

EAQ221CD/CONCURXP

For Patients with HR+ HER2- Metastatic Breast Cancer

EAQ221CD Available Through ECOG-ACRIN Cancer Research Group

Improving Medication Adherence in Metastatic Breast Cancer Using a CONnected CUstomized Treatment Platform (CONCURXP)

Patient Population

See Section 3 for Complete Eligibility Details

Non-Patient Participants: Subset of oncology healthcare providers (n=~20) from a minimum of 10 participating NCORP sub-affiliates

- Must have taken care of at least one patient randomized to Arm B who had less than 85% adherence rate at 12 months as measured by the WiseBag
- Must speak English
- Must be employed at an NCORP site for at least 6 months

Patient Eligibility- Step 0 (Screening Registration):

- Must be ≥ 18 years of age; must not have an ECOG PS
 ≥ 3 (or must not be deemed medically unable to participate)
- Must be fluent in written and spoken English <u>or Spanish</u>
- Must be present with new or established pathologically proven HR+ HER2- metastatic breast cancer
- Must have initiated any of the CKD4/6 inhibitors within 30 days prior to consenting to Step 0 or have received a prescription order with stated intent to initiate within 30 days following Step 0 consent
 - Must not have been previously treated with Palbociclib/Ibrance, Ribociclib/Kisqali, Abemaciclib/ Verzenio
- Must not be enrolled in 1) a therapeutic clinical trial that monitors CDK4/6 inhibitors, or 2) a symptom science clinical trial that monitors/intervenes on symptoms related to CDK4/6i, or 3) other trials offering financial assistance
- Must have a personal mobile phone in which they are able and willing to send and receive text messages; must have an email address

Patient Eligibility- Step I (Randomization):

 Must have completed Baseline Survey and initiated their CDK4/6 inhibitors within 30 days of OPEN screening registration (Step 0)

Methodology Plan

See Section 5 for Methodology Details

- After consent/OPEN registration, NCORP CRA will:
 - $\lozenge \qquad \text{Ask the patient to complete the Patient Contact} \\ \text{Sheet} \\$
 - Encourage the patient to activate their EASEE-PRO account and complete the baseline survey as soon as possible (within 30 days)
 - Enter patient contact info, incentive choice and method of delivery, and CRA contact data into UCI REDCap Project Enrollment Log
 - ♦ Complete subject randomization (after baseline survey); send info via REDCap to UCI/Emory

Baseline study procedures (both arms):

- Prescription schedule for CDK4/6 inhibitors and internal financial counseling documented in RED-Cap (by NCORP CRA)
- WiseBag activation (by UCI Coordinator)
- ♦ Study package distribution (by UCI Coordinator)
- Creation of patient's schedule on Wisepill portal (by UCI Coordinator)

Enhanced Usual Care (EUC)- Arm A:

- Use WiseBag exclusively with prescribed CDK4/6i for ~ 12 months
- Participants will receive a link every 4 weeks with educational materials (any patient-reported changes in their CDK4/6i prescription/schedule will be confirmed with the NCORP CRA)

• CONCURXP Intervention- Arm B:

- Personalized and interactive bidirectional text messaging (smart reminders and missed dose questions specifically per drug); over-adherence confirmation/addressing; financial barrier addressing; patients able to view history of doses taken
- See protocol for follow-up procedures for both arms (i.e., surveys at 3, 6, and 12 months)

Patient Enrollment

All Sites: Oncology Patient Enrollment Network (OPEN) https://open.ctsu.org/open

Protocol Information

ECOG-ACRIN Operations-Boston: 857-504-2900, http://ecog-acrin.org (Member Login)

Please Enroll Your Eligible Patients!

Study Chair: Gelareh Sadigh, MD

Study Co-Chair: Ilana Graetz, PhD

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* Site staff refers to NCORP site staff/CRA



A. Identify Site Population Review patient lists, medical records, and/or HER reports to identify potentially eligible patients: ≥ 18 years old, having new or established HR+ HER2-metastatic breast cancer and a new prescription for CDK4/6i, who are not enrolled in any non-therapeutic triple offering finengial expirated. Screen Log (REDCap survey hosted by UCI) trials offering financial assistance Site Staff: Enter a monthly summary into REDCap survey: • Total number of eligible patients approached Total number of patients who refused to participate and reasons (optional) B. Screen/Approach Patients Site Staff: Approach all potentially eligible patients (identified in **A** above): **OPEN Registration (Step 0)** · Verify (screen for) study eligibility Site Staff: Register patients who sign consent into OPEN (Step 0) Recruit eligible subjects for participation Complete Registration to Study Site Staff: C. Consent Patients • Have the participant complete the Patient Contact sheet and fax it to the Brown OEAU (Tel: 855-404-3278 Fax: 401-863-9635) Site Staff: Encourage the patient to either activate their EASEE-PRO account Obtain written informed consent and complete the baseline survey as soon as possible by logging into to the pride portal at https://pride.stat.brown.edu/Participant-Login (or by administering the baseline survey on paper in clinic) or complete paper surveys in the clinic and return to NCORP site staff/CRA D. Baseline Procedures • Submit a REDCap form with consented patient information, choice Site Staff: of incentive and preferred method of receipt of incentive, and CRA Fax PC to OEAU contact Encourage participant to either complete EASEE-PRO setup ASAP and complete baseline survey online or complete paper survey in the clinic Email participant's provider a notification of their patient's enrollment in the study Submit a REDCap form with consented patient info Email provider of patient's enrollment Obtain Randomization & WiseBag Setup Patients: Site Staff: Complete baseline survey • Register patients who complete baseline survey into OPEN (Step 1) · Receive arm assignment UCI Staff: E. OPEN Randomization (Step 1) Mail patients' incentives • Mail WiseBag & Manual to participant Site Staff: Register patients to Step 1 in OPEN, upon completion of baseline survey · Contact participant and assist with setup • Create Wisepill schedule based on patient's treatment cycle and Sample size in Step 1 = 390 preferences Arm B: CONCURXP **Emory**: Monitor patient responses imbedded in intervention and refer to site staff when oncology referral or PAF referral needed Arm A: Enhanced Usual Care **Emory:** Text and email patients (monthly) educational materials and ensure patients' use of WiseBag **PAF:** Document services offered to patient in REDCap **Site:** Document results of oncology provider contact in REDCap F. Follow-Up (3, 6, and 12 months) EASEE-PRO/OEAU: Send patient email notifications and reminders that post-intervention survey is able to complete or mail paper survey. Provide patient outreach to recover delinquent surveys. Send UCI notification when survey has been completed. Document and send appropriate incentives to patient for survey completion Sites: Review medical records Update RAVE (vital status, treatment changes, pathology, etc.) Document unexpected ED/hospitalization Update REDCap (treatment, CDK4/6is changes, etc.) **NOTE: See protocol for the Non-**G. Qualitative Interview (12 months Post-Randomization) **Patient Participant Intervention** Select and interview 30 patients and 20 providers from Arm B (CONCURXP) Schema