S2210 Kickoff Meeting

Join us in Seattle, or virtually, to learn about opening and enrolling to this new phase II prostate cancer trial.

FRIDAY, APRIL 5 • 8 – 9 AM, PT • BECKLER ROOM (302), LEVEL 3

S2210 is enrolling patients with high-risk localized prostate cancer who carry an inherited BRCA1 or BRCA2 pathogenic mutation.

Hypothesis: these patients will benefit from neoadjuvant carboplatin chemotherapy before surgery to remove their prostate cancer. Carboplatin is a DNA-damaging drug that is especially effective against other cancers with BRCA1 or BRCA2 mutations.

An estimated 5% of high-risk patients carry an inherited BRCA1 or BRCA2 mutation, although having a strong family history of cancer (especially breast cancers at an early age, ovarian, pancreas and metastatic prostate cancers) increases the likelihood of being a carrier.

Genetic testing for an inherited BRCA1 or BRCA2 mutation is now clinically indicated for patients with high-risk prostate cancer (NCCN guidelines). So timely workflows will help your site implement guideline-concordant genetic testing and counseling, and help you find candidates for S2210.

CHAIRS

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Daniel W. Lin, MD
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PATIENT ADVOCATE

Lee Moultrie

CATHY TANGEN, DrPH

BIOSTATISTICIANS

Sam Callis, MS

AGENDA

• Study background and rationale
• Review of protocol and trial workflow
• Patient advocate on genetic testing experience
• Review of first patient experience
• Genetic screening workflows, logistics, helpful resources

S2210: A Phase II Study of Neoadjuvant Carboplatin for Localized, High-Risk Prostate Cancer with Germline BRCA1/2 Mutations