Epigenetic diversity of Kaposi’s sarcoma–associated herpesvirus

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ABSTRACT

Spontaneous lytic reactivation of Kaposi’s sarcoma–associated herpesvirus (KSHV) occurs at a low rate in latently infected cells in disease and culture. This suggests imperfect epigenetic maintenance of viral transcription programs, perhaps due to variability in chromatin structure at specific loci across the population of KSHV episomal genomes. To characterize this locus-specific chromatin structural diversity, we used MAPit single-molecule footprinting, which simultaneously maps endogenous CG methylation and accessibility to M.CviPI at GC sites. Diverse chromatin structures were detected at the LANA, RTA and vIL6 promoters. At each locus, chromatin ranged from fully closed to fully open across the population. This diversity has not previously been reported in a virus. Phorbol ester and RTA transgene induction were used to identify chromatin conformations associated with reactivation of lytic transcription, which only a fraction of episomes had. Moreover, certain chromatin conformations correlated with CG methylation patterns at the RTA and vIL6 promoters. This indicated that some of the diverse chromatin conformations at these loci were epigenetically distinct. Finally, by comparing chromatin structures from a cell line infected with constitutively latent virus, we identified products of lytic replication. Our findings show that epigenetic drift can restrict viral propagation by chromatin compaction at latent and lytic promoters.

INTRODUCTION

Kaposi’s sarcoma–associated herpesvirus (KSHV) infection causes sarcoma and lymphoproliferative disorders in immunocompromised individuals [reviewed in (1)]. In the latent phase, KSHV genomes reside within the nucleus as multicopy circular extrachromosomal elements or episomes. Both micrococcal nuclease cleavage patterns and immunoprecipitation of histone proteins demonstrate that episomal DNA is incorporated into nucleosomes (2–4). Lytic reactivation, thought to be triggered by multiple cellular stress pathways in vivo, is associated with changes in chromatin structure (3,4). Furthermore, histone deacetylase and DNA methyltransferase (DNMT) inhibitors can reactivate the lytic cycle in latently infected cell culture, as can phorbol esters, e.g. 12-O-tetradecanoylphorbol-13-acetate (TPA), which are protein kinase C agonists [reviewed in (5)]. Thus, episomal chromatin contributes to maintenance of the latent state and the repression of lytic transcription.

Chromatin intrinsically regulates gene expression, as many transcription factors exhibit reduced access to binding sites incorporated into nucleosomes [reviewed in (6)]. Furthermore, chromatin components often have post-synthesis chemical modifications (DNA methylation and acetylation, methylation, etc. of histones) that can activate or repress transcription locally, as well as recruit their own modifying enzymes. Chromatin modifications can thus constitute a self-perpetuating state. Such states are termed epigenetic if maintained throughout the cell cycle, as in the case of DNA methylation [reviewed in (7)]. Epigenetics underpin the differentiation of isogenic cells during development, as well as inactivation of one of two X chromosomes in female mammals. Imperfect

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The authors wish it to be known that, in their opinion, the first two authors should be regarded as joint First Authors.

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maintenance of chromatin states, i.e. epigenetic drift, may contribute to aging and cancer [reviewed in (8)].

Epigenetic drift could be important for viral biology as well. DNA methylation gradually builds over much of the KSHV genome after de novo infection (3), indicating that epigenetic changes occur. However, the phenotypic consequences of this drift are unknown. Generally, DNA methylation is linked to transcriptional silencing [reviewed in (9)], suggesting that it could inactivate the virus. On the other hand, it might protect against the host innate immune response to unmethylated DNA (10).

Recently, it has become possible to characterize heterogeneity of chromatin structure at select loci within populations of mammalian nuclei by treatment with recombinant M.CviPI DNMT [reviewed in (11–13)]. This enzyme, which we cloned from Chlorella virus, methylates cytosine in GC dinucleotides to G-m5CG (14). The methyl marks can be read by bisulfite genomic sequencing [BGS; (15)] and unequivocally distinguished from endogenous mammalian methylation at H-m5CG (by convention, H equals A, C, or T). As nucleosomes and DNA-bound factors block access of exogenous DNMT probes to DNA (16–19), the resultant GC methylation pattern is used to infer chromatin structure. The combination of high-resolution probing by M.CviPI and BGS is termed MAPit, for methylation accessibility probing for individual templates, developed by our laboratory (20). By sequencing clonally expanded individual molecules, MAPit allows detection of multiple chromatin states at selected loci within a cell population. This is in contrast to techniques such as chromatin immunoprecipitation, which report the population average [reviewed in (11)]. Although new, MAPit is rapidly being adapted to the study of diverse genomic loci (20–26). Related studies support the ability of MAPit to accurately map gaps between nucleosomes across the genome (27) and track heterogeneity of chromatin structure (28,29).

Hypothesizing that KSHV populations would exhibit epigenetic drift, we examined promoters and 5’ ends of genes of interest in latently infected cells by MAPit. The degree of diversity seen was surprising. Both fully closed and fully open conformations were present at three different KSHV promoters. Treatment with TPA and induced expression of the replication and transcription activator (RTA) were used to test which of the diverse chromatin structures found were linked to reactivation during lytic transcription. At the time point tested, only a subset of episomes appeared to have remodeled chromatin in response to lytic reactivation by both means. We also observed a subset of episomes associated with lytic replication, and identified a link between chromatin compaction, CG methylation and inhibition of reactivation. These findings suggest that latent virus is subject to progressive epigenetic inactivation.

MATERIALS AND METHODS

Cell culture

Body-cavity-based lymphoma cell line 1 (BCBL1) cells (30) were cultured in RPMI 1640 medium supplemented with 10% (v/v) fetal calf serum (FCS), 2 mM glutamine, 1 mM sodium pyruvate, 100 U/ml penicillin and 100 µg/ml streptomycin at 37°C under 5% CO2. TREx BCBL1-RTA cells for doxycycline-induced expression of RTA (31) were cultured in RPMI 1640 medium supplemented with 10% (v/v) FCS or tetracycline-free FCS, 2 mM glutamine, 1 mM sodium pyruvate, 100 U/ml penicillin and 100 µg/ml streptomycin at 37°C under 5% CO2. Telomerase-immortalized vein endothelial long-term–infected (TIVE-LTC) cells (32) were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% (v/v) FCS, 4.5 g/l glucose, 2 mM glutamine, 1 mM sodium pyruvate, 100 U/ml penicillin and 100 µg/ml streptomycin at 37°C under 5% CO2.

For TPA-induced reactivation, BCBL1 cells were grown to a density of 2–3 × 10^5 cells/ml. TPA was added to a final concentration of 20 ng/ml and cells were incubated at standard conditions (37°C, 5% CO2) for 6 h. Cells were pelleted at 247 g for 5 min and resuspended in the same volume of fresh RPMI medium and grown for another 6 h before nuclei isolation for MAPit. For reactivation by doxycycline-inducible RTA, TREx BCBL1-RTA cells were cultured for at least 2 days in tetracycline-free medium before performing the induction. Cells at a density of 4–5 × 10^5 per milliliter were then treated with 1 µg/ml doxycycline and 250 µM foscarnet for 6 h. Foscarnet was used to suppress viral lytic replication; quantification of viral load by real-time polymerase chain reaction (PCR) indicated that viral replication was largely blocked (not shown). Cells were pelleted at 247 g for 5 min and resuspended in the same volume of fresh RPMI medium with 250 µM foscarnet and grown for another 6 h before nuclei isolation for MAPit. Uninduced BCBL1 or TREx BCBL1-RTA cells (treated with 250 µM foscarnet), respectively, served as the 0 h time point.

MAPit single-molecule footprinting

MAPit was performed as described previously (12,13). Briefly, BCBL1 or TREx BCBL1-RTA cells were harvested at a density of 3–6 × 10^5 cells/ml, washed twice with ice-cold phosphate-buffered saline (PBS), counted and resuspended at 10^6 cells/ml in PBS. Four to five million cells were stored as pellet at −80°C for later RNA extraction (see below). For each MAPit assay, 10^6 cells were pelleted and resuspended in 200 µl cold Buffer A [(10 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES)-KOH, pH 7.5, 2 mM MgCl2, 10 mM KCl, 2 mM dithiothreitol (DTT), 0.2 mM phenylmethanesulfonyl fluoride (PMSF)]. Cells were incubated on ice for 10 min to release nuclei by hypotonic lysis, then vortexed 10 s at medium velocity and centrifuged for 10 s at 16,000g. To check for complete cellular lysis, an aliquot of nuclei in 0.2% (w/v) trypan blue were examined by light microscopy. Nuclei were only used for MAPit if the lysis was complete, and nuclei were evenly round and granular in appearance. Nuclear integrity was also confirmed in parallel by Hoechst staining, which dyes DNA, and
fluorescence microscopy. Nuclei were washed twice with 500 μl cell resuspension buffer [20 mM HEPES-KOH, pH 7.5, 70 mM NaCl, 0.25 mM ethylenediaminetetraacetic acid (EDTA), 0.5 mM ethylene glycol tetraacetic acid (EGTA), 0.5% (v/v) glycerol, 10 mM DTT, 0.25 mM PMSF] and resuspended in 90 μl methylation buffer (cell resuspension buffer supplemented with 160 μM S-adenosyl-L-methionine).

TIVE-LTC cells were harvested at 100% confluency, washed twice with ice cold PBS, counted and resuspended at 10^6 cells/ml in PBS. Cells were pelleted (5 × 10^6 cells for each MAPit assay), washed once in 250 μl cell resuspension buffer and lysed in 100 μl cell lysis buffer [cell resuspension buffer plus 0.25% (v/v) Nonidet P-40] for 10 min on ice. Nuclei were pelleted at 1000g for 5 min at 4°C, then washed twice with 500 μl cell resuspension buffer and resuspended in 90 μl methylation buffer.

A 30 U recombinant M.CviPI-maltose binding protein (MBP) fusion [New England Biolabs, hereafter M.CviPI; (14)] were added per 10^9 nuclei (based on cells counted after all washes completed), which were then incubated at 37°C for 15 min. Reactions were stopped by addition of an equal volume of 1% (v/v) sodium dodecyl sulfate (SDS), 100 mM NaCl, 10 mM EDTA and incubated overnight at 50°C with 100 μg/ml proteinase K. DNA was then phenol extracted (partitioned with an equal volume phenol:chloroform:isoamyl alcohol:25:24:1) and ethanol precipitated. A 0 U M.CviPI control reaction was also performed to gauge endogenous GC methylation (not shown; none was observed at any locus).

To examine comparable results between experiments, M.CviPI activity was calibrated between aliquots of enzyme using methylation of a HaeIII site in the GRP78 promoter in BCBL1 cells as a standard. MAPit experiments were also conducted at 10 rather than 70 mM NaCl, with 3-fold higher and lower effective M.CviPI concentration, and over a 3-fold range in M.CviPI incubation time (Supplementary Figures S1 and S2). A non-linear relationship was seen between M.CviPI concentration and degree of methylation. This matches a digital methylation maps for both CG and GC methylation

**Bisulfite genomic sequencing**

To deaminate, 2–4 μg DNA was denatured for 5 min at 95°C in freshly made 0.3 N NaOH, then incubated for 6 h at 50°C in a saturated metabisulfite solution as previously described (33). Bisulfite-converted DNA was desulfonated and concentrated with a commercial kit (Zymo Research catalog no. D5026). Target loci were amplified from 20–100 ng bisulfite-treated DNA with 3 U HotStar Taq DNA polymerase (Qiagen) and 250 nM primers in 10 mM tris(hydroxymethyl)aminomethane (Tris)–HCl, pH 8.6, 50 mM KCl, 3 mM MgCl2 and 200 μM of each 2′-deoxynucleoside 5′-triphosphates (dNTP). Primers sequences are given in Supplementary Table S1. Amplification was performed in 2–3 separate reactions to minimize stochastic variation in PCR amplification, then pooled for cloning. Cycling conditions were 5 min at 95°C, followed by 40 cycles of 1 min at 95°C, 1 min at 56°C–60°C, (optimized for each primer pair), 3 min at 72°C, and a final extension of 5 min at 72°C.

Amplified DNA was separated from unincorporated primers by electrophoresis in 1% (w/v) agarose buffered with Tris-acetate-EDTA, excised and purified from the gel and ligated into a cloning vector using topoisomerase (TOPO TA; Invitrogen). Calcium-competent DH10b or TOP10 cells were transformed with 10–50 ng plasmid and plated on lysogeny broth (LB) medium with 100 μg/ml ampicillin and 40 μg/ml 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside (X-gal) for blue/white screening. Colony PCR with M13 forward and reverse primers was also used to test cloning efficiency. Colonies containing the insert were suspended in 100 μl LB plus 100 μg/ml ampicillin and grown at 37°C overnight in 96-well plates without shaking. Glycerol was then added to 10% (v/v) and the plates were incubated at −80°C for at least 2 h, whereupon plasmid was subjected to TempliPhi™ rolling circle amplification and automated BigDye sequencing (Applied Biosystems) at the University of Florida Interdisciplinary Center for Biotechnology Research (http://www.biotech.ufl.edu).

**Sequence analysis**

The genomic sequences corresponding to BGS amplicons are listed in Supplementary Table S2. Sequence files in ABI format were aligned in Sequencher™ (Genecodes) and are available on request. MethylViewer [http://dna.leeds.ac.uk/methylviewer; (22)] was used to ascertain percent cytosine conversion and tabulate methylation status of each site in each sequence read (i.e. molecule). Molecules with <95% conversion of cytosines outside potential methylation sites were discarded. Molecules from the same initial DNA sample that had identical patterns of both methylation and unconverted cytosines were presumed to be duplicate clones, and all copies but one were then discarded. Molecules with either zero methylation or 100% conversion of cytosines were not screened in this way, as there were insufficient grounds to infer duplication. Duplicates made up 0–10% of molecules in each dataset.

The tables of methylation status were used to generate digital methylation maps for both CG and GC methylation
for each molecule in Excel (Microsoft). Within the LANA, RTA and vIL6 promoters, CG methylation was rare; thus, GGC sites were counted as GC sites to increase resolution. For the ampiclon overlapping the LANA open reading frame (ORF), GGC sites were excluded from the analysis. Digital methylation maps were clustered hierarchically using Cluster [http://rana.lbl.gov/EisenSoftware.htm; (34)], and printed as color maps with MapleTree (http://mapletree.sourceforge.net). Dendrograms were split by eye into clusters of interest. Molecules in each cluster originating from each experiment were counted. Expectation values for the observed distributions were computed as two-tailed Fisher’s exact tests. To generate the accessibility profile (Figure 2D), the fraction of sequence reads bearing accessible spans of 1–100 or 101–400 bp was computed at each base pair position. The difference in accessibility after TPA treatment was then subtracted from that before treatment. Likewise, to generate occupancy profiles (Figures 3D, 5D and 8D), the fraction of sequence reads bearing footprints of 4–39, 40–100 or 101–200 bp was computed at each base pair position. Footprints extending to the ends of the amplicons were not counted. To get the difference in occupancy, the fraction for one experimental condition was then subtracted from the fraction for another.

**RNA preparation for real-time reverse transcription PCR (RT-PCR)**

BCBL1 and TREx BCBL1-RTA cells were treated as described above in two independent experiments (including different stocks of cells). From 4 to 5 × 10⁶ induced and uninduced cells, RNA was extracted using RNA Bee (TEL-TEST, Inc.) according to the manufacturer’s instructions. RNA was DNase I treated [10 μg RNA, 1 μl RNasin® (Promega), 1 × DNase I buffer, 2 μl (4 U) RNase-free DNase I (New England Biolabs) in 50 μl total volume] for 25 min at 37°C and recovered by ethanol precipitation after ~16 h at −20°C. An aliquot of DNase I-digested RNA was tested for RNA integrity by agarose gel electrophoresis and the rest stored at −80°C. RNA (1–3 μg) was reverse transcribed using oligo dT primer and Invitrogen SuperScript III reverse transcriptase (RT) in a 20 μl reaction mixture as indicated by the manufacturer (20 μl reaction mixture per 1 μg RNA; for more RNA, the assay was scaled up accordingly). A reverse-transcription reaction without RT enzyme served as control (no RT).

**Quantitative real-time PCR**

According to the MIQE guidelines for quantitative real-time PCR [qPCR; (35)], we refer to reference genes instead of housekeeping genes and Cq (quantification cycle) instead of Ct (threshold cycle). All primers (Supplementary Table S1) were purchased from Integrated DNA Technologies, Inc. and tested for their efficiency (between 90 and 110%) and specificity (melting curve analysis).

Per 12 μl qPCR reaction, 0.5 μl complementary DNA was used. Primers were used at a final concentration of 200 nM. Two-step qPCR was performed with the Fast SYBR Green Master Mix and a StepOnePlus real-time PCR instrument (both Applied Biosystems). Cycle conditions were initially 95°C for 10 min, 40 cycles of 95°C for 15 s, 60°C for 1 min, followed by melting curve analysis.

For quantification of M.CviPI-mediated protection from R.HaeIII digest, 1 μg undeaminated DNA was digested with 10 U R.HaeIII, as per the enzyme manufacturer’s directions, or mock-treated with an equal volume of 50% (v/v) glycerol. For each sample, three wells with 10 ng DNA apiece were quantified by real-time PCR, as above.

**RESULTS**

**Multiple chromatin conformations in the KSHV latency-associated region**

LANA, vFLIP, vCyclin, Kaposin and 12 pre-microRNAs are highly expressed off polycistronic transcripts arising from a single KSHV latency-associated region (KLAR) in tumor cells. During latency, host RNA polymerase II initiates transcription of KLAR from the upstream latent transcription start site (TSS; Figure 1B). Additionally, a second downstream TSS is used during lytic reactivation (36). Three CCCTC-binding factor (CTCF) binding sites are located between these two TSSs and the LANA open reading frame (ORF73); their occupancy by the chromatin insulator CTCF (37) contributes to LANA expression, mediates chromosomal looping and plays a role in episonal maintenance (38,39). All KSHV-infected cells express LANA, which is required for replication and maintenance of viral episomes in latently infected cells [reviewed in (40)]. Yet, not all of the ~80 episodes present in latently infected lymphoma cells need express LANA. We hypothesized that a subset of episodes might provide LANA function for the rest, which could be epigenetically inactivated.

As a single-molecule technique, MAPit provided a way to test this hypothesis directly. A key advantage of MAPit, in contrast to population average readouts, is the ability to discover distinct subpopulations without enrichment. Nuclei from the KSHV-infected pleural effusion lymphoma (PEL) cell line BCBL1 were incubated with 0 or 30 U M.CviPI, following the protocol outlined in Figure 1A. We cloned and sequenced 10 bisulfite-converted molecules encompassing the LANA promoter from each sample. The patterns of CG and GC methylation were drawn by MethylViewer (22), which represents methylated cytosines as ball-and-stick symbols (Figure 1B). Efficiency of conversion was estimated as percent HCH (H is any base but G) conversion in each sequence read (i.e. cytosine outside potential methylation sites; blue vertical ticks mark unconverted HCH). As expected, without M.CviPI addition, methylation of GCH sites was not above the background given by incomplete conversion. The M.CviPI-probed sample exhibited prominent accessibility above background between and adjacent to three known CTCF sites. Some molecules were also accessible near the latent TSS.

Although MethylViewer is excellent for comparing different methylation types on single molecules, such detailed
Figure 1. MAPit analysis of chromatin structural diversity. (A) MAPit workflow diagram, adapted from Darst et al. (13). Chromatin (e.g. nucleosomes) modulates accessibility of DNA sites to soluble proteins, such as (i) transcription factors (represented by shaded rectangle) in live cells, or (ii) exogenous M.CviPI DNMT in nuclei. Bisulfite treatment (iii) records M.CviPI accessibility pattern in the DNA sequence by conversion of unmethylated cytosine to uracil (cyan). Subsequent amplification and cloning (iv) allows recovery of this information from individual molecules. Note that endogenous and probe methylation sites are distinguished by sequence specificity of the respective DNMTs (respectively, cyan and magenta). The DNA sequences obtained are used to generate maps (v) of DNA methylation and accessibility. (B) MAPit at the KSHV gene LANA detected endogenous methylation and sporadic accessibility of the latent promoter. MAPit was performed in nuclei of BCBL1 cells. Top: map of genetic elements within the MAPit amplicon: LANA ORF (ORF73), three CTCF sites (ovals), lytic and latent TSSs (bent arrows) and TATA boxes (triangles). Middle: MethylViewer (22) display of MAPit results from control nuclei probed as indicated with 0 U or 30 U M.CviPI. Blue
graphs are unwieldy for large-scale analysis. We therefore devised a three-color map scheme (Figure 1C), used throughout the rest of this work. Patches of accessibility to DNMT (M.CviPI and endogenous DNMTs) received bright color (yellow and red, respectively), and patches of protection were colored black. Gray color denoted borders between methylated and unmethylated patches. We used unsupervised hierarchical clustering to order molecules, facilitating pattern recognition. Color maps for both GC and CG methylation status were clustered together. The 16 GCG sites were disregarded in this analysis, leaving 52 sites for GC and 36 for CG.

The data showed ‘banded’ CG methylation across almost all molecules in the LANA ORF. By contrast, the 10 molecules varied in degree and pattern of M.CviPI accessibility. This could reflect variation in chromatin structure between episomes. An alternate explanation would be a poor nuclear preparation, such that the M.CviPI probe did not enter all nuclei. However, this seems unlikely as nuclei appeared intact and homogenous by vital (trypan blue) and Hoechst staining (not shown).

Moreover, all 10 clones of the LANA amplicon exhibited GC methylation above background unconverted HCH (Figure 1C, compare lower panels in 0 versus 30 U M.CviPI). In addition, we performed BGS at select host loci from the same sample. Two representative loci are shown (Figure 1D). At the more open locus, the promoter of GRP78, all 17 molecules sequenced had GC methylation. This locus was previously found to be highly accessible to a DNMT probe in a colorectal line (28,29).

Because MAPit requires amplification and cloning of bisulfite-converted DNA, it can be affected by sequence-specific biases in either step. We therefore measured M.CviPI methylation by R.HaeIII digestion of deaminated DNA from the same samples used for MAPit (Figure 1E and Supplementary Figure S2). R.HaeIII does not digest its site GGCC when the central cytosine is methylated, the concordance between R.HaeIII protection and BGS showed the latter to be unbiased. In turn, this meant that M.CviPI efficiently permeated nuclei and accessed the GRP78 promoter. We also probed nuclear DNA directly with R.HaeIII, without prior M.CviPI treatment (Supplementary Figure S3). The restriction enzyme, which binds DNA cooperatively as a dimer and thus is similar in size to the monomeric M.CviPI fusion protein, respectively, 74.2 and 83.7 kDa, was able to access and cut its sites on most molecules. Thus, variation at LANA was not due to uneven probing of nuclei.

As indicated by the second locus shown, the GAPDH promoter, heterogeneity of promoter chromatin structure was not unique to LANA. Variation in chromatin state between copies of the same gene can arise from several causes. Epigenetic variation occurs in X inactivation, imprinting and mononucleic expression. Dynamic variation occurs in periodic and sporadic cycles of chromatin remodeling and transcription [for instance, (41)]. However, none of these phenomena have been described for a viral gene such as LANA. In any case, the implication would be that the promoters in the closed conformation do not support transcription. Thus, as hypothesized, not all episomes continuously express LANA. Because all infected cells express LANA (32), this result implies heterogeneity of episomes within individual nuclei.

MAPit survey of LANA promoter architecture

In addition to the latent TSS, the LANA promoter uses a lytic TSS on reactivation (36). Furthermore, transcription starting at the latent TSS overlaps the lytic K14 promoter on the opposite strand. We asked whether MAPit could visualize changes in chromatin structure within this complex regulatory region during lytic reactivation induced by phorbol ester treatment. To test this, we cloned and sequenced 102 amplified deaminated LANA promoter molecules from M.CviPI-treated BCBL1 nuclei (0 h or uninduced), and 110 molecules from M.CviPI-treated nuclei harvested after 12 h of TPA treatment of cells (12 h or induced). Induced transcription of two early lytic genes, RTA (ORF50) and vIL6 (ORF K2), was demonstrated by real-time RT-PCR (Supplementary Figure S4). An early time point was chosen to obtain incomplete lytic reactivation. Otherwise, histone-depleted products of viral DNA replication would out-compete cloning of BGS products derived from chromatinized viral episomes (4,42). Hence, these induction conditions did not have an obvious effect on M.CviPI accessibility averaged across the population (Figure 2A and B).

We therefore asked whether differences might be detected in a single-molecule analysis. To determine which methylation patterns were enriched in one dataset relative to the other, we pooled all reads from the 0- and 12-h TPA treatment time points and clustered them hierarchically as GC methylation plots (Figure 2C). We

Figure 1. Continued

vertical ticks indicate positions of unconverted cytosines, excluding GC and CG sites (i.e. HCH); percent conversion of all HCH cytosines for each sequence clone is given on the right. Bottom (key): Circles and triangles, respectively, indicate CG and GC sites. Methylated sites are shaded. Methylated GCG sites are shaded gray to indicate their ambiguity; however, comparison of sequences from nuclei with and without M.CviPI treatment (middle, lower and upper panels) indicated that GCG sites within the LANA ORF were endogenously methylated. (C) Condensed view of the data in B, to scale with locus map at top. Each row of pixels represents one cloned sequence, i.e. molecule. Each plot tracks only GC or CG methylation; GCG sites are ignored. As indicated by key at bottom, spans of color mark ≥2 contiguous methylated sites; black marks ≥2 contiguous unmethylated sites; gray marks spans between methylated and unmethylated sites; white indicates missing or unaligned sequence. Positions of sites are indicated by circles and triangles (HCG and GCH, respectively). (D) Host promoters varied in degree of accessibility to M.CviPI. Maps of GC methylation at two representative promoters are shown, to scale with LANA amplicon. (E) MAPit at the three loci shown was semi-quantitative. At the HaeIII sites indicated in red at GAPDH, GRP78 and LANA, 10 of 20, 15 of 17 and 1 of 10 molecules had a methylated GC site, respectively (blue bars). R.HaeIII digestion followed by quantitative real-time PCR was used to measure bulk methylation of the same three sites in genomic DNA purified from M.CviPI-treated nuclei (red bars). Under these conditions, ~95% of DNA from untreated nuclei (i.e. unmethylated) was digested (not shown). The concordance between BGS and R.HaeIII digestion indicated that representation of methylated and unmethylated molecules in MAPit was unbiased.
plotted accessibility to M.CviPI in each molecule as blue (0 h) or yellow (12 h) to mark its source and split the dendrogram into clusters of interest. To better visualize differences in chromatin structure between clusters, each was rescaled vertically. The fraction of molecules present in each cluster from each time point is given on the right. The KLAR amplicon in this experiment was positioned farther upstream from the LANA gene than that analyzed in Figure 1, allowing better characterization of accessibility at the LANA latent TSS. Strikingly, the three CTCF sites were seen to be occupied by footprints in almost all molecules. Even in cluster i, which had reduced M.CviPI methylation overall, the three CTCF sites were dimly outlined. Importantly, this demonstrates that M.CviPI permeated these nuclei, but was preferentially excluded from the latent TSS (especially compared with clusters iii and iv).

For each cluster, we found the probability P of having so skewed a distribution of molecules from the two samples by chance. Although clustering revealed several distinct chromatin conformations, TPA treatment increased representation of molecules only in cluster iii (by 13%) with borderline significance (P = 0.05). To further examine TPA-induced changes in LANA promoter chromatin structure, we calculated accessibility of lengths 1–100 and 101–400 bp occurring at each position, and subtracted accessibility before TPA treatment (0 h) from that after (12 h) (Figure 2D). TPA treatment decreased accessibility adjacent to the CTCF sites (1–100 bp, red curve) and just downstream of the LANA latent TSS (101–400 bp, blue curve) in ∼10–15% of the molecules, corresponding well with the number of molecules TPA increased in cluster iii. We conclude that TPA increased protection at the K14 TATA element, mainly in cluster iii.

Clustering also highlighted the large degree of diversity of chromatin structure at this locus. Both fully protected and almost fully accessible molecules were detected. Cluster ix was of particular interest as it appeared to be nearly nucleosome free. Complete chromatin unfolding and nucleosome displacement across a region of this size has not been observed at any human chromosomal locus studied by MAPit. We hypothesize that molecules of cluster ix originated from spontaneous lytic reactivation of KSHV, which occurs within 1–3% of PEL cells (30,43,44). As is the case for other herpesviruses, the KSHV genome is depleted for histones during lytic replication before being packaged into virions (4,42). These results indicated that MAPit could discover chromatin conformations comprising <∼5% of a complex population.

A subset of episomes undergoes changes in chromatin structure during reactivation

The master regulator of lytic transcription is the replication and transcription activator RTA (45). RTA directly activates several downstream viral genes, such as vIL6, as well as transcription of its own gene (46). We hypothesized that a subset of latent episomes would be poised to transcribe RTA (ORF50) and that these would have a distinct chromatin structure visible by MAPit. Induced reactivation would be predicted to shift these molecules to the actively transcribing state, which might involve nucleosome repositioning. We therefore cloned and sequenced 114 and 112 molecules from M.CviPI-treated nuclei harvested after 0- or 12-h TPA treatment, respectively (Figure 3).

Accessibility to CG and GC DNMTs was computed at each position. Within ORF48 (5' of RTA), a peak of CG methylation was observed upstream (313–284 bp upstream of the RTA TSS) of a positioned nucleosome, hereafter N−1 (Figure 3A and B). The same position had been previously shown to be endogenously methylated in BCBL1, although in a larger proportion of the population (53). A second notable feature of this plot was a broad peak in GC accessibility centered ∼100-bp upstream of the RTA TSS. An open chromatin structure was not expected to be so common in the latent state. Moreover, there was no obvious difference between GC accessibility profiles before and after TPA treatment (Figure 3A). Thus, the bulk of the population was unchanged. As expected, the GAPDH (Supplementary Figure S2C) and GRP78 control loci also showed no change in chromatin architecture in response to TPA (compare Supplementary Figure S5 with Figure 1D, upper).

As before, molecules from the two time points were clustered together to identify chromatin conformations enriched in either (Figure 3C). Cluster i, comprising ∼50% of all molecules, was, as at KLAR, largely (but not completely) inaccessible to M.CviPI and presumably transcriptionally inactive. Clusters ii–vi all had an N−1 footprint. Clusters ii and iii appeared similar to the eye: both featured three evenly spaced positioned nucleosome-sized footprints. However, cluster iii was more closely related to cluster v than ii in the original dendrogram (Supplementary Figure S6). The central N+1 nucleosome in cluster ii covered several previously identified transcription factor binding sites upstream of the RTA TSS. The N+1 and N+2 nucleosomes seemed to shift slightly downstream in cluster iii versus ii. Cluster iv had a variably positioned N−1 footprint, followed downstream by a large open gap, then 2–3 variably positioned nucleosome-sized footprints. Cluster v molecules shared a large M.CviPI-accessible linker upstream of N−1, with chromatin that appeared to be denser downstream over the RTA transcribed region. Cluster vi lacked N−1. Lastly, cluster vii was, just as cluster v found at KLAR, almost entirely accessible and again comprised ∼3% of all molecules.

Clusters ii, v and vi had P values below the usual threshold for significance (<0.05). Because clusters ii and vi decreased in response to TPA, they may represent transcriptionally poised states, primed to respond to activating stimuli. Cluster iv would then correspond to a transcriptionally active state or a thermodynamically stable stage in the cycle of RNA polymerase recruitment and release. Although it had too few molecules for statistical significance, cluster iii was also enriched for molecules from TPA-treated cells and might also represent a stage in the transcription cycle. Comparing the increase in cluster v to the decrease in clusters ii and vi, we conclude that at least...
Figure 2. TPA treatment altered KLR chromatin structure in ~10% of BCBL1 episomes. Note that this KLR amplicon, although overlapping the amplicon of Figure 1, was centered upstream to better observe accessible DNA at the latent TSS. (A) Profile of endogenous CG (0-h TPA treatment, red; 12-h TPA treatment, green) and exogenous GC methylation (0-h TPA treatment, black; 12-h TPA treatment, blue) across the locus. GCG sites were not counted. Note locus of highest CG methylation coincided with a peak of GC methylation; elsewhere, CG methylation was minimal. (B) Schematic of the analyzed amplicon. The LANA ORF begins left of the amplicon. The LANA latent and lytic TSSs are indicated. TATA boxes are indicated by triangles (upside down, latent and lytic TSSs are indicated. TATA (T). A second RTA TSS was recently reported roughly 500-bp upstream of this amplicon (52). Below, inferred nucleosome positions are indicated as two overlapping gray ovals, each 150-bp long. (C) Clustered GC methylation maps from 0-h (blue) and 12-h (yellow) treatment with TPA. Methylation of GC sites (including GCG, as m^5CG was rare) was plotted as in Figure 1, to better observe accessible DNA at the latent TSS. (D) Difference in occupancy from 0 to 12 h of TPA treatment. Footprints in two size ranges, 4–39 and 121–200 bp, were tallied at every position in each dataset, then the 0-h profile was subtracted from the 12-h profile.
10% of episomes remodeled RTA chromatin in response to TPA. By the same token, a large fraction of molecules appeared not to have responded to TPA at this time point, including those of cluster i, which were highly compacted.

Several small footprints covering 1–2 GC sites occurred at 93, 86 and 74 bp upstream of the TSS (Supplementary Figure S6), which might represent some of the transcription factors reported to bind this area. To determine which footprints were more abundant in each dataset, we counted footprints of lengths 4–39 and 121–200 bp occurring at each position and subtracted footprint counts before TPA treatment from those after (Figure 3D). The result is difference in occupancy of factors within each footprint size range at each base pair position. This analysis revealed greater definition of the N/C0 footprint (broad peak in black spanning from 300 to 100 bp relative to the RTA TSS), repositioning of nucleosomes inside the transcribed region of RTA (sharp peaks on right), and changes in some sub-nucleosomal footprints flanking these areas (in red), but not changes in the putative transcription factor footprints. Nucleosome repositioning in the gene was consistent with the hypothesized shift from cluster ii to iii.

Together, these results suggested that only a subset of KSHV episomes had responded to TPA by reorganizing RTA promoter chromatin by 12 h of TPA induction. From this, one would predict a similar proportion of episomes to be expressing a downstream target of RTA, an immediate-early gene such as vIL6. We cloned 96 and 71 molecules amplified from bisulfite-converted DNA isolated from TPA-treated and untreated cells, respectively. Unlike at RTA, at the vIL6 promoter (within ORF2), there was a wide peak of accessibility to M.CviPI (Figure 4A and B). As with RTA, no change in bulk accessibility was seen on TPA treatment from the methylation plots alone. However, clustering of molecules once again showed otherwise. As at KLAR and RTA, there was both an abundant, highly inaccessible, likely silenced cluster (Figure 4C, cluster i) and a rare, highly accessible cluster (v), suggesting that the underlying chromatin behavior was episome wide. Again, neither of these two clusters changed significantly on TPA treatment. However, cluster iv, lacking two or more nucleosomes within the vIL6 promoter, did increase significantly from 8 to 24% of molecules after induction ($P = 0.01$). This suggests chromatin remodeling in response to activation by RTA. Moreover, these numbers were consistent with the estimate that ~10% of episomes were transcribing RTA at this time point. The observation that 8% of molecules showed the same vIL6 chromatin accessibility during latency is in congruence with a recent report of low levels of vIL6 expression during latency (54). As at RTA, results were consistent with a model of heterogeneous reactivation of the lytic pathway, in which an
open chromatin structure such as in Figure 3C, cluster vi is prerequisite. We conclude that chromatin structure reflects ability of episomes to respond to inducers of lytic transcription.

As a protein kinase C agonist, TPA induces weak signaling that may resemble a pathway for spontaneous KSHV reactivation in vivo. To further examine the extent to which the fraction of KSHV episomes with relatively inaccessible chromatin (i.e. in cluster i) was resistant to lytic reactivation, we performed an experiment with TREx BCBL1-RTA cells. These cells, containing an integrated doxycycline-inducible RTA transgene, overexpress RTA after 12 h of treatment with doxycycline and hence undergo efficient reactivation (31,55). We collected cells after 12 h of induction and performed MAPit, using primers specific to the wild-type KSHV RTA promoter. We cloned 51 and 45 molecules amplified from bisulfite-converted DNA isolated from doxycycline-treated and untreated cells, respectively (Figure 5).

Several differences in RTA promoter chromatin architecture were also observed between the cell lines. In the TREx BCBL1-RTA cell line, the upstream peak of CG methylation was absent (compare Figures 5A and 3B, red and green curves), and a footprint spanned the two GC sites within the Sp1 binding site in 10–20% of molecules [Figure 5B and C, under triangle labeled ‘S’; (2)]. Although this footprint was slightly more common on RTA induction, the increase did not pass the threshold for significance ($P = 0.4$).

As before, we noted a subset of episomes with highly accessible chromatin, cluster v. These were more abundant than we had observed in the parent cell line BCBL1, possibly due to higher basal expression of RTA in the untreated TREx BCBL1-RTA cells. Consistent with this, 5.6-fold more RTA transcript was observed in uninduced (tetracycline-free FCS) TREx BCBL1-RTA versus parental BCBL1 cells treated with TPA (Supplementary Figure S4).

As in the BCBL1 cells assayed in Figure 3, a large fraction of cloned sequences showed promoters with compact chromatin, inaccessible to M.CviPI (Figure 5C, cluster i). The fraction of KSHV episomes with closed chromatin did not significantly decrease in response to RTA induction in trans despite 30-fold increased RTA expression as compared with TPA induction. Thus, MAPit indicated heterogeneous reactivation of RTA in highly induced TREx BCBL1-RTA cells, just as in TPA-treated BCBL1 cells.

![figure](image)

**Figure 5.** Expression of a doxycycline-inducible RTA transgene altered chromatin structure at the endogenous RTA promoter of 30–40% of episomes. (A) Profile of endogenous CG (uninduced, red; 12 h of doxycycline, green) and exogenous GC methylation (uninduced, black; 12 h of doxycycline, blue). GC sites were not counted. (B) Schematic of the RTA locus, as in Figure 3B. (C) GC methylation plots for 0- and 12-h RTA induction (blue and yellow, respectively). GC sites are indicated above; plot is to scale with other panels. Dendrogram was split into five fractions of interest. On right, proportion of sequence reads from each data set present in each cluster, and expectation value $P$ calculated, as in Figure 2. (D) Difference in footprint occupancy from 0 to 12 h of TPA treatment. Footprints in two size ranges, 4–39 and 121–200 bp, were counted at every position in each dataset, then the 0-h profile was subtracted from the 12-h profile.
Epigenetic diversity in latent KSHV

Our results suggested that the promoters of \( \text{LANA}, \ RTA \) and \( v\i\!l6 \) existed in at least two states across the population of episomes. The first had compact chromatin and was non-responsive for lytic reactivation, and the second exhibited various open conformations. However, it was not yet clear whether these states were epigenetic or reflected periodic or sporadic cycles of chromatin remodeling and transcription. Heritability of chromatin conformations identified by MAPit can be inferred by correlations between GC methylation and CG methylation, which is epigenetic. Indeed, a patch of endogenous CG methylation occurred upstream of the \( RTA \) promoter in 25% of BCBL1 molecules sequenced (Figures 3A and 6A and B). Note that clusters shown in Figure 6B were based solely on HCG methylation, i.e. excluding all GCG sites. Thus, they do not correspond to the GCH methylation clusters in Figure 3C. To test whether the latter were epigenetically distinct, we counted molecules with upstream m\(^5\)CG patches in each M.CviPI methylation-defined cluster (Figure 6C). This feature was defined as methylation of 2–3 of the CG sites 284–313 bp upstream of the \( RTA \) TSS.

From both TPA-treated and untreated cells, 30% of cluster \( i \) molecules in Figure 3C had upstream m\(^5\)CG patches. However, only 15% of molecules from clusters \( ii–vi \) and 0% from cluster \( vii \) had upstream m\(^5\)CG patches. Clearly, lytic induction did not affect abundance of upstream m\(^5\)CG patches in either group of molecules. Pooling the two experimental conditions, we computed \( P < 0.01 \) that CG methylation would differ so much by chance between clusters \( i \) and \( ii–vi \). Hence, the presence of m\(^5\)CG patches upstream of \( RTA \) was anti-correlated with openness of nearby chromatin to GC methylation. Furthermore, epigenetic ‘fingerprinting’ showed that changes in cluster abundance on TPA treatment were primarily due to flux in molecules lacking the upstream CG methylation (Supplementary Table S3). This indicated that upstream-methylated \( RTA \) promoters were not significantly remodeled in response to TPA. Thus, CG methylation fingerprinting indicates that different MAPit patterns arose from epigenetically distinct subpopulations at \( RTA \). These data indicate an epigenetic basis for the chromatin structural diversity observed at \( RTA \), implying long-term stability.

Altered chromatin dynamics on constitutively latent episomes

Unlike BCBL1, TIVE-LTC do not reactivate either spontaneously or in response to TPA (32). We therefore decided to study chromatin conformation in TIVE-LTC as a model for purely latent episomes, which are transcriptionally silenced except for the \( \text{LANA} \) region. For comparison with 94 TPA-untreated \( \text{LANA} \) reads from latent BCBL1, we cloned and sequenced 107 from TIVE-LTC cells. To determine the extent to which TIVE-LTC and BCBL1 episomes had chromatin structures in common, we pooled all GC methylation plots from the two datasets and clustered them hierarchically (Figure 7).

We divided the dendrogram into clusters of interest as before and calculated \( P \) values for under- or overrepresentation of molecules in each cluster from each cell type. There was a significant increase in locus-wide protection (46–78%) of \( \text{LANA} \) in TIVE-LTC compared with BCBL1 cells (cluster \( i, P = 10^{-8} \)). This was not caused by reduced nuclear permeation by M.CviPI, as the control locus \( GAPDH \) was, if anything, more open in TIVE-LTC cells (Supplementary Figure S2C). TIVE-LTC cells also produced fewer molecules in which nucleosomes were shifted upstream to cover the latent TSS (Figure 7B, cluster \( ii \)). In addition, there was a significant reduction from 37 to 16% in the number of molecules with accessible chromatin over the latent TSS in BCBL1 cells (cluster \( iii, P = 7 \times 10^{-6} \)). However, CTCF occupancy was unaltered between the cell lines, as observed from molecules exhibiting accessible DNA around the CTCF sites (clusters \( ii \) and \( iii \)). Finally, we observed no highly accessible molecules in the TIVE-LTC cells (cluster \( iv \)). Combining data from both the \( \text{LANA} \) and \( RTA \) loci (below) yielded \( P = 0.01 \) for the presence of highly open...
presumably lytic molecules in BCBL1, but not in TIVE-LTC, cells.

We also compared the 114 TPA-untreated RTA reads from BCBL1 to 105 from M.CviPI-treated TIVE-LTC nuclei, and plotted the percent methylation at each site in the amplicon (Figure 8A). Surprisingly, KSHV episomes from TIVE-LTC cells, which except for the LANA region are transcriptionally silenced (32), had less overall endogenous CG methylation than did those from BCBL1 cells at the RTA locus, and lacked the upstream peak (Figure 8A, –313 to –284). The GC accessibility profile was different as well. TIVE-LTC episomes had a different nucleosome organization with less accessible chromatin at the RTA promoter (Figure 8A, –200 to +100 relative to the RTA TSS), but slightly more observed downstream (Figure 8A, between +100 and +200 bp from the RTA TSS). Subtracting BCBL1 footprints from TIVE-LTC footprints (Figure 8D), we found differences in frequency at which nucleosomes were footprinted in the RTA promoter (broad black peaks between –250 and +100 bp). Increased positioning of the N + 1 nucleosome centered upstream of the RTA TSS in TIVE-LTC cells corresponded with an increase in an accessible linker downstream, which was occupied by sub-nucleosomal footprints of variable size (red peak centered at +109, cf. cluster iii in panel C). Thus, the bulk accessibility data suggested that the RTA locus was more compact in TIVE-LTC than in BCBL1 cells. This compaction of the RTA promoter may account for the inability of TIVE-LTC cells to spontaneously reactivate.

**Figure 7.** Chromatin structure at KLMR in TIVE-LTC compared with BCBL1 cells. (A) Schematic of the locus. The LANA ORF begins left of the ampiclon. The latent and lytic TSSs are indicated. TATA boxes are indicated by triangles with orientations as in Figure 1A. (B) All GC methylation plots for BCBL1 and TIVE-LTC (yellow and blue, respectively, including GCG) were organized by unsupervised hierarchical clustering. GC sites are indicated above; plot is to scale with the other panel. Dendrogram was split into four fractions of interest. On right, proportion of sequence reads from each data set present in each cluster, and estimate of expectation value P calculated as in previous figures. Column for TIVE-LTC cells does not sum to 100% owing to rounding.

**Figure 8.** Silent chromatin at RTA promoter in TIVE-LTC cells. TPA-untreated BCBL1 MAPit reads from Figure 3 were compared with MAPit reads from TIVE-LTC cells. (A) Profile across RTA promoter of average endogenous CG (green) and exogenous GC (blue) methylation in TIVE-LTC cells, as well as CG (red) and exogenous GC (black) methylation in BCBL1 cells from Figure 2A for comparison. Note absence of “linker” H.m.CG peak at –300 bp in TIVE-LTC cells, despite the presence of an overlapping G.m.CH peak, on left side of graph. (B) Diagram of RTA locus amplicon and inferred nucleosome positions, as in Figure 2, to scale with other panels. (C) Clustered GC (including GCG) methylation maps from BCBL1 (yellow) and TIVE-LTC (blue) presented as in Figure 2. Five clusters of interest are presented. On right, proportion of sequence reads from each data set present in each cluster, and estimate of expectation value P calculated as in Figure 2. (D) Difference in frequency of footprints of two size ranges from 0 to 12 h of TPA calculated as in Figure 2.

**DISCUSSION**

We applied single-molecule MAPit footprinting to interrogate the chromatin structures of viral episomes in KSHV-infected cells of lymphoid and endothelial origin, which contain ~80 and 5 copies per cell, respectively. Three loci that differed with respect to expression were analyzed: the highly transcriptionally active LANA promoter, the inactive but poised RTA promoter, and the moderately active but highly RTA-responsive vIL6 promoter (58). At all three loci, MAPit revealed that KSHV episomes within latently infected PEL cells exist in a complex mix of chromatin conformations. Our data
suggest that, as evidenced by chromatin architecture, only a fraction of episomes participate in latent gene expression and in reactivation at any given time. Links between nucleosome positioning and CG methylation provided novel evidence that epigenetic drift restricts ability of virus to reactivate lytic transcription. We discuss how this drift may affect important steps within the γ-herpesvirus life cycle, such as long-term maintenance of latency and the rate and ability to reactivate lytic replication.

**Heterogeneity in chromatin accessibility**

MAPit was originally developed as a probe of heterogeneity in chromatin remodeling (28). We have now detected diverse chromatin states at two human and three viral loci by MAPit. These results were reproducible, verified by restriction endonuclease digestion and consistent with previous single-molecule studies of promoter function (12,28,29).

Heterogeneity in chromatin architecture can occur at even constitutively active promoters, as transcription factors dissociate, bind, recruit remodelers and so forth. However, molecules without accessibility to M.CviPI are presumably inaccessible to other soluble factors as well. Thus, it is difficult to envision how the transcriptional machinery could be recruited to the M.CviPI-inaccessible copies of the promoters GAPDH, LANA and so forth. On the other hand, the half-life of this compacted state in live cells is unknown. It may or may not in all cases be due to long-term silencing. However, as discussed below, at the RTA promoter, at least some episomes were in an epigenetically distinct state with reduced local accessibility to M.CviPI.

Broad peaks of accessibility to M.CviPI, i.e. nucleosome-free regions typically associated with transcriptionally active or poised promoters, were identified near each of the TSSs. In latently infected cells, the NFR at the LANA promoter co-localizes with active histone modifications (H3K9/K14ac and H3K4me3), but lacks detectable repressive marks (H3K27me3 and H3K9me3) (3,4), (R.R., unpublished data). In the absence of stable repressive marks, it is surprising that the NFR was essentially absent from 40 to 50% of episomes in BCBL1 (Figure 2C) and a larger fraction (~80%) in TIVE-LTC cells (Figure 7B). However, at present, it is not clear if the inaccessible fraction either lacked positive marks or, alternatively, possessed positive marks and lacked an NFR needed to support assembly of the preinitiation complex.

**Mechanism of lytic reactivation**

Spontaneous viral reactivation occurs as a result of leaky RTA expression. We analyzed chromatin structures during lytic reactivation by both TPA and induction of a conditional RTA transgene. Both latent and lytic episomes were observed. The latter, comprising ~3% (16% in TREx BCBL1-RTA cells with higher basal RTA expression) of molecules sequenced, were distinguished by their high chromatin accessibility and were absent in TIVE-LTC, in which KSHV cannot reactivate. We hypothesize that these highly GC-methylated reads derive from products of lytic replication and therefore largely lack nucleosomes. Although certain related viruses have been reported to incorporate nucleosomes during lytic replication (59,60), our results suggest that lytically replicated KSHV DNA is largely protein free.

Some of the other clusters might correspond to chromatin conformations present earlier in the lytic cycle. Indeed, MAPit revealed open chromatin at the RTA promoter in latently infected BCBL1 cells. Nucleosomes N−1 and N+1 were both occasionally displaced (Figure 3C, clusters iv and vi; Figure 5, clusters iii and iv). To link any of these observed conformations with transcription, RTA was induced either by TPA or by RTA overexression and resulting changes in chromatin states were determined. Under the two conditions, different classes of intermediate-open promoter molecules were observed to change in abundance at RTA. Those that decreased might have been poised for reactivation, whereas those that increased may represent stages in cycles of active transcription.

In both induction experiments, a pool of molecules with a partly accessible RTA promoter was depleted in response to lytic reactivation (cluster ii in Figures 3 and 5). Promoters with this chromatin architecture were therefore probably primed or poised to respond to activators of RTA. Such molecules comprised 10–20% of the total population. As spontaneous reactivation is rare, it follows that many of the intermediates we observed required some additional event to progress to reactivation. Interestingly, the promoters for RTA and vIL6, both targets of RTA, have been shown to have trimethylation of both H3K4 and H3K27 (3,4) (R.R., unpublished data), the bivalent mark associated with developmentally poised genes (61). Trimethylation of H3K27, which is associated with transcriptional repression, is lost quickly on RTA (and vIL6) promoter activation (3,4). Loss of this mark may allow episomes with an open RTA promoter to progress to lytic reactivation. Alternatively, the mark may be ancillary. In either case, our results show that reactivation as monitored by increased chromatin accessibility is limited to a subset of episomes, even when induced by overexpression of RTA. Increased accessibility in a limited subset of KSHV copies is consistent with the finding that histone H3 binding was not depleted at the RTA and vIL6 promoters under identical conditions of reactivation (4). An uneven distribution of episomes potent for reactivation between cells might contribute to the low rate of spontaneous reactivation, as seen both in BCBL1 cells and in endothelial cells within KS tumors (62).

**Cytosine methylation on linker DNA in KSHV**

Bands of CG methylation separated by a nucleosome particle length of DNA occurred across nearly all episomes in the population downstream of the three CTCF sites in the LANA gene (Figure 1B and C). It was unlikely coincidence that GC methylation peaked in the same locations (our unpublished observations). CTCF is known to organize arrays of nucleosomes (63). We therefore propose that nucleosomes downstream of CTCF in
the LANA ORF are highly positioned. Another potential example would be the methylation patch just upstream of the N–1 nucleosome in the RTA promoter. A corollary conclusion is that DNMTs methylate linker, but not nucleosomal DNA, in KSHV. This conclusion contradicts a report of preferential methylation of nucleosomal DNA within Arabidopsis (64). Curiously, only linkers between nucleosomes were methylated. No patch of CG methylation appeared in an open TSS or between CTCF sites. Nor were all linkers sites of CG methylation: although the 5′ end of RTA ORF was occupied by nucleosomes at varying translational positions, little CG methylation occurred there. Finally, TIVE-LTC cells, which appeared to have the highest nucleosome density (i.e. least accessibility to M.CviPI) did not develop patches of CG methylation. Evidently, although CG methylation was restricted to linker DNA, it had other unknown determinants as well, post-translational modifications of histones perhaps. Indeed, several DNMTs interact with histone tails [reviewed in (65)].

Epigenetic diversity in KSHV

The discovery of partial CG methylation upstream of RTA provided a means to test whether the diverse chromatin states that we observed in BCBL1 episomes were epigenetic. Günther and Grundoff (3) showed that repression of RTA in latency was independent of DNA methylation in PEL and SLK cells. We have now observed the same in TIVE-LTC cells (Figure 8A). Thus, we do not conclude that methylation of 2–3 sites upstream of the RTA promoter affects transcription of the locus. However, we hypothesize that, as DNA methylation is epigenetic, the sites can be used to ‘fingerprint’ distinct epigenetic lineages within the BCBL1 viral population.

At the RTA promoter, the degree of upstream CG methylation correlated with the chromatin conformation identified by MAPit. Whereas 31% of episomes in the most compact chromatin state (Figure 3C, cluster 0) had the upstream patch of CG methylation, only 15% of molecules in the intermediate-open conformations (clusters ii–vii) did (Figure 6B). This uneven distribution suggested that there were at least two epigenetic states within latently infected BCBL1 cells. Furthermore, these states likely differed in their propensity to reactivate lytic expression, as cluster 0 molecules were unchanged by addition of TPA.

Because the TREx BCBL1-RTA cells, cloned from BCBL1, lacked the upstream patch of CG methylation, we could not test whether resistance to direct RTA induction was epigenetic. However, it was interesting to note that ∼40% of TREx BCBL1-RTA endogenous RTA promoters were in a compact state, and the proportion in this state was not significantly affected by induction of RTA in trans.

Several latent states—L1, L2 and L3—have been defined in a close relative to KSHV, the Epstein–Barr virus (EBV). These states are distinguished by expression of different subsets of the latent genes. In EBV, CTCF restricts spread of DNA methylation (66). Rare events in which CTCF function is compromised could allow EBV to

Figure 9. Model for three distinct epigenetic states of KSHV defined by MAPit at RTA promoter. (A) In the most open latent state, Latent 2 (L2), unfolding of higher-order chromatin structure transiently exposes the TSS and an upstream linker. With induced or spontaneous reactivation, transcription factors (polygon labeled TF) replace repressors (not shown) near the TSS. This leads to lytic replication, producing nucleosome-free DNA (far right). (B) The exposed linker in L2 can be CG methylated (triangles) by endogenous DNMTs. This methylation correlates with a shift to another epigenetic state, Latent 1 (L1), in which chromatin tends to be more closed. L1 virus responds to phorbol ester induction of lytic transcription weakly or not at all. (C) A third state, Latent 0 (L0), occurred in TIVE-LTC cells. Although in these cells, different stages of chromatin folding were observed, neither DNMTs nor transcription factors were able to access internucleosomal linker DNA, perhaps due to the presence (or absence) of specific histone modifications. L0 cannot be induced to reactivate by phorbol ester.
open structure at RTA, with the promoter poised for reactivation. We have not yet determined the range of chromatin conformations accessed by each epigenetic state, only that the L1 state is more likely to have a closed chromatin configuration at RTA. It seems likely that the same episomes have compact chromatin at the LANA and vIL6 promoters, given that the proportions in the closed state were the same at all three loci.

We hypothesize that infrequent methylation of RTA promoter DNA accompanied chromatin compaction and the transition from L2 to L1. Our findings suggest that the gradual methylation of episomes following infection (3) is linked to phenotypic changes. DNA methylation, at least at the RTA promoter, was associated with loss of both chromatin plasticity (Supplementary Table S3) and accessibility to soluble factors (e.g. M.CviPI). Progressive silencing of KSHV episomes would serve as a natural defense for the host cell. It could eventually lead to repression of LANA and loss of infection, a phenomenon observed in vitro (68,69). These results support a model originally proposed by Grundhoff and Gamem (2004) (68) in which lytic reactivation, albeit infrequent, is required for life-long persistence of KSHV infection. Recently, Kobiler et al. (70), using the Brainbow cassette to trace the lineage of viral genomes during lytic replication, demonstrated that only 6–10 genomes within each cell (i.e. <1%) gave rise to progeny virus. Our observations suggest that a similar restriction may apply to KSHV reactivation from latency.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online: Supplementary Tables 1–3, Supplementary Figures 1–6 and Supplementary References [71–73].

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