Treatment Naïve and Treatment Experienced

# Ledipasvir-Sofosbuvir in HCV Genotype 4 NIAID SYNERGY (Genotype 4)

Source: Kohli A, et al. Lancet Infect Dis. 2015:15:1049-54.



### Ledipasvir-Sofosbuvir in Genotype 4 NIAID SYNERGY (GT4) Trial: Features

#### **NIAID SYNERGY Trial**

- Design: Open-label, phase 2a trial using fixed dose ledipasvir-sofosbuvir for treatment-naïve and interferon treatment-experienced patients with chronic HCV genotype 4
- Setting: single center (Clinical Center at NIH, United States)
- Entry Criteria
  - Age 18 years or older
  - Chronic HCV genotype 4
  - Treatment naïve or prior interferon treatment failure
  - HCV RNA ≥2,000 IU/mL
  - Exclusions: HBV, HIV, or decompensated liver disease
- Primary End-Point: SVR12



### Ledipasvir-Sofosbuvir in Genotype 4 NIAID SYNERGY (GT4) Trial: Features

Week 0 12 24

#### **Genotype 4**

Treatment Naïve (n = 13)
Treatment Experienced (n = 8)

n = 21 **Ledipasvir-Sofosbuvir** SVR12

#### **Drug Dosing**

Ledipasvir-sofosbuvir (90/400 mg): fixed dose combination; one pill once daily



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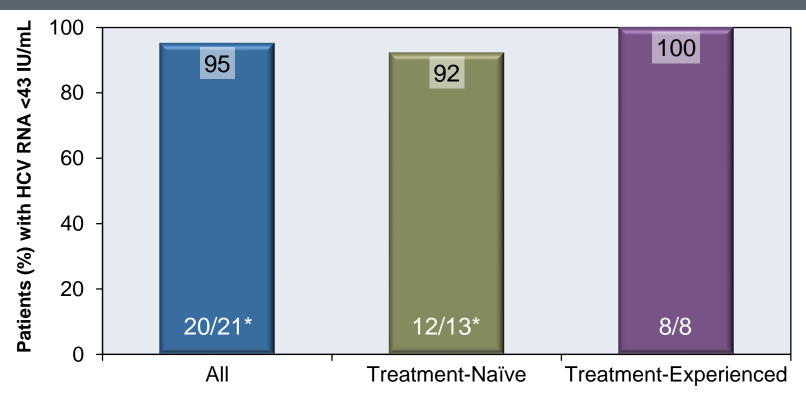
### Ledipasvir-Sofosbuvir in Genotype 4 NIAID SYNERGY (GT4) Trial: Key Baseline Characteristics

- Sex: Male 67%
- Race: 43% Black; 52% White; 5% Native American
- Country of Origin: 29% Egypt; 24% United States
- Treatment Experience: 62% naïve; 38% experienced
- HCV RNA >800,000 IU/mL: 62%



## Ledipasvir-Sofosbuvir in Genotype 4 NIAID SYNERGY (GT4) Trial: Results

NIH SYNERGY: SVR 12, Intent to Treat Analysis



**Ledipasvir-Sofosbuvir Treated Patients** 



<sup>\*1</sup> patient did not complete 12 weeks of treatment due to drug non-adherence

### Ledipasvir-Sofosbuvir in Genotype 4 NIAID SYNERGY (GT4) Trial: Interpretation

Interpretation: "Ledipasvir and sofosbuvir treatment for 12 weeks was well tolerated by patients with HCV genotype 4 and resulted in 100% SVR for all patients who received all 12 weeks of study drugs, irrespective of previous treatment status and underlying liver fibrosis. This is the first report of a single-pill, all-oral, interferon-free, ribavirin-free treatment for patients with HCV genotype 4."

