Immunoglobulin M.
¶ Antibody to hepatitis B surface antigen.
** Transient HBsAg positivity (lasting ≤21 days) might be detected in some patients during vaccination.

BOX 6. Prophylaxis after exposure to hepatitis B virus (HBV)

**Type of exposure**
- Perinatal
- Sexual — acute case
- Sexual — chronic HBV infection
- Household (e.g., cell or dormitory) contact — to person with chronic HGV infection
- Household (e.g., cell or dormitory) contact — acute case

**Type of immunoprophylaxis**
- Vaccination* + HBIG†
- Vaccination* + HBIG
- Vaccination
- Vaccination. If not previously vaccinated, also administer HBIG if known exposure§
- Vaccination. If not previously vaccinated, also administer HBIG if known exposure§
- Vaccination +/- HBIG¶

* See Table 4.
† HBIG = hepatitis B immune globulin. Dosages: perinatal = 0.5 mL intramuscular; all other = 0.06 L/kg, intramuscular.
§ Identifiable blood exposure to infected contact (e.g., by sharing toothbrushes or razors).
¶ See Table 5.

Detection of HBV Infection

Acute HBV Infection. Acute HBV infection is asymptomatic among 60%–70% of patients, but can have symptoms and signs associated with acute viral hepatitis (e.g., loss of appetite, nausea, vomiting, fever, abdominal pain, or jaundice), and must be confirmed by serologic testing (Table 2, Box 2). Treatment for acute hepatitis B is supportive, consisting of rest, hydration, and symptomatic relief as needed. Identification of an inmate with acute HBV infection, especially one that has been incarcerated >6 months, requires an epidemiologic investigation by correctional officials, in collaboration with the appropriate health authorities, to identify the source of infection. Depending on the results, vaccination of sexual, prison cell, dormitory and household (e.g., conjugal and other family members) can be indicated.

Chronic HBV Infection. Chronic HBV infection can be distinguished from acute infection by serologic testing (Table 2). Inmates identified with chronic HBV infection require evaluation to determine the extent of liver disease, virus replication, indications for antiviral therapy (64), and need for vaccination of contacts to prevent HBV transmission.

Management of Chronic HBV Infection

Initial evaluation of patients with chronic HBV infection includes biochemical tests for liver disease (e.g., alanine transaminase, alkaline phosphatase, and γ-glutamyl transferase) to assess the extent of liver injury and the potential for disease progression. Antiviral therapy is indicated for patients with chronic hepatitis B who have evidence of active liver disease, as determined by liver biopsy or serologic markers of viral replication (64). The choice of antiviral therapy should be based on the results of the initial evaluation and the specific needs of the patient. Antiviral therapy typically involves the use of nucleoside or nucleotide analogs, which target the viral polymerase and inhibit viral replication. These medications are usually administered for a prolonged period of time, and require close monitoring of liver function and side effects. Inmates identified with chronic HBV infection require periodic monitoring of liver function and serologic markers of viral replication, as well as close collaboration with correctional and health authorities to ensure appropriate follow-up and management.
aminotransferase [ALT], aspartate aminotransferase [AST]), for the extent of liver disease (e.g., serum albumin, prothrombin time), and status of HBV replication (e.g., HBeAg, antibody to HBeAg [anti-HBe], and HBV DNA). Alpha interferon, lamivudine, or adefovir dipivoxil are approved by the Food and Drug Administration (FDA) for treatment of chronic hepatitis B (64,155). Therapy can be appropriate for patients who have abnormal levels of liver enzymes, active virus replication (HBeAg-positive or high levels of HBV DNA), and a liver biopsy indicating presence of moderate disease activity and fibrosis. (64)

Treatment with interferon, administered by injection 3 times/week, substantially decreases HBV DNA levels and clears HBeAg among >50% of patients with ALT levels >5 times the upper limit of normal and among 20%–35% of patients with ALT levels 2–5 times the upper limit of normal. Among patients with ALT levels <2 times the upper limit of normal, response is poor and therapy should be deferred. Long-term follow-up of treated patients suggests remission of chronic hepatitis induced by alpha interferon is of long duration (64). Patient characteristics associated with positive response to interferon therapy include low pretherapy HBV DNA levels, high pretherapy ALT levels, short duration of infection, acquisition of disease in adulthood, and histology indicative of active inflammation.

Lamivudine, administered orally daily, has been as effective as interferon at clearing HBeAg. Although a majority of patients taking lamivudine demonstrate improved liver histology, development of lamivudine-resistant HBV mutants is common, especially with prolonged use, and diminishes the effectiveness of treatment. Studies of lamivudine in combination with interferon have not been demonstrated to be superior to monotherapy (64).

The newest therapy to be approved is adefovir, which also is administered orally daily. Patients treated with adefovir exhibited substantial improvements in liver histology and decreased levels of HBV DNA; however, durability of the response has not been determined (156). Adefovir has been demonstrated to be effective in patients with chronic hepatitis B who have experienced resistance to lamivudine (156).

Treatment of persons coinfected with HIV and HBV requires additional monitoring. After initiation of highly active antiretroviral therapy (HAART) for treatment of HIV infection, reactivation of HBV replication with development of acute hepatitis has been observed among persons thought to have resolved HBV infection. While interferon treatment is not as effective for patients coinfected with HIV, HBV and HIV can be simultaneously treated (157).

Inmates identified with chronic HBV infection can benefit from counseling regarding ways to reduce the likelihood of transmitting HBV infection to others. In addition, vaccination of sexual and nonsexual contacts (e.g., cellmates) can prevent transmission (9).


Juveniles. Recommendations exist for universal vaccination of adolescents, and juveniles in the justice system have been found to have increased risk for HBV infection (125). In 2001, a national survey of state juvenile correctional systems reported that 36 (86%) of 42 responding systems had a hepatitis B prevention program in place; 78% used the VFC program to pay for vaccine; and 85% considered vaccinations to be a corrections responsibility while a juvenile is in custody. Written hepatitis B prevention policies were in place in 65% of states, and 27% used a vaccine tracking system or immunization registry (CDC, unpublished data).

In states with immunization registries and VFC participation, vaccination coverage among incarcerated juveniles has reached levels of over 90% (G. Treden, B.S.N., Wisconsin Department of Corrections, personal communication, 2002). However, when the correctional system does not have legal guardianship of the detained juvenile, the need for parental consent can pose a barrier to vaccination. In states with laws that enable minors to consent to their own STD-related treatment and prevention, hepatitis B vaccination has been considered part of such a consultation and has facilitated implementation of vaccination programs (M. Staples-Horne, M.D., Georgia Department of Juvenile Justice, personal communication, 2002).

Adults. Hepatitis B vaccination is recommended for adults in correctional settings because of their increased risk for infection, both inside and outside of prisons and jails (9,33,34,100). Vaccinating inmates in prisons has been demonstrated feasible and cost-saving from both a societal and prison perspective (158) (CDC, unpublished data). Approximately 25 state correctional systems and the Federal Bureau of Prisons have implemented hepatitis B immunization programs, which vary in scope and are often limited by funding or staffing resources. System policies include immunization of 1) all incoming inmates; 2) inmates of certain ages; 3) inmates with certain lengths of sentences; 4) inmates with HCV infection; or 5) inmates who request vaccination. In certain correctional systems, inmates must pay for vaccination (137,159). Among inmates in three systems (in Massachusetts, Michigan, and Texas) that offer hepatitis B vaccine, 60%–80% accept vaccination (T. Lincoln, M.D., Hampden County Correctional Center, Ludlow, Massachusetts; D. Thelen, Michigan State
Department of Health; and M. Kelley, M.D., Texas Department of Corrections, Huntsville, Texas; personal communications, 2001). Successful hepatitis B vaccination programs, like other successful adult vaccination programs (e.g., influenza) include establishment of policies for vaccination and a source of payment for vaccine (160–163). Among states, Hawaii, Michigan, Texas and Wisconsin have extensive experience in offering vaccine to inmates.

The Texas Department of Criminal Justice has 105 adult facilities with approximately 145,000 inmates. In 1999, funds were appropriated for hepatitis B vaccination of all offenders. A cost analysis indicated prevaccination testing would be cost-effective if prior HBV infection rates were >25%. However, a seroprevalence study identified an HBV prevalence of 17.8% and a history of vaccination among another 5.5%. Medical records are reviewed for a history of hepatitis B vaccination or evidence of HBV infection from prior clinical testing.

All inmates are offered vaccine, and the central pharmacy delivers second and third doses of vaccine to the appropriate housing units on a 0-, 2-, and 4-month vaccination schedule. Scheduled vaccine doses are listed in each inmate’s medical record to serve as an additional reminder to complete the vaccination series.

In the first 18 months of the program, 115,627 previously incarcerated inmates initiated the vaccine series, and since November 2001, the program has vaccinated all inmates at entry, an estimated 35,000/year. The estimated cost for vaccination of 121,000 inmates during the first 18 months of the program was $8 million, with an expected recurring annual cost of $2.6 million to vaccinate incoming inmates. (M. Kelley, M.D., Texas Department of Criminal Justice, personal communication, 2001).

Prevention of HCV Infection

Strategy to Prevent HCV Infection

CDC’s national strategy to prevent HCV infection includes 1) prevention of transmission during high-risk activities (e.g., injection-drug use and unprotected sex with multiple partners) through risk-reduction counseling, testing, and appropriate medical management of infected persons; 2) donor screening and product inactivation procedures to eliminate transmission from blood, blood products, donor organs, and tissue; and 3) improved infection control practices to further reduce the risk of transmission during medical procedures † (10). Primary prevention is directed at lowering the incidence of HCV infection. Of the estimated 25,000–40,000 persons newly infected with HCV annually during the past 5 years, approximately 60% acquired their infection through injection-drug use (45,111). Because no vaccine exists to prevent HCV infection, prevention must focus on risk reduction through counseling of persons who have admitted to or are at risk for illicit drug use or high-risk sexual practices. Counseling and testing to prevent HCV infection should be conducted in settings where persons at high risk are identified, including STD, HIV/AIDS, and substance-abuse treatment and prevention clinics, and correctional health programs (10) (Box 7).

The high prevalence of HCV infection and risk associated with HCV infection among inmates indicates the need to include HCV prevention activities in correctional settings. Inmates often have multiple substance abuse, as well as physical and mental health problems. To be effective, risk reduction among this population often requires a multidisciplinary approach to address drug use and other medical, psychological, social, vocational, and legal problems (164).

Identification of HCV-infected persons is required to initiate secondary and tertiary prevention activities, to reduce the risk of HCV transmission by the infected person and the risk for chronic liver disease (10). The substantial number of undiagnosed persons with chronic HCV infection makes identification of these persons a major focus of hepatitis C

BOX 7. Persons for whom routine hepatitis C virus (HCV) testing is recommended*

<table>
<thead>
<tr>
<th>Based on risk for infection, persons who</th>
</tr>
</thead>
<tbody>
<tr>
<td>• ever injected illegal drugs;</td>
</tr>
<tr>
<td>• received clotting factor concentrate produced before 1987;</td>
</tr>
<tr>
<td>• ever were on long-term hemodialysis;</td>
</tr>
<tr>
<td>• have evidence of chronic liver disease including persistently abnormal levels of alanine aminotransferase (ALT); or</td>
</tr>
<tr>
<td>• received a transfusion of blood or blood components or an organ transplant before July 1992.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Based on a recognized exposure,</th>
</tr>
</thead>
<tbody>
<tr>
<td>• health-care, emergency medical, public safety, and correctional workers after needle sticks, sharps, or mucosal exposure to HCV-positive blood; or</td>
</tr>
<tr>
<td>• children born to HCV-positive women.</td>
</tr>
</tbody>
</table>


* Screening testing for antibody to HCV (anti-HCV) followed by appropriate confirmatory testing for persons found to be screening test positive.

prevention programs. Comprehensive testing programs for persons at risk for HCV infection should include risk- and harm-reduction counseling, and substance-abuse treatment where appropriate, to reduce the risk of acquiring HCV infection. Anti-HCV–positive persons require further evaluation for chronic HCV infection and liver disease, and persons with chronic hepatitis C require evaluation for possible antiviral therapy and the need for further medical management. In addition, HCV-infected persons benefit from counseling to reduce their risk for transmitting HCV infection to others or developing further liver injury.

Persons with chronic hepatitis C are at risk for increased morbidity from additional hepatic insults. Fulminant hepatitis caused by hepatitis A can be prevented by vaccination (50). HCV-infected persons often have risk factors for HBV infection; therefore, hepatitis B vaccination is also recommended (10). Persons with hepatitis C should be counseled to not use alcohol, because its use (>10g/day for women and >20g/day for men) has been associated with more rapid progression of cirrhosis, which puts persons at higher risk for HCC (10,165,166).

Persons at risk for HCV infection or those chronically infected with HCV can benefit from health education to reduce or eliminate the risk for acquiring or transmitting HCV infection, including: 1) substance abuse treatment where appropriate, 2) clean needle and syringe use, 3) not sharing drug paraphernalia, and 4) condom use (10). Counseling and educational materials should include information concerning reducing further liver damage, as well as treatment options for those with chronic liver disease. Release planning should include substance-abuse–treatment referrals for IDUs and medical referrals to specialists for future medical management and treatment (see juvenile and adult sections on Health Education and Release Planning).

Testing for HCV Infection

Anti-HCV testing is recommended to identify infected persons and should include use of both an antibody screening assay and supplemental or confirmatory testing with an additional, more specific assay (e.g., recombinant immunoblot assay [RIBA,® Chiron Corporation, Emeryville, California] for anti-HCV or nucleic acid testing for HCV RNA) to prevent reporting of false positive results. However, substantial variation exists among laboratories regarding the extent to which more specific testing is performed. The level of the screening test signal-to-cut-off ratio has been shown to predict a true antibody-positive result. Use of the signal-to-cut-off ratio limits supplemental testing to those samples for which the ratio is low (167).

Detection of HCV Infection

Acute Hepatitis C. Acute HCV infection is usually asymptomatic (80%). However, acute hepatitis C should be included in the differential diagnosis of inmates who have signs and symptoms of acute hepatitis (Box 2). Confirmation of acute hepatitis C requires negative test results for IgM anti-HAV and IgM anti-HBc and a positive screening test result for anti-HCV, verified by supplemental testing or a high signal-to-cut-off ratio. Among a limited number of patients, onset of symptoms may precede anti-HCV seroconversion, and follow-up antibody testing might be necessary to make the diagnosis.

Identification of an inmate with acute hepatitis C, especially a person incarcerated for >6 months, requires initiation of an epidemiologic investigation by correctional officials, in collaboration with the appropriate health authorities, to identify the source of the infection. Depending on the results, testing of contacts might be indicated.

Chronic HCV Infection. Anti-HCV alone does not distinguish between acute, chronic, or resolved infection. In persons testing positive for anti-HCV, chronic HCV infection can be distinguished by persistence of HCV RNA for >6 months.

Management of HCV Infection

HCV-positive persons benefit from evaluation for the presence and severity of chronic liver disease. Antiviral therapy is recommended for persons with persistently elevated ALT levels, detectable HCV RNA, and a liver biopsy that indicates either portal or bridging fibrosis or moderate degrees of inflammation and necrosis. No clear consensus exists on whether to treat patients with persistently normal serum transaminases. Information is available on the National Institute of Health (NIH) website§ regarding regimens with proven efficacy approved by the FDA for the treatment of chronic hepatitis C (168). The FDA has approved three antiviral therapies for treatment of chronic hepatitis C in persons aged >18 years: alpha interferon, pegylated interferon, and alpha or pegylated interferon in combination with ribavirin. All can be administered for 52 weeks. Among persons with HCV genotype 1, the most common genotype in the United States, the response rate to either of the interferons given alone is <20%, but the response rate to the combination of alpha interferon and ribavirin is 30%–40%, and to pegylated interferon and ribavirin, 40%–50%. Both the alpha and pegylated interferons are administered by injection; ribavirin is taken orally. All of these drug regimens have certain side effects, some of which

can be serious. Successful treatment eliminates viremia and the potential for HCV transmission and further chronic liver disease (168,169).

In persons with both HCV and HIV infection, benefits of therapy for chronic hepatitis C have only recently been evaluated. The decision to treat persons coinfect ed with HIV must take into consideration the concurrent medications and medical conditions, including hyperthyroidism, renal transplant, and evidence of autoimmune disease. If CD4 counts are normal or minimally abnormal (>500/mL), treatment responses to interferon monotherapy are similar to non-HIV–infected persons (106,170,171). The efficacy of ribavirin/interferon combination therapy among HIV-infected persons has been tested in only a limited number of patients. Ribavirin can have substantial interactions with other antiretroviral drugs (168). Each patient should be evaluated by a physician familiar with the treatment of patients with HCV infection and HIV infections when appropriate, and indications for therapy should be reassessed at regular intervals.


Testing persons in populations with high proportions of IDUs (e.g., correctional institutions, HIV counseling and testing sites, and drug treatment programs) is an efficient strategy for identifying HCV-positive persons (10). However, in the correctional setting, a limited number of studies have examined concerns surrounding willingness to be tested, treatment options, compliance, and outcomes among those offered therapy (122,172). In assessments of other screening programs in prisons (e.g., those for HIV and STDs), a relatively high rate (approximately 50%) of refusal has been reported (173–175).

Limited data from studies in Rhode Island and Pennsylvania indicate approximately 7%–27% of all inmates identified with HCV infection ultimately begin treatment (122,172) (F. Maue, M.D., Pennsylvania State Department of Corrections, personal communication, 2001). The majority of inmates were excluded from treatment because of clinical contraindications, short lengths of prison stay, and drug and alcohol use (122,172) (F. Maue, M.D., Pennsylvania State Department of Corrections, personal communication, 2001). Less-restrictive criteria might increase numbers of inmates eligible for treatment (168). However, factors that contribute to acceptance and completion of treatment regimens among this population need to be identified to improve outcomes.

Health Education

Health education directed toward prevention of viral hepatitis includes information related to the disease, routes of transmission, risk factors for infection, methods of prevention, disease outcomes and treatment options. During incarceration, numerous opportunities exist for educational interventions, including at facility entry, HIV-education classes, school curricula, and peer education courses. Education can take different forms, including videos, brochures, and formal classroom presentations. However, repeated face-to-face sessions have been determined to be the most effective means to transmit information to be retained by students (Box 8) (176–178). Model programs use peer health educators in workshops for incoming inmates and community educators to discuss risk assessment, risk reduction, and referrals for soon-to-be released inmates.¶

Health education programs aimed at reducing the risk of infection with hepatitis viruses include a discussion of hepatitis A prevention, including risk factors for prevention, hygiene practices, and the significance of vaccination for persons at risk for infection. Curricula addressing prevention of HBV and HCV infection include information concerning the similar modes of transmission and means for prevention, and information about hepatitis B vaccination and risk reduction. This information can be incorporated into health-education programs for the prevention of HIV/AIDS.

Release Planning

Release planning is a relatively new component of health-care management for incarcerated persons. The majority of medical release and discharge planning programs in prison

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**Box 8. Elements of viral hepatitis health education**

<table>
<thead>
<tr>
<th>Health education programs and curricula should include</th>
</tr>
</thead>
<tbody>
<tr>
<td>• routes of transmission;</td>
</tr>
<tr>
<td>• risk factors for infection;</td>
</tr>
<tr>
<td>• disease outcomes, the need for medical management</td>
</tr>
<tr>
<td>and treatment options;</td>
</tr>
<tr>
<td>• methods to prevent infection, including immunization</td>
</tr>
<tr>
<td>and harm and risk reduction;</td>
</tr>
<tr>
<td>• the importance of substance abuse treatment, when</td>
</tr>
<tr>
<td>appropriate;</td>
</tr>
<tr>
<td>• sexual precautions including abstinence counseling</td>
</tr>
<tr>
<td>and condom use;</td>
</tr>
<tr>
<td>• risk-reduction counseling, including not sharing</td>
</tr>
<tr>
<td>drug paraphernalia; and</td>
</tr>
<tr>
<td>• resources in the community available to support</td>
</tr>
<tr>
<td>and sustain a reduction in risk behaviors.</td>
</tr>
</tbody>
</table>

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¶ Available at http://www.hepprograms.org.
facilities have focused on HIV aftercare (179,180), but management of other chronic infections can result in the same beneficial outcomes for HIV-infected inmates.

Comprehensive release planning includes transitional housing, continued access to discharge medications and immunizations, and coordination and case-management for long-term specialized care for persons with chronic conditions. Persons diagnosed with chronic HBV infection can benefit from counseling about preventing transmission to household, sexual and drug-use contacts. Susceptible contacts of persons diagnosed with chronic HBV infection benefit from hepatitis B vaccination. Persons with chronic hepatitis B or chronic hepatitis C can benefit from 1) counseling regarding ways to reduce further liver damage, 2) referrals to substance-abuse–treatment and other programs for IDUs if indicated, and 3) medical referrals to specialists for future treatment.

**Rationale for Prevention and Control of Viral Hepatitis in Correctional Settings**

The high prevalence of chronic HBV and HCV infection and risk factors for transmission of these infections in the correctional system make their prevention and control high priorities for correctional health programs. In addition, because a substantial proportion of incarcerated persons return to the community and continue to transmit or acquire these infections at a high rate, prevention and control of viral hepatitis in the correctional setting should become part of community prevention and control efforts.

Highly effective and safe vaccines are available to prevent HAV and HBV infections. Identification of risk factors and infection status, combined with harm- and risk-reduction counseling, and substance-abuse treatment, have the potential to prevent HCV infections in the same manner they have reduced the risk of HIV/AIDS. In addition, identification of persons with chronic HBV and HCV infection provides the opportunity for medical evaluation and treatment of chronic liver disease, and to initiate measures to prevent further transmission.

The feasibility of including viral hepatitis prevention activities in existing prevention programs has been demonstrated. However, the challenges to integration of a comprehensive viral hepatitis prevention and control program in correctional health settings are substantial. They include budgetary and staffing constraints, priorities that compete with preventive health care, and lack of communication among the correctional health, public health, and private health-care systems. Nonetheless, the correctional health setting has demonstrated that it provides a vehicle to deliver viral hepatitis prevention and control activities.

The recommendations for prevention and control of viral hepatitis that follow are specifically adapted to the correctional setting. The objective of these recommendations is to reduce transmission of hepatitis virus infections both during and after incarceration. Implementation of these recommendations can 1) reduce transmission of HAV infection in the community by immunizing incarcerated persons at highest risk for infection; 2) eliminate transmission of HBV infection among the inmate population through immunization; 3) reduce the number of new HCV infections by testing, harm- and risk-reduction counseling, and substance-abuse treatment and prevention; 4) reduce the burden of viral hepatitis-related chronic liver disease through appropriate medical management; and 5) prevent HBV and HCV infections among correctional employees.

**Rating the Recommendations**

The following recommendations are rated, where applicable, on the basis of the strength of evidence indicating changes in outcomes attributable to the interventions. Where formal recommendations previously have been published, they are cited as supporting evidence and can be referred to for the original studies. Ratings have been assigned using a modification of criteria published by the Guide to Community Preventive Services (181). No rating was assigned to a recommendation considered standard practice.

- **Strongly recommended** (on the basis of >2 consistent, well-conceived, well-executed studies with control groups or longitudinal measurements).
- **Recommended** (on the basis of >1 well-conceived, well-executed, controlled or time-series study; or >3 studies with more limited execution).
- **Indicated** (on the basis of previous scientific observation and theoretic rationale, but case-controlled or prospective studies do not exist).
- **Not recommended** (on the basis of published literature recommending against a practice).

**Recommendations for Juvenile Correctional Facilities — Hepatitis A Virus Infection**

- Hepatitis A vaccination is recommended for all juveniles in those states where routine vaccination is recommended (Box 1) (11). Strongly recommended.
- In all other states, hepatitis A vaccination of all juveniles should be considered because of the high prevalence of risk factors for HAV infection among this population. However, if routine vaccination is not implemented, vaccination is recommended for juveniles with risk factors for HAV infection (Box 1) or for those at risk for severe adverse outcomes of infection (e.g., persons with chronic liver disease) (11,57). Strongly recommended.

- Vaccination should be initiated as soon as possible after entry into incarceration or detention using the appropriate dosage and schedule (Standard practice) (Table 3).

- Tracking systems to ensure completion of vaccine series within the correctional system should be established, and systems should be established to facilitate completion of the vaccine series in the community. (Standard practice).

- Vaccination information, including date of administration, dose, and manufacturer should be included in the medical record, and an immunization record should be given to juvenile, or their parents or guardians, upon release. (Standard practice).

- Routine screening or prevaccination testing of juveniles for markers of HAV infection is not recommended. Prevaccination testing should be considered for older adolescents (e.g., >15 years) in certain population groups (i.e., American Indians, Alaska Natives, and Hispanics) because of higher prevalence of infection or previous infection (11). Indicated.

- Juveniles with signs or symptoms suggestive of acute hepatitis should have appropriate diagnostic testing to differentiate acute hepatitis A, hepatitis B, or hepatitis C and to determine if the patients have chronic HBV or HCV infection (Box 2). (Standard practice).

- Cases of acute hepatitis A should be reported to the appropriate public health jurisdiction (e.g., county or state health department). (Standard practice).

- Identification of a case of hepatitis A in a correctional facility should prompt an epidemiologic investigation by correctional officials, in collaboration with the appropriate health authorities, to identify the source of infection and contacts who might have been exposed. (Standard practice).

- Unvaccinated close contacts of a confirmed case of hepatitis A should be administered postexposure prophylaxis with 1 dose of IG (0.02 mL/kg body weight, intramuscular) as soon as possible, but not >2 weeks after the last exposure. If the contact has indications for hepatitis A vaccination, vaccine should be administered either at the same or a later time (Box 3) (11). Strongly recommended.

### Recommendations for Juvenile Correctional Facilities — Hepatitis B Virus Infection

#### Preventing Perinatal HBV Infection

- All pregnant incarcerated juveniles should be tested for HBsAg after their pregnancy is recognized, even if previously vaccinated or tested. Because of the high risk of HBV infection among this population, testing should be performed even if the female was tested before incarceration. The HBsAg status of incarcerated pregnant juveniles should be reported to the hospital where the juvenile will deliver her infant, along with other prenatal medical information. HBsAg-positive females should also be reported to the appropriate public health authority (9). Strongly recommended.

- Infants born to HBsAg-positive mothers should receive HBIG (0.5 mL) and the first dose of hepatitis B vaccine <12 hours of birth (Table 4) (9). Strongly recommended.

- Females admitted for delivery without HBsAg test results should have blood drawn for testing. While test results are pending, the infant should receive hepatitis B vaccine without HBIG within 12 hours of birth (Table 4). (Standard practice).

- If the mother is later found to be HBsAg positive, her infant should receive HBIG as soon as possible, but ≤7 days after birth. If the infant does not receive HBIG,

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### TABLE 3. Hepatitis A vaccination dosages and schedule

<table>
<thead>
<tr>
<th>Vaccine and recipient ages (yrs)</th>
<th>Dose</th>
<th>Volume (mL)</th>
<th>No. of doses</th>
<th>Schedule (mos)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Havrix™</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2–18</td>
<td>720 EL.U.†</td>
<td>0.5</td>
<td>2</td>
<td>0 and 6–12</td>
</tr>
<tr>
<td>&gt;18</td>
<td>1,440 EL.U.†</td>
<td>1.0</td>
<td>2</td>
<td>0 and 6–12</td>
</tr>
<tr>
<td>VAQTA®§</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2–18</td>
<td>25 U</td>
<td>0.5</td>
<td>2</td>
<td>0 and 6–18</td>
</tr>
<tr>
<td>&gt;18</td>
<td>50 U</td>
<td>1.0</td>
<td>2</td>
<td>0 and 6</td>
</tr>
<tr>
<td>Twinrix®¶</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;17</td>
<td>720 EL.U.†</td>
<td>1.0</td>
<td>3</td>
<td>0, 1, and 6</td>
</tr>
</tbody>
</table>


* Manufactured by GlaxoSmithKline Biologicals, Rixensart, Belgium.
† Enzyme-linked immunosorbent assay (ELISA) units.
§ Manufactured by Merck & Co., Inc., Whitehouse Station, New Jersey.
¶ Manufactured by GlaxoSmithKline Biologicals, Rixensart, Belgium.
** Twinrix also contains hepatitis B vaccine antigen.
TABLE 4. Recommended dosages of licensed hepatitis B vaccines

<table>
<thead>
<tr>
<th>Age group (yrs)</th>
<th>Recombivax HB®†</th>
<th>Engerix-B®§</th>
<th>Twinrix®¶</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>µg/mL</td>
<td>µg/mL</td>
<td>µg/mL</td>
</tr>
<tr>
<td>Persons &lt;19 (including infants born to HBsAg mothers)</td>
<td>5  0.5</td>
<td>10  0.5</td>
<td>—</td>
</tr>
<tr>
<td>Persons 11–15</td>
<td>10  1.0**</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Persons ≥20</td>
<td>10  1.0</td>
<td>20  1.0</td>
<td>20††  1.0</td>
</tr>
<tr>
<td>Dialysis patients and other immunocompromised persons</td>
<td>40  1.0§§</td>
<td>40  2.0¶¶</td>
<td>—</td>
</tr>
</tbody>
</table>

* Both vaccines are routinely administered in a 3-dose series, which includes schedules of 0, 1, and 6 months; 0, 2, and 4 months; 0, 2, and 6 months; and, for adolescents, 0, 12, and 24 months.
† Manufactured by Merck & Co. Inc., Whitehouse Station, New Jersey.
§ Manufactured by GlaxoSmithKline Biologicals, Rixensart, Belgium.
¶ Manufactured by GlaxoSmithKline Biologicals, Rixensart, Belgium.
** Administered in a 2-dose schedule at 0 and 4–6 months.
†† Twinrix is only licensed for persons aged >17 years, and contains both hepatitis A and hepatitis B vaccine antigens administered as a 3-dose schedule.
§§ Special formulation.
¶¶ Two 1.0-mL doses administered at one site, in a 4-dose schedule at 0, 1, 2, and 6 months.

it is important that the second dose of vaccine be administered at 1 month of age. The final dose should be given at age 6 months (Table 4). Strongly recommended.

— If the mother is found to be HBsAg negative, her infant should continue to receive hepatitis B vaccine as part of the routine vaccination schedule (Table 4). Strongly recommended.

— If the mother is never tested to determine her HBsAg status, the infant should continue to receive hepatitis B vaccine as part of the routine vaccination schedule (Table 4). Strongly recommended.

• Case management should be established to ensure appropriate postexposure prophylaxis and follow-up for children born to incarcerated or recently released HBsAg-positive mothers, including completion of the vaccine series at age 6 months and postvaccination testing during ages 9–15 months (9,182). Recommended.

• Infants born to HBsAg-negative mothers should receive the first dose of hepatitis B vaccine before release from the hospital (9,142,143). Strongly recommended.

• Previously unvaccinated HBsAg-negative pregnant juveniles should be vaccinated; pregnancy is not a contraindication to vaccination (9,183–185). Strongly recommended.

• Discharge planning for pregnant HBsAg-positive juveniles should include transfer of appropriate medical records to the hospital where the juvenile plans to deliver her infant, along with other prenatal medical information. Test results should also be provided to the patient and her parent or guardian. (Standard practice).

Hepatitis B Vaccination

Preexposure

• All juveniles who receive a medical evaluation in a correctional facility should be administered hepatitis B vaccine, unless they have proof of completion of the vaccine series or serologic evidence of immunity to infection. The vaccine series should be started for those juveniles who have never been vaccinated, irrespective of their length of stay, and the series should be completed for those incompletely immunized (9,142,186,187). Strongly recommended.

— For juveniles who do not receive medical evaluation upon entry into correctional custody, vaccination should be considered for those who lack proof of previous vaccination (125). (Standard practice).

— Catch-up vaccination of previously unvaccinated, already incarcerated juveniles should be considered in facilities in which routine hepatitis B vaccination of entering inmates is established (33). (Standard practice).

• An appropriate vaccination dose and schedule should be selected to facilitate completion of the vaccine series while the juvenile is in custody. For previously unvaccinated juveniles held in a correctional facility for <6 months, the vaccine series should be initiated and completed by using a 4-month schedule (0, 1–2, and 4 months) (Table 4) (186–189). Recommended.

• Vaccination information, including date of administration, dose, and manufacturer should be included in the medical record, and an immunization record should be given to juveniles, or to their parents or guardians, upon release. (Standard practice).

• Discharge planning should include transfer of immunization records to the person’s medical home to ensure completion of the vaccine series for those juveniles not
fully vaccinated while in the correctional facility, and for all fully vaccinated persons as well. (Standard practice).

**Prevaccination and Postvaccination Testing**

- Prevaccination serologic testing is not indicated (148). Indicated.
  - As hepatitis B vaccination coverage among adolescents increases, validation of immunization records or serologic testing might become a cost-effective means to minimize overvaccination. Indicated.
  - Knowledge of state middle school hepatitis B vaccination requirements and performance of periodic vaccine coverage serologic surveys to determine the proportion of vaccinated or immune adolescents entering juvenile facilities should be used to define prevaccination screening policies (e.g., history or serologic testing) and the need for hepatitis B immunization among specific age groups. (Standard practice).
- Postvaccination testing is not indicated for healthy juveniles (9,142,143,190). However, for juveniles with special conditions (e.g., immunocompromised or HIV-infected), postvaccination testing for anti-HBs is recommended 1–2 months after completion of the vaccine series. Nonresponders in this category should be revaccinated (149,150). Strongly recommended.

<table>
<thead>
<tr>
<th>TABLE 5. Postexposure prophylaxis for exposure to hepatitis B virus</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Vaccination and antibody response status of exposed person</strong></td>
</tr>
<tr>
<td><strong>Positive</strong></td>
</tr>
<tr>
<td>Unvaccinated</td>
</tr>
<tr>
<td>Previously vaccinated</td>
</tr>
<tr>
<td>Known responder¶</td>
</tr>
<tr>
<td>Known nonresponder**</td>
</tr>
<tr>
<td>Antibody response unknown</td>
</tr>
<tr>
<td>1. If adequate, no treatment is necessary.¶¶</td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>

**Source:** CDC. Updated U.S. Public Health Service guidelines for the management of occupational exposures to HBV, HCV, and HIV and recommendations for postexposure prophylaxis. MMWR 2001;50(No. RR-11):1-52.

* Persons who have previously been infected with HBV are immune to reinfection and do not require postexposure prophylaxis.
† Hepatitis B immune globulin; dose is 0.06 mL/kg body weight intramuscularly.
§ Hepatitis B vaccine.
¶ A responder is a person with adequate levels of serum antibody to HBsAg (i.e., anti-HBs >10 mIU/mL).
** A nonresponder is a person with inadequate response to vaccination (i.e., anti-HBs <10 mIU/mL).
†† The option of administering one dose of HBIG and reintitling the vaccine series is preferred for nonresponders who have not completed a second 3-dose vaccine series. For persons who previously completed a second vaccine series but failed to respond, 2 doses of HBIG are preferred.
§§ Hepatitis B surface antigen.
¶¶ Antibody to HBsAg.

**Postexposure Prophylaxis**

- After any percutaneous (e.g., sharing injection-drug equipment, human bite) or mucosal (e.g., sexual) exposure to blood, unvaccinated juveniles should begin the vaccine series, and the exposure incident should be evaluated to determine if additional postexposure prophylaxis (i.e., HBIG) is required (Table 5) (9,47). Strongly recommended.
  - The first dose of hepatitis B vaccine should be administered immediately, and the remaining doses, 1 and 6 months later. (Standard practice).
  - For an exposed juvenile who has begun but not completed the vaccine series, subsequent vaccine doses should be administered as scheduled. (Standard practice).
  - The person who was the source of the exposure should be tested for HBsAg, even if this person was previously vaccinated. If the source person is HBsAg-positive, HBIG should be administered as soon as possible and ≤7 days after the exposure. (Standard practice).
  - Postexposure prophylaxis is not necessary for a fully vaccinated juvenile after exposure to HBV (9,47). Not recommended.
Serologic Testing for Hepatitis B Virus Infection

- Routine testing of juveniles for markers of HBV infection (e.g., HBsAg, anti-HBs, anti-HBc) is not recommended (5,76,81,148,191). Not recommended.
- Juveniles with signs or symptoms suggestive of viral hepatitis should have appropriate diagnostic testing to differentiate acute hepatitis A, hepatitis B, or hepatitis C and to determine if the patient has chronic HBV or HCV infection (Box 2). (Standard practice).
  — Cases of acute hepatitis B should be reported to the appropriate public health authority. (Standard practice).
  — Cases of chronic HBV infection should be reported in those states that require reporting. (Standard practice).
- Identification of a case of acute hepatitis B should prompt an epidemiologic investigation by correctional officials, in collaboration with the appropriate health authorities, to identify the source of infection and provide appropriate postexposure prophylaxis (Table 5) (Box 6) to nonimmunized contacts at risk for infection. (Standard practice).

Chronic Hepatitis B Treatment

- Juveniles identified as having, or who are known to have chronic HBV infection during routine medical screening should be evaluated to determine the presence and extent of chronic liver disease and candidacy for antiviral therapy. Recommended.
  — Lamivudine can be used to treat patients aged >2 years. Recommended.
  — The safety and efficacy of interferon and adefovir in pediatric patients has not been established. Recommended.
  — Treatment of patients with chronic hepatitis B should be conducted in consultation with a pediatric specialist experienced with these treatment regimens (64,192,193). Recommended. Recommended.
- All long-term correctional facilities should establish criteria for identification of inmates who might benefit from treatment, based on the latest treatment guidelines. (Standard practice).
- Discharge planning for persons with chronic HBV infection should include referral to medical care, risk-reduction programs, and social services necessary to maintain behavior changes; vaccination of contacts should also be arranged before patient discharge. (Standard practice).

Recommendations for Juvenile Correctional Facilities — Hepatitis C Virus Infection

Testing for Hepatitis C Virus Infection

- A history of risk factors for HCV infection should be obtained from juveniles undergoing medical evaluations, and those with risk factors should be tested for anti-HCV (Box 7). Routine testing of all juveniles for anti-HCV is not recommended (10). Recommended.
- Juveniles with signs or symptoms suggestive of viral hepatitis should have appropriate diagnostic testing to differentiate acute hepatitis A, hepatitis B, or hepatitis C and to determine if the patient has chronic HBV or HCV infection (Box 2). (Standard practice).
- Cases of acute hepatitis C should be reported to the appropriate public health authority. (Standard practice).
- Anti-HCV–positive persons should be reported if required by state regulations. (Standard practice).
- Identification of juveniles with acute hepatitis C, including ones who have been incarcerated for >6 months, should prompt an epidemiologic investigation by correctional officials, in collaboration with the appropriate health authorities, to identify the source of the infection. Depending on the results of the investigation, testing of contacts might be indicated (Box 7). (Standard practice).
- Juveniles who are anti-HCV–positive should receive further medical evaluation to determine if chronically infected (Box 2). (Standard practice).

Postexposure Management for HCV

- After a percutaneous or percutaneous exposure to blood, the source person should be tested for anti-HCV. If the source person is anti-HCV–positive, the exposed person should be tested for anti-HCV and ALT activity at baseline and 4–6 months later. For earlier diagnosis, testing for HCV RNA can be performed in 4–6 weeks (10). Recommended.
- Immune globulin and antiviral agents are not recommended for postexposure prophylaxis of hepatitis C (10). Not recommended.

Chronic Hepatitis C Treatment

- Juveniles identified as having chronic HCV infection should be evaluated to determine the presence and extent of chronic liver disease. FDA-approved antiviral agents for treatment of hepatitis C are not indicated for persons aged <18 years, although participation in clinical trials
might be possible. Although HCV infection in juveniles can result in less severe disease, infected juveniles should be monitored by a specialist familiar with this disease. Discharge planning should also include drug and alcohol abuse treatment, risk-reduction programs, and social services necessary to maintain behavior changes. (Standard practice).

Juvenile Health Education and Release Planning (see Box 8)

- Prevention of HAV, HBV, and HCV infections should be incorporated into health education programs (e.g., programs for preventing HIV/AIDS) and include information concerning modes of disease transmission and means for prevention, including risk-reduction and immunization. (Standard practice).
- An integrated health education and risk reduction program should be established in each facility and include a written plan of health instruction completed by each inmate. (Standard practice).
- Such instruction should address a range of issues relevant to the diverse developmental and cultural composition of correctional populations, and should include basic skill development, literacy, and home economics, as well as tools needed to avoid behaviors that result in acquisition of HIV, hepatitis, and other bloodborne and sexually transmitted infections. (Standard practice).
- Teachers should be trained professionals or inmate peers with specific training to teach comprehensive life-skills programs, including health education. (Standard practice).
- A system for periodic evaluation, updating and improvement should exist. (Standard practice).
- Documentation of hepatitis A or hepatitis B vaccination should be included in the medical record retained within the correctional system, as well as in any medical record provided to other health-care providers. In addition, vaccinated persons or their parents or guardians should be provided a personal immunization record. (Standard practice).
- Juvenile correctional health facilities should establish links with community and public health facilities, and where available, with immunization registries, to ensure tracking and completion of hepatitis A and hepatitis B vaccine series. (Standard practice).
- Juveniles with chronic HBV or HCV infection should be — counseled regarding ways to reduce further liver damage, including limiting alcohol and drug use, and afforded substance-abuse treatment when appropriate; and — provided aftercare that includes medical follow-up. (Standard practices).

Recommendations for Adult Correctional Facilities — Hepatitis A Virus Infection

- Hepatitis A vaccination is recommended for adults in groups at risk for HAV infection (e.g., men who have sex with men or drug users) or who are likely to experience severe adverse outcomes of infection (e.g., persons with chronic liver disease) (Box 1). Recommended.
- The vaccination series should be initiated as soon as possible after incarceration using the appropriate dosage and schedule (Table 3). Tracking systems to ensure completion of the vaccine series within the correctional system should be established, and systems should be developed to facilitate completion of the second vaccine dose for those inmates who return to the community (11). Strongly recommended.
- Prevaccination serologic testing to identify susceptible persons should be considered if determined to be cost-effective (Box 5), and it does not compromise initiation of vaccination. Inmates aged >40 years and those from regions of high endemicity (see Appendix) should be considered for prevaccination testing because of the high prevalence of past HAV infection among these groups (11). Indicated.
- Routine screening of adults for anti-HAV is not recommended, except when used to identify susceptible persons for vaccination (11). Indicated.
- Vaccination information, including date of administration, dose, and manufacturer should be included in the medical record, and an immunization record should be given to the inmate upon release. (Standard practice).
- Adults with signs or symptoms suggestive of acute hepatitis should have appropriate diagnostic testing to differentiate acute hepatitis A, hepatitis B, or hepatitis C, and to determine if the patient has chronic HBV or HCV infection (Box 2). (Standard practice).
- Cases of hepatitis A should be reported to the appropriate public health authority. (Standard practice).
- Identification of a case of hepatitis A in a correctional facility should prompt an epidemiologic investigation by correctional officials, in collaboration with the appropriate health authorities, to identify the source
of infection and contacts that might have been exposed. (Standard practice).

— Unvaccinated or known susceptible close contacts of a confirmed case of hepatitis A should be administered postexposure prophylaxis with a single dose of IG (0.02 mL/kg body weight, intramuscular) as soon as possible, but not >2 weeks after the last exposure (Box 3) (11). Strongly recommended.

Recommendations for Adult Correctional Facilities — Hepatitis B Virus Infection

Preventing Perinatal HBV Infection

- All pregnant women should be tested for HBsAg after their pregnancy is recognized, even if previously vaccinated or tested. Because of the high risk for HBV infection among this incarcerated population, testing should be performed even if the woman was tested before incarceration. The HBsAg status of a pregnant woman should be reported to the hospital where she will deliver her infant, along with other prenatal medical information. HBsAg-positive women should also be reported to the appropriate public health authority (9). Strongly recommended.
- Infants born to HBsAg-positive mothers should receive HBIG (0.5 mL) and the first dose of hepatitis B vaccine <12 hours after birth (Table 4) (9). Strongly recommended.
- Females admitted for delivery without HBsAg test results should have blood drawn for testing. While test results are pending, the infant should receive hepatitis B vaccine without HBIG within 12 hours of birth (Table 4) (Standard practice).
  — If the mother is later found to be HBsAG positive, her infant should receive HBIG as soon as possible, but ≤7 days after birth. If the infant does not receive HBIG, it is important that the second dose of vaccine be administered at 1 month of age. The final dose should be given at age 6 months (Table 4). Strongly recommended.
  — If the mother is found to be HBsAg negative, her infant should continue to receive hepatitis B vaccine as part of the routine vaccination schedule (Table 4). Strongly recommended.
  — If the mother is never tested to determine her HBsAg status, the infant should continue to receive hepatitis B vaccine as part of the routine vaccination schedule (Table 4). Strongly recommended.

- Case management should be established to ensure appropriate postexposure prophylaxis and follow-up for children born to incarcerated or recently released HBsAg-positive mothers, including completion of the vaccine series at age 6 months and postvaccination testing during ages 9–15 months (9,182). Recommended.
- Infants born to HBsAg-negative mothers should receive the first dose of hepatitis B vaccine before release from the hospital (9,130). Strongly recommended.
- Previously unvaccinated HBsAg-negative pregnant women should be vaccinated; pregnancy is not a contraindication to vaccination (9,183–185). Strongly recommended.
- Discharge planning for pregnant HBsAg-positive women should include transfer of appropriate medical records to the hospital where the woman plans to deliver her infant, along with other prenatal medical information. Test results should also be provided to the patient. (Standard practice).

Hepatitis B Vaccination

Preexposure

- All persons who receive a medical evaluation in a correctional facility should be administered hepatitis B vaccine, unless they have proof of completion of the vaccine series or serologic evidence of immunity to infection. The vaccine series should be started for those who have never been vaccinated, irrespective of the length of their stay, and the series should be completed for those incompletely immunized (9,142,143,190). Strongly recommended.
  — For persons who did not receive medical evaluation upon entry into correctional custody, vaccination should be considered for those who lack proof of previous vaccination or immunity (9,142,143). Recommended.
  — Catch-up vaccination of already incarcerated, previously unvaccinated persons, or persons known to be susceptible to infection, should be considered in facilities in which routine hepatitis B vaccination of entering inmates is being established. Priority should be given to vaccination of contacts of known HBsAg-positive persons (e.g., cellmates or persons living in the same cell block or dormitory) (33,101,102). Recommended.
  — An appropriate vaccination dose and schedule should be selected to facilitate completion of the vaccine series while the person is in custody. For previously unvaccinated persons held in a correctional facility for <6 months, the vaccine series should be initiated and completed by using
a 4-month schedule (0, 1–2, and 4 months) (Table 4) (186–189). Recommended.
• Vaccination information, including date of administration, dose, and manufacturer should be included in the medical record, and an immunization record should be given to the incarcerated person upon release. (Standard practice).
• Discharge planning should include transfer of immunization records to the person’s medical home to ensure completion of the vaccine series for persons not fully vaccinated while in the correctional facility, and for all fully vaccinated persons as well. (Standard practice).

**Prevaccination and Postvaccination Testing**

• Prevaccination serologic testing should be considered for adult incarcerated populations and is likely to be cost-effective when the prevalence of immunity from prior infection and vaccination exceeds 25%–30% (Box 5) (148). Indicated.
  — To assist correctional facilities in determining whether to conduct prevaccination testing, periodic serologic surveys of incoming inmates can be used to determine the prevalence of markers of immunity to HBV infection. (Standard practice).
  — Testing for anti-HBs provides the best measure of immunity to HBV infection, because it detects infection or vaccine-induced immunity. (Standard practice).
  — When prevaccination testing is done, the first dose of vaccine should be administered at the same time the blood sample is obtained to ensure optimal vaccination coverage (Box 5) (9). Recommended.

• Postvaccination testing is not indicated for healthy adults (9,142,143). Not recommended.
• For persons with special conditions (e.g., immunodeficiency, HIV infection, or chronic hemodialysis), or who are likely to be exposed to HBV (e.g., sex partner of HBsAg positive person, health-care worker), postvaccination testing for anti-HBs is recommended 1–2 months after completion of the vaccination series. Nonresponders in this category should be revaccinated (149,150). Strongly recommended.

**Postexposure Prophylaxis**

• After any percutaneous (e.g., sharing injection-drug equipment or human bite) or mucosal (e.g., sexual) exposure to blood, an unvaccinated person should begin the vaccine series, and the exposure incident should be evaluated to determine if additional postexposure prophylaxis (i.e., HBIG) is required (Table 5) (9,47). Strongly recommended.
  — The first dose of hepatitis B vaccine should be administered immediately, and the remaining doses 1 and 6 months later (Table 4). (Standard practice).
  — For an exposed person who has begun but not completed the vaccine series, subsequent vaccine doses should be administered as scheduled. (Standard practice).
  — The person who was the source of the exposure should be tested for HBsAg, even if that person was previously vaccinated. If the source person is HBsAg-positive, HBIG (0.06 mL/kg body weight intramuscular) should be administered to the exposed person as soon as possible and ≤7 days after the exposure. (Standard practice).
  — Postexposure prophylaxis is not necessary for a fully vaccinated person after exposure to HBV (9,47,138). Not recommended.

**Serologic Testing for Hepatitis B Virus Infection**

• Correctional facilities should consider routine testing of long-term inmates for chronic HBV infection (Box 2, Table 2), to facilitate rapid vaccination of contacts, direct counseling for preventing secondary transmission, and ensure medical evaluation of infected persons. If routine testing is not performed, testing should be considered for inmates in groups with risk factors for chronic HBV infection (e.g., injection-drug use, men who have sex with men, foreign-born persons from countries with high rate of infection). Indicated.
  — Cases of acute hepatitis B should be reported to the appropriate public health authority. (Standard practice).
  — If an inmate is identified as having chronic HBV infection, the case should be reported in those states where reporting is required. (Standard practice).
  — Identification of acute hepatitis B should prompt an epidemiologic investigation by correctional officials, in collaboration with the appropriate health authorities, to identify the source of infection and provide appropriate postexposure prophylaxis (Box 6, Table 4) to nonimmunized contacts at risk for infection. (Standard practice).
  — Persons diagnosed with acute hepatitis B should be observed for progressive liver dysfunction and evidence of acute liver failure. (Standard practice).
Chronic Hepatitis B Treatment

- Inmates identified as having chronic HBV infection during medical screening should be evaluated to determine the presence and extent of chronic liver disease and the potential benefit of antiviral therapy. Therapies for treatment of hepatitis B include interferon, alpha, lamivudine, and adefovir. Treatment of patients with chronic hepatitis B should be conducted in consultation with a specialist experienced with these treatment regimens. (Standard practice).
- All long-term correctional facilities should establish criteria for identifying prisoners who might benefit from treatment, on the basis of the latest treatment guidelines. (Standard practice).
- Discharge planning for persons with chronic HBV infection should include referral to medical care, risk-reduction programs, and social services necessary to maintain behavior changes; vaccination of contacts should also be arranged before patient discharge. (Standard practice).

Recommendations for Adult Correctional Facilities — Hepatitis C Virus Infection

Testing for Hepatitis C Virus Infection

- All inmates should be asked questions regarding risk factors for HCV infection during their entry medical evaluations, and all inmates reporting risk factors for HC. J. Pfister, M.S., Wisconsin State Laboratory of Hygiene; T. Lincoln, M.D., Hampden County Correctional Center, Ludlow, Massachusetts; personal communications; 2001). Recommended.
- The sensitivity of risk factor-based screening should be periodically determined by seroprevalence surveys, in combination with ascertainment of demographic and risk-factor information. Serologic testing of expanded groups of inmates or all inmates is recommended when
  - self-reported history of risk factors alone identifies <75% of anti-HCV positive inmates; or
  - the prevalence of risk factors for HCV infection, including injection-drug use, is known to be high (>75%), and a high prevalence exists (>20%) of HCV infection among inmates who deny risk factors. (Standard practice).
- Anti-HCV–positive persons should be reported if required by state regulations. (Standard practice).
- Adults with signs or symptoms suggestive of viral hepatitis should have appropriate diagnostic testing to differentiate acute hepatitis A, hepatitis B, or hepatitis C and to determine if the patient has chronic HBV or HCV infection (Box 2). (Standard practice).
- Cases of acute hepatitis C should be reported to the appropriate public health authority. (Standard practice).
- Identification of an inmate with acute hepatitis C, including ones who have been incarcerated for >6 months, should prompt an epidemiologic investigation by correctional officials, in collaboration with the appropriate health authorities, to identify the source of the infection. Depending on the results of the investigation, testing of contacts might be indicated (Box 7). (Standard practice).
- Adults who test positive for anti-HCV should receive further medical evaluation to determine chronic infection and liver disease. (Standard practice).

Postexposure Management for HCV

- After a percutaneous or permucosal exposure to blood, the source person should be tested for anti-HCV. If the source person is anti-HCV–positive, the exposed person should be tested for anti-HCV and ALT activity at baseline and 4–6 months later. For earlier diagnosis, testing for HCV RNA can be performed at 4–6 weeks (10). Recommended.
- Immune globulin and antiviral agents are not recommended for postexposure prophylaxis of hepatitis C (10). Not recommended.

Chronic Hepatitis C Treatment

- All anti-HCV–positive inmates should be evaluated for evidence of chronic HCV infection, including the presence and extent of chronic liver disease and candidacy for antiviral therapy. Treatment of patients with chronic hepatitis C should be conducted in consultation with a specialist familiar with these treatment regimens. (Standard practice).
- Inmates with chronic hepatitis C should receive hepatitis B vaccination and hepatitis A vaccination, if not previously immunized or known to be susceptible to infection (9–11,50). Recommended.
- Correctional facilities or systems should establish criteria based on the latest treatment guidelines for the identification of prisoners who might benefit from antiviral treatment. For HCV-infected patients who are actively abusing substances (e.g., drugs or alcohol), appropriate substance-abuse treatment should be initiated to limit disease transmission, reinfection, and liver disease progression (10,168,194–197). Recommended.
Adult Health Education and Release Planning (see Box 8)

- Prevention of HAV, HBV, and HCV infection should be incorporated into health education programs (e.g., programs for preventing HIV/AIDS) and include information concerning modes of disease transmission, methods for prevention, including risk reduction and immunization, disease outcomes, and options for treatment. (Standard practice).
- An integrated health education and risk reduction program should be established in each facility and include a written plan of health instruction completed by each inmate. (Standard practice).
- Such instruction should address a range of issues relevant to the diverse developmental and cultural composition of correctional populations, and should include basic skill development, literacy, and home economics, as well as tools needed to avoid behaviors that result in acquisition of HIV, hepatitis, and other bloodborne and sexually transmitted infections. (Standard practice).
- Teachers should be trained professionals or inmate peers with specific training to teach comprehensive life-skills programs, including health education. (Standard practice).
- A system for periodic evaluation, updating and improvement should exist. (Standard practice).
- Documentation of hepatitis A or hepatitis B vaccination should be included in the medical record retained within the correctional system, as well as in any medical record provided to other health-care providers. In addition, the vaccinated person should be provided a personal immunization record. (Standard practice).
- Correctional health facilities should establish links with community and public health facilities, and where available, with immunization registries, to ensure tracking and completion of hepatitis A and hepatitis B vaccine series. (Standard practice).
- Persons with chronic HBV or HCV infection should be — counseled regarding preventing transmission to household, sexual, and drug-use contacts, including risk reduction and condom use; — provided referral for hepatitis B vaccination of contacts; — counseled regarding ways to reduce further liver damage, including limiting alcohol and drug use, and afforded substance-abuse treatment when appropriate; and — provided aftercare that includes medical follow-up. (Standard practices).

Preventing and Controlling Hepatitis Virus Infections among Correctional Staff

Hepatitis A Virus Infection

- Hepatitis A is not occupationally acquired in the healthcare or correctional setting, and neither routine screening nor routine vaccination of staff is indicated. Indicated.

Infection Control Plan for HBV and HCV Prevention

- Measures to prevent occupational exposure to HBV and HCV among correctional workers should be integrated into each facility’s bloodborne pathogen and infection control plan according to the requirements of the Occupational Safety and Health Administration (OSHA) or the respective state OSHA. Elements of this plan should be coordinated with the infection control plan for correctional workers for all other infectious agents (e.g., HIV and Mycobacterium tuberculosis). (Standard practice).
- The plan should cover all employees (including inmates who are assigned work duties at a correctional facility) who could be reasonably anticipated, as the result of job duties, to be exposed to blood, bodily fluids, or other materials that might contain HBV or HCV. (Standard practice).
- The plan should identify tasks, procedures, and job classifications in which occupational exposure to potentially infectious material occurs — without regard to personal protective clothing and equipment. The plan must be accessible to employees and employee representatives. The employer should review and update the plan at least annually — more often if necessary to accommodate changes or recommendations from appropriate agencies. (Standard practice).
- The plan should mandate standard (i.e., universal) precautions for all contact with blood or body fluids. This should include procedures used to prevent needle sticks, including use of safer needle devices (139), to minimize splashing and spraying of potentially infectious material, and to ensure appropriate disinfection and decontamination of potentially contaminated surfaces and equipment, and appropriate disinfection and disposal of infectious material and contaminated clothing (198). As a part of the plan, correctional facilities should require employees to use appropriate personal protective equipment (e.g., gloves, gowns, masks, mouthpieces, and resuscitation bags) that are provided by the employer. (Standard practice).
• The plan should ensure that all workers are familiar with all aspects of infection control, including bloodborne pathogens and their transmission, the written exposure control plan, engineering and work practice controls, personal protective equipment, hepatitis B vaccine, response to emergencies involving blood, how to handle exposure incidents, the postexposure evaluation and follow-up program, and signs/labels/color-coding to alert persons to potentially infectious material. (Standard practice).
• Plan administrators should consider strategies to overcome the unique barriers to an effective infection control plan in a correctional environment (41). For example, potential inaccessibility of sharps disposal containers might necessitate using specific safe-needle devices and other strategies to minimize needle-stick injuries in correctional health-care settings. (Standard practice).
• A work practices program should be established that includes standard operating procedures for all activities having exposure potential. No worker should engage in such tasks or activities before receiving training pertaining to the procedures, work practices, and protective equipment required for that task. (Standard practice).

Preexposure Hepatitis B Vaccination and Postexposure Management for HBV and HCV
• Hepatitis B vaccination is recommended for all previously unvaccinated persons (e.g., correctional and medical staff) whose work duties involve exposure to blood or other potentially infectious body fluids (9,138,199). Strongly recommended.
• Prevaccination serologic screening is not indicated for persons being vaccinated because of occupational risk, unless the hospital or health-care organization considers screening cost-effective. Indicated.
• Staff with continued contact with patients or blood and who are at ongoing risk for percutaneous injuries should be tested for anti-HBs 1–2 months after completion of the 3-dose vaccination series. Staff who do not respond to a primary vaccine series should complete a second 3-dose vaccine series or be evaluated to determine if they are HBsAg-positive (47,199). (Standard practice).
• For correctional workers who have no contact with inmates and no routine exposure to blood and body fluids in the correctional setting, timely postexposure prophylaxis should be provided if an exposure occurs, rather than routine vaccination (47,199). (Standard practice).
• Evaluation for appropriate postexposure prophylaxis for an employee who has had an exposure incident should be performed in a timely fashion according to recommendations for HBV and HCV (47). Strongly recommended.
• When an exposure to potentially infectious blood or body fluid has occurred, a blood sample from the source person should be tested for HBsAg and anti-HCV. If the source person cannot be identified or tested, the respective postexposure protocol (i.e., HBV or HCV) should be followed to evaluate the need for postexposure prophylaxis or follow-up. (Standard practice).
• Appropriate postexposure prophylaxis and follow-up for HBV infection after exposure is dependent on the HBsAg status of the source person, as well as the immunization status of the exposed person (Tables 2 and 4) (47) (see Recommendations for Adult Inmates). (Standard practice).
• If the source person is anti-HCV positive, CDC guidelines for postexposure follow-up should be followed (10,47) (see Recommendations for Adult Inmates). (Standard practice).

HBV or HCV Serologic Testing
• Routine testing for HBV or HCV infection is not recommended for correctional workers, except as described for hepatitis B vaccination or postexposure management (10,48,123,200). Indicated.

Implementation of Recommendations
• The unique nature of correctional institution populations necessitates close collaboration between corrections and public health personnel at state and local levels for effective implementation of the recommendations contained in this report. Preventing and controlling viral hepatitis among incarcerated and released persons, and among persons in the communities to which they return, requires defining specific roles for each public agency.
• Correctional officials should review these recommendations and develop written policies for their implementation. Policies should include implementation by contractors where correctional health care is provided by the private sector. Correctional officials should also evaluate the implementation of these recommendations to monitor 1) the proportion of inmates (adults and juveniles) who begin and complete the hepatitis B vaccine series; 2) prevalence of immunity to HBV infection among incoming inmates; 3) vaccine-series–completion rates for released prisoners; 4) proportion of inmates tested for HCV infection and reasons that inmates decline testing;
• Correctional officials should establish close working relationships with state and local health departments to ensure awareness of the correctional system’s viral hepatitis prevention and control activities. Written agreements can better ensure that all agencies participate in such activities as 1) reporting and investigating acute cases of viral hepatitis among inmates; 2) reporting of inmates with chronic HBV and HCV infection in states where this is a requirement; 3) vaccination of contacts of inmates with chronic HBV infection; and 4) follow-up of inmates released before completing the hepatitis A or hepatitis B vaccine series, or before completing treatment for chronic HBV or HCV infection. Correctional officials should also collaborate with health department staff to provide hepatitis education and counseling to inmates and correctional employees.

• Public health departments should work closely with correctional systems and facilities to develop community-based strategies for preventing and controlling viral hepatitis. Integration of correctional health care into such strategies can be facilitated through designation of health department personnel to provide epidemiologic and programmatic assistance to correctional facilities. Other activities might include 1) development of record-keeping systems that facilitate hepatitis B vaccination; 2) case management of persons on antiviral therapy for chronic hepatitis C or hepatitis B; 3) substance-abuse treatment where appropriate; and 4) development of training courses for correctional facility staff.

• Public health departments should be considered resources for consultation on all aspects of viral hepatitis prevention and control, including quality assurance of laboratory testing services. Development of training and educational programs for correctional staff should include such topics as diagnosis of viral hepatitis and interpretation of laboratory test results, vaccination delivery and assessment of vaccination programs, disease reporting, and health education. Health department officials should provide educational information to senior-level prison and jail officials and to county and other elected officials.

• Public health departments should develop mechanisms that encourage reporting of viral hepatitis cases identified in correctional facilities. In addition, mechanisms should be established to provide epidemiologic consultation for investigations of acute disease in the complex setting of the correctional facility. Other areas for which mechanisms should be established include follow-up of persons with chronic HBV and HCV infection for vaccination of contacts (HBV) and appropriate counseling and referral for medical follow-up and treatment.

**Internet Resources**

The following Internet sites provide additional information (listed by source, topic, and website):


**Published Resources**

• CDC. Immunization of adolescents: recommendations of the Advisory Committee on Immunization Practices, the American Academy of Pediatrics, the American Academy of Family Physicians, and the American Medical Association. MMWR 1996;45 (No. RR-13):1–16

References
91. Kaufman ML, Faiver KL, Harness JK. Hepatitis B markers among
90. Bader T. Hepatitis B carriers in the prison population [Letter]. New
89. Kibby T, Devine J, Love C. Prevalence of hepatitis B among men


139. Occupational Safety and Health Standards. [29 CFR § 1910.1030].


149. CDC. Recommendations for preventing transmission of infections among chronic hemodialysis patients. MMWR 2001;50 (No. RR-5):1–43.
160. Lofgren RP, Paul JM, Kefalos SG, Nichol KL. A multifaceted influenza vaccination program can be exported successfully to a different clinical site. Clinical Research 1990;38:864A.
167. CDC. Guidelines for laboratory testing and result reporting of antibody to hepatitis C virus. MMWR 2003 (In press).


Hepatitis A Vaccine

Long-term protection from hepatitis A virus (HAV) infection can be achieved through active, preexposure vaccination with hepatitis A vaccine. Inactivated hepatitis A vaccines licensed for use in the United States are Havrix® (GlaxoSmithKline Biologicals, Rixensart, Belgium), VAQTA® (Merck & Co., Inc., Whitehouse Station, New Jersey), and Twinrix® (GlaxoSmithKline Biologicals), a combined hepatitis A and hepatitis B vaccine (1,2). All are produced from HAV grown in cell culture, inactivated with formalin, and formulated with alum adjuvant in pediatric and adult dosages that are 94%–100% effective in preventing clinical disease among juveniles and adults when administered according to recommended schedules (1,2) (Table 3).

Protective levels of antibody to HAV (anti-HAV) develop among 94%–100% of vaccinated persons within 1 month after administration of the first dose. A second dose results in protective levels of antibody among all persons vaccinated, and is considered necessary for long-term protection. Estimates of antibody persistence suggest protective levels of anti-HAV persist for ≥20 years (1).

A delay in administration of the second vaccine dose does not result in lowered final antibody levels or seroconversion rates, and restarting the vaccine series if the second dose is delayed is not needed. Vaccination begun with vaccine from one manufacturer can be completed with vaccine from the other (3,4). Hepatitis A vaccine can be administered at the same time as other vaccines, including hepatitis B vaccine, without affecting immunogenicity or increasing the frequency of adverse events.

Adverse Reactions

The most frequently reported adverse reactions occurring ≤3 days after vaccination are soreness at the injection site (53%–56%), headache (14–16%), and malaise (7%). Reviews of data from multiple sources have not identified any serious adverse events among juveniles or adults associated with hepatitis A vaccination (1). Any adverse event occurring after hepatitis A vaccination should be reported to the Vaccine Adverse Events Reporting System (VAERS). Reporting forms can be obtained by calling 1-800-822-7967.

Contraindications

Hepatitis A vaccine should not be administered to persons with a history of hypersensitivity reactions to alum, or for Havrix or Twinrix, to the preservative 2-phenoxyethanol. The safety of hepatitis A vaccination during pregnancy has not been determined. However, because this is an inactivated vaccine, the theoretical risk to the developing fetus is low. The risk associated with vaccination should be weighed against the risk for hepatitis A among women who might be at high risk for exposure to HAV infection. No special precautions are needed when vaccinating immunocompromised persons.

Serologic Testing for HAV Infection

Antibody produced after HAV infection results in lifelong immunity. Among adult populations with high rates of prior HAV infection, prevaccination testing can reduce costs by avoiding the vaccination of persons with prior immunity. However, the vaccination of an immune person does not increase the risk for adverse events. The decision to test should be based on 1) expected prevalence of immunity; 2) cost of vaccination compared with cost of serologic testing; and 3) likelihood that testing will not interfere with initiating vaccination.

Prevaccination testing of younger juveniles (ages <15 years) is not indicated because of their low prevalence of infection. Prevaccination testing is most likely to be cost-effective for older juveniles and adults born in countries, or who have been residents for extensive periods in countries, with a high endemicity of HAV infection (e.g., Mexico, South and Central America, Africa, and all of Asia except Japan), populations with historically high rates of infection (e.g., American Indians or Alaska Natives), and those engaging in behaviors that place them at high risk for infection (e.g., drug users or men who have sex with men). Because anti-HAV prevalence increases with age, prevaccination testing of any person aged ≥40 years would likely be cost-effective (1). Commercially available tests for total anti-HAV can be used for prevaccination testing. Postvaccination testing is not indicated because of high rates of vaccine response among both adults and juveniles. In addition, no Food and Drug Administration (FDA)-approved testing method exists that has the sensitivity to detect low anti-HAV concentrations after vaccination.
Hepatitis B Vaccine

Vaccines available in the United States use hepatitis B surface antigen (HBsAg) produced in yeast cells by recombinant deoxyribonucleic acid (DNA) technology, and are formulated to contain 5–40 µg HBsAg protein/mL and 0.5 mg/mL aluminum hydroxide as the adjuvant. The two available single antigen hepatitis B vaccines are Recombivax HB® (Merck & Co., Inc., Whitehouse Station, New Jersey) and Engerix-B® (GlaxoSmithKline Biologicals, Rixensart, Belgium) (5). A combination hepatitis A and hepatitis B vaccine, Twinrix, is also licensed for persons aged ≥18 years old (2) (Table 4).

Antibody Response to Vaccination

Licensed formulations for both vaccines produce high (≥95%) rates of protective antibody (anti-HBs >10 mIU/mL) when the complete series is administered in different schedules to infants, juveniles, and adults aged <40 years (5). Among healthy adults, 30%–50% develop a protective antibody response after the first vaccine dose, 75% after the second dose, and >95% after the third dose (5–9). Increasing the interval between the first and second dose of vaccine has little effect on immunogenicity or final antibody titer, although data are limited regarding intervals >2 months among adults (5,8). The third dose confers the maximum rate of seroprotection; it primarily acts as a booster and confers optimal long-term protection through the induction of maximum immune memory (5,9). Both licensed vaccines administered on a 0-, 1-, and 6-month schedule produce a >95% final seroprotection rate among adolescents and healthy young adults, and studies indicate that vaccination of adolescents and adults on a 0-, 2-, and 4-month, and adolescents on a 0-, 12-, and 24-month schedule, achieved final seroprotection rates similar to the 0-, 1-, and 6-month schedule (8–10). In addition, a 2-dose vaccination series using Recombivax HB® at the adult dosage has been demonstrated among adolescents aged 11–15 years to produce protective antibody responses equivalent to that of the 3-dose series, although the long-term protection afforded from this schedule is not known (8,11).

The duration of vaccine-induced antibody and protection from hepatitis B virus (HBV) infection has been evaluated among vaccinated infants, juveniles, and adults (5,12–14). Studies indicate that although loss of detectable anti-HBs has ranged from 13% to 60% by 9–15 years after vaccination, immune memory provides protection from HBV infection, and protection remains intact for ≥15 years, the longest period for which follow-up data are available (5,12–14). Because of the long duration of protection afforded by the 3-dose vaccine series, booster doses of vaccine are not needed among vaccinated immunocompetent juveniles or adults.

Adverse Reactions

Adverse reactions associated with hepatitis B vaccine include pain at the injection site (3%–29%) and a temperature ≥37.7°C (1%–6%), although these effects are reported no more frequently among vaccine recipients than among placebo recipients in controlled trials (5). Anaphylaxis has been reported in 1/600,000 vaccine recipients; however, no deaths have been attributed to vaccination. A number of case reports and case series have claimed an association between hepatitis B vaccination and serious adverse health events (e.g., multiple sclerosis) (15,16); however, these have not been proven by other epidemiologic studies (17–22). Adverse events suspected to be associated with hepatitis B vaccination should be reported to VAERS, and reporting forms can be obtained by calling 1-800-822-7967.

References

Prevention and Control of Infections with Hepatitis Viruses in Correctional Settings

External Consultants: Katie Arnold, M.D., Georgia Department of Human Resources, Atlanta, Georgia; Jack Beck, Legal Aid Society of New York, New York, New York; Ellen Bentz, National Minority AIDS Council, Washington, D.C.; David Bergmire-Sweat, M.P.H., American Social Health Association, Research Triangle Park, North Carolina; Joseph Bick, M.D., California Medical Facility, Vacaville, California; Kitty Bradley, Hepatitis Foundation International, Cedar Grove, New Jersey; Marie Bresnahan, M.P.H., American Liver Foundation, New York, New York; Judy Byrnes, Georgia Department of Human Resources, Atlanta, Georgia; William Cassidy, M.D., Louisiana State University Health Science Center, Baton Rouge, Louisiana; Grace Chao, M.D., University of Texas Medical Branch, Galveston, Texas; Robert L. Cohen, M.D., New York, New York; Thomas Conklin, M.D., Hampden County Correctional Center, Ludlow, Massachusetts; Anne S. DeGroot, M.D., Brown University School of Medicine, Providence, Rhode Island; Carmen C. Deseda, M.D., Puerto Rico Department of Health, San Juan, Puerto Rico; David Doolen, National Minority AIDS Council, Washington, D.C.; Ronald Feinstein, M.D., The Society for Adolescent Medicine, Birmingham, Alabama; Felicia Fong, Minnesota Department of Health, Minneapolis, Minnesota; Peter M. Ford, M.D., Queen's University, Ontario, Canada; Alan Franciscus, Hepatitis C Support Project, San Francisco, California; Joe Goldenson, M.D., Jail Health Services, San Francisco, California; Brenda Goldhammer, M.P.H., California STD/HIV Prevention Training Center, Berkeley, California; Betty Gondles, Ph.D., American Correctional Association, Arlington, Virginia; James A. Gondles, Jr., American Correctional Association, Lanham, Maryland; Betty Adams Greene, Lanham, Maryland; Robert B. Greifinger, M.D., The Bromeen Group, Dubbs Ferry, New York; Ted Hammet, Ph.D., Atr Associates, Cambridge, Massachusetts; Edward Harrison, National Commission on Correctional Health Care, Chicago, Illinois; Alicia Herman, Association of State and Territorial Health Officials, Washington, D.C.; Evalyn Horowitz, M.D., California Department of Corrections, Elk Grove, California; Judy Hubbard, Cobb County Board of Health, Atlanta, Georgia; Bert Hurowitz, Florida Department of Corrections, Tallahassee, Florida; Bethanne Jenks, M.D., Atlanta, Georgia; Glenn Johnson, M.D., Wackenhut Correctional Corporation, Austin, Texas; Renee Kanan, M.D., Correct Care Management, Seattle, Washington; Mike Kelley, M.D., Texas Department of Corrections, Huntsville, Texas; Steve Koester, Ph.D., University of Colorado at Denver, Denver, Colorado; Ronald Koretz, M.D., UCLA Medical School, Los Angeles, California; Madie LaMarre, Georgia Department of Corrections, Atlanta, Georgia; Daryl Lau, M.D., University of Texas Medical Branch, Galveston, Texas; Thomas Lincoln, M.D., Hampden County Correctional Center, Hampden City, Massachusetts; Leigh Lipson, National Association of County and City Health Officials, Washington, D.C.; Grace Macalino, M.D., Brown University Department of Community Health; Fred Maue, M.D., Pennsylvania State Department of Corrections, Camp Hill, PA; Joseph McGarry, M.D., Colorado Department of Corrections, Denver, Colorado; Lynn Mercedes, Minnesota Department of Health, Minneapolis, Minnesota; Amy B. Middleman, M.D., Baylor College of Medicine, Houston, Texas; J. Tom Morgan, J.D., National District Attorney Association, Decatur, Georgia; Joseph E. Paris, M.D., Georgia Department of Corrections, Atlanta, Georgia; David Parr, M.D., University of Texas Medical Branch, Galveston, Texas; John R. Pfister, M.S., Wisconsin State Laboratory of Hygiene, Madison, Wisconsin; Randy Pope, National Association of State and Territorial AIDS Directors, Washington, D.C.; Michael Puisis, D.O., Evanston, Illinois; Aaron Roome, Ph.D., Connecticut Department of Public Health, Hartford, Connecticut; David W. Roush, Ph.D., National Juvenile Detention Association, Albion, Michigan; Tom Saari, M.D., University of Wisconsin Medical School, Madison, Wisconsin; Tamara Serwer, J.D., Southern Center for Human Rights, Atlanta, Georgia; Ronald M. Shansky, M.D., Society of Correctional Physicians, Chicago, Illinois; Frederick E. Shaw, M.D, J.D., Texas Department of Health, Austin, Texas; Gary Shostak, M.P.H., Massachusetts Department of Youth Services, Boston, Massachusetts; Richard Staller, Louisiana Department of Public Safety and Corrections, Baton Rouge, Louisiana; Jason Stanford, Georgia Department of Human Resources, Atlanta, Georgia; Michelle Staples-Horne, M.D., Georgia Department of Corrections, Atlanta, Georgia; Leah Stockett, Association of State and Territorial Health Officials, Washington, D.C.; David Stoff, Ph.D., National Institutes of Health, Washington, D.C.; Sara Straub, Florida Department of Corrections, Tallahassee, Florida; Betsy Stubblefield, HIV Education Prison Project, Providence, Rhode Island; Thelma Thiel, R.N., Hepatitis Foundation International, Cedar Grove, New Jersey; David Thomas, M.D., J.D., Florida Department of Corrections, Tallahassee, Florida; Ann Thomas, M.D., Oregon Health Division, Portland, Oregon; Louis Tripoli, M.D., Spectrum Healthcare Services, St. Louis, Missouri; Theresa Turski, Georgia Department of Human Resources, Atlanta, Georgia; R.J. Verdeny, American Correctional Association, Lanham, Maryland; John Vertefeuille, Maryland AIDS Administration, Baltimore, Maryland; John Vierling, M.D., Cedars-Sinai Medical Center, Los Angeles, California; Deborah Wexler, M.D., Immunization Action Coalition, St. Paul, Minnesota; Steven Wiersma, M.D., Florida Department of Health, Tallahassee, Florida; Lester N. Wright, M.D., New York State Department of Corrections, Rensselaer, New York; Barry Zack, M.P.H., Centerforce Health Programs, San Quentin, California; Jacqueline Zalumas, Ph.D., Emory University, Atlanta, Georgia.

CDC Consultants: Miriam J. Alter, Ph.D., Beth P. Bell, M.D.; Amy Khan, M.D.; Linda A. Moyer; Ian A. Williams, Ph.D., Division of Viral Hepatitis, National Center for Infectious Diseases; Douglas Trout, M.D., National Institute for Occupational Safety and Health; Steve Hadler M.D., National Immunization Program; Anne Spaulding, M.D., National Center for HIV, STD and TB Prevention.


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Goal and Objectives

This MMWR provides information and recommendations regarding hepatitis prevention and control in correctional settings. The recommendations were prepared by CDC staff from the National Center for Infectious Diseases, in consultation with external consultants. The goal of this report is to provide information and recommendations for physicians, health-care delivery staff, corrections staff, administrators, and other public health professionals who deliver services to incarcerated persons. After completing this educational activity, the reader should be able to 1) identify the risk factors for transmission of viral hepatitis among incarcerated persons; 2) describe the epidemiology of viral hepatitis in the United States; 3) describe the outcome of infection with hepatitis viruses; 4) describe methods used to prevent and control viral hepatitis in juvenile and adult correctional settings; and 5) identify infection control measures to prevent occupational exposure to hepatitis B and C viruses among correctional workers.

To receive continuing education credit, please answer all of the following questions.

1. Which drugs are currently approved by the Food and Drug Administration (FDA) for treatment of persons with chronic hepatitis B? (A) Alpha interferon, lamivudine, and adefovir dipivoxil. (B) Ribavirin and pegylated interferon. (C) Alpha interferon and ribavirin. (D) Milk thistle and ribavirin.

2. Where should adverse events be reported that are thought to be associated with vaccination? (A) CDC. (B) The National Institutes of Health. (C) The Vaccine Adverse Events Reporting System. (D) FDA.

3. Which of the following are risk factors for viral hepatitis among inmates? (A) Injection-drug use. (B) Sex among inmates and between inmates and staff. (C) Percutaneous exposures. (D) Occupational exposures. (E) All of the above. (F) None of the above.

4. Chronic hepatitis B virus (HBV) infection might result in . . . (A) an asymptomatic carrier state. (B) chronic persistent hepatitis. (C) chronic active hepatitis which progresses to cirrhosis. (D) chronic active hepatitis which progresses to liver cancer. (E) all of the above. (F) none of the above.

5. No association has been found with hepatitis C virus (HCV) infection and military service or exposures resulting from medical, surgical, or dental procedures, tattooing, acupuncture, ear piercing, or foreign travel. (A) True. (B) False.

6. Routine HCV testing is recommended for . . . (A) persons who ever injected illegal drugs. (B) persons who received clotting factor concentrate produced before 1987. (C) persons ever on long-term hemodialysis. (D) persons with persistently abnormal alanine aminotransferase (ALT). (E) prior recipients of transfusions or organ transplants including those who received blood or blood components or received an organ transplant before July 1992. (F) all of the above.

7. A delay in the administration of the second hepatitis A vaccine dose results in lowered final antibody levels or seroconversion, and if the second dose is delayed, the series should be restarted. (A) True. (B) False.

8. Adults and adolescents for whom hepatitis A vaccination is recommended include . . . (Indicate all that apply.) (A) travelers or workers in countries with high rates of infection. (B) men who have sex with men. (C) illegal drug users. (D) persons with chronic liver disease. (E) adults living in Texas. (F) all of the above.

9. Hepatitis B vaccination should be initiated even when completion of a series cannot be ensured, because relatively high levels of immunity are provided by one or two doses of vaccine. Administration of even a partial vaccine series (i.e., 1–2 doses) during incarceration might avert new infections. (A) True. (B) False.

10. Combined with routine infant vaccination, catch-up vaccination of adolescents is not considered an effective means to rapidly eliminate HBV transmission in the United States. (A) True. (B) False.

11. Screening for antibody to HCV is done with enzyme immunoassay (EIA) testing, which detects >97% of infected patients, but does not distinguish between acute, chronic, or resolved infection. Supplemental testing with a more specific assay (i.e., recombinant immunoblot assay [RIBA]) prevents reporting of false-positives. (A) True. (B) False.

12. Which of the following drugs are currently FDA-approved for treatment of persons with hepatitis C? (Indicate all that apply.) (A) Interferon. (B) Pegylated interferon. (C) Ribavirin. (D) Lamivudine.

13. Postexposure prophylaxis is necessary for a fully vaccinated juvenile after a percutaneous or sexual exposure that might contain HBV. (A) True. (B) False.

14. Juveniles with chronic HBV or HCV infections, and their parents or guardians, should be counseled regarding which of the following? (Indicate all that apply.) (A) Preventing transmission to household, sexual, and drug-use contacts. (B) Referral for hepatitis B vaccination of susceptible contacts. (C) Ways to reduce further liver damage, including limiting alcohol and drug use. (D) Postexposure prophylaxis.
15. Previously unvaccinated hepatitis B surface antigen-negative pregnant women should not be vaccinated; pregnancy is a contraindication to vaccination.
   A. True.
   B. False.

16. CDC recommends that all persons who receive a medical evaluation in a correctional facility be administered hepatitis B vaccine, unless they have proof of completion of the vaccine series.
   A. True.
   B. False.

17. For which of the following persons is routine testing for HCV infection recommended? (Indicate all that apply.)
   A. Persons who have been tattooed.
   B. Persons who have a history of sexually transmitted disease.
   C. Persons who have ever injected illegal drugs.
   D. Persons who have had a transfusion of blood or blood components before July 1992.
   E. Persons who have received clotting factor concentrates made before 1987.

18. Indicate your work setting.
   A. State/local health department.
   B. Other public health setting.
   C. Hospital clinic/private practice.
   D. Managed care organization.
   E. Academic institution.
   F. Other.

19. Which best describes your professional activities?
   A. Patient care — emergency/urgent care department.
   B. Patient care — inpatient.
   C. Patient care — primary-care clinic or office.
   D. Laboratory/Pharmacy.
   E. Public health.
   F. Other.

20. I plan to use these recommendations as the basis for . . . (Indicate all that apply.)
   A. health education materials.
   B. insurance reimbursement policies.
   C. local practice guidelines.
   D. public policy.
   E. other.

21. Each month, approximately how many patients do you see who have hepatitis?
   A. None.
   B. 1–10.
   C. 11–100.
   D. 101–1,000.
   E. >1,000.

22. How much time did you spend reading this report and completing the exam?
   A. <2.0 hours.
   B. >2.0 hours but <3.0 hours.
   C. >3.0 but <4.0 hours.
   D. >4.0 hours.
   E. other.

\[\text{Fill in the appropriate blocks to indicate your answers. Remember, you must answer all of the questions to receive continuing education credit!}\]
23. After reading this report, I am confident I can identify the risk factors for transmission of viral hepatitis among incarcerated persons.
   A. Strongly agree.
   B. Agree.
   C. Neither agree nor disagree.
   D. Disagree.
   E. Strongly disagree.

24. After reading this report, I am confident I can describe the epidemiology of viral hepatitis in the United States.
   A. Strongly agree.
   B. Agree.
   C. Neither agree nor disagree.
   D. Disagree.
   E. Strongly disagree.

25. After reading this report, I am confident I can describe the outcome of infections with hepatitis viruses.
   A. Strongly agree.
   B. Agree.
   C. Neither agree nor disagree.
   D. Disagree.
   E. Strongly disagree.

26. After reading this report, I am confident I can describe methods used to prevent and control viral hepatitis in juvenile and adult correctional settings.
   A. Strongly agree.
   B. Agree.
   C. Neither agree nor disagree.
   D. Disagree.
   E. Strongly disagree.

27. After reading this report, I am confident I can identify infection control measures to prevent occupational exposure to HBV and HCV among correctional workers.
   A. Strongly agree.
   B. Agree.
   C. Neither agree nor disagree.
   D. Disagree.
   E. Strongly disagree.

28. The objectives are relevant to the goal of this report.
   A. Strongly agree.
   B. Agree.
   C. Neither agree nor disagree.
   D. Disagree.
   E. Strongly disagree.

29. The tables and boxes are useful.
   A. Strongly agree.
   B. Agree.
   C. Neither agree nor disagree.
   D. Disagree.
   E. Strongly disagree.
30. Overall, the presentation of the report enhanced my ability to understand the material.
   A. Strongly agree.
   B. Agree.
   C. Neither agree nor disagree.
   D. Disagree.
   E. Strongly disagree.

31. These recommendations will affect my practice.
   A. Strongly agree.
   B. Agree.
   C. Neither agree nor disagree.
   D. Disagree.
   E. Strongly disagree.

32. The availability of continuing education credit influenced my decision to read this report.
   A. Strongly agree.
   B. Agree.
   C. Neither agree nor disagree.
   D. Disagree.
   E. Strongly disagree.

33. How did you learn about this continuing education activity?
   A. Internet.
   B. Advertisement (e.g., fact sheet, MMWR cover, newsletter, or journal).
   C. Coworker/supervisor.
   D. Conference presentation.
   E. MMWR subscription.
   F. Other.