

Safety and Efficacy of Bezuclastinib (CGT9486), a Novel, Highly Selective, Potent KIT D816V Tyrosine Kinase Inhibitor, in Patients with Advanced Systemic Mastocytosis (AdvSM): Results From Part 1 of the Phase 2 Apex Trial

Pankit Vachhani¹, MD; Tsewang Tashi², MD; Gary Schiller³, MD; Stephanie Lee⁴, MD, MSc; Miguel Piris-Villaespesa⁵, MD, PhD; Cristina Bulai Livideanu⁷; Jonathan Lambert⁸, PhD, BSc, BMBS, FRCP, FRCPath; Anthony M. Hunter⁹, MD; Tracy I. George¹⁰, MD; Cristina Papayannidis¹¹, MD; Khalid Shoumariyeh¹², MD; Lei Sun¹³, PhD; Rita Petroro¹³, Jenna Zhang¹³, PhD; LouAnn Cable¹³; Amanda Pilla^{13;} Hina A. Jolin¹³, PharmD; Rachael Easton¹³, MD; Vinod Pullarkat¹⁴, MD, MRCP ¹University of Alabama Birmingham, ²Huntsman Cancer Institute, University of Utah, Division of Hematologic Malignancies, Salt Lake City, UT; ³David Geffen School of Medicine at UCLA, Los Angeles, ⁴St. Michael's Hospital, Toronto, ⁵Hospital, Toronto, ⁵Hospital, Toronto, ⁵Hospital, Toulouse, CHU Toulouse, CHU Toulouse, CHU Toulouse, CHU Toulouse, CHU Toulouse, CHU Toulouse, Salt Lake City, UT; ³David Geffen School of Medicine at UCLA, Los Angeles, ⁴St. Michael's Hospital, Toronto, ⁵Hospital, Toronto, ⁵Hospital, Toulouse, CHU Toulouse, C of Medicine, Atlanta; ¹⁰ARUP Laboratories, University of Utah School of Medicine, Salt Lake City; ¹¹IRCCS Azienda Ospedaliero-University of Freiburg; ¹³Cogent Biosciences, Inc., Waltham, MA; ¹⁴City of Hope Medical Center, Duarte, CA

INTRODUCTION

Unmet Need Remains for Advanced Systemic Mastocytosis Patients

- Advanced Systemic Mastocytosis (AdvSM) is an aggressive and life-threatening form of systemic mastocytosis (SM) that is primarily driven by KIT D816V mutation and leads to uncontrolled proliferation of mast cells (MC)^{1,2}
- Subtypes: aggressive SM (ASM); SM with associated hematologic neoplasm (SM-AHN); mast cell leukemia (MCL)
- Based on subtype, the median overall survival ranges from <6 months to 3-4 years^{3,4}
- Unmet need remains for approved therapies without associated clinically significant toxicities – Reported toxicities for marketed tyrosine kinase inhibitor (TKI) therapies include nausea, vomiting, diarrhea, edema, intracranial bleeding, and cognitive effects⁵⁻⁷

Bezuclastinib: Highly Selective and Potent KIT D816V Inhibitor

- Oral, selective type I tyrosine kinase inhibitor (TKI) with potent activity against KIT D816V, the driving mutation in 95% of SM
- Preclinically, highly active with specificity for mutations in KIT exons 9, 11, 17, and 18
- Spares closely related kinases and has minimal brain penetration and favorable PK properties⁸ – Inhibition of closely related kinases have been linked to off-target toxicities, such as bleeding, edema, and pleural effusions^{9,1}

Nonclinical Models Provide Strong Support for Bezuclastinib and Azacitidine^a Concomitant Therapy

- Colony-forming assays (CFU-Mk) were performed to determine the likelihood of drug-induced thrombocytopenia in patients when combining bezuclastinib with azacitidine
- Bezuclastinib plus azacitidine is **non-interactive** based on Bliss Synergy Score (-5.98), supporting clinical investigation as a concomitant therapy for SM-AHN patients¹¹



Figure 1. CFU-Mk Assay Predicts No Synergistic Drug-Induced Thrombocytopenia

	Human CFU-Mk	Fold Change Above	Azacitidine (μΜ)		M)									
Drug	Single Agent IC ₅₀ (nM)	Primary Target KIT D816V			-0.003;	-0.01	-0.033	-0.1	-0.33		-3.3	-10		
Bezuclastinib	5,000	357x		0.0033-									 100	K)
Azacitidine	1,200	n/a	í,	0.01-										s (M
Avapritinib	330	25x	ju) di	0.033-										ocyte bility
The effect of bezuclastinib combined with azacitidine was assessed using			stini	0.1-										aryo Via
CFU-Mk assays. The table reflects single agent IC ₅₀ s for each drug tested alone in the CFU-Mk assay. The heatmap represents human megakaryocyte viability			ucla	0.33-									50	∍gak ative
following a dose matrix of increasing concentrations of bezuclastinib and			Bez	1-										U-Me Relå
(ReachBio, Seattle, WA). Colors in the dose inhibition heatmap indicate low				3.3-										CFI
viability (red) and high viability (green).				10-									0	%

^aRefer to the Additional Planned Cohorts on the Apex Study Design for information on this cohort that is now enrolling

STUDY DESIGN

Figure 2. Apex (NCT04996875): A Phase 2 Open-Label, Multicenter Clinical Study of **Bezuclastinib in Patients with Advanced Systemic Mastocytosis**

FORMULATION A

KEY ENTRY CRITERIA

- ASM, SM-AHN, or MCL per
- WHO 2022 Classification Confirmed measurable
- disease per mIWG-MRT-
- ECNM (mIWG) No restrictions on prior
- therapy
- Platelet count ≥50 x 10^{9} /L

Primary Endpoint

• **Part 1:** Incidence of AEs/SAEs, laboratory changes, PK,

50 mg BID

100 mg BID

200 mg BID

400 mg QD

- biomarkers, ORR • Part 2: ORR (confirmed CR, CRh, PR and CI) per mIWG MRT-ECNM and assessed by Central Response Review Committee
- **Other Endpoints** • Safety/Tolerability: Incidence of AEs leading to dose modification, changes in Patient Reported Outcomes
- Efficacy: DOR, TTR, PFS, OS, pure pathologic response • PK/PD: plasma concentration of bezuclastinib, serum tryptase, KIT D816V burden, BM mast cells



RESULTS

Table 1. Patient Demographics and Characteristics

33 patients enrolled[§]: median age: 68 years: range: 33-87

Male, n (%) ECOG PS 0-1, n (%)	21 (65.6) 27 (84.4)	6 (75.0)	4 (57.1)		
ECOG PS 0-1, n (%)	27 (84.4)			5 (62.5)	6 (66.7)
		8 (100)	5 (71.4)	7 (87.5)	7 (77.8)
AdvSM Subtype per Central Eligibility Review, n (%)					
ASM	7 (21.9)	2 (25)	0	0	5 (55.6)
SM-AHN	23 (71.9)	5 (62.5)	6 (85.7)	8 (100)	4 (44.4)
MCL	2 (6.3)	1 (12.5)	1 (14.3)	0	0
Prior therapy for AdvSM, n (%) [∫]					
TKI Naïve*	22 (69)	7 (88)	4 (57)	6 (75)	5 (56)
Avapritinib	5 (16)	0	2 (29)	2 (25)	1 (11)
Midostaurin	10 (31)	1 (13)	3 (43)	2 (25)	4 (44)
SRSF2/ASXL1/RUNX1 Mutation in Peripheral Blood	19 (59.4)	5 (62.5)	5 (71.4)	5 (62.5)	4 (44.4)
<i>KIT</i> D816V in Whole Blood, Positive, n (%)	29 (90.6)	8 (100)	6 (85.7)	7 (87.5)	8 (88.9)
Median KIT D816V VAF, % (range)	6.1 (0-47.2)	3.4 (0-39.0)	29.2 (0-38.9)	2.9 (0-47.2)	1.9 (0-42.2)
Median Bone Marrow MC Burden, % (range)	30 (5-90)	50 (20-70)	70 (5-90)	10 (5-30)	40 (10-80)
Median Serum Tryptase, ng/mL (range)	153.5 (35.0-1578.0)	178.0 (130.0- 605.0)	233.0 (53.6- 1578.0)	97.1 (35.0- 131.0)	182.0 (50.2- 370.0)

^sOne patient never dosed was excluded

Additional therapies included cytoreductives and biologics Patients who have received no prior SM-directed therapy with midostaurin and/or avapritinit Data as of: 25Sep2023

Data as of: 25Sep2023

Figure 3. Dose Dependent Increase in Bezuclastinib Steady State Exposure with 100 mg BID (200 mg per day) Identified as Optimal





[¥]Figure was derived from mIWG population with PK available at C2D1 (n=25) *Overall response rate (ORR; CR+CRh+PR+CI) was assessed in all mIWG evaluable patients (n=27) Data as of: 25Sep2023

Bezuclastinib Continues to Demonstrate a Differentiated Safety Profile

Table 2. Treatment Related Adverse Events in > 10% Patients

	Total (n=32) n (%)		50 mg BID (n=8) n (%)	100 mg BID (n=7) n (%)	200 mg BID (n=8) n (%)	400 mg QD (n=9) n (%)	
Preferred Term	All grade	Grade ≥3	All grade	All grade	All grade	All grade	
Hair color changes	11 (34)	0	0	4 (57)	3 (38)	4 (44)	
Thrombocytopenia [*]	7 (22)	2 (6)	0	4 (57)	1 (13)	2 (22)	
Transaminase increased*	7 (22)	1 (3)	3 (38)	2 (29)	1 (13)	1 (11)	
Neutropenia [*]	6 (19)	3 (9)	1 (13)	2 (29)	1 (13)	2 (22)	
Taste disorder [*]	6 (19)	0	1 (13)	1 (14)	1 (13)	3 (33)	
Peripheral edema	4 (13)	0	0	1 (14)	1 (13)	2 (22)	
Periorbital edema	4 (13)	1 (3)	0	0	3 (38)	1 (11)	

*Includes pooled preferred terms

The majority of adverse events were of low grade and reversible.

• No related cognitive impairment or bleeding events reported.

• The majority of hematological adverse events were of low grade, reversible and did not require dose reduction

• Related SAEs reported in 4 patients including Gr4 Thrombocytopenia, Gr3 Hypersensitivity (mediator flare), Gr3 Leishmaniasis, and Gr3 DILI (presented with late onset [day 488] and mixed cholestatic pattern of injury and subject was subsequently found to have biliary outflow tract obstruction).

• 9/32 patients required dose reduction due to adverse events, 6 of whom were at 400 mg; 3/32

patients discontinued due to adverse events.

Bezuclastinib Demonstrates Deep Reductions in Markers of Mast Cell Burden



• 94% (30/32) of patients achieved $a \ge 50\%$ reduction

- 100% (29/29) of patients with at least 2 cycles of treatment achieved $a \ge 50\%$ reduction
- 53% (17/32) achieved below 20 ng/mL
- Median time to first serum tryptase <20 ng/mL was 4.0 weeks (range: 1.1-66.9)



[§]Five patients excluded: Three (3) were KIT D816V negative at baseline and two (2) had no post-baseline data *Central lab lower limit of detection of KIT D816V VAF by ddPCR is 0.03% mutated alleles

• 93% (26/28) of patients achieved $a \ge 50\%$ reduction

§Four patients without post-baseline data were excluded

Median time to first VAF below 1% was 9.0 weeks (range: 6.0-85.3)





• 97% (28/29) of patients with baseline and at least 1 post-baseline assessment achieved $a \ge 50\%$ reduction

- 79% (23/29) achieved complete clearance of mast cell aggregates by central review
- Median time to first clearance of mast cell aggregates was 9.0 weeks (range: 7.3-34.3)









Data as of: 25Sep2023

Data as of: 25Sep2023



safety outcomes

REFERENCES: 1. Pardanani A. Am J Hematol. 2021;96(4):508-525. 2. DeAngelo DJ et al. Nat Med. 2021;27(12):2183-2191. 3. Ustun C et al. Haematologica. 2016;101(10):1133-1143. 4. Lim K-H

Early and Sustained Responses Observed by mIWG-MRT-ECNM and PPR

Table 3. Apex Part 1: Responses Observed by mIWG-MRT-ECNM

Total* Confirmed and unconfirmed mIWG-MRT-ECNM Responses per CRRC Assessment (n=27)	Confirmed mIWG-MRT-ECNM Responses per CRRC Assessment (n=27)	mIWG-MRT-ECNM per CRRC Assessment* (TKI [‡] Therapy Naïve) (n=18)	mIWG-MRT-ECNM per CRRC Assessment* (Prior TKI [‡] Exposure) (n=9)
15 (56)	12 (44)	11 (61)	4 (44)
14 (52)	10 (37)	10 (56)	4 (44)
6 (22)	6 (22)	6 (33)	O (O)
8 (30)	4 (15)	4 (22)	4 (44)
1 (4)	2 (7)	1 (6)	0 (0)
9 (33)	12 (44)	6 (33)	3 (33)
3 (11)	3 (11)	1 (6)	2 (22)
	Total* Confirmed and unconfirmed mIWG-MRT-ECNM Responses per CRRC Assessment (n=27) 15 (56) 14 (52) 6 (22) 8 (30) 1 (4) 9 (33) 3 (11)	Total* Confirmed and unconfirmed mIWG-MRT-ECNM Responses per CRRC Assessment (n=27)Confirmed mIWG-MRT-ECNM Responses per CRRC Assessment (n=27)15 (56)12 (44)15 (56)12 (44)14 (52)10 (37)6 (22)6 (22)8 (30)4 (15)1 (4)2 (7)9 (33)12 (44)3 (11)3 (11)	Total* Confirmed mIWG-MRT-ECNM mIWG-MRT-ECNM MIWG-MRT-ECNM Responses per CRRC Assessment (n=27) Assessment (n=27) mIWG-MRT-ECNM 15 (56) 12 (44) 11 (61) 14 (52) 10 (37) 10 (56) 6 (22) 6 (22) 6 (33) 8 (30) 4 (15) 4 (22) 1 (4) 2 (7) 1 (6) 9 (33) 12 (44) 6 (33) 3 (11) 3 (11) 1 (6)

Data as of: 25Sep2023

Data as of: 25Sep2023

finding at baseline were Not mIWG-MRT-ECNM Evaluable (NE) and therefore are excluded: one additional patient was excluded due to discontinuation prior to first dose (Not Dosed [ND]) *4 patients who remain on therapy but have not vet reached the 12-week confirmation duration for partial response (PR) are included [‡] SM-directed therapy with midostaurin and/or avapritinib

PTable 4. Apex Part 1: Responses Observed by PPR Criteria

Best Response, n (%) ^α	Total (n=32)	PPR per Investigator Assessment (TKI‡ Therapy Naïve) (n=22)	PPR per Investigator Assessment (Prior TKI [‡] Therapy) (n=10)			
all response rate (CR + PR)	24 (75)	19 (86)	5 (50)			
lete Response (CR)	13 (41)	12 (55)	2 (20)			
al Response (PR)	11 (34)	7 (32)	3 (33)			
e Disease (SD)	5 (16)	2 (9)	3 (33)			
valuable	3 (9)	1 (5)	2 (20)			

^a One patient was excluded due to discontinuation prior to first dose (Not Dosed [ND]) [‡] SM-directed therapy with midostaurin and/or avapritinib

Figure 7. Early and Sustained Responses Observed by mIWG-MRT-ECNM Criteria

• Median duration on study = 34.7 (range: 0.43-89) weeks

• Median time to confirmed response (CR+CRh+PR+CI) = 2.02 (range 1.9-4.8) months • First confirmed CRh by mIWG documented as early as 8 weeks and first confirmed CR as early as 20 weeks

CONCLUSIONS

Bezuclastinib continues to demonstrate a differentiated safety profile

- The majority of adverse events reported were of low grade and reversible

No related cognitive impairment or bleeding events reported

– 28% of patients required dose reduction due to adverse events; 9% of patients discontinued due to adverse events

• Treatment with bezuclastinib resulted in encouraging signs of clinical activity

– 56% overall response rate (CR + CRh + PR + CI; confirmed and unconfirmed) per mIWG-MRT-ECNM and 75% ORR (CR +PR) per PPR criteria

- Deep reductions demonstrated across commonly used biomarkers of mast cell activity:

≥50% Serum Tryptase	≥50% KIT D816V VAF	≥50% Bone Marrow MC Burden
94% of patients	93% of patients	97% of patients

• Exposure achieved with 100 mg BID (200 mg per day) dose resulted in optimal efficacy and

– All patients receiving 100mg BID achieved PR or better and remain on trial with 3 patients at ≥20 cycles

– Dose of 100mg BID was well tolerated; majority of dose reductions occurred in patients receiving 400mg total daily dose (200 mg BID or 400 mg QD)

• Enrollment to Part 2 is ongoing

– 150 mg QD of the optimized formulation expected to deliver exposures consistent with 200 mg total daily dose of the original formulation

– A cohort evaluating bezuclastinib with concomitant AHN therapy, which is supported by nonclinical data, is open for enrollment