

EHA2022 HYBRID ## JUNE 9-17 # VIENNA



TAKEAIM LEUKEMIA- A PHASE 1/2A STUDY OF THE IRAK4 INHIBITOR EMAVUSERTIB (CA-4948) AS MONOTHERAPY OR IN COMBINATION WITH AZACITIDINE OR VENETOCLAX IN RELAPSED/REFRACTORY AML OR MDS

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Emavusertib, An Oral IRAK4 Inhibitor

Emavusertib:

- Selective, small molecule inhibitor of IRAK4
- ATP-competitive, type 1 inhibitor, reversible
- Excellent drug-like properties:
 - Orally bioavailable (>100% dog/mouse)
 - Moderate plasma binding (77% human)
 - Stable in plasma, liver microsomes, hepatocytes
 - No inhibition of 7 major CYP450s
 - No significant metabolism in vitro
 - \circ Humans: rapid absorption/clearance, T_{1/2} 6 hr, no accumulation with QD dosing

IRAK4/Emavusertib Co-crystal Structure



2.4Å resolution



Emavusertib: Introduction



- Emavusertib (CA-4948), a novel oral IRAK4 inhibitor has potential anti-leukemia activity
- Specific genetic mutations (*SF3B1*, *U2AF1*) in the spliceosome drive overexpression of IRAK4 long isoform (IRAK4-L)
- IRAK4-L then causes constitutive activation of the myddosome, leading to overactivity of NF-κB
- Therefore, this drug can target patients with splicing mutations
- Emavusertib also targets FLT3 and has shown potential synergetic activity with other drugs



Emavusertib: Preclinical Activity in AML and MDS





both azacitidine and venetoclax in THP-1 model²



Emavusertib: Study Design

TakeAim Leukemia (NCT #04278768): Open-label, single arm, Phase 1/2 dose escalation and expansion 3+3 study design in R/R AML or high-risk MDS (HR-MDS)



Study Objectives

- 1º: Determine maximum tolerated dose Determine recommended Phase 2 dose
- 2º: Pharmacokinetic (PK) profile Preliminary anti-cancer activity

Study Population

- Relapsed/Refractory AML or high-risk MDS
- ECOG performance Status of ≤ 2
- Age ≥ 18 years

Dosing

- Oral, BID Dosing
- 28-day cycles

All the data was extracted on Dec 16, 2021. Patients began enrollment into the combination therapy portion of the study in November 2021.

- 1. These are non-targeted patients, due to lack of spliceosome or *FLT3* mutation, this population will be addressed in the combination therapy study
- 2. One patient was not response evaluable because of discontinuation due to patient decision
- 3. Two AML patients have both a spliceosome and *FLT3* mutation and are included in both populations (there are 13 total evaluable patients with spliceosome or *FLT3* mutation)
- 4. Six patients did not start treatment by September 30th, 2021, which did not allow 2 on-study disease assessments



Emavusertib: Baseline Characteristics

			AML/MDS Subsets				
		All patients (n=49)	AML Spliceosome ¹ (n=6)	MDS Spliceosome (n=7)	AML FLT3 ¹ (n=3)		
Female n (%) : Male n (%)		16 (33) : 33 (67)	0 (0) : 6 (100)	5 (71) : 2 (29)	0 (0) : 3 (100)		
Age (yrs): median (range)		74 (32, 87)	76 (60, 84)	74 (61, 80)	80 (78, 87)		
ECOG: n 0/1/2		11/30/8	0/4/2	2/5/0	0/1/2		
Median platelets (10 ³ /mm ³) (range)		30 (4, 275)	28 (21, 80)	16 (7, 146)	21 (9, 23)		
Median ANC (10 ³ /mm ³) (range)		0.64 (0, 14.75)	0.23 (0, 3.3)	1.85 (0.15, 11.0)	0.05 (0, 0.11)		
Median bone marrow blasts (%) (range)		-	33 (20, 95)	8 (3, 12)	60 (39, 95)		
Median lines of prior therapy (range)		2 (1, 5)	2.5 (1, 4)	2 (1, 4)	2 (1, 4)		
	HMA ²	-	6 (100)	7 (100)	3 (100)		
Prior therapy, n (%)	Chemotherapy ³	-	3 (50)	0 (0)	1 (33)		
	Venetoclax	-	4 (67)	1 (14)	3 (100)		

1. Two AML patients have both a spliceosome and FLT3 mutation and are included in both populations (there are 13 total evaluable patients with spliceosome or FLT3 mutation)

2. HMA includes azacitidine, decitabine, and guadecitabine

3. Chemotherapy includes cytarabine



Emavusertib: Toxicities Profile

• During the initial dose escalation phase, no DLT was observed in 200-400 mg BID dose levels. Additional patients were enrolled at 300 mg and 400 mg BID to further explore the safety profile.

Grade 3+ Treatment-Related Adverse Event	200 mg BID (N = 3)	300 mg BID (N = 26) ¹	400 mg BID (N = 17)	500 mg BID (N = 3)	
	n (%)	n (%)	n (%)	n (%)	
Number of patients having grade 3+ TRAEs	1 (33.3)	6 (23.1)	6 (35.3)	2 (66.7)	
Alanine aminotransferase increased	1 (33.3)				
Blood creatine phosphokinase increased		1 (3.8)			
Dizziness	1 (33.3)				
Dyspnoea			1 (5.9)		
Enterobacter infection			1 (5.9)		
Fatigue			1 (5.9)		
Gastrointestinal haemorrhage		1 (3.8)			
Hypophosphataemia		1 (3.8)			
Hypotension		1 (3.8)			
Lipase increased		2 (7.7)			
Platelet count decreased		1 (3.8)			
Presyncope			1 (5.9)		
Rhabdomyolysis		1 (3.8)	2 (11.8)	1 (33.3)	
Syncope				1 (33.3)	

1. Data for the two patients that have escalated from 300 mg BID to 400 mg BID were included in the 400 mg BID dose group.

One death occurred after the data extraction date, currently under review.



Emavusertib: Single-agent Activity in AML and HR-MDS



Only evaluable patients with baseline and post-treatment bone marrow blast counts are included in the waterfall plot; among the patients w/o targeted mutations (SF3B1 / U2AF1 / FLT3 mutation), 1 reached CR and 2 PR

Subset of patients with targeted mutations (SF3B1 / U2AF1 / FLT3 mutation)

Best Response	Efficacy		
Population #1: AML Spliceosome Patients ^{1, 2}			
CR/CRh Rate	2/5 (40%)		
CR	1/5 (20%)		
CRh	1/5 (20%)		
Population #2: MDS Spliceosome Patients			
Objective Response Rate (ORR)	4/7 (57%)		
CR	0/7 (0%)		
mCR	4/7 (57%)		
Population #3: AML FLT3 Patients ¹			
CR/CRh Rate	1/3 (33%)		
CR	1/3 (33%)		
CRh	0/3 (0%)		

1. Two AML patients have both a spliceosome and *FLT3* mutation and are included in both populations (there are 13 total evaluable patients with spliceosome or *FLT3* mutation)

2. One patient was not response evaluable because of discontinuation due to patient decision

* Indicates the best percentage change from baseline >100%



CR

Best Response

CRh

Mar

0%

Baseline

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Emavusertib: Single-agent Activity in R/R AML with Spliceosome Mutation



Data extraction date: Dec 16, 2021; "+" in Duration of Treatment indicates the patient remains on treatment as of the date of data extraction. 1. Two AML patients have both a spliceosome and *FLT3* mutation and are included in both populations (there are 13 total evaluable patients with spliceosome or *FLT3* mutation).

Emavusertib achieved 40% CR/CRh rate, despite transformed AML being historically highly resistant to treatment AML Spliceosome Mutation



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Emavusertib: Single-Agent Activity in R/R HR-MDS with Spliceosome Mutation

MDS								
	Dose (BID)	IPSS-R	Baseline Molecular Mutations	# of Prior Therapies	Duration on emavusertib (mos)	Blasts Baseline	Blasts Best Response	% Change
15%	200 mg	Very High Risk	U2AF1 ,ASXL1, NF1, PHF6, GFI1, KDM6A, TET2	1	5.7	11	2	-82% (mCR)
Spliceosome	300 mg	Very High Risk	U2AF1, DNMT3A, BCOR, STAG2, BCORL1, ETV6, SETBP1	1	3.3+	12	5	-58% (mCR)
10% Spliceosome	400 mg	Very High Risk	SF3B1, RUNX1, NFE2	2	4.3	7	3	-57% (mCR)
Spliceosome Spliceosome	300 mg	High Risk	SF3B1, DNMT3A, ASXL1, TET2, EZH2	2	0.9	8	4	-50% (mCR)
5% mCR	300 mg	High Risk	U2AF1, ASXL1	4	5.3+	3	2	-33%
Spliceosome mcR mCR	300 mg	Very High Risk	SF3B1, ASXL1, NF1, SH2B3, RUNX1, PHF6, CBL, GFI1, EZH2	3	1.6	8	9	13%
0% Baseline Best Response	400 mg	Very High Risk	U2AF1, ASXL1, BCOR, DNMTA, GATA2, SETBP1	1	1.2	9	62	>100%
Patient went to SCT $ -$	Data extraction date: Dec 16, 2021; "+" in Duration of Treatment indicates the patient remains on treatment as of the date of data extraction.							

Emavusertib achieved 57% ORR, including one patient who was able to proceed to transplant

MDS Spliceosome Mutation



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Emavusertib: Single-agent Activity in R/R AML with *FLT3* Mutation

1000/	FLT3								
100% -	FLT3	Dose (BID)	Risk (ELN)	Baseline Molecular Mutations	# of Prior Therapies	Duration on emavusertib (mos)	Blasts Baseline	Blasts Best Response ¹	% Change
75% st Keduction 50%	FLT3	400 mg	Adverse	FLT3 (eradicated at C3D1), ASXL1, BCOR, CEBPA (eradicated at C3D1), CSF3R, EZH2, NRAS, RUNX1 (X3), STAG2, TET2(X2,1) (eradicated at C3D1)	2	5.1	60	5	-92%
Z5%	pliceosome and FLT3	300 mg	Intermediate	FLT3 (eradicated at C4D1), BCOR (eradicated at C4D1), U2AF1 (decreased to 1.3 VAF at C4D1), WT1 (eradicated at C4D1)	1	6.2+	39	4	-90% (CR)
	CR	300 mg	Adverse	FLT3, SF3B1, NRAS, PTPN11, RAD21, RUNX1, TET2, GATA, STAT3	4	2.6	95	77	-19%
0%	Baseline Best Response	Data extraction	n date: Dec 16, 2021; tients have both a spl	"+" in Duration of Treatment indicates the patient re iceosome and <i>FLT3</i> mutation and are included in bo	mains on treatment	t as of the date of data re are 13 total evaluabl	extraction.	liceosome or FLT3 r	nutation).

Emavusertib achieved 33% CR rate, and FLT3 mutation eradicated in 2 out of 3 patients AML FLT3 Mutation



Summary



- Emavusertib has a manageable safety profile
- Demonstrates oral, single-agent, anti-cancer activity in heavily pretreated AML and HR-MDS patients with targeted mutations (*U2AF1*, *SF3B1*, or *FLT3*)
- Potential candidate for use in combination therapy for all AML/HR-MDS patients, including patients without a targeted mutation

Next Steps:

- Correlative analysis ongoing
- Trials in lymphoma and solid tumors are being explored

We would like to thank the patients, their families and caregivers for their invaluable contribution and participation in this study.



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Q & A