REVIEW

Opportunities and Challenges in Implementation of Multiparameter Single Cell Analysis Platforms for Clinical Translation

Susan M. Keating1, D. Lansing Taylor2, Anne L. Plant3, E. David Litwack4, Peter Kuhn5, Emily J. Greenspan6, Christopher M. Hartshorn7, Caroline C. Sigman1, Gary J. Kelloff8, David D. Chang9, Gregory Friberg9, Jerry S. H. Lee6,∗ and Keisuke Kuida10,∗

The high-content interrogation of single cells with platforms optimized for the multiparameter characterization of cells in liquid and solid biopsy samples can enable characterization of heterogeneous populations of cells ex vivo. Doing so will advance the diagnosis, prognosis, and treatment of cancer and other diseases. However, it is important to understand the unique issues in resolving heterogeneity and variability at the single cell level before navigating the validation and regulatory requirements in order for these technologies to impact patient care. Since 2013, leading experts representing industry, academia, and government have been brought together as part of the Foundation for the National Institutes of Health (FNIH) Biomarkers Consortium to foster the potential of high-content data integration for clinical translation.

Clin Transl Sci (2018) 11, 267–276; doi:10.1111/cts.12536; published online on 2 March 2018.

Precision medicine is moving cancer treatment from etiological and histological parameter-based treatments to those that target specific key molecular drivers of disease in a time-resolved fashion. To arrive at actionable end points, representative subamples of the disease tissue and multiparameter measurements are required to accurately profile tissues at the single cell level to describe the relevant biological unit of disease. Bulk measurements must assume a normal distribution of data points. Thus, given the possible spatial and temporal heterogeneity of tumors with morphological, biochemical, physical, and genetically different subpopulations, coupled with the capacity for cell evolution, the true variation of cellular features in the population may be masked if the measured responses from cells are averaged across cell populations, limiting the ability to detect and target relevant biomarkers. For example, in solid tumors, the tumor is a “system” within the organ that includes normal cells, cancer cells exhibiting a range of genetic alterations, stromal cells, including fibroblasts, vasculature cells, and immune cells, such as dendritic cells, granulocytes, macrophages, and lymphocytes that are critical to the development, progression, metastasis, and response to therapy, for both standard cytotoxic chemotherapy and immunotherapy. The value in analyzing single cells as biological units of disease to more accurately record the individual and range of cellular responses is increasingly recognized, as is the value of multiparameter single cell analysis that can be obtained using next-generation technologies. This includes identification of rare genetic mutations within a tumor, better understanding of signaling and metabolic pathways, and prediction of the optimal treatment regimens to prevent tumor regrowth. Next-generation technologies are defined here as methods that permit medium to high throughput, and multiparameter analyses of either fixed or living, single cell morphometric, genomic, proteomic, and/or metabolomics characteristics. Significant challenges include the deconvolution of biological variation from technical uncertainty, the computational analysis of biological variations over time and space, and the integration of single cell genomic, proteomic, and metabolic data.

High-content platforms were initially built over 2 decades ago for drug discovery screening with a focus on the temporal-spatial dynamics and heterogeneity among cells by making measurements on each cell in a population. Since they were first introduced, high-content systems have been further developed to incorporate advances in optical contrast physics, automated microscopy systems, image analysis software, fluorescence-based reagents, and computational methods. They are often coupled with other platforms, such as next-generation sequencing for genomic profiling of cells and/or imaging mass cytometry, and are deployed as large-scale information-based tools.

1CCS Associates, San Jose, California, USA; 2University of Pittsburgh Drug Discovery Institute, University of Pittsburgh, Pittsburgh, Pennsylvania, USA; 3Biosystems and Biomaterials Division Materials Measurement Laboratory, National Institute of Standards and Technology, Gaithersburg, Maryland, USA; 4Office of In Vitro Diagnostics and Radiological Health, Center for Devices and Radiological Health, Food and Drug Administration, Silver Spring, Maryland, USA; 5Center for Strategic Scientific Initiatives, National Cancer Institute, Bethesda, Maryland, USA; 6University of Southern California, Los Angeles, California, USA; 7Center for Strategic Scientific Initiatives, National Cancer Institute, Bethesda, Maryland, USA; 8Kite Pharma, Inc., Santa Monica, California, USA; 9Amgen, Thousand Oaks, California, USA; 10Deciphera Pharmaceuticals, LLC, Waltham, Massachusetts, USA.

∗Correspondence: Jerry S. H. Lee (leejerry@usc.edu); Keisuke Kuida (kkuida@deciphera.com)

Received 25 October 2017; accepted 19 December 2017; published online on 2 March 2018. doi:10.1111/cts.12536
multiparameter information available in these systems is starting to be more fully exploited in in vitro and ex vivo settings to understand the biology of clinical samples. Furthermore, the definition of high-content has evolved to include the methods defined above as next-generation technologies that apply to genomic, proteomic, and/or metabolomic measurements that are based on single cell, usually extensive, multiparameter measurements.

The High-Content Data Integration (HCDI) working group was formed in early 2013 out of the Foundation for the National Institutes of Health (FNIH) Biomarkers Consortium Cancer Steering Committee (a public-private partnership) to evaluate emerging technologies (other than those for sequencing) that use advanced acquisition and analysis of single and bulk cell populations for development of tailored signatures that can assist in patient stratification and identification of patient responses to treatment. The HCDI working group has discussed platforms for patient sample single-cell analysis, such as those listed in Table 1, while considering key challenges and critical questions concerning real-world development and translational potential of these types of platforms, as listed in Table 2.

In this article, the challenges encountered with newly evolving high-content measurement systems are illustrated with examples of two platforms, the first, the High Definition Single Cell Analysis (HD-SCA) workflow, optimized for identifying and characterizing rare cells in liquid biopsy samples and the second, the MultiOmyx Immunofluorescence (MxIF), for enabling “hyperplexed” measurements of proteins (Fluidigm, South San Francisco, CA),52

### Table 1 Examples of new technologies capable of clinical single cell analysis

| Technology platform | Developer / sponsor | Description | Data elements |
|---------------------|---------------------|-------------|---------------|
| HD-SCA              | Peter Kuhn, University of Southern California / Commercialization: Epic Sciences, San Diego, CA | Imaging platform for multiplex single cell measurements, imaging, immunofluorescence labelling. Individual cells can be picked for DNA sequencing. Slides can also be subjected to laser ablation for imaging mass cytometry for spatial resolution of proteins (Fluidigm, South San Francisco, CA).52 | Cell images:  
- Cell and nuclear morphology  
- Selected cell and nuclear protein expression (2–3 immunofluorescence label intensity)  
- Multiplex protein expression (4–37)  
- Single cell DNA sequencing |
| SMR                 | Scott Manalis / Massachusetts Institute of Technology | Single-cell mass accumulation rate real-time measurements over short time frames (20 min). Ex vivo evaluation of patient samples provides evidence as to the range in responses from a patient’s tumor cells to a drug, including development of resistance.53 | Time-stamped single-cell mass (changes)  
- Additional, protein expression!4 |
| MxIF                | GE Global Research and Lans Taylor & Chakra Chennubhotla / University of Pittsburgh | Sequential fluorescent labeling of slides with antibodies, DNA and RNA probes, imaging for “hyperplexed” (>7 biomarkers up to ca. 60) fluorescence imaging for quantitative, single-cell, and subcellular characterization of analytes in formalin-fixed paraffin-embedded tissue coupled with spatial statistical methods to define microdomains.24,25 | Cell images (immunofluorescence label intensity):  
- Protein expression  
- RNA  
- DNA  
- In situ hybridization |
| SCBC                | James Heath / California Institute of Technology | Multiplex quantitative protein expression, secretion, and intracellular signaling, from single cells. Dissociated cells are introduced into microchambers containing miniature antibody-DNA-barcoded microarrays. Analyte detection using miniature ELISA measurement and quantitation methods.42,55,56 | Cell-based (immunofluorescence label intensity) >20 protein expression |
| Mass spectrometry imaging | Garry Nolan / Stanford University | Single-cell analysis utilizing mass spectrometric measurement of metal elements tagged to antibodies. Individual antibody-bound cell is vaporized, ionized, and analyzed on a mass spectrometer.57 | Simultaneous quantification of 50 mass tags (markers) |
| CAFE MiCells        | David Andrews / Sunnybrook Research Institute | Automated high-content image analysis using nontoxic, cell permeable dyes58 | Visualization of cell states and outcomes of treatment |

CAFÉ MiCells, classification and automated feature extraction of micrographs of cells; ELISA, enzyme-linked immunosorbent assay; HD-SCA, high definition single cell analysis; MxIF, MultiOmyx; SCBC, single-cell barcode chip; SMR, suspended microchannel resonator.
MULTIPARAMETER CHARACTERIZATION OF CIRCULATING VS. SOLID TUMOR CELLS

Circulating tumor cells (CTCs) are viable tumor-derived cells that exist in the peripheral blood of patients with cancer in very low concentrations (as low as ~10^4/mL). The CTCs extravasate into the bloodstream and circulate; they may form secondary metastases, self-seed, or remain in the circulatory system until clearance. These cells are an accessible source of nonhematological tumor cells and, along with circulating nucleic acids, are components of what are termed liquid biopsies, which are increasingly being recognized as potentially valuable noninvasive tools for temporal characterization of a patient’s tumor (including the constituent heterogeneity), because they can be isolated and characterized from a noninvasive blood draw. One significant barrier to translation of CTC detection platforms to clinical use is the understanding of the relationship of biological signatures found in CTCs to those of solid tissue-derived tumor cells to support the use of liquid biopsies in place of the more invasive and expensive solid tissue biopsies. One approach to overcome this hurdle is the analysis and comparison of biomarkers measured from isolated single cells from solid tissue and CTCs using a common platform. The HCDI working group worked with the Kuhn group at the University of Southern California and FNHI to design and fund a project, the High Definition Single Cell Analysis of Blood and Tissue Biopsies in Patients with Colorectal Cancer undergoing Hepatitis Metastasectomy (HD-SCA), to investigate the correlation between single-cell-based biological signatures of liquid biopsies, particularly CTCs and solid tumor tissue. One of the outcomes of this project will be to demonstrate the advantage of single cell, CTC-based biomarkers to assist in laying the foundation for subsequent studies to define clinical utility of liquid biopsy-based biomarkers.

The HD-SCA platform developed by Kuhn and colleagues is a single-cell direct imaging analysis system that can be used for morphometric, proteomic, and downstream genomic characterization of any rare cell from either the fluid phase or solid tissues with the flexibility to probe for any combination of markers needed to characterize specific populations of cells. This workflow then can be used to generate highly multiplexed data points derived from any combination of the morphologic, genomic, and proteomic (4–37 proteins with copy number variant and single nucleotide variation) measurements. As shown in Figure 1, all nucleated cells from a liquid biopsy are immunofluorescently labeled for high resolution digital imaging, providing detailed nuclear and cytoplasmic features for morphological and protein expression measurements. The location of a specific cell of interest on a slide is registered so that the cell can be selected for molecular analysis, including genomic sequencing, targeted proteomics, and other measurements. Moreover, a number of slides are generated from each patient sample so that each sample can be stained multiple times for analysis of different parameters.

HYPERPLEXED FLUORESCENCE MEASUREMENTS OF THE CELLULAR AND SUBCELLULAR EXPRESSION LEVELS AND SPATIAL RELATIONSHIPS BETWEEN TUMOR BIOMARKERS IN TISSUE SECTIONS

A goal of high-content and all image cytometry methods is to increase the number of biomarkers that can be measured within the same cells and tissues so that the complexity of intracellular interactions and intratumoral heterogeneities can be defined in greater molecular detail within the physical context or architecture of the tumor “system.” A number of approaches have been developed to address this goal from multiplexed fluorescence, with up to seven distinct fluorescent probes applied to the same sample, to hyperplexed fluorescence, using more than seven separate images of biomarkers in the same sample (reviewed in ref. 9). Extensive literature exists on the development of multiplexed and more recently hyperplexed fluorescence methods for tissue analyses, including instrumentation, software, and bioinformatics tools that can be used in the development of diagnostic tests. The ability to measure many parameters in large numbers of tissue samples has propelled these assays into the realm of big data challenges for the future.

One platform that exemplifies this approach evolved in collaboration among General Electric Global Research and the Taylor/Chennubhotla Laboratories at The Drug Discovery Institute, and the Department of Computational and Systems Biology at the University of Pittsburgh (Figure 2). MxIF is a fluorescence imaging technology that allows up to 60 fluorescent protein biomarkers, typically antibodies, to be analyzed in situ on a single formalin-fixed, paraffin-embedded tissue section. Using single or multiplexed (from 2–7 biomarkers) or hyperplexed (>7 biomarkers), the expression levels and subcellular locations of proteins can be quantified in tissue sections. DNA and RNA fluorescence in situ hybridization (FISH) may also be conducted on the same sample, thus providing additional -omic measurements. MxIF involves a series of sequential labeling, imaging, and dye inactivation steps so that a large number of different biomarkers can be measured with only a few fluorophores. Spatial statistical methods are then
Opportunities and Challenges in Single Cell Analysis for Clinical Translation
Keating et al.

Figure 1 The high definition single cell analysis (HD-SCA) generic temporal and spatial analysis. The HD-SCA workflow is a single-cell analysis system generating morphometric, proteomic, and genomic characterization of any rare cell from either liquid (blood draw or bone marrow aspirate) or solid tissue biopsy. Representative circulating tumor cells (CTCs) and solid tissue samples from patients with cancer are isolated and imaged using the same HD-SCA system. Blood cells after red blood cell depletion and tissue cells obtained from touch preparations of either metastases or primary tumor are plated. Slides with nucleated blood cells and cell monolayers from touch preparations are immunofluorescently labeled in the same batch in three wavelengths, and the resultant stained slides are imaged at 40× magnification to generate high-resolution digital images with detailed nuclear and cytoplasmic features for morphological cellular characteristics and protein expression. Captured CTCs are classified as CK+ (red), CD45− (green) cells of epithelial origin with an intact, nonapoptotic-appearing nucleus by DAPI (blue) imaging, morphologically distinct from surrounding white blood cells by shape and/or size. Cells of interest can be picked individually and isolated for single-cell genomic copy number alteration (CNA) or targeted proteomic analysis via imaging mass cytometry.

applied to the digital images to define the extent of heterogeneity and to identify tumor microdomains. The single cell analysis capability was originally demonstrated in the analysis of 700 patients with colorectal cancer with differing phosphorylation of ribosomal protein S6. Mutually exclusive phosphorylation patterns of the two canonical substrates of mTORC1 (S6K1 and 4E-BP1) were observed in individual cells, large regions of most tumors, and in distinct cell lineages, demonstrating differential pathway activation. In the research setting, MxIF has also been applied to evaluate signaling pathway changes in mouse salivary glands, estrogen receptor (ER), progesterone receptor (PR), human epidermal growth factor receptor 2 (HER2), and Ki67 colocalization in breast cancer, and other single-cell analyses.

The future of targeted multiplexed and hyperplexed fluorescence using platforms, such as MxIF, is very promising for basic research, including defining pathways of cancer evolution using “spatial proteomics” to directly map pathways inferred from genomics and proteomics approaches, and defining the microdomains of cancer within the native architecture, stromal and immune cell interactions that influence drug treatments, drug discovery/development, and diagnostics (Figure 2). The platform is valuable for identifying the optimal/minimal combination of biomarkers that could be incorporated into a smaller number of biomarkers for multiplexed tissue-based diagnostic tests and biomarkers for drug development. Hyperplexed fluorescence with the MxIF using panels of biomarkers tailored to the cell types will be an important platform to define the functional interplay between the cells in the tumor microenvironment and to define the response to therapeutics when combined with automated machine learning software tools to identify and to quantify spatial patterns of biomarkers that reflect the heterogeneity of disease.

An alternative approach is imaging mass cytometry, as implemented on Fluidigm’s Hyperion Imaging System. Antibodies are mass-tagged with metal labels and, after incubation of liquid or solid biopsy samples on standard microscope slides, a region of interest is selected, laser ablated, and the atomized sample injected into a time of flight mass spectrometer. By calculating the slide position, this approach yields a 1 micron spatial resolution of the sample with 4–37 or more proteins simultaneously. This concept is currently being
Figure 2  Pointwise mutual information (PMI) for quantifying spatial heterogeneity. (a) A pseudo-colored multichannel fluorescence image labeled iteratively by the MultiOmyx platform is shown for an estrogen receptor (ER)+ invasive ductal carcinoma from a tissue microarray. Three biomarker channels were used to demonstrate the approach: HER2 (red), ER (blue), and PR (green), although this method can be scaled for >50 biomarkers. Areas of PR/ER co-activation will appear in cyan, HER2/ER co-activation in magenta, and PR/HER2 co-activation in yellow. The upper and lower arrows indicate heterogeneous tumor microdomains with higher than average ER+/PR+ phenotyped cells and mostly ER+ cells, respectively. (b) Machine learning methods can be used to identify dominant cellular phenotypes from biomarker expression patterns over an entire tissue microarray, which in this case were eight. Each cell is then classified with the most similar dominant phenotype. (c) In order to represent the tumor topology, a spatial network of the cells in each tissue microarray spot or whole tissue section is constructed, in which each cell has the ability to communicate with nearby cells up to a certain limit, 250 μm, and the communication propensity is assumed to be inversely proportional to the cellular distance. (d) PMI quantifies the statistical associations, both linear and non-linear, between each pair of cellular phenotypes. In particular, PMI calculates the logarithmic joint probability of finding a particular pair of cellular phenotypes occurring in close proximity, relative to the probability of these phenotypes co-occurring at random. (e) By referencing a specific interaction pair in the PMI plot, one can interrogate the network subgraphs/motifs that contribute to the PMI dependencies. A PMI map with strong diagonal entries and weak off-diagonal entries describes a globally heterogeneous but locally homogeneous tumor. On the contrary, a PMI map with strong off-diagonal entries describes a tumor that is locally heterogeneous. (f) An example TMA spot with three locally heterogeneous tumor microdomains denoted by the off-diagonal entries in the PMI map, containing phenotypes 1 and 6, 2 and 4, and 3 and 8. PMI maps can also portray anti-associations (e.g., if phenotype 1 never occurs spatially near phenotype 3). The ensemble of associations and anti-associations of varying intensities along or off the diagonal represent the true complexity of tumor images in a format that can be summarized and interrogated. PMI maps are predicted to become diagnostic and prognostic biomarkers.

integrated into various basic science and clinical research workflows.

UNIQUE CHALLENGES AND OPPORTUNITIES POSED BY SINGLE-CELL ANALYSIS VARIABILITY

Single-cell science attempts to measure the biological heterogeneity across cell populations to provide a quantitative description of the overall biological state or disease state. Biological heterogeneity can be manifested in a population of cells by the observation of a range of phenotypes even when the population of cells and the environment are nominally homogeneous. This heterogeneity can be due to stochastic fluctuations in molecular events, as well as extrinsic effects, including the effect of nearby cells. Biological heterogeneity, and the dynamics of how heterogeneity arises in a population, can be used to develop theoretical constructs for understanding control mechanisms and predicting population dynamics, and provide a better understanding of intracellular pathways, control systems, and mechanisms that determine disease progression. Biological heterogeneity cannot be assessed unless the technical variability in the measurement can be independently determined. Analytical methods, such as Western blots or enzyme-linked immunosorbent assays or other solution-based assays, provide a single value that represents an average over all the cells in the population. When using such techniques, single-cell data that report on biological heterogeneity are lost, although replicates of the experiment can provide information about measurement variability. Single-cell methods for transcriptomics can provide information about biological variability between cells, but the data are convoluted with measurement variability unless duplicate or triplicate measurements can be made on each cell. This is often impossible for single cell transcriptomics; replicate measurements are not typically accessible because of the quantity of samples and the limits on the sensitivity of the technique. Determining how to best address measurement uncertainties under this situation becomes a challenge. A number of approaches are being considered for assuring confidence in these measurements, including the use of the External RNA Controls Consortium (ERCC) spike-in material to provide a metric of accuracy for those known sequences. Other approaches include the use of Bayesian statistics to assess real differences in the presence of dropout events that occur due to the low amount of RNA
in cell samples and result in detection of the gene in some cells and not in others. When the apparent heterogeneity in gene expression is simply due to technical issues, the data can lead to erroneous conclusions of biological heterogeneity. Other methods include the use of unique molecular identifiers in the primers for counting individual transcripts. In cases in which replicate measurements can be performed, such as has been shown with single cell proteomics, duplicate proteomic measurements from individual cells can be performed. Such replicate measurements can result in quantification of the technical error in the measurement, and unambiguous determination of whether the values measured from one cell are significantly different from the values measured for another cell.

Technologies, such as nondestructive microscopic imaging of individual cells, can have an advantage over other analytical methods, because it is possible to generate multiple samplings of the same field of view to provide a measure of instrument noise, such as shot noise, lamp fluctuations, and photobleaching. An automated routine that operates within open source software provides an easy way to qualify instrument performance to assess such operational figures of merit and to provide instrument calibration. The relative contribution of these sources of uncertainty can be compared with other aspects of the measurement and determined to be significant contributors to be quantified, or not significant contributors and can be safely ignored. Other aspects of the measurement that can add significantly to uncertainty include sampling, handling, and reagents. The aggregate of these contributions can be determined by comparison of the distribution of responses provided by replicate measurements as described in ref. 40, in which the width of the distribution of cell phenotype reflects biological heterogeneity and the variation between replicate measurements of the population distribution reflects measurement uncertainty. Using sufficient care to quantify measurement uncertainty allows deconvolution of biological heterogeneity and measurement uncertainty, and biological heterogeneity can be unambiguously determined. Accurate quantification of heterogeneity requires sampling of the appropriate number of cells; the more disperse the population phenotype the larger the number of cells that must be sampled to accurately represent the population.

Time-resolved dynamic data on individual cells within a living cell population are uniquely accessible through imaging. Examining differences in dynamics of processes, such as promoter activation in a large number of individual cells, provides determination of variations in rates of fluctuations in cellular responses and epigenetic effects, and assists in choosing appropriate theoretical treatments. Dynamic data can also confirm stability in gene expression, such as in stem cell colonies, and the spatial location of the expressed gene within colonies. Such analysis can also allow detection of rare events. Longer-term time-dependent measurements of cell populations by imaging or flow cytometry can indicate whether population heterogeneity is the result of stochastic fluctuations within individual cells or is due to genomic differences within the population. A distribution in cellular responses within a population due to stochastic fluctuations will be stable or stationary over time. In this case, a selected “subpopulation” will relax back to the original broad distribution after a number of passages because each cell belongs to the same distribution of probable phenotypes. A population that is composed of true subpopulations that are genetically distinct will diverge according to the relative rates of proliferation of the genetically distinct cell populations.

Although measurement uncertainty cannot be eliminated, it can be mitigated by good experimental design and protocols, as discussed in Table 3. Mitigation of measurement uncertainty can be achieved in single-cell measurements by appropriate control experiments in the evaluation of the single-cell barcode chip for proteomics analysis. This group utilized well-characterized recombinant proteins for calibration and defining the dynamic range of different protein assays and simulation based on known physical principles applied to the device to estimate measurement errors from locations of cells.

VALIDATION AND REGULATORY CONSIDERATIONS

Although tests that leverage CTC platforms (i.e., platforms or systems that identify and isolate CTCs; the entire in vitro diagnostic (IVD) will be referred to as a “CTC test”) remain largely in the research and development phase, their diagnostic potential is clearly on the horizon. Widespread adoption of these tests in the clinic will require studies to demonstrate analytical and clinical validity for appropriate marketing authorization, as well as clinical utility studies for adoption and reimbursement. Early consideration of regulatory requirements may make a large difference in reducing later potential regulatory challenges, many of which may be related to the platforms used to isolate CTCs or, if applicable, the high content methodology used for CTC analysis.

When reviewing a premarket submission for an IVD, the US Food and Drug Administration (FDA) typically evaluates data establishing the analytical and clinical validity of the IVD in the context of its proposed intended use. The types of studies necessary for marketing authorization will depend on the details of the diagnostic platform that is eventually developed. An open question is: what role will high-content systems play in future diagnostic tests?

Currently, research using high-content assays that use CTC platforms aim to identify new biomarkers for CTCs. If a discrete number of biomarkers are identified, IVDs that use more traditional techniques (e.g., antibody staining, polymerase chain reaction, etc) to detect those same biomarkers may be developed for a specific clinical use. In this scenario, the analytical and clinical validity of the assay could be demonstrated for each analyte that is detected. To date, the only CTC test that has been authorized by the FDA is the CellSearch Circulating Tumor Cell Kit, which is intended for the enumeration of CD45–EpCAM+ CK+ CTCs in whole blood for monitoring patients with metastatic breast, prostate, and colorectal cancer (see ref. 46 and related IVDs). The review of this product involved evaluation of the analytical and clinical validity of a well-defined analyte (e.g., a specific biomolecule or set of biomolecules) for specific clinical conditions.
Another possibility may be the development of a clinical test that produces as its reportable output high content data from single cells with too many targets to validate independently. For instance, genome sequencing of single CTCs may help address issues of tumor heterogeneity in a patient with cancer when selecting an appropriate therapy (e.g., identifying rare cells that carry actionable somatic mutations). In such cases, special challenges related to the unique features of high-content tests will need to be addressed, including the demonstration of analytical and clinical validation of high-density data from rapidly evolving technologies; the possibility of unlimited (unique) results, with unknown clinical significance that are, therefore, open to interpretation; and the identification and availability of appropriate reference methods and clinical samples for demonstration of analytical reliability and variation across all possible outputs.

In either case, it will be important for developers to plan ahead on how to demonstrate analytical and clinical performance for regulatory clearance or approval. Below, we outline some of the key issues that should be considered as early as possible.

**Analytical validity**

Analytical validity is the ability of a test to measure or detect a particular analyte (e.g., analytical specificity, limit of detection, analytical accuracy, precision, and robustness). Traditionally, for any type of assay that simultaneously measures multiple analytes, manufacturers must demonstrate that the test is analytically valid for each analyte. Because high-content tests are designed to simultaneously assess a very large number of analytes, a complete demonstration of analytical validity may not be feasible. In some such cases, the FDA has allowed representative analytical performance encompassing the type of variants and analytical challenges that could be encountered during use of the instrument or assay, and inferred overall performance from this representative subset (e.g., Illumina MiSeqDx Cystic Fibrosis Clinical Sequencing Assay).  

**Clinical validity**

Clinical validity is the association between the test result and a disease or condition, and is commonly described in terms of diagnostic accuracy. Test performance is commonly reported as clinical sensitivity and specificity, and/or positive
or negative predictive value. Clinical validity for IVDs can be established using a variety of approaches, such as clinical studies or from evidence in the literature. This is often not possible for high-content assays. Because they can simultaneously detect a large number of markers, such assays frequently detect rare or novel biological variation. Therefore, high-content data can often be used to divide patients with a given diagnosis into small, or even private, subsets defined by specific markers (e.g., rare genetic variants). With very low prevalence markers, it is difficult to perform clinical studies to definitively demonstrate clinical validity, and published evidence may be limited or not available. In such cases, other sources of evidence may be considered, provided the evidence is appropriate for the intended use of the IVD. For instance, one strategy that was successful in the clearance of the Illumina MiSeqDX Cystic Fibrosis 139-Variant Assay\(^48\) was to leverage information from the cystic fibrosis transmembrane conductance regulator CFTR2 database\(^49\) which is a high quality, well curated database that aggregates evidence from patients around the globe to establish the link between specific variants in the CFTR gene and the disease. The applicability of this strategy to other IVDs will depend on a number of factors, and may not be suitable for every situation.

The following additional issues should be considered for regulatory clearance or approval of IVDs incorporating single-cell platforms, whether or not they included a high-content component.

**Pre-analytical variables**

The term “pre-analytical” variables refers to all factors that may affect a specimen or sample before it enters the analytical process. No matter how complete analytical and clinical validation for an assay is, if there is not confidence that the sample being analyzed actually reflects what is happening in the patient, then the results will be meaningless. Often times, especially in oncology and related fields, specimen collection, handling, and processing (CHP) variables are the most important of these pre-analytical variables to consider. For example, what conditions (e.g., ischemia time) will be present at the time of surgery/resection? What will be the temperature range used for sample processing, shipping, and storage? In certain cases, it may be beneficial to undergo certain biospecimen-based or pre-analytical variable experiments to define the optimal collection, handling, and processing variables for the system, and ultimately produce best-practice guidelines that the rest of the field can use.

**Specimens**

Early planning should also take into account the need for adequate specimens that represent the analytes to be detected by the test. Given that high-content assays will likely identify rare markers, it is often not possible to obtain specimens representing all possible analytes that could be detected. In such cases, it may be possible to use contrived samples in lieu of some patient specimens, provided there are data to demonstrate that the operational behavior of the contrived samples is similar to that of clinical specimens. Where applicable, the need for separate training and validation sets should be factored in.

**Reference materials and performance standards**

One of the common challenges when developing new high-content tests is the lack of available reference materials and performance standards for assessing analytical and clinical error rates. For this reason, collaborative efforts involving multiple stakeholders, including the FDA, National Institutes of Health, the National Institute of Standards and Technology (NIST), the Centers for Disease Control and Prevention (CDC), academia, and industry are critical for development of performance standards and reference materials for all types of next-generation platforms, including both imaging and nonimaging methods, and for standardizing file formats and other aspects of informatics associated with these platforms. Such efforts are already beginning to yield results. For instance, the high-confidence calls produced by the National Institute of Standards and Technology through the Genome in a Bottle (GIAB) Consortium\(^50\) are widely used as standard reference materials for validation of next-generation sequencing-based tests. However, those sequences do not currently cover many of the medically relevant variants that will be detected using genomic tests and additional work is being done by the GIAB Consortium, the FNIH, and other efforts to address this. For other technologies, it is important for developers to assess whether reference materials and standards are available and, if not, what alternative strategies can be used to establish analytical validity in their absence. In the absence of reference materials, confirmation of results by a validated orthogonal method may be used to benchmark test performance.

**DISCUSSION**

It is becoming apparent that cancer is an evolutionary process of genetic, epigenetic, molecular, and physical-based alterations that manifests as heterogeneity across a tissue or development of resistance to therapeutic interventions over time. Clinical treatment decisions are often made based upon bulk measures of bone marrow blasts or size of a tumor mass on a computed tomography scan, leaving a vast amount of biologic information on the table that could theoretically be leveraged for the good of the patient. The ability to sample the cancer cells throughout the course of treatment and disease progression, and assay multiple functional changes simultaneously with the platforms, such as those described in this paper, may enable better and quicker detection of these changes and the ability to improve therapeutic interventions for disease management. The platforms profiled here are examples of instruments that translate high-content information out of the research laboratory to the clinical research setting for characterization and monitoring of single cells in tissue or blood. They go beyond enumeration of assay end points to enable profiling of heterogeneous cell populations. As such, the systems could be exploited in the drug development and clinical arenas not only for *in vitro* evaluations guiding clinical decisions, but also *ex vivo* studies, such as assessing target activity, probing for clinically relevant targets, and characterizing cancer cell and tumor heterogeneity.

Because these multiparameter systems can involve complicated analytic signatures, one of the keys to translation
of these new tools into the clinic will be in the careful attention to deconvolution of biological and instrument variability, and demonstration of analytical performance. In addition, demonstration of the biological significance and usefulness of an analysis is key. Furthermore, it is important to pay thoughtful attention to a developmental regulatory strategy, including early determination of the intended use of the platform, which will lead to successful development, validation, regulatory authorization, and clinical use. The process of making such multi-analyte platforms and complex analysis more generally available, such as through commercial clinical assay systems, may present interesting challenges. An understanding of the path to market for next-generation sequencing platforms and other complex clinical systems\(^{51}\) can inform that process, as more of these multiparameter, complex technological platforms will undoubtedly make their way into the clinic.

**Acknowledgments.** We thank the High Content Data Integration Working Group for the ongoing discussion of the platforms and clinical impact reviewed over the course of the meetings and Ms Rona Ackerman for her help editing the manuscript. Certain commercial equipment, instruments, or materials are identified in this paper to foster understanding. Such identification does not imply recommendation or endorsement by the National Institute of Standards and Technology, nor does it imply that the materials or equipment identified are necessarily the best available for the purpose.

**Conflict of Interest.** D.L.T. is a co-founder of Spintellec, a computational pathology company. P.K. is a shareholder, royalty recipient, and advisor of Epic Sciences, which has licensed the HD-SCA technology from The Scripps Research Institute.

**Source of Funding.** A project utilizing the HD-SCA led by P.K. is supported by the NHGRI Biomarkers Consortium.

1. Gerlinger, M. et al. Intratumor heterogeneity and branched evolution revealed by multi-region sequencing. N. Engl. J. Med. 366, 886–892 (2012).
2. Sutherland, K.D. & Visvader, J.E. Cellular mechanisms underlying intertumoral heterogeneity. Trends Cancer 1, 15–23 (2015).
3. Axlrod, D.E., Miller, N. & Chapman, J.A. Avoiding pitfalls in the statistical analysis of heterogeneous tumors. Biomed. Intrum. Inform. 2, 11–18 (2009).
4. Weaver, W.M. et al. Advances in high-throughput single-cell microtechnologies. Curr. Opin. Biotechnol. 25, 114–123 (2014).
5. Heath, J.R., Ribas, A. & Mishell, PS. Single-cell analysis tools for drug discovery and development. Nat. Rev. Drug Discov. 15, 204–216 (2016).
6. Korey, T., Sarkar, S., Sabhachandani, P. & Cohen, N. Innovative tools and technology for analysis of single cells and cell-cell interaction. Annu. Rev. Biomed. Eng. 18, 259–284 (2016).
7. Taylor, D.L. Past, present, and future of high content screening (HCS): from the beginning. Nat. Methods 15, 720–725 (2010).
8. Gough, A. et al. High-content analysis with cellular and tissue systems biology: a bridge between cancer cell biology and tissue-based diagnostics. In: The Molecular Basis of Cancer. 4th ed. J. Mendelsohn, et al. eds. 369–392 (Elsevier, New York, NY, 2015).
9. Ashworth, T.R. A case of cancer in which cells similar to those in the tumours were seen in the blood after death. Aust. Med. J. 14, 146–149 (1869).
10. Pantel, K., Brakenhoff, R.H. & Brandt, B. Detection, clinical relevance and specific biological properties of disseminating tumour cells. Nat. Rev. Cancer 8, 329–340 (2008).
11. Parkinson, D.R. et al. Considerations in the development of circulating tumor cell technology for clinical use. J. Transl. Med. 10, 138 (2012).
12. Aalto, M., Toner, M., Maheswaran, S. & Haber, D.A. En route to metastasis: circulating tumor cell clusters and epithelial-to-mesenchymal transition. Trends Cancer 1, 44–52 (2015).
13. van de Stolpe, A., Pantel, K., Steijger, S., Terstappen, L.W. & den Toonder, J.M. Circulating tumor cell isolation and diagnostics: toward routine clinical use. Cancer Res. 71, 5955–5960 (2011).
14. Scherer, M.T. & Antonarakis, E.S. Liquid biopsy: clues on prostate cancer drug resistance. Sci. Transl. Med. 7, 312fs45 (2015).
15. Fischer, A.H. Circulating tumor cells: seeing is believing. Arch. Pathol. Lab. Med. 133, 1367–1369 (2009).
16. Marrucci, D. et al. Fluid biopsy in patients with metastatic prostate, pancreatic and breast cancers. Phys. Biol. 9, 016003 (2012).
17. Rodriguez-Lee, M., Kuhn, P. & Webb, D.R. Advanced cancer patient care by integrating circulating tumor cell technology to understand the spatial and temporal dynamics of cancer. Drug. Dev. Res. 75, 384–392 (2014).
18. Dago, A.E. et al. Rapid phenotypic and genomic change in response to therapeutic pressure in prostate cancer inferred by high content analysis of single circulating tumor cells. PLoS One 9, e101777 (2014).
19. Camp, R.L., Chung, G.G. & Rimm, D.L. Automated subcellular localization and quantification of protein expression in tissue microarrays. Nat. Med. 8, 1323–1327 (2002).
20. Critchley-Thorne, R.J., Miller, S.M., Taylor, D.L. & Lingle, W.L. Applications of cellular systems biology in breast cancer patient stratification and diagnostics. Comb. Chem. High Throughput Screen. 12, 860–869 (2009).
21. Barbaud, A. et al. Molecular classification of nonsmall cell lung cancer using a 4–protein quantitative assay. Cancer 118, 1607–1615 (2012).
22. Schubert, W. et al. Analyzing proteome topology and function by automated multidimensional fluorescence microscopy. Nat. Biotechnol. 24, 1270–1278 (2006).
23. Gerdes, M.J. et al. Highly multiplexed single-cell analysis of formalin-fixed, paraffin-embedded cancer tissue. Proc. Natl. Acad. Sci. USA 110, 11982–11987 (2013).
24. Spagnolo, D.M. et al. Pointwise mutual information quantifies intratumor heterogeneity in tissue sections labeled with multiple fluorescent biomarkers. J. Pathol. Inform. 7, 47 (2016).
25. Nelson, D.A. et al. Quantitative single cell analysis of cell population dynamics during submandibular salivary gland development and differentiation. Biol. Open. 2, 439–447 (2013).
26. Clarke, G.M. et al. A novel, automated technology for multiplex biomarker imaging and application to breast cancer. Histopathology 64, 242–255 (2014).
27. Li, C. et al. Excess PLAC8 promotes an unconventional ERK2-dependent EMT in colon cancer. J. Clin. Investig. 124, 2172–2187 (2014).
28. Santagata, S. et al. Taxonomy of breast cancer based on normal cell phenotype predicts outcome. J. Clin. Investig. 124, 859–870 (2014).
29. Critchley-Thorne, R.J., Miller, S.M. et al. A tissue systems pathology assay for high-risk Barrett’s esophagus. Cancer Epidemiol. Biomarkers Prev. 25, 958–968 (2016).
30. Huang, S. Non-genetic heterogeneity of cells in development: more than just noise. Development 136, 3853–3862 (2009).
31. Chang, H.H., Hemberg, M., Barahona, M., Ingber, D.E. & Huang, S. Transcriptome-wide noise controls lineage choice in mammalian progenitor cells. Nature 453, 544–547 (2008).
32. Sisan, D.R., Halter, M., Hubbard, J.B. & Plant, A.L. Predicting rates of cell state change caused by stochastic fluctuations using a data-driven landscape model. Proc. Natl. Acad. Sci. USA 109, 19362–19367 (2012).
33. Brennecke, P. et al. Accounting for technical noise in single-cell RNA-seq experiments. Nat. Methods 10, 1093–1095 (2013).
34. Wu, A.R. et al. Quantitative assessment of single-cell RNA-sequencing methods. Nat. Methods 11, 41–46 (2014).
35. Kharchenko, P., Silberstein, L. & Scadden, D.T. Bayesian approach to single-cell differential expression analysis. Nat. Methods 11, 740–742 (2014).
36. Islam, S. et al. Quantitative single-cell RNA-seq with unique molecular identifiers. Nat. Methods 11, 163–166 (2014).
37. Bailey, R.C., Kwong, G.A., Radu, C.G., Witte, O.N. & Heath, J.R. DNA-encoded antibody libraries: a unified platform for multiplexed cell sorting and detection of genes and proteins. J. Am. Chem. Soc. 129, 1959–1967 (2007).
38. Halter, M. et al. An automated protocol for performance benchmarking a widefield fluorescence microscope. Cytometry A. 85, 978–985 (2014).
39. Langenbach, K.J., Elliott, J.T., Tona, A., McDaniel, D. & Plant, A.L. Thin films of type 1 collagen for cell by cell analysis of morphology and tenascin-C promoter activity. BMC Biotechnol. 6, 14 (2006).
40. Bhadriraju, K. et al. Large-scale time-lapse microscopy of O24 expression in human embryonic stem cell colonies. Stem Cell Res. 17, 122–129 (2016).
41. Shi, Q. et al. Single-cell proteomic chip for profiling intracellular signaling pathways in single tumor cells. Proc. Natl. Acad. Sci. USA 109, 419–424 (2012).
42. Tekab, Z., Kondratovich, M.V. & Mansfield, E. US FDA and personalized medicine: In vitro diagnostic regulatory perspective. Per. Med. 7, 517–530 (2010).
43. Woodcock, J. Assessing the clinical utility of diagnostics used in drug therapy. Clin. Pharmacol. Ther. 88, 765–773 (2010).
44. Parkinson, D.R. et al. Evidence of clinical utility an unmet need in molecular diagnostics for patients with cancer. Clin. Cancer Res. 20, 1429–1444 (2014).
45. Decision Summary: CellSearch Circulating Tumor Cell Kit Model 7900001. US Food and Drug Administration. https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpml/ pmln.cfm?ID=K103502 (2010). Accessed 29 September 2017.
Opportunities and Challenges in Single Cell Analysis for Clinical Translation
Keating et al.

47. Decision Summary: Illumina MiSeqDx Cystic Fibrosis Clinical Sequencing Assay. US Food and Drug Administration. https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfPMN/cfPMN.cfm?ID=K132750 (2013). Accessed 29 September 2017.

48. Decision Summary: Illumina MiSeqDx Cystic Fibrosis 139-Variant Assay. US Food and Drug Administration. https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfPMN/cfPMN.cfm?ID=K124006 (2013). Accessed 29 September 2017.

49. The Clinical and functional translation of CFTR (CFTR2). http://cftr2.org (2011). Accessed 29 September 2017.

50. Genome in a Bottle. National Institute of Standards and Technology. https://www.nist.gov/programs-projects/genome-bottle (2015). Accessed 29 September 2017.

51. Collins, F.S. & Hamburg, M.A. First FDA authorization for next-generation sequencer. N. Engl. J. Med. 369, 2369–2371 (2013).

52. Lazar, D.C. et al. Cytometric comparisons between circulating tumor cells from prostate cancer patients and the prostate-tumor-derived LNCaP cell line. Phys. Biol. 9, 016002 (2012).

53. Cermak, N. et al. High-throughput measurement of single-cell growth rates using serial microfluidic mass sensor arrays. Nat. Biotechnol. 34, 1052–1059 (2016).

54. Stevens, M.M. et al. Drug sensitivity of single cancer cells is predicted by changes in mass accumulation rate. Nat. Biotechnol. 34, 1161–1167 (2016).

55. Ma, C. et al. A clinical microchip for evaluation of single immune cells reveals high functional heterogeneity in phenotypically similar T cells. Nat. Med. 17, 738–743 (2011).

56. Xue, M. et al. Chemical methods for the simultaneous quantitation of metabolites and proteins from single cells. J. Am. Chem. Soc. 137, 4066–4069 (2015).

57. Giesen, C. et al. Highly multiplexed imaging of tumor tissues with subcellular resolution by mass cytometry. Nat. Methods 11, 417–422 (2014).

58. Oppermann, S. et al. High-content screening identifies kinase inhibitors that overcome venetoclax resistance in activated CLL cells. Blood 128, 934–947 (2016).

59. Francis, K. & Palsson, B.O. Effective intercellular communication distances are determined by the relative time constants for cytokine/chemokine secretion and diffusion. Proc. Natl. Acad. Sci. USA 94, 12255–12262 (1997).

© 2018 The Authors. Clinical and Translational Science published by Wiley Periodicals, Inc. on behalf of American Society for Clinical Pharmacology and Therapeutics. This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.