TLR-TRIF Pathway Enhances the Expression of KSHV Replication and Transcription Activator*

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Background: Host innate immunity is against virus infection and replication.

Results: Toll-like receptor 3 activation leads to enhanced expression of a key Kaposi’s sarcoma-associated herpesvirus (KSHV) protein.

Conclusion: KSHV uses host Toll-like receptor pathway to augment its critical gene expression.

Significance: A virus may usurp host innate immunity for its own benefits.

Kaposi’s sarcoma-associated herpesvirus (KSHV) is a human γ-herpesvirus. KSHV replication and transcription activator (RTA) is necessary and sufficient for KSHV reactivation from latency. Toll-like receptors (TLRs) recognize pathogen-associated molecular patterns, act through adaptors, and initiate innate and adaptive immune responses against pathogens. Toll/interleukin-1 receptor domain containing adaptor protein producing interferon-β (TRIF) is an adaptor associated with TLR3 and TLR4 signaling, and is closely related to antiviral signaling to activate type I interferon (IFN). We previously found that KSHV RTA degrades TRIF indirectly and blocks TLR3 pathways. In this report, we find that TRIF, as well as TLR3 activation, enhances KSHV RTA protein expression. The C-terminal region of the RTA is involved in the responding TRIF-mediated enhancement. The degradation of TRIF and the enhancement of RTA expression are using two different pathways. The enhancement by TLR-TRIF is at least partially via promoting translational efficiency of RTA mRNA. Finally, the receptor-interacting protein 1 (RIP1) may be involved in TRIF-mediated enhancement of RTA expression, but not in the RTA-mediated degradation of TRIF. Therefore, the activation of TLR-TRIF pathway enhances KSHV RTA protein expression, and KSHV RTA in turn degrades TRIF to block innate immunity. The putative KHSV-TLR-adaptor-interacting loop may be a critical element to evade and usurp host innate immunity in KSHV life-cycle.

Toll-like receptors (TLRs) are a family of evolutionarily conserved receptors that recognize molecular patterns unique to pathogens and activate host innate immunity against the pathogen (1, 2). One of the major products from TLR activation is production of inflammatory cytokines such as tumor necrosis factor-α (TNF-α), interleukin 1 (IL1), and type I interferons (IFN). IFN is key component to mount a proper and robust immune response to a viral infection (3, 4). Toll-IL-1 receptor (TIR) domain-containing adaptor inducing IFN-β (TRIF), also called TIR domain-containing adaptor molecule-1 (TICAM-1), is an adaptor protein involved in signal transduction during the activation of TLR3 and 4 leading to activation of nuclear factor κB (NF-κB) and type I IFN (5–7). TLR3 seems to be dependent on TRIF for its downstream cascades (5, 8).

Kaposi’s Sarcoma-Associated Herpesvirus (KSHV), also known as human herpesvirus 8, is a member of the human γ-herpesviruses family. KSHV is believed to be an etiological factor for Kaposi’s Sarcoma (KS), and associated with several other B lymphocytes malignancies such as primary effusion lymphoma and multicentric Castleman’s disease. As other herpesviruses, KSHV consists of two distinct phases: latent and lytic replication during its life cycle. During latency, the virus establishes persistent infection and only a small subset of genes are typically expressed. Under conditions of lytic replication, all the viral genes are activated in cascade mode and new viruses are packaged and released from cells (9–12). KSHV replication and transcription activator (RTA) is an immediate early gene and highly conserved among γ-herpesviruses (13–15). RTA is apparently necessary and sufficient for the switch from KSHV latency to lytic replication (12, 16). Beyond functioning in initiating viral lytic replication, RTA is involved in the induction of cellular IL6 (17), degradation of TRIF, IFN regulatory factor 7 (IRF7), K-RBP, and Hey1 through proteasome pathway (18–21), and blockage of p53-mediated apoptosis by competing for binding to CBP (22). RTA interacts with other factors to modulate its transcription potential and other cellular activities (22–26).

KSHV mainly infects endothelial cells and B lymphocytes, and those cells express multiple TLRs (27–30). TLR4 is identified as an important barrier against KSHV infection, and KSHV has developed a mechanism to rapidly suppress TLR4 expression (31). KSHV infection activates TLR3 and TLR9 pathways, and TLR7 signaling may lead to lytic replication of KSHV (31–34). Also, murine γ-herpesvirus 68 (MHV68) is a herpesvirus with significant similarities to KSHV. Activation of the TLR3/4

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4 The abbreviations used are: TLR, Toll-like receptor; KSHV, Kaposi’s sarcoma-associated herpesvirus; RTA, replication and transcription activator; TRIF, Toll/interleukin-1-receptor domain containing adaptor protein inducing interferon-β; TIR, Toll-IL-1 receptor; RIP, receptor-interacting protein.
TRIF Enhances KSHV RTA Expression

pathway potently inhibits the replication of MHV68 (35). However, other reports suggest that TLR3 activation increases MHV68 viral replication in vivo (36).

Previously, we found that KSHV RTA degrades TRIF protein through a proteasome pathway (21). In this report, we have found that TRIF up-regulates the expression of RTA protein. The enhanced RTA protein expression by TRIF is at the translational efficiency of its mRNA. The downstream target of TRIF, receptor-interacting protein 1 (RIP1), may be involved in the process. Those data strongly suggest that KSHV usurps host innate immune system for its own benefits.

MATERIALS AND METHODS

Plasmids, Antibodies, and dsRNA—Expression plasmids of KSHV RTA and its mutant (RTA-K152E), EBV RTA, pcDNA3.1-myc-TRIF, TRIF mutants (TRIF-Del, TRIF-N, and TRIF-C) were described previously (21, 37–42). RTA-ΔC plasmid (aa 1–527) was a gift from Dr. Charles Wood (43). Human FLAG-tagged-RIP1 expression plasmid was obtained from Dr. Ning Shunbin. RTA antibody was described (44). Pan-luc, fetal bovine serum (FBS; Invitrogen) and 1% penicillin-streptomycin (46). Those cells were grown in Dulbecco’s modified Eagle’s medium (DMEM; Invitrogen). LGH4930 (5’-TTGGCCTGTAGAAGAAGC-3’) and LGH4929 (5’-GATTCGGAAGCTTCACTCGAGAATTAACG-3’) primers were used for detection of RTA expression, and actin1 (5’-TCAATGAGCTCGGTTG-3’) and actin2 (5’-GAGGGGAGACGACTCTGGTGG-3’) primers were used for actin control.

Translational Efficiency Assays—293T cells were transfected with various plasmids. 24 h after transfection, cells were washed, cultured in methionine-free DMEM ( Gibco) supplemented with 10% of dialyzed FBS for 30 min, and then labeled with of [35S]methionine (Perkin Elmer, 100 μCi/ml) for 30 min. Cells were lysed in 1% Nonidet P-40 buffer (100 mM Tris, pH 8.0, 1 mM EDTA, 100 mM NaCl, 1% Nonidet P-40, 1 mM PMSF plus one tablet of protease inhibitor (Roche) per 10 ml) for 30 min on ice. Supernatants were immunoprecipitated with FLAG antibody. Proteins were transferred to a PVDF membrane and allowing to air dry. The membrane was exposed overnight to a Kodak film to capture the 35S signal. The membrane was re-hybridized in 100% methanol, and Western blot was carried out with RTA antibody. The signal strengths were enumerated by the use of the Bio-Rad Software Quantity One (version 4.6.7).

Transient Transfection and Reporter Assays—Effectene (Qiagen) was used for the transfection of 293T cells. The luciferase assays were performed using the assay kit from Promega according to the manufacturer’s recommendation.

RESULTS

TRIF Increases Steady State Levels of RTA Protein—To examine an effect of TRIF on RTA expression, equal amounts of RTA along with various amounts of KSHV plasmids were transfected into 293T cells. As shown in Fig. 1A, TRIF protein expression is gradually decreased as reported (21). However, the steady state of RTA protein expression was increased upon TRIF expression. Of note, the expression of TRIF could be detected with longer exposure and more lysates (data not shown). Epstein-Barr virus (EBV), the prototype of human γ-herpesvirus, has a RTA homologue (EBV-R). We further tested if EBV-R was degraded by RTA. As shown in Fig. 1B, EBV-R protein expression was not increased by TRIF expression, and EBV-R did not degrade TRIF (Fig. 1B). In addition, the same phenomenon can be observed in other human cell lines, such as MCF7 and U2OS (data not shown). Because RTA degrades TRIF, whether the same pathway was used by TRIF to enhance the expression of RTA was examined. RTA-K152E weakly degraded TRIF, but the expression of the mutant was enhanced by TRIF (Fig. 1C). Furthermore, we tested if TRIF enhanced the RTA-mediated transactivation of reporter gene constructs. As shown in Fig. 1D, TRIF increased the activation of Pan as well as K14 promoter reporter constructs, probably through the enhancing the expression of RTA. Therefore, TRIF activation increased steady state levels of KSHV RTA protein, and the pathways for TRIF degradation and RTA enhancement seem to be different.
TRIF Enhances KSHV RTA Expression—TRIF is required for TLR-3 signaling (5, 8). Whether the activation of TLR-3 increases RTA protein expression in TLR-3-expressing cells (WT11) was tested (45). The RTA expression plasmid was transfected into the cells, and the cells were treated with dsRNA for the activation of TLR3. The activation of TLR3 led to the enhancement of the RTA expression (Fig. 2A). This enhancement was specific for KSHV, as EBV-R could not be enhanced by TLR3 activation (Fig. 2A). Therefore, TLR3 signal specifically enhanced KSHV RTA expression.

TRIF Enhances Endogenous RTA Expression—To examine whether TRIF could enhance RTA under physiological conditions, KSHV latently infected PEL cell line, BC3, were transfected with various expression plasmids along with CD4 selection. The cells were treated with sodium butyrate, and the transfected cells were enriched by CD4-magnetic bead selection. The cells were treated with sodium butyrate, and the transfected cells were enriched by CD4-magnetic bead selection as described before (50–54). The concentrations of sodium butyrate were adjusted to the levels that just barely induced the lytic replication of KSHV. Following the transfection of TRIF, the expression of RTA was clearly increased (Fig. 2A). Therefore, TLR3 signal specifically enhanced KSHV RTA expression.

TRIF Increases the Translation Efficiency of RTA mRNA—To determine the mechanism by which TRIF enhances the expression of RTA, whether the RNA translational efficiency was involved was examined. Basically, the cells were transfected with various plasmids, pulse labeled with [S35]methionine for 30 min, and cell lysates were used for immunoprecipitation with FLAG antibody, followed by Western blot analysis with RTA antibody. The newly synthesized proteins were metabolically labeled with [S35]methionine, and the relative translational efficiency was measured by the ratio of the newly synthesized proteins versus total proteins. As shown in Fig. 3, A and B, the translation efficiency of RTA mRNA was enhanced at least 50% in the presence of TRIF. In addition, TRIF did not have an effect on the half-life of RTA protein in the 24 h time period, during which RTA expression was enhanced (Fig. 3C). And the protein synthesis inhibitor was working properly because IRF1 has anticipated half-life (55). Next, the mRNA levels of RTA were not changed dramatically in the presence of TRIF (Fig. 3D). All those data collectively supports that notion that TRIF increased the translation efficiency of RTA mRNA, which consequently increased the levels of RTA proteins.

Multiple Regions of TRIF Stimulate RTA Expression—To narrow down the region(s) of TRIF for the enhancement of RTA expression, we have used several mutants as shown in Fig. 4A (21). Those deletion mutants were transfected into 293T cells, and whether the TRIF mutants were enhancing the RTA expression was examined. As shown in Fig. 4B, all mutants were able to enhance the RTA expression. In addition, all those TRIF mutants were degraded by RTA as reported (21). These data suggest that the multiple regions of TRIF targeted RTA for its enhancement.

A RTA Deletion Mutant Failed to Be Enhanced by TRIF—To determine the region of RTA responsive to TRIF expression, a serial of mutants had been examined, and one mutant in particular (RTA-ΔC) was different from others (Fig. 4C). TRIF failed to enhance the expression of the RTA-ΔC mutant, however, TRIF significantly enhanced the expression of wtRTA in the same assay (Fig. 4D). The expression of TRIF is also shown. Therefore, the C-terminal region of RTA was responsible for the enhancement of RTA expression by TRIF.

FIGURE 1. TRIF increases the expression of RTA. A, 293T cells were transfected with cDNA, RTA (0.1 μg), and various TRIF plasmid (0.01, 0.05, 0.1 μg). The cell lysates were collected 1 day later, and the expression was examined by Western blot. B, same as A, but TRIF (0.1 μg) and EBV-R (0.05, 0.1 μg) were used. C, 293T cells were transfected with various plasmids as shown on the top. RTA (0.1 μg) and RTA-K152E (0.05 μg) were used for transfection. The images in the same box indicate that they are derived from the same membranes. The identity of proteins is as shown. D, 293T cells were transfected with various reporter constructs along with the CMV-β-gal, TRIF, or RTA expression plasmid as shown on the top. Both Pan and K14 promoters are well-known for their responsiveness to RTA. Luciferase activity was normalized to β-galactosidase activity. The fold activation of each promoter construct is shown with standard deviation. One representative of three independent experiments is shown.
Receptor Interacting Protein-1 (RIP1) May Be Involved in the Enhancement of RTA Expression—Because we failed to detect physical interactions between TRIF and RTA (21), it is very likely that TRIF degradation and RTA expression enhancements occur indirectly. To examine the cellular factors involved in the degradation and enhancement, we used specific cell lines that lack critical components for TLR3 signaling, especially the downstream mediator of TRIF. RIP1 is critically involved in the TRIF-mediated NF-κB activation (56). In the RIP1(−/−) mouse embryonic fibroblast (MEF) line, TRIF failed to increase the expression of RTA; however, the degradation of TRIF was still exist (Fig. 5A). In the corresponding RIP1(+/−) MEFs, TRIF was still enhancing the expression of RTA (Fig. 5B). In addition, overexpression for human RIP1 alone was sufficient to increase the RTA expression in human 293T cells. RTA did not obviously degrade RIP1 because the relative ratios of RIP1 to tubulin were similar (Fig. 5C). All those data collectively suggested that RIP1 might be involved in TRIF-mediated enhancement of RTA expression.

DISCUSSION

Innate immunity is important to control viral infection, and a successful counteraction of innate signaling may be a necessity for the survival of a virus in vivo. It has been shown that KSHV encodes several genes to counteract the innate system. KSHV uses at least two viral gene products to nullify the function of IRF7, a master gene for type I IFN production (20, 47). Latency-associated nuclear antigen blocks the activation of IFN through IRF-3 (48). ORF10 blocks the IFN signaling in KSHV-infected cells (49). On the other hands, it has been shown that virus may use molecules involved in the innate immunity for its own benefits. KSHV uses TLR7 signaling pathway for its reactivation process (33), EBV uses the same signaling pathway for its viral protein expression (57). MHV68 uses cellular Mavs adaptor.
molecular pathway to modify its gene products for the maximum replication (58). MyD88 may be involved in the establishment of MHV68 latency in vivo (59).

In this report, we studied the effects of TRIF on the regulation of KSHV. We find that: 1) TRIF specifically increases steady state levels of KSHV RTA protein and the increased RTA was functional (Fig. 1); 2) TLR3 activation leads to higher levels of steady-state RTA protein expression (Fig. 2A). Because TRIF is required for TLR3 signaling, the results suggest that the endogenous TRIF activation would contribute to RTA expression; 3) the ectopic expression of TRIF leads to enhanced expression of RTA from the viral genome upon induction of lytic replication (Fig. 2B). All those data collectively indicates that TLR-TRIF pathway may enhance the expression of RTA under native environments. Therefore, there is a potential regulatory loop: the activation of TRIF leads to enhanced KSHV RTA protein expression, and RTA degrades TRIF to block innate immunity induced by TLR activations (Fig. 6).

We have extensively addressed the biological function of the regulatory loop. We thought the loop might suggest that TLR3 was a positive regulator of KSHV lytic replication as RTA is a key modulator of lytic replication. We have concentrated on the detection of TLR3 agonist (dsRNA) on the effect of KSHV for the following reasons: (A) KSHV infection activates TLR3 pathway. (B) TRIF is specifically involved in the TLR3 pathway; and (C) TLR4 already shown to be a negative regulator of KSHV (31). We had examined: (A) if TLR3 enhances spontaneous KSHV replication; (B) whether TLR3 activation plus chemical treatments lead to greater KSHV lytic replication; (C) If TLR3 activation at different times of lytic replication leads to KSHV replication enhancements. (D) Whether overexpression of TRIF in KSHV-infected cells, enhances the KSHV lytic replication. Of note, virus production in the supernatant was used as readout for the detection of viral replication, and all our data suggest that TLR3 activation and TRIF per se were negative regulators of KSHV lytic replication (data not shown). Therefore, the only evidence that TLR3 may enhance viral replication is from murine γ-herpesvirus 68 (MHV68), a murine KSHV homologue. It is reported that TLR3 activation increases MHV68 virus titers in vivo (36). However, other report argues against this notion (35). Based on all the results, it is apparent that TRIF-mediated RTA enhancement will facilitate the deg-
radiation of the TRIF itself, and thus blocks the inhibitory effects of TLR3 (Fig. 6).

TRIF is a multifunctional adaptor protein, mediating activation of several transcription factors including NF-κB (60), and trigger apoptosis. Our work has been added another novel function of TRIF, i.e. to enhance a viral gene expression for the benefits of viruses. The mechanism of the enhancement is mainly by promoting translational efficiency of RTA mRNA (Fig. 3), although other mechanisms, such as transcriptional control as well as protein stability, might also present. Furthermore, RIP1, which is a downstream molecule of TRIF activation pathway, might be involved in the TRIF-mediated enhancement of RTA expression (Fig. 5). However, the results seem to contradict to the fact that TRIF interacts with RIP1 using its C-terminal region. It is clear that the capability of TRIF to enhance RTA expression varies among different cell lines. In 293T cells, TRIF seems to have highest efficiency (Fig. 1 and data not shown). In addition, different TRIF fragments also have different efficiency for enhancing RTA expression. We summon that the different cell lines may be responsible for the apparent discrepancy. As RIP1 expression in 293T cells enhancing RTA expression (Fig. 5), RIP1 seems to be one of the mediators involved in the enhancement of RTA.

So far the exact responsive domain(s) for RTA to respond to TLR3/TRIF activation is not completely identified (Fig. 4). We narrowed the region down to the C-terminal (Fig. 4). Based on the results in 293T cells, a fine mapping of the RTA domain responsive to TRIF had been proven to be difficult (data not shown). RTAΔC is unique that it did not respond to TRIF. We suspect that multiple regions of RTA may response to different regions of TRIF for the enhancements.

The detailed molecular mechanism for the enhancement of translation efficiency of RTA mRNA is currently unknown. It is known that TRIF signaling stimulates translation efficiency of TNF-α mRNA via prolonged activation of protein kinase MK2 in macrophages (61). Also, TLR3-TRIF signaling has been shown to suppress endoplasmic reticulum stress-induced translation inhibition through activation of eIF2B (62, 63). Whether a similar mechanism is also used for TRIF-mediated RTA enhancement is under investigation. The specific enhancement of KSHV RTA may have a unique mechanism because the responsive sequences are located in the C-terminus of the molecule (Fig. 4).

We propose the following scenario upon primary KSHV infection: KSHV infection of cells would lead to the activation of TLR3, which may result in IFN production as well as the apoptosis of the infected cells for the suppression of viral replication. KSHV RTA is an immediate early gene that expressed upon viral infection, and RTA may block TLR3-mediated innate immunity by degrading TRIF. Furthermore, the activation of TLR3 pathway leads to the enhancement of RTA protein expression, which would accelerate the degradation process of TRIF, which would attenuate the TRIF signaling (Fig. 6). In addition, RTA may indirectly affect TLR4 signaling by activating vIRF1 that inhibits TLR4 mRNA expression. Because TLR4 is involved in the KSHV pathogenesis in KS and exerts innate immunity against KSHV (31), and TRIF is involved in TLR3, 4 signaling (60), the putative regulatory loop may play an important role in KSHV life cycle.

In sum, we have identified a novel method that KSHV uses for its gene expression. Together with our previous findings, a regulatory loop is apparent: KSHV degrades TRIF, a TLR adaptor, to block TLR-mediated antiviral effects; and TRIF enhances expression of a KSHV critical protein to block TLR3-mediated repressive effects. This report provides evidence that a virus not only blocks the host innate immunity, but also harnesses an immune pathway for the benefits of virus infection.

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