Viral Hepatitis Resource And Services Directory

Indiana State Department of Health
Office of HIV/AIDS/Viral Hepatitis
2 N. Meridian St. 6-C
Indianapolis, IN 46202
Phone: (317) 233-1325
Fax: (317) 233-7663
www.in.gov/isdh/programs/hivstd
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It is estimated that more than 18,000 Hoosiers have a hepatitis C infection. Many who are infected do not even know it.

**HEALTH CARE PROVIDERS** - are critical links in the prevention, diagnosis, and treatment of viral hepatitis, including those specializing in STD/HIV, family planning, and substance abuse. This resource guide and Service Directory will help you integrate important viral hepatitis messages into your regular patient/client education practices. Both youth and adults are equally at risk for hepatitis.

**WHAT IS VIRAL HEPATITIS?**
Viral hepatitis is an inflammation of the liver that is caused by a viral infection, but can also be caused by bacteria, drugs, toxins, or excessive alcohol intake. At this time, there are five viruses known to affect the liver and cause hepatitis: A, B, C, D and E.

- An estimated 30% of the population has been infected with Hepatitis A at some point in time.
- In the United States, more than 1 million people are infected with Hepatitis B.
- Nearly 4 million people nationwide are infected with Hepatitis C.
How to Use This Directory Effectively

The Indiana State Department of Health values all providers who care for Hoosiers who may have an infectious disease and/or co-occurring disorders. Thank you for choosing to use this directory as an informational and reference guide to assist you in your work.

The challenge we all have is to properly assess the patients we are working with and then provide appropriate assurances for their care. Often referrals are a necessary part of this care and this directory is intended to assist you and your staff in making those referrals.

The complexities associated with individuals infected with viral hepatitis are well founded. Add to that the real possibility that they may not have adequate insurance and/or may have multiple conditions. Clearly, you as the provider caring for these individuals are greatly challenged.

Inside these pages is information and resources that will assist you in the following areas:

- Referrals for Hepatitis C Virus (HCV) testing
- A list of medical providers who treat HCV
- IN HCV Support Groups
- Local and national help lines
- Web sites
- Substance abuse treatment services
- Fact sheets
- Frequently asked questions
How to Integrate Hepatitis Prevention into Patient/Client Education
Integrating Hepatitis Education

Talking about contraception? STDs? HIV? Drug use? Hygiene? Educating patients/clients who are at high risk about viral hepatitis can easily become part of your current practice. Here are some examples:

<table>
<thead>
<tr>
<th>HEPATITIS A VIRUS</th>
<th>HEPATITIS B VIRUS</th>
<th>HEPATITIS C VIRUS</th>
</tr>
</thead>
</table>
| • Good hygiene (being clean) is important to your overall health.  
  • Washing your hands with soap and water helps kill the germs that cause infections including hepatitis A.  
  • Indiana offers free Hepatitis A vaccines for people under the age of 19. | • Use a condom every time you have sex to prevent the spread of STDs such as gonorrhea, hepatitis B, or HIV.  
  • Hepatitis B is 100 times more contagious than HIV, and can lead to liver cancer, cirrhosis, and even death.  
  • Indiana offers free Hepatitis B vaccines for people under the age of 19. | • If you shoot or snort drugs, don’t share works (needles, cottons, cookers). Sharing works puts you at risk of getting HIV and hepatitis C.  
  • Hepatitis C can cause liver cancer, liver scarring, and in some cases death.  
  • There is no vaccine available for hepatitis C. |

Tips on Starting a Sensitive Conversation

To begin a conversation about viral hepatitis (or other STDs or HIV) ask:

“Have you ever heard about hepatitis? ...”  
“What have you heard?”  
“Do you know anyone who has hepatitis?”  
“Do you know that hepatitis B is 100 times more contagious than HIV?”

Normalizing risky behaviors can help. Start your conversation with:

“People your age sometimes experiment with drugs and alcohol.”  
“Have you ever tried drugs?”  
“Have you ever shared injection drug equipment or a straw to snort drugs?”  
“Do you think people your age are practicing safe sex, for example, by using condoms to protect themselves from sexually transmitted diseases and pregnancy?”
Tips on Counseling

Sex practices and drug use are difficult topics to discuss—especially with youth. But it is important to provide everyone with information on how to protect themselves. Here are some tips to keep in mind when talking with patients/clients about practicing safe sex and reducing or stopping substance use.

- Assure patient confidentiality.
- Listen to your patient.
- Accept that your patient/client’s values may be different from your own.
- Avoid judging a patient/client’s personal behaviors.
- Be sensitive to expressions and gestures (both yours and your patient/client’s).
- Help your patients explore their options for reducing or stopping unsafe sexual or substance-using behaviors.

Assessing Patient Risk

Ask patients/clients to respond to the following statements to quickly assess their hepatitis risk level:

I wash my hands with soap and water before eating and after going to the bathroom. (Hepatitis A)  
Yes  No

I use a condom every time I have sex. (Hepatitis B or C)  
Yes  No

I have never injected or snorted drugs. (Hepatitis C)  
Yes  No

Patients who answer “no” to any of these questions may be at risk for viral hepatitis.
What to Tell Patients/Clients Who are at Risk

All patients/clients, but especially those who are at risk, should learn about viral hepatitis. Our conversations with teens and others showed they want to know what viral hepatitis is and how to prevent it. When talking to patients/clients, the use of meaningful statistics can have a powerful effect.

Example: *Hepatitis B is 100 times more contagious than HIV.*

Tell patients/clients who are at risk to:
- Get immunized against *both* hepatitis A and B.
- Use a latex condom every time you have sex.
- Never share works with anyone if they shoot or snort drugs.
- Wash their hands with soap and water often and thoroughly.
- Only get tattoos or ear, tongue, and other body piercings from licensed facilities.
- Your local health department can tell you if a facility is licensed.
- Avoid contact with someone else’s blood (e.g. cuts or scrapes), and don’t share razors, toothbrushes, or anything else that may have blood on it.
- Avoid eating raw shellfish.

Testing Information

Once a patient/client has been identified as being at high risk for viral hepatitis infection, testing should be considered. The CDC recommends testing for hepatitis C, and immunizing against hepatitis A and B. For more information, visit www.cdc.gov/hepatitis. Testing for hepatitis C in Indiana is confidential. To help your patients locate a testing site in their area, call your local health department.
**Immunization Information**

- Hepatitis A and B immunizations are recommended for all who are at risk.

- People infected with hepatitis C should receive hepatitis A and B immunizations to prevent further liver damage.

- There is no immunization available against hepatitis C.

Most youth have received the hepatitis B immunization as a newborn, or in school through Indiana’s Vaccinate Before You Graduate Program. However, there are still many youth and adults who have not been immunized against hepatitis B.

Ask the patient/client you serve if they know their hepatitis immunization status, and check their immunization record. If an immunization record does not exist, start one for them.

**Cost and availability**

Indiana offers hepatitis A and B immunizations to at-risk clients seeking STD services in all Indiana STD clinics. An STD exam may be required and a small administrative fee may be charged.*

*Charges are waived for inability to pay.
Sites Where Hepatitis C Anti-Body Testing, Hepatitis A and B Vaccine, and STD Exams are Available for Uninsured and Underinsured (STD Exam may be Required)
CENTRAL INDIANA

Bartholomew County Health Dept
1971 State Street
Columbus, IN 47201-6965
812-379-1555

Bell Flower Clinic
1101 W. 10th St.
Indianapolis, IN 46202
317-221-8347

Boone County Health Dept
116 W. Washington St.
Lebanon, IN 46052
765-973-9243

Madison County Health Dept
206 E. 9th St.
Anderson, IN 46016
765-646-9206

Wayne County Health Dept
301 E. Main St. - Old Courthouse
Richmond, IN 47374
812-379-1555
### NORTHERN INDIANA

<table>
<thead>
<tr>
<th>Health Department</th>
<th>Address</th>
<th>City</th>
<th>State</th>
<th>Zip Code</th>
<th>Phone</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allen County Health Dept</td>
<td>1 E. Main St. 5th Fl.</td>
<td>Ft. Wayne</td>
<td>IN</td>
<td>46802-1810</td>
<td>260-449-7054</td>
</tr>
<tr>
<td>Healthy Children/Healthy Teens</td>
<td>3500 S. LaFountain</td>
<td>Kokomo</td>
<td>IN</td>
<td>46902</td>
<td>765-854-2440</td>
</tr>
<tr>
<td>East Chicago Health Dept</td>
<td>100 W. Chicago Ave.</td>
<td>East Chicago</td>
<td>IN</td>
<td>46132</td>
<td>219-391-8258</td>
</tr>
<tr>
<td>Howard County Health Dept</td>
<td>120 E. Mulberry</td>
<td>Kokomo</td>
<td>IN</td>
<td>46901</td>
<td>765-456-2408</td>
</tr>
<tr>
<td>Elkhart County Health Dept</td>
<td>608 Oakland Ave.</td>
<td>Elkhart</td>
<td>IN</td>
<td>46516</td>
<td>574-523-2128</td>
</tr>
<tr>
<td>Lafayette Planned Parenthood</td>
<td>1016 E. Main St.</td>
<td>Lafayette</td>
<td>IN</td>
<td>47901</td>
<td>765-742-7524 ext.1</td>
</tr>
<tr>
<td>Gary City Health Dept</td>
<td>1145 W. 5th Ave.</td>
<td>Gary</td>
<td>IN</td>
<td>46402</td>
<td>219-882-5565</td>
</tr>
<tr>
<td>La Porte County Health Dept</td>
<td>302 W. 8th St.</td>
<td>Michigan City</td>
<td>IN</td>
<td>46360</td>
<td>219-326-6808 ext. 2000</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>219-873-7001</td>
</tr>
</tbody>
</table>
SOUTHERN INDIANA

Clark County Health Dept
1403 Spring St.
Jeffersonville, IN 47130
812-288-2706;
1-800-828-5624

Vigo County Health Dept
175 Oak St.
Terre Haute, IN 47802
812-462-3431

Vanderburgh County Health Dept
420 Mulberry St
Evansville, IN 47708-1888
812-435-5683
Indiana Healthcare Providers for Uninsured and Underinsured
CENTRAL INDIANA

Citizens Primary Care Health Center
1650 N. College Ave.
Indianapolis
317-396-0211
Counties Served: Marion

Open Door/BMH Health Center
905 S. Walnut St.
Muncie, IN 47308
765-289-5928
Counties Served: Delaware

Edinberg/Trafalgar Family Health Centers, Inc.
14 Trafalgar Square
Trafalgar, IN 46181
317-878-2301
Counties Served: Johnson

Raphael Health Center
401 E. 34th St.
Indianapolis
317-926-1507
Counties Served: Marion

HealthNet Inc.
3401 E. Raymond St.
Indianapolis
317-781-3175
Counties Served: Marion

Shalom Health Care Center, Inc.
3737 N. Meridian St. STE 402
Indianapolis
317-923-4915 ext 202 or 317-923 4915
Counties Served: Marion

HealthNet Centers
Barrington, Peoples,
Southeast, Southwest,
Martindale Brightwood,
Homeless Initiative Program

Marion County Health Department
3838 N. Rural St.
Indianapolis
317-221-2037
Counties Served: Marion

Madison County Community Health Center, Inc.
1547 Ohio Ave.
Anderson, IN 46015
765-641-7499
Counties Served: Madison
<table>
<thead>
<tr>
<th>Location</th>
<th>Address</th>
<th>Phone Number</th>
<th>Counties Served</th>
</tr>
</thead>
<tbody>
<tr>
<td>East Chicago Community Health Center</td>
<td>100 W. Chicago Ave.</td>
<td>219-392-4900</td>
<td>Lake</td>
</tr>
<tr>
<td>Indiana Health Center at South Bend</td>
<td>1901 #B Western Avenue</td>
<td>574-234-9033</td>
<td>St. Joseph</td>
</tr>
<tr>
<td>Gary Community Health Center Inc.</td>
<td>1021 West 5th Ave.</td>
<td>219-880-1190</td>
<td>Lake</td>
</tr>
<tr>
<td>LaPorte Indigent Drug Program</td>
<td>1007 Lincolnway,</td>
<td>317-326-2504</td>
<td>LaPorte</td>
</tr>
<tr>
<td>Heart City Health Center</td>
<td>236 Simpson Ave.</td>
<td>574-293-0052</td>
<td>Elkhart</td>
</tr>
<tr>
<td>Memorial Health System</td>
<td>707 N. Michigan St.</td>
<td>574-237-6640</td>
<td>St. Joseph</td>
</tr>
<tr>
<td>Hilltop Community Health Center Inc.</td>
<td>460 S. College Ave.</td>
<td>219-462-7173</td>
<td>Lake, Porter, Starke, Jasper, Marshall, St. Joseph, Newton, LaPorte</td>
</tr>
<tr>
<td>Neighborhood Health Clinics, Inc.</td>
<td>P.O. Box 11949,</td>
<td>260-458-2644</td>
<td>Allen</td>
</tr>
<tr>
<td>Indiana Health Center at Kokomo</td>
<td>3118 S. LaFountain St.</td>
<td>317-864-4160</td>
<td>Howard</td>
</tr>
<tr>
<td>North Shore Health Centers</td>
<td>6450 U.S. Highway 6</td>
<td>219-763-8112</td>
<td>Northeast part of Lake (Hobart, Lake Station, Miller)</td>
</tr>
<tr>
<td>Indiana Health Center at Marion</td>
<td>925 S. Nebraska St.</td>
<td>765-664-7492</td>
<td>Grant</td>
</tr>
<tr>
<td>Tippecanoe Community Health Clinic</td>
<td>1716 Hartford St.</td>
<td>765-742-1567</td>
<td>Tippecanoe</td>
</tr>
</tbody>
</table>
SOUTHERN INDIANA

Echo Community Health Care, Inc.
501 John St., Suite 12
Evansville, IN 47713
812-436-0214
Counties Served: Vanderburgh Posey,
Gibson, Warick

Jackson County Community
Health Center, Inc.
c/o: Indiana Health Centers
317-632-1231
Counties Served: Jackson

Tulip Tree Family Health Services
Gibson Co. Inc
123 N. McCreary Street,
Ft. Branch, IN 47670
812-753-1039
County Served: Gibson
Gastroenterologists
CENTRAL INDIANA

Brian Clarke M.D.  
7250 Clearvista Dr., Ste. 390  
Indianapolis, IN 46256  
317-621-2100

Central Indiana  
Gastroenterology Group  
Christopher Foster M.D.  
2020 Meridian St. Ste. 340  
Anderson, IN 46016  
765-646-8477

Central Indiana  
Gastroenterology Group  
Jon Maier M.D.  
2020 Meridian St. Ste. 340  
Anderson, IN 46016  
765-646-8477

Central Indiana  
Gastroenterology Group  
Rod Nisi M.D.  
2020 Meridian St. Ste. 340  
Anderson, IN 46016  
765-646-8477

Digestive Diseases Center  
Richard Weddle M.D.  
2920 McIntire Dr. #310  
Bloomington, IN 47403  
812-331-0233

Gastroenterology Associates  
Joseph Henderson M.D.  
1400 N. Ritter Ave., Ste. 370  
Indianapolis, IN 46219  
317-355-1144

Gastroenterology Associates  
Eldred Mac Donel M.D.  
7950 N. Shadeland Ave. Ste. 350  
Indianapolis, IN 46250  
317-578-2600

Gastroenterology Associates  
David Hollander M.D.  
1400 N. Ritter Ave. Ste. 370  
Indianapolis, IN 46219  
317-355-1144

Gastroenterology Associates  
Stephen Carlson M.D.  
7950 N. Shadeland Ave. Ste. 350  
Indianapolis, IN 46250  
317-578-2600

Gastroenterology Associates  
Matthew Harrison M.D.  
7950 N. Shadeland Ave. Ste. 350  
Indianapolis, IN 46250  
317-578-2600

Gastroenterology Associates  
Donato Chiccia M.D.  
7950 N. Shadeland Ave. Ste. 350  
Indianapolis, IN 46250  
317-578-2600

Gastroenterology Hepatology  
Lawrence Lumeng M.D.  
1481 W. 10th St., Room-C 7117  
Indianapolis, IN 46202  
317-554-0000
Gastroenterology Hepatology
Naga Chalasani M.D.
975 W. Walnut St., Rm 424
Indianapolis, IN 46202
317-274-3090

Indiana Gastroenterology
Paul Frederick M.D.
8424 Naab Rd., Ste. 1L
Indianapolis, IN 46260
317-872-1161

Indianapolis Gastroenterology
& Hepatology
Brian Speri M.D.
8051 S. Emerson Ave., Ste. 200
Indianapolis, IN 46237
317-865-2955

Indianapolis Gastroenterology
& Hepatology
David Pound M.D.
8051 S. Emerson Ave., Ste. 200
Indianapolis, IN 46237
317-865-2955

Indianapolis Gastroenterology
& Hepatology
Earl Brown M.D.
8051 S. Emerson Ave., Ste. 200
Indianapolis, IN 46237
317-865-2955

Internal Medicine Associates
Prodyot Ghosh M.D.
550 Landmark Ave.
Bloomington, IN 47402
812-333-5973

IU Medical Center
Paul Kwo M.D.
975 W. Walnut St. 1B 327
Indianapolis, IN 46202
317-274-3090

Indiana Gastroenterology
William Fecht Jr. M.D.
8424 Naab Rd. 1L
Indianapolis, IN 46260
317-872-1161

Indianapolis Gastroenterology
& Hepatology
A. Thompson Colley M.D.
8051 S. Emerson Ave., Ste. 200
Indianapolis, IN 46237
317-865-2955

Indianapolis Gastroenterology
& Hepatology
Linda Ritchison M.D.
8051 S. Emerson
Indianapolis, IN 46237
317-865-2955

Indianapolis Gastroenterology
& Hepatology
Gregory Lemmel M.D.
8051 S. Emerson Ave., Ste. 200
Indianapolis, IN 46237
317-865-2955

Internal Medicine Associates
Jitender Bhandari M.D.
550 Landmark Ave.
Bloomington, IN 47402
812-333-5973

IU Medical Center
Emad Rahmani M.D.
550 University Blvd., Ste. 2300
Indianapolis, IN 46202
317-274-5000

Medical Consultants
Steven Kaiser M.D.
800 S. Tillotson Ave.
Muncie, IN 17304
765-281-2000
Meridian Medical Group
Martin Meisenheimer M.D.
1801 N. Senate Ave. #400
Indianapolis, IN 46202
317-962-6300

Northside Gastroenterology
Lawrence Born M.D.
8424 Naab Rd. Ste 3-J
Indianapolis, IN 46260
317-872-7396

Medical Consultants
Steven Kaiser M.D.
800 S. Tillotson Ave.
Muncie, IN 17304
765-281-2000

Northside Gastroenterology
Robert Callon M.D.
8424 Naab Rd. Ste., 3-J
Indianapolis, IN 46260
317-872-7396

Northside Gastroenterology
Arthur Baluyut M.D.
8424 Naab Rd. Ste., 3-J
Indianapolis, IN 46260
317-872-7396

Northside Gastroenterology
Daryl Daugherty M.D.
8424 Naab Rd. Ste., 3-J
Indianapolis, IN 46260
317-872-7396
NORTHERN INDIANA

Dr. Kaleen Ahmed  
1900 Carew St. Ste. 1  
Ft. Wayne, IN 46805  
260-484-2524

Dr. Imad Horani  
1900 Carew St. Ste. 1  
Ft. Wayne, IN 46805  
260-484-2524

Dr. Ricky Meyer  
1900 Carew St. Ste. 1  
Ft. Wayne, IN 46805  
260-484-2524

Dr. Andrew Katz  
7836 Jefferson Blvd.  
Ft. Wayne, IN 46804  
260-432-2297

Dr. Sushi Jain  
7836 Jefferson Blvd.  
Ft. Wayne, IN 46804  
260-432-2297

Dr. Carl Wrobleski  
7836 Jefferson Blvd.  
Ft. Wayne, IN 46804- 
260-432-2297

Dr. Binoy Ouseph  
1 Cayor Nickel Square  
Bluffton, IN 46714

Dr. Navinchandra Barot  
Barot & Associates  
7550 Hohman Ave. Ste. 1200 B  
Munster, IN 46321  
219-836-7890

Dr. Joel Cahan  
Consultants Gastroenterology  
701 Superior Ave. #G  
Munster, IN 46321  
219-922-3040

Dr. Benjamin Schmid  
Hammond Clinic Specialty Center  
7905 Calumet Ave.  
Munster, IN 46321  
219-836-5800

Dr. John Mirro/  
Internal Medicine Associates  
8895 Broadway  
Merrillville, IN 46410  
219-738-2081

Dr. Rao Ramachandra  
1400 S Lake Park Ave  
Hobart, IN 46342  
219-942-2185
SOUTHERN INDIANA

Dr. Ramesh Kalari
2520 Q. St.
Bedford, IN 47421
812-277-0977

Gastroenterology Associates
Dr. Cesar Bello
801 St. Mary's Dr. Ste. 110
Evansville, IN 47714
812-477-6103

Dr. Imad Koj
4757 S. 7th St.
Terre Haute, IN 47802
812-234-2289

Gastroenterology Associates
Dr. Gardar Gislason
801 St. Mary's Dr. Ste. 110
Evansville, IN 47714
812-477-6103

Dr. Jason Samuel
PO Box 16022
Evansville, IN 47714
812-473-1470

Gastroenterology Associates
Dr. J. Dennis Guletz
801 St. Mary's Dr. Ste. 110
Evansville, IN 47714
812-477-6103

AP & S Clinic
Dr. John Morse
221 S. 6th St.
Terre Haute, IN 47807
812-232-0564

Gastroenterology Associates
Dr. Joseph Haseman
801 St. Mary's Dr. Ste. 110
Evansville, IN 47714
812-477-6103

A P & S Clinic
Dr. Martin Schmidt
221 S. 6th St.
Terre Haute, IN 47807
812-232-0564

Gastroenterology Associates
Dr. William Johnson
801 St. Mary's Dr. Ste. 110
Evansville, IN 47714
812-477-6103

Gastroenterology Associates
Dr. Donald Bailey
801 St. Mary's Dr. Ste. 110
Evansville, IN 47714
812-477-6103

Gastroenterology Associates
Dr. Gregory McCord
801 St. Mary's Dr. Ste. 110
Evansville, IN 47714
812-477-6103
Gastroenterology Associates
Dr. Vajravel Prasad
801 St. Mary's Dr. Ste. 110
Evansville, IN 47714
812-477-6103

Gastroenterology Associates
Dr. Herman Rusche
801 St. Mary's Dr. Ste. 110
Evansville, IN 47714
812-477-6103

Gastroenterology of Southern Indiana
Dr. Stewart Coleman
825 University Woods Dr.
New Albany, IN 47150
812-945-0145

Dr. Atam Mesdiratta
Welborn Downtown
421 Chestnut
Evansville, IN 47713
812-426-9545

Dr. Bruce Schneider
Welborn Downtown
421 Chestnut
Evansville, IN 47713
812-426-9545

Gastroenterology of Southern Indiana
Dr. David Dresner
825 University Woods Dr.
New Albany, IN 47150
812-945-0145
Infectious Disease Doctors
## CENTRAL INDIANA

<table>
<thead>
<tr>
<th>Practice Name</th>
<th>Address 1</th>
<th>Address 2</th>
<th>City, State, Zip</th>
<th>Phone 1</th>
<th>Phone 2</th>
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<tbody>
<tr>
<td>Dr. Jesus Aguilar</td>
<td>5508 E. 16th St. Ste. C37</td>
<td></td>
<td>Indianapolis, IN 46218</td>
<td>317-355-3266</td>
<td></td>
</tr>
<tr>
<td>Comprehensive Infectious Disease of Indiana</td>
<td>998 E. Main St.</td>
<td>Danville, IN 46122</td>
<td>317-745-8686</td>
<td></td>
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<tr>
<td>Dr. Renita Brown</td>
<td>1633 N. Capitol Ave.</td>
<td></td>
<td>Indianapolis, IN 46202</td>
<td>317-962-2424</td>
<td></td>
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<tr>
<td>Infectious Disease of Indiana</td>
<td>Dr. Robert Baker</td>
<td></td>
<td>7430 N. Shadeland Ave., Ste. 230, Indianapolis, IN 46250</td>
<td>317-216-3626</td>
<td></td>
</tr>
<tr>
<td>Dr. Thomas Slama</td>
<td>8240 Naab Rd.</td>
<td></td>
<td>Indianapolis, IN 46260</td>
<td>317-870-1970</td>
<td></td>
</tr>
<tr>
<td>Infectious Disease of Indiana</td>
<td>Dr. Steven Norris</td>
<td></td>
<td>7430 N. Shadeland Ave. Ste. 230, Indianapolis, IN 46250</td>
<td>317-16-3626</td>
<td></td>
</tr>
<tr>
<td>Advanced Healthcare Associates</td>
<td>Dr. Hope Chema</td>
<td></td>
<td>5150 Shelbyville Rd., Indianapolis, IN 46237</td>
<td>317-782-1577</td>
<td></td>
</tr>
<tr>
<td>Infectious Disease of Indiana</td>
<td></td>
<td></td>
<td>10610 N. Pennsylvania Ste. #A, Indianapolis, IN 46280</td>
<td>317-587-2300</td>
<td></td>
</tr>
<tr>
<td>Beech Grove Internal Medicine</td>
<td>2030 Churchman Ave.</td>
<td></td>
<td>Beech Grove, IN 46107</td>
<td>317-781-2100</td>
<td></td>
</tr>
<tr>
<td>IU Medical Group</td>
<td></td>
<td></td>
<td>550 N. University Blvd., Indianapolis, IN 46202</td>
<td>317-630-6579</td>
<td></td>
</tr>
<tr>
<td>Circle City Internal Medicine</td>
<td>1500 Albany St. Ste. 1001</td>
<td></td>
<td>Indianapolis, IN 46107</td>
<td>317-82-6725</td>
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</tr>
<tr>
<td>IU Medical Group</td>
<td></td>
<td></td>
<td>550 N. University Blvd., Indianapolis, IN 46202</td>
<td>317-630-6579</td>
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</tr>
</tbody>
</table>
IU Medicine Infectious Disease
545 Barnhill Dr.
Indianapolis, IN 46202
317-554-4146

Riley Hospital for Children
Pediatric Infectious Diseases
702 Barnhill Dr.
Indianapolis, IN 46202
317-274-7260

LifeCare
1633 N. Capitol Ave. Ste. # 700
Indianapolis, IN 46202
317-962-3724

University Hospital Clinic
550 N. University Blvd.
Indianapolis, IN 46202
317-274-8660

Medical Specialists of Madison County
2101 Jackson St # 110
Anderson, IN 46016
765-43-6012

People's Health Center
2340 E. 10th St.
Indianapolis, IN 46201
317-33-736
NORTHERN INDIANA

Dr. Robert Hunt
919 E Jefferson Blvd # 400
South Bend, IN 46117
574-233-4083

Medical Associates of Grant County
Dr. Peter-Benson So
610 N River Rd
Marion, IN 46952
765-662-9870

Dr. Sheree Peglow
720 E. Cedar St. # 420
South Bend, IN 46601
574-234-5938

Medical Specialists Munster
Dr. Matthew Meyer
761 45th Ave # 103
Munster, IN 46321
219-922-3002

Arnett Health System
Dr. Barbara Beilska
1500 Salem St.
Lafayette, IN 47904
765-448-8488

Medical Specialists Munster
Dr. Alexander Stemer
761 45th Ave # 103
Munster, IN 46321
219-922-3002
219-922-3002
219-922-3002

Infectious Disease Associates
Dr. Barbara Nohinek
7910 W Jefferson Blvd Bldg 2
Fort Wayne, IN 46804
260-435-7590

Medical Specialists Munster
Dr. Matthew Meyer
761 45th Ave # 103
Munster, IN 46321
219-922-3002

Infectious Disease Specialist
Dr. Richard Yu
3229 Broadway # 151
Gary, IN 46409
219-985-0673

Medical Specialists Munster
Dr. Patricia Wangerin
761 45th Ave # 103
Munster, IN 46321
219-922-3000
SOUTHERN INDIANA

Dr. Jose Salgado
3801 Bellemeade Ave # 200 C
Evansville, IN 47714
812-485-1788

Matthew 25
100 Washington St
Evansville, IN 47713
812-423-5192

A P & S Clinic
221 S 6th St
Terre Haute, IN 47807
812-232-0564

University Physician Group
530 S Jackson St
Louisville, KY 40202
502-852-4929

Deaconess Medical Group Specialty
519 Harriet St
Evansville, IN 47710
812-423-9699
Research Hospitals
And
Veterans Affairs Hospitals
And Clinic
RESEARCH HOSPITALS

Indiana University Medical Center
550 University Blvd.
Indianapolis, IN 46202
317-278-1187

VETERAN FACILITIES

CENTRAL INDIANA

Bloomington Clinic
200 E. Winslow Rd.
Bloomington, IN 47701
812-353-2600

Muncie/Anderson Clinic
3500 W. Purdue Ave.
Muncie, IN 47304
765-284-8860

Richard L. Roudebush
VA Medical Center
1481 W. 10th St. 118
Indianapolis, IN 46202
317-554-0000

Richmond Clinic
4351 South A. St.
Richmond, IN 47346
765-973-6915
NORTHERN INDIANA

Crown Point Clinic
9330 S. Broadway
Crown Point, IN 46037-8602
219-662-0001

South Bend Clinic
17615 State Rd. 23
South Bend, IN 46635-1718
574-251-2819

Northern Indiana VA
Healthcare System
1700 E. 38th St.
Marion, IN 46953-4568
765-674-3321

West Lafayette Clinic
3851 N. River Rd.
Lafayette, IN 47906-3762
765-464-2280

Northern Indiana VA Healthcare System
2121 Lake Avenue
Ft. Wayne, IN 46805-5100
260-426-5431
# SOUTHERN INDIANA

<table>
<thead>
<tr>
<th>Clinic</th>
<th>Address</th>
<th>Location</th>
<th>Phone</th>
</tr>
</thead>
<tbody>
<tr>
<td>Evansville Clinic</td>
<td>500 E. Walnut</td>
<td>Evansville, IN</td>
<td>812-465-6202</td>
</tr>
<tr>
<td>Terre Haute Clinic</td>
<td>1635 N. Third St.</td>
<td>Terre Haute, IN</td>
<td>812-232-2890</td>
</tr>
<tr>
<td>New Albany Clinic</td>
<td>811 Northgate</td>
<td>New Albany, IN</td>
<td>502-894-6188</td>
</tr>
</tbody>
</table>
Drug Treatment Facilities
## CENTRAL INDIANA

<table>
<thead>
<tr>
<th>Organization</th>
<th>Address</th>
<th>Phone Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adult &amp; Child Mental Health Center Inc</td>
<td>8320 Madison Ave, Indianapolis, IN 46227</td>
<td>317-882-5122</td>
</tr>
<tr>
<td>Comprehensive Mental Health Services Inc</td>
<td>240 N Tillotson Ave, Muncie, IN 47304</td>
<td>765-254-5182</td>
</tr>
<tr>
<td>Amethyst House</td>
<td>645 North Walnut, Bloomington, IN 47402</td>
<td>812-336-3570</td>
</tr>
<tr>
<td>Cummins Behavioral Health Systems</td>
<td>6655 E US 36, Avon, IN 46123</td>
<td>317-272-3330</td>
</tr>
<tr>
<td>Aurora-Reid Hospital</td>
<td>1401 Chester Blvd., Richmond, IN 47374</td>
<td>800-232-3150</td>
</tr>
<tr>
<td>Dunn Mental Health Center Inc</td>
<td>6655 E US 36, Avon, IN 46123</td>
<td>317-272-3330</td>
</tr>
<tr>
<td>Center for Behavioral Health</td>
<td>645 S Rogers St, Bloomington, IN 47403</td>
<td>812-337-2215</td>
</tr>
<tr>
<td>Family Service of Central Indiana, Inc.</td>
<td>615 N. Alabama Ste. 320, Indianapolis, IN 46204</td>
<td>317-634-6341</td>
</tr>
<tr>
<td>Richard L. Roudebush Medical Center (VA)</td>
<td>1481 W. 10th St.118 - Psychiatry, Indianapolis, IN 46202</td>
<td>317-54-0000 ext. 5743</td>
</tr>
<tr>
<td>Gallahue Mental Health Center</td>
<td>6950 Hillsdale Ct, Indianapolis, IN 46250</td>
<td>317-621-5700</td>
</tr>
<tr>
<td>Midtown Narcotics Treatment Program</td>
<td>964 N. Pennsylvania St., Indianapolis, IN 46204</td>
<td>317-30-8800 ext.</td>
</tr>
<tr>
<td>Harbor Light Center</td>
<td>2400 N Tibbs Ave, Indianapolis, IN 46222</td>
<td>317-972-1450 ext. 332</td>
</tr>
</tbody>
</table>
Indianapolis Treatment Center, Inc.  
2626 E. 46th St.  
Indianapolis, IN 46205  
317-475-9066

Richmond Treatment Center, Inc.  
4265 South A St.  
Richmond, IN 47374  
765-962-8843

Midtown Narcotics Treatment Program  
694 N. Pennsylvania st.  
Indianapolis, IN 46204  
317-630-8800

Washington House  
2720 Culbertson St.  
Indianapolis, IN 46802  
317-432-8684

Richard L. Roudebush  
VA Medical Center  
1481 W. 10th St.  
Indianapolis, IN  
317-554-0000 ext. 5743
<table>
<thead>
<tr>
<th>Organization</th>
<th>Address</th>
<th>City, State, Zip</th>
<th>Phone</th>
</tr>
</thead>
<tbody>
<tr>
<td>Center For Behavioral Health of Indiana, Inc.</td>
<td>1414 Wells St.</td>
<td>Fort Wayne, IN 46802-2796</td>
<td>260-420-6010</td>
</tr>
<tr>
<td>Grant Blackford Mental Health Inc</td>
<td>505 Wabash Ave</td>
<td>Marion, IN 46952-2608</td>
<td>765-662-3971</td>
</tr>
<tr>
<td>Discovery House, Inc.</td>
<td>4195 Cleveland St.</td>
<td>Gary, IN 46408-2427</td>
<td>219-985-1668</td>
</tr>
<tr>
<td>Madison Center Inc</td>
<td>PO Box 80</td>
<td>South Bend, IN 46624-0080</td>
<td>574-234-0061</td>
</tr>
<tr>
<td>Metro Treatment of Gary, LP</td>
<td>d/b/a Semoran Treatment Center</td>
<td>Gary, IN 46403-3114</td>
<td>219-938-4651</td>
</tr>
<tr>
<td>Four County Counseling Center</td>
<td>1015 Michigan Ave</td>
<td>Logansport, IN 46947-1526</td>
<td>574-722-5151</td>
</tr>
<tr>
<td>Northeastern Center</td>
<td>PO Box 817</td>
<td>Kendallville, IN 46755-0817</td>
<td>260-347-2453</td>
</tr>
<tr>
<td>Geminus</td>
<td>8400 Louisiana St.</td>
<td>Merrillville, IN 46410</td>
<td>219-757-1905</td>
</tr>
<tr>
<td>PSI Services III, Inc.</td>
<td>3229 Broadway, Suite 207</td>
<td>Gary</td>
<td>219-884-0185</td>
</tr>
<tr>
<td>Grant Blackford Mental Health Inc</td>
<td>505 Wabash Ave</td>
<td>Marion, IN 46952-2608</td>
<td>765-662-3971</td>
</tr>
<tr>
<td>St Joseph Hospital &amp; Health Center of Kokomo</td>
<td>1907 W Sycamore</td>
<td>Kokomo</td>
<td>7654565900</td>
</tr>
</tbody>
</table>
St Margaret Mercy Healthcare Centers Inc
24 Joliet St
Dyer
219-865-2141

Swanson Center
450 St John Rd Ste 501
Michigan City
219-879-4621

Tri City
Comprehensive CMHC Inc
3903 Indianapolis Blvd
East Chicago
219-398-7050

Wabash Valley Hospital Inc
2900 N River Rd
West Lafayette
765-463-2555

Howard Community Hospital Psychiatric Services
PO Box 9011
Kokomo, IN 46904-9011
765-453-8555

Madison Center Inc
PO Box 80
South Bend, IN 46624-0080
574-234-0061

Metro Treatment of Gary, LP
d/b/a Semoran Treatment Center
8060 Melton Rd.
Gary, IN 46403-3114
219-938-4651

Northeastern Center
PO Box 817
Kendallville, IN 46755-0817
260-347-2453
## SOUTHERN INDIANA

<table>
<thead>
<tr>
<th>Name</th>
<th>Address</th>
<th>Phone</th>
</tr>
</thead>
<tbody>
<tr>
<td>Community Mental Health Center Inc</td>
<td>285 Bielby Rd, Lawrenceburg, IN 47025</td>
<td>812-537-1302</td>
</tr>
<tr>
<td>Samaritan Center</td>
<td>515 Bayou St, Vincennes, IN 47591</td>
<td>812-886-6800</td>
</tr>
<tr>
<td>East Indiana Treatment Center, Inc.</td>
<td>816 Rudolph Way, Lawrenceburg, IN 47205</td>
<td>812-537-1668 ext.</td>
</tr>
<tr>
<td>Southern Hills Counseling Center</td>
<td>PO Box 769, Jasper, IN 47547-0769</td>
<td>812-482-3020</td>
</tr>
<tr>
<td>Evansville Treatment Center</td>
<td>1510 W. Franklin St, Evansville, IN 47710</td>
<td>812-424-0223 ext.</td>
</tr>
<tr>
<td>Southern Indiana Treatment Center</td>
<td>1713 E. 10th St, Jeffersonville, IN 47130</td>
<td>812-283-4844 ext. 0000</td>
</tr>
<tr>
<td>East Indiana Treatment Center, Inc.</td>
<td>816 Rudolph Way, Lawrenceburg, IN 47205</td>
<td>812-537-1668 ext.</td>
</tr>
<tr>
<td>Southwestern Indiana Mental Health Center Inc</td>
<td>415 Mulberry St, Evansville, IN 47713</td>
<td>812-423-7791</td>
</tr>
<tr>
<td>Hamilton Center Inc</td>
<td>PO Box 4323, Terre Haute, IN 47804</td>
<td>812-231-8323</td>
</tr>
<tr>
<td>Lifespring Inc.</td>
<td>460 Spring Street, Jeffersonville, IN 47130</td>
<td>812-206-1234</td>
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</table>
HIV Care
And Services
Bloomington Hospital Positive Link
333 E. Miller Dr.
Bloomington, IN 47401
812-353-3250; 800-245-0261

LifeCare Program of Clarian Health
1633 N. Capitol Ave. Ste. 700
Indianapolis, IN 46202
317-962-3724

Center for Mental Health – Central
PO Box 304
Elwood, IN 46036
765-552-5009; 866-264-2020

Wishard Health Services
1050 Wishard Blvd. RHC Room 229
Indianapolis, IN 46202
317-630-7645

Center For Mental Health – Southeast
1401 Chester Blvd.
Jenkins Hall, Room 310
Richmond, IN 47374
765-983-7913

Concord Center Association
1310 S. Meridian St.
Indianapolis, IN 46225
317-637-4376

Comprehensive Mental Health
Services, Inc.
240 N. Tillotson Ave.
Muncie, IN 47304
765-288-1928

Damien Center
1350 N. Pennsylvania St.
Indianapolis, IN 46202
317-632-4376
NORTHERN INDIANA

AIDS Ministries/AIDS Assist
201 S. William Street
South Bend, IN 46601
South Bend, IN 46601
574-234-2870; 800-388-2437

Imani Unidad
914 Lincolnway West
South Bend, IN 46616
574-288-2887

AIDS Task Force of Northeast Indiana
525 Oxford Street
Ft. Wayne, IN 46806
260-744-1144; 800-417-3085

Ft. Wayne/Allen County Health Dept.
One E. Main St.
Ft. Wayne, IN 46802
260-449-3021

Aliveness Project of Northwest Indiana
PO Box 64568
Gary, IN 46401
219-985-6170; 800-489-1561

Center For Mental Health
12 N. 9th St.
Lafayette, IN 47901
765-742-4481

Elkhart County Health Department
608 Oakland Ave.
Elkhart, IN 46516
812-523-2127
SOUTHERN INDIANA

AIDS Resource Group
of Evansville
201 NW 4th Street Ste. B-7
Evansville, IN 47708
812-421-0059; 800-423-6255

Clark County Health Department
1403 Spring St. Ste. 200
Jeffersonville, IN 47130
812-288-2706; 1-800-828-5624

Area VII Agency on Aging
and the Disabled
PO Box 359
Terre Haute, IN 47808
812-238-1561; 800-489-1561

Matthew 25 AIDS Services
411 Letcher St.
Henderson, KY 42420
270-826-0200
Drug Assistance Programs And Relevant Websites
DRUG ASSISTANCE PROGRAMS

Roche Pharmaceuticals
Drug Name Pegasys
1-(877) 737-2797
Website: http://www.pegasysinfo.net

Schering-Plough Pharmaceuticals
Drug Name Peg-Intron
1-(800) 521-7157
Website: http://www.pegintron.com

RELEVANT WEBSITES

Centers for Disease Control Division of Viral Hepatitis
Resources Offered
Free Educational Materials; Publications; Training; Surveillance Information

Hepatitis Foundation International
http://www.hepfi.org/
Resources Offered
Hepatitis Information; News and Research; Educational Resources

Immunization Action Coalition
http://www.immunize.org/
Resources Offered
Vaccine Information; Vaccine Rules; Vaccine Screening Information; Hepatitis Test Result Interpretation

Harm Reduction Coalition
http://www.harmreduction.org/
Resources Offered
Harm Reduction Programs and Information; Publications; Resources

American Liver Foundation
http://www.liverfoundation.org/
Resources Offered
Quick Fact Sheets; Local Chapter Information; Support Group Development Information; Clinical Trials

VA National Hepatitis C Program
http://www.hepatitis.va.gov/
Resources Offered
Treatment guidelines, in-depth information, clinician tools, journal articles, patient handouts; Basic information on hepatitis C

HCV Advocate
http://www.hcvadvocate.org/
Resources Offered
Hepatitis C Information; Clinical Trials; News Updates
INDIANA
HEPATITIS C
SUPPORT GROUPS
CENTRAL INDIANA

A.R.K.
2020 N. Girls School Rd.
Indianapolis, IN 46214
rickeymikee@aol.com
Chapel Rock Christian Church
3rd Saturday of the Month 12:00 PM

S.T.A.R.S. VA Medical Center
1481 W. 10TH St.
Indianapolis, IN 46202
Phyllis.Baker@med.va.gov
VA Medical Center Room C-1192
1st and 3rd Tuesday of the Month
9:00 AM - 10:30 PM

NORTHERN INDIANA

DeKalb County Support Group
824 Martz Dr.
Auburn, IN 46706
ladytrucker62@netzero.com
DeKalb Medical Arts Building Room 303
3rd Friday of the Month 6:00 PM

SOUTHERN INDIANA

E.A.S.E. Hep C
415 W. Columbia St.
Evansville, IN
easehepc@aol.com
Heart Group Auditorium Deaconess Hospital
2nd Monday of the Month 6:00 PM
Hepatitis A: is a liver disease caused by the hepatitis A virus (HAV). Hepatitis A can affect anyone. In the United States, hepatitis A can occur in situations ranging from isolated cases of disease to widespread epidemics.

Hepatitis B: is a serious disease caused by a virus that attacks the liver. The virus, which is called hepatitis B virus (HBV), can cause lifelong infection, cirrhosis (scarring) of the liver, liver cancer, liver failure, and death.

Hepatitis C: is a liver disease caused by the hepatitis C virus (HCV), which is found in the blood of persons who have the disease. HCV is spread by contact with the blood of an infected person.

Hepatitis D: is a liver disease caused by the hepatitis D virus (HDV), a defective virus that needs the hepatitis B virus to exist. Hepatitis D virus (HDV) is found in the blood of persons infected with the virus.

Hepatitis E: is a liver disease caused by the hepatitis E virus (HEV) transmitted in much the same way as hepatitis A virus. Hepatitis E, however, does not occur often in the United States.
### Hepatitis A Virus Fact Sheet

<table>
<thead>
<tr>
<th><strong>SIGNS &amp; SYMPTOMS</strong></th>
<th>Adults will have signs and symptoms more often than children.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Jaundice</td>
</tr>
<tr>
<td></td>
<td>• Fatigue</td>
</tr>
<tr>
<td></td>
<td>• Abdominal pain</td>
</tr>
<tr>
<td></td>
<td>• Loss of appetite</td>
</tr>
<tr>
<td></td>
<td>• Nausea</td>
</tr>
<tr>
<td></td>
<td>• Diarrhea</td>
</tr>
<tr>
<td></td>
<td>• Fever</td>
</tr>
</tbody>
</table>

| **CAUSE**            | • Hepatitis A virus (HAV)                                   |

| **LONG-TERM EFFECTS**| • There is no chronic (long-term) infection.                 |
|                      | • Once you have had hepatitis A you cannot get it again.     |
|                      | • About 15% of people infected with HAV will have prolonged or relapsing symptoms over a 6-9 month period. |

| **TRANSMISSION**     | • HAV is found in the stool (feces) of persons with hepatitis A. |
|                      | • HAV is usually spread from person to person by putting something in the mouth (even though it may look clean) that has been contaminated with the stool of a person with hepatitis A. |

| **PERSONS AT RISK OF INFECTION** | • Household contacts of infected persons                     |
|                                  | • Sex contacts of infected persons                           |
|                                  | • Persons, especially children, living in areas with increased rates of hepatitis A during the baseline period from 1987-1997. (view map) |
|                                  | • Persons traveling to countries where hepatitis A is common (view map) |
|                                  | • Men who have sex with men                                  |
|                                  | • Injecting and non-injecting drug users                     |

| **VACCINE RECOMMENDATIONS** | Vaccine is recommended for the following persons 2 years of age and older: |
|                            | • Travelers to areas with increased rates of hepatitis A (view map) |
|                            | • Men who have sex with men                                   |
|                            | • Injecting and non-injecting drug users                      |
|                            | • Persons with clotting-factor disorders (e.g. hemophilia)    |
|                            | • Persons with chronic liver disease                          |
|                            | • Children living in areas with increased rates of hepatitis A during the baseline period from 1987-1997. (view map) |
# Hepatitis B Virus – Fact Sheet

<table>
<thead>
<tr>
<th>SIGNS &amp; SYMPTOMS</th>
<th>About 30% of persons have no signs or symptoms. Signs and symptoms are less common in children than adults.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Jaundice</td>
</tr>
<tr>
<td></td>
<td>• Fatigue</td>
</tr>
<tr>
<td></td>
<td>• Abdominal pain</td>
</tr>
<tr>
<td></td>
<td>• Loss of appetite</td>
</tr>
<tr>
<td></td>
<td>• Nausea, vomiting</td>
</tr>
<tr>
<td></td>
<td>• Joint pain</td>
</tr>
</tbody>
</table>

| CAUSE            | • Hepatitis B virus (HBV)                                                                              |

<table>
<thead>
<tr>
<th>LONG-TERM EFFECTS WITHOUT VACCINATION</th>
<th>Chronic infection occurs in:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• 90% of infants infected at birth</td>
</tr>
<tr>
<td></td>
<td>• 30% of children infected at age 1 - 5 years</td>
</tr>
<tr>
<td></td>
<td>• 6% of persons infected after age 5 years</td>
</tr>
</tbody>
</table>

Death from chronic liver disease occurs in:

- 15-25% of chronically infected persons

<table>
<thead>
<tr>
<th>TRANSMISSION</th>
<th>Occurs when blood or body fluids from an infected person enters the body of a person who is not immune.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HBV is spread through having sex with an infected person without using a condom (the efficacy of latex condoms in preventing infection with HBV is unknown, but their proper use may reduce transmission), by sharing drugs, needles, or &quot;works&quot; when &quot;shooting&quot; drugs, through needlesticks or sharps exposures on the job, or from an infected mother to her baby during birth. Persons at risk for HBV infection might also be at risk for infection with hepatitis C virus (HCV) or HIV.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>RISK GROUPS</th>
<th>Persons with multiple sex partners or diagnosis of a sexually transmitted disease</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Men who have sex with men</td>
</tr>
<tr>
<td></td>
<td>Sex contacts of infected persons</td>
</tr>
<tr>
<td></td>
<td>Injection drug users</td>
</tr>
<tr>
<td></td>
<td>Household contacts of chronically infected persons</td>
</tr>
<tr>
<td></td>
<td>Infants born to infected mothers</td>
</tr>
<tr>
<td></td>
<td>Infants/children of immigrants from areas with high rates of HBV infection (view map)</td>
</tr>
<tr>
<td></td>
<td>Health care and public safety workers</td>
</tr>
<tr>
<td></td>
<td>Hemodialysis patients</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PREVENTION</th>
<th>Hepatitis B vaccine is the best protection.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>If you are having sex, but not with one steady partner, use latex condoms correctly and every time you have sex. The efficacy of latex condoms in preventing infection with HBV is unknown, but their proper use may reduce transmission.</td>
</tr>
<tr>
<td></td>
<td>If you are pregnant, you should get a blood test for hepatitis B; Infants born to HBV-infected mothers should be given HBIG (hepatitis B immune globulin) and vaccine within 12 hours after birth.</td>
</tr>
</tbody>
</table>
| **VACCINE RECOMMENDATIONS** | • Hepatitis B vaccine available since 1982  
• Routine vaccination of 0-18 year olds  
• Vaccination of risk groups of all ages. |
| **TREATMENT & MEDICAL MANAGEMENT** | • HBV infected persons should be evaluated by their doctor for liver disease.  
• Adefovir dipivoxil, alpha interferon, and lamivudine are three drugs licensed for the treatment of persons with chronic hepatitis B.  
• Pregnant women should not use these drugs.  
• Drinking alcohol can make your liver disease worse. |
| **TRENDS & STATISTICS** | • Number of new infections per year has declined from an average of 260,000 in the 1980s to about 73,000 in 2003.  
• Highest rate of disease occurs in 20-49-year-olds.  
• Greatest decline has happened among children and adolescents due to routine hepatitis B vaccination.  
• Estimated 1.25 million chronically infected Americans, of whom 20-30% acquired their infection in childhood. |
Hepatitis B Vaccine – Fact Sheet

First Anti-cancer Vaccine

- Hepatitis B vaccine prevents hepatitis B disease and its serious consequences like hepatocellular carcinoma (liver cancer). Therefore, this is the first anti-cancer vaccine.

Safe and Effective

- Medical, scientific and public health communities strongly endorse using hepatitis B vaccine as a safe and effective way to prevent disease and death.
- Scientific data show that hepatitis B vaccines are very safe for infants, children, and adults.
- There is no confirmed evidence, which indicates that hepatitis B vaccine can cause chronic illnesses.
- To assure a high standard of safety with vaccines, several federal agencies continually assess and research possible or potential health effects that could be associated with vaccines.

Vaccine Schedule

- National Immunization Program, CDC 2005 recommends that all infants should receive the first dose of HepB vaccine soon after birth and before hospital discharge; the first dose may also be administered by age 2 months if the mother is hepatitis B surface antigen (HbsAg) negative. The second dose should be administered at least 4 weeks after the first dose, except for combination vaccines, which cannot be administered before 6 weeks. The third dose should be given at least 16 weeks after the first dose and at least 8 weeks after the second dose.
- If the vaccination series is interrupted after the first dose, the second dose should be administered as soon as possible. The second and third doses should be separated by an interval of at least 2 months. If only the third dose is delayed, it should be administered when convenient.

Booster Doses

- Current data show that vaccine-induced hepatitis B surface antibody (anti-HBs) levels may decline over time; however, immune memory (anamnestic anti-HBs response) remains intact indefinitely following immunization. Persons with declining antibody levels are still protected against clinical illness and chronic disease.
- For health care workers with normal immune status who have demonstrated an anti-HBs response following vaccination, booster doses of vaccine are not recommended nor are periodic anti-HBs testing.

Post-vaccination Testing

- After routine vaccination of infants, children, adolescents, or adults post-vaccination testing for adequate antibody response is not necessary.
- Post-vaccination testing IS recommended for persons whose medical management will depend on knowledge of their immune status.
This includes persons who:

- are immunocompromised (e.g., hemodialysis patients)
- received the vaccine in the buttock
- are infants born to HBsAg (hepatitis B surface antigen)-positive mothers
- are healthcare workers who have contact with blood
- are sex partners of persons with chronic hepatitis B virus infection
- Post-vaccination testing should be completed 1-2 months after the third vaccine dose for results to be meaningful. A protective antibody response is 10 or more milliinternational units (≥10mIU/mL).

**Adverse Events**

- Case reports of unusual illnesses following vaccines are most often related to other causes and not related to a vaccine. Whenever large numbers of vaccines are given, some adverse events will occur coincidentally after vaccination and be falsely attributed to the vaccine.
- Anyone believing they have had a possible reaction or adverse health effect from a vaccine should report it to their health care provider. The Vaccine Adverse Events Reporting System (1-800-822-7967) receives reports from health care providers and others about vaccine side effects.
Hepatitis C Virus – Fact Sheet

<table>
<thead>
<tr>
<th>SIGNS &amp; SYMPTOMS</th>
<th>80% of persons have no signs or symptoms.</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Jaundice</td>
<td>• Abdominal pain</td>
</tr>
<tr>
<td>• Fatigue</td>
<td>• Loss of appetite</td>
</tr>
<tr>
<td>• Dark urine</td>
<td>• Nausea</td>
</tr>
</tbody>
</table>

| CAUSE            | • Hepatitis C virus (HCV)                 |

| LONG-TERM EFFECTS | • Chronic infection: 55%-85% of infected persons  |
|                  | • Chronic liver disease: 70% of chronically infected persons  |
|                  | • Deaths from chronic liver disease: 1%-5% of infected persons may die  |
|                  | • Leading indication for liver transplant     |

| TRANSMISSION     | • Occurs when blood or body fluids from an infected person enters the body of a person who is not infected.  |
|                 | • HCV is spread through sharing needles or "works" when "shooting" drugs, through needlesticks or sharps exposures on the job, or from an infected mother to her baby during birth.  |

Persons at risk for HCV infection might also be at risk for infection with hepatitis B virus (HBV) or HIV.

Recommendations for Testing Based on Risk for HCV Infection

<table>
<thead>
<tr>
<th>PERSONS</th>
<th>RISK OF INFECTION</th>
<th>TESTING RECOMMENDED?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Injecting drug users</td>
<td>High</td>
<td>Yes</td>
</tr>
<tr>
<td>Recipients of clotting factors made before 1987</td>
<td>High</td>
<td>Yes</td>
</tr>
<tr>
<td>Hemodialysis patients</td>
<td>Intermediate</td>
<td>Yes</td>
</tr>
<tr>
<td>Recipients of blood and/or solid organs before 1992</td>
<td>Intermediate</td>
<td>Yes</td>
</tr>
<tr>
<td>People with undiagnosed liver problems</td>
<td>Intermediate</td>
<td>Yes</td>
</tr>
<tr>
<td>Infants born to infected mothers</td>
<td>Intermediate</td>
<td>After 12-18 mos. old</td>
</tr>
<tr>
<td>Healthcare/public safety workers</td>
<td>Intermediate</td>
<td>Only after known exposure</td>
</tr>
<tr>
<td>People having sex with multiple partners</td>
<td>Low</td>
<td>No*</td>
</tr>
<tr>
<td>People having sex with an infected steady partner</td>
<td>Low</td>
<td>No*</td>
</tr>
</tbody>
</table>

*Anyone who wants to get tested should ask their doctor.
<table>
<thead>
<tr>
<th>PREVENTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>• There is no vaccine to prevent hepatitis C.</td>
</tr>
<tr>
<td>• Do not shoot drugs; if you shoot drugs, stop and get into a treatment</td>
</tr>
<tr>
<td>program; if you can't stop, never share needles, syringes, water, or</td>
</tr>
<tr>
<td>&quot;works&quot;, and get vaccinated against hepatitis A &amp; B.</td>
</tr>
<tr>
<td>• Do not share personal care items that might have blood on them (razors,</td>
</tr>
<tr>
<td>toothbrushes).</td>
</tr>
<tr>
<td>• If you are a health care or public safety worker, always follow routine</td>
</tr>
<tr>
<td>barrier precautions and safely handle needles and other sharps; get</td>
</tr>
<tr>
<td>vaccinated against hepatitis B.</td>
</tr>
<tr>
<td>• Consider the risks if you are thinking about getting a tattoo or body</td>
</tr>
<tr>
<td>piercing. You might get infected if the tools have someone else's blood</td>
</tr>
<tr>
<td>on them or if the artist or piercer does not follow good health</td>
</tr>
<tr>
<td>practices.</td>
</tr>
<tr>
<td>• HCV can be spread by sex, but this is rare. If you are having sex with</td>
</tr>
<tr>
<td>more than one steady sex partner, use latex condoms* correctly and</td>
</tr>
<tr>
<td>every time to prevent the spread of sexually transmitted diseases. You</td>
</tr>
<tr>
<td>should also get vaccinated against hepatitis B.</td>
</tr>
<tr>
<td>• If you are HCV positive, do not donate blood, organs, or tissue.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>TREATMENT &amp; MEDICAL MANAGEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>• HCV positive persons should be evaluated by their doctor for liver</td>
</tr>
<tr>
<td>disease.</td>
</tr>
<tr>
<td>• Interferon and ribavirin are two drugs licensed for the treatment of</td>
</tr>
<tr>
<td>persons with chronic hepatitis C.</td>
</tr>
<tr>
<td>• Interferon can be taken alone or in combination with ribavirin.</td>
</tr>
<tr>
<td>Combination therapy, using pegylated interferon and ribavirin, is</td>
</tr>
<tr>
<td>currently the treatment of choice.</td>
</tr>
<tr>
<td>• Combination therapy can get rid of the virus in up to 5 out of 10</td>
</tr>
<tr>
<td>persons for genotype 1 and in up to 8 out of 10 persons for genotype 2</td>
</tr>
<tr>
<td>and 3.</td>
</tr>
<tr>
<td>• Drinking alcohol can make your liver disease worse.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>STATISTICS &amp; TRENDS</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Number of new infections per year has declined from an average of</td>
</tr>
<tr>
<td>240,000 in the 1980s to about 25,000 in 2001.</td>
</tr>
<tr>
<td>• Most infections are due to illegal injection drug use.</td>
</tr>
<tr>
<td>• Transfusion-associated cases occurred prior to blood donor screening;</td>
</tr>
<tr>
<td>now occurs in less than one per million transfused unit of blood.</td>
</tr>
<tr>
<td>• Estimated 3.9 million (1.8%) Americans have been infected with HCV,</td>
</tr>
<tr>
<td>of whom 2.7 million are chronically infected.</td>
</tr>
</tbody>
</table>
## Hepatitis D Virus – Fact Sheet

### SIGNS & SYMPTOMS
- jaundice
- fatigue
- abdominal pain
- loss of appetite
- nausea, vomiting
- joint pain
- dark (tea colored) urine

### CAUSE
- Hepatitis D virus (HDV)

### LONG-TERM EFFECTS WITHOUT VACCINATION
- HDV can be acquired either as
  - a co-infection (occurs simultaneously) with hepatitis B virus (HBV) or
  - as a superinfection in persons with existing chronic HBV infection.
- HBV-HDV co-infection:
  - may have more severe acute disease and a higher risk (2%-20%) of developing acute liver failure compared with those infected with HBV alone
- HBV-HDV superinfection
  - chronic HBV carriers who acquire HDV superinfection usually develop chronic HDV infection
    - progression to cirrhosis is believed to be more common with HBV/HDV chronic infections

### TRANSMISSION
- Occurs when blood or body fluids from an infected person enters the body of a person who is not immune.
- HBV is spread through having sex with an infected person without using a condom (the efficacy of latex condoms in preventing infection with HBV is unknown, but their proper use may reduce transmission);
- By sharing drugs, needles, or "works" when "shooting" drugs;
- Through needlesticks or sharps exposures on the job; or
- From an infected mother to her baby during birth.

### RISK GROUPS
- Injection drug users
- Men who have sex with men
- Hemodialysis patients
- Sex contacts of infected persons
- Health care and public safety workers
- Infants born to infected mothers (very rare)

### PREVENTION
- Hepatitis B vaccination
- HBV-HDV co-infection
  - pre- or post-exposure prophylaxis (hepatitis B immune globulin or vaccine) to prevent HBV infection
- HBV-HDV superinfection
  - education to reduce risk behaviors among persons with chronic HBV infection

### VACCINE RECOMMENDATIONS
- Hepatitis B vaccine should be given to prevent HBV/HDV co-infection

### TREATMENT & MEDICAL
- Acute HDV infection
| MANAGEMENT     | • Supportive care  
|               | • Chronic HDV infection  
|               |   • interferon-alfa  
|               |   • liver transplant  
| TRENDS & STATISTICS | • Routine surveillance data are not available.  |
## Hepatitis E Virus – Fact Sheet

<table>
<thead>
<tr>
<th>SIGNS &amp; SYMPTOMS</th>
<th>Highest attack rate among persons aged 15-40 years</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>jaundice</td>
</tr>
<tr>
<td></td>
<td>fatigue</td>
</tr>
<tr>
<td></td>
<td>abdominal pain</td>
</tr>
<tr>
<td></td>
<td>loss of appetite</td>
</tr>
<tr>
<td></td>
<td>nausea, vomiting</td>
</tr>
<tr>
<td></td>
<td>dark (tea colored) urine</td>
</tr>
</tbody>
</table>

| CAUSE            | Hepatitis E virus (HEV)                           |

| LONG-TERM EFFECTS WITHOUT VACCINATION | There is no chronic (long-term) infection |
|                                      | Hepatitis E is more severe among pregnant women, especially in third trimester |

| TRANSMISSION | HEV is found in the stool (feces) of persons and animals with hepatitis E. |
|             | HEV is spread by eating or drinking contaminated food or water. |
|             | Transmission from person to person occurs less commonly than with hepatitis A virus |
|             | Most outbreaks in developing countries have been associated with contaminated drinking water. |

| RISK GROUPS | Travelers to developing countries, particularly in South Asia and North Africa |
|            | Rare cases have occurred in the United States among persons with no history of travel to endemic countries |

| PREVENTION | Always wash your hands with soap and water after using the bathroom, changing a diaper, and before preparing and eating food |
|           | Avoid drinking water (and beverages with ice) of unknown purity, uncooked shellfish, and uncooked fruits or vegetables that are not peeled or prepared by the traveler. |

| TREATMENT & MEDICAL MANAGEMENT | Treatment is supportive |

| TRENDS & STATISTICS | Hepatitis E remains uncommon in the United States. Routine surveillance data are not available. |
HIV & HCV COINFECTION

HIV, the human immunodeficiency virus is a virus that causes acquired immunodeficiency syndrome (AIDS).

HIV and HCV (Hepatitis C Virus) share an important mode of transmission: both viruses are blood borne pathogens and are transmitted by blood-to-blood contact. Because of this, professionals working in HIV and HCV prevention share a concern for the risk behaviors associated with injecting drug use (IDU) such as syringe sharing. While it is generally accepted that bleaching syringes will sufficiently clean syringes of HIV, this has not been proven to be the case with HCV. This has resulted in harm reduction programs promoting a total abstinence of any type of paraphernalia sharing including syringes and works such as cookers and cotton. Syringe exchange clients are now being told to use a new syringe every time they inject rather than to clean them.

Presently HIV and HCV are both tested for in the blood supply. This means that the public can be assured that blood transfusions were without HIV after 1985. HCV testing in the blood supply did not start until 1990. In both cases, there are people who became infected with a virus from transfusions, transplants and other blood products before testing of the blood supply began. It is estimated that 10% of people living with HCV were previously infected from transfusions and blood products. This is not the case with HIV. People infected with HIV from transfusions, transplants and other blood products before testing of the blood supply was significantly less and in many cases are no longer living.

The effect of co-infection with HIV and HCV is not well understood. It is estimated that up to 40% of people infected with HIV are also infected with HCV. Many co-infected people have the additional public health issues of substance abuse and histories of incarceration, which complicate their access to services and adherence to consistent treatment.

Treating HIV in People Living with HCV

The United Public Health Service and Infectious Disease Society of America recommend that all HIV infected people be tested for HCV. It is generally recommended that HIV be under control or treated first before treating HCV. The good news is that HIV can be successfully treated in individuals co-infected with HIV and HCV.

It appears that people infected with HIV and HCV may have more rapid progression of their liver disease. Having HCV may increase the incidence of hepatotoxicity or liver damage from HIV treatments therapies. The adverse side effects of HIV medical therapies may be more severe and accelerate the distress on already impaired liver function. Most people with HCV can tolerate HIV medications as long as they are closely monitored for potential liver toxicity. While the potential for HIV medications to produce liver damage is very real as demonstrated with an increase in liver enzyme leveling and HCV viral load, these laboratory results will usually stabilize over time. HIV medications do not seem to have a direct effect on HCV. However, some experts believe that when HIV is under control, HCV disease progression is slowed.*
On the other hand, it is unclear if HCV makes HIV worse. The introduction of combination antiretroviral therapy has greatly improved and extended the life for many people living with HIV. The majority of studies have not been able to correlate a more aggressive HIV disease progression to being infected with HCV.

**Treating HCV in the People Living with HIV**
Individuals with HIV who have been diagnosed with HCV should be evaluated and considered for HCV treatment. The same treatment guidelines for treating HCV can generally be applied to people living with HIV. However, HIV positive individuals with CD4 counts of less than 200, or a concurrent opportunistic illness, are not considered good candidates for HCV treatment, until the CD4 count goes up and/or the opportunistic illness is treated.

Studies have shown that people living with HIV and HCV will have similar response rates to HCV treatment as HCV positive individuals without HIV. Patients should be monitored closely for possible side effects associated with interferon and ribavirin.

Support groups for co-infected individuals are highly recommended due to the emotional complexities of living with these two life-threatening diseases. Additionally, support groups can be a good resource for information sharing since there is so much misinformation regarding these two diseases.

*Adapted from Hepatitis C Support Project, San Francisco. HIV/HCV Coinfection, What you need to know. HCV Advocate. August 2001; Version 1.1; Recommendations for Prevention and Control of Hepatitis C Virus (HCV) Infection and HCV-Related Chronic Disease October 16, 1998 / 47(RR19): 1-39*
FREQUENTLY ASKED QUESTIONS
Hepatitis A Virus

What is hepatitis A?
Hepatitis A is a liver disease caused by hepatitis A virus.

How is hepatitis A virus transmitted?
Hepatitis A virus is spread from person to person by putting something in the mouth that has been contaminated with the stool of a person with hepatitis A. This type of transmission is called "fecal-oral." For this reason, the virus is more easily spread in areas where there are poor sanitary conditions or where good personal hygiene is not observed.

Most infections result from contact with a household member or sex partner who has hepatitis A. Casual contact, as in the usual office, factory, or school setting, does not spread the virus.

What are the signs and symptoms of hepatitis A?
Persons with hepatitis A virus infection may not have any signs or symptoms of the disease. Older persons are more likely to have symptoms than children. If symptoms are present, they usually occur abruptly and may include fever, tiredness, loss of appetite, nausea, abdominal discomfort, dark urine, and jaundice (yellowing of the skin and eyes). Symptoms usually last less than 2 months; a few persons are ill for as long as 6 months. The average incubation period for hepatitis A is 28 days (range: 15–50 days).

How do you know if you have hepatitis A?
A blood test (IgM anti-HAV) is needed to diagnose hepatitis A. Talk to your doctor or someone from your local health department if you suspect that you have been exposed to hepatitis A or any type of viral hepatitis.

How can you prevent hepatitis A?
Always wash your hands after using the bathroom, changing a diaper, or before preparing or eating food.

Two products are used to prevent hepatitis A virus infection: immune globulin and hepatitis A vaccine.

1. Immune globulin is a preparation of antibodies that can be given before exposure for short-term protection against hepatitis A and for persons who have already been exposed to hepatitis A virus. Immune globulin must be given within 2 weeks after exposure to hepatitis A virus for maximum protection.
2. Hepatitis A vaccine has been licensed in the United States for use in persons 2 years of age and older. The vaccine is recommended (before exposure to hepatitis A virus) for persons who are more likely to get hepatitis A virus infection or is more likely to get seriously ill if they do get hepatitis A. The vaccines currently licensed in the United States are HAVRIX® (manufactured by GlaxoSmithKline) and VAQTA® (manufactured by Merck & Co., Inc).
Hepatitis A Vaccine

**What are the dosages and schedules for hepatitis A vaccines?**

<table>
<thead>
<tr>
<th>Vaccinee's age (years)</th>
<th>Dose (EL.U.)²</th>
<th>Volume (mL)</th>
<th>No. doses</th>
<th>Schedule (mos.)³</th>
</tr>
</thead>
<tbody>
<tr>
<td>2-18</td>
<td>720</td>
<td>0.5</td>
<td>2</td>
<td>0,6-12</td>
</tr>
<tr>
<td>&gt;18</td>
<td>1,440</td>
<td>1.0</td>
<td>2</td>
<td>0,6-12</td>
</tr>
</tbody>
</table>

Recommended dosages of HAVRIX®¹

<table>
<thead>
<tr>
<th>Vaccinee's age (years)</th>
<th>Dose (U)²</th>
<th>Volume (mL)</th>
<th>No. doses</th>
<th>Schedule (mos.)³</th>
</tr>
</thead>
<tbody>
<tr>
<td>2-18</td>
<td>25</td>
<td>0.5</td>
<td>2</td>
<td>0,6-18</td>
</tr>
<tr>
<td>&gt;18</td>
<td>50</td>
<td>1.0</td>
<td>2</td>
<td>0,6-12</td>
</tr>
</tbody>
</table>

1 Hepatitis A vaccine, inactivated, Merck & Co., Inc.
² Units.
³ 0 months represents timing of the initial dose; subsequent numbers represent months after the initial dose.

**Can a patient receive the first dose of hepatitis A vaccine from one manufacturer and the second (last) dose from another manufacturer?**
Yes. Although studies have not been done to look at this issue, there is no reason to believe that this would be a problem.

**What should be done if the second (last) dose of hepatitis A vaccine is delayed?**
The second dose should be administered as soon as possible. There is no need to repeat the first dose.

**Can other vaccines be given at the same time that hepatitis A vaccine is given?**
Yes. Hepatitis B, diphtheria, poliovirus (oral and inactivated), tetanus, oral typhoid, cholera, Japanese encephalitis, rabies, yellow fever vaccine or immune globulin can be given at the same time that hepatitis A vaccine is given, but at a different injection site.
Is hepatitis A vaccine safe?
Yes, hepatitis A vaccine has an excellent safety profile. No serious adverse events have been attributed definitively to hepatitis A vaccine. Soreness at the injection site is the most frequently reported side effect.

Any adverse event suspected to be associated with hepatitis A vaccination should be reported to the Vaccine Adverse Events Reporting System (VAERS). VAERS forms can be obtained by calling 1-800-822-7967.

How is hepatitis A vaccines made?
There is no live virus in hepatitis A vaccines. The virus is inactivated during production of the vaccines, similar to Salk-type inactivated polio vaccine.

How long does hepatitis A vaccine protect you?
Although data on long-term protection are limited, estimates based on modeling techniques suggest that protection will last for at least 20 years.

When are persons protected after receiving hepatitis A vaccine?
Protection against hepatitis A begins four weeks after the first dose of hepatitis A vaccine.

Can hepatitis A vaccine be given after exposure to hepatitis A virus?
No, hepatitis A vaccine is not licensed for use after exposure to hepatitis A virus. In this situation, immune globulin should be used.

Should pre-vaccination testing be done?
Pre-vaccination testing is done only in specific instances to control cost (e.g., persons who were likely to have had hepatitis A in the past). This includes persons who were born in countries with high levels of hepatitis A virus infection, elderly persons, and persons who have clotting factor disorders and may have received factor concentrates in the past.

Should post-vaccination testing be done?
No.

Can hepatitis A vaccine be given during pregnancy or lactation?
We don't know for sure, but because vaccine is produced from inactivated hepatitis A virus, the theoretical risk to the developing fetus is expected to be low. The risk associated with vaccination, however, should be weighed against the risk for hepatitis A in women who may be at high risk for exposure to hepatitis A virus.

Can hepatitis A vaccine be given to immunocompromised persons? (e.g., persons on hemodialysis or persons with HIV/AIDS)
Yes.
What is Twinrix®?

It is a combined hepatitis A and hepatitis B vaccine for use in persons aged 18 years and older. Primary vaccination consists of three doses, given on a 0-, 1-, and 6-month schedule, the same schedule as that used for hepatitis B vaccine alone.

Immune Globulin

What is immune globulin?
Immune globulin is a preparation of antibodies that can be given before exposure for short-term protection against hepatitis A and for persons who have already been exposed to hepatitis A virus. Immune globulin must be given within 2 weeks after exposure to hepatitis A virus for maximum protection.

Is immune globulin safe?
Yes. No instance of transmission of HIV (the virus that causes AIDS) or other viruses has been observed with the use of immune globulin administered by the intramuscular route. Immune globulin can be administered during pregnancy and breast-feeding.

WHO SHOULD GET VACCINATED AGAINST HEPATITIS A?

Hepatitis A vaccination provides protection before one is exposed to hepatitis A virus. Hepatitis A vaccination is recommended for the following groups who are at increased risk for infection and for any person wishing to obtain immunity.

Persons traveling to or working in countries that have high or intermediate rates of hepatitis A.
All susceptible persons traveling to or working in countries that have high or intermediate rates of hepatitis A virus should be vaccinated or receive immune globulin before traveling. Persons from developed countries who travel to developing countries are at high risk for hepatitis A. Such persons include tourists, military personnel, missionaries, and others who work or study abroad in countries that have high or intermediate levels of hepatitis A. The risk for hepatitis A exists even for travelers to urban areas, those who stay in luxury hotels, and those who report that they have good hygiene and that they are careful about what they drink and eat.

Children in states, counties, and communities where rates of hepatitis A were/are at least twice the national average during the baseline period of 1987-1997.
Children living in states, counties, and communities where rates of hepatitis A are at least twice the national average (≥ 20 cases/1000,000) in baseline period should be routinely vaccinated beginning at 2 years of age. High rates of hepatitis A have been found in these populations, both in urban and rural...
settings. In addition, to effectively prevent epidemics of hepatitis A, vaccination of previously unvaccinated older children is recommended within 5 years of initiation of routine childhood vaccination programs. Although rates differ among areas, available data indicate that a reasonable cutoff age in many areas is 10-15 years of age because older persons have often already had hepatitis A. Vaccination of children before they enter school should receive highest priority, followed by vaccination of older children who have not been vaccinated.

Men who have sex with men
Sexually active men (both adolescents and adults) who have sex with men should be vaccinated.

Hepatitis A outbreaks among men who have sex with men have been reported frequently. Recent outbreaks have occurred in urban areas in the United States, Canada, and Australia.

Illegal-drug users
Vaccination is recommended for injecting and non-injecting illegal-drug users.

Persons who have occupational risk for infection
Persons who work with hepatitis A virus-infected primates or with hepatitis A virus in a research laboratory setting should be vaccinated. No other groups have been shown to be at increased risk for hepatitis A virus infection because of occupational exposure.

Outbreaks of hepatitis A have been reported among persons working with non-human primates that are susceptible to hepatitis A virus infection, including several Old World and New World species. Primates that were infected were those that had been born in the wild, not those that had been born and raised in captivity.

Persons who have chronic liver disease
Persons with chronic liver disease who have never had hepatitis A should be vaccinated, as there is a higher rate of fulminant (rapid onset of liver failure, often leading to death) hepatitis A among persons with chronic liver disease. Persons who are either awaiting or have received liver transplants also should be vaccinated.

Persons who have clotting-factor disorders
Persons who have never had hepatitis A and who are administered clotting-factor concentrates, especially solvent detergent-treated preparations, should be given hepatitis A vaccine.

All persons with hemophilia (Factor VIII, Factor IX) who receive replacement therapy should be vaccinated because there appears to be an increased risk of transmission from clotting-factor concentrates that are not heat inactivated.
WHICH GROUPS DO NOT ROUTINELEY NEED HEPATITIS A VACCINE?

**Food service workers**
Food borne hepatitis A outbreaks are relatively uncommon in the United States; however, when they occur, intensive public health efforts are required for their control.

Although persons who work as food handlers have a critical role in common-source food borne outbreaks, they are not at increased risk for hepatitis A because of their occupation. Consideration may be given to vaccination of employees who work in areas where community-wide outbreaks are occurring and where state and local health authorities or private employers determine that such vaccination is cost-effective.

**Sewerage workers**
In the United States, no work-related outbreaks of hepatitis A have been reported among workers exposed to sewage.

**Health-care workers**
Health-care workers are not at increased risk for hepatitis A. If a patient with hepatitis A is admitted to the hospital, routine infection control precautions will prevent transmission to hospital staff.

**Children under 2 years of age**
Because of the limited experience with hepatitis A vaccination among children under 2 years of age, the vaccine is not currently licensed for this age group.

**Day-care attendees**
The frequency of outbreaks of hepatitis A is not high enough in this setting to warrant routine hepatitis A vaccination. In some communities, however, day-care centers play a role in sustaining community-wide outbreaks. In this situation, consideration should be given to adding hepatitis A vaccine to the prevention plan for children and staff in the involved center(s).

**Residents of institutions for developmentally disabled persons**
Historically, hepatitis A virus infections were common among persons with developmental disabilities living in institutions. Currently, the occurrence of hepatitis A virus infections have diminished.
Who should receive protection against hepatitis A before travel?
All susceptible persons traveling to or working in countries that have high or intermediate rates of hepatitis A should be vaccinated or receive immune globulin before traveling. Persons from developed countries who travel to developing countries are at high risk for hepatitis A. Such persons include tourists, military personnel, missionaries, and others who work or study abroad in countries that have high or intermediate levels of hepatitis A. The risk for hepatitis A exists even for travelers to urban areas, those who stay in luxury hotels, and those who report that they have good hygiene and that they are careful about what they drink and eat.

How soon before travel should the first dose of hepatitis A vaccine be given?
For optimal protection, the first dose of hepatitis A vaccine should be given at least 4 weeks prior to travel. Check with your doctor about when the next dose is due.

What should be done if a person cannot receive hepatitis A vaccine?
Travelers who are allergic to a vaccine component or who elect not to receive vaccine should receive a single dose of immune globulin (0.02 mL/kg), which provides effective protection against hepatitis A virus infection for up to 3 months. Travelers whose travel period exceeds 2 months should be administered immune globulin at 0.06 mL/kg; administration must be repeated if the travel period exceeds 5 months.
If travel starts sooner than 4 weeks prior to the first vaccine dose, what should be done?
Because protection might not be optimal until 4 weeks after vaccination, persons traveling to a high-risk area less than 4 weeks after the initial dose of hepatitis A vaccine should also be given immune globulin (0.02 mL/kg), but at a different injection site. Therefore, the first dose of hepatitis A vaccine should be administered as soon as travel to a high-risk area is planned.

What should be done for travelers who are less than 2 years of age to protect them from hepatitis A virus infection?
Immune globulin is recommended for travelers less than 2 years of age because the vaccine is currently not licensed for use in this age group.

Hepatitis B Virus

What is hepatitis B?
Hepatitis B is caused by a virus that attacks the liver. The virus, which is called hepatitis B virus (HBV), can cause lifelong infection, cirrhosis (scarring) of the liver, liver cancer, liver failure, and death.

How do you know if you have hepatitis B?
Only a blood test can tell for sure.

How is HBV spread?
HBV is spread when blood or body fluids from an infected person enters the body of a person who is not infected. For example, HBV is spread through having sex with an infected person without using a condom (the efficacy of latex condoms in preventing infection with HBV is unknown, but their proper use might reduce transmission), by sharing drugs, needles, or "works" when "shooting" drugs, through needlesticks or sharps exposures on the job, or from an infected mother to her baby during birth.

Hepatitis B is not spread through food or water, sharing eating utensils, breastfeeding, hugging, kissing, coughing, sneezing or by casual contact.

What are the symptoms of hepatitis B?
Sometimes a person with HBV infection has no symptoms at all. The older you are, the more apt you are to have symptoms. You might be infected with HBV (and be spreading the virus) and not know it.

If you have symptoms, they might include:

- Yellow skin or yellowing of the whites of your eyes (jaundice)
- Tiredness
- Loss of appetite
- Nausea
- Abdominal discomfort
• Dark urine
• Clay-colored bowel movements
• Joint pain

**What are the risk factors for hepatitis B?**
You are at increased risk of HBV infection if you:

- Have sex with someone infected with HBV
- Have sex with more than one partner
- Shoot drugs
- Are a man and have sex with a man
- Live in the same house with someone who has chronic (long-term) HBV infection
- Have a job that involves contact with human blood
- Are a client in a home for the developmentally disabled
- Have hemophilia
- Travel to areas where hepatitis B is common ([view map](#))

One out of 20 people in the United States will get infected with HBV some time during their lives. Your risk is higher if your parents were born in Southeast Asia, Africa, and the Amazon Basin in South America, the Pacific Islands, or the Middle East.

**Is there a cure for hepatitis B?**
There are no medications available for recently acquired (acute) HBV infection. Hepatitis B vaccine is available for the prevention of HBV infection. There are antiviral drugs available for the treatment of chronic HBV infection.

**How common is HBV infection in the U.S.?**
In 2003, an estimated 73,000 people were infected with HBV. People of all ages get hepatitis B and about 5,000 die per year of sickness caused by HBV.

**If you are pregnant, should you worry about hepatitis B?**
Yes, you should get a blood test to check for HBV infection early in your pregnancy. This test is called hepatitis B surface antigen (HBsAg). If you test HBsAg-negative early in pregnancy, but continue behaviors that put you at risk for HBV infection (e.g., multiple sex partners, injection drug use), you should be retested for HBsAg close to delivery. If your HBsAg test is positive, this means you are infected with HBV and can give the virus to your baby. Babies who get HBV at birth might develop chronic HBV infection that can lead to cirrhosis of the liver or liver cancer.

If your blood test is positive, your baby should receive the first dose of hepatitis B vaccine, along with another shot, hepatitis B immune globulin (called HBIG), at birth. The second dose of vaccine should be given at aged 1-2 months and the third dose at aged 6 months (but not before aged 24 weeks).

**Can I donate blood if I have had any type of viral hepatitis?**
If you had any type of viral hepatitis since aged 11 years, you are not eligible to donate blood. In addition, if you ever tested positive for hepatitis B or hepatitis C, at any age, you are not eligible to donate, even if you were never sick or jaundiced from the infection.

**How long can HBV survive outside the body?**
HBV can survive outside the body at least 7 days and still be capable of causing infection.
Hepatitis B Vaccine Information

Who should get vaccinated?

- All babies, at birth
- All children 0-18 years of age who have not been vaccinated
- People of any age whose behavior or job puts them at high risk for HBV infection (see risk factors under general information)

What are the dosages and schedules for hepatitis B vaccines?
The vaccination schedule most often used for adults and children has been three intramuscular injections, the second and third administered 1 and 6 months after the first. Recombivax HB® has been approved as a two dose schedule for aged 11-15 years. Engerix-B® has also been approved as a four dose accelerated schedule.

Can you receive one dose of hepatitis B vaccine from one manufacturer and the other doses from another manufacturer?
Yes. The immune response when one or two doses of a vaccine produced by one manufacturer are followed by subsequent doses from a different manufacturer has been shown to be comparable with that resulting from a full course of vaccination from one manufacturer.

What should be done if there is an interruption between doses of hepatitis B vaccine?
If the vaccination series is interrupted after the first dose, the second dose should be administered as soon as possible. The second and third doses should be separated by an interval of at least 2 months. If only the third dose is delayed, it should be administered when convenient.

Can other vaccines be given at the same time that hepatitis B vaccine is given?
Yes. When hepatitis B vaccine has been administered at the same time as other vaccines, no interference with the antibody response of the other vaccines has been demonstrated.

Are hepatitis B vaccines safe?
Yes. Hepatitis B vaccines have been shown to be safe when administered to both adults and children. Over 4 million adults have been vaccinated in the U.S., and at least that many children have received hepatitis B vaccine worldwide.

How long does hepatitis B vaccine protect you?
Long-term studies of healthy adults and children, who have developed adequate antibody to hepatitis B surface antigen (anti-HBs), indicate that immunologic memory remains intact for at least 15 years and confers protection against clinical illness and chronic HBV infection, even though anti-HBs levels might become low or decline below detectable levels.

Can hepatitis B vaccine be given after exposure to HBV?
Yes. After a person has been exposed to HBV, appropriate treatment, given in an appropriate time frame, can effectively prevent infection. The mainstay of post exposure immunoprophylaxis is hepatitis B vaccine, but in some settings the addition of HBIG will provide some increase in protection.

Should pre-vaccination testing be done?
Pre-vaccination testing is not routinely recommended. The decision to do pre-vaccination testing is usually based on cost. To avoid vaccinating persons who have already had or have HBV infection, testing for prior infection should be considered for adults in risk groups with high rates of HBV infection (e.g., injecting drug users, men who have sex with men and household contacts of persons with chronic HBV infection). Pre-vaccination testing is not indicated for immunization programs for children or adolescents because of the low rate of HBV infection and the relatively low cost of vaccine.

Who should get post-vaccination testing?
Testing for immunity is advised only for persons whose subsequent clinical management depends on knowledge of their immune status (e.g., infants born to HBsAg-positive mothers, immune compromised persons, healthcare workers, and sex partners of persons with chronic HBV infection).

When should post-vaccination testing be done?
When necessary, post-vaccination testing, using the anti-HBs test, should be performed 1 to 2 months after completion of the vaccine series – EXCEPT for post-vaccination testing of infants born to HBsAg-positive mothers. Testing of these infants should be performed 3 to 9 months after the completion of the vaccination series.

For how long is hepatitis B vaccine effective?
Long-term studies of healthy adults and children indicate that hepatitis B vaccine protects against chronic HBV infection for at least 15 years, even though antibody levels might decline below detectable levels.

Are booster doses of hepatitis B vaccine needed routinely?
No, booster doses of hepatitis B vaccine are not recommended routinely for persons who are not immune compromised. Data show that vaccine-induced anti-HBs levels might decline over time; however, immune memory remains intact indefinitely following immunization. Immune competent people with declining antibody levels are still protected against clinical illness and chronic disease.

Can hepatitis B vaccine be given during pregnancy or when breastfeeding?
Yes, neither pregnancy nor breastfeeding should be considered a contraindication to vaccination of women. On the basis of limited experience, there is no apparent risk of adverse effects to developing fetuses when hepatitis B vaccine is administered to pregnant women. The vaccine contains noninfectious HBsAg particles and should cause no risk to the fetus. HBV infection affecting a pregnant woman might result in severe disease for the mother and chronic HBV infection for the newborn.

Can hepatitis B vaccine be given to immune compromised people? (e.g., people on hemodialysis or people with HIV/AIDS)
Yes, however larger vaccine doses or an increased number of doses are required to induce protective antibody in a high proportion of hemodialysis patients and might also be necessary for other immune compromised people (e.g., those who take immunosuppressive drugs or who have AIDS). For immune compromised people, it is important that post vaccination testing, using the anti-HBs test, be done 1-2 months after the last dose of vaccine to check that the vaccine worked. In addition, immune compromised people need periodic testing and possibly booster doses of hepatitis B vaccine to assure that anti-HBs is still adequate.

What is the rationale for recommending the hepatitis B vaccination of children and other groups mentioned above?

- In the United States, hepatitis B virus (HBV) transmission occurs in all age groups and a comprehensive strategy is needed to provide widespread immunity and to effectively prevent HBV-related chronic liver disease. Beginning in the late 1980s, the Advisory Committee on Immunization Practices to the U.S. Public Health Service
developed a comprehensive strategy to eliminate HBV transmission in the United States. This strategy includes 1) screening of all pregnant woman for hepatitis B surface antigen (HbsAg) and providing postexposure immunoprophylaxis beginning at birth to infants of HbsAg-positive mothers; 2) routine infant vaccination; 3) catch-up vaccination of previously unvaccinated children and adolescents; and 4) vaccination of adults in high risk groups.

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**If You Are Living With Chronic Hepatitis B**

**What does the term "chronic hepatitis B" mean?**

Chronic infection with HBV means that you have a long-term HBV infection; your body did not get rid of the virus when you were first infected with HBV. Two percent to 6% of people over aged 5 years; 30% of children aged 1-5 years; and up to 90% of infants develop chronic infection. People with chronic infection can infect others and are at increased risk of serious liver disease including cirrhosis and liver cancer. In the United States, an estimated 1.25 million people are chronically infected with HBV.

**What is the treatment for chronic hepatitis B?**

There are three drugs licensed for the treatment of people with chronic hepatitis B: Adefovir dipivoxil, alpha interferon, and lamivudine. These drugs work in up to 40% of people.

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**Traveler's Health Information**

**What is the risk of getting HBV infection while traveling in other countries?**

The risk of HBV infection for international travelers is generally low, except for certain travelers in countries where the prevalence of chronic HBV infection is high or intermediate (see map).

Factors to consider in assessing risk include 1) the prevalence of chronic HBV infection in the local population, 2) the extent of direct contact with blood or other body fluids or of sex contact with potentially infected people, and 3) the duration of travel.

Modes of HBV transmission in areas with high or intermediate prevalence of chronic HBV infection that are important for travelers to consider are contaminated injection and other equipment used for health care-related procedures and blood transfusions from unscreened donors. However, unprotected sex and sharing illegal drug injection equipment are also risks for HBV infection in these areas.
1. **High (8% or more) in**
   - Alaska and extreme northern Canada and southern Greenland;
   - A band crossing South America, including northern Chile, southern Colombia, extreme southern Venezuela, northwestern Brazil and northern Bolivia;
   - Parts of Eastern Europe, including Moldova, Bulgaria, Georgia, Armenia, and Azerbaijan
   - All of Africa except northern Morocco, Algeria, Tunisia, Libya, and Egypt;
   - Saudi Arabia, Lebanon, Israel, and Jordan;
   - Turkmenistan, Uzbekistan, Kazakhstan, Kyrgyzstan, Tajikistan, Mongolia;
   - South Asia and Southeast Asia.

2. **Intermediate (2% to 7%) in**
   - Parts of Central America, including Guatemala, El Salvador, Honduras, Haiti and the Dominican Republic;
   - Parts of South America, including northern Venezuela, Guyana and Suriname, and central and southern Brazil;
   - Northern Africa, including northern Morocco, Algeria, Tunisia, Libya, and Egypt;
   - The Middle East except Saudi Arabia, Lebanon, Israel, and Jordan;
   - Southern sections of Eastern Europe except Moldova, Bulgaria, Georgia, Armenia, and Azerbaijan; and Poland;
   - Portions of Western Europe including Italy, Sardinia, Spain, Portugal;
   - Russia;
   - Japan.

3. **Low (under 2%) in all areas not already listed**
## Serology

### How do I interpret serological lab results?

#### Interpretation of the Hepatitis B Panel

<table>
<thead>
<tr>
<th>Tests</th>
<th>Results</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>HBsAg</td>
<td>negative</td>
<td>Susceptible</td>
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<tr>
<td>anti-HBc</td>
<td>negative</td>
<td></td>
</tr>
<tr>
<td>anti-HBs</td>
<td>negative</td>
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</tr>
<tr>
<td>HBsAg</td>
<td>negative</td>
<td>Immune due to natural infection</td>
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<tr>
<td>anti-HBc</td>
<td>positive</td>
<td></td>
</tr>
<tr>
<td>anti-HBs</td>
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<td>HBsAg</td>
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<td>HBsAg</td>
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<td>IgM anti-HBc</td>
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<td>anti-HBs</td>
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<td>HBsAg</td>
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<td>Chronically infected</td>
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<tr>
<td>anti-HBc</td>
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<td>anti-HBs</td>
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<tr>
<td>HBsAg</td>
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<td>Four interpretations possible</td>
</tr>
<tr>
<td>anti-HBc</td>
<td>positive</td>
<td></td>
</tr>
<tr>
<td>anti-HBs</td>
<td>negative</td>
<td></td>
</tr>
</tbody>
</table>

* Four Interpretations:

1. Might be recovering from acute HBV infection.
2. Might be distantly immune and test not sensitive enough to detect very low level of anti-HBs in serum.
3. Might be susceptible with a false positive anti-HBc.
4. Might be undetectable level of HBsAg present in the serum and the person is actually chronically infected.

### What do the different abbreviations on the lab results mean?

- **Hepatitis B Surface Antigen (HBsAg):** A serologic marker on the surface of HBV. It can be detected in high levels in serum during acute or chronic hepatitis. The presence of HBsAg indicates that the person is infectious. The body normally produces antibodies to HBsAg as part of the normal immune response to infection.

- **Hepatitis B Surface Antibody (anti-HBs):** The presence of anti-HBs is generally interpreted as indicating recovery and immunity from HBV infection. Anti-HBs also develops in a person who has been successfully vaccinated against hepatitis B.

- **Hepatitis B Core Antibody (anti-HBc):** Appears at the onset of symptoms in acute hepatitis B and persists for life. The presence of anti-HBc indicates previous or ongoing infection with hepatitis B virus (HBV) in an undefined time frame.
Hepatitis B e Antigen (HBeAg): A secreted product of the nucleocapsid gene of HBV and is found in serum during acute and chronic hepatitis B. Its presence indicates that the virus is replicating and the infected individual has high levels of HBV.

Hepatitis B e Antibody (HBeAb or anti-HBe): produced by the immune system temporarily during acute HBV infection or consistently during or after a burst in viral replication. Spontaneous conversion from e antigen to e antibody (a change known as seroconversion) is a predictor of long-term clearance of HBV in patients undergoing antiviral therapy and indicates lower levels of HBV.

Hepatitis B Immune Globulin (HBIG): A product available for prophylaxis against HBV infection. HBIG is prepared from plasma containing high titers of anti-HBs and provides short-term protection (3 - 6 months).

Hepatitis C Virus

What is hepatitis C?
Hepatitis C is a liver disease caused by the hepatitis C virus (HCV), which is found in the blood of persons who have this disease. HCV is spread by contact with the blood of an infected person.

Is there a vaccine for the prevention of HCV infection?
No.

What blood tests are available to check for hepatitis C?
There are several blood tests that can be done to determine if you have been infected with HCV. Your doctor may order just one or a combination of these tests. The following are the types of tests your doctor may order and the purpose for each:

a) Anti-HCV (antibody to HCV)
- EIA (enzyme immunoassay) or CIA (enhanced chemiluminescence immunoassay)
  Test is usually done first. If positive, it should be confirmed
- RIBA (recombinant immunoblot assay)
  A supplemental test used to confirm a positive EIA test

Anti-HCV does not tell whether the infection is new (acute), chronic (long-term) or is no longer present.

b) Qualitative tests to detect presence or absence of virus (HCV RNA)

c) Quantitative tests to detect amount (titer) of virus (HCV RNA)

A single positive PCR test indicates infection with HCV. A single negative test does not prove that a person is not infected. Virus may be present in the blood and just not found by PCR. Also, a person infected in the past who has
recovered may have a negative test. When hepatitis C is suspected and PCR is negative, PCR should be repeated.

Can you have a "false positive" anti-HCV test result?
Yes. A false positive test means the test looks as if it is positive, but it is really negative. This happens more often in persons who have a low risk for the disease for which they are being tested. For example, false positive anti-HCV tests happen more often in persons such as blood donors who are at low risk for hepatitis C. Therefore, it is important to confirm a positive anti-HCV test with a supplemental test as most false positive anti-HCV tests are reported as negative on supplemental testing. Click here for more information on Guidelines for Laboratory Testing and Result Reporting of Antibody to Hepatitis C Virus.

Can you have a "false negative" anti-HCV test result?
Yes. Persons with early infection may not as yet have developed antibody levels high enough that the test can measure. In addition, some persons may lack the (immune) response necessary for the test to work well. In these persons, research-based tests such as PCR may be considered.

How long after exposure to HCV does it take to test positive for anti-HCV?
Anti-HCV can be found in 7 out of 10 persons when symptoms begin and in about 9 out of 10 persons within 3 months after symptoms begin. However, it is important to note that many persons who have hepatitis C have no symptoms.

How long after exposure to HCV does it take to test positive with PCR?
It is possible to find HCV within 1 to 2 weeks after being infected with the virus.

Who should get tested for hepatitis C?

- persons who ever injected illegal drugs, including those who injected once or a few times many years ago
- persons who were treated for clotting problems with a blood product made before 1987 when more advanced methods for manufacturing the products were developed
- persons who were notified that they received blood from a donor who later tested positive for hepatitis C
- persons who received a blood transfusion or solid organ transplant before July 1992 when better testing of blood donors became available
- long-term hemodialysis patients
- persons who have signs or symptoms of liver disease (e.g., abnormal liver enzyme tests)
- healthcare workers after exposures (e.g., needle sticks or splashes to the eye) to HCV-positive blood on the job
- children born to HCV-positive women

What is the next step if you have a confirmed positive anti-HCV test?
Measure the level of ALT (alanine aminotransferase, a liver enzyme) in the
blood. An elevated ALT indicates inflammation of the liver and you should be checked further for chronic (long-term) liver disease and possible treatment. The evaluation should be done by a healthcare professional familiar with chronic hepatitis C.

**Can you have a normal liver enzyme (e.g., ALT) level and still have chronic hepatitis C?**

Yes. It is common for persons with chronic hepatitis C to have a liver enzyme level that goes up and down, with periodic returns to normal or near normal. Some persons have a liver enzyme level that is normal for over a year but they still have chronic liver disease. If the liver enzyme level is normal, persons should have their enzyme level re-checked several times over a 6 to 12 month period. If the liver enzyme level remains normal, your doctor may check it less frequently, such as once a year.

**How is HCV spread from one person to another?**

**How could a person have gotten hepatitis C?**

HCV is spread primarily by direct contact with human blood. For example, you may have gotten infected with HCV if:

- you ever injected or snorted street drugs, as the needles and/or other drug "works" used to prepare or inject the drug(s) may have had someone else's blood that contained HCV on them.
- you received blood, blood products, or solid organs from a donor whose blood contained HCV.
- you were ever on long-term kidney dialysis as you may have unknowingly shared supplies/equipment that had someone else's blood on them.
- you were ever a healthcare worker and had frequent contact with blood on the job, especially accidental needlesticks.
- your mother had hepatitis C at the time she gave birth to you. During the birth her blood may have gotten into your body.
- you ever had sex with a person infected with HCV.
- you lived with someone who was infected with HCV and shared items such as razors or toothbrushes that might have had his/her blood on them.

**How long can HCV live outside the body and transmit infection?** Recent studies suggest that HCV may survive on environmental surfaces at room temperature at least 16 hours, but no longer than 4 days.

**Is there any evidence that HCV has been spread during medical or dental procedures done in the United States?** Medical and dental procedures done in the United States generally do not pose a risk for the spread of HCV. However, there have been a few situations in which HCV has been spread between patients when supplies or equipment were shared between them.
Can HCV be spread by sexual activity?
Yes, but this does not occur very often. See section below on counseling for more information on hepatitis C and sexual activity.

Can HCV be spread by oral sex?
There is no evidence that HCV has been spread by oral sex. See section on counseling for more information on hepatitis C and sexual activity.

Can HCV be spread within a household?
Yes, but this does not occur very often. If HCV is spread within a household, it is most likely due to direct exposure to the blood of an infected household member.

Since more advanced tests have been developed for use in blood banks, what is the chance now that a person can get HCV infection from transfused blood or blood products?
Less than 1 chance per million units transfused.

Pregnancy and Breast feeding

Should pregnant women be routinely tested for anti-HCV? No. Pregnant women have no greater risk of being infected with HCV than non-pregnant women. If pregnant women have risk factors for hepatitis C, they should be tested for anti-HCV.

What is the risk that HCV infected women will spread HCV to their newborn infants?
About 5 out of every 100 infants born to HCV infected women become infected. This occurs at the time of birth, and there is no treatment that can prevent this from happening. Most infants infected with HCV at the time of birth have no symptoms and do well during childhood. More studies are needed to find out if these children will have problems from the infection as they grow older. There are no licensed treatments or guidelines for the treatment of infants or children infected with HCV. Children with elevated ALT (liver enzyme) levels should be referred for evaluation to a specialist familiar with the management of children with HCV-related disease.

Should a woman with hepatitis C be advised against breast-feeding?
No. There is no evidence that breast-feeding spreads HCV. HCV-positive mothers should consider abstaining from breast-feeding if their nipples are cracked or bleeding.

When should babies born to mothers with hepatitis C be tested to see if they were infected at birth?
Children should not be tested for anti-HCV before 18 months of age as anti-HCV from the mother might last until this age. If diagnosis is desired prior to 18 months of age, testing for HCV RNA could be performed at or after an infant's first well-child visit at age 1-2 months. HCV RNA testing should then
be repeated at a subsequent visit independent of the initial HCV RNA test result.

## Counseling

**How can persons infected with HCV prevent spreading HCV to others?**

- Do not donate blood, body organs, other tissue, or semen.
- Do not share personal items that might have your blood on them, such as toothbrushes, dental appliances, nail-grooming equipment or razors.
- Cover your cuts and skin sores to keep from spreading HCV.

**How can a person protect themselves from getting hepatitis C and other diseases spread by contact with human blood?**

- Don't ever shoot or snort drugs. If you shoot drugs, stop and get into a treatment program. If you can't stop, never reuse or share syringes, water, or drug works, and get vaccinated against hepatitis A and hepatitis B.
- Do not share toothbrushes, razors, or other personal care articles. They might have blood on them.
- If you are a healthcare worker, always follow routine barrier precautions and safely handle needles and other sharps. Get vaccinated against hepatitis B.
- Consider the health risks if you are thinking about getting a tattoo or body piercing: You can get infected if:
  - the tools that are used have someone else's blood on them.
  - the artist or piercer doesn't follow good health practices, such as washing hands and using disposable gloves.

HCV can be spread by sex, but this does not occur very often. If you are having sex, but not with one steady partner:

- You and your partners can get other diseases spread by having sex (e.g., AIDS, hepatitis B, gonorrhea or chlamydia).
- You should use latex condoms correctly and every time. The efficacy of latex condoms in preventing infection with HCV is unknown, but their proper use may reduce transmission.
- You should get vaccinated against hepatitis B.

**Should patients with hepatitis C change their sexual practices if they have only one long-term steady sex partner?** No. There is a very low chance of spreading HCV to that partner through sexual activity. If you want to lower the small chance of spreading HCV to your sex partner, you may decide to use barrier precautions such as latex condoms. The efficacy of latex condoms...
in preventing infection with HCV is unknown, but their proper use may reduce transmission. Ask your doctor about having your sex partner tested.

**What can persons with HCV infection do to protect their liver?**

- Stop using alcohol.
- See your doctor regularly.
- Don't start any new medicines or use over-the-counter, herbal, and other medicines without a physician's knowledge.
- Get vaccinated against hepatitis A if liver damage is present.

**What other information should patients with hepatitis C be aware of?**

- HCV is not spread by sneezing, hugging, coughing, food or water, sharing eating utensils or drinking glasses, or casual contact.
- Persons should not be excluded from work, school, play, child-care or other settings on the basis of their HCV infection status.
- Involvement with a support group may help patients cope with hepatitis C.

**Should persons with chronic hepatitis C be vaccinated against hepatitis B?**

If persons are in risk groups for whom hepatitis B vaccine is recommended, they should be vaccinated.

### Long-term Consequences of HCV Infection

**What are the chances of persons with HCV infection developing long term infection, chronic liver disease, cirrhosis, liver cancer, or dying as a result of hepatitis C?**

Of every 100 persons infected with HCV about:

- 55-85 of persons might develop long-term infection
- 70 persons might develop chronic liver disease
- 5-20 persons might develop cirrhosis over a period of 20 to 30 years
- 1-5 of persons might die from the consequences of long term infection (liver cancer or cirrhosis)

Hepatitis C is a leading indication for liver transplants.

**Do medical conditions outside the liver occur in persons with chronic hepatitis C?**

A small percentage of persons with chronic hepatitis C develop medical conditions outside the liver (this is called extrahepatic). These conditions are thought to occur due to the body's natural immune system fighting against itself. Such conditions include: glomerulonephritis, essential mixed cryoglobulinemia, and porphyria cutanea tarda.
Management and Treatment of Chronic Hepatitis C

When might a specialist (gastroenterologist, infectious disease physician, or hepatologist) be consulted in the management of HCV-infected persons?
A referral to or consultation with a specialist for further evaluation and possible treatment may be considered if a person is anti-HCV positive and has elevated liver enzyme levels. Any physician who manages a person with hepatitis C should be knowledgeable and current on all aspects of the care of a person with hepatitis C.

What is the treatment for chronic hepatitis C?
Combination therapy with pegylated interferon and ribavirin is the treatment of choice resulting in sustained response rates of 40%-80%. (up to 50% for patients infected with the most common genotype found in the U.S. [genotype 1] and up to 80% for patients infected with genotypes 2 or 3). Interferon monotherapy is generally reserved for patients in whom ribavirin is contraindicated. Ribavirin, when used alone, does not work. Combination therapy using interferon and ribavirin is now FDA approved for the use in children aged 3-17 years.

What are the side effects of interferon therapy?
Most persons have flu-like symptoms (fever, chills, headache, muscle and joint aches, fast heart rate) early in treatment, but these lessen with continued treatment. Later side effects may include tiredness, hair loss, low blood count, trouble with thinking, moodiness, and depression. Severe side effects are rare (seen in less than 2 out of 100 persons). These include thyroid disease, depression with suicidal thoughts, seizures, acute heart or kidney failure, eye and lung problems, hearing loss, and blood infection. Although rare, deaths have occurred due to liver failure or blood infection, mostly in persons with cirrhosis. An important side effect of interferon is worsening of liver disease with treatment, which can be severe and even fatal. Interferon dosage must be reduced in up to 40 out of 100 persons because of severity of side effects, and treatment must be stopped in up to 15 out of 100 persons. Pregnant women should not be treated with interferon.

What are the side effects of combination (ribavirin + interferon) treatment?
In addition to the side effects due to interferon described above, ribavirin can cause serious anemia (low red blood cell count) and can be a serious problem for persons with conditions that cause anemia, such as kidney failure. In these persons, combination therapy should be avoided or attempts should be made to correct the anemia. Anemia caused by ribavirin can be life-threatening for persons with certain types of heart or blood vessel disease. Ribavirin causes birth defects and pregnancy should be avoided during treatment. Patients and their healthcare providers should carefully review the product manufacturer information prior to treatment.
Can anything be done to reduce symptoms or side effects due to antiviral treatment?
You should report what you are feeling to your doctor. Some side effects may be reduced by giving interferon at night or lowering the dosage of the drug. In addition, flu-like symptoms can be reduced by taking acetaminophen before treatment.

Can children receive interferon therapy for chronic hepatitis C?
The Food and Drug Administration has approved the use of the combination anti-viral therapy for the treatment of hepatitis C in children 3 to 17 years old. For details please refer to page 11 of AASLD Practice Guideline: Diagnosis, Treatment, and Management of Hepatitis C.

Genotype

What does the term genotype mean?
Genotype refers to the genetic make-up of an organism or a virus. There are at least 6 distinct HCV genotypes identified. Genotype 1 is the most common genotype seen in the United States.

Is it necessary to do genotyping when managing a person with chronic hepatitis C?
Yes, as there are 6 known genotypes and more than 50 subtypes of HCV, and genotype information is helpful in defining the epidemiology of hepatitis C. Knowing the genotype or serotype (genotype-specific antibodies) of HCV is helpful in making recommendations and counseling regarding therapy. Patients with genotypes 2 and 3 are almost three times more likely than patients with genotype 1 to respond to therapy with alpha interferon or the combination of alpha interferon and ribavirin. Furthermore, when using combination therapy, the recommended duration of treatment depends on the genotype. For patients with genotypes 2 and 3, a 24-week course of combination treatment is adequate, whereas for patients with genotype 1, a 48-week course is recommended. For these reasons, testing for HCV genotype is often clinically helpful. Once the genotype is identified, it need not be tested again; genotypes do not change during the course of infection.

Why do most persons remain infected?
Persons infected with HCV mount an antibody response to parts of the virus, but changes in the virus during infection result in changes that are not recognized by preexisting antibodies. This appears to be how the virus establishes and maintains long-lasting infection.

Can persons become infected with different genotypes?
Yes. Because of the ineffective immune response described above, prior infection does not protect against reinfection with the same or different genotypes of the virus. For the same reason, there is no effective pre- or postexposure prophylaxis (i.e., immune globulin) available.
**Hepatitis C and Healthcare Workers**

**What is the risk for HCV infection from a needle-stick exposure to HCV contaminated blood?**
After needle stick or sharps exposure to HCV positive blood, about 2 (1.8%) healthcare workers out of 100 will get infected with HCV (range 0%-10%).

**What are the recommendations for follow-up of healthcare workers after exposure to HCV positive blood?**
Anti-viral agents (e.g., interferon) or immune globulin should not be used for postexposure prophylaxis.

1. For the source, baseline testing for anti-HCV.
2. For the person exposed to an HCV-positive source, baseline and follow-up testing including baseline testing for anti-HCV and ALT activity; and follow-up testing for anti-HCV (e.g., at 4-6 months) and ALT activity. (If earlier diagnosis of HCV infection is desired, testing for HCV RNA may be performed at 4-6 weeks.)
3. Confirmation by supplemental anti-HCV testing of all anti-HCV results reported as positive by enzyme immunoassay.

**Should HCV-infected healthcare workers be restricted in their work?**
No, there are no recommendations to restrict a healthcare worker who is infected with HCV. The risk of transmission from an infected healthcare worker to a patient appears to be very low. As recommended for all healthcare workers, those who are HCV positive should follow strict aseptic technique and standard precautions, including appropriate use of hand washing, protective barriers, and care in the use and disposal of needles and other sharp instruments.

**HIV and Hepatitis C Virus Co-infection**

**Why should HIV-infected persons be concerned about co-infection with HCV?**
About one quarter of HIV-infected persons in the United States are also infected with hepatitis C virus (HCV). HCV is one of the most important causes of chronic liver disease in the United States and HCV infection progresses more rapidly to liver damage in HIV-infected persons. HCV infection may also impact the course and management of HIV infection. The latest U.S. Public Health Service/Infectious Diseases Society of America (USPHS/IDSA) guidelines recommend that all HIV-infected persons should be screened for HCV infection. Prevention of HCV infection for those not already infected and reducing chronic liver disease in those who are infected are important concerns for HIV-infected individuals and their health care providers.
Who is likely to have HIV-HCV co-infection?
The hepatitis C virus (HCV) is transmitted primarily by large or repeated direct percutaneous (i.e., passage through the skin by puncture) exposures to contaminated blood. Therefore, co-infection with HIV and HCV is common (50%-90%) among HIV-infected injection drug users (IDUs). Co-infection is also common among persons with hemophilia who received clotting factor concentrates before concentrates were effectively treated to inactivate both viruses (i.e., products made before 1987). The risk for acquiring infection through perinatal or sexual exposures is much lower for HCV than for HIV. For persons infected with HIV through sexual exposure (e.g., male-to-male sexual activity), co-infection with HCV is no more common than among similarly aged adults in the general population (3%-5%).

What are the effects of co-infection on disease progression of HCV and HIV?
Chronic HCV infection develops in 75%-85% of infected persons and leads to chronic liver disease in 70% of these chronically infected persons. HIV-HCV co-infection has been associated with higher titers of HCV, more rapid progression to HCV-related liver disease, and an increased risk for HCV-related cirrhosis (scarring) of the liver. Because of this, HCV infection has been viewed as an opportunistic infection in HIV-infected persons and was included in the 1999 USPHS/IDSA Guidelines for the Prevention of Opportunistic Infections in Persons Infected with Human Immunodeficiency Virus. It is not, however, considered an AIDS-defining illness. As highly active antiretroviral therapy (HAART) and prophylaxis of opportunistic infections increase the life span of persons living with HIV, HCV-related liver disease has become a major cause of hospital admissions and deaths among HIV-infected persons.
The effects of HCV co-infection on HIV disease progression are less certain. Some studies have suggested that infection with certain HCV genotypes is associated with more rapid progression to AIDS or death. However, the subject remains controversial. Since co-infected patients are living longer on HAART, more data are needed to determine if HCV infection influences the long-term natural history of HIV infection.

How can co-infection with HCV be prevented?
Persons living with HIV who are not already co-infected with HCV can adopt measures to prevent acquiring HCV. Such measures will also reduce the chance of transmitting their HIV infection to others.
Not injecting or stopping injection drug use would eliminate the chief route of HCV transmission; substance-abuse treatment and relapse-prevention programs should be recommended. If patients continue to inject, they should be counseled about safer injection practices; that is, to use new, sterile syringes every time they inject drugs and never reuse or share syringes, needles, water, or drug preparation equipment.
Toothbrushes, razors, and other personal care items that might be contaminated with blood should not be shared. Although there are no data from the United States indicating that tattooing and body piercing place persons at increased risk for HCV infection, these procedures may be a source for infection with any bloodborne pathogen if proper infection control practices are not followed.
Although consistent data are lacking regarding the extent to which sexual activity contributes to HCV transmission, persons having multiple sex partners are at risk for other sexually transmitted diseases (STDs) as well as for transmitting HIV to others. They should be counseled accordingly.
How should patients co-infected with HIV and HCV be managed?

General guidelines
Patients co-infected with HIV and HCV should be encouraged to adopt safe behaviors (as described in the previous section) to prevent transmission of HIV and HCV to others. Individuals with evidence of HCV infection should be given information about prevention of liver damage, undergo evaluation for chronic liver disease and, if indicated, be considered for treatment. Persons co-infected with HIV and HCV should be advised not to drink excessive amounts of alcohol. Avoiding alcohol altogether might be wise because the effects of even moderate or low amounts of alcohol (e.g., 12 oz. of beer, 5 oz. of wine or 1.5 oz. hard liquor per day) on disease progression are unknown. When appropriate, referral should be made to alcohol treatment and relapse-prevention programs. Because of possible effects on the liver, HCV-infected patients should consult with their health care professional before taking any new medicines, including over-the-counter, alternative or herbal medicines. Susceptible co-infected patients should receive hepatitis A vaccine because the risk for fulminant hepatitis associated with hepatitis A is increased in persons with chronic liver disease. Susceptible patients should receive hepatitis B vaccine because most HIV-infected persons are at risk for HBV infection. The vaccines appear safe for these patients and more than two-thirds of those vaccinated develop antibody responses. Pre-vaccination screening for antibodies against hepatitis A and hepatitis B in this high-prevalence population is generally cost-effective. Post-vaccination testing for hepatitis A is not recommended, but testing for antibody to hepatitis B surface antigen (anti-HBs) should be performed 1-2 months after completion of the primary series of hepatitis B vaccine. Persons who fail to respond should be revaccinated with up to three additional doses.

HAART has no significant effect on HCV. However, co-infected persons may be at increased risk for HAART-associated liver toxicity and should be closely monitored during antiretroviral therapy. Data suggest that the majority of these persons do not appear to develop significant and/or symptomatic hepatitis after initiation of antiretroviral therapy.

Treatment for HCV Infection
A Consensus Development Conference Panel convened by The National Institutes of Health in 1997 recommended antiviral therapy for patients with chronic hepatitis C who are at the greatest risk for progression to cirrhosis. These persons include anti-HCV positive patients with persistently elevated liver enzymes, detectable HCV RNA, and a liver biopsy that indicates either portal or bridging fibrosis or at least moderate degrees of inflammation and necrosis. Patients with less severe histological disease should be managed on an individual basis. In the United States, three different regimens have been approved as therapy for chronic hepatitis C in mono-infected patients: monotherapy with alpha interferon and combination therapy with alpha interferon and ribavirin. Among HIV-negative persons with chronic hepatitis C, combination therapy consistently yields higher rates of sustained virologic response than monotherapy. Viral genotypes 2 and 3 require a shorter course of treatment. However, viral genotype 1 is the most common among U.S. patients. Combination therapy is associated with more side effects than monotherapy, but, in most situations, it is preferable. At present, interferon monotherapy is reserved for patients who have contraindications to the use of ribavirin.

Studies thus far, although not extensive, have indicated that response rates in HIV-infected patients to alpha interferon monotherapy for HCV were lower than in non-HIV-infected patients. On February 25, 2005 one regimen became FDA approved for treatment of HCV in HIV co-infected patients, pegylated interferon alpha 2-a and ribavirin tablets. A 40% sustained viral rate was demonstrated in this population; 29% sustained viral rate in genotype 1 and 62% sustained viral rate in genotypes 2 and 3.
The decision to treat people co-infected with HIV and HCV must also take into consideration their concurrent medications and medical conditions. If CD4 counts are normal or minimally abnormal (> 400/ul), there is little difference in treatment success rates between those who are co-infected and those who are infected with HCV alone.

**Other Treatment Considerations**

Persons with chronic hepatitis C who continue to use alcohol are at risk for ongoing liver injury, and antiviral therapy may be ineffective. Therefore, strict abstinence from alcohol is recommended during antiviral therapy, and interferon should be given with caution to a patient who has only recently stopped alcohol abuse. Typically, a 6-month abstinence is recommended for alcohol abusers before starting therapy; such patients should be treated with the support and collaboration of alcohol abuse treatment programs.

Although there is limited experience with antiviral treatment for chronic hepatitis C of persons who are recovering from long-term injection drug use, there are concerns that interferon therapy could be associated with relapse into drug use, both because of its side effects and because it is administered by injection. There is even less experience with treatment of persons who are active injection drug users, and an additional concern for this group is the risk for reinfection with HCV. Although a 6-month abstinence before starting therapy also has been recommended for injection drug users, additional research is needed on the benefits and drawbacks of treating these patients. Regardless, when patients with past or continuing problems of substance abuse are being considered for treatment, such patients should be treated only in collaboration with substance abuse specialists or counselors. Patients can be successfully treated while on methadone maintenance treatment of addiction.

Because many co-infected patients have conditions or factors (such as major depression or active illicit drug or alcohol use) that may prevent or complicate antiviral therapy, treatment for chronic hepatitis C in HIV-infected patients should be coordinated by health care providers with experience in treating co-infected patients or in clinical trials. It is not known if maintenance therapy is needed after successful therapy, but patients should be counseled to avoid injection drug use and other behaviors that could lead to re-infection with HCV and should continue to abstain from alcohol.

**Infections in Infants and Children**

The average rate of HCV infection among infants born to women co-infected with HCV and HIV is 14% to 17%, higher than among infants born to women infected with HCV alone. Data are limited on the natural history of HCV infection in children, and antiviral drugs for chronic hepatitis C are not FDA-approved for use in children under aged 18 years. Therefore, children should be referred to a pediatric hepatologist or similar specialist for management and for determination for eligibility in clinical trials.
TESTING GUIDELINES
Testing Algorithm
# Reference for Interpretation of HCV Test Results

<table>
<thead>
<tr>
<th>If Your HCV Test Result Is:</th>
<th>Supplemental Test</th>
<th>Interpretation</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Anti-HCV Screening Test</strong></td>
<td><strong>RIBA†</strong> or <strong>HCV RNA</strong></td>
<td><strong>Anti-HCV</strong></td>
<td><strong>HCV Infection</strong></td>
</tr>
<tr>
<td>Negative</td>
<td>Not Needed</td>
<td>Negative</td>
<td>None</td>
</tr>
<tr>
<td>Positive</td>
<td>Not Done</td>
<td>Not Done</td>
<td>Not Known</td>
</tr>
<tr>
<td>Positive</td>
<td>Not Done</td>
<td>Negative</td>
<td>Not Known*</td>
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<td>Not Done</td>
<td>Positive</td>
</tr>
<tr>
<td>Positive</td>
<td>Positive</td>
<td>Not Done</td>
<td>Past/Current</td>
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<tr>
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<td>Negative</td>
<td>Current</td>
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<td>Positive/not done</td>
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</tr>
<tr>
<td><strong>Positive</strong></td>
<td>Indeterminate</td>
<td>Negative</td>
<td>None</td>
</tr>
</tbody>
</table>

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* EIA - enzyme immunoassay or CIA - enhanced chemiluminesence immunoassay
† Recombinant immunoblot assay, a more specific anti-HCV assay
* Single negative HCV RNA result cannot determine infection status as persons might have intermittent viremia.
§ Samples with high s/co ratios usually (>95%) confirm positive, but supplemental serologic testing was not performed. Less than 5 of every 100 might represent false-positives; more specific testing should be requested, if indicated.