Evidence linking APOBEC3B genesis and evolution of innate immune antagonism by gamma-herpesvirus ribonucleotide reductases

Short title: APOBEC3B antagonism by gamma-herpesviruses

Sofia N. Moraes¹, Jordan T. Becker¹, Seyed Arad Moghadasi¹, Nadine M. Shaban¹, Ashley A. Auerbach¹,2, Adam Z. Cheng¹, Reuben S. Harris¹,2,3,*

¹ Department of Biochemistry, Molecular Biology, and Biophysics, Institute for Molecular Virology, Masonic Cancer Center, University of Minnesota, Minneapolis, MN 55455, USA.
² Department of Biochemistry and Structural Biology, University of Texas Health San Antonio, San Antonio, TX 78229, USA.
³ Howard Hughes Medical Institute, University of Texas Health San Antonio, San Antonio, TX 78229, USA.

* Corresponding author. Email: rsh@uthscsa.edu

Key words: APOBEC3B; evolutionary arms race; herpesvirus; host-pathogen interaction; innate immunity; virus restriction factor; ribonucleotide reductase; virus evolution
Abstract

Viruses have evolved diverse mechanisms to antagonize host immunity such as direct inhibition and relocalization of cellular APOBEC3B (A3B) by the ribonucleotide reductase (RNR) of Epstein-Barr virus. Here, we investigate the mechanistic conservation and evolutionary origin of this innate immune counteraction strategy. First, we find that human gamma-herpesvirus RNRs engage A3B via largely distinct surfaces. Second, we show that RNR-mediated enzymatic inhibition and relocalization of A3B depend upon binding to different regions of the catalytic domain. Third, we show that the capability of viral RNRs to antagonize A3B is conserved among gamma-herpesviruses that infect humans and Old World monkeys that encode this enzyme but absent in homologous viruses that infect New World monkeys that naturally lack the A3B gene. Finally, we reconstruct the ancestral primate A3B protein and demonstrate that it is active and similarly engaged by the RNRs from viruses that infect humans and Old World monkeys but not by the RNRs from viruses that infect New World monkeys. These results combine to indicate that the birth of A3B at a critical branchpoint in primate evolution may have been a driving force in selecting for an ancestral gamma-herpesvirus with an expanded RNR functionality through counteraction of this antiviral enzyme.
Introduction

Cell-intrinsic innate immune proteins, or ‘restriction factors’, contribute to the first line of defenses against incoming viruses (reviewed by [1]). A prototypical example is the APOBEC3 (A3) family of single-stranded (ss)DNA cytosine deaminases, which provide broad and overlapping protection against viral infections (reviewed by [2, 3]). Classically, A3s exert antiviral activity by catalyzing cytosine deamination in exposed ssDNA replication intermediates, often resulting in lethal mutagenesis of viral genomes. Humans encode seven A3 enzymes (A3A-D, A3F-H), which have been implicated in the restriction of retroviruses (HIV and HTLV) [4-8], endogenous retroviruses (HERV) [9, 10], hepadnaviruses (HBV) [11, 12], papillomaviruses (HPV) [13], polyomaviruses (BK) [14], and most recently the gamma-herpesvirus Epstein-Barr virus (EBV [15]). It is difficult to predict which subset of A3 enzymes has the potential to restrict a given virus, but differences in the subcellular localization properties of these enzymes provide useful insights into their antiviral targets. For instance, cytoplasmic A3s like A3G and A3F potently restrict HIV-1 by binding to viral genomic RNAs in the cytoplasm, packaging into budding virions at the cell membrane, and deaminating viral ssDNA intermediates during reverse transcription [16-18]. It thus follows that A3B—the only constitutively nuclear human A3 enzyme—may pose a threat to the genomic integrity of viruses that undergo replication in the nucleus, such as EBV and related herpesviruses.

Viruses that are susceptible to restriction by cellular A3 enzymes have evolved equally potent counteraction mechanisms to ensure successful replication (reviewed by [2, 3, 19]). The most well-studied A3 antagonist is the lentiviral accessory protein Vif, which nucleates the formation of an E3 ubiquitin ligase complex to promote the selective degradation of up to five different cytoplasmic A3 enzymes: A3C, A3D, A3F, A3G, and A3H (reviewed by [2, 20, 21]).
Extensive mutagenesis analyses revealed that HIV-1 Vif evolved to use distinct binding surfaces in order to recognize different A3s, as underscored by separation-of-function mutants of Vif that selectively degrade only a subset of these enzymes (reviewed by [20, 22, 23]). These studies have been essential to our understanding of how viruses adapt to overcome innate host defenses and, collectively, highlight the importance of elucidating structure-function relationships between proteins at the host-pathogen interface.

HIV-1 and related retroviruses are thought to be primary targets of A3 enzymes due to obligate ssDNA replication intermediates, abundant evidence for A3-catalyzed hypermutation of viral genomes, and a potent A3 counteraction mechanism. Reports of DNA viruses exhibiting signatures of A3-mediated mutagenesis suggest that these viruses may also be susceptible to restriction by A3 enzymes [2, 24], although the identification of viral antagonists has been challenging. We recently discovered that the large double-stranded (ds)DNA herpesvirus EBV uses a two-pronged approach to counteract the antiviral activity of A3B. The large subunit of the EBV ribonucleotide reductase (RNR), BORF2, directly inhibits A3B catalytic activity and relocates this enzyme from the nucleus to cytoplasmic aggregates—most likely to protect viral genomes from deamination during lytic replication [15]. Consistent with this interpretation, BORF2-null EBV accumulates A3B signature C/G to T/A mutations upon lytic reactivation and exhibits significantly lower infectivity [15].

The deamination activity of all A3 family enzymes requires a zinc-coordinating motif, which forms the core of the catalytic site and is surrounded by three loop regions (L1, L3, and L7) that contribute to binding of ssDNA substrates [25-27]. These loops vary in size and amino acid composition and account for differences in A3 catalytic rates and local target sequence preferences (reviewed by [28, 29]). The neutralization of A3B by EBV BORF2 is the first
example of direct inhibition of A3 ssDNA deaminase activity by another protein, and our recently published cryo-EM structure of EBV BORF2 in complex with the catalytic domain of A3B (A3Bctd) provides critical insights into the structural basis and molecular mechanism of this unanticipated host-pathogen interaction [30]. Specifically, our cryo-EM studies revealed that EBV BORF2 binds to the L1 and L7 regions of A3Bctd, resulting in a high-affinity interaction that effectively blocks the A3B active site from binding to ssDNA substrates and catalyzing deamination. Recently, we and others have shown that, in addition to EBV BORF2, the large RNR subunit from related human herpesviruses can also bind and relocalize A3B—as well as the highly similar A3A enzyme—suggesting that this A3 counteraction mechanism may be conserved evolutionarily [31-33].

Here, we investigate the molecular and functional conservation of A3-antagonism by the RNR subunits of different herpesviruses using structurally and evolutionarily informed approaches. We find that the RNR subunits of EBV and the related human gamma-herpesvirus Kaposi’s sarcoma-associated herpesvirus (KSHV) interact with A3B and A3A by binding to distinct loop regions near the catalytic site, which accounts for differences in binding selectivity and capacity to inhibit A3 enzymatic activity. We also uncover potential coevolutionary interactions between A3 enzymes and primate gamma-herpesviruses by discovering a link between the generation of the A3B gene in primates and the evolution of A3-neutralization functionalities in viral RNRs. Last, we reconstruct an ancestral primate A3B enzyme and show that it is bound, relocalized, and inhibited by the RNRs of gamma-herpesviruses that infect primate species that encode A3B but not by the RNRs from primate viruses that infect species that naturally lack this gene. Taken together, our data provide evidence of a conserved and likely ancient A3 antagonism mechanism in primate gamma-herpesviruses and offer insights into how
genetic conflicts with host restriction factors may have affected—and likely continue to affect—the structure and function of viral proteins such as the herpesvirus RNR.

**Results**

**Structural differences and similarities between EBV and KSHV RNR large subunits**

Our recent cryo-EM studies of EBV BORF2-A3B complexes revealed a large binding surface composed of multiple structural elements from each protein [30]. Specifically, loops 1 (L1) and 7 (L7) of the A3B catalytic domain (A3Bctd) are sequestered by multiple interactions with BORF2 including contacts with residues L133 and Y134 of a novel short helix insertion (SHI) and residues R484 and Y481 of an additional helical loop structure (HLS). Y134 additionally forms a network of stabilizing interactions with other BORF2 residues including Q476 and R484 ([30]; depicted in **Figure 1A**). The BORF2 homolog from related gamma-herpesvirus KSHV (ORF61) has also been shown to bind A3B [15, 32]. However, an amino acid sequence alignment of EBV BORF2 and KSHV ORF61 indicates limited identity in the SHI and HLS regions (**Figure 1B**; complete sequence alignment in **Figure 1—figure supplement 1A**). We therefore used RoseTTAFold [34] to generate a structural model of KSHV ORF61 for 3-dimensional comparisons (**Figure 1C**). Although sharing only ~42% primary amino acid identity, the predicted model of KSHV ORF61 overlays well with the cryo-EM structure of EBV BORF2 with root mean square deviation (rmsd) of 1.26 Å. Interestingly, in addition to similar HLS regions, EBV BORF2 and the predicted structure of KSHV ORF61 share a similar sized SHI region (depicted in red in **Figure 1C**), which is lacking in nonviral class Ia RNRs such as those from *E. coli* and humans [30].
To gain further insights into the potential role of KSHV ORF61 SHI and HLS regions in binding to A3B, we used *in silico* docking to investigate how ORF61 and A3Bctd might interact. RoseTTAFold models of KSHV ORF61 and A3Bctd were aligned to the cryo-EM BORF2-A3Bctd complex using PyMOL [35] to create an initial docking pose followed by local protein-protein docking using RosettaDock 4.0 [36, 37]. Models were ranked by CAPRI-score and the best structure was selected based on overall interface energy score (Methods and Figure 1—figure supplement 1B). Several high-scoring models suggested that the ORF61 SHI region may interact with the loop 3 (L3) region of A3B, instead of L1 and L7 as for BORF2 (Figure 1D-E). Additionally, the predicted HLS region of KSHV ORF61 appears to be positioned further away from the RNR-A3Bctd binding interface and may not contribute to the interaction (Figure 1E). Taken together, these *in silico* predictions suggest that EBV BORF2 and KSHV ORF61 may differentially engage the catalytic domain of A3B by interacting with distinct loop regions. This possibility is tested in the following sections.

**Involvement of the A3B and A3A loop 7 regions in binding to gamma-herpesvirus RNRs**

Recent biochemical and structural studies demonstrated that direct contacts between EBV BORF2 and the L7 region of A3Bctd are essential for this host-pathogen interaction [15, 30]. A3Bctd is closely related to A3A (>90% amino acid identity including identical L7 regions; Figure 2A), suggesting that a similar interaction with L7 could explain EBV BORF2 ability to engage A3B and A3A, but not other A3 family members. Recent work also revealed that KSHV ORF61 is similarly capable of binding and relocalizing A3B and A3A, but not other A3 enzymes [32].
To test whether binding to L7 is a conserved requirement for viral RNR-A3 interaction, we generated a series of chimeric A3 proteins with reciprocal L7 swaps and tested their ability to bind to EBV BORF2 and KSHV ORF61 in co-immunoprecipitation (co-IP) experiments (Figure 2B). 293T cells were transfected with FLAG-tagged viral RNRs and EGFP-tagged A3s, subjected to anti-FLAG affinity purification, and analyzed by immunoblotting. As expected from our prior studies, replacing L7 of A3Bctd or A3A with L7 of A3Gctd completely abolishes interaction with EBV BORF2 (Figure 2B, left panel, lanes 1-4) and, reciprocally, replacing L7 of A3Gctd with L7 of A3A/Bctd endows binding (Figure 2B, left panel, lanes 5-6). However, we were surprised to find that replacing L7 of A3Bctd or A3A with L7 of A3Gctd does not affect binding by KSHV ORF61 (Figure 2B, right panel, lanes 1-4). Accordingly, replacing L7 of A3Gctd with L7 of A3A/Bctd does not enable pull-down by KSHV ORF61 (Figure 2B, right panel, lanes 5-6)

An additional defining feature of the interaction between A3B/A and viral RNRs is relocalization from the nucleus to the cytoplasm [15, 31, 32]. To evaluate the subcellular localization phenotypes of the L7 chimeras described above, we performed quantitative fluorescence microscopy experiments using HeLa cells co-transfected with EGFP-tagged A3s and FLAG-tagged viral RNRs (Figure 2C, representative images). In agreement with our co-IP results, replacing L7 of A3Bctd or A3A with L7 of A3Gctd markedly decreases EBV BORF2-mediated relocalization, although low levels of A3Bctd L7G relocalization can still be detected in a subset of cells. Reciprocally, replacing L7 of A3Gctd with L7 of A3A/Bctd enables effective relocalization (Figure 2C, representative images; Figure 2D-E, quantification). In further agreement with our co-IP results, swapping L7 does not decrease KSHV ORF61-mediated relocalization of A3Bctd or A3A, nor does it increase A3Gctd relocalization (Figure 2C,
representative images; Figure 2D-E, quantification). Collectively, our experimental data support
the in silico protein docking predictions and indicate that, despite an overall high degree of
predicted structural similarity, the RNR subunits from EBV and KSHV engage A3Bctd and A3A
through at least partially distinct surfaces.

Role of A3B and A3A loop 3 in binding to gamma-herpesvirus RNRs

L3 is one of three loops that surround the A3 active site (Figure 2A). In our recent cryo-
EM studies of the EBV BORF2-A3Bctd complex, we were unable to model A3Bctd L3 residues
due to the flexibility of this region ([30]; dashed region in Figure 3A). In silico protein-protein
docking predictions in Figure 1D-E suggest that this loop may be important for KSHV ORF61
binding. Therefore, to examine whether L3 participates in viral RNR-A3 interactions, we
performed co-IP experiments using a panel of chimeric A3 proteins with reciprocal L3 swaps
(Figure 3B).

Once again, we observed markedly distinct phenotypes in co-IPs with EBV BORF2 and
KSHV ORF61. For EBV BORF2, similar pull-down levels are observed for WT versus L3-
swapped A3s (Figure 3B, left panel, lanes 1-4), whereas replacing L3 of A3Bctd or A3A with
L3 of A3Gctd nearly abolishes the interaction with KSHV ORF61 (Figure 3B, right panel, lanes
1-4). Additionally, replacing L3 of A3Gctd with L3 of A3A/Bctd is sufficient to promote modest
immunoprecipitation with KSHV ORF61 (Figure 3B, right panel, lanes 5-6). These results
indicate that the L3 region is critical for KSHV ORF61 binding and likely to have little or no role
in binding to EBV BORF2.

To evaluate the subcellular localization phenotypes of the L3 chimeras described above,
we performed quantitative fluorescent microscopy experiments using HeLa cells (Figure 3C,
representative images). Similar to our L7 results, microscopy observations are largely concordant with co-IP phenotypes. For EBV BORF2, L3 swaps have no substantial effect on A3 relocalization, whereas replacing L3 of A3Bctd or A3A with the corresponding region of A3Gctd causes a marked decrease in relocalization by KSHV ORF61 (Figure 3C, representative images; Figure 3D-E, quantification). Notably, KSHV ORF61 is able to relocalize the chimeric A3Gctd L3A/Bctd protein, albeit to considerably lesser extents compared to WT A3Bctd or A3A. These data provide further support for the in silico predictions above by demonstrating the importance of the L3 region for binding to KSHV ORF61 and promoting cytoplasmic aggregate formation in living cells.

Role of A3B and A3A loop 1 in binding to gamma-herpesvirus RNRs

An unanticipated discovery made possible by our recent EBV BORF2-A3Bctd cryo-EM structure is the existence of an extensive interaction between the short HLS region of BORF2 and L1 of A3Bctd [30]. This L1 interaction accounts for EBV BORF2 binding preferentially to A3B as opposed to A3A. We therefore compared the role of the L1 region of A3Bctd versus A3A in the interaction of these enzymes with EBV BORF2 and KSHV ORF61. First, we confirmed our recent report by showing that replacing L1 of A3A with L1 of A3Bctd leads to a marked increase in EBV BORF2 co-IP (Figure 4A, left panel, lanes 1-2). A strong increase in pull-down is also visible when L1 of A3A is replaced with L1 of A3Gctd, possibly due to the similarities between the L1 regions of A3Bctd and A3Gctd, where each enzyme has a 3-residue motif that is absent in A3A (PLV and PWV, respectively; Figure 2A). Reciprocally, replacing L1 of A3Bctd with L1 of A3A—but not with L1 of A3Gctd—dramatically diminishes the interaction with EBV BORF2 (Figure 4A, left panel, lanes 4-5).
In contrast, KSHV ORF61 does not appear to bind more strongly to A3Bctd compared to A3A (Figure 4A, right panel, compare lanes 1 and 4). Moreover, L1 swaps between A3Bctd and A3A have no effect on KSHV ORF61 co-IPs (Figure 4A, right panel, lanes 1, 2, 4, and 5). However, interestingly, replacing L1 of either A3Bctd or A3A with residues from A3Gctd decreases interaction with KSHV ORF61 (Figure 4A, right panel, lanes 3 and 6). This may be attributable to the presence of a unique bulky tryptophan residue in A3Gctd L1 potentially clashing with a nearby interacting surface (Figure 1 and Figure 2A).

We next assessed the subcellular localization phenotypes of these L1 chimeras using quantitative fluorescence microscopy (Figure 4B, representative images; Figure 4C-D, quantification). In contrast to the co-IP results above, swapping the L1 regions of A3Bctd and A3A does not overtly affect relocalization by EBV BORF2. This agrees with our published results [30] and supports a model in which compensatory avidity interactions during cytoplasmic aggregate formation are likely to account for the similar relocalization phenotypes of A3Bctd and A3A despite BORF2 interacting more weakly with the latter protein. In comparison, for KSHV ORF61, swapping L1 of A3Bctd with L1 of A3Gctd slightly decreases relocalization, whereas the remaining L1 swaps have no significant effect (Figure 4B, representative images; Figure 4C-D, quantification).

EBV BORF2 interacts with A3Bctd loop 1 to inhibit DNA deaminase activity

In addition to causing protein relocalization, direct binding by EBV BORF2 potently inhibits A3B catalytic activity in vitro [15, 30]. This supports a model in which EBV BORF2 simultaneously inhibits A3B activity and moves the complex away from viral replication centers to protect EBV genomes from deamination during lytic reactivation. Given results here showing
that EBV BORF2 and KSHV ORF61 bind to both A3B and A3A, we next asked whether direct enzymatic inhibition is a conserved feature of this pathogen-host interaction.

HeLa T-Rex cells stably expressing doxycycline-inducible A3B-EGFP or A3A-EGFP were transfected with an empty vector control or FLAG-tagged EBV BORF2 and KSHV ORF61. After 24 hours of doxycycline induction, whole cell extracts were incubated with a fluorescently labelled ssDNA oligonucleotide substrate containing a single A3B/A deamination motif (5′-TC). A3B/A-mediated deamination of cytosine-to-uracil followed by uracil excision by UDG and abasic site cleavage by sodium hydroxide (NaOH) yields a shorter product that can be detected by SDS-PAGE and fluorescence scanning (see Methods). As expected, EBV BORF2 strongly inhibits A3B deaminase activity in cell lysates as indicated by minimal product accumulation (Figure 5A, left panel, compare lanes 1 and 2). KSHV ORF61 also inhibits A3B-catalyzed deamination when compared to a no-RNR control, albeit to a considerably lesser extent compared to EBV BORF2 (Figure 5A, left panel, compare lanes 1 and 3).

Surprisingly, neither EBV BORF2 nor KSHV ORF61 substantially inhibits the deaminase activity of A3A (Figure 5A, right panel). This stark difference is supported by our cryo-EM structure showing extensive contacts between BORF2 and the L1 and L7 regions of A3Bctd, which results in a high-affinity interaction that blocks the A3B active site from binding to ssDNA substrates and catalyzing deamination [30]. Specifically, A3B L1 residues 206-PLV-208 contribute to the formation of a hydrophobic pocket in which EBV BORF2 residue Y481 is inserted (depicted in Figure 5B). In contrast, this 3-residue motif is absent in A3A L1 (Figure 2A). We therefore hypothesized that inhibition of deaminase activity by EBV BORF2 requires not only binding to the catalytic domain through L7, but also a significant interaction with L1.
To further test this idea, we first generated mutants of EBV BORF2 Y481 and examined their ability to inhibit A3B-catalyzed deamination using the HeLa T-REx A3B-EGFP system described above. Mutating EBV BORF2 Y481 to an alanine or a proline effectively prevents inhibition of A3B-catalyzed deamination, likely by weakening the interaction with the L1 region (Figure 5C). Next, we compared the ability of EBV BORF2 to inhibit the deaminase activity of WT versus L1 swapped A3B and A3A in HeLa cells transfected with FLAG-tagged BORF2 and EGFP-tagged A3s. In agreement with results from Figure 5A, WT A3B deaminase activity is potently inhibited by EBV BORF2 (Figure 5D, left, lanes 1 and 2). In contrast, when L1 of A3B is replaced with L1 of A3A, EBV BORF2 completely loses its capacity to inhibit ssDNA deamination (Figure 5D, left, lanes 3 and 4). In further agreement with data in Figure 5A, WT A3A deaminase activity is not inhibited by EBV BORF2 (Figure 5D, right, lanes 1 and 2). Unfortunately, the chimeric A3A L1B construct exhibits very low levels of ssDNA deaminase activity, and it is therefore not possible to determine the susceptibility to inhibition by EBV BORF2 (Figure 5D, right, lanes 3 and 4). Nevertheless, these results combine to indicate that the EBV BORF2 interaction with A3B L1 is a requirement for inhibiting deamination and also help explain why A3A activity is not inhibited similarly.

A structural overlay of the cryo-EM structure of the EBV BORF2-A3B complex and the RoseTTAFold-predicted model of KSHV ORF61 indicates that a residue analogous to the interlocking Y481 of BORF2 is lacking in ORF61 (Figure 5E). Instead, KSHV ORF61 has a proline (P488) immediately downstream of shared proline and threonine residues. This difference suggests that L1 may be allowed to remain flexible and engage ssDNA substrates when A3B is bound to KSHV ORF61. Alternatively—and perhaps more likely—a lack of interaction with L1 may result in RNR-A3B complexes with lower binding affinity (i.e., lower
association rate and/or higher dissociation rate), thereby rendering KSHV ORF61 unable to inhibit A3B-catalyzed deamination to the same extent as EBV BORF2.

The genesis of $A3B$ may have shaped gamma-herpesvirus RNR evolution in primates

$A3$ genes are unique to placental mammals and the evolutionary history of the $A3$ locus is marked by frequent expansions and contractions resulting from independent events of gene duplication, loss, and fusion [38-41]. For instance, in the primate lineage leading to humans the $A3$ locus evolved from an ancestral 3 deaminase domain cluster to the present day 11 domain (7 gene) cluster. Interestingly, phylogenetic and comparative genomics studies indicate that $A3B$ may be the youngest $A3$ family member in primates [39, 42]. Specifically, $A3B$ was likely generated by duplication of ancestral deaminase domains in the common ancestor of Catarrhini primates (i.e., hominoids and Old World monkeys) and is consequently absent in Platyrhini primates (i.e., New World monkeys) (depicted in Figure 6A). We therefore hypothesized that the function of an ancestral gamma-herpesvirus RNR as an innate immune antagonist may have been selected by the birth of $A3B$ during this critical period of primate evolution (i.e., 43 to 29 million years ago, after the split of Catarrhini and Platyrhini primates).

To test this idea, we cloned the RNR large subunit genes from representative gamma-herpesviruses that infect primates with $A3B$ (EBV and KSHV, which infect humans, and McHV-4 and McHV-5, which infect rhesus macaques) and related viruses that infect primates without $A3B$ (CalHV-3 and SaHV-2, which infect New World marmosets and squirrel monkeys, respectively) (Figure 6B). Of note, these RNRs represent gamma-herpesviruses from two evolutionarily diverged genera—Lymphocryptovirus and Rhadinovirus (Figure 6B). We then examined the capacity of these viral RNRs to bind to human A3Bctd and A3A in co-IP
experiments (Figure 6C). In support of our hypothesis, only the RNRS from viruses that infect A3B-encoding primates (humans and rhesus macaques) pull-down human A3Bctd and A3A (Figure 6C, lanes 1-4, left and right panels). In contrast, the RNRS from viruses that infect primates without A3B (marmosets and squirrel monkeys) are unable to pull-down either enzyme (Figure 6C, lanes 5-6, left and right panels).

We next asked if these viral RNRS can relocalize human A3B and A3A. HeLa T-REx cells were transfected with FLAG-tagged viral RNRS and incubated with doxycycline to induce A3B-EGFP or A3A-EGFP expression (Figure 6D). In agreement with our co-IP results, only the RNRS from viruses that infect A3B-encoding primates are able to relocalize human A3B and A3A from the nucleus to the cytoplasmic compartment. Together, these results suggest that the ability to bind and relocalize human A3 enzymes is conserved among RNRS from gamma-herpesviruses that infect Catarrhini primates.

We additionally used HeLa T-Rex cells stably expressing doxycycline-inducible human A3B-EGFP or human A3A-EGFP to ask whether these viral RNRS are capable of inhibiting the ssDNA deaminase activity of human A3B and A3A (Figure 6E). Consistent with our previous results, EBV BORF2 potently inhibits A3B (Figure 6E, top panel, lane 2). A similarly strong inhibition is seen with the RNR of rhesus macaque Lymphocryptovirus McHV-4 (Figure 6E, top panel, lane 3). The RNRS from KSHV and rhesus macaque Rhadinovirus McHV-5 did not overtly affect the formation of a product band but appeared to slow down A3B-catalyzed deamination, as suggested by a stronger substrate band compared to the no-RNR control (Figure 6E, top panel, lanes 5 and 6). Most importantly, the RNRS from marmoset Lymphocryptovirus CalHV-3 and squirrel monkey Rhadinovirus SaHV-2 are both defective in inhibiting A3B activity (Figure 6E, top panel, lanes 4 and 7). Additionally, none of the six different viral RNRS
inhibits the activity of human A3A (Figure 6E, bottom panel). Collectively, our results suggest a possible link between the birth of A3B in the common ancestor of Catarrhini primates and the evolution of A3B-counteraction functionalities in an ancestral gamma-herpesvirus RNR large subunit (i.e., direct binding, subcellular relocalization, and enzymatic inhibition).

A short helical loop structure from EBV BORF2 enables the marmoset CalHV-3 RNR to bind to human A3B and A3A

A3 enzymes are composed of single or double zinc-coordinating (Z) domains that can be grouped into three classes (Z1, Z2, and Z3) based on sequence similarity [43]. In humans, A3A, A3C, and A3H are single Z domain proteins (Z1, Z2, and Z3, respectively), whereas A3B, A3G, A3D, and A3F harbor two Z domains (Z2-Z1 for A3B and A3G, Z2-Z2 for A3D and A3F). Although New World monkeys lack A3B, the genomes of these species—as well as all primates—encode a single-domain Z1 enzyme homologous to human A3A. Therefore, it is possible that the RNRs of gamma-herpesviruses that infect New World monkeys may have evolved to antagonize the A3A enzymes of their natural host species.

To investigate this possibility, we performed co-IP experiments using FLAG-tagged viral RNRs and host-matched EGFP-tagged A3As (Figure 7A). In line with our results above with human A3Bctd and A3A (Figure 6), all four RNRs from viruses that infect Catarrhini primates bind to the A3A enzymes of their respective host species (Figure 7A, lanes 1, 2, 4, and 5). In contrast, the RNRs from viruses infecting New World monkeys are unable to bind to their hosts’ A3As (Figure 7A, lanes 3 and 6). We next asked if these RNRs can relocalize the A3A enzymes encoded by their respective host species (Figure 7B). In agreement with the co-IP results, the RNRs from EBV/KSHV and McHV-4/McHV-5 relocalize human and rhesus A3A, respectively.
(Figure 7B, left and center). In contrast, the RNRS from CalHV-3 and SaHV-2 do not alter the localization of marmoset and squirrel monkey A3A, respectively (Figure 7B, right). These results further support the idea that A3 antagonism by gamma-herpesvirus RNRS is likely to be limited to viruses that infect primates with A3B and, additionally, suggest that binding to A3A may be a consequence of similarity to A3Bctd (identical loop 3 and loop 7 regions and >90% overall identity).

Our recent cryo-EM structure of the EBV BORF2-A3Bctd complex revealed that loops 1 and 7 of A3B are sequestered by multiple interactions with BORF2 including contacts with a short helical loop structure (HLS) involving the critical BORF2 residues Y481 and R484 ([30]; region depicted in red in Figure 7C and alternative poses in Figure 1 and Figure 5). Amino acid sequence alignments of the RNRS from human EBV, rhesus macaque McHV-4, and marmoset CalHV-3, which are all members of the Lymphocryptovirus genus, show that the former two RNRS share identical amino acid sequences in the HLS region, whereas the latter protein has multiple differences (HLS region alignment in Figure 7C). Given the overall similarity between the 3-dimensional architecture of EBV BORF2 and the predicted structure of CalHV-3 RNR (Figure 7—figure supplement 1), we next asked whether replacing the HLS region of the CalHV-3 RNR with the corresponding motif from EBV BORF2 may be sufficient to enable binding to A3B/A enzymes in co-IP experiments. Interestingly, although the chimeric CalHV-3 RNR construct is still unable to bind marmoset A3A (Figure 7D, lane 2), this protein is now able to efficiently bind to both human A3Bctd and A3A (Figure 7D, lanes 3 and 4). These results indicate that the HLS region of EBV BORF2 is sufficient to endow a normally non-binding Lymphocryptovirus RNR with a capacity to engage both human A3Bctd and A3A, likely by
enabling a direct physical interaction with the identical loop 7 region of these enzymes (L7 region alignment in Figure 7E).

Ancestral A3B is active and antagonized by present-day RNRs from gamma-herpesviruses that infect A3B-encoding primates

To gain deeper insights into the interaction between A3B and gamma-herpesvirus RNRs, we generated a phylogenetic tree of all publicly available primate A3B sequences and used the GRASP-suite [44] to reconstruct the amino acid sequence of ancestral A3B (Figure 8A and Figure 8—figure supplement 1). We then examined an expanded panel of primate gamma-herpesvirus RNRs (i.e., 11 different viral proteins) for a capacity to bind human and ancestral A3Bctd in co-IP experiments (Figure 8B). In agreement with results above, only the RNRs from gamma-herpesviruses that infect primates with A3B are capable of binding to human A3Bctd (Figure 8B, left). Similarly, nearly all RNRs from viruses that infect primates with A3B are able to bind the ancestral protein and, importantly, all viruses that infect primates which naturally lack the A3B gene are unable to bind the ancestral enzyme (Figure 8B, right).

To further explore the properties of the ancestral A3B protein, we performed a comprehensive series of subcellular localization experiments and ssDNA deaminase activity assays using this expanded panel of primate gamma-herpesvirus RNRs (Figure 8C-D and Figure 8—figure supplement 2A). As above, human A3B is completely relocalized from the nucleus to the cytoplasmic compartment by the RNRs of viruses that infect humans and other primates with an A3B gene (Figure 8C, top; Figure 8—figure supplement 2A). Similar results were obtained with ancestral A3B, although only partial relocalization is observed with the KSHV ORF61 (Figure 8C, bottom; Figure 8—figure supplement 2A). KSHV ORF61 also
exhibit a decreased ability to inhibit the deaminase activity of ancestral A3B compared to human A3B (Figure 8D, compare lane 5 from the top and bottom panels). These relocalization and ssDNA deaminase activity results correlate with the co-IP observations in Figure 8B, where the RNRs from the two human gamma-herpesviruses showed decreased binding to ancestral A3Bctd compared to the human enzyme. Such differences in binding to human versus ancestral A3B may be an indication of adaptive changes in present day viral RNRs driven by host-pathogen conflicts.

A phylogenetic tree based on the amino acid sequences of available gamma-herpesvirus RNRs indicates that the sequences from human, rhesus macaque, and marmoset Lymphocryptoviruses cluster together and the sequences from the different Rhadinoviruses cluster according to viral lineage (Figure 8E). Interestingly, despite its similarity to KSHV ORF61, the RNR from ColHV-1—a novel KSHV-related virus identified in African Colobinae monkeys [45]—differs from the RNRs of all other viruses that infect Catarrhini primates tested here in that is shows no ability to bind, relocalize, or inhibit present-day human or ancestral A3B (Figure 8B, Figure 8—figure supplement 2A, and Figure 8D) or interact with its host’s A3A (Figure 8—figure supplement 3A). This apparent incongruity may be explained by the fact that African Colobinae monkeys appear to have lost the A3B gene following the Colobinae subfamily split into the African and Asian tribes 10 to 14 million years ago (Figure 8—figure supplement 3B-C), which presumably relieved the pressure for ColHV-1 to maintain an A3B-neutralization activity.

Differential binding of viral RNRS to present day and ancestral A3B is likely a reflection of past—and possibly still ongoing—genetic conflicts. To further explore this idea, we analyzed a data set of primate A3B DNA sequences for evidence of codon sites under diversifying/positive
selection using the mixed effects model of evolution algorithm (MEME) [46]. We detected
evidence for diversifying selection at multiple sites, including clusters of positively selected
residues in loops 1 and 3 (Figure 8F, top; Figure 8—figure supplement 4A-B). This suggests
that the amino acid composition of these regions may have been influenced by recurrent
diversifying selection, possibly as a mean to escape binding by gamma-herpesvirus RNRs, which
in turn would select for virus adaptation to restore binding (depicted in Figure 8F, bottom). In
line with this interpretation and together with results in Figure 3 and Figure 4, it is possible that
differences in positively selected residues in loops 1 and 3 may respectively account for the
apparent increased ability of EBV BORF2 and KSHV ORF61 to antagonize present day human
A3B compared to ancestral A3B.

Discussion

Viruses use a plethora of mechanisms to avoid and antagonize host immune responses.
We previously reported an unexpected role for herpesvirus RNRs in counteracting innate
immunity by revealing that EBV repurposes its RNR subunit BORF2 to neutralize the antiviral
activity of cellular A3B [15]. The importance of this A3B counteraction mechanism is evidenced
by BORF2-null EBV exhibiting lower infectivity and an accumulation of A3B signature C/G-to-
T/A mutations [15]. Subsequent studies revealed that the RNR subunits from related human
herpesviruses also bind and relocalize A3B, as well as the highly similar A3A enzyme [31, 32].
Here, we leverage knowledge gained from our recent cryo-EM structure of EBV BORF2 in
complex with the catalytic domain of A3B [30] to investigate the mechanistic conservation of A3
antagonism by herpesviruses. We find that EBV BORF2 and KSHV ORF61 bind to cellular A3
enzymes via distinct surfaces and show that direct inhibition of ssDNA deaminase activity by
EBV BORF2 requires specific interactions with the L1 of A3B in addition to the L7 region. We further find that across different primate gamma-herpesvirus lineages, the existence of an RNR-mediated mechanism to counteract cellular A3 enzymes appears to be confined to viral species that infect A3B-expressing primates. Together, these results suggest that the genesis of A3B by duplication of ancestral Z2 and Z1 deaminase domains in the common ancestor of present-day hominoids and Old World monkeys may have selected for an ancestral gamma-herpesviruses with a capacity to antagonize this new restriction factor through its large RNR subunit. To our knowledge, this is the first study to link the birth of an antiviral host factor and the evolution of an equally potent viral escape mechanism.

Our data here highlight the overall conservation of A3B antagonism by gamma-herpesvirus large RNR subunits but also reveal new and unexpected molecular differences. In agreement with our recent cryo-EM studies [30], we show that A3B L7 is a critical determinant of EBV BORF2 binding, whereas A3B L1 serves to help strengthen this interaction and confer selective binding over A3A (Figure 2 and Figure 4). Surprisingly, these same rules do not apply to KSHV ORF61, which appears to mainly rely on L3 to bind and relocalize cellular A3B/A (Figure 3). Together, our results suggest a model where interactions with L7 and/or L3—which are identical in A3B and A3A, but not in any other members of the A3 family—enable gamma-herpesvirus RNRs to bind and relocalize both enzymes, whereas additional interacting regions (i.e., EBV BORF2 HLS and A3B L1) may help confer selectivity and additional neutralization activity (e.g., potent inhibition of A3B ssDNA deaminase activity). It is additionally interesting that this may be a new manifestation of a pathogen-host wobble mechanism [47] in which ongoing small changes over evolutionary time in the viral RNRs can manifest molecularly in the present-day target protein as distinct binding surfaces (e.g., EBV BORF2 requires A3Bctd L7...
and KSHV ORF61 requires L3).

Using a panel of 11 primate gamma-herpesvirus RNRs, we show that, in addition to binding and relocalization, all RNRs derived from viruses that infect hosts that encode A3B show some capacity to inhibit the ssDNA deamination activity of human A3B, but none of these viral proteins could inhibit the deaminase activity of human A3A (Figure 8 and Figure 8—figure supplement 2B). This is consistent with a model in which the antiviral activity of A3B may have forced these viruses to evolve and maintain a potent counteraction mechanism, whereas interaction with A3A may be a consequence of sequence homology to A3B and less likely a true physiologic function. This interpretation is further supported by our observation that the RNRs from viruses that infect A3B-null primates are incapable of interacting with the A3A enzymes of their natural host species (Figure 7).

The A3B-selective nature of this host-pathogen interaction is orthogonally supported by previous binding studies showing that the EBV BORF2-A3Bctd interaction has a biolayer interferometry dissociation constant of ~1 nM, whereas the interaction with the >90% identical A3A enzyme is over 10-fold weaker [30]. Our results here reveal that, in addition to providing selective binding to A3B, EBV BORF2 interaction with the L1 region of A3B is required for potently inhibiting DNA deaminase activity (Figure 5). It is thus possible that the direct binding of EBV BORF2 to two loop regions near the A3B catalytic site—L1 and L7—helps stabilize the BORF2-A3B complex, resulting in a high affinity interaction and consequently blocking the A3B active site from accessing ssDNA substrates and catalyzing deamination. Importantly, the potential adaptability of this A3B counteraction mechanism in Lymphocryptoviruses is highlighted by results showing that grafting the HLS region from EBV BORF2—which contains residues that directly contact A3B L1 and L7—into the RNR from marmoset CalHV-3 enables
binding to human A3B (Figure 7). In comparison, KSHV ORF61 binding to A3B largely through L3 may result in a lower affinity interaction that more weakly inhibits ssDNA deamination. These inferences support a model in which binding to the A3 active site by viral RNRs is responsible for both protein relocalization and enzymatic inhibition, yet substantial deaminase activity inhibition is only observed above a certain affinity threshold. We therefore postulate that preferential binding to A3B might have been an evolutionary adaptation resulting from selective pressures disproportionately affecting gamma-herpesviruses within the Lymphocryptovirus genus, such as EBV and MeHV-4. For instance, EBV latency is limited to B lymphocytes [48], which can express high levels of endogenous A3B [49, 50], strongly implying that the neutralization of this enzyme is required to preserve viral genome integrity upon lytic reactivation. On the other hand, the broader tropism of KSHV suggests that lytic replication might not always occur in the presence of A3B [51-55], which may result in lower exposure to A3B-mediated deamination.

Evolutionary analyses of host-pathogen antagonistic relationships are powerful tools for understanding protein functions. The link between A3B generation by duplication of ancestral Z2 and Z1 domains and the evolution of A3-neutralization functionalities in herpesvirus RNRs described here suggests that the double domain architecture of A3B may be a requirement for herpesvirus restriction. Notably, the non-catalytic Z2 domain of A3B has been shown to account for this enzyme’s distinct nuclear localization [56] and association with multiple RNA binding proteins, in addition to allosterically regulating the catalytic activity of the Z1 domain [57]. It is therefore plausible that cooperation between Z2 and Z1 domains accounts for A3B antiviral function, possibly by facilitating access to herpesvirus transcription complexes inside the nucleus (e.g., ssDNA exposed in viral R-loop structures [58]). Future studies dissecting the molecular
mechanisms that allow A3B to recognize and target foreign nucleic acids such as herpesvirus
genomes might reveal insights into the regulation of ssDNA deaminase activity and how the loss
of such regulatory mechanisms can at times result in cellular chromosomal DNA mutations and
cancer development (reviewed by [59, 60]). Future work will likely benefit from interrogating
the RNRs and A3 enzymes from a larger and more diverse panel of virus and host species, which
is currently prevented by the relatively small number of herpesvirus complete genomes deposited
in public databases. Bat species may be particularly interesting given they encode the largest
known A3 repertoire in mammals [61, 62] and are also known to host a variety of different
herpesviruses [63-73].

Materials and methods

DNA constructs

The large RNR subunit from EBV (HHV-4), McHV-4, CalHV-3, KSHV (HHV-8),
McHV-5, McHV-11, McHV-12, AtHV-3, SaHV-2, ColHV-1, and SgHV-1 were obtained as
gBlocks purchased from Integrated DNA Technologies and cloned into pcDNA4/TO (Invitrogen
#V102020) with a C-terminal 3×FLAG-tag (DYKDDDDK) by restriction digestion and ligation.
EBV BORF2Y481A, EBV BORF2Y481P, CalHV-3 RNRWEOARDV490YDSRDM constructs were generated
by site-directed mutagenesis using Q5 High-Fidelity DNA Polymerase (NEB #M0491). Wild-type
constructs match the following GenBank accessions: EBV BORF2 (V01555.2), McHV-4
BORF2 (AY037858.1), CalHV-3 ORF55 (AF319782.2), KSHV ORF61 (U75698.1), McHV-5
ORF61 (AF083501.3), McHV-11 ORF61 (AY528864.1), McHV-12 ORF61 (KP265674.2),
AtHV-3 ORF61 (AF083424.1), SaHV-2 ORF61 (X64346.1), ColHV-1 ORF61 (MH932584.1), and SgHV-1 ORF26 (OK337614.1).

APOBEC3 enzymes were obtained as gBlocks purchased from Integrated DNA Technologies and cloned into pcDNA5/TO (Invitrogen #V103320) with a C-terminal EGFP-tag by restriction digestion and ligation. The loop swap constructs A3B_L1A3A_{25-30}, A3Bctd_{193-382}_L1A3A_{25-30}, A3Bctd_{193-382}_L1A3G_{209-217}, A3Bctd_{193-382}_L3A3G_{247-254}, A3Bctd_{193-382}_L7A3G_{317-322}, A3A_L1A3B_{205-213}, A3A_L1A3G_{209-217}, A3A_L3A3G_{247-254}, A3A_L7A3G_{317-322}, A3Gctd_{197-384}_L3A3A_{60-67}, and A3Gctd_{197-384}_L7A3A_{132-137} were generated by site-directed mutagenesis using Q5 High-Fidelity DNA Polymerase (NEB #M0491). Wild-type constructs match the following GenBank accessions: Homo sapiens A3B (NM_004900), Homo sapiens A3A (NM_145699), Homo sapiens A3G (NM_021822), Macaca mulatta A3A (NM_001246231.1), Macaca mulatta A3B (NM_001246230.2), Callithrix jacchus A3A (NM_001301845.1), Saimiri boliviensis A3A (XM_039462978.1), Colobus angolensis palliatus A3A (XM_011964212.1).

**Human cell culture**

Unless indicated, cell lines were derived from established laboratory collections. All cell cultures were supplemented with 10% fetal bovine serum (Gibco #26140-079), 1× penicillin-streptomycin (Gibco #15140-122), and periodically tested for mycoplasma contamination using MycoAlert PLUS Mycoplasma Detection Kit (Lonza #LT07-710). 293T cells were cultured in high glucose DMEM (HyClone #SH30022.01) and HeLa cells were cultured in RPMI 1640 (Gibco #11875-093).
To generate HeLa T-REx A3A-EGFP and A3B-EGFP cell lines, HeLa cells were transduced with pLENTI6/TR (Invitrogen #V480-20), selected with 5 μg/ml blasticidin (GoldBio #B-800-25), and subcloned by serial dilution to obtain single cell clones. Clonal populations were stably transfected with pcDNA5/TO-A3A-EGFP or pcDNA5/TO-A3B-EGFP, selected with 200 μg/ml hygromycin (GoldBio #H-270-1), and subcloned by serial dilution. Representative clones were used for DNA deaminase activity assays (Figure 5A, Figure 5C, Figure 6E) and IF microscopy experiments (Figure 6D).

Co-immunoprecipitation experiments

Approximately 250,000 293T cells were seeded in 6-well plates and transfected the next day with 100 ng pcDNA5/TO-A3-EGFP and 200 ng pcDNA4/TO-RNR-3×FLAG using TransIT-LT1 (Mirus #2304). Cells were harvested ~24 hours after transfection and resuspended in 500 μl of lysis buffer (50 mM Tris-HCl pH 7.4, 150 mM NaCl, 10% glycerol, 0.5% IGEPAL, 100 μg/ml RNaseA, 1 tablet of Sigma-Aldrich cOmplete Protease Inhibitor Cocktail). Resuspensions were vortexed, sonicated on ice-bath for 10 seconds at the lowest setting using a Branson sonifier, and centrifuged at 15,000 × g at 4 °C for 15 minutes. A 50 μl aliquot was removed from each sample for input detection and the remaining clarified whole cell lysate was incubated with 20 μl packed gel volume of anti-FLAG M2 magnetic beads (Sigma-Aldrich #M8823) at 4 °C overnight with gentle rotation. Samples were washed 3 times with 500 μL of lysis buffer and bound protein was eluted with 50 μl of elution buffer (100 μg/mL FLAG peptide (Sigma-Aldrich #F3290), 50 mM Tris-HCl pH 7.4, 150 mM NaCl). Input and elution samples were analyzed by immunoblot using mouse anti-GFP (1:5,000, Clontech #632381) to detect A3-EGFP, rabbit anti-FLAG (1:5,000, Sigma-Aldrich #F7425) to detect RNR-3×FLAG, and rabbit
anti-GAPDH (1:5,000, Proteintech #60004) to detect GAPDH (loading control). Secondary antibodies used were IRDye 800CW anti-rabbit (LI-COR #926-32211) and HRP-linked anti-mouse (Cell Signaling #7076), both at 1:10,000 dilution.

**Immunofluorescence microscopy experiments**

In Figure 6D, approximately 5,000 engineered HeLa T-REx cells (described above) were seeded into 96-well imaging plates (Corning #3904) and transfected the next day with 30 ng pcDNA4/TO-RNR-3×FLAG. A3B-EGFP or A3A-EGFP expression was induced 6 hours after transfection with 50 ng/mL doxycycline. Approximately 24 hours after induction, cells were fixed in 4% formaldehyde (ThermoFisher #28906), permeabilized in 0.2% Triton X-100 (Sigma-Aldrich #T8787) for 10 minutes and incubated in blocking buffer (2.8 mM KH2PO4, 7.2 mM K2HPO4, 5% goat serum (Gibco #16210064), 5% glycerol, 1% cold water fish gelatin (Sigma-Aldrich #G7041), 0.04% sodium azide, pH 7.2) for 1 hour. Cells were then incubated with primary antibody mouse anti-FLAG (1:1,000, Sigma #F1804) overnight at 4 °C with gentle rocking to detect FLAG-tagged viral RNRs. The next day, cells were incubated with secondary antibody anti-mouse Alexa Fluor 594 (1:1,000, Invitrogen #A-11005) for 2 hours at room temperature protected from light and nuclei were counterstained with Hoechst 33342 (ThermoFisher #62249) for 10 minutes. Representative images were acquired using a Cytation 5 Cell Imaging Multi-Mode Reader (BioTek) at 20× magnification.

In Figures 2-4, approximately 5,000 HeLa cells were seeded into 96-well imaging plates (Corning #3904) and transfected the next day with 15 ng pcDNA5/TO-A3-EGFP and 30 ng pcDNA4/TO-RNR-3×FLAG. Cells were fixed approximately 24 hours after transfection. Permeabilization, antibody staining, and nuclei counterstaining with Hoechst 33342 were
performed as described above. Plates were imaged using a Cytation 5 Cell Imaging Multi-Mode Reader (BioTek) at 20× magnification using the automated imaging function to capture 50 non-overlapping images per well. Background subtraction was performed using Fiji [74] and images were analyzed using CellProfiler v4.2 [75]. A pipeline was generated to identify individual nuclei, trace cell boundaries, segment the cytoplasmic/nuclear compartments, and take measurements of intensity, texture, and correlation for the Hoechst (nuclear stain), RNR, and A3 channels within each subcellular compartment. At least 100 cells were measured for each well. All measurements were exported to an SQLite database and CellProfiler Analyst v3.0 [76] was used for scoring phenotypes by machine learning in order to quantify the number of cells showing relocalized versus non-relocalized A3 within the population of RNR-expressing cells in each well. The percentages of cells with relocalized A3 in Figure 2D, Figure 3D, and Figure 4C were calculated for n = 3 independent experimental replicates (i.e., data generated from the analysis of cells seeded, transfected, and imaged on three different days). The ridgeline plots in Figure 2E, Figure 3E, and Figure 4D were generated in RStudio with the ggplot2 package [77] using the Pearson correlation coefficient measurements for the A3 and Hoechst channels within each segment cell nucleus.

In Figure 7B and Figure 8C, approximately 5,000 HeLa cells were seeded into 96-well imaging plates (Ibidi #89626) and transfected the next day with 15 ng pcDNA5/TO-A3-EGFP and 30 ng pcDNA4/TO-RNR-3×FLAG. Cells were fixed approximately 24 hours after transfection. Permeabilization, antibody staining, and nuclei counterstaining with Hoechst 33342 were performed as described above. Plates were imaged using an Eclipse Ti2-E inverted microscope (Nikon) at 20× magnification.
**ssDNA deaminase activity assays**

In *Figure 5A, Figure 5C,* and *Figure 6E,* approximately 250,000 HeLa T-REX cells stably expressing doxycycline-inducible A3B-EGFP or A3A-EGFP (described above) were seeded into 6-well plates and transfected the next day with 200 ng pcDNA4/TO-RNR-3×FLAG using TransIT-LT1 (Mirus #2304). A3B-EGFP or A3A-EGFP expression was induced 6 hours after transfection with 50 ng/mL doxycycline. Approximately 24 hours after induction, cells were harvested, resuspended in 100 µL of reaction buffer (25 mM HEPES, 15 mM EDTA, 10% glycerol, 1 tablet of Sigma-Aldrich cOmplete Protease Inhibitor Cocktail), and subjected to freeze-thaw. Whole cell lysates were then centrifuged at 15,000 × g for 15 minutes and the clarified supernatant was transferred to a new tube. Soluble lysates were incubated with 100 µg/mL RNAse A at room temperature for 1 hour. A 5 µl aliquot was removed from each sample for immunoblots using mouse anti-GFP (1:5,000, Clontech #632381) to detect A3-EGFP, rabbit anti-FLAG (1:5,000, Sigma-Aldrich #F7425) to detect RNR-3×FLAG, and rabbit anti-GAPDH (1:5,000, Proteintech #60004) to detect GAPDH (loading control). Secondary antibodies used were IRDye 800CW anti-rabbit (LI-COR #926-32211) and HRP-linked anti-mouse (Cell Signaling #7076), both at 1:10,000 dilution. 5 µl of soluble lysate were incubated at 37 °C for 1 hour with 0.7 µM of a fluorescent ssDNA substrate (5′-ATT ATT A TT ATT CAA ATG GAT TTA TTT ATT TAT TTA TTT ATT T-fluorescein-3′) and 2.5 U of UDG (NEB #M0280) in a total reaction volume of 10 µl (diluted in reaction buffer). NaOH was then added to the reaction mix to a final concentration of 100 mM and samples were incubated at 98 °C for 10 minutes. The reaction was then mixed with 11 µl 2× formamide buffer (80% formamide, 1×TBE, bromophenol blue, and xylene cyanol) and reaction products were separated on a 15% TBE-urea
PAGE gel. Separated DNA fragments were visualized on a Typhoon FLA 7000 scanner (GE Healthcare) on fluorescence mode.

In Figure 5D and Figure 8D, approximately 250,000 293T cells were seeded in 6-well plates and transfected the next day with 100 ng pcDNA5/TO-A3-EGFP and 200 ng pcDNA4/TO-EBV-BORF2-3×FLAG. Cells were harvested approximately 24 hours after transfection and the DNA deaminase activity assay proceeded as described above.

**Protein structure modeling**

Protein structure models were generated with RoseTTAFold [34] using the ROBETTA server followed by simple all-atom refinement of the top-ranked models using the Relax application in the ROSIE 2 server [36]. To model the structure of the ORF61-A3Bctd complex (Figure 1), RoseTTAFold-predicted models of KSHV ORF61 and A3Bctd were aligned to the cryo-EM BORF2-A3Bctd complex (pdb: 7RW6) using PyMOL [35] to arrange the starting docking pose and local protein-protein docking was performed using the RosettaDock 4.0 server [36, 37]. The 10,000 generated structures were sorted based on Critical Assessment of Predicted Interactions (CAPRI) score [78] and subsequently ranked according to their interface energy score (Figure 1—figure supplement 1B). All protein structure figures were made using PyMOL [35].

**Herpesvirus RNR phylogenetic trees generation**

Amino acid sequences of the large RNR subunits from the following herpesviruses were obtained from the NCBI Protein RefSeq database: HSV-1 (YP_009137114.1), HSV-2 (YP_009137191.1), VZV (NP_040142.1), EBV (YP_401655.1), HCMV (YP_081503.1), HHV-
6A (NP_042921.1), HHV-6B (NP_050209.1), HHV-7 (YP_073768.1), KSHV (YP_001129418.1), McHV-4 (YP_067953.1), CalHV-3 (NP_733909.1), McHV-5 (NP_570809.1), McHV-11 (AAT00096.1), McHV-12 (AJE29717.1), AtHV-3 (AAC95585.1), SaHV-2 (NP_040263.1), ColHV-1 (QDQ69272.1), and SgHV-1 (UNP64460.1). To generate the phylogenetic trees in Figure 6B and Figure 8E, protein sequences were aligned using MUSCLE [79] and maximum likelihood phylogenies were inferred using PhyML 3.0 [80] using the Smart Model Selection (SMS) tool [81] and 100 bootstraps.

Ancestral APOBEC3B sequence reconstruction

To generate the phylogenetic tree in Figure 8A, primate A3B amino acid sequences were aligned using MUSCLE [79] (Figure 8—figure supplement 1) and a maximum likelihood phylogeny was inferred using PhyML 3.0 [80] using the Smart Model Selection (SMS) tool [81] and 100 bootstraps. Reconstruction of the ancestral A3B protein sequence was performed with GRASP [44] using the sequence alignment and phylogenetic tree above.

Positive Selection Analysis

Primate A3B DNA sequences were aligned using ClustalOmega [82] and a maximum likelihood phylogeny was inferred using PhyML 3.0 [80] using the Smart Model Selection (SMS) tool [81] and 100 bootstraps. Codon sites under episodic diversifying selection were identified with the HyPhy 2.5 software package [83] using the mixed effects model of evolution algorithm (MEME) [46] with 100 bootstraps and p-value threshold of 0.1 (Figure 8—figure supplement 4A-B).
Acknowledgements

We thank members of the Harris lab for thoughtful comments. These studies were supported in part by grants to RSH from the National Institute for Allergy and Infectious Diseases (NIAID) R37-AI064046 and the National Cancer Institute (NCI) P01-CA234228. Salary support for SNM was provided by NIAID F31-AI161910 and subsequently by a grant to the University of Minnesota (SNM and RSH) from the Howard Hughes Medical Institute through the James H. Gilliam Fellowships for Advanced Study program. Salary support for JTB was provided by NIAID F32-AI147813. Salary support for NMS and AAA was provided in part by NIAID R56-AI150402 and NIH T32-AI83196, respectively. RSH is the Ewing Halsell President’s Council Distinguished Chair and an Investigator of the Howard Hughes Medical Institute. The authors have no competing interests to declare.

References


26. Kouno T, Silvas TV, Hilbert BJ, Shandilya S, Bohn MF, Kelch BA, Royer WE, Somasundaran M, Kurt Yilmaz N, Matsuo H. Crystal structure of APOBEC3A bound to single-


Zheng X-y, Qiu M, Chen S-w, Xiao J-p, Ma L-z, Liu S, Zhou J-h, Zhang Q-h, Li X, Chen Z. High prevalence and diversity of viruses of the subfamily Gammaherpesvirinae, family


Figure 1. Structural differences and similarities between EBV and KSHV RNR large subunits

(A) Zoom-in of the EBV BORF2-A3Bctd interface (pdb 7rw6, chains A and B) showing a network of interactions between BORF2 (blue) and A3Bctd L7 and L1 regions (gray). BORF2 short helix insertion (SHI), helical loop structure (HLS), and interacting residue Q476 are shown in red. Gray dashed line represents the L3 region not visible by cryo-EM.

(B) Amino acid sequence alignment of the SHI and HLS regions of EBV BORF2 and KSHV ORF61 (complete alignment in Figure 1—figure supplement 1A).

(C) RoseTTAFold-predicted structure of KSHV ORF61 (orange, above) overlayed with EBV BORF2 (pdb 7rw6, chain A; blue, below). HLS and SHI regions in both proteins and residue G482 in ORF61 are shaded red.

(D-E) Cryo-EM structure of EBV BORF2-A3Bctd (pdb 7rw6, chains A and B) and highest-ranked protein-protein docking model of KSHV ORF61-A3Bctd generated using RosettaDock (Figure 1—figure supplement 1B). Panel D shows the full complexes with key regions labeled. Panel E shows a zoom-in of the RNR-A3Bctd interface (BORF2, blue; ORF61, orange; A3Bctd gray; SHI and HLS regions, red). A3Bctd loops are labeled L1, L3, and L7 (gray dashed line represents the L3 region not visible by cryo-EM).

Figure 1—figure supplement 1

(A) Amino acid sequence alignment of EBV BORF2 and KSHV ORF61 with identical residues shaded yellow and the SHI and HLS regions designated by red lines.
A superposition of the three highest-ranked protein-protein docking models of the KSHV ORF61-A3Bctd complex generated using RosettaDock (left). The CAPRI model rankings computed by RosettaDock range from 0 (incorrect model) to 3 (highest quality model; detailed in [78] with additional metrics shown to right).

**Figure 2. Involvement of the A3B and A3A loop 7 regions in binding to gamma-herpesvirus RNRs**

(A) Surface-filled representation of ssDNA bound by an A3 (pdb 5sww, chains A and E). L1, L3, and L7 regions are colored in yellow with amino acid alignments to the right.

(B) Co-IP of A3 L7 chimeras with EBV BORF2 and KSHV ORF61. FLAG-tagged RNR subunits were co-expressed with the indicated A3-EGFP constructs in 293T cells, affinity purified, and analyzed by immunoblotting to detect co-purifying A3 proteins.

(C) Representative IF microscopy images of HeLa cells co-transfected with FLAG-tagged EBV BORF2 or KSHV ORF61 (red) and the indicated A3-EGFP constructs (green). Scale = 10 μm.

(D-E) Quantification of A3-EGFP relocalization in HeLa cells expressing EBV BORF2 or KSHV ORF61. Panel D shows the percentage of cells exhibiting relocalized A3-EGFP (mean ± standard deviation, n ≥ 100 cells per condition). Statistical analyses were performed using unpaired t-tests to determine significant changes in the relocalization of L7 swapped A3s (darker shade) relative to WT (lighter shade; n = 3 independent experimental replicates; ns, not significant P > 0.5; * P ≤ 0.5; ** P ≤ 0.01; *** P ≤ 0.001). Panel E shows Pearson correlation coefficient (PCC) values for A3-EGFP (or EGFP alone as a control) and Hoechst (nuclear stain).

Density curves show the PCC value distribution in each condition, and the white vertical lines indicate the median PCC value (n ≥ 100 cells per condition).
Figure 2—source data 1
File contains original immunoblots for Figure 2B.

Figure 3. Role of A3B and A3A loop 3 in binding to gamma-herpesvirus RNRs
(A) Ribbon schematic of the EBV BORF2-A3Bctd complex (pdb 7rw6, chains A and B) with BORF2 in blue and A3Bctd in gray. Gray dashed line represents the L3 region not visible by cryo-EM.
(B) Co-IP of A3 L3 chimeras with EBV BORF2 and KSHV ORF61. FLAG-tagged RNR subunits were co-expressed with the indicated A3-EGFP constructs in 293T cells, affinity purified, and analyzed by immunoblotting to detect co-purifying A3 proteins.
(C) Representative IF microscopy images of HeLa cells co-transfected with FLAG-tagged EBV BORF2 or KSHV ORF61 (red) and the indicated A3-EGFP constructs (green). Scale = 10 μm.
(D-E) Quantification of A3-EGFP relocalization in HeLa cells expressing EBV BORF2 or KSHV ORF61. Panel D shows the percentage of cells exhibiting relocalized A3-EGFP (mean ± standard deviation, n ≥ 100 cells per condition). Statistical analyses were performed using unpaired t-tests to determine significant changes in the relocalization of L3 swapped A3s (darker shade) relative to WT (lighter shade; n = 3 independent experimental replicates; ns, not significant P > 0.5; * P ≤ 0.5; ** P ≤ 0.01; *** P ≤ 0.001). Panel E shows Pearson correlation coefficient (PCC) values for A3-EGFP (or EGFP alone as a control) and Hoechst (nuclear stain). Density curves show the PCC value distribution in each condition, and the white vertical lines indicate the median PCC value (n ≥ 100 cells per condition).
**Figure 3—source data 1**

File contains original immunoblots for Figure 3B.

**Figure 4. Role of A3B and A3A loop 1 in binding to gamma-herpesvirus RNRs**

**(A)** Co-IP of A3 L1 chimeras with EBV BORF2 and KSHV ORF61. FLAG-tagged RNR subunits were co-expressed with the indicated A3-EGFP constructs in 293T cells, affinity purified, and analyzed by immunoblotting to detect co-purifying A3 proteins.

**(B)** Representative IF microscopy images of HeLa cells co-transfected with FLAG-tagged EBV BORF2 or KSHV ORF61 (red) and the indicated A3-EGFP constructs (green). Scale = 10 μm.

**(C-D)** Quantification of A3-EGFP relocalization in HeLa cells expressing EBV BORF2 or KSHV ORF61. Panel C shows the percentage of cells exhibiting relocalized A3-EGFP (mean ± standard deviation, n ≥ 100 cells per condition). Statistical analyses were performed using unpaired t-tests to determine significant changes in the relocalization of L1 swapped A3s (darker shade) relative to WT (lighter shade; n = 3 independent experimental replicates; ns, not significant P > 0.5; * P ≤ 0.5; ** P ≤ 0.01; *** P ≤ 0.001). Panel D shows Pearson correlation coefficient (PCC) values for A3-EGFP (or EGFP alone as a control) and Hoechst (nuclear stain).

Density curves show the PCC value distribution in each condition, and the white vertical lines indicate the median PCC value (n ≥ 100 cells per condition).

**Figure 4—source data 1**

File contains original immunoblots for Figure 4A.

**Figure 5. EBV BORF2 interacts with A3Bctd loop 1 to inhibit DNA deaminase activity.**
(A) Upper panels show ssDNA deaminase activity of extracts from HeLa T-REx cells transfected with the indicated FLAG-tagged RNR subunits and incubated with doxycycline to induce the expression of A3B-EGFP or A3A-EGFP. Lower panels show protein expression levels in the same extracts by immunoblotting.

(B) Zoom-in of the EBV BORF2-A3Bctd interface (pdb 7rw6, chains A and B) showing a network of interactions between BORF2 (blue, ribbon representation) and A3Bctd L7 and L1 regions (gray, surface-filled representation). BORF2 SHI, HLS, and Q476 are shown in red.

(C) Upper panels show ssDNA deaminase activity of extracts from HeLa T-REx cells transfected the indicated FLAG-tagged EBV BORF2 constructs and incubated with doxycycline to induce the expression of A3B-EGFP. Lower panels show protein expression levels in the same extracts by immunoblotting.

(D) Upper panels show ssDNA deaminase activity of extracts from HeLa cells co-transfected with FLAG-tagged EBV BORF2 and the indicated A3-EGFP constructs. Lower panels show protein expression levels in the same extracts by immunoblotting.

(E) Ribbon schematic of the EBV BORF2-A3Bctd interface (pdb 7rw6, chains A and B) highlighting BORF2 tyrosine 481 and A3Bctd L1. BORF2 is shown in blue, the HLS region in red, and A3Bctd in gray. The right schematic shows the same view with the predicted structure of KSHV ORF61 (orange) overlayed and BORF2 residues hidden to allow better visualization (full model in Figure 1C).

Figure 5—source data 1

File contains original deaminase activity assay gels and immunoblots for Figure 5A.
Figure 5—source data 2
File contains original deaminase activity assay gels and immunoblots for Figure 5C.

Figure 5—source data 3
File contains original deaminase activity assay gels and immunoblots for Figure 5D.

Figure 6. The genesis of A3B may have shaped gamma-herpesvirus RNRs evolution.
(A) Chronogram of primates depicting the generation of A3B in the common ancestor of hominoids and Old World monkeys. Primate groups shown in blue are composed of species that encode A3B and those shown in red lack A3B. Primate phylogeny and divergence intervals were derived from timetree.org [84]. Mya = million years ago.

(B) Top: Phylogenetic tree of human herpesvirus and indicated non-human primate herpesvirus RNR large subunits. Shaded boxes indicate alpha-, beta-, and gamma-herpesvirus subfamilies. Genus classification is shown for gamma-herpesviruses. Gamma-herpesviruses that infect host species that encode A3B are shown in blue and those that infect host species that lack A3B are shown in red. Bottom: Host species infected by the indicated gamma-herpesviruses. Virus-host pairs are colored according to A3B status, as described for panel A.

(C) Co-IP of human A3A and A3Bctd with the RNR large subunits from the indicated viruses. FLAG-tagged RNR subunits were co-expressed with A3A-EGFP or A3Bctd-EGFP in 293T cells, affinity purified, and analyzed by immunoblotting to detect co-purifying A3 proteins.

(D) Representative IF microscopy images of HeLa T-REx cells transfected with the indicated FLAG-tagged RNR subunits (red) and incubated with doxycycline to induce the expression of human A3B-EGFP or A3A-EGFP (green). Scale = 10 μm.
Upper panels show ssDNA deaminase activity of extracts from HeLa T-REx cells transfected with the indicated FLAG-tagged RNR subunits and incubated with doxycycline to induce the expression of A3B-EGFP or A3A-EGFP. Lower panels show protein expression levels in the same extracts by immunoblotting.

**Figure 6—source data 1**

File contains original immunoblots for Figure 6C.

**Figure 6—source data 2**

File contains original deaminase activity assay gels and immunoblots for Figure 6E.

**Figure 7. A short helical loop structure from EBV BORF2 enables the marmoset CalHV-3 RNR to bind to human A3B and A3A**

(A) Co-IP of human, rhesus macaque, marmoset, and squirrel monkey A3As with the indicated gamma-herpesvirus RNR subunits. FLAG-tagged RNR subunits were co-expressed with the indicated A3-EGFP constructs in 293T cells, affinity purified, and analyzed by immunoblotting to detect co-purifying A3 proteins.

(B) Representative IF microscopy images of HeLa cells co-transfected with the indicated FLAG-tagged viral RNRS (red) and A3A-EGFP constructs (green). Scale = 10 μm.

(C) Ribbon schematic of the EBV BORF2-A3Bctd complex (pdb 7rw6, chains A and B) with BORF2 in blue and A3Bctd in gray. The HLS of EBV BORF2 is depicted in red. Bottom-right: zoom-in of the interacting surfaces highlighting BORF2 Y481 and R484 interactions with A3B.
loops 1 and 7, respectively. Top-right: amino acid alignment of the HLS regions of the EBV, McHV-4, and CalHV-3 RNRs (red boxes) and adjacent residues.

(D) Co-IP of marmoset A3A, human A3A, and human A3Bctd with WT or chimeric CalHV-3 RNR containing the HLS region of EBV BORF2. FLAG-tagged RNR subunits were co-expressed with the indicated A3-EGFP constructs in 293T cells, affinity purified, and analyzed by immunoblotting to detect co-purifying A3 proteins.

(E) Amino acid alignment of the loop 7 regions of marmoset A3A, human A3A, and human A3B (gray boxes) with non-identical loop 7 residues highlighted in red.

Figure 7—source data 1
File contains original immunoblots for Figure 7A.

Figure 7—source data 2
File contains original deaminase activity assay gels and immunoblots for Figure 7D.

Figure 7—figure supplement 1
Ribbon schematics of the EBV BORF2 cryo-EM structure (pdb 7rw6, chain A; blue) and the RoseTTAFold-predicted CalHV-3 RNR structure (green). The HLS region of each protein is shaded red and a structural overlay is shown for comparison.

Figure 8. Ancestral A3B is active and antagonized by the RNRs from gamma-herpesviruses that infect A3B-encoding primates
(A) Phylogenetic tree of primate A3B sequences used in this study to reconstruct ancestral A3B amino acid sequence alignment in **Figure 8—figure supplement 1**.

(B) Co-IP of human and ancestral A3Bctd with the indicated gamma-herpesvirus RNR subunits. FLAG-tagged RNR subunits were co-expressed with the indicated A3-EGFP constructs in 293T cells, affinity purified, and analyzed by immunoblotting to detect co-purifying A3 proteins. RNRs from viruses that infect host species that encode A3B are shown in blue, and RNRs from viruses that infect host species that lack A3B are shown in red. LCV = *Lymphocryptovirus*; RV = *Rhadinovirus*; UN = unclassified gamma-herpesvirus (no official genus classification from the International Committee on Taxonomy of Viruses [85]).

(C) Representative IF microscopy images of HeLa cells co-transfected human or ancestral A3B-EGFP (green) and the indicated FLAG-tagged viral RNRs (red). Scale = 10 μm. See **Figure 8—figure supplement 2A** for additional RNR data.

(D) Upper panels show ssDNA deaminase activity of extracts from HeLa cells co-transfected with human or ancestral A3B-EGFP and the indicated FLAG-tagged viral RNRs. Lower panels show protein expression levels in the same extracts by immunoblotting.

(E) Phylogenetic tree of primate gamma-herpesviruses RNR large subunits. *Rhadinovirus* RV1, RV2, and NW refer to primate *Rhadinovirus* lineages [86-90] (UN, unclassified).

(F) Top: positively selected sites in A3Bctd identified using MEME [46] with p < 0.1 (red triangles in schematic and red residues in loop region alignments; see **Figure 8—figure supplement 4** for details). Bottom: schematic of the proposed genetic conflict between primate gamma-herpesvirus RNRs and cellular A3B.

**Figure 8—source data 1**
Figure 8—source data 2
File contains original deaminase activity assay gels and immunoblots for Figure 8D.

Figure 8—figure supplement 1
Amino acid sequence alignment of the indicated primate A3Bs. A3Bctd regions are shaded yellow and the L1, L3, and L7 regions are highlighted with blue boxes. The predicted ancestral A3B protein sequence is positioned at the bottom of the group.

Figure 8—figure supplement 2
(A) Representative IF microscopy images of HeLa cells co-transfected human or ancestral A3B-EGFP (green) and the indicated FLAG-tagged viral RNRs (red). Scale = 10 μm.
(B) Upper panels show ssDNA deaminase activity of extracts from HeLa cells co-transfected with human A3A-EGFP and the indicated FLAG-tagged viral RNRs. Lower panels show protein expression levels in the same extracts by immunoblotting.

Figure 8—figure supplement 2—source data 1
File contains original deaminase activity assay gels and immunoblots for Figure 8—figure supplement 2, panel B.

Figure 8—figure supplement 3
(A) Co-IP of human, rhesus macaque, and Angola colobus A3A enzymes with the ColHV-1 RNR. FLAG-tagged ColHV-1 RNR was co-expressed with the indicated A3A-EGFP constructs in 293T cells, affinity purified, and analyzed by immunoblotting to detect co-purifying A3 proteins. A3A enzymes from primate species that encode A3B are shown in blue and A3A enzymes from primate species that lack A3B are shown in red.

(B) Schematic diagram depicting the APOBEC3 loci from the indicated primate species representative of Hominoidea (human, genome assembly GRCh38.p13), Cercopithecinae (rhesus macaque, genome assembly Mmul_10), African Colobinae (Angola colobus, genome assembly Cang.pa_1.0; Ugandan red colobus, genome assembly ASM277652v2), and Asian Colobinae (golden snub-nosed monkey, genome assembly Rrox_v1). The African Colobinae monkey genomes available via ENSEMBL [91] indicate that the A3B gene has been lost, likely after the Colobinae subfamily split into the African and Asian tribes (10 to 14 million years ago) because its closest relatives, the Asian Colobinae (e.g., Golden snub-nosed monkey), still encode A3B. APOBEC3 domains are colored according to Z domain classification (gold = Z1, blue = Z2, red = Z3).

(C) Chronogram of primates depicting the loss of A3B following the split of the Colobinae subfamily into the African and Asian tribes (indicated by delta symbol). Primate groups shown in blue are composed of species that encode A3B and primates groups shown in red lack A3B. Primate phylogeny and divergence intervals were obtained from timetree.org [84]. Mya = million years ago.

Figure 8—figure supplement 3—source data 1

File contains original immunoblots for Figure 8—figure supplement 3, panel A.
Figure 8—figure supplement 4

(A) Amino acid sequence alignment of the C-terminal catalytic domain of the indicated primate A3B proteins with sites under diversifying positive selection identified using MEME [46] with p < 0.1 highlighted in red. L1, L3, and L7 regions are colored with blue boxes.

(B) Summary of MEME analysis results showing the p-values in dN/dS ratio test and the most common codon substitutions at each codon site.
### A
- **A3B**
  - Empty
  - BORF2
  - ORF61
- **A3A**
  - Empty
  - BORF2
  - ORF61

<table>
<thead>
<tr>
<th>Protein</th>
<th>Empty</th>
<th>BORF2</th>
<th>ORF61</th>
</tr>
</thead>
<tbody>
<tr>
<td>Substrate</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Product</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GFP (A3)</td>
<td>70</td>
<td>55</td>
<td>100</td>
</tr>
<tr>
<td>FLAG (RNR)</td>
<td>100</td>
<td>55</td>
<td>100</td>
</tr>
<tr>
<td>GAPDH</td>
<td>35</td>
<td>35</td>
<td></td>
</tr>
</tbody>
</table>

### B
- EBV BORF2
- Proteins: L133, Y134, R484, L7, Y481, L1, P479, T480

### C
- **BORF2**
  - Empty
  - WT
  - Y481A
  - Y481P

<table>
<thead>
<tr>
<th>Protein</th>
<th>Empty</th>
<th>WT</th>
<th>Y481A</th>
<th>Y481P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Substrate</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Product</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GFP (A3B)</td>
<td>70</td>
<td>55</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>FLAG (BORF2)</td>
<td>100</td>
<td>55</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>GAPDH</td>
<td>35</td>
<td>35</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### D
- **A3B**
  - -
  - +
- **A3B L1A**
  - -
  - +
- **A3A**
  - -
  - +
- **A3A L1B**
  - -
  - +

<table>
<thead>
<tr>
<th>Protein</th>
<th>-</th>
<th>+</th>
<th>-</th>
<th>+</th>
</tr>
</thead>
<tbody>
<tr>
<td>Substrate</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Product</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GFP (A3)</td>
<td>55</td>
<td></td>
<td>40</td>
<td></td>
</tr>
<tr>
<td>FLAG (BORF2)</td>
<td>100</td>
<td></td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>GAPDH</td>
<td>35</td>
<td></td>
<td>35</td>
<td></td>
</tr>
</tbody>
</table>

### E
- EBV BORF2
- KSHV ORF61 (predicted)

<table>
<thead>
<tr>
<th>Protein</th>
<th>EBV BORF2</th>
<th>KSHV ORF61 (predicted)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HLS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>T487</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L207</td>
<td></td>
<td></td>
</tr>
<tr>
<td>V208</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P486</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P206</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Y481</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P479</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L133</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Y134</td>
<td></td>
<td></td>
</tr>
<tr>
<td>R484</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Y481</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P479</td>
<td></td>
<td></td>
</tr>
<tr>
<td>T480</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>NMPQNSFRPEM RMYRDFYDN FEMEPFLXGR SWTFLCVRK VRGRGNLNIQ DTQVFRQXIV FQXQRSAMFC FLSWFRGQL</td>
<td></td>
</tr>
<tr>
<td>---</td>
<td>-----------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>Gorilla (XM 001300506)</td>
<td>Human (NM 004900.5)</td>
<td></td>
</tr>
<tr>
<td>Bonobo (NM 001301766)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chimpanzee (NM 001289639)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Orangutan (XM 009234378.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gibbon (NM 001345972.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Green monkey (NM 001345954.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sooty mangabey (NM 001200202.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Drill (XM 001197845)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baboon (NM 001346001)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rhesus mac (NM 001246230.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cynomolgus mac (XM 005567246.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pig-tailed mac. (KX538362.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Snub-nosed monkey (XM 030915811.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ancestral</td>
<td></td>
<td></td>
</tr>
<tr>
<td>81</td>
<td>PAYCFCQITW FYSWTCPDCE VAXLAEFLSE HPWTVLITA ARLYVGYTFD YRALCRSLQ AGAVRTYVNY EEFAYCWENF</td>
<td></td>
</tr>
<tr>
<td>Gorilla (XM 030100506)</td>
<td>Human (NM 004900.5)</td>
<td></td>
</tr>
<tr>
<td>Bonobo (NM 001301766)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chimpanzee (NM 001289639)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Orangutan (XM 009234378.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gibbon (NM 001345972.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Green monkey (NM 001345954.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sooty mangabey (NM 001200202.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Drill (XM 001197845)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baboon (NM 001346001)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rhesus mac (NM 001246230.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cynomolgus mac (XM 005567246.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pig-tailed mac. (KX538362.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Snub-nosed monkey (XM 030915811.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ancestral</td>
<td></td>
<td></td>
</tr>
<tr>
<td>161</td>
<td>VYNNEQGPMP WYKFDENAYA ------- F LHRKKEILR LUMDFTFTF NFPFELLVR FTQYLCRVEY ERLNQKTVVL</td>
<td></td>
</tr>
<tr>
<td>Gorilla (XM 030100506)</td>
<td>Human (NM 004900.5)</td>
<td></td>
</tr>
<tr>
<td>Bonobo (NM 001301766)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chimpanzee (NM 001289639)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Orangutan (XM 009234378.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gibbon (NM 001345972.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Green monkey (NM 001345954.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sooty mangabey (NM 001200202.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Drill (XM 001197845)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baboon (NM 001346001)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rhesus mac (NM 001246230.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cynomolgus mac (XM 005567246.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pig-tailed mac. (KX538362.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Snub-nosed monkey (XM 030915811.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ancestral</td>
<td></td>
<td></td>
</tr>
<tr>
<td>241</td>
<td>MQEHHGFLCH EAALKLLGGFY QRBAELRFDL LVPSLQLDPA QYHRVTWTFIS WSPCFSWQCA GEVRAFLQEN THVRLRIFAA</td>
<td></td>
</tr>
<tr>
<td>Gorilla (XM 030100506)</td>
<td>Human (NM 004900.5)</td>
<td></td>
</tr>
<tr>
<td>Bonobo (NM 001301766)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chimpanzee (NM 001289639)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Orangutan (XM 009234378.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gibbon (NM 001345972.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Green monkey (NM 001345954.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sooty mangabey (NM 001200202.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Drill (XM 001197845)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baboon (NM 001346001)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rhesus mac (NM 001246230.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cynomolgus mac (XM 005567246.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pig-tailed mac. (KX538362.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Snub-nosed monkey (XM 030915811.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ancestral</td>
<td></td>
<td></td>
</tr>
<tr>
<td>321</td>
<td>RITYDYESX BALQMLRADK AQUSSIMTGYD FEYKCDPWY QROCFQPDMD GLEBSQAL GRLAILQ0QN GN</td>
<td></td>
</tr>
<tr>
<td>Gorilla (XM 030100506)</td>
<td>Human (NM 004900.5)</td>
<td></td>
</tr>
<tr>
<td>Bonobo (NM 001301766)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chimpanzee (NM 001289639)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Orangutan (XM 009234378.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gibbon (NM 001345972.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Green monkey (NM 001345954.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sooty mangabey (NM 001200202.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Drill (XM 001197845)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baboon (NM 001346001)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rhesus mac (NM 001246230.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cynomolgus mac (XM 005567246.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pig-tailed mac. (KX538362.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Snub-nosed monkey (XM 030915811.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ancestral</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### A

<table>
<thead>
<tr>
<th>Gene</th>
<th>Codon Substitution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human (NM_004900.5)</td>
<td></td>
</tr>
<tr>
<td>Gorilla (XM_031005506.1)</td>
<td></td>
</tr>
<tr>
<td>Bonobo (NM_001301766.1)</td>
<td></td>
</tr>
<tr>
<td>Chimpanzee (NM_001289639.1)</td>
<td></td>
</tr>
<tr>
<td>Orangutan (XM_009234378.2)</td>
<td></td>
</tr>
<tr>
<td>Gibbon (NM_001345972.1)</td>
<td></td>
</tr>
<tr>
<td>Green monkey (NM_001345954.1)</td>
<td></td>
</tr>
<tr>
<td>Sooty mangabey (XM_012032024.1)</td>
<td></td>
</tr>
<tr>
<td>Drill (XM_011978466.1)</td>
<td></td>
</tr>
<tr>
<td>Baboon (NM_001346001.1)</td>
<td></td>
</tr>
<tr>
<td>Rhesus mac. (NM_001246230.2)</td>
<td></td>
</tr>
<tr>
<td>Cynomolgus mac. (NM_005567246.2)</td>
<td></td>
</tr>
<tr>
<td>Pig-tailed mac. (KX583652.1)</td>
<td></td>
</tr>
<tr>
<td>Snub-nosed monkey (XM_030915811.1)</td>
<td></td>
</tr>
<tr>
<td>Ancestral</td>
<td></td>
</tr>
</tbody>
</table>

### B

<table>
<thead>
<tr>
<th>Site</th>
<th>p-value</th>
<th>Most common codon substitution</th>
</tr>
</thead>
<tbody>
<tr>
<td>216</td>
<td>0.099</td>
<td>CCT&gt;CTT, CTT&gt;CTT</td>
</tr>
<tr>
<td>219</td>
<td>0.0693</td>
<td>CCT&gt;CTT, CTT&gt;CTT, CTT&gt;TTT</td>
</tr>
<tr>
<td>220</td>
<td>0.0891</td>
<td>CGA&gt;GGA</td>
</tr>
<tr>
<td>233</td>
<td>0.0693</td>
<td>CTG&gt;CGG</td>
</tr>
<tr>
<td>240</td>
<td>0.0594</td>
<td>CGG&gt;CAG, CTG&gt;AAG, CTG&gt;CGG</td>
</tr>
<tr>
<td>249</td>
<td>0.0099</td>
<td>TGG&gt;CAC</td>
</tr>
<tr>
<td>255</td>
<td>0.0594</td>
<td>CCT&gt;CTT, CTT&gt;CTT</td>
</tr>
<tr>
<td>257</td>
<td>0.0891</td>
<td>CAT&gt;CGT, TAT&gt;CAT, TAT&gt;TGT</td>
</tr>
<tr>
<td>259</td>
<td>0.0297</td>
<td>GGT&gt;GAT, GTT&gt;GTT, TTT&gt;TGG</td>
</tr>
<tr>
<td>271</td>
<td>0.0099</td>
<td>CAG&gt;TGG, CAG&gt;CAG, CAG&gt;TGG, TGG&gt;AAG</td>
</tr>
</tbody>
</table>