Molecular and cellular heterogeneity: the hallmark of glioblastoma

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Glioblastoma is the most common primary malignant brain tumor in adults and continues to portend poor prognosis despite decades of research. Even with aggressive resection followed by concomitant chemotherapy and radiation, there is a high recurrence rate with median survival of less than 15 months. Indeed, fewer than 5% of patients survive 5 years. Extensive investigation of the cellular and molecular biology of glioblastoma over the last decade has identified several histopathological and chromosomal hallmarks that have enhanced diagnosis and clinical stratification without a significant improvement in overall outcome. Nevertheless, a deepening focus on the molecular characteristics of glioblastoma has revealed that this histopathological term comprises a range of diseases, some of which have divergent natural histories. Indeed, it is becoming clear that malignant glioma exhibits striking cellular and molecular heterogeneity not only across but also within glioblastomas. In fact, a patient with glioblastoma likely harbors several different diseases within the same tumor such that each tumor is actually a complex and dynamic constellation of cellular and molecular changes that contribute to disease pathobiology.

Cellular and molecular heterogeneity is not unique to glioblastoma and has been extensively studied in several other solid tumors. Next-generation sequencing of subregions and even single cells within breast, pancreas, prostate, and renal cancer have demonstrated extensive regional clonality within primary tumors, where only one-third of mutations, known as founding mutations, occur ubiquitously in all cells. Remarkably, even minor subclones from a primary tumor may become treatment-refractory clones that dominate the recurrence. This concept of clonal evolution has been firmly established in acute myeloid leukemia, where deep sequencing has revealed that subclones representing as little as 5% of the total tumor cells can survive treatment, acquire new mutations, and lead to relapse. In the related context of metastatic disease, deep sequencing of driver mutations identified in genomic studies of medulloblastoma revealed that while distinct metastatic clones within a given patient are genetically similar, they are divergent from the primary tumor, suggesting clonal enrichment and/or evolution.

Key Words • glioblastoma • glioma • genomic • heterogeneity • IDH1

Abbreviations used in this paper: G-CIMP = glioma CpG island methylator phenotype; H3K27Ac = histone H3 Lysine27 acetylation; RTK = receptor tyrosine kinase; SCNA = somatic copy number alteration; TCGA = The Cancer Genome Atlas; TIC = tumor-initiating cell; 2-HG = 2-hydroxyglutarate.

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It is clear from these studies in other cancers and a quickly accumulating body of literature in glioblastoma that an improved understanding of glioblastoma heterogeneity has important implications not only for the direction of future scientific investigation, but also for clinical diagnosis and patient management. Herein, we review the salient themes of tumor heterogeneity on the molecular and cellular levels and highlight the critical studies germane to neurosurgeons and neuro-oncologists that are driving our growing understanding that diversity and heterogeneity in malignant glioma are the rules, rather than the exceptions (Table 1).

Molecular Heterogeneity

Molecular Classification: Diversity Across Tumors at the RNA Level

Transcriptomic Subtypes. Commitment by cancer biologists to understanding the genomic basis of glioblastoma has yielded ever higher-resolution pictures of the types of alterations that glioblastomas harbor. These studies have been central to the ongoing revolution in cancer genomics that is revealing the molecular origins of every cancer type and establishing a new disease taxonomy founded on shared and/or recurrent genomic lesions. Broadly speaking, this sea change in cancer genomics has been possible due to our ability to perform genome-wide profiling of tumor DNA and RNA in a systematic fashion. In the case of glioblastoma, perhaps one of the greatest influences in our early understanding of glioma heterogeneity was the application of transcriptional profiling to brain tumor classification. Transcriptional profiling, pioneered by Golub et al. and Alizadeh and colleagues, has provided powerful evidence demonstrating the striking heterogeneity that exists among glioblastomas arising in different patients. Among many, 2 studies in particular have provided our working foundation of glioblastoma subtypes defined by transcriptomic profile. Phillips et al. identified 3 transcriptional subtypes—proneural, mesenchymal, and proliferative—based on expressed genes that were most strongly correlated with survival. Complementing this work, Verhaak et al. employed unsupervised clustering approaches to classify 200 glioblastomas into 4 subtypes—proneural, mesenchymal, classical, and neural. As discussed below, these subtypes also harbor distinct DNA alterations, further emphasizing intertumoral heterogeneity and distinct pathways of tumor evolution. Importantly, although the specific number of subtypes defined by these 2 studies differs, the proneural and mesenchymal classifications appear to be the most robust and concordant. Specifically, both studies identified expression of proneural genes such as DLL3 and OLIG2 as well as mesenchymal class expression of CD40 and CHI3L1/YKL-40. Further work has attempted to develop prognostic assays based on a limited set of genes that correlates with survival, which is compelling but has not yet yielded completely overlapping gene sets across studies. Nevertheless, the study of gene expression programs and signatures has been pivotal to the evolution of a conceptual framework that glioblastoma development is variable from patient to patient—i.e., the “subtype” heuristic—and surely will continue to drive key biological insights. Although beyond the scope of this review, there has also been a surge of interest in the epigenetic landscape of glioblastoma, and further understanding of these structural epigenetic phenotypes will likely yield similarly important insights into intertumoral heterogeneity and natural history.

Molecular Classification: Diversity Across Tumors at the DNA Level

The landmark glioblastoma expression mRNA profiling studies have been complemented by equally pivotal DNA profiling studies. To date, the glioblastoma genome is one of the most comprehensively annotated among all cancer types, and the extensive work in this area has highlighted many of the recurrent pathways that are altered in these tumors. At the DNA level, glioblastoma is similar to other cancers in that there are gains and losses of gene regions and even whole chromosomes; these somatic copy number alterations (SCNAs) have been cataloged by a number of groups using sophisticated computational methodologies such as Genomic Identification of Significant Targets in Cancer (CISTIC) and Genomic Topography Scan. Together, these studies identified commonly amplified genes such as EGFR, MET, PDGFR, MDM2, PIK3CA, CDK4, and CDK6 as commonly deleted genes including CDKN2A/B, PTEN, and RBL. In addition to clarifying the gains and losses of DNA segments, more recent work has focused on the alterations in glioblastoma at the single nucleotide level using sequencing approaches. There is no question that the application of next-generation sequencing approaches to the study of cancer has fueled a punctuated leap in our understanding of the genomic landscape across all cancer types. Work from the Vogelstein group and The Cancer Genome Atlas (TCGA) first demonstrated the types and frequencies of mutations seen in glioma, albeit using distinct approaches. While both provided significantly overlapping information, Parsons et al. identified recurrent mutations in the IDH1 gene, which was a seismic finding to be discussed further below, while work from the TCGA paper also provided a proof-of-principle showing that systematic, multiparametric profiling of tumor types was possible. These studies were a prelude to the comprehensive, nearly encyclopedic anthology of the glioblastoma genome published by TCGA working group in 2013. This study, analyzing more than 500 glioblastomas, integrated information from sequencing, SCNA analysis, transcriptomic analysis, epigenetic profiling, and rearrangement studies to provide a breathtaking molecular picture of these tumors. Recurrent mutations were identified in genes such as PTEN (31%), TP53 (29%), EGFR (26%), PIK3R1 (11%), PIK3CA (11%), and IDH1 (5%). In addition, TERT promoter mutations were identified in the majority of tumors assessed by whole genome sequencing, corroborating previous findings. Novel mutations in genes such as LITF, SPTA1, and TCHH were also identified. Most importantly, the systematic TCGA profiling of a large number of tumors enabled high-level integration of structural genomic information that provided a more
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The IDH1 Story: Not all Gliomas Are Created Equal

If there is a single mutation that underscores the idea that not all glioblastomas are created equally, it is the mutation found in the IDH1 gene.22,23 IDH1 encodes isocitrate dehydrogenase 1, a cytoplasmic enzyme that catalyzes the oxidative decarboxylation of isocitrate to α-ketoglutarate and NAD+ to NADPH.50 Recurrent mutations in IDH1 were first identified by Parsons et al. in 2008 in a study in which 5 of 22 tumors in the study’s discovery set harbored a single mutation encoding the IDH1 R132H mutant protein. Extensive work by many groups has since demonstrated how important the distinction between “IDH1 mutant” and “IDH1 wild type” may be. In brain tumors, IDH1 mutations are typically R132H (90%), although other substitutions at position R132 are also found.69 IDH1 mutations are identified in 60%–80% of Grade II and III astrocytomas, oligodendrogliomas, and oligoastrocytomas.23,28,29 Whereas 5%–6% of primary glioblastomas harbor IDH1 mutations, the majority of secondary glioblastomas carry this mutation. Similar mutations in the IDH2 gene often seen in acute myeloid leukemia are rarely found in glioma. Interestingly, mutations in the IDH genes have also been identified in cholangiocarcinomas and cartilaginous tumors.23

The foundation of all efforts to understand IDH1 mutant protein biology is the work of Dang and colleagues.20 This study used a liquid chromatography/mass spectrometry metabolomics approach to discover that IDH1-R132H mutant proteins acquire a neomorphic function that leads to the production of high levels of the “R” enantiomer of 2-hydroxyglutarate (2-HG) that are never seen at comparable levels in an IDH1 wild-type setting and can be detected using MR spectroscopy.2 Because tumor cells typically harbor one copy of IDH1 mutation and do not exhibit loss of heterozygosity at that locus, these mutations are likely gain of function in nature. Significant effort is underway to understand the mechanistic underpinnings of the IDH1 mutant state. Although 2-HG

TABLE 1: Molecular and cellular heterogeneity in glioblastoma* (continued)

| Francis et al., 2014 | Structural DNA alterations between cells. Single-cell whole genome sequencing analysis of multiple EGFR aberrations within the same tumor identified different clonal events. EGFR truncation variants segregated into nonoverlapping subclonal populations. Patel et al., 2014 | Glioblastoma subtype classifiers were variably expressed across individual cells within a single tumor. Soda et al., 2011, & Turcan et al., 2012 | RTK-amplified subpopulations share a common early precursor with genetic alterations such as homozygous deletion of CDKN2A. A CD133+ cell subpopulation from human brain tumors exhibited stem cell-like properties and initiated tumors in vivo. Singh et al., 2004 | Bhat et al., 2013 | The mesenchymal TIC signature, CD44 expression, and NF-κB activation correlate with relative radioresistance and shorter survival. Suvà et al., 2014 | TIC population program that defines TIC identity. A core set of neurodevelopmental transcription factors (POU3F2, SOX2, SALL2, & OLIG2) coordinately activate an epigenetic information set harbored a single mutation encoding the IDH1 R132H mutant protein. Extensive work by many groups has since demonstrated how important the distinction between “IDH1 mutant” and “IDH1 wild type” may be. In brain tumors, IDH1 mutations are typically R132H (90%), although other substitutions at position R132 are also found.69 IDH1 mutations are identified in 60%–80% of Grade II and III astrocytomas, oligodendrogliomas, and oligoastrocytomas.23,28,29 Whereas 5%–6% of primary glioblastomas harbor IDH1 mutations, the majority of secondary glioblastomas carry this mutation. Similar mutations in the IDH2 gene often seen in acute myeloid leukemia are rarely found in glioma. Interestingly, mutations in the IDH genes have also been identified in cholangiocarcinomas and cartilaginous tumors.23 |
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has been shown to inhibit the α-ketoglutarate-dependent superfamily of dioxygenases affecting histone and DNA methylation,\textsuperscript{18,67} likely contributes to the G-CIMP phenotype,\textsuperscript{16} and also alters HIF-1α biology.\textsuperscript{23,35} Further work is necessary to clarify the causal mechanistic basis of mutant IDH1 biology.

Accumulating evidence suggests that IDH1 mutant and IDH1 wild-type tumors should perhaps be considered different tumors, albeit with similar histopathology. Patients with IDH1 mutant tumors are nearly 2 decades younger than patients with IDH1 wild-type tumors and exhibit an overall survival that can be up to 3-fold greater than patients with wild-type tumors.\textsuperscript{35,46} This survival advantage is even more amplified in lower-grade astrocytomas. Additionally, IDH1 mutant tumors appear to harbor distinct imaging characteristics: they often enhance less with gadolinium contrast, harbor less peritumoral edema, exhibit increased cystic components, and have an increased incidence of frontal lobe involvement.\textsuperscript{14,37} It is likely that the IDH1 mutation is a primordial alteration in tumor development. In recent work from the Costello group, all IDH1 mutant low-grade gliomas that recurred still harbored IDH1 mutations despite substantial differences in other tumor-associated genes,\textsuperscript{35} suggesting that this alteration occurs in the tumor’s cell of origin. Ultimately, these distinct clinical characteristics have raised the possibility that IDH1 status may influence how mutant tumors are treated. Indeed, Beiko et al. showed that resection of total IDH1 mutant malignant astrocytoma volume—i.e., both T1 enhancing and T2 hyperintense areas—was associated with a median survival longer than 9 years; in comparison, the only survival benefit in IDH1 wild-type malignant astrocytomas was associated with resection of T1 enhancing disease.\textsuperscript{3} This study is perhaps the first to demonstrate that an understanding of tumor heterogeneity in glioblastoma is not just a basic science interest but may significantly influence how we surgically manage glioblastomas from patient to patient.

**Intratumoral Heterogeneity: Tumors Within Tumors**

The realization that tumors can vary structurally from patient to patient has been complemented by additional recent work demonstrating that there is much more to tumor heterogeneity than we had anticipated. Specifically, recent studies have demonstrated that there is substantial subclonality within each tumor in a manner that suggests that patients harbor glioblastomas that resemble admixtures of molecularly distinct tumors rather than a uniform disease. In particular, studies that have focused on the distribution of alterations of receptor tyrosine kinases have provided a window into the intratumoral diversity seen in glioblastoma. Simply stated, it is becoming clear that glioblastoma is a composite of multiple tumors within the same tumor, and the extent to which we deeply understand this intrapatient heterogeneity will have significant therapeutic implications.

**Receptor Tyrosine Kinase Heterogeneity: The Subclonal Landscape.** The study of both the activation states and the distribution of SCNA in receptor tyrosine kinases (RTKs) in glioblastoma has provided powerful evidence of the subclonal landscape within these tumors. Receptor tyrosine kinases are highly dysregulated in glioma, with high-level amplification observed in RTKs such as EGFR (60%–70%), PDGFRA (12%–15%), and MET (5%).\textsuperscript{11} However, monotherapy with RTK inhibitors has been inefficacious to date in this tumor type.\textsuperscript{22} It has been demonstrated previously that the EGFRvIII variant, which is found in up to 20% of glioblastomas,\textsuperscript{11} is heterogeneously expressed across EGFRvIII-positive tumors,\textsuperscript{43} suggesting that its true contribution to gliomagenesis stems from paracrine mechanisms.\textsuperscript{31,42} In addition, Stommel et al. demonstrated that RTKs likely cooperate to drive mitogenic signaling in glioblastoma.\textsuperscript{32} Specifically, the majority of glioblastoma cell lines analyzed exhibited coactivation of 3 or more RTKs. Moreover, immunofluorescence staining with phosphospecific antibodies for multiple RTKs revealed variable coexpression of activated RTKs among individual cells derived from a single primary glioblastoma tumor. Thus, there is promiscuous RTK activation within a single glioblastoma tumor, although the etiology for this coactivation state is not known.

Recent work provided further compelling evidence of clonal heterogeneity within a single tumor on the structural genomics level. Both Snuderl and Szerlip and their colleagues discovered that nearly 13% of glioblastomas carrying amplifications of EGFR, MET, or PDGFRA harbored multiple copy number gains in these genes.\textsuperscript{34,61} Exclusivity analysis revealed that the predominant pattern was mosaic amplification of these RTKs rather than coamplification within a single cell. This mosaic pattern of RTK amplification was graphically illustrated in a remarkable figure in which fluorescence in situ hybridization analysis for EGFR and PDGFRA was performed on a coronal section of a whole brain autopsy specimen from a patient with infiltrative left parietal glioblastoma. Whereas the large left parietal mass harbored EGFR-amplified and PDGFRA-amplified cells, the majority of cells within the infiltrated right contralateral tumor were EGFR amplified. These studies therefore provide clear examples of the subclonal architecture in glioblastoma and suggest that there may be a high potential for greater diversity when we begin to explore other molecular features beyond RTK alterations.

**Heterogeneity Identified by Analyzing Single Cells.** The future of heterogeneity analysis will be conducted at the single cell level. Although TCGA and other genomic studies to date have been incredibly informative, they were conducted on bulk tumor DNA samples and therefore were not able to capture cell-to-cell variability. Two recent studies isolated and analyzed single cells, thereby providing not only proof-of-principle of this type of approach in glioblastoma but important information about the nature of heterogeneity in a single nucleus. It is important to note here that the type of sequencing possible from single cell nucleic acids is not yet at the same high coverage as that performed from larger amounts of DNA, and the types of whole genome amplification necessary to do this kind of analysis can still be prone to error. Nevertheless, this field is developing rapidly. Patel et al. used single-cell RNA sequencing to profile 430 cells from 5...
different primary glioblastomas and provided a stunning demonstration of intratumoral heterogeneity. Cells from the same tumor expressed different RTKs, and there was a gradient of “stemness” capability from cell to cell within each tumor. Strikingly, using the transcriptomic subtypes from TCGA, each tumor was found to harbor within each tumor. Strikingly, using the transcriptomic from the same tumor expressed different RTKs, and there

An additional study using single cell analysis focused on DNA, rather than RNA, analysis. Francis et al. isolated nuclei from 2 glioblastomas using flow cytometry and amplified single cell genomes for next-generation sequencing. Multiple distinct alterations can lead to the coexistence of various EGFR aberrations within tumors, including chromosomal break points, EGFR variants, extracellular domain missense mutations, and carboxy-terminal truncation. Among several interesting findings, this study showed that there was heterogeneous expression of oncogenic EGFR variants within single cells of the same tumor, all of which were mutually exclusive, implying that these variants evolved independently in a convergent manner. Further work using the elegant approach described will undoubtedly reveal novel insights into oncogene cooperativity, mechanisms of subclonal evolution, and more detailed analysis of the types of alterations found in single cells, especially as DNA sequencing coverage potential increases.

Taken together, it is becoming clear that, next to invasion, the hallmark of glioblastoma is heterogeneity—both between individual tumors and within each tumor (Fig. 1). Further study on the single cell level will be crucial to understand how subclonal ensembles behave, what the nature of cell state transitions might be, and how populations of subclones can be targeted therapeutically.

Cellular Heterogeneity
Identification of a Subpopulation of Tumor-Initiating Cells

Our conceptual framework for tumor biology has been greatly influenced by the introduction of the cancer stem cell hypothesis, which posits a “developmental” tumor hierarchy maintained by a rare fraction of cells with stem cell–like properties called “cancer stem cells” or “tumor-initiating cells” (TICs). Although the identity of a true “cancer stem cell” in glioblastoma and its contribution to the overall disease process remain somewhat controversial, there has been intensive investigation of glioblastoma cell subpopulations with the biological properties of self-renewal in vitro and tumor formation in xenografts in vivo. In glioblastoma, reasonable human data and very rigorous transgenic mouse data exist, which together support the notion that TICs represent a key and influential component of the cellular heterogeneity observed in glioblastoma and is thought to be responsible for overall tumor growth, therapy resistance, and tumor recurrence.

One of the main reasons for dissent regarding the identity of TICs in glioblastoma is the lack of reliable markers that definitively and prospectively identify glioblastoma cells as TICs. Singh and colleagues originally reported that human brain tumor cells expressing cell surface antigen CD133 were capable of self-renewal in vitro and tumor development in the brains of immunodeficient mice. In contrast, CD133-negative cells engrafted to the injection site but failed to form tumors. However, others have found that CD133-negative tumors can also initiate aggressive tumors in xenograft models. Among the markers used for identification of TICs (some not applicable in a prospective manner) are Nestin, SOX2, ID1, CD15, CD44, Integrin alpha 6, and the ability to extrude Hoechst dye (known as side-population analysis). In fact, part of the issue with identifying a reliable marker for TICs is that more than one type of TIC likely exists in glioblastoma, a concept most recently supported by the observation that TICs grown in sphere culture cluster predominantly into two transcriptomic subtypes—proneural and mesenchymal—with transcriptional parallels to the respective TCGA subtypes. Importantly, compared with proneural TICs, mesenchymal TICs were relatively radioresistant, in a manner dependent on NF-κB activity. Moreover, these subtypes themselves were found to be dynamic and plastic, as proneural TICs could be converted into mesenchymal TICs in culture through exposure to tumor necrosis–α, a factor elaborated by microglia, suggesting that the microenvironment might be capable of controlling the transcriptional signature and consequently the phenotype of TICs potentially in vivo. Given the discovery of diverse and shifting transcriptional profiles in TICs, it is perhaps not surprising that a consistent and ubiquitous TIC marker has been challenging to identify.

Besides cell extrinsic factors, recent exciting data have highlighted the importance of epigenetic regulation in defining the TIC phenotype. Using global profiling of the histone H3 Lysine27 acetylation (H3K27Ac), which marks genomic loci with active enhancers and promoters, Suva and colleagues uncovered an epigenetic signature that defines the self-renewal and tumorigenic potential of TICs. Conceptually similar to work done on induced pluripotent stem cells (iPSCs), this epigenetic profile provided a roadmap to identifying the master regulator transcription factors required for tumor initiation and moreover were sufficient to “reprogram” differentiated and nontumorigenic glioblastoma cells into TIC-like cells, which they named “induced tumor-propagating cells” (iTPCs). The transcription factors identified were POUF2, SOX2, SALL2, and OLG2, the last of which could be interchanged with the epigenetic regulator ROR2. Remarkably, sequential gene “add-back” of the 4 identified transcription factors in differentiated glioblastoma cells recapitulated the H3K27Ac epigenetic

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signature observed in TICs, suggesting that these genes in collaboration could sculpt the chromatin state of glioblastoma cells. These results also raise the intriguing idea that an epigenetic profile might represent the underlying abnormality in TICs. Together, it appears that the TIC, rather than representing one particular cell type arising from a particular transformed stem/progenitor cell, more likely represents a dynamic—rather than fixed—cell state in flux, shaped by genetic, epigenetic, and microenvironmental events (Fig. 2).

**TIC Involvement in Treatment Resistance and Recurrence**

Glioblastoma remains one of the most lethal human malignancies in part due to resistance to radiation therapy. TICs in glioblastomas are hypothesized to contribute to resistance mechanisms. Bao and colleagues demonstrated that ionizing radiation enriched the TIC subpopulation, as shown through increased CD133 positivity, in culture and in xenografts, likely due to diminished apoptosis. Mechanistically, the authors suggested that CD133-positive cells exhibited higher phosphorylation and therefore activation of regulatory proteins that respond to DNA damage. TICs have also been implicated in tumor recurrence following treatment with the alkylating agent temozolomide. Although the relative resistance of TICs to temozolomide in comparison with their non-TIC counterparts remains unclear, clever transgenic mouse experiments have clearly demonstrated that TICs contribute to recurrence following temozolomide. Specifically, Chen and colleagues used a genetic mouse model of glioblastoma with conditional loss of major tumor suppressors PTEN, NF1, and P53 in neural stem cells and crossed these tumor-prone mice with a transgenic mouse that marks Nestin-positive TICs. Temozolomide eradicated the majority of proliferative cells in tumors in vivo, which was then followed by reemergence of cell division and tumor growth via the Nestin-positive TIC subpopulation. Thus, the distinct responses of TICs versus non-TICs to standard chemoradiation therapies highlights the clinical importance of cellular heterogeneity and motivates further investigation into the mechanisms underlying these divergent treatment responses.

**Trans-Differentiation of TICs**

Although it is well established that TICs can undergo aberrant differentiation along restricted neural lineages, emerging evidence suggests that the stem cell–like properties of TICs might also render TICs capable of adopting nonneural fates. Several groups have reported that TICs have the capacity to transdifferentiate into components of vessels, namely endothelial cells and pericytes, suggesting the possibility that glioblastoma tumors may sculpt their own environment. The tumor perivascular niche exhibits striking parallels with the developmental niche seen with normal neural stem/progenitor cells and has been shown to act as a sanctuary for TICs. Three groups first reported that glioblastoma TICs can transdifferentiate into endothelial cells. As a proof of concept that tumor-derived endothelial cells contribute to the growth of glioblastoma tumors, disruption of endothelial cells arising from TICs in a xenograft model reduced tumor volume. Additionally, Soda and colleagues demonstrated that the development of tumor-derived endothelial cells were resistant to anti–vascular endothelial growth factor treatment in culture and in a glioblastoma mouse model. Therefore, the presence of tumor-derived...
endothelial cells in xenografts and clinical specimens from glioblastoma patients highlights the clinical value of considering TIC transdifferentiation in therapy development. Cheng and colleagues used rigorous in vivo cell lineage tracing and a large number of human glioblastoma specimens to demonstrate that TICs can also differentiate into vascular pericytes. Interestingly, in the pericyte study, the proportion of tumor-derived pericytes in human samples appeared much more significant than that of tumor-derived endothelial cells. Targeting of tumor-derived pericytes disrupted tumor vasculature and impacted tumor growth in xenografts. Together, the cellular heterogeneity provided by transdifferentiation of TICs into specific vascular cells supports the idea that glioblastomas actively establish and remodel a favorable tumor environment.

Discussion

We have used key examples in the literature to illustrate our growing understanding that heterogeneity—both molecular and cellular—is a defining hallmark of this disease (Table 1). Clearly, heterogeneity analysis is a rapidly expanding field, and with further technological advances in genomic profiling approaches at the single cell level, there will surely be additional insights obtained in a short period of time.

A remaining and intriguing question regarding both molecular and cellular heterogeneity is the relationship between structural alterations and cellular phenotypes within a single tumor. Might certain cellular phenotypes including the TIC state be enriched for particular structural alterations? Similarly, as suggested by the findings of Snuderl and colleagues, are particular niches or domains within a tumor enriched for distinct mutations that either establish or select for that tumor niche? Moreover, what is the relationship between cell-intrinsic tumor heterogeneity and cells of the brain tumor microenvironment? Finally, are the fixed molecular alterations or biological programs that drive the dynamic and fluid interchange that may occur between cellular phenotypes/cell states/transcriptional subtypes?

We have just scratched the molecular surface of this complex brain cancer and are already confronted with the challenging, if not disturbing, reality that glioblastomas are not only molecularly diverse across different patients...
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but that they also consist of multiple molecularly distinct tumors with different cellular phenotypes within one tumor. Our challenge is to understand the principles underlying this massive amount of data on tumor heterogeneity to offer insight into prognosis and guide our management of this devastating disease. The goal of translating these scientific observations into effective clinical practice will undoubtedly continue to require discovery that can only come from rigorous basic science. Indeed, the data suggest that the kind of profiling efforts conducted through TCGA may need to be applied to 1000–10,000 individual cells within a similar number of patient tumors. Ultimately, there is still much to learn about how our emerging appreciation of tumor heterogeneity can be harnessed to predict patient prognosis and direct treatment.

Disclosure

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

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