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Single-cell analysis of clonal maintenance of transcriptional and epigenetic states in cancer cells

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#### **ABSTRACT**

Propagation of clonal regulatory programs contributes to cancer development. It is poorly understood how epigenetic mechanisms interact with genetic drivers to shape this process. Here we combine single-cell analysis of transcription and DNA methylation with a Luria-Delbrück experimental design to demonstrate the existence of clonally stable epigenetic memory in multiple types of cancer cells. Longitudinal transcriptional and genetic analysis of clonal colon cancer cell populations reveals a slowly drifting spectrum of epithelial-to-mesenchymal transcriptional identities that is seemingly independent of genetic variation. DNA methylation landscapes correlate with these identities but also reflect an independent clock-like methylation loss process. Methylation variation can be explained as an effect of global *trans*-acting factors in most cases. However, for a specific class of promoters, in particular cancer testis antigens (CTA), derepression is correlated with and likely driven by loss of methylation in *cis*. This study indicates how genetic sub-clonal structure in cancer cells can be diversified by epigenetic memory.

#### INTRODUCTION

The ability of cells to maintain their molecular identity through mitotic cell divisions is essential for the establishment of functionally coherent and stable clonal cell populations. During carcinogenic transformation, selection for beneficial driver mutations contributes to the basic clonal and subclonal structure of the evolving tumor<sup>1</sup>. However, this selective process provides limited flexibility for rapid adaptation and diversification in the context of a dynamic stromal environment, immune interactions or following treatment. Non-genetic mechanisms can modulate cellular states and enhance such flexibility if their diversification can persist and form an epigenetic memory. Pioneer systematic screenings for transcriptional memory<sup>2-4</sup> demonstrated the potential for clonally stable transcriptional phenotypes in mammalian systems. Nonetheless, it remains difficult to measure the extent of such transcriptional memory and to distinguish transient transcriptional heterogeneity within cell populations from stable and clonally transmitted states.

Theoretical and experimental models of commitment and memory through specific gene network architecture in unicellular organisms<sup>5</sup>, and systems for analyzing stable mitotic transmission of epigenetic states in mammals<sup>6-8</sup> provide a basis for understanding the mechanisms underlying the formation and maintenance of epigenetic memory. DNA methylation is the best studied epigenetic mechanism for stable memory formation, and the ability of cells to copy their methylation makeup to daughter cells is well established 9-10. The correlation between specific DNA methylation patterns and cell-type-specific transcriptional programs has been also demonstrated<sup>11</sup>. But the role of DNA methylation in regulating clonal heterogeneity within diversifying cell populations is still unclear. Methylation changes with ageing<sup>12</sup>, cellular senescence<sup>13</sup>, and transformation<sup>14-15</sup>. Errors in replicating methylation marks, which cause epimutations, can accumulate to create global, replication-dependent reduction in methylation levels that is associated with clock-like dynamics<sup>6</sup>. Such changes are unlikely to have a direct functional impact, and are observed as a background process in both normal and cancer tissues. Other cancer-linked methylation changes may be driven by modulation in the activity of transacting factors, including recurrent genetic mutations in the methylation machinery itself<sup>16</sup>. Here again, the role of DNA methylation as a carrier of molecular memory in cis is limited since it is only indicating the activity of an aberrant epigenetic modulator in trans. Given these broad background dynamics and trans-acting effects, it remains difficult to identify cases in which methylation is effectively maintaining repressed/de-repressed switches in cis at gene promoters or distal regulatory elements.

To address some of these difficulties, we implemented a Luria-Delbrück experimental design and compared single-cell transcriptional and epigenetic distributions in cancer cells to the distributions of mean gene expression and methylation across clones originating from the same cell populations. Using high-throughput and precise single-cell genomic technology, this resulted in thousands of single-cell profiles and hundreds of matching clonal profiles, revealing broad clonally stable transcriptional diversity in immortalized fibroblasts and in lung and colon cancer cells. Genetic profiling and longitudinal analysis indicate that the observed clonal diversity represents epigenetic memory. Analysis of DNA methylation characterizes the epigenetic makeup underlying repression and de-repression of key clonal gene modules, in particular along a spectrum of epithelial-to-mesenchymal transition (EMT) identities in colon cancer cells. DNA methylation in cis cannot be implicated unambiguously with a causal role in maintaining (and drifting) gene expression state for most clonal genes. Nevertheless, we observe a class of promoters, including several cancer testis antigens (CTA), for which correlation indicates a causal cis-regulatory role for DNA methylation. In conclusion, our in vitro Luria-Delbrück assays suggest that epigenetic memory in cancer cell populations operates pervasively and in parallel to genetic drivers, to diversify transcriptional programs and channel cells toward EMT and other tumorigenic transcriptional dynamics.

#### **RESULTS**

# Luria-Delbrück assays identify clonally stable transcriptional memory in cancer cells

To facilitate detailed exploration of clonal or transient transcriptional and methylation states in cancer cells we followed the classic Luria-Delbrück scheme (**Fig. 1a**). Clonally stable transcriptional programs or epigenetic states are expected to propagate from founder cells to give rise to homogeneous clones. In this case, inter-clonal variance should recapitulate precisely single-cell heterogeneity in the founding population. In contrast, cell-cycle signatures or other transient fluctuations are averaged out in clones so as to eliminate inter-clonal variance (**Extended Data Fig. 1a**). We FACS-sorted and cultured single cells of two lung cancer cell lines (NCI-H1299 and A549), one colorectal cancer (HCT116) and the non-cancerous WI38 fibroblast cell line. We analyzed 841 clones (median coverage of 103,000 UMIs), that were expanded for 9-10 doublings (**Extended Data Fig. 1b**) and compared those to single-cell RNA-seq profiles obtained from the matching founding populations (**Extended Data Fig. 1c-f**). Clonal populations (500 to 1,000 cells) allowed sampling of 50 cells in replicates, ensuring quantitative estimation of

clonal RNA concentrations (**Extended Data Fig. 1g-n**). Comparison of pooled RNA from cells and clones showed high degree of concordance and suggested the bias or selective constraint associated with the single-cell cloning procedure itself was limited (**Extended Data Fig. 2a-c**).

As expected, cell-cycle-linked single-cell variation was observed pervasively in all studied cell lines (Fig. 1b-d, Extended Data Fig. 2d-g and Supplementary Table 1). This prompted us to implement a strategy for discovery of cell-cycle independent gene-gene correlation modules, by comparing correlations in the raw count matrix and a randomized matrix obtained by shuffling RNA counts over cells with similar cell-cycle characteristics (see Methods and Extended Data Fig. 2h,i). Interestingly, we found strong cell-cycle independent transcriptional variation in all cell lines, suggesting the existence of either transient transcriptional dynamics, genetic sub-clonal population structure, and/or non-genetic clonal population structure. Our Luria-Delbrück scheme allowed us to distinguish transient from clonal population structure (Fig. 1e-g), confirming that the S-phase and M-phase gene modules observed in single-cell analysis were indeed transient, but suggesting other gene modules to be clonally stable (Extended Data Fig. 2j-o and Supplementary Table 2). For example, in HCT116 we discovered a clonally stable epithelial gene module, marked by EpCAM expression, and an anti-correlated EMT-like gene module, marked by ZEB1 and VIM expression (Extended Data Fig. 3 and Supplementary Fig. 1). H1299 cells showed variable expression of a module including ID1, ID2 and ID3, WI38 fibroblasts variably expressed a Collagen/fibronectin gene module, and A549 cells showed continuous variation of several additional gene modules (Extended Data Fig. 4 and Supplementary Table 3). Taken together, these findings showed that at least within 9-10 cell divisions, all examined cell populations featured a strong component of clonal and cell-cycle independent transcriptional stability.

## Characterizing a continuum of clonal epithelial identities in HCT116 cells

Both single cells and clones of HCT116 cells showed a consistent distribution of epithelial gene expression (**Fig. 1e**) and expression of genes linked with EMT (**Fig. 1h**), adhering to the Luria-Delbrück principle of clonal memory. This suggested this system can be an effective model to test the possible genetic or epigenetic basis for clonal transcriptional memory in our cancer cells. Genes linked with high or low *EpCAM* expression (**Fig. 1i**, **Supplementary Fig. 1d** and **Supplementary Table 4**) defined two continuous and anti-correlated spectra of epithelial and EMT-like identities across all clones (**Fig. 1j** and **Supplementary Fig. 1e**). At the extreme lowend of this spectrum we identified 4% of the clones (and consistently 4% of the single cells) with particularly high expression of *VIM* (defining a population we denote below as VIM-high clones).

The continuum of epithelial and EMT-like transcriptional states were both correlated with specific transcription factors (TFs), suggestive of possible regulatory networks diversifying the memorized clone states (**Fig. 1k**).

To elicit specific molecular mechanism that support such memory, we performed genetic, transcriptional and epigenetic analysis on single-cell clones propagated for up to 168 days. We first generated hundreds of clones that were profiled after 10 and 18 days at low resolution (**Fig. 2a,b**). We then selected six clones representing high and low *EpCAM*-expressing states, as well as a VIM-high state, and followed them for additional 150 days. For these clones we performed exome sequencing in two time points (d = 78, 168) and discovered that the high mutational load in the mutagenic HCT116 system gave rise to polymorphisms that are not shared between clones, even if their epithelial signatures are similar (**Fig. 2c**). Exome analysis of clones selected using a similar strategy from two additional cancer cell lines (**Extended Data Fig. 5**) revealed highly robust genetic subclones underlying one transcriptional module in H1299 cells, but no evidence for genetic basis for variation in all other transcriptional modules detected.

To further characterize clonal transcriptional dynamics, we next sampled 7,590 single cells from the six selected HCT116 clones after 33, 62, 98 and 148 days. We modeled the transcriptional space of single cells in these clones (**Fig. 2d**) and tracked evolving variation over this space within the clonal populations. This first reconfirmed the clonality of epithelial identity in the HCT116 system, showing some clones (1d12, 4e1, 7b11) maintain a distribution of high and low *EpCAM*-expressing states across many dozens of cell divisions (**Fig. 2e,f**). Clonal stability was however imperfect, as expected from a non-genetic mechanism, and we observed decrease in epithelial gene expression progressively in clone 3b3 and a reciprocal effect in clone 7a2 (**Fig. 2g**). Together, these experiments support the idea that epithelial transcriptional heterogeneity in the system has a non-genetic basis, permitting (slowly) drifting identities, and convergence of multiple clones toward similar transcriptional fates.

# Replication-linked loss of methylation underlies genome-wide clonal epigenetic diversity

We next performed single-cell analysis of global DNA methylation profiles (1,022 cells at low depth) and targeted methylation profiling (enriching over 70-fold the coverage for 318,783 select sites) of 251 9-days old HCT116 clones (**Extended Data Fig. 6a-d**) for which transcriptional data were also available (**Supplementary Table 5**). We focused initially on global genome methylation dynamics at low- and high-CpG content loci (LCG and HCG). Interestingly, LCG sites, that represent the majority of genomic territories and are highly methylated in normal cells, showed

surprising quantitative variation in average methylation for both clones and single cells (**Fig. 3a** and **Extended Data Fig. 6e-k**, 72-83.5% for single cells, 68.3-78.9% for clones). A specific small subpopulation of cells and clones (3.7% and 4.7%, respectively) showed major LCG hypomethylation (46.5-62%), and was shown to coincide with the VIM-high clone population we defined based on gene expression (**Fig. 3b**). But beyond this small subpopulation, we identified only weak negative correlations between gene expression and LCG methylation (**Fig. 3c** and **Supplementary Table 6**). In contrast to the lack of expression correlation, we observed lower clonal methylation (**Fig. 3d**) and higher inter-clonal variance (**Fig. 3e**) for LCG sites associated with late time of replication, compared to early replicating LCG sites. Overall, these data are consistent with several recent reports<sup>17</sup>, suggesting variation in LCG methylation in cancer may originate from the accumulation of replication-linked epimutations. Such epimutations are more frequent in late-replicating domains and may cause progressive loss of methylation in loci that are originally highly methylated. The process is pervasive, and is initially unlinked to transcriptional perturbation or transcriptional memory. It can however predispose specific genes to stochastic de-repression as we demonstrate below.

## High CpG content sites show clonal instability and turnover linked with cell proliferation

High CpG-content sites (HCG) are normally protected from methylation and are observed at CpG islands within gene promoters and distal regulatory elements. Similarly to LCG sites, we observed significantly high inter-clonal variation in HCG methylation levels (**Fig. 3f**). But unexpectedly the two global methylation trends were uncorrelated (**Fig. 3g**) and HCG methylation correlated extensively to gene expression (**Fig. 3h**). HCG methylation lacked association with genomic time of replication at regions with overall low methylation (**Fig. 3i,j**). Many of the top HCG correlated genes, both negative and positive, were associated with cell proliferation (**Supplementary Table 6**), suggesting possible linkage between clone proliferation rate and poor maintenance of HCG methylation.

We next used our target set of enhancers and promoters (**Extended Data Fig. 6l-n**) to study the intra- and inter-clonal distribution of methylation at higher depth. We contrasted two models of methylation transmission per locus (**Fig. 3k**): one assuming continuous methylation turnover within a growing clone, leading to convergence to a clonal distribution that is independent from the founding cell's epigenome, and the other assuming single cells are sampling two epi-alleles and transmitting them to all offsprings. Using data on all clones, we assayed the degree of clonal coherence of each locus in our target set (**Fig. 3l**) defining "hot" (marked red) and "cold" loci (marked blue) as those showing high turnover or high degree of clonal persistence, respectively.

Remarkably, we observe average methylation in "cold" sites to be correlated with LCG methylation, and average methylation in "hot" sites to be correlated with HCG methylation (**Fig. 3m**).

Importantly, the unexpected variation and independence of the HCG and LCG methylation signatures we characterize here are observed pervasively in TCGA tumor methylation data (Extended Data Fig. 6o-q). In summary, we demonstrated two types of methylation regimes that globally affect most CpG loci in the genome. The first involves replication-linked progressive hypomethylation in LCG sites that are normally methylated. The second involves epigenetic instability at high CpG content sites that are normally unmethylated, in correlation with perturbed expression of many cell-cycle genes.

## Clonal DNA methylation signatures are correlated with epithelial transcriptional identity

To search further for an epigenetic basis for the broad transcriptional epithelial identity spectrum we observed above, we normalized methylation of enriched enhancers and promoters given clones' global HCG and LCG methylation levels (Extended Data Fig. 7a-c) and clustered the normalized profiles to reveal additional inter-clonal epigenomic structure (Extended Data Fig. 7d). Two of the observed gene clusters were associated with the epithelial identity spectrum (Extended Data Fig. 7e,f). We therefore screened directly for promoters with differential methylation between high and low *EpCAM*-expressing clones (**Fig. 3n**). We detected strong anticorrelation between expression and methylation in EpCAM-high and EpCAM-low promoters (red and blue points, Fig. 3n,  $P = 8 \times 10^{-5}$ , D = 0.3, Kolmogorov-Smirnov (KS) two-tailed test). Weak hypomethylation was observed in the *EpCAM* locus itself (P = 0.01,  $X^2 = 6.5$ , chi-squared test), and stronger reduction in methylation was shown for additional promoters of induced genes (Fig. 30). We expanded this analysis to putative enhancer loci (Extended Data Fig. 7g), identifying 53 and 30 hypomethylated enhancers in EpCAM-high and EpCAM-low clones, respectively (Supplementary Table 7). For example, a hypomethylation hotspot observed for putative enhancer in chromosome 20 is correlated with EpCAM-high expression for at least 5 genes within 400 kb around it (Extended Data Fig. 8a,b). Interestingly, the dynamics of CpGs in this enhancer are classified as "cold" by the model described above, supporting their possible role as epigenetic memory carriers (Extended Data Fig. 8c,d, see Supplementary Table 8 for hot/cold classification of all EpCAM-linked enhancer CpGs).

## **Genes that Escaped Mitotically Inherited Inhibition (GEMINIS)**

We next developed a screen for genes with high clonal expression in one or more clones, but very low expression in at least 90% of the other clones. These represented loci that escaped repression rarely, in a clonally stable fashion, and not within the context of a larger co-regulated gene module (see **Methods** and **Extended Data Fig. 8e**). We reasoned such genes may be strong candidates for *cis*-acting epigenetic control. We detected 206 rare clonal de-repression events of 98 different genes in 97 distinct clones and define these as Genes that Escaped Mitotically INherited Inhibition (GEMINIs) (**Fig. 4a**). We found GEMINIs are encoding for different functions, including transcription factors (RUNX2, GRHL2, ELF3, TCF4 and ATOH8), long noncoding RNAs (LOC100287225, BC037861), transmembrane proteins (TM4SF18, CHODL), phosphatases (CCDC8) and more. Interestingly, five of the GEMINIs (PAGE1, PAGE4, MAGEA1, MAGEA11 and MAGEB1) belong to a subset of Cancer-Testis Antigens (CTAs), previously annotated as germline restricted genes that are de-repressed in different cancers<sup>18</sup>. Average expression of GEMINIs in clones and cells is generally consistent (**Fig. 4b**), but GEMINIs are noisier in single cells compared to control genes with similar expression levels (*P* = 0.01, KS test, **Fig. 4c**).

GEMINIs constitute a set of de-repression events that are uncorrelated to each other and uncorrelated to observed *trans*-factors regulatory changes. We hypothesized that the regulation of their clonal expression signature may therefore be mediated by *cis*- epigenetic effects. Promoters of GEMINIs were found to be natively more methylated in comparison to matched controls with similar CpG content and overall expression levels ( $P = 2 \times 10^{-6}$ , D = 0.35, KS test). GEMINIs tended to be positioned away from constitutively expressed genes ( $P = 8 \times 10^{-8}$ , D = 0.22, KS test) (**Fig. 4d**). Most importantly, data from clones showed nearly 50% reduction in promoter methylation in *cis* for de-repressed loci ( $P = 9 \times 10^{-13}$ ,  $X^2 = 51$ , chi-squared test, **Fig. 4e,f**), compatible with a mono-allelic loss of methylation and suggesting allele-specific derepression. In addition, we observed GEMINIs expression is more associated with the "cold" LCG clonal methylation trend (**Fig. 4g**, left) and show lower correlation with the "hot" HCG trend (**Fig. 4g**, right, see also **Extended Data Fig. 8f**). These data identify a class of loci in which DNA methylation is not only tightly correlated to de-repression in *cis*, but is likely driving it. Stochastic loss of methylation at these sites (and subsequent clonal maintenance of this hypomethylation) stabilize a de-repressed state in specific clones.

## **GEMINIs** are induced in methylation-impaired cells

We next profiled 4,523 single cells and 319 short-term clonal populations derived from methylation-impaired HCT116 cells (double knockout of DNMT1 and DNMT3B DNAmethyltransferases or DKO). As expected, DKO cells displayed severe whole-genome reduction in DNA methylation when compared to parental HCT116 single cells (Extended Data Fig. 8g-m). We observed a high degree of concordance of cell-cycle dependent transcriptional states between the DKO system and the wild-type (Extended Data Fig. 8n-p), and also identified cellcycle independent co-variating gene modules in DKO cells and clones, including the same epithelial gene module observed in wild-type cells (Extended Data Fig. 9a-c). DKO cells were biased toward lower EpCAM module expression compared to wild-type cells, but the distribution in single cells and clones was conserved (Extended Data Fig. 9d-f and Supplementary Fig. 2). This suggested the EpCAM module gene expression remains largely coordinated even when methylation is impaired. We also find that DKO cells maintain a gene module that includes ZEB1 and several HOXB genes, which was anti-correlated to expression of the epithelial module, as well as instability of Interferon type-I (IFN-I) and DNA damage response (DDR) gene modules (Supplementary Fig. 3). Together the data show transcriptional memory for co-regulated gene modules in HCT116 can be independent from a fully functional DNA methylation machinery.

Using loci identified as GEMINI in the wild-type, we next tested if repression of GEMINI is maintained in the DKO system. Strikingly, 40 out of the 98 GEMINIs found in WT were pervasively expressed in DKO clonal populations (FDR corrected q-value < 0.001, KS test, Fig. 4h, right panel). De-repression was significant compared to control genes that were repressed in the wild type ( $P = 2 \times 10^{-3}$ , D = 0.2, two-tailed KS test, **Fig. 4i**). Notably, de-repression was in general observed more often for low CpG content repressed promoters that were methylated in the wildtype, compared to controls that were not methylated ( $P = 4 \times 10^{-3}$ ,  $X^2 = 8.1$ , chi-squared test, Fig. 4j). We profiled 251 additional clones on the background of a single DNMT3b knockout (Supplementary Fig. 4a), showing milder, but still noticeable de-repression in some of the GEMINIs (Supplementary Fig. 4b). We hypothesize that the in cis de-repression we have observed in our clonal populations applies for other systems as well, and specifically may be occurring in colon tumors. We thereby analyzed bulk RNA-seq experiments of nearly 400 colorectal adenocarcinoma tumors from the TCGA database, focusing on repressed genes and computing the ratio between maximally observed de-repression of each gene to its expression in the top five percentile of the cohort. Interestingly, we found that rare de-repression is indeed enriched for genes identified in our screen as GEMINIs ( $P = 5 \times 10^{-5}$ , D = 0.28, KS test, Fig. 4k).

## Correlated clonal de-repression within topological associating domains

Some of the clonally stable gene modules we detected above were defined through co-expression of spatially linked genes within a single topologically associating domain (TAD). We therefore screened systematically for clonal co-expression of genes within TADs in the HCT116 system (Methods and Extended Data Fig. 10a-c), using shuffled controls and accounting for possible chromosomal dosage effects using pBAT coverage statistics (Methods and Supplementary Fig. 5). This approach identified 149 genes in 89 TADs with statistical support (FDR < 0.25) for intra-TAD co-expression, including the embryonic globin genes *HBE1* and *HBG* (Extended Data Fig. 10d-f and Supplementary Fig. 6a,b). Similar analysis in WI38 and A549 cells identified additional putative cases for TAD-linked clonal co-expression (Extended Data Fig. 10g-I, Supplementary Fig. 6c-f and Supplementary Table 9). In-depth analysis of the co-expression around the betaglobin genes in HCT116 cells (Fig. 5a), suggested a bimodal clonal distribution of transcription in these loci (denoted as HB-high and HB-low). We observed weak de-repression of several OR genes in HB-high clones (Fig. 5b,c). De-repression in the HB TAD was uncorrelated with the EpCAM expression signature (Fig. 5d), but showed remarkable long-term stability (Fig. 5e). Another notable spatial clonal expression pattern involved 11 keratin associated proteins (KRTAP) organized in a 130-kb cluster on chromosome 17 (Fig. 5f-j). HCT116 DKO clones showed conserved bimodal HB and KRTAP TAD expression (Supplementary Fig. 6g,h). Importantly, de-repression of the globin and KRTAP clusters was observed in a large fraction of the clones (47% and 23%, respectively). Nevertheless the transcriptional output per single cell was low for most cells even in de-repressed clones (Fig. 5k,l). Moreover, in some cases TAD clonal output was also correlated with the expression of an additional factor in trans (as for TCF25 and the HB expression, Extended Data Fig. 10m,n), suggesting de-repression requires a permissive TAD state but also additional driving factors. In summary, spatially correlated clonal expression patterns raise the hypothesis that TADs can toggle between clonally inactive and active states, where in the active state multiple genes within a TAD are predisposed for derepression, and in the inactive state complete repression is secured. This mechanism can thereby diversify clonal expression patterns of spatially (and functionally) linked genes.

#### **DISCUSSION**

We studied the clonal propagation of transcriptional and epigenetic identity in cancer cells using a Luria-Delbrück design in which single-cell distributions of RNA expression and DNA methylation are compared to the analogous distributions in hundreds of short-term clones. We characterized stable transcriptional heterogeneity in all studied cell types, which in all except for one case appeared to be unlinked with a sub-clonal genetic structure. In colon cancer cells, we identified clonally stable variation in epithelial gene expression and a reciprocal variation in expression of EMT genes. Longitudinal analysis of select clones using additional single-cell analysis in four time points showed that slow drift in the epithelial/EMT identity is observed for up to 150 days. These data pave the way to analysis of the dynamics underlying non-genetic clonal memory, and the role of epigenetic mechanisms implementing it.

Non-genetic clonal memory is first (and perhaps foremost) implemented by gene regulatory circuits using feedback to enable semi-stable transcriptional states, but our studies here show that in addition to these mechanisms, stable epigenetic variation between cells contribute to clonal memory. DNA methylation is the best understood clonally persistent epigenetic mark. It is generally assumed methylation propagates from mother to daughter cells through the housekeeping methylation machinery. Our data on single cells and clonal methylation distributions suggest more complex modes of methylation dynamics. First, most of the genome, being rarely targeted by epigenetic modulators or other trans-factors, indeed shows high degree of epigenetic persistence that is driven by the housekeeping machinery. Despite this persistence, we show here variability in average methylation between single cells and clones at such loci. We hypothesize this is the result of the accumulation of epimutations with replicative age<sup>17</sup>. Second, and unexpectedly, we show that methylation of CpG islands (or high CpG content loci in general) is governed by a different and much more dynamic process. Clones and single cells are shown to be variable in their ability to protect CpG islands from methylation accumulation. We define this molecular phenotype as "epigenetic instability" and show it to be uncorrelated with the replicationage effects that are observed in the rest of the genome.

We previously suggested that methylation dynamics can be governed by clonal persistence in somatic cells and turn-over in embryonic stem cells<sup>6</sup>. This model can now be generalized, predicting that in cancer cells protection from DNA methylation at promoters and regulatory elements primarily involves turnover. CpG island hypermethylation can then result from change in the efficiency of the protective turnover mechanism, rather than from progressive accumulation of epimutations with time.

The regulatory consequences of the two types of proposed methylation dynamics are global – affecting thousands of loci together rather than tinkering gene expression of individual genes or programs. But the combination of stochastic and regulated changes in methylation at loci with low turnover rate gives rise to an epigenetic landscape that is specific to each clone, resulting in many opportunities to diversify transcription in a clonally stable fashion. One example demonstrated here is the methylation change at "cold" promoters and enhancers of epithelial/EMT genes, that may participate in regulating the spectrum of epithelial identities in colon cancer cells. A second example involve GEMINIs - cases of rare, but clonally stable promoter hypomethylation and gene de-repression events that are enriched for CTA loci. A similar mechanism of clonal epigenetic change predisposing transcriptional dynamics is suggested by analysis of correlated de-repression of genes residing in the same TAD. In this case, TADs can repress effectively their associated genes when present in their "epigenetically closed" state, but are allowing their occasional de-repression in clones that switch to the "epigenetically open" state.

Together, the data here highlight the ability of non-genetic mechanisms to stably diversify transcription in cancer cells. This in turn can facilitate opportunities for adaptation that run in parallel (and in coordination) to the genetic evolutionary process driving carcinogenic transformation. An immediate challenge is to evaluate the impact of such non-genetic mechanisms in vivo. Initial analysis of TCGA data<sup>19</sup> suggests DNA methylation dynamics can be generalized from the ex vivo systems we analyzed here to tumors in vivo. Additional profiling of tumors using new single-cell epigenomic and multi-omic strategies will be essential for fully appreciating how clonally stable epigenetic changes drive long-term regulatory programs and how such changes may affect tumor function, response to therapy, or metastasis.

#### **DATA AVAILABILITY**

Raw and transformed data are available at GEO and SRA (Accession GSE144357).

#### **CODE AVAILABILITY**

Metacell<sup>20</sup> code is available in <a href="https://tanaylab.github.io/metacell/">https://tanaylab.github.io/metacell/</a>. Source code, additional metadata and a vignette exemplifying cell-cycle normalization with MetaCell are available at <a href="https://github.com/tanaylab/Meir\_et\_al\_nat\_gen\_2020\_clonemem">https://github.com/tanaylab/Meir\_et\_al\_nat\_gen\_2020\_clonemem</a>.

#### **AUTHORS CONTRIBUTION**

ZMe and AT designed the study with help from ZMu. ZMe performed experiments with help from ZMu. EC helped with MARS-seq and automation. ZMe and AT analyzed data with help from AL. ZMe and AT wrote the manuscript with input from all authors.

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#### **CONFLICT OF INTEREST**

The authors declare no conflict of interest.

## **METHODS** (including online methods)

#### **Cell lines**

HCT116 parental colorectal cell line (HD PAR-033), as well as KO (DNMT3B<sup>-/-</sup>, HD R02-023) and DKO (DNMT1<sup>exons3-5/exons3-5</sup>; DNMT3B<sup>-/-</sup>, HD R02-022) were obtained from Horizon Discovery Ltd., Cambridge, UK. A549 cells were obtained from the NCI-60 cell panel, NCI-H1299 cells were purchased from ATCC (ATCC<sup>®</sup> CRL-5803<sup>™</sup>) and WI38-hTERT embryonic lung fibroblasts (WI38) were generated as described in Milyavsky et al.<sup>21</sup>. All Cells were cultured on 100 × 20 mm culture dishes (Corning, 353063) in heat-inactivated medium and split at a ratio of 1:10 every 2–3 days using 0.05% trypsin-EDTA solution C (Biological Industries, Israel; 03-053-1B). McCoy's 5A medium (Biological Industries, Israel; 01-075-1A) was used for HCT116 WT, KO and DKO, DMEM medium (Dulbecco's Modified Eagle Medium, Biological Industries, Israel; 01-050-1A) was used for WI38, DMEM/F-12 (HAM) medium (Biological Industries, Israel; 01-170-1A)) was used for A549 cells and RPMI medium (Biological Industries, Israel; 01-100-1A) was used for NCI-H1299

cells. All media were supplemented with 10% Fetal Bovine Serum (GibcoTm FBS, 10270-106), 0.4% Penicillin-Streptomycin (Biological Industries, Israel; 01-031-1C) and 1% L-glutamine (Biological Industries, Israel; 01-020-1A). Modified medium was filtered through a 0.22-µm filter (Corning, 430769) prior to culture.

#### Single-cell isolation and clonal expansion

In parallel to sorting of single cells for MARS-seq analysis, individual cells were selected by FACS (SORP FACSAriall cell sorter) using a 70-µm diameter nozzle into 96-well culture plates (Corning-Costar, 3596), which already contained 100 µl of conditioned media in each well. Conditioned media were taken from cycling populations of the respective cell line, centrifuged and filtered through a 0.22-µm filter (Corning, 430769). Subsequent to sorting, plates were transferred immediately to a 37 °C incubator for culture. 48 hours post sorting, 100 µl of conditioned media were added to each well. Clonal expansion was terminated when populations reached approximately 500 cells (8-10 cell divisions), in average after 9-10 days for HCT116 parental, NCI-H1299 and A549 cells, 12 days for parental and DKO cells and 21 days for WI38 cells. All clonal populations were examined by microscope prior to their harvesting. Small clones (~200 cells or less) were discarded, as well as wells that were suspected of having more than one founder cell in them.

#### Harvesting clonal populations

In order to process each clonal population for MARS-seq<sup>22</sup> (transcriptome analysis) and PBAT-capture (methylome analysis), clones were detached by 30 µl trypsin-incubation for 2 min. (HCT116 parental, A549, H1299 and WI38 cells) or 4 min. (HCT116 DKO cells) in 37 °C, washed by 100 µl PBS (Biological Industries, 02-023-1A) and suspended in 10 µl ultra-pure water (Biological Industries, 01-866-1A) and 0.005% RNase inhibitor (RNasin plus, Promega, N2611). The 10 µl suspension of each clone was then transferred to a skirted twin.tec 96-well PCR plate, (Eppendorf, 0030 128.648) on dry ice. When a 96-well plate was filled, replicates of 1 µl from each clone (two for HCT116 parental and HCT116 DKO, and four replicates for H1299 and A549 cells) were transferred into a barcoded (8 nM oligo-dT barcodes concentration) twin.tec 384-well (Eppendorf, 0030 128.508) MARS-seq plate that already contained 2 µl of lysis-buffer (0.1% Triton, 0.005% RNasin and ultra-pure water). The 384-well plates were then transferred into

80°C until its cDNA library preparation by MARS-seq. The remaining 8 μl from each HCT116 clone were kept in -20 °C for further methylome profiling by PBAT-capture. For WI38 clones, the 10-μl suspension of each clone was used for 6 technical replicates in 384-wells barcoded MARS-seq plates. For A549 and H1299 clones, out of the 10 μl of each clone – 5 μl were immediately transferred into 96-wells culturing plates (Corning-Costar, 3596) that contained 200 μl of the respective growth media and the rest were used for MARS-seq. A549 and H1299 clones that were further growing in 96-well culture plates were then routinely split until selection (according to their transcriptome profiles obtained by MARS-seq), resulting in six A549 and seven NCI-H1299 clonal populations assayed by whole-exome sequencing.

#### Single-cell transcriptome profiling

For MARS-seq scRNA protocol, single cells were sorted into 384-well plates containing 2 µl of barcoded RT primers in concentration of 8 nM in each well. Downstream library preparation was done according to Jaitin et al. 2014, and by using randomized UMI sequence of 8 base-pairs (allowing maximal count of ~65k UMIs per gene per well). For NCI-H1299 scRNA, we used data of '5 10x libraries described in Brocks et al. 2019<sup>23</sup>. HCT116 WT and HCT116 DKO scRNA libraries based on 10x Genomic platform, were generated using Chromium™ Single Cell 3' Library & Gel Bead Kit v2 (PN-120237), Chromium Single cell 3' Chip Kit V2 (PN-120236) and Chromium i7 Multiplex Kit (PM-120262), following the manufacturer's instructions (10x Genomics®, Inc.). Two samples of 5,000 cells were loaded per each, HCT116 PAR and HCT116 DKO-033 libraries were sequenced paired-end 150 bp on Nextseq 500 to mean depth of 68,257 and 45,259 reads per cell for PAR and DKO cells, respectively.

## Multiplexed transcriptional analysis of clonal populations

Lysed clonal populations were manually transferred in two replicates of 1 µl into MARS-seq barcoded plates. Library preparation was identical to that of single cells, with the single exception of extending the initial evaporation time on 95°C before RT1 from 3 min to 4 min, in order to compensate for the additional volume in each well.

## Single-cell methylation data

We used data on HCT116 single-cell DNA methylation that was derived as controls for analyzing tumor samples (Mukamel et al., in preparation), using a variant of a previously described approach<sup>24</sup>. To quantify the distribution of average methylation in the founding single-cell population, we generated an ultra-low-depth library of 1,022 cells (1,045 – 52,740 uniquely mapped molecules per cell). For QC, we filtered verified rate of incomplete Cytosine conversion (or alternatively CHH methylation) is lower than 2%. We then binned CpG loci into groups according to the CpG content in the 500 bp around the locus, and estimated average methylation for independent bins (as shown in **Extended Data Fig. 9e**). The correlation between such bins allow some validation on the estimation noise of the average, which is verified to be robust as expected from the sampling depth of each cell. We note that for quantitative methylation analysis we rely on targeted analysis of clonal populations as described below.

## **Targeted methylation analysis**

We used the PBAT-capture protocol combining Post-Bisulfite Adaptor Tagging (PBAT) and hybridization to an RNA probe library (capture) as described in Mukamel et al. (in preparation), which we briefly summarize here. The remaining 8  $\mu$ l of each clonal population was treated with 4  $\mu$ l RNase-A (20 mg/ml, ThermoFisher Scientific 1910121, diluted 1:3 in water, 30 min. in 37 °C), 3  $\mu$ l Proteinase-K (20 mg/ml, ThermoFlsher Scientific 25530-49, diluted 1:1 in water, 30 min. in 65 °C), then stored in -20 °C in 96-well plate.

65 μl of Lightning Conversion Reagent (ZYMO REASEARCH, D5032-1) was added to 15 μl of each clone in each well, and bisulfite conversion was performed according to the EZ-96 DNA Methylation-Lightning MagPrep kit (D5046, Zymo) manufacturer's protocol, where 100 μl of M-Binding buffer and MagnaBeads mix in a ratio of 9:1 was added to each sample. Converted DNA samples were eluted in 38 μl elution buffer and were subjected to the first tagging step (PBAT1).

First-strand synthesis was performed in a 50  $\mu$ l reaction that contained 5  $\mu$ l NEB buffer 2, 2  $\mu$ l dNTPs (10 mM) and 4  $\mu$ l PBAT1 oligo (4  $\mu$ M, **Supplementary Table 10**). Prior to addition of enzyme, the reaction mix was incubated at 65 °C for 3 min. followed by 4 min. at 4 °C and pause break to add 2  $\mu$ l Klenow exo– (M0212L, NEB). Graduate increase of temperature +1 °C/15 sec. to 37 °C for 90 min, followed by heat inactivation of the enzyme at 70 °C for 10 min. Removing excess of oligonucleotides was done by adding 1.5  $\mu$ l exonuclease I (M0293L, NEB) for 45 min.

at 37 °C, followed by DNA purification by 0.8× Agencourt Ampure XP beads (A63881, Beckman Coulter) and elution in 38 µl.

Second strand synthesis was performed similarly to the first strand synthesis, using 4  $\mu$ I PBAT2 oligo (4  $\mu$ M, **Supplementary Table 10**), incubating at 95 °C for 45 sec, followed by 4 min at 4 °C and pause at 4 °C for adding 2  $\mu$ I Klenow exo- during incubation at 4 °C for 5 min., +1 °C/15 sec. to 37 °C, 37 °C for 90 min., 70 °C for 10 min. Excess of oligonucleotides removed by adding 1.5  $\mu$ I exonuclease I, cleaning with 0.8× beads and elution in 22  $\mu$ I elution buffer. Tagged products were then amplified for library preparation in 14 PCR cycles in 25  $\mu$ I Kapa HiFi Hot start ready mix kit (KK2601, KAPABIOSYSTEMS) following the manufacturer's protocol, and by using 3  $\mu$ I (from a 10  $\mu$ M stock) of 1:1 PBAT\_PCR\_For and PBAT\_PCR\_Rev primers mix (**Supplementary Table 10**). The reaction mix was then cleaned with 0.7x beads and eluted in 25  $\mu$ I 10 mM Tris pH 8.

Pools of indexed libraries in same molarity (30-40 clones in each pool) were concentrated by 1x Agencourt Ampure XP beads cleanup to a volume of 10 µl (~75 ng from each clone, to 2.5 µg in pool). Each 10 µl pool was subjected to Mybait capture reaction (MYcroarray) according to the manufacturer's instructions and by adding the following modifications: The amount of baits in the hybridization mix was replaced by 2 µl baits and therefore the hybridization mix was in aliquots of 16 µl. Captured products were washed 4 times for 10 min. with washing buffer 2.2. Following amplification, the product was purified by 0.7x beads, eluted in 10 µl elution buffer and was subjected to an additional round of capture, 4 washes and 14 cycles of amplification. Final libraries were pooled and sequenced on an Illumina Nextseq system using the 150 bp high output sequencing kit.

For PBAT-capture we used a probe library designed for colon cancer analysis (Mukamel et al., in preparation), targeting 47,871 loci, out of which 9,275 were designed to enrich for promoters, 17,120 for enhancers and 21,476 sampled different non-functional epigenomic context for control.

#### Mapping and low-level analysis of Capture-PBAT reads

Sequenced reads were aligned to the GRCh37 (hg19) reference genome (UCSC, February 2009) using an in-house script based on Bowtie2. In cases where the two reads ends were not aligned in a concordant manner, the reads were discarded. Reads that were mapped to the same genome coordinates where marked as duplicates and were used only once. The level of incomplete

conversion was estimated from loci in CHH contexts, and we have validated all libraries show lower than 2%.

#### Identifying copy number aberrations using PBAT data

To assess copy number trends in the data we used the total number of PBAT reads that were not mapped to on-target loci. For global comparison of HCT116 single cells to normal tissues, or of subset of the clones as shown in **Supplementary Figure 5**, we collected coverage statistics on bins of 1 Mb bins that showed less than 100 mapped reads in both pools were discarded. The ratio between normalized coverage was then computed and visualized over the chromosome coordinates. For analysis of individual clones, we used binning to entire chromosomal arms and compared the  $log_2$  ratio between the fractions of reads mapped to each arm in each clone, to the median of the coverage of all clones. This analysis was done separately for the two batches of clones we processed.

## Mapping and low-level analysis of single cells and clonal RNA-seq data

Mapping MARS-seq reads was performed using the standard MARS-seq pipeline as described<sup>22</sup>. We filtered wells with less than 1,000 UMIs from further processing and verified the estimated level of empty well contamination was less than 4% in all cases. For MARS-seq libraries of clonal populations, we pooled UMIs from replicate experiments (wells) for 300 HCT116 WT, 319 DKO, 251 KO, 266 NCI-H1299, 208 A549 and 67 WI38 clones, according to the 384-well plate design of each. For 10X single-cell analysis we used the CellRanger software for de-multiplexing, barcode processing and alignment of 10X reads. The total UMI threshold was determine by CellRanger at 4,053 for HCT116-WT cells and 2,995 for DKO cells. Further analysis of the 10X data was performed using Metacell as described below.

# scRNA cell-cycle modelling

We used the MetaCell package (Baran et al.<sup>20</sup>), to generate cell-cycle models for the HCT116 WT, H1299, WI38 and HCT116 DKO data. For HCT116 as an example, we selected 722 candidate genes with high correlation to either MKI67, PTTG1, GINS2 or ACTB and clustered them using analysis of the correlation of UMI for single cells following downsampling to 6,422

UMIs. Supervised analysis of the derived metacells identified 320 genes correlated with either S or M-phase genes (See summary of S and M core genes of all cell lines in Supplementary Table 1). We generated a restricted count matrix including only these 320 cell-cycle genes and used the Metacell pipeline with parameters K=100 and minimal meta-cell size=50, without outlier filtering, to generate cell-cycle metacells. These were annotated using analysis of the total expression of the S-phase and M-phase gene modules. A similar protocol was used for all other cell lines. Full assignment of cells to Metacells throughout this manuscript can be found in the source code companion to this paper.

## Identification of cell-cycle independent gene correlation

In order to identify cell-cycle independent transcriptional dynamics in single cells, while avoiding various biases emerging when normalizing gene counts, we developed a randomization approach aiming at estimation of the degree of correlation between any gene pair, given the cell-cycle trend alone. This was implemented by first constructing the Metacell balanced KNN similarity graph based on expression of cell-cycle genes only. This was followed by randomization of a downsampled UMI matrix by drawing the molecule count for each cell and gene from one of its 20 most similar neighbors (which were defined based on cell-cycle genes alone). We identified only genes that had at least one high correlation prior to randomization (>0.1 in HCT116 WT, >0.12 in WI38, >0.16 in H1299 and >0.15 HCT116 DKO), but had lower maximal correlation following cell-cycle randomization (at least 0.02 reduction in HCT116 WT and WI38, 0.03 in HCT116 DKO and 0.1 in H1299 cells). See **Extended Data Figures 2k,m,o, and 9a,** and a full summary of all cell-cycle independent genes in **Supplementary Table 2**.

We then performed further normalization to consider sampling depth. Given the sparse nature of the scRNA data, the probability of any pair of genes to display high correlation score inversely depends on their sampling variance, which can in turn be predicted by the total number of UMIs captured for the gene. To take this into account, we analyzed for each gene g the correlations of its original (un-normalized) UMI vector with all other genes' UMI vectors. We sorted all genes based on their total number of UMIs, and computed the empirical trend predicting correlation to gene g from total number of UMIs in any other gene. This was done using a simple rolling mean analysis with window size of 101 genes. We subtracted this empirical trend to report for each gene pair (g, g') two version of a normalized correlation value (based on g and g' empirical trends). This analysis is exemplified for the EpCAM gene in HCT116 WT cells in Extended Data

**Figure 2i**. Depth-adjusted correlation of cell-cycle independent genes in cells and clones are summarized in **Supplementary Table 3**.

#### Metacell analysis of long-term clones scRNA-seq

7,590 HCT116 single-cell profiles were obtained by MARS-seq for six selected clones in four different time points. We generated a Metacell model for these cells using 68 gene features that were selected to have normalized variance higher than 0.3 and at least 50 UMIs in a downsampled matrix. We filtered from the feature list any of the 320 cell-cycle genes described above. Metacell was then applied using K=100 to create cell graph and minimal metacell size = 30 cells, using 600 bootstrap iterations and generating 55 metacells ranging in size between 85 to 281 cells. These were visualized using the Metacell 2D projection as shown in the text.

# Whole-exome sequencing (WES) of single-cell derived clonal populations

DNA from approximately 5-10 × 10<sup>6</sup> cells of the seven H1299, six A549, five HCT116 (at two different time-points) single-cell derived clonal populations, as well as of polyclonal cell population that was grown in parallel to clones, was extracted by Quick-DNA™ Universal Kit (Zymo, D4069). Exome capture sequencing of extracted DNA was done by IDTxGen lockdown human panel (Admera Health, LLC South Plainfield, NJ). Initial processing of the PE 150-bp reads was done by GATK v3.5 and mapped to hg19 reference genome, following GATK best practices (<a href="https://software.broadinstitute.org/gatk/best-practices/">https://software.broadinstitute.org/gatk/best-practices/</a>), and using Mutect2 module for SNP detection in each one of the clones in comparison to matching polyclonal population, when screening only homozygous sites in polyclonal samples (allowing default parameter of up to 3% variant allele frequency in each polyclonal reference population).

## **Defining genomic and epigenomic features**

For all analyses, we used the fraction of CG in the closest 500 bp as a measure for CG content. Promoters were defined as regions spanning up to 500 bp of a gene's transcription start site (TSS). For definition of enhancers, we used ChIP-seq data of H3K4me1 in HCT116 cells (GSM945858), and classified them as regions that reside up to 500 bp from a peak of H3K4me1 (95<sup>th</sup> percentile) and located at least 2 kb away from TSS to classify genomic time-of-replication

we used ENCODE data of Repli-seq experiment performed on MCF7 cells (GSM923442), relying on the considerable conservation of this effect between cell lines. For **Figure 3e,j**, we considered CpGs with value lower than 60 (28<sup>th</sup> quantile of CpGs) as CpGs of late-replicating regions. Genomic regions that reside in a distance of 100 bp or greater from our PBAT probe-set intervals were classified as off-target regions, for example in Extended Data Figure 6b (see the source code companion for detailed information on probe-set intervals).

# Comparing models for persistent methylation patterns of individual loci in clonal populations (Fig. 3k,I)

We analyzed 12,536 CpGs covered at least 6 molecules in at least 60 clones. These were downsampled to retain exactly 6 methylation calls for 60 clones, such that the inter-clonal methylation variance could be computed robustly without coverage bias. We modeled the expected variance in methylation for loci govern by clonal dynamics involving no memory by assuming independent sampling from a variable that is methylated with probability p (i.e.  $V^{mix}(\hat{p}) = 6\hat{p}*(1-\hat{p})$ , where  $\hat{p}$  is the empirical average methylation). A model assuming perfect memory for a locus with empirical methylation average  $\hat{p}$  was estimated by sampling methylation of two founding epi-alleles independently with probability  $\hat{p}$  (i.e. assuming Hardy-Weinberg equilibrium), and then sample six methylation calls for the clone randomly from the selected two epi-alleles. The variance of this two-step process (computed by exhaustive summation on the number of reads sampled from the first epi-allele) was defined as  $V^{pers}(\hat{p})$ .

Given the two models, we computed for each CpG a *deviation from persistency* value by subtracting its empirical inter-clonal variance from its expected  $V^{pers}(\hat{p})$ . As shown in Figure 3I, we defined "cold" CpGs as those that show minor deviation from the expected inter-clonal variance according to the perfect memory model (absolute deviation is lower than 0.3), and "hot" CpGs as those showing lower variance than expected variance according to this model (difference higher than 3). We limited this screen to partially methylated CpGs, where we have the most power to distinct between the methylation dynamics regimes (average methylation in the population p between 0.3 and 0.7).

## Testing for differential expression and identifying GEMINIs

GEMINIs were defined as genes that are almost always repressed but show one or few clones with significantly high level of expression. To screen for GEMINIs we used clone RNA profiles that were downsampled to 10k UMIs for WT, DKO and KO clones. We identified for each gene the maximal expression level over clones and computed its fold-change to the 90% expression percentile, assigning a "GEMINI score" to every gene (as shown in **Fig. 4a**). GEMINIs were restricted to basal expression of up to 1 UMI / 10k UMIs, where basal expression of each gene computed as the average expression across all clones, except the 10% of clones with highest expression of it. GEMINIs also required to have at least 4 UMIs in their maximal clone, and at least 5 UMIs overall. In practice, genes with the highest ratio showed total expression of 5-590 UMIs (median of 46), and maximal expression of 4-106 (median 7). We then assigned a "repressed" or "de-repressed" states for each clone regarding each GEMINI, requiring at least 50% of maximal expression level for a particular GEMINI to define a "de-repressed" state in clone. The number of clones showing de-repression was 1 for 69 genes, 2 for 11 genes and over 2 for 18 genes.

# Hi-C analysis and Shaman normalization.

We re-analyzed HCT116 data from Rao et al. 2017, using the Shaman package as described (Olivares-Chauvet et al. 2016, Bonev et al. 2017). TADs were called using computation of insulation profiles as described before (Bonev et al. 2017, see example in Extended Data Fig. 10c). Overall 3,690 TADs were defined ranging in size between 51 kb - 38 Mb (median = 397 kb), and we focused our analysis to TADs with at least four annotated TSSs. Each TAD's total expression was defined as the total UMIs per clones from genes with their TSSs within it. We computed correlation between each gene's expression to all TADs total expression across all clones. To compute the correlation between a gene to the TAD it resides in ("TAD Auto-correlation") we first subtracted the gene's expression from its total TAD's output. To screen for surprising gene-TAD expression correlations, we subtracted from the TAD Auto-correlation score the 20th highest correlation of the gene with any TAD:

$$TAD_{Auto_{correlation}score_{gene}} = TAD \; Auto \; correlation - \; TAD \; top \; correlation\_20_{gene}$$

To visualize normalized contact maps of specific genomic regions (as in **Fig. 5f**), we used SHAMAN package to first normalize local cis-decay over observed contacts by using the shaman\_shuffle\_and\_score\_hic\_mat() function, and then to plot the resulted map with function shaman\_plot\_map\_score\_with\_annotations (). SHAMAN source code is available at <a href="https://github.com/tanaylab/shaman">https://github.com/tanaylab/shaman</a>.

## TCGA analysis

All RNA-seq and methylome datasets that were downloaded from the TCGA repository (<a href="http://cancergenome.nih.gov/">http://cancergenome.nih.gov/</a>) and re-analyzed in this manuscript are listed in **Supplementary Table 11**.

#### FIGURE LEGENDS

# Figure 1 | A Luria-Delbrück design for testing transcriptional and epigenetic memory.

a, Schematic overview of our experimental design. Green dashed arrows: culturing steps. Black dashed arrows: sorting of single cells into conditioned media. Non-dashed arrows/lines: processing steps. WES: whole-exome sequencing. b, Left: Expression of selected genes over 3,255 HCT116 single cells (columns) grouped into metacells (top labels) according to similarity in cell-cycle gene expression. Center: 2D Projection of the cell-cycle metacell model. Metacells (large ovals) are color coded according to the expression intensity of the cell-cycle marker MKi67, cells are shown as small gray dots. Right: Comparing expression of M-phase and S-phase genes (Supplementary Table 1) for cells and metacells. c, As b, for 3,584 NCI-H1299 single cells. d, As in b, for 1,172 WI38 single cells. e, Normalized expression (UMI per 100k UMIs) distribution in HCT116 cells and clones of epithelial (EpCAM) and S-phase gene modules (as detailed in Supplementary Table 1 and Supplementary Table 3). f, As e, for ID module and M-phase in NCI-H1299 cells. q, As e, for Collagen module and M-phase in WI38 cells. h, Distribution of VIM expression (log<sub>2</sub> of UMI per 10k UMIs) in HCT116 single cells (left) and clones (right). i, log<sub>2</sub> expression fold changes for genes enriched in EpCAM high clones (blue, top 30% of clones, n = 51) and EpCAM low clones (red, lowest 30% of clones, n = 51), after exclusion of 11 VIM-high clones. For all genes shown here, FDR corrected q-value << 0.001, chi-squared test. j, Shown is the total UMIs for genes positively (blue) and negatively (red) enriched in EpCAM-high clones, in clones grouped based on expression of the *EpCAM* gene module. **k**, Density plots of expression in EpCAM-high (black line), EpCAM-low (red line) and 11 VIM-high clones (orange line) for selected genes.

## Figure 2 | Long-term clonal maintenance of Epithelial and VIM-high transcriptional states.

a, Expression (log<sub>2</sub> count per 10k UMI), of VIM (left panel) and the EpCAM module (right panel) for clonal populations that were sampled twice, after 10 and 18 days (> 15 k UMI per sample in both). b, Expression (note linear scale) of VIM and the EpCAM module in six clones selected for further longitudinal analysis. Dashed horizontal line represents median expression over all clones sampled after 10 (grey, n = 59) and 18 (dark grey, n = 59) days. c, Analysis of genetic kinship between HCT116 clones. Text in each cell shows the absolute count of shared SNPs between two clones. Upper bars show VIM (red) and EpCAM gene module (blue) expression (pooled single cells RNA in the closest time point). d, Metacell 2D projection for single-cell RNA-seq data from longitudinal analysis of six clones. Colors represent the level of EpCAM expression. e, Average expression of VIM (upper panel) and epithelial genes (lower panel) in tracked clones over six time-points (clonal RNA-seg at day = 10 and 18; pool of single-cell RNA-seg at day = 33, 62, 98 and 148). See panel g below for the number of cells at each time-point. Error-bars represent SE of binomial sampling, based on total sampled UMIs per clone per time-point. f-h, For each clone (row), showing total epithelial and VIM transcriptional output per cell by time-point (f); 2D projection of clones' single cells, coloring according to the EpCAM module expression intensity (g) and the changes in VIM and the EpCAM gene module expression distributions over time (h).

#### Figure 3 | Hot and cold dynamics of clonal methylation.

**a,** Distribution of clonal average methylation in low CpG content (LCG, 0-3% CG content) loci (observed vs. shuffled control, excluding VIM-high clones as defined in Fig. 1h). **b,** *VIM* expression by clonal LCG methylation. VIM-high clones are colored in red. **c,** Distribution of Spearman correlations between LCG methylation and gene expression over all genes. Controls are based on shuffling clonal LCG values. **d,** Clonal LCG methylation in early- and late-replicating genomic domains. **e,** Distributions of early and late replicating loci methylation over clones, indicating by s the standard deviations. **f,** As in **a,** for high CpG content (HCG, 7-15% CG content) loci. **g,** clonal LCG vs HCG average methylation, indicating lack of correlation. **h,** As in **c,** for HCG

methylation. i, As in d, for HCG methylation (but excluding loci with overall average methylation higher than 0.3). j, As in e, for HCG methylation. k, We simulated two alternative methylation dynamics in clonal population assuming zero memory (left, mixture model) and perfect memory (right, persistency model). I, Shown are inter-clonal methylation variance vs. average methylation across well covered loci (running median is defined by a gray curve). Blue and orange lines depict the variance predicted by the persistency and mixture models (see Methods). Red and Blue dots mark partially methylated loci showing empirically "hot" (high turnover, mixture model) and "cold" (low turnover, persistency model) dynamics, respectively. m, We grouped 192 clones with sufficient coverage by their LCG (left) or HCG (right) methylation (minimal group size = 54, excluding VIM-high clones). Boxplots depict distribution of average methylation in hot and cold loci across the groups. In all boxplots throughout this manuscript we used R version 3.5.3 defaults for boxplot() function – where middle line indicates median, box limits are quantiles, and whiskers are 1.5 x IQR. Kolmogorov-Smirnov two-tailed test, LCG-high to LCG-mid: D = 0.39,  $P = 3 \times 10^{-5}$ <sup>5</sup>. LCG-low to LCG-mid: D = 0.49,  $P = 9 \times 10^{-7}$ . HCG-high to HCG-mid, D = 0.31,  $P = 3 \times 10^{-3}$ . HCG-low to HCG-mid: D = 0.29, P = 0.01. HCG-high to HCG-low: D = 0.39,  $P = 1 \times 10^{-4}$ . n, Pooled average methylation of promoter CpGs in EpCAM-high (n = 51) and -low (n = 51) clones, highlighting promoters with up-regulated (blue) or down-regulated (red) expression in EpCAMhigh clones (D = 0.3,  $P = 8 \times 10^{-5}$ , KS two-tailed,  $n_{blue} = 257$ ,  $n_{red} = 88$ ). **o**, Promoter methylation in clones showing high (n = 51) and low (n = 51) EpCAM expression. Bars showing average expression and error-bars represent SE of binomial sampling. Chi-squared test, all P values < 2 x 10<sup>-15</sup>. Panels below bars indicate chromosomal coordinates and show average methylation of covered CpGs in EpCAM-high (blue dots) and -low (light blue dots).

## Figure 4 | Screening for Genes that Escaped Mitotically Inherited Inhibition (GEMINIs).

**a,** 98 GEMINIs were selected based on genes with low basal expression (less than 1 UMI / 10k UMIs) and high maximum de-repression (red dots, see Methods). For each gene, showing averaged basal expression across HCT116 clones (x-axis, see Methods), and GEMINI score indicating rare de-repression in few clones (y-axis, see Methods). **b,** Density plot of overall GEMINI pooled expression per cell (red line) and clone (black line). **c,** Distributions of normalized variances in single cells ( $\log_2$  of variance-to-mean ratio) for 98 GEMINIs and 969 randomized matched controls with similar expression levels and promoter CG content. Two-tailed KS test, D = 0.21,  $P = 9 \times 10^{-4}$ . **d,** Distributions of genomic features of GEMINIs and randomized controls matched for expression levels. High-exp gene: TSS within top 20 expression percentiles. Two-

tailed KS test, from left to right: CG (D = 0.13;  $P = 8 \times 10^{-3}$ , compared to matched-controls of expression only), Distance (0.22;  $8 \times 10^{-8}$ ), Methylation (0.35;  $2 \times 10^{-6}$ ). **e**, We annotated each clone with a "repressed", or "de-repressed" state regarding each one of the GEMINIS (see Methods). Bars showing average methylation of pooled GEMINI promoters in their repressed (black bar, n = 20,797) and de-repressed clones (grey bar, n = 201). Error-bars represent sampling SE, based on total methylation calls in de-repressed or repressed clones. Chi-squared test,  $\chi^2 = 51$ ,  $P = 9 \times 10^{-13}$ . f, Average methylation of selected GEMINIs in their de-repressed clone. g, Distribution of gene expression correlation to clonal LCG (blue boxplots, as in Fig. 3c), and HCG (red boxplot, as in Fig. 3h) for GEMINIs and for matched controls. Two-sided KS test, LCG (D = 0.19;  $P = 5 \times 10^{-3}$ ), HCG (D = 0.16; P = 0.035). h, Left panel: GEMINIs (rows) expression in 300 HCT116 WT clones (columns). Expression levels are normalized by maximal expression value of each of the 98 GEMINIs. Order of columns is determined by clonal LCG average methylation. Right panel: Expression of GEMINIs in DKO clones. Expression is normalized by the maximal expression of each GEMINI in its de-repressed WT clone (\*\*\* FDR corrected q-value < 0.001, KS test for comparison of GEMINIs expression in HCT116 WT and HCT116 DKO clones). i, Cumulative distribution for the fraction of repressed DKO clones for GEMINIs (red line) and for the matched randomized subset of control genes with similar CG-content and expression levels in WT (black line). De-repression threshold was set to half of the maximal normalized expression in WT clone. Two-sided KS test, D = 0.2,  $P = 2 \times 10^{-3}$ . j, Comparison of de-repression in DKO for methylated and unmethylated genes. Showing data only on genes that were considered repressed in HCT116 WT (< 0.25 UMIs / 10k UMIs in at least 80% of the clones). Pooled methylation data on clones used to define methylated (> 0.9, n = 365) and unmethylated (< 0.9, n = 4,160) promoters. Chi-squared test: GEMINI/meth:  $\chi^2$  = 5.9 P = 0.015; GEMINIS/unmeth (32;  $2 \times 10^{-8}$ ); meth/unmeth (8.1;  $4 \times 10^{-3}$ ). **k**, Cumulative distribution of genes showing rare derepression (see Methods for definition of GEMINI score) in 399 colon adenocarcinoma RNA-seq samples (obtained from TCGA). GEMINI score compared expression in maximal tumor to 95'th expression quantile. Showing only genes that were generally repressed in both HCT116 clones and TCGA samples. CTAs represent 77 annotated Cancer-Testis Antigens. Two-tailed KS test, Geminis (D = 0.28,  $P = 5 \times 10^{-5}$ ), CTA (0.42; 2 × 10<sup>-9</sup>).

## Figure 5 | Evidence for in-TAD memory.

a, Normalized expression of genes spanning the beta-globin TAD on chromosome 11. Heatmap is showing gene (rows) expression over clones (columns), normalized to the gene's median

expression across clones. Top bars annotate clones as HB-high and -low, (VIM-high clones are excluded). Spatial map using one letter encoding (left of heatmap) is shown at the bottom, also indicating TAD borders as grey dashed vertical lines. b, Pooled average expression in HBE1-high (n = 76, blue) and HBE1-low (n = 77, black) clones defined in a. Error bars represent SE of sampling a binomial distribution. Chi-squared P values: HBG1,OR51B5  $< 2 \times 10^{-16}$ , OR51B4 = 5  $\times$  10<sup>-12</sup>, OR51B2 = 0.018, OR51I1 = 9  $\times$  10<sup>-8</sup>. **c**, Expression of embryonic (HBE1) and fetal (HBG2, HBG1) beta-globin genes (x-axis) vs. expression of adjacent cluster of olfactory receptors (yaxis). Dots represent clones and Spearman correlation is indicated here and in other figure panels. n = 168 clones covered by >100k UMIs. d, Expression of the HB (x-axis) and EpCAM (yaxis) modules, indicating lack of correlation. e, Temporal change in HB module expression for six clones, similar to Figure 2e. Error bars represent sampling SE, based on total sampled UMIs per clone per time-point. f, As in a, for KRTAP region on chromosome 17. Spatial map of genes is drawn on top, and the SHAMAN-normalized contact frequency map is depicted as a triangle respective to the region's linear coordinates (HCT116 Hi-C data obtained from Rao et al. 2017, see Methods). g, As in b, Bars indicate average pooled expression in LOC730755-high (brown, n = 50) and LOC730755-low (orange n = 55) clones defined in f. Error bars represent SE of binomial sampling. Chi-squared P values: KRTAP3-1, KRTAP3-2 < 2 x 10<sup>-16</sup>, KRTAP2-2 = 0.03, KRTAP4-12 =  $3 \times 10^{-16}$ . h, Comparing total clonal expression of four genes in the KRTAP2 subcluster (x-axis) and 8 genes in the KRTAP3, 4 sub-cluster (y-axis). Sample size as in c. i, Comparing clonal expression of KRTAP-associated genes (x-axis) and the EpCAM module (yaxis). j, Similar to e, for KRTAP-associated genes. k, Distribution of HB region expression (genes E-L in a) in clones and single cells. I, Distribution of KRTAP region expression (genes E-O in f) in clones and single cells. Vertical dashed lines in k-I indicate the overall normalized expression in clones (black) and cells (red).

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Figure 1 | A Luria-Delbrück design for testing transcriptional and epigenetic memory.

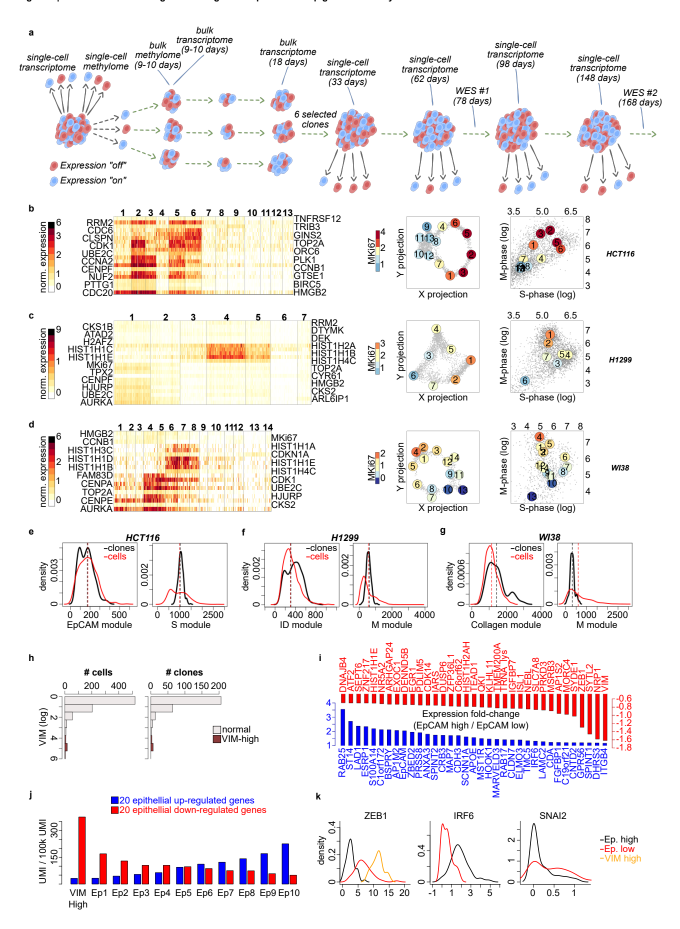


Figure 2 | Long-term clonal maintenance of epithelial and VIM-high transcriptional states.

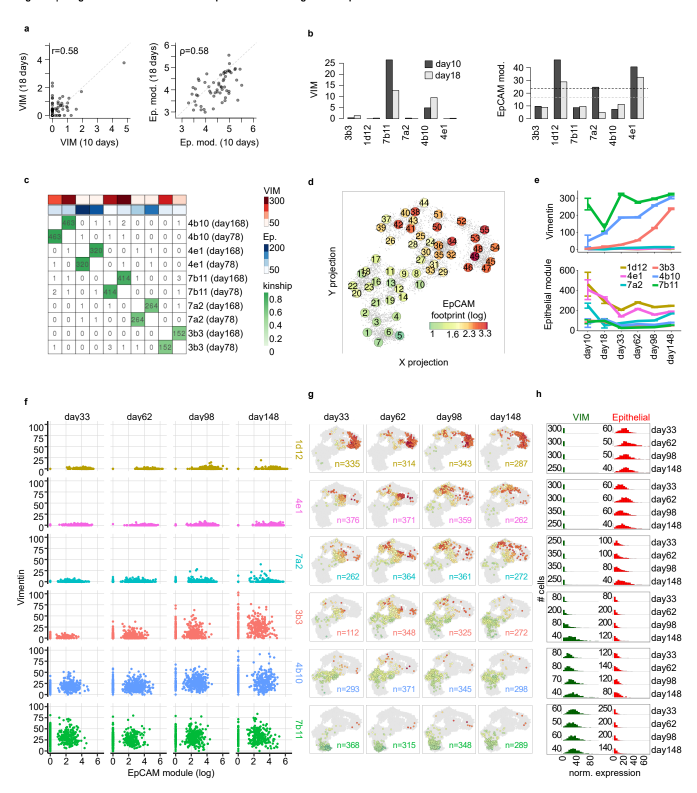


Figure 3 | Hot and cold dynamics of clonal methylation.

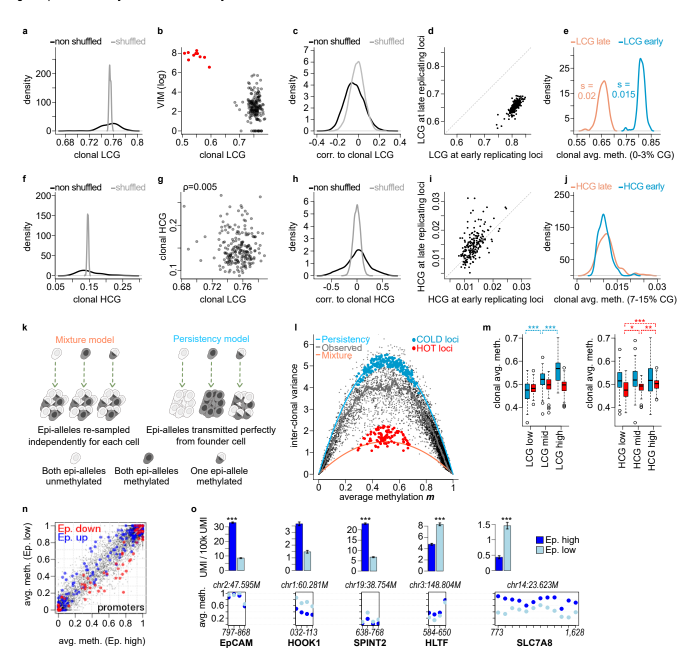


Figure 4 | Screening for GEMINIs.

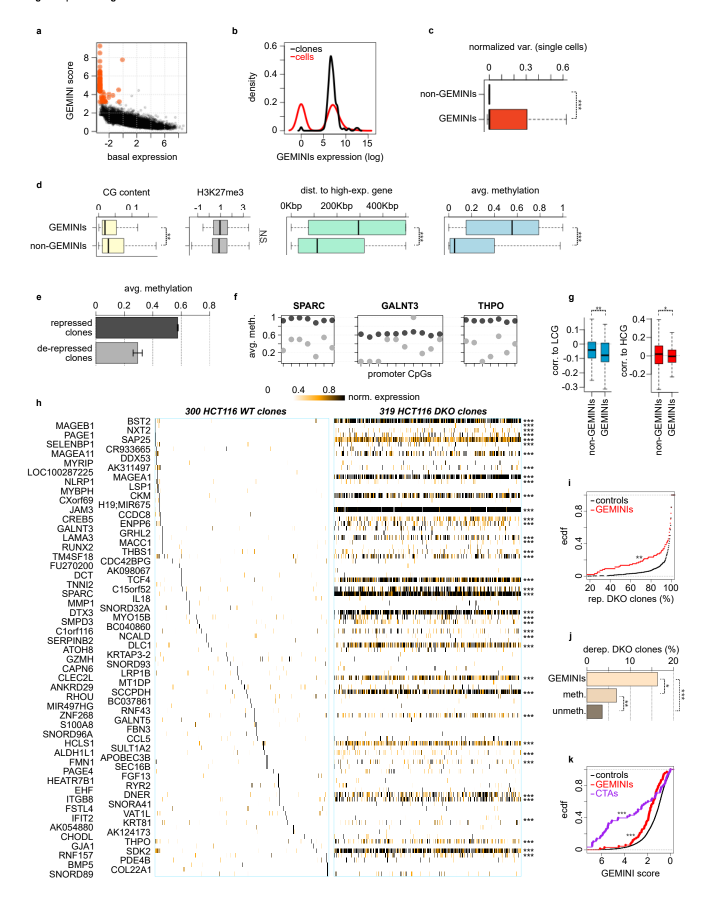
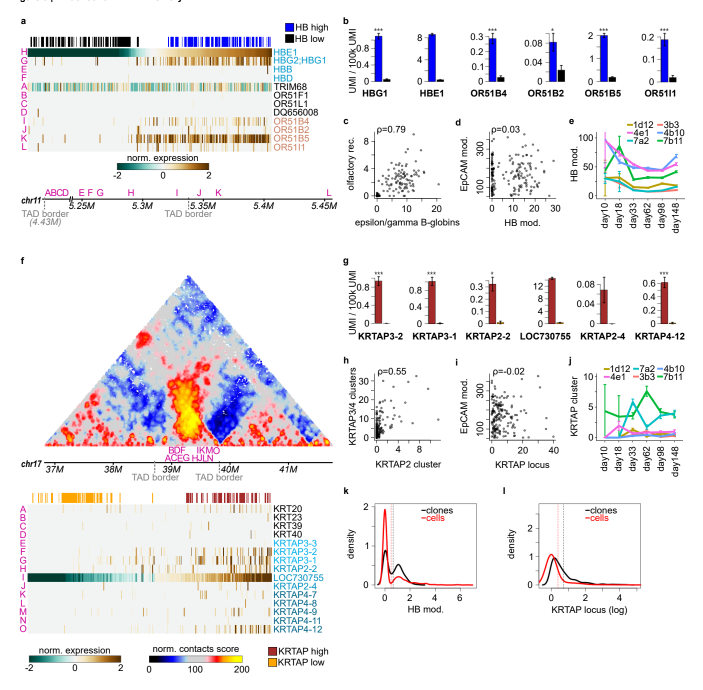
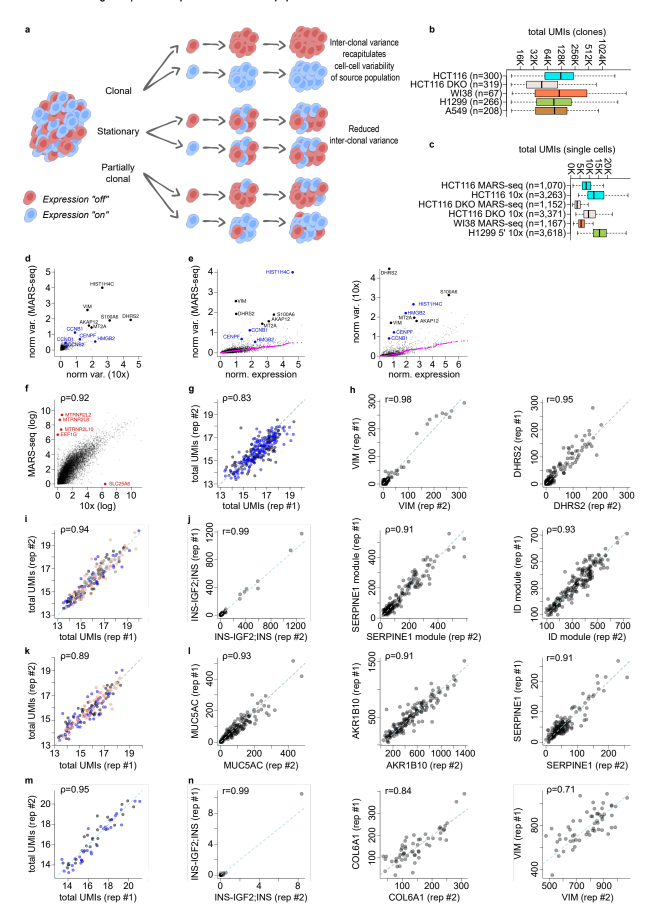
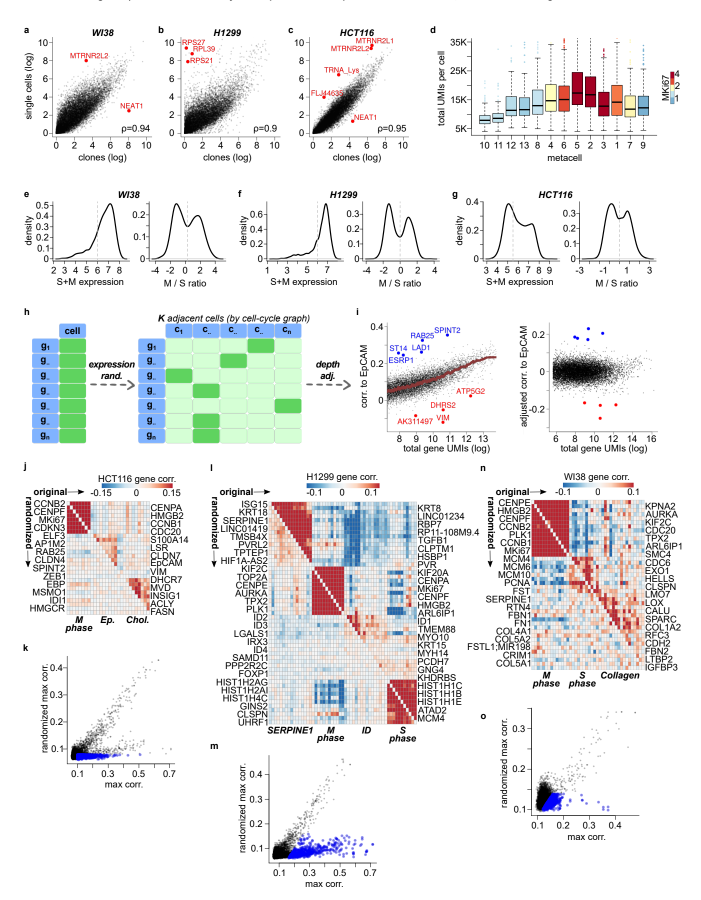
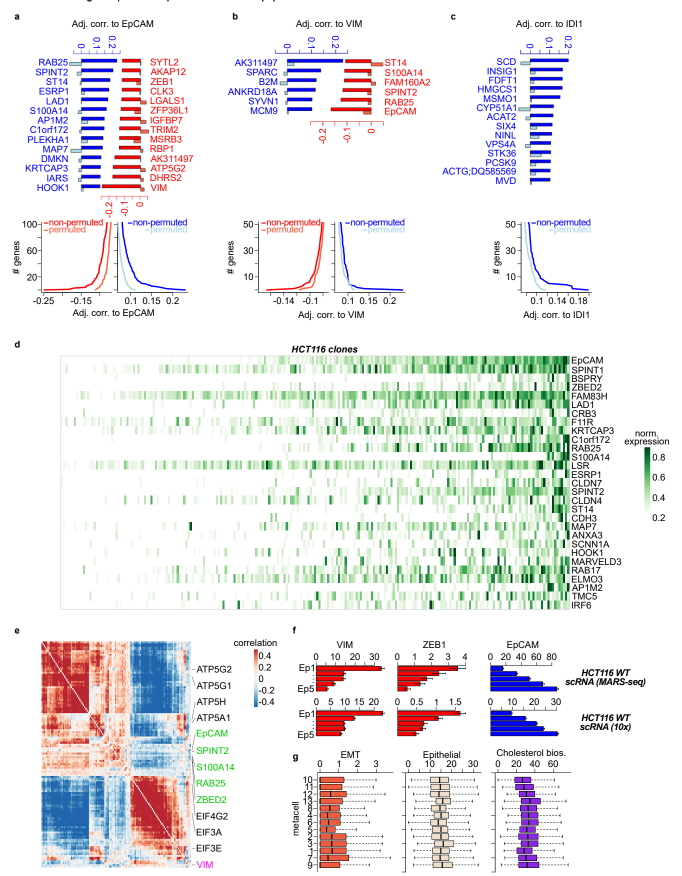


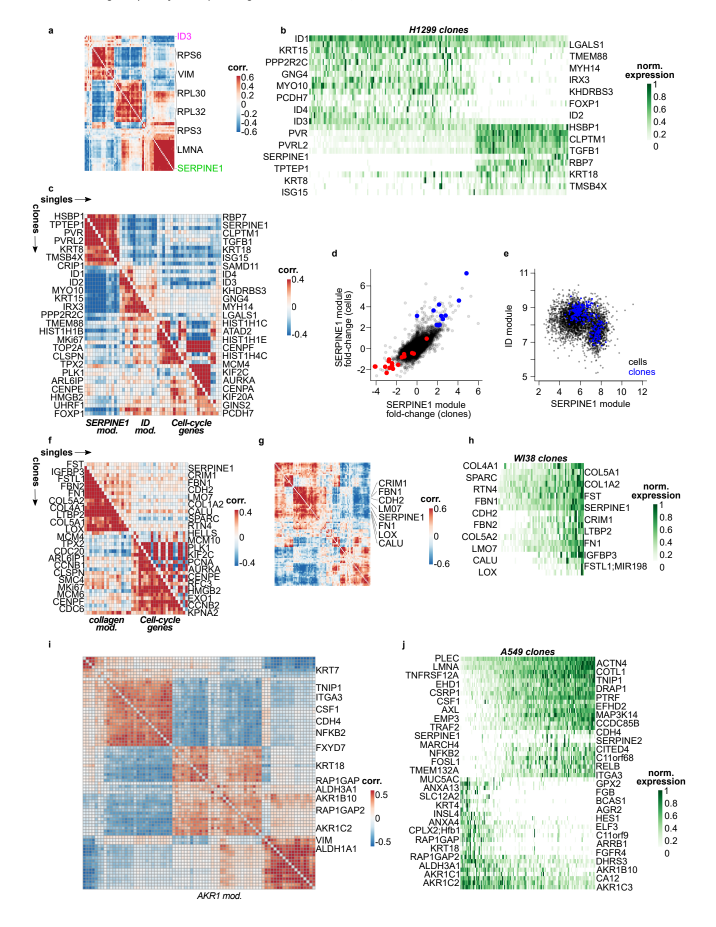
Figure 5 | Evidence for in-TAD memory.

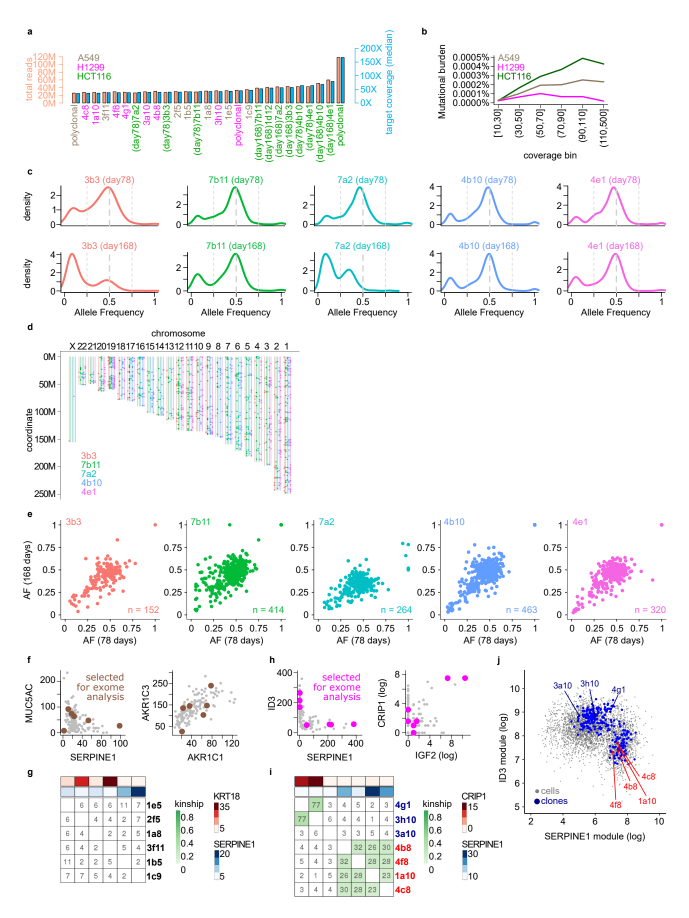


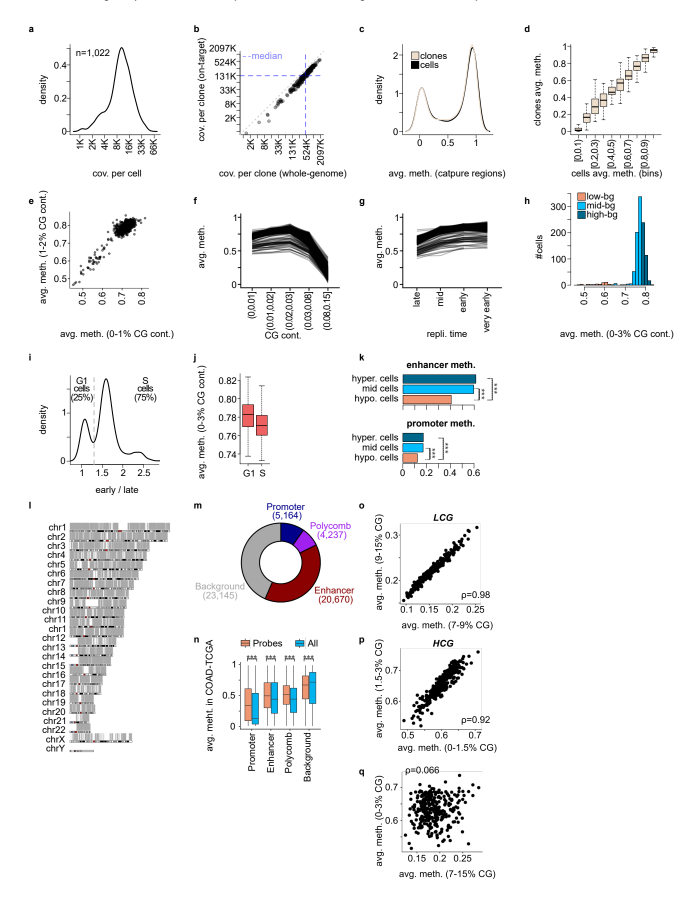


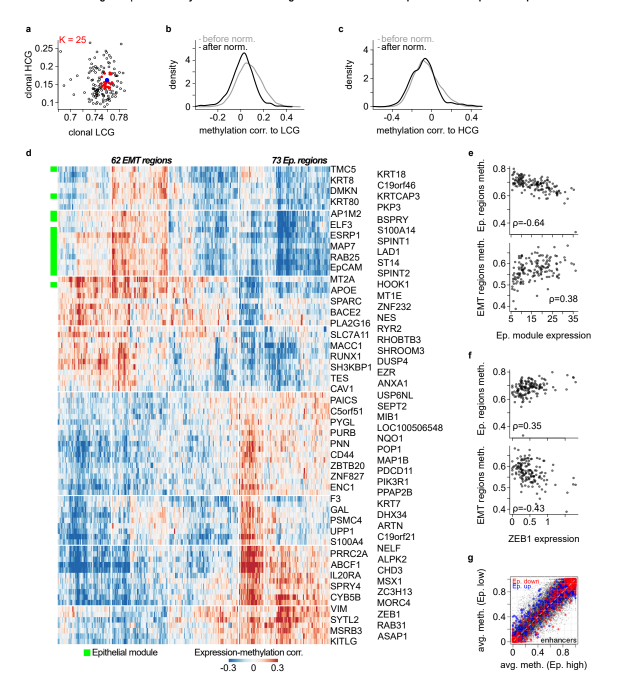


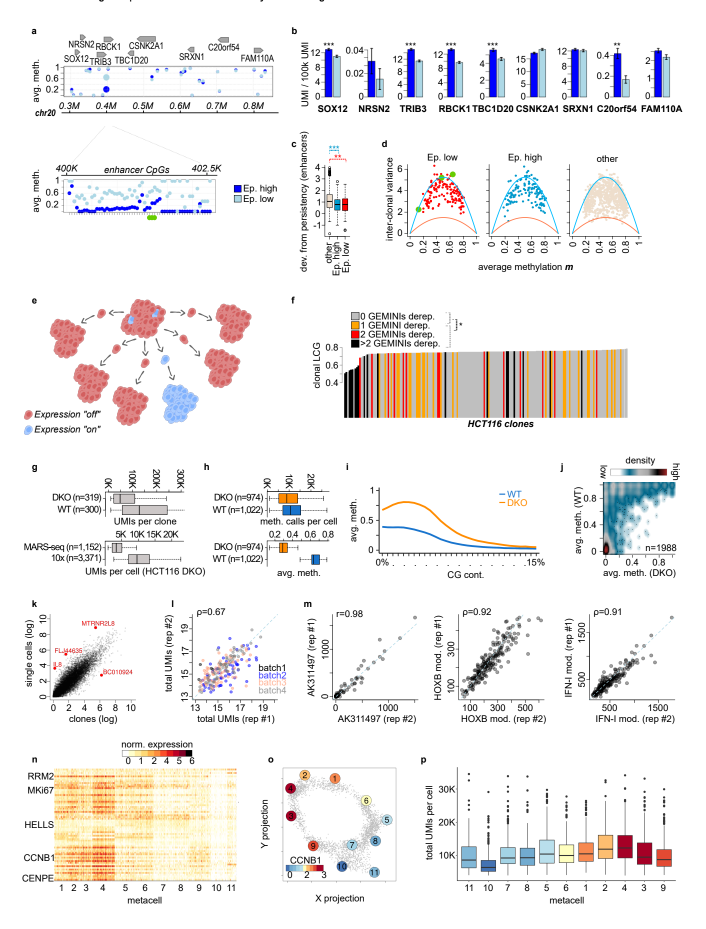




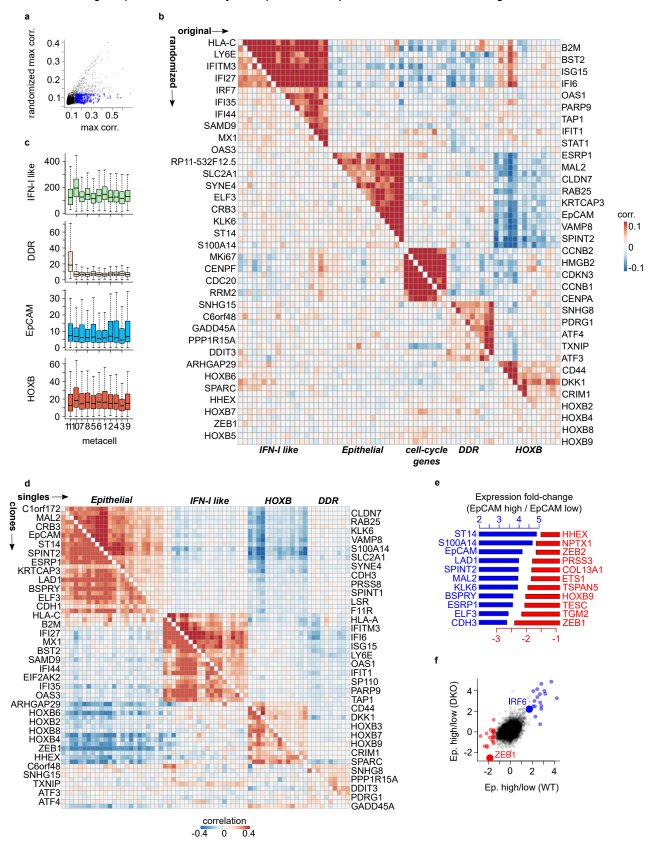


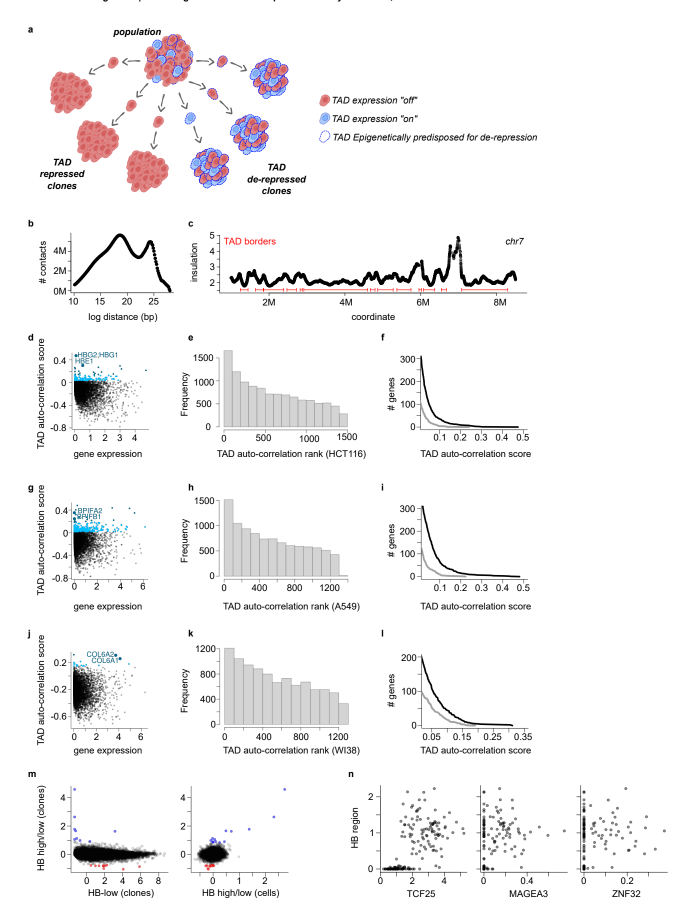


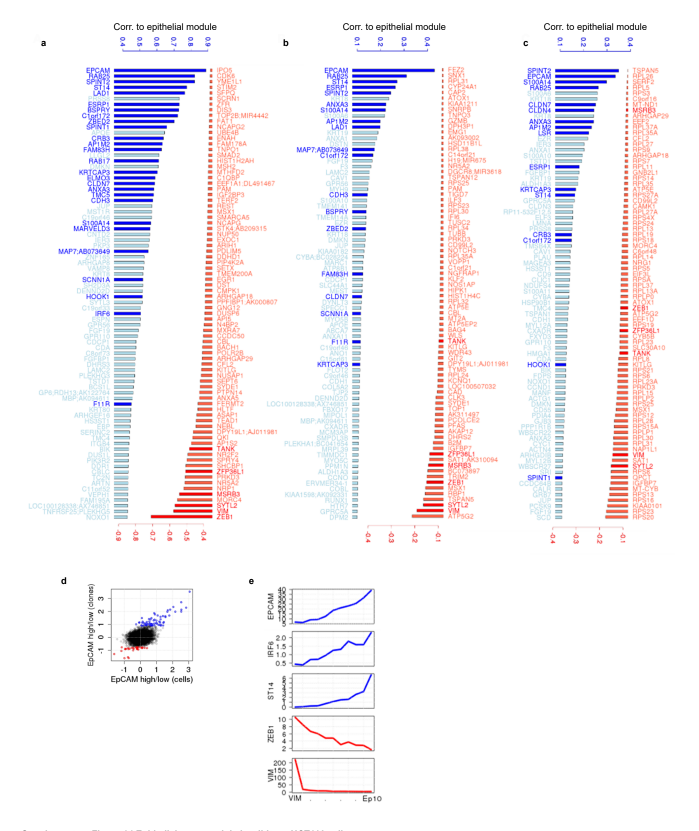




## Extended Data Figure 9 | Identification of cell-cycle independent transcriptional variance in DKO HCT116 single cells.

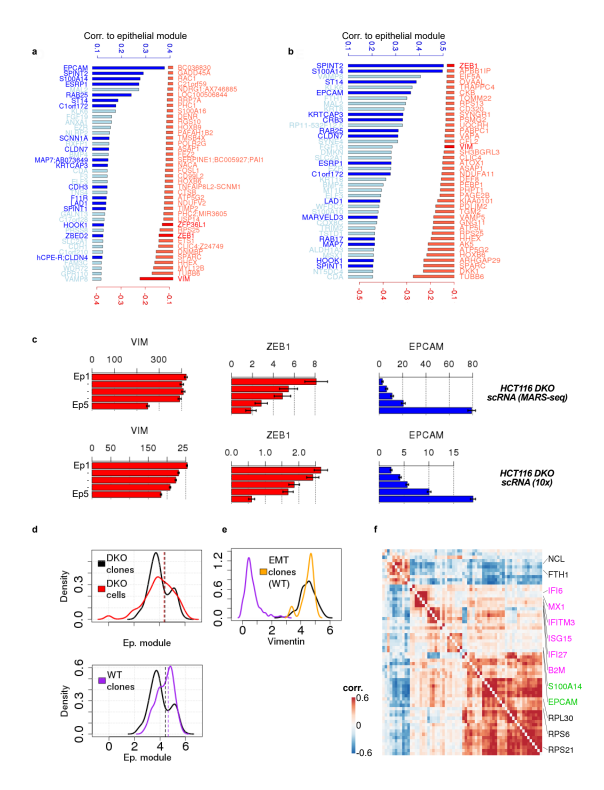






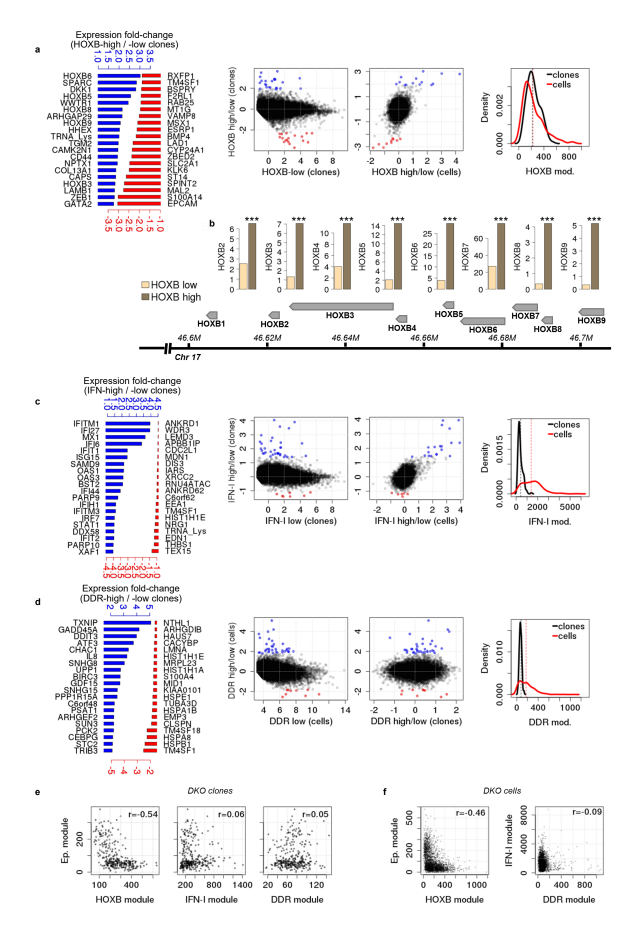
## Supplementary Figure 1 $\mid$ Epithelial gene module in wildtype HCT116 cells.

a-c, Showing highest and lowest Spearman's correlation of all genes to total expression coming from the epithelial gene module in 300 HCT116 clones (a), 1,070 single cell transcriptomes derived by MARS-seq (b) and 3,263 single cell transcriptomes derived by 10x (c). Genes annotated as part of the epithelial gene module (Supplementary Table 3) are labeled in dark blue, and their correlation to the gene module was computed after their exclusion from the module. Genes with consistent anti-correlation expression to epithelial genes are highlighted in red. d, For each gene, showing the log2-ratio of relative expression in high EpCAM (top 25% cells and 30% clones) and low EpCAM (lowest 25% cells and 30% clones) cells (x-axis) and clones (y-axis). e, Similar to Fig. 1j but showing line graph for specific genes in clones, binned by VIM-high (leftmost) and 10 groups of epithelial expression.



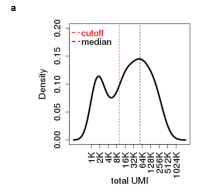
Supplementary Figure 2 | Epithelial gene module in DKO HCT116 cells.

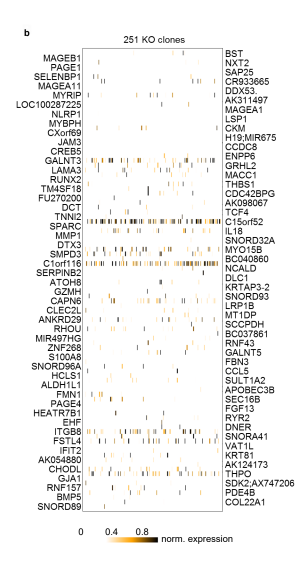
**a-b**, As in supplementary Fig. 1a-c, showing highest and lowest gene correlation to expression of the epithelial gene module in 1,152 HCT116 DKO single cells obtained by MARS-seq (a) and 3,371 single cell transcriptomes derived by 10x (b). Genes annotated as part of the epithelial gene module are labeled in dark blue, and their correlation to the gene module was computed after their exclusion from the module. Genes with consistent anti-correlation expression to epithelial genes are highlighted in red. **c**, As in Extended Data Fig. 3f, we grouped DKO cells obtained by MARS-seq (top) and 10x (bottom) into five bins based on expression of the EpCAM gene module (Ep5 consisting of cells with highest module expression). Bars showing mean expression of each bin for EpCAM gene (blue) and for genes negatively enriched in EpCAM high cells (red). Error-bars represents standard error of sampling a binomial distribution. **d**, Comparing distribution of EpCAM gene module expression between DKO clones and cells (top), and between in DKO and WT clones (bottom). **e**, Distribution of VIM expression in DKO clones (black line), VIM-high WT clones (orange line) and non VIM-high WT clones (purple). **f**, As shown in Extended Data Fig. 3e for HCT116 WT, clustering gene-gene correlations of all HCT116 DKO cell-cycle independent genes labeled in Extended Data Fig. 9a. Epithelial module genes are colored in green and IFN-I related genes are colored in magenta.



Supplementary Figure 3 | Additional profiling of cell-cycle independent gene modules in HCT116 DKO cells.

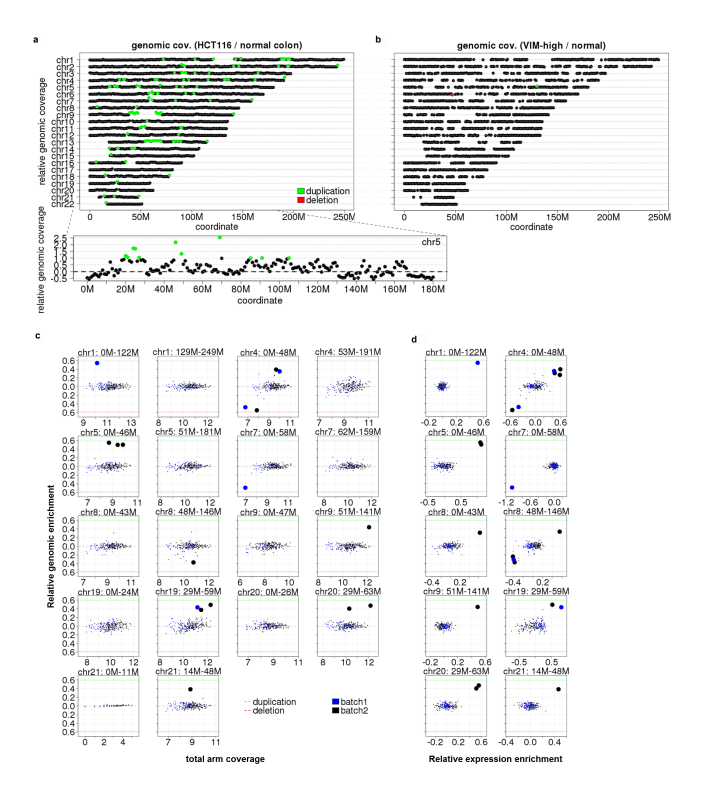
a, Clonal stability of HOXB gene module expression in DKO clones and cells. For differentially expressed genes shown in these figures FDR corrected q-value < 0.001, chi-squared test. nHOXB-high = 30, nHOXB-low = 28 biologically independent HCT116 DKO clones. b, Spatial map and expression in the HOXB genes cluster on chromosome 17. Chi-squared test, in all instances P < 2\*10-16. c, Interferon-I like gene module expression in DKO clones and cells. As done in A for HOXB gene module. nIFN-high = 51, nIFN-low = 51. d, DNA-Damage Response genes expression in DKO clones and cells. nDDR-high = 30, nDDR-low = 61. e, Co-variation of gene modules in DKO clones. f, Co-variation of gene modules in DKO cells.





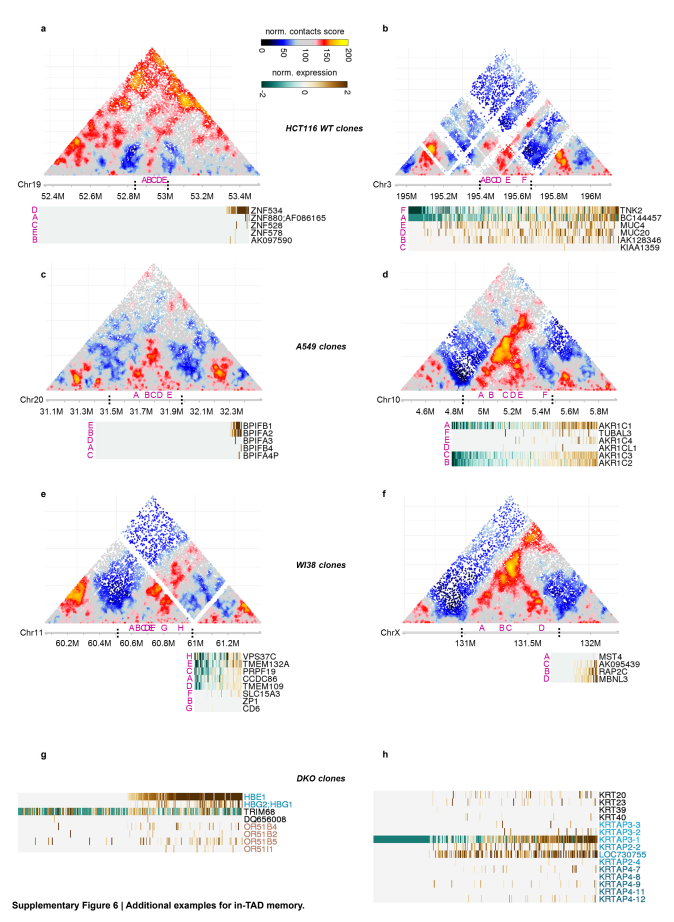
## Supplementary Figure 4 | GEMINIs expression in KO HCT116 cells.

a, Density plot of total UMIs per KO clone. The vertical black dashed line indicates median coverage per clone, and the vertical red dashed line represents the minimal coverage threshold for analysis. b, Expression of GEMINIs in 251 KO clones. GEMINIs are ordered as in Fig. 4h. Expression is normalized by the maximal expression of each GEMINI in WT clones.



Supplementary Figure 5 | PBAT-based copy number variation analysis in HCT116 cells.

a, Spatial map of relative genomic coverage in pooled HCT116 single cells and in normal colon tissues, divided into bins of 1Mbp. Showing only bins with at least X100 coverage in at least one of the pooled datasets. Relative genomic coverage is computed as log2 of the ratio between normalized coverage of the two downsampled datasets. Green dots indicate genomic bins with coverage ratio of 2 or higher. Lower panel: magnified example of chromosome 5. b, Same as a, comparing pooled genomic coverage in VIM-high clones and non VIM-high clones. Red dots indicate genomic bins with coverage ratio of 0.5 or lower. c, Aggregated relative genomic enrichment in individual clones over whole chromosomal arms. Showing only chromosomes for which at least one clone had evidence for large-scale duplication or loss, indicated by larger dots. Green and red dashed lines indicated the expected values of relative enrichment following complete chromosomal arm duplication or loss, respectively. d, Relative expression emerging from all genes in each chromosomal arm in each clone (x-axis) compared to the total arm relative genomic coverage, as plotted in c and using same y-axis.



**a-f**, Additional examples for in-TAD transcriptional memory. Following the same spatial scheme of Fig. 5f, showing expression of genes spanning selected TADs in HCT116 clones (**a-b**), A549 clones (**c-d**) and Wl38 (**e-f**). Heatmap is showing log-ratio of expression for each gene in each clone relative to the gene's median expression across all clones. Each gene position is indexed by one letter in purple. TAD borders suggested by our initial screen shown as black dashed vertical lines. **g**, Expression of beta-globin region, as defined in Fig. 5a, in HCT116 DKO clones. **h**, Expression of KRTAP region, as defined in Fig. 5f, in HCT116 DKO clones.