Translational Implications of Tumor Heterogeneity
Mariam Jamal-Hanjani1,2, Sergio A. Quezada1, James Larkin3, and Charles Swanton1,2

Abstract
Advances in next-generation sequencing and bioinformatics have led to an unprecedented view of the cancer genome and its evolution. Genomic studies have demonstrated the complex and heterogeneous clonal landscape of tumors of different origins and the potential impact of intratumor heterogeneity on treatment response and resistance, cancer progression, and the risk of disease relapse. However, the significance of subclonal mutations, in particular mutations in driver genes, and their evolution through time and their dynamics in response to cancer therapies, is yet to be determined. The necessary tools are now available to prospectively determine whether clonal heterogeneity can be used as a biomarker of clinical outcome and to what extent subclonal somatic alterations might influence clinical outcome. Studies that use longitudinal tissue sampling, integrating both genomic and clinical data, have the potential to reveal the subclonal composition and track the evolution of tumors to address these questions and to begin to define the breadth of genetic diversity in different tumor types and its relevance to patient outcome. Such studies may provide further evidence for drug-resistance mechanisms informing combinatorial, adaptive, and tumor immune therapies placed within the context of tumor evolution.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

Editor's Disclosures
The following editor(s) reported relevant financial relationships: J.R. Grandis—None.

CME Staff Planners’ Disclosures
The members of the planning committee have no real or apparent conflicts of interest to disclose.

Learning Objectives
Upon completion of this activity, the participant should have a better understanding of the concept of intratumor heterogeneity and the existing evidence supporting it, as well as its potential implications in clinical practice.

Acknowledgment of Financial or Other Support
This activity does not receive commercial support.

Introduction
Over the past few years, genomic studies have demonstrated the complex and heterogeneous landscape in cancer (1, 2) and its potential implications for treatment response and prognosis (2–4). Using whole-genome or whole-exome sequencing, these studies have demonstrated that tumors consist of somatic events, defined by mutations and copy number alterations (CNA), occurring early in tumor evolution, and somatic events present in some cells, but not all, occurring later in tumor evolution (5). In addition, these studies have shown spatial heterogeneity, branched evolution, and mutational patterns that can vary over time (6, 7) and in response to cancer therapies (8–10). Given the existence of such heterogeneity in tumors in advanced metastatic disease, the efficacy of therapies targeting somatic driver aberrations, even in combination, may be limited in terms of achieving disease cure and contribute in some health economies to a prohibitive health economic burden (11). The relevance of subclonal mutations to cancer outcome and therapeutic response requires further investigation (12). Although this review focuses on genetic intratumor heterogeneity, both stochastic and epigenetic factors are known to play a role in shaping the tumor landscape and are reviewed in-depth elsewhere (13).
Types of Tumor Heterogeneity

There are several types of genetic heterogeneity in cancer biology. The most well-known is interpatient tumor heterogeneity (Fig. 1A), wherein no two patients with the same subtype of tumor behave the same clinically, with or without treatment. This may be related to host factors, such as tumor microenvironment and germline variants influencing treatment response, together with the unique somatic mutations that can occur within the tumor of each individual patient (14). Intratumor heterogeneity (Fig. 1B) describes the existence of distinct cellular populations with specific genetic, epigenetic, and phenotypic features within tumors and has long been recognized (15, 16). It has been described in several tumor types, including lung (7, 17), breast (6, 18, 19), ovarian (20), pancreatic (21, 22), kidney (23, 24), colorectal (25), brain (26, 27), and prostate cancers (28, 29), as well as hematologic malignancies, such as chronic lymphoblastic leukemia (CLL; ref. 30) and acute lymphoblastic leukemia (31). Metastatic lesions at different secondary sites can arise from different cellular populations within a primary tumor, resulting in heterogeneity among metastases, known as intermetastatic heterogeneity (refs. 32–36; Fig. 1C). In addition, since metastatic lesions can acquire new mutations and evolve independently with each cell division, heterogeneity within a metastasis can also exist, known as intrametastatic heterogeneity (ref. 37; Fig. 1D). This can be associated with multiple mechanisms of acquired drug resistance in the same patient with metastatic disease (38–40).

Clonal Evolution and Phylogenetic Analyses

The clonality of somatic mutations within a tumor can be estimated with bioinformatics methods using variables such as tumor purity, allelic copy number, and mutation variant allele frequency (41). The analysis of multiple biopsies from the same tumor can reveal the spatial composition and the evolutionary trajectory of subclones. The clonal and subclonal composition of each tumor can be used to construct distance-based phylogenetic trees, wherein clonal mutations present in all tumor regions occur early in tumorigenesis representing the most recent common ancestor (truncal events on the evolutionary tree) and subclonal mutations present in only a subset of regions, or cells within a single biopsy, occur later in tumorigenesis (branched events on the evolutionary tree; Fig. 2). The accuracy of such clonality analyses is limited to the number of regions sampled and therefore is likely only based on a subset of tumor cells from the entire tumor. An alternative to this approach may be single-cell sequencing, which has the potential to determine the exact number of tumor cells with distinct mutations and copy number variations, and reveal the
known Marusyk and colleagues demonstrated that subclones, without a growth and malignant potential. Using a mouse xenograft model, not always be the case that the dominant clone dictates tumor heterogeneity in predicting adverse outcomes and the evolution of treatment. Studies like this indicate the likely relevance of tumor evolution, such that the extent of heterogeneity evolved during risk factor for disease progression, and those patients treated with found that the presence of a subclonal driver was an independent driver mutations corresponding to CLL evolution (30). They exome sequencing and CNA to identify clonal and subclonal but not all (B and C) occur later in tumorigenesis, represented by the yellow branches of the tree; and private alterations (D–F) present in only one region of the tumor also occur later in tumorigenesis, represented by the red branches of the tree.

subclonal composition of a tumor although such methods are still limited by allelic drop out and sequencing artifacts.

**Tumor Heterogeneity and Cancer Progression**

In a recent study of CLL, Landau and colleagues used whole-exome sequencing and CNA to identify clonal and subclonal driver mutations corresponding to CLL evolution (30). They found that the presence of a subclonal driver was an independent risk factor for disease progression, and those patients treated with cytotoxic chemotherapy were more likely to undergo clonal evolution, such that the extent of heterogeneity evolved during treatment. Studies like this indicate the likely relevance of tumor heterogeneity in predicting adverse outcomes and the evolution of tumor subclonal composition during treatment. Therefore, it may not always be the case that the dominant clone dictates tumor growth and malignant potential. Using a mouse xenograft model, Marusyk and colleagues demonstrated that subclones, without a known fitness advantage, could drive tumor growth in a non–cell-autonomous manner by inducing tumor-promoting changes in the microenvironment (42). Furthermore, non–cell-autonomous subclones could be outcompeted by other subclones with greater proliferative potential resulting in tumor collapse. Observations where minority subclones influence progression of the tumor mass suggest challenges for predictive and prognostic biomarker discovery efforts that have traditionally focused on identifying genomic alterations in the dominant clone as well as the need to fully understand subclonal interactions within a complex ecological framework. The mechanisms by which subclonal alterations have an impact on tumor biology and phenotype are yet to be determined and will require further functional genomic studies. Such studies will need to investigate the interactions, both synergistic and antagonistic, between subclones during tumor evolution and specifically how these relationships may promote or impede cancer progression and contribute to therapeutic failure and drug resistance (12). Heterogeneity in the tumor microenvironment may also influence the evolution and progression of tumors. Interactions between tumor and stromal cells, changes in the level of hypoxia or acidity, increased or decreased inflammatory cell infiltrate, and remodeling of the extracellular matrix may act as selection pressures and lead to increased phenotypic heterogeneity, potentially influencing treatment response and therefore tumor evolution (for a review, please refer to ref. 13).

**Therapeutic Failure and Resistance**

Heterogeneous tumors are composed of multiple subclones and under selection pressures, such as chemotherapy, subclones with either intrinsic or acquired resistance can be selected for, allowing these subclones to dominate a tumor mass and potentially drive disease progression (refs. 43, 44; Fig. 3). The selection of resistant subclonal populations as a result of therapy leading to treatment resistance has been shown in several tumor types, including lung (45, 46), colorectal (10), gastrointestinal (47), and brain (48) tumors as well as chronic myeloid leukemia, among others (49).

The mechanisms by which a tumor develops resistance may involve multiple somatic events affecting distinct signal transduction pathways, resulting in multiple distinct drug resistance events within the same patient’s tumor burden (50), which poses considerable challenges for the design and selection of effective drug combinations. However, there is evidence for phenotypic convergence within and across tumor types, suggesting that genetic events driving resistance and disease progression may focus on either one or several signaling pathways that may be therapeutically targetable (5, 23, 24, 48, 51–53).

Alternative therapeutic strategies may have the potential to overcome the challenges posed by clonal heterogeneity. Gatenby and colleagues have proposed the concept of adaptive therapy, whereby cancer treatment is continuously adapted in order to maintain a fixed population of drug-sensitive cells, which can in turn suppress the growth of drug-resistant cells, and therefore allow the overall
Intratumor Heterogeneity

Figure 3.
Intratumor heterogeneity and clonal evolution. Primary tumors consisting of different subclones may be subjected to various selection pressures, including chemotherapy and microenvironmental factors such as hypoxia, infiltrating stromal, and immune cells. Under the influence of such selection pressures, subclones with intrinsic resistance (green subclone) can outgrow a tumor mass, potentially leading to disease progression, and/or subclones can acquire somatic alterations (purple subclone), promoting cell survival, proliferation, and metastatic tumor formation. The outgrowth of some subclones (red subclone) may be constrained by selection pressures that they are sensitive to, for example, targeted therapy against a tumor subclone with a somatic alteration sensitive to therapy. TILs may recognize neoantigens presented on the surface of tumor cells as nonself, promoting enhanced T-cell activation and immune cell tumor infiltration.

tumor burden to remain stable (54). Knowledge of the existence of low-frequency resistant subclones at diagnosis may allow for the use of combined therapeutic regimens targeting mechanisms of resistance. Evidence for the parallel evolution of subclones during tumor evolution suggests that there exist constraints to tumor growth that might be exploitable. If evolutionary “rule books” of tumor progression could be deciphered, by predicting the likely next step in the evolution of the tumor, one might be able to devise preventative measures and potentially delay the onset of cancer progression (5). Such tactics will require a good evidence base from studies in different tumor types involving longitudinal tissue sampling, and therapeutic trial designs will need to take into account alternative approaches, such as adaptive therapy.

Host Immunity and Tumor Neo-antigens

The interplay between a tumor and the host immune system can help determine the immunogenicity of a tumor (55). The process of immunosurveillance can select for subclones lacking immunogenic antigens (56) or subclones with reduced sensitivity to immune attack (57) allowing for clonal evolution and tumor progression, a process termed immunoeediting (58). Tumors can also create a microenvironment that actively counteracts the host antitumor immune response by using suppressive mechanisms such as the expression of programmed cell death ligand 1 (PD-L1), the expression of indolamine 2,3-dioxygenase, secretion of anti-inflammatory mediators (i.e., TGFβ) or the induction of T-cell anergy (59). Alternatively, tumor cells can attract cell types that suppress invading immune cells and support tumor growth and survival, such as regulatory T cells, tumor-associated macrophages (TAM), cancer-associated fibroblasts (CAF), and myeloid-derived suppressor cells (MDSC). Immunotherapeutic drugs, such as monoclonal antibodies and cancer vaccines, have become a relevant player in the landscape of cancer therapies (60), and compared with conventional therapies, immunotherapy benefits from its ability to induce epitope spreading and immunologic memory, which may significantly contribute to preventing disease relapse. In particular, the modulation of immunoregulatory checkpoints has recently been illustrated in clinical trials in which antibody-mediated blockade of the immune inhibitory receptors programmed cell death protein-1 (PD-1), programmed cell death ligand-1 (PD-L1), or cytotoxic T-lymphocyte–associated protein 4 (CTLA-4) produced significant clinical benefits in a variety of cancers, including metastatic melanoma (61–63).

Although evidence suggests that intratumor heterogeneity may limit the efficacy of conventional and targeted therapies, on account of factors such as sampling bias and clonal evolution (53), increased mutational diversity may lead to increased tumor neo-antigen production, and therefore potential new targets for immunotherapeutic drugs, such as immune checkpoint inhibitors (53, 64–66). Somatic tumor mutations resulting in mutant short peptide fragments (neo-antigens) are presented on the tumor cell surface by MHC, promoting activation and expansion of tumor-infiltrating lymphocytes (TIL; Fig. 3). TILs, in particular CD8+ T cells, have been shown to be associated with improved treatment response and prognosis in several tumor types (67), including breast (68), ovarian (69–71), melanoma (72), and lung (73). The immunogenicity of a mutant peptide depends on its affinity for binding MHC class I ligands so that it can be presented to, and recognized by, CD8+ T cells (74). Conceivably, the greater the extent of intratumor heterogeneity and mutational burden, the greater the repertoire of potentially exploitable neo-antigens.
within a tumor (75). Using RNA-sequencing data for several tumor types from The Cancer Genome Atlas, Brown and colleagues identified immunogenic mutations predicted to result in mutational epitopes with high affinity for MHC class I ligands and found that patients with predicted immunogenic mutations compared with those without had significantly improved overall survival (76). In a study of patients with melanoma treated with CTLA-4 blockade (ipilimumab or tremelimumab), Snyder and colleagues identified candidate tumor neo-antigens for each patient and validated a neo-antigen signature associated with a strong treatment response (77). Such studies have the potential to use the neo-antigen landscape of heterogeneous tumors to predict immunotherapy response and therefore aid treatment stratification.

**Insight into Tumor Heterogeneity Using "Liquid Biopsies"**

Serial tumor sampling poses practical challenges and is currently not a standard practice. An alternative approach may be the use of “liquid biopsies,” whereby circulating cell-free tumor DNA (cfDNA) or circulating tumor cells (CTC) are analyzed in the peripheral blood of patients with cancer. Such sampling may offer a relatively easy and noninvasive method for the analysis of primary and metastatic tumors (78). Several studies have demonstrated that genetic alterations in cfDNA are also present in matched tumors and can be used to track mutational burden over time (10, 28, 79–82). Structural genomic rearrangements, such as translocations (83), as well as gene amplifications (84), have also been identified in cfDNA. In a study by Murtaza and colleagues, serial sampling and exome sequencing of plasma in 6 patients with advanced-stage breast, ovarian, and lung cancer was performed during several lines of therapy (79). By quantifying the allele frequencies of mutations in the plasma, they were able to correlate an increased representation of mutant alleles with emerging therapy resistance, including a T790M mutation following gefitinib therapy, a truncating mutation in mediator complex subunit 1 (MED1) following treatment with tamoxifen and trastuzumab, and a truncating mutation in RBI (retinoblas- toma 1) following treatment with cisplatin chemotherapy. Bettegowda and colleagues detected cfDNA in a broad spectrum of both early- and late-stage cancers in a total of 640 patients (10, 85). Forty-seven percent of patients with stage I cancers of any type had detectable levels of cfDNA, indicating the potential use of cfDNA in early detection of cancer. In a subset of patients with colorectal cancer receiving anti-EGFR antibody therapy who subsequently relapsed, they were able to detect the emergence of several mutations in genes involved in the MAPK pathway, and therefore potential mechanisms of resistance, by genetically profiling cfDNA using digital PCR-based technologies. In a study by Diehl and colleagues, cfDNA was used to follow the course of therapy in patients with colorectal cancer (80). Fluctuations in cfDNA after surgery corresponded with the extent of surgical resection, and patients with a detectable cfDNA after surgery tended to relapse within 1 year. Compared with the standard biomarker carcinoembryonic antigen (CEA), cfDNA was found to be more reliable. Similarly, in a study by Dawson and colleagues, cfDNA was found to be a more reliable indicator of tumor burden compared with the standard biomarker CA 15-3 (carcinoma antigen 15-3; ref. 82). Rothé and colleagues used next-generation sequencing of cancer gene panels to analyze matched primary and metastatic tumors as well as cfDNA samples in patients with breast cancer (86). In 76% (13 of 17) of cases, mutations found in the cfDNA were concordant with the tumor, indicating that in some cases, the cfDNA could be used as an alternative to metastatic tumor biopsies.

Similar to the studies in cfDNA, CTCs have also been used to study mutations and CNAs (87–91) and have been shown to correlate with clinical outcome. In a large prospective study by Rack and colleagues, CTC enumeration in patients with breast cancer was found to be an independent prognostic marker for disease-free survival (DFS) and overall survival (OS; ref. 92). The worst prognosis was seen in patients with at least 5 CTCs per 30-mL blood sample, and the presence of CTCs after cytotoxic chemotherapy was associated with a poorer DFS and OS. One of the advantages of studying CTCs is that next-generation sequencing can allow for the detection of intratumor heterogeneity in terms of driver mutations at single-cell resolution, for example, heterogeneity in EGFR expression between CTCs in the same patient (93). Another advantage is the ability to study CTCs in the context of patient-derived mouse models. In a study by Hodkinson and colleagues, CTCs were found to be tumorigenic in immunocompromised mice (90). CTCs derived from patients with small cell lung cancer were implanted into mice resulting in CTC-derived explants (CDX; ref. 90). The genomic profiles of CTCs derived from patients and CDXs showed considerable similarity. Interestingly, CDXs reflected the donor patient’s response to platinum and etoposide chemotherapy, suggesting that such mouse models could be used for therapy testing and increase our understanding of potential resistance mechanisms.

However, although CDXs, as well as patient-derived tumor xenografts, may be effective tools in studying intratumor heterogeneity, there are caveats to consider (for a review, please refer to ref. 94). In a study by Klc and colleagues, subclones derived from peripheral blood in patients with acute myeloid leukemia were injected into immunocompromised mice (95). To assess the xenograft subclonal composition, human leukemic cells were purified from bone marrow xenografts using flow cytometry for subsequent whole-genome sequencing. All xenografts had a skewed subclonal composition where discreet subclones, despite a low frequency in the injected sample, defined the engrafted cell population resulting in unequal engraftment and/or growth potential, such that none of the xenografts had a subclonal composition that was identical to the input sample. The engrafted cell population may also be dependent on the mouse strain used, as immunophenotypic variability, based on flow cytometry using human antibodies to CD45, CD33, or CD34, was observed between different strains of mice in e18.1 identical acute myeloid leukemia samples were injected. It therefore follows that the interpretation of the clonal composition in PDXs requires some caution, as these models may not represent the genetic heterogeneity of the human-derived cell population and may not mirror the patient’s tumor clonal evolution in parallel longitudinal studies. In addition, one must take into account the potential for acquired mutations during the passage of human cells into immunocompromised mice (96).

Liquid biopsies have the potential to inform early detection of cancer, detect minimal residual disease, and track evolution of resistant disease and therefore detect early relapse (for a review, please refer to ref. 97). Further evidence supporting the use of liquid biopsies to detect early-stage cancer may inform cancer.
screening strategies and allow the detection of tumors in early evolution. Such strategies could aid early therapeutic intervention before the onset of increased intratumor heterogeneity. In addition, they may be of use in circumstances where the biopsy of certain relapse sites, such as brain or bone, is not easily accessible. However, as cfDNA and CTCs may represent cancer genomes derived from multiple metastatic sites within the body, the interpretation of such biopsies may be confounded by intratumor heterogeneity (78). Whether whole-genome or -exome sequencing compared with targeted deep sequencing (98) is the preferred method, or exactly how representative liquid biopsies are of the underlying tumor landscape, and how sensitive they are in identifying clonal and subclonal alterations, is yet to be fully determined (99). Although not discussed in this review, it is worth noting that tumor DNA may also be detected and analyzed in samples such as urine, stool, and circulating cerebrospinal fluid.

Implications for Precision Medicine and Clinical Trial Development

Given the evidence for spatial intratumor heterogeneity and the evolution of tumor subclones with time and in response to cancer therapies, the potential sampling bias from a single tumor biopsy representing a snapshot in time may limit the ability to identify and qualify biomarkers for clinical use. Ideally, predictive biomarkers should be identified on the basis of the genetic and phenotypic profile of each tumor, alongside the development of new therapies. Several recent studies are modeling such an approach, including the BATTLE-2 (NCT01248247) and Lung-MAP (NCT02154490) studies. To determine the true impact of intratumor heterogeneity on clinical outcome, future therapeutic studies aimed at delivering precision medicine may need to identify biomarkers in the context of tumor spatial heterogeneity. Because potential drivers of disease may vary during tumor evolution and treatment, and the presence of subclonal drivers may affect the efficacy of targeted therapies, potential drug combinations may need to address efficacy in the context of tumor clonal composition (100). In addition, bioinformatics analyses will need to capture low-frequency driver events, which may predict for disease relapse and progression.

Taking multiple tumor biopsies to truly determine the clonal composition of tumors is not a simple or practical solution outside the context of clinical studies, but it is an effective approach to understand how tumors evolve during the disease course and in response to treatment. Longitudinal studies involving both tissue and liquid biopsies may allow us to develop cancer evolutionary “rule books” in an attempt to predict the disease course and likely beneficial therapeutic interventions. Such studies can also give us the opportunity to characterize exceptional or poor responders to treatment, which may be of benefit to patients with similar genetic or phenotypic patterns.

While precision medicine has the potential to improve clinical outcome, reduce toxicity, and increase cost effectiveness, it is not without its challenges, including access to sequencing equipment and its associated cost implications, the development of clinical trials involving adequate tumor profiling to identify actionable alterations, the availability of targeted drugs, and the validation of predictive and prognostic biomarkers. The appropriate infrastructure for such studies needs to be in place providing adequate computing resources and approved sequencing technologies. Clinical practice will need to move toward the implementation of genomics and detailed molecular characterization of a patient’s tumor to enable tailored therapies and attempt to improve patient outcomes. Furthermore, clinicians will need to adapt to the challenges in analyzing such data and the ethics of communicating increased risk of false-positive and false-negative results (101). Ultimately, our aim should be to develop economically feasible predictive and prognostic biomarkers to aid clinical decision making and improve risk stratification and to work toward significant, as opposed to marginal, improvements in outcomes using patient-centered measures such as quality-adjusted lifetimes. Possible approaches in achieving such goals have been reviewed in depth elsewhere (11).

Conclusions

Increasing evidence for intratumor heterogeneity and its potential impact on clinical outcomes requires further exploration of the molecular mechanisms driving genomic instability and clonal evolution in cancer (102). Prospective genomic studies have the potential to allow us to gain a deeper insight into the true extent of tumor heterogeneity and how this evolves during the disease course and in response to therapeutic intervention. Furthermore, understanding the phylogenetic relationship between primary and metastatic tumors, and the significance of subclonal driver mutations, may allow clinicians to make informed therapeutic choices treating patients with drugs tailored to their tumor profile, taking into account tumor clonal composition. An example of such a study is TRACERx [TRArrcking non-small cell lung Cancer Evolution through therapy (Rx)], NCT01888601, in which multiregion and longitudinal tumor sampling and sequencing is performed from diagnosis to relapse, to define the genomic landscape of tumors and to understand the impact of tumor clonal heterogeneity on therapeutic and survival outcome (103). Tumors within TRACERx are genetically profiled to identify the most immunogenic and targetable mutational epitopes and to determine the relationship between the extent of intratumor heterogeneity and TILs and clinical outcomes. In determining whether circulating biomarkers can truly represent the tumor clonal composition, this study also involves the serial sampling of cfDNA and CTCs at different disease time points and in doing so explores the use of circulating biomarkers in tracking clonal evolution, detecting minimal residual disease and early relapse. In light of emerging evidence that driver mutations may be subclonal in some tumors, there is a need to define the relationship between driver clonality and benefit from targeted therapy exposure in prospective clinical studies, such as the DARWIN (Deciphering Anti-tumour Response With InTumour Heterogeneity; NCT02183883) clinical trials program, which may help determine the impact of subclonal drivers on therapeutic response, disease relapse, and progression (32, 103).

Collaborations between industry, scientists, and clinicians are required to meet the challenges posed by future clinical trials designed to address intratumor heterogeneity. Such studies are likely to require repeated tissue and blood sampling and infrastructural support addressing financial, technological, and regulatory aspects to trial design in light of complex subclonal architectures. With the integration of large cohort clinical and genomic data and the development of bioinformatics analyses of such large
datasets, the delivery of precision medicine has the potential to become a reality with the ultimate aim of improving patient outcomes.

Grant Support

M. Jamal-Hanjani and S.A. Quezada are Cancer Research UK Career Development Fellows. J. Larkin is supported by the Royal Marsden Hospital/Institute of Cancer Research and the National Institute for Health Research Biomedical Research Centre for Cancer. C. Swanton is a senior Cancer Research UK Clinical Research Fellow and is supported by Cancer Research UK, the National Institute for Health Research University College London Hospitals Biomedical Research Centre for Cancer, the Rosetrees Trust, EU FP7 (projects PREDICT and RESPONSIFY, ID: 259303), the Prostate Cancer Foundation, the European Research Council, and the Breast Cancer Research Foundation.

Received November 5, 2014; revised January 5, 2015; accepted January 5, 2015; published online March 13, 2015.
Intratumor Heterogeneity

Translational Implications of Tumor Heterogeneity

Access the most recent version of this article at:
http://clincancerres.aacrjournals.org/content/21/6/1258

This article cites 102 articles, 28 of which you can access for free at:
http://clincancerres.aacrjournals.org/content/21/6/1258.full#ref-list-1

This article has been cited by 6 HighWire-hosted articles. Access the articles at:
http://clincancerres.aacrjournals.org/content/21/6/1258.full#related-urls

Sign up to receive free email-alerts related to this article or journal.
To order reprints of this article or to subscribe to the journal, contact the AACR Publications Department at pubs@aacr.org.
To request permission to re-use all or part of this article, contact the AACR Publications Department at permissions@aacr.org.