Unlocking New Treatment Options for Patients with Metastatic Prostate Cancer

NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines®) Recommend Molecular Testing in Metastatic Prostate Cancer

1. Strongly advocates a metastatic biopsy for histologic and molecular evaluation. When this is not possible, plasma ctDNA (circular tumor DNA) assay is an option, preferably at time of biochemical PSA (prostate-specific antigen) or radiographic progression in order to maximize yield.

2. Homologous recombination (HRD) pathway genes: BRCA1, BRCA2, ATM, PALB2, FANCA, RAD51D, CHEK2 and CDK12 for genetic counseling, early use of platinum chemotherapy, olaparib, or rucaparib and/or eligibility for clinical trials (e.g., PARP inhibitors).

3. Microsatellite instability (MSI) or mismatch repair deficiency (dMMR): If MSI is used, testing using an NGS assay validated for prostate cancer is preferred. MSI-High or dMMR indicate eligibility for pembrolizumab in later lines of treatment for CRPC (castrate-resistant prostate cancer).†

Our portfolio of tests analyzes all guideline recommended genes and biomarkers for relevant alterations in patients with prostate cancer including: BRCA1, BRCA2, ATM, PALB2, FANCA, RAD51D, CHEK2, CDK12 and MSI*.

Advancing Therapy Options for Metastatic Prostate Cancer Patients

<table>
<thead>
<tr>
<th>BIOMARKER</th>
<th>FDA-APPROVED THERAPY</th>
<th>FOUNDATION MEDICINE COMPANION DIAGNOSTIC</th>
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</thead>
<tbody>
<tr>
<td>Homologous Recombination Repair</td>
<td>Lynparza® (olaparib)</td>
<td>FoundationOne®CDx</td>
</tr>
<tr>
<td>(HRR) gene (BRCA1, BRCA2, ATM,</td>
<td></td>
<td>FoundationOne®Liquid CDx (BRCA1/2, ATM only)</td>
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<tr>
<td>BARD1, BRIP1, CDK12, CHEK1, CHEK2, PALB2, RAD51B, RAD51C, RAD51D and RAD54L alterations</td>
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<tr>
<td>BRCA1, BRCA2</td>
<td>Rubraca® (rucaparib)</td>
<td>FoundationOne®Liquid CDx</td>
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<td>NTRK1/2/3 fusions</td>
<td>Rozytrek® (entrectinib)</td>
<td>FoundationOne®CDx</td>
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<td>TMB ≥ 10 mutations per megabase</td>
<td>Keytruda® (pembrolizumab)</td>
<td>FoundationOne®CDx</td>
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<tr>
<td>Microsatellite Instability-High</td>
<td>Keytruda® (pembrolizumab)</td>
<td>FoundationOne®CDx</td>
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As of March 2021

The Value of Comprehensive Genomic Profiling with Foundation Medicine:

FoundationOne Liquid CDx is the only blood-based comprehensive genomic profiling test FDA-approved to identify prostate cancer patients likely to respond to Lynparza® (olaparib) or Rubraca® (rucaparib).

By testing for all mutations, whether germline or somatic in origin, our portfolio of tests can identify more than twice as many men who may benefit from PARP inhibitor therapy than conventional germline-only testing.²

A blood-based comprehensive genomic profiling test is an important option to consider for mCRPC patients due to tissue sample issues, as demonstrated in the TRITON2 trial for rucaparib where liquid biopsy identified an additional 22.5% of patients with a BRCA1/2 alteration for whom tissue was sample unavailable³, and in the PROfound trial for olaparib where 31% of tissue samples were unable to be successfully sequenced for comprehensive genomic profiling testing.⁴

Due to genomic variation in metastatic sites, liquid biopsies may have advantages over individual metastatic site biopsies in their ability to capture the entire range of potential therapeutic opportunities for prostate cancer patients.⁵

* FoundationOne Liquid CDx only reports MSI when determined to be high
† MSI/dMMR testing should be considered in patients with metastatic castration-naive prostate cancer and is recommended in patients with metastatic CRPC.
FDA-approved portfolio of tests to help identify more treatment options:

**FOUNDATIONONE® CDx**

**TISSUE BIOPSY**
- Analyzes 300+ genes
- Reports Tumor Mutational Burden (TMB) and Microsatellite Instability (MSI)
- Covered by Medicare for qualifying patients†

Total: **10 SLIDES or 1 FFPE BLOCK**

IHC Testing for PD-L1
- Optional add-on with 4 additional slides

**FOUNDATIONONE® LIQUID CDx**

**LIQUID BIOPSY**
- Analyzes 300+ genes†
- Blood Tumor Mutational Burden (bTMB), MSI-High, and tumor fraction as a professional service, which has not been reviewed or approved by the FDA††
- Covered by Medicare for qualifying patients§

Total: **TWO 8.5mL TUBES of PERIPHERAL WHOLE BLOOD**

† FoundationOne® Liquid CDx is FDA-approved to report substitutions and indels in 311 genes, including rearrangements in ALK and BRCA1/2 and copy number alterations in BRCA1/2 and ERBB2 (HER2).
†† Comprehensive results across all 324 genes, including bTMB, MSI-H status, and tumor fraction are reported as a laboratory professional service which is not reviewed or approved by the FDA.
§ FoundationOne®CDx and FoundationOne® Liquid CDx are qualitative next-generation sequencing based in vitro diagnostic tests for advanced cancer patients with solid tumors and are for prescription use only. FoundationOne CDx utilizes FFPE tissue and analyzes 324 genes as well as genomic signatures. FoundationOne Liquid CDx analyzes 324 genes utilizing circulating cell-free DNA and is FDA-approved to report short variants in 311 genes. The tests are companion diagnostics to identify patients who may benefit from treatment with specific therapies in accordance with the therapeutic product labeling. Additional genomic findings may be reported and are not prescriptive or conclusive for labeled use of any specific therapeutic product. Use of the tests does not guarantee a patient will be matched to a treatment. A negative result does not rule out the presence of an alteration. Some patients may require a biopsy for testing with FoundationOne CDx when archival tissue is not available which may pose a risk. Patients who are tested with FoundationOne Liquid CDx and are negative for companion diagnostic mutations should be reflexed to tumor tissue testing and mutation status confirmed using an FDA-approved tumor tissue test, if feasible. For the complete label, including companion diagnostic indications and important risk information, please visit www.F1CDxLabel.com and www.F1LCDxLabel.com.

To order a test, go to [www.foundationmedicine.com/order](http://www.foundationmedicine.com/order)

Lynparza® is a registered trademark of the AstraZeneca group of companies. Rubraca® is a registered trademark of Clovis Oncology. Vitrakvi® is a registered trademark of Bayer Aktiengesellschaft. Rozlytrek® is a registered trademark of Keytruda® is a registered trademark of Merck Sharp & Dohme Corp.

References:
1. Referenced with permission from the NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines®) for Prostate Cancer V2.2021. National Comprehensive Cancer Network, Inc. 2021. All rights reserved. Accessed March 25, 2021. To view the most recent and complete version of the guideline, go online to NCCN.org. NCCN makes no warranties of any kind whatsoever regarding their content, use or application and disclaims any responsibility for their application or use in any way.
5. Necchi, Andrea et al. Comprehensive genomic profiling (CGP) in post-systemic treatment (Post) metastatic sites (MET) and pretreatment (Pre) primary tumors (PT) of metastatic prostate cancer (mPC). Journal of Clinical Oncology 38, no. 6_suppl (February 20, 2020) 175S-175. DOI: 10.1200/JCO.2020.38.6_suppl175
6. The Centers for Medicare & Medicaid Services (CMS) Decision Memo for Next Generation Sequencing (NGS) for Medicare Beneficiaries with Advanced Cancer (CAG-00450R) see Appendix B.

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