Viral Hepatitis A-E

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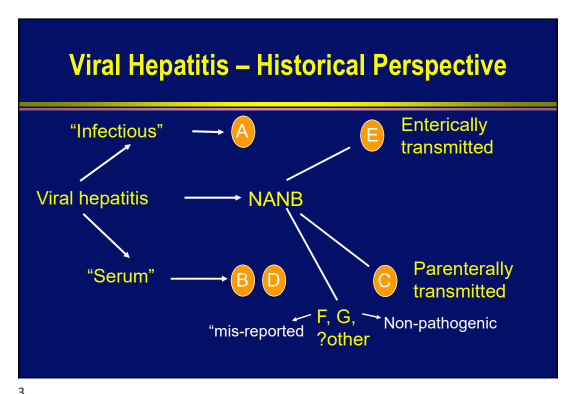
Former Director, Hospital Epidemiology, Occupational Health and Safety, UNC Hospitals, Chapel Hill, NC (1979-2017)

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Viral Hepatitis - Overview

- Primary infection of the liver caused by at least five unrelated viruses: A, B, C, D, E
- HAV and HEV
 - Fecal-oral route
 - Acute self-limited disease; no chronic infection
- HBV, HCV, HDV
 - Percutaneous or mucosal exposures to blood
 - Chronic infection major causes of cirrhosis and hepatocellular carcinoma worldwide



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		Туре	of Hepa	titis	
	A	В	С	D	E
Source of virus	feces	blood/ blood-derived body fluids	blood/ blood-derived body fluids	blood/ blood-derived body fluids	feces
Route of transmission	fecal-oral	percutaneous permucosal	percutaneous permucosal	percutaneous permucosal	fecal-oral
Chronic infection	no	yes	yes	yes	no
Prevention	pre/post- exposure immunization	pre/post- exposure immunization	blood donor screening; risk behavior modification	pre/post- exposure immunization; risk behavior modification	ensure safe drinking water

Clinical Features of Hepatitis

Common

- malaise
- anorexia
- nausea & vomiting
- fever

Less Common

- diarrhea
- arthralgias

- jaundice
- abdominal pain
- hepatomegaly
 - pruritis
 - rash

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Enterically Transmitted Viral Hepatitis

Hepatitis A—Highlights

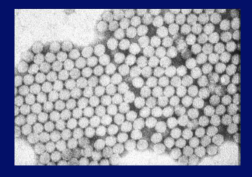
- Estimated 1.4 million clinical cases of hepatitis A annually worldwide
- Tens of millions of hepatitis A virus infections occur each year
- Universal childhood vaccination effective in countries with varying endemic rates
 - Reduces morbidity and mortality
 - However, incidence in US stable 2011-2016, then increasing

Wasley A, Epidemiologic Reviews 2006; https://www.cdc.gov/hepatitis/hav/havfaq.htm#general

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Hepatitis A Virus

- RNA Picornavirus
- Single serotype worldwide
- Acute disease and asymptomatic infection
- No chronic infection
- Protective antibodies develop in response to infection - confers lifelong immunity
- Vaccine preventable



Transmission of Hepatitis A Virus

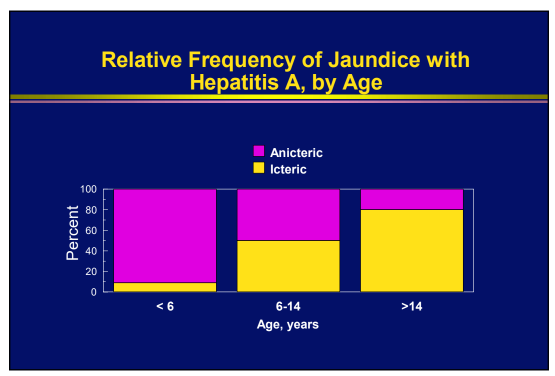
- Fecal-oral transmission leads to spread between close contacts
- Incubation period, 15-50 days (average: 28 days)
- Greatest period of communicability: 2 weeks before onset of jaundice
- Stable in environment for months

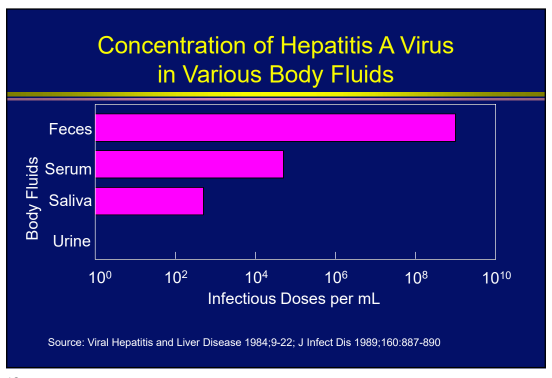
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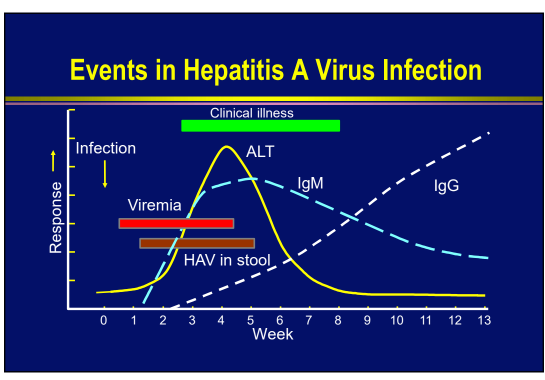
Transmission of Hepatitis A Virus

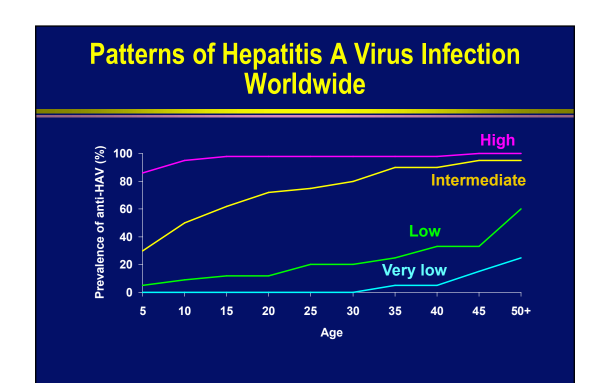
- Close personal contact
 (e.g., household contact, sex contact)
- Contaminated food (water)
 (e.g., infected food handlers, produce)
- Blood exposure (uncommon)
 (e.g., injecting drug use, transfusion)

Hepatiti	s A - Clinical Features
Incubation period	Average 30 days Range 15-50 days
Jaundice by age	<6 yrs <10% 6-14 yrs 40%-50% >14 yrs 70%-80%
Case fatality rate	0.3% (0.2%-2.0%)
Complications	Fulminant; cholestatic; relapsing
Chronic sequelae	None (prolonged shedding in neonates and immunocompromised)

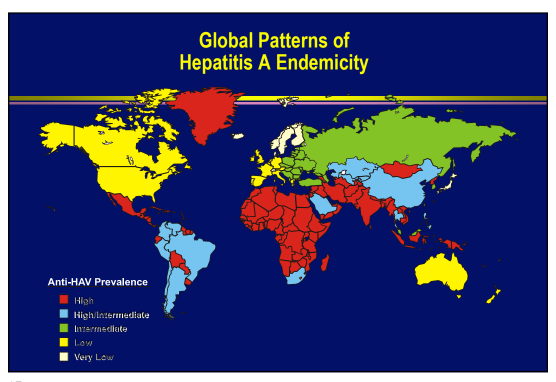




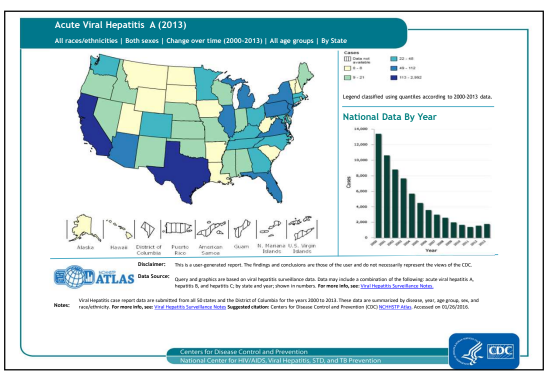




Hepatitis A Virus Transmission Global Patterns			
Endemicity High	Disease <u>Rate</u> Low	Age at <u>Infection</u> Early childhood	Transmission patterns Person to person; outbreaks uncommon
Intermediate	High	Late childhood/ young adults	Person to person; food and waterborne outbreaks
Low	Low to high	Late childhood/ young adults	Person to person; food and waterborne outbreaks
Very low	Very low	Adults	Travelers; outbreaks uncommon







Epidemiologic Features of Hepatitis A in the United States

- Wide geographic and temporal variation in incidence
 - Areas with highest rates account for most reported cases
- Most disease occurs in the context of community-wide outbreaks
- Infection transmitted from person to person in households and extended family settings
 - Facilitated by asymptomatic infection among children
- Groups at increased risk can be identified
 - Specific factor varies
 - Do not account for majority of cases
- No risk factor identified for 30%-40% of cases

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Type of risk	Risk category	Examples
Increased risk for HAV infection	Close personal contacts of persons with HAV infection [†]	Household contacts Caretakers Sexual contacts Persons who anticipate close personal contact with an international adoptee
	Occupational risk	Persons working with nonhuman primates Persons working with clinical or nonclinical material containing HAV in a research laboratory
	Persons who use drugs Persons in settings where services to adults are provided	Persons who use injection or noninjection drugs (i.e., all those who use illegal drugs) Group settings for persons with developmental disabilities Homeless shelters Syringe services programs Correctional facilities during outbreaks
	International travelers	Persons traveling to or working in countries with high or intermediate HAV endemicity
Increased risk for severe disease from HAV infection	Immunocompromised persons	Congenital or acquired immunodeficiency HIV infection Chronic renal failure, undergoing dialysis Solid organ, bone marrow, or stem cell transplant recipients Persons with diseases requiring treatment with immunosuppressive drugs/biologics (e.g., tumor necrosis alpha inhibitors), long-term systemic corticosteroids, radiation therapy
	Persons with chronic liver disease	Hepatitis B virus infection Hepatitis C virus infection Cirrhosis (any etiology) Fatty liver disease (hepatic steatosis) Alcoholic liver disease Autoimmune hepatitis Alanine aminotransferase or aspartate amino transferase level more than twice the upper limit of normal or persistently elevated for 6 months
	Age	Adults aged >40 years

ACIP 2020; www.cdc.gov/mmwr/volumes/69/rr/pdfs/rr6905a1-H.pdf

SAFETY OF HEPATITIS A VACCINE

- Most common side effects
 - Soreness/tenderness at injection site 50%
 - Headache 15%
 - Malaise 7%
- No severe adverse reactions attributed to vaccine
- Safety in pregnancy not determined risk likely low
- Contraindications severe adverse reaction to previous dose or allergy to a vaccine component
- No special precautions for immunocompromised persons

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Immunogenicity of Inactivated Hepatitis A Vaccines

- 2-dose series 6-18 months apart (dosage varies by manufacturer)
- 95%-100% had protective levels of antibody one month after receiving one dose
- 100% had protective levels of antibody one month after receiving second dose with high geometric mean titers

HEPATITIS A VACCINES

TARIF 1	Jacolnos usod	to prevent	honatitis A	virus Infection
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Vaccine	Trade name (manufacturer)	Age group (yrs)	Dosage	Route	Schedule	Booster
HepA, inactivated	Havrix (GlaxoSmithKline)	1–18	0.5 mL (720 ELISA units inactivated HAV)	IM	0, 6-12 months	None
(2 doses)		≥19	1 mL (1,440 ELISA units inactivated HAV)	IM	0, 6-12 months	None
HepA, inactivated	Vagta (Merck)	1-18	0.5 mL (25 units HAV antigen)	IM	0, 6-18 months	None
(2 doses)	1.87	≥19	1 mL (50 units HAV antigen)	IM	0, 6-18 months	None
Combined HepA	Twinrix (GlaxoSmithKline)	≥18 (primary)	1 mL (720 ELISA units inactivated HAV + 20 µg HBsAg)	IM	0, 1, 6 months	None
and HepB* (3 doses)		≥18 (accelerated)	1 mL (720 ELISA units inactivated HAV + 20 μ g HBsAg)	IM	0, 7, 21–30 days	12 months

Abbrevlations: ELISA = enzyme-linked immunosorbent assay; HAV = hepatitis A virus; HBsAg = hepatitis B surface antigen; HepA = hepatitis A; HepB = hepatitis B; IM = intramuscular.

ACIP 2020: https://www.cdc.gov/mmwr/volumes/69/rr/pdfs/rr6905a1-H.pdf

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IMMUNE GLOBULIN-PROPHYLAXIS

TABLE 2. Dosage recommendations for GamaSTAN human immune globulin for preexposure and postexposure prophylaxis against hepatitis A infection

Indication	Time	Dose*	Route
Preexposure prophylaxis	Up to 1 month duration of travel	0.1 mL/kg	IM
Preexposure prophylaxis	Up to 2 months duration of travel	0.2 mL/kg	IM
Preexposure prophylaxis	≥2 months duration of travel	0.2 mL/kg (repeat every 2 months)	IM
Postexposure prophylaxis	Within 2 weeks of exposure	0.1 mL/kg	IM

Abbreviation: IM = intramuscular.

ACIP 2020: https://www.cdc.gov/mmwr/volumes/69/rr/pdfs/rr6905a1-H.pdf

^{*} Combined HepA and HepB vaccine (Twinrix) should not be used for postexposure prophylaxis.

^{*}The dosage of immune globulin is based on weight for all ages and does not have a maximum dose for protection against hepatitis A (Source: Grifols, Treating with GamaSTAN [immune globulin (human)] Los Angeles, CA: Grifols, 2019. https://www.hypermunes.com/en/hcp/gamastan-hepatitis-a).

Prevention of Hepatitis A

- Worldwide
 - Hygiene and sanitation
 - Handwashing and clean water sources
 - Prevent contamination of fresh produce
- Low endemic countries
 - Preexposure vaccination
 - All children aged ≥1 years
 - Risk groups (travelers, MSM, illegal drug users, persons with chronic liver disease)
 - Postexposure prophylaxis (vaccine or IG)
 - Household and sex contacts
 - Common source exposure (prepared food by infected handler)

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Recommendations for Hepatitis A Vaccine and Immune Globulin for Preexposure **Prophylaxis**

BOX. Prevention of hepatitis A virus infection in the United States: Recommendations of the Advisory Committee on Immunization

All children aged 12–23 months Unvaccinated children and adolescents aged 2–18 years

Persons at increased risk for HAV infection International travelers

Men who have sex with men

Persons who use injection or noninjection drugs (i.e., all

those who use illegal drugs) Persons with occupational risk for exposure

Persons who anticipate close personal contact with an international adoptee
Persons experiencing homelessness

Persons at increased risk for severe disease from HAV

Persons with chronic liver disease

Persons with human immunodeficiency virus infection

Other persons recommended for vaccination
Pregnant women at risk for HAV infection or severe
outcome from HAV infection
Any person who requests vaccination

Vaccination during outbreaks

Unvaccinated persons in outbreak settings who are at risk for HAV infection or at risk for severe disease from HAV

Implementation strategies for settings providing services to adults

Persons in settings that provide services to adults in which a high proportion of those persons have risk factors for HAV infection

Persons who receive blood products for clotting disorders (e.g., hemophilia)

*See the Recommendations for Hepatitis A Vaccine and Immune Globulin for Preexposure Prophylaxis section in this report for additional information.

ACIP 2020: https://www.cdc.gov/m mwr/volumes/69/rr/pdf s/rr6905a1-H.pdf

Hepatitis E Virus (HEV)

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Hepatitis E—Highlights

Like HAV

- Fecal-oral
- Acute self-limiting infection
 - Exception—may persist in immunologically compromised host
- Vaccine preventable

Unlike HAV

- Incubation period ~10 days longer
- Clinical response dose-dependent
 - Not age dependent
 - Infection rare in children and household contacts
- Higher mortality overall
 - Much higher in pregnant women
- Sporadic disease very rare in developed countries, Ab relatively common
 - Not always associated with travel*
 - Zoonotic source

Purcell RH, Emerson SU, J Hepatology 2008

* Autochthonous

First Recognized Outbreak of Hepatitis E New Delhi, India (1956)

- 35,000 cases acute jaundice
- City water system contaminated by sewage
- Highest attack rate persons 15-40 years old
- Case-fatality rate 10.5% among pregnant women
- Originally thought to be hepatitis A

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History

- 1983 human challenge experiment
 - Ingested pooled stool from outbreak-related cases
 - Developed acute non-A, non-B hepatitis
 - Excretion of virus-like particles
 - Stool infectious for non-human primates
- Classified enterically-transmitted NANB hepatitis
- 1990 HEV genome cloned
 - Classified as calicivirus



Diagnosis of Acute Hepatitis E

- Clinical illness indistinguishable from other types of acute viral hepatitis (A, B, C, D,....)
- Definitive diagnosis requires laboratory confirmation
- Serology:
 - IgM anti-HEV, IgG anti-HEV
 - Acute infection good sensitivity and specificity
 - Prevalent infection discordance between tests
- Research labs
 - HEV RNA by PCR (serum, stool, liver)
 - HEVAg by immunofluorescent probe (liver)

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Hepatitis E - Clinical Features

Incubation period Average 40 days; Range 15-60 days

Clinical illness Case/infection ratio and severity increase with

age

Chronic sequelae None ("chronic" viremia recently reported in

transplant patients)

Case-fatality rate Overall 1-3%

Pregnant women 15-20%

Factors related to increased severity Chronic liver disease, large inoculum, pregnancy

Hepatitis E in Pregnancy

- Reasons for poor outcomes are unclear
- Most severe in 3rd trimester
 - 1/2: asymptomatic or mild HEV infection
 - 1/2: acute HE
 - 1/3 have FHF (fulminant hepatic failure, in resource-poor settings: high mortality)
 - 2/3 preterm delivery
 - High rates of obstetric complications
- Vertical transmission is common with 3rd trimester
 - Rate 33%-100%
 - Clinical outcome in infants is highly variable
 - Asymptomatic infection to hepatic necrosis
 - Hypoglycemia and hypothermia associated with mortality

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Treatment

- Supportive
- No specific antiviral therapy
- Acute HE in pregnant women
 - No published data on potential benefits of early delivery
- No U.S. approved vaccine

Epidemiologic Features of HEV

- Responsible for sporadic cases of acute hepatitis and outbreaks
- Mode of transmission fecal-oral
 - Recent report of solid organ transplant-related
- Pattern
 - Outbreaks fecally contaminated drinking water
 - Sporadic cases not known
- Highest attack rate in young adults
- Minimal person-to-person transmission
- Animal reservoir

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Outbreaks or Confirmed Infection in >25% of Sporadic Non-ABC Hepatitis HEV may be most common etiology of acute viral hepatitis

"Endemic" Countries

- Outbreaks
 - Waterborne
 - Flooding, disruptions in water systems
- Sporadic Hepatitis E
 - HEV accounts for a variable proportion of acute viral hepatitis
 - Source of transmission not clear

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"Non-Endemic" Countries

- No outbreaks
- Sporadic HE
 - Travel-related
 - Most commonly after travel to Asia, especially India and China
 - Domestically acquired
 - Rare
 - Source unknown

Prevention of Hepatitis E

- Treatment None
- Vaccine (not approved in US) effective, high-risk populations
- Immune globulin Not effective
- Clean and reliable water supply
 - Virus probably inactivated by boiling
 - Effect of chlorination not known
- Proper sanitation (e.g., safe disposal of human and animal sewage)

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Bloodborne Viral Hepatitis

Hepatitis B Virus

Hepatitis C Virus

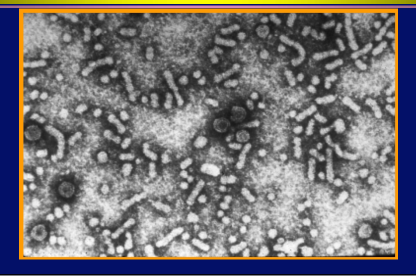
Hepatitis D Virus

Bloodborne Viral Hepatitis HBV and HCV—Highlights

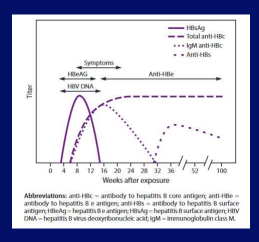
- HBV
 - Status of universal childhood immunization
- HBV and HCV
 - Transfusion Safety
 - Injection Safety
 - Changing epidemiology
 - Chronic disease burden

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Hepatitis B Virus



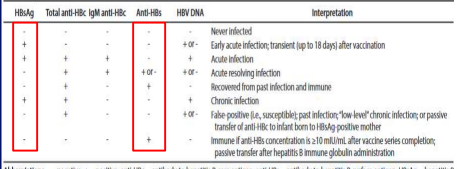
Acute hepatitis B virus infection with recovery



ACIP, 2018: https://www.cdc.gov/mmwr/volumes/67/rr/pdfs/rr6701-H.pdf

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Typical interpretation of test results for hepatitis B virus infection



Abbreviations: -= negative; += positive; anti-HBc = antibody to hepatitis B core antigen; anti-HBs = antibody to hepatitis B surface antigen; HBsAg = hepatitis

ACIP, 2018: https://www.cdc.gov/mmwr/volumes/67/rr/pdfs/rr6701-H.pdf

Ten Leading Causes of Infectious Disease Deaths Worldwide (2000)

Disease Deaths per Year Lower resp tract infections ~3.5 million **HIV/AIDS** ~3.0 million Diarrheal diseases ~2.2 million **Tuberculosis** ~2.0 million Malaria ~1-3 million Measles ~888,000 Hepatitis B ~750.000 **Pertussis** ~355,000 Neonatal tetanus ~300,000 Hepatitis C ~ 250,000

Source: CDC, WHO, UNICEF, UNAIDS

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Global and US Disease Burden from Bloodborne Viral Infections

Estimated No. Chronic Infections Global US 370 million 1.25 million **HBV** 130 million 3-4 million **HCV** HIV 40 million 1 million (3–4 million) HIV / HBV (250,000)(4–5 million) (40-50,000)HIV / HCV Sources: WHO and CDC, unpublished data.

Features of HBV & HCV Infection				
	<u>HBV</u>	<u>HCV</u>		
Virus Classification	DNA	RNA		
	Hepadnavirus	Flavivirus		
Incubation period – average	8–12 wks	6–7 wks		
– range	6–26 wks	2–26 wks		
Specific serologic markers				
acute infection	Yes	No		
active infection	Yes	No		
chronic infection	Yes	No		
Clinical illness (jaundice)	30%–50%	20%		
Chronic infection	90% (infants) 5–10% (adults)	~70%		
Mortality from CLD, cirrhosis, HCC	25%	1-5%		

Relative Efficiency of Transmission by Type of Exposure

Type of exposure	Efficienc	y of transr	nission
to infected source	<u>HBV</u>	<u>HCV</u>	<u>HIV</u>
Transfusion	++++	++++	++++
Injecting drug use	++++	++++	++++
Unsafe injections	+++	+++	++
Needlestick	+++	+	<+
Sexual	+++	+	+++
Perinatal	++++	++	+++
Non-intact skin	++	+/-	+/-

Relative Infectivity of HBV, HCV, HIV

<u>HBV</u>	<u>HCV</u>	HIV
108-9	10 ⁵	10 ³
++++	++	-
<u>></u> 7d	≥16h (<4d)	0
	10 ⁸⁻⁹	10 ⁸⁻⁹ 10 ⁵ ++++ ++ ≥7d ≥16h

Bond WW, Lancet 1981;1:550-51; Kamili S, Infect Control Hosp Epi 2006.

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Environmental Stability of HBV and HCV Facilitates Their Transmission

- More rapid acquisition among IDUs
 - Clean needles and syringes alone insufficient to interrupt transmission because virus can live on contaminated drug preparation equipment (i.e., cookers and cotton)
- latrogenic transmission
 - Virus can live in contaminated multi-dose vials and on needles and syringes

Routes of HBV Transmission

Age Group Routes of Infection

Newborn Mother to infant (perinatal)

Childhood Household (non-intact skin)

Adolescent/Adult Sexual contact

Injecting drug use equipment

Occupational exposures

All ages Unsafe injections

Transfusion from unscreened donors

Other health care related

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Global Differences in HBV Transmission Patterns

Chronic

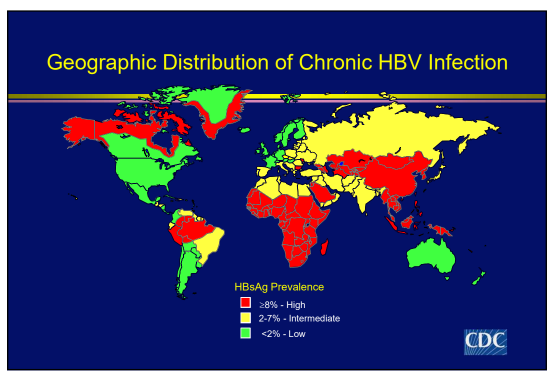
infection Primary Age Primary Modes (% immune) <u>at Infection</u> <u>of Transmission</u>

High ≥8% Infants Perinatal, horizontal, unsafe (≥60%) Young children injections, unscreened blood

Intermediate 2-7% All age groups Perinatal, horizontal, unsafe injections, sexual, IDU

(20-60%)

Low <2% Adolescents Sexual, IDU Adults



Prevalence of chronic hepatitis B virus infection, by country

High (28% prevalence): Angola, Benin, Burkina Faso, Burundi, Cameroon, Central African Republic, Congo, Côte d'Ivoire, Djibouti, Equatorial Guinea, Gabon, Gambia, Ghana, Guinea, Haiti, Kiribati, Kyrgyxstan, Laos, Liberia, Malawi, Mali, Mauritania, Mongolia, Mozambique, Namibia, Nauru, Niger, Nigeria, Niue, Papua New Guinea, Senegal, Sierra Leone, Solomon Islands, Somalia, South Sudan, Sudan, Swaziland, Togo, Tonga, Uganda, Vanuatu, Vietnam, Yemen, and Zimbabwe.

Intermediate (5%–7.9% prevalence): Albania, Bhutan, Cape Verde, China, Democratic Republic of the Congo, Ethiopia, Kazakhstan, Kenya, Marshall Islands, Moldova, Oman, Romania, Rwanda, Samoa, South Africa, Tajikistan, Tanzania, Thailand, Tunisia, Tuvalu, Uzbekistan, and Zambia.

Low Intermediate (2%–4.9% prevalence): Algeria, Azerbaijan, Bangladesh, Belarus, Belize, Brunei Darussalam, Bulgaria, Cambodia, Colombia, Cyprus, Dominican Republic, Ecuador, Eritrea, Federated States of Micronesia, Fiji, Georgia, Italy, Jamaica, Kosovo, Libya, Madagascar, Myanmar, New Zealand, Pakistan, Palau, Philippines, Peru,

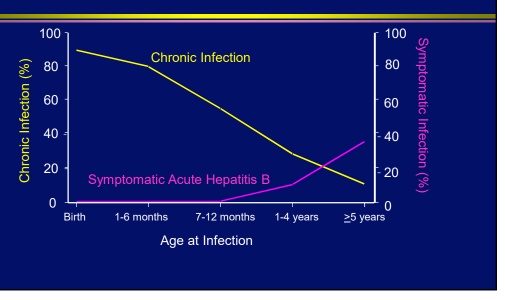
Russia, Saudi Arabia, Singapore, South Korea, Sri Lanka, Suriname, Syria, Tahiti, and Turkey.

Low (s1.9% prevalence): Afghanistan, Argentina, Australia, Australia, Australia, Alastria, Bahrain, Barbados, Belgium, Bolivia, Bosnia and Herzegovina, Brazil, Canada, Chile, Costa Rica, Croatia, Cuba, Czech Republic, Denmark, Egypt, France, Germany, Greece, Guatemala, Hungary, Iceland, India, Indonesia, Iran, Iraq, Ireland, Israel, Japan, Jordan, Kuwait, Lebanon, Lithuania, Malaysia, Mexico, Morocco, Nepal, Netherlands, Nicaragua, Norway, Palestine, Panama, Poland, Portugal, Qatar, Serbia, Seychelles, Slovakia, Slovenia, Spain, Sweden, Switzerland, Ukraine, UK, Unitted Arab Emirates, United States of America, and Venezuela.

No data: Andorra, Antigua and Barbuda, Armenia, The Bahamas, Botswana, Chad, Comoros, Cook Islands, Dominica, El Salvador, Finland, Grenada, Guinea Bissau, Guyana, Honduras, Latvia, Lesotho, Lithuania, Luxembourg, Macedonia, Maldives, Malta, Mauritius, Monaco, Montenegro, North Korea, Paraguay, Saint Kitts and Nevis, Saint Lucia, Saint Vincent and the Grenadines, San Marino, Sao Tome and Principe, Timor-Leste, Trinidad and Tobago, Turkmenistan, and Uruguay.

^{*}Source: CDC. Travelers health: infectious diseases related to travel. Atlanta, GA: US Department of Health and Human Services, CDC; 2017.

Outcome of HBV Infection by Age at Infection



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Global Strategy to Prevent HBV Transmission

- Hepatitis B Vaccination
 - Routine infant vaccination (all countries)
 - Catch-up vaccination of older children/adolescents
 - Vaccination of high-risk groups
- Prevention of latrogenic transmission
 - Routine screening of transfused blood
 - developed countries -100% screen
 - least developed countries 35% screen (?)
 - Safe injection practices
 - Proper infection control practices





* Other - Household contact, occupational exposure, hemodialysis, institutionalization, transfusion

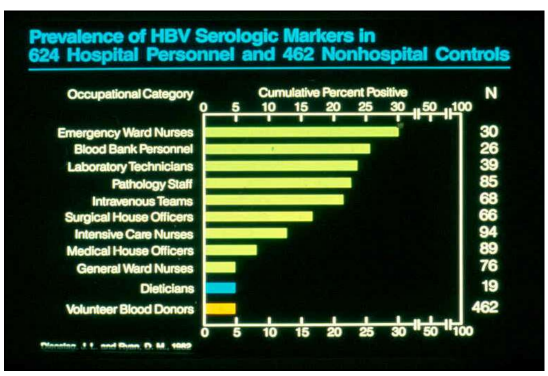
Source: Adapted from Sentinel Counties and NNDSS, CDC

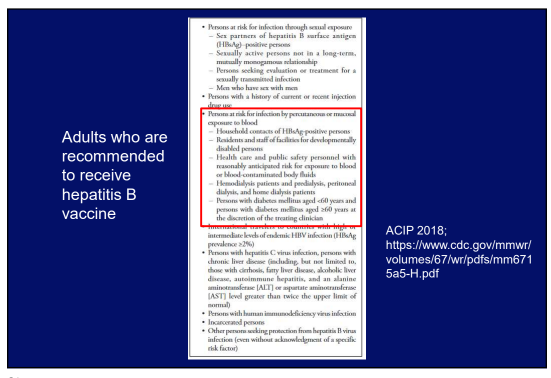
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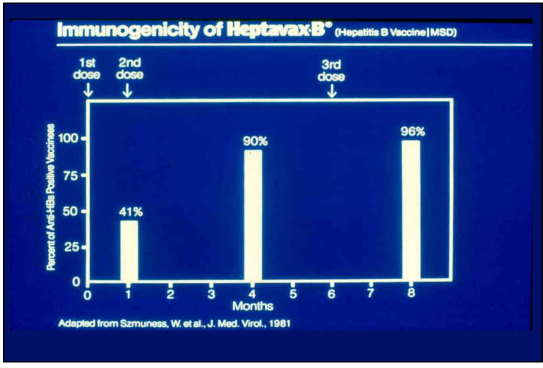
Recent HBV Outbreaks Associated with Blood Glucose Monitoring

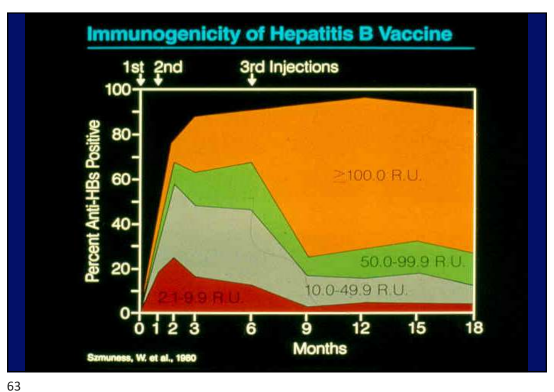
- Shared fingerstick devices
 - 1999 VA Assisted Living Centers 4 cases
 - 1999 CA Nursing Home 4 cases
 - 2003 MS Nursing Home 15 cases
 - 2003 CA Assisted Living Center 8 cases
 - 2005 VA Assisted Living Centers (2) 11 cases
- Dedicated fingerstick devices
 - 1999 CA Skilled Nursing Facility 5 cases
 - 2002 CA Subacute Hospital 3 cases
 - 2003 NC Nursing Home 11 cases

Hepatitis B in Health-Care Personnel









Recommended doses of hepatitis B vaccine, by group and vaccine type

Age group	Schedule* (Interval represents tim In months from first dose)	
Children (1-10 yrs)	0, 1, and 6 mos 0, 1, 2, and 12 mos	
Adolescents (11–19 yrs)	0, 1, and 6 mos 0, 12, and 24 mos 0 and 4–6 mos [†] 0, 1, 2, and 12 mos 0, 7 days, 21–30 days, 12 mos [§]	
Adults (≥20 yrs)	0, 1, and 6 mos 0, 1, 2, and 12 mos 0, 1, 2, and 6 mos [¶] 0, 7 days, 21–30 days, 12 mos [§]	

ACIP 2018: https://www.cdc.gov/mmwr/volumes/67/rr/pdfs/rr6701-H.pdf

Hepatitis B Vaccine

Factor	Response	Factor	Response
Age 20-29	95%	Diabetes	70-80%
Age 30-39	90%	Liver disease	60-70%
Age 40-49	86%	Gender	Female>male
Age 50-59	71%	Obesity	Decreased
Age <u>≥</u> 60	47%	Smokers	Decreased
Renal Failure	50-80%	Gluteal injection	Decreased
HIV infection	50-70%		
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Recommended doses of hepatitis B vaccine, by group and vaccine type

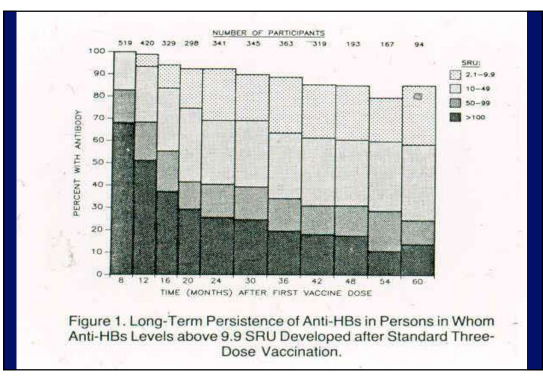
	Single-antigen vaccine				Combination vaccine			
	Recom	nbivax	Eng	erix	Pedi	arix*	Twir	nrix [†]
Age group (yrs)	Dose (μg)	Vol (mL)	Dose (μg)	Vol (mL)	Dose (μg)	Vol (mL)	Dose (μg)	Vol (mL)
Birth-10	5	0.5	10	0.5	10*	0.5	N/A	N/A
11-15	10 ⁵	1	N/A	N/A	N/A	N/A	N/A	N/A
11-19	5	0.5	10	0.5	N/A	N/A	N/A	N/A
≥20	10	1	20	1	N/A	N/A	20†	1
Hemodialysis patients	and other immune-c	compromised po	ersons					
<20	5	0.5	10	0.5	N/A	N/A	N/A	N/A
≥20	40	1	40	2	N/A	N/A	N/A	N/A

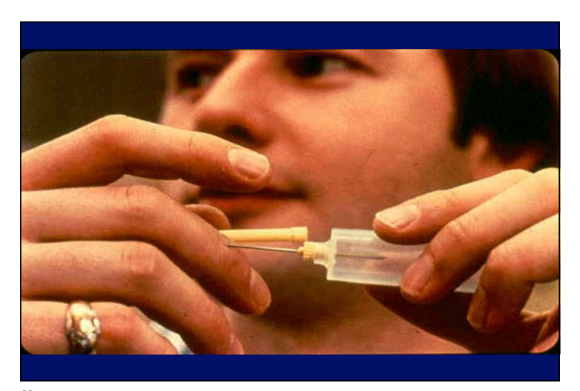
obreviation: N/A = not applicative. Pediarix is approved for use in persons aged 6 weeks through 6 years (prior to the 7th birthday). Twinrix is approved for use in persons aged ≥18 years. Adult formulation administered on a 2-dose schedule.

ACIP 2018: https://www.cdc.gov/mmwr/volumes/67/rr/pdfs/rr6701-H.pdf

Hepatitis B Vaccine: Administration 2

- Schedule
 - 0, 1, 6 mo
 - 0, 1, 2, 12 mo (more rapid antibody rise) (Engerix)
- Pre-exposure
 - ACP: Three doses, obtain titer (1-6 mo). If antibody negative, provide up to 3 additional doses (titer 1-2 mo after each dose)
 - If no response after 6 doses, provide HBIG for exposures





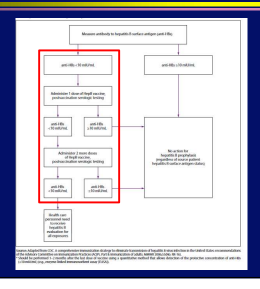
Postexposure management of health care personnel after occupational percutaneous or mucosal exposure to blood or body fluids, by health care personnel HepB vaccination and response status

	Postexpo	sure testing	Postexposur			
HCP status	Source patient (HBsAg)	HCP testing (anti-HBs)	HBIG	Vaccination	Postvaccination serologic testing	
Documented responder after complete series			No action needed			
Documented nonresponder after two complete series	Positive/unknown	_*	HBIG x2 separated by 1 month	77.	N/A	
55 (c. 5) (c. 4) No. (c. 5) (c	Negative		No actio			
Response unknown after complete series	Positive/unknown	<10 mIU/mL	HBIG x1	Initiate revaccination	Yes	
100 45 \$ 100 00 \$ 100 \$ 100 00 00 00 00 00 00 00 00 00 00 00 00	Negative	<10 mIU/mL	None	Initiate revaccination	Yes	
	Any result	lt ≥10 mIU/mL No action needed		n needed		
Unvaccinated/incompletely vaccinated or	Positive/unknown	_	HBIG x1	Complete vaccination	Yes	
vaccine refusers	Negative		None	Complete vaccination	Yes	

Abbreviations: anti HBs = antibody to hepatitis B surface antigen; HBIG = hepatitis B immune globulin; HBsAg = hepatitis B surface antigen; HCP = health care personnel; N/A = not applicable.
*Not indicated.

ACIP, 2018: https://www.cdc.gov/mmwr/volumes/67/rr/pdfs/rr6701-H.pdf

Pre-exposure evaluation for health care personnel previously vaccinated with complete, ≥3-dose HepB vaccine series who have not had postvaccination serologic testing*



https://www.cdc.gov/mmwr/volu mes/67/rr/pdfs/rr6701-H.pdf

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Serology Testing and Vaccinations

BOX 6. Persons recommended to receive serologic testing prior to vaccination⁴ Household, sexual, or neelle contacts of hepatitis B surface antigen (HBsAp.) positive penons³ HIV-positive perons³ Persons with elevated alanine aminotransferase/ aspartate aminotransferase of unknown etiology³ Hemodialysis patients³ Penons born in countries of unknown etiology³ Persons born in countries of high and intermediate hepatitis B virus (HBV) endemicity (HBsAg prevalence 22%) U.S.-born persons not vaccinated as infants whose parents were born in countries with high HBV endemicity (28%) U.S.-born persons not vaccinated as infants whose parents were born in countries with high HBV endemicity (28%) Persons needing immunosuppressive therapy, including chemotherapy, immunosuppressive to organ transplantation, and immunosuppression for the unautologic of gastroenterologic disorders Donors of blood, plasma, organs, tissues, or semen

- Infants born to hepatitis B surface antigen (HBsAg)— positive mothers or mothers whose HBsAg starus remains unknown (e.g., when a parent or person with lawful custody safely surrenders an infant confidentially deathy after birth infants safely memodered as or
- shortly after birth)*

 Health care personnel and public safety workers

 Hemodialysis patients and others who might require ourportions boundinlysis (e.g., predialysis, personnel)
- output beneathate for purificate potential dialysis, and home dialysis.

 HIV-infected persons

 Other immunocompromised persons (e.g., hematopoietic stem-cell transplant recipients or persons receiving chemotherapy)

 Sex partners of HBsAg-positive persons

ACIP 2018: https://www.cdc.gov/mmwr/volumes/67/rr/pdfs/rr6701-H.pdf

HBV: POST-EXPOSURE PROPHYLAXIS

Exposed person	Source HBsAg+	Source HBsAg-	Source unknown
Unvaccinated	HBIG x 1 HBV vaccine	HBV vaccine	HBV vaccine
Vaccinated, Responder	No therapy	No therapy	No therapy
Vaccinated, Nonresponder	HBIG x 2 or HBIG x 1 & HBV vaccine	No therapy	If known high-risk source, treat as if source HBsAg+
Vaccinated, Response unknown	Obtain anti-HBs * If ok, no therapy * If low, HBIG x 1 & vaccine	No therapy	Obtain anti-HBs * If OK, no therapy * If low, vaccine booster

Adequate anti-HBs is >10 mIU/mL; HBIG = 0.06 mg/kg IM

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Hepatitis C Virus

- RNA Flavivirus (Hepacivirus)
 - Discovery using recombinant DNA technology reported in 1989
 - Clinical entity (non-A, non-B hepatitis) in transfused patients reported late 1960s
 - Target organ liver
- Bloodborne (primarily) and sexually-transmitted
- No vaccine
 - Mutations occur during viral replication
 - Substantial heterogeneity (quasi species) prevents effective neutralization
- Treatable and curable (most people free of virus in months)

Features of Hepatitis C Virus Infection

Incubation period Average 6-7 weeks

Range 2-26 weeks

Acute illness (jaundice) Mild (20%-30%)

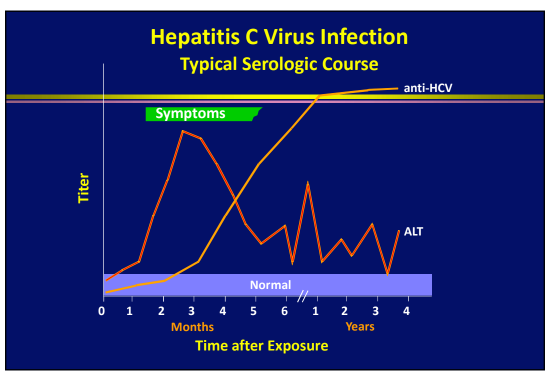
Case fatality rate Low

Chronic infection 75%-85%

Chronic hepatitis 70%

Mortality from CLD 1%-5%

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Hepatitis C Virus Infection United States

New infections per year 1985-89 242,000

2006 20,000

Deaths from acute liver failure Rare

Persons ever infected (1.6%) 4.1 million (3.4-4.9)*

Persons with chronic infection 3.1 million (2.5-3.7)*

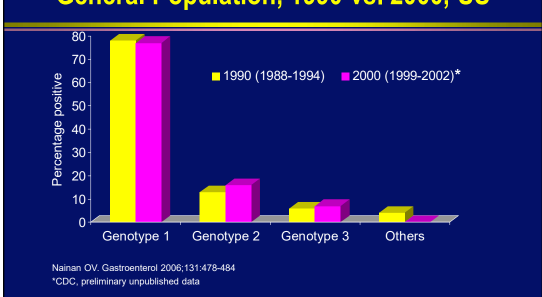
HCV-related chronic liver disease 40% - 60%

Deaths from chronic disease/year 8,000-10,000

* 95% confidence interval (data from 1999-2002)

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Distribution of HCV Genotypes in the General Population, 1990 vs. 2000, US

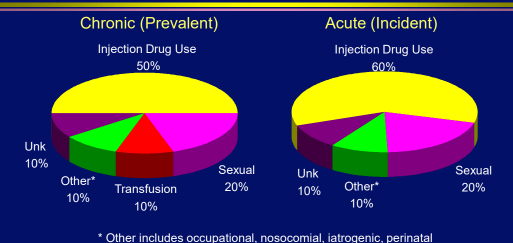


Transmission of HCV

- Percutaneous
 - Injecting drug use
 - Clotting factors before viral inactivation
 - Transfusion, transplant from infected donor
 - Therapeutic (contaminated equipment, unsafe injection practices)
 - Occupational (needlestick)
- Permucosal
 - Perinatal
 - Sexual

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Risk Factors For Persons with Acute or Chronic Hepatitis C 1999-2002, U.S.



latrogenic-Related Outbreaks of HCV Infections in **Developed Countries**

- In- and outpatient care
 - Chronic hemodialysis, surgery, endoscopy, pain management clinic, oncology clinic, in-patient ward
- Most due to unsafe injection practices, i.e., failure to use aseptic techniques
 - Contamination of multi-dose medication vials and IV solutions
 - Reuse of syringes/needles
 - Contamination of finger stick devices
- Infected HCW rarely source
 - Usually due to self-injection of patients' narcotics

Williams IT, Clin Infect Dis 2004;38:1592-1598

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Health-Care Related HCV Transmission

- Blood transfusion from unscreened donors
 - including plasma-derived products not inactivated
- Unsafe injection practices
 - inadequate sterilization of reusable needles and syringes
 - sharing of disposable needles and syringes
- Contaminated equipment
 - inadequate cleaning and disinfection
 health care settings

 - alternative medicine practices, rituals

Global Burden of Disease Associated with Unsafe Injections

- Estimated annual incidence, 2000
 - > 20 million HBV infections
 - 30% of new infections
 - > 2 million HCV infections
 - 40% of new infections
 - > 250,000 HIV infections
 - 5% of new infections

Ezzati M et al. Lancet. 360(9343):1347-60, 2002.

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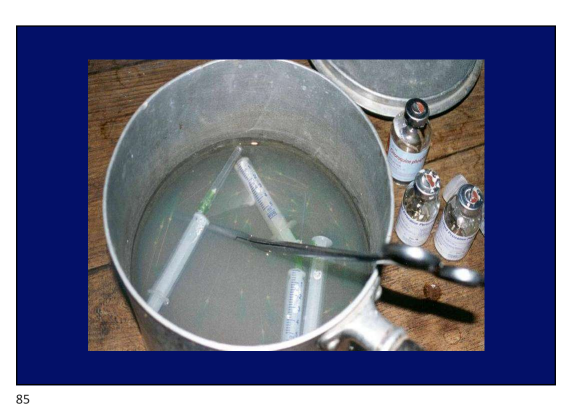
Unsafe Injection Practices

Developing Countries

- Inadequate supplies of sterile syringes
- Inadequate sterilization of reusable syringes and needles
- Administration at home by nonprofessionals
- Syringes shared with others (family, neighbors)
- Overuse of therapeutic injections

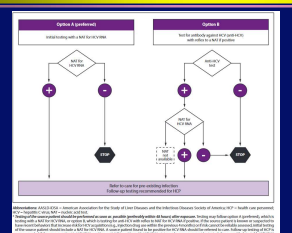
Developed Countries

- Failure to use aseptic techniques
 - Reuse of same syringe and needle to administer meds to multiple patients
 - Medication preparation and blood sample handling in same area
 - Contamination of multiple dose medication vials
- Infected HCW rarely source
 - Usually due to self-injection of patients' narcotics



Children Handling Medical Waste, Bangladesh

Testing of source patients after potential exposure of health care personnel to hepatitis C virus — CDC guidance, United States, 2020*



ACIP 2020: https://www.cdc.gov/mmwr/v olumes/69/rr/pdfs/rr6906a1-H.pdf

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Testing of source patients after potential exposure of health care personnel to hepatitis C virus — CDC guidance, United States, 2020*

BOX. Testing of source patients and health care personnel potentially exposed to hepatitis C virus — CDC guidance, United States, 2020

Source-patient testing

• Tiesting of the source patient may follow option A (preferred), which is resting with a nucleic acid test (PAT) for hepatitis C virus (HCV) RNA, or option B, which is testing for anti-HCV with reflex to a NAT if positive.

• If a source patient is known or suspected to have recent behaviors that increase risk for HCV acquisition (e.g., injection drug use within the previous 4 months) or if risk cannot be reliably assessed, initial testing should include a NAT.

• Follow-up testing of health care personnel (HCP) is recommended if the source patient is HCV RNA positive, anti-HCV positive with RNA status unknown, or cannot be tested.

HCP testing

• Baseline testing of HCP for anti-HCV with reflex to a NAT if positive should be conducted as soon as possible (preferably within 48 hours) after the exposure and may be simultaneous with source-patient testing.

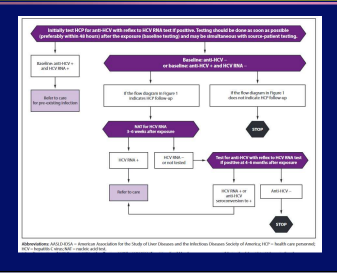
• If follow-up testing of HCP is recommended based on the source-patient status, test with a NAT at 3-6 weeks postexposure.

• If the HCP is NAT negative at 3-6 weeks postexposure.

• If the HCP is NAT negative at 3-6 weeks postexposure.

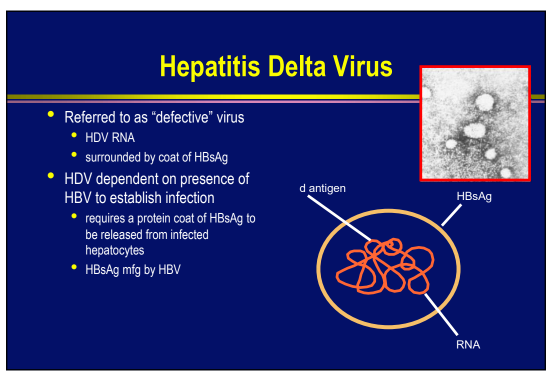
ACIP 2020: https://www.cdc.gov/mmwr/v olumes/69/rr/pdfs/rr6906a1-H.pdf

Testing of source patients after potential exposure of health care personnel to hepatitis C virus — CDC guidance, United States, 2020*



ACIP 2020: https://www.cdc.gov/mmwr/v olumes/69/rr/pdfs/rr6906a1-H.pdf

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Infection with HDV

HBV-HDV Coinfection

- Simultaneous infection with HBV and HDV in a person susceptible to HBV
- Low risk of death from chronic liver disease

HBV-HDV Superinfection

- Infection with HDV in a person with chronic HBV
- High risk of severe chronic liver disease and death

Persons immune from HBV infection – either through vaccination or resolved infection – cannot become infected with HDV

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Epidemiologic Features of HDV

- Transmission similar to HBV
 - Percutaneous highly efficient
 - SexualPerinatalMuch less efficient than HBV
- Uncommon in U.S. seen mainly in IDU's
- Worldwide, endemic in Amazon,
 Mediterranean, Central Asia, Africa

Prevention of HDV

HBV-HDV coinfection

Hepatitis B vaccination

HBV-HDV superinfection

- Prevent exposure to HBV
 - screen blood for HBV (need HBV for HDV)
- Reduce high-risk behaviors
 - safer sex practices
 - safer injection practices

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Current and Future Issues

- Identification of infected persons
 - Screening and testing not routinely performed
 - Lack effective methods for reaching those whose risk was in the remote past
 - Risk factor ascertainment in routine healthcare visits is rare
- Therapy regimens less than ideal, especially those with genotype 1
 - In US, treatment offered to low % of HCV-positives
- Implications of multiple co-factors on liver disease progression and response to therapies not well understood
 - Impact likely to grow creating an even greater challenge
- Need to be alert to changes in epidemiology

Viral Hepatitis - Overview

- Primary infection of the liver caused by at least five unrelated viruses: A, B, C, D, E
- HAV and HEV
 - Fecal-oral route
 - Acute self-limited disease; no chronic infection
- HBV, HCV, HDV
 - Percutaneous or mucosal exposures to blood
 - Chronic infection major causes of cirrhosis and hepatocellular carcinoma worldwide

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Viral Hepatitis in US: Trends

HHS.gov

- ~2M living with HCV in US
- ~1M Living with HBV in US
- More than half of persons with hepatitis do not know they have the virus
 - 67% of persons with HBV
 - 51% of persons with HCV
- Baby boomers (mid-50s to early 70s) made up 36.3% chronic HCV
- HCV increasing in the US
- · Viral hepatitis is leading cause of liver cancer

Acknowledgment

 Most slides provided by Miriam J. Alter, Ph.D. University of Texas, Galveston

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Geographic Differences in HCV Transmission Patterns

Exposures among	Contribution of exposures to disease burden by HCV prevalence			
prevalent infections	Low	<u>Moderate</u>	<u>High</u>	
Injecting drug use	++++	++	+	
Transfusions (before testing	ng) +++	+++	+++	
Unsafe therapeutic injection	ons +	++++	++++	
Occupational	+	+	+	
Perinatal	+	+	+	
High-risk sex	++	+	+/-	