

Tucatinib and trastuzumab in *HER2*-mutated metastatic breast cancer: a phase 2 basket trial

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A list of authors and their affiliations appears at the end of the paper

Human epidermal growth factor receptor 2 (*HER2*, also known as *ERBB2*) signaling promotes cell growth and differentiation, and is overexpressed in several tumor types, including breast, gastric and colorectal cancer. *HER2*-targeted therapies have shown clinical activity against these tumor types, resulting in regulatory approvals. However, the efficacy of *HER2* therapies in tumors with *HER2* mutations has not been widely investigated. SGNTUC-019 is an open-label, phase 2 basket study evaluating tucatinib, a *HER2*-targeted tyrosine kinase inhibitor, in combination with trastuzumab in patients with *HER2*-altered solid tumors. The study included a cohort of 31 heavily pretreated female patients with *HER2*-mutated metastatic breast cancer who were also *HER2* negative per local testing. Hormone receptor (HR)-positive patients also received fulvestrant. The overall response rate (primary endpoint) was 41.9% (90% confidence interval (CI): 26.9–58.2). Secondary endpoints of duration of response and progression-free survival were 12.6 months (90% CI: 4.7 to not estimable) and 9.5 months (90% CI: 5.4–13.8), respectively. No new safety signals were detected. Responses were observed across various *HER2* mutations, including mutations in the tyrosine kinase and extracellular domains. The chemotherapy-free regimen of tucatinib and trastuzumab showed clinically meaningful antitumor activity with durable responses and favorable tolerability in heavily pretreated patients with *HER2* mutations. These data support further investigation of *HER2*-targeted therapies in this patient population. ClinicalTrials.gov registration: [NCT04579380](https://clinicaltrials.gov/ct2/show/study/NCT04579380).

HER2 is a member of the epidermal growth factor receptor family of four related tyrosine kinases. When activated, *HER2* signals through the phosphoinositide 3-kinase (PI3K) and mitogen-activated protein kinase (MAPK) pathways to promote cell growth and differentiation. In tumors in which *HER2* is overexpressed or amplified (*HER2*-positive, *HER2*⁺), activation results in uncontrolled cell proliferation, inhibition of apoptosis and metastasis^{1–6}. *HER2* is a validated, actionable target, with *HER2*-targeted therapies showing clinical activity that resulted in regulatory approvals for the treatment of patients with *HER2*⁺ breast, gastric and colorectal cancers^{7–15}.

Recently, with the increased utilization of genomic profiling through next-generation sequencing (NGS), there has been a growing interest in *HER2* (also known as *ERBB2*) somatic mutations as oncogenic drivers, in the absence of *HER2*⁺ disease, in multiple solid tumors. In breast cancer, approximately 2–5% of tumors harbor *HER2* mutations^{16–21}. *HER2*-mutated (*HER2*-mut) breast cancer is more frequently associated with lobular histology and heavily pretreated patients with metastatic disease^{22–24}. *HER2*-targeted agents have shown clinical activity in *HER2*-mut metastatic breast cancer (MBC) in basket trials and case reports^{25–27}. However, there are currently no treatments

✉ e-mail: prpohlmann@mdanderson.org

that have been approved by the US Food and Drug Administration (FDA) or the European Medicines Agency specifically for patients with *HER2*-mut MBC.

Tucatinib is an oral tyrosine kinase inhibitor that is highly selective for *HER2* (ref. 28) and approved in multiple geographic regions for treatment of previously treated *HER2*⁺ MBC (combined with trastuzumab and capecitabine)^{11,29–31} and previously treated *HER2*⁺ metastatic colorectal cancer (combined with trastuzumab)¹². In addition, preclinical data have shown that tucatinib in combination with trastuzumab has antitumor activity in *HER2*-mut patient-derived xenograft tumor models of multiple solid tumors, suggesting that patients with activating *HER2* mutations may derive clinical benefit from the combination³². SGNTUC-019 (NCT04579380) is an open-label phase 2 basket study evaluating tucatinib in combination with trastuzumab in patients with *HER2*-altered solid tumors. Here we report results of the efficacy, safety and exploratory biomarker analyses from the *HER2*-mut MBC cohort of tucatinib in combination with trastuzumab (and fulvestrant in patients who have hormone-receptor-positive (HR⁺) disease).

Results

Patient characteristics

The first patient was enrolled in the basket study on 18 January 2021, and the last patient enrolled on 14 April 2023. Thirty-one patients were enrolled in the *HER2*-mut MBC cohort from September 2021 to December 2022. Patients with *HER2*-mut MBC were enrolled if they had measurable disease, had progressed during or after at least one previous line of systemic therapy in the locally advanced or metastatic setting, and had progressed during or were intolerant of the most recent line of systemic therapy; patients with *HER2*⁺ MBC were excluded. Demographics and baseline characteristics of patients with *HER2*-mut MBC are shown in Table 1. The median age was 64.0 years (range 43–76 years). Tumor histology was 58% lobular and 42% ductal. Most patients were HR⁺ (87%). Seven patients (23%) had stage IV disease at initial diagnosis. The median number of previous lines of therapy in any setting was 4 (range 1–10) and 3 (range 1–7) in the locally advanced or metastatic setting. Seventeen patients (55%) had previously received fulvestrant before enrollment.

Patient disposition is summarized in Fig. 1. As of the data cutoff (1 November 2023), the median duration of follow-up was 15.0 months (90% CI: 13.1–17.5). Nine patients remain on study treatment, 10 are in long-term follow-up and 12 are off study. Of the 22 patients who discontinued study treatment, 20 discontinued because of progressive disease and one patient each stopped treatment because of investigator decision or patient decision. No patient discontinued all study treatments because of a treatment-emergent adverse event (TEAE). Of the 12 patients that were off study, 11 were off study owing to death.

Efficacy

The confirmed objective response rate (cORR) per investigator assessment was 41.9% (90% CI: 26.9–58.2; Table 2), including 2 confirmed complete responses. Patients with HR⁺ or HR[–] MBC had a cORR of 44.4% (12 out of 27; 90% CI: 28.0–61.8) and 25.0% (1 out of 4; 90% CI: 1.3–75.1), respectively. Of the 31 patients, 13 were classified with ductal histology and 18 were classified with lobular histology. cORR was similar in both groups: 5 out of 13 for ductal histology (38.5% (90% CI: 16.6–64.5)) and 8 out of 18 for lobular histology (44.4% (90% CI: 24.4–65.9)). The median duration of response (DOR) was 12.6 months (90% CI: 4.7 to not estimable), and the median time to first response was 1.4 months (range 1.2–6.2). The disease control rate was 80.6% (90% CI: 65.3–91.2). Tumor size decreased in 22 patients (73%) of the 30 with a baseline and at least one post-baseline assessment (Fig. 2).

The median progression-free survival (PFS) was 9.5 months (90% CI: 5.4–13.8), with an estimated 6 month PFS of 65.0% (90% CI: 48.3–77.5) and an estimated 12 month PFS of 45.0% (90% CI: 28.8–60.0; Fig. 3a). The median overall survival (OS) was 20.1 months

Table 1 | Patient demographics and baseline characteristics of the *HER2*-mut MBC cohort

Characteristics	Total (N=31)
Median age, years (range)	64.0 (43, 76)
Age category, n (%)	
<65 years	16 (52)
≥65 years	15 (48)
Sex, n (%)	
Male	0
Female	31 (100)
Ethnicity, n (%)	
Hispanic, Latino or Latina, or of Spanish origin	2 (6)
Not Hispanic, Latino or Latina, or of Spanish origin	24 (77)
Unknown	1 (3)
Not reportable	4 (13)
Race, n (%)	
Asian	11 (35)
Black or African American	1 (3)
White	15 (48)
Not reportable	4 (13)
ECOG performance status score, n (%)	
0	19 (61)
1	12 (39)
Hormone receptor status	
Negative	4 (13)
Positive	27 (87)
Tumor type histology	
Ductal	13 (42)
Lobular	18 (58)
TNM stage at time of diagnosis	
I	5 (16)
II	6 (19)
III	13 (42)
IV	7 (23)
History of metastatic disease	
Yes	31 (100)
No	0
Number of previous systemic therapy regimens, median (range)	4 (1, 10)
Number of previous systemic therapy regimens in the locally advanced or metastatic setting, median (range)	3 (1, 7)

(90% CI: 15.9 to not estimable; Fig. 3b), with an estimated 12 month OS of 74.2% (90% CI: 58.6–84.7).

Safety

The median treatment duration with tucatinib was 9.0 months (range 0.7–24.4); with trastuzumab, 9.0 months (range 0.7–24.4); and with fulvestrant, 7.4 months (range 0.9–24.4). TEAEs were reported in all patients (Table 3 and Extended Data Table 1). The most common TEAEs were diarrhea (20 patients, 65%), nausea (11, 35%), vomiting and pruritus (9, 29%), and infusion-related reaction (8, 26%; Table 3). Among the 20 patients reported to have diarrhea, 4 patients (13%) had maximum

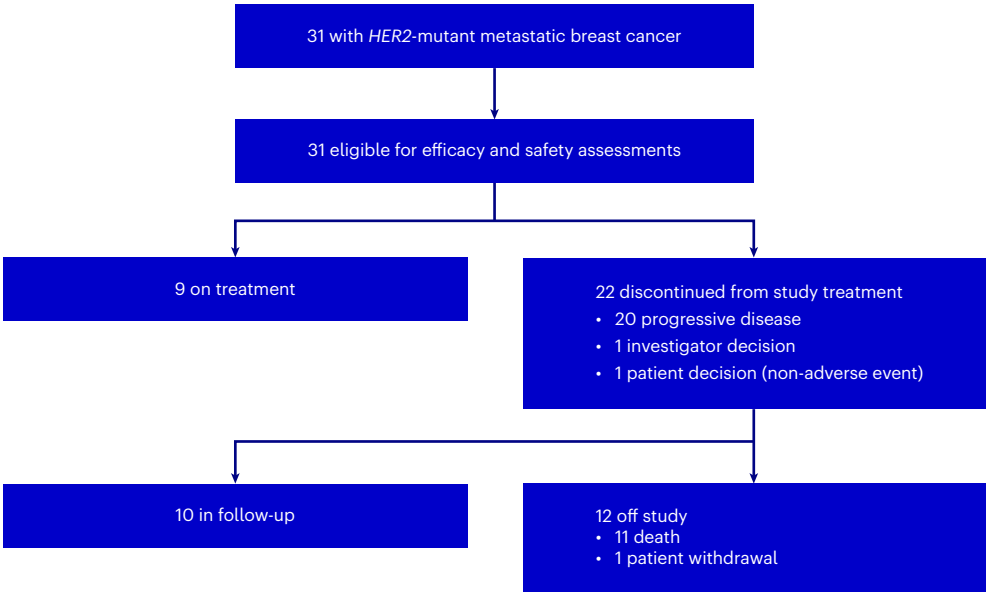


Fig. 1 | CONSORT diagram of the SGNTUC-019 study. Flow diagram detailing patients enrolled in the SGNTUC-019 study (NCT04579380) *HER2*-mutated MBC cohort.

severity of grade 3, while 12 patients (39%) and 4 patients (13%) had grade 1 and grade 2 diarrhea, respectively. The median time to resolution was 7.0 days (range 1–530) for any-grade diarrhea and 2.0 days (range 1–4) for grade ≥ 3 diarrhea. The other most common grade ≥ 3 events were increased levels of alanine aminotransferase and hypertension, with 3 patients (10%) reported for each. A total of 8 patients (26%) had serious TEAEs, of which 3 (10%) were related to tucatinib and none were related to either trastuzumab or fulvestrant.

No TEAEs resulted in the discontinuation of all study treatments. Tucatinib was discontinued because of TEAEs in 2 patients (6%), including an event of increased aspartate aminotransferase (grade 4) and an event of pseudocirrhosis (grade 2). There were no discontinuations of trastuzumab or fulvestrant due to TEAEs. TEAEs resulted in dose reduction of tucatinib in 7 patients (23%) and fulvestrant in 1 patient (3%). No TEAEs resulted in death. All 11 deaths (35%) were related to disease progression.

Exploratory biomarker analyses

Among the 31 patients enrolled in the *HER2*-mut MBC cohort, 28 had evaluable central immunohistochemistry (IHC) results, 27 had evaluable central fluorescence in situ hybridization (FISH) results, 24 had evaluable central tissue-based NGS results and 29 had evaluable blood-based NGS results. Responses were observed in patients with a range of *HER2* mutations, including mutations in both the tyrosine kinase and the extracellular domains (Fig. 2a). A listing of individual responses and *HER2* mutations as determined by local and central testing is found in Extended Data Table 2.

***HER2* alterations and response.** Among patients with a mutation identified by local testing ($n = 28$), central blood-based NGS ($n = 23$) or central tissue-based NGS ($n = 20$), *HER2* mutations (both single nucleotide variants and insertions and deletions (indels)) were most often located exclusively in the tyrosine kinase domain or exclusively in the extracellular domain. Depending on the type of assay used, 69.6–85.0% of patients had mutations exclusively in the tyrosine kinase domain and 15.0–21.4% of patients had mutations exclusively in the extracellular domain (Fig. 2a and Extended Data Table 3). The range of response rates across assays was 36.4–41.2% in patients with mutations exclusively in the tyrosine kinase domain and 50.0–66.7% in patients with mutations exclusively in the extracellular domain (Fig. 2a and Extended Data Table 3).

Table 2 | Summary of responses in the *HER2*-mut MBC cohort

	Total (N=31)
Best overall response ^a , n (%)	
CR	2 (6.5)
PR	11 (35.5)
SD	12 (38.7)
PD	4 (12.9)
Not available ^b	2 (6.5)
cORR, n (%)	13 (41.9)
90% CI ^c for cORR	(26.9, 58.2)
Median duration of objective response ^d (months) (90% CI) ^e	12.6 (4.7, –)
Disease control rate ^a , n (%)	25 (80.6)
90% CI ^c for disease control rate	(65.3, 91.2)
Median PFS (months) (90% CI) ^f	9.5 (5.4, 13.8)
Median OS (months) (90% CI) ^f	20.1 (15.9, –)

CI, confidence interval; cORR, confirmed objective response rate; CR, complete response; OS, overall survival; PFS, progression-free survival; PR, partial response; SD, stable disease; PD, progressive disease. In the case of SD, follow-up measurements must have met the SD criteria at least once after study entry at a minimum interval of 5 weeks. ^aConfirmed best overall response assessed per Response Evaluation Criteria in Solid Tumors (RECIST) v1.1. ^bIncludes patients whose disease assessments were not evaluable and patients with no post-baseline response assessment. ^cTwo-sided 90% exact CI, computed using the Clopper–Pearson method⁴⁰. ^dAs estimated using Kaplan–Meier methods. ^eDisease control defined as confirmed CR or confirmed PR or SD. ^fCalculated using the complementary log–log transformation method⁴¹.

The most commonly mutated amino acid residues in the *HER2*-mut MBC cohort were at amino acid residues Leu755 (a tyrosine kinase domain residue) and Ser310 (an extracellular domain residue). Depending on the type of assay used, 40.0–47.8% of patients had a mutation at Leu755, and 15.0–26.1% of patients had a mutation at Ser310 (Extended Data Fig. 1 and Extended Data Table 4). As this analysis was based on the presence or absence of an individual mutation, some patients had co-occurring *HER2* mutations. The range of response rates across assays was 33.3–37.5% in patients with a mutation in Leu755 and 50.0–66.7% in patients with a mutation in Ser310 (Extended Data Fig. 1 and Extended Data Table 4).

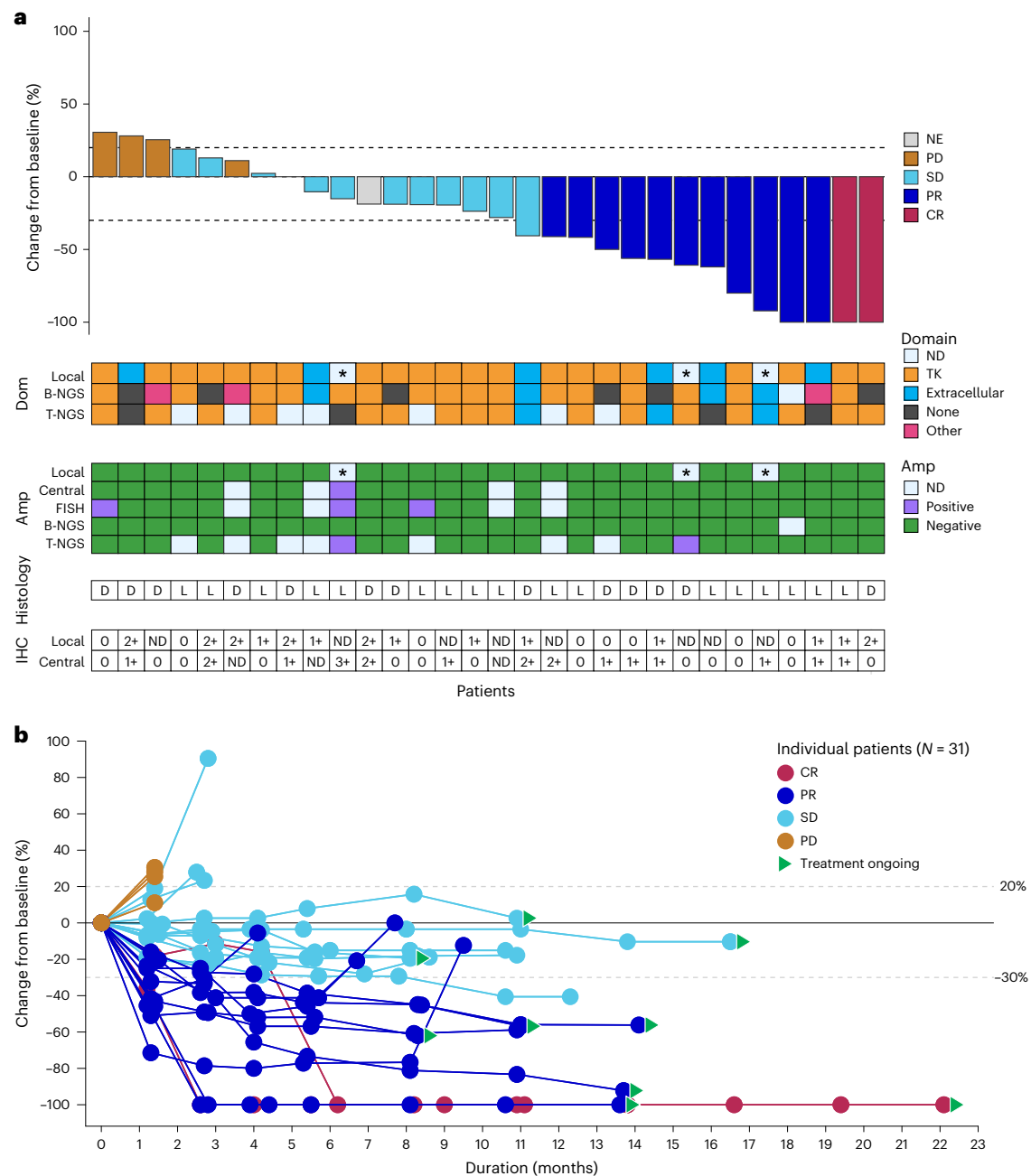


Fig. 2 | Response to study treatment and HER2 biomarker assessments.

a, Maximum percentage reduction in the sum of the diameters of target lesions per investigator and *HER2* biomarker assessments. Only patients who had baseline and post-baseline target lesion measurements are included ($n = 30$). The asterisks indicate patients that were excluded from any analysis of local results because their eligibility (*HER2*-negative status and *HER2* mutations) was determined by central blood-based NGS (B-NGS). T-NGS denotes tissue-based NGS testing by central lab assay. In all assays, 'ND' (not determined) indicates that there was no assay result, which may have been due to failed assay, failed quality

control or no assay run due to sample issues, and 'NE' denotes non-evaluable. In the 'Amp' (amplification) panel, 'central' indicates overall *HER2*-amplification status as determined by IHC with FISH reflex, both by central lab testing; whereas 'local' denotes local testing results. In the 'Domain' panel, patients had mutations exclusively in the domains indicated (that is, the tyrosine kinase (TK) or extracellular domains of *HER2*). The dashed line at 20% indicates progressive disease (PD). The dashed line at -30% indicates the threshold for a partial response (PR). D, ductal; L, lobular. **b**, Duration of treatment of patients in the *HER2*-mut MBC cohort.

Results showed a roughly equal distribution of patients who were *HER2*-low (IHC 1+ or IHC 2+/FISH-negative, $n = 12$) and those without *HER2* expression (IHC 0, $n = 14$). Based on central testing, the response rate was 50.0% (6 out of 12; 90% CI: 24.5–75.5) in *HER2*-low patients and 42.9% (6 out of 14; 90% CI: 20.6–67.5) in patients without *HER2* expression.

HER2 assay agreement. In current practice, multiple types of assay are used to identify *HER2* alterations. To determine whether patient

outcomes were impacted by this variability, results were analyzed among patients with sufficient samples available for standardized central testing. The percentage agreement for detection of *HER2* mutations (detected versus not detected) was 76.9% (90% CI: 59.5–89.4, $n = 26$) between local and central blood-based NGS, and 85.7% (90% CI: 67.1–96.0, $n = 21$) between local and central tissue-based NGS (Fig. 2a, Extended Data Fig. 1 and Extended Data Table 2). There was 100% agreement between any local test for *HER2* overexpression or amplification status versus central blood-based

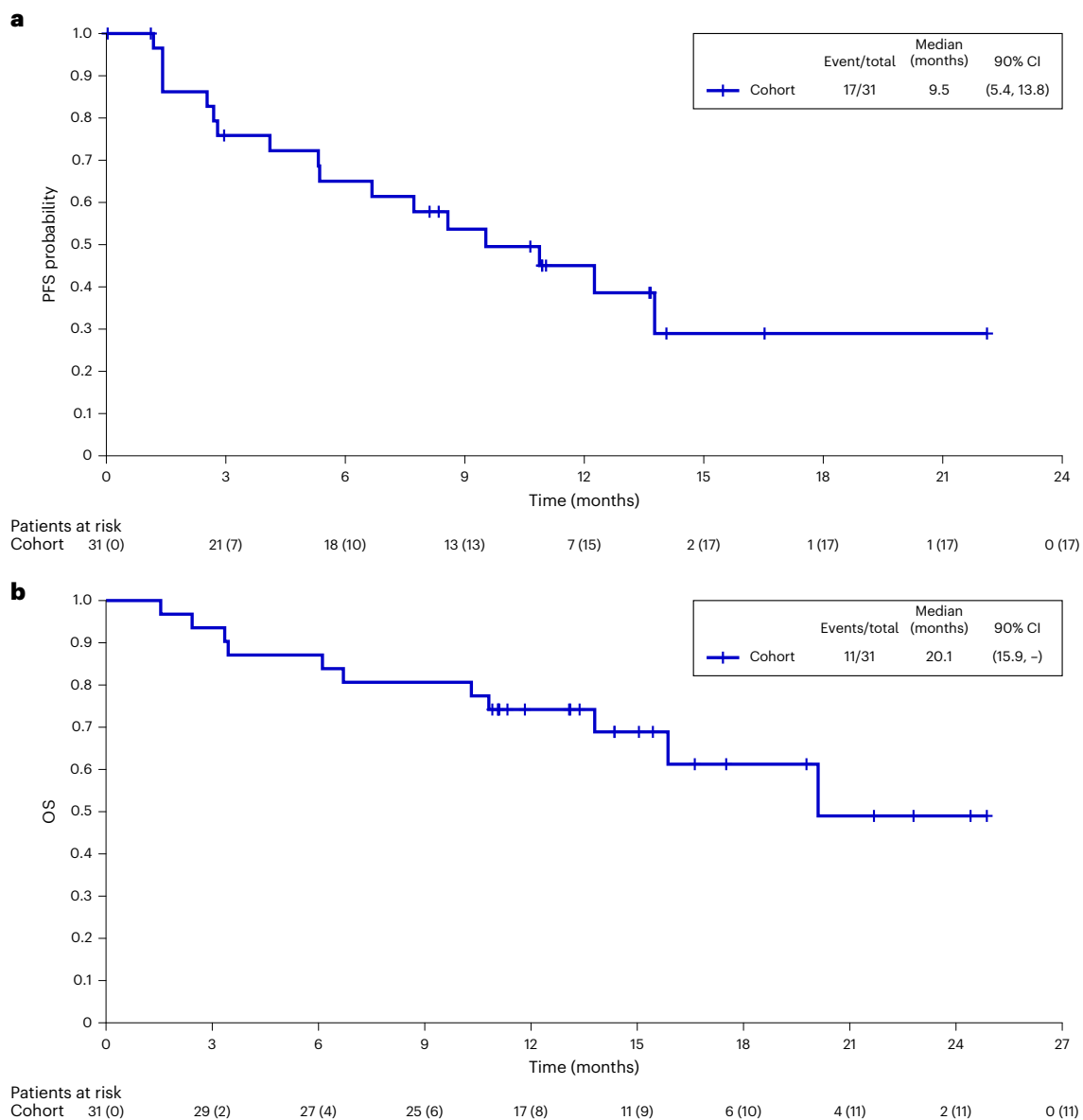


Fig. 3 | Kaplan–Meier curves for PFS and OS per investigator in the *HER2*-mut MBC cohort. The vertical lines indicate censoring.

NGS (90% CI: 89.1–100, $n = 26$) and central tissue-based NGS (90% CI: 86.7–100, $n = 21$) (Fig. 2a).

Concomitant non-*HER2* alterations. Additional concomitant non-*HER2* alterations (that is, amplified and/or with mutations or indels) identified using central tissue-based and blood-based NGS are shown in Extended Data Figs. 2 and 3. Owing to the small dataset (24 and 29 evaluable results in the tissue-based and blood-based NGS data, respectively), we examined genes that were altered in at least 25% of patients in a given assay. In the tissue-based NGS dataset, *CDH1*, characteristic of invasive lobular carcinoma, and *PIK3CA* were the only genes altered in at least 25% of patients. Responses were observed in patients with and without alterations in *CDH1* and *PIK3CA*. Response rates among patients with and without *CDH1* alterations were 62.5% (9 out of 15; 90% CI: 36.0–80.9) and 22.2% (2 out of 9; 90% CI: 4.1–55.0), respectively. Response rates among patients with and without *PIK3CA* alterations were 25.0% (2 out of 8; 90% CI: 4.6–60.0) and 56.3% (9 out of 16; 90% CI: 33.3–77.3), respectively. In the blood-based NGS dataset, only *PIK3CA* was altered in at least 25% of patients. Patients with

and without alterations in *PIK3CA* showed responses of 36.4% (4 out of 11; 90% CI: 13.5–65.0) and 44.4% (8 out of 18; 90% CI: 24.4–65.9), respectively.

Discussion

Over the past 25 years, monoclonal antibodies, antibody–drug conjugates and tyrosine kinase inhibitors targeting *HER2* have been developed that have resulted in notable improvements in PFS and OS for patients with *HER2*⁺ MBC^{7,8,11,13,33,34}. In the past decade, the use of NGS assays to molecularly profile and identify other potential targetable alterations in patients with MBC has increased substantially³⁵. *HER2* somatic mutations occur in about 2–5% of all patients with breast cancer, with a higher prevalence in the advanced disease setting^{16–21,35}. However, the benefit of using *HER2*-targeted therapies in patients with *HER2*-mut MBC, in the absence of *HER2*⁺ disease, is not well established; early data using *HER2*-targeted agents in this patient population have shown evidence of clinical activity²⁵.

In the SGNTUC-019 study, the combination of tucatinib and trastuzumab (with fulvestrant in patients with HR⁺ disease) is active

Table 3 | The most common TEAEs in the *HER2*-mut MBC cohort

Preferred term	Total (N=31)	
	Any grade, n (%)	Grade ≥3 ^a , n (%)
Any TEAE	31 (100)	15 (48)
Diarrhea	20 (65)	4 (13)
Nausea	11 (35)	0
Vomiting	9 (29)	0
Pruritus	9 (29)	0
Infusion-related reaction	8 (26)	0
Back pain	7 (23)	2 (6)
Cough	7 (23)	0
Edema peripheral	7 (23)	0
Chills	6 (19)	0
Pyrexia	6 (19)	0
Appetite decreased	6 (19)	2 (6)
Fatigue	6 (19)	1 (3)
Arthralgia	6 (19)	1 (3)
Dry skin	6 (19)	0
Alanine aminotransferase increased	5 (16)	3 (10)
Aspartate aminotransferase increased	5 (16)	2 (6)
Hypertension	5 (16)	3 (10)
Constipation	4 (13)	0
Insomnia	4 (13)	0
Blood creatinine increased	3 (10)	0
Anemia	3 (10)	0
Abdominal pain	3 (10)	0
Dyspnea	3 (10)	0
Headache	3 (10)	0
COVID-19	3 (10)	0
Stomatitis	3 (10)	0
Dizziness	3 (10)	0
Gastroesophageal reflux disease	3 (10)	0
Blood bilirubin increased	3 (10)	0
Myalgia	3 (10)	0
Onychoclasis	3 (10)	0

TEAEs reported in ≥10.0% of the patients who received at least a single dose of the study drug are listed here. The events are reported as per the preferred terms in the *Medical Dictionary for Regulatory Activities*. ^aGrade ≥3 TEAEs not listed here included acute kidney injury, cellulitis, chronic kidney disease, hepatic function abnormal, hydronephrosis, hypokalemia, influenza, sepsis and systemic inflammatory response syndrome; each was reported in one patient (3%).

in patients with heavily pretreated *HER2*-mut MBC, with a cORR of 41.9%. Moreover, the responses were durable, with a median DOR of 12.6 months. The response to study treatment occurred early, with the median time to first response of 1.4 months corresponding with the first tumor assessment on treatment. In addition, the median PFS and OS were 9.5 months (90% CI: 5.4–13.8) and 20.1 months (90% CI: 15.9 to not estimable), respectively. This clinically meaningful activity was seen despite patients having been heavily pretreated with a median of four previous lines of systemic therapy in any setting and three previous lines in the locally advanced or metastatic setting. This further adds to the data from the SUMMIT trial, which showed that *HER2* mutations are potentially actionable targets in MBC; treatment with neratinib in combination with fulvestrant and trastuzumab for patients with HR⁺,

HER2-mut MBC (*n* = 57) showed similar clinical activity (cORR, 39%, and median PFS, 8.3 months)²⁷.

In patients with tumor tissue available, central testing showed that almost all patients either were *HER2*-low (IHC 2+/FISH-negative or IHC 1+) or lacked expression of *HER2* (IHC 0). Responses to study treatment were similar in both patient populations, suggesting that the *HER2* mutation is an oncogenic driver in those patients.

The combination of tucatinib and trastuzumab, with fulvestrant for HR⁺ patients, was well tolerated and consistent with the established safety profile of the combination^{36,37}. Diarrhea was the most commonly reported adverse event; however, most events were grade 1 in severity and manageable with standard clinical care, with a grade 3 occurrence of 13%. Furthermore, antidiarrheal prophylaxis was not required on the study and no discontinuations of tucatinib occurred because of diarrhea. By contrast, grade 3 diarrhea occurred in 53% (30 out of 57) of patients receiving the combination of neratinib, fulvestrant and trastuzumab in the SUMMIT trial, despite the patients receiving loperamide prophylaxis²⁷. Only two patients (6%) discontinued tucatinib because of TEAEs, but remained on trastuzumab. No patients discontinued all study treatments because of an adverse event, highlighting the tolerability of this chemotherapy-free approach in patients with previous exposure to multiple lines of therapy and associated toxicities.

Exploratory biomarker analyses showed responses in patients harboring a variety of *HER2* mutations. There was evidence of clinical activity in patients with mutations in the tyrosine kinase domain and extracellular domain of *HER2*, specifically with mutations at Leu755 and Ser310, two of the most frequently mutated amino acids in *HER2* in multiple cancers, including breast cancers^{38,39}. Moderate-to-strong agreement between local and central test methods was observed, suggesting that both tissue-based and blood-based NGS may be used to identify patients harboring *HER2* mutations who may respond to treatment with tucatinib and trastuzumab. Finally, a variety of concomitant non-*HER2* genomic alterations were observed, including alterations in *CDH1* and *PIK3CA*. Clinical activity was observed across patients with a wide variety of alterations; however, larger datasets are required to draw any conclusions about the impact of concomitant non-*HER2* mutations on response.

One limitation of this analysis is the small size of the cohort (31 patients in total) and lack of a comparator arm. In addition, the duration of follow-up was short; therefore, the OS data are immature with relatively few events and should be interpreted with caution. Conclusions based on the exploratory biomarker data are also limited by the relatively small size of the cohort and the availability of data across multiple central testing methods (24 patients with evaluable tissue-based NGS and 29 with blood-based NGS). Conclusions may also be affected by assay differences between tissue-based and blood-based NGS, notably by differences in the set of genes included in each assay panel, as well as differential timing of tissue and blood collection in many cases.

In summary, the combination of tucatinib and trastuzumab (with fulvestrant in HR⁺ disease) was well tolerated and clinically active in patients with previously treated *HER2*-mut MBC. These results support further investigation of *HER2*-directed therapies in patients with *HER2* mutations.

Online content

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41591-024-03462-0>.

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Alicia F. C. Okines¹, Giuseppe Curigliano^{2,3}, Nobumasa Mizuno⁴, Do-Youn Oh⁵, Andree Rorive⁶, Hatem Soliman⁷, Shunji Takahashi⁸, Tanios Bekaii-Saab⁹, Mark E. Burkard¹⁰, Ki Y. Chung¹¹, Philip R. Debruyne^{12,13,14}, Jenny R. Fox¹⁵, Valentina Gambardella¹⁶, Marta Gil-Martin¹⁷, Erika P. Hamilton¹⁸, Bradley J. Monk¹⁹, Yoshiaki Nakamura²⁰, Danny Nguyen²¹, David M. O'Malley²², Alexander B. Olawaiye²³, Bhavana Pothuri²⁴, Martin Reck²⁵, Kazuki Sudo²⁶, Yu Sunakawa²⁷, Cedric Van Marcke²⁸, Evan Y. Yu²⁹, Jorge Ramos³⁰, Sherry Tan³⁰, Mark Bieda³⁰, Thomas E. Stinchcombe³¹ & Paula R. Pohlmann³²✉

¹The Royal Marsden NHS Foundation Trust, London, UK. ²Istituto Europeo di Oncologia, IRCCS, Milan, Italy. ³University of Milano, Milan, Italy. ⁴Aichi Cancer Center Hospital, Nagoya, Japan. ⁵Seoul National University Hospital, Cancer Research Institute, Seoul National University College of Medicine, Integrated Major in Innovative Medical Science, Seoul National University Graduate School, Seoul, South Korea. ⁶CHU Sart Tilman Liège, University of Liège, Liège, Belgium. ⁷H. Lee Moffitt Cancer Center and Research Institute, Tampa, FL, USA. ⁸Cancer Institute Hospital of JFCR, Tokyo, Japan. ⁹Mayo Clinic, Scottsdale, AZ, USA. ¹⁰UW Carbone Cancer Center, University of Wisconsin, Madison, WI, USA. ¹¹Prisma Health Institute, Greenville, SC, USA. ¹²Kortrijk Cancer Centre, General Hospital AZ Groeninge, Kortrijk, Belgium. ¹³Medical Technology Research Centre (MTRC), School of Life Sciences, Anglia Ruskin University, Cambridge, UK. ¹⁴School of Nursing and Midwifery, University of Plymouth, Plymouth, UK. ¹⁵Rocky Mountain Cancer Center, Boulder, CO, USA. ¹⁶Hospital Clínico Universitario de Valencia, Valencia, Spain. ¹⁷Institut Català d'Oncologia L'Hospitalet-IDIBELL, Hospitalet de Llobregat, Spain. ¹⁸Sarah Cannon Research Institute, Nashville, TN, USA. ¹⁹Florida Cancer Specialists and Research Institute, West Palm Beach, FL, USA. ²⁰National Cancer Center Hospital East, Kashiwa, Japan. ²¹City of Hope National Medical Center, Duarte, CA, USA. ²²The Ohio State University and James Comprehensive Cancer Center, Columbus, OH, USA. ²³University of Pittsburgh Medical Center, Pittsburgh, PA, USA. ²⁴Laura & Isaac Perlmutter Cancer Center, NYU Langone Health, New York, NY, USA. ²⁵Department of Thoracic Oncology, Airway Research Center North, Germany Center for Lung Disease, Grosshansdorf, Germany. ²⁶National Cancer Center, Tokyo, Japan. ²⁷St. Marianna University Hospital, Kawasaki, Japan. ²⁸Cliniques Universitaires Saint-Luc, Brussels, Belgium. ²⁹Fred Hutchinson Cancer Center/University of Washington, Seattle, WA, USA. ³⁰Pfizer, Bothell, WA, USA. ³¹Duke Cancer Institute, Durham, NC, USA. ³²University of Texas MD Anderson Cancer Center, Houston, TX, USA. ✉e-mail: prpohlmann@mdanderson.org

Methods

Inclusion and ethics

The study was conducted in accordance with regulatory requirements, the International Council for Harmonisation Good Clinical Practice guidelines and the Declaration of Helsinki. All patients provided written informed consent. The protocol was approved by institutional review boards and ethics committees according to the practice at each participating study site (Supplementary Information).

Patients and study design

SGNTUC-019 (NCT04579380) is an open-label, phase 2 basket study of patients with previously treated, locally advanced, unresectable or metastatic *HER2*-altered solid tumors. The breast cancer cohort comprised patients with *HER2*-mut locally advanced or metastatic disease, without evidence of overexpression or amplification of *HER2*, and enrolled patients at academic hospitals or cancer centers in North America, Europe and Asia. Patients must have progressed during or after at least one previous line of systemic therapy in the locally advanced or metastatic disease setting. In addition, patients must have progressed or be intolerant of the most recent line of systemic therapy. Patients with HR⁺ disease must have received a cyclin-dependent kinase (CDK4/6) inhibitor in the metastatic setting. Patients with previous exposure to fulvestrant were eligible. All patients were required to have evidence of measurable disease as per Response Evaluation Criteria in Solid Tumors (RECIST) v1.1 on baseline imaging. Patients with an Eastern Cooperative Oncology Group (ECOG) performance status of 0 or 1 and adequate baseline cardiac, hepatic, renal and hematologic function were eligible. Patients previously treated with any systemic anticancer therapy, radiation therapy, major surgery or experimental agent within 3 weeks of the first dose of study treatment were excluded. In addition, patients must not have received previous *HER2*-directed therapy. Study enrollment criteria did not include any requirements regarding sex and/or gender. Patient sex was based on self-identification at baseline. Full inclusion and exclusion criteria are available in the protocol, which has been previously published³⁷.

Eligible patients had *HER2* mutations identified by local or central NGS. Local testing for identification of *HER2* mutations could be via archival or fresh tumor tissue or blood. Central testing was through blood-based NGS (Guardant360 CDx Test, Guardant). *HER2* mutations eligible for inclusion in the cohort were prespecified. Patients with *HER2* mutations that were not prespecified were also potentially eligible, if supported by scientific literature.

Procedures

Patients in the *HER2*-mut MBC cohort were treated with tucatinib 300 mg orally twice daily and trastuzumab 8 mg kg⁻¹ intravenously then 6 mg kg⁻¹ every 3 weeks in a 21 day cycle. Patients with HR⁺ disease also received fulvestrant 500 mg intramuscularly once every 4 weeks starting from cycle 1 day 1, and once on cycle 1 day 15. Disease response to the study treatment and the occurrence of disease progression were determined according to RECIST v1.1, as assessed by the investigator. Disease assessments were performed at baseline, every 6 weeks for the first 24 weeks and then every 12 weeks until the occurrence of documented disease progression per RECIST v1.1, death, withdrawal of consent, loss to follow-up or study closure.

Safety was assessed by the incidence of TEAEs, graded according to the National Cancer Institute Common Terminology Criteria for Adverse Events v5.0, recording of concomitant medications, physical examination findings, vital signs, laboratory tests, pregnancy testing and cardiac function. Cardiac ejection fraction was assessed via echocardiogram or a multigated acquisition scan at screening and every 12 weeks thereafter.

For the exploratory biomarker assessments, central *HER2* testing was performed in a laboratory accredited by Clinical Laboratory Improvement Amendments. Patients' blood samples were collected

during prescreening or on day 1 of cycle 1 for central testing using a 74-gene blood-based NGS assay (Guardant360 CDx Test, Guardant). Tumor tissue biopsies, archival or fresh (if available), were collected during screening. Central *HER2* testing of tumor tissue was performed using IHC (PATHWAY *HER2*, Roche), FISH (*HER2* IQFISH pharmDx Assay, Agilent) and 505-gene tissue-based NGS assay (Elio Tissue Complete, PGDx). IHC and FISH results were evaluated using the American Society of Clinical Oncology and College of American Pathologists Breast Scoring criteria⁴².

Assessments

The primary endpoint was cORR, defined as the proportion of patients with confirmed complete response or partial response according to RECIST v1.1, per investigator assessment. Secondary endpoints included disease control rate, DOR, PFS by investigator assessment, OS and safety. The exploratory endpoints included biomarker assessments as described in the section 'Exploratory biomarker data assessment and analyses'.

Statistical analysis

The *HER2*-mut MBC cohort aimed to enroll 30 response-evaluable patients, a number calculated per the 90% exact CI given a range of expected cORR of 10–30%. One more patient was accrued than planned because one patient was initially deemed not response evaluable but was later found to be response evaluable. An interim analysis was not planned for the *HER2*-mut MBC cohort. All enrolled patients received at least one dose of tucatinib and trastuzumab and were included in the evaluation for efficacy and safety. Two-sided 90% exact CIs for response rates were calculated using the Clopper–Pearson method. Median PFS and OS were estimated using the Kaplan–Meier method; the associated 90% CI was calculated on the basis of the complementary log–log transformation. Safety and concordance of local versus central *HER2* testing results were assessed by descriptive statistics. All analyses except the biomarker analyses were performed with SAS, version 9.4 (SAS Institute). Biomarker analyses were performed with R, version 4.0.2 (R Core Team and the R Foundation for Statistical Computing). Because all patients in the study cohort were female, no sex- or gender-based analyses have been performed.

Exploratory biomarker data assessment and analyses

Central testing methods. Blood-based NGS analyses of amplifications and sequence alterations (mutations and indels) were from the standard analysis pipelines used by Guardant (Guardant360 CDx Test). Similarly, tissue-based NGS determination of amplifications and sequence alterations were from internal pipelines used by PGDx (Elio Tissue Complete). A further analysis step was the removal of synonymous mutations, except those indicated as potentially affecting a splice region. For blood-based NGS testing, we removed *TP53* from the analyses as this gene's mutations may be from clonal hematopoiesis of indeterminate potential as opposed to reflecting tumor state.

Assignment of domains for *HER2* mutation. Consistent with previous approaches, the following definitions were used to assign mutations by amino acid residue number to protein in *HER2*: 52–643 for the extracellular domain and 694–883 for the tyrosine kinase domain³⁹.

***HER2* assay agreement.** In the analyses of local versus central testing, local refers to any test to determine *HER2* mutations, or overexpression or amplification status, performed at the local site. Three patients used results from central blood-based NGS rather than local testing to establish *HER2* mutational and amplification status for eligibility. They were excluded from these analyses, and their local results are labeled as not determined when data are shown.

HER2 status, defined as the final *HER2* overexpression or amplification status, could be either positive, negative or not determined.

For central IHC and FISH testing, HER2 status was determined as follows: HER2-negative status (IHC 0, IHC 1 or IHC 2+ with FISH negative), HER2-positive status (IHC 2+ with FISH-positive or IHC 3+ with any FISH result) and HER2 status ‘not determined’ (IHC 2+ without evaluable FISH results).

To determine the HER2 overexpression or amplification status across assays, pairwise comparisons were performed using data from patients who had results (that is, positive or negative) from both assays. The results are reported as percentage agreement, with two-sided 90% exact CIs calculated using the Clopper–Pearson method⁴⁰.

HER2 mutation detection agreement was calculated using the same methods described above for the HER2 overexpression or amplification status. These comparisons were based on simple HER2 ‘mutation detected’ versus HER2 ‘mutation not detected’.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

Following requests and subject to review, Pfizer will provide the data that support the findings of this study. Subject to certain criteria, conditions and exceptions, Pfizer may also provide access to the related individual de-identified participant data. See <https://www.pfizer.com/science/clinical-trials/trial-data-and-results> for more information.

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Author contributions

A.F.C.O., T.B.-S., B.J.M., Y.N., D.M.O., M.R., E.Y.Y., J.R., S. Tan, M.B., T.E.S. and P.R.P. contributed to the study conception or design. All authors contributed to the acquisition, analysis or interpretation of the data and the drafting or substantial revision of the paper.

Competing interests

A.F.C.O. reports research funding from Pfizer and Roche; honoraria for presentations from AstraZeneca, Esai, Gilead, Lilly, Pfizer, Roche and Seagen; advisory board fees from AstraZeneca, Pfizer, Roche and Seagen; and conference support from AstraZeneca, Lilly, Novartis and Roche. G.C. reports consulting or advisory roles with AstraZeneca, Celucity, Daiichi Sankyo, Ellipsis, Exact Sciences, Gilead, Lilly, MBS, Menarini, Merck, Pfizer, Roche and Veracyte; institutional research funding from Astellas, AstraZeneca, Blueprint Medicine, BMS, Daiichi Sankyo, Kymab, Merck, Novartis, Philogen,

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Additional information

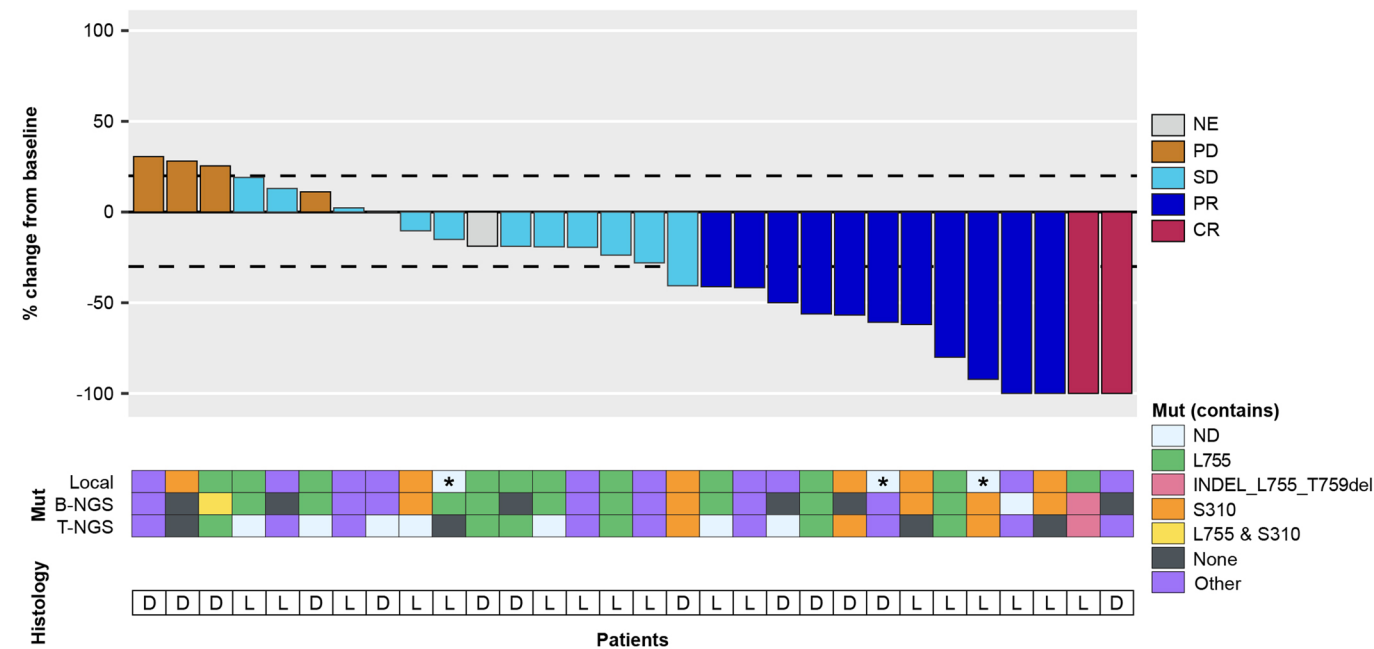
Extended data is available for this paper at <https://doi.org/10.1038/s41591-024-03462-0>.

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Correspondence and requests for materials should be addressed to Paula R. Pohlmann.

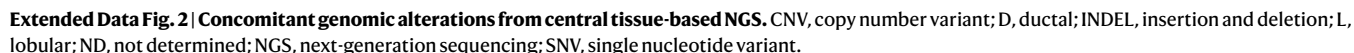
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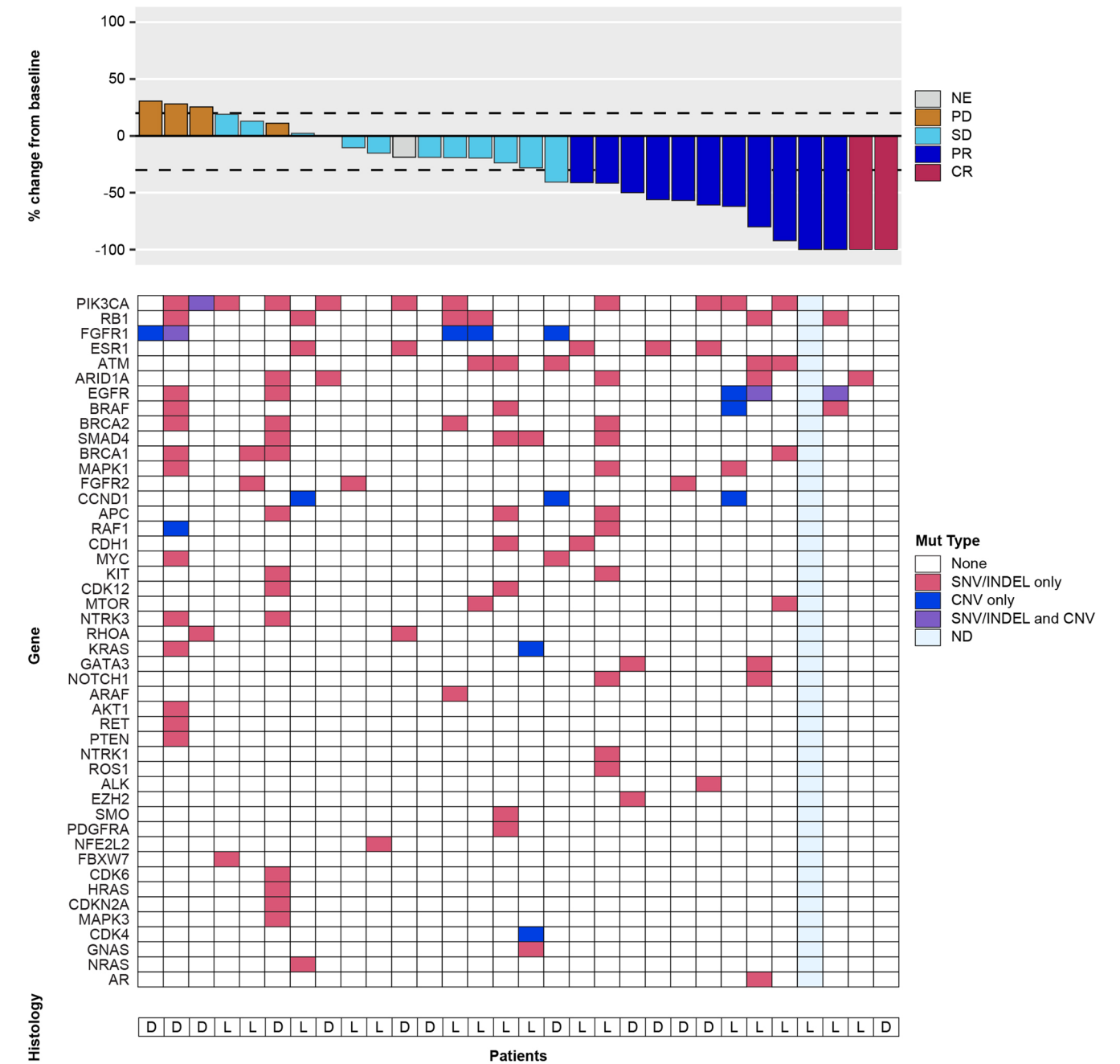
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Extended Data Fig. 1 | Response versus presence of L755 and/or S310F mutations. B-NGS, blood-based NGS testing by central lab assay; CR, complete response; D, ductal; L, lobular; Local, local testing results; Mut, Mutation; ND, not determined; NE, non-evaluable; NGS, next-generation sequencing; PD, progressive disease; PR, partial response; SD, stable disease; T-NGS, tissue-based

NGS testing by central lab assay. Only subjects who had baseline and post-baseline target lesion measurements are included (n = 30). * indicates patients that were excluded from any analysis of local results because their eligibility (HER2-negative status and *HER2* mutations) was determined by central blood-based NGS.





Extended Data Table 1 | Summary of safety events in the *HER2*-mut MBC cohort

TEAE	Total (N = 31) n (%)
Any TEAE	31 (100)
Treatment-related	30 (97)
Grade ≥ 3 TEAE	15 (48)
Tucatinib-related	9 (29)
Trastuzumab-related	1 (3)
Fulvestrant-related	0
Any serious TEAE	8 (26)
Tucatinib-related	3 (10)
Trastuzumab-related	0
Fulvestrant-related	0
TEAEs leading to discontinuation	2 (6)
Tucatinib	2 (6)
Trastuzumab	0
Fulvestrant	0
TEAEs leading to death	0

HER2, human epidermal growth factor receptor 2; TEAE, treatment emergent adverse event.

Extended Data Table 2 | Listing of responses, histology, and *HER2* mutations by local and central testing

Response	Histology	Local Testing	Central Blood-Based NGS	Central Tissue-Based NGS
CR	Ductal	G778 mutation	no mutation found	INDEL_V777_G778insGSP
CR	Lobular	L755P/S mutation	INDEL_L755_T759del	INDEL_L755_T759del
PR	Ductal	Central blood-based NGS used for enrollment	INDEL_P780_Y781insGSP, SNV_V773M	INDEL_V777_G778insGSP
PR	Lobular	Central blood-based NGS used for enrollment	SNV_S310F	SNV_S310F
PR	Lobular	L755P/S mutation	SNV_L755S	no valid results
PR	Lobular	L755P/S mutation	SNV_L755S	SNV_L755S
PR	Ductal	L755P/S mutation	SNV_L755S	SNV_L755S
PR	Ductal	L869R mutation	no mutation found	no valid results
PR	Ductal	S310F/Y mutation	no mutation found	SNV_S310Y
PR	Lobular	S310F/Y mutation	SNV_D211H, SNV_E698V, SNV_S310Y, SNV_T733I, SNV_V697L	no mutation found
PR	Lobular	S310F/Y mutation	SNV_S310F	no mutation found
PR	Lobular	T733I	no valid results	SNV_T733I
PR	Lobular	T862S	SNV_E717K, SNV_T733I, SNV_T862S	SNV_T862S
SD	Lobular	Central blood-based NGS used for enrollment	SNV_L755S	no mutation found
SD	Lobular	D769Y, V777L/M mutation	SNV_D769Y, SNV_V777L	SNV_D769Y, SNV_V777L
SD	Ductal	L755P/S mutation	no mutation found	SNV_L755S
SD	Lobular	L755P/S mutation	SNV_L755S	no valid results
SD	Lobular	L755P/S mutation	SNV_L755S	no valid results
SD	Lobular	S310F/Y mutation	SNV_S310F	no valid results
SD	Ductal	S310F/Y mutation	SNV_S310F	SNV_S310F
SD	Lobular	T733I	SNV_T733I	SNV_T733I
SD	Lobular	V777L mutation, L755P/S mutation	SNV_L755S, SNV_V777L, SNV_V777M	SNV_L755S, SNV_V777L
SD	Lobular	V777L/M mutation	no mutation found	SNV_G727A, SNV_V777L
SD	Ductal	V777L/M mutation	SNV_V777L	no valid results
SD	Lobular	V777L/M mutation	SNV_V777L	SNV_V777L
PD	Ductal	A775, Y772 mutation	INDEL_A775_G776insYVMA	INDEL_E770_A771insAYVM
PD	Ductal	L755P/S mutation	SNV_E717D, SNV_E992K, SNV_L726I, SNV_L755S, SNV_T733I, SNV_T733R	no valid results
PD	Ductal	L755P/S mutation	SNV_L755S, SNV_S310Y	SNV_L755S
PD	Ductal	S310F/Y mutation	no mutation found	no mutation found
NE	Ductal	L755P/S mutation	SNV_L755S	SNV_L755S
NV	Lobular	L755P/S mutation	no valid results	SNV_L755S

CR, complete response; del, deletion; *HER2*, human epidermal growth factor receptor 2; INDEL, insertion and deletion; ins, insertion; ND, not determined; NE, non-evaluable (post-baseline assessment is not evaluable); NGS, next-generation sequencing; NV, not available (patient has no post-baseline response assessment); PD, progressive disease; PR, partial response; SD, stable disease; SNV, single nucleotide variant.

Each row includes the best response from one patient and mutations detected by local and central testing.

Extended Data Table 3 | *HER2* altered domains: prevalence and response rates

Assay	Prevalence % (n/N)		Response Rate % (n/N)	
	Mutations Limited to TyrKin Domain	Mutations Limited to Extracell Domain	Mutations Limited to TyrKin Domain	Mutations Limited to Extracell Domain
Local	78.6% (22/28) 90% CI: 62.0-90.2%	21.4% (6/28) 90% CI: 9.8-38.0%	36.4% (8/22) 90% CI: 19.6-56.1%	50.0% (3/6) 90% CI: 15.3-84.7%
B-NGS	69.6% (16/23) 90% CI: 50.4-84.8%	17.4% (4/23) 90% CI: 6.2-35.5%	37.5% (6/16) 90% CI: 17.8-60.9%	50.0% (2/4) 90% CI: 9.8-90.2%
T-NGS	85.0% (17/20) 90% CI: 65.6-95.8%	15.0% (3/20) 90% CI: 4.2-34.4%	41.2% (7/17) 90% CI: 21.2-63.6%	66.7% (2/3) 90% CI: 13.5-98.3%

B-NGS, blood-based NGS testing by central lab assay; Extracell, extracellular domain of *HER2*; *HER2*, human epidermal growth factor receptor 2; Local, local testing results; NGS, next-generation sequencing; T-NGS, tissue-based NGS testing by central lab assay; TyrKin; tyrosine kinase domain of *HER2*.

Extended Data Table 4 | *HER2* L755 and S310 mutations: prevalence and response rates

Assay	Prevalence % (n/N)		Response Rate % (n/N)	
	L755 Mutation(s)	S310 Mutations(s)	L755 Mutation(s)	S310 Mutations(s)
Local	42.9% (12/28) 90% CI: 26.9-60.0%	21.4% (6/28) 90% CI: 9.8-38.0%	33.3% (4/12) 90% CI: 12.3-60.9%	50.0% (3/6) 90% CI: 15.3-84.7%
B-NGS	47.8% (11/23) 90% CI: 29.6-66.5%	26.1% (6/23) 90% CI: 12.0-45.1%	36.4% (4/11) 90% CI: 13.5-65.0%	50.0% (3/6) 90% CI: 15.3-84.7%
T-NGS	40.0% (8/20) 90% CI: 21.7-60.6%	15.0% (3/20) 90% CI: 4.2-34.4%	37.5% (3/8) 90% CI: 11.1-71.1%	66.7% (2/3) 90% CI: 13.5-98.3%

B-NGS, blood-based NGS testing by central lab assay; *HER2*, human epidermal growth factor receptor 2; Local, local testing results; NGS, next-generation sequencing; T-NGS, tissue-based NGS testing by central lab assay.

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☐	☒ A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
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Software and code

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Data collection	Clinical data: Medidata Classic Rave 2024.1.1.(electronic data collection), Suvoda (clinical supply management); Biomarker analyses: Guardant360® CDx Test [Guardant, Redwood City, CA]), IHC (PATHWAY HER-2 [Roche, Tucson, AZ]), FISH (HER2 IQFISH pharmDx Assay [Agilent, Singapore, Singapore]), tissue-based NGS assay (Elio Tissue Complete [PGDx, Baltimore, MD])
Data analysis	All analyses except the biomarker analyses were performed with SAS, version 9.4 (SAS Institute, Cary, NC). Biomarker analyses were performed with R, version 4.0.2 (R Core Team and the R Foundation for Statistical Computing, Vienna, Austria).

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Reporting on sex and gender	Sex data were collected and 100% of the patients participating in the HER2-mut BC cohort were female (31/31).
Reporting on race, ethnicity, or other socially relevant groupings	Ethnicity data were collected as part of baseline patient characteristics with the majority of patients not Hispanic, Latino/a, or of Spanish origin and 48% White (15/31), 35% Asian (11/31), 3% Black (1/31) and 13% (4/31) not reportable.
Population characteristics	No covariates are discussed in the protocol or SAP. Given the small sample size in each cohort, no subgroup analyses were performed.
Recruitment	All 31 subjects in the cohort were prospectively identified in alignment with the inclusion and exclusion criteria by PIs, Sub-Is, and participating clinicians at 20 sites in 7 countries. All subjects signed informed consent prior to study participation. Subjects were not paid for their participation in the trial. Protocol compliance may impact the results. Protocol deviations are summarized in the manuscript.
Ethics oversight	The protocol was approved by institutional review boards and ethics committees according to the practice at each participating study site (as listed in the Supplementary Information).

Note that full information on the approval of the study protocol must also be provided in the manuscript.

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Life sciences study design

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Sample size	The HER2-mut MBC cohort aimed to enroll 30 response-evaluable patients, a number calculated per the 90% exact CI given a range of expected cORR of 10% to 30%. One more patient was accrued than planned because one patient was initially deemed not response-evaluable but was later found to be response-evaluable.
Data exclusions	Patients enrolled using central blood-based NGS rather than local testing to establish HER2 mutational and amplification status were excluded from HER2 assay agreement.
Replication	Not applicable. Analyses were performed during clinical investigations and as such were performed once.
Randomization	Open-label study. No randomization was utilized.
Blinding	Open-label study. Blinding was not performed.

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n/a	Involved in the study
☒	☐ ChIP-seq
☒	☐ Flow cytometry
☒	☐ MRI-based neuroimaging

Clinical data

Policy information about [clinical studies](#)

All manuscripts should comply with the ICMJE [guidelines for publication of clinical research](#) and a completed [CONSORT checklist](#) must be included with all submissions.

Clinical trial registration	NCT04579380
Study protocol	The SGNTUC-019 protocol has been previously published (DOI: 10.1200/JCO.23.00606) and is available at this link: https://ascopubs.org/action/downloadSupplement?doi=10.1200%2FJCO.23.00606&file=protocol_JCO.23.00606.pdf
Data collection	Data were collected from September 2021 to December 2022 at 67 academic research or cancer treatment centers in North America, Europe and Asia.
Outcomes	The primary endpoint was confirmed objective response rate, defined as the proportion of patients with confirmed complete response or partial response according to RECIST v1.1, per investigator assessment. Secondary endpoints included disease control rate (confirmed complete or partial response, or stable disease per investigator assessment), duration of response (confirmed complete or partial response per investigator assessment) progression-free survival by investigator assessment, and overall survival. Safety endpoints included type, incidence, severity, seriousness and relatedness of treatment-related adverse events, and were graded according to the NCI-CTCAE 5.0. Treatment duration and discontinuations were also assessed. Exploratory endpoints included concordance of HER2 alterations as detected by different testing methodologies and identification of tumor-specific alterations associated with resistance to tucatinib.

Plants

Seed stocks	<i>Report on the source of all seed stocks or other plant material used. If applicable, state the seed stock centre and catalogue number. If plant specimens were collected from the field, describe the collection location, date and sampling procedures.</i>
Novel plant genotypes	<i>Describe the methods by which all novel plant genotypes were produced. This includes those generated by transgenic approaches, gene editing, chemical/radiation-based mutagenesis and hybridization. For transgenic lines, describe the transformation method, the number of independent lines analyzed and the generation upon which experiments were performed. For gene-edited lines, describe the editor used, the endogenous sequence targeted for editing, the targeting guide RNA sequence (if applicable) and how the editor was applied.</i>
Authentication	<i>Describe any authentication procedures for each seed stock used or novel genotype generated. Describe any experiments used to assess the effect of a mutation and, where applicable, how potential secondary effects (e.g. second site T-DNA insertions, mosaicism, off-target gene editing) were examined.</i>