ΔNp63α promotes Epstein-Barr virus latency in undifferentiated epithelial cells

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Abstract

Epstein-Barr virus (EBV) is a human herpesvirus that causes infectious mononucleosis and contributes to both B-cell and epithelial-cell malignancies. EBV-infected epithelial cell tumors, including nasopharyngeal carcinoma (NPC), are largely composed of latently infected cells, but the mechanism(s) maintaining viral latency are poorly understood. Expression of the EBV BZLF1 (Z) and BRLF1 (R) encoded immediate-early (IE) proteins induces lytic infection, and these IE proteins activate each other’s promoters. ΔNp63α (a p53 family member) is required for proliferation and survival of basal epithelial cells and is over-expressed in NPC tumors. Here we show that ΔNp63α promotes EBV latency by inhibiting activation of the BZLF1 IE promoter (Zp). Furthermore, we find that another p63 gene splice variant, TAp63α, which is expressed in some Burkitt and diffuse large B cell lymphomas, also represses EBV lytic reactivation. We demonstrate that ΔNp63α inhibits the Z promoter indirectly by preventing the ability of other transcription factors, including the viral IE R protein and the cellular KLF4 protein, to activate Zp. Mechanistically, we show that ΔNp63α promotes viral latency in undifferentiated epithelial cells both by enhancing expression of a known Zp repressor protein, c-myc, and by decreasing cellular p38 kinase activity. Furthermore, we find that the ability of cis-platinum chemotherapy to degrade ΔNp63α contributes to the lytic-inducing effect of this agent in EBV-infected epithelial cells. Together these findings demonstrate that the loss of ΔNp63α expression, in conjunction with enhanced expression of differentiation-dependent transcription factors such as BLIMP1 and KLF4, induces lytic EBV reactivation during normal epithelial cell differentiation. Conversely, expression of ΔNp63α in undifferentiated nasopharyngeal carcinoma cells and TAp63α in Burkitt lymphoma promotes EBV latency in these malignancies.

Author summary

Epstein-Barr virus (EBV) is an important cause of both epithelial cell and B cell human cancers. EBV-infected tumors have predominantly latent viral infection, allowing them to
escape the cell killing that occurs during lytic viral infection. EBV is highly lytic in normal
differentiated oral epithelial cells. Thus, an important question is how the virus maintains
the latent form of viral infection in EBV-associated epithelial cell tumors such as undifferen-
tiated nasopharyngeal carcinoma (NPC). This study demonstrates that the cellular tran-
scription factor ΔNp63α, which is specifically expressed in undifferentiated basal
epithelial cells and is over-expressed in NPC tumors, maintains EBV latency by inhibiting
the activity of the viral immediate-early (IE) promoter (Zp) that drives expression of the
BZLF1 IE protein. A related splice variant, TAp63α, found in some EBV+ lymphomas,
has a similar inhibitory effect. Our findings reveal that ΔNp63α and TAp63α contribute
to EBV latency in both epithelial and B cell tumors. Furthermore, since differentiation
results in loss of ΔNp63α expression, our results help to explain why lytic EBV reactiva-
tion is promoted by epithelial cell differentiation.

Introduction
Epstein-Barr virus (EBV) is a gamma-herpesvirus that infects over 90% of the human popula-
tion by adulthood and causes the clinical syndrome, infectious mononucleosis. EBV primarily
infects B cells and epithelial cells and is associated with both B-cell and epithelial-cell malign-
nancies, including Burkitt lymphoma (BL), diffuse large B cell lymphoma, Hodgkin lym-
phoma, gastric carcinoma, and nasopharyngeal carcinoma (NPC) [1]. Like all herpesviruses,
EBV can infect host cells in either latent or lytic forms and persists in the host for life. In
latently infected cells, the virus expresses only a small subset of viral genes and replicates once
per cell cycle using the host cell DNA polymerase. In lytically infected cells, the full viral gene
complement is expressed, the virus is replicated using the virally-encoded DNA polymerase
and infectious virions are produced. The major site of persistent latent EBV infection in
humans is memory B cells, although the lytic form of infection can be induced by B-cell recep-
tor stimulation, plasma cell differentiation or various other stimuli in B cells [2–6].

In contrast, normal oropharyngeal epithelial cells support the lytic form of EBV infection,
and it remains uncertain whether latent EBV infection of epithelial cells normally occurs in
humans [7–9]. Lytically-infected epithelial cells are thought to be the major source of infec-
tious EBV particles in saliva [10] and thus are essential for the spread of the virus from host to
host. EBV infection of non-transformed tongue epithelial cells causes the clinical syndrome,
oral hairy leukoplakia (OHL), in immunocompromised patients [7,11]. Analysis of OHL
lesions has revealed that EBV infection is confined to the more differentiated epithelial cell lay-
ers, and that the infection is completely lytic without evidence of concomitant latent infection
[7,11]. Nevertheless, the EBV-associated epithelial tumor, undifferentiated nasopharyngeal
carcinoma, contains largely latent EBV infection, and the ability of the virus to maintain
latency in the tumor cells is likely required for NPC development. Thus, understanding how
EBV stays latent in a cell type where it is usually lytic is important for understanding NPC.

Lytic EBV infection is initiated by expression of the two EBV immediate-early (IE) proteins,
BZLF1 (Z) and BRLF1 (R) [12–14]. The Z and R IE genes are driven by the Z and R promotors,
respectively, and activation (or repression) of these promoters by cellular transcription factors
serves as the major control point in determining if EBV infection is latent versus lytic in cells
[15–22]. Once expressed, the Z and R proteins function as viral transcription factors that acti-
vate each other’s promotors, as well as their own promotors, in a positive feedback loop
[12,23–25]. Z and R then synergistically activate early lytic viral gene promotors to induce
early gene expression and lytic viral DNA replication.
Z is a part of the bZip protein family and binds to AP-1-like sites called Z-responsive elements (ZREs), while R has no cellular homolog and binds to R-responsive elements (RREs) [12,26,27]. R can also activate some promoters, including Zp, indirectly through non-DNA binding mechanisms that are still not totally understood [25,28]. Many ZREs are preferentially bound by Z in the methylated form, helping Z to induce lytic viral reactivation even when the viral genome becomes highly methylated (as occurs normally in EBV-infected B cells and in EBV-positive B cell and epithelial cell tumors) [29]. In contrast, R preferentially activates unmethylated lytic EBV promoters [29], and in EBV-infected hTERT-immortalized normal oral keratinocyte cells (NOKs), where the viral genome remains hypomethylated, over-expression of R, but not Z, induces lytic reactivation [12,13,29–32]. Regardless of the viral genome methylation state, Z and R synergistically activate many early lytic gene promoters and expression of both proteins is required for the virus to lytically replicate and complete the lytic cycle [29,33].

We and others have previously demonstrated that epithelial cell differentiation induces EBV lytic reactivation in stably infected oral keratinocyte cells [15,16,34–39]. Similarly, EBV infection of primary oral epithelial cells grown in stratified “raft” cultures resulted in no detectable latent or lytic infection in undifferentiated layers and a robust lytic infection in the differentiated cell layers [8]. We show the cellular transcription factors KLF4 and BLIMP1, which are preferentially expressed in differentiated epithelial cells, synergistically activate the Z, R, and LMP1 EBV promoters, at least partly explaining why EBV reactivation is induced by epithelial cell differentiation [15,16,36,40,41]. However, given the robust latency of EBV infection in undifferentiated epithelial cell models [16,34–36], we hypothesized that additional factors expressed in undifferentiated (but not differentiated) epithelial cells may also contribute to the differentiation-dependent phenotype by inhibiting lytic viral reactivation in undifferentiated cells.

Here we show that expression of the master-regulator of epithelial cell identity, p63, in undifferentiated epithelial cells is a potent negative regulator of lytic EBV reactivation. The cellular p63 protein is a member of the p53 family that transcriptionally regulates many different epithelial cell-specific genes [42]. The major epithelial cell p63 isoform, ΔNp63α, is a critical regulator of proliferation and differentiation in keratinocytes and a marker for the epithelial stem cell compartment [42–47]. ΔNp63α mutations result in developmental defects in limb development, while deletion of ΔNp63α in mice is lethal after birth due to a lack of a stratified epidermis [42,46,48]. Like p53, p63 contains a DNA binding domain and, through this domain, can bind to thousands of sites on the human genome to regulate transcriptional function [49,50]. This DNA binding domain retains homology to p53, and p63 can bind to some of the same promoter sites as p53 [51]. However, unlike p53, ΔNp63α is rarely mutated in cancer and instead has been reported to be over-expressed in several epithelial tumor types, including NPC [52,53]. We find that ΔNp63α inhibits lytic EBV reactivation in epithelial cells by decreasing activity of the Z IE promoter and show that this effect is mediated both by reduced cellular p38 kinase activity and increased c-myc expression. In addition, we show that another isoform of p63, TAp63α, which is expressed in some EBV+ B cell lymphomas, likewise inhibits EBV reactivation. These findings not only further elucidate how EBV uses differentiation-dependent transcription factors (ΔNp63α, KLF4 and BLIMP1) to ensure differentiation-dependent lytic reactivation in epithelial cells, but also reveal an important mechanism for promoting viral latency in EBV-infected tumor cells.

Results

ΔNp63α and immediate-early protein Z are expressed in distinct sections of organotypic raft cultures of NOKs-Akata cells

Since ΔNp63α is the master regulator of stem cell identity in undifferentiated basal epithelial cells, and inhibits epithelial cell differentiation [43,54], we hypothesized that it also blocks lytic...
EBV reactivation. NOKs-Akata is an EBV-infected telomerase-immortalized normal oral keratinocyte cell line that retains the ability to differentiate. We previously showed that lytic reactivation of EBV occurs only in the differentiated layers of rafted NOKs-Akata cells [16,34,36]. In contrast, ΔNp63α is primarily expressed in the basal epithelial layer of normal stratified epithelium. ΔNp63α expression is lost during epithelial cell differentiation due to IRF6-mediated protein degradation as well as the antagonistic effect of the differentiation-dependent miR-203 [55–58]. To determine if expression of the IE lytic EBV protein Z overlaps that of ΔNp63α in rafted NOKs-Akata cells, we performed immunohistochemistry (IHC) using anti-Z or anti-p63α antibodies. ΔNp63α was expressed primarily in the basal layer and immediate suprabasal layers of the NOKs-Akata cell raft cultures, and ΔNp63α expression ceased in the more differentiated layers, congruent with previous literature [55–57] (Fig 1A). In contrast, Z expression was observed exclusively in the raft’s more differentiated layers (Fig 1B) with no overlap in expression of the ΔNp63α and Z proteins in the raft cultures. These observations are consistent with the hypothesis that ΔNp63α impairs the ability of EBV to lytically reactivate in epithelial cells.

Knockdown of ΔNp63α expression results in EBV lytic reactivation in EBV + carcinoma and immortalized keratinocyte cell lines

To determine if ΔNp63α expression inhibits EBV lytic reactivation in the context of undifferentiated epithelial cells, lentivirus vectors expressing shRNAs targeting p63 (or control vectors) were used to infect the CNE-2-Akata cell line (a carcinoma cell line incapable of differentiation) and stably infected cells were selected with puromycin. As shown in Fig 2A, ΔNp63α expression was successfully decreased with the p63 targeting shRNAs. Furthermore, loss of ΔNp63α expression resulted in increased expression of EBV IE proteins Z and R as well as the early lytic protein BMRF1 in comparison to control lentivirus infections (Fig 2A). Importantly, we confirmed that depletion of ΔNp63α in an authentic EBV-infected nasopharyngeal cell line, NPC43 [59], also induced expression of the lytic viral proteins Z and BMRF1 (Fig 2B).

Additionally, siRNAs targeting ΔNp63α were delivered into a third EBV-infected epithelial cell line, NOKs-Akata cells, which retain the ability to differentiate [16]. As shown in Fig 2C, knockdown of ΔNp63α expression also increased expression of EBV IE proteins, Z and R, in NOKs-Akata cells. These results indicate ΔNp63α expression represses the EBV lytic cascade and that this repression can occur even in carcinoma cell lines that are incapable of differentiation.
ΔNp63α inhibits lytic reactivation during epithelial cell differentiation

EBV lytically reactivates during epithelial cell differentiation, and this effect is at least partially mediated through enhanced expression of the transcription factors KLF4 and BLIMP1 [15,16,36]. ΔNp63α is known to impede keratinocyte differentiation through transcriptional repression of cellular genes required for differentiation [54,60,61]. To determine if ΔNp63α inhibits lytic EBV reactivation during epithelial cell differentiation.

Fig 2. ΔNp63α depletion results in EBV lytic reactivation in EBV+ carcinomas (CNE-2 and NPC43) and telomerase-immortalized oral keratinocyte (NOKs) cell lines. A) CNE-2-Akata cells were infected with lentiviruses expressing shRNAs against p63 or a control sequence, selected with puromycin, and harvested for immunoblot analyses to measure expression levels of the EBV Z, R, and BMRF1 lytic proteins, ΔNp63α, and tubulin proteins as indicated. B) NPC43 cells were infected with lentiviruses expressing shRNAs that target p63, or a control sequence. Two days post-infection, cells were harvested for immunoblot analysis to assess the expression of Z, BMRF1, ΔNp63α, and the loading control tubulin. C) NOKs-Akata cells were transfected with siRNAs against p63 or a control siRNA. Two days after transfection, the cells were harvested for immunoblot analyses and expression of the Z, R, ΔNp63α and tubulin proteins was examined as indicated. The black line indicates where irrelevant lane(s) were removed. The original western blots used to construct this figure are shown in supplemental Fig 1. All figures shown are representative of eight independent biological replicates.

https://doi.org/10.1371/journal.ppat.1010045.g002
over-expression reduces EBV lytic reactivation during epithelial cell differentiation, we stably infected NOKs-Akata cells with either a lentivirus over-expressing ΔNp63α or with a control lentivirus vector, and then seeded the cells onto collagen-treated membranes in air-liquid interface culture for three days in the presence of media containing vitamin C, calcium, and 10% serum. The cells were then harvested for an immunoblot and expression levels of the R, Z, BMRF1, ΔNp63α, involucrin, BLIMP1, and tubulin proteins was determined as indicated. 

https://doi.org/10.1371/journal.ppat.1010045.g003

over-expression reduces EBV lytic reactivation during epithelial cell differentiation, we stably infected NOKs-Akata cells with either a lentivirus over-expressing ΔNp63α or with a control lentivirus vector, and then seeded the cells onto collagen-treated membranes in air-liquid interface culture conditions. After three days of differentiating the cells, immunoblot analysis was performed to assess EBV lytic reactivation and cellular differentiation. While EBV lytically reactivated in the NOKs-Akata cells infected with the control lentivirus, cells infected with the ΔNp63α expressing lentivirus had reduced expression of the lytic EBV proteins, R, Z, and BMRF1 (Figs 3 and S3). Interestingly, the levels of the differentiation-induced cellular proteins, involucrin and BLIMP1, did not change between cells infected with the control and ΔNp63α-expressing lentiviruses, indicating that ΔNp63α repression of EBV lytic reactivation may be directly impeding viral processes as opposed to reducing cellular differentiation in these experiments.

ΔNp63α over-expression inhibits R- but not Z- mediated induction of lytic reactivation

To begin to examine the mechanism(s) for the ΔNp63α effect on lytic EBV reactivation, we next compared the effect of co-transfected ΔNp63α protein on the ability of the Z versus R EBV IE proteins to induce lytic EBV reactivation when transfected into EBV-infected epithelial cell lines that can be reactivated by either Z or R transfection. In the gastric carcinoma cell line AGS-Akata, which lacks endogenous ΔNp63α expression, co-transfected ΔNp63α inhibited the ability of transfected R protein to induce expression of endogenously expressed lytic viral proteins, Z and BMRF1 (Fig 4A). In contrast, co-transfected ΔNp63α protein did not
inhibit the ability of transfected Z protein to induce expression of endogenous lytic proteins R and BMRF1 (Fig 4A) even though more ΔNp63α was present in this condition. A repeat experiment confirmed that ΔNp63α does not inhibit the ability of transfected Z to induce lytic protein expression in AGS-Akata cells (S5 Fig). This result suggests that ΔNp63α may inhibit lytic EBV reactivation through effects on the R IE protein function and/or the BZLF1 Z promoter, since R must first induce Z expression in order to turn on early lytic genes such as BMRF1 [33].

To confirm that ΔNp63α inhibits R-mediated lytic reactivation, we examined its effect in NOKs-Akata cells, in which R transfection, but not Z transfection, can initiate the lytic cascade

https://doi.org/10.1371/journal.ppat.1010045.g004
Similar to the results in AGS-Akata cells, ΔNp63α co-transfection inhibited the ability of transfected R protein to induce expression of the Z and BMRF1 proteins, as well as the late viral capsid protein, p18 (Fig 4B and 4C). Co-transfected ΔNp63α likewise decreased the ability of transfected R to induce expression of the Z and BMRF1 lytic proteins in CNE-2-Akata cells (Fig 4D).

While R activates many lytic EBV promoter targets through a direct DNA binding mechanism [62], it is not known to bind directly to the IE Z promoter and instead is thought to activate this promoter through indirect effects on cellular transcription factors [24,25,63,64]. In addition, while some early lytic EBV promoters (such as the BMRF1 promoter) contain both Z and R binding sites and require both Z and R for efficient expression [29,65,66], maximal expression of the SM early lytic promoter has been shown to only require R binding to the promoter, and is not further enhanced by Z expression [67,68]. Interestingly, we found that while ΔNp63α inhibits the ability of co-transfected R to activate expression of the Z and BMRF1 proteins in CNE-2-Akata cells, it does not affect R activation of the SM protein (Fig 4D) [62,67]. Finally, we found that co-transfection of R with ΔNp63α in CNE-2-Akata cells decreased the production of infectious virions compared to R transfected alone (Fig 4E). These results suggest that ΔNp63α primarily inhibits lytic EBV reactivation by preventing R’s ability to induce the Z IE promoter, since the concomitant decreased expression of the BMRF1 and VCA p18 proteins, as well as the decreased titer of infectious virions, would both be the expected outcome from loss of Z expression.

**ΔNp63α repression of EBV lytic reactivation is independent of p53**

ΔNp63α is a member of the p53 protein family, and approximately 60% of ΔNp63α’s DNA binding domain is homologous to the DNA binding domain of p53 [55]. Furthermore, p53 and ΔNp63α bind to some of the same promoters, and ΔNp63α can repress p53 activation at these promoters [51,55,69]. Since p53 was previously reported to be a positive regulator of EBV lytic reactivation and binds to the Z promoter via the Sp1 transcription factor and HIF-1α [70–73], we determined if ΔNp63α’s repression of EBV lytic reactivation is due to competition with p53. For these experiments, ΔNp63α expression was inhibited by siRNAs in a NOKs-Akata cell line that had CRISPR-Cas9 mediated knockout of p53 (Figs 5A and S6). As shown in Fig 5B, immunoblot analysis revealed that loss of ΔNp63α expression upregulates Z and R expression even in the absence of p53 expression. These results suggest that ΔNp63α represses EBV lytic reactivation independently of competitive inhibition of p53 in EBV infected cells.

**ΔNp63α does not directly bind to the EBV immediate-early promoters**

ΔNp63α can function as either a transcriptional repressor or transcriptional activator, and is known to bind to promoters of cellular genes such as KLF4, p21, and p16 to prevent their expression and maintain proliferation of the keratinocyte stem cell compartment [61,69,74]. To determine if ΔNp63α inhibits lytic EBV replication by binding directly to the Zp and/or Rp IE viral promoters, we conducted chromatin-immunoprecipitation quantitative PCR (ChIP-qPCR) assays. AGS-Akata cells were transfected with a FLAG-tagged ΔNp63α vector or vector control and ChIP-qPCR assays were performed two days later using an anti-FLAG antibody as described in the methods. As shown in Fig 6A, we found that ΔNp63α binds to the cellular promoter for NECTIN1, a known ΔNp63α binding target [75], but we did not detect ΔNp63α binding to either the Z promoter or the R promoter on the EBV genome (Fig 6A). This result suggests that ΔNp63α does not bind to the immediate-early gene promoters and inhibits EBV reactivation through a different mechanism.
ΔNp63α does not prevent R binding to the EBV genome

R binds to RREs on the EBV genome during lytic reactivation to induce lytic gene expression [62]. To determine if ΔNp63α prevents R binding to RREs on the EBV genome during lytic reactivation, CNE-2-Akata cells were transfected with a FLAG-tagged R vector with or without ΔNp63α. We then performed ChIP-qPCR on known RRE sites using primers against the SM (BMLF1) promoter, BMRF1 promoter, BARF1 promoter, and the Z and R promoters (which do not contain RRE sites) to determine if R binding is affected by co-transfection with ΔNp63α [62]. Interestingly, we found that R binds to the EBV genome RRE sites in the SM, BMRF1 and BARF1 promoters equivalently with or without ΔNp63α (Fig 6B). These results are consistent with our finding that R activates SM gene expression in the presence or absence of co-transfected ΔNp63α (Fig 4D), and again suggest that a major ΔNp63α inhibitory effect may be mediated by decreasing the ability of R to indirectly activate Zp.

The C-terminal domain of TAp63 and ΔNp63 inhibits R-induced lytic reactivation

There are two primary p63 isoforms, ΔN and TA, which are defined by their N-terminal regions. The isoforms are derived from alternative transcription start sites, which either lack or contain the N-terminal transactivation domain, respectively [42]. An additional five isoform subtypes (α, β, δ, ε, and γ) can also arise from alternative splicing (or in the case of ε, a premature stop codon in codon 10) [55,76,77]. The ΔNp63α isoform is the primary isoform expressed in epithelial cells. Interestingly, TAp63α is expressed in some B-cell lymphomas, including Burkitt lymphoma and diffuse large B cell lymphomas (DLBCL), and is a potential marker for poor patient prognosis [42,55,78–81].
To determine the region(s) of ΔNp63 responsible for the suppression of EBV lytic reactivation, we transfected human ΔNp63α, mouse ΔNp63α, and mouse ΔNp63β isoforms into NOKs-Akata cells with or without R. We found that, like human ΔNp63α, mouse ΔNp63α inhibits R-mediated initiation of the lytic replication cycle. However, the mouse ΔNp63β isoform, which is missing 121 amino acids of the C-terminal domain of ΔNp63α, failed to do so (Fig 7). Additionally, TAp63α also inhibited R-induced lytic reactivation similarly to ΔNp63α, indicating that TAp63α may also be an inhibitor of lytic reactivation. ΔNp63α and TAp63α are identical at the C-terminal domains, which contain sterile alpha motif (SAM) and the post-sterile alpha motif domains, features that ΔNp63β lacks. SAM motifs in p63 isoforms provide docking sites for proteins containing SAM motifs, indicating that a protein-protein interaction could be mediating ΔNp63α and TAp63α lytic repression [82].
TAp63α acts as a repressor of EBV lytic reactivation in the Akata Burkitt lymphoma cell line

Since TAp63α is expressed in some Burkitt lymphomas and diffuse large B cell lymphomas [79,80] and TAp63α over-expression inhibits lytic reactivation in NOKs-Akata cells, we asked whether TAp63α helps maintain EBV latency in a lymphoma cell line that expresses TAp63α.

The level of TAp63α protein expression in various EBV-infected lymphoma cell lines or lymphoblastoid cell lines (including the ILB1 DLBCL line, the Akata BL cell line, an Akata EBV strain virus transformed lymphoblastoid cell line (LCL), and an AG876 EBV strain virus transformed LCL) was examined by immunoblot analysis. We found that the ILB1 DLBCL and Akata BL cell lines express TAp63α, which has a higher molecular weight compared to the ΔNp63α protein in the control NOKs-Akata epithelial cell sample (Fig 8A).

To determine if TAp63α represses EBV lytic reactivation in the B cell environment, we knocked down TAp63α expression in the Burkitt Akata cell line using an shRNA against p63 or infected the cells with two separate control shRNAs. EBV lytic protein expression was increased in cells expressing the shRNAs against TAp63α compared to the control conditions (Fig 8B). These results suggest that expression of TAp63α, like ΔNp63α, inhibits EBV lytic reactivation.
ΔNp63α inhibits constitutive Zp activity in AGS gastric cells and prevents KLF4-mediated activation of Zp

Since R activates the Zp through indirect mechanism(s) involving poorly defined cellular transcription factors [25], we next asked if the constitutive activity of Zp in the gastric AGS cell line is also inhibited by ΔNp63α. EBV-negative AGS cells were chosen for these studies since Zp activity is very high in these cells (in contrast to most cell lines) due to high expression of positive regulators of the Z promoter such as c-jun, and a lack of negative regulators such as ZEB1/2 [83]. Luciferase reporter constructs containing either no promoter, or various portions of the Z promoter, were transfected into AGS cells with or without a ΔNp63α expression vector, and the amount of luciferase activity was quantified two days later. As shown in Fig 9A, ΔNp63α decreased the activity of each of the Zp reporter constructs tested, including a construct containing only 83 base pairs upstream of the BZLF1 transcription initiation site. In contrast, ΔNp63α did not repress activity of the promoterless luciferase vector, and, as previously reported [84], increased the activity of the EBV BARF1 promoter in the same reporter vector (Fig 9B).

We also examined the effect of ΔNp63α on the ability of co-transfected KLF4 to activate Zp in reporter gene assays. KLF4 is thought to activate Zp through a direct binding mechanism [16]. ΔNp63α decreased the ability of co-transfected KLF4 to activate Zp-driven luciferase activity but did not affect the total level of transfected KLF4 protein (Fig 9B and 9C). Together, these results strongly suggest that ΔNp63α inhibits Z promoter activation by positively acting regulators, including the EBV R protein and KLF4.
Fig 9. ΔNp63α inhibits Z promoter activity in reporter gene assays. A) Luciferase vectors containing various amounts of the Z promoter sequence, or the pCpG (promoterless) negative control vector, were transfected into EBV-negative AGS gastric carcinoma cells with or without an ΔNp63α expression vector. Luciferase activity was measured two days later. The average fold change in luciferase activity in ΔNp63α transfected cells versus control vector transfected cells for each promoter construct is shown, with error bars indicating standard error. Analysis of statistical significance between ΔNp63α and control conditions in the pCpG vector, Zp-346 vector, Zp-169 vector, and Zp-83 vector was performed using a two-sample t-test as indicated. B) AGS cells were transfected with the pCpG negative control luciferase vector, or luciferase vectors driven by the EBV early lytic BARF1 promoter, or the Z (Zp-346) promoter, in the presence or absence of a KLF4 expression vector, with or without co-transfected ΔNp63α, as indicated. The average fold change in luciferase activity for each condition (relative to each promoter construct transfected with control vector) is shown, as well as the standard error. Analysis of statistical significance between ΔNp63α and the control or KLF4 conditions in the pCpG vector, BARF1p vector, and Zp-346 vector was performed using a two-sample t-test as indicated. C) Immunoblot analysis of the Zp-luciferase assay samples shown in Fig 9B.
Lytic repressor c-myc is increased by ΔNp63α

Since our results are consistent with a model in which ΔNp63α indirectly inhibits the activity of the Z promoter by upregulating expression of a cellular protein that inhibits Zp activity, we examined the effect of ΔNp63α over-expression, or ΔNp63α knock-down, on the level of c-myc in NOKs-Akata or CNE-2-Akata cells. C-myc was recently shown to be a potent inhibitor of EBV reactivation via repressive effects on Zp activity [19]. We observed that knockdown of ΔNp63α expression is associated with a concomitant decrease in c-myc expression (Fig 10A), while over-expression of ΔNp63α increases c-myc expression (Fig 10B), consistent with previous reports that Np63α helps maintain c-myc expression in basal epithelial cells [19,85].

To determine if this alteration in c-myc expression is sufficient to completely explain the inhibitory effect of ΔNp63α on lytic reactivation, we knocked down c-myc expression using siRNAs in NOKs-Akata cells, and then transfected cells with an R expression vector in the presence or absence of a ΔNp63α expression vector. As shown in Fig 10C, although we confirmed that knock-down of c-myc alone does increase R-mediated lytic reactivation, we also found that ΔNp63α still inhibits R-mediated lytic reactivation even when c-myc expression is knocked down. Similar results were obtained in a second experiment (S8 Fig). Thus, while alterations in c-myc expression may well contribute to the ΔNp63α inhibitory effect, additional inhibitory mechanisms are also likely involved.

ΔNp63α inhibits p38 MAPK activity, which also contributes to ΔNp63α suppression of lytic EBV reactivation

ΔNp63α was recently reported to inhibit activity of cellular p38 MAP kinase by increasing expression of the cellular DUSP6 phosphatase [86]. Since ΔNp63α inhibits the ability of transfected R, but not Z, to induce lytic reactivation in epithelial cells (Fig 4), and we previously showed that the cellular p38 MAP kinase is required for R-mediated (but not Z-mediated) activation of the BZLF1 and BMRF1 lytic promoters in latently infected epithelial cells [25], we next asked whether ΔNp63α at least partially inhibits R-mediated activation of BZLF1 and BMRF1 expression by decreasing p38 kinase activity. To examine this possibility, we over-expressed ΔNp63α in NOKs-Akata cells (using a lentivirus vector) and examined the effect on p38 kinase activity. We confirmed that phosphorylated (activated) p38 is substantially decreased in ΔNp63α over-expressing NOKs-Akata cells compared to control cells (Fig 11A). Additionally, when we knocked down ΔNp63α in the context of the authentic NPC cell line NPC43, we found that this depletion increased phosphorylated p38 (Fig 11B).

Cellular p38 kinase is encoded by four different highly homologous cellular genes (MAPK11–14). As shown in Fig 11C, knock-down of MAPK14 (p38α) expression (the most highly transcribed p38 encoding gene in NOKs-Akata cells [35]) inhibited constitutive Z and BMRF1 expression in NOKS-Akata cells, while R expression was only slightly affected. To examine how loss of p38 kinase activity affects the ability of R to induce various different lytic EBV promoters, NOKs-Akata cells were transfected with R in the presence or absence of a p38 kinase inhibitor (SB 202190, which affects both MAPK11 and MAPK14 p38 kinase activity), or an siRNA directed against the MAPK14 p38α kinase. As shown in Fig 11D, similar to the effect of ΔNp63α, treatment of cells with a p38 inhibitor greatly decreased R’s ability to activate expression of both the BZLF1 IE protein and BMRF1 early lytic proteins. As expected,
inhibition of p38 kinase activity decreased expression of its known downstream target, TTP (ZFP36) [87]. Specific knock-down of MAPK14 (p38α) expression inhibited R’s ability to activate BMRF1 expression while having a lesser effect on BZLF1 expression (Fig 11E).

https://doi.org/10.1371/journal.ppat.1010045.g010

Fig 10. ΔNp63α expression increases the expression of the lytic repressor c-myc, but c-myc is not required for ΔNp63α mediated repression of lytic reactivation. A) NOKs-Akata cells were transfected with either ΔNp63α or c-myc targeting siRNAs or a control nonspecific siRNA. Two days after siRNA transfection, the cells were harvested for an immunoblot to determine the expression levels of ΔNp63α, c-myc, and tubulin (as a loading control). B) CNE-2-Akata cells were infected with a ΔNp63α expressing lentivirus or empty control. After five days of puromycin selection, the cells were harvested and an immunoblot performed to assess expression of the ΔNp63α, c-myc and tubulin proteins. C) NOKs-Akata cells were transfected with siRNAs against c-myc or a control siRNA for two days, and then transfected with or without an R expression vector in the presence or absence of ΔNp63α. Immunoblot was performed one day later to examine expression of c-myc, ΔNp63α, R, BMRF1, Z, and tubulin proteins. Black lines indicate where irrelevant lanes in the blot were removed; the original western blots used to construct this figure are shown in supplemental Fig 8. All experiments are representative of two independent experiments.
ΔNp63α over-expression decreases p38 kinase activity, which is required for R mediated lytic reactivation.

A) Immunoblot analysis was performed on protein lysates harvested from NOKs-Akata cells infected with a lentivirus vector expressing ΔNp63α, or a control lentivirus vector, and expression of ΔNp63α, phosphorylated (activated) p38, total p38α (MAPK14), and tubulin (a loading control) was assessed.

B) NPC43 cells were infected with a lentivirus expressing shRNA targeting p63 or a control sequence, then harvested for western blots to examine the expression of ΔNp63α, phosphorylated p38, total p38α and tubulin. Note that these samples were also used in Fig 2B.

C) NOKs-Akata cells were transfected with either a control siRNA or an siRNA targeting p38α (MAPK14), and two days later immunoblot analysis was performed to examine expression of p38α (MAPK14), R, BMRF1, Z, and tubulin as indicated. Quantitation of R, BMRF1 and Z expression was determined through ImageJ, normalized to tubulin, and the siRNA control sample was set as 1.

D) NOKs-Akata cells were transfected with or without an R expression vector in the presence or absence of 10 μM p38 MAPK inhibitor SB202190. Immunoblot analysis was performed two days later to examine expression levels of Z, R, BMRF1, TTP and tubulin as indicated. Expression of TTP was quantitated with ImageJ, normalized to tubulin with the first lane set as 1.

E) NOKs-Akata cells were transfected with either a
Interestingly, the finding that specific knock-down of the p38 α form of p38 kinase affected BZLF1 expression less than the p38 kinase inhibitor suggests that additional p38 kinase protein(s) may contribute to inhibition of BZLF1 expression, although this effect could also be due to incomplete knockdown. Together, these results reveal that p38 kinase activity is crucial for R-mediated lytic viral reactivation in NOKs-Akata cells and suggest that ΔNp63α inhibition of p38 kinase activity is an important mechanism by which it inhibits lytic reactivation.

### P38 kinase activation of lytic EBV infection does not require activation of the MK2 pathway

Although we previously showed that one mechanism by which p38 MAP kinase enhances lytic EBV reactivation is via phosphorylation and activation of the cellular ATF2 transcription factor (which directly binds to and activates the BZLF1 (Zp) promoter)[25], growing evidence suggests that p38 kinase commonly activates gene expression not only by enhancing transcription factor activity, but also by increasing the stability of ARE-containing RNA transcripts [87]. Since the major mechanism by which the p38 kinase promotes RNA stability is through phosphorylation and activation of the MK2 kinase (MAPKAPK2) [88], we examined the ability of a MK2 inhibitor to affect R-mediated lytic reactivation in NOKs-Akata cells. As shown in Fig 11F, inhibition of MK2 activity using a chemical inhibitor did not suppress the ability of transfected R protein to induce BZLF1 or BMRF1 expression. Loss of TTP (ZFP36) expression (which requires MK2 activity for high level expression [87]) in this experiment confirmed that MK2 activity was decreased (Fig 11G) [87].

### ΔNp63α over-expression during cis-platinum treatment of CNE-2-Akata and NOKs-Akata cells inhibits EBV lytic reactivation

Chemotherapy agents such as cis-platinum have been shown to potently induce EBV lytic reactivation [71,89,90], and the treatment of cells with cis-platinum is known to degrade ΔNp63α [91]. We hypothesized that ΔNp63α over-expression may therefore inhibit cis-platinum-induced lytic reactivation in epithelial cells. To examine this, we over-expressed ΔNp63α in CNE-2-Akata cells and then treated with 10 μM cis-platinum for two days to induce lytic reactivation. Over-expression of ΔNp63α resulted in a striking complete loss of lytic induction in response to cis-platinum (Fig 12A); similar results were obtained in NOKs-Akata cells (Fig 12B). These results indicate that ΔNp63α is a major repressor of cis-platinum-induced lytic reactivation and conversely, loss of intact ΔNp63α protein expression contributes to the lytic inducing effect of cis-platinum.

### Discussion

Infection of oropharyngeal epithelial cells plays an essential role in the EBV life cycle, particularly since these cells are the major source of transmissible virus. However, while EBV infection of normal oropharyngeal epithelial cells is highly lytic, and likely confined to the more
differentiated cell layers, EBV infection of nasopharyngeal carcinoma cells is largely latent. Furthermore, the ability of NPC tumor cells to maintain EBV latency is likely required for their growth and survival. A major difference between NPC cells and the non-transformed oropharyngeal epithelial cells normally infected by EBV is that NPC cells are undifferentiated basal cells that express high levels of the basal cell transcription factor, ΔNp63α, which is lost when stratified epithelial cells differentiate. Here we demonstrate that both the basal epithelial cell specific ΔNp63α transcription factor, and the related isoform TAp63α (expressed in some EBV-positive lymphomas), act as potent negative regulators of EBV lytic reactivation. These findings help to explain how EBV can achieve latent infection in undifferentiated NPC and B cell lymphomas.

EBV has two different IE proteins, Z and R, which encode viral transcription factors that cooperate to induce lytic reactivation of EBV. Our studies here show that the major inhibitory effect of ΔNp63α on lytic EBV reactivation occurs via negative regulation of the BZLF1 gene promoter (Zp). We found that ΔNp63α prevents the ability of transfected R protein to induce Z protein expression in latently infected epithelial cells, while not inhibiting the ability of transfected Z protein to induce R expression and downstream early lytic EBV proteins (Fig 4). These results indicate that Z over-expression (under the control of a heterologous promoter) is sufficient to bypass the inhibitory effect of ΔNp63α. Interestingly, in contrast to the effect on R-mediated early lytic BMRF1 protein expression, we found that ΔNp63α does not inhibit R-mediated activation of the early lytic SM protein (Fig 4D). This difference likely reflects the fact that the SM lytic gene can be turned on by R alone (even in the absence of Z expression [67,92]), while activation of BMRF1 gene expression requires both the Z and R proteins [29,65,66]. Our finding that ΔNp63α also inhibits late lytic protein expression in R-transfected cells reflects the known requirement for Z to complete the lytic form of viral DNA replication [29,33] and to initiate subsequent late viral gene expression. Thus, loss of sufficient Z expression is the key defect impeding lytic viral reactivation in ΔNp63α expressing cells.

Although ΔNp63α is an essential regulator of epithelial cell differentiation and is well known to inhibit differentiation of epithelial cells by repressing expression of key cellular
transcription factors such as KLF4 [61] that are required for epithelial cell differentiation, we found here that ΔNp63α inhibits Zp activity even in cells that cannot differentiate such as AGS cells and CNE-2 cells (Figs 4A, 4D and 9). Thus, although EBV uses differentiation-dependent cellular transcription factors such as KLF4 and BLIMP1 to ensure that Zp and Rp are both activated when EBV-infected epithelial cells differentiate [15,16], it has usurped the functions of the basal cell specific transcription factor ΔNp63α to inhibit Zp activity even in the absence of KLF4 and BLIMP1. We show here that ΔNp63α inhibits the constitutive activity, as well as KLF4-induced activity, of Zp-driven reporter gene constructs in EBV-negative AGS gastric carcinoma cells (Fig 9).

EBV-negative AGS cells do not normally express ΔNp63α and in contrast to most other cell types are remarkable for their high level of constitutive Zp activity [83]. We found that ΔNp63α strongly inhibits constitutive Zp activity in AGS cells and this effect was observed even with the smallest Z promoter construct studied (which contains only 83 base pairs upstream of the transcriptional initiation site). Of note, each of the Zp luciferase constructs used in this study contains the AP-1-like “ZII” motif, which binds to the AP-1 and ATF2 [25] transcription factors and is required for constitutive Zp activity. ΔNp63α also inhibits the ability of transfected KLF4 protein to enhance Zp activity in AGS cells, without affecting the level of transfected KLF4 (Fig 9). We previously showed that KLF4 binds directly to Zp and contributes to differentiation-dependent lytic EBV reactivation in epithelial cells [16]. Together, these results suggest that ΔNp63α inhibits the ability of multiple different stimuli to induce Zp activity.

To examine whether ΔNp63α inhibits lytic EBV reactivation by binding directly to the Zp or Rp IE promoters, we performed ChIP assays using a FLAG-tagged ΔNp63α expression vector (Fig 6A). Although we confirmed that ΔNp63α bound to a positive control cellular promoter in these assays, we did not observe binding of ΔNp63α to the Zp or Rp EBV IE promoters. To determine whether ΔNp63α blocks R binding to known RRE sites in the early lytic BMRF1 and SM promoters, we also performed ChIP assays using a FLAG-tagged R expression vector to compare R binding to these promoters in the presence and absence of co-transfected ΔNp63α (Fig 6B). ΔNp63α co-transfection did not affect R binding to either the BMRF1 or SM promoters. Together, these results suggest that the inhibitory effect(s) of ΔNp63α on Zp activity do not require either direct ΔNp63α binding to the Zp or Rp IE promoters, or inhibition of R binding to other EBV promoters.

To further examine potential mechanism(s) for the inhibitory effect of ΔNp63α on Zp activity, we compared the ability of different p63 isoforms to block R-mediated lytic EBV reactivation in NOKs-Akata cells (Fig 7). These studies revealed that the C-terminal domain of ΔNp63α is required for the inhibitory effect. Since the TAp63α isoform (which is expressed in some EBV+ lymphomas) shares this inhibitory C-terminal domain, we also asked whether this isoform can block lytic viral reactivation. Importantly, we found that the Burkitt Akata lymphoma line expresses TAp63α, and that knockdown of TAp63α in these cells increases constitutive lytic EBV protein expression (Fig 8). Thus, other p63 family members may also play a role in maintaining EBV latency in some EBV-infected lymphomas.

Our results here suggest that one mechanism by which ΔNp63α promotes viral latency is by increasing expression of c-myc (Fig 10). ΔNp63α is known to enhance c-myc expression in basal epithelial cells [93], and c-myc was recently shown to potently inhibit Z expression and lytic reactivation in EBV-infected cells [19]. We confirmed that ΔNp63α activates c-myc expression in NOKs-Akata cells and showed that siRNA-mediated knock-down of c-myc increases the ability of transfected R protein to induce Z and BMRF1 expression (Fig 10). Nevertheless, since we found that ΔNp63α still inhibits R-mediated lytic reactivation in NOKs-Akata cells even when c-myc expression is greatly decreased using siRNA (Fig 10), the
\(\Delta Np63\alpha\) effect on c-myc expression is unlikely to be the only mechanism by which \(\Delta Np63\alpha\) promotes viral latency.

Instead, our experiments here suggest that another major mechanism by which \(\Delta Np63\alpha\) promotes viral latency in epithelial cells is by decreasing p38 kinase activity. We found that \(\Delta Np63\alpha\) expression inhibits p38 kinase activity in NOKs-Akata cells, and that inhibition of p38 kinase activity reduces both constitutive, and R-mediated activation of the Z and BMRF1 proteins in NOKs-Akata cells (Fig 11). Although the mechanism(s) by which R increases Zp activity in the context of the intact viral genome are not completely understood, direct DNA binding of R to the Z promoter does not appear to be involved, and instead R activation of Zp is thought to be mediated through indirect effects of R on cellular transcription factors [28,63,64,71]. We previously showed that inhibition of p38 kinase activity blocks the ability of over-expressed R protein, but not over-expressed Z protein, to induce lytic EBV reactivation in latently infected epithelial cells, and correlated this effect with the ability of p38-phosphorylated ATF2 transcription factor to bind to and activate the BZLF1 promoter [25]. More recently, the ability of the late lytic BGLF2 tegument protein to promote lytic reactivation was also shown to be at least partially mediated through p38 kinase activation [94,95].

In addition to its requirement for R-mediated and BGLF2-mediated lytic EBV reactivation in epithelial cells, a growing literature indicates that p38 kinase activity is required for the ability of multiple different types of stimuli to induce lytic EBV reactivation in both epithelial cells and B cells. Indeed, p38 kinase inhibitors have been shown to block viral reactivation in response to such diverse stimuli as B-cell receptor activation, chemotherapy agents, phorbol esters, reactive oxygen species, and TGF-\(\beta\) [96–101]. Given the large number of direct p38 kinase substrates, and the multiple downstream pathways that are indirectly affected by this kinase, we speculate that p38 kinase contributes to lytic EBV reactivation through multiple different mechanisms. Our results here suggest that one of the primary effects of this kinase on EBV reactivation is to enhance BZLF1 promoter activation in response to both viral and cellular transcription factors. Interestingly, a number of different cellular transcription factors that have previously been shown to bind to and activate the BZLF1 promoter (including ATF1, ATF2, c-jun, c-fos, XBP1, HIF-1\(\alpha\), and p53) are known to be directly phosphorylated and activated by p38 kinase [102]. Furthermore, p38 kinase more globally enhances transcription by activating downstream kinases such as MSK1 and MSK4 that directly phosphorylate histone sites such as H3 S28; this phosphorylation then reduces inhibitory modifications on H3 K27 that recruit the polycomb complex [103–105]. Of note, p38\(\gamma\) activity has previously been shown to increase during epithelial cell differentiation and to be required for efficient induction of differentiation genes such as involucrin [106].

In addition, p38 kinase activity is increasingly recognized to exert many of its cellular effects by increasing stability of a subset of cellular RNAs [88]. This effect is mediated largely through activation of the MK2/MKK3 kinases, which phosphorylate and inactivate a cellular protein, ZFP36 (TTP), that induces degradation of RNAs containing AU-rich 3’ LTRs [87]. However, treatment of NOKS-Akata with an MK2 inhibitor did not impact R-induced lytic reactivation, indicating that p38 may facilitate lytic reactivation through pathways independent of effects on RNA stability (Fig 11F). Interestingly, p38 kinase was also recently reported to inhibit nonsense mediated decay (NMD) of RNA in cells with DNA damage [107]. Since NMD was recently shown to promote degradation of the transcript encoding the BZLF1 and BRLF1 genes in latently infected cells [108], decreasing NMD of this transcript in the context of the intact viral genome could be another post-transcriptional mechanism by which p38 kinase promotes lytic EBV reactivation.

Finally, we show here that the recently described ability of the chemotherapy agent cis-platinum to induce \(\Delta Np63\alpha\) degradation in epithelial cells [91] can potentially be used to potentiate
“lytic induction” therapy for EBV-positive epithelial tumors. Lytic induction therapy of latently infected EBV-positive tumor seeks to identify small molecules that can reactivate the lytic form of viral infection in tumors, thereby using the virus itself to help kill the tumor cell. Interestingly, we previously showed that p38 kinase activity is required for chemotherapy to induce lytic EBV reactivation in certain epithelial cell lines [89]. Here we confirmed that cis-platinum treatment of EBV-infected epithelial cells does, indeed, result in loss of intact ΔNp63α expression and showed that restoration of ΔNp63α expression in cis-platinum treated cells reduces the amount of lytic EBV reactivation (Fig 12A and 12B). Thus, the use of chemotherapy drugs such as cis-platinum that can not only induce DNA damage and p53/ATM activation (known activators of lytic EBV reactivation [71,73]), but can simultaneously degrade ΔNp63α, is predicted to be the most effective approach for achieving efficient lytic induction therapy in EBV-positive NPC tumors.

Materials and methods

Cell culture

The normal oral keratinocytes (NOKs) cell line (a generous gift from Karl Munger of Tufts University via Paul Lambert of the University of Wisconsin) is a telomerase-immortalized normal oral keratinocyte cell line, grown in keratinocyte serum-free media supplemented with 12.5 mg bovine pituitary extract, and 0.1 μg epidermal growth factor per 500ml of media (KSFM, Lifetech). NOKs were derived as previously described [109]. p53KO NOKs were derived by CRISPR-Cas9 mutagenesis of the p53 gene as previously described [73]. In brief, these NOKs- cells were infected with a lentivirus containing a guide RNA (5'-TCGACGC TAGGATCTGACTG-3') targeting the second exon of p53 and selected with puromycin. EBV-infected NOKs-Akata cells were created as previously described [29] and were maintained with 50 μg/ml G418 antibiotic selection in addition to the media/growth supplements used to grow NOKs cells. The CNE-2 cell line was derived initially from NPC tumors but has been subsequently shown to have HeLa cell genome contamination and HPV infection [110,111]. CNE-2-Akata cells (a generous gift from K.W. Lo of the Chinese University of Hong Kong via Diane Hayward) are stably EBV-infected with the Akata strain of EBV (containing a G418 resistance gene cassette and GFP gene inserted into the EBV BXLF1 gene) and were grown in DMEM with 10% fetal bovine serum, 1% pen-strep, and 400 μg/ml G418 antibiotic selection. The NPC43 cell line is an authentic EBV-infected nasopharyngeal carcinoma cell line described previously [59] and was grown in RPMI media supplemented with 10% fetal bovine serum, 1% pen-strep, and 4 μM Y-27632. The AGS cell line is an EBV-negative gastric carcinoma cell line that was obtained from the ATCC and was grown in F12 media with 10% fetal bovine serum and 1% pen-strep. The AGS-Akata cell line is an AGS cell line stably infected with the Akata strain of EBV (containing a G418 resistance gene cassette and GFP gene inserted into the EBV BXLF1 gene) was derived as previously described [112] and was a kind gift from Lindsay Hutt-Fletcher. The AGS-Akata cell line was grown in F-12 media with 10% fetal bovine serum, 1% pen-strep, and 400 μg/ml G418 antibiotic selection. Both the uninfected and EBV-infected AGS were cured (in the Hutt-Fletcher lab) of the contaminating SV5 virus present in most AGS lines. The Burkitt lymphoma cell line Akata (Burkitt Akata) (a gift from Lindsay Hutt-Fletcher) was derived as previously described by super-infecting an EBV-negative Akata Burkitt lymphoma cell clone with the Akata strain of EBV (containing a G418 resistance gene cassette and GFP gene inserted into the EBV BXLF1 gene)[112] and was maintained with RPMI media with 10% fetal bovine serum, 1% pen-strep and 500 μg/ml G418 antibiotic selection. EBV-negative Akata cells were a kind gift from Kenzo Takada of Hokkaido University, Japan, via Bill Sugden of the University of Wisconsin and have been previously
derived as described [113]. The IBL1 diffuse large B cell lymphoma line is a gift from Ethel Cesarman and was derived as previously described [114]. The AG876 and Akata lymphoblastoid cell lines (LCLs) were derived from peripheral blood B cells transformed with either the AG876 or Akata strains of EBV, respectively, as previously described [115] and were maintained with RPMI supplemented with 10% fetal bovine serum and 1% pen-strep.

Collagen membrane differentiation

Collagen membrane differentiation was performed as previously described [116]. In brief, approximately 5x10^5 total NOKs-Akata cells were seeded onto a collagen-treated transwell membrane (Corning #3460) with KSFM media on both the basal and apical surfaces of the membrane. After 24 hours, when cells were 100% confluent, apical media was removed and the basal media exchanged for Epilife media (Thermofisher # MEPICF500) supplemented with 10% FBS, 1.4 mM CaCl_2, and 5 μg/ml ascorbic acid. Cell extracts on the collagen were harvested with sumo lysis buffer for immunoblot analysis after three days of differentiation.

Immunoblots

Immunoblots were performed as previously described [117]. Briefly, cell lysates were harvested with sumo lysis buffer (1:3 mixture of buffer I (5% sodium dodecyl sulfate (SDS), 0.15 M Tris-HCl (pH 6.8), 30% glycerol) and buffer II (25 mM Tris-HCl (pH 8.3), 50 mM NaCl, 0.5% NP-40, 0.5% deoxycholate, 0.1% SDS) with protease inhibitors (cOmplete, Roche). Quantitation of protein concentration was conducted with a DC Bio-rad protein assay. The lysates were separated using a 10% polyacrylamide gel and then transferred onto a nitrocellulose membrane. The membranes were subsequently blocked with 5% milk consisting of 0.1% Tween 20 and 1X PBS for one hour. Membranes were then incubated with primary antibody overnight. The following day the antibodies were removed and the membrane was washed with wash buffer (1X PBS, 0.1% Tween 20) three times for 5 minutes. The membrane was then incubated with secondary antibody suspended in 5% milk for 30 minutes. DAB was added for a period of approximately 15 seconds, and then horseradish peroxidase for approximately 10 seconds. Hematoxylin was used as a counterstain and the slides were dehydrated in alcohols before mounting. Z antibody (Santa Cruz, catalog # sc-53904) and p63α antibody (Cell Signaling Technologies catalog # 13109) were both used at a 1:200 dilution.

Immunohistochemistry

Immunohistochemistry (IHC) was conducted as previously described [118]. To summarize, paraffin-embedded slides were initially heated on 70°C heat blocks and deparaffinized in xylene two times for 10 minutes. The slides were then hydrated in a series of alcohols (100%, 90%, 70%) and subsequently water for a period of 5 minutes each. The slides were then boiled in 1X sodium citrate buffer for 20 minutes and allowed to cool for an hour before blocking with horse serum. Primary antibodies were added to the slides for an hour, and then the primary was washed off in 1X PBS before adding secondary for thirty minutes. DAB was added for a period of approximately 15 seconds, and then horseradish peroxidase for approximately 10 seconds. Hematoxylin was used as a counterstain and the slides were dehydrated in alcohols before mounting. Z antibody (Santa Cruz, catalog # sc-53904) and p63α antibody (Cell Signaling Technologies catalog # 13109) were both used at a 1:200 dilution.

Luciferase assays

Luciferase assays were conducted as previously described [16]. To summarize, 48 hours after transfection EBV-negative AGS cells were washed once with PBS and suspended in 200 μL of reporter lysis buffer (Promega). The lysate was flash-frozen once and pelleted by
centrifugation. The supernatant of the lysate was used for the luciferase assays according to the manufacturer’s instruction using a BD Monolight 3010 luminometer (BD Biosciences).

**Chemicals**

Cisplatin (Sigma, catalog #1134357) was made up at 10 mM and used at 10 μM. SB202190 p38 kinase inhibitor was purchased from Santa Cruz (sc-202334) was used at 10 μM. Phorbol 12-myristate 13-acetate (TPA) was purchased from Sigma (catalog #P8139) used at 20 ng/ml. MK2 inhibitor PF-364402 hydrate was purchased from Millipore-Sigma (catalog #PZ0188) and used at 10 μM. Y-27632 was purchased from R&D systems (catalog #1254/10) and used at 4 μM. Control conditions were treated equal amounts of the solvent.

**Antibodies**

The following antibodies were used for immunoblot analyses in this study: anti-R rabbit polyclonal antibody directed against the R peptide (peptide sequence EDPDEETSSQAVKALRE-MAD 1: 2500), anti-p63α (Cell Signaling Technologies catalog # 13109 1:1000), anti-ΔNp63 (Biolegend, catalog #619001), anti-KLF4 (Cell Signaling Technologies, catalog #4038 1: 1000), anti-BZLF1 (Santa Cruz, catalog # sc-53904, 1:500), anti-BMRF1 (Millpore, catalog # MAB8186, 1: 2500), anti-p18 (Thermo-Fisher Scientific, # PA1-73003, 1:2000), anti-BLIMP1 (Cell Signaling Technologies catalog # 9115S 1:1000), anti-involucrin (Sigma, catalog #19018, 1:3000), anti-c-myc (Abcam catalog# ab32072, 1:10000), p38 MAPK14 antibody (Cell Signaling Technologies catalog#9218 1:1000) and phospho-p38 MAPK (Cell Signaling Technologies catalog #9211 1:1000), a generous gift from Sankar Swaminathan of the University of Utah, anti-Tristetraprolin (Cell Signaling Technologies catalog# #71632 1:1000). The secondary antibodies used were Horseradish peroxide (HRP)- labeled goat anti-mouse antibody (Thermo Scientific# 31430, 1:5000), HRP- labeled donkey anti-goat antibody (Santa Cruz#sc-2056, 1:1000), and HRP- labeled goat anti-rabbit antibody (Fisher Scientific 1:10000).

**Plasmids**

All plasmid DNA was prepared using the Qiagen Maxi-prep kit according to the manufacturer’s instructions. The plasmid pSG5 was purchased from Stratagene. pSG5-R and pSG5-Z (kind gifts from Diane Hayward of John Hopkins University) contain the BZLF1 (Z) and BRLF1 (R) immediate-early genes driven by the SV40 promoter as previously described [14,119]. The pCPG, BARF1p, and the Zp luciferase expression vectors have been described previously [16,29]. ΔNp63α-FLAG (Addgene # 26979, a gift from David Sidransky of John Hopkins University) encodes the ΔNp63α gene with a FLAG tag. ΔNp63α pReceiver-Lv105 (Genecopeia product # Z57540-Lv105) is a ΔNp63α lentiviral expression vector, and plenti is the lentiviral empty control (Addgene #9481, a gift from Ie-Ming Shih of John Hopkins University). ΔNp63α-CMV was purchased from Genecopeia (catalog # EX-Z5740-M02). pCMV3FC and BRLF1-pCMV3FC (containing a FLAG-tagged BRLF1 protein) are both kind and generous gifts from Lori Frappier of the University of Toronto [120]. Mouse ΔNp63α-myc, TAp63α-myc, and ΔNp63β-myc plasmids are generous gifts of Xiaohua Su at MD Anderson and Elsa Flores at the Moffitt Cancer Center.

**siRNAs**

siRNAs against ΔNp63α (sc-36161A, sc-36161B), c-myc (sc-29226, sc-44248), MAPK14 (sc-29433), and control siRNAs A and C (sc-37007, sc-44231) were purchased from Santa Cruz.
siRNAs at 20 pM final concentration were transfected in 12 well plates using RNAiMAX (Invitrogen) according to the manufacturer’s protocol. After two days, the cells were harvested with sumo lysis buffer for immunoblot analysis.

### Transient transfections

DNA was transfected into NOKs-Akata, CNE-2-Akata, AGS-, and AGS-Akata using the Lipofectamine 2000 (Thermo Fisher #11668019) system according to the manufacturer’s protocol. Generally, 500 ng of total DNA with 1.5 μL Lipofectamine 2000 was used per condition to transfect epithelial cells that were approximately 70% confluent in a 12 well plate.

### shRNAs and lentivirus packaging

shRNAs against ΔNp63α purchased from Horizon (catalog#RHS4533-EG8626). Lentivirus packaging was done as previously described [16]. Packaging components pCMV-VSV-G (Plasmid #8454, a gift from Bob Weinberg of the Massachusetts Institute of Technology) and pSPAX2 (Plasmid #12260, a gift from Didier Trono from the École Polytechnique Fédérale de Lausanne) were purchased from Addgene. To package the lentiviruses, 4 μg of the vector, 0.6 μg of VSV-G, and 1.4 μg of pSPAX2 were transfected into 293T cells plated in a 10 cm dish. The cell’s media was changed 24 hours post-transfection and viral supernatant was harvested on days 2 and 3 to infect cells. Cells were selected with puromycin three days post-infection to derive a stable cell line.

### Organotypic rafting

Cells were stratified by organotypic rafting as described previously [16]. To briefly summarize, a dermal equivalent was created using transwell inserts (24 mm diameter, 0.4 μM pore Costar) that were coated with 1ml collagen mix (3 mg/ml Wako) also containing F-media, 10% FBS and 1% pen-strep. This layer was coated with an additional 2.5 ml collagen mix containing F-12 media, 10% FBS, 1% pen-strep, and 4.5 X 10^5 early- passage human fibroblasts (EF-1-F). This dermal equivalent was suspended in F-12 medium supplemented with 10% FBS and 1% pen-strep. Four days later, 2.1 x 10^5 NOKs-Akata cells were plated on the dermal equivalent suspended in keratinocyte plating media (F-medium [1.88 mM Ca^2+]) supplemented with 0.5% FBS, adenine (24 μg/ml), cholera toxin (8.4 ng/ml), hydrocortisone (2.4 μg/ml), and insulin (5 μg/ml). The cells were allowed to grow for four days to reach confluence, and at this point the media was exchanged for cornification media (keratinocyte plating medium containing 5% FBS and 10 μM C8:0), and the cells were lifted to the air liquid interface so that only the basal cells were supplied with media. After 11 days of differentiation in cornification media that was replaced every other day, the cells were harvested and embedded in 2% agar-1% formalin, and fixed in 10% neutral buffered formalin overnight. The raft tissues were subsequently embedded in paraffin and sectioned in 4 μM cross sections.

### Chromatin immunoprecipitation (ChIP) and quantitative PCR

ChIP assays were performed as described previously [121]. For p63 ChIP assays, AGS-Akata cells (using three 10cm dishes per condition) were transfected with a pCDNA empty vector control or a p63-FLAG expression vector. For the R-FLAG ChIP assays, CNE-2-Akata cells were transfected with BRLF1-pCMV3FC (which contains a FLAG tag), BRLF1-pCMV3FC with ΔNp63α-CMV, or pCDNA empty vector control. 24 hours post-transfection, the cells were fixed with 1% paraformaldehyde for 10 minutes and quenched with 125 mM glycine for 5 minutes. The cells were then pelleted by centrifugation at 1500 rpm for 10 minutes, washed
with PBS. The cell pellets were then lysed in cell lysis buffer (10 mM Tris pH 8.0, 10 mM NaCl, 0.2% NP40) with protease inhibitors (cOmplete, Roche) to remove cytoplasmic protein. Nuclear pellets were lysed in nuclei lysis buffer (50 mM Tris-SC pH 8.0, 10 mM EDTA pH 8.0, 1% SDS) with protease inhibitors and the mixture was left on ice for 10 minutes. The nuclear lysates were then diluted in IP dilution buffer (20 mM Tris-SC pH 8.0, 2 mM EDTA, 150 mM NaCl, 1% Triton X100, and 0.01% SDS) and sonicated 4 cycles (30 seconds ON at 10 watts/90 seconds OFF) (Fisher Scientific, Sonic Dismembrator Model 100). Approximately 25 μg DNA were diluted with IP dilution buffer, blocked with magnetic A/G beads (Thermo-Fisher, 88802) for one hour, and immunoprecipitated with M2 FLAG (Sigma-Aldrich, M8823-1 ml) beads overnight. The samples were then washed with low salt, high salt, and lithium chloride wash buffer for 15 minutes each, followed by two 15-minute washes with T₁₀E buffer. Protein/chromatin complexes were eluted with elution buffer (0.1M NaHCO₃, 1% SDS). Crosslinked protein DNA complexes were reverse cross-linked with 0.3M NaCl for 4 hours at 65˚C with RNase A, then proteinase K were added to all samples. Sample DNA was purified with phenol-chloroform, precipitated with EtOH, and resuspended in T₁₀E for real-time PCR. Purified DNA was quantified using following primers (Z promoter: 5' CCGGCA AGGTGCAATGTATTAG/ 3' CATCACAGAGGGAGCTGTGTG, R promoter: 5' TGCCGGC TGCATCGGATTACT/ 3' GATGCTGATGCAGAGTCGCC, BMRF1 promoter: 5' CACTG CGGTGGAGGTAGAG/ 3' GGTGGTGTGCCATACAAGG, NECTIN1, 5' TGAGCCTGTA GGACCAGAATCA/ 3' TTTCCCACTCAAGC TGTGTCTCT) and iTaq universal SYBR green supermix (Bio-Rad) using a CFX96 touch real-time PCR detection system (Bio-Rad). Purified input DNAs were used in real-time PCR for standardization. The experiments shown are representative of two independent experiments and error bar indicating standard error of the mean within experiments.

Viral titer assay

Media containing infectious virus was harvested from CNE-2-Akata cells three days after a transfection of either R, R with ΔNp63α, or a vector control. The media was filtered with a 0.8 μm filter (Millipore-Sigma), and 10 μL of virus prep was used to infect EBV negative Akata Burkitt cells. 24 hours after infection, these cells were treated with 20 ng/ml of TPA, and 3 mM sodium butyrate. After an additional 24 hours, GFP-positive cells were counted to determine quantity of infectious virus. The experiment shown is the average of three independent viral titers, with an error bar indicating standard error of the mean, and two-sample T-test used to determine significance.

Supporting information

S1 Fig. The original western blots used to construct Fig 2A are shown. The lanes used in the blot shown in Fig 2A are labelled. Lanes not used in the final figure are indicated by a “I” symbol. (TIF)

S2 Fig. NOKs-Akata were transfected with siRNAs against ΔNp63α or a control sequence. Two days after transfection the cells were harvested for immunoblot analysis and the expression of ΔNp63α, Z, R, and tubulin was determined. (TIF)

S3 Fig. A) NOKs-Akata infected with either a lentivirus expressing ΔNp63α or a control lentivirus were differentiated on a collagen membrane or untreated for three days before harvesting for immunoblots examining the expression of ΔNp63α, Z, BMRF1, involucrin, BLIMP1,
and tubulin. B) The original western blots used to construct supplemental Fig 3A are shown; the lanes used for the figure are labeled 1 through 4, and lanes not used are indicated by a “I” symbol.

(TIF)

S4 Fig. The original western blots used to construct Fig 4A are shown in the right panel. The lanes used in the blot shown in Fig 4A are labelled 1–6 as indicated. Lanes not used in the final figure are indicated by a “I” symbol. Different western blots were used to derive the ΔNp63α and R expression levels whereas the same three western blots were used to derive the Z, BMRF1, and tubulin expression levels. “Blank” refers to lanes where no protein was loaded. Note that the same protein lysates were used in each of the western blots shown.

(TIF)

S5 Fig. A) AGS-Akata cells were transfected with a vector control or Z expression vector in the presence of absence of ΔNp63α expression vector, as indicated. Western blots were performed to examine the expression level of transfected Z and ΔNp63α proteins, R, BMRF1, and tubulin. B) The original western blots used to generate the supplemental Fig 5A are shown; lanes are numbered to indicate their position in the Fig A. Lanes not used are indicated by an “I”.

(TIF)

S6 Fig. P53KO NOKs-Akata were transfected with siRNAs targeting either ΔNp63α or a control sequence. After two days the cells were harvested for immunoblot analysis and the expression of ΔNp63α, Z, R, and tubulin was assessed.

(TIF)

S7 Fig. The original western blots used to construct Fig 7 are shown. The lanes used in the blot shown in Fig 7 are labelled. Different western blots were used to derive the ΔNp63α, p63α, and R expression levels, and another western blot was used to derive the Z, BMRF1, and tubulin expression levels. “Blank” (B) refers to lanes where no protein was loaded. Note that the same protein lysates were used in each of the western blots shown.

(TIF)

S8 Fig. A) The original western blots used to construct Fig 10A are shown. The lanes used in the blot shown in Fig 10A are labelled. Lanes not used in the final figure are indicated by a “I” symbol. B) The original western blots used to construct Fig 10C are shown on the right panel, along with Fig 10C on the left panel with lanes number 1 through 6. The lanes used in the original blots shown in Fig 10C are numbered 1 through 6. Lanes not used in the final Fig are indicated by a “I” symbol. C) Left panel: NOKs-Akata cells were transfected with siRNAs against c-myc or a control siRNA for two days, and then transfected with or without an R expression vector in the presence or absence of ΔNp63α. Immunoblot was performed one day later to examine expression of c-myc, R, ΔNp63α, BMRF1, and tubulin. Black lines indicate where irrelevant lanes in the blot were removed. Right panel: In the original western blot used to make the left panel figure, the lanes used in the figure are numbered 1 through 6, and lanes not used are indicated by a “I” symbol.

(TIF)

Acknowledgments

We would like to thank Xiaohua Su and Elsa Flores for providing the mouse p63 expression vectors, Lori Frappier for the BRLF1-pCMV3FC vector (containing a FLAG-tagged BRLF1
protein), Kathy Shair and Ben Gewurz for sending us NPC43 cells, and Janet Mertz for reviewing the manuscript.

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References
1. Khan G, Hashim MJ. Global burden of deaths from Epstein-Barr virus attributable malignancies 1990–2010. Infect Agent Cancer. 2014; 9: 38. https://doi.org/10.1186/1750-9378-9-38 PMID: 25473414
2. Young LS, Yap LF, Murray PG. Epstein–Barr virus: more than 50 years old and still providing surprises. Nat Rev Cancer. 2016; 16: 789–802. https://doi.org/10.1038/nrc.2016.92 PMID: 27687982
3. Laichalk LL, Thorley-Lawson DA. Terminal Differentiation into Plasma Cells Initiates the Replicative Cycle of Epstein-Barr Virus In Vivo. J Virol. 2005; 79: 1296–1307. https://doi.org/10.1128/JVI.79.2.1296-1307.2005 PMID: 15613356
4. Sun CC, Thorley-Lawson DA. Plasma Cell-Specific Transcription Factor XBP-1s Binds to and Trans-activates the Epstein-Barr Virus BZLF1 Promoter. J Virol. 2007; 81: 13566–13577. https://doi.org/10.1128/JVI.01055-07 PMID: 17898050
5. Bhende PM, Dickerson SJ, Sun X, Feng W-H, Kenney SC. X-Box-Binding Protein 1 Activates Lytic Epstein-Barr Virus Gene Expression in Combination with Protein Kinase D. J Virol. 2007; 81: 7363–7370. https://doi.org/10.1128/JVI.00154-07 PMID: 17494074
6. Takada K. Cross-linking of cell surface immunoglobulins induces Epstein-barr virus in Burkitt lymphoma lines. Int J Cancer. 1984; 33: 27–32. https://doi.org/10.1002/ijc.2910330106 PMID: 6319296
7. Greenspan JS, Greenspan D, Lennette ET, Abrams DI, Conant MA, Petersen V, et al. Replication of Epstein–Barr Virus within the Epithelial Cells of Oral Hairy Leukoplaikia, an AIDS-Associated Lesion. N Engl J Med. 1985; 313: 1564–1571. https://doi.org/10.1056/NEJM198512193132502 PMID: 2999595
8. Temple RM, Zhu J, Budgeon L, Christensen ND, Meyers C, Sample CE. Efficient replication of Epstein–Barr virus in stratified epithelium in vitro. Proc Natl Acad Sci U S A. 2014; 111: 16544–16549. https://doi.org/10.1073/pnas.1406818111 PMID: 25313069
9. Hadinoto V, Shapiro M, Sun CC, Thorley-Lawson DA. The Dynamics of EBV Shedding Implicate a Central Role for Epstein Cells in Amplifying Viral Output. PLOS Pathog. 2009; 5: e1000496. https://doi.org/10.1371/journal.ppat.1000496 PMID: 19578433
10. Borza CM, Hutt-Fletcher LM. Alternate replication in B cells and epithelial cells switches tropism of Epstein–Barr virus. Nat Med. 2002; 8: 594–599. https://doi.org/10.1038/nm0602-594 PMID: 12042810

11. Niedobitek G, Young LS, Lau R, Brooks L, Greenspan D, Greenspan JS, et al. Epstein–Barr Virus Infection in Oral Hairy Leukoplakia: Virus Replication in the Absence of a Detectable Latent Phase. J Gen Virol. 1991; 72: 3035–3046. https://doi.org/10.1099/0022-1317-72-12-3035 PMID: 16626955

12. Kenney SC, Mertz JE. Regulation of the latent-lytic switch in Epstein–Barr virus. Semin Cancer Biol. 2014; 26: 60–68. https://doi.org/10.1016/j.semcancer.2014.01.002 PMID: 24457012

13. Counytryman J, Miller G. Activation of expression of latent Epstein-Barr herpesvirus after gene transfer with a small cloned subfragment of heterogeneous viral DNA. Proc Natl Acad Sci. 1985; 82: 4085–4089. https://doi.org/10.1073/pnas.82.12.4085 PMID: 2987963

14. Reusch JA, Nawandar DM, Wright KL, Kenney SC, Mertz JE. Cellular Differentiation Regulator BLIMP1 Induces Epstein-Barr Virus Lytic Reactivation in Epithelial and B Cells by Activating Transcription from both the R and Z Promoters. J Virol. 2015; 89: 1731–1743. https://doi.org/10.1128/JVI.02781-14 PMID: 25410866

15. Countryman J, Miller G. Activation of expression of latent Epstein-Barr herpesvirus after gene transfer with a small cloned subfragment of heterogeneous viral DNA. Proc Natl Acad Sci. 1985; 82: 4085–4089. https://doi.org/10.1073/pnas.82.12.4085 PMID: 2987963

16. Zhang Y, Jiang C, Trudeau SJ, Narita Y, Zhao B, Teng M, et al. Histone Loaders CAF1 and HIRA Restrict Epstein-Barr Virus B-Cell Lytic Reactivation. mBio. 2020; 11. https://doi.org/10.1128/mBio.01063-20 PMID: 33109754

17. Zhang K, Lv D-W, Li R. B Cell Receptor Activation and Chemical Induction Trigger Caspase-Mediated Cleavage of Pias1 to Facilitate Epstein-Barr Virus Reactivation. Cell Rep. 2017; 21: 3445–3457. https://doi.org/10.1016/j.celrep.2017.11.071 PMID: 29263252

18. Guo R, Jiang C, Zhang Y, Govande A, Trudeau SJ, Chen F, et al. MYC Controls the Epstein-Barr Virus Lytic Switch. Mol Cell. 2020. https://doi.org/10.1016/j.molcel.2020.03.025 PMID: 32315601

19. Vogel EM, Goldbach-Mansky R, Bhaduri-McIntosh S. A promiscuous inflammasome sparks replication of a common tumor virus. Proc Natl Acad Sci U S A. 2020; 117: 1722–1730. https://doi.org/10.1073/pnas.1919133117 PMID: 31919284

20. Einsatz C, Perrigoue JG, Mertz JE. ZEB Negatively Regulates the Lytic-Switch BZLF1 Gene Promoter of Epstein-Barr Virus. J Virol. 2003; 77: 199–207. https://doi.org/10.1128/JVI.77.1.199-207.2003 PMID: 12477825

21. Adamson AL, Darr D, Holley-Guthrie E, Johnson RA, Mauser A, Swenson J, et al. Epstein-Barr Virus Immediate-Early Proteins BZLF1 and BRLF1 Activate the ATF2 Transcription Factor by Increasing the Levels of Phosphorylated p38 and c-Jun N-Terminal Kinases. J Virol. 2000; 74: 1224–1233. https://doi.org/10.1128/jvi.74.3.1224-1233.2000 PMID: 10627532

22. Ragozy T, Miller G. Autostimulation of the Epstein-Barr Virus BRLF1 Promoter Is Mediated through Consensus Sp1 and Sp3 Binding Sites. J Virol. 2001; 75: 5240–5251. https://doi.org/10.1128/JVI.75.11.5240-5251.2001 PMID: 11339061

23. Swenson JJ, Houghton PM, Kenney SC. Epstein-Barr Virus Immediate-Early Protein BRLF1 Interacts with CBP, Promoting Enhanced BRLF1 Transactivation. J Virol. 2001; 75: 6228–6234. https://doi.org/10.1128/JVI.75.13.6228-6234.2001 PMID: 11390628

24. Wille CK, Nawandar DM, Panfil AR, Ko MM, Hagemeier SR, Kenney SC. Viral Genome Methylation Differentially Affects the Ability of BZLF1 versus BRLF1 To Activate Epstein-Barr Virus Lytic Gene Expression and Viral Replication. J Virol. 2013; 87: 935–950. https://doi.org/10.1128/JVI.01790-12 PMID: 23135711
30. Bhende PM, Seaman WT, Delecluse H-J, Kenney SC. The EBV lytic switch protein, Z, preferentially binds to and activates the methylated viral genome. Nat Genet. 2004; 36: 1099–1104. https://doi.org/10.1038/ng1424 PMID: 15361873

31. Zalani S, Holley-Guthrie E, Kenney S. Epstein-Barr viral latency is disrupted by the immediate-early BRLF1 protein through a cell-specific mechanism. Proc Natl Acad Sci. 1996; 93: 9194–9199. https://doi.org/10.1073/pnas.93.17.9194 PMID: 8799177

32. Bhende PM, Seaman WT, Delecluse H-J, Kenney SC. BZLF1 Activation of the Methylated Form of the BRLF1 Immediate-Early Promoter Is Regulated by BZLF1 Residue 186. J Virol. 2005; 79: 7338–7348. https://doi.org/10.1128/JVI.79.12.7338-7348.2005 PMID: 15919888

33. Feederle R, Kost M, Baumann M, Janz A, Drouet E, Hammer schmidt W, et al. The Epstein–Barr virus lytic program is controlled by the co-operative functions of two transactivators. EMBO J. 2000; 19: 3080–3089. https://doi.org/10.1093/emboj/19.12.3080 PMID: 10856251

34. Makielski KR, Lee D, Lorenz LD, Nawandar DM, Chiu Y-F, Kenney SC, et al. Human papillomavirus promotes Epstein-Barr virus maintenance and lytic reactivation in immortalized oral keratinocytes. Virology. 2016; 495: 52–62. https://doi.org/10.1016/j.virol.2016.05.005 PMID: 27179345

35. Eichelberg MR, Welch R, Guidry JT, Ali A, Ohashi M, Makielski KR, et al. Epstein-Barr Virus Infection Promotes Epithelial Cell Growth by Attenuating Differentiation-Dependent Exit from the Cell Cycle. mBio. 2019; 10. https://doi.org/10.1128/mBio.01332-19 PMID: 31431547

36. Nawandar DM, Ohashi M, Djavadian R, Barlow E, Makielski K, Ali A, et al. Differentiation-Dependent LMP1 Expression Is Required for Efficient Lytic Epstein-Barr Virus Reactivation in Epithelial Cells. J Virol. 2017; 91: e02438–16. https://doi.org/10.1128/JVI.02438-16 PMID: 28179525

37. Caves EA, Cook SA, Lee N, Stoltz D, Watkins S, Shair KHY. Air-Liquid Interface Method To Study Epstein-Barr Virus Pathogenesis in Nasopharyngeal Epithelial Cells. mSphere. 2018; 3. https://doi.org/10.1128/mSphere.00453-17 PMID: 29134204

38. Caves EA, Butch RM, Cook SA, Wasil LR, Chen C, Di YP, et al. Latent Membrane Protein 1 Is a Novel Determinant of Epstein-Barr Virus Genome Persistence and Reactivation. mSphere. 2017; 2. https://doi.org/10.1128/mSphereDirect.00453-17 PMID: 29134204

39. Ziegler P, Tian Y, Bai Y, Abrahamsson S, Bäckerholm A, Reznik AS, et al. A primary nasopharyngeal three-dimensional air-liquid interface cell culture model of the pseudostratified epithium reveals differential donor- and cell type-specific susceptibility to Epstein-Barr virus infection. PLOS Pathog. 2021; 17: e1009041. https://doi.org/10.1371/journal.ppat.1009041 PMID: 33914843

40. Chiang M-F, Yang S-Y, Lin I-Y, Hong J-B, Lin S-J, Ying H-Y, et al. Inducible deletion of the Blimp-1 gene in adult epidermis causes granulocyte-dominated chronic skin inflammation in mice. Proc Natl Acad Sci. 2013; 110: 356–360. https://doi.org/10.1073/pnas.1219462110 PMID: 23576729

41. Segre JA, Bauer C, Fuchs E. Klf4 is a transcription factor required for establishing the barrier function of the skin. Nat Genet. 1999; 22: 356–360. https://doi.org/10.1038/11926 PMID: 10431239

42. Crum CP, McKeon FD. p63 in Epithelial Survival, Germ Cell Surveillance, and Neoplasia. Annu Rev Pathol Mech Dis. 2010; 5: 349–371. https://doi.org/10.1146/annurev-pathol-121808-102117 PMID: 20078223

43. Pellegrini G, Dellambra E, Golisano O, Martinelli E, Fantozzi I, Bondanza S, et al. p63 identifies keratinocyte stem cells. Proc Natl Acad Sci. 2001; 98: 3156–3161. https://doi.org/10.1073/pnas.061032098 PMID: 11248048

44. Parsa R, Yang A, McKeon F, Green H. Association of p63 with Proliferative Potential in Normal and Neoplastic Human Keratinocytes. J Invest Dermatol. 1999; 113: 1099–1105. https://doi.org/10.1046/j.1523-1747.1999.00780.x PMID: 10594758

45. Ince TA, Cviko AP, Quade BJ, Yang A, McKeon FD, Mutter GL, et al. p63 Coordinates Anogenital Modeling and Epithelial Cell Differentiation in the Developing Female Urogenital Tract. Am J Pathol. 2002; 161: 1111–1117. https://doi.org/10.1016/S0002-9440(10)64387-8 PMID: 12368184

46. Yang A, Schweitzer R, Sun D, Kaghad M, Walker N, Bronson RT, et al. p63 is essential for regenerative proliferation in limb, craniofacial and epithelial development. Nature. 1999; 398: 714–718. https://doi.org/10.1038/19539 PMID: 10227294

47. Keyes WM, Wu Y, Vogel H, Guo X, Lowe SW, Mills AA. p63 deficiency activates a program of cellular senescence and leads to accelerated aging. Genes Dev. 2005; 19: 1986–1999. https://doi.org/10.1101/gad.342305 PMID: 16107615

48. Celli J, Duif P, Hamel BJC, Bamshad M, Kramer B, Smits APT, et al. Heterozygous Germline Mutations in the p53 Homolog p63 Are the Cause of EEC Syndrome. Cell. 1999; 99: 143–153. https://doi.org/10.1016/s0092-8674(00)81646-3 PMID: 10535733
49. Yang A, Zhu Z, Kapranov P, McKeon F, Church GM, Gingeras TR, et al. Relationships between p63 Binding, DNA Sequence, Transcription Activity, and Biological Function in Human Cells. Mol Cell. 2006; 24: 593–602. https://doi.org/10.1016/j.molcel.2006.10.018 PMID: 17188034

50. Kouwenhoven EN, van Heeringen SJ, Tena JJ, Otí M, Dutilha BE, Alonso ME, et al. Genome-Wide Profiling of p63 DNA–Binding Sites Identifies an Element that Regulates Gene Expression during Limb Development in the 7q21 SHFM1 Locus. PLoS Genet. 2010; 6. https://doi.org/10.1371/journal.pgen.1001065 PMID: 20808887

51. Flores ER, Tsai KY, Crowley D, Sengupta S, Yang A, McKeon F, et al. p63 and p73 are required for p53-dependent apoptosis in response to DNA damage. Nature. 2002; 416: 560–564. https://doi.org/10.1038/416560a PMID: 11932750

52. Crook T, Nicholls JM, Brooks L, O’Nions J, Allday MJ. High level expression of ΔN-p63: a mechanism for the inactivation of p53 in undifferentiated nasopharyngeal carcinoma (NPC)? Oncogene. 2000; 19: 3439–3444. https://doi.org/10.1038/sj.onc.1203656 PMID: 10918601

53. Botchkarev VA, Flores ER. p53/p63/p73 in the Epidermis in Health and Disease. Cold Spring Harb Perspect Med. 2014; 4: a015248. https://doi.org/10.1101/cshperspect.a015248 PMID: 25085956

54. Botchkarev VA, Flores ER, Kouwenhoven EN, Dotto GP, Restivo G, Nguyen B-C, et al. Cooperation between the transcription factors p63 and IRF6 is essential to prevent cleft palate in mice. J Clin Invest. 2010; 120: 1561–1569. https://doi.org/10.1172/JCI40266 PMID: 20424327

55. Moretti F, Marinari B, Iacono NL, Botti E, Giunta A, Spallone G, et al. A regulatory feedback loop involving p63 and IRF6 links the pathogenesis of 2 genetically different human ectodermal dysplasias. J Clin Invest. 2010; 120: 1570–1577. https://doi.org/10.1172/JCI40267 PMID: 20424325

56. Liu C, Sista ND, Pagano JS. Activation of the Epstein-Barr virus DNA polymerase promoter by the BRLF1 immediate-early protein is mediated through USF and E2F. J Virol. 1996; 70: 2545–2555. https://doi.org/10.1128/JVI.70.4.2545-2555.1996 PMID: 8642684

57. Robinson AR, Kwek SS, Hagemeier SR, Willie CK, Kenney SC. Cellular Transcription Factor Oct-1 Interacts with the Epstein-Barr Virus BRLF1 Protein To Promote Disruption of Viral Latency. J Virol. 2011; 85: 8940–8953. https://doi.org/10.1128/JVI.00569-11 PMID: 21974976

58. Holley-Guthrie EA, Quinlivan EB, Mar EC, Kenney S. The Epstein-Barr virus (EBV) BRLF1 promoter for early antigen (EA-D) is regulated by the EBV transactivators, BRLF1 and BZLF1, in a cell-specific manner. J Virol. 1990; 64: 3753–3759. https://doi.org/10.1128/JVI.64.8.3753-3759.1990 PMID: 2164595

59. Quinlivan EB, Holley-Guthrie E, Norris M, Gutsch D, Bachenheimer SL, Kenney SC. Direct BRLF1 binding is required for cooperative BZLF1/BRLF1 activation of the Epstein-Barr virus early promoter, BRLF1. Nucleic Acids Res. 1993; 21: 1999–2007. https://doi.org/10.1093/nar/21.8.1999 PMID: 8395364

60. Gruffat H, Duran N, Buisson M, Wild F, Buckland R, Sergeant A. Characterization of an R-binding site mediating the R-induced activation of the Epstein-Barr virus BMLF1 promoter. J Virol. 1992; 66: 46–52. https://doi.org/10.1128/JVI.66.1.46-52.1992 PMID: 1309256
68. Adamson AL, Kenney SC. Rescue of the Epstein–Barr Virus BZLF1 Mutant, Z(S186A), Early Gene Activation Defect by the BRLF1 Gene Product. Virology. 1998; 251: 187–197. https://doi.org/10.1006/viro.1998.9396 PMID: 9813214

69. Su X, Cho MS, Gi Y-J, Ayanga BA, Sherr CJ, Flores ER. Rescue of key features of the p63-null epithelial phenotype by inactivation of Ink4a and Arf. EMBO J. 2009; 28: 1904–1915. https://doi.org/10.1038/emboj.2009.151 PMID: 19494829

70. Hagemeier SR, Barlow EA, Kleman AA, Kenney SC. The Epstein-Barr Virus BRRF1 Protein, Na, Induces Lytic Infection in a TRAF2- and p53-Dependent Manner. J Virol. 2011; 85: 4318–4329. https://doi.org/10.1128/JVI.01856-10 PMID: 21325409

71. Hagemeier SR, Barlow EA, Meng Q, Kenney SC. The Cellular Ataxia Telangiectasia-Mutated Kinase Promotes Epstein-Barr Virus Lytic Reactivation in Response to Multiple Different Types of Lytic Reactivation-Inducing Stimuli. J Virol. 2012; 86: 13360–13370. https://doi.org/10.1128/JVI.01850-12 PMID: 23015717

72. Chua H-H, Chiu H-Y, Lin S-J, Weng P-L, Lin J-H, Wu S-W, et al. p53 and Sp1 cooperate to regulate the expression of epstein–barr viral Zta protein. J Med Virol. 2012; 84: 1279–1288. https://doi.org/10.1002/jmv.23316 PMID: 22711357

73. Kraus RJ, Cordes BA, Sathiamoorthi S, Patel P, Yuan X, Iempridee T, et al. Reactivation of Epstein-Barr Virus by HIF-1α Requires p53. J Virol. 2020; 94. https://doi.org/10.1128/JVI.00722-20 PMID: 32641480

74. Westfall MD, Mays DJ, Sniezek JC, Pietenpol JA. The σσ promoters In Vivo and Has Transcriptional Repressor Activity That Is Reduced by Hay-Wells Syndrome-Derived Mutations. Mol Cell Biol. 2003; 23: 2264–2276. https://doi.org/10.1128/MCB.23.7.2264-2276.2003 PMID: 12640112

75. Mollo Maria Rosaria Antonini Dario, Karen Mitchell, Paola Fortugno, Antonio Costanzo, Jill Dixon, et al. p63-dependent and independent mechanisms of nectin-1 and nectin-4 regulation in the epidermis. Exp Dermatol. 2015; 24: 114–119. https://doi.org/10.1111/exd.12593 PMID: 25387952

76. Mangiulli M, Valletti A, Caratozzolo MF, Tullo A, Sbisà E, Pesole G, et al. Identification and functional characterization of two new transcriptional variants of the human p63 gene. Nucleic Acids Res. 2009; 37: 6092–6104. https://doi.org/10.1093/nar/gkp674 PMID: 19700772

77. Su X, Chakravarti D, Flores ER. p63 steps into the limelight: crucial roles in the suppression of tumorigenesis and metastasis. Nat Rev Cancer. 2013; 13: 136–143. https://doi.org/10.1038/nrc3446 PMID: 23344544

78. Sethi I, Romano R-A, Gluck C, Smalley K, Vojtesek B, Buck MJ, et al. A global analysis of the complex landscape of isoforms and regulatory networks of p63 in human cells and tissues. BMC Genomics. 2015; 16: 584. https://doi.org/10.1186/s12864-015-1793-9 PMID: 26251276

79. Fukushima N, Satoh T, Sueoka N, Sato A, Ide M, Hisatomi T, et al. Clinico-pathological characteristics of p63 expression in B-cell lymphoma. Cancer Sci. 2006; 97: 1050–1055. https://doi.org/10.1111/j.1349-7006.2006.00284.x PMID: 16918993

80. Zou H, Zou R, Chen K, Zhu C, Tian X, You Y, et al. miR-129 targets CDK1 and iASPP to modulate Burkitt lymphoma cell proliferation in a Tap63-dependent manner. J Cell Biochem. 2018; 119: 9217–9228. https://doi.org/10.1002/jcb.27189 PMID: 30105797

81. Hedvat CV, Teruya-Feldstein J, Puig P, Capodici P, Dudas M, Pica N, et al. Expression of p63 in Diffuse Large B-Cell Lymphoma. Appl Immunohistochem Mol Morphol. 2005; 13: 237–242. https://doi.org/10.1097/01.pai.0000142169.52670.ce PMID: 16082248

82. Schultz J, Bork P, Ponting CP, Hofmann K. SAM as a protein interaction domain involved in developmental regulation. Protein Sci. 1997; 6: 249–253. https://doi.org/10.1002/pro.5560060128 PMID: 9007998

83. Feng W, Kraus RJ, Dickerson SJ, Lim HJ, Jones RJ, Yu X, et al. ZEB1 and c-Jun Levels Contribute to the Establishment of Highly Lycy Epstein-Barr Virus Infection in Gastric AGS Cells. J Virol. 2007; 81: 10113–10122. https://doi.org/10.1128/JVI.00692-07 PMID: 17629078

84. Hove E, Ville C, Hagemeier S, Kenney S, Greijer A, Middeldorp J. Epstein–Barr Virus Gene BARF1 Expression Is Regulated by the Epithelial Differentiation Factor ΔΔNp63a in Undifferentiated Nasopharyngeal Carcinoma. Cancers. 2018; 10: 76. https://doi.org/10.3390/cancers10030076 PMID: 29562599

85. Lin Z, Yin Q, Flemington E. Identification of a Negative Regulatory Element in the Epstein-Barr Virus Zta Transactivation Domain That Is Regulated by the Cell Cycle Control Factors c-Myc and E2F1. J Virol. 2004; 78: 11962–11971. https://doi.org/10.1128/JVI.78.21.11962-11971.2004 PMID: 15479836

86. Wang L, Xia W, Chen H, Xiao Z-X. ΔΔNp63α modulates phosphorylation of p38 MAP kinase in regulation of cell cycle progression and cell growth. Biochem Biophys Res Commun. 2019; 509: 784–789. https://doi.org/10.1016/j.bbr.2018.12.185 PMID: 30635119
87. O’Neil JD, Ammit AJ, Clark AR. MAPK p38 regulates inflammatory gene expression via tristetraprolin: Doing good by stealth. Int J Biochem Cell Biol. 2018; 94: 6–9. https://doi.org/10.1016/j.biocel.2017.11.003 PMID: 29128684

88. Soni S, Anand P, Padwad YS. MAPKAPK2: the master regulator of RNA-binding proteins modulates transcript stability and tumor progression. J Exp Clin Cancer Res. 2019; 38: 121. https://doi.org/10.1186/s13046-019-1115-1 PMID: 30850014

89. Feng W, Israel B, Raab-Traub N, Busson P, Kenney SC. Chemotherapy Induces Lytic EBV Replication and Confers Ganciclovir Susceptibility to EBV-positive Epithelial Cell Tumors. Cancer Res. 2002; 62: 1920–1926. PMID: 11912175

90. Feng W, Hong G, Delecluse H-J, Kenney SC. Lytic Induction Therapy for Epstein-Barr Virus-Positive B-Cell Lymphomas. J Virol. 2004; 78: 1893–1902. https://doi.org/10.1128/jvi.78.4.1893-1902.2004 PMID: 14747554

91. Fomenkov A, Zangen R, Huang Y-P, Osada M, Guo Z, Fomenkov T, et al. RACK1 and Stratifin Target ΔNp63α for a Proteasome Degradation in Head and Neck Squamous Cell Carcinoma Cells upon DNA Damage. Cell Cycle. 2004; 3: 1285–1295. https://doi.org/10.4161/cc.3.10.1155 PMID: 15467455

92. Kenney S, Holley-Guthrie E, Mar EC, Smith M. The Epstein-Barr virus BMLF1 promoter contains an enhancer element that is responsive to the BZLF1 and BRLF1 transactivators. J Virol. 1989; 63: 3878–3883. https://doi.org/10.1128/JVI.63.9.3878-3883.1989 PMID: 2548003

93. Chen Y, Li Y, Peng Y, Zheng X, Fan S, Yi Y, et al. ΔNp63α down-regulates c-Myc modulator MM1 via E3 ligase HERC3 in the regulation of cell senescence. Cell Death Differ. 2018; 25: 2118–2129. https://doi.org/10.1038/s41418-018-0132-5 PMID: 29880857

94. Konishi N, Narita Y, Hijioka F, Masud HMAA, Sato Y, Kimura H, et al. BGLF2 Increases Infectivity of Epstein-Barr Virus by Activating AP-1 upon De Novo Infection. mSphere. 3: e00138–18. https://doi.org/10.1128/mSphere.00138-18 PMID: 29695622

95. Liu Y-R, Huang S-Y, Chen J-Y, Wang LH-C. Microtubule depolymerization activates the Epstein-Barr virus lytic cycle through protein kinase C pathways in nasopharyngeal carcinoma cells. J Gen Virol. 2013; 94: 2750–2758. https://doi.org/10.1099/jvi.0.058040-0 PMID: 24062531

96. Huang S-Y, Fang C-Y, Wu C-C, Tsai C-H, Lin S-F, Chen J-Y. Reactive oxygen species mediate Epstein-Barr virus reactivation by N-methyl-N'-nitro-N-nitrosoguanidine. PloS One. 2013; 8: e84919. https://doi.org/10.1371/journal.pone.0084919 PMID: 24367863

97. Gao X, Wang H, Sairenji T. Inhibition of Epstein-Barr virus (EBV) reactivation by short interfering RNAs targeting p38 mitogen-activated protein kinase or c-myc in EBV-positive epithelial cells. J Virol. 2004; 78: 11798–11806. https://doi.org/10.1128/JVI.78.21.11798-11806.2004 PMID: 15479821

98. Han J, Wu J, Silke J. An overview of mammalian p38 mitogen-activated protein kinases, central regulators of cell stress and receptor signaling. F1000Research. 2020; 9: F1000 Faculty Rev-653. https://doi.org/10.12688/f1000research.22092.1 PMID: 32612808

99. Lee J, Sun C, Zhou Y, Lee J, Gokaipal D, Herrera H, et al. p38 MAPK-mediated regulation of Xbp1s is crucial for glucose homeostasis. Nat Med. 2011; 17: 1251–1260. https://doi.org/10.1038/nm.2449 PMID: 21892182

100. Klein AM, Zaganae Jr, Cobb MH. Chromatin-tethered MAPKs. Curr Opin Cell Biol. 2013; 25: 272–277. https://doi.org/10.1016/jceb.2013.01.002 PMID: 23430467

101. Gehani SS, Agrawal-Singh S, Dietrich N, Christophersen NS, Helin K, Hansen K. Polycomb Group Protein Displacement and Gene Activation through MSK-Dependent H3K27me3S28 phosphorylation. Mol Cell. 2010; 39: 886–900. https://doi.org/10.1016/j.molcel.2010.08.020 PMID: 20864036
106. Efimova T, LaCelle P, Welter JF, Eckert RL. Regulation of Human Involucrin Promoter Activity by a Protein Kinase C, Ras, MEKK1, MEK3, p38/RK, AP1 Signal Transduction Pathway. J Biol Chem. 1998; 273: 24387–24395. doi:10.1074/jbc.273.38.24387 PMID: 9733728

107. Nickless A, Cheruiyot A, Flanagan KC, Piwnica-Worms D, Stewart SA, You Z. p38 MAPK inhibits nonsense-mediated RNA decay in response to persistent DNA damage in noncycling cells. J Biol Chem. 2017; 292: 15266–15276. doi:10.1074/jbc.M117.787846 PMID: 28765281

108. Gent M van, Reich A, Velu SE, Gack MU. Nonsense-mediated decay controls the reactivation of the oncogenic herpesviruses EBV and KSHV. PLOS Biol. 2021; 19: e3001097. doi:10.1371/journal.pbio.3001097 PMID: 33596193

109. Glaser R, Zhang HY, Yao KT, Zhu HC, Wang FX, Li GY, et al. Two epithelial tumor cell lines (HNE-1 and HONE-1) latently infected with Epstein-Barr virus that were derived from nasopharyngeal carcinomas. Proc Natl Acad Sci. 1989; 86: 9524–9528. doi:10.1073/pnas.86.23.9524 PMID: 2556716

110. Strong MJ, Baddoo M, Nanbo A, Xu M, Puettner A, Lin Z. Comprehensive High-Throughput RNA Sequencing Analysis Reveals Contamination of Multiple Nasopharyngeal Carcinoma Cell Lines with HeLa Cell Genomes. J Virol. 2014; 88: 10696–10704. doi:10.1128/JVI.01457-14 PMID: 24991015

111. Molesworth SJ, Lake CM, Porzio CM, Turk SM, Hutt-Fletcher LM, Epstein-Barr Virus gH Is Essential for Penetration of B Cells but Also Plays a Role in Attachment of Virus to Epithelial Cells. J Virol. 2000; 74: 6324–6332. doi:10.1128/JVI.74.14.6324-6332.2000 PMID: 10864642

112. Shimizu N, Yoshiyama H, Takada K. Clonal propagation of Epstein-Barr virus (EBV) recombinants in EBV-negative Akata cells. J Virol. 1996; 70: 7260–7263. doi:10.1128/JVI.70.10.7260-7263.1996 PMID: 8794379

113. Guasparri I, Bubman D, Cesarman E. EBV LMP2A affects LMP1-mediated NF-κB signaling and survival of lymphoma cells by regulating TRAF2 expression. Blood. 2008; 111: 3813–3820. doi:10.1182/blood-2007-03-080309 PMID: 18230756

114. Romero-Masters JC, Huebner SM, Ohashi M, Bristol JA, Benner BE, Barlow EA, et al. B cells infected with Type 2 Epstein-Barr virus (EBV) have increased NFATc1/NFATc2 activity and enhanced lytic gene expression in comparison to Type 1 EBV infection. PLOS Pathog. 2020; 16: e1008365. doi:10.1371/journal.ppat.1008365 PMID: 32059024

115. Sciver NV, Ohashi M, Pauly NP, Bristol JA, Nelson SE, Johannsen EC, et al. Hippo signaling effectors YAP and TAZ induce Epstein-Barr Virus (EBV) reactivants in EBV-negative Akata cells. J Virol. 2019; 93: 7260–7263. doi:10.1128/JVI.01079-19 PMID: 30711843

116. Sarisky RT, Gao Z, Fixman ED, Haywood GS, Hayden SD. A replication function associated with the activation domain of the Epstein-Barr virus Zta transactivator. J Virol. 1996; 70: 8340–8347. doi:10.1128/JVI.70.12.8340-8347.1996 PMID: 8970953

117. Cruz-Herrera CFDL, Shire K, Siddiqi UZ, Frappier L. A genome-wide screen of Epstein-Barr virus proteins that modulate host SUMOylation identifies a SUMO E3 ligase conserved in herpesviruses. PLOS Pathog. 2018; 14: e1007176. doi:10.1371/journal.ppat.1007176 PMID: 29979787

118. Ohashi M, Hayes M, McChesney K, Johannsen E. Epstein-Barr virus nuclear antigen 3C (EBNA3C) interacts with the metabolism sensing C-terminal binding protein (CtBP) repressor to upregulate host genes. PLOS Pathog. 2021; 17: e1009419. doi:10.1371/journal.ppat.1009419 PMID: 33720992