



# Advancing Life-Changing Therapies for Patients with Serious Diseases

**September 2026**

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The information in this presentation contains forward-looking statements that are subject to certain risks and uncertainties that could cause actual results to materially differ. These risks and uncertainties include: Assembly Bio's ability to realize the potential benefits of its collaboration with Gilead Sciences, Inc. (Gilead), including all financial aspects of the collaboration and equity investments; Assembly Bio's ability to initiate and complete clinical studies involving its therapeutic product candidates, including studies contemplated by Assembly Bio's collaboration with Gilead, including studies conducted by Gilead, in the currently anticipated timeframes or at all; safety and efficacy data from clinical or nonclinical studies may not warrant further development of product candidates; clinical and nonclinical data may not differentiate product candidates from other companies' candidates; Assembly Bio's ability to maintain financial resources and secure additional funding necessary to continue its research activities, clinical studies, and other business operations; potential effects of changes in government regulation; results of nonclinical studies may not be representative of disease behavior in a clinical setting and may not be predictive of the outcomes of clinical studies; and other risks identified from time to time in Assembly Bio's reports filed with the U.S. Securities and Exchange Commission (the SEC). You are urged to consider statements that include the words may, will, would, could, should, might, believes, hopes, estimates, projects, potential, expects, plans, anticipates, intends, continues, forecast, designed, goal or the negative of those words or other comparable words to be uncertain and forward-looking. Assembly Bio intends such forward-looking statements to be covered by the safe harbor provisions contained in Section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended. More information about Assembly Bio's risks and uncertainties are more fully detailed under the heading "Risk Factors" in Assembly Bio's filings with the SEC, including its most recent Annual Report on Form 10-K, Quarterly Reports on Form 10-Q and Current Reports on Form 8-K. Except as required by law, Assembly Bio assumes no obligation to update publicly any forward-looking statements, whether as a result of new information, future events or otherwise.





### PIPELINE INCLUDES MULTIPLE CLINICAL STAGE INVESTIGATIONAL THERAPIES

- Focused on areas with high unmet medical need and significant market opportunity
- Three Phase 2 studies initiating across the portfolio in 2026 and early 2027



### EXPERIENCED LEADERSHIP AND R&D ORGANIZATION

- R&D team with over 15 approved drugs in viral and liver diseases



### INDUSTRY LEADING PARTNER IN GILEAD

- Collaboration brings together the teams' expertise in virology and provides funding opportunities and an established partner for late-stage development and commercialization

# Developing Differentiated Approaches for Treating Serious Diseases

| Program     | Indication                                 | Mechanism      | Preclinical | IND/CTA enabling | Phase 1 | Phase 2 | Partner              |
|-------------|--------------------------------------------|----------------|-------------|------------------|---------|---------|----------------------|
| ABI-6250    | Hepatitis D                                | NTCP inhibitor |             |                  |         |         | Gilead opt-in rights |
|             | PBC and PSC                                |                |             |                  |         |         |                      |
| ABI-7272    | Transplant associated herpesviruses        | NNPI           |             |                  |         |         | Gilead opt-in rights |
| Undisclosed | Research programs against multiple targets |                |             |                  |         |         | Gilead opt-in rights |

## Partner Directed – GS-1179 selected to advance

|         |                      |                 |             |  |  |  |                                                                                                                                                       |
|---------|----------------------|-----------------|-------------|--|--|--|-------------------------------------------------------------------------------------------------------------------------------------------------------|
| GS-1179 | Herpes simplex virus | Long-acting HPI | <div></div> |  |  |  | <div> <b>GILEAD</b><br/>ASMB: 40%<br/>U.S. profit/cost share</div> |
| GS-5366 |                      |                 | <div></div> |  |  |  |                                                                                                                                                       |

## Exploring Partnering Opportunities

|          |             |                     |             |  |  |  |  |
|----------|-------------|---------------------|-------------|--|--|--|--|
| ABI-4334 | Hepatitis B | Next-generation CAM | <div></div> |  |  |  |  |
|----------|-------------|---------------------|-------------|--|--|--|--|



# Projected Clinical Milestones and Data Readouts

## LIVER DISEASES

Phase 2 initiation  
expected **4Q 2026**

Data expected  
**4Q 2027**

**ABI-6250**

**HDV**  
NTCP inhibitor

**ABI-6250**

**PBC/PSC**  
NTCP inhibitor

Phase 2 initiation  
expected **1Q 2027**

Data expected  
**1H 2028**

## HERPES SIMPLEX VIRUS

**GS-1179**

Long-acting  
helicase-primase  
inhibitor

Advancing under collaboration with Gilead with  
Phase 2 initiation in recurrent genital herpes  
expected **4Q 2026**

**Cash Runway into 2029** – backed by a ~\$320.4M cash position at June 30, 2026, plus \$75M  
Gilead collaboration extension payment due Q4 2026



# Gilead Collaboration

## OVERVIEW

- Long-term collaboration with Gilead entered into in October 2023
- Brings together two teams' knowledge and expertise in antiviral research, clinical development and commercialization
- Gilead holds option rights on programs at end of Phase 1 or Phase 2
- Assembly Bio may advance internally or partner externally programs upon Gilead opt-out

## OPTION STRUCTURE

- Assembly Bio primarily responsible for R&D before opt-in
- Upon opt-in, Gilead leads all development and commercialization at its cost
- Global milestones and royalties or 40% U.S. profit/cost share option and ex-U.S. milestones and royalties

*Gilead has exercised its option on the HSV program and Assembly Bio has elected the 40% U.S. profit/cost share*

## FUTURE ECONOMICS

- \$75M collaboration extension payments due 4Q 2026, 2028 and 2030
- $\geq$  \$45M opt-in fee per program (depends on stage at opt-in)
- Eligible for up to ~\$330M milestones per program (up to ~\$280M ex-U.S. when 40% U.S. profit/cost share elected)
- Royalties: high single-digits to high-teens per program, no profit/cost share; high single-digits to low teens ex-U.S. for HSV

## GS-1179 and GS-5366 (formerly ABI-1179 and ABI-5366)

Long-Acting HSV Helicase-Primase Inhibitors (HPIs)  
for Recurrent Genital Herpes

**GS-1179 and GS-5366 – Phase 1b completed dosing and follow up  
Gilead selected GS-1179 to advance; Phase 2 anticipated by year-end 2026**

# Genital Herpes is a Serious Condition that Impacts Millions of Individuals in the US/EU

## MILLIONS AFFECTED IN US/EU5



**4M+**

recurrent (3+/yr)  
genital herpes<sup>1,2</sup>

**8M+**

diagnosed with  
genital herpes<sup>3</sup>

**60M+**

people living  
with HSV-2<sup>4,5</sup>

## SERIOUS HEALTH IMPACTS



### PROLONGED PAIN AND SYMPTOMS

Painful lesions, lymphadenopathy and urinary problems that can persist 2-3 weeks<sup>6</sup>



### FREQUENT RECURRENCES

Most people with an initial symptomatic genital HSV-2 infection experience frequent recurrences (3-15 times in a year)<sup>1,2</sup>



### PSYCHOSOCIAL IMPACT

Significant impairment to quality of life through anxiety, concerns about transmission, depression, and social stigma<sup>7</sup>



### INCREASED RISK OF HIV ACQUISITION

30% of incident HIV infections acquired via sexual transmission attributable to HSV-2 infection<sup>8</sup>



# Recurrent Genital Herpes: Urgent Need for Innovative Therapies

## CURRENT STANDARD OF CARE

- Daily chronic suppressive therapy with viral polymerase inhibitors (e.g., acyclovir, valacyclovir)
- No new therapies approved since 1995<sup>1</sup>
- Wide treatment pattern variability seen in claims data<sup>4</sup>

### LIMITED EFFICACY



Only 1/3 with frequent outbreaks achieve recurrence prevention<sup>1</sup>

### HIGH TRANSMISSION



Less than 50% transmission reduction<sup>2</sup>

### HIGH PILL BURDEN



Lifelong daily treatment: Up to 1 gram, 1-3x/day<sup>1,3</sup>

### TREATMENT VARIABILITY



Many seeking care may not receive suppressive therapy consistently<sup>4</sup>

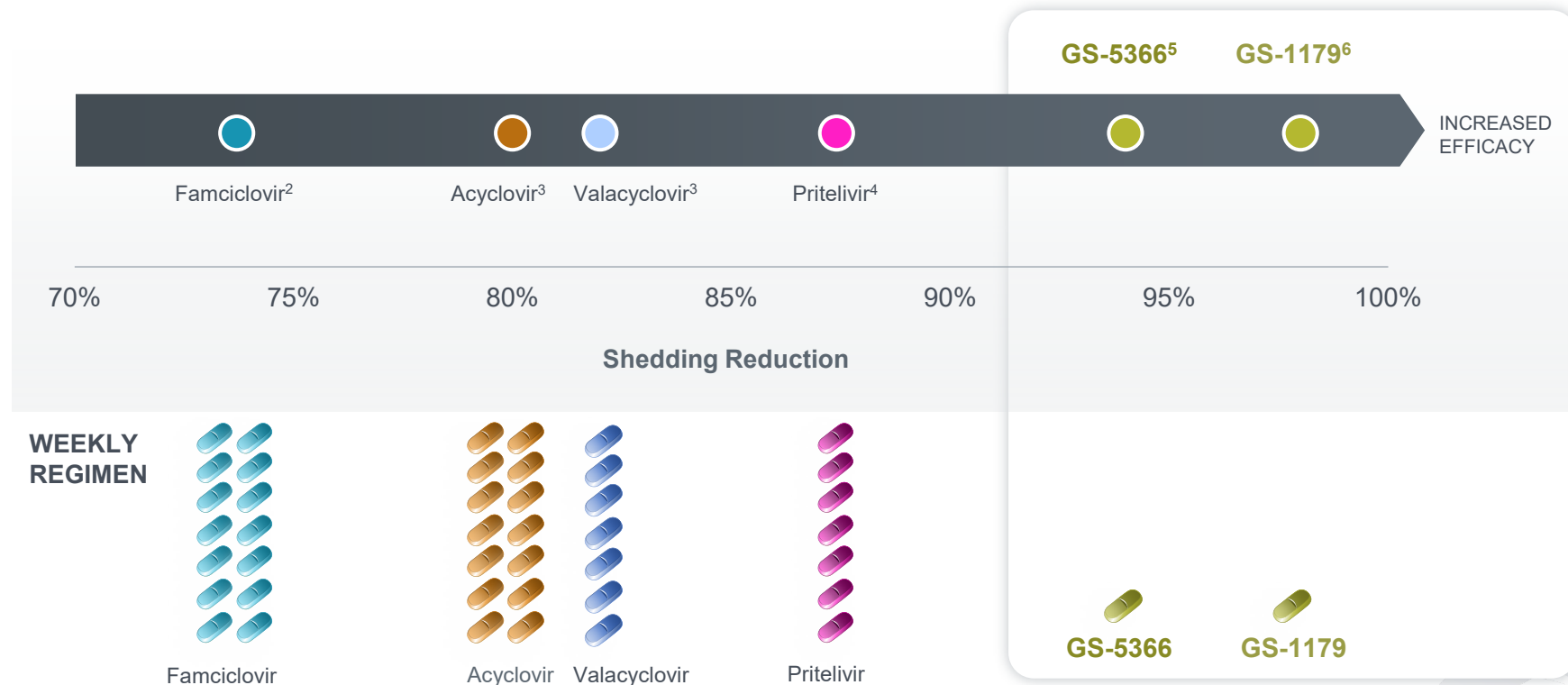
## GS-1179: INNOVATIVE POTENTIAL

- ✓ **Superior efficacy**  
Targeting superior efficacy to SOC; much greater potency demonstrated preclinically
- ✓ **Long-acting**  
Evaluating weekly oral dosing, with the goal of improving efficacy, adherence, and clinical outcomes
- ✓ **>\$2 billion**  
Market opportunity for recurrent genital herpes for profile of weekly dosing with superior efficacy to SOC

**Additional opportunities:** Combination strategies with HIV pre-exposure prophylaxis (PrEP), transmission prevention, patients currently treated episodically, oro-facial herpes, injectable formulations

**THERE IS AN  
URGENT NEED  
FOR INNOVATIVE  
THERAPIES**  
that offer  
improved efficacy  
and greater  
convenience

# GS-1179 & GS-5366 Phase 1b Efficacy: Comparisons to Historical Placebo-Controlled Phase 1b Studies<sup>1</sup>



Note: Length of evaluation of studies differs by compound. Famciclovir=14 days, Acyclovir/Valacyclovir=42 days, Pritelivir/GS-1179/GS-5366=29 days

1. Not head-to-head studies; 2. Leone P et al. Sexually Transmitted Diseases, 34 (11), 2007; 3. Gupta et al. JID 190, 2004; 4. Wald A et al. NEJM 370 (3) 2014; 5. Sasadeusz, et al. ESCMID, 2026. Interim Phase 1b data as of November 25, 2025; 6. Edwards, et al. ESCMID, 2026. Interim Phase 1b data as of November 25, 2025

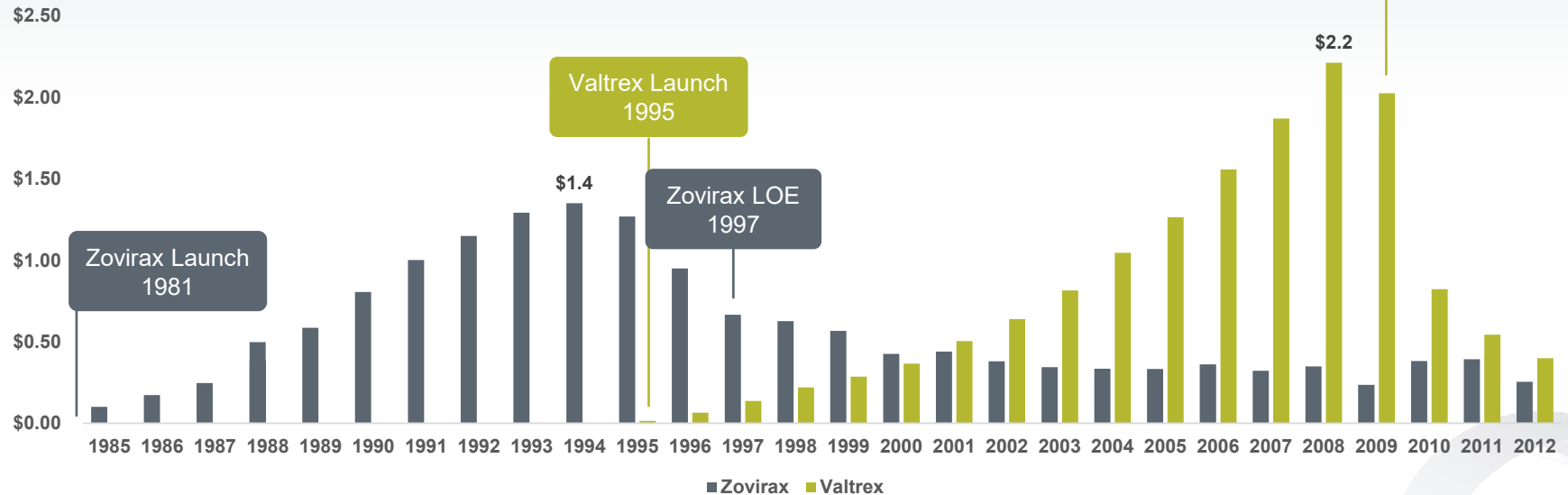
# Valtrex Launch in HSV Shows the Market Potential of a Novel, Longer-Acting Medicine

VALTREX TOOK CONSIDERABLE SHARE OVER TIME DESPITE GENERIC ACYCLOVIR AS CONVENIENCE AND ACCEPTANCE OF CHRONIC THERAPY DROVE ADOPTION

Sales in \$B

## Zovirax and Valtrex WW Sales

Branded Product Sales Only



# GS-1179-101 & GS-5366-101 Phase 1b Study Design

## Two Separate Studies Conducted Concurrently

- Double-blind, placebo-controlled sequential cohorts
- All participants seropositive for HSV-2 with recurrent genital herpes
- Each cohort enrolled 20 patients on active drug and 5 on placebo
- Treatment period: 29 days; follow-up period: 29 days for GS-1179, 98 days for GS-5366

### GS-1179-101

| COHORT | REGIMEN | WEEKLY DOSE |
|--------|---------|-------------|
| B1     | Weekly  | 50 mg       |
| B2     | Weekly  | 20 mg       |
| B3     | Weekly  | 10 mg       |

### GS-5366-101

| COHORT | REGIMEN | LOADING DOSE | WEEKLY DOSE |
|--------|---------|--------------|-------------|
| B1     | Weekly  | 150 mg       | 30 mg       |
| B2     | Weekly  | 350 mg       | 350 mg      |
| B3     | Monthly | 350 mg x5    | -           |

### KEY EFFICACY ASSESSMENTS

- Anogenital swabs collected twice daily over 4-week period<sup>1</sup>; e.g., viral shedding rate
- Daily diary of symptoms; e.g., days with lesions

### DATA IN CURRENT ANALYSIS

- GS-1179 – diary through D35; safety through D57 for first two cohorts
- GS-5366 – diary through D36; complete on-treatment safety data for all cohorts
- 100% shedding data available for both studies



# Executive Summary:

## GS-1179 and GS-5366 Showed Strong Phase 1b Antiviral Activity

### PHASE 1B TRIAL GOALS

- **80 to 85% reduction** in HSV-2 shedding vs placebo
- **Significant reduction** in high viral load swabs<sup>1</sup>
- **Directional reduction** in genital lesions
- **Clean safety profile**

### GS-1179 COHORT B1 (50 MG, QW)

- ✓ **98% reduction** in HSV-2 shedding ( $p < 0.01$ )
- ✓ **>99% reduction** in high viral load swabs
- ✓ **91% reduction** in virologically confirmed<sup>2</sup> genital lesions ( $p < 0.01$ )
- ✓ **No safety signals identified to date**
  - Chronic toxicology studies underway

### GS-5366 COHORT B2 (350 MG, QW)

- ✓ **94% reduction** in HSV-2 shedding ( $p < 0.01$ )
- ✓ **98% reduction** in high viral load ( $p < 0.05$ )
- ✓ **97% reduction** in virologically confirmed<sup>2</sup> genital lesions ( $p < 0.05$ )
- ✓ **No safety signals identified to date**
  - Chronic toxicology studies complete



# GS-1179 Phase 1b:

## Safety Summary – Adverse Events Cohorts B1 & B2

| PARAMETER                                                                            | GS-1179<br>20mg weekly/PBO<br>N=24 | GS-1179<br>50mg weekly/ PBO<br>N=25 |
|--------------------------------------------------------------------------------------|------------------------------------|-------------------------------------|
| <b>Subjects with any Treatment Emergent Adverse Events (TEAE) (max grade), N (%)</b> | 17 (71%)                           | 23 (92%)                            |
| Grade 1, N (%)                                                                       | 12 (50%)                           | 9 (36%)                             |
| Grade 2, N (%)                                                                       | 5 (21%)                            | 13 (52%)                            |
| Grade 3, N (%)                                                                       | 0                                  | 1 (4%) <sup>1</sup>                 |
| Grade 4, N (%)                                                                       | 0                                  | 0                                   |
| <b>TEAE Related to Study Drug, N (%)</b>                                             | 8 (33%)                            | 10 (40%)                            |
| <b>TEAE Leading to Study Drug Discontinuation, N (%)</b>                             | 0                                  | 0                                   |
| <b>Serious Adverse Event</b>                                                         | 0                                  | 0                                   |
| <b>Death</b>                                                                         | 0                                  | 0                                   |
| <b>Treatment Emergent Lab Abnormalities, N (%)</b>                                   | 9 (38%)                            | 8 (32%)                             |
| Grade 1, N (%)                                                                       | 8 (33%)                            | 6 (24%)                             |
| Grade 2, N (%)                                                                       | 2 (8%)                             | 3 (12%)                             |
| Grade 3, N (%)                                                                       | 0                                  | 0                                   |
| Grade 4, N (%)                                                                       | 0                                  | 0                                   |

Safety data includes patients through Day 57



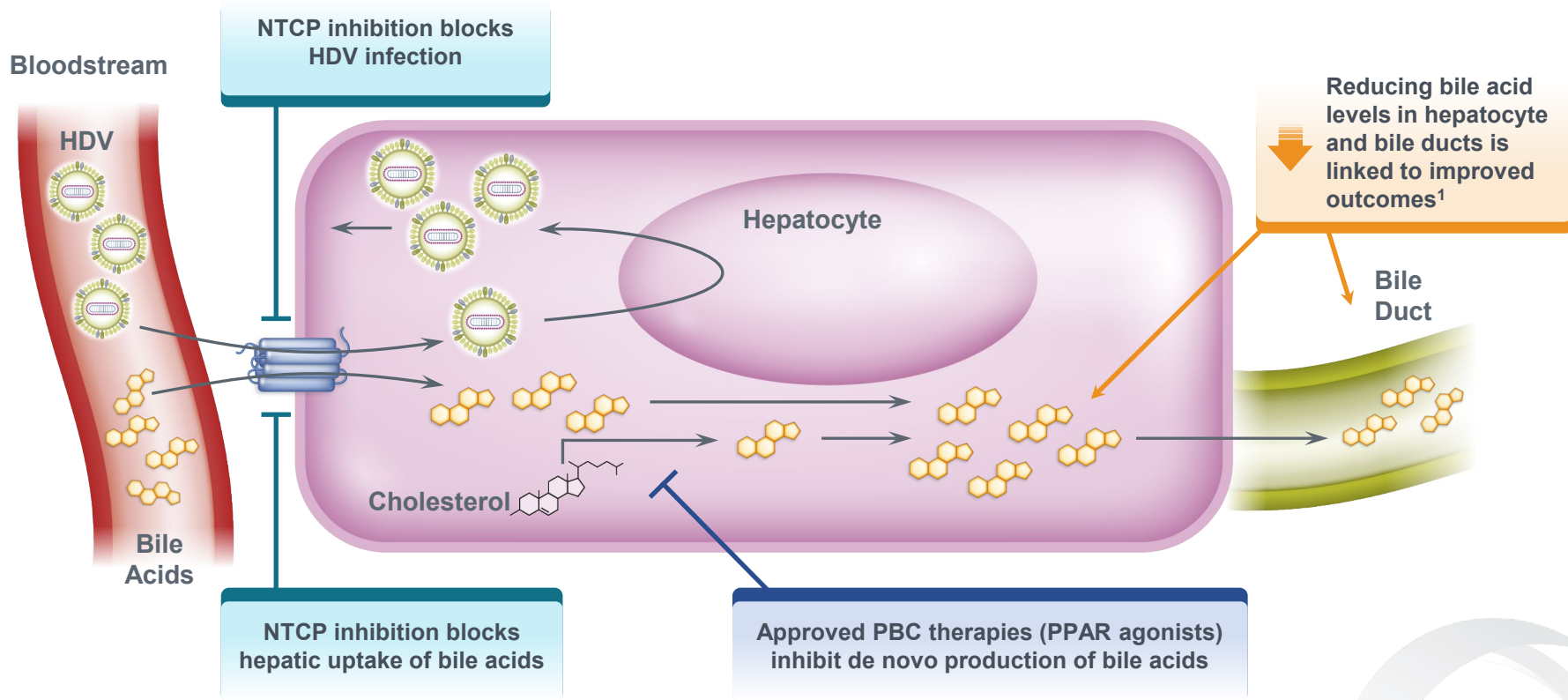
## ABI-6250

### Oral NTCP Inhibitor for Hepatitis D and Cholestatic Liver Diseases (CLD)

**Phase 2 initiation in HDV anticipated 4Q 2026**

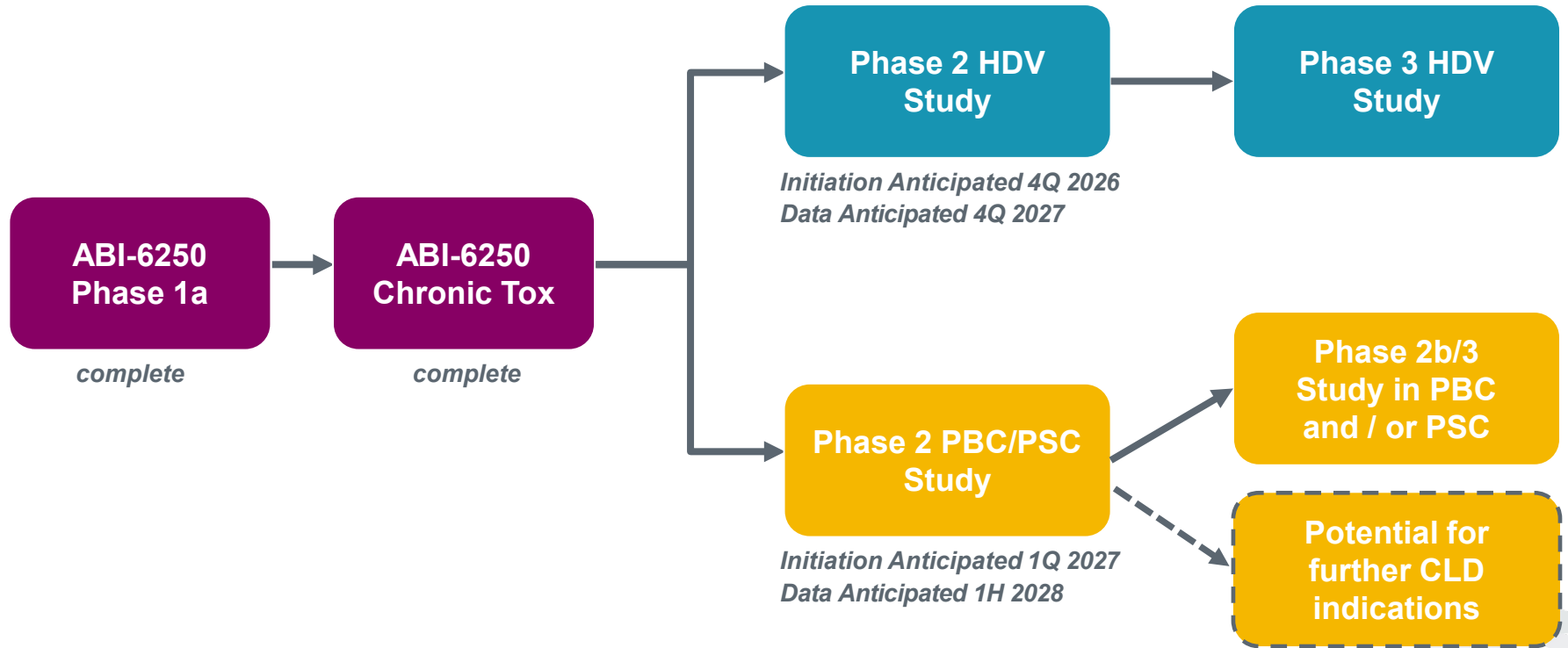
**Phase 2 initiation in PBC/PSC anticipated 1Q 2027**

# NTCP: A Shared Target Across HDV and CLD



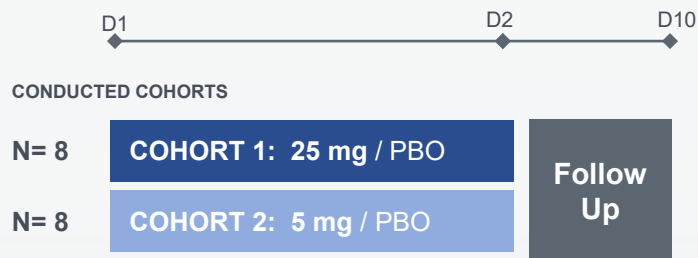
# ABI-6250 Overall Strategy:

## Parallel Development Strategy Across HDV and CLD

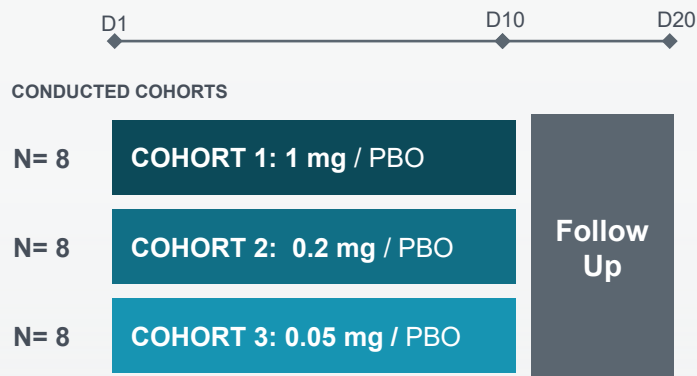


# ABI-6250-101 Phase 1a Study Design

## SINGLE-ASCENDING DOSE



## MULTIPLE-ASCENDING DOSE

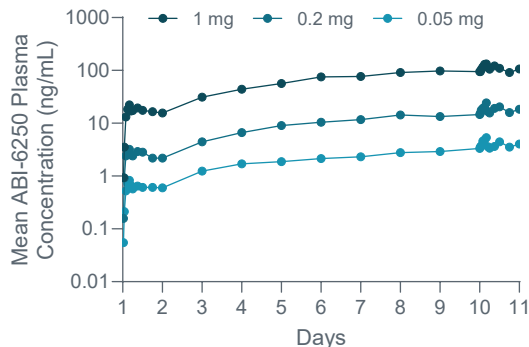


## KEY OUTCOMES

- Safety and pharmacokinetics
- Biomarker of target engagement (serum bile acid levels) with single and multiple doses

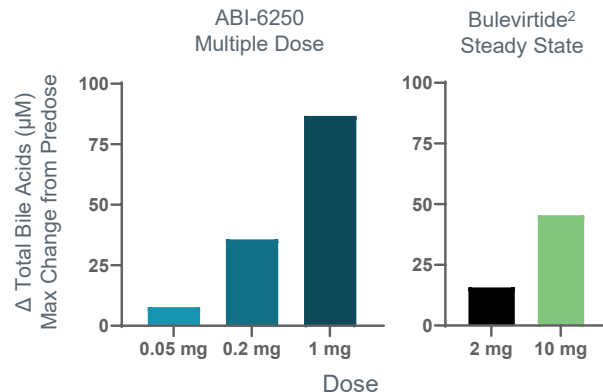
# ABI-6250 Shows PK/PD/Safety in Phase 1a to Support Phase 2

## Pharmacokinetics<sup>1</sup>



- Half-life estimate of 4 days
- ~7-fold accumulation with repeated oral daily dosing

## Pharmacodynamics<sup>1</sup>



- Dose dependent increase in serum bile acids
- Fold increase in bile acids matched or exceeded those by bulevirtide (BLV)

## Safety

- Grade 1-2 ALT elevations observed at similar frequency across all dosing levels, including in placebo subjects
  - Similar findings observed in early BLV studies
- No AEs of pruritis observed



1. Adapted from Gane et al. EASL 2026;

2. BLV is approved at a 2 mg once-daily dose in the EU, AU and Canada and 8.5mg in the US. The 10 mg dose has been evaluated in clinical studies but is not an approved dose in any jurisdiction

# ABI-6250 Phase 1a: Safety Data

|                                                                                | PBO SD<br>(N=4) | 5 mg SD<br>(N=6)       | 25 mg SD<br>(N=6) | PBO MD<br>(N=6) | 0.05 mg MD<br>(N=6) | 0.2 mg MD<br>(N=6) | 1 mg MD<br>(N=6)       |
|--------------------------------------------------------------------------------|-----------------|------------------------|-------------------|-----------------|---------------------|--------------------|------------------------|
| <b>Subjects with any TEAE, N (%)</b>                                           | 0               | 4 (66.7%)              | 2 (33.3%)         | 4 (66.7%)       | 6 (100%)            | 4 (66.7%)          | 4 (66.7%)              |
| Grade 1, N (%)                                                                 | 0               | 4 (66.7%)              | 2 (33.3%)         | 4 (66.7%)       | 6 (100%)            | 3 (50.0%)          | 4 (66.7%)              |
| Grade 2, N (%)                                                                 | 0               | 0                      | 0                 | 0               | 0                   | 2 (33.3%)          | 0                      |
| Grade 3, N (%)                                                                 | 0               | 0                      | 0                 | 0               | 0                   | 0                  | 0                      |
| Grade 4, N (%)                                                                 | 0               | 0                      | 0                 | 0               | 0                   | 0                  | 0                      |
| <b>TEAE related to study drug, N (%)</b>                                       | 0               | 0                      | 0                 | 0               | 1 (16.7%)           | 0                  | 2 (33.3%)              |
| <b>Serious TEAE, N (%)</b>                                                     | 0               | 0                      | 0                 | 0               | 0                   | 0                  | 0                      |
| <b>TEAE leading to study drug discontinuation, N(%)</b>                        | 0               | 0                      | 0                 | 0               | 0                   | 0                  | 1 (16.7%) <sup>1</sup> |
| <b>Death</b>                                                                   | 0               | 0                      | 0                 | 0               | 0                   | 0                  | 0                      |
| <b>Number (%) of subjects with any graded TE lab abnormalities<sup>3</sup></b> | 2 (50%)         | 4 (66.7%)              | 3 (50.0%)         | 5 (83.3%)       | 3 (50%)             | 5 (83.3%)          | 3 (50.0%)              |
| Grade 1, N (%)                                                                 | 2 (50%)         | 4 (66.7%)              | 2 (33.3%)         | 4 (66.7%)       | 2 (33.3%)           | 5 (83.3%)          | 3 (50.0%)              |
| Grade 2, N (%)                                                                 | 0               | 0                      | 1 (16.7%)         | 3 (50.0%)       | 1 (16.7%)           | 1 (16.7%)          | 0                      |
| Grade 3, N (%)                                                                 | 0               | 0                      | 0                 | 0               | 0                   | 0                  | 0                      |
| Grade 4, N (%)                                                                 | 0               | 1 (16.7%) <sup>2</sup> | 0                 | 0               | 0                   | 0                  | 0                      |

**Chronic toxicology studies complete; Preparation for Phase 2 studies underway**



# ABI-6250: Oral Hepatitis D Virus Entry Inhibitor

**Phase 2 initiation anticipated by end of 2026**

# Chronic HDV is a Serious Life-Threatening Disease and Major Unmet Need with Limited Treatment Options



**12 million**

**PEOPLE ESTIMATED TO BE CHRONICALLY INFECTED WITH HDV GLOBALLY<sup>1</sup>**

Considered the most severe form of chronic viral hepatitis, with more rapid progression to liver cancer and liver-related death



**Very limited treatment options**

**BULEVIRTIDE, LARGE MOLECULE ENTRY INHIBITOR, ONLY APPROVED DRUG (EU, AU, and Canada; US approved 2026)**

Shown to be safe and highly effective in long-term clinical trials, but requires daily injection and cold storage



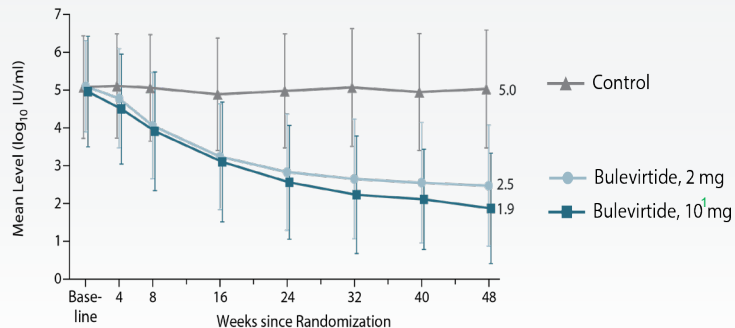
**ABI-6250, an opportunity to simplify treatment**

**SMALL MOLECULE TARGETING SAME MECHANISM AS BULEVIRTIDE**

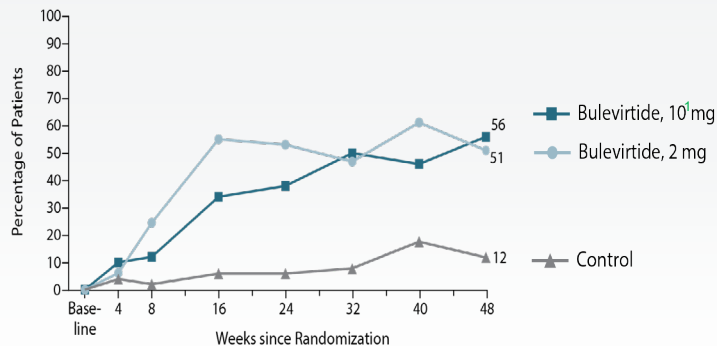
An oral treatment is expected to further enhance treatment uptake and diagnosis rates

# Inhibition of HDV Entry by Blocking NTCP is Clinically Validated to Lower Viral Load and Normalize ALT

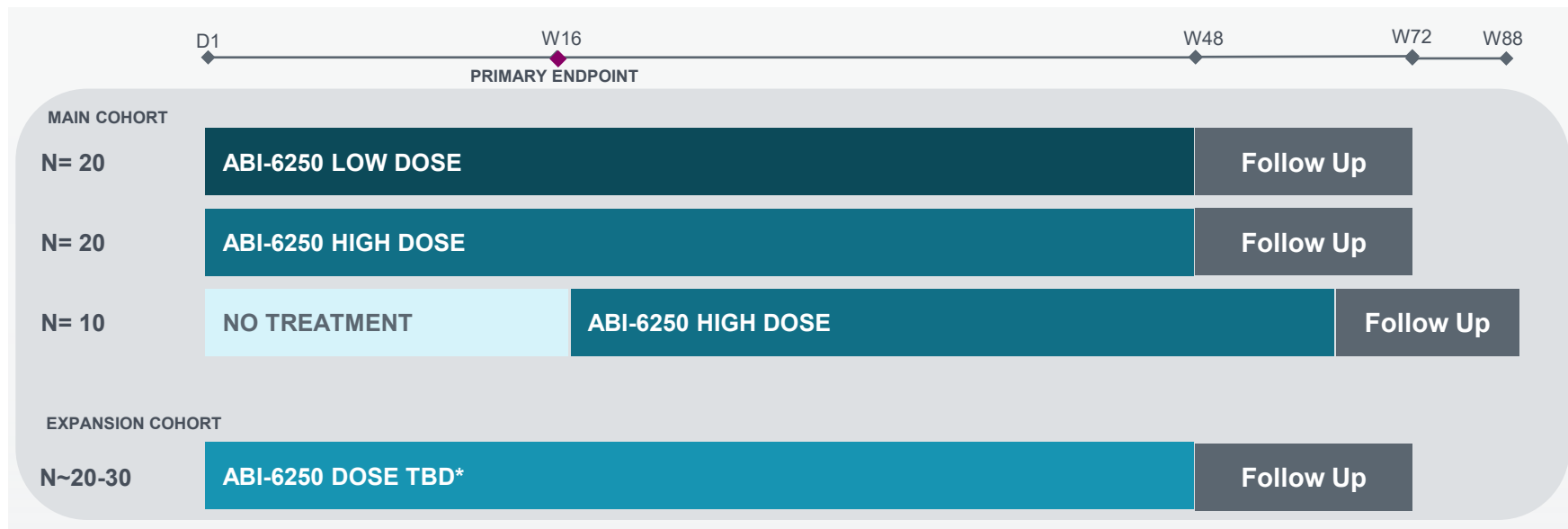
## Viral Load Reductions



## ALT Normalization



# ABI-6250 HDV Phase 2 Design



## TRIAL CHARACTERISTICS

- Open-label
- Expansion cohort for additional safety data to support Phase 3
- Key efficacy endpoints of HDV RNA change from baseline and ALT change from baseline

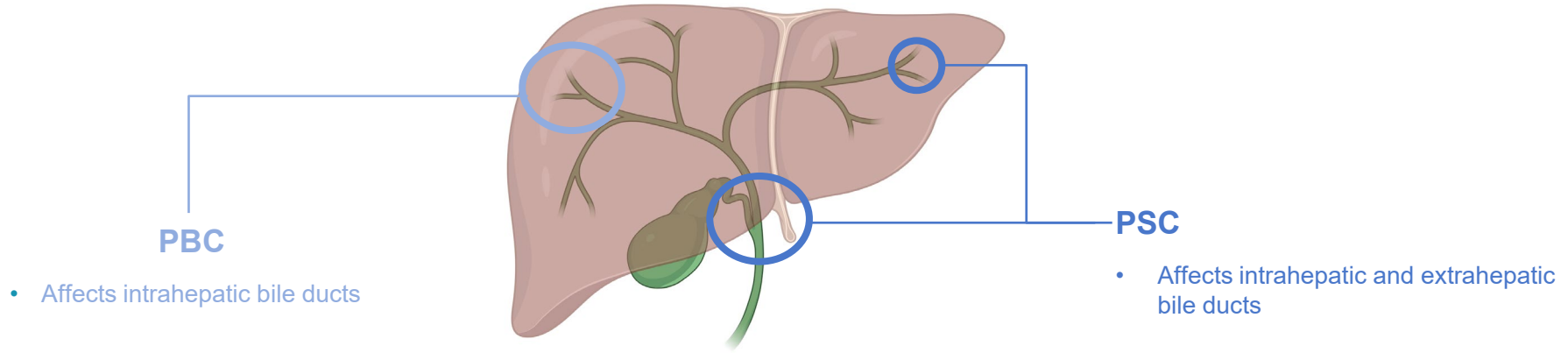


\*Dose selection for expansion cohort to occur following primary endpoint analysis of the main cohort at Week 16

# ABI-6250: Expanding Beyond HDV into Cholestatic Liver Diseases (CLD)

Phase 2 initiation anticipated Q1 2027

# PBC and PSC Share a Common Pathology of Bile Acid Accumulation and Progressive Liver Injury

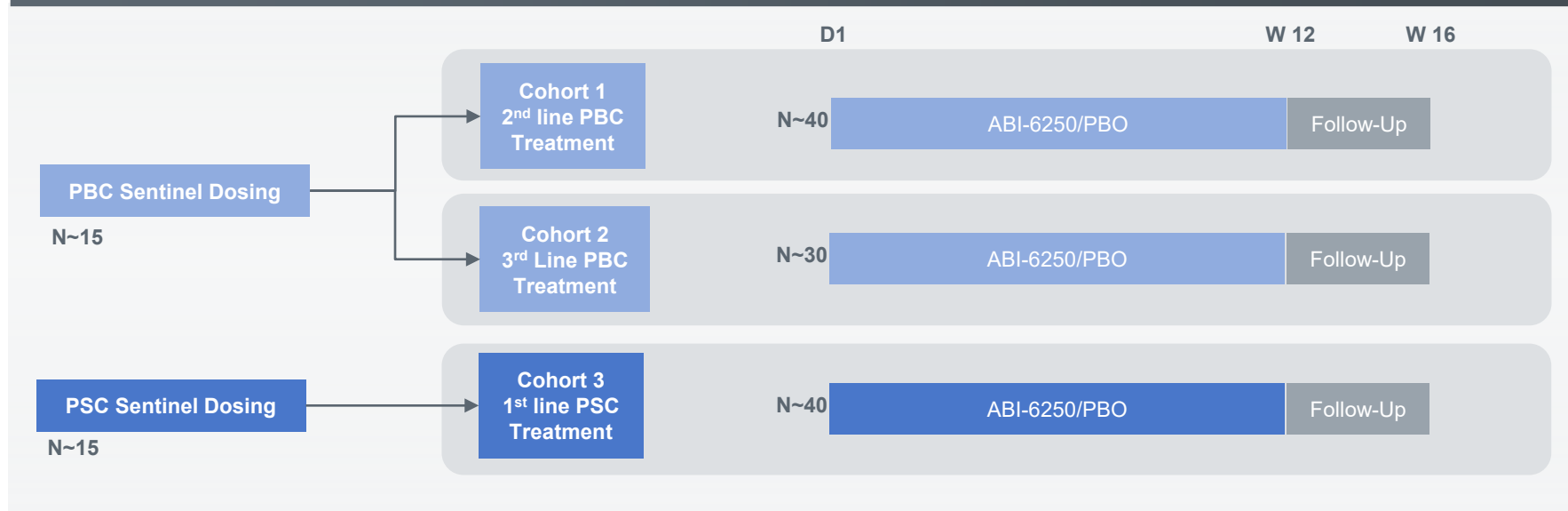


|                       | PBC                                                                              | PSC                                                                                 |
|-----------------------|----------------------------------------------------------------------------------|-------------------------------------------------------------------------------------|
| Associated Conditions | Sjogren's Syndrome                                                               | Inflammatory Bowel Disease                                                          |
| Complications         | Cirrhosis, hepatocellular carcinoma                                              | Cirrhosis, Cholangiocarcinoma, Cholangitis                                          |
| Treatments            | 1 <sup>st</sup> Line: Ursodeoxycholic Acid<br>2 <sup>nd</sup> Line: PPAR agonist | No medical treatments available<br>Disease progression may require liver transplant |



# ABI-6250 Cholestatic Liver Disease Preliminary Phase 2 Design

## Basket Study Design Exploring Efficacy and Safety in PBC and PSC in a Single Protocol



### TRIAL CHARACTERISTICS

- Double-blind, placebo controlled
- Sentinel dosing will help select dose(s) for PBC/PSC cohorts
- Key endpoints of biochemical, pruritis and health-related quality of life improvements will be evaluated



# ABI-6250 Represents a Multi-Billion-Dollar Opportunity Across PBC and PSC

## PBC

**~\$2B+**

**U.S. estimated peak-year revenue**

**POPULATION** ~150,000+ diagnosed adults in the U.S.

**UNMET NEED** ~40% of patients fail 1st line therapy

## PSC

**~\$1B+**

**U.S. estimated peak-year revenue**

**POPULATION** ~30,000+ diagnosed adults in the U.S.

**UNMET NEED** No approved therapy  
~35% 10-year overall mortality rate in U.S.

Expansion into CLD adds meaningful upside to an established core asset

- Multiple optional pathways: PBC, PSC, both or more
- HDV execution remains unchanged
- Opportunity to generate additional long-term value from a single molecule

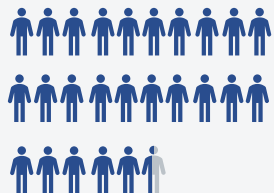


# ABI-4334: Next-Generation CAM for Hepatitis B

**Phase 1b topline data reported June 2025**

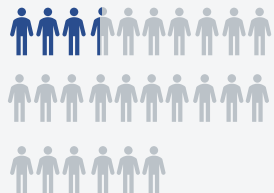
**Positioned for next-phase development upon securing partner**

# HBV is a Major Unmet Medical Need Globally



HBV PREVALENCE:

**240M<sup>1</sup>**



DIAGNOSED:

**65M<sup>1</sup>**



TREATED:

**10M<sup>1</sup>**

**Up to 1,100,000 people died in 2024<sup>1</sup>**

FROM HBV-RELATED CAUSES

**Treatments are life-long**

INHIBIT VIRUS BUT CURE RATES VERY LOW

**Opportunity to improve patient outcomes**

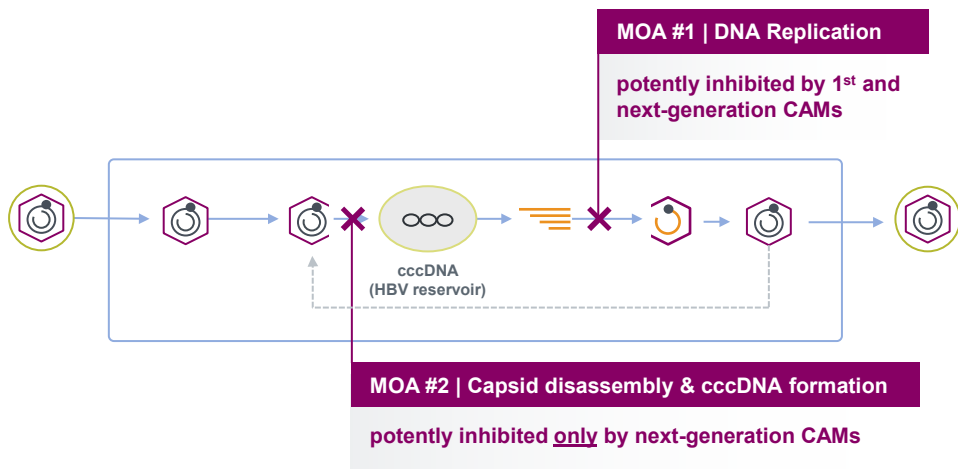
AND INCREASE NUMBER DIAGNOSED AND TREATED,  
with development of finite and curative therapies

**No new MOAs approved for HBV in  
>25 years**

# ABI-4334 is a Next-Generation Capsid Assembly Modulator Designed to Target Both MOAs for the Class

## CAPSID ASSEMBLY MODULATORS (CAMs)

Direct-acting antivirals with two distinct mechanisms of action



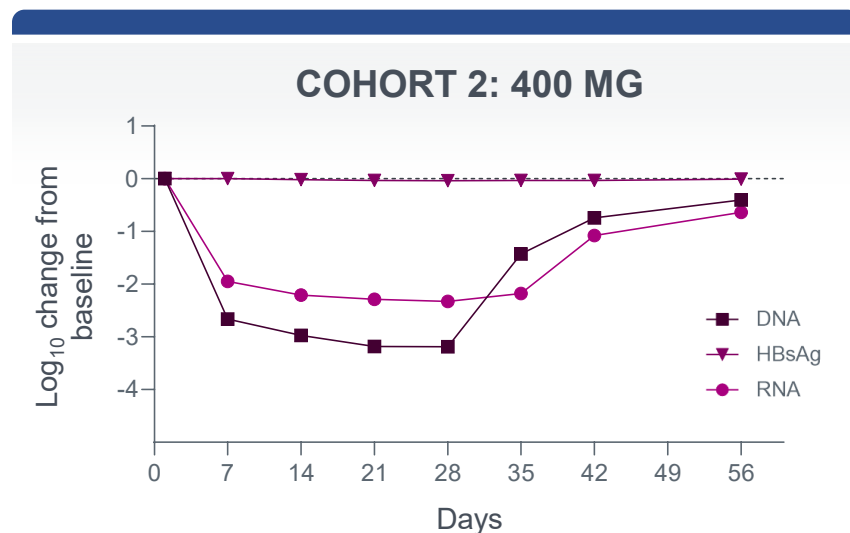
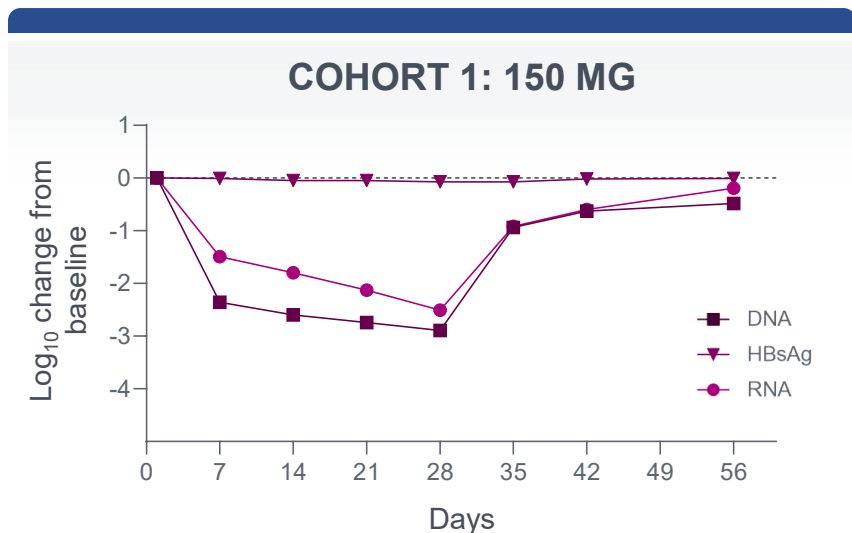
## ABI-4334 PHASE 1B PK

Supportive of the ability to potentially achieve double-digit multiples over  $\text{paEC}_{50}$

| 4334 Phase 1b Cohorts <sup>1</sup>                        |                    |                    |
|-----------------------------------------------------------|--------------------|--------------------|
|                                                           | 150mg <sup>2</sup> | 400mg <sup>2</sup> |
| Fold of $C_{\min}/\text{paEC}_{50}$<br>MOA #1 (antiviral) | 62x                | 269x               |
| Fold of $C_{\min}/\text{paEC}_{50}$<br>MOA #2 (cccDNA)    | 12x                | 52x                |



# Positive Phase 1b Data: ABI-4334 Demonstrates Potent Antiviral Activity



HBeAg-positive or -negative cHBV infected participants not on NrtI, randomized 8:2 active:placebo per cohort

- 2.9 and 3.2 log<sub>10</sub> IU/mL mean decline in HBV DNA over 28 days observed in 150 mg and 400 mg cohorts, respectively, supporting ability of lower dose to potentially saturate antiviral mechanism of action
- Limited changes in HBsAg observed as expected given 28-day treatment period



# Topline Safety Data Supports Flexibility in Dose Range

|                                                                    | Placebo<br>(N=4) | ABI-4334 150mg<br>(N=8) | ABI-4334 400mg<br>(N=8) |
|--------------------------------------------------------------------|------------------|-------------------------|-------------------------|
| <b>Subjects with any TEAE (max toxicity), N (%)</b>                | 2 (50%)          | 5 (62.5%)               | 6 (75%)                 |
| Grade 1, N (%)                                                     | 0                | 2 (25%)                 | 1 (12.5%)               |
| Grade 2, N (%)                                                     | 2 (50%)          | 2 (25%)                 | 5 (62.5%)               |
| Grade 3, N (%)                                                     | 0                | 1 (12.5%)*              | 0                       |
| Grade 4, N (%)                                                     | 0                | 0                       | 0                       |
| <b>TEAE related to study drug, N (%)</b>                           | 1 (25%)          | 5 (62.5%)               | 1 (12.5%)               |
| <b>Serious TEAE, N (%)</b>                                         | 0                | 0                       | 0                       |
| <b>TEAE leading to study drug discontinuation, N (%)</b>           | 0                | 0                       | 0                       |
| <b>Death</b>                                                       | 0                | 0                       | 0                       |
| <b>Number (%) of subjects with any graded TE lab abnormalities</b> | 3 (75%)          | 6 (75%)                 | 8 (100%)                |
| Grade 1, N (%)                                                     | 3 (75%)          | 5 (62.5%)               | 8 (100%)                |
| Grade 2, N (%)                                                     | 3 (75%)          | 3 (37.5%)               | 4 (50%)                 |
| Grade 3, N (%)                                                     | 1 (25%)**        | 1 (12.5%)*              | 0                       |
| Grade 4, N (%)                                                     | 0                | 0                       | 0                       |

\*ALT elevation; resolved by Day 28 with continued dosing of ABI-4334

\*\* Total Bilirubin Increased

# ABI-7272:

## Oral Broad-Spectrum Non-Nucleoside Polymerase Inhibitor (NNPI) for Transplant-Associated Herpesviruses

IND/CTA-enabling studies

# Multiple Herpesviruses Can Cause Significant Morbidity and Mortality in Immunocompromised Transplant Recipients

**95,000** PATIENTS AFFECTED<sup>1</sup>

## AMONG TRANSPLANT PATIENTS:

 ~60% are CMV positive

 ~60% are HSV positive

 ~80% are VZV positive

 ~45% are EBV positive

## Lifelong latent infections

FREQUENTLY REACTIVATE DURING IMMUNOSUPPRESSION

## Uncontrolled viral replication

AND SEVERE DISEASE DURING REACTIVATION

## Risk of graft loss and death

## SOC antivirals are:

- PARTIALLY EFFICACIOUS
- NOT BROAD-SPECTRUM
- HAVE TOLERABILITY AND DRUG INTERACTION LIMITATIONS

**An oral broad-spectrum herpesvirus antiviral could improve efficacy and greatly simplify treatment**

Patel and Paya. Clin. Microbiol. Rev. 1997; Breuer, et al. Mol. Diagn. Ther. 2012; Clark, et al. Semin. Respir. Crit. Care Med. 2013; Haider and Singh. Curr. Opin. Infect. Dis. 2019; Beyar-Katz et al. Clin. Microbiol. Infect. 2020; Kwon et al. Transp. Infect. Dis. 2021; Wutzler et al. Vaccine 2001; Bauer et al. BMC Infect. Dis. 2010; Reynolds et al. Public Health Rep. 2010; Lanzieri et al. Int. J. Gynaecol. Obstet. 2016; Lachmann et al. PLoS One 2018; Patton et al. Clin. Infect. Dis. 2018; Ayoub et al. BMC Med. 2019; Zuhair et al. Rev. Med. Virol. 2019; Zhang et al. Virol. J. 2022; Marty et al. NEJM 2017; Limaye et al. JAMA 2023; Witzke et al. Transp. 2012; Witzke et al. Transp. 2018; Höcker et al. Clin. Infect. Dis. 2012; Cho et al. Am. J. Clin. Pathol. (2014); Holman et al. Clin. Transplant (2012); Bamouli et al. Am. J. Transplant. (2013); Verghese et al. Transplant. (2015).

1. EBMT, OPTN, UNOS, and IRODAT (estimate for transplanted-associated herpesvirus reflects US and EU only)

# ABI-7272 is Designed to Provide Significant Innovation Over Current Standard of Care

## CONSERVED VIRAL POLYMERASES PROVIDE OPPORTUNITY FOR BROAD-SPECTRUM HERPESVIRUS INHIBITION

### OPPORTUNITY TO ADVANCE CURRENT STANDARD OF CARE

- Improve efficacy and broaden spectrum of antiviral activity
- Simplify treatment (1 agent to target 5 viruses)
- Improve tolerability and reduce drug-drug interactions



HSV-1



HSV-2



CMV



VZV



EBV

**IND/CTA  
ENABLING**  
studies ongoing





Nasdaq: ASMB