Antitumor Activity of Zanidatamab with Palbociclib and Fulvestrant in HER2+ Hormone-Receptor Positive (HR+) Metastatic Breast Cancer

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BACKGROUND

Figure 1: Unique Binding

Properties of Zanidatamab

ER2+ metastatic breast cancer (mBC) remains incurable e ongoing clinical need to develop new HER2-targeted therapies, including chemotherapy-free regimens Zanidatamab is a novel, bispecific, monoclonal antibody targetin 2 nonoverlapping extracellular domains (ECD4 and ECD2) on HER2 (Figure 1)1

Zanidatamab's unique binding properties result in:1 - Receptor clustering, internalization, and downregulation Inhibition of growth factor-dependent and

-independent tumor cell proliferation Antibody-dependent cellular cytotoxicity and



Approximately 50% of HER2+8C is also positive for the estrogen receptor (ER+)⁴. Signaling through HER2 and ER, along with cyclin D-CDK4 kinase, contributes to tumor growth and resistance to therapy. Targeting all 3 pathways may be of benefit to patients

Iriving

Here we explore the combination of zanidatamab with palbociclib (a CDK 4/6 inhibitor) plus fulvestrant (a sele down-regulator) as a novel chemotherapy-free treatment option

METHODS

 Study ZWI-ZW25-202 (NCT04224272) is an ongoing single-arm, open-label, international (Canada, Spain & USA), Phase 2 study of zanidatamab combined with palbociclib plus fulvestrant to treat patients with HER2+ HR+ mBC (Figure 2) · Part 1 of the study evaluated safety and tolerability of zanidatamab in combination with palbociclib plus fulvestrant. A safety monitoring committee (SMC) reviewed safety data and assessed dose-limiting toxicities (DLTs) in Part 1 to confirm recomm

doses (RDs) for Part 2 Part 2 of the study is evaluating anti-tumor activity of the combination at the RD level determined in Part 1

Per protocol, patients treated at the RD in Part 1 were included in the efficacy analysis

Total enrollment will be up to approximately 58 patients

Figure 2: ZWI-ZW25-202 Study Design (Parts 1 and 2 combined)

SELECT ELIGIBILITY CRITERIA		SINGLE ARM		ENDPOINTS
Unresectable, locally advanced and/or metastatic HER2+HR+BC ECOG PS ≤ 1 Prior treatment with tractivesiab performable and	→	Zanidatamab 20 mg/kg IV Q2W + Palboci(lib 125 mg PO QD (Doy 1-21) +	+	Primary Endpoint: • Part 1: Safety • Part 2: PF56 ^b
T-DM1 required No prior treatment with CDK4/6 inhibitor(s) Previously treated, stable brain metastases allowed		Fullvestrant SOD mg/MQGW ⁴ Roudy treatment cartinuu with disase progression, unacceptable tradition, pr other discontinuation ortheria are met		Select Secondary Endpoints: • ORR ^c • OS ^c • DCR ^d • DOR ^e • PFS ^b
		Tumor assessments QBW ^a		
breast cancer, CDK = cyclin-dependent kinase, CR = comp = homan epiderma growth factor 2; H4 = homane roce kod; P0 = by mouth; P4 = partial response; 50 = stable di	plete req gitor; ili sease; Q	pontar; DCR = disease control rate; BOR = duration of respo = instrumenoular; IV = instrumenous; DRR = objective respon (b = daily; Q2W = every 2 weeks; Q8W = every 4 weeks; Q8W	nse; ECOG se rate; OS III = every 8	PS = Eastern Cooperative Oncology Group performance eventil survival; PO = progressive disease, PFS = prog Reeeks; T-OMS = ado-trastazamb entransine.

Data were extracted on 31 August 2022 from an unloci 45 patients have been enrolled and treated in the stud

19 (42%) patients continue on zanidatamab treatment 2 (92-n) parents consists consists of an annual measurement of the second se

Table 1: Demographics and Baseline Characteristics	Patients (N=45
Median age (range), years	54 (36 - 77)
Female sex, n (%)	43 (96)
Race, n (%) White Aslan Other ⁴	37 (82) 2 (4) 6 (13)
ECOG PS, n (%) 0 1	23 (51) 22 (49)
HER2 ^b +/HR+, n (%) HER2 ^k -Pby central testing HR+ by local testing	25 (56) 25 (56) ^c 45 (100)
Prior history of brain metastases, n (%)	7 (16)
De novo metastatic disease, n (%)	16 (36)
Prior systemic anti-cancer therapy regimens in metastatic setting ⁴ , median (range) Prior systemic HER2 therapy in metastatic setting ⁴ , median (range)	4 (1-10) 3 (1-9)
Percent of patients with prior endocrine therapy in metastatic setting ^d , n (%) Median (range) Percent of patients with prior fulvestrant therapy, n (%)	35 (78) 1 (0-5) 9 (20)
Percent of patients with prior HER2 therapy , n(k) Taduumab Taduumab Lapatinb Total Total Total Total	45 (100) 45 (100) 44 (98) 36 (80)* 13 (29) 11 (24) 7 (16)

ECOG PS + Eastern Coopera a. American (srd), Not Reported (srd), Urknown (srd), and Other (srd). It Patients were allowed to enable based on local HER2 results. Central laboratory HER2 analysis arcentral testion. It contacts that seed HER2 meaning are to be for early that sees increasing for evaluation for analysis.

Safety

- Part 1: A DLT of Grade 4 neutropenia lasting > 7 days occurred in 1 of 7 DLT-evaluable patients, assessed by the investigator as related to palbociclib. All 3 agents were held; zanidatamab and fulvestrant resumed after 3 weeks, whereas palbociclib was resumed after 4 weeks at a reduced dose and when the neutropenia resolved to Grade \$ 2
- The SMC recommended that the starting doses (Figure 2) administered in Part 1 be used in Part 2 of the study
- Parts 1 and 2 (N=45): tment-related (related to zanidatamab, palbociclib and/or fulvestrant) adverse events (AEs) were Grade ≤ 2 The most common were diarrhea and neutronenia/neutronhil count decreased
- 34 (76%) patients experienced diarrhea of any grade vs 4 (9%) patients experienced Grade 3 diarrhea 28 (62%) patients experienced any grade of neutropenia/neutrophil count decreased, and 25 (56%) experienced a Grade 3 or Grade 4 event
- One zanidatamab- and palbociclib-related serious adverse event (SAE) of Grade 3 increased tra
- This event led to discontinuation of palbociclib There were no directionations of applications has fulwartrant due to an AF
- Dose reduction of zanidatamab due to an AE occurred in 2 patients: one had Grade 3 diarrhea and the other had Grade 3 creased transaminases
- Six deaths occurred and none were related to treatment: 5 due to disease progression (for 2 patients, cause of death was
- updated from "unknown" and "other" (respectively) to disease progression after the data cut off date) and 1 due to an unrelated AE of COVID 19



RESULTS

36 patients with measurable disease (defined as patients with at least 1 measurable target lesion at baseline and at least 1 post-baseline disease assessment or discontinued study due to death, progressive disease, or an AE) were evaluable for response

Median follow-up time was 14 months

Number of efficacy-evaluable patients with measurable disease still on treatment: 19 (53%) 9 of 12 confirmed responses ongoing; with response duration range of 2.1+ to 14.9+ months

Figure 3: Majority of Patients had Reduction in Target Lesion



C = tracativity Fale = fulvetrant; L = lagativity NE = not evaluable; P = persoannals; T-6MS = ado-trastaurenab eestassine; Trt = treatment; X = trastaurenab devanteran; cDH confirmed complete response; CH = confirmed partial response; CR = complete response; PD = progressive disease; PR = partial response; SD = trable disease.

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The primary endpoint of PFS6 could not be determined in this interim data analysis with study enrol interim median PFS (n=40 for mITT*) was 9.6 months (95% CI: 7.2, 16.6), range 1.58 to 16.72+ Iment ongoing

CONCLUSIONS

•	Interim analyses of activity and safety data of zanidatamab combined with palbociclib plus fulvestrant in heavily pretreated patients with advanced/metastatic HER2+ HER+ BC are encouraging
•	The regimen shows encouraging antitumor activity; in patients with response evaluable disease:
	 Most demonstrated reduction in the size of their target lesions
	 33% achieved a confirmed response
•	Durable disease control was demonstrated with a median PFS of 9.6 months in the mITT data set:
	9 of 12 patients with a confirmed response remained progression-free on therapy for > 1 year, including 7 still on treatment
	at the time of data cut
	 Of patients with a best response of stable disease, 3 remained progression free on therapy for > 1 year, and an additional 5 remained progression free on therapy for > 6 months
•	Zanidatamab in combination with palbociclib plus fulvestrant is well tolerated and shows a manageable safety profile
•	These results support further investigation of zanidatamab in combination with palbociclib plus fulvestrant, and enrollment in
	the study is continuing

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