# Results from a phase 1 study to assess the safety, tolerability, pharmacokinetics, and pharmacodynamics of tebapivat (AG-946) in patients with sickle cell disease

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## BACKGROUND

- In sickle cell disease (SCD), pyruvate kinase activation increases adenosine triphosphate (ATP), leading to improved membrane integrity and survival of red blood cells (RBCs), and decreases 2,3-diphosphoglycerate (DPG), preventing the polymerization of sickle hemoglobin (HbS) in its deoxygenated state<sup>1</sup>
- Mitapivat, an allosteric activator of the RBC-specific (PKR) and M2 (PKM2) isoforms of pyruvate kinase, demonstrated clinically meaningful improvements in hemoglobin (Hb) response and improvements in markers of hemolysis and erythropoiesis in phase 2 trials in SCD<sup>2,3</sup>
- Mitapivat is currently being evaluated in a phase 3 trial in patients with SCD<sup>4</sup>
- Tebapivat (formerly AG-946) is an oral, once daily (QD), potent, allosteric activator of PKR and PKM2 (**Figure 1**)<sup>5</sup>; results from the randomized, double-blind, placebocontrolled single ascending dose (SAD) and multiple ascending dose (MAD) parts of a phase 1 study of tebapivat in healthy volunteers (HVs; NCT04536792) have been previously reported<sup>6</sup>



ADP, adenosine diphosphate; ATP, adenosine triphosphate; DPG, diphosphoglycerate; FBP, fructose bisphosphate; Hb, hemoglobin; HbS, sickle hemoglobin; PEP, phosphoenolpyruvate; PG, phosphoglycerate; RBC, red blood cell

## **OBJECTIVE**

• To understand the safety, tolerability, and pharmacokinetics/pharmacodynamics (PK/PD) of tebapivat in the non-randomized, open-label, third part of a phase 1 study in adult patients with SCD

## METHODS

## **Study design**

- Adult patients (aged 18–70 years) with sickle cell anemia (homozygous for HbS [HbSS] or HbS/ß<sup>o</sup>-thalassemia) and adequate organ function received 2 mg or 5 mg tebapivat QD for 28 days, with a further 28-day observational safety follow-up<sup>14</sup> (**Figure 2**)
- Further details of the study eligibility criteria can be found via the QR code



## **Study endpoints**

- The primary endpoints were: - Relationships between tebapivat dose, concentration, and safety endpoints - Relationships between tebapivat dose, concentration,
- Type, severity, and relationship of adverse events (AEs) and serious AEs (SAEs) – Plasma pharmacokinetic parameters after both single
- Change over time in the whole blood concentrations of 2,3-DPG and ATP
- Change from baseline in Hb
- Change from baseline in markers of hemolysis (including total bilirubin and lactate dehydrogenase [LDH] levels) and erythropoiesis (including reticulocyte percentage
- and erythropoietin [EPO])

## RESULTS

- Sixteen adult patients with SCD received ≥1 dose of either 2 mg QD (N=8) or 5 mg QD (N=8) oral tebapivat
- Fourteen patients (87.5%) completed the 28-day dosing period
- analysis

## Safety

- the Investigator

## Figure 2. Study design<sup>14</sup> Part 3 (SCD [QD × 28 davs]) 100% of ma AD level fro HVs Dose with maximum pharmacologic activity (↑ATP, ↓2,3-DPG) Focus of this poste

ATP, adenosine triphosphate; DPG, diphosphoglycerate; HV, healthy volunteer; MAD, multiple ascending dose; PAD, pharmacologically active dose; PK/PD, pharmacokinetic/pharmacodynamic; QD, once daily; SAD, single ascending dose; SCD, sickle cell disease

- and pharmacodynamic endpoints
- Secondary endpoints included:
  - and multiple oral dose administration of tebapivat

- One patient in the 2 mg QD cohort discontinued tebapivat due to an AE (sickle cell anemia with crisis), and 1 patient in the 5 mg QD cohort discontinued tebapivat due to increased Hb (but completed the study) – All 16 patients were included in the intention-to-treat

• Two (25.0%) patients in the 2 mg QD cohort reported an SAE of sickle cell anemia with crisis; one patient reported two events, one during the treatment period and one during the safety follow-up, and the other patient experienced one event during the safety follow-up (**Table 2**)

• No TEAEs of sickle cell anemia with crisis were reported during the 5 mg QD treatment period; 1 (12.5%) patient in the 5 mg QD cohort reported an SAE of sickle cell anemia with crisis during the safety follow-up, which was the only AE/SAE of sickle cell anemia with crisis considered treatment-related by

Table 1. Baseline demographics and disease characteristics			
Demographics and disease characteristics	Tebapivat 2 mg QD (N=8)	Tebapivat 5 mg QD (N=8)	
Age, median (range), years	28.0 (19.0–48.0)	37.5 (25.0–51.0)	
Male, n (%)	4 (50.0)	4 (50.0)	
Race, n (%)			
Black or African American	7 (87.5)	6 (75.0)	
White	0 (0.0)	1 (12.5)	
Multiracial	0 (0.0)	1 (12.5)	
Not reported	1 (12.5)	0 (0.0)	
Hb concentration, mean (SD), g/dL	7.8 (1.0)	8.1 (1.1)	
VOC in the prior 12 months, <sup>a</sup> n (%)	4 (50.0)	3 (37.5)	
Recieved prior SCD-related therapies, <sup>b</sup> n, (%)	4 (50.0)	6 (75.0)	
$\frac{1}{2}$			

Prior disease-modifying SCD-related therapies included hydroxyurea, crizanlizumab, L-glutamine, and voxelotor Hb, hemoglobin: QD, once daily: SCD, sickle cell disease: SD, standard deviation: VOC, vaso-occlusive crisis

Patients, n (%)	Tebapivat 2 mg QD (N=8)	Tebapivat 5 mg QD (N=8)
Any TEAEs	8 (100.0)	8 (100.0)
Grade ≥3 TEAEs	3 (37.5)	3 (37.5)
Freatment-related TEAEs	1 (12.5)	2 (25.0)
Grade ≥3 treatment-related TEAEs	0 (0.0)	1 (12.5)
Serious TEAEs	3 (37.5)	1 (12.5)
Serious treatment-related TEAEs	0 (0.0)	1 (12.5)
<b>FEAEs leading to discontinuation of study drug</b>	0(0.0)	0 (0.0)
<b>FEAEs leading to dose reduction</b>	0 (0.0)	0 (0.0)
FEAEs leading to interruption of study drug	2 (25.0)	0 (0.0)
FEAEs leading to death	0 (0.0)	0 (0.0)
Most frequently reported (≥10%) TEAEs		
Sickle cell anemia with crisis		
Any grade	4 (50.0)	3 (37.5)
Grade ≥3	2 (25.0)	1 (12.5)
Jpper respiratory tract infection		
Any grade	1 (12.5)	2 (25.0)
Grade ≥3	0 (0.0)	1 (12.5)

AE, adverse event; QD, once daily; TEAE, treatment-emergent adverse ever

## • All pain crises occurred in the setting of known triggers

- Hb and markers of hemolysis and erythropoiesis
- the 2 mg QD cohort and 1.9 g/dL (0.7) in the 5 mg QD cohort (**Figure 3**)
- Overall decreases in markers of hemolysis (total bilirubin and LDH) and erythropoiesis (reticulocyte both cohorts (**Figure 4A–C**)



Hb, hemoglobin; QD, once daily; SD, standard deviation

• At the end of the 28-day treatment period, the mean (SD) change from baseline for Hb was 1.2 g/dL (0.4) in

percentage) from baseline were observed at Day 28 in



## **Pharmacokinetics**

- Overall, tebapivat exposure increased with a higher dose (2 mg QD vs 5 mg QD)
- Tebapivat exposures in patients with SCD on both Day 1 (2 mg QD: 58 h·ng/mL; 5 mg QD: 197 h·ng/mL) and Day 15 (2 mg QD: 157 h·ng/mL; 5 mg QD: 447 h·ng/mL) were comparable to exposures in HVs<sup>15</sup>

## **Pharmacodynamics**

- Dose-dependent pharmacodynamic effects on 2,3-DPG and ATP levels were demonstrated with tebapivat, with higher doses resulting in greater changes from baseline
- 2,3-DPG and ATP concentrations reached steady state after 2 weeks of QD dosing
- At Day 28 (pre-dose [sample collected ≤60 minutes before the administration of tebapivat]), mean (SD) percent reduction in 2,3-DPG from baseline was 20.9% (7.1) and 29.4% (12.7) for the 2 mg and 5 mg cohorts, respectively (**Figure 5A**)
- At Day 28 (pre-dose), mean (SD) percent increase in ATP from baseline was 46.3% (29.1) and 67.8% (30.9) for the 2 mg and 5 mg cohorts, respectively (Figure 5B)

## • A sustained pharmacodynamic effect was observed up to 4 weeks after 28 days of QD dosing



aseline was defined as the pre-dose concentration on study Day 1. "Pre-dose" refers to samples collected  $\leq 60$ inutes before the administration of tebapivat. "Anytime" refers to samples collected at any point during that day ATP, adenosine triphosphate; DPG, diphosphoglycerate; QD, once daily; SD, standard deviation

## CONCLUSIONS

- Tebapivat was well tolerated in patients with SCD receiving either 2 mg or 5 mg QD for 28 days
- Increases in Hb and trends towards improvements in hemolytic and erythropoietic markers were observed, and there was a sustained effect after tebapivat was stopped
- ATP levels were increased and 2,3-DPG levels decreased during the study, consistent with the proposed mechanism of action of tebapivat
- A sustained pharmacodynamic effect was observed up to 4 weeks after the last dose
- Tebapivat will be further evaluated in different clinical studies

Tebapivat is a potent pyruvate kinase activator with the potential to provide benefit in SCD

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