

Prediction of PARP inhibitor response and resistance utilizing a CTC phenotypic classifier in patients with metastatic castration-resistant prostate cancer (mCRPC):Results from the NCI 9012 trial

Felix Y. Feng¹, Stephanie Daignault-Newton², Adam Jendrisak³, Yipeng Wang³, Stephanie Greene³, Angel Rodriguez³, Jerry Lee³, Lyndsey Dugan³, Javed Siddiqui², Jessica Louw³, Chassidy Johnson³, Przemyslaw Twardowski⁴, Costantine Albany⁵, Mark Stein⁶, Walter M. Stadler⁷, Lakshmi Kunju², Arul M. Chinnaiyan², Mark Landers³, Ryan Dittamore³, Maha Hussain²

¹ UCSF, Diller Comprehensive Cancer Center, San Francisco, CA; ² University of Michigan Comprehensive Cancer Center, Ann Arbor, MI; ³ Epic Sciences, Inc., San Diego, CA; ⁴ City of Hope, Duarte, CA; ⁵ Indiana University Health, Indianapolis, IN; ⁶ Rutgers Cancer Institute of New Jersey, Rutgers, NJ; ⁷ University of Chicago Medicine, Chicago, IL

Background

PARP inhibitors (PARPi) have efficacy in mCRPC harboring homologous recombination DNA repair deficiencies (HRD), but there is no non-invasive assay to predict PARPi response.¹ Biomarkers of HRD, including genomic scarring or alterations of HRD related genes, have demonstrated an ability to predict PARPi patient response.² However, despite HRD related genomics, not all patients respond, while many patients without HRD genomics do. In addition, the identification of secondary mutations identify cells ability to become HRD competent, PARP reversion.³ Previous work characterizing single CTCs from mCRPC patients has identified subclonal populations of CTCs with unique phenotypes and somatic genomic instability consistent with HRD and resulting in worse outcomes following Abiraterone treatment.⁴ NCI 9012 evaluated Abiraterone alone with or without the PARPi Veliparib in mCRPC patients. We now determine if we can identify PARPi resistant CTC phenotypic subtypes and when combined with previously trained HRD CTC phenotypes can develop a biomarker which identifies patients who will have improved outcomes with Veliparib + Abiraterone vs. Abiraterone alone.



-) Epic CTC identification, enumeration and analysis: Nucleated cells from blood sample placed onto slides and stored in a -80°C biorepository. Slides are stained with cytokeratin (CK), CD45, DAPI, AR N-term and scanned. CTC candidates are detected by a multi-parametric digital pathology algorithm followed by human reader confirmation of CTCs and quantification of biomarker expression
- 2) Segmentation and image analysis: CTCs are segmented within the DAPI, CK, AR channels and ~20 nuclear and morphological features are extracted. 3) Single Cell Sequencing: Individual CTCs representing unique morphological phenotypes are recovered from the surface of the slide post morphological analysis. As per previously presented⁵, single CTCs are lysed, whole genome amplified and constructed into shotgun libraries. Each library is sequenced by low pass whole genome (0.3X) Whole genome CNV categorizes 1Mb segments for amplifications or deletions. Regions >10Mb combined gained or loss are scored as a single large scale transition (LST)⁶.
- 4) Phenotype to genotype analysis: As previously described⁴, quantitative and qualitative digital pathology features were correlated with the magnitude of quantified, sequenced-derived <u>actual LST</u> (aLST) per CTC using regression modeling. The output of this model is dichotomized and identified as an HRD+ phenotype CTC.

Patient Demographics & CTC Classifier Development

NCI 9012 Clinical Trial Demographics		В	lood Sample
NCI 9012 Demographics	Number (%)		NCI 9012
Patients registered (12 sites)	190		samples f
Race of Patient			
White	164 (86%)		↓
Black/Other	26 (14%)	Arı	m A
PSA/Age Demographics		Abirat	terone
Median Age (Min, Max)	68 (45-90)	(40 Pa	atients)
Median PSA (Min, Max)	35.5 (0.04-1558)		
Віорѕу Туре			
Patients who underwent metastatic biopsy	185	Baseline Samples	On-therapy Samples (24
Bone	96 (52%)	(42 samples, 707 CTCs)	samples, 1153 CTCs)
Soft Tissue	89 (48%)	* 2/84 base	line sample not utilized

Components of the CTC Classifier HRD+ CTC Phenotypes





HRD- CTC Phenotypes

Biomarker	Bi
classification	
Biomarker +	Pr
	&
	C٦
Biomarker -	Pr
	or

Training of **PARP resistant CTC phenotype** Developed from analysis and comparisons of ARM A & ARM B blood samples using regression analysis of phenotypic features

, Joaquin, et al. New England Journal of Medicine 373.18 (2015): 1697-1708. (2) Fong, Peter C., et al. New England Journal of Medicine 361.2 (2009): 123-134. (3) Cruz, C., et al. Cancer Research 76.4 Supplement (2016): P4-07. (4) Scher, Howard I., et al. ASCO Annual Meeting Proceedings. Vol. 34. No. 15_suppl. 2016. (5) Landers, Mark, et al. ASCO Annual Meeting Proceedings. Vol. 33. No. 15_suppl. 2015. (6) Popova, Tatiana, et al. Cancer research 72.21 (2012): 5454-5462.



PARPi CTC Biomarker Classifier Definition

marker Definition

resence of HRD+ CTCs **Absence of PARP resistant**

resence of PARP resistant CTCs absence of HRD+ CTCs

HRD- CTCs Phenotypes

- Low nuclear/cytoplasm ratio
- High eccentricity
- Low nuclear texture





Phenotype \rightarrow Genotype Association HRD+ Phenotypic CTC scores have increased Genomic Instability

434 CTCs from 35 patients were measured for HRD Phenotype utilizing a previous trained algorithm in an independent cohort (Scher et al.)⁴



Visualized are the 434 single CTC CNV genomic alterations (green=deletion; red=amplification) of known oncogenes and tumor suppressors, ranked from highest (left) to lowest (right).



mic

Ο

LST



Patient Baseline Incidence of CTC Classifier in Cohort

	Arm A (Abiraterone only)	Arm B (Abiraterone + Veliparib)	Total
Biomarker Positive	23% (9/39)	35% (15/43)	29% (24/82)
Biomarker Negative	77% (30/39)	65% (28/43)	71% (58/82)

Changes	
---------	--

Therapy Arm	Baseline HRD+ CTCs (of patient samples)
Arm A (Abiraterone)	9 HRD+ CTCs/patien (40 samples)
Arm B (Abiraterone + Veliparib)	17 HRD+ CTCs/patien (44 samples)

- compared to Abiraterone alone.
- of efficacy of novel HRD targeted therapies.

Support: NCI Cancer Therapy Evaluation Program

• Patients with the PARP sensitive CTC biomarker and HRD+ CTCs have improved PSA ORR (93% vs. 22%) and trend to improved time to PSA progression (HR=0.45; p=0.12) with Abiraterone + Veleparib treatment

The CTC HRD+ phenotype increases in the total population in patients receiving Abiraterone only (+166%), but decrease in patients receiving Abiraterone + Veliparib (-82%), supporting cellular PARPi sensitivity.

• Further development of single CTC phenotypic classifiers could help with patient selection, and monitoring