

# **Akutní a chronické virové hepatitidy**

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# **Discovery of aetiological agents of VH**

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- **1965 Hepatitis B surface antigen (HBsAg) was isolated (Blumberg)**
- **1973 Hepatitis A virus (HAV) was identified (Feinstone)**
- **1977 Delta agent (HDV) was identified (Mario Rizzetto)**
- **1985 recombinant DNA technology**
- **1989 HCV was identified (Houghton)**
- **1990 HEV was identified (Balayan)**
- **1994 HFV, but existence of HFV as well as HF was not confirmed later**
- **1995 HGV was identified (later renamed on GBV-C)**

Původce	HAV = enterovirus 72	HBV	HCV
Čeleď (Family)	Picornaviridae	Hepadnaviridae	Flaviviridae
Rod (Genus)	Hepatovirus	Orthohepadnavirus	Hepacivirus
Velikost	27-28 nm	42 nm	30-60 nm
Genom	RNA	DNA	RNA
Počet genotypů	tři (I, II a III) se subtypy A a B pouze 1 sérotyp	9 (A až I), několik subgenotypů V Evropě nejčastěji A a C V Asii nejčastěji B a C	> 6 genotypů > 120 subtypů
Ink. doba (dny)	15-50	30-180	15-150
Přenos: - enterálně	ano	ne	ne
- krví	vzácně	ano	ano
- sexuálně	vzácně	ano	vzácně
- vertikálně	ne	ano	vzácně
Přechod do chronicity	ne	novor. > 90 %, děti do 5 let 25-50 %, dospělí: imunokompetentní < 5 %, imunokompromitovaní > 50 %	50 - 90 %
Laboratorní diagnostika akutní infekce	anti-HAV IgM (pozitivita také po akt. imunizaci)	HBsAg HBeAg anti-HBc IgM	anti-HCV HCV RNA
Fulminantní průběh (rozvoj akutního jaterního selhání do 8 týdnů)	0,1 %	0,1 - 1 %	krajně vzácně
Akt. a pas. imunizace	ano	ano	ne

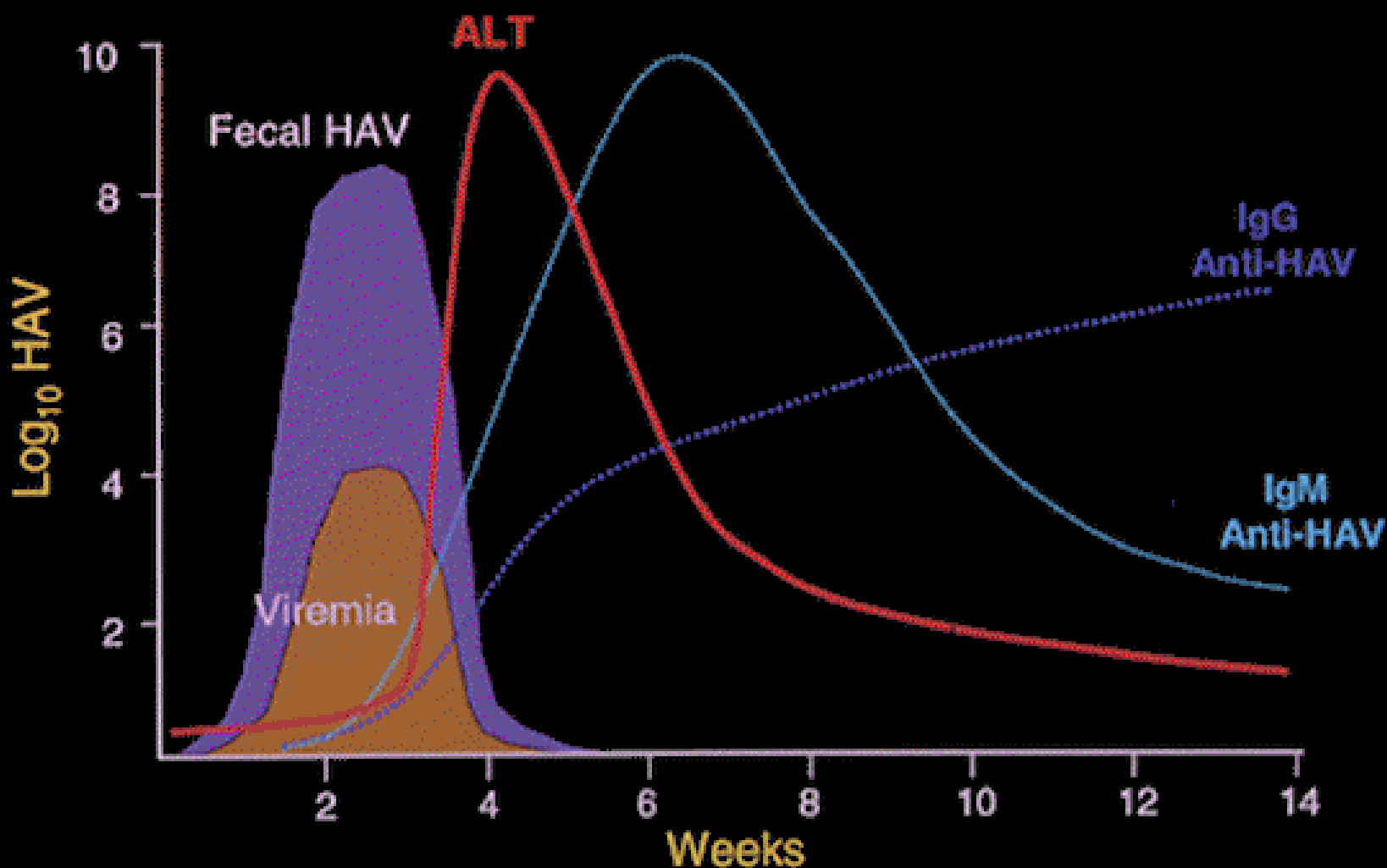
Původce	HDV (delta agens)	HEV swHEV (swine kmeny HEV)	HGV (nyní GBV-C) Nezpůsobuje hepatitidu
Čeleď (Family)		Hepeviridae	Flaviviridae
Rod (Genus)	Deltavirus	Orthohepevirus	
Velikost	35-37 nm	27-34 nm	
Genom	RNA	RNA	RNA
Počet genotypů (GT)	nejméně 8	Osm, ale GT1, 2, 3 a 4 jsou nejdůležitější	
Ink. doba (dny)	30-180	15-60	
Přenos: - enterálně	ne	ANO GT 1 a GT 2 kontaminovanou vodou GT 3 a GT 4 zoonotický přenos (vepři, kanci, jelenovití)	ne
- krví	ano	vzácně	ano
- sexuálně	ano	vzácně	
- vertikálně	ano	ano	
Přechod do chronicity	koinfekce < 5 % superinfekce 80-90 %	NE. Chron. průběh bývá u pac. po transplantacích, u malignit, u HIV pozit.	Inhibice HIV replikace - zpomalení progrese
Lab. diagnostika akutní infekce	anti-HDV IgM, HBsAg	anti-HEV IgM a/nebo HEV RNA (ze séra nebo stolice)	anti-HGV, HGV RNA
Fulminantní průběh	u koinfekce 1 % u superinfekce 5 %	1-2 %, 3. trim. grav. kolem 22 % (GT 1 a GT 2)	
Vakcína	proti HB	Rekombinantní; zatím dostupná jen v Číně	ne
Imunoglobulin	proti HB	ne	ne

# Virus hepatitidy F

- V prosinci 1994 publikována zpráva o přenosu agens, který způsobuje non-A, non-E hepatitidu u opic
- Virus-like particles o průměru 27-37 nm) byly detekovány ve stolici francouzských pacientů a v játrech a stolicích infikovaných zvířat
- Agens označeno jako hepatitis French virus (HFV) a analýzou virového genomu zjištěna dvojvláknitá DNA  
*(Journal of Gastroenterology and Hepatology 2001;16(2):124-131)*
- Existence tohoto viru později nebyla potvrzena!!

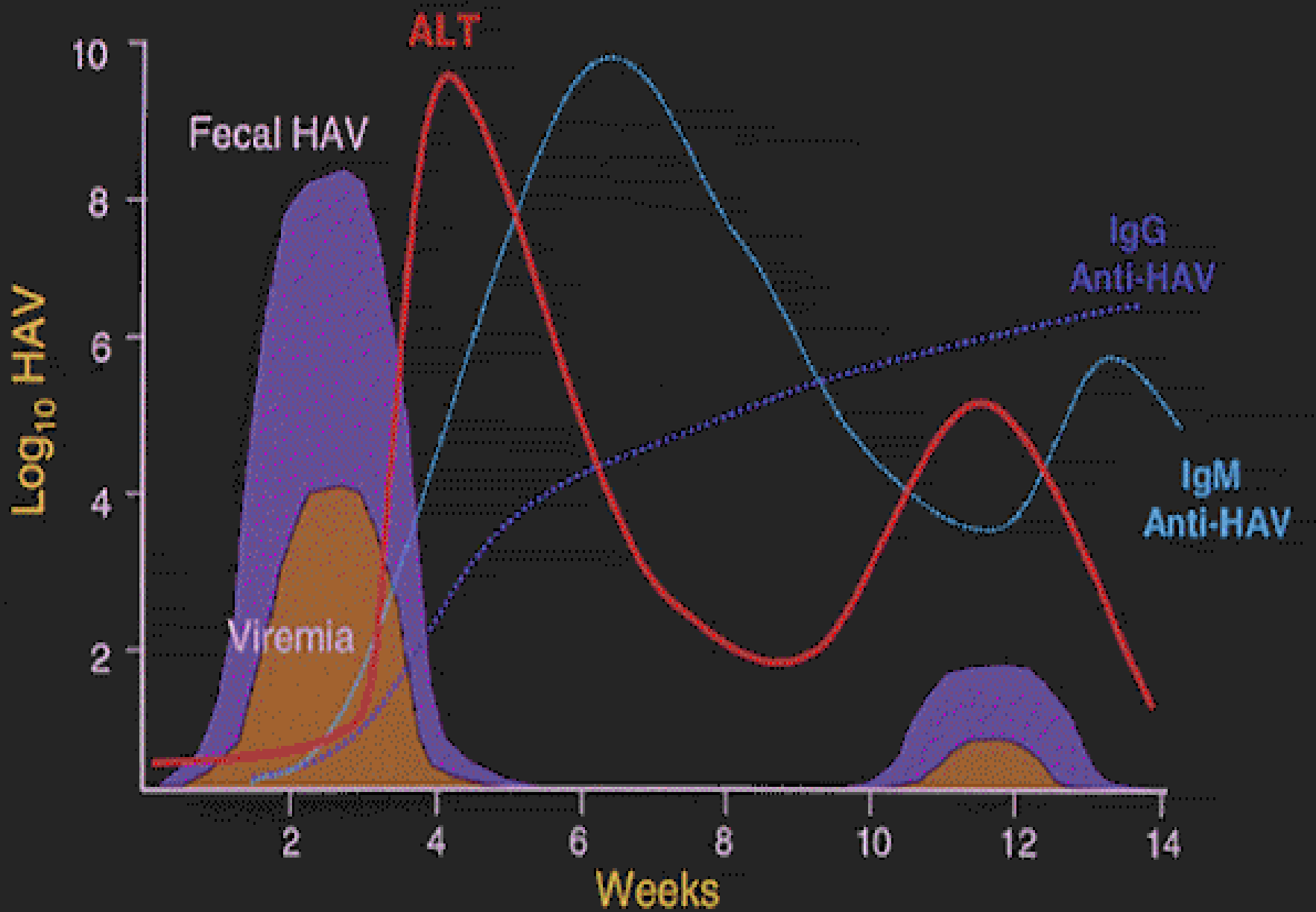
	Anti HAV IgM	HBs Ag	HBe Ag	Anti HBc IgG	Anti HBc IgM	Anti HBs	Anti HCV	Anti HBe	Anti HEV IgM
<b>HA akutní</b>	+	-	-	-	-	-	-	-	-
<b>HB akutní</b>	-	+	+	-	+	-	-	-	-
<b>HB chronická</b>	-	+	+	+	-	-	-	-	-
<b>HBeAg pozitivní</b> (dříve infekce divokým typem viru)	-	+	-	+	-	-	-	-	-
<b>HB chronická</b> <b>HBeAg negativní</b> (dříve infekce e-minus mutantou)	-	+	-	+	-	-	-	+	-
<b>St. po vakcinaci proti HB</b>	-	-	-	-	-	+	-	-	-
<b>HC akutní nebo chronická</b>	-	-	-	-	-	-	+	-	-
							a		
							HCV RNA		
<b>HE akutní</b>	-	-	-	-	-	-	-	-	+
							a/nebo		
							HEV RNA		

## Virology of Acute Hepatitis A



Source: AASLD, 2000

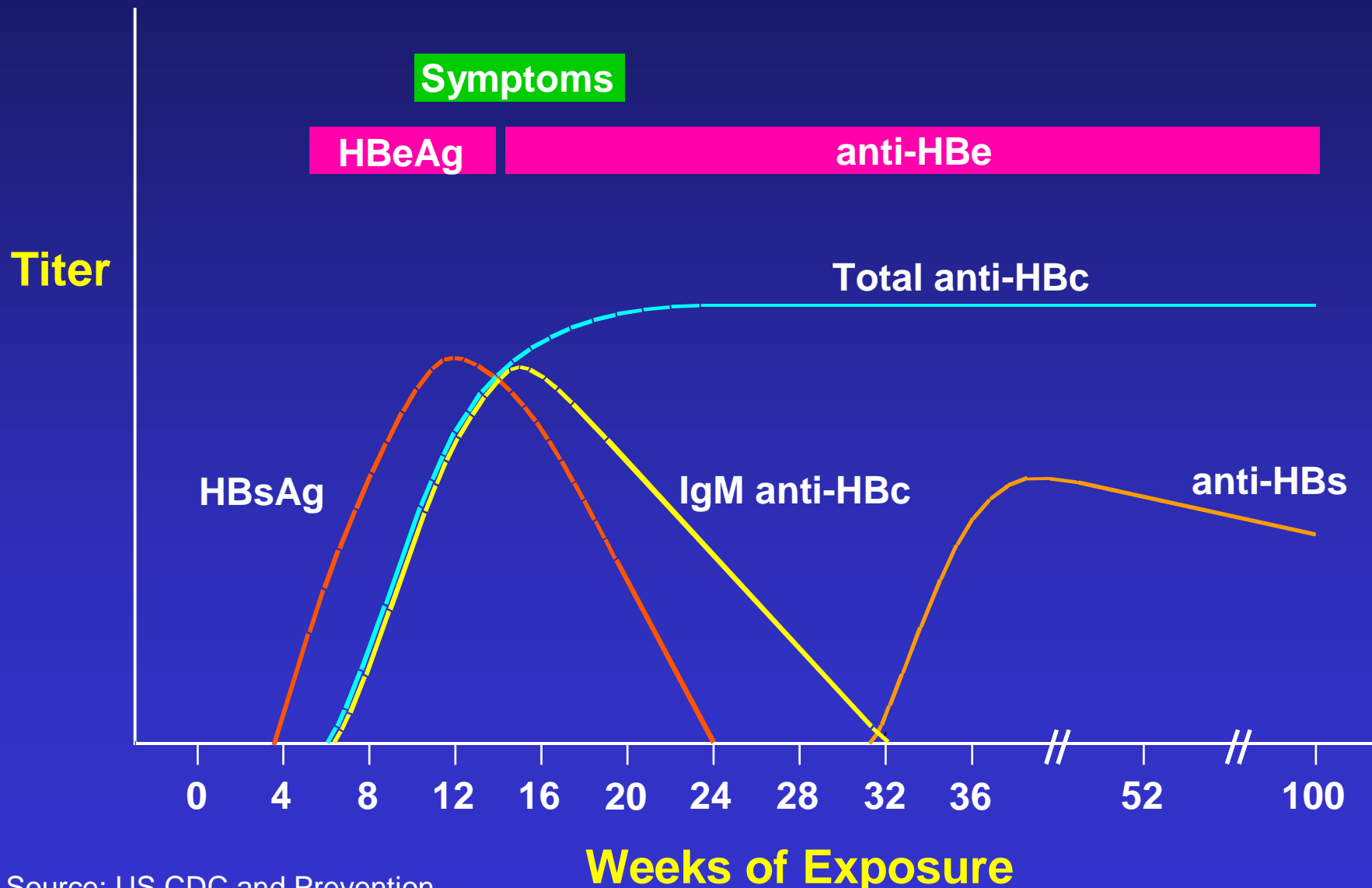
# Relapsing Hepatitis A



# Acute HBV Infection with Recovery

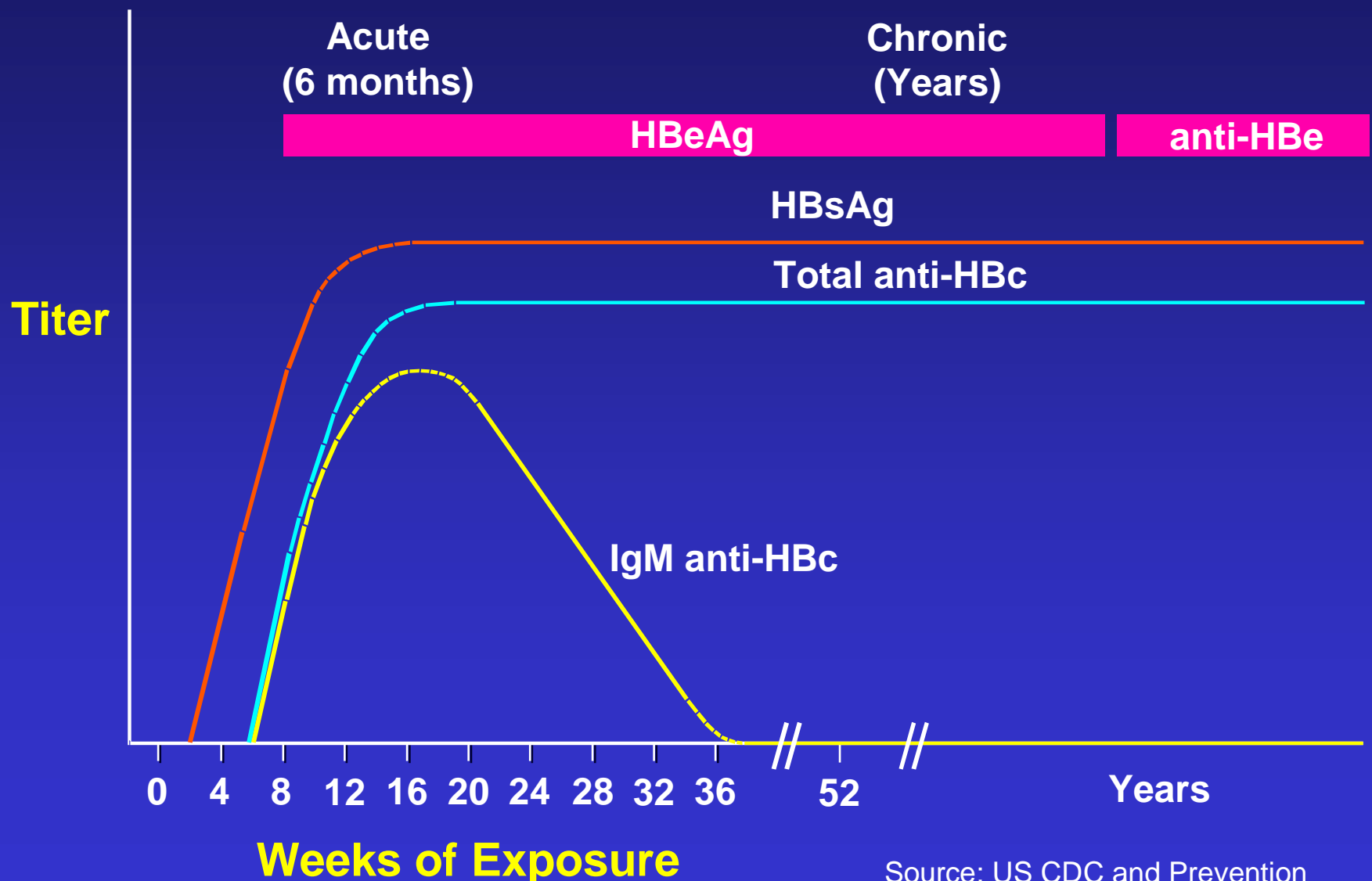
## Typical Serologic Course

HBVMSLS09



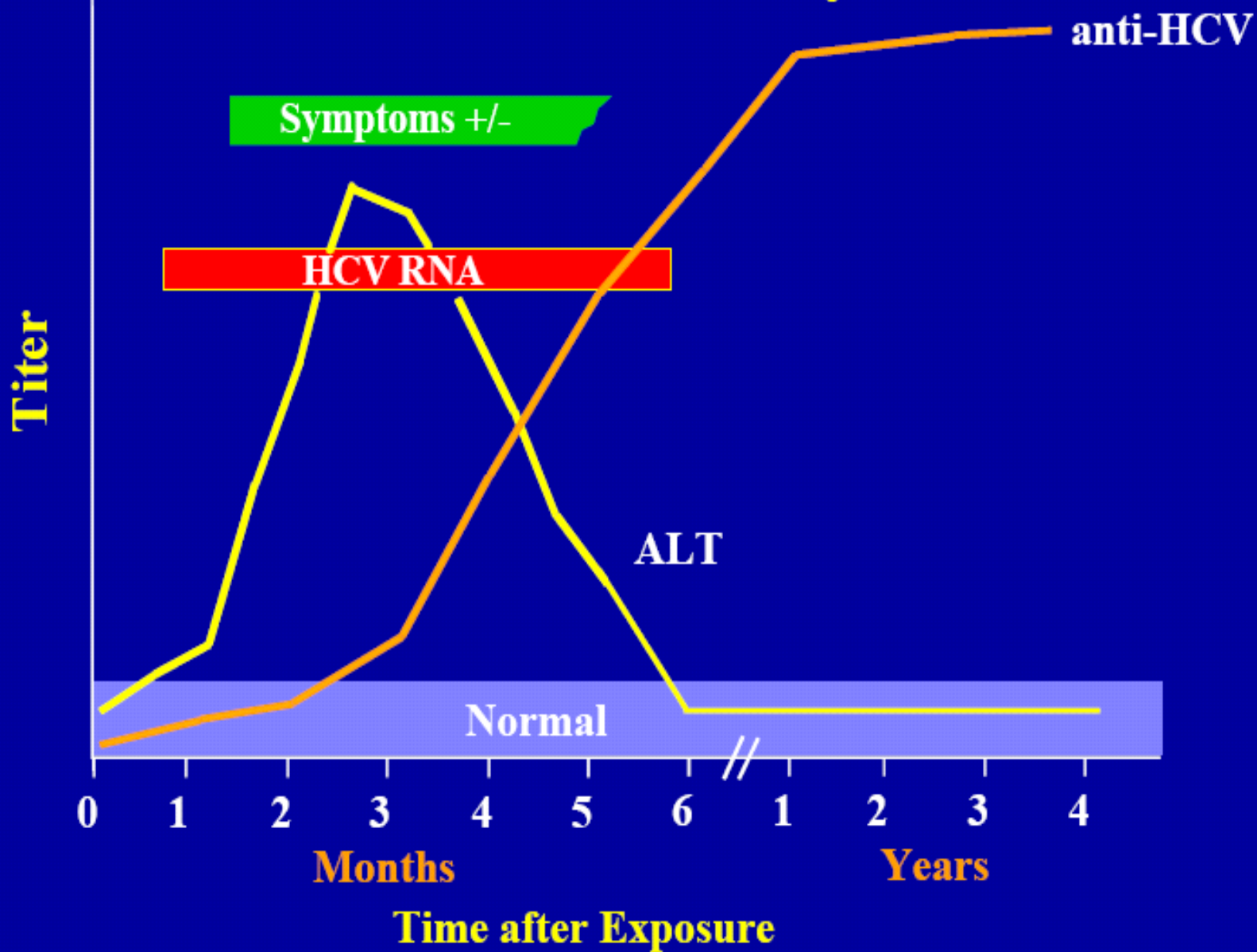
# Progression to Chronic HBV Infection

## Typical Serologic Course

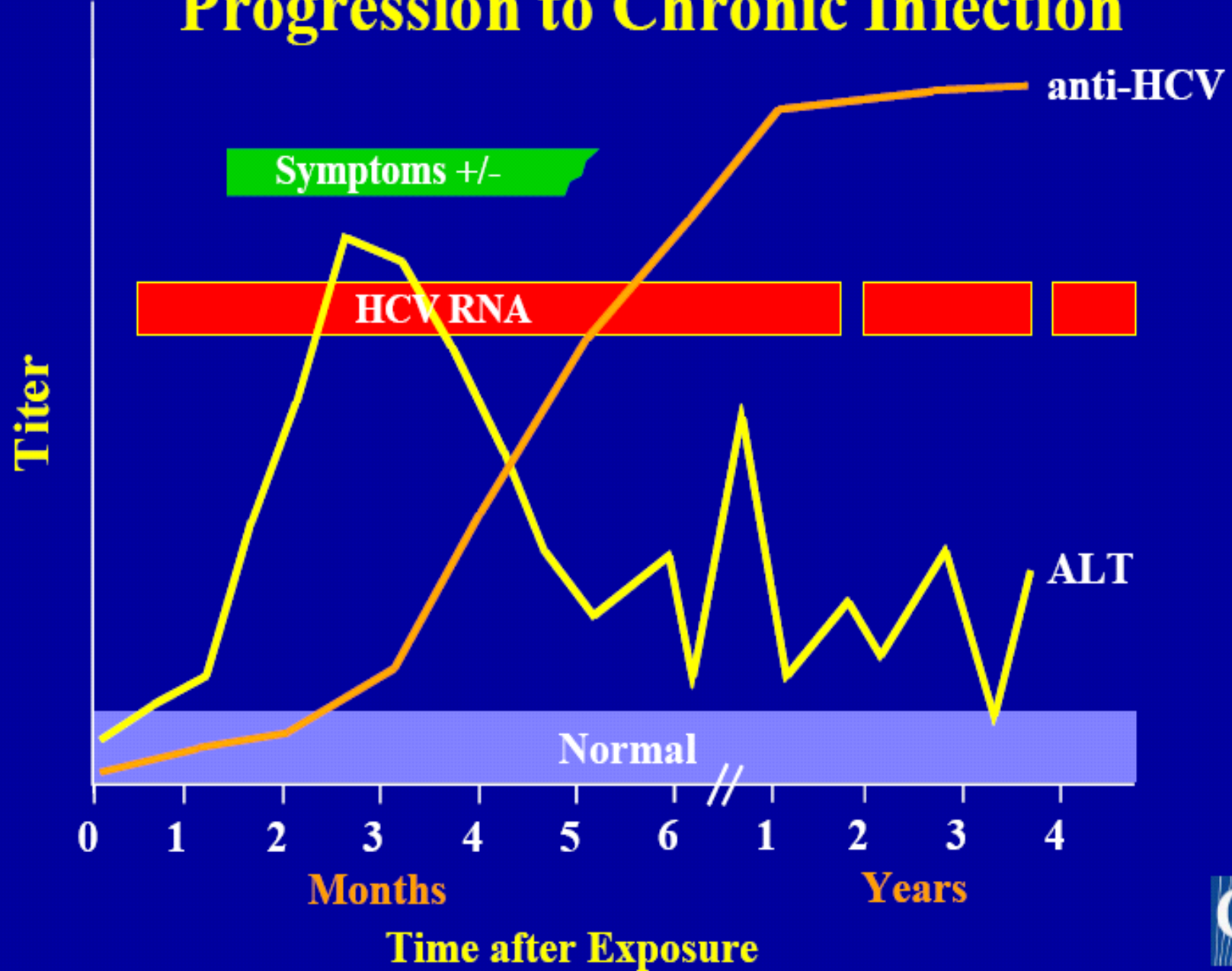


Source: US CDC and Prevention

# Serologic Pattern of Acute HCV Infection with Recovery

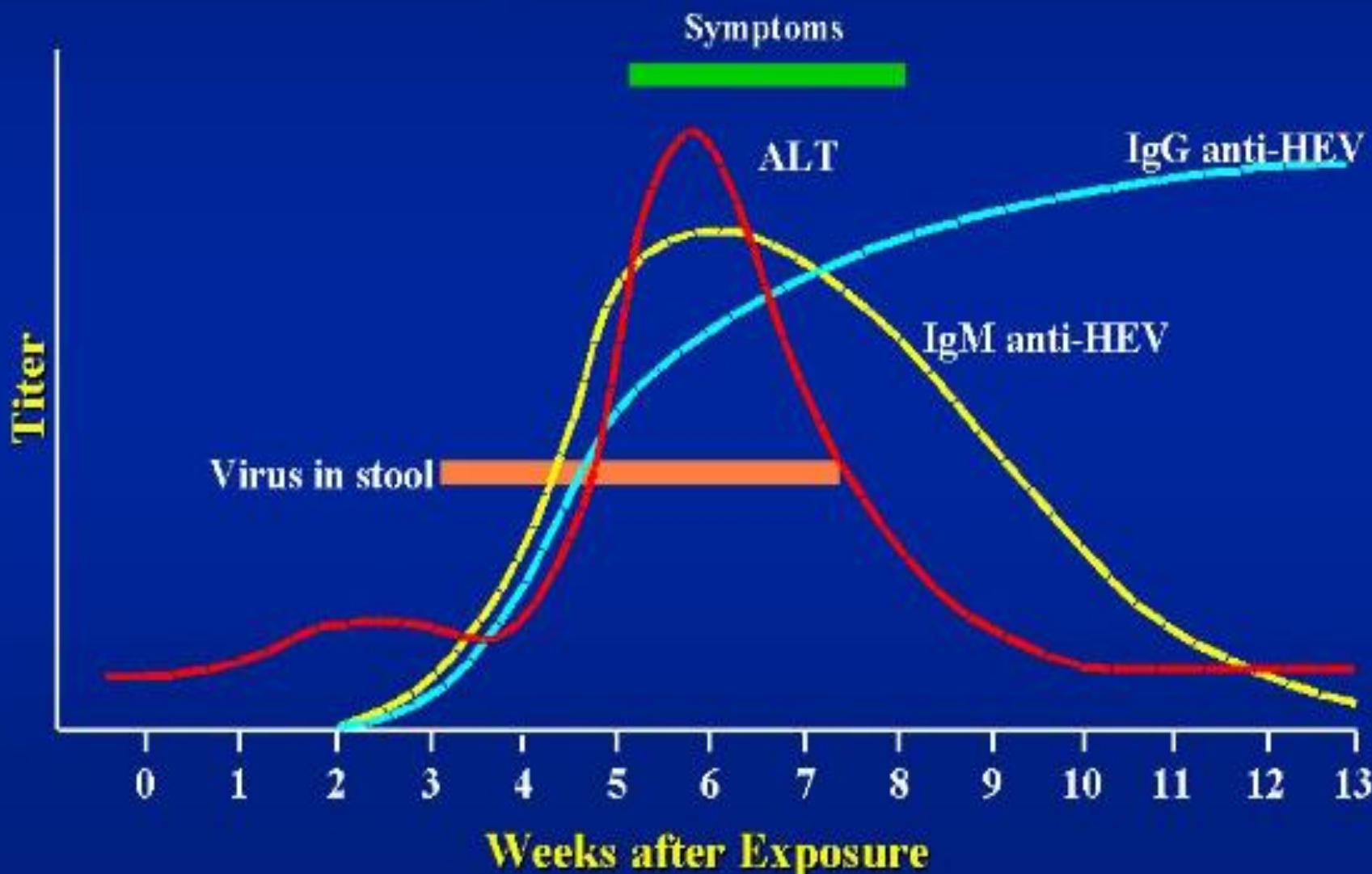


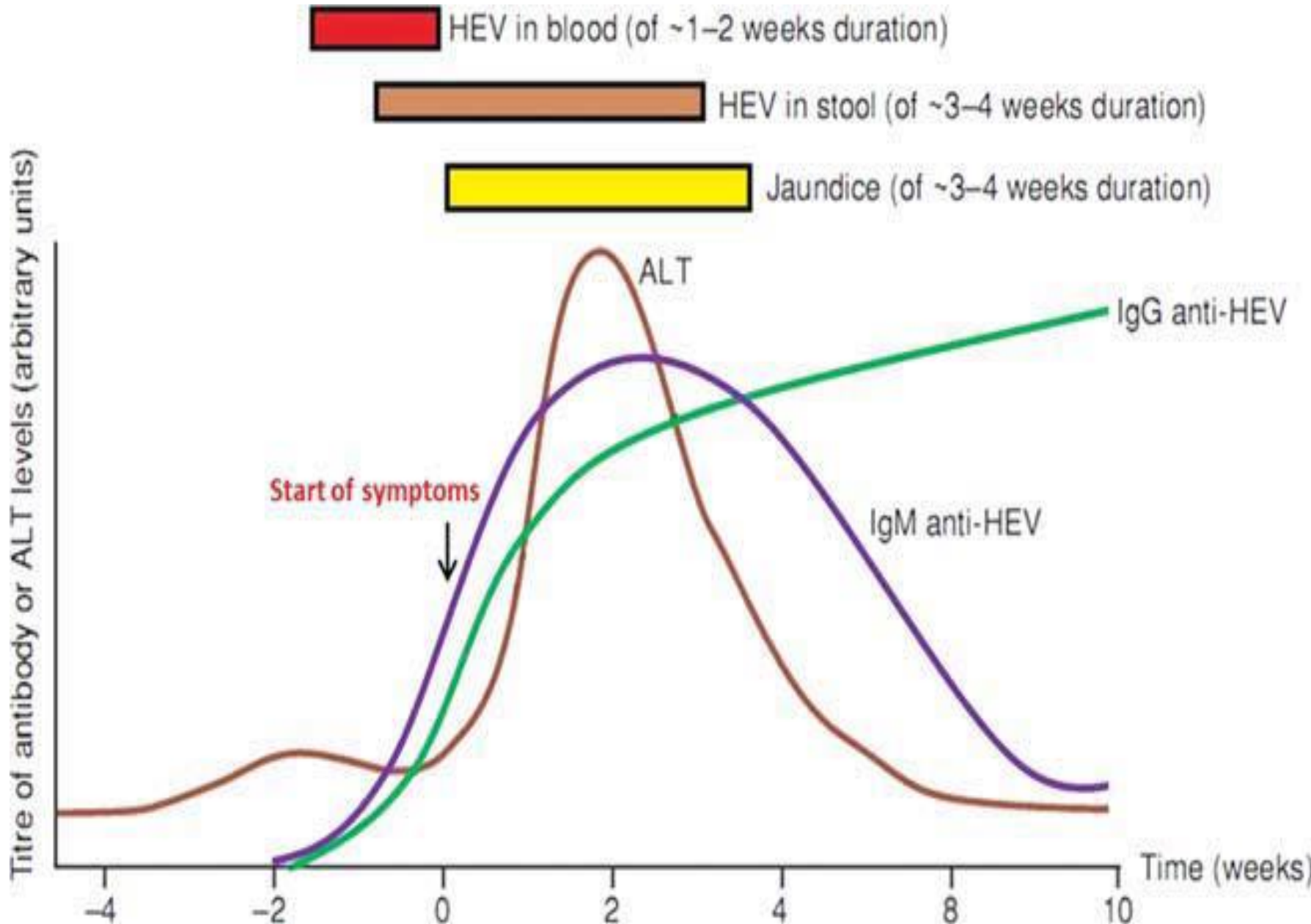
# Serologic Pattern of Acute HCV Infection with Progression to Chronic Infection



# Hepatitis E Virus Infection

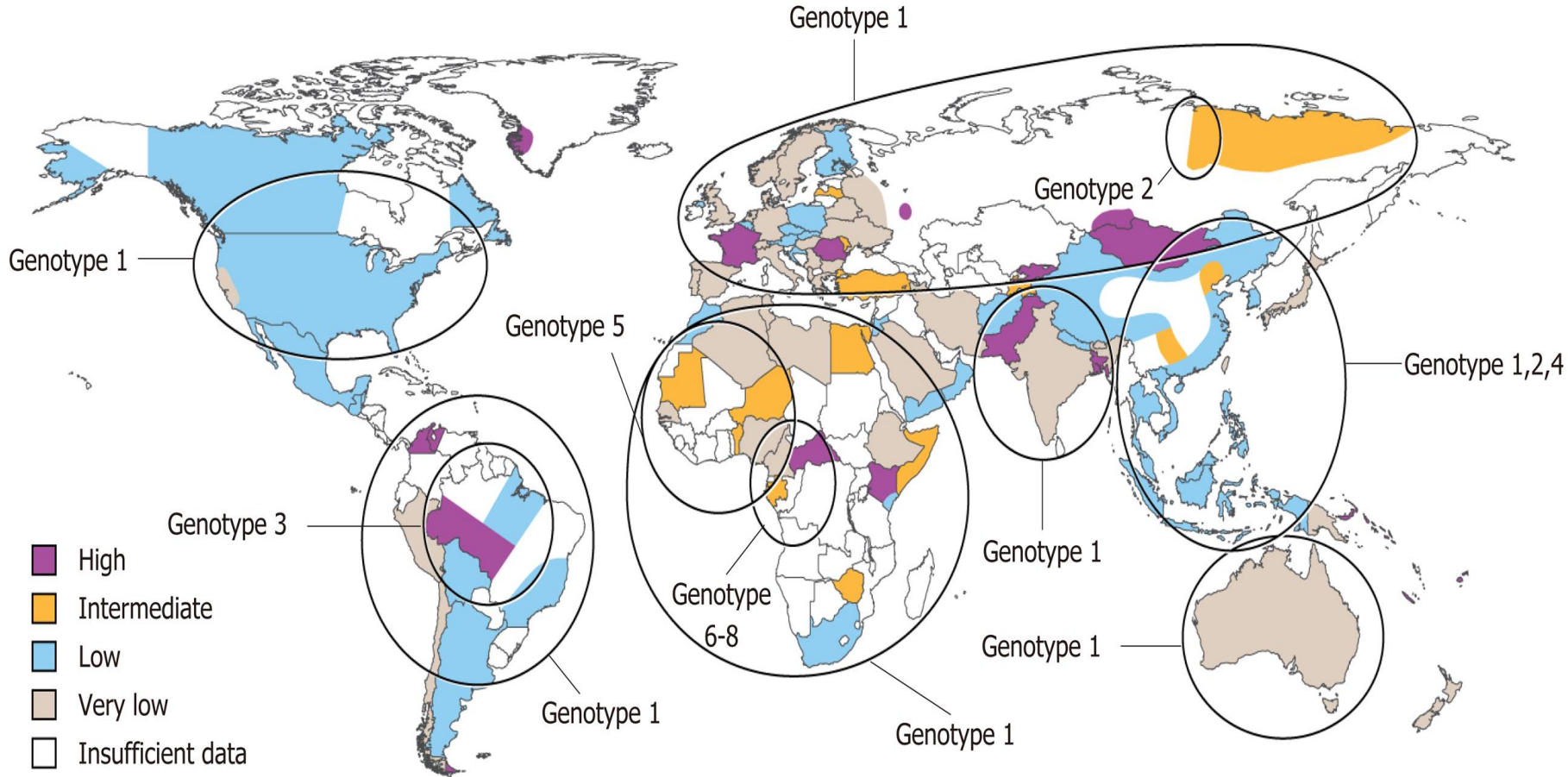
## Typical Serologic Course





# Odhadovaná prevalence HDV infekce

(*World J Gastroenterol* 2019;25(32):4580-4597)



# Acute VH in CR reported to Epidat and ISIN

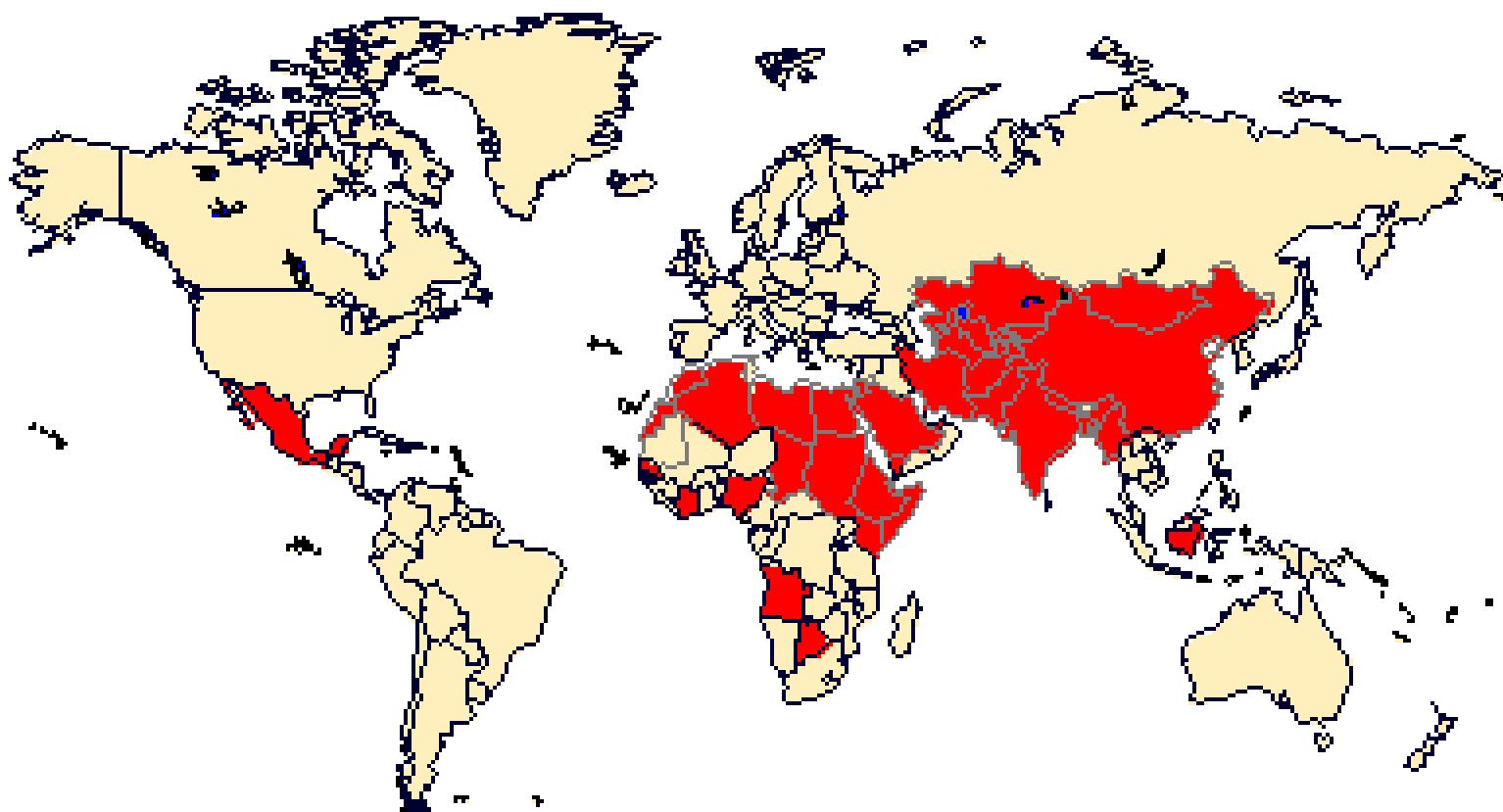
(HC = acute HC + chronic HC firstly diagnosed together)

	1997	1998	1999	2000	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011	2012	2013	2014	2015	2016	2017	2018	2019
H A	1195	904	933	614	325	127	114	70	322	132	128	1648	1104	862	264	284	348	673	724	930	772	211	240
H B	564	575	636	604	457	413	370	392	361	307	307	306	247	244	192	154	133	105	89	73	85	54	41
H C	273	448	634	637	798	858	846	868	844	1022	980	974	836	709	812	794	873	867	956	1104	992	1050	1138
H E	5	17	5	12	13	12	21	36	37	35	43	65	99	72	163	258	218	299	412	339	344	272	268

# Previous theory about HE prevalance

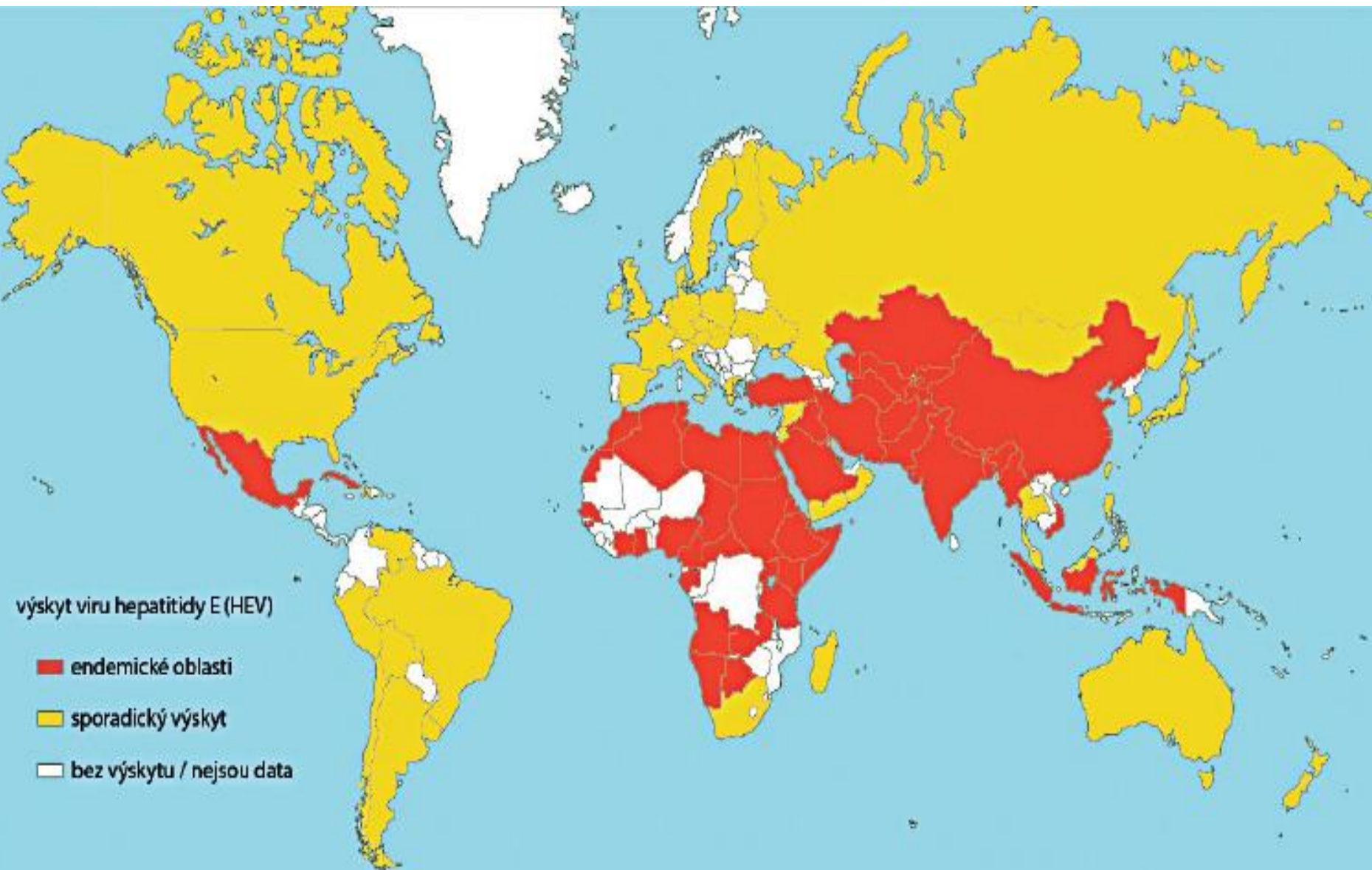
## Geographic Distribution of Hepatitis E

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



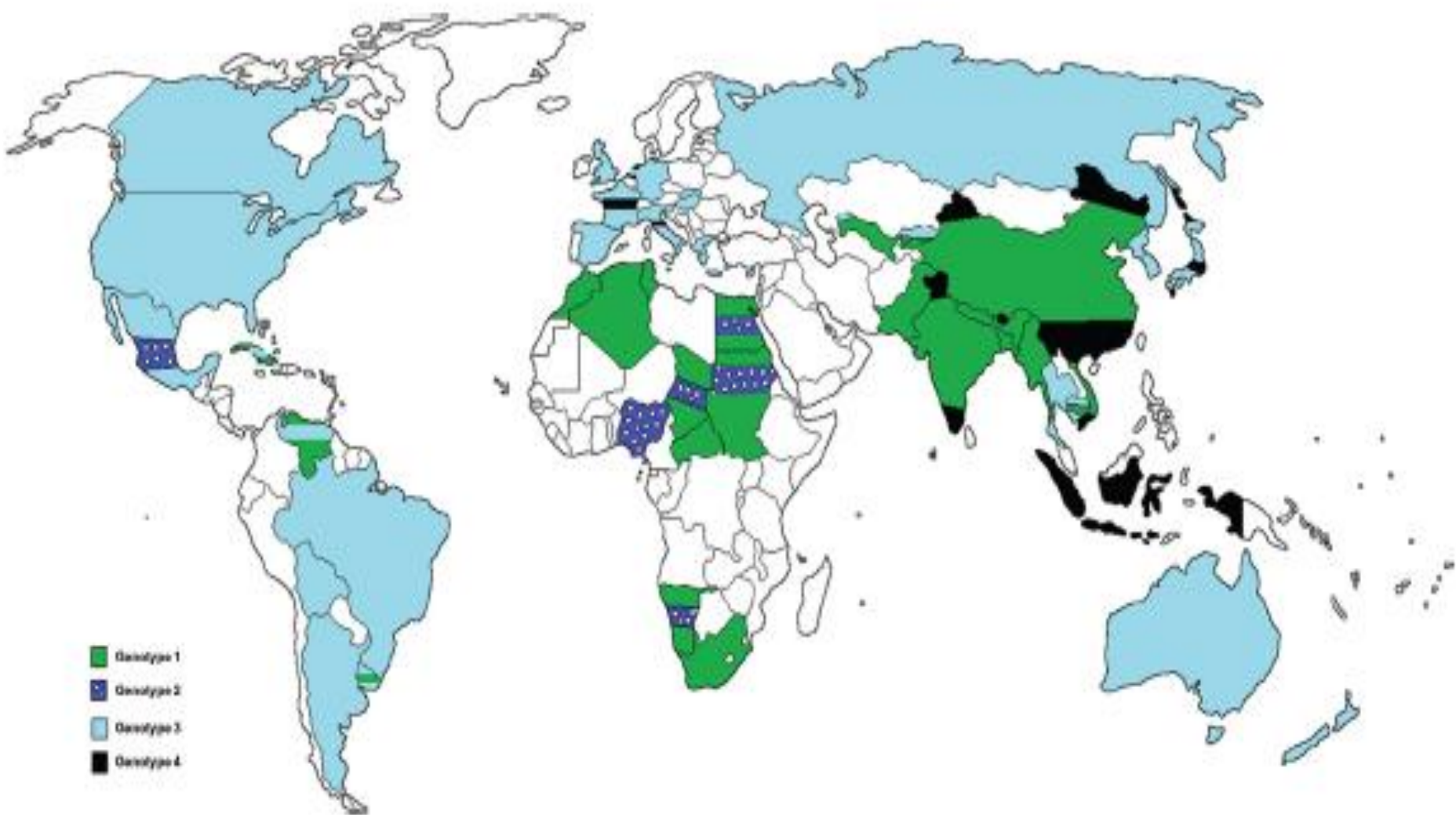
# Výskyt HE ve světě

(Epidem. Mikrobiol. Imunol. 2015;64(2):72-78)

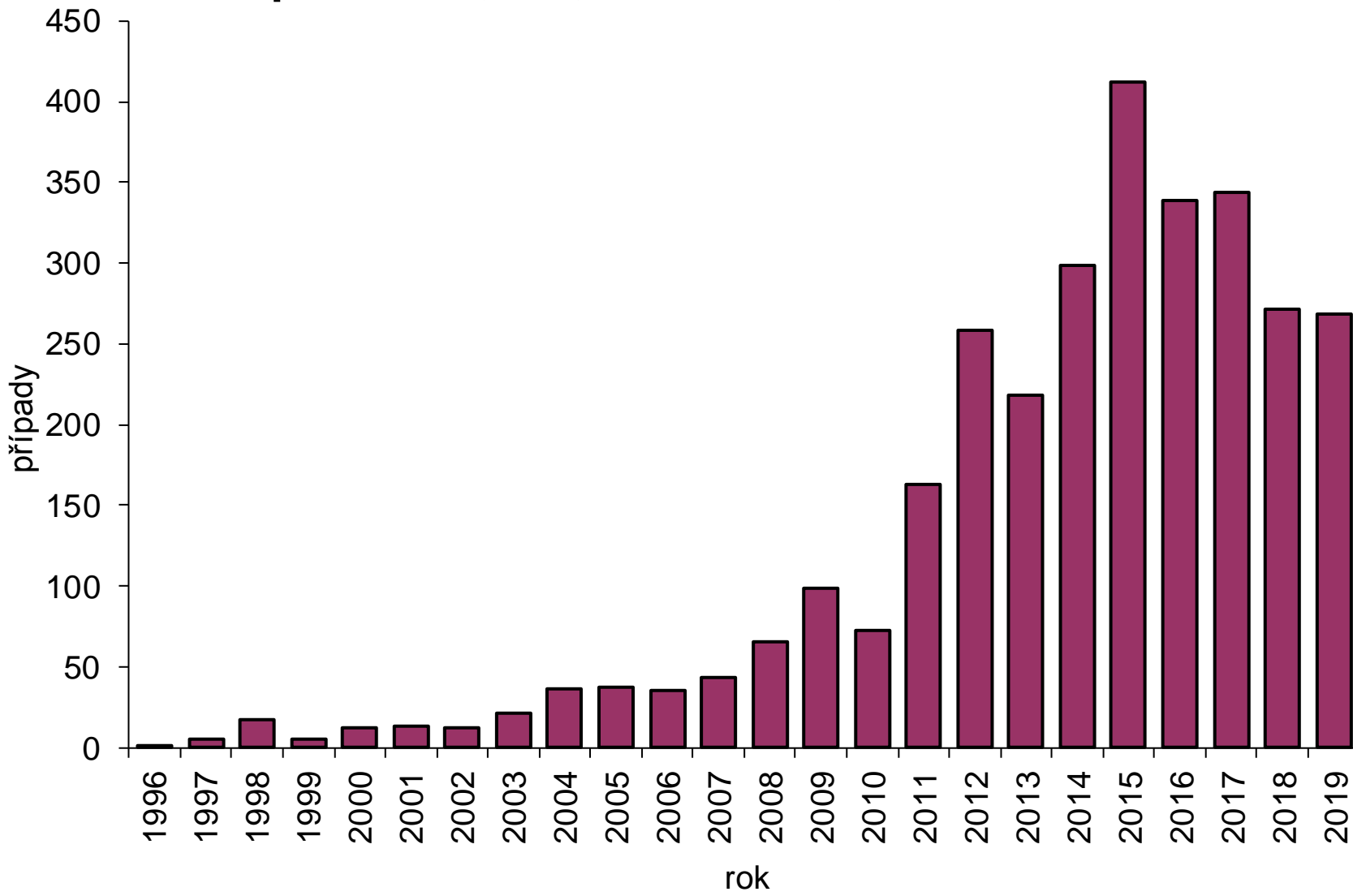


# Genotypes of HE in the world

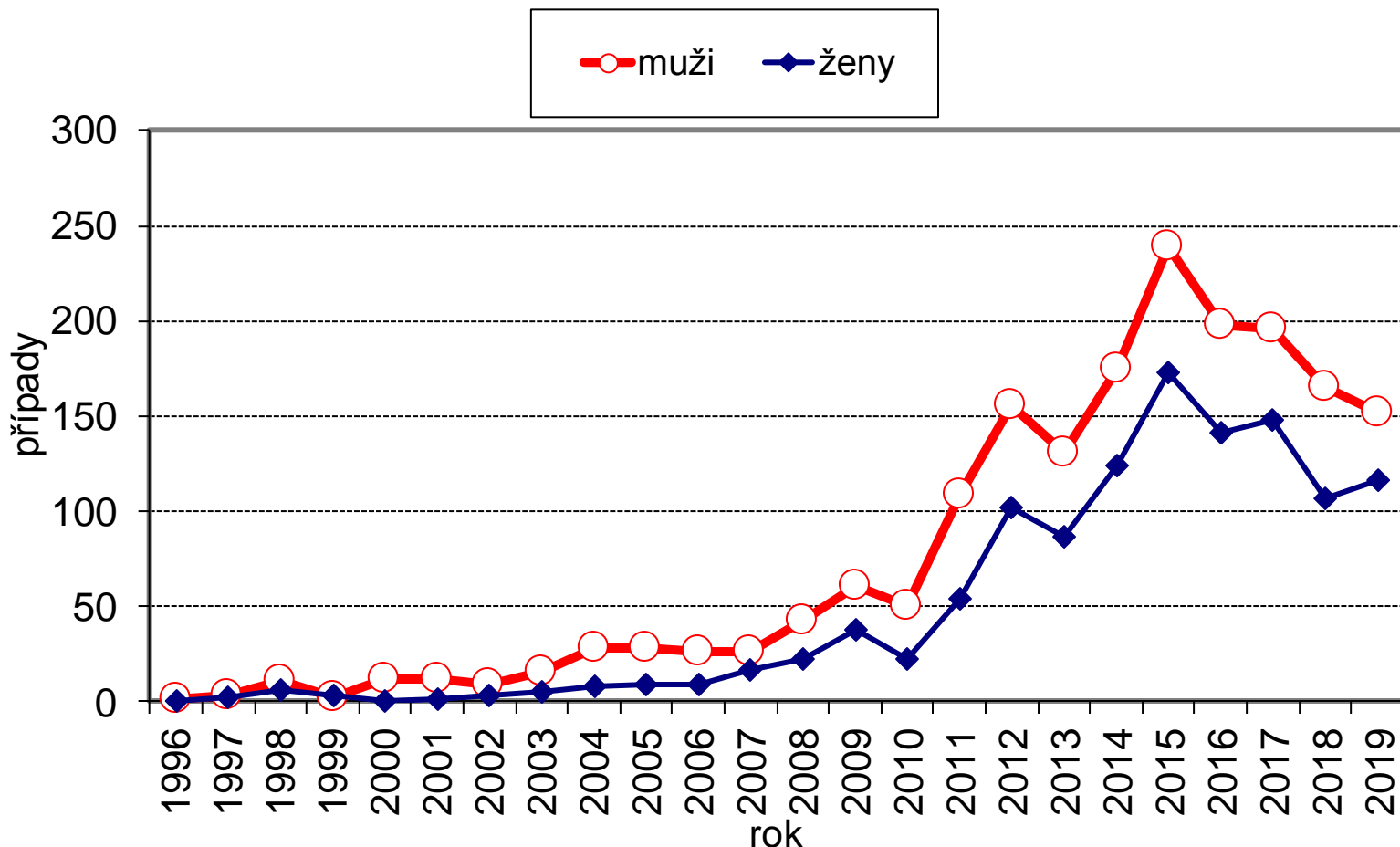
(Hepatic Medicine:Evidence and Research 2014;6:45-59)



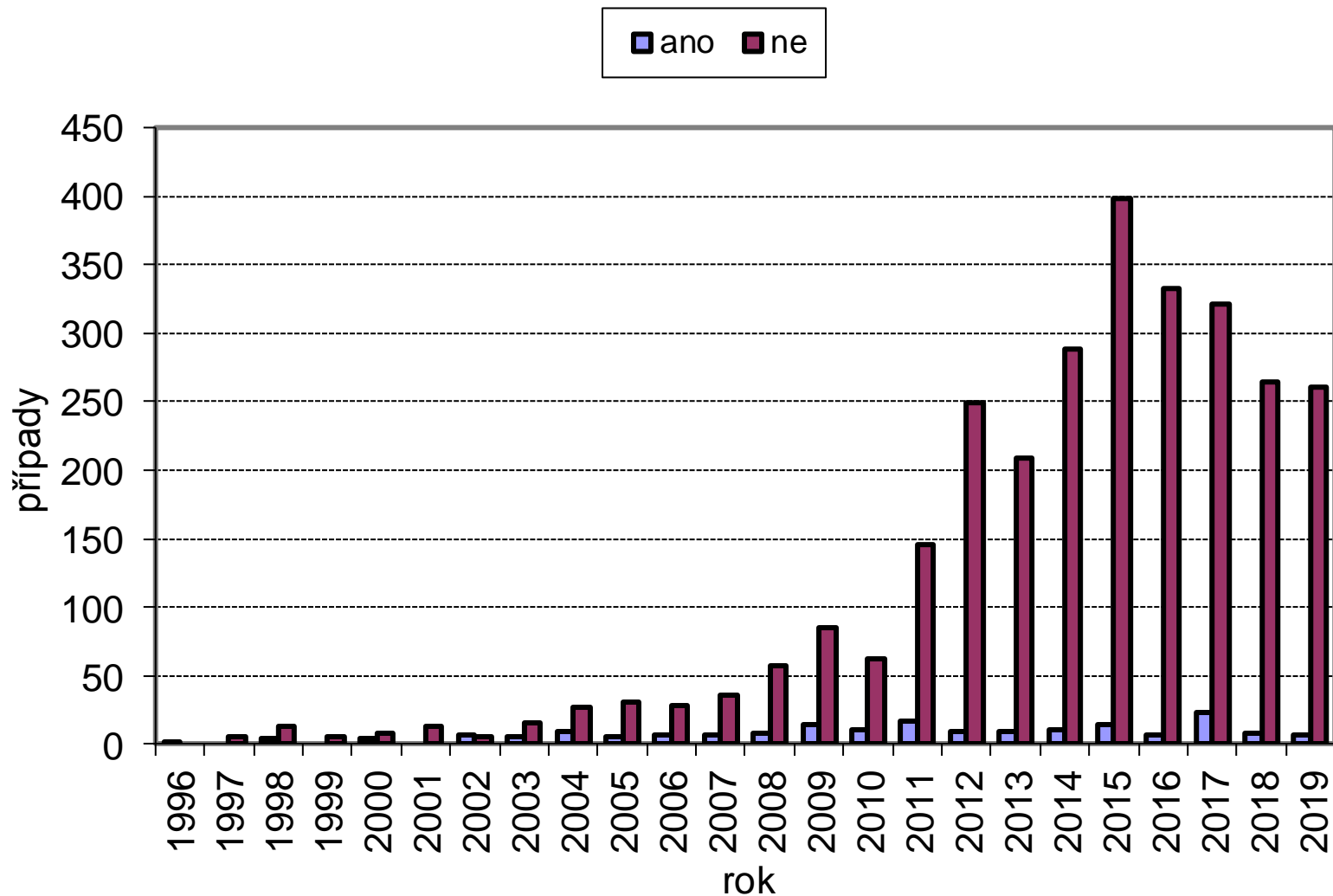
# Hepatitida E, ČR, 1996–2019, hlášená onemocnění



## Hepatitida E, ČR, 1996–2019, muži-ženy, hlášené případy



# Hepatitida E, ČR, 1996–2019, podle importu



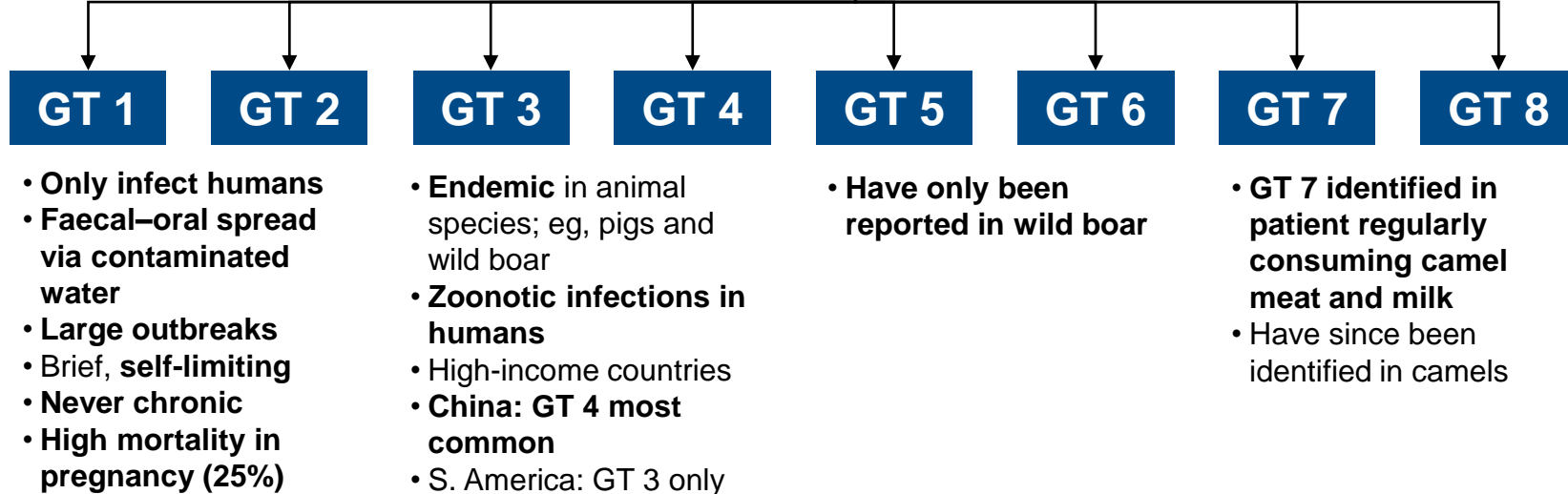
# Virology of HEV



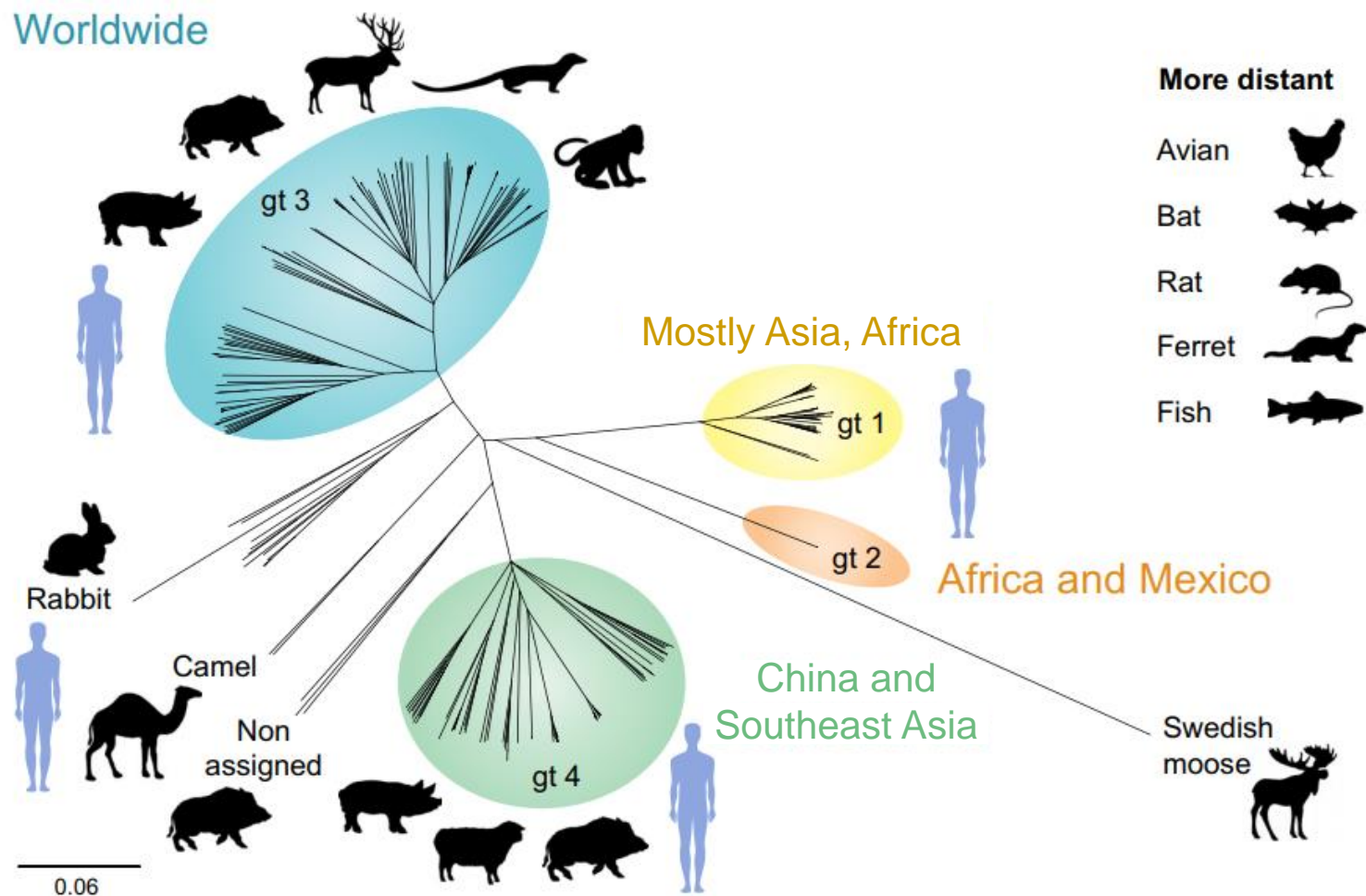
***Hepeviridae* viruses infect mammals, birds and fish**

**Strains infecting humans belong to the *Orthohepevirus* genus, species A**

Species A comprises  
**8 genotypes**



# Phylogenetic relationship of hepeviruses identified in various hosts



# Promyka mungo či promyka indická



Promyka mungo (*Herpestes edwardsii*, angl. mongoose) je promykovitá šelma vyskytující se na jihu Asie. Loví i velké a prudce jedovaté hady. Indové ji s oblibou dávají dětem a nechavají ji přespávat v dětských pokojích (jako domácího mazlíčka, ale též jako ochranu pred hady).

# HEV GT 1 and 2 in brief



- ~20 million infections worldwide
  - 3 million symptomatic cases and 70,000 deaths/year\*
  - WHO guidelines should be consulted for management of outbreaks of acute HEV in resource-limited settings
- Brief, self-limiting, usually in young adults
- **Never chronic**
  - Acute-on-chronic liver failure possible
- **High mortality in pregnancy (25%)**

## Recommendations

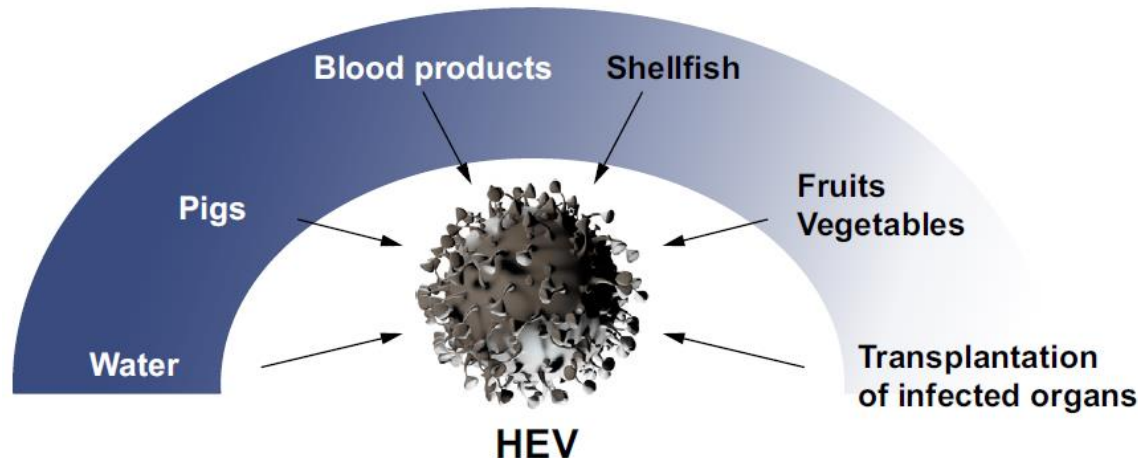
- |  |  |  |
|--|--|--|
| <ul style="list-style-type: none"><li>• Travellers with hepatitis returning from areas endemic for HEV GT 1 or 2 should be tested for HEV</li></ul>          |  |  |
| <ul style="list-style-type: none"><li>• Pregnant women with HEV GT 1 or 2 should be transferred to a liver transplant unit if liver failure occurs</li></ul> |  |  |

# HEV GT 3 and 4: epidemiology

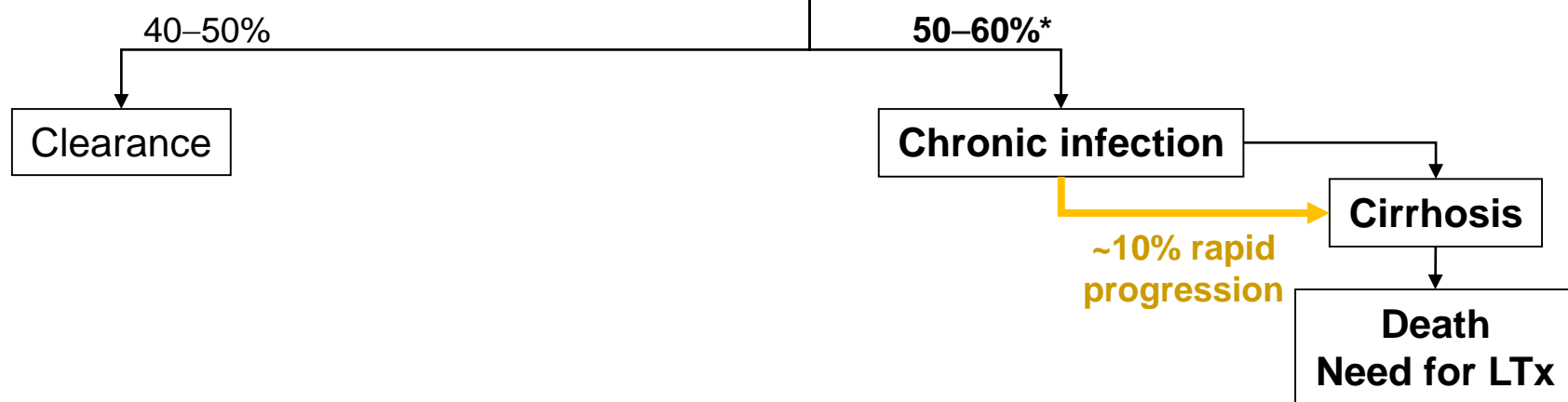


- Endemic in some developing countries, as well as most high-income countries
- **Most common cause of acute viral hepatitis in many European countries**
- Estimated that  $\geq 2$  million locally acquired HEV infections/year
  - Most as a result of zoonotic infection
    - Primary hosts are pigs
- HEV GT 3 and 4 tend to affect older males
  - In an English study, male:female ratio was 3:1; median age, 63 years<sup>1</sup>
- Incidence varies between and within countries, and over time
  - Multiple ‘hotspots’ of HEV infection in Europe

# Transmission and disease progression in transplanted individuals



## Solid organ transplanted individuals with HEV infection



# Treatment of acute HEV infection



- **Acute HEV infection does not usually require antiviral therapy\***
- Most cases of HEV infection are spontaneously cleared
  - Some patients may progress to liver failure
  - Ribavirin
    - Early therapy of acute HEV may shorten course of disease and reduce overall morbidity

Recommendation	□ Grade of evidence	□ Grade of recommendation
<ul style="list-style-type: none"><li>• <b>Ribavirin treatment may be considered in cases of severe acute hepatitis or acute-on-chronic liver failure</b></li></ul>	C	2

# Management of HEV infection



Recommendations	Grade of evidence	Grade of recommendation
<ul style="list-style-type: none"><li><b>Decrease immunosuppression at diagnosis of chronic HEV infection in solid organ transplant recipients, if possible</b></li></ul>	B	1
<ul style="list-style-type: none"><li><b>Give ribavirin for 12 weeks in patients with persisting HEV replication 3 months after detection of HEV RNA</b></li></ul>	B	1
<b>Monitoring of HEV RNA</b> <ul style="list-style-type: none"><li>Assess in serum and stool at the end of scheduled period of ribavirin therapy</li><li>Stop ribavirin if undetectable in both serum and stool</li></ul>	B	1
	C	2

# Management of HEV infection



- Optimal treatment duration in patients who test HEV RNA positive after 4 or 8 weeks of therapy and who are HEV RNA negative after 12 weeks of therapy is unknown\*
- Optimal therapeutic approach unknown in patients who show no response to ribavirin and/or who relapse after retreatment\*

Recommendation	Grade of evidence	Grade of recommendation
<ul style="list-style-type: none"><li>• If HEV RNA is still detectable in serum and/or stool after 12 weeks, ribavirin monotherapy may be continued for an additional 3 months (6 months therapy overall)</li></ul>	C	2
<ul style="list-style-type: none"><li>• Liver transplant recipients who show no response to ribavirin can be considered for treatment with pegylated interferon-<math>\alpha</math></li></ul>	C	2



\*Grade of evidence C

EASL CPG HEV. J Hepatol 2018;doi: 10.1016/j.jhep.2018.03.005 [Epub ahead of print]

# HEV resistance on heat

**Heating the food to an internal temperature  
of 71 °C for 20 min is necessary to  
completely inactivate HEV**

(Barnaud E. et al.: Thermal inactivation of infectious hepatitis E virus in experimentally contaminated food. *Applied and Environmental Microbiology* 2012;78(15):5153-9)

Onemocnění	Název vakcíny	Počet dávek	Množství antigenu	Schéma	Indikace	Objem
VHA	<b>HAVRIX</b>	2	A: 720 EU A:1440 EU	M 0-(6-12)	1– 5 let ≥ 16 let	0,5 ml 1,0 ml
	<b>AVAXIM</b>	2	A: 160 AU	M 0-(6-18)	≥ 2 let	0,5 ml
	<b>VAQTA</b>	2	A: 25 IU A: 50 IU	M 0-(6-18)	2-17 let ≥ 18 let	0,5 ml 1,0 ml
VHB	<b>ENGERIX -B</b>	3	B: 10 µg B: 20 µg	M 0-1-6 (M 0-1-2-12)	0-15 let ≥ 16 let	0,5 ml 1,0 ml
	<b>HB VAX PRO</b>	3	B: 5 µg B: 10 µg	M 0-1-6 (M 0-1-2-12)	0-15 let ≥ 16 let	0,5 ml 1,0 ml
	<b>FENDRIX</b>	4	B: 20 µg	M 0-1-2-6	≥ 15 let	0,5 ml
VHA+VHB	<b>TWINRIX</b>	3	A/B:360 EU/10 µg A/B:720 EU/20 µg	M 0-1-6 (D 0-7-21 + M 12)	1-15 let ≥ 16 let	0,5 ml 1,0 ml

D – den, M – měsíc

**Tab. I** Přehled vakcín proti VHA a VHB registrovaných v ČR

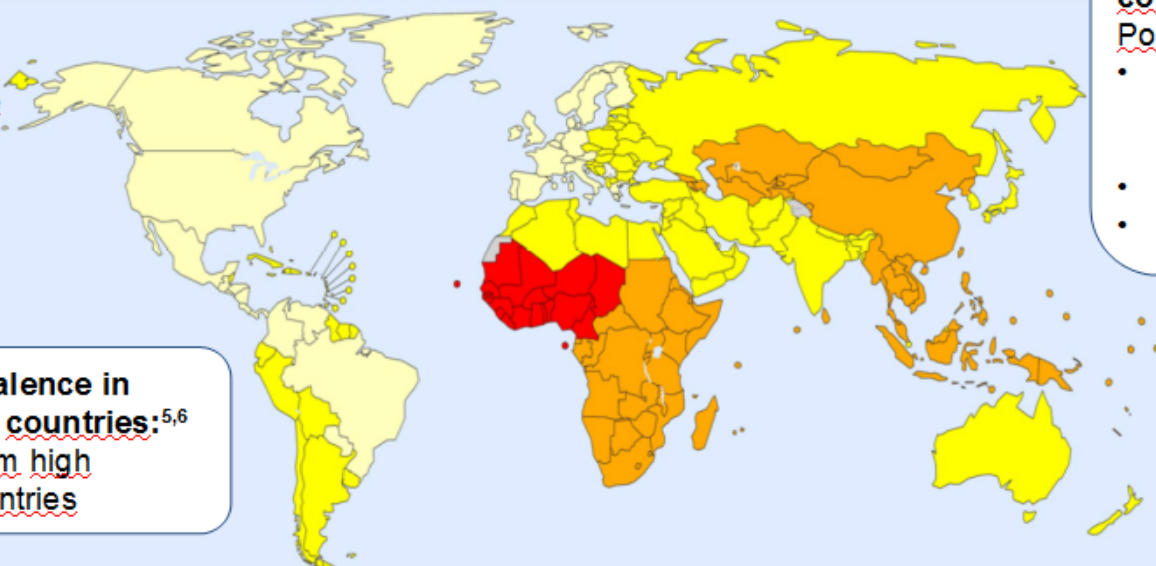
# Epidemiology and public health burden<sup>1</sup>



- Worldwide ≈250 million chronic HBsAg carriers<sup>2,3</sup>
- 686,000 deaths from HBV-related liver disease and HCC in 2013<sup>4</sup>

HBsAg prevalence, adults (19–49 years), 2005<sup>3</sup>

- <2%
- 2–4%
- 5–7%
- ≥8%
- Not applicable



Increasing prevalence in some European countries:<sup>5,6</sup>

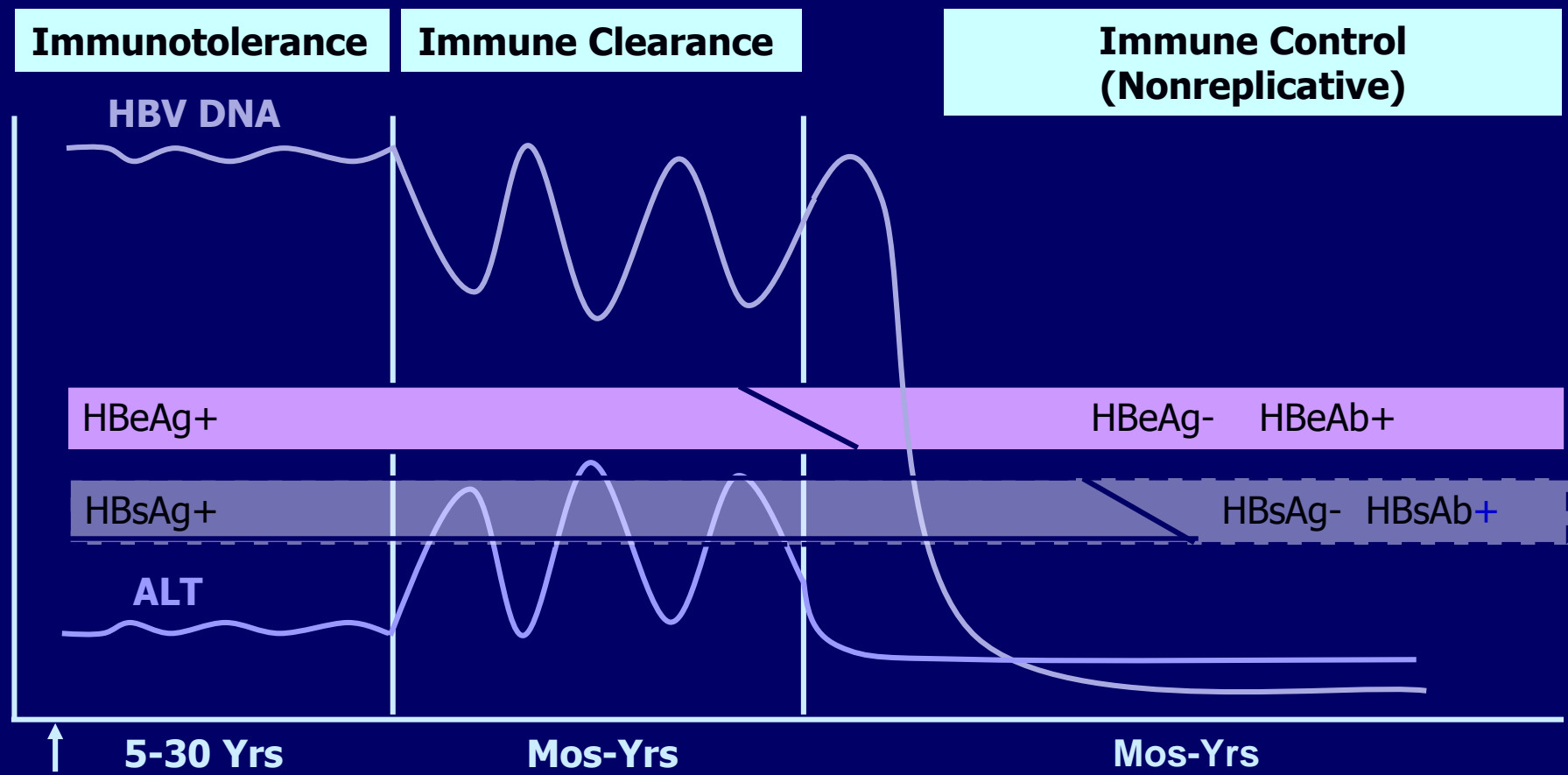
- Migration from high endemic countries

Decreasing prevalence in some endemic countries, e.g. Taiwan<sup>7</sup>

- Possible reasons:
- Improved socioeconomic status
  - Vaccination
  - Effective treatments

1. EASL CPG HBV. J Hepatol 2017;67:370–98; 2. Schweitzer A, et al. Lancet 2015;386:1546–55;  
3. Ott JJ, et al. Vaccine 2012;30:2212–9; 4. GBD 2013 Mortality and Causes of Death Collaborators. Lancet 2015;385:117–71;  
5. Coppola N, et al. Euro Surveill 2015;20:30009; 6. Hampel A, et al. Bundesgesundheitsblatt Gesundheitsforschung  
Gesundheitsschutz 2016;59:578–83; 7. Chen C-L, et al. J Hepatol 2015;63:354–63.

# Natural History of Chronic HBV Infection

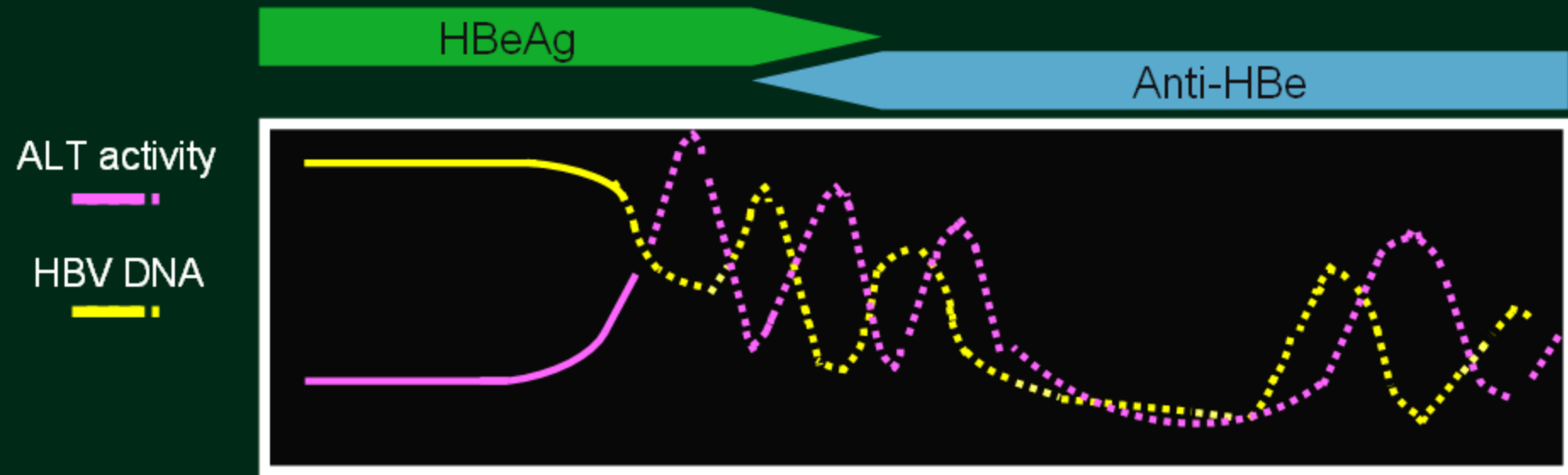


**Infection**

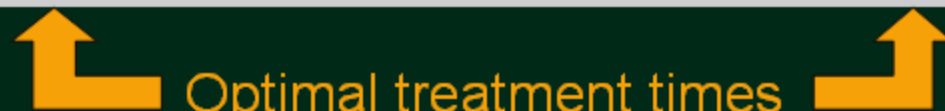
Yim HJ, et al. Hepatology. 2006;43:S173-S181.

# 4 Phases of Chronic HBV Infection

## Current Understanding of HBV Infection



Phase	Immune Tolerant	Immune Clearance	Inactive Carrier State	Reactivation
Liver	Minimal inflammation and fibrosis	Chronic active inflammation	Mild hepatitis and minimal fibrosis	Active inflammation



# Subset of Agents Reported to Cause HBV Reactivation

Class	Agents
Corticosteroids	Dexamethasone, methylprednisolone, prednisolone
Antitumor antibiotics	Actinomycin D, bleomycin, daunorubicin, doxorubicin, epirubicin, mitomycin-C
Plant alkaloids	Vinblastine, vincristine
Alkylating agents	Carboplatin, chlorambucil, cisplatin, cyclophosphamide, ifosfamide
Antimetabolites	Azauridine, cytarabine, fluorouracil, gemcitabine, mercaptopurine, methotrexate, thioguanine
Monoclonal antibodies	Alemtuzumab, rituximab
Others	Colaspase, docetaxel, etoposide, fludarabine, folinic acid, interferon, procarbazine

# HBV Reactivation: Overview

- Clinical syndrome characterized by an increase in HBV DNA and ALT/AST with or without symptoms or jaundice
- Occurs in pts with active (HBsAg+) and resolved (HBsAg-, anti-HBc+) HBV infection
- Wide clinical spectrum  
Ranges from silent to acute liver failure (fibrosing cholestatic hepatitis)
- Can occur during treatment with many immunosuppressive agents  
**May also occur up to 12 mos after treatment**
- Preventable by antiviral prophylaxis

Di Bisceglie AM, et al. Hepatology. 2015;61:703-711.

Perrillo RP, et al. Gastroenterology. 2015;148:221-244.



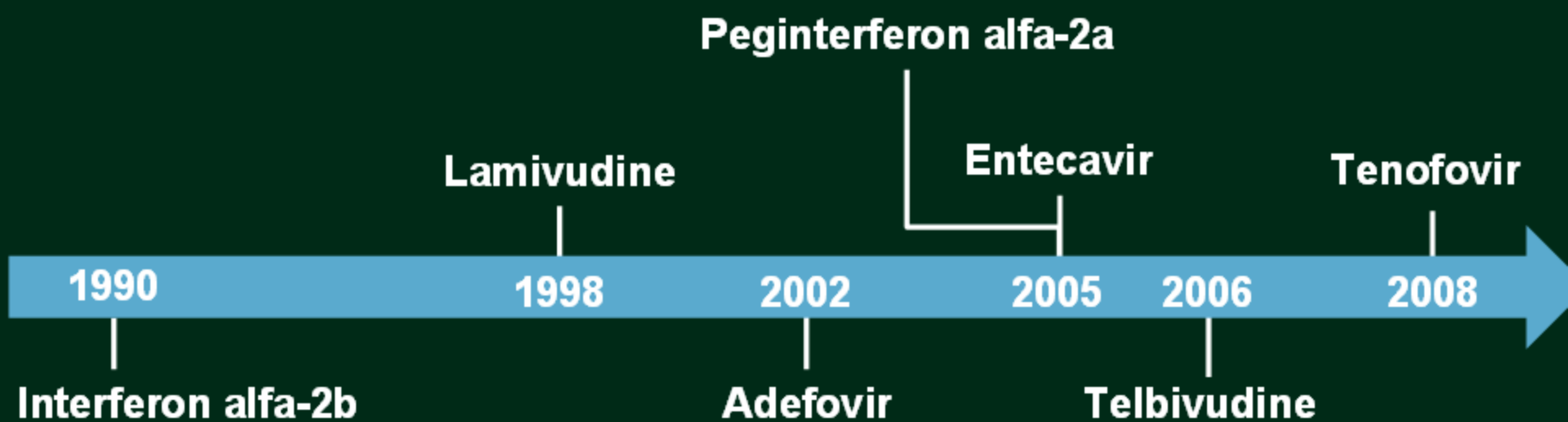
Slide credit: [clinicaloptions.com](http://clinicaloptions.com)

# Nová nomenklatura chronické HBV infekce

- Chronická HBV infekce = dynamický proces odrážející interakce mezi replikací HBV a imunitním systémem
- Ne všichni pacienti s chronickou HBV infekcí mají chronickou hepatitidu B

1. HBeAg pozitivní chronická HBV infekce  
(dříve „imunotolerantní fáze“)
2. HBeAg pozitivní chronická hepatitida B
3. HBeAg negativní chronická HBV infekce  
(dříve „inaktivní nosičství“)
4. HBeAg negativní chronická hepatitida B
5. HBsAg negativní fáze  
(také „okultní HBV infekce“)

# Evolution of Chronic HBV Therapy Over Time



# Nucleos(t)ide analogues (NA)

\*famciklovir – *analogue of deoxyguanosin* (*Famvir*™)

\*lamivudine (3TC, LAM) – *analogue of deoxycytidin* (*Zeffix*™)

\*adefovir dipivoxil (ADV) – *analogue of dAMP* (*Hepsera*™)

\*tenofovir – *similar as ADV*

TDF = *tenofovir disoproxil fumarate* (*Viread*™)

TAF = *tenofovir alafenamide* (*Vemlidy*™)

\*entecavir (ETC) – *analogue of deoxyguanin* (*Baraclude*™)

\*emtricitabine (FTC) – *analogue of cytosinu* (*Emtriva*™)

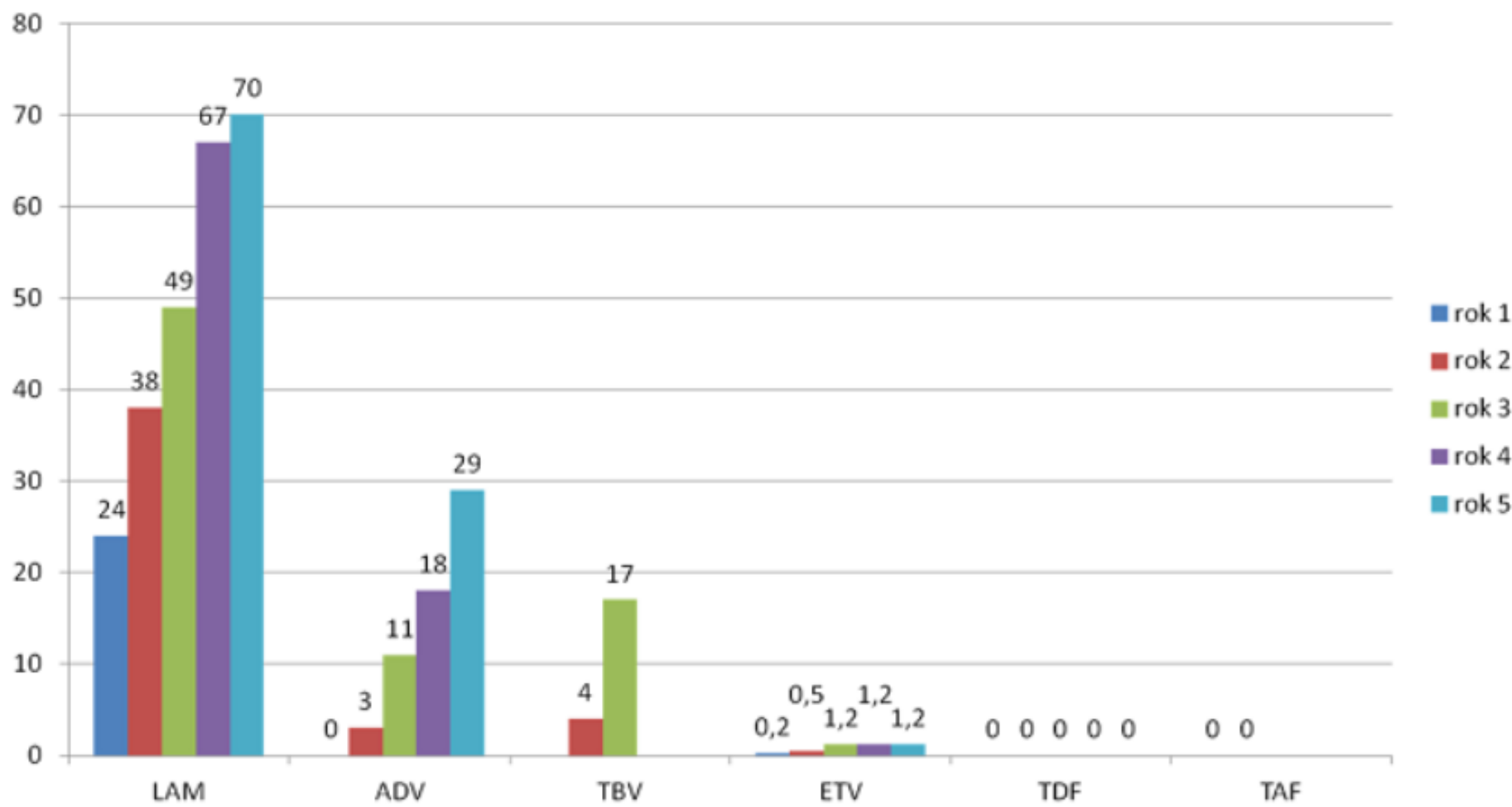
\*clevudine (LFMAU) – *analogue of pyrimidin* (*Levoril*™, *Revovir*™)

\*telbivudine (LdT) – *analogue of L-deoxythymidin* (*Sebivo*™)

# Duration of the therapy of chronic HB

	HBeAg pos.	HBeAg neg.
<b>PEG-IFN alfa</b>	<b>48 wks</b>	<b>48 wks</b>
<b>Nucleos(t)ide analogues (TDF, ETC, TAF)</b>	<i>Pts without CIH:</i>  To HBeAg disappearance and next 12 months	To HBsAg disappearance  X  HBsAg persistency very often to the end of the life
<b>Initiation of antiviral therapy</b>	<b>HBV DNA &gt;2 000 IU/mL (&gt;<math>10^4</math> copies/mL)</b>	<b>HBV DNA &gt;2 000 IU/mL (&gt;<math>10^4</math> copies/mL)</b>

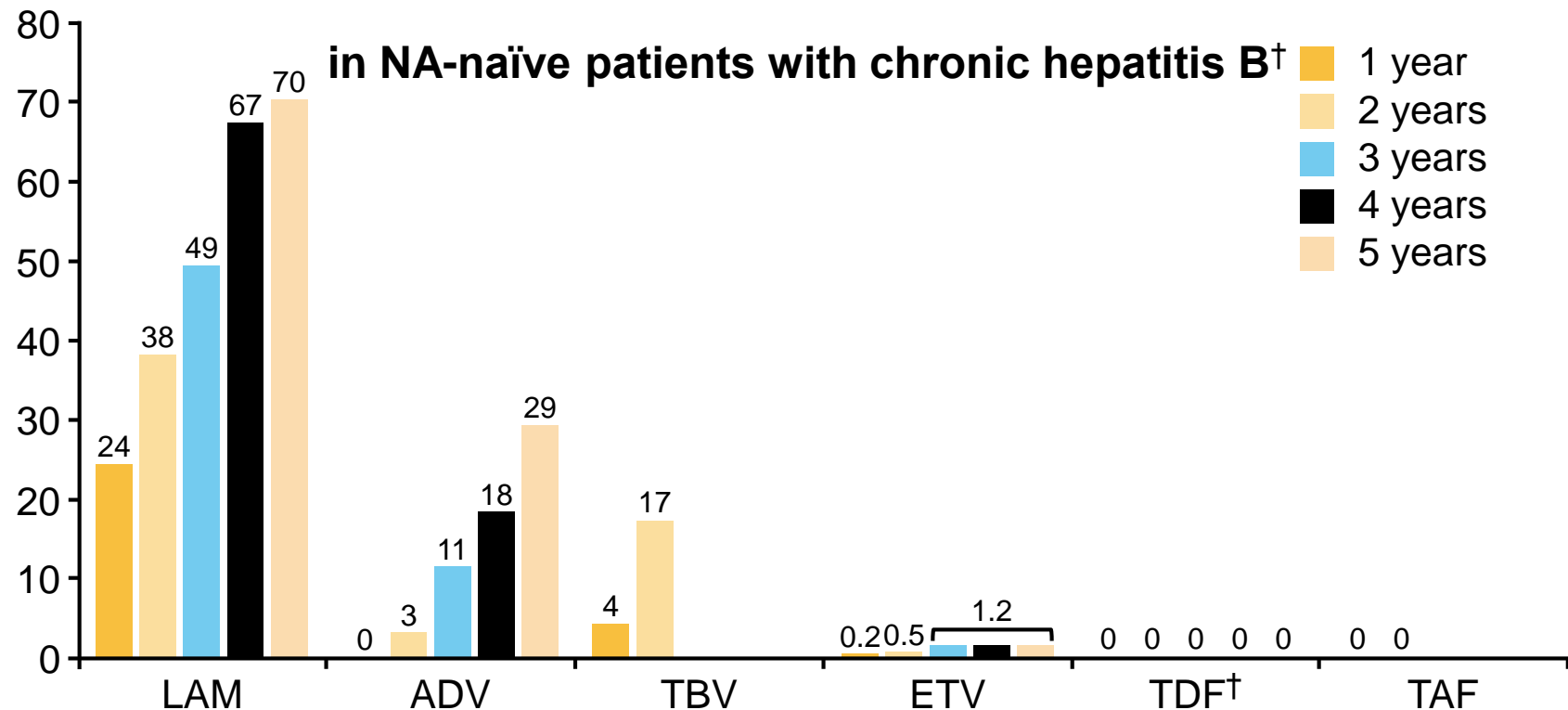
**Cumulative incidence of HBV resistance for lamivudine (LAM), adefovir (ADV), telbivudine (TBV), entecavir (ETV), tenofovir (TDF) and tenofovir alafenamide (TAF) in pivotal trials in nucleos(t)ide-naïve patients with chronic HB (*Hepatology* 2014;60:313A-317A)**



Prevention of resistance should rely on the use of first-line NAs with a high barrier to resistance\*



### Cumulative incidence of HBV resistance to NAs in pivotal trials



\*Evidence level I, grade of recommendation 1; †Collation of currently available data – not from head-to-head studies;

**#No evidence of resistance has been shown after 8 years of TDF treatment**

# PegIFN vs Nucleos(t)ide Analogues

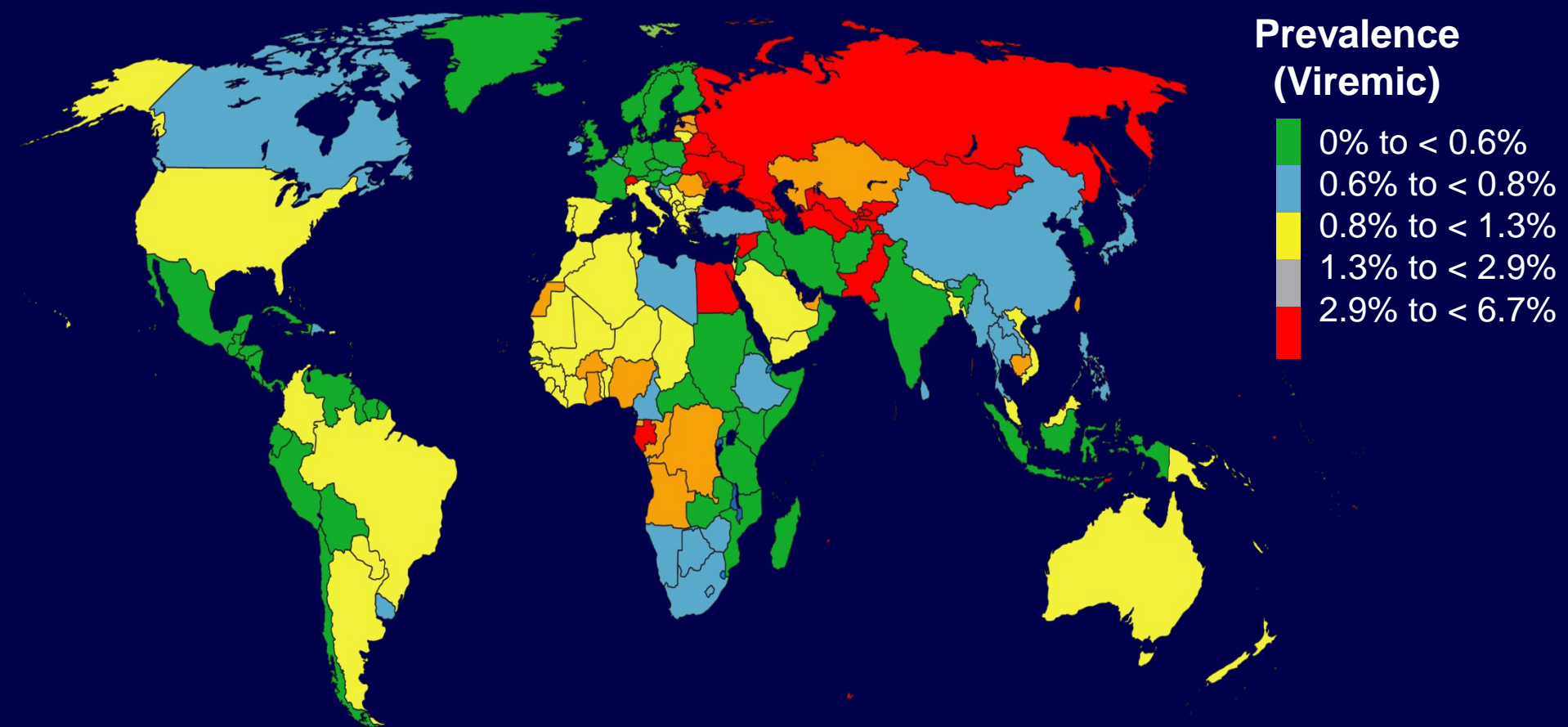
PegIFN		Nucleos(t)ide Analogues	
Pro	Con	Pro	Con
<ul style="list-style-type: none"> <li>▪ Finite course of therapy</li> <li>▪ No resistance</li> <li>▪ Higher rate of HBeAg loss in 1 yr</li> <li>▪ Higher rate of HBsAg loss with short duration therapy*</li> </ul>	<ul style="list-style-type: none"> <li>▪ SQ administration</li> <li>▪ Frequent AEs</li> <li>▪ Contraindicated in patients with cirrhosis, in pregnancy, with acute hepatitis B, and who are immunosuppressed</li> </ul>	<ul style="list-style-type: none"> <li>▪ PO administration</li> <li>▪ Infrequent AEs</li> <li>▪ Safe at all stages of disease, including decompensated cirrhosis†</li> <li>▪ Safe in immunocompromised populations</li> <li>▪ Selected drugs probably safe in pregnancy</li> </ul>	<ul style="list-style-type: none"> <li>▪ Need for long-term or indefinite therapy</li> <li>▪ Potential for drug resistance</li> </ul>

\*Particularly for HBeAg-positive patients with genotype A infection.

†Recent case report of lactic acidosis in severe liver failure.

Lok AS, et al. Hepatology. 2007;45:507-539. Lok AS, et al. Hepatology. 2009;50:661-662. Lok AS. Hepatology. 2010;52:743-747. Buster EH, et al. Gastroenterology. 2008;135:459-467. Lange CM, et al. Hepatology. 2009;50:2001-2006.

# Estimated 70 Million Persons Living With HCV

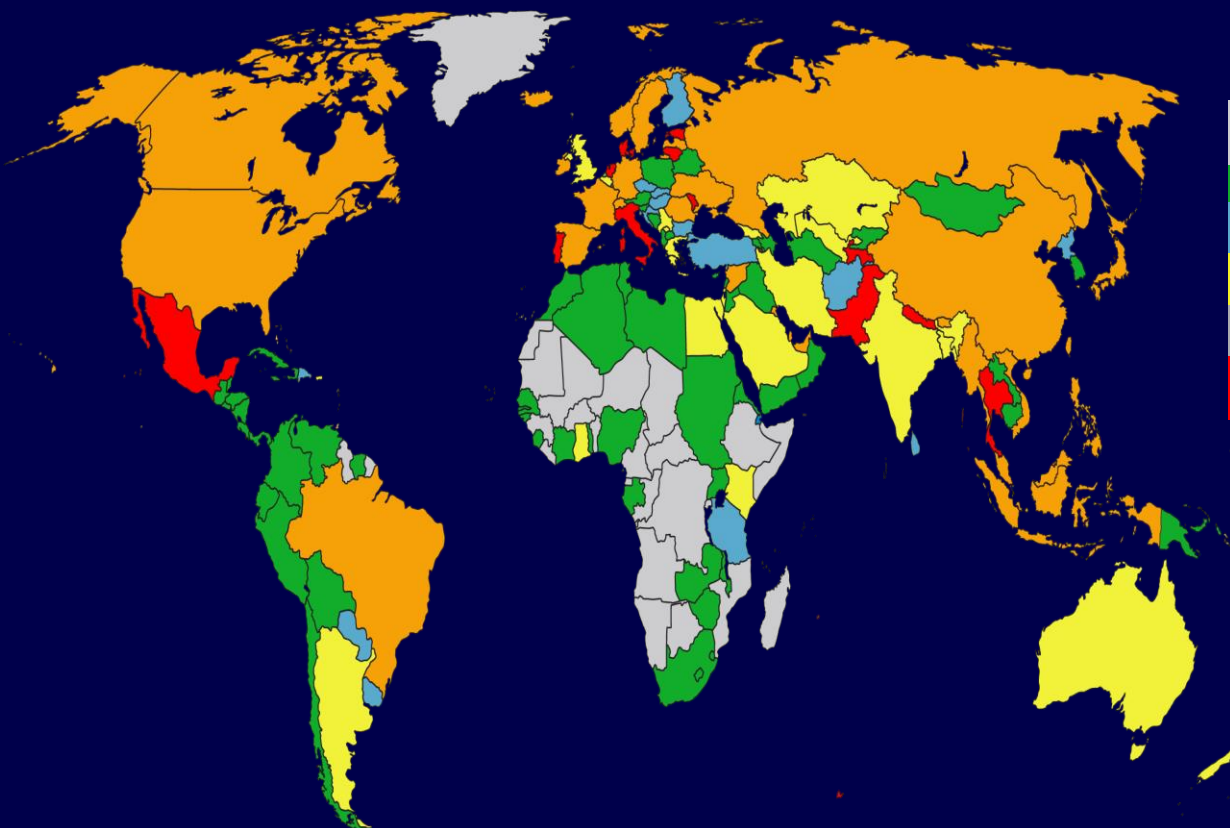


Polaris Observatory HCV Collaborators. Lancet Gastroenterol Hepatol.  
2017;2:161-176.



Slide credit: [clinicaloptions.com](https://clinicaloptions.com)

# HCV Infection Among Persons Who Inject Drugs



Anti-HCV Prevalence in PWID

- No evidence of injecting drug use
- No eligible report (74 countries)
- < 40% (16 countries)
- 40% to < 60% (24 countries)
- 60% to < 80% (25 countries)
- ≥ 80% (12 countries)

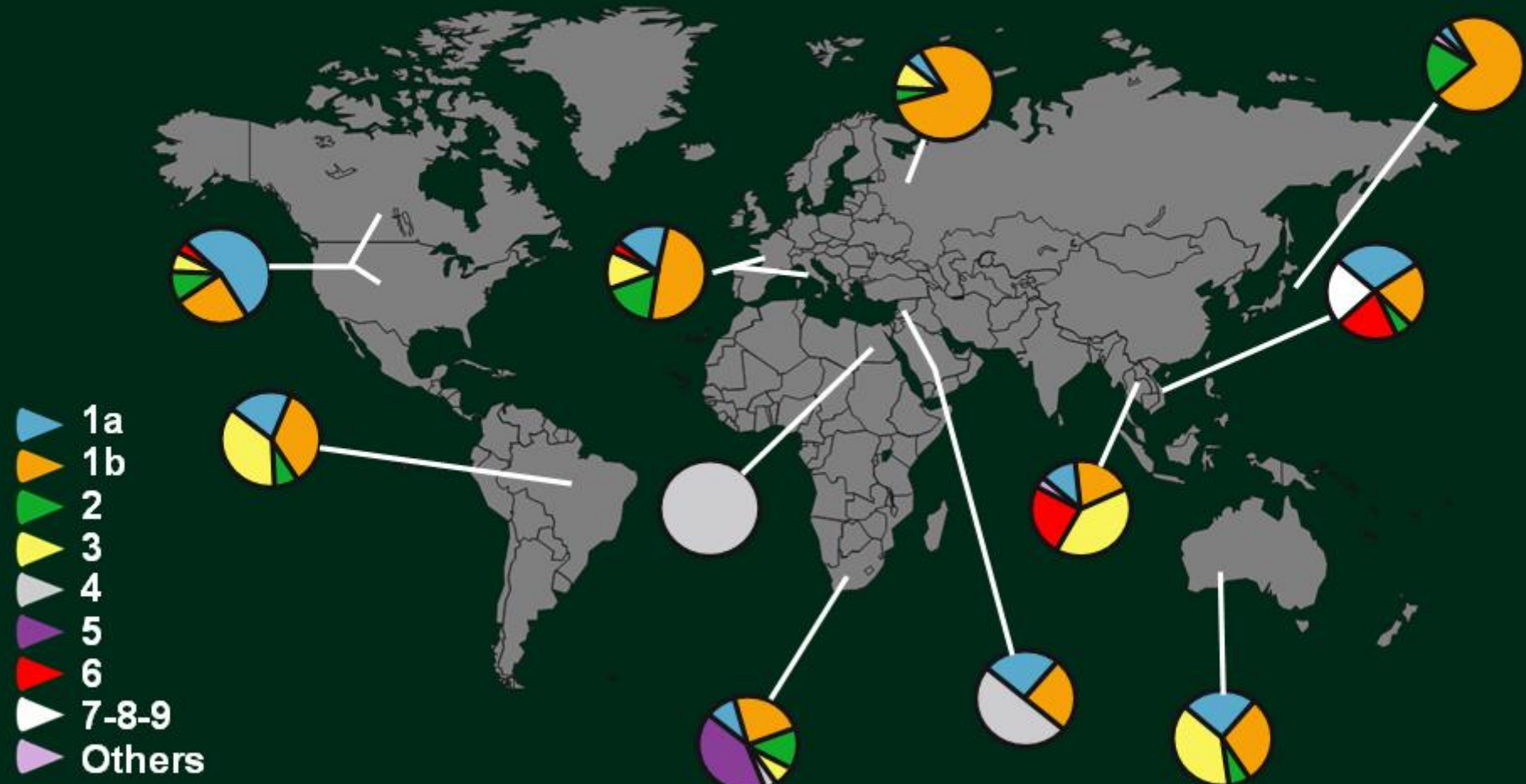
- In most developing countries, injection drug use is primary source of new infections
- HCV treatment must be coupled with harm reduction measures to reduce total infections

Nelson P, et al. Lancet 2011;378:571-583.



Slide credit: [clinicaloptions.com](#)

# Worldwide Distribution of HCV Genotypes



Zein N. Clin Microbiol Rev. 2000;13:223-235. Reproduced with permission.  
<http://cmr.asm.org/cgi/content/full/13/2/223?view=long&pmid=10755999>

# Chronic Hepatitis C Is a Progressive Disease



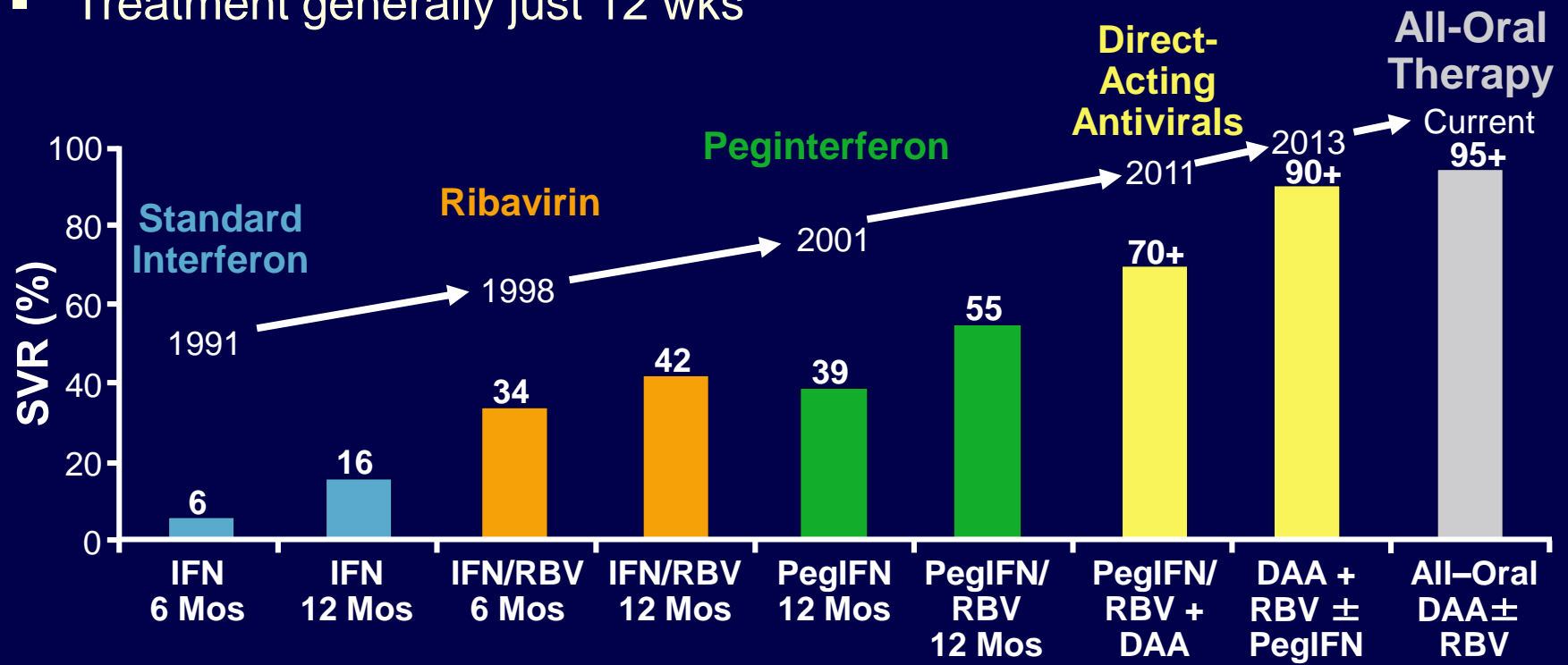
- Few or no symptoms; can progress without signs for decades<sup>[1]</sup>
- Most pts asymptomatic until serious liver complications arise<sup>[2]</sup>

1. CDC. MMWR Morb Mortal Wkly Rep. 1998;47(RR-19):1-39.

2. Heidelbaugh JJ, et al. Am Fam Physician. 2006;74:756-762.

# Nearly Everyone With HCV Can Now Be Treated Successfully

- Very high SVR rates; therapies highly tolerable
- All-oral therapy for almost every pt
- Treatment generally just 12 wks

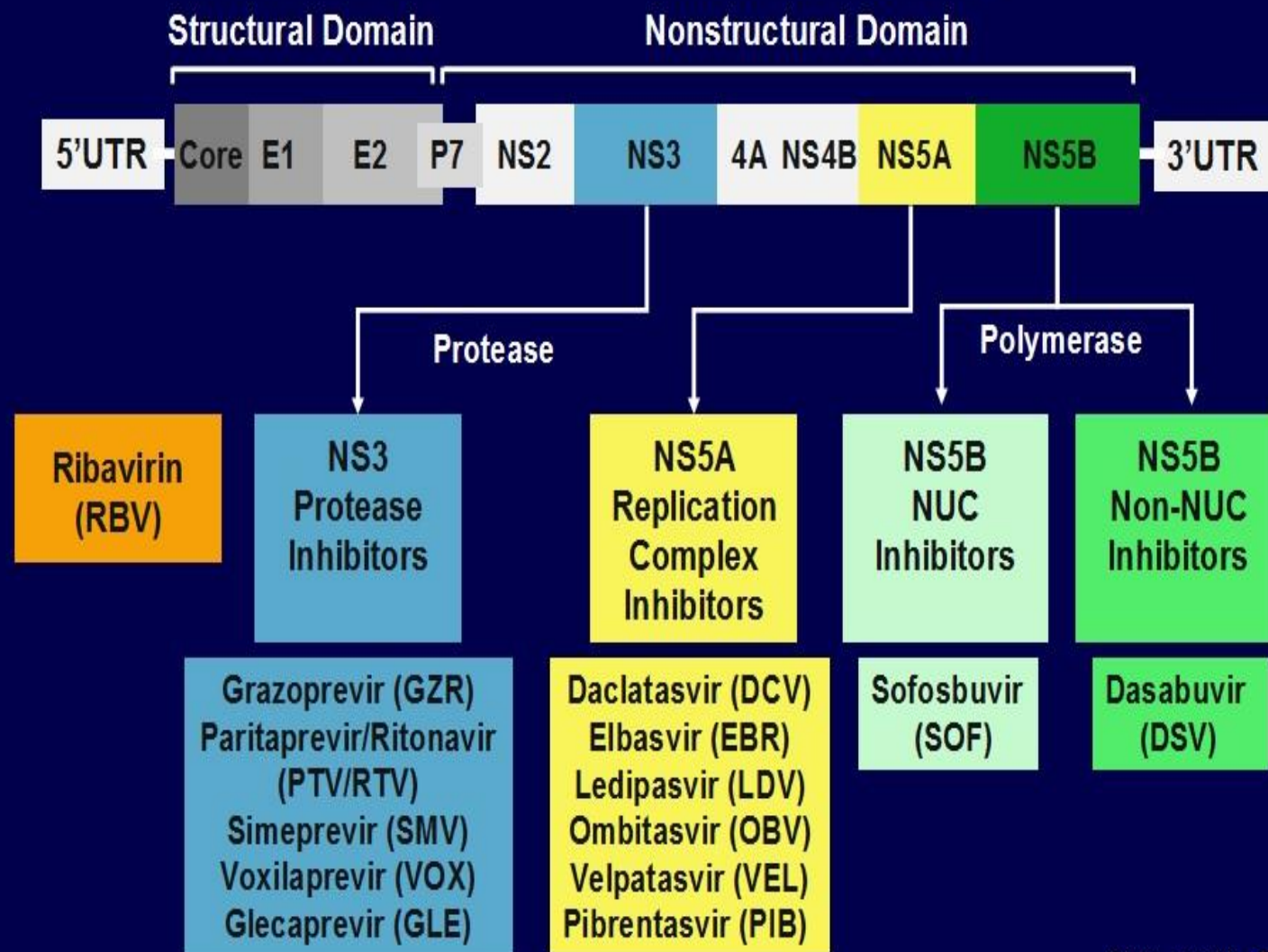


References in slidenotes.

Slide credit: [clinicaloptions.com](http://clinicaloptions.com)



# Approved DAAs From Multiple Classes: Basis of 2018 Combination HCV Regimens



# Dostupné varianty IFN-free režimů pro jednotlivé genotypy HCV (Doporučený postup ČHS ČLS JEP 2017)

kombinace	genotypy HCV				
	GT 1	GT 2	GT 3	GT 4	GT 5 a 6
sofosbuvir/ledipasivir ± ribavirin	ano	ne	ne	ano	ano
sofosbuvir/velpatasvir ± ribavirin	ano	ano	ano	ano	ano
ombitasvir/paritaprevir/(r) + dasabuvir ± ribavirin *	ano	ne	ne	ne	ne
ombitasvir/paritaprevir/(r) ± ribavirin *	ne	ne	ne	ano	ne
grazoprevir/elbasvir ± ribavirin	ano	ne	ne	ano	ne
sofosbuvir + daclatasvir ± ribavirin	ano	ano	ano	ano	
sofosbuvir + simeprevir ± ribavirin	suboptimální	ne	ne	ano	ne

**sofosbuvir+ledipasvir = HARVONIT™ (Gilead)**  
**sofosbuvir+velpatasvir = EPCLUSA™ (Gilead)**  
**ombitasvir+paritaprevir = VIEKIRAX™ (AbbVie)**  
**dasabuvir = EXVIERA™ (AbbVie)**  
**(r) = ritonavir**  
**grazoprevir+elbasvir = ZEPATIER™ (MSD)**  
**asunaprevir = SUNVEPRA™ (BMS)**

**sofosbuvir = SOVALDI™ (Gilead)**  
**daclatasvir = DAKLINZA™ (BMS)**  
**simeprevir = OLYSIO™ (Janssen)**  
**sofosbuvir+velpatasvir+voxilaprevir = VOSEVI™ (Gilead)**  
**glecaprevir+pibrentasvir = MAVYRET™ (AbbVie) - US**  
**glecaprevir+pibrentasvir = MAVIRET™ (AbbVie)-EU, Japan**

# EASL 2018 HCV Treatment Guidelines

- Recommendations for treatment-naive patients and patients previously treated with pegIFN/RBV ± SOF or SOF + RBV, ± compensated cirrhosis

HCV Genotype	SOF/VEL (Epclusa™)	GLE/PIB (Maviret™)	SOF/VEL/ VOX (Vosevi™)	LDV/SOF (Harvoni™)	EBR/GZR (Zepatier™)	OBV/PTV/ RTV + DSV (Viekirax™ + Exviera™)
1a	Yes	Yes	No*	Yes†	Yes‡	No
1b	Yes	Yes	No*	Yes	Yes	Yes
2	Yes	Yes	No*	No	No	No
3	Yes¶	Yes	Yes§	No	No	No
4	Yes	Yes	No*	Yes†	Yes#	No
5 or 6	Yes	Yes	No*	Yes†	No	No

\*Triple combination effective but not needed because double combinations comparably effective. †Tx naive ± compensated cirrhosis. ‡Tx naive or tx experienced without cirrhosis or with compensated cirrhosis and HCV RNA ≤ 800,000 IU/mL. ¶Tx naive or tx experienced without cirrhosis. §Tx naive or tx experienced with compensated cirrhosis.

#Treatment naive without cirrhosis or with compensated cirrhosis and HCV RNA ≤ 800,000 IU/mL.

# Dostupné varianty IFN-free režimů pro jednotlivé genotypy HCV (Doporučený postup ČHS ČLS JEP leden 2019)

HCV GT	Pangenotypové režimy			Genotypově-specifické režimy		
	SOF/VEL Epclusa™	GLE/PIB Maviret™	SOF/VEL/VOX Vosevi™	SOF/LDV Harvoni™	GZR/EBR Zepatier™	OBV/PTV/r Viekirax™ + DSV Exviera™
1a	Ano	Ano	Ne*	Ano <sup>a</sup>	Ano <sup>b</sup>	Ne
1b	Ano	Ano	Ne*	Ano	Ano	Ano
2	Ano	Ano	Ne*	Ne	Ne	Ne
3	Ano <sup>c</sup>	Ano	Ano <sup>d</sup>	Ne	Ne	Ne
4	Ano	Ano	Ne*	Ano <sup>a</sup>	Ano <sup>e</sup>	Ne
5	Ano	Ano	Ne*	Ano <sup>a</sup>	Ne	Ne
6	Ano	Ano	Ne*	Ano <sup>a</sup>	Ne	Ne

\* V této indikaci je kombinace SOF/VEL/VOX účinná, nicméně upřednostněno podání dvojkombinačních režimů

<sup>a</sup> Dosud neléčení bez CIH nebo s kompenzovanou CIH

<sup>b</sup> Dosud neléčení i v minulosti léčení bez CIH nebo s komp. CIH a HCV RNA ≤ 800 000 IU/ml

<sup>c</sup> Dosud neléčení i v minulosti léčení bez CIH

<sup>d</sup> Dosud neléčení nebo v minulosti léčení s kompenzovanou CIH

<sup>e</sup> Dosud neléčení bez CIH nebo s komp. CIH a HCV RNA ≤ 800 000 IU/ml

sofosbuvir+velpatasvir = EPCLUSA™  
glecaprevir+pibrentasvir = MAVIRET™  
sofosbuvir+velpatasvir+voxilaprevir = VOSEVI™  
sofosbuvir+ledipasvir = HARVONI™  
grazoprevir+elbasvir = ZEPATIER™  
ombitasvir+paritaprevir = VIEKIRAX™  
r = ritonavir (k potencování účinku)  
dasabuvir = EXVIERA™

## **SVR12 a SVR24**

**Cílem terapie je vyléčení HCV infekce = dosažení trvalé eliminace viru. Eliminace viru brání rozvoji jaterních i mimojaterních komplikací HCV infekce, včetně pokročilé jaterní fibrózy, CIH, dekompenzované CIH a HCC.**

**Eliminací infekce je myšleno dosažení setrvalé virologické odpovědi (SVR, sustained viral response) = dosažení negativní HCV RNA v séru ve 12. nebo 24. týdnu po skončení protivirové léčby (SVR12 a SVR24).**

**SVR12 a SVR24 spolu korelují v 99 % případů.**

**Podle dlouhodobých studií znamená dosažení SVR v 99 % případů trvalé vyléčení HCV infekce, tj. u osob se SVR nedochází k pozdním relapsům onemocnění.**

# Pangenotypální režimy

**Epclusa™ (Gilead)**

= sofosbuvir + velpatasvir

**Vosevi™ (Gilead)** – vhodný pro případy, kde po IFN-free léčbě nedojde k SVR

= sofosbuvir + velpatasvir + voxilaprevir

**Maviret™ (AbbVie)** – EU, Japan

**Mavyret™ (AbbVie)** - US

= glecaprevir + pibrentasvir

# Management of Pts With HCV Who Achieved SVR

- SVR associated with myriad clinical benefits
- Mandatory to continue HCC surveillance post SVR in pts with F3/F4 fibrosis; ultrasound every 6 mos
  - Consider assessing AFP levels as well for these pts
  - Less evidence to support continued screening of F0-F2 but remains a theoretical concern
- Indefinite screening for varices not warranted if varices absent at baseline: 1 more exam after SVR?
  - Surveillance of small varices if no other liver disease present requires further study but advisable
  - Large varices require ongoing management and surveillance
- Routine monitoring for fibrosis regression not the standard of care; further studies may change this
- For pts with ongoing risk for HCV infection after SVR, counsel and test HCV RNA annually



# Main extrahepatic manifestations in pts with HCV infection

(Ther Adv Infect Dis 2016;3(1):3-14)

## Immune-related extrahepatic manifestations

- Mixed cryoglobulinemia
- Cryoglobulinemic vasculitis
- B-cell non-Hodgkin's lymphoma
- Sicca syndrome (Sjögren's syndrome)
- Arthralgia/myalgia
- Autoantibody production (i.e. cryoglobulins, rheumatoid factor, and antinuclear, anticardiolipin, antithyroid and anti-smooth muscle antibodies)
- Polyarteritis nodosa
- Monoclonal gammopathies
- Immune thrombocytopenia

## Inflammatory-related extrahepatic manifestations

- Type 2 diabetes mellitus type 2
- Insulin resistance
- Glomerulonephritis (membranoproliferative)
- Renal insufficiency
- Fatigue
- Cognitive impairment
- Depression
- Impaired quality of life
- Polyarthritis/fibromyalgia
- Cardiovascular disorders (i.e. stroke, ischemic heart disease)

**Děkuji Vám  
za pozornost**

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