

J.P. Morgan Healthcare Conference Agios Pharmaceuticals

Brian Goff, Chief Executive Officer January 10, 2024



Forward-looking statements

This presentation and various remarks we make during this presentation contain forward-looking statements within the meaning of The Private Securities Litigation Reform Act of 1995. Such forward-looking statements include those regarding the potential benefits of PYRUKYND® (mitapivat), AG-946, TMPRSS6 siRNA and its PAH stabilizer: Agios' plans. strategies and expectations for its preclinical, clinical and commercial advancement of its drug development, including PYRUKYND®, AG-946 and its PAH stabilizer; Agios' strategic vision and goals, including its key milestones for 2024 and potential catalysts through 2026; and the potential benefits of Agios' strategic plans and focus. The words "anticipate," "expect," "goal," "hope," "milestone," "opportunity," "plan," "potential," "possible," "strategy," "will," "vision," and similar expressions are intended to identify forward-looking statements, although not all forward-looking statements contain these identifying words. Such statements are subject to numerous important factors, risks and uncertainties that may cause actual events or results to differ materially from Agios' current expectations and beliefs. For example, there can be no guarantee that any product candidate Agios is developing will successfully commence or complete necessary preclinical and clinical development phases, or that development of any of Agios' product candidates will successfully continue. There can be no guarantee that any positive developments in Agios' business will result in stock price appreciation. Management's expectations and, therefore, any forward-looking statements in this presentation and various remarks we make during this presentation could also be affected by risks and uncertainties relating to a number of other important factors, including, without limitation: risks and uncertainties related to the impact of pandemics or other public health emergencies to Agios' business, operations, strategy, goals and anticipated milestones, including its ongoing and planned research activities, ability to conduct ongoing and planned clinical trials, clinical supply of current or future drug candidates, commercial supply of current or future approved products, and launching, marketing and selling current or future approved products; Agios' results of clinical trials and preclinical studies, including subsequent analysis of existing data and new data received from ongoing and future studies; the content and timing of decisions made by the U.S. FDA, the EMA or other regulatory authorities, investigational review boards at clinical trial sites and publication review bodies; Agios' ability to obtain and maintain requisite regulatory approvals and to enroll patients in its planned clinical trials; unplanned cash requirements and expenditures; Agios' ability to obtain, maintain and enforce patent and other intellectual property protection for any product candidates it is developing; Agios' ability to establish and maintain key collaborations; uncertainty regarding any milestone or royalty payments related to the sale of Agios' oncology business or its in-licensing of TMPRSS6 siRNA, and the uncertainty of the timing of any such payments; uncertainty of the results and effectiveness of the use of proceeds from the transaction with Servier; competitive factors; and general economic and market conditions. These and other risks are described in greater detail under the caption "Risk Factors" included in Agios' public filings with the Securities and Exchange Commission. Any forward-looking statements contained in this presentation and various remarks we make during this presentation speak only as of the date hereof, and Agios expressly disclaims any obligation to update any forward-looking statements, whether as a result of new information, future events or otherwise, except as required by law.



Our Mission

Our Vision

Our Values



Develop and deliver transformative medicines that elevate and extend the lives of patients





To become a leading rare disease company providing first-in-class and/or best-in-class new therapies for diseases with high unmet need





Blaze New Trails

Fueled by Connections to Transform Rare Diseases



Well-positioned with multiple near-term catalysts to enter multi-billion dollar markets and deliver significant value

PKa franchise with multi-billion dollar potential

Large opportunities
with substantial value potential for two
additional first and
best-in-class
indications for
PYRUKYND® by 2026

Differentiated mechanism of action

Clearly differentiated
PK activation franchise
targeting red blood cell
health beyond
hemoglobin increase

Increasing probability of success

Proven track record supported by compelling and consistent data to date

Growing pipeline

Diversified pipeline
addressing the
underlying
pathophysiology of
rare diseases
with high unmet need



PYRUKYND® expansion into diseases with larger patient populations provides significant near-term growth potential for first- and best-in-class therapies



3-8K patients in the U.S./EU5

PK deficiency 2022

Approved for adults in the U.S., EU and Great Britain

OUR GOAL
Deliver the first
approved therapy for
pediatric PK deficiency

18-23K patients in the U.S./EU5

~70K patients in GCC

>1M patients worldwide

Thalassemia 2025

Potential U.S. approval

OUR GOAL Deliver the first therapy approved for all thalassemia subtypes

120-135K patients in the U.S./EU5

~150K patients in GCC

>3M patients worldwide

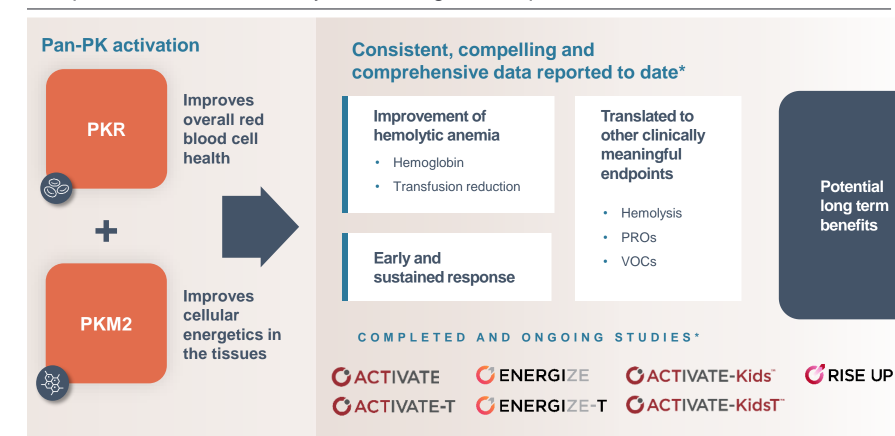
Sickle cell disease 2026

Potential U.S. approval

OUR GOAL Deliver a novel oral therapy that improves anemia and reduces VOCs



Unique PK activation mechanism has demonstrated comprehensive benefits beyond hemoglobin improvement





2020 Pipeline: the beginning of a rare disease portfolio leveraging our expertise in cellular metabolism

COMPOUND	INDICATION	PRECLINICAL EARLY-STAGE LATE-STAGE REGULATORY APPROVAL CLINICAL DEVELOPMENT SUBMISSION
PYRUKYND® First-in-class PK activator	Pyruvate Kinase Deficiency (PKD)	ACTIVATE AND ACTIVATE- T
	α- and β-Thalassemia	PHASE 2
AG-946 Novel PK activator	Healthy Volunteers / Sickle Cell Disease (SCD)	PHASE 1

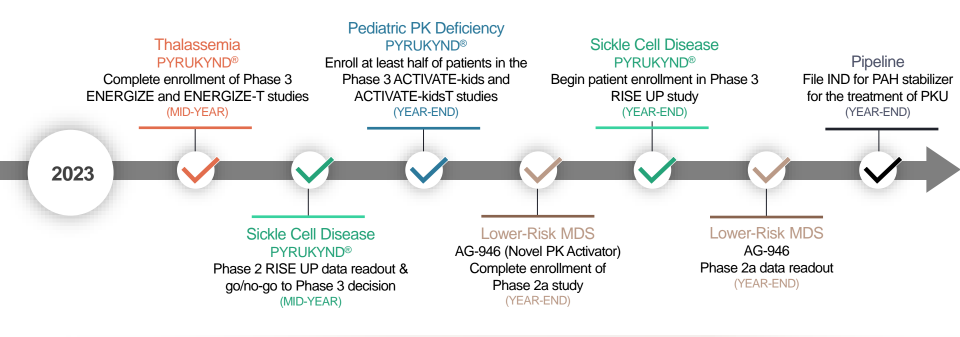


2024 Pipeline: significant advancement building depth and breadth in our rare disease pipeline

COMPOUND	INDICATION	PRECLINICAL EARLY-STAGE LATE-STAGE REGULATORY APPROVAL CLINICAL DEVELOPMENT SUBMISSION
	Pyruvate Kinase Deficiency (PKD)	US, EU, GB
		ACTIVATE KIDS - T
PYRUKYND® First-in-class		ACTIVATE KIDS
PK activator	α- and β-Thalassemia	ENERGIZE
		ENERGIZE - T
	Sickle Cell Disease (SCD)	RISE UP
AG-946 Novel PK	Healthy Volunteers / Sickle Cell Disease	
activator	Lower Risk Myelodysplastic Syndrome (LR-MDS)	
Phenylalanine hydroxylase (PAH) stabilizer	Phenylketonuria (PKU)	
siRNA Targeting TMPRSS6	Polycythemia Vera (PV)	



Momentum building as we delivered on all 2023 goals to expand and advance our pipeline and strengthened clinical evidence across our PKa franchise



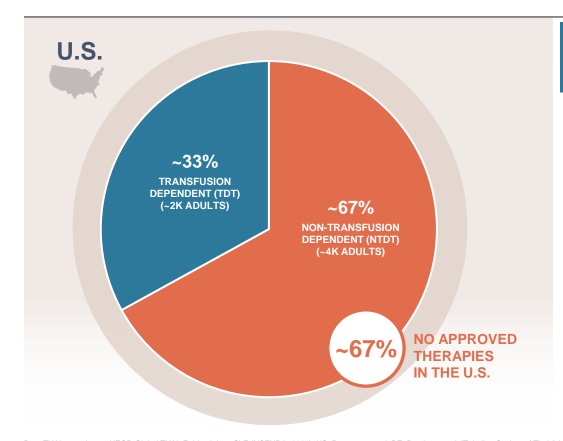
Evaluate business development opportunities to expand pipeline and build commercial capabilities to efficiently launch additional indications



Pipeline

License Agreement with Alnylam for novel siRNA for potential treatment of PV (Q3 2023)

Agios aims to deliver the first therapy approved for all thalassemia subtypes



Mitapivat Thalassemia Phase 3 program

CENERGIZE

- Alpha- and Beta-thalassemia Nontransfusion dependent patients
- Primary endpoint: Hemoglobin (Hb) response
- Data announced January 2024

CENERGIZE-T

- Alpha- and Beta-thalassemia
 Transfusion dependent patients
- Primary endpoint: Transfusion Reduction Response
- Data readout mid-2024



Thalassemia in all forms remains an area of high unmet need, high morbidity and significant impact on quality of life

18-23K patients in the U.S./EU5

~70K patients
in GCC

>1M patients
worldwide

Disease Impact



Increased Mortality

Survival is lower than the general population and is significantly worse in those who remain non-regularly transfused than those who are regularly transfused



Serious, Irreversible Morbidities

High rates of morbidities (even compared to TDT – e.g., thrombosis); frequency of complications increasing as patients age



Poor Quality of Life

Adult patients with NTDT may have similar or worse Healthcare Related QoL compared with patients with TDT



Healthcare Resource Utilization & Cost

A 1g/dL decrease in average Hb levels is associated with increased inpatient, outpatient and ER visits/costs, Rx costs, and total healthcare costs in patients with NTDT



ENERGIZE: achieved significance across both primary and all key secondary endpoints



Key findings in ENERGIZE trial

- Total of 194 patients were randomized 2:1 to 100 mg mitapivat (n=130) or placebo (n=64)
- Statistically significant increase in hemoglobin response rate (42.3%) compared to patients on placebo (1.6%)
- Statistically significant change from baseline in average FACIT-Fatigue score and average hemoglobin concentration
- During the 24-week double-blind period, 4 subjects in the mitapivat arm experienced adverse events (AEs) leading to discontinuation; no AEs in the placebo arm leading to discontinuation
- All pre-specified subgroup analyses favored the mitapivat treatment arm compared to placebo

Next Steps

- Full data set to be presented at an upcoming medical meeting
- ENERGIZE T readout expected by mid-year
- Data to be submitted together to FDA by year end
- Potential US launch in 2025



Thalassemia: an attractive global rare disease opportunity with limited or no treatment options and geographic concentration

US Thalassemia Market Opportunity

- 6,000 diagnosed adult patients in US
- Most patients diagnosed before adulthood
- Nearly 70% of patients have no treatment options
- Concentrated providers: ~50% of active adult patients in <150 hem/onc practices
- Recognized high unmet need
- Well-established ICD-10 codes
- Established patient advocacy organizations

+ GCC prevalence ~8-9x the US

PYRUKYND

- All-inclusive global trial design
- ENERGIZE demonstrated statistically significant results on primary and all key secondary endpoints
- ENERGIZE-T readout in mid-2024
- Opportunity to deliver best-in-class therapy and transform treatment for the full range of thalassemia patients
- Potential US launch in 2025



Sickle Cell Disease remains an area of significant need for innovative therapies that can demonstrate meaningful benefits beyond hemoglobin increase

Disease Overview

Genetic blood disorder that causes **sickling of red blood cells**

Caused by mutations in HBB gene leading to anemia, hemolysis and sickle cell pain crises

Market Opportunity



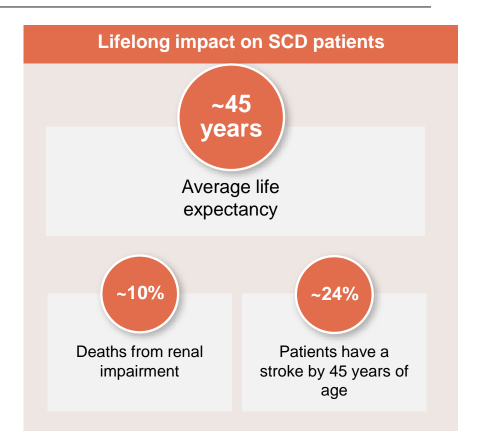
No novel oral therapy improves anemia and reduces sickle cell pain crises (SCPC)



Significant global opportunity

~100,000 patients in the U.S.

>3 million worldwide





PYRUKYND: a novel oral therapy with potential to be best-in-class improving anemia, reducing SCPCs and improving how patients feel and function



Phase 2 Data

- Statistically significant increase in hemoglobin response rate observed in both doses compared to placebo
- Improvements in markers of hemolysis and erythropoiesis observed at both doses compared to placebo
- A trend in sickle cell pain crises reduction was observed at both doses compared to placebo
- No adverse events (AEs) leading to discontinuation

Phase 3 Design⁽²⁾

- Phase 3 primary endpoints:
 Hb response⁽³⁾ and annualized rate of SCPCs
- N = 198 with a 2:1 randomization (100 mg mitapivat and placebo)
- 52-week double blinded period followed by 216-week open label extension

PYRUKYND

- Seamless Phase 2/3 global study designed with community input
- Potential for mitapivat to:
 - · improve anemia
 - reduce sickle cell pain crises
 - improve how patients feel and function
- Expected data readout in 2025
- Potential US launch in 2026



Abbreviations: BID = twice daily; Hb = hemoglobin; SCPC = sickle cell pain crises

^{(1) 100}mg was selected for Phase 3 portion of the study

⁽²⁾ Phase 2 and phase 3 components are part of a single study/protocol

⁽³⁾ Hb response is defined as a ≥ 1.0 g/dL increase in average Hb concentration over Weeks 24–52 compared with baseline

Fueling growth beyond 2026, an early-stage pipeline addressing the underlying pathophysiology of rare diseases with high unmet need

Lower-Risk MDS

~75-80k patients in the U.S. and EU5

No oral therapy addresses ineffective erythropoiesis

40% Achieved Transfusion Independence endpoint in open-label Phase 2a study



Expect to initiate Phase 2b study of AG-946 in mid-2024

Phenylketonuria (PKU)

~35-40k patients in the U.S. and EU5

Limited treatment options; patients consume high restricted diet

IND filed December 2023



Expect to initiate Phase 1 study of oral PAH stabilizer in 2024

Polycythemia Vera (PV)

~100k patients in the U.S.

Phlebotomy is the standard of care

In-licensed siRNA from Alnylam in 2023



Advancing to pre-IND studies



Continuing clinical and regulatory milestone momentum into 2024

2024

EARLY

MID-YEAR

YEAR-END



Phase 3 data readout for ENERGIZE study (COMPLETED)

Pipeline

Begin Phase 1 dosing for PAH stabilizer for the treatment of PKU

Thalassemia PYRUKYND®

Phase 3 data readout for ENERGIZE-T study

Pediatric PK Deficiency PYRUKYND®

Complete enrollment Phase 3 ACTIVATE kids study

Lower-Risk MDS

AG-946 (Novel PK Activator)

Regin patient enrollment of

Begin patient enrollment of Phase 2b study Thalassemia PYRUKYND®

Filing for FDA Approval

Sickle Cell Disease PYRUKYND®

Complete Phase 3 enrollment

Pediatric PK Deficiency PYRUKYND®

Phase 3 data readout ACTIVATE kids-T study



Strong beginning of 2024 with positive Thalassemia Phase 3 ENERGIZE readout and four additional Phase 3 readouts expected by the end of 2025

2024

2025

2026



Phase 3 ENERGIZE readout (completed)

Thalassemia PYRUKYND®

Phase 3 ENERGIZE-T readout

Pediatric PK Deficiency PYRUKYND®

Phase 3 ACTIVATE kids-T readout

Sickle Cell Disease PYRUKYND®

Phase 3 RISE UP readout

Thalassemia PYRUKYND®

Potential approval

Pediatric PK Deficiency PYRUKYND®

Phase 3 ACTIVATE kids readout

Sickle Cell Disease PYRUKYND®

Potential approval

Pediatric PK Deficiency PYRUKYND®

Potential approval



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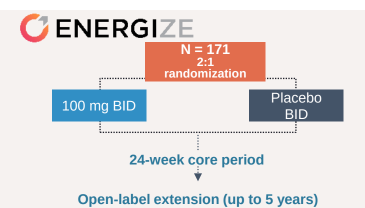
agios

Thank you



Appendix

PYRUKYND®: first Phase 3 program to encompass full range of thalassemia patients



Primary endpoint

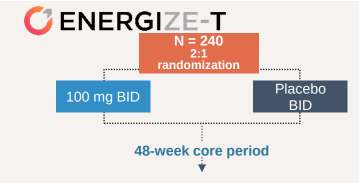
Mean Hb ↑
 ≥ 1 g/dL from baseline

Secondary endpoints

 Fatigue, additional measures of Hb \(^1\), hemolysis, patientreported outcomes, physical activity, iron metabolism, safety, PK/PD

Key inclusion criteria

- ≥ 18 years
- β-thalassemia ± α-globin mutations, HbE β-thalassemia, or α-thalassemia (HbH disease)
- Non-transfusion-dependent defined as ≤5 RBC units during the 24-week period before randomization and no RBC transfusions ≤8 weeks prior
- Hb ≤ 10.0 g/dL



Open-label extension (up to 5 years)

Primary endpoint

 50% reduction in transfusion burden in any 12-week rolling period

Secondary endpoints

 Additional measures of transfusion reduction, safety, PK/PD

Key inclusion criteria

- ≥ 18 years
- β-thalassemia ± α-globin mutations, HbE β-thalassemia, or α-thalassemia (HbH disease)
- Transfusion-dependent defined as 6 to 20 RBC units transfused and ≤6-week transfusion-free period during the 24-week period before randomization



Phase 3 ENERGIZE study: primary endpoint achieved



- Total of 194 patients were randomized 2:1 to 100 mg mitapivat (n=130) or placebo (n=64)
- Hemoglobin response is defined as ≥1.0 g/dL (10 g/L) increase in average Hb concentrations from Week 12 through Week 24 compared with baseline.
- Treatment with mitapivat demonstrated a statistically significant increase in hemoglobin response rate compared to placebo

Primary Endpoint	Placebo N=64	Mitapivat 100 mg BID N=130
Hemoglobin responders, n (%)	1 (1.6)	55 (42.3)
Adjusted difference of response rate (Mitapivat-Placebo), %		40.9
95% CI		(32.0, 49.8)
2-sided p-value		<0.0001

Abbreviations: RBC = red blood cell; Hb = hemoglobin. Subjects who do not have at least 2 on-treatment Hb concentration assessments between Week 12 and Week 24 are considered non-responders.

Baseline is defined as the average of all assessments within 42 days before randomization for subjects randomized and not dosed or within 42 days before the start of study treatment for subjects randomized and dosed.







Key secondary endpoints: change from baseline in both hemoglobin concentration and FACIT-Fatigue Score

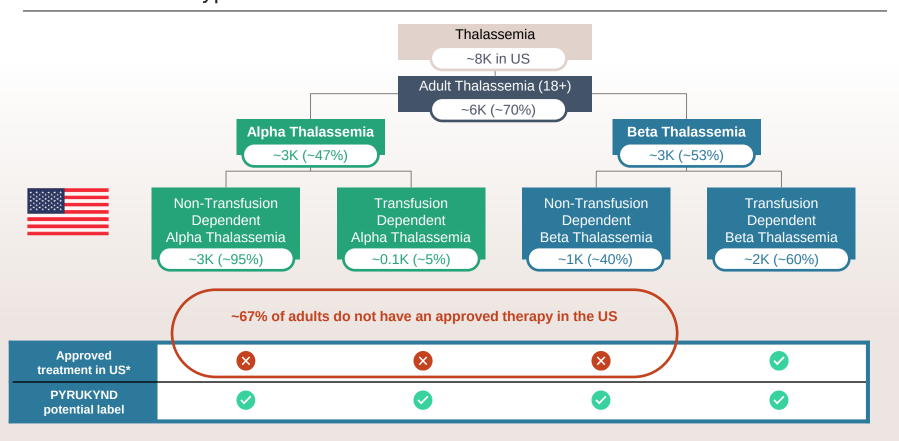
- Change from baseline in average FACIT-Fatigue (Functional Assessment of Chronic Illness Therapy-Fatigue)
 subscale score from Week 12 to Week 24
- Change from baseline in average hemoglobin concentration from Week 12 to Week 24
- Treatment with 100 mg mitapivat demonstrated statistically significant improvements on both key secondary endpoints compared to placebo

Safety

- Overall, incidence of adverse events was similar across mitapivat and placebo arms.
- During the 24-week double-blind period, 4 (3.1%) subjects in the mitapivat arm experienced adverse events (AEs) leading to discontinuation; there were no AEs in the placebo arm leading to discontinuation



PYRUKYND® has the potential to become the first therapy approved for all thalassemia subtypes





Advancing RISE UP Phase 3 Study of PYRUKYND® in sickle cell disease with expected readout in 2025

Phase 3 primary endpoints (1):

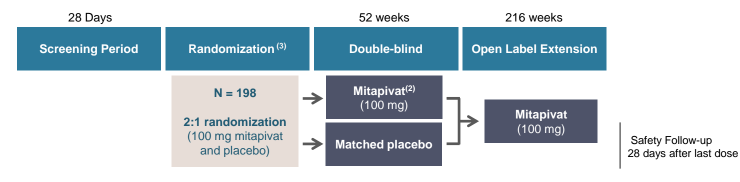
Hb response, defined as a ≥ 1.0 g/dL increase in average Hb concentration over Weeks 24–52 compared with baseline, and annualized rate of SCPCs

Key inclusion criteria

- ≥ 16 years of age
- Documented SCD (HbSS, HbSC, HbSβ0/HbSβ+ thalassemia, other SCD variants)
- Recurrent VOCs (vaso-occlusive crises) defined as the occurrence of 2–10 SCPCs (acute pain needing medical contact, acute chest syndrome, priapism, hepatic or splenic sequestration) in the prior 12 months
- Anemia defined as a Hb level of 5.5–10.5 g/dL
- If taking HU, the dose must be stable for ≥ 90 days before starting study drug

Key exclusion criteria

- Receiving regularly scheduled blood transfusions
- · Severe kidney disease or hepatobiliary disorders
- Currently receiving treatment with SCD therapies (excluding HU)
- Prior exposure to gene therapy, or prior bone marrow or stem cell transplantation





Treatment with mitapivat demonstrated a statistically significant increase in hemoglobin response rate compared to placebo



	Placebo N=27	Mitapivat 50 mg BID N=26	Mitapivat 100 mg BID N=26
Hemoglobin responders, n (%)	1 (3.7)	12 (46.2)	13 (50.0)
Difference of response rate (Mitapivat-Placebo), %		42.5	46.3
95% CI ⁽¹⁾		(18.8, 63.4)	(22.0, 66.8)
2-sided p-value ⁽²⁾		0.0003	0.0001

Abbreviation: RBC = red blood cell

Hemoglobin response is defined as ≥1.0 g/dL (10 g/L) increase in average Hb concentrations from Week 10 through Week 12 compared to baseline.

Assessments collected within 8 weeks after an RBC transfusion are excluded from the analysis.

Subjects who do not have any Hb concentration assessments from Week 10 through Week 12 are considered nonresponders.

(1) Exact 95% C

(2) The p-value is based on the Fisher's exact test



Annualized rates of sickle cell pain crises for patients in the mitapivat arms were lower compared to patients in the placebo arm



CRC Adjudicated Data

Negative Binomial Regression Model

	Placebo N=27	Mitapivat 50 mg BID N=26	Mitapivat 100 mg BID N=26
Annualized Rate of SCPC	1.71	0.83	0.51
95% CI	(0.95, 3.08)	(0.34, 1.99)	(0.16, 1.59)
Rate ratio (Mitapivat/Placebo)		0.48	0.30
95% CI		(0.17, 1.39)	(0.08, 1.07)

Abbreviations: CRC = crisis review committee; SCPC = sickle cell pain crisis

SCPC events that occur within 7 days of a prior SCPC onset are not counted as a separate event. Each subject time in the Double-blind Period is defined as (end date – date of randomization + 1), where end date is last dose of study drug during the Double-blind Period for subjects randomized and dosed, or the randomization date for subjects randomized and not dosed.



The estimates and 95% CIs are based on a negative binomial regression model with natural log link. The model included the number of SCPC events during the Double-blind Period of the study as the response variable and treatment arm as the independent variable. The natural log of time on study was used as the offset to account for the varying lengths of subjects' time in the Double-blind Period of the study.