Hepatitis Delta Update



Hepatitis Delta

- Caused by the hepatitis delta virus (HDV)
- "Satellite virus"
 - Infects only those with hepatitis B (HBV)
- Can lead to acute or chronic disease
- May be acquired simultaneously with HBV as co-infection
- May be acquired by chronically infected HBV patients

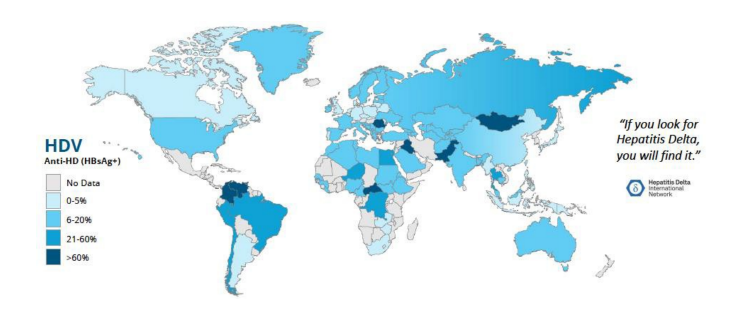
Epidemiology

HDV Worldwide Prevalence: 15 – 20 Million

- Common in Eastern Europe, Southern Europe, Mediterranean region, Middle East, West and Central Africa, East Asia, Amazon Basin
 - > 100 K in US
 - > 200K in EU
 - > 2M in China
- In US, at least 4% of HBV infected patients have HDV

HDV Worldwide Prevalence: 15 - 20 Million

6% of HBV population coinfected with HDV



HDV Has 8 Genotypes

Genotype	Commonly Found		
1	North America, Europe, Middle East, North Africa		
2 and 4	East Asia		
3	Amazon Basin of South America		
5,6,7,8	West and Central Africa		

HDV Geographic Footprint Is Growing

U.S. major metro hotspots identified



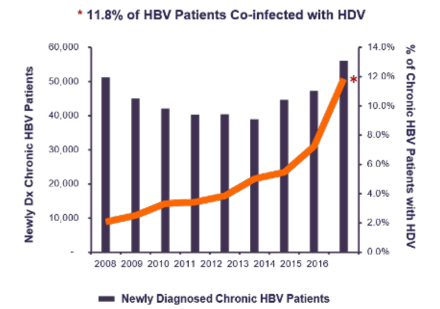
Top 10 U.S. Cities in 2016 1. Chicago, Illinois 2. Berwyn, Illinois 3. Brooklyn, New York 4. Corona, New York 5. Waukegan, Illinois 6. New York, New York 7. Bronx, New York 8. Jamaica, New York 9. Lombard, New York 10. Aurora, Illinois

Martins, et al. DDW. 2017.

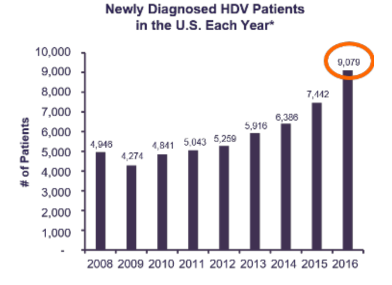
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U.S. HDV Prevalence in 2018: ~110,000

Increased screening leads to increased HBV and HDV diagnosis

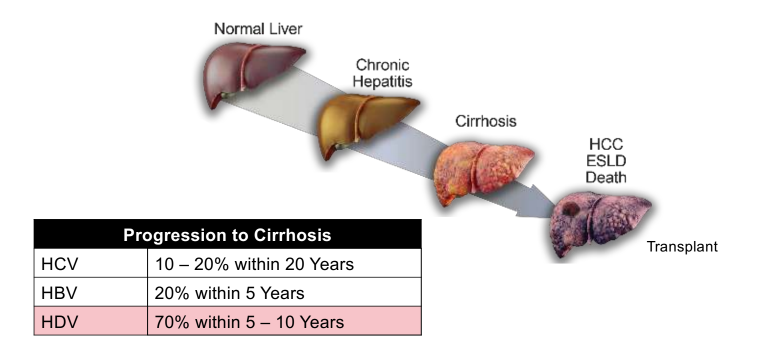


— % of Chronic HBV Patients with HDV



Martins, et al. DDW. 2017.

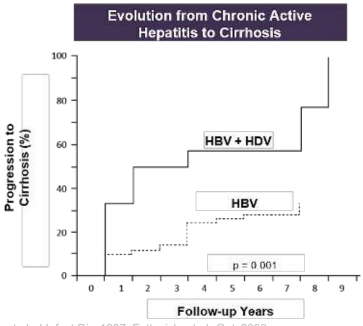
HDV: Most Rapid Progression of Viral Hepatitis

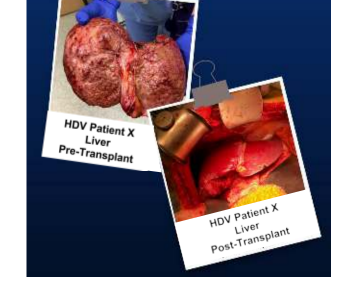


Westbrook, et al. J Hepatology. 2014; Fattovich, et al, Seminars in Liver Diseases. 2003; Nourredin, et al. Curr Gasterol Rep. 2013.

HDV Causes Most Rapid Disease Progression

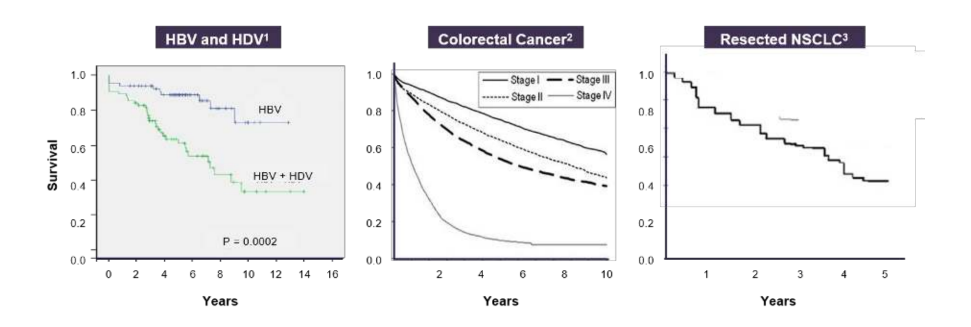
At diagnosis, > 50% of HDV patients are cirrhotic





Fattovich, et al. J Infect Dis. 1987; Fattovich, et al. Gut. 2000.

Survival: HDV vs Cancer



¹Serrano, et al. EASL. 2011; ²Cancer Causes Control. 2012. 23:1421–1428; ³Cerfolio, et al. *Ann Thorac Surg.* 2007. 84:182–90.

Risk Factors

Hepatitis Delta Risk Factors

- Birth to an infected mother
- Intravenous drug use
- Sex with an infected partner
- Contact with blood from or open sores of an infected person
- Needle sticks or exposures to sharp instruments
- Sharing items (razors, toothbrushes, etc.) with an infected person
- HDV endemic country of origin
- Elevated ALT despite suppressed HBV DNA

Populations at Risk For HDV Infection

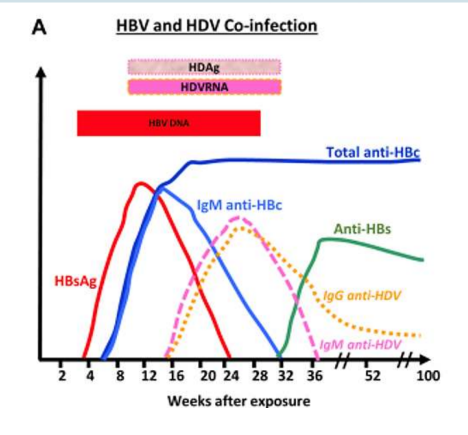
- People chronically infected with HBV
- Infants born to mothers with HDV
- Sex partners of persons infected with HDV
- MSM
- Injection drug users
- Household contacts of people with HDV
- Health care and public safety workers at risk for occupational exposure to blood or blood-contaminated fluids
- Hemodialysis patients

How Hepatitis Delta is Not Spread

- Through food or water
- Sharing eating utensils
- Breastfeeding
- Hugging
- Kissing
- Hand holding
- Coughing or sneezing

HBV/HDV Coinfection

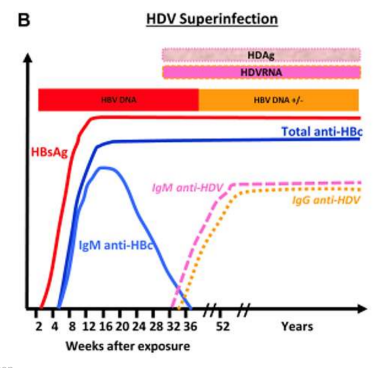
- Simultaneous infection with HBV and HDV
- Usually resolves
- > 5% develop chronic disease
- Acute Liver Failure more common with HDV/HBV than HBV alone



WHO 2019 Smedile et al. Lancet 1982.

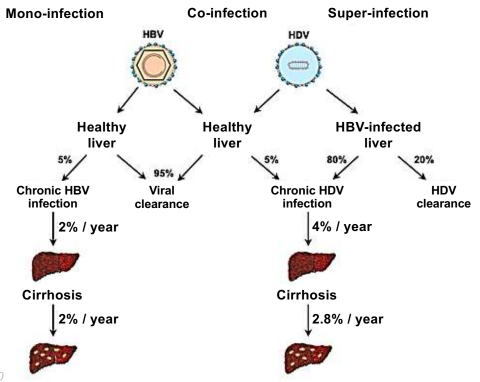
HDV Superinfection

- Occurs when a person already with chronic HBV acquires HDV
- Results in rapid progression to cirrhosis and liver failure
 - 15% in 1-2 years
 - 70-80% in 5-10 years
- Higher risk of HCC
 - 3-6 fold increase
- Higher risk of mortality
 - 2 fold increase



National Institute for Health. National Institute of Diabetes and Digestive and Kidney Diseases. United States 2017. Department of Health and Human Services. Available at https://www.niddk.nih.gov/health-information/liver-disease/viral-hepatitis/hepatitis-d#common Da, et al. Gastro Report. 2019

Natural History of HBV and HDV Infection



Turon Lagot, et al. J Clin Med. 2020

HDV: Signs/Symptoms

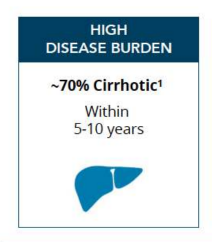
- Fever
- Fatigue
- Loss of appetite
- Nausea
- Vomiting

- Abdominal Pain
- Dark urine
- Clay-colored stools
- Joint pain
- Jaundice

These signs typically appear 3-7 weeks after infection.

HDV Clinical Course and Outcomes

HDV: A devastating disease with no approved treatment







25% of People on Waiting List Die Each Year Before Receiving a Liver Transplant¹

¹Nourredin, et al. Curr Gasterol Rep. 2013; ²Serrano, et al. EASL. 2011; ³UPMC Health Beat. 2018. US liver transplant cost.

Prevention

- Risk factor avoidance
- Hepatitis B vaccine
- No vaccine available for HDV

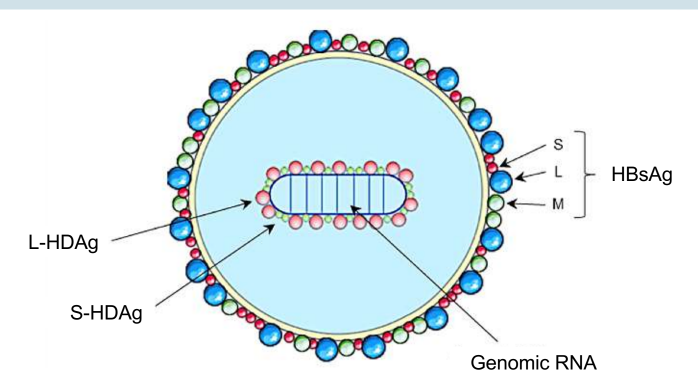
Molecular Biology

HDV Structure

- Positive single stranded circular RNA genome of ~1700 nucleotides
- HDV RNA genome is ~74% of paired bases giving it a rod-like structure
- Encodes HDAg, in 2 forms
 - Small (S-HDAg)
 - Large (L-HDAg)
 - HDAg associated with HDV genome forming a ribonucleoprotein (RNP)
 - RNP is enveloped by HBsAg

Turon Lagot, et al. J Clin Med. 2020

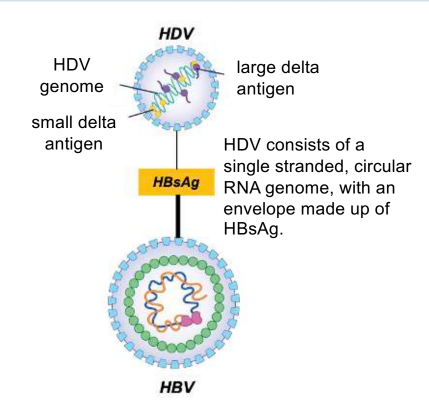
HDV Structure



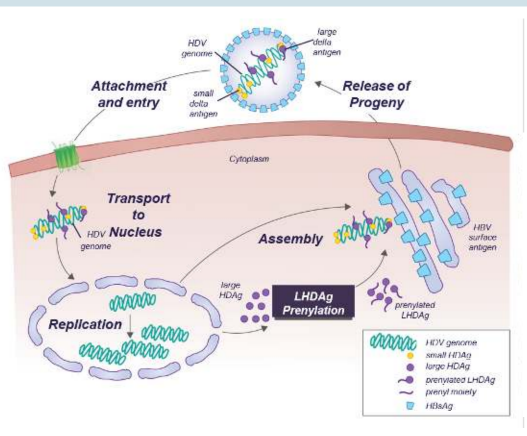
Turon Lagot, et al. J Clin Med. 2020

HDV Requires HBsAg to Complete Virus Assembly

 Assembly with HBsAg is mediated by large delta antigen prenylation



HDV Life Cycle



- HDV genome encodes for a single protein, the hepatitis delta antigen.
- HDV relies on host cell machinery for replication.
- New virions can be assembled only in the presence of hepatitis B virus.

Diagnosis & Natural History

HDV: Diagnostic Tests

- HDV antibodies (IgG, IgM)
- HDV RNA
- HBsAg

HDV RNA Quantification is the Gold Standard in HDV Diagnosis and Management

- A leading provider of diagnostic services
- Over 2,200 patient service centers across the US
- Highly targeted patient and physician outreach
- HDV testing program for HBV-positive patients
- HDV RNA quantification and HBV/HDV reflex testing



Treatment

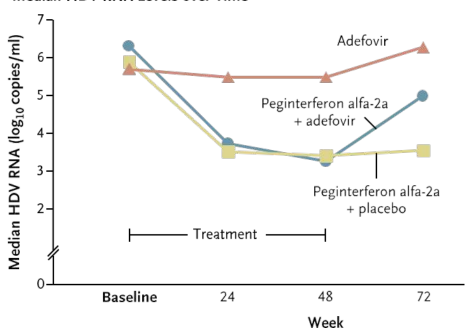
HDV Treatment

- No FDA approved therapy for HDV
- PEG IFN alpha is only drug with any demonstrated efficacy
- 20-25% response rate and rarely achieve sustained viral clearance
- Suboptimal due to:
 - Significant side effects (including cytopenias)
 - Limited efficacy
 - High long-term relapse rates

Turon Lagot, et al. J Clin Med. 2020

HDV Treatment

Median HDV RNA Levels over Time

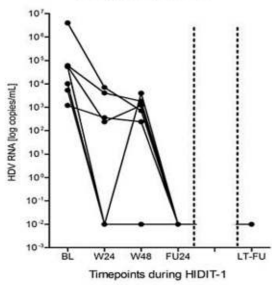


- ~2.5 log decline in median HDV RNA at EOT
- 25% neg 24 wks post Rx in patients receiving PEG-IFN
- 0% with nucleoside analog alone (no effect on HBsAg)

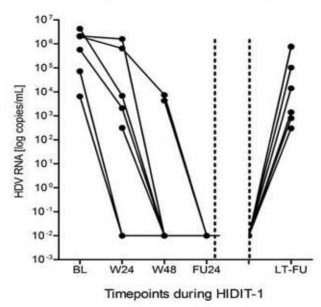
Late Relapse is Common with PEG IFN alpha

Late Kelapse is Common with Let in it diplie





HDV RNA of patients with late relapse

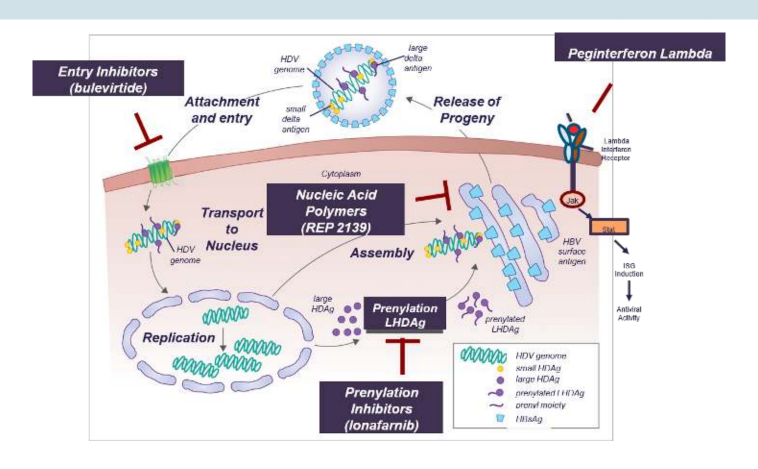


- Long term f/u of HIDIT-1
- Of 16 patients that were negative for HDV RNA at 6 months after therapy ended, 9 will test positive during long term f/u

HBV Functional Cures Will Not Eradicate HDV

- Approved HBV nucleos(t)ide treatments only suppress HBV DNA
 - They do not affect HBsAg and have no effect on HDV
- Investigational HBV treatments target functional cure
 - Not expected to completely eliminate HBsAg needed by HDV

HDV Treatments in Development



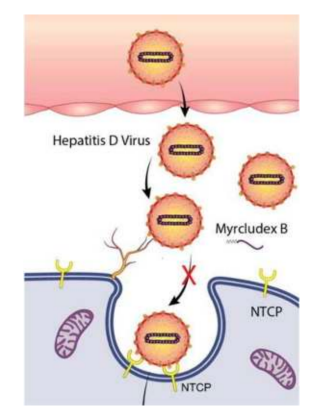
HDV Treatments in Development

Treatment	Company	Stage of Development	
		Phase 2	Phase 3
Lonafarnib	Fr EIGER	Daily Oral	
Peginterferon Lambda	EIGER ANNIMALIE	Weekly SQ injection	
Bulevirtide	Gilead	Daily SQ injection	
Rep-2139	replicor	Weekly IV infusion	19

Bulevirtide: Entry Inhibitor Targeting NTCP

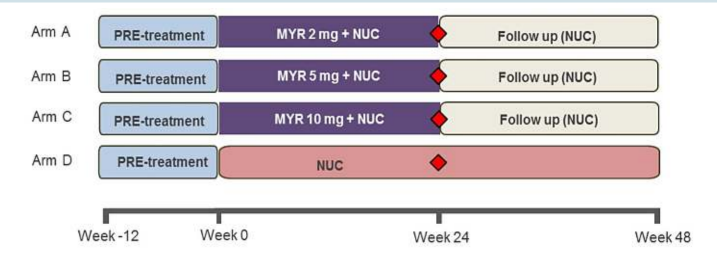
- Daily, subcutaneous injections
- Synthetic 47 amino acid, N-acylated preS1 lipopeptide
- Targets Na-taurocholate cotransporting polypeptide (NTCP)
 - On surface of hepatocytes
- Blocks receptor functions of NTCP and

HBV/HDV virus entry



Bogomolov et al. J Hep 2016

Phase 2b, Open-Label Bulevirtide / TDF Study



Primary Endpoint

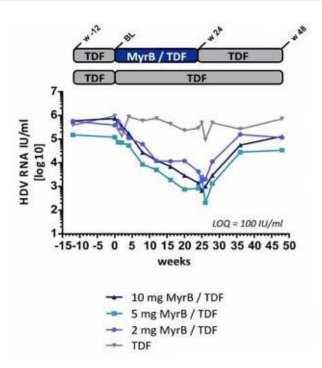
 HDV RNA < LLOD at 24 weeks post-tx (Week 72)

Secondary Endpoint

- HDV RNA < LLOD at Week 48
- ALT nl at Week 48, 72
- HDV RNA > 2 log decline and ALT nl at Week 48, 72
- HBsAg undetectable or > 1 log decline at Week 48, 72

Wedemeyer, et al. EASL. 2018.

Phase 2b, Open-Label Bulevirtide / TDF Study

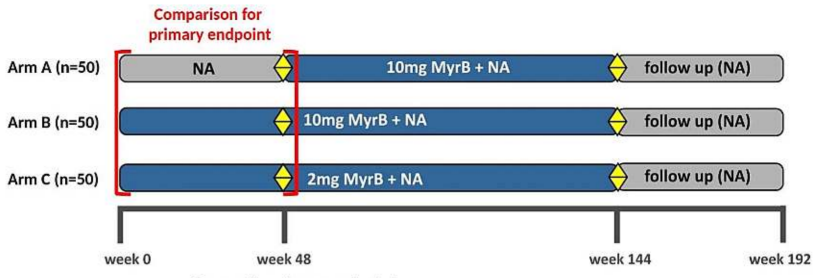


Regimen	Median HDV RNA Change from Baseline at Week 24		
2mg MyrB/TDF	-1.75		
5mg MyrB/TDF	-1.60		
10mg MyrB/TDF	-2.70		
TDF	-0.18		

Modeling suggested 2-3 years of continuous treatment might be needed to achieve HDV RNA negativity. Wedemeyer, et al. EASL. 2018.

Phase 3 Bulevirtide Study

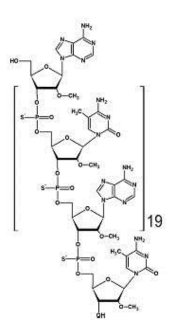
Across 7 Sites in Russia



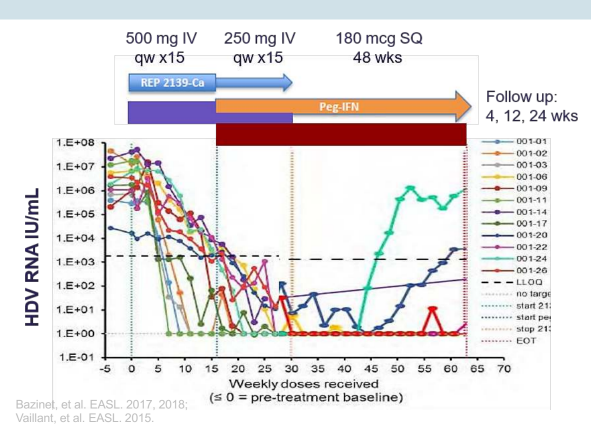
Composite primary endpoint: HDV RNA negativation or >2log decline as well as ALT normalization

REP 2139

- Nucleic acid polymers (NAPs) are oligonucleotides with broad spectrum in vitro antiviral activities
- Reported to act via entry inhibition in other viruses
- Also proposed to bind to amphipathic protein structures
- These amphipathic protein structures are common in viral proteins, but are also found in key host cell proteins
- REP 2139 inhibits secretion of HBsAg from cells



Phase 2 REP 301 Study

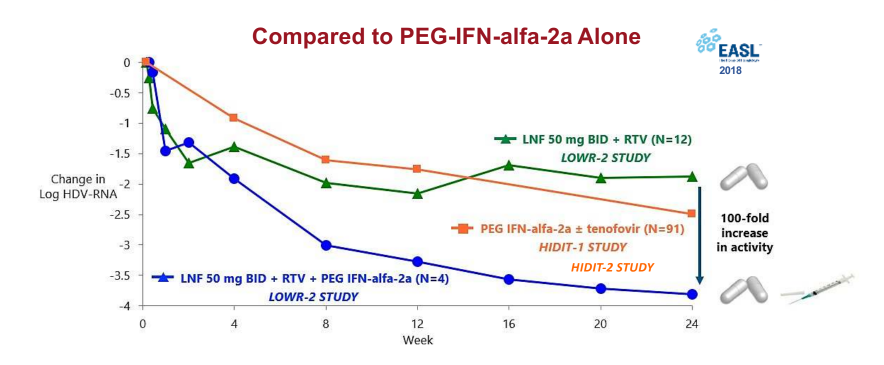


- 12 HDV pts in Moldova
- Anti-HDV/RNA +
- HBsAg > 1000
- Non-cirrhotic
- Responses mostly maintained on interferon
- 5 patients rebound with cessation of IFN (EASL 2017)
- Responses maintained to date (EASL 2018)

Lonafarnib

- Small molecule, first-in-class, oral prenylation inhibitor
- Well-characterized through Phase 3
 - > 2,000 subjects dosed in oncology program by Merck (Schering)
 - Dose limiting toxicity is GI (class effect)
- Over 120 HDV patients dosed across international sites
- US & EU Orphan Designation, FDA Breakthrough and EMA PRIME Designation
- Broad range of lonafarnib + ritonavir doses and durations studied
- US and multiple international sites

LOWR: Phase 2 Lonafarnib Study



Yurdaydin, et al. EASL. 2018. LOWR-2; Wedemeyer. et al. 2019. HIDIT-2.

Peg IFN Lambda: A Better Tolerated Interferon

- A novel, first-in-class Type III interferon
- Binds to a unique receptor versus Type I interferons
 - Highly expressed on hepatocytes
 - Limited expression on hematopoietic cells and CNS cells
- Uses similar downstream signaling pathway as Type I interferons
- Studied in > 3,000 subjects across 17 clinical trials (HCV / HBV)
- Comparable antiviral activity with less of the typical IFN alfa related side effects

IFN-a receptors widely distributed throughout body.

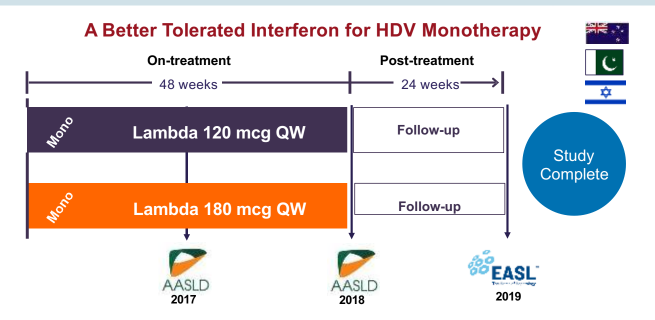


Lambda receptors NOT widely distributed throughout body.



Chan, et al. *J Hepatol*. 2016 May;64(5):1011-1019.

LIMT: Phase 2 Lambda Study



Primary Endpoint:

 Evaluate Safety, Tolerability, Efficacy

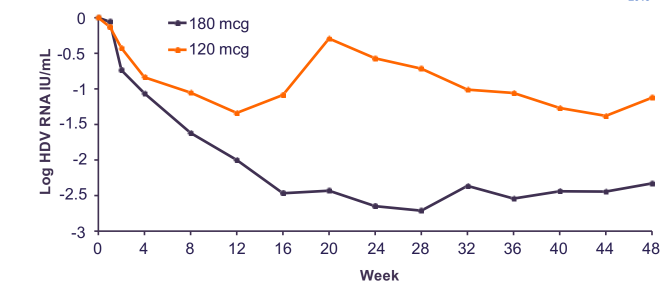
Secondary Endpoint:

 Proportion of Patients with HDV RNA BLQ 24 weeks after EOT

Peginterferon Lambda Activity Through Week 48

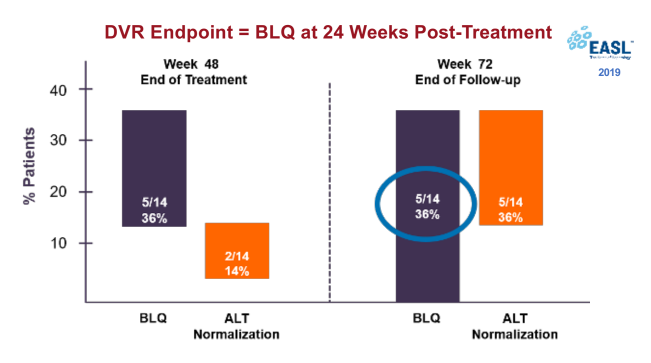
Lambda 180 mcg has Comparable Antiviral Activity to Alfa 180 mcg with Improved Tolerability





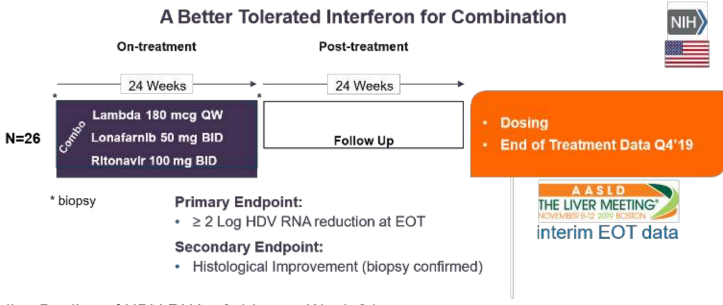
Etzion O, et al. 2017. Robogene® 2.0 HDV RNA PCR assay. LLOQ = 14 IU/mL.

Peginterferon Lambda: 36% Durable Virologic Response



Etzion et al, EASL 2019, LIMT Phase 2 Study; Robogene® 2.0 HDV RNA PCR assay, LLOQ = 14 IU/mL; DVR = BLQ at 24 Weeks Post-Treatment

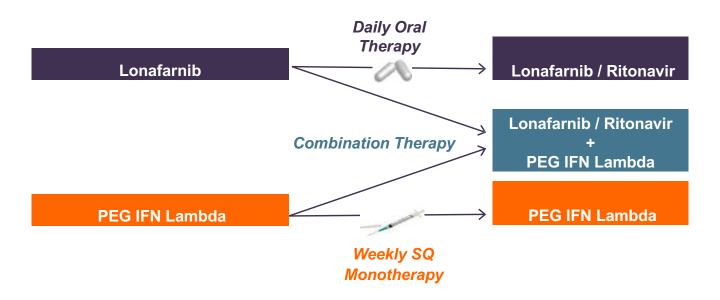
LIFT: Phase 2 Lambda & Lonafarnib Combination Study



- Median Decline of HDV RNA: -3.4 Log at Week 24
- 95% of Patients Achieve > 2 Log Decline in HDV RNA at Week 24
- > 50% of Patients Achieve Undetectable or BLOQ HDV RNA at Week 24

First-in-Class Treatments in Development for HDV

Multiple Options to Treat HDV



Lonafarnib, Ritonavir, and Lambda Interferon for HDV: Interim End-of-Treatment Results – the LIFT Study

Aims:

Evaluate the safety and antiviral effects of therapy with lonafarnib (LNF), ritonavir (RTV), and lambda interferon (LMD) in patients with chronic hepatitis D

Methods:

Phase 2a, open-label, prospective treatment trial in 26 patients for 24 weeks, with 24 weeks of post-therapy follow-up

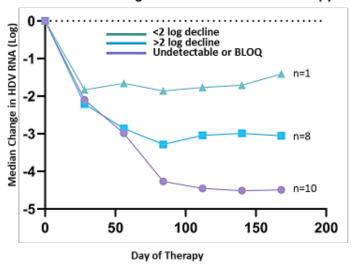
Main Findings:

At the end of therapy (19 of 26 patients), the median HDV RNA decline was 3.4 log IU/ml (p<0.0001) with 10 (53%) patients achieving undetectable or BLOQ HDV RNA in serum.

Conclusions:

Triple combination therapy with LNF/RTV/LMD in chronic HDV patients appears to be safe and tolerable for up to 6 months in most patients.

HDV RNA Change from Baseline to End of Therapy

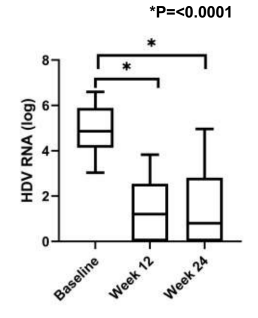


58% of patients achieved undetectable or BLOQ* HDV RNA by the end of therapy

Koh C, et al., Abstract LO8

LIFT Treatment Response at Week 12 and 24

Treatment Duration	n	Mean Change in log HDV RNA (IU/mL)	Confidence Interval	p-value
12 Weeks	26	3.36	(2.86-3.85)	<0.0001
24 Weeks	22	3.21	(2.50-3.93)	<0.0001



Summary

- HDV is the most severe form of viral hepatitis
- HDV remains underdiagnosed
- HBV vaccination is protective against acquiring HDV
- HDV/HBV infection causes more rapid disease progression as compared to HBV monoinfection
- New therapies being developed based upon HDV life cycle
- Several HDV therapies currently in phase 2 and 3 development
 - Phase 2 Bulevirtide and Ionafarnib
 - Phase 3 Lonafarnib