Protein Arginine Methyltransferase 1-directed Methylation of Kaposi Sarcoma-associated Herpesvirus Latency-associated Nuclear Antigen*1″

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Background: Post-translational modifications generate functional heterogeneity of viral regulatory factors.

Results: Viral chromatin association by Kaposi sarcoma-associated herpesvirus (KSHV) latency-associated nuclear antigen (LANA) is modulated by protein arginine methyltransferase 1 (PRMT1)-directed methylation.

Conclusion: Methylation of KSHV LANA antagonizes viral reactivation.

Significance: Protein methylation contributes to the functional properties of viral regulatory proteins, including KSHV LANA.

The Kaposi sarcoma-associated herpesvirus (KSHV) latency-associated nuclear antigen (LANA) is a multifunctional protein with roles in gene regulation and maintenance of viral latency. Post-translational modification of LANA is important for functional diversification. Here, we report that LANA is subject to arginine methylation by protein arginine methyltransferase 1 in vitro and in vivo. The major arginine methylation site in LANA was mapped to arginine 20. This site was mutated to either phenylalanine (bulk hydrophobic, constitutive methylated mimetic) or lysine (positively charged, non-arginine methylatable) residues. The significance of the methylation in LANA function was examined in both the isolated form and in the context of the viral genome through the generation of recombinant KSHV. In addition, authentic LANA binding sites on the KSHV episome in naturally infected cells were identified using a whole genome KSHV tiling array. Although mutation of the methylation site resulted in no significant difference in KSHV LANA subcellular localization, we found that the methylation mimic mutation resulted in augmented histone binding in vitro and increased LANA occupancy at identified LANA target promoters in vivo. Moreover, a cell line carrying the methylation mimic mutant KSHV showed reduced viral gene expression relative to controls both in latency and in the course of reactivation. These results suggest that residue 20 is important for modulation of a subset of LANA functions and properties of this residue, including the hydrophobic character induced by arginine methylation, may contribute to the observed effects.

Kaposi sarcoma-associated herpesvirus (KSHV3/human herpesvirus 8) is a γ-herpesvirus linked to Kaposi sarcoma (KS) and at least two rare lymphoproliferative disorders, primary effusion lymphoma (or body cavity-based lymphoma (BCBL)) and a subset of Multicentric Castleman disease (1–4). Like all herpesviruses, KSHV has distinct latent and lytic phases whose transition is regulated primarily at the transcriptional level by transcription, silencing, and chromatin remodeling of the viral genome (5–10). KSHV latency-associated nuclear antigen (LANA/ORF73) is a key regulatory protein that is essential for the establishment and maintenance of viral latency. LANA is a DNA-binding protein that binds to the viral latent origin of replication located at the terminal repeat sequence of the KSHV genome, is highly expressed in all KSHV-associated disorders (11–13), and is a functional homologue of Epstein-Barr virus transcription factor EBNA1 (14–17). LANA functions as both a transcriptional activator and a repressor depending on the context of promoters and cell line interrogated (18, 19). Accordingly, LANA has been shown to associate with a broad range of transcriptional regulators such as RBP-Jκ, CBP, Daxx, BRD2, RBp53, and Sp-1 (20–26). In addition to influences via protein partner interaction, LANA function has also been reported to be regulated by post-translational modifications, including phosphorylation (27, 28), acetylation (29), poly(ADP-ribosyl)ation (30), and sumoylation.4 Among these post-translational modifications, phosphorylation, acetylation, and poly(ADP-ribosyl)ation have been reported to antagonize LANA function, measured by either genomic copy number alterations or derepression of LANA transcriptional targets such as ORF50 (28–30).

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1 The abbreviations used are: KSHV, Kaposi sarcoma (KS)-associated herpesvirus; LANA, latency-associated nuclear antigen; BCBL-1, body cavity-based lymphoma; PRMT, protein arginine methyltransferase; BAC, bacterial artificial chromosome; FRT, Flp recombinase target; FLPe, enhanced Flp recombinase; qPCR, quantitative PCR.

4 Y. Izumiya, unpublished results.
Protein arginine methylation is a post-translational modification frequently observed in nucleic acid-binding proteins. The mammalian PRMT family consists of nine highly conserved members that catalyze distinct types of methyl group addition to the guanidino group of protein arginine residues (31). Substrates include many proteins involved in RNA metabolism (32); however, the list of targets has become increasingly broad, encompassing transcriptional coactivators, corepressors, DNA repair factors, and signaling molecules (31, 33). Viral proteins are also modified by PRMTs. This class of substrates includes a variety of viral proteins including adenovirus type 5 100K (34, 35), Epstein-Barr virus EBNA1 (36), herpes simplex virus type 1 ICP27 (37, 38), human immunodeficiency virus type 1 Tat (39), hepatitis C virus NS3 (40), and hepatitis δ virus small form antigen (41). The consequences imparted by arginine methylation on viral protein function include alterations in target localization (35, 36, 38), transcriptional activation (39), viral gene expression (38), and viral replication (34, 38, 39, 42).

In this report we demonstrate that KSHV LANA is arginine-methylated in vitro and in vivo by PRMT1. Using a strategy of amino acid substitution mutants of the primary arginine methylation site in LANA in the context of the KSHV genome as well as in its isolated form, our results suggest that methylation of LANA modulates the transcriptional control of viral gene targets, mediated in part by a mechanism that involves modulation of the histone binding function of LANA.

EXPERIMENTAL PROCEDURES

Plasmids—The cloning strategy employed throughout these studies involved cloning of fragments of interest into several vectors that contain a Cpol site inserted into each polyclinker. This strategy has been described previously (6, 43–46). Fragments were digested with Cpol, desalted (Qiagen) and cloned into Cpol-digested vectors for expression in mammalian cells (pCDNA3-Cpol), lentiviral transduction (pLTRexBGHPACpol), baculoviral expression (pFastBac-Cpol), and for GST pulldown assays (pGEX-2T-Cpol). The resultant clones from all procedures possess an N-terminal FLAG, HA, or GST tag. A full-length wt-LANA expression vector was constructed as described above using cDNA prepared from BCBL-1 cells. This plasmid served as the template for mutagenesis and for construction of truncated LANA fragments by PCR. Oligonucleotide sequences used throughout this study are available upon request.

Protein Methylsome Library—Details of the methylase/demethylase library will be described elsewhere.5 Pertinent to this publication, PRMTs 1–8 were cloned by PCR using Pfu Turbo (Agilent Technologies), and cDNA was prepared from a pool of RNAs from several cell lines.

Mutagenesis—Mutations were introduced into wt-LANA using QuickChange II kit (Agilent Technologies). LANA amino acids 1–70 was inserted into plasmid pGex-2T-Cpol. This plasmid served as the template for mutagenesis of the individual N-terminal arginine residues of LANA.

Cell Lines—293T (SV40 large T/human embryonic kidney epithelial) and SLK cells (human endothelial cells) (47, 48) were grown in monolayer culture in DMEM, supplemented with 10% fetal bovine serum (complete DMEM) in the presence of 5% CO2. 293T BAC stable cell lines were selected and maintained in complete DMEM containing 100 μg/ml hygromycin. SLK lentiviral stable cells were selected and maintained in complete DMEM containing 1 μg/ml puromycin.

Immunoprecipitation and Immunoblotting—The procedures used have been described (43). Immunoprecipitations (IP) utilized 2 μg of preimmune IgG or 2 μg of specific IgG per immunoprecipitation. For immunoblotting, the final concentration of all primary antibodies was 1 μg/ml. Primary antibodies used were anti-FLAG-M2 (Sigma), anti-HA (Covance), antilamin A/C (Cell Signaling Technology), anti-β actin (Sigma), anti-PRMT1 (Sigma), or anti-H2B (Millipore). Secondary antibodies were obtained from Santa Cruz Biotechnologies.

Methylation Assays—Purified recombinant proteins were incubated for 3 h at 37 °C with 500 nm recombinant PRMT in 30 μl of methylation buffer (20 mM Tris (pH 7.5), 150 mM NaCl, 1 mM EDTA, and 1 mM PMSF) supplemented with 1 μl (0.5 μCi) of S-adenosyl-L-[methyl-3H]methionine (Amersham Biosciences). Reactions were stopped by the addition of 2× SDS-PAGE sample buffer and heating. Samples were analyzed by SDS-PAGE and autoradiography. For in vivo methylation, cells were labeled for 3 h at 37 °C with 50 μCi of S-adenosyl-L-[methyl-3H]methionine in the presence of 100 μg/ml cycloheximide. After labeling, the cells were processed for immunoprecipitation using anti-FLAG antibody. After electrophoresis of the immunoprecipitates, the gels were subjected to fluoroigraphy with Amplify (GE Healthcare) and exposed to film at −80 °C.

Gene Knockdown—The shRNA directed against PRMT1 was obtained from Open Biosystems. Cells were transfected using FuGENE HD (Roche Applied Science). The shRNA expression plasmid was cotransfected with FLAG-LANA, and the knockdown was started 72 h before the methylation labeling procedure. After the labeling step, cell lysates were prepared and analyzed for LANA methylation as described above.

Cellular Fractionation—SLK, SLK-LANA, and SLK-LANA Arg-20 mutants were fractionated into nuclear and cytoplasmic pools using the NE-PER kit (Pierce). The nuclear matrix fractionation protocol has been described (49).

Recombinant Protein Production—Spodoptera frugiperda Sf9 cells were maintained in EX-CELL 420 medium (JRH Biosciences), and recombinant baculoviruses were generated as previously described (43). Recombinant baculovirus bacmid DNA was transfected into Sf9 cells by using FuGENE 6 (Roche Applied Science), and recombinant viruses were subsequently amplified twice. Expression of recombinant proteins was confirmed by immunoblotting with anti-FLAG monoclonal antibody. Large scale cultures of Sf9 cells (100 ml) were infected with recombinant baculovirus at a multiplicity of infection of 0.1–1.0, and cells were harvested 48 h after infection. Recombinant proteins were purified as described previously (6). The purity and amount of protein was measured by SDS-PAGE and Coomassie Blue staining with BSA as a standard. The preparation and purification of GST-fusion proteins has been described (6).

Recombineering Protocols—Mutagenesis of a bacterial artificial chromosome containing the entire KSHV genome (BAC-
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16) was performed by using a recombineering system (58). The strains and vectors used in the recombineering procedure were obtained from the National Cancer Institute, National Institutes of Health. The host strain for λ-mediated recombination was a BAC-16/SW105 transformant. A LANA targeting vector was constructed using modified pCD-LANA that contains a 5′ KSHV —200-bp homology arm followed by an ampicillin resistance (Bla) cassette upstream of the LANA coding region. The Bla cassette is flanked by site-specific recognition (FRT) sequences for the FLPe recombinase (50). The linearized targeting cassette contains LANA homology arms of 0.2 and 1 kb on the 5′ and 3′ sides of the ampicillin resistance marker. The targeting cassette was released from the vector sequences by digestion with PstI and BamHI. The fragment was gel-purified and electroporated into induced (recombination +) SW105/BAC-16 cells. Ampicillin-resistant transformants were selected and analyzed by PCR for insertion of the targeting cassette. Two or three positive clones of each intermediate were then grown at 32 °C with removal of the targeting cassette accomplished by arabinose induction of the FLPe recombinase. Ampicillin-sensitive clones were screened by PCR for removal of the targeting cassette. Clones positive by PCR for removal of the cassette were then further verified by sequencing the targeted region of LANA. A similar approach was used for the Arg-20 mutants that were cloned into the targeting construct. The introduced Arg-20 mutations were confirmed by sequencing, and the integrity of each recombinant BAC was examined by restriction enzyme digestions and Southern blotting.

LANA Wild-type Revertant Constructs—LANA Arg-20 mutant alleles in BAC-16 were reverted to wild-type using the recombineering protocols described above. For these constructions, the mutant LANA allele in the KSHV bacmid was replaced with wild-type LANA using the wt-LANA targeting cassette. Reversion of each mutant LANA allele to wild-type was confirmed by sequencing. 293T BAC stable cell lines containing each revertant bacmid were then created as described below.

Construction of BAC Stable Cell Cultures—293T cells were transfected with 5 μg of recombinant BAC DNA using HEKfectin (Bio-Rad). After 2 days of culture, the cells were expanded to a single T175 flask, and hygromycin (100 μg/ml) selection was started. After approximately 2 weeks, the hygromycin-resistant colonies that emerged from a single flask were trypsinized and pooled to establish each culture. Early passage pools were used throughout the experiments described herein.

ChIP, KSHV Tiling Array, and ChIP-on-chip—ChIP assays were performed following the protocol described by the Farnham laboratory at the University of California-Davis. 293T BAC stable cell cultures were used in the ChIP experiments, and Dox-inducible K-Rta-BCBL-1 cells were used in ChIP-on-chip experiments (51). Antibodies used for LANA immunoprecipitation were rat anti-LANA (Advanced Biotechnologies, Inc.) and rat non-immune serum IgG (Alpha Diagnostic International). The KSHV tiling array was designed across KSHV genome sequence (NC_009333.1) and manufactured by Agilent Technologies. The probes were spotted on 8 × 15 K array format. The details of this array have been described recently (10). Labeling, hybridization, scanning, and analyzing of the arrays were done at UC Davis Cancer Center Genomic Shared Resource. The enrichment of wt-LANA binding to the KSHV genome was calculated by the intensity ratio of immunoprecipitated and input DNA for each array spot, and the probe signals were further normalized by blank subtraction and the Whitehead error model using Genomic Workbench software. Each ChIP-on-chip experiment was performed in triplicate. For real-time PCR analysis of ChIP samples, the quantity of DNA amplified corresponding to specific KSHV loci was compared with the matching input samples. Results are expressed as the ratio of the ChIP sample amplification signal normalized to the signal obtained using the sample input control.

In Vitro Histone Binding—Binding reactions were carried as described (6, 52). Binding buffer contained 20 mM Hepes (pH 7.9), 150 mM NaCl, 1 mM EDTA, 4 mM MgCl₂, 0.1% Nonidet P-40, 10% glycerol, and 1 mM PMSF. Histone octamers were formed by resuspension of lyophilized calf thymus histones (Roche Applied Science) in 2 M NaCl followed by stepwise dialysis against PBS at 4 °C overnight. Binding reactions (20 μl) contained LANA GST fusion protein and 0.5 μg/μl histone octamer. Reactions were incubated at 4 °C for 90 min. Wash buffers contained binding buffer supplemented with 125–500 mM NaCl. After extensive washing, the GST beads were resuspended in 2× sample buffer, boiled, and loaded on SDS-PAGE-15% gels. The gels were subjected to immunoblotting as described above and probed for H2B (43).

RESULTS

LANA Is Arginine-methylated in Vitro—Our laboratory has developed a methylase/demethylase library that contains nearly the entire repertoire of human PRMTs, lysine methyltransferases, and lysine demethylases.5 Included in this library are eight of the nine mammalian PRMTs described to date. As part of a search for novel protein modifications directed by PRMTs, we screened a panel of recombinant viral proteins for their ability to serve as substrates for methylation in vitro. This panel included several KSHV proteins including LANA, ORF K8 (K-bZIP), ORF50 (K-Rta), and viral DNA replication enzymes. Although several important KSHV regulatory factors such as K-bZIP and K-Rta did not score in this assay, one strong positive hit that emerged from this screen was LANA. An example of the methylation reactions utilized in this screen is shown in Fig. 1, B and C. Each reaction contained purified, recombinant substrates (i.e. LANA), individual PRMTs (Fig. 1A), and S-adenosyl-1-[methyl-³H]methionine. A methylated
protein with a molecular weight consistent with the migration of LANA was only observed in reactions containing PRMT1 and PRMT8 as the enzymatic activity (Fig. 1B, lanes 1 and 8). As PRMT8 expression is largely restricted to the brain and is localized to the cell membrane (53), we have focused our studies on the relationship between PRMT1 and LANA. Control reactions using histone substrates demonstrated that all of the PRMTs were enzymatically active (data not shown), although there were differences in specific activity. Using a series of LANA deletion proteins as substrates, we narrowed the methylation site(s) to the N-terminal 1/11011 350 amino acids of LANA (data not shown). Further deletion analysis of this region of LANA mapped the methylation site(s) to the N-terminal 20 amino acid residues of LANA (Fig. 2A). This region of LANA contains a single RG motif, characteristic of the arginine methylation site glycine and arginine-rich patches (54). The first 5 N-terminal arginine residues of LANA (amino acids 1–70) were individually mutated to lysine, and the mutant proteins were purified as GST fusions (Fig. 2B). Using these substrates, a single PRMT1 methylation site was mapped in vitro to LANA residue Arg-20 (Fig. 2, B and C). Taken together, these results map the major in vitro PRMT1 methylation site in LANA to residue Arg-20 (Fig. 2, B and C).

LANA Is Arginine-methylated in Vivo—As a next step, arginine methylation of LANA was evaluated in vivo. 293T cells were transfected with expression vectors expressing full-length FLAG-tagged wt-LANA or LANA R20K. Two days post-transfection, the cells were labeled for 3 h with [3H]methionine in the presence of a high concentration (100 μg/ml) of cycloheximide. Lysates were prepared, and LANA was immunoprecipitated using anti-FLAG antibody. Methylation of wt-LANA was readily detectable (Fig. 2D, lanes 2 and 4). In contrast, the signal was markedly reduced in immunoprecipitates prepared from LANA R20K lysates (Fig. 2D, lane 3). These results suggest that Arg-20 is the major methylation site in LANA but that there may be additional, minor methylation sites yet to be defined. Additional experiments demonstrated that partial depletion of cellular PRMT1 expression by shPRMT1, but not shControl, greatly reduced the LANA methylation signal (Fig. 2D, lanes 4 and 5). Conversely, transient overexpression of PRMT1 significantly increased LANA methylation (Fig. 2D, lane 6). Control blots confirm the partial knockdown and overexpression of PRMT1 in these experiments (Fig. 2E). Importantly, LANA physically associated with PRMT1 in transfected 293T cells, suggesting the specificity of the methylation (Fig. 2F). These results established that PRMT1 interacts with and methylates LANA in vivo. A summary of the methylation site mapping

FIGURE 1. LANA is arginine-methylated in vitro. A, purified PRMTs produced in baculovirus-infected cells were electrophoresed on 8% SDS-PAGE and stained with Coomassie Brilliant Blue. Numbers at the top of each lane of the gel signify PRMT designation. B, PRMT screening against LANA is shown. Numbers at the top of each lane of the gel signify PRMT designation. Reaction products were electrophoresed on 8% SDS-PAGE and processed as described under “Experimental Procedures.” The arrow denotes arginine-methylated LANA. C, confirmatory methylation reactions demonstrate methyltransferase activity of PRMT1 and PRMT8 on LANA substrate.
experiments in relationship to the N-terminal LANA histone binding domain, nuclear localization signal (NLS), and potential phosphorylation sites is shown in Fig. 2 (52, 55).

LANA Arginine Methylation and Subcellular Localization—Our mapping studies positioned the arginine methylation site adjacent to (i) the LANA major histone binding domain and (ii) the LANA NLS (52, 55). Arginine methylation has been shown to regulate subcellular localization of proteins (36). Therefore, the effect of methylation on LANA localization was evaluated using cellular fractionation procedures on lysates prepared from SLK cells engineered to overexpress wt-LANA or LANA Arg-20 mutants. We substituted the positively charged arginine 20 methylation site residue of LANA with either of two amino acids in an attempt to infer the effect of methylation on LANA function. In one case, a bulky hydrophobic residue (Phe) was introduced; this residue has been proposed to mimic the constitutively arginine-methylated state (56).

SLK cells were transduced with lentiviral vectors encoding FLAG-tagged versions of LANA or mutant LANA and selected with puromycin. To avoid clonal variation, mass cultures of puromycin-resistant pools of cells were screened for LANA wild-type or Arg-20 mutant expression by Western blotting. The immunoblots shown in Fig. 3 demonstrate that the steady-state level of LANA protein was similar among the cell lines, suggesting similar protein stability among the mutants. Lysates from these cell lines were then fractionated into cytoplasmic and nuclear pools, and each fraction was probed for LANA by Western blotting. The results showed that LANA protein is entirely nuclear regardless of the status of Arg-20 (Fig. 3B). Moreover, probing of nuclear matrix fractions from this same series of SLK cells demonstrated that LANA or mutant LANA was preferentially associated with the nuclear matrix fraction (Fig. 3B). Taken together, these results suggest that methylation is not involved in either LANA protein stabil-
ity or nuclear localization, effects attributed to arginine methylation that have been observed with other viral proteins (34, 40).

**LANA Arg-20 Mutation Shows Increased LANA-Histone Octamer Interaction**—As a next step we examined effects on histone binding. In contrast to the lack of effect of the Arg-20 methylation site on LANA subcellular localization (Fig. 3), this site did influence LANA-histone interaction. An elevated degree of *in vitro* histone octamer binding by the R20F methylation mimic LANA protein was observed under our standard binding conditions (Fig. 4Aa, lane 2). Moreover, when preformed LANA-histone octamer complexes were washed with binding buffers containing increasing amounts of sodium chloride, the LANA R20F mutant protein showed increased binding stability at higher salt concentrations relative to wild-type LANA (Fig. 4Aa, a, lanes 3–5, and b). In addition, prior methylation of wild type, but not R20F mutant LANA, by PRMT1 also resulted in increased histone binding *in vitro* (Fig. 4B). Note that in this latter series of experiments, PRMT1 was removed from the reactions subsequent to methylation by washes with high salt buffer (500 mM NaCl) with 5% glycerol before histone binding. As the protein substrates used in these experiments were produced in *Escherichia coli*, which do not encode arginine methylases, GST-wt LANA represents a purely non-methylated protein. These results suggest that methylation of LANA increased its affinity to histone octamers *in vitro*.

**Generation of LANA Arg-20 Mutants Using Recombineering**—The influence of the LANA arginine methylation site was examined in the context of the KSHV genome. This newly developed bacmid is called KSHV BAC-16. BAC-16 is derived from rKSHV.219; this is a recombinant KSHV in which GFP is constitutively expressed (*e.g.* in latency) (57). BAC-16 also contains the selectable resistance marker hygromycin. Details of the construction of the parental KSHV BAC-16 will be published elsewhere. Using this new BAC system in conjunction with recombineering protocols (58), we incorporated the LANA Arg-20 mutants into the KSHV genome, individually replacing the endogenous wt-LANA allele with the Arg-20 substitutions. A LANA wild-type control was also constructed in which the wt-LANA allele on the targeting vector was used to replace the corresponding LANA wild-type allele on BAC-16. In addition, revertant LANA alleles in each Arg-20 substitution BAC were constructed in which the wt-LANA allele on the targeting vector was used to replace the mutant LANA Arg-20 allele that had been previously introduced into BAC-16. These constructs served as additional controls for potential influences of unintended mutations introduced into the bacmids during the course of recombineering. Construction of each recombinant BAC utilized a two-step procedure as described under “Experimental Procedures” and is shown schematically in Fig. 6

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**FIGURE 3. Arginine methylation does not influence LANA cellular localization.** A, Western analysis (WB) of total wt-LANA/mutant LANA expression in SLK lentiviral stable cell lines is shown. B, Western analysis of cytoplasmic/nuclear and nuclear matrix-associated LANA fractionation from extracts prepared from SLK/lentiviral LANA/mutant LANA stable cell lines is shown. Blots were probed with the indicated antibodies.

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6 J. U. Jung, personal communication.
Recombineering at the LANA locus was carried out using a SW105 transformant carrying BAC-16. This strain carries the BAC of interest as well as inducible systems for /H9261-mediated homologous recombination (step 1) and FLPe-mediated excision (step 2) of the targeting cassette. In step one, a linearized targeting cassette containing the LANA substitution mutation of interest and a selectable marker (ampicillin) was introduced into the recipient BAC cells by electroporation. Recombination between the linear targeting cassette and the LANA locus on the BAC proceeded using /H9261-mediated homologous recombination machinery that is present in SW105 (Fig. 5A). Drug-resistant intermediates were then screened by PCR for the presence of the targeting cassette. In step two, the targeting cassette was excised from 1–3 individual positive intermediates using induced FLPe recombinase (Fig. 5A). Colonies (now ampicillin-sensitive) were screened by PCR for FRT-dependent excision of the selectable marker. The introduced mutation in the BAC was then sequence-verified. Final validation of the recombineered BAC was confirmed by restriction enzyme digestions and Southern blotting. Revertant LANA alleles were constructed in a similar fashion, except that recombination utilized individual SW105 strains carrying LANA Arg-20 mutant BACs and a wt-LANA targeting cassette. A Southern blot analysis of the LANA R20K mutant construction is shown in Fig. 5C. Probing recombinant BAC DNA clones with a LANA probe generated a band pattern consistent with an insertion of the targeting cassette at the LANA locus. BamHI-digested DNA purified from BAC intermediates was predicted to show a /H11011.1-kb band shift upon successful recombination of the targeting cassette (Fig. 5C, right panel, lanes B and C) and, similarly, return to the position of the BAC-16 parental band (Fig. 5C, right panel, lane A) upon FLPe-mediated removal of the targeting cassette (Fig. 5C, right panel, lanes D', E', and F'). When probed with an FRT-Bla fragment, only the BAC intermediates (Fig. 5C, left panel, lanes B and C) and not the final recombinants (Fig. 5C, left panel, lanes D, E, and F) or parental BAC (Fig. 5C, left panel, lane A) contain the corresponding fragment, as predicted. The band patterns obtained by Southern blot analysis of R20F and wt-LANA control bacmid constructs were identical to those listed in Fig. 5C. Final confirmation of each allele replacement strategy was conducted by sequence analysis of PCR products generated using primer pairs that bracket the mutation site in the

FIGURE 4. Enhanced histone octamer binding by methylation mimic LANA R20F and by arginine methylation of wt-LANA. Aa, in vitro histone binding by LANA and LANA R20F mutant is shown. H2B immunoblot analysis (WB) of in vitro histone binding reactions is shown. GST-LANA 1–70, GST-LANA 1–70 R20F mutant, or GST control beads were incubated with histone octamer as described under “Experimental Procedures.” Binding reactions were then washed with binding buffer (lane 2) or binding buffer supplemented with 125 mM (lane 3), 250 mM (lane 4), or 500 mM (lane 5) NaCl. 5% of input histone octamer is shown for each. Loading controls for each reaction are shown by GST immunoblot. Coomassie Brilliant Blue staining of each GST-LANA species used in the reactions is also shown. b, shown is quantitation of the H2B binding immunoblots in a (circles, wt-LANA; diamonds, LANA R20F; triangles, GST). B, coupled in vitro methylation/histone binding reactions are shown. GST-LANA 1–70 and GST-LANA 1–70 R20F mutant were incubated with recombinant PRMT1 (2 μM) and S-adenosyl-L-methymethionine (4 mM) for 3 h at 37°C. Control reactions were similarly set up without PRMT1. After incubation, the beads were washed several times with PBS containing 500 mM NaCl plus 5% glycerol. The beads were then equilibrated with histone binding buffer and assayed for histone octamer binding under standard conditions as described under “Experimental Procedures”.
recombinant BAC DNA. Fig. 5D depicts a small portion of the nucleotide sequence of the 5′ end of the LANA locus near the LANA ATG and compares this sequence with the residual nucleotides that remain in the final BAC recombinants. The recombinant BAC DNAs were then transfected into 293T cells, and mass cultures of hygromycin-resistant BAC stable cells (Fig. 5B) were created containing each of the Arg-20 mutants, the wt-LANA control, or the LANA revertants. After Hirt DNA extraction (59) and qPCR using LANA-specific primers, each culture was found to contain 1–5 genome copies per cell using a BAC-16 bacmid standard curve (data not shown). The BAC cultures were then used to study the effect of LANA methylation at position Arg-20 on KSHV viral gene expression.

LANA Occupancy of KSHV Genome during Latency—We wished to examine the effects of Arg-20 methylation on KSHV gene expression profiles of several select viral genes during latency and viral reactivation using the BAC stable cell lines. Before undertaking these studies, we needed to determine the LANA target viral genes throughout the KSHV genome, as we envisioned using the stable cell lines to measure KSHV gene expression of viral ORFs, whose gene expression was directly regulated by LANA. To achieve this, we constructed a KSHV tiling array and conducted a LANA ChIP on KSHV chip analyses with K-Rta-inducible BCBL-1 cells (51). Surprisingly, although there are peaks and troughs in the LANA binding pattern, the array depicts LANA occupancy throughout the entire latent KSHV genome when we considered 2-fold enrichment as significant (Fig. 6A). ChIP–KSHV-chip analyses were verified by qPCR with primers specific for the LANA promoter region (Fig. 6Ba). Interestingly, during viral reactivation there is an immediate (i.e. within 4 h) disassociation of LANA from the KSHV genome after K-Rta induction (supplemental Fig. 1). These results propose a model in which LANA may play a key role in maintaining the entire latent KSHV epigenome during latency and rapidly disassociates from the genome during the early stages of reactivation.

Increased Chromatin Interaction in Vivo by LANA R20F Mutation during Latency—After major LANA binding sites were in hand, we examined the effects of arginine methylation on recruitment of LANA to select target regions. LANA ChIP analysis was used to examine the degree of LANA wt or Arg-20 mutant association with the KSHV genome. We examined several regions where LANA was significantly enriched in naturally infected cells. Two regions containing lower levels of LANA enrichment (vIRF3 and ORF64) were also included in the analysis. As shown in Fig. 6Bb, the ChIP analysis demonstrated an increase in LANA R20F occupancy within 7/7 LANA recruitment sites in viral genes analyzed during latency, which
LANA Arginine Methylation and KSHV Gene Expression—If LANA plays a role in gene repression during reactivation (60) and LANA R20F differs in recruitment, one may expect to see the effects on viral gene expression at LANA target genes. RNA was prepared from 293T BAC stable cells harboring BAC-16 episomes with the LANA R20F allele were examined, KSHV gene expression was reduced under both latent and reactivated conditions relative to 293T wt-LANA cells (Fig. 7A and data not shown). When viewed as -fold changes in response to reactivation, the gene expression results in the R20F culture were markedly reduced relative to the response of the wt control or R20K cell lines (Fig. 7B). Importantly, this profile of gene expression correlated with recruitment of LANA, where higher enrichment of LANA showed reduced target gene expression. ORF73 expression was modestly stimulated in LANA R20F Bac stable cells, whereas wt- and R20K LANA expression declined slightly during reactivation (Fig. 7A). Similar results to those described above were obtained when we conducted this experiment using (i) K-Rta

is consistent with the observed increase in affinity toward histone octamers in vitro. In contrast, the decrease in LANA R20K occupancy at this same panel of viral genes was reduced (5/7 genes) or equivalent (2/7 genes) to wt-LANA binding (Fig. 6Bb). Consistent with the LANA ChIP-KSHV-chip results, using Bac stable LANA ChIP DNA, non-enriched genes (vIRF3 and ORF64) also showed low levels of LANA recruitment during latency when analyzed by qPCR (Fig. 6Bb).

**FIGURE 6. LANA Arg-20 mutations affects KSHV chromatin binding during latency.** 

A, KSHV wt-LANA whole genome tiling array is shown. LANA ChIP DNAs were prepared from K-Rta-inducible BCBL-1 cells during latency. The immunoprecipitated DNA was probed against a KSHV tiling array as described under “Experimental Procedures.” The graph depicts the ratio of LANA/input signal across the KSHV genome. The entire KSHV genome is annotated at the bottom of the figure. The arrows denote ORFs evaluated by ChIP and/or RT-qPCR in the 293T BAC stable cell cultures. Several regions of the KSHV genome are indicated above the graph for reference. B, validation of the anti-LANA antibody in LANA ChIP assay is shown. ChIP DNA samples were prepared as described under “Experimental Procedures” and immunoprecipitated with anti-LANA antibody or normal rat IgG. DNA associated with each immunoprecipitate was analyzed by qPCR using primer pairs against the ORF73 promoter. Values are reported as the ratio of the Ct values of the amplified DNA immunoprecipitated relative to the amount of input DNA amplified. b, ChIP analysis is shown. ChIP DNA was prepared from each BAC stable cell line during latency. ChIP DNA associated with each immunoprecipitate was analyzed by qPCR using primer pairs directed against the ORFs listed. Values are reported as the ratio of the Ct values of the amplified DNA immunoprecipitated with anti-LANA antibody relative to the amount of input DNA amplified.
overexpression as a means of reactivation and (ii) chemical reactivation using an independently derived series of 293T BAC stable cells (data not shown).

Finally, to show that the phenotype we observed is not due to nonspecific mutations, we constructed wild-type revertant alleles of each LANA Arg-20 mutant (Fig. 7C). 293T BAC stable cell lines of each revertant were established, and the gene expression phenotype of each culture was examined as described in the previous section. The revertant cultures contained amounts of KSHV genomic DNA copy numbers similar to the Arg-20 mutant cultures (data not shown). When compared with the gene expression results for the Arg-20 mutant cultures, all revertant BAC-containing cultures exhibited changes in KSHV gene expression (Fig. 7D) that approached the values obtained with the wt-LANA control BAC culture.

**DISCUSSION**

In this report we demonstrate that the KSHV LANA protein is subject to arginine methylation. The major arginine methylation site in LANA was mapped to residue Arg-20. This modification further expands the LANA repertoire of post-translational modifications including phosphorylation (28), acetylation (29), poly(ADP-ribosyl)ation (30), and sumoylation. Importantly, we could detect arginine methylated LANA in vivo, and we have preliminary estimates of this value as \( \frac{H_1}{H_11349} 2\% \) LANA methylation relative to the total amount of immunoprecipitated LANA present in latent BCBL-1 cell extracts (supplemental Fig. 2A). A higher value of \( \frac{H_1}{H_11011} 5\% \) methylation was obtained analyzing lysates from LANA-transfected 293T cells (supplemental Fig. 2B). One issue that remains is when and where LANA arginine methylation occurs and the relative levels of arginine-methylated versus unmethylated LANA under different conditions. Indeed, the question of the relative levels of modified versus unmodified species applies to all post-translational protein modifications, including LANA modifications.

It is tempting to speculate that methylation of LANA may be a signal for recruiting protein complexes, which recognizes pro-
protein methylation as seen in cellular histone binding proteins (for a recent review, see Ref. 61). Specific tools such as modified LANA-specific antibodies may be needed to answer those questions. The exact nature and location of the methyl mark(s) on LANA molecules present in BCBL-1 cells is still not known. However, because PRMT1 is classified as a type I PRMT (31) and our knockdown data indicate that PRMT1 is the predominant enzymatic activity responsible for methylation of transfected LANA in 293T cells, the arginine methyl mark on LANA at position Arg-20 is most likely to be asymmetric dimethylation. Our detailed mutagenesis studies, which focused on the N terminus of LANA, suggest that Arg-20 is the major arginine methylation site. However, our results using the full-length LANA R20K mutant indicate that there may be other minor methylation sites in LANA. The role and nature of these additional methylation sites remain to be determined. We have conducted large-scale LANA immunoprecipitations using nuclear extracts prepared from HeLa-LANA-overexpressing cells and have also identified PRMT1 as a LANA interacting partner.5 This observation taken together with the ability of LANA to be methylated in vitro and in vivo by PRMT1 suggests that LANA is a genuine PRMT1 substrate and not a physically interacting but non-methyl accepting partner, as has been observed for other PRMT-interacting partners (56, 62, 63).

Our mapping of the LANA arginine methylation site to residue Arg-20 is very intriguing given the fact that this site is interposed between the dominant LANA histone binding domain (52) and the LANA NLS (55). Moreover, this region of LANA contains several predicted protein phosphorylation sites, suggesting the possibility for modulation of LANA function via phospho-methyl modification cassettes or switches as has been seen in other proteins (64, 65). Mutation of LANA residue Arg-20 did not influence LANA protein stability, nuclear localization, or nuclear matrix association; however, this residue did affect LANA histone binding activity in vitro and LANA-KSHV chromatin association in vivo. Importantly, cell lines harboring LANA R20F mutant BACs responded to reactivation signals very poorly. Inspection of the ChIP and gene expression data for a given KSHV locus suggests an inverse correlation between LANA association during latency and the -fold response to chemical reactivation. A high relative level of LANA binding within a specific KSHV gene resulted in a lower -fold response to chemical reactivation, whereas a low or intermediate level of binding had the opposite effect. ORF73 was an exception to this trend, where expression was modestly stimulated in LANA R20F BAC stable cells, possibly the result of hyperstimulation of the LANA promoter (66) by increased R20F binding. Taken together, we interpret these results to suggest that arginine methylation of LANA results in a LANA molecule that is more effective at KSHV chromatin binding and subsequent silencing of certain regions of the KSHV genome. Importantly, the arginine methylation site is conserved in Rhesus monkey rhadinovirus ORF73. In fact, Rhesus monkey rhadinovirus ORF73 contains five RG motifs in the N-terminal region of ORF73, indicating a possible conserved function of arginine methylation. Consistent with our data suggesting that Arg-20 methylation strengthens the LANA-chromatin interaction, LANA R20K (a non-arginine-methylatable residue) exhibited the lowest relative LANA occupancy at seven of nine KSHV loci examined and a higher-fold response at two loci where this parameter was assayed (K2 and K12). These results suggest that arginine methylation, in addition to other factors such as target DNA sequence and other DNA binding proteins, may influence LANA chromatin binding. Interestingly, a recent report showed that methylation of EBNA2 is required for interaction with DNA-bound RBP-Jκ to modulate gene expression (67). Because LANA has been shown to associate with RBP-Jκ, this function may be conserved between Epstein-Barr virus and KSHV (20).

The replacement of arginine with a lysine residue does preserve the positive charge of the position and renders this site refractory to arginine methylation; however, this introduction might be potentially complicated due to the ability of lysine to be subjected to other post-translational modifications such as lysine methylation, acetylation, and ubiquitylation. Importantly, the analysis of LANA Arg-20 revertant alleles suggests that the observed gene expression phenotype among the LANA Arg-20 mutants is LANA-specific and not due to the introduction of nonspecific mutations outside of the targeted region. Nonetheless, we acknowledge that the notion of whether or not the R20F mutation accurately mimics methylation is subject to debate, and therefore, it can be argued that if Arg-20 plays an important role in LANA function in the unmethylated state, then any mutation of this residue would impact LANA function.

Another key function of LANA is episomal maintenance during latency. Because protein methylation has been reported to affect both protein-protein and protein-nucleic acid interactions (31, 54), it is possible that arginine methylation of LANA may influence KSHV episomal maintenance. However, our experiments suggest that this may not be the case. After transient transfections of BAC DNA into 293T cells without incorporation of a selection procedure, GFP fluorescence declined precipitously for all BAC transfectants regardless of the LANA Arg-20 status (supplemental Fig. 3).

When combined with recombineering protocols, the KSHV bacmid used in these experiments allows one to readily construct and evaluate mutations and the corresponding revertant alleles within the context of the KSHV genome. The procedure adapted herein is very efficient and yields a very low background of non-recombinants. We have utilized our current mutagenesis protocol to introduce mutations into multiple KSHV loci including K-Rta and K-bZIP. The BAC-16 system could also be amenable to recombineering procedures that utilize single-stranded oligonucleotides (68) or short denatured PCR fragments as targeting vectors (69) to further facilitate BAC-16 manipulation.

In the course of conducting these experiments, we carried out a LANA ChIP-chip experiment using a customized, whole genome KSHV tiling array that we designed. Our results demonstrate for the first time that LANA is associated with essentially the entire latent KSHV genome in BCBL-1 cells with several highly enriched regions. This may suggest that LANA has additional roles as a structural element coordinating the binding, arrangement, and modulation of factors that interact with KSHV chromatin. This is consistent with the finding that
LANA function can be partially replaced by histone H1 (70). Interestingly, the LANA–episome interaction rapidly dissociates upon K-Rta-mediated reactivation such that at 4 h post-reactivation, LANA association with the KSHV episome is markedly reduced (supplemental Fig. 1). Because this study utilized samples derived from early time points, this phenotype is not likely due to encapsidation of the KSHV genome. A previous report had observed the rapid loss of LANA from the ORF50 region during reactivation (29); our results now expand this loss to the entire episome. We are currently investigating mechanisms of LANA dissociation from the KSHV genome. Importantly, our LANA ChIP-chip result complements recent reports that have provided comprehensive maps of active and repressive histone marks along the entire KSHV chromosome (8, 9), as well as a map of the binding sites across the KSHV genome of an associated cellular histone demethylase (10). These findings together with our LANA tiling array data may provide new insights into epigenetic regulation via viral proteins. Moreover, the results reported herein underscore the potential of relatively understudied methylation/demethylation events to influence the function of viral proteins.

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