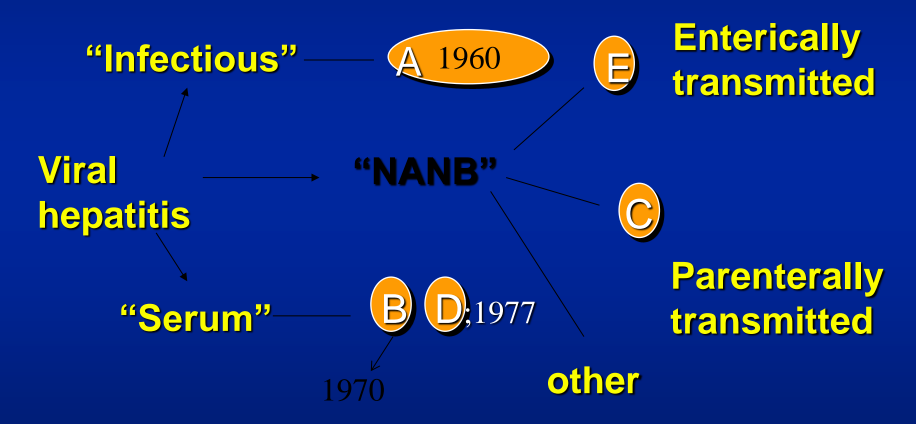
VIRAL HEPATITIS

Kolářová M., Autumn 2017



VIRAL HEPATITIS HISTORICAL PERSPECTIVE

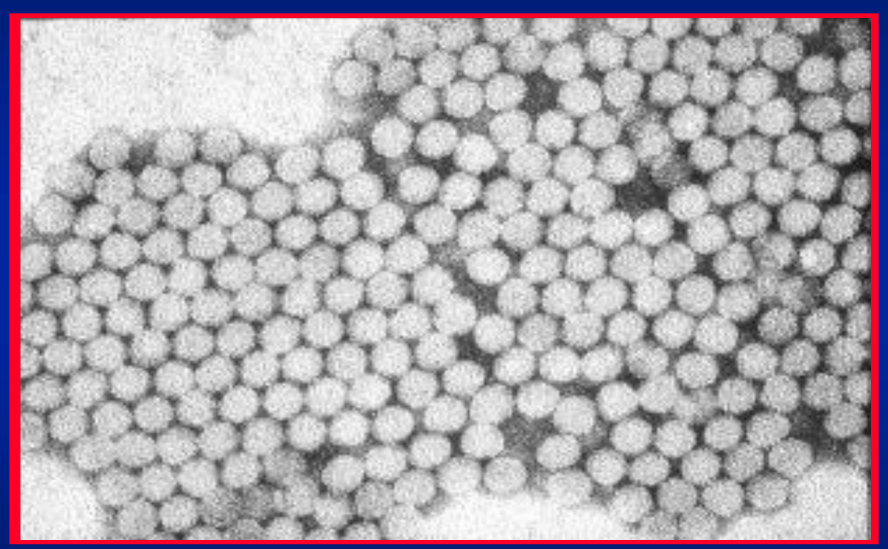




VIRAL HEPATITIS A



HEPATITIS A VIRUS





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



HEPATITIS A - CLINICAL FEATURES

Jaundice by <6 yrs <10%

age group: 6-14 yrs 40%-50%

>14 yrs 70%-80%

Rare complications: Fulminant hepatitis

Cholestatic hepatitis

Relapsing hepatitis

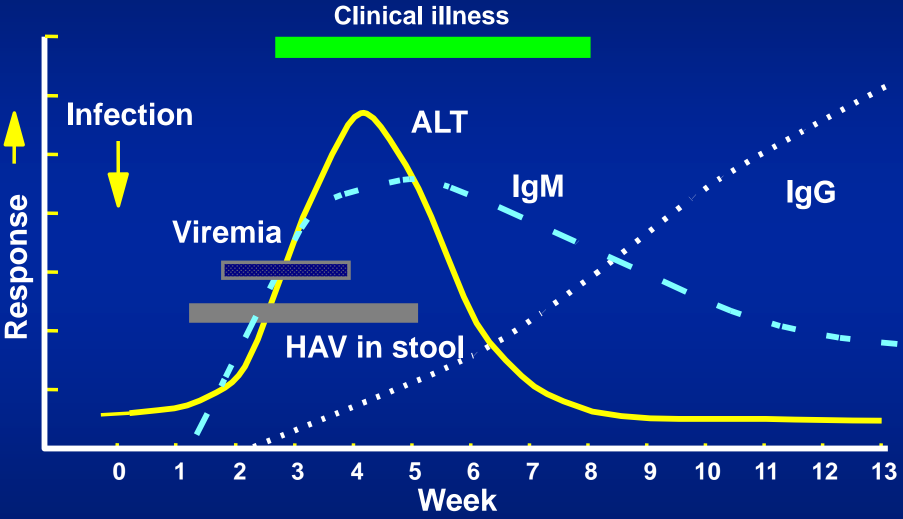
Incubation period: Average 30 days

Range 15-50 days

Chronic sequelae: None



EVENTS IN HEPATITIS A VIRUS INFECTION





CONCENTRATION OF HEPATITIS A VIRUS IN VARIOUS BODY FLUIDS



Infectious Doses per mL

Source: Viral Hepatitis and Liver Disease 1984;9-22

J Infect Dis 1989;160:887-890



Is estimated that in 2013 HAV caused 14 900 deaths.

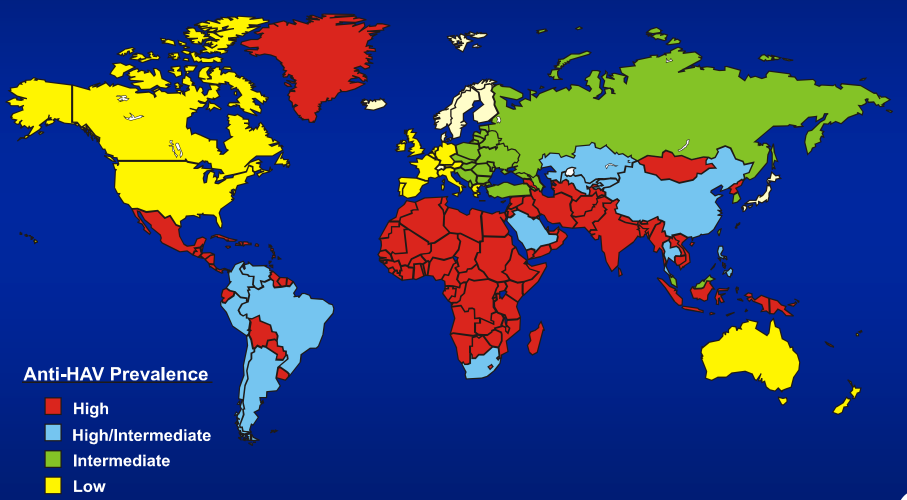
The severity of clinical disease associated with HAV infection increases with increasing age; jaundice occurs among less than:

- □ 10% of children younger than 6 years of age,
- □ 40%-50% of older children, and
- □ 70%-80% of adults.

Complications of hepatitis A include fulminant hepatitis, in which the case fatality rate can be greater than 50% despite medical interventions such as liver transplantation; cholestatic hepatitis, with very high bilirubin levels that can persist for months; and relapsing hepatitis, in which exacerbations can occur weeks to months after apparent recovery. Chronic infection does not occur following HAV infection.



GEOGRAPHIC DISTRIBUTION OF HEPATITIS A VIRUS INFECTION



Very Low



ACUTE HEPATITIS A CASE DEFINITION FOR SURVEILLANCE

Clinical criteria

- Any person with a discrete onset of symptoms (e.g. fatigue, abdominal pain, loss of appetite, intermittent nausea and vomiting)
- AND at least one of the following three:
- Fever;
- Jaundice;
- Elevated serum aminotransferase levels.

Laboratory criteria

- At least one of the following three:
- Detection of hepatitis A virus nucleic acid in serum or stool;
- Hepatitis A virus specific antibody response;
- Detection of hepatitis A virus antigen in stool.

Epidemiological criteria

- At least one of the following four:
- Human-to-human transmission;
- Exposure to a common source;
- Exposure to contaminated food/drinking water;
- Environmental exposure.

Case classification

- A. Probable case: Any person meeting the clinical criteria with an epidemiological link
- B. Confirmed case: Any person meeting the clinical and the laboratory criteria

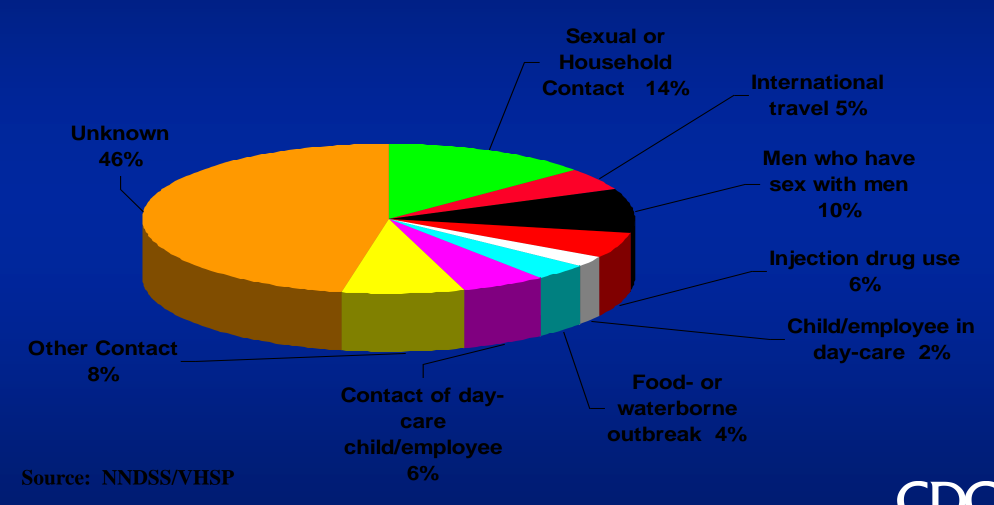


HEPATITIS A VIRUS TRANSMISSION

- Close personal contact
 (e.g., household contact, sex contact, child day-care centers)
- Contaminated food, water
 (e.g., infected food handlers)
- Blood exposure (rare)
 (e.g., injection drug use, rarely by transfusion)



RISK FACTORS ASSOCIATED WITH REPORTED HEPATITIS A, 1990-2000, UNITED STATES



PREVENTING HEPATITIS A

- Hygiene (e.g., hand washing)
- Sanitation (e.g., clean water sources)
- Hepatitis A vaccine (pre-exposure)
- Immune globulin (pre- and post-exposure)



PREPARATION OF INACTIVATED HEPATITIS A VACCINES

 Cell culture adapted virus grown in human fibroblasts

Purified product inactivated with formalin

Adsorbed to aluminum hydroxide adjuvant



HEPATITIS A VACCINES

- Highly immunogenic
 - 97%-100% of children, adolescents, and adults have protective levels of antibody within 1 month of receiving first dose; essentially 100% have protective levels after second dose
- Highly efficacious
 - In published studies, 94%-100% of children protected against clinical hepatitis A after equivalent of one dose



DURATION OF PROTECTION AFTER HEPATITIS A VACCINATION

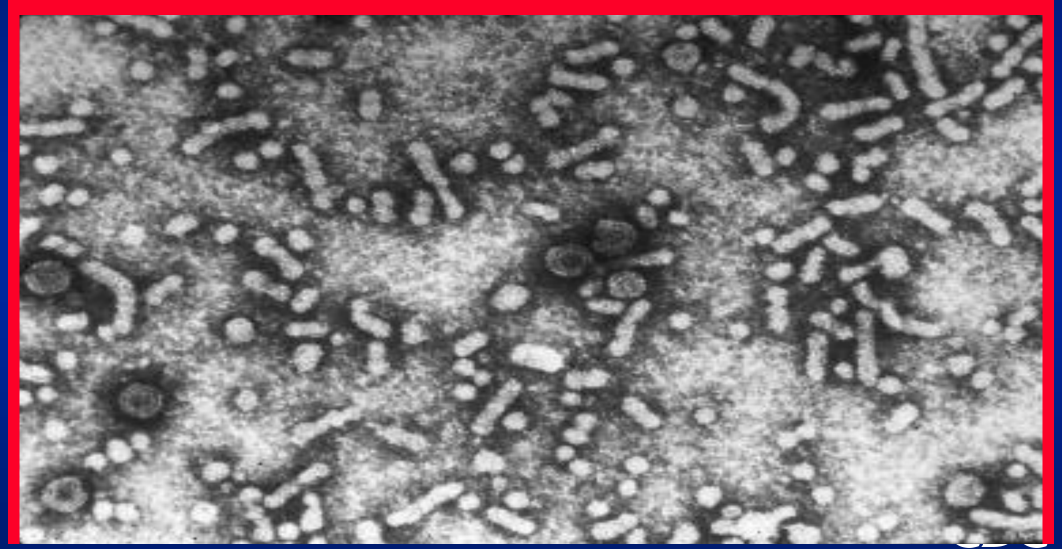
- Persistence of antibody
 - At least 5-8 years among adults and children
- Efficacy
 - No cases in vaccinated children at 5-6 years of follow-up
- Mathematical models of antibody decline suggest protective antibody levels persist for at least 20 years
- Other mechanisms, such as cellular memory, may contribute



VIRAL HEPATITIS B



The hepatitis_ B virus is a DNA virus belonging to the Hepadnaviridae family of viruses.



VIRAL HEPATITIS TYPE B

Etiology:

Hepatitis B virus, HBV, Hepadnavirus, the so-called Dane particle with a core (formed by DNA, DNA polymerase, and a nucleocapsid protein with the hepatitis B core antigen (HBcAg) and a coat of hepatitis B surface antigen (HBsAg)). The whole virus is infectious with a diameter of 42 nm.

The source of infection

Two months in the ende of incubation period, the sick or carriers.

Parenteral transmission - blood, blood products and inoculation of the infectious material are of principal significance in the transmission.

Route of transmission

Professional risk to medical personnel (injury by needle - transmission in 7 - 30 %, contaminated instruments, blood transfusions - transmission in 90 %).

i.v. drug addicts - injury during tattooing, possibly other minute injuries of the skin and mucosa.

By **sexual intercourse** in homosexuals, bisexuals, and prostitutes.

Vertical - perinatal transmission from mother to child when the mother is the virus carrier or the sick person. About 95 % of newborns infect intranatally and 5 % intrauterinely.

Susceptibility

General

Preventive measures:





VIRAL HEPATITIS TYPE B

Preventive measures:

Health education - to emphasize the extent of risk Observance of epidemic measures in medical establishments. Handling biological material and contaminated instruments, consistent disinfection and sterilization. application of single-use needles and syringes, use of closed hemodialysis systems, smoking and drinking in workplaces with biological material is forbidden. Postexposure prophylaxis - passive and active immunization (newborns). Examination of blood-donors - exclusion of HBsAg carriers from blood donation Designation and inspection of sanitary-epidemic measures in non-medical establishments (hair-dressing salons, barber shops, etc.) Active immunization in persons with a high risk of infection (stated by public notice)

.

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The disease occurs worldwide.

In 2013, HBV caused:

- *686 000 deaths, including 68 600 deaths from fulminant hepatitis,
- 300 000 deaths from hepatocellular carcinoma
- 317 400 deaths from cirrhosis
- with a very high burden among an estimated 280 million carriers (prevalence 3,7 %).



The symptoms can vary greatly and many of those infected with HBV never develop any symptoms at all.

Those who do get symptoms (30-50% of cases) usually suffer from tiredness, loss of appetite, abdominal discomfort, nausea, vomiting and fever. The vast majority of healthy adults who get acute hepatitis B will recover with no liver damage in 4–12 weeks but the death rate can reach 2% in the elderly.

Chronic infection is most likely to develop in young babies.

Hepatitis B – Clinical Features

* Incubation period:

Average 60-90 days

Range 45-180 days

* Clinical illness (jaundice):

<5 yrs, <10%

>5 yrs, 30%-50%

* Acute case-fatality rate:

0.5%-1%

* Chronic infection:

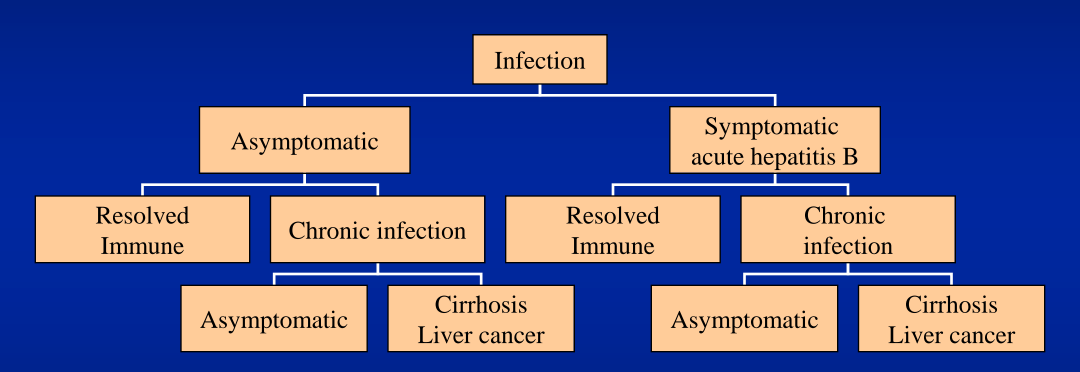
<5 yrs, 30%-90%

>5 yrs, 2%-10%

* Premature mortality from chronic liver disease:



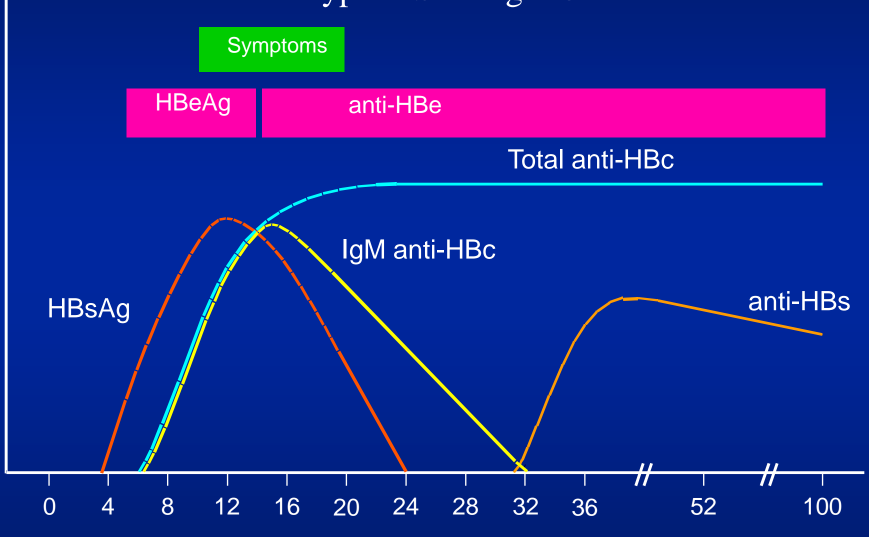
Outcome of HBV Infection





Acute Hepatitis B Virus Infection with Recovery

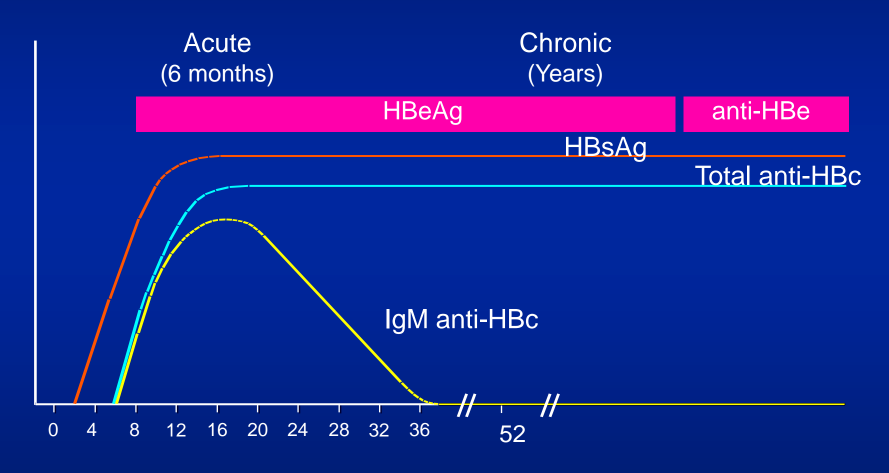




Weeks after Exposure



Progression to Chronic Hepatitis B Virus Infection Typical Serologic Course





HBV Modes of Transmission

- Sexual
- Parenteral
- Perinatal







Concentration of HBV in Various Body Fluids

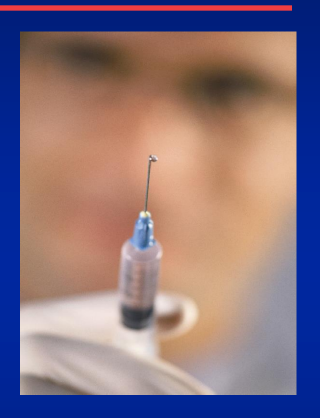
High		Moderate	Low/Not Detectable
blood		semen vaginal fluid	urine feces
wound exuda	udates	breast milk	sweat
			tears



Elimination of HBV Transmission,

Strategy

- Prevent perinatal HBV transmission
- Routine vaccination of all infants
- Vaccination of children in high-risk groups
- Vaccination of adolescents
 - all children up through age 18
- Vaccination of adults in high-risk groups





Hepatitis B Vaccine

- Licensed in 1982; currently recombinant (in US)
- 3 dose series, typical schedule 0, 1-2, 4-6 months no maximum time between doses (no need to repeat missed doses or restart)

- Protection ~30-50% dose 1; 75% 2; 96% 3;
- lower in older, immunosuppressive illnesses (e.g., HIV, chronic liver diseases, diabetes), obese, smokers



Hepatitis B Vaccination ACIP Recommendations

- Routine infant
- Ages 11-15 "catch up", and through age 18(VFC eligible)
- Over 18 high risk
 - Occupational risk (HCWs)
 - Hemodyalisis patients
 - All STD clinic clients
 - Multiple sex partners or prior STD
 - Inmates in Correctional settings
 - MSM
 - IDU
 - Institution for developmental disability
- Pre-vaccination testing if cost effective
- Post-vaccination testing 1-2 months after last shot, if establishing response critical (HCW)



VIRAL HEPATITIS C



VIRAL HEPATITIS TYPE C

Etiology:

Hepatitis C virus is a RNA-virus measuring 50 nm. It is classed into a separate genus, Hepacavirus of the Flaviviridae family.

The source of infection

Long-term in viremia (in the ende IP), chronic infections.

Route of transmission

Parenteral transmission. Sporadically, vertical and sexual transmissions were reported carrier or the sick person.

Susceptibility

Susceptibility is general.

Preventive measures:

The same as for HBV, exclusive immunization.



Features of Hepatitis C Virus Infection

Incubation period

Acute illness (jaundice)

Case fatality rate

Chronic infection

Chronic hepatitis

Cirrhosis

Agerelated Average 6-7 weeks

Range 2-26 weeks

Mild (<u><</u>20%)

Low

60%-85%

10%-70% (most asx)

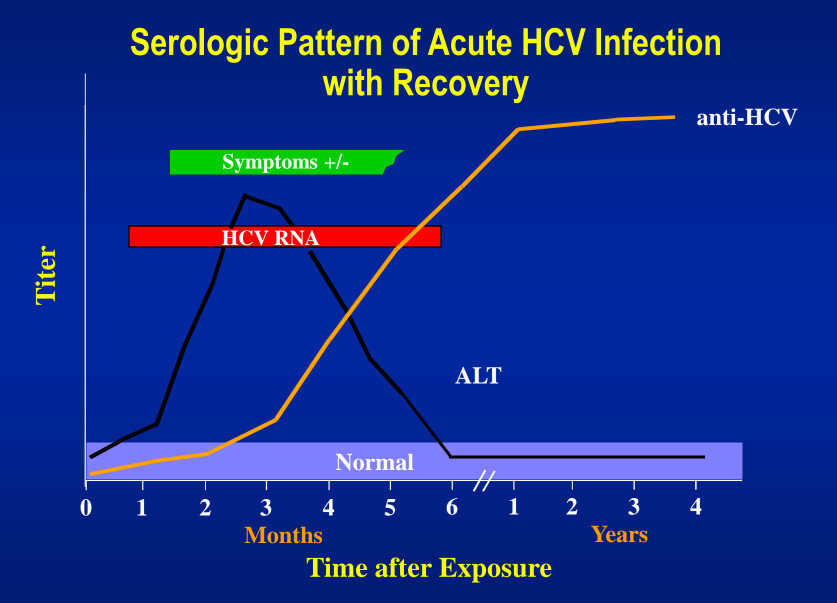
<5%-20%



Chronic Hepatitis C Factors Promoting Progression or Severity

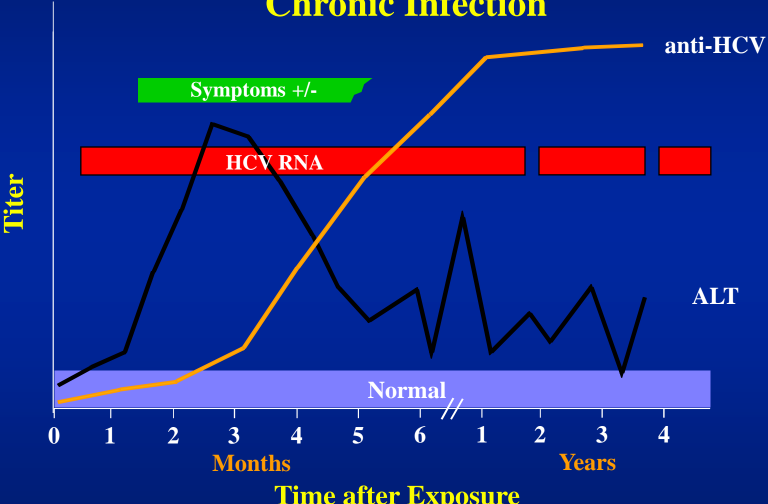
- Increased alcohol intake
- Age > 40 years at time of infection
- HIV co-infection
- Other
 - Male gender
 - Chronic HBV co-infection







Serologic Pattern of Acute HCV Infection with Progression to **Chronic Infection**



Time after Exposure



Exposures Known to Be Associated With HCV Infection

- Injecting drug use
- Transfusion, transplant from infected donor
- Occupational exposure to blood
 - Mostly needle sticks
- latrogenic (unsafe injections)
- Birth to HCV-infected mother
- Sex with infected partner
 - Multiple sex partners

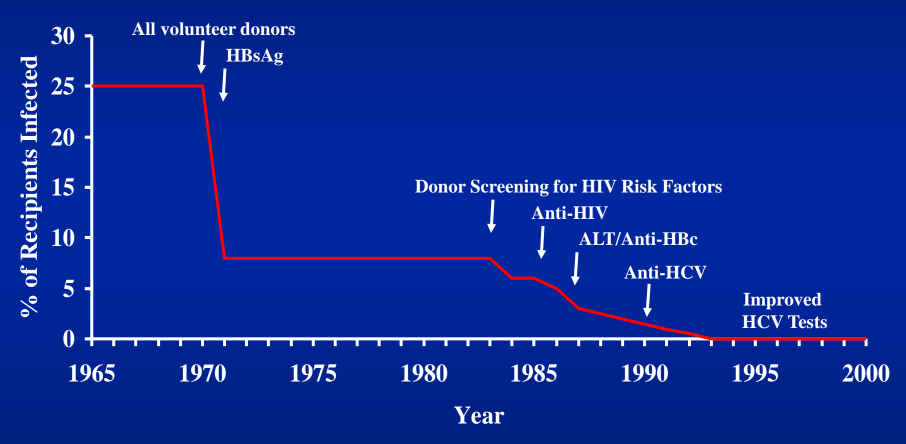


Injecting Drug Use and HCV Transmission

- Highly efficient
 - Contamination of drug paraphernalia, not just needles and syringes
- Rapidly acquired after initiation
 - 30% prevalence after 3 years
 - ->50% after 5 years
- Four times more common than HIV



Posttransfusion Hepatitis C



Adapted from HJ Alter and Tobler and Busch, Clin Chem 1997



Occupational Transmission of HCV

- Inefficient by occupational exposures
- Average incidence 1.8% following needle stick from HCV-positive source
 - Associated with hollow-bore needles
- Case reports of transmission from blood splash to eye; one from exposure to non-intact skin
- Prevalence 1-2% among health care workers
 - Lower than adults in the general population
 - 10 times lower than for HBV infection



HCV Related to Health Care Procedures

- Recognized primarily in context of outbreaks
 - Chronic hemodialysis
 - Hospital inpatient setting
 - Private practice setting
 - Home therapy
- Unsafe injection practices
 - Reuse of syringes and needles
 - Contaminated multiple dose medication vials



Perinatal Transmission of HCV

- Transmission only from women HCV-RNA positive at delivery
 - Average rate of infection 6%
 - Higher (17%) if woman co-infected with HIV
 - Role of viral titer unclear
- No association with
 - Delivery method
 - Breastfeeding
- Infected infants do well
 - Severe hepatitis is rare



Sexual Transmission of HCV

- Case-control, cross sectional studies
 - Infected partner, multiple partners, early sex, non-use of condoms, other STDs, sex with trauma, BUT
 - MSM no higher risk than heterosexuals
- Partner studies
 - Low prevalence (1.5%) among long-term partners
 - infections might be due to common percutaneous exposures (e.g., drug use), BUT
 - Male to female transmission more efficient
 - more indicative of sexual transmission



Sexual Transmission of HCV

- Occurs, but efficiency is low
 - Rare between long-term steady partners
 - Factors that facilitate transmission between partners unknown (e.g., viral titer)
- Accounts for 15-20% of acute and chronic infections
- Sex is a common behavior
 - Large chronic reservoir provides multiple opportunities for exposure to potentially infectious partners



Household Transmission of HCV

- Rare but not absent
- Could occur through percutaneous/mucosal exposures to blood
 - Contaminated equipment used for home therapies
 - IV therapy, injections
 - Theoretically through sharing of contaminated personal articles (razors, toothbrushes)

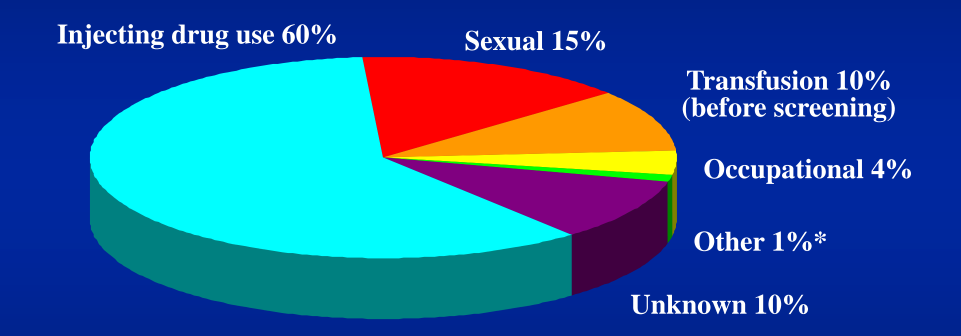


Other Potential Exposures to Blood

- No or insufficient data showing increased risk
 - intranasal cocaine use, tattooing, body piercing, acupuncture, military service
- No associations in acute case-control or population-based studies
- Cross-sectional studies in highly selected groups with inconsistent results
 - Temporal relationship between exposure and infection usually unknown
 - Biologically plausible, but association or causal relationship not established



Sources of Infection for Persons With Hepatitis C



Source: Centers for Disease Control and Prevention



^{*} Nosocomial; iatrogenic; perinatal

Reduce or Eliminate Risks for Acquiring HCV Infection

- Screen and test donors
- Virus inactivation of plasma-derived products
- Risk-reduction counseling and services
 - Obtain history of high-risk drug and sex behaviors
 - Provide information on minimizing risky behavior, including referral to other services
 - Vaccinate against hepatitis A and/or hepatitis B
- Safe injection and infection control practices

MMWR 1998;47 (No. RR-19)



Reduce Risks for Disease Progression and Further Transmission

- Identify persons at risk for HCV and test to determine infection status
 - Routinely identify at risk persons through history, record review
- Provide HCV-positive persons
 - Medical evaluation and management
 - Counseling
 - Prevent further liver damage
 - Prevent transmission to others



HCV Testing Routinely Recommended

Based on increased risk for infection

- Ever injected illegal drugs
- Received clotting factors made before 1987
- Received blood/organs before July 1992
- Ever on chronic hemodialysis
- Evidence of liver disease

Based on need for exposure management

- Healthcare, emergency, public safety workers after needle stick/mucosal exposures to HCV-positive blood
- Children born to HCV-positive women



Postexposure Management for HCV

- IG, antivirals not recommended for prophylaxis
- Follow-up after needlesticks, sharps, or mucosal exposures to HCV-positive blood
 - Test source for anti-HCV
 - Test worker if source anti-HCV positive
 - Anti-HCV and ALT at baseline and 4-6 months later
 - For earlier diagnosis, HCV RNA at 4-6 weeks
 - Confirm all anti-HCV results with RIBA
- Refer infected worker to specialist for medical evaluation and management



Routine HCV Testing of Uncertain Need

Not confirmed as risk factor/prevalence low or unknown

- Recipients of transplanted tissue
- Intranasal cocaine or other non-injecting illegal drug users
- History of tattooing, body piercing

Confirmed risk factor but prevalence of infection low

- History of STDs or multiple sex partners
- Long-term steady sex partners of HCV-positive persons



Mother-to-Infant Transmission of HCV

- Postexposure prophylaxis not available
- No need to avoid pregnancy or breastfeeding
 - Consider bottle feeding if nipples cracked/bleeding
- No need to determine mode of delivery based on HCV infection status
- Test infants born to HCV-positive women
 - ->15-18 months old
 - Consider testing any children born since woman became infected
 - Evaluate infected children for CLD



Sexual Transmission of HCV

Persons with One Long-Term Steady Sex Partner

- Do not need to change their sexual practices
- Should discuss with their partner.
 - Risk (low but not absent) of sexual transmission
 - Counseling and testing of partner should be individualized
 - May provide couple with reassurance
 - Some couples might decide to use barrier precautions to lower limited risk further



Sexual Transmission of HCV

Persons with High-Risk Sexual Behaviors

- At risk for sexually transmitted diseases, e.g.,
 HIV, HBV, gonorrhea, chlamydia, etc.
- Reduce risk
 - Limit number of partners
 - Use latex condoms
 - Get vaccinated against hepatitis B
 - MSMs also get vaccinated against hepatitis A



Other Transmission Issues

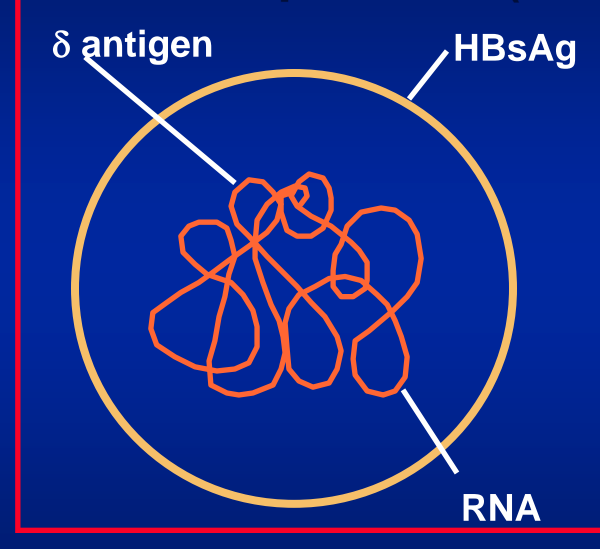
- HCV not spread by kissing, hugging, sneezing, coughing, food or water, sharing eating utensils or drinking glasses, or casual contact
- Do not exclude from work, school, play, childcare or other settings based on HCV infection status



VIRAL HEPATITIS D



Hepatitis D (Delta) Virus







Hepatitis D - Clinical Features

Coinfection

- -severe acute disease
- -low risk of chronic infection
- Superinfection
 - -usually develop chronic HDV infection
 - -high risk of severe chronic liver disease



Hepatitis D Virus Modes of Transmission

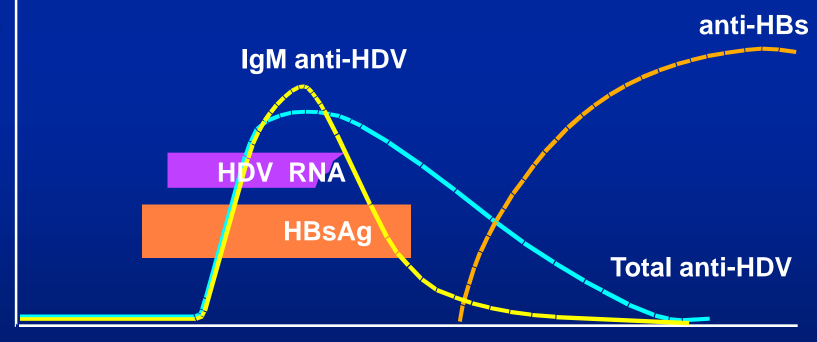
- Percutanous exposures
 - -injecting drug use
- Permucosal exposures
 - -sex contact



HBV - HDV Coinfection Typical Serologic Course

Symptoms

ALT Elevated

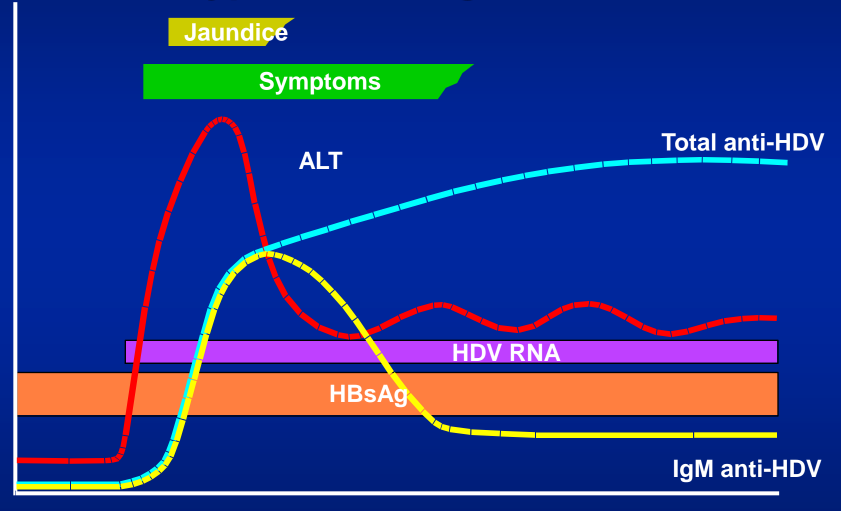


Time after Exposure



Tel

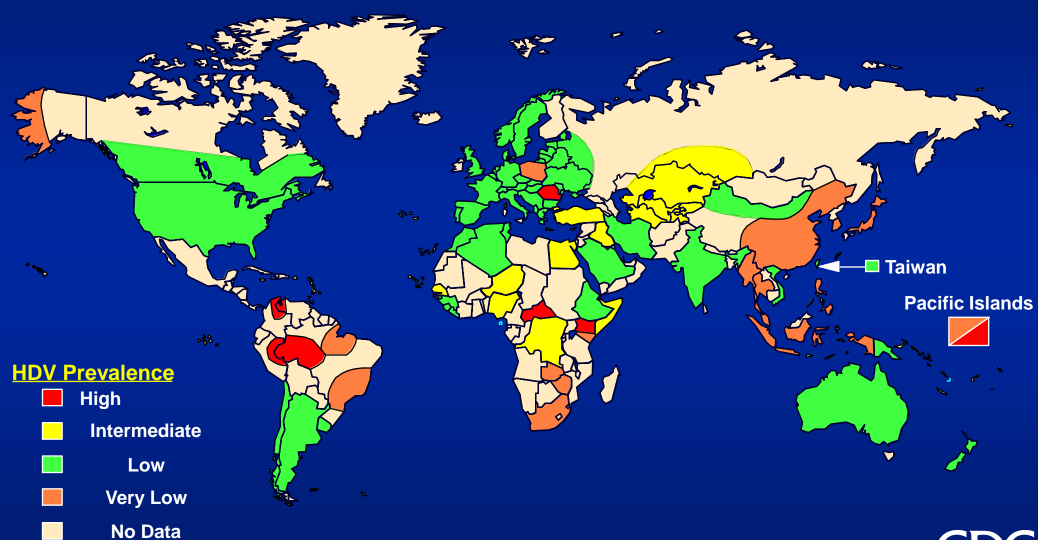
HBV - HDV Superinfection Typical Serologic Course



Time after Exposure



Geographic Distribution of HDV Infection





Hepatitis D - Prevention

HBV-HDV Coinfection

Pre or postexposure prophylaxis to prevent HBV infection

HBV-HDV Superinfection

Education to reduce risk behaviors among persons with chronic HBV infection



VIRAL HEPATITIS E



Hepatitis E Virus



Hepatitis E - Clinical Features

Incubation period: Average 40 days

Range 15-60 days

Case-fatality rate: Overall, 1%-3%

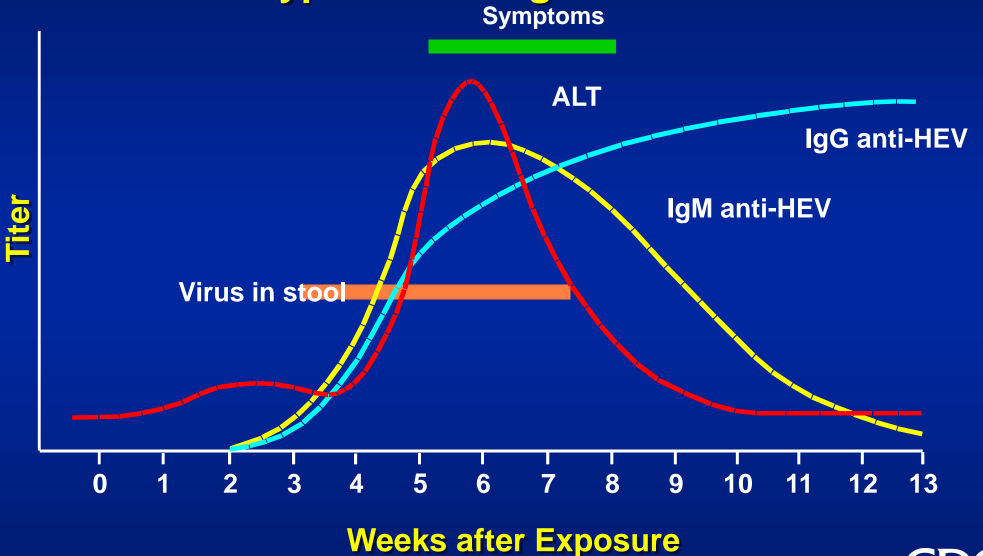
Pregnant women, 15%-25%

Illness severity: Increased with age

Chronic sequelae: None identified



Hepatitis E Virus Infection Typical Serologic Course





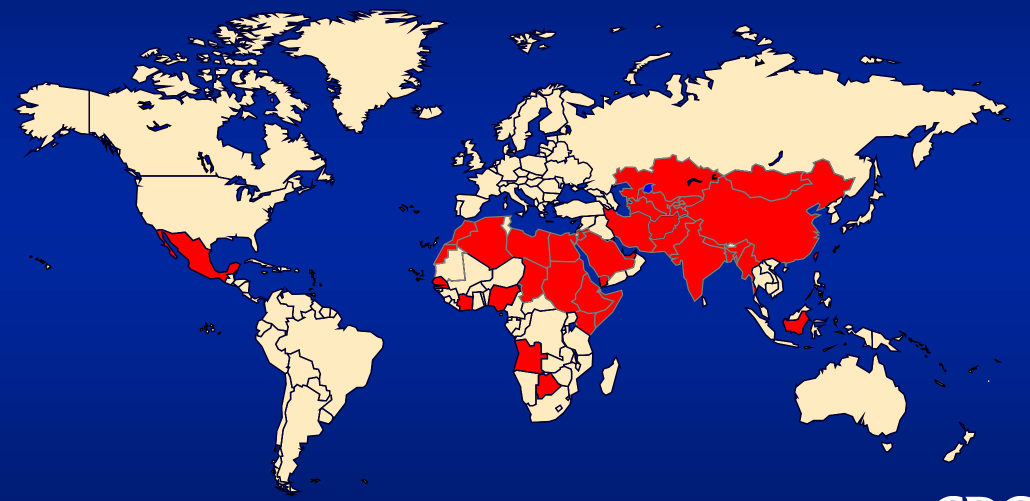
Hepatitis E - Epidemiologic Features

- Most outbreaks associated with fecally contaminated drinking water
- Minimal person-to-person transmission
- U.S. cases usually have history of travel to HEV-endemic areas



Geographic Distribution of Hepatitis E

Outbreaks or Confirmed Infection in >25% of Sporadic Non-ABC Hepatitis





Prevention and Control Measures for Travelers to HEV-Endemic Regions

- Avoid drinking water (and beverages with ice) of unknown purity, uncooked shellfish, and uncooked fruit/vegetables not peeled or prepared by traveler
- IG prepared from donors in Western countries does not prevent infection
- Unknown efficacy of IG prepared from donors in endemic areas
- Vaccine?



Key characteristics of HAV, HBV, HCV, HDV, HEV A Picornaviridae Hepadnaviridae Deltaviridae Raviviridae **RNA** DNA **RNA RNA**

2 - 6 months

Acute hepatitis more

common in adults

IgM anti-HBc

Yes: 0,1 -1,0 % are

fulminant

period

chronic disease, carriers

blood

genital secretions

Perinatal

Bloodborne

Sexual

Active (recombinant

Chronic infection leading to Chronic infection leading to

last 2 months of incubation last 2 months of incubation

entire period of acute stage entire period of acute stage

Causative agent

sequelae

2-6 weeks

Case fatality increases with age

IgM anti-HAV

none

No

last 2 weeks of incubation period

first day of acute stage

faeces

viremia - 1. day of illnes

Person-to person

Foodborne

Waterborne

Inactivated hepatitis A vaccines are

Incubation period

Characteristic of

Biomarker of recent

Chronic infection

hepatocelular Ca

Infectious biological

Cirrhosis and

The period of

infectivity

material

Mode of

transmission

acute hepatitis

infection

2 - 6 months

almost never fulminant

None

seguelae

Yes: 50 % can be fulminant

period

chronic disease, carriers

blood

genital secretions

Blodborne

Perinatal

Sexual

Acute hepatitis uncommon, HDV in chronic heptitis

Ε

Hepeviridae

RNA

2 - 10 weks

High case fatality in

pregnant women -10-20 %;

other 1 -2 %

IgM anti-HEV

Very rare

NO

??

faeces

meat of animals

Waterborne

Foodborne

Person-to person

3-7 weeks Superinfection with

B may lead to fulminnat

disease

IgM anti-HDV

Chronic hepatitis that

copmlicated chronic

hepatitis B

Yes: 5 - 20 % can be

fulminant

??

blood

Blodborne