Analysis of Circulating Tumor Cells (CTCs) in Patients Across Multiple Metastatic Breast Cancer (mBCa) Cohorts Identifies Marked Inter- and Intra-Patient Heterogeneity in CTC Size, Shape, and Overall Morphology

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Background
- The choice between hormonal therapies and chemotherapy is a frequent decision in the care of metastatic breast cancer (mBCa) patients.
- We previously developed quantitative measures of phenotypic CTC heterogeneity in metastatic castration-resistant prostate cancer (mCRPC) and found higher heterogeneity was associated with better survival on chemotherapy vs. targeted hormonal therapies, and the reverse was true in low heterogeneity patients (Scher et al., 2017 Cancer Research).
- We apply our previous heterogeneity methodologies to a cohort of mBCa patient CTCs to ascertain feasibility in mBCa.

Methods
- 165 blood samples from mBCa patients were processed for CTC analysis utilizing the Epicon Sciences platform. Following enumeration, multi-dimensional phenotypic characterization analysis was performed utilizing protein expression and digital pathology features.
- Features from each CTC (1590 CTCs from 144 patients, 81 HR+, 12 HER2+; 4 HR+/HER2+, 47 TNBC) were compared by unsupervised clustering. Shannon index and intra-patient variance analyses to assess the intrapatient heterogeneity among mBCa CTC phenotypes.

Schematic of Epic CTC Platform CTC enumeration, morphology, biomarker analyses and single cell sequencing workflow:
1. Nucleated cells from blood sample placed onto slides and stored in a -80°C freezer.
2. Slides are stained with cytokeratin (CK), CD45, DAPI and scanned. CTC candidates are detected by a multiparametric digital pathology algorithm followed by manual reader confirmation of CTCs and quantification of biomarker expression.
3. CTCs are segmented from the DAPI and CK channels, and single cell features are extracted.
4. CTCs undergo Principle Component Analysis (PCA) removing noise and redundant dimensions, and weighting features with more variance. Machine learning clustering algorithms found 7 CTC subtypes from macro trends in high-dimensional biomarkers across all CTCs from all samples in cohort, and assigned each CTC to 1 of 7 subtypes. Heterogeneity is quantified by counting CTCs per “Cell Type” in each sample, then using a standard Shannon Index to quantify CTC phenotypic diversity per patient sample.
5. Single cells are identified, relocated, captured, whole genome amplified (WGA), library prepared and low pass whole genome sequenced for Large Scale Transitions (LST, a surrogate of chromosomal instability) and gene copy number alterations (CNA) (Green et al., 2016 PLoSOne).

Conclusions
- Distinct CTC phenotypes are visible across mBCa patients and clustered into 7 CTC phenotypic subtypes.
- A wide range of phenotypic CTC heterogeneity is observed between and within patients.
- mBCaphenotypic CTC subtypes are associated with unique genomic profiles.
- We seek to determine if patients with high heterogeneity might be better candidates for hormonal therapy. Studies linking heterogeneity to therapeutic efficacy and patient outcome are ongoing.