

A Phase 3 study of gedatolisib plus fulvestrant with and without palbociclib in patients with HR+/HER2- advanced breast cancer previously treated with a CDK4/6 inhibitor plus a non-steroidal aromatase inhibitor (VIKTORIA-1)

Sara Hurvitz¹, Fabrice Andre², Massimo Cristofanilli³, Giuseppe Curigliano^{4,5}, Antonio Giordano⁶, Hyo Han⁷, Miguel Martin⁸, Barbara Pistilli², Hope Rugo⁹, Robert Wesolowski¹⁰, Samuel Suzuki¹¹, Sarah C. Mutka¹¹, Igor Gorbachevsky¹¹, Sibylle Loibl¹²

¹David Geffen School of Medicine, University of California Los Angeles, Los Angeles, CA; ²Gustave Roussy Cancer Center, Villejuif, France; ³Weill Department of Medicine, Cornell University, New York, NY; ⁴Istituto Europeo di Oncologia, IRCCS, Milano, Italy; ⁵University of Milano, Milan, Italy; ⁶Dana Farber Cancer Institute, Harvard University, Boston, MA; ⁷Moffitt Cancer Center Department of Breast Oncology, Florida; ⁸Hospital General Universitario Gregorio Marañón, Universidad Complutense, Madrid, Spain; ⁹University of California San Francisco Comprehensive Cancer Center, San Francisco, CA; ¹⁰James Cancer Hospital and the Ohio State University Comprehensive Cancer Center, Columbus, OH; ¹¹Celcuity, Minneapolis, MN; ¹²German Breast Group (GBG), Neu-Isenburg, Germany

celcuity

EXPANDING TREATMENT OPTIONS

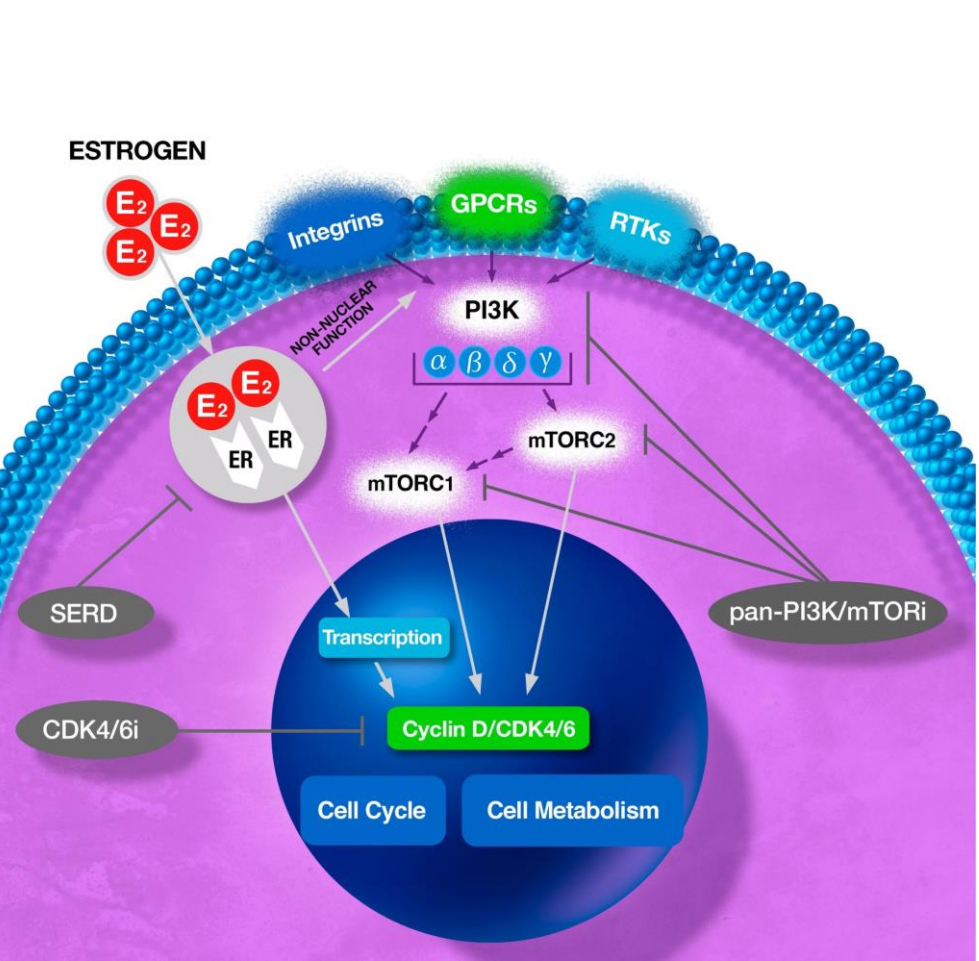
Abstract CT063

BACKGROUND

Gedatolisib is a Potential First-in-Class PI3K/mTOR Inhibitor

Mechanism of Action	• First small molecule dual inhibitor of the PI3K/mTOR pathway administered intravenously • Inhibits all isoforms of PI3K and mTOR at low or sub-nanomolar concentrations
Efficacy	• Compelling efficacy relative to 1st & 2nd line SOC with HR+/HER2- ABC with gedatolisib + ET + CDK4/6i – Phase 1b trial (N=103) reported 63% ORR in 95 response evaluable patients across four expansion arms (this includes 7 unconfirmed PR) – Median PFS 42.3 months in 1L arm; 12.9 months in 2L arm with Phase 3 dosing schedule – NCT02684032
Tolerability	• Addition of gedatolisib to palbociclib and fulvestrant in the Phase 1b trial was shown to be well-tolerated with manageable TEAEs • Few patients discontinued treatment due to an AE, with only one (4%) discontinuation in cohort with Phase 3 dosing • Low incidence of the Grade 3/4 adverse events that are generally associated with the PI3K/mTOR class of inhibitors: hyperglycemia (7%), diarrhea (6%), AST/ALT increase (4%), and no Grade 3/4 colitis

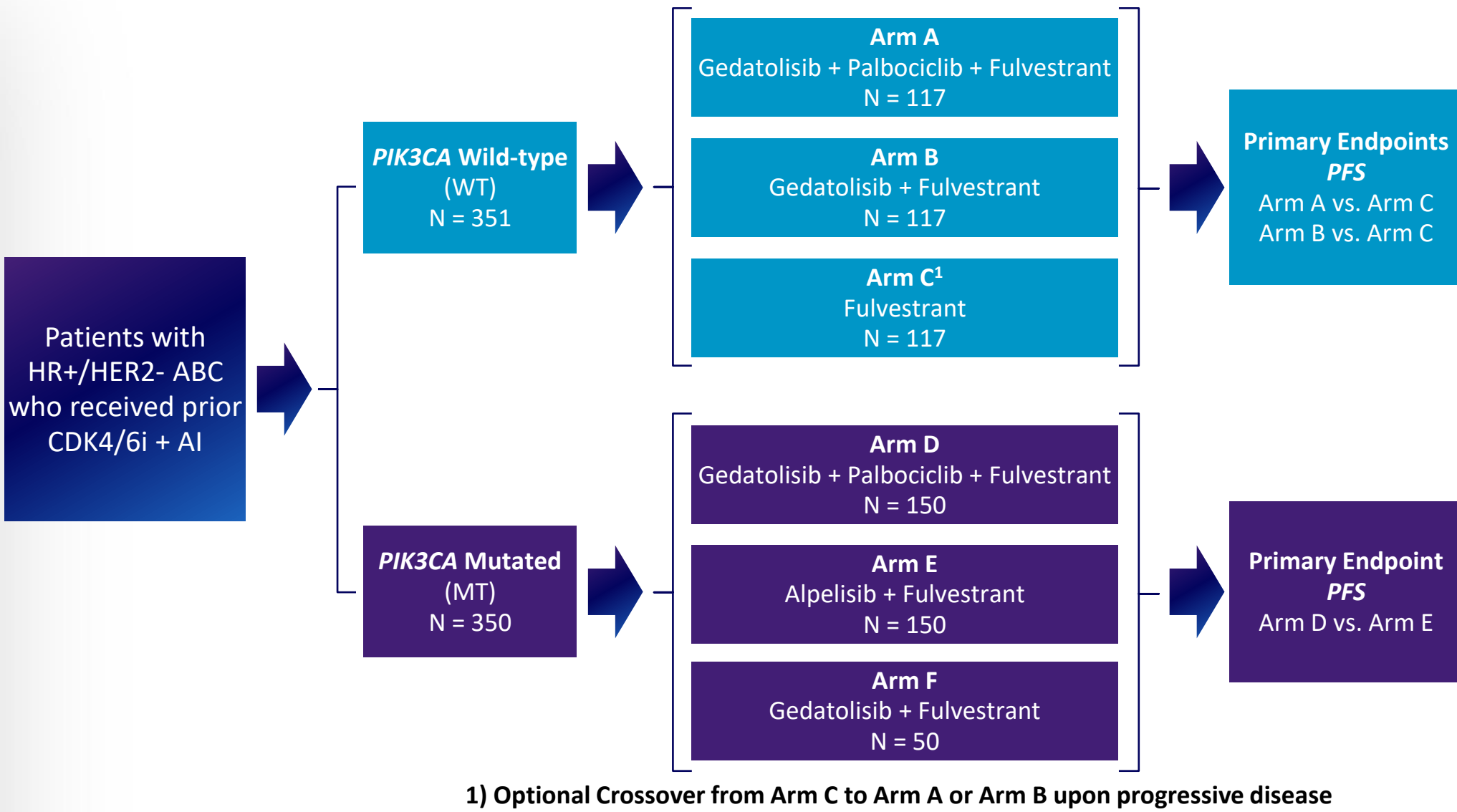
TREATMENT STRATEGY



Simultaneously blockading PI3K/mTOR, CDK4/6, and ER signaling pathways disrupts complex cooperation between these pathways to inhibit tumor growth

- The upregulation of the PI3K/AKT/mTOR pathway promotes hormone dependent and independent estrogen receptor (ER) transcriptional activity.
- This contributes to endocrine resistance, leading to tumor cell growth, survival, motility, and metabolism.
- Available evidence indicates that resistance to CDK4/6 inhibition is a transient adaptive mechanism, most likely involving the PI3K/mTOR pathway.
- These data indicate that continuing CDK4/6 inhibitor treatment in combination with a PI3K/mTOR inhibitor in patients who progressed on their prior CDK4/6 inhibitor, would both block the reactivated CDK4/6 pathway and prevent adaptive activation of the PI3K/mTOR pathway.
- This suggests that patients whose disease progressed on a CDK4/6 inhibitor may benefit from continued treatment with a CDK4/6 inhibitor when it is combined with a PI3K/mTOR inhibitor as their next line of therapy.

VIKTORIA-1 is a Phase 3, open-label, randomized, two-part clinical trial to evaluate the efficacy and safety of gedatolisib in combination with fulvestrant with or without palbociclib. Two studies based on *PIK3CA* mutation status are included in the trial. *PIK3CA* mutation status will be assessed centrally using an FDA approved *PIK3CA* test. According to confirmed *PIK3CA* mutation status, subjects will be manually assigned to Study 1 (*PIK3CA* WT) or Study 2 (*PIK3CA* MT). The two studies will be randomized separately



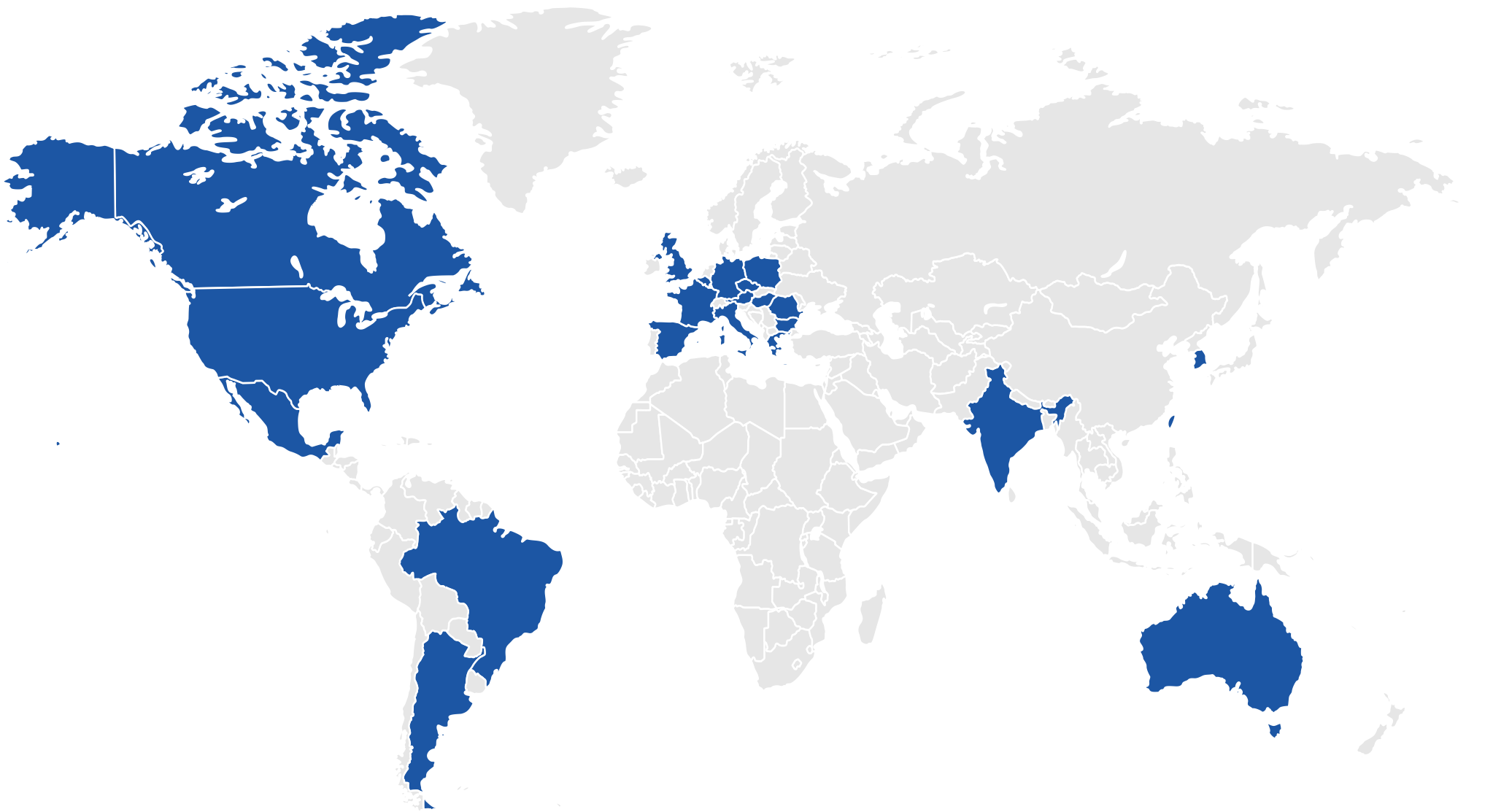
OBJECTIVES AND ENDPOINTS

VIKTORIA-1 will evaluate the efficacy and safety of gedatolisib and fulvestrant with or without palbociclib in patients with HR+/HER2- ABC previously treated with any CDK4/6i in combination with non-steroidal AI therapy.

Study 1 (<i>PIK3CA</i> WT)	Study 2 (<i>PIK3CA</i> MT)
Primary Objectives	
• Compare efficacy, as measured by progression-free survival, of gedatolisib in combination with palbociclib and fulvestrant (Arm A) to fulvestrant (Arm C) • Compare efficacy as measured by PFS, of gedatolisib in combination with fulvestrant (Arm B) to Arm C	• Compare the efficacy, as measured by PFS, of gedatolisib in combination with palbociclib and fulvestrant (Arm D) to alpelisib with fulvestrant (Arm E)
Key Secondary Objectives	
• Compare efficacy as measured by PFS of Arm A to Arm B • Compare the efficacy as measured by OS of Arm A to Arm C, Arm B to Arm C, and Arm A to Arm B • Compare safety & tolerability between treatment arms	• Compare the efficacy, as measured by PFS, of Arm D to Arm F (gedatolisib with fulvestrant) • Compare the efficacy, as measured by OS, of Arm D to Arm E • Compare the efficacy, as measured by OS, of Arm D to Arm F • Compare safety & tolerability between treatment arms
Additional Secondary Objectives	
• Evaluate contributory treatment effect of gedatolisib, palbociclib, & combined treatment effect of gedatolisib and palbociclib in the stratified Cox proportional hazard model • Estimate and compare PFS and OS based on HER2 status (HER2-low, defined as an IHC score of 1+ or IHC 2+ with a negative ISH score, and HER-negative status, defined as an IHC score of 0) • Compare efficacy, as measured by ORR, DOR, TTR, & CBR of Arm A to Arm C, Arm B to Arm C, & Arm A to Arm B. • Compare change in health status/QOL of Arm A to Arm C, Arm B to Arm C, and Arm A to Arm B. • PK of gedatolisib	• Compare the efficacy, as measured by PFS, of Arm E to Arm F • Estimate and compare PFS and OS based on HER2 status (HER2-low, defined as an IHC score of 1+ or IHC 2+ with a negative ISH score, and HER-negative status, defined as an IHC score of 0) • Compare the efficacy, as measured by ORR, DOR, TTR, and CBR, of Arm D to Arm E • Compare change in health status/QOL of Arm D to Arm E • PK of gedatolisib

Trial Status

- VIKTORIA-1 is enrolling
- Approximately 701 subjects are expected to be enrolled at sites in the Americas, Europe, and Asia-Pacific
- The primary completion date is estimated to occur in the second half of 2024



ELIGIBILITY CRITERIA

Key Inclusion Criteria	Key Exclusion Criteria
• Adults ≥ 18 years of age • Confirmed diagnosis of ER+ and/or PR+ as per ASCO-CAP 2020 guidelines • Documented HER2- as per ASCO-CAP 2018 guidelines • Adequate archival or fresh tumor tissue specimen for analysis of <i>PIK3CA</i> mutational status by central lab using an FDA approved test • Radiologically evaluable disease according to RECIST v1.1 • Progressed during or after CDK4/6i combination treatment with non-steroidal AI • Adequate bone marrow, hepatic, renal and coagulation function as defined by acceptable laboratory parameters	• Prior treatment with PI3K, Akt, or mTOR inhibitors • Prior chemotherapy for advanced disease • More than 2 prior lines of endocrine therapy treatment • Bone only disease with no soft tissue component • Type 1 diabetes or uncontrolled type 2 diabetes • History of drug induced pneumonitis or interstitial lung disease • Pregnant or breast-feeding women

Acknowledgements

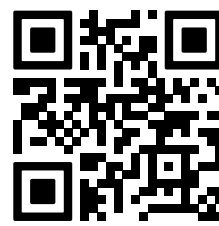
The authors would like to acknowledge the patients and their families/ caregivers, investigators and their support staff for participating in this trial and Celcuity Inc. for sponsoring the study.

These data were first presented at the San Antonio Breast Cancer Symposium® December 6–10, 2022

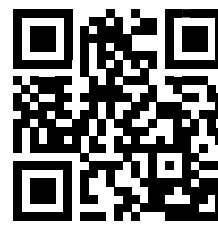
Clinical Trial Registry Number

ClinicalTrials.gov Identifier: NCT05501886
EU CT 2022-502145-10-00

Copies of this poster obtained through Quick Response (QR) Code are for personal use only and may not be reproduced without permission from AACR® and the author of this poster.



Poster



Study Site