The Epstein-Barr Virus BRRF1 Protein, Na, Induces Lytic Infection in a TRAF2- and p53-Dependent Manner

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The Epstein-Barr virus (EBV) BRRF1 lytic gene product (Na) is encoded within the same immediate-early region as the BZLF1 (Z) and BRLF1 (R) gene products, but its role during EBV infection has not been well defined. We previously showed that Na cooperates with the R protein to induce lytic gene expression in latently infected EBV-positive 293 cells, and in some EBV-negative cell lines it can activate the Z promoter in reporter gene assays. Here we show that overexpression of Na alone is sufficient to induce lytic gene expression in several different latently infected epithelial cell lines (Hone-Akata, CNE2-Akata, and AGS-Akata), while knockdown of endogenous Na expression reduces lytic gene expression. Consistent with its ability to interact with tumor necrosis factor receptor-associated factor 2 (TRAF2) in a yeast two-hybrid assay, we demonstrate that Na interacts with TRAF2 in cells. Furthermore, we show that TRAF2 is required for Na induction of lytic gene expression, that Na induces Jun N-terminal protein kinase (JNK) activation in a TRAF2-dependent manner, and that a JNK inhibitor abolishes the ability of Na to disrupt viral latency. Additionally, we show that Na and the tumor suppressor protein p53 cooperate to induce lytic gene expression in epithelial cells (including the C666-1 nasopharyngeal carcinoma cell line), although Na does not appear to affect p53 function. Together these data suggest that Na plays an important role in regulating the switch between latent and lytic infection in epithelial cells and that this effect requires both the TRAF2 and p53 cellular proteins.

Epstein-Barr virus (EBV) is a gammaherpesvirus that causes infectious mononucleosis and infects a large percentage of the human population (51, 66). EBV is associated with several epithelial cell cancers, such as nasopharyngeal carcinoma (NPC) and gastric carcinoma, and B-cell cancers, including Burkitt lymphoma and Hodgkin lymphoma (51, 66). EBV normally establishes a life-long latent infection in the memory B cells of the host but is reactivated periodically to the lytic form of infection. Lytic reactivation in B cells occurs following B-cell receptor engagement and/or plasma cell differentiation (40). Although EBV infection of normal oral epithelial cells commonly results in lytic infection (51), EBV-positive epithelial tumors are composed primarily of cells with the latent form of viral infection (37, 51).

The switch between latent and lytic infection is induced by the EBV immediate-early (IE) proteins BZLF1 (Z; also known as Zta, Zebra, or EB1) and BRLF1 (R) (13, 15, 37, 61, 65). Z and R serve as transcriptional activators and function to trans-activate one another’s promoters and to activate early lytic gene expression (3, 13, 14, 16, 19, 22, 29, 32, 35, 43, 50, 52, 61, 65). The cellular and viral proteins that regulate transcription of the Z and R gene products thus play a key role in determining if EBV infection is latent or lytic.

The Z promoter (Zp) is regulated by a number of cellular factors. In latently infected cells, several repressors have been shown to inhibit Zp activity, including the ZEB1, ZEB2, and MEF2D cellular proteins (21, 30, 39, 45, 57). During lytic infection, a variety of cellular transcription factors, including CREB, ATF-1, ATF-2, c-Jun, and XBP-1, bind to the Zp ZII motif (a CRE binding site), leading to Zp activation (1, 6, 44, 47, 58, 63). Phosphorylation of the c-Jun transcription factor by the c-Jun N-terminal protein kinase (JNK) greatly enhances its activity (17), and activation of JNK activity has been shown to be important for lytic EBV induction mediated by a variety of different agents (9, 23, 24).

The EBV BRRF1 open reading frame, which encodes the Na protein, is contained within the same locus as the Z and R IE genes. Our lab previously showed that Na enhances R-mediated disruption of viral latency in 293 cells that are stably infected with an EBV mutant that does not express either R or Na (293RKO cells) (33). Likewise, the murine gammaherpesvirus 68 (MHV68) Na homologue (ORF49) has been reported to enhance the ability of the R homologue (Rta) to induce lytic viral gene expression (41). Furthermore, we showed that Na activates a Zp-driven reporter gene in EBV-negative HeLa cells through the Zp ZII motif (a c-Jun binding site), and that Na induces c-Jun phosphorylation and c-Jun transcriptional activity (33). However, the mechanism by which Na induces c-Jun activation and the role that Na plays during the switch between latent and lytic infection in epithelial cell lines remains unclear.

The Na protein was previously demonstrated to interact with the cellular protein tumor necrosis factor (TNF) receptor-associated factor 2 (TRAF2) in a yeast two-hybrid analysis (10), although the possible functions(s) of this interaction and whether it occurs in vivo remain unknown. TRAF2 is a signal transducer for the TNF receptor superfamily members, including TNFR1/2, CD40, and IRE1 (4, 8). TRAF2 signaling me-
diates several major signaling pathways following receptor activation, including those of mitogen-activated protein kinase, classical and noncanonical NF-κB, and JNK (4, 8). In EBV-infected B cells, TRAF2 interacts with latent membrane protein 1 (LMP1), and this interaction is thought to be important for LMP1 activation of NF-κB (18, 34, 56). Although the role of TRAF2 during latent infection of B cells has been well studied, the role of TRAF2 during EBV lytic infection remains unclear.

In this study, we demonstrate that overexpression of the Na protein induces lytic gene expression in several different latently infected epithelial cell lines and that, conversely, knockdown of endogenous Na expression inhibits lytic gene expression. We confirm that Na interacts with TRAF2 in vivo and show that TRAF2 expression is required for Na disruption of latency. We also demonstrate that Na induces JNK phosphorylation in a TRAF2-dependent manner and that JNK activation is required for Na-induced lytic gene expression. Furthermore, we demonstrate that the ability of Na to disrupt viral latency also requires the p53 cellular protein, consistent with a recent report showing that p53 is required for histone deacetylase (HDAC) inhibitor (HDACi)-induced EBV lytic infection in a latently infected NPC cell line (12). Together these results suggest that Na plays an important role in promoting EBV lytic reactivation in epithelial cells, and the results also reveal unexpected roles for TRAF2, as well as for p53, in Na-induced lytic reactivation.

**MATERIALS AND METHODS**

**EBV cell lines.** Hone-Akata (a gift from Lawrence Young) and CNE2-Akata (a gift from K. W. Lo at The Chinese University of Hong Kong [received via Diane Hayward]) are NPC epithelial cell lines superinfected with the Akata strain of EBV (25, 46), and they were maintained in RPMI medium supplemented with 10% fetal bovine serum (FBS), penicillin-streptomycin, and G418 (400 μg/ml). CNE2-Akata cells contain an arginine-to-threonine mutation at codon 280 of p53 (20) as well as wild-type p53 (11). C666-1 cells, an NPC line, were maintained in RPMI medium supplemented with 10% FBS and penicillin-streptomycin and were grown on plates treated with fibronectin (Sigma). C666-1 cells (a gift from Dolly Huang) contain a deletion at codon 249 of p53 (64). 293 cells (a gift from Diane Hayward) are NPC epithelial cell lines superinfected with the Akata strain of EBV and were maintained in F-12 medium supplemented with 10% FBS, penicillin-streptomycin, and G418 (400 μg/ml). Raji cells (ATCC) were maintained in Dulbecco's modified Eagle's medium supplemented with 10% FBS and penicillin-streptomycin. AGS gastric carcinoma cells were maintained in F-12 medium supplemented with 10% FBS and penicillin-streptomycin. AGS-Akata cells are gastric carcinoma cells superinfected with the Akata strain of EBV and were maintained in F-12 medium supplemented with 10% FBS, penicillin-streptomycin, and G418 (400 μg/ml). Raji cells (ATCC) were maintained in RPMI medium supplemented with 10% FBS and penicillin-streptomycin.

**Plasmids.** Plasmid DNA was purified using Qiagen maxi-prep columns according to the manufacturer's protocol. pSG5 and pCDNA3.1 were obtained from Stratagene and Invitrogen, respectively. pSG5-FLAGNa was generated by PCR amplification of the Na open reading frame from EBV B95.8 viral DNA with 10% FBS, penicillin-streptomycin, and G418 (400 μg/ml). Raji cells (ATCC) were maintained in RPMI medium supplemented with 10% FBS and penicillin-streptomycin.

**Immunoblot analysis.** Immunoblotting was performed as previously described (33). CNE2-Akata cells were transfected with control or FLAG-Na expression vectors. Medium was changed 2 days posttransfection, and supernatant was harvested and filtered through a 0.8-μm-pore-size filter 3 days posttransfection. Raji cells (2 × 10^6 cells) were infected with various amounts of virus and incubated at 37°C, Phorbol-12-myristate-13-acetate (TPA; 20 ng/ml) and sodium butyrate (3 mM, final concentration) were added 24 h after infection. Green fluorescent protein (GFP)-positive Raji cells were counted 48 h postinfection to determine viral titer.

**Immunoprecipitation analysis.** 293 cells were transfected with the indicated expression plasmids, and cell lysates were harvested in NP-40 lysis buffer (50 mM Tris [pH 7.4], 150 mM NaCl, 0.5% NP-40, 0.75% IGEPAL) containing protease inhibitor cocktail (Roche) and quantified in a SUMO protein assay (Bio-Rad). Equivalent amounts of protein were separated in sodium dodecyl sulfate–10% polyacrylamide gels and transferred to membranes. Membranes were blocked in phosphate-buffered saline (PBS) containing 5% milk and 0.1% Tween 20 solution and incubated with primary antibodies. Immunoblots were probed with the following antibodies: anti-FLAG (1:2,000; Sigma), anti-R (1:250; Argene), anti-Z (1:250; BZ-1; Santa Cruz Biotechnology), anti-EAD (1:250; BMRF1; Vector), anti-β1 (1:500; Sigma), anti-α (1:500; Santa Cruz Biotechnology), anti-T (1:500; Santa Cruz Biotechnology), anti-phospho-JNK (1:4,000; Cell Signaling), anti-JNK (1:1,000; Cell Signaling), anti-p53 (1:250; Santa Cruz Biotechnology), anti-IκBα (1:250; Santa Cruz Biotechnology), anti-p52 (1:500; Milipore), anti-α (1:500; a kind gift from Pierre Charton; 55), anti-Tubulin (1:2,000; Sigma), and anti-lamin A/C (1:500; Santa Cruz Biotechnology).
the supernatant was saved as the cytosolic fraction. The pellet was washed in PBS–0.1%NP-40 and pelleted, and the supernatant was discarded. The remaining pellet was then resuspended in sample buffer and saved as the nuclear fraction.

**Fluorescent microscopy.** Hone-Akata cells plated on glass coverslips were transfected with various yellow fluorescent protein (YFP) expression plasmids and fixed 24 h after transfection with 0.4% paraformaldehyde in PBS. Cell nuclei were stained with Hoechst 33342 (Sigma), and coverslips were mounted onto slides by using Vectashield hard set. Images were visualized using a Nikon A1R confocal microscope.

**CAT reporter assay.** CAT assays were performed as previously described (27). CNE2-Akata cells were transfected with the indicated expression vectors and harvested in 0.25 M Tris (pH 7.5) and SUMO buffers 48 h after transfection. Tris buffer lysates were incubated with acetyl coenzyme A (CoA) and [14C]chloramphenicol, and acetyltransferase activity was determined following thin-layer chromatography. Activity was quantified on a Storm 840 PhosphorImager (Molecular Dynamics). Western blot analysis was performed on the SUMO buffer lysates to determine protein expression.

**RESULTS**

**Na induces lytic gene expression in EBV-infected epithelial cells.** Although the Na protein was previously shown to induce c-Jun phosphorylation and activate the Z promoter in EBV-negative HeLa cells (33), overexpression of Na has not previously been shown to induce lytic EBV gene expression in the context of the intact viral genome. To examine this further, we transfected Hone-Akata cells (a nasopharyngeal carcinoma line superinfected with the Akata strain of EBV) with two different Na expression vectors (FLAG tagged), or the appropriate control vectors, and examined the expression levels of various lytic viral proteins released into the supernatant. As shown in Fig. 1A, overexpression of the Na protein clearly induced expression of the EBV lytic proteins BMRF1 and Z. These results indicate that activation of Na expression is sufficient to disrupt viral latency in the Hone-Akata cell line.

To ensure Na disruption of viral latency was not specific to the Hone-Akata line, the effect of Na overexpression was also examined in two additional EBV superinfected epithelial cell lines, CNE2-Akata (an NPC line) (Fig. 1B) and AGS-Akata (a gastric carcinoma cell line) (Fig. 1C). Similar to the Na effect in Hone-Akata cells, Na also induced expression of lytic EBV proteins (BMRF1, Z, and R) in CNE2-Akata and AGS-Akata cells. Together, these results indicate that overexpression of Na alone is sufficient to induce lytic gene expression in at least a subset of EBV-infected epithelial cell lines.

To determine if Na expression also results in enhanced lytic viral replication, CNE2-Akata cells were transfected with control or Na expression vectors, and the amount (titer) of infectious viral particles released into the supernatant was determined.
the ability of the lysis-inducing agents TPA and sodium butyrate (NaBut) to activate lytic protein expression (Fig. 2A). These results demonstrate that Na derived from the endogenous viral genome contributes to both constitutive and chemically induced lytic viral protein expression in Hone-Akata cells.

To determine if the level of Na protein obtained following Na plasmid transfection was physiologic with regard to the level of endogenous Na protein expressed during normal lytic infection, we transfected CNE2-Akata cells with different amounts of the FLAG-Na construct and used an anti-Na antibody to compare the level of transfected Na versus that expressed in TPA-sodium butyrate-treated MutuI cells (Fig. 2B). The amount of Na expressed in the lytically induced MutuI Burkitt lymphoma cells was comparable to the amount of expressed in the CNE2-Akata cells transfected with 0.2 μg Na expression vector (in a 12-well plate). Since most of the transfection experiments presented in this study used 0.2 μg (or less) of the Na vector, we conclude that the amount of Na expressed is physiologic.

Na induction of JNK activity is required for Na disruption of viral latency. Our lab previously showed that Na induces c-Jun phosphorylation and enhances c-Jun transactivator function (33). Furthermore, expression of the Kaposi’s sarcoma-associated herpesvirus (KSHV) homologue of Na, Orf49, in 293 and CV-1 cells induces phosphorylation of both JNK and c-Jun (26). To determine if Na likewise induces JNK phosphorylation, Hone-Akata cells were transfected with Na or control expression plasmids, and the level of JNK activation was examined by immunoblot analysis using antibodies directed against phosphorylated (p-JNK) versus total JNK. As shown in Fig. 3A, p-JNK levels increased in the presence of Na, while the total JNK level was not affected. Similar results were obtained in CNE2-Akata cells (Fig. 3B). These results suggest that Na, like the KSHV Orf49 protein, induces c-Jun activation by promoting JNK phosphorylation.

To determine if JNK activation is required for Na-mediated disruption of viral latency, CNE2-Akata cells were transfected with control or Na expression plasmids in the presence or absence of the JNK inhibitor SP600125 (10 μM). The ability of Na to induce expression of the lytic EBV proteins BMRF1, Z, and R was reduced in the presence of the JNK inhibitor (Fig. 3B). These results suggest that Na activation of JNK plays an important role in Na-induced disruption of viral latency.

Na interacts with the cellular protein TRAF2 in vivo. Since Na has been reported to interact directly with the cellular protein TRAF2 by yeast two-hybrid analysis (10) and TRAF2 is involved in activating JNK during certain forms of signaling, these observations suggest that the ability of Na to activate JNK may involve a direct interaction with the TRAF2 protein. Nevertheless, the TRAF2 protein is not thought to enter the nucleus in most cell types, while Na has been reported to be a nuclear protein (55). To determine whether the Na and TRAF2 proteins can interact in vivo, 293 cells were transfected with FLAG-Na and HA-TRAF2 expression plasmids, immunoprecipitation assays were performed using antibodies directed against the FLAG or HA tags, and then immunoblot analyses were conducted to detect coprecipitated FLAG-Na and HA-TRAF2 proteins. As shown in Fig. 4, an interaction between Na and TRAF2 was detected following immunopre-

**FIG. 2.** Knockdown of Na inhibits lytic protein expression. (A) CNE-2 Akata cells were transfected with siRNAs targeting control or Na sequences. Cells were then treated with either dimethyl sulfoxide (DMSO) or TPA (20 ng/ml) plus sodium butyrate (3 mM; NaBut). Cell lysates were harvested 48 h after induction, and immunoblot analysis was performed using antibodies against BMRF1, Na, R, Z, and β-actin. (B) CNE2-Akata cells were transfected with various amounts of the FLAG-Na vector as indicated (in a 12-well plate). At 48 h posttransfection, an immunoblot assay was performed (using an anti-Na antibody) to compare the level of Na expressed in the transfected CNE-Akata cells versus that expressed from the endogenous viral genome in MutuI cells treated with TPA/NaBut. β-Actin levels were examined as a protein loading control.
cipitation with either FLAG or HA antibodies. These data demonstrate that Na and TRAF2 can interact when expressed in a cellular environment.

Na is present in both the nucleus and cytoplasm. The finding that Na interacts with TRAF2 in cells suggests that a portion of Na may be located within the cytoplasm, similar to what has been described for the KSHV Orf49 homologue (26).

To investigate the subcellular localization of both Na and TRAF2, we constructed plasmids in which the Na or TRAF2 proteins were fused to a C-terminal YFP tag and examined the cellular localization of each protein by confocal fluorescent microscopy in transfected Hone-Akata cells. As shown in Fig. 5A, the Na-YFP protein, similar to the Orf49 protein, can be detected both in the nucleus and the cytoplasm of cells. TRAF2-YFP localized primarily in the cytoplasm but was also detected in the nucleus of some Hone-Akata cells. Although TRAF2 is predominantly localized within the cell cytoplasm, other reports have also detected TRAF2 in the nucleus of the cell (38).

FIG. 3. Na induction of JNK is required for Na to activate lytic protein expression. (A) Hone-Akata cells were transfected with control vector or FLAG-Na expression plasmids, and immunoblot analysis was performed using antibodies against phosphorylated JNK (p-JNK), total JNK, FLAG (to detect transfected Na), and β-actin. (B) CNE2-Akata cells were transfected with control vector or FLAG-Na expression plasmids in the presence or absence of the JNK inhibitor SP600125 (10 μM). Cell lysates were harvested, and immunoblot analysis was performed to examine p-JNK, total JNK, BMRF1, Z, R, FLAG (Na), and β-actin expression levels.

FIG. 4. Na interacts with TRAF2. 293 cells were cotransfected with the indicated combinations of vector control, FLAG-Na, or HA-TRAF2. Cell lysates were harvested in NP-40 buffer 48 h posttransfection. Immunoprecipitation analysis was performed using antibodies against the FLAG or HA tags of transfected FLAG-Na and HA-TRAF2. Immunoblot analysis was then performed using antibodies against FLAG or HA to detect the immunoprecipitated proteins. Direct load samples were also used for immunoblot analysis as a control.
viral latency, Hone-Akata cells were pretreated with either control siRNA or TRAF2 siRNA and then transfected with control or Na expression vectors. As expected, Na induced BMRF1 and Z expression in cells transfected with control siRNA (Fig. 6A). However, following knockdown of TRAF2 expression, Na lost the ability to induce lytic viral protein expression. In addition, the low level of constitutive lytic protein expression in Hone-Akata cells was also reduced by loss of TRAF2 expression. Similar results were obtained when lentivirus vectors expressing control or short hairpin sequences against TRAF2 were used to reduce TRAF2 expression in CNE2-Akata cells (Fig. 6B). Together these data indicate that TRAF2 is required for Na-mediated disruption of EBV latency and suggest that the interaction between Na and TRAF2 is important for induction of lytic protein expression.

**TRAF2 is required for Na-induced JNK activation.** To determine if TRAF2 is also involved in Na-mediated JNK activation, Hone-Akata cells were transduced with lentiviruses expressing TRAF2 shRNAs or control shRNAs, stably selected, and then transfected with control or Na expression plasmids. Na induced JNK phosphorylation as expected in cells expressing the control shRNAs (Fig. 6C). However, knockdown of TRAF2 expression greatly inhibited the ability of Na to induce JNK phosphorylation, as well as decreasing the constitutive level of p-JNK expression in cells transfected with control vector. As expected, TRAF2 knockdown also reduced the ability of Na to increase lytic BMRF1 protein expression. Together, these data suggest that TRAF2 is required both for the ability of Na to induce JNK phosphorylation and for its ability to initiate lytic viral protein expression.

**Na induces constitutive p100/p52 processing but does not affect TNF-α signaling.** Since TRAF2 is involved in multiple different aspects of TNF-α-induced signaling, including activation of the classical NF-κB pathways, as well as JNK, and inhibition of the alternative NF-κB pathway, we also examined whether Na affects TNF-α-induced signaling. CNE2-Akata cells were transfected with control or Na expression vectors and treated with TNF-α (10 ng/ml) for 20 or 60 min prior to harvesting (Fig. 7A). Cell lysates were examined for IκBα, which becomes phosphorylated and rapidly degraded through classical NF-κB activation, as well as for the level of p100/p52 processing (which is increased during activation of the alternative NF-κB pathway) (Fig. 7B). Overexpression of Na did not affect the ability of TNF-α to decrease expression of IκBα (Fig. 7A) or to increase the level of JNK activation (data not shown). Interestingly, overexpression of Na consistently increased the level of constitutive p100/p52 processing (Fig. 7B). Since the CNE2-Akata cells have some level of constitutive p100/p52 processing even in the absence of Na, the effects of Na on p100 processing were also examined in the EBV-negative AGS cell line, which has a lower baseline level of constitutive p100/p52 processing. Again, overexpression of Na led to an increase in p52 accumulation in the nucleus, as did transfected LMP1 protein (which is known to induce p100/p52 processing [49] and served as a positive control). These results suggest Na does not affect TNF-α signaling through the classical NF-κB pathway or the JNK pathway, but that it increases the level of p100/p52 processing in the alternative NF-κB pathway, even in the absence of TNF-α signaling.

**Na does not affect p53 function.** Since a recent study suggested that p53 is important for HDACi-mediated lytic reac-
tivation in EBV-positive epithelial cell lines (12), and our results here indicate that Na is also important for viral reactivation by HDAC inhibitors (Fig. 2), we hypothesized that Na may enhance p53 transcriptional activity. To examine this, CNE2-Akata cells were transfected with control or FLAG-Na expression plasmids and treated with TNF-α (10 ng/ml) for 20 or 60 min prior to harvesting. (A) Immunoblot analysis was performed using antibodies against IκBα, FLAG (Na), and β-actin. (B) Immunoblot analysis was performed using antibodies against p52, FLAG (Na), and β-actin. (C) AGS cells were transfected with control, LMP1, or FLAG-Na expression plasmids and harvested 48 h posttransfection. Biochemical fractionation was used to obtain nuclear (Nuc) and cytosolic (Cyto) extracts. Immunoblot analysis was performed to examine p100 processing by using a p52 antibody. Tubulin and lamin A/C antibodies were used as controls for fractionation.

FIG. 6. TRAF2 is required for Na-mediated induction of lytic protein expression. (A) Hone-Akata cells were transfected with control or TRAF2 siRNAs. At 48 h after transfection, the cells were retransfected with the control or TRAF2 siRNAs along with control or FLAG-Na expression plasmids. Immunoblot analysis was performed 2 days later using antibodies against BMRF1 and Z to examine lytic protein expression, or with TRAF2 and FLAG to detect TRAF2 and Na, respectively. β-Actin was used as a loading control. (B) CNE2-Akata cells were transfected with lentiviruses expressing shRNAs against either a control sequence or TRAF2. At 48 h posttransfection, cells were retransfected with the shRNA vectors along with either control or FLAG-Na expression plasmids. Cell lysates were harvested, and immunoblot analyses were performed using antibodies against TRAF2, BMRF1, FLAG (Na), R, Z, and β-actin. The data shown represent samples run on the same gel; irrelevant lanes were cropped from the figure. (C) Stable selected Hone-Akata cell lines transduced with lentiviruses expressing shRNAs against TRAF2, BMRF1, FLAG (Na), R, Z, and β-actin. The data shown represent samples run on the same gel; irrelevant lanes were cropped from the figure. Immunoblot analysis was performed using antibodies against phosphorylated JNK (p-JNK), total JNK, BMRF1, TRAF2, FLAG (Na), and β-actin.
enhanced the effect of the FLAG-Na expression vector. These altered residue 248 (which inhibits p53 DNA binding) and residue 175 (which affects p53 conformation) nor a p53 mutant with an expression, neither a p53 mutant with an altered residue 175 p53-wt construct cooperated with Na to induce lytic protein expression in this cell type, but that the combination of p53 and Na expression together was clearly more effective than either protein alone. Furthermore, in the tightly latent C666-1 NPC cell line (the only NPC cell line which has remained persistently EBV positive in culture without selection), we found that only the combination of both p53 and Na was sufficient to induce lytic EBV gene expression (Fig. 2).

Since p53 is mutated in many carcinoma cell lines, we next tested the ability of two different p53 mutant constructs to synergistically induce lytic protein expression with cotransfected Na in CNE2-Akata cells. As shown in Fig. 9C, while the p53-wt construct cooperated with Na to induce lytic protein expression, neither a p53 mutant with an altered residue 175 (which affects p53 conformation) nor a p53 mutant with an altered residue 248 (which inhibits p53 DNA binding) enhanced the effect of the FLAG-Na expression vector. These results indicate that wild-type p53 function is important for enhancing Na-induced lytic protein expression.

**p53 is required for Na-mediated disruption of viral latency.** Finally, to examine whether endogenous p53 expression is required for the ability of Na to induce lytic viral protein expression, CNE2-Akata cells were transfected with lentivirus vectors expressing control shRNA or p53-directed shRNA (shp53) and then transfected with control or Na expression vectors. As shown in Fig. 10A, knockdown of endogenous p53 expression greatly reduced the ability of transfected Na to induce lytic EBV protein expression. In addition, knockdown of endogenous p53 expression reduced the level of constitutive lytic EBV protein expression found in CNE2-Akata cells (Fig. 10A) and reduced the ability of transfected R protein to activate lytic EBV protein expression (Fig. 10B). In contrast, knockdown of p53 expression did not affect the ability of transfected Z protein to induce lytic gene expression (Fig. 10B). These results suggest that endogenous p53 activity is required for the ability of both Na and R (but not Z) to induce lytic EBV gene expression, and the results furthermore suggest that a key aspect of the p53 effect may be to help the Z/R proteins to activate Zp in the context of the intact viral genome.

**DISCUSSION**

The switch between latent and lytic EBV infection is known to be activated by expression of either the BZLF1 (Z) and/or BRLF1 (R) IE proteins (13, 15, 37, 61, 65). Z and R activate one another’s promoters (Zp and Rp) and cooperatively activate the early lytic viral promoters (3, 13, 14, 16, 19, 22, 29, 32, 35, 43, 50, 52, 65). Although the early lytic viral protein Na (the BRRF1 gene product) was previously reported to activate the Z promoter in HeLa cells and to cooperate with R to activate lytic gene expression in EBV-infected 293 cells, overexpression of Na by itself was not found to activate lytic gene transcription in 293 cells (33). Here we show that overexpression of Na is sufficient to induce lytic gene expression in a number of EBV-infected epithelial cell lines (Fig. 1) and that, conversely, knockdown of endogenous Na expression inhibits low-level constitutive lytic protein expression (Fig. 2).
We have demonstrated that Na interacts directly with TRAF2 in cells (Fig. 4) and that TRAF2 expression is required for the lysis-inducing effect of Na (Fig. 6). Furthermore, similar to a recent report indicating that HDAC inhibitors require wild-type p53 to activate lytic viral protein expression in NPC cell lines (12), we likewise found that wild-type p53 (but not mutant p53) enhanced both Na-induced as well as constitutive lytic viral protein expression in EBV-infected epithelial cell lines (Fig. 9 and 10). Together these results suggest that Na plays an important regulatory role in promoting the switch between latent and lytic EBV infection in epithelial cells, and cellular factors that activate Na expression may in fact be sufficient to induce the whole lytic viral protein cascade expression in some cell lines.

Our results suggest that the ability of Na to induce lytic EBV protein expression is at least partially mediated through activation of the JNK pathway. We have shown that Na, like the KSHV homologue Orf49, induces JNK activation in cells, and that furthermore JNK activation is required for the ability of Na to activate lytic viral protein expression (Fig. 3). These results are consistent with our previous findings that Na induces c-Jun phosphorylation and activates c-Jun transcriptional function (33). Since c-Jun is a known activator of the Z promoter (binding to the Zp ZII motif), our results support a model in which Na expression leads to lytic EBV expression via its ability to activate the Z promoter in certain cell lines.

Although the Na and TRAF2 proteins were previously shown by yeast two-hybrid assays to directly interact (10), the potential significance of this interaction has been unclear, particularly since TRAF2 is thought to be a cytoplasmic protein, whereas Na has been reported to be a nuclear protein (55). However, since TRAF2 is known to be essential for activation of the JNK pathway following TNFR activation (42) and overexpression of TRAF2 alone has been reported to activate JNK in certain cell lines (31), these previous reports suggested the possibility that one or more Na effects may be mediated through TRAF2. We have demonstrated that Na interacts with TRAF2 in cells (Fig. 4), and we have shown that Na (similar to KSHV Orf49) is located within both the nucleus and cytoplasm (Fig. 5). Most importantly, we have demonstrated that constitutive cellular TRAF2 expression is required not only for Na-induced JNK activation but also for Na-induced lytic viral protein expression (Fig. 6).

Together, these results strongly suggest that the Na effect is at least partially mediated through its direct interaction with TRAF2 and that this Na-TRAF2 interaction mediates JNK activation. Nevertheless, the proof of this hypothesis will ultimately require the identification of a Na mutant that is specifically deficient in the ability to interact with TRAF2. Unfortunately, to date we have found that Na mutants unable to interact with TRAF2 are also unstable (data not shown). Additionally, due to the limitations of interaction studies using overexpressed proteins, the interaction between endogenous Na and TRAF2 should be confirmed. Also unclear at present is whether Na affects any other TRAF2 functions, for example, the ability of TRAF2 to regulate NF-κB signaling in response to the TNF-α ligand. While we did not observe any affect of Na on 1κBα expression following TNF-α treatment of CNE2-Akata cells (Fig. 7A), suggesting that Na does not alter the canonical NF-κB signaling pathway, we routinely observed an
increase in the level of constitutive p100/p52 processing in cells expressing Na (Fig. 7B and C). Since TRAF2 knockout mouse embryonic fibroblast cells have been reported to have an increased level of constitutive p100/p52 processing (which is thought to reflect the loss of TRAF2-mediated degradation of NIK) (28, 62), the effect of Na on p100/p52 processing may reflect its ability to inhibit this particular TRAF2 function. Although the mechanism(s) by which Na induces p100/p52 processing remains to be explored, since this Na effect was observed in both EBV-negative as well as EBV-positive cell types, it does not seem to require any other EBV-encoded proteins, such as LMP1. Whether Na-induced p100/p52 processing enhances or inhibits its ability to induce lytic viral gene expression will be an important focus for future studies.

In this study we have also demonstrated that p53 contributes to the ability of Na to induce lytic EBV protein expression. We show that coexpression of wild-type p53, but not mutant p53, enhances the ability of Na to induce lytic EBV protein expression in CNE2-Akata and Hone-Akata cells (Fig. 9A and C) and that Na cannot induce lytic gene expression in the tightly latent C666-1 NPC cell line unless it is coexpressed with a p53 expression vector (Fig. 9B). Furthermore, we have shown that knockdown of endogenous p53 expression in CNE2-Akata cells diminishes the ability of both Na and R to induce lytic gene expression (Fig. 10) while having no effect on the ability of transfected Z protein to disrupt viral latency (Fig. 10B). These results suggest that the effect of p53 is mediated at the level of Z promoter activation (and thereby effectively bypassed when Z is expressed under the control of a strong heterologous promoter). Similar to our results here, loss of p53 expression was recently shown to inhibit HDACi-induced lytic reactivation (12), while overexpression of p53 was reported to enhance early EBV lytic infection (54). Our finding that endogenous p53 expression appears to cooperate with Na to induce lytic EBV expression in CNE2-Akata and Hone-Akata cells, but not in C666-1 cells, is possibly explained by the fact that C666-1 cells contain only a mutant form of p53 (64), whereas the CNE2-Akata and Hone-Akata lines, although originally described as NPC lines, are at least partially derived from HeLa cells (11) and thus may express some wild-type p53 as well as mutant p53. Alternatively, the p53 mutant found in CNE2-Akata and Hone-Akata cells (p53 Thr280) retains the ability to cooperate with Na to disrupt viral latency.

Together, the results presented here suggest a model in which Na interacts with TRAF2 to activate JNK signaling, leading to transactivation of Zp, and wild-type p53 enhances Na-mediated activation of Zp (Fig. 11). Although we demonstrated that wild-type p53 enhances Na induction of lytic protein expression (Fig. 9), the mechanism by which p53 and Na synergistically activate lytic protein expression remains to be determined. Since we did not find that Na activates p53 transcriptional function (Fig. 8) or consistently affects endogenous p53 expression, we speculate that Na and p53 enhance BZLF1 transcription in the context of the intact viral genome through independent mechanisms (Fig. 11). To date, we have been unable to demonstrate this synergistic activation of the BZLF1 promoter by Na and p53 using Zp-driven reporter gene assays (data not shown). This result suggests that the synergistic effect

FIG. 10. p53 is required for Na activation of lytic protein expression. CNE2-Akata cells were transfected with lentivirus vectors expressing shRNAs against either control or p53 sequences. At 48 h posttransfection, the cells were cotransfected with the same shRNAs and control, FLAG-Na (A), or Z or R (B) expression plasmids. Immunoblot analysis was performed 2 days later using antibodies against BMRF1, Z, and R to examine EBV lytic gene expression and using antibody against FLAG or p53 to detect transfected Na and endogenous p53 expression. β-Actin was included as an internal loading control.

FIG. 11. Hypothesized model for Na transactivation of the Z promoter. A potential model for Na-induced lytic protein expression is shown. Na interacts with TRAF2 to activate the JNK signaling cascade, leading to phosphorylation of c-Jun and activation of Zp. p53 independently induces activation of Zp and has a synergistic effect when coexpressed with Na. The mechanism by which p53 enhances Zp activity is currently unknown.
moting lytic viral protein expression during normal EBV infection suggests that Na has an important and essential role in promoting lytic protein expression from the endogenous viral genome in Hone-Akata cells not only decreased chemically induced lytic protein expression but also activation of the Z protein in all cell lines tested. Thus, activation of the Z and/or R proteins induce lytic protein expression more efficiently than the Na protein in all cell lines tested. Furthermore, the Z and/or R promoters by cellular transcription factors likely plays a more important role than activation of the Na promoter in promoting lytic infection in virally infected epithelial cells. Since the EBV-infected epithelial cell lines that respond to the lytic-inducing effect of Na alone can constitutively express lytic EBV proteins at a very low level, it remains possible that an important aspect of the Na lytic induction effect is to enhance R-mediated and/or Z-mediated transcriptional effects. In any event, our finding that knockdown of Na expression derived from an EBV early promoter. EMBO J. 5:3243–3249.


Expression of LMP1 in epithelial cells leads to the activation of a select subset of NF-kappa B/Rel family proteins. J. Virol. 69:4572–4576.
56. Seguin-R *