Ivosidenib (IVO) prior to hematopoietic cell transplant for patients with IDH1-mutant relapsed or refractory acute myeloid leukemia (R/R AML)

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IVO 500 mg QD, R/R AML

BACKGROUND

- · Allogeneic hematopoietic cell transplantation (HCT) provides a potentially curative option for patients with relapsed or refractory (R/R) acute myeloid leukemia (AML)¹
- Pre-HCT remission status is a major determinant of long-term prognosis^{2,3}
- Older and/or heavily pretreated patients frequently cannot tolerate intensive salvage chemotherapy to obtain adequate disease control prior to HCT⁴
- Mutations in the metabolic enzyme isocitrate dehydrogenase 1 (IDH1) are detected in approximately 6-10% of patients with AML⁵⁻⁷ and result in the production of D-2-hydroxyglutarate (2-HG)
- 2-HG production is suppressed through targeted inhibition of the mutant IDH1 (mIDH1) enzyme, which restores cell differentiation8
- Ivosidenib (IVO) is approved in the US for the treatment of AML with a susceptible IDH1 mutation as detected by an FDA-approved test in:
- adults with newly diagnosed AML who are ≥ 75 years of age or who have comorbidities that preclude the use of intensive induction chemotherapy adults with R/R AML

OBJECTIVE

 To assess HCT outcomes in 18 patients with mIDH1 R/R AML who proceeded to HCT after responding to treatment with IVO in the AG120-C-001 phase 1 study

METHODS

- Here we report outcomes in patients with mIDH1 R/R AML from the phase 1 study who received a starting dose of IVO 500 mg once daily (QD), responded to treatment, and then proceeded to HCT
- This was a multicenter, open-label, dose-escalation and expansion study enrolling patients \geq 18 years of age with an advanced m/DH1 hematologic malignancy (ClinicalTrials.gov NCT02074839)9
- IVO monotherapy was administered orally, daily, in continuous 28-day cycles (Figure 1) During dose escalation, IVO was administered at doses of 200–1200 mg daily: 500 mg QD was selected for expansion
- · Per protocol, patients with R/R AML achieving an adequate response to IVO and meeting other criteria required for transplant could proceed to HCT after discontinuation of IVO

Figure 1. Study design Single-arm, open-label, phase 1, multicenter trial (ClinicalTrials.gov NCT02074839) Dose expansion (n = 180) Dose escalation (n = 78) Enrollment complete: 500 mg QD in continuous 28-day cycles R/R AML in 2nd+ relapse, relapse after HCT, refractory to Patients with m/DH1+ induction or reinduction, or relapse within 1 year, n = 126 advanced hematologic malignancies Untreated AML not eligible for standard of care, n = 25 Oral ivosidenib daily in continuous Other non-AML m/DH1 R/R advanced hematologic 28-day cycles malignancies, n = 11 Doses included 100 mg BID. 300, 500, 800, 1200 mg QD 4 Other R/R AML not eligible for Arm 1. n = 18 HCT subgroup: R/R AML, responded to IVO 500 mg QD, and proceeded to HCT, n = 18 From N Engl J Med. DiNardo CD et al. Durable Remissions with lyosidenib in IDH1-Mutated Relapsed or Refractory AML, 378., Supplementary Appendix

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- mIDH1 variant allele frequency (VAF) from bone marrow mononuclear cells was assessed using BEAMing digital PCR (0.02-0.04% VAF detection limit)9
- · Baseline co-mutation analysis was performed by next-generation sequencing on bone marrow samples9
- The data cutoff date for this analysis was 02Nov2018

Patients who underwent HCT Overall cohor (n = 18) (n = 179)^a Median (range) age, years 61.5 (36-68) 67.0 (18-87) 89/90 Female/male n 8/10 Prior history of MDS, n (% 1(5.6)29 (16.2) AML classification, n (%) 120 (67.0) De novo 15 (83.3) Secondary 3 (16.7) 59 (33.0) ECOG PS, n (%) 7 (38.9) 36 (20.1) 9 (50.0) 99 (55.3) 2 (11.1) 42 (23.5) 2 (1.1) Prior regimens, n (%) 2 (1.1)° 10 (55.6) 75 (41.9) 5 (27.8) 52 (29.1) ≥ 3 3 (16.7) 50 (27.9) Prior therapy type,d n (%) Intensive chemotherapy 18 100.0 127 (70.9) Nonintensive therapy 5 (27.8) 115 (64.2) Investigational 4 (22.2) 55 (30.7) Prior HCT for AML n (%) 43 (24.0) 2 (11.1) Cytogenetic risk status, n (%) Intermediate 12 (66.7) 105 (58.7) Poor 3 (16.7) 50 (27.9) Unknown 5(2.8)Missing 3 (16.7) 19 (10.6) Baseline cytogenetic results, n (%) Normal 10 (55.6) 60 (33.5) Abnormal 5 (27.8) 100 (55.9) Missing 3 (16.7) 19 (10.6) Prior AML therapy outcomes, en (%) Relapsed after transplant 2 (11.1) 43 (24.0) In second or later relapse 2 (11.1) 26 (14.5) Refractory to initial induction/reinduction therapy 13 (72.2) 106 (59.2) Relapsed ≤ 1 year of initial therapy^f 1 (5.6) 17 (9.5) Other 2 (11.1) 20 (11.2)

Table 1. Baseline demographic and disease characteristics

RESULTS

Baseline characteristic

*The overall cohort of 179 patients with R/R AML treated with IVO in the phase 1 study. *Patients met eligibility criteria at screening but had a decline in ECOG PS at The observation of the second se second sec ve Cancer Network guidelines. ECOG PS = Eastern Cooperative Oncology Group Performance Status; MDS = myelodyspl

Table 2. BOR, duration on IVO, and last response prior to HCT for patients who

	underwen	it HCT (n = 18)	
Patient	BOR on IVO	Duration on IVO, days	Time from last IVO dose to HCT, days	Last response evaluation prior to HCT

CRh	227	1	CRi	12.7ª
CR	63	18	CR	6.5
CR	105	18	CR	28.5 ^a
CR	113	35	CR	2.6
CR	190	13	CR	7.3
MLFS	107	9	NE	15.3
CR	130	12	CR	29.4ª
CRi/CRp	72	15	CRp	7.6
	68	31	MLFS	6.0
CRi/CRp	90	23	CRp	3.4
	67	8	MLFS	0.8
CR	462	50	RL	17.2ª
CR	320	10	CRp	31.1ª
CR	125	8	CR	35.7ª
CR	195	5	CR	4.5
CR	196	14	NE	14.2ª
CR	195	13	CR	7.7
CR	86	14	CR	15.8 ^a
	CR CR CR MLFS CR CR//CRp CR//CRp CR//CRp CR//CRp CR//CRp CR//CRp CR CR CR CR CR CR CR CR CR CR	CR 63 CR 105 CR 113 CR 190 MLFS 107 CR 130 CR//CRp 72 CR//CRp 68 CR//CRp 69 CR//CRp 67 CR 462 CR 320 CR 125 CR 195 CR 196 CR 196	CR 63 18 CR 105 18 CR 113 35 CR 190 13 MLFS 107 9 CR 130 12 CR/CRp 72 15 CR/CRp 68 31 CR/CRp 67 8 CR 422 50 CR 320 10 CR 125 8 CR 195 5 CR 196 14 CR 195 13	CR 63 18 CR CR 105 18 CR CR 113 35 CR CR 190 13 CR CR 190 13 CR CR 130 12 CR CR 130 12 CR CR/CRP 72 15 CRp CR/CRP 72 15 CRp CR/CRP 72 15 CRp CR/CRP 68 31 MLFS CR/CRP 67 8 MLFS CR 320 10 CRp CR 320 10 CR CR 125 8 CR CR 195 5 CR CR 196 14 NE CR 195 13 CR

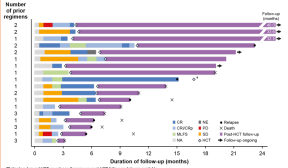
Indicates censored observatio BOR = best overall response: CR = complete remission: CRb = complete remission with partial hematologic recovery: CRi = complete remission with incomplete hematologic recovery; CRp = complete n survival; RL = relapse on with incomplete platelet recovery; MLFS = morphologic leul

· Baseline demographic and disease characteristics are reported in Table 1

- For patients who underwent HCT (n = 18), median (range) duration of IVO treatment prior to HCT was 3.9 (2.1-15.2) months
- In the HCT subgroup, the BOR on IVO prior to HCT was CR in 66.7% (12 / 18) of patients, and last response prior to HCT was CR in 50% (9 / 18) of patients (Figure 2, Table 2)

The median (range) time from last IVO dose to HCT was 13.5 (1–50) days





atient underwent HCT as salvage therapy; no post-HCT follow-up data are available = not assessed; PD = progressive disease; SD = stable disease

In the HCT subgroup:

- Median (95% CI) OS was 16.8 months (9.2, NE), calculated from the start of IVO treatment, compared with 9.0 months (7.1, 10.2) in the overall R/R AML study cohort (Table 3)
- 6-month OS was 94.4% and 12-month OS was 61.1% (Table 3)
- Median (range) duration of follow-up was 33.2 months (3.2–41.9)

Table 3, OS and RFS outcomes

Outcome	IVO 500 mg QD, R/R AML		
	Patients who underwent HCT (n = 18)	Overall cohort (n = 179) ^a	
OS ^b			
Median (95% CI), months	16.8 (9.2, NE)	9.0 (7.1, 10.2)	
Censored, c n (%)	8 (44.4)	32 (17.9)	
Survival rates, %			
6 months	94.4	61.9	
12 months	61.1	37.5	
OS post HCT ^d			
Median (95% CI), months	11.5 (6.0, NE)	-	
Censored, n (%)	8 (44.4)	-	
Survival rates, %			
6 months	77.8	-	
12 months	50.0	-	
RFS post HCT ^e			
Median (95% CI), months	7.3 (2.6, NE)		
Censored, n (%)	6 (35.3)		
Survival rates, %			
6 months	58.8	-	
12 months	47.1		

cause. Five patients in remission, two relapsed and in survival following, and one total to following. "Calculated as the time from the date of death due to any cause. "Five patients in remission, two relapsed and in survival following, and one total to following. "Calculated as the time from the date of death due to any cause." "Five patients in termission, two relapsed and in survival following. "Calculated as the time from the date of death due to any cause." "Five patients in termission, two relapsed and the date of death due to any cause." "Five patients in termission to the date of death due to following. "Calculated as the time from the date of death due to following." Calculated as the time from the date of death due to following. "Calculated as the time from the date of death due to following." Calculated as the time from the date of death due to following. "Calculated as the time from the date of death due to following." Calculated as the time from the date of death due to following. "Calculated as the time from the date of death due to following." Calculated as the time from the date of death due to following. The date of the dat

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· For patients achieving a BOR of CR, median (95% CI) OS was:

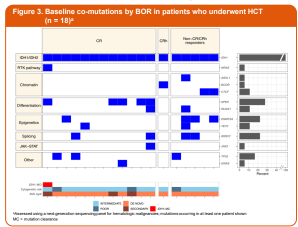
- NE (9.1, NE) in the HCT subgroup (n = 12)
- 20.5 months (16.4, NE) in those who did not undergo HCT (n = 31)
- Survival post HCT (Table 3);
- Median (95% CI) RFS post HCT was 7.3 months (2.6, NE); 6- and 12-month RFS rates post HCT were 58.8% and 47.1%, respectively
- 6- and 12-month post-HCT OS rates were 77.8% and 50.0%, respectively
- In the HCT subgroup, mIDH1 clearance occurred in 1 of 12 (8.3%) patients with BOR of CR, and in 0 of 1 patient with BOR of CRh (Table 4)

Table 4. IDH1 mutation clearance status at any assessment prior to HCT

IDH1 mutation clearance, n / N (%)	IVO 500 mg QD, R/R AML		
	Patients who underwent HCT (n = 18)	Overall cohort (n = 179) ^a	
Detection limit 0.02–0.04% ^b			
All patients	1 / 18 (5.6)	14 / 145 (9.7)	
CR	1 / 12 (8.3)	12 / 43 (27.9)	
CRh	0 / 1 (0)	2 / 14 (14.3)	

rall cohort of 179 patients with R/R AML treated with IVO in the phase 1 study. ∜When ≤ 1% VAF cutoff was applied. /DH1 mutation cleara in 6 of 18 (33.3%) patients in the HCT subgroup, including 6 of 12 (50.0%) with CR

Baseline co-mutation profiles by BOR are shown in Figure 3



CONCLUSIONS

- IVO monotherapy is a potential treatment option to induce remissions prior to HCT for patients with m/DH1 R/R AML who were not previously considered candidates for intensive salvage therapy
- Post-transplant survival rates are encouraging and warrant further investigation of IVO monotherapy or combination salvage therapies prior to HCT
- The molecular clearance of mIDH1 before HCT does not appear to be a prerequisite for successful HCT
- The potential of IVO is being assessed in other HCT settings
- An ongoing phase 1 study (ClinicalTrials.gov NCT03564821) is assessing IVO in post-HCT maintenance in patients with mIDH1 myeloid neoplasms

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