

## MOLECULAR CHARACTERIZATION OF CLINICAL RESPONSE IN RELAPSED/REFRACTORY ACUTE MYELOID LEUKEMIA AND HIGH-RISK MYELODYSPLASTIC SYNDROME PATIENTS TREATED WITH SINGLE AGENT EMAVUSERTIB

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# INTRODUCTION

- Acute myeloid leukemia (AML) and myelodysplastic syndrome (MDS) are myeloid malignancies that exhibit a dynamic mutational landscape as the disease progresses.
- Genetic mutations in splicing factors SF3B1 and *U2AF1* drive overexpression of a highly active long isoform of interleukin-1 receptorassociated kinase 4 (IRAK4), which is critical in triggering inflammation, oncogenesis, and survival of cancer cells.<sup>1, 2</sup>
- Emavusertib dual targeting of IRAK4 and FLT3 (ITD and TKD) confers a potential efficacy advantage.
- As of October 12, 2022, the ongoing TakeAim Leukemia trial (NCT04278768) has 71 patients treated with emavusertib monotherapy. The safety profile remains well tolerated without significant cumulative side effects.

# AIM

Sub-analysis of the monotherapy cohort (45 AML and 26 high-risk MDS [HR-MDS]) aims to:

- Investigate the mutational landscape at baseline and on-treatment
- Assess the association of targeted biomarkers and gene expression signatures with clinical activity

## METHOD

- Bone marrow and peripheral blood of enrolled patients were collected at the baseline and on treatment.
- Targeted next generation sequencing (NGS) and RNA sequencing were performed on bone marrow (BMMC) or peripheral blood mononuclear cells (PBMC).
- Mutations were also documented based on patients' molecular pathology reports provided by trial sites.

# RESULTS Table 1: Patient demographics Female n (%) Age (yrs): med Race n (%) Median ANC Median lines of Figure 1: Single-agent activity in R/R AML and HR-MDS All patients Without Targeted Mutation Response evaluable patients with baseline and post-treatment bone marrow blast counts are included. Responses assessed by the investigators were shown above. \* indicates the graphic cutoff as 100% Table 2: Characteristics of sequenced samples Total patien otal patien emale n Age (yrs): Diagnosi Cell type

- and resistance.

	All nationts	AML/MDS Subsets <sup>2</sup>		
	(n=71)	AML Spliceosome <sup>1</sup> (n=12)	MDS Spliceosome (n=12)	AML FLT3 <sup>1</sup> (n=8)
ale n (%)	21 (29.6) : 50 (70.4)	1 (8.3) : 11 (91.7)	5 (41.7) : 7 (58.3)	3 (37.5) : 5 (62.5)
(range)	74 (32, 87)	71 (60, 84)	75 (61, 80)	78 (61, 87)
n	1 (1.4)	0 (0)	1 (8.3)	0 (0)
k or African American	2 (2.8)	0 (0)	0 (0)	0 (0)
te	61 (85.9)	11 (91.7)	9 (75)	8 (100)
ers	7 (9.9)	1 (8.3)	2 (16.7)	0 (0)
(10 <sup>3</sup> /mm <sup>3</sup> ) (range)	26 (1, 275)	22 (1, 80)	11 (1, 146)	22 (1, 38)
<sup>3</sup> /mm <sup>3</sup> ) (range)	0.56 (0, 14.75)	0.3 (0, 3.3)	1.33 (0.15, 11.02)	0.13 (0, 0.88)
ior therapy (range)	2 (1, 6)	2.5 (1, 5)	2 (1, 4)	2.5 (1, 5)

. Three AML patients have both a spliceosome (U2AF1 or SF3B1) and FLT3 mutation and are included in both populations. 2. There are 29 total patients with spliceosome or FLT3 mutation, of which 24 are response evaluable subjects, 27 have received prior HMAs.

Targeted DNA sequencing					
amples (n)		106			
(n)		33			
: Male n (%)		12 (36%) : 21 (64%)			
dian (range)		74 (32 - 87)			
	AML	19			
	HR-MDS	14			
	BMMC	37			
	PBMC	69			

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



Two additional AML patients with spliceosome mutation had no posttreatment bone marrow blast count but reported progressive disease and were also considered as response evaluable. \* indicates the graphic cutoff as 10%

† indicates three AML patients with both a spliceosome and *FLT3* mutation that are included in both populations

RNA sequencing					
Total patient samples (n)		32			
Total patients (n)		20			
Female n (%) : Male n (%)		7 (35%) ; 13 (65%)			
Age (yrs): median (range)		74.5 (32 – 87)			
Diagnosis	AML	13			
	HR-MDS	7			
Cell type	BMMC	0			
	PBMC	32			

# CONCLUSIONS

Emavusertib monotherapy demonstrated anti-cancer activity in R/R AML and HR-MDS patients, especially in those with *FLT3*, *U2AF1*, or *SF3B1* mutations.

The preliminary NGS data is suggestive of molecular responses and disease-modifying activity of emavusertib.

Patients with targeted mutations responded to emavusertib, even in the presence of co-mutations (e.g., ASXL1, RUNX1) that are typically associated with poor prognosis. We will continue to explore potential genomic biomarkers of emavusertib sensitivity



### REFERENCES

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