

# RESECTABLE & BORDERLINE RESECTABLE

**Clinical Trial Name:** Adaptive Modification of Neoadjuvant Therapy Based on Clinical Response in Patients with Localized Pancreatic Cancer (PANC Trial)

**Study Design:** This is a single arm, Phase II clinical trial utilizing neoadjuvant therapy and surgery for patients with resectable and borderline resectable pancreatic adenocarcinoma which utilizes a total neoadjuvant therapy approach with adaptive modification of the chemotherapy regimen based on radiographic response (CT scan), biochemical response (CA19-9 decline), and performance status (as measured by a short physical performance battery).

**NCT#:** [NCT03322995](#)

**Key Inclusion:**

- ECOG performance status of < 2
- Histologically confirmed adenocarcinoma of the pancreas
- Clinical stage resectable or borderline resectable pancreatic adenocarcinoma
- Must be CA19-9 producer (pretreatment CA19-9 > 35 U/mL when total bilirubin ≤ 2 mg/dL)

**Study PI:**  
Dr. Kathleen Christians

**Clinical Research Coordinator:**  
Lauren Schmitz  
**Phone:** 414-805-5175

**Key Exclusion:**

- Received chemotherapy and/or radiation within 3 years prior to study enrollment
- History of prior malignancy except for adequately treated in situ cancer of the cervix or basal cell or squamous cell skin cancer or localized prostate cancer with a normal PSA within the last 3 years

**Clinical Trial Name:** Promoting CT Engagement for Pancreatic Cancer With App (PROCLAIM)

**Study Design:** To develop a culturally tailored informational mobile application and test whether it will increase participation among Black pancreatic cancer subjects in clinical trial discussions with their care team. This project aims to identify and address barriers to enrollment of Black subjects in pancreatic cancer clinical trials using a culturally informed mobile health application to promote participation.

**NCT#:** [NCT06252545](#)

**Key Inclusion:**

- Participants must meet the following inclusion criteria in order to participate in the mobile health application randomized clinical trial component of the study:
  - Informed consent obtained to participate in the study
  - 18 years or older
  - English speaking
  - Able and willing to participate in the trial
  - Newly diagnosed or progressive pancreatic cancer
  - Have access to a mobile device
  - Identify as Black or African American

**Study PI:**  
Dr. Ugwuji Maduekwe

**Clinical Research Coordinator:**  
Elizabeth Jeanes  
**Phone:** 414-955-6806

**Key Exclusion:**

- Inability to read and speak English
- Dementia altered mental status, or any psychiatric condition that would prohibit understanding or rendering of informed consent as determined by the study physician.
- Active participation in a therapeutic clinical trial

**Clinical Trial Name:** Molecular Profile-related Individualized Targeted Therapy in Resected Pancreatic Cancer with High-Risk of Cancer Recurrence (PROTECT-PANC)

**Study Design:** This is a prospective, open-label therapeutic interventional investigation designed to interrogate the efficacy and safety of individualized matched therapies in patients with pancreatic cancer at high risk of disease recurrence post-surgery.

**NCT#:** [NCT06228599](#)

**Key Inclusion:**

- Pathologically confirmed pancreatic cancer (excluding neuroendocrine histology).
- Pancreatic tumor is surgically removed and
  - Patient has received multimodal therapy (neoadjuvant, sandwich or adjuvant chemotherapy  $\pm$  radiation) or
  - Patient is ineligible for or refuses multimodal therapy
- Patient has one of the following:
  - Post-surgical cancer antigen (CA) 19-9 elevation ( $> 35$  U/mL at least 6 weeks post-surgical resection) in the setting of bilirubin  $< 2$  mg/dL (unless bilirubin elevation is consistent with Gilbert's syndrome) OR
  - High-risk pathological features, defined as positive surgical margin or lymph node involvement in cancer.
- Patient has no definitive measurable disease recurrence or metastatic disease at the time of first post-surgical imaging (in those with high-risk pathological features) or within four weeks of elevated CA 19-9 value as evidenced by appropriate imaging
- Laboratory values:
  - Absolute neutrophil count (ANC)  $\geq 1.0 \times 10^9/L$
  - Platelet count  $\geq 75,000/mm^3$  ( $125 \times 10^9/L$ )
  - Hemoglobin (Hgb)  $\geq 8$  g/dL
  - aspartate aminotransferase (AST) serum glutamic-oxaloacetic transaminase (SGOT), alanine transaminase (ALT) serum glutamate-pyruvate transaminase (SGPT)  $\leq 5 \times$  upper limit of normal range (ULN)
  - ECOG Performance Status  $< 3$
  - At the time of treatment, patient should be off other anti-tumor agents for at least five half-lives of the agent or three weeks from the last day of treatment, whichever is shorter
  - Patient must be presented at the Molecular Tumor Board (MTB) and agree to receive the MTB-recommended therapy
- Key Exclusion:
  - CA 19-9 non-producers, unless high-risk pathological features present.
  - Receiving concomitant investigational agent(s) for pancreatic ductal adenocarcinoma (PDAC)
  - Radiographic evidence of metastatic disease
  - Inability to ingest study drugs by mouth
  - Diarrheal bowel movements  $> 6$  per day postoperatively on maximal medical therapy
  - Patient has active, untreated, or uncontrolled bacterial, viral, or fungal infection(s) requiring systemic intravenous therapy
  - Patient has undergone or planned major surgery other than diagnostic surgery (i.e., surgery done to obtain a biopsy for diagnosis without removal of an organ) within four weeks prior to Day 1 of study therapy
  - Uncontrolled concurrent illness, including, but not limited to, unstable angina pectoris, uncontrolled and clinically significant cardiac arrhythmia, or psychiatric illness/social situations that would limit compliance with study requirements

**Study PI:**  
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