Multiplex mapping of chromatin accessibility and DNA methylation within targeted single molecules identifies epigenetic heterogeneity in neural stem cells and glioblastoma

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Human tumors are comprised of heterogeneous cell populations that display diverse molecular and phenotypic features. To examine the extent to which epigenetic differences contribute to intratumoral cellular heterogeneity, we have developed a high-throughput method, termed MAPit-patch. The method uses multiplexed amplification of targeted sequences from submicrogram quantities of genomic DNA followed by next generation bisulfite sequencing. This provides highly scalable and simultaneous mapping of chromatin accessibility and DNA methylation on single molecules at high resolution. Long sequencing reads from targeted regions maintain the structural integrity of epigenetic information and provide substantial depth of coverage, detecting for the first time minority subpopulations of epigenetic configurations formerly obscured by existing genome-wide and population-ensemble methodologies. Analyzing a cohort of 71 promoters of genes with exons commonly mutated in cancer, MAPit-patch uncovered several differentially accessible and methylated promoters that are associated with altered gene expression between neural stem cell (NSC) and glioblastoma (GBM) cell populations. In addition, considering each promoter individually, substantial epigenetic heterogeneity was observed across the sequenced molecules, indicating the presence of epigenetically distinct cellular subpopulations. At the divergent MLH1/EPM2AIP1 promoter, a locus with three well-defined, nucleosome-depleted regions (NDRs), a fraction of promoter copies with inaccessible chromatin was detected and enriched upon selection of temozolomide-tolerant GBM cells. These results illustrate the biological relevance of epigenetically distinct subpopulations that in part underlie the phenotypic heterogeneity of tumor cell populations. Furthermore, these findings show that alterations in chromatin accessibility without accompanying changes in DNA methylation may constitute a novel class of epigenetic biomarker.

[Supplemental material is available for this article.]

Human tumors often display substantial intratumoral heterogeneity in both phenotypic and molecular features. Cells that are drug-tolerant or have tumor-initiating capabilities are of high biological interest and are estimated to represent 1%–20% of bulk tumor cells (for review, see Visvader and Lindeman 2008). This cellular heterogeneity represents a formidable challenge to the discovery of effective cancer treatments. The frequency and degree of tumor heterogeneity cannot be explained solely by genetic determinants. Additionally, the reversible nature of cancer cell proliferative potential and drug tolerance suggests mechanisms that invoke plasticity (Sharma et al. 2010).

Dynamic control of gene expression is exerted by various epigenetic mechanisms, including DNA methylation, histone post-translational modifications, and nucleosome positioning and occupancy; although the latter three features have not been rigorously proven to be heritable (Schreiber and Bernstein 2002; Fuks 2005; Esteller 2007). Aberrant DNA methylation of CpG (hereafter, CG) dinucleotides is a well-documented phenomenon in cancer (Baylin and Jones 2011). It is widely accepted that DNA methylation near transcriptional start sites (TSSs) is associated with gene silencing. Hypermethylation of tumor-suppressive genes and hypomethylation of tumor-promoting genes is commonly observed, even in early stages of carcinogenesis (Herman and Baylin 2003). Although often evaluated separately, DNA methylation exerts control over gene expression within the context of chromatin. Expressed and poised genes are usually unmethylated and depleted of nucleosomes near their TSSs, thereby exhibiting increased accessibility to trans-activating factors (for review, see Jiang and Pugh 2009). Conversely, the TSSs of transcriptionally inactive genes tend to be associated with increased nucleosome occupancy, conferring chromatin inaccessibility, but can be either unmethylated or methylated. Thus, integrated evaluation of DNA methylation within the context of chromatin accessibility is more in-
formative than evaluating each epigenetic feature separately (Pardo et al. 2011b; You et al. 2011; Kelly et al. 2012). Notably, the extent of cell-to-cell heterogeneity in chromatin accessibility at gene promoters in either disease-free or tumor cells remains ill-defined.

Assessing intratumoral epigenetic heterogeneity necessitates the use of methods able to query chromatin structure at the level of single molecules. Our laboratory has developed a high-resolution footprinting technique, termed MAPit (methyltransferase accessibility protocol for individual templates). MAPit exploits exogenous addition of DNA methyltransferases (DNMTs), to probe accessibility of DNA in chromatin (Kladde et al. 1996; Xu et al. 1998b; Kilgore et al. 2007; Pardo et al. 2009). This technique has been used to simultaneously map DNA methylation and nucleosome positions on single molecules in many gene-specific studies (Kilgore et al. 2007; Wolff et al. 2010; Delmas et al. 2011; Pardo et al. 2011a; You et al. 2011; Yang et al. 2012; Darst et al. 2013), and more recently, genome wide (Kelly et al. 2012).

The identification and study of minority epigenetic subpopulations at multiple loci by gene-specific or genome-wide bisulfite genomic sequencing (BGS) is currently precluded due to requirements for large amounts of input DNA and prohibitive costs associated with obtaining the needed depth in sequencing coverage. To circumvent these limitations, we have adapted bisulfite patch PCR (Varley and Mitra 2010), a highly multiplexed approach to prepare targeted DNA for next-generation sequencing, to accommodate DNA obtained from chromatin probed for accessibility. We applied the resulting method, termed MAPit-patch, to determine the extent to which epigenetic heterogeneity exists in human GBM and control NSC. We concurrently profiled DNA methylation and chromatin accessibility at 71 promoters and identified several classes of epigenetic heterogeneity as well as 29 promoters that were differentially methylated and/or differentially accessible between GBM and NSC. Strikingly, a subpopulation of cells exhibiting inaccessible, but unmethylated, MLH1 promoter chromatin was negative for MLH1 immunostaining and enriched in TMZ-tolerant GBM cells. Epigenetic heterogeneity is therefore a common feature within a given GBM and NSC cell line and may contribute to diverse cellular phenotypes, including drug tolerance.

Results

MAPit-patch, a multiplexed, targeted method for simultaneous mapping of chromatin accessibility and DNA methylation on single molecules

To obtain combined DNA methylation and chromatin accessibility data on individual DNA strands or molecules, nuclei are probed with M.CviPI, which methylates cytosine in accessible GC dinucleotides (Xu et al. 1998a). GC sites within nucleosomes or those occupied by nonhistone proteins impair accessibility to M.CviPI and remain unmethylated (Kladde et al. 1996). For MAPit-BGS (Fig. 1A), genomic DNA is then bisulfite treated such that unmethylated C is deaminated to U, whereas methylated C (mC) remains mC. Bisulfite-treated DNA is then amplified using locus-specific primers; then reaction products are purified and cloned. Individual clones are sequenced and data are analyzed to map the methylation status of CG and GC sites. (B) For MAPit-patch, genomic DNA is fragmented using a GC and CG methylation-insensitive enzyme, such as R.AluI. Fragmented DNA is then denatured and subjected to target selection, whereby left and right patch oligonucleotides hybridize to each end of one strand of each locus and “patch” complementary oligonucleotides for universal priming (U1 and U2) by ligation (step not shown). The U2 oligonucleotide contains exonuclease-resistant modifications at its 3' end (black oval). Therefore, subsequent (7) 3’ to 5’ exonuclease digestion leaves targeted DNA strands intact and removes unhybridized oligonucleotides as well as nontargeted genomic DNA. Enriched DNA is (8) bisulfite converted and (9) amplified using universal primers that comprise sequences of U1 or complementary to U2, 5-bp barcodes to facilitate multiplexing, and adapter sequences specific for a sequencing platform.

Figure 1. MAPit-BGS and MAPit-patch workflow. Both assays begin with (1) preparation of nuclei and (2) incubation with M.CviPI. Upon termination of the chromatin probing reaction, (3) genomic DNA is extracted and processed. (4) For MAPit-BGS, genomic DNA is (4) bisulfite treated such that unmethylated C is deaminated to U, whereas methylated C (mC) remains mC. Bisulfite-treated DNA is then (5) PCR amplified using locus-specific primers; then reaction products are (6) purified and cloned. Individual clones are sequenced and data are analyzed to map the methylation status of CG and GC sites. (B) For MAPit-patch, genomic DNA is fragmented using a GC and CG methylation-insensitive enzyme, such as R.AluI. Fragmented DNA is then (5) denatured and (6) subjected to target selection, whereby left and right patch oligonucleotides hybridize to each end of one strand of each locus and “patch” complementary oligonucleotides for universal priming (U1 and U2) by ligation (step not shown). The U2 oligonucleotide contains exonuclease-resistant modifications at its 3’ end (black oval). Therefore, subsequent (7) 3’ to 5’ exonuclease digestion leaves targeted DNA strands intact and removes unhybridized oligonucleotides as well as nontargeted genomic DNA. Enriched DNA is (8) bisulfite converted and (9) amplified using universal primers that comprise sequences of U1 or complementary to U2, 5-bp barcodes to facilitate multiplexing, and adapter sequences specific for a sequencing platform.

The bisulfite patch PCR study targeted 94 loci in a single reaction using the restriction (R) enzyme R.AluI (recognizes AGCT) for fragmentation of genomic DNA. R.AluI is well suited for patch selection as it occurs frequently in CG islands, which are present in as many as 70% of mammalian promoters (Gardiner-Garden and Frommer 1987; Takai and Jones 2002). However, as digestion by R.AluI is blocked by C-5 methylation, its use is not compatible with M.CviPI-modified DNA. The isoschizomer, R.AluBI, also recognizes AGCT sites, but is not affected by C-5 methylation (Sibenzyme.com; Supplemental Fig. S1), allowing use of the original bisulfite patch PCR oligonu-
To identify probing conditions that would allow for detection of different chromatin states, nuclei from NSC were probed with 0, 30, and 100 U M.CviPI and analyzed by MAPit-BGS (Fig. 2). NSC were harvested from serum-free, suspension culture (hereafter, spheroid culture), which maintains these cells in an undifferentiated state and preserves their phenotypic heterogeneity (Deleyrolle and Reynolds 2009). To facilitate pattern recognition, aligned sequences were uploaded into a web-based hierarchical clustering program called MethylMapper (http://genome.ufl.edu:8080/methyl) (Darst et al. 2012). MethylMapper generates 3-color images of clustered CG methylation (Fig. 2, left panels) or GC accessibility (Fig. 2, right panels). Each row represents one sequenced molecule. CG and GC information was clustered end-to-end, so the top-to-bottom presentation order of the molecules is linked in the left and right panels.

In NSC, MAPit-BGS of the proximal promoter of MLH1, an expressed gene, showed that this region is unmethylated and highly accessible around its two TSSs (Fig. 2A), defining two NDRs. A protected region (footprint) of 16 bp within the downstream NDR likely corresponds to a DNA-bound protein or protein complex. The protected region between the two accessible regions is consistent with the size of a nucleosome core particle (147 bp). Conversely, the promoter of PYCARD, a silenced gene, is hypermethylated and largely inaccessible (Fig. 2B). Though both 30 units and 100 units M.CviPI showed equivalent levels of probing at both the MLH1 and PYCARD promoters, we opted to use 100 units in all further reactions to ensure saturation. To determine if epigenetic heterogeneity is also observed, we amplified the promoter of PROM1 (Fig. 2C), which encodes the cell surface antigen CD133, expressed in up to 40% of cultured NSC (Piao et al. 2006; Sun et al. 2009). All analyzed PROM1 sequences had low or no methylation (Fig. 2C, left panel). However, heterogeneity in chromatin accessibility was observed across the locus, especially nearby TSS1b (Fig. 2C, right panel), where transcription initiates in neural tissue (Shmelkov et al. 2004). Approximately 50% of the promoters in the NSC population exhibited substantial accessibility at TSS1b, indicating nucleosome depletion. The remaining half of promoter molecules was inaccessible, and exhibited a nucleosome-sized or larger footprint that encompassed TSS1b. In conclusion, probing with 100 units of M.CviPI enables interrogation of diverse, heterogeneous chromatin states in spheroid cultured cells.

We sought to ensure that M.CviPI probing of chromatin structure and hence GC methylation would not affect the coverage and reproducibility of results obtained by bisulfite patch PCR. Target enrichment was therefore performed using the published patch oligonucleotide library. This library targets promoters within 700 bp of the TSS of 90 genes that are commonly mutated in breast and/or colon cancer (“CAN genes”) (Varley and Mitra 2010). Four control loci

**Figure 2.** MAPit identifies expected epigenetic patterns and detects heterogeneous chromatin structures at specific loci. MAPit-BGS in NSC nuclei of the promoters of (A) MLH1 (expressed), (B) PYCARD (silenced), and (C) PROM1 (heterogeneously expressed) probed with the indicated units (U) of M.CviPI activity. A schematic of each promoter is indicated at the very top: bent arrows, TSSs; ellipses, length of a 147-bp nucleosome core particle; map coordinates relative to the first TSS are indicated for the most upstream and downstream CG (left) or GC (right) sites. Data were plotted with MethylMapper; each row of pixels represents one sequenced DNA strand or molecule, with the same top-to-bottom presentation order in each panel. Vertical hashes demarcate individual CG (left) or GC (right) sites (CCG excluded), and an additional site density plot is shown below each promoter schematic. The key for CG or GC methylation status is shown in A, bottom. Two or more consecutively methylated CG and GC sites are connected by red and yellow, respectively, whereas two or more consecutively unmethylated CG and GC sites are connected by black. Gray connects the borders between methylated and unmethylated sites. White at either end of a molecule indicates missing or unaligned sequence. Note a variably positioned NDR associated with the neural-specific TSS1b in the PROM1 promoter in C, whereas both TSSs at MLH1 in A colocalize with an NDR, the downstream from which is occupied by a DNA-binding factor(s) (labeled footprint).
were also included and we added 19 additional cancer-associated loci targeted within 600 bp of the TSS (Supplemental Tables S1, S2). MAPit-patch was performed using DNA from NSC and GBM L0 spheroid cultures probed with 0 or 100 units M.CviPI. Bar-coded reactions were pooled and sequenced using one-eighth of a plate on a 454 FLX Life Sciences sequencer. After removing sequences with bisulfite conversion efficiencies of <95%, and sequencing reads <100 bp, we obtained 22,356 sequences. Of these, 100% aligned to 104 of the 113 targeted loci (92% of targets), indicating a high sensitivity of the technique. Mean coverage of each promoter was 215 reads (range of 1–1039 reads; median, 99 reads), and the sequencing depth of 87% of the targeted promoters was within 10-fold of the median. Consistent with the published bisulfite patch PCR results (Varley and Mitra 2010), we observed a significant inverse correlation between amplicon length and read coverage with MAPit-patch (Fig. 3A) (P < 0.0001; Pearson’s correlation). To determine if targeted loci were reproducibly amplified, the number of reads per locus was plotted for each sample and correlation coefficients between all pairs of samples were calculated (data not shown). The mean correlation coefficient was 0.94, comparable to the value of 0.91 obtained by bisulfite patch PCR. As previously reported by Varley and Mitra (2010), these data indicate that the coverage of each promoter is not stochastic between samples, but is reproducible and affected by amplicon length. Thus, chromatin probing with M.CviPI and R.AluBI substitution in MAPit-patch do not affect the performance of bisulfite patch PCR.

To determine if probing with M.CviPI would affect accurate quantification of CG methylation, the fraction of methylated CGs (excluding GCGs in all analyses) at each promoter was calculated and compared between the 0- and 100-unit samples. The fraction of CG methylation correlated significantly (P < 0.0001) between the 0- and 100-unit samples, r = 0.99 (Fig. 3B). To confirm that modification by M.CviPI did not alter the ability of bisulfite patch PCR to amplify methylated and unmethylated molecules with equal efficiency, we examined the DNA methylation profile of the imprinted locus H19 in NSC. The 0- and 100-unit M.CviPI-treated samples showed indistinguishable levels of CG methylation (0 units = 48%, 100 units = 50%; P = 0.483) and amplified methylated and unmethylated molecules with equivalent efficiencies (Fig. 3C,D, left panels). In conclusion, MAPit-patch does not introduce bias in quantification of CG methylation nor does it alter the ability to equivalently amplify methylated and unmethylated molecules. In addition, MAPit-patch accurately profiles the expected copy-specific inverse relationship between DNA methylation and chromatin accessibility at the imprinted H19 locus (P = 0.0015) (Fig. 3D, cf. right and left panels).

**Figure 3.** Probing chromatin with M.CviPI neither alters bisulfite patch PCR performance nor detection of CG methylation. (A) Number of sequencing reads decreases as a function of amplicon size. (B) Linear regression and Pearson’s correlation plotted for CG methylation levels in NSC treated with 0-unit versus 100-unit M.CviPI DNA methylation and chromatin accessibility at the imprinted H19 locus for NSC treated with (C) 0 units or (D) 100 units of M.CviPI. Symbols and the key for methylation status at right of C are as defined in Figure 1.

**Figure 4.** Overall distribution in NSC and GBM L0 of patterns of promoter CG methylation is similar and inversely associated with GC accessibility. (A) Distribution of promoters by methylation status in NSC (left) and GBM L0 (right). Dot plots of fraction of GC accessibility within each promoter methylation class in (B) NSC and (C) GBM L0. The mean fractions of GC accessibility ± one standard error of the mean (SEM) are indicated. (***) P < 0.001 compared to GC accessibility from unmethylated promoters for each sample.

**CAN gene promoter methylation and chromatin accessibility in NSC and GBM L0**

We quantified the fraction of methylated CGs at each promoter for which we obtained at least 10× sequencing coverage in NSC and GBM (71 promoters) (Supplemental Tables S3, S4). Promoters exhibiting ≤20% CG methylation were classified as “unmethylated,” those with ≥80% methylation were classified as “methylated,” and those with 21%–79% CG methylation were considered
Table 1. Differentially methylated regions (DMRs) between NSC and GBM L0

| Locus    | NSC  | GBM  | P-value | Direction |
|----------|------|------|---------|-----------|
| CDH1     | 0.05 | 0.52 | 0.0001  | Hyper     |
| DBN1     | 0.59 | 0.80 | 0.0001  | Hyper     |
| EPHB6    | 0.01 | 0.03 | 0.0001  | Hyper     |
| LRRFIP1  | 0.03 | 0.43 | 0.0001  | Hyper     |
| PCDH9A   | 0.32 | 0.87 | 0.0001  | Hyper     |
| RASSF1   | 0.34 | 0.74 | 0.0001  | Hyper     |
| SORL1    | 0.81 | 0.96 | 0.0001  | Hyper     |
| SOX10    | 0.91 | 0.97 | 0.0001  | Hyper     |
| VEPH1    | 0.70 | 0.84 | 0.0001  | Hyper     |
| ACSL5    | 0.73 | 0.51 | 0.0001  | Hypo      |
| AGAP2    | 0.73 | 0.53 | 0.0001  | Hypo      |
| CD93     | 0.39 | 0.03 | 0.0001  | Hypo      |
| COL19A1  | 0.47 | 0.30 | 0.0001  | Hypo      |

*Frequency of methylated CG sites.

**Hyper** Methylation; **Hypo** Hypomethylation.

differentially methylated and/or accessible genes, although known to be associated with colon and/or breast cancer, may also be generally affected in GBMs.

Heterogeneity in chromatin accessibility and DNA methylation at multiple CAN gene promoters

To identify cell-to-cell heterogeneity, we examined patterns of GC accessibility in target gene promoters that sequenced with ≥20× coverage (54 promoters from NSC; 67 promoters from GBM L0) (Supplemental Tables S3, S4). Two parameters were counted: (1) the number of reads per locus that exhibited ≥126 bp of inaccessible GC sites (i.e., minimal protection consistent with nucleosome occupancy) divided by the total number of reads and subtracted from a value of 1; and (2) the number of reads per locus that contain a nucleosome-free region (three or more consecutively accessible GC sites) divided by the total number of reads. The average of these two values gives the GC accessibility score, reflecting the fraction of molecules that are nucleosome-free and "accessible" at each locus (Supplemental Tables S3, S4). Promoters were stratified into chromatin class quintiles as follows: (1) accessible (≥81% molecules accessible); (2) mostly accessible (61%–80% molecules accessible); (3) half accessible (40%–60% molecules accessible); (4) mostly inaccessible (20%–39% molecules accessible); and (5) inaccessible (≤19% molecules accessible) (Fig. 6A). To provide independent assessment that MAPit-patch accurately assesses the degree of accessibility characteristic of each class of promoters, we performed quantitative restriction enzyme accessibility assays (Fig. 6B; Supplemental Fig. S2, assay optimization). We then identified four promoters that contain a SacI site within 350 bp of a TSS and for which we obtained ≥20× coverage by MAPit-patch. After incubation of GBM L0 nuclei with R.Sacl, accessibility was quantitatively measured by qPCR with convergent primers spanning each SacI site. The accessibility profiles of all four promoters in Figure 6B corresponded well with those determined by MAPit-patch (Supplemental Table S4). Confirmation of accessibility by this independent, quantitative approach indicates that the heterogeneous accessibility patterns identified by MAPit-patch reflect biological diversity in chromatin accessibility, not only among the interrogated promoters but also across the cohort of sequenced molecules for each promoter. The distribution of promoter amplicons among the five different accessibility classes was similar for both NSC and GBM L0 (Fig. 6C). GC methylation was inversely correlated with GC accessibility for both samples (Fig. 4B,C). In contrast to unmethylated promoters, a similar decrease in GC accessibility was observed among promoters that were methylated and variably methylated, suggesting that decreased accessibility can occur independent of high levels of DNA methylation.

Comparing DNA methylation and chromatin accessibility between NSC and GBM L0 cells, MAPit-patch identified 13 promoters with differential CG methylation (differentially methylated region [DMR]) (Table 1), seven with differential GC accessibility (differentially accessible region [DAR]) (Table 2), and nine with differential CG methylation and GC accessibility (differentially methylated and accessible region [DMAR]) (Table 3). These genes exhibited reproducible differences (P < 0.01; NSC 0 units M.CviPI versus GBM 0 units, and NSC 100 units versus GBM 100 units) and no statistically significant differences in CG methylation between replicates (NSC 0 units versus NSC 100 units, and GBM 0 units versus GBM 100 units).

We selected 15 promoters (5 DMR, 4 DAR, and 6 DMAR) and measured gene expression in the NSC culture as well as two primary GBM cultures derived from different patients (L0 and L2), using quantitative reverse transcription PCR (qRT-PCR). Ten of 15 tested promoters exhibited the expected correlations between altered CG methylation, chromatin accessibility, and gene expression (Tables 1–3; Fig. 5). Two genes (AGAP2 and TAF1) showed no expression change in one GBM culture, but showed the expected change in the other GBM culture. Finally, three promoters showed unexpected changes in gene expression (ICAMS, NKX2-5, and ABCB8; NKX2-5 expression data not graphed due to 6500- and 2800-fold increases in GBM L0 and GBM L2, respectively, compared to NSC). For ICAMS, the increase in expression correlated with the increase in GC accessibility rather than the small site-specific increase in CG methylation. These results indicate that the differential epigenetic features identified by MAPit-patch are reflective of differential gene expression in most cases. Importantly, most of the genes that are differentially expressed between NSC and GBM L0 were also differentially expressed in GBM L2. This indicates that these

to have variable methylation. Overall, we observed that both NSC and GBM L0 had a similar distribution of promoters by methylation status (Fig. 4A). For each promoter, we calculated the fraction of accessible GCs. As expected, GC accessibility was inversely correlated with CG methylation for both samples (Fig. 4B,C). Interestingly, compared to unmethylated promoters, a similar decrease in GC accessibility was observed among promoters that were methylated and variably methylated, suggesting that decreased accessibility can occur independent of high levels of DNA methylation.

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Differentially methylated and accessible regions (DMARs) between NSC and GBM L0

| Locus  | NSC*  | GBM*  | P-value  | Direction* | NSC*  | GBM*  | P-value  | Direction* |
|--------|--------|--------|----------|------------|--------|--------|----------|------------|
| ICAMS  | 0.01   | 0.05   | 0.0001   | Hyper      | 0.23   | 0.28   | 0.0003   | Hyper      |
| IGFBP3 | 0.03   | 0.95   | 0.0001   | Hyper      | 0.13   | 0.06   | 0.0004   | Hypo       |
| NKB2L3 | 0.02   | 0.44   | 0.0001   | Hyper      | 0.17   | 0.12   | 0.0017   | Hypo       |
| H19    | 0.07   | 0.79   | 0.0001   | Hyper      | 0.08   | 0.04   | 0.0035   | Hypo       |
| SH3TC1 | 0.09   | 0.04   | 0.0001   | Hypo       | 0.48   | 0.68   | 0.0001   | Hypo       |
| SLCTC1 | 0.91   | 0.75   | 0.0001   | Hypo       | 0.07   | 0.11   | 0.0001   | Hypo       |
| TAF1   | 0.29   | 0.01   | 0.0001   | Hypo       | 0.32   | 0.50   | 0.0001   | Hypo       |
| TNN    | 0.86   | 0.79   | 0.0074   | Hypo       | 0.04   | 0.07   | 0.0005   | Hyper      |

*Frequency of methylated CG sites.
*Hyper* Hypermethylation; *Hypo* Hypomethylation.

A subpopulation of drug-tolerant cells is associated with increased promoter nucleosome occupancy

We wanted to determine if epigenetic subpopulations observed in GBM L0 were associated with a disease-relevant phenotype. Molecules from the divergent MLH1/EPM2AIP1 promoter were unmethylated and mostly accessible in GBM at both the distal and proximal promoter region (Fig. 7A–C). There was, however, a subpopulation of promoter copies that were almost completely inaccessible at both target amplicons (Fig. 7B,C, enclosed by cyan rectangles). The protein product of MLH1 is involved in DNA mismatch repair and is considered a tumor suppressor (Bronner et al. 1994; Prolla et al. 1998). Importantly, it is known that loss of MLH1 protein expression renders cells insensitive to treatment with DNA-alkylating agents such as temozolomide (TMZ) (Taverna et al. 2000), which is the frontline chemotherapeutic treatment for GBM (Hegi et al. 2005; Stupp et al. 2005). We hypothesized that the subpopulation of MLH1 promoter copies with inaccessible chromatin observed in GBM may reflect a cellular subpopulation that does not express MLH1 and therefore is tolerant of TMZ.

We conducted MAPit-BGS to confirm the presence of the inaccessible subpopulation of MLH1 promoter copies in GBM that was identified by MAPit-patch. As in Figure 7B and C, we observed that the status of MLH1 was mostly accessible, with an inaccessible subpopulation at both the distal (Fig. 7D) and proximal (Supplemental Fig. S3A, lower panel) promoter. This was observed in GBM L0 as well as in GBM L2 but not in NSC (Fig. 2A; Supplemental Fig. S3B). Notably, this inaccessible subpopulation was specific to MLH1 and was not observed at the completely accessible promoter of the PMS2 gene, which encodes the mismatch repair binding partner of MLH1 (Fig. 7E). Immunostaining for MLH1 followed by flow cytometry showed that both GBM lines contained a subpopulation of MLH1-negative or low-expressing cells (Fig. 7F).

To determine if the MLH1-low or -negative phenotype was associated with cells harboring copies of MLH1 promoter chromatin that were inaccessible, GBM cells were treated with TMZ for 4 d to counter select against cells expressing MLH1 protein. Flow cytometry of immunostained cells confirmed dose-dependent enrichment for MLH1-negative/low cells upon treatment with TMZ (Fig. 8A). Surviving TMZ-tolerant cells were assayed for accessibility of distal MLH1 promoter chromatin by MAPit-BGS (Fig. 8B, cf. middle and top panels) and R.SacI digestion (Fig. 8C, MLH1, cf. middle and left blue bars) and found to be enriched for inaccessible chromatin (both experiments, $P < 0.0001$). TMZ-tolerant cells from the same experiment were outgrown in the absence of TMZ for 10 additional passages and assayed for chromatin accessibility. Compared to the starting TMZ-tolerant cells, the outgrown cells exhibited a significant increase in accessibility of distal MLH1 promoter chromatin to M.CviPI (Fig. 8B, cf. bot-
Integrated determination of epigenetic features is important for understanding how epigenetic mechanisms contribute to tumorigenesis and how to effectively target them for cancer treatment. Single-molecule-level technologies that preserve the heterogeneity inherent within human tumors are essential to fully understand the contribution of these factors to disease progression and resistance to treatment. We have described a novel deep sequencing approach, termed MAPit-patch, which simultaneously determines chromatin structure and DNA methylation with single-molecule resolution, thus preserving sample heterogeneity. The method is highly scalable and cost effective, which will facilitate the screening of multiple tissue samples. Also, the targeted (rather than genome-wide) approach is within reach for translating these technologies for assessment of particular disease biomarkers. Using MAPit-patch, we have shown that epigenetic heterogeneity at a given locus is commonly observed in both NSC and GBMs. Furthermore, our results indicate that, in addition to DNA hypo- or hypermethylation, changes in chromatin accessibility alone are associated with tumor-specific alterations in gene expression. Finally, we have shown for the first time that a small subpopulation of cells with inaccessible chromatin at the promoter of a tumor suppressor is associated with drug tolerance. These results show that epigenetic heterogeneity may underlie some of the phenotypic diversity observed in human GBM and has broad implications for molecular profiling of tumors in general.

We identified a number of genes that were differentially methylated and/or accessible and differentially expressed between NSC and GBM L0. The identification of hypomethylation/hyperaccessibility at tumor-promoting genes and hypermethylation/hypoaccessibility at tumor-suppressive genes suggests that these epigenetic features are not random, but represent biologically relevant events. However, additional studies are required to determine the functional significance of DNA methylation, if any, at these genes. Our finding of differentially methylated/accessible genes previously shown to harbor mutations in GBM—COL19A1 (Sumiyoshi et al. 1997), CD93 (Dieterich et al. 2012), AGAP2 (Knobbe et al. 2005), and ACSL5 (Mashima et al. 2009)—supports the validity of this approach in identifying GBM-relevant epigenetic perturbations. Interestingly, most of the genes that were identified as differentially methylated between GBM and NSC were classified as variably methylated in NSC (Table 4, lower). It has been reported that loci exhibiting variability in methylation status between different types of normal tissues are more often aberrantly methylated in tumors (Feinberg and Irizarry 2010; Hansen et al. 2011). It was proposed that these regions exhibit the greatest amount of epigenetic plasticity and are therefore more susceptible to perturbations during tumorigenesis. Our results suggest that this same phenomenon may be observed within...
a given sample, whereby loci that are variably methylated within the cellular population in a normal tissue are more amenable to aberrant hypo- or hypermethylation in tumors. However, assessment of additional loci in several normal and tumor samples is necessary to support this premise.

We found that a subpopulation of molecules with inaccessible chromatin at MLH1 was associated with a cellular subpopulation of MLH1-negative/low cells. This MLH1-negative/low subpopulation with inaccessible chromatin is enriched upon treatment with TMZ (Fig. 8B,C). These results are significant for several reasons. Although a biomarker exists to predict sensitivity to TMZ treatment, a small number of cells with nucleosome-depleted MLH1 expression in sensitivity to TMZ (Liu et al. 1996; von Bueren et al. 2012). Second, two studies have shown by immunohistochemistry that small subpopulations of MLH1-negative cells commonly arise in primary GBM. One study showed that MLH1-negative cells were enriched after TMZ treatment in recurrent tumors (Stark et al. 2010). The second study showed that cells lacking PMS2 were enriched in recurrent tumors (Felsberg et al. 2011). Thus, loss of mismatch repair protein expression appears to be associated with clinical relapse of GMB and further studies to test these genes as biomarkers of treatment resistance is of high interest.

Importantly, DNA methylation was tested and excluded as the mechanism driving mismatch repair gene silencing in Felsberg et al. (2011). This is consistent with our results showing that aberrant chromatin inaccessibility, not DNA methylation, is associated with MLH1-negative GBM cells. This is relevant because studies evaluating epigenetic biomarkers often exclusively query DNA methylation. Our results suggest that chromatin accessibility may also be a useful feature to identify disease biomarkers. Furthermore, DNA methylation is considered to be a more stable biomarker than repairs DNA lesions (Taverna et al. 2000). Thus, loss of expression of MLH1 presumably supersedes the effects of aberrant MGMT expression in sensitivity to TMZ (Liu et al. 1996; von Bueren et al. 2012). Second, two studies have shown by immunohistochemistry that small subpopulations of MLH1-negative cells commonly arise in primary GBM. One study showed that MLH1-negative cells were enriched after TMZ treatment in recurrent tumors (Stark et al. 2010). The second study showed that cells lacking PMS2 were enriched in recurrent tumors (Felsberg et al. 2011). Thus, loss of mismatch repair protein expression appears to be associated with clinical relapse of GMB and further studies to test these genes as biomarkers of treatment resistance is of high interest.

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contains a 3-carbon spacer was also synthesized with five phosphorothioate bonds to further protect target loci from exonuclease digestion. Reactions were treated with exonucleases and bisulfite converted as described (Varley and Mitra 2010). Amplification of target loci was carried out in 50-μL reactions with the following components: all recovered bisulfite-converted DNA (10 μL), 1× HotStarTaq buffer (Qiagen), 500 μM MgCl₂, 50 μM each dNTP, 250 nM each barcoded primer, and 10 units HotStarTaq DNA polymerase (Qiagen). Reaction products were pooled and PCR purified, then gel purified. Purified products were sequenced at the University of Florida Interdisciplinary Center for Biotechnology Research using the Roche 454 GS FLX+ instrument according to manufacturer protocols.

**Sequencing data analysis**

Data was processed with custom Python code. Sequences were divided by barcode, using FASTools (http://genome.ufl.edu/rivalab/fastools/), then aligned to the MAPit-patch reference library by BLAST. To prevent bias, C residues in both read and reference sequences were fully converted to T in silico before alignment. After restoration of cytosine information, sequences were scored for percent deamination of HCH, i.e., cytosines neither CG nor GC. Sequences with <95% conversion of HCH to HTH and those covering <50% of the reference sequence were discarded. All GCG sites were removed from analysis. Genome-wide, GCGs represent only 5.6% of all GC dinucleotides, and removal of these sites does not strongly affect chromatin accessibility information (Kelly et al. 2012). For our promoter targets, GCGs represent 30.6% of all CG and 22.2% of all GC dinucleotides. Removal of these sites does not strongly affect DNA methylation or chromatin accessibility information (Supplemental Fig. S4).

**R.SacI accessibility assay**

Nuclei were prepared exactly as for MAPit, except that ethylenediaminetetraacetic acid and glycerol were omitted and 5 mM MgCl₂ was included.

**Statistical analysis**

All statistical analyses were performed using GraphPad Prism software. Pearson’s correlation was used to determine correlations and coefficients between samples and between amplicon length and abundance. For reproducibility measures, reads per locus were plotted in a correlation matrix for pairwise comparisons as previously described (Varley and Mitra 2010). Comparisons between groups were tested using two-way ANOVA followed by a Bonferroni ad hoc test. Significance values for differentially regulated promoters, copy-restricted signatures, and enrichment for methylation states were obtained using a two-sided Fisher’s exact test.

**Immunostaining and flow cytometry**

Cells were seeded in spheroid culture conditions, grown for 4 d, then treated with the indicated doses of TMZ. Three days after drug treatment, cells were harvested. Intact cells that excluded propidium iodide were then recovered by fluorescence-activated cell sorting for both downstream immunolabeling using antibody against MLH1 (sc-11442, 1/500 dilution, Santa Cruz) and chromatin accessibility information (Supplemental Fig. S4).
matin accessibility assays. MLH1 staining was quantified by flow cytometry on a Becton Dickinson LSRII instrument.

Data access
Raw 454 sequence reads have been submitted to the NCBI Gene Expression Omnibus (GEO; http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE50047.

Competing interest statement
M.P.K. shares royalties with coinventors for the M.CviPI enzyme used as a chromatin probe.

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Figure 8. Cells with inaccessible chromatin at MLH1 are enriched upon treatment with TMZ. (A) Immunostaining with anti-MLH1 antibody and flow cytometry were conducted on GBM L0 cells after 72-h treatment with the indicated doses of TMZ. Chromatin accessibility at MLH1 was measured in control (top), TMZ-treated (+TMZ, middle), and TMZ-treated cells subsequently propagated in drug-free media (+TMZ, then outgrown–TMZ, bottom) by (B) MAPIT-BGS (key at right) and by (C) protection from R.SacI activity for each locus. Bars represent the mean protection from R.SacI activity for each locus ± SEM (control and +TMZ, n = 5; outgrown, n = 3), normalized to a control locus lacking a SacI site. A second control locus, SEMA3B, which contains a SacI site, but is inaccessible in GBM L0, was also assayed. (***) P < 0.001.
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