Variola virus E3L Zα domain, but not its Z-DNA binding activity, is required for PKR inhibition

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ABSTRACT

Responding to viral infection, the interferon-induced, double-stranded RNA (dsRNA)–activated protein kinase PKR phosphorylates translation initiation factor eIF2α to inhibit cellular and viral protein synthesis. To overcome this host defense mechanism, many poxviruses express the protein E3L, containing an N-terminal Z-DNA binding (Zα) domain and a C-terminal dsRNA-binding domain (dsRBD). While E3L is thought to inhibit PKR activation by sequestering dsRNA activators and by directly binding the kinase, the role of the Zα domain in PKR inhibition remains unclear. Here, we show that the E3L Zα domain is required to suppress the growth-inhibitory properties associated with expression of human PKR in yeast, to inhibit PKR kinase activity in vitro, and to reverse the inhibitory effects of PKR on reporter gene expression in mammalian cells treated with dsRNA. Whereas previous studies revealed that the Z-DNA binding activity of E3L is critical for viral pathogenesis, we identified point mutations in E3L that functionally uncouple Z-DNA binding and PKR inhibition. Thus, our studies reveal a molecular distinction between the nucleic acid binding and PKR inhibitory functions of the E3L Zα domain, and they support the notion that E3L contributes to viral pathogenesis by targeting PKR and other components of the cellular anti-viral defense pathway.

Keywords: E3L; PKR; eIF2; translation

INTRODUCTION

Viruses rely on the host cell translational machinery for the synthesis of viral proteins. One strategy that host cells use to limit viral replication is to inhibit cellular protein synthesis. Virus infection initiates an anti-viral immune response that results in the production of interferons, which, in turn, induce expression of a variety of anti-viral proteins including the double-stranded RNA (dsRNA)–dependent protein kinase PKR. PKR consists of two dsRNA-binding domains (dsRBD) at the N terminus and a Ser/Thr kinase domain at the C terminus (Fig. 2A, right). Binding of dsRNA, a product of viral replication or gene expression, to the dsRBDs promotes oligomerization of PKR, leading to kinase domain dimerization and autophosphorylation. Following this activation, PKR is competent to phosphorylate its substrate, the α subunit of eukaryotic translation initiation factor eIF2 on Ser51 (Dever et al. 1993; Dar et al. 2005; Dey et al. 2005a). Phosphorylation of eIF2α converts eIF2 from a substrate to an inhibitor of its guanine nucleotide exchange factor eIF2B, thereby preventing the recycling of inactive eIF2–GDP to functional eIF2–GTP. The resulting inhibition of translation initiation impairs both cellular and viral protein synthesis, and thus blocks virus replication.

To enhance their success, viruses have developed a variety of mechanisms to subvert the host cell anti-viral defenses. Viral inhibitors of PKR target various stages in PKR activation and function including dsRNA binding, kinase dimerization and autophosphorylation, and eIF2α phosphorylation (for review, see Garcia et al. 2007). The adenovirus VA RNAs and Epstein–Barr virus EBER RNAs bind to the dsRBDs of PKR and block kinase activation (Kitajewski et al. 1986; Mathews and Shenk 1991; Sharp et al. 1993), while dsRNA-binding proteins including the poxvirus E3L protein (Chang et al. 1992), the rotavirus NSP3 protein (Langland et al. 1994), and the reovirus virus σ3 protein (Beattie et al. 1995; Yue and Shatkin 1997) are thought to sequester dsRNA and thereby prevent PKR activation. In addition to the E3L protein, many poxviruses also encode a pseudosubstrate inhibitor of PKR (Bratke et al. 2013). For example, the vaccinia virus K3L protein (Davies et al. 1992; Carroll et al. 1993; Kawagishi-Kobayashi et al. 1997; Dar and Sicheri 2002; Seo et al. 2008), variola virus C3L protein (Seo et al. 2008),
swinepox virus C8L protein (Kawagishi-Kobayashi et al. 2000), and ranavirus vIF2α protein (Rothenburg et al. 2011) resemble the N-terminal OB-fold domain of vIF2α (Dar and Sicheri 2002; Dar et al. 2005). Binding of these proteins to activated PKR (Dar et al. 2005) is thought to prevent access to vIF2α and thus block vIF2α phosphorylation.

Whereas vaccinia virus mutants lacking K3L remain competent to infect cells as well as mice, vaccinia virus mutants lacking E3L are nonpathogenic in wild-type mice (Brandt and Jacobs 2001; Rice et al. 2011). Consistent with the sensitivity of pkr−/− mice to infection with ΔE3L virus (Rice et al. 2011), replication of the mutant virus in HeLa cells was rescued by knockdown of PKR expression by RNA interference (Zhang et al. 2008). It is noteworthy that the vaccinia virus E3L protein has been reported to inhibit PKR by sequestering dsRNA activators of the kinase and by directly binding to PKR and inhibiting kinase activation (Chang et al. 1992; Davies et al. 1993; Romano et al. 1998; Sharp et al. 1998). The E3L protein consists of two functional domains: an N-terminal domain (NTD) containing a Z-DNA/RNA binding domain (ZBD), and a C-terminal dsRBD (Fig. 2A, left). The dsRBD shares a dsRNA-binding motif (dsRBM) found in many proteins that interact with dsRNA, including PKR, Staufen, RNase III, and XbrpA (St Johnston et al. 1992; Ryter and Schultz 1998; Saunders and Barber 2003). The dsRBD is required for E3L protein binding to dsRNA in vitro (Chang and Jacobs 1993), and point mutations in the E3L dsRBD that severely impair its ability to bind dsRNA in vitro (Ho and Shuman 1996) not only abolish its ability to inhibit PKR in yeast (Romano et al. 1998), but also eliminate its ability to prevent dsRNA-, 5′-triphosphate single-stranded RNA-, and DNA-induced activation of the IFN-α gene in human cell lines (Marq et al. 2009). In addition to its role in nucleic acid–induced cellular responses, the dsRNA-binding activity of E3L is also critical for vaccinia virus pathogenesis in mouse as well as virus replication in cell culture (Chang et al. 1995; Shors et al. 1997; Brandt and Jacobs 2001).

In support of the dsRNA sequestration model for E3L inhibition of PKR, heterologous dsRNA-binding proteins such as reovirus σ3 (Beattie et al. 1995), rotavirus NSP3 (Langland et al. 1994), human cytomegalovirus TRS1 (Bierle et al. 2013), TAR RNA-binding protein TRBP (Park et al. 1994), and *Escherichia coli* RNase III (Shors and Jacobs 1997) were shown to functionally complement an E3L-deleted virus. However, E3L was also reported to directly bind to PKR. It was proposed that in the presence of dsRNA, the dsRBDs of PKR and E3L heterodimerize, and that this interaction enables the NTD of E3L to interact with the PKR kinase domain (Romano et al. 1998; Sharp et al. 1998). Furthermore, mutations that disrupt dsRNA binding to E3L were proposed to block PKR inhibition by preventing both dsRNA sequestration and PKR–E3L heterodimer formation (Chang and Jacobs 1993; Davies et al. 1993; Shors and Jacobs 1997; Romano et al. 1998). In contrast, mutations in the E3L NTD that do not impair dsRNA binding were proposed to weaken PKR inhibition by interfering with the interaction between the E3L NTD and the PKR kinase domain (Romano et al. 1998).

While the nucleic acid–binding activity of the E3L dsRBD plays an important role for the virus in sequestering dsRNA and in mediating dsRNA-dependent heterodimerization of E3L with PKR, the function of the N-terminal ZBD of E3L is not as well understood. Moreover, it is not clear whether the Z-DNA binding function of the ZBD is important for E3L inhibition of PKR. The E3L ZBD belongs to the functional Zα family of Z-DNA binding domains based on its sequence similarity with the first (Zα) domain of two ZBDs found in the enzyme adenosine deaminase acting on RNA1 (ADAR1). In addition to ADAR1, ZBDs are also found in DNA-dependent activator of interferon regulatory factor DAI (also known as DLM-1 or ZBP1) (Schwartz et al. 2001), and the protein kinase PKZ (Rothenburg et al. 2005). Interestingly, the ZBD of E3L can be functionally replaced by the heterologous ZBDs from ADAR1 (ZαADAR1) or DAI (ZαDAI) in mouse infection assays (Kim et al. 2003).

The crystal structures of ZBDs from human ADAR1 (Schwartz et al. 1999), mouse DLM-1 (Schwartz et al. 2001), the yatapoxvirus E3L ortholog (Ha et al. 2004), the cyripinid herpesvirus 3 ORF112 protein (Tome et al. 2013), human DAI (ZBP1) (Ha et al. 2008), and zebrafish PKZ (de Rosa et al. 2013) have been solved either free or in complex with Z-DNA. Consistent with amino acid sequence identities of ∼30%, the ZBDs fold into similar α/β structures with three α-helices bundled on one side of a triple-stranded β-sheet (Schwartz et al. 2001; Ha et al. 2004, 2008; de Rosa et al. 2013). Moreover, the solution structure of vaccinia virus E3L revealed a similar topology (Kahmann et al. 2004). Notably, five of the nine residues that contact Z-DNA are highly conserved across the family of E3L proteins and related ZBDs (Fig. 1; Kim et al. 2003; Ha et al. 2004). Mutation of these five residues substantially impaired or abolished the Z-DNA binding activity of the isolated first ZBD (Zα domain) from ADAR1 (Schade et al. 1999; Kim et al. 2003). Similarly, mutation of these same residues impaired the ability of an ADAR-E3L chimera, in which the vaccinia E3L ZBD was replaced by the Zα domain of ADAR1, to functionally substitute for E3L and maintain vaccinia virus pathogenicity in mice (Kim et al. 2003). Likewise, substitution of alanine in place of vaccinia E3L residues Y48 and P63 markedly reduced viral lethality following intracranial inoculation in mice (Kim et al. 2003). In contrast to these studies revealing the importance of the E3L Z-DNA binding activity, the ability of E3L to antagonize the innate immune response in plasmacytid dendritic cells is dependent on the ZBD, but not its Z-DNA binding activity (Dai et al. 2011; Cao et al. 2012). Taken together, these data reveal important roles for the E3L ZBD-and its Z-DNA binding activity; however, the contribution of Z-DNA binding to E3L inhibition of PKR is not known.

Here, we use yeast, mammalian cell, and in vitro assays to further examine the inhibition of PKR by a poxviral E3L...
**TABLE 1.** Multiple sequence alignment of the N-terminal domains of the indicated poxvirus E3L proteins and the first Z-DNA binding (ZBD or Zα) domains from mouse ADAR1 and mouse DLM1 (ZBD1). Sequences were aligned using the program PROMALS3D (Pei et al. 2008). (Red) Consensus:究竟…

Contact Z-DNA: **...**

| Protein                     | Consensus: | Sites of E3L-L36R and -L50P mutations that impair PKR inhibitory, but not Z-DNA binding | Sites of E3L-N44D,Y48A & W66A mutations that impair Z-DNA binding |  |
|-----------------------------|------------|------------------------------------------------------------------------------------------|-----------------------------------------------------------------|----|
| E3L variola virus           | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L vaccinia-WR virus       | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L bovine Papul. Stem. v.  | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L ectromelia virus        | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L cowpox virus            | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L lataropox virus         | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L camelpox virus          | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L cowpox-G9 virus         | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L rabbitpox virus         | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L horepox virus           | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L Yaba monkey tumor v.    | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L Yaba-like disease v.    | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L tanapox virus           | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L deerpox-84 virus        | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L goatpox virus           | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L Lumpy skin disease v.   | **...**    | **...**                                                                                  | **...**                                                         |  |
| E3L anigenpox virus         | **...**    | **...**                                                                                  | **...**                                                         |  |
| ADAR1 Zα M. musculus       | **...**    | **...**                                                                                  | **...**                                                         |  |
| DLM1 Zα M. musculus        | **...**    | **...**                                                                                  | **...**                                                         |  |

**FIGURE 1.** Multiple sequence alignment of the N-terminal domains of the indicated poxvirus E3L proteins and the first Z-DNA binding (ZBD or Zα) domains from mouse ADAR1 and mouse DLM1 (ZBD1). Sequences were aligned using the program PROMALS3D (Pei et al. 2008). (Red) Consensus:究竟…

Contact Z-DNA: **...**

protein. As previous studies revealed that PKR has evolved under selection by viral inhibitors (Elde et al. 2009; Rothenburg et al. 2009) and that various poxvirus E3L proteins differ in their ability to inhibit human PKR (Myskiw et al. 2011), we focused our studies on variola virus E3L and its natural target human PKR. Our selection of variola E3L was also spurred by the critical biodefense importance of variola virus. We show that variola virus E3L protein, hereafter referred to as E3L, readily suppresses PKR toxicity in yeast, PKR inhibition of reporter gene expression in mammalian cells, and PKR phosphorylation of eIF2α in vitro. Whereas deletion of the NTD abolishes E3L inhibition of PKR in yeast, in vitro, and in mammalian cells treated with dsRNA, point mutations that substantially reduce Z-DNA binding to E3L do not impair PKR inhibition. Thus, these results support a nucleic acid–binding-independent function of the E3L NTD required for inhibition of PKR.

**RESULTS**

**NTD of E3L is required for inhibition of PKR in yeast and in vitro**

Variola and vaccinia virus E3L proteins differ at 10 out of the 190 residues in these proteins. To test the ability of variola virus E3L to inhibit PKR, we generated a variola virus E3L expression vector by introducing the 10 required mutations (see Materials and Methods) in a vaccinia virus E3L expression vector. Yeast strain H2544, which expresses human PKR under the control of a galactose-inducible GAL–CYC1 hybrid promoter, was transformed with either empty vector or plasmids that express N-terminally HA-tagged vaccinia or variola E3L also under the control of the GAL–CYC1 promoter. High-level expression of PKR is toxic in yeast due to phosphorylation of eIF2α on Ser51 (Chong et al. 1992; Vazquez de Aldana et al. 1993; Kagawa-Kobayashi et al. 1997). Consistently, the empty vector transformant of H2544 failed to grow when PKR expression was induced on galactose medium (Fig. 2B, right panel, row 1). While high-level expression of E3L was slightly toxic in yeast even in the absence of PKR (Fig. 2B, left panel, row 2 vs. 1), expression of variola E3L suppressed the lethal phenotype associated with high-level expression of PKR (Fig. 2B, row 2). Deletion of the NTD (residues 1–80) from E3L, generating E3L–Δ80, abolished its ability to suppress PKR toxicity in yeast (Fig. 2B, row 3). Previous studies have shown that a C-terminal truncated version of PKR, retaining residues 1–258 and consisting of the two dsRBDs (PKR-ΔKD), functioned in a dominant-negative manner and blocked PKR toxicity in yeast (Fig. 2B, row 4). This inhibitory function was attributed to formation of heterodimers between PKR-ΔKD and the full-length kinase (Romano et al. 1995). We generated a corresponding version of E3L in which the E3L dsRBD was substituted for the NTD, generating a protein...
with two dsRBDS (E3L-R:R) (Fig. 2B, row 5). In contrast to PKR-ΔKD, E3L-R:R failed to inhibit PKR toxicity in yeast (Fig. 2B, row 5). Although we cannot assess whether the dsRBDS are folding properly in E3L-R:R, this result suggests that simply linking two dsRBDS in a single protein, as found in E3L-R:R, is not sufficient to confer PKR inhibitory function. Western analyses demonstrated that both E3L-Δ80 and E3L-R:R were readily expressed in yeast, albeit at lower levels than E3L (Fig. 2C). While the reduced expression of the E3L mutants limits the strength of the conclusion that a second copy of the E3L dsRBD cannot functionally replace the E3L NTD, the in vitro experiments described below support the conclusion that the NTD of E3L is critical for the inhibition of PKR.

To directly examine E3L inhibition of PKR, we carried out in vitro kinase assays. Flag-tagged PKR was purified from yeast and tested for the ability to phosphorylate recombinant Histag-eIF2α (residues 1–200) purified from E. coli. As the PKR expressed in yeast is purified in a pre-activated form, already phosphorylated on Thr446 (Dey et al. 2005a,b), it was unnecessary to add dsRNA to activate the kinase. Moreover, inhibitors that simply sequester dsRNA or other activators are unlikely to inhibit this pre-activated kinase. The in vitro kinase assays were conducted in the presence or absence of GST, GST-E3L, or GST-E3L-Δ80, and eIF2α phosphorylation was monitored by immunoblot analysis using antibodies that specifically detect phospho-Ser51 on eIF2α. As shown in Figure 3A, PKR readily phosphorylated eIF2α on Ser51 (lane 1). Pre-incubation of PKR with a 10-fold molar excess of GST-E3L (lane 5), but not GST (lane 3), substantially reduced eIF2α phosphorylation. In contrast, pre-incubation of Flag-PKR with GST-E3L-Δ80 failed to inhibit eIF2α phosphorylation (Fig. 3A, lanes 6, 7). Thus, deletion of the NTD abolished the ability of E3L to inhibit PKR both in yeast (Fig. 2B) and in vitro (Fig. 3A). Immunoblot analyses confirmed that equivalent amounts of PKR, eIF2α, and the various GST or GST-E3L fusion proteins were used in each assay (Fig. 3A). Moreover, unlike GST, which as shown in Figure 3B was largely localized in the unbound fraction, both GST-E3L and GST-E3L-Δ80 readily bound to an agarose affinity resin containing poly(I:C) (polynosinic–polycytidylic acid, a double-stranded RNA homopolymer known to activate PKR) (Minks et al. 1979).

Thus, the loss of PKR inhibitory activity by GST-E3L-Δ80 was not due to impaired dsRNA-binding ability. Taken together, these in vitro experiments and yeast studies demonstrate that the E3L NTD plays a critical role in PKR inhibition, and they indicate that E3L can inhibit PKR following kinase activation.

**Abundance of dsRNA alters the requirement of the E3L NTD for PKR inhibition in mammalian cells**

PKR knockdown HeLa cells (PKRkd), which express an shRNA that targets the PKR mRNA and efficiently abolishes endogenous PKR expression and blocks dsRNA-induced eIF2α phosphorylation (Zhang and Samuel 2007), were used to test the role of the E3L NTD for inhibition of PKR in mammalian cells. Previous studies demonstrated that PKR expression could be restored in the PKRkd cells using PKR expression constructs in which the shRNA target sequence was altered (Rothenburg et al. 2009). The PKRkd cells were transiently transfected with a luciferase reporter plasmid, the knockdown-resistant PKR construct, and various C-terminally HA-tagged E3L expression vectors. As expected, expression of PKR impaired luciferase expression (Fig. 4A) and increased eIF2α phosphorylation (Fig. 4C, lane 2 vs. 1). Cotransfection of E3L with PKR increased luciferase activity (Fig. 4A) and lowered eIF2α phosphorylation (Fig. 4C, lane 3), indicating that E3L was able to inhibit PKR in mammalian cells. Surprisingly, unlike the yeast and in vitro studies, E3L-Δ80 functioned like wild-type (WT) E3L in the mammalian cell transfection assays, increasing luciferase activity (Fig. 4A) and lowering eIF2α phosphorylation levels in the transfected cells (Fig. 4C, lane 4). Western analyses revealed that WT E3L, which, as previously reported (Watson et al.
Abundance of dsRNA alters the requirement of the E3L ZBD for PKR inhibition in mammalian cells. (A) Human HeLa PKRΔ80 cells were cotransfected with expression vectors for luciferase, an empty vector or knockdown-resistant human PKR (100 ng), and C-terminally HA-tagged variola E3L-HA or E3L-Δ80-HA (400 ng). After 40 h, cells were harvested, lysed, and extracts were assayed for luciferase activity. Luciferase activity was normalized to the transfections containing PKR but lacking E3L. Error bars indicate the standard deviation for three independent transfections. (B) Analysis of luciferase activity following cotransfection of PKRΔ80 cells with poly(I:C) (500 ng) and the expression vectors for luciferase, vector or PKR (100 ng), and E3L-HA or E3L-Δ80-HA (400 ng) as described in panel A. (C) Immunoblot analyses of WCEs from cells transfected as described in panels A and B with or without poly(I:C) and the indicated expression vectors for PKR, E3L, or E3L-Δ80. Blots were probed with antibodies specific for phosphorylated Ser51 in eIF2α (top panel), polyclonal antisera against human eIF2α (second panel), monoclonal anti-PKR antibody (third panel), and monoclonal anti-HA antibody to detect HA-tagged E3L (fourth panel). Note that E3L-HA resolves as two species on SDS-PAGE (see text). (*) E3L-Δ80 comigrates with a nonspecific protein that cross-reacts with the anti-HA antibody. (Bottom panel) Relative level of eIF2α-P to total eIF2α was determined for three independent experiments using quantitative densitometry and Image J software (NIH) and normalized to the vector transformant.
Consistent with the notion that PKR activators are abundant in yeast, the kinase-dead variant PKR-K296R, which functions in a dominant-negative manner in mammalian cells (Barber et al. 1993), fails to inhibit PKR in yeast (Romano et al. 1995). It is presumed that PKR-K296R effectively sequesters the low-abundance activators in mammalian cells, but is unable to sequester the activators present in yeast cells.

To test the hypothesis that the E3L NTD is critical for inhibition of PKR when activators are abundant, we increased the levels of PKR activators in the PKRkd cells by cotransfecting poly(I:C) together with PKR and E3L. As shown in Figure 4B, the ability of E3L-Δ80 to suppress the inhibition of luciferase expression by PKR was diminished in the presence of added poly(I:C) dsRNA. E3L readily restored luciferase expression both in the presence (Fig. 4B) and absence (Fig. 4A) of added dsRNA, and likewise lowered eIF2α phosphorylation levels (Fig. 4C, lane 3 vs. 2, lane 9 vs. 8). In contrast, E3L-Δ80 only partially restored luciferase expression (Fig. 4B) and failed to lower eIF2α phosphorylation when poly(I:C) was cotransfected with PKR (lane 10). Thus, increasing the abundance of PKR activators in mammalian cells mimics the conditions in yeast, or in vitro with pre-activated PKR, and reveals the importance of the NTD for E3L inhibition of PKR.

**ZBD functions independent of Z-DNA binding in PKR inhibition**

As the best-characterized function of a ZBD is to bind Z-DNA/RNA, it was intriguing to determine whether this nucleic acid–binding activity is required for E3L inhibition of PKR. It is noteworthy that the dsRNA–binding activity of E3L is critical for inhibition of PKR in vitro (Chang and Jacobs 1993). Moreover, mutation of Lys167 in the E3L dsRBD impairs dsRNA binding (Chang and Jacobs 1993; Ho and Shuman 1996; Romano et al. 1998) and the ability of E3L to inhibit PKR toxicity in yeast (Romano et al. 1998), and to block PKR phosphorylation of eIF2α and inhibition of luciferase expression in mammalian cells (data not shown). Analysis of the sequence and structure of ZBDs from viral E3L proteins as well as from mouse ADAR1 and DLM-1 (Fig. 1) identified seven perfectly conserved residues including the variola E3L residues Asn44, Tyr48, and Pro63. These three residues are among the set of residues that directly contact Z-DNA in the co-crystal structure of the ADAR1 ZBD bound to Z-DNA (Schwartz et al. 1999), and mutation of the corresponding residues (N173D, Y177A, P192A) in the Za domain of ADAR1 impaired Z-DNA binding (Schade et al. 1999; Kim et al. 2003). Furthermore, similar mutations significantly reduced the ability of a ZaΔADAR1–RBDΔ3AL chimera, consisting of the ADAR1 Za domain fused to the E3L RBD to substitute for E3L in a viral pathogenesis assay in mice (Kim et al. 2003). Since these three Z-DNA binding residues are identical in E3L and ADAR1, we introduced the corresponding N44D, Y48A, and P63A mutations in the E3L ZBD and examined the ability of the mutant proteins to inhibit PKR in yeast. As shown in Figure 5A, expression of WT E3L or the N44D, Y48A, or P63A mutants of E3L suppressed the lethal effects associated with high-level expression of PKR in yeast. Combining these mutations to create double (N44D,Y48A [DM] and Y48A,P63A) or triple (N44D,Y48A, P63A) mutants failed to abolish the ability of E3L to suppress PKR toxicity in yeast (Fig. 5A). Consistent with their potent activity to inhibit PKR, the E3L mutants were all well expressed in yeast (Fig. 5B).

To directly examine the impact of the ZBD mutations on the Z-DNA binding activity of E3L, recombinant GST-E3L fusion proteins were purified from E. coli and tested for Z-DNA binding ability. Electrophoretic gel mobility shift assays were used to monitor the binding of the E3L proteins with [32P]–labeled partially brominated synthetic poly(dG:dC), which was previously shown to be stable in the Z conformation (Zhang et al. 1992; Kim et al. 2006). As shown in Figure 6A, incubation of the poly(dG:dC) with increasing amounts of GST-E3L resulted in the appearance of multiple slower-migrating species consistent with the binding of GST-E3L to Z-DNA. The exact number of complexes observed varied...
in different experiments (see Fig. 6A,B), perhaps reflecting heterogeneity in the Z-DNA or the proteins. However, the appearance of the upper complex in Figure 6A, and to a lesser extent the lower complex, was reduced when unlabeled Z-DNA (lanes 6,7), but not unlabeled B-DNA (lanes 8,9), was added as a competitor in the binding reaction. These results demonstrate that E3L binds specifically to Z-DNA. Next, this assay was used to monitor the Z-DNA binding activity of the E3L-N44D,Y48A double mutant (E3L-DM). Note that the corresponding single mutations in the Zα domain of ADAR1 were shown to impair Z-DNA binding activity (Schade et al. 1999; Kim et al. 2003). Consistent with the results of these previous studies, the Z-DNA binding ability of GST-E3L (Fig. 6B, lane 1) was severely impaired, with reduced detection of all Z-DNA–protein complexes, following deletion of the ZBD (E3L-Δ80, lane 2) or by the N44D,Y48A double mutation (E3L-DM, lane 5). In contrast to these ZBD mutations, the K167A mutation in the E3L dsRBD, which abolishes dsRNA binding (Romano et al. 1998), did not impair Z-DNA binding (lane 3). These data demonstrate that the E3L-N44D,Y48A double mutant is impaired in binding to Z-DNA.

Having shown that E3L-DM fails to bind Z-DNA, yet retains PKR inhibitory activity in yeast, we next examined the ability of this mutant to directly inhibit PKR using in vitro kinase assays. PKR purified from yeast was incubated with GST-E3L, GST-E3L-DM, or GST-E3L-K167A and then tested for eIF2α phosphorylation. As shown in Figure 6C, GST-E3L and GST-E3L-DM, but not GST-E3L-K167A, substantially reduced the ability of PKR to phosphorylate eIF2α (lanes 2–4). Thus, the Z-DNA binding ability of E3L, but not its dsRNA-binding ability, is dispensable for PKR inhibition in vitro.

Next, we examined the importance of Z-DNA binding for E3L inhibition of PKR in mammalian cells. Consistent with our previous data showing that deletion of the NTD did not impair Z-DNA binding (lanes 3), these data demonstrate that the E3L-N44D,Y48A double mutant is impaired in binding to Z-DNA.

FIGURE 6. E3L inhibition of PKR in vitro and in mammalian cells treated with dsRNA is independent of Z-DNA binding activity. (A) Electrophoretic mobility shift assay of GST-E3L binding to the Z-DNA substrate [32P]-labeled d(G:5BrC)20. Indicated concentrations of GST-E3L were incubated with labeled d(G:5BrC)20 in the presence of 30 nM or 300 nM unlabeled Z-DNA (lanes 6,7) or B-DNA (lanes 8,9). Positions of the GST-E3L-Z-DNA complex and free probe are indicated. (B) Z-DNA binding activity of E3L mutants. GST or the indicated GST-E3L derivative (5 μM) was incubated with the Z-DNA substrate and analyzed as described in A. (C) Z-DNA binding mutations do not impair E3L inhibition of PKR in vitro. The indicated GST or GST-E3L derivatives were incubated with purified Flag-Hisα-PKR and then mixed with Hisα-eIF2α1–200 and ATP. Reactions were resolved by SDS-PAGE and subjected to immunoblot analysis using antibodies specific for the Ser51-phosphorylated form of eIF2α (top panel) or using polyclonal antiserum against total yeast eIF2α (bottom panel). The relative level of eIF2α-P to total eIF2α (2α-P/2α) was determined for three independent experiments using quantitative densitometry and Image J software (NIH) and normalized to the vector transformant.

in different experiments (see Fig. 6A,B), perhaps reflecting heterogeneity in the Z-DNA or the proteins. However, the appearance of the upper complex in Figure 6A, and to a lesser extent the lower complex, was reduced when unlabeled Z-DNA (lanes 6,7), but not unlabeled B-DNA (lanes 8,9), was added as a competitor in the binding reaction. These results demonstrate that E3L binds specifically to Z-DNA. Next, this assay was used to monitor the Z-DNA binding activity of the E3L-N44D,Y48A double mutant (E3L-DM). Note that the corresponding single mutations in the Zα domain of ADAR1 were shown to impair Z-DNA binding activity (Schade et al. 1999; Kim et al. 2003). Consistent with the results of these previous studies, the Z-DNA binding ability of GST-E3L (Fig. 6B, lane 1) was severely impaired, with reduced detection of all Z-DNA–protein complexes, following deletion of the ZBD (E3L-Δ80, lane 2) or by the N44D,Y48A double mutation (E3L-DM, lane 5). In contrast to these ZBD mutations, the K167A mutation in the E3L dsRBD, which abolishes dsRNA binding (Romano et al. 1998), did not impair Z-DNA binding (lane 3). These data demonstrate that the E3L-N44D,Y48A double mutant is impaired in binding to Z-DNA.

Having shown that E3L-DM fails to bind Z-DNA, yet retains PKR inhibitory activity in yeast, we next examined the ability of this mutant to directly inhibit PKR using in vitro kinase assays. PKR purified from yeast was incubated with GST-E3L, GST-E3L-DM, or GST-E3L-K167A and then tested for eIF2α phosphorylation. As shown in Figure 6C, GST-E3L and GST-E3L-DM, but not GST-E3L-K167A, substantially reduced the ability of PKR to phosphorylate eIF2α (lanes 2–4). Thus, the Z-DNA binding ability of E3L, but not its dsRNA-binding ability, is dispensable for PKR inhibition in vitro.

Next, we examined the importance of Z-DNA binding for E3L inhibition of PKR in mammalian cells. Consistent with our previous data showing that deletion of the NTD did...
not impair PKR inhibition in this assay system (Fig. 4A), E3L-DM (N44D,Y48A) functioned like WT E3L and lowered eIF2α phosphorylation levels and restored luciferase expression in cells cotransfected with the PKR expression vector (data not shown). Importantly, unlike E3L-A80, but similar to WT E3L, E3L-DM retained the ability to suppress the PKR inhibition of luciferase reporter expression in cells treated with excess dsRNA (Fig. 6D). Likewise, both WT E3L and E3L-DM lowered the level of eIF2α phosphorylation in the cells treated with dsRNA (Fig. 6E, lanes 3,4 vs. 2), although for unknown reasons E3L-DM was expressed at higher levels than WT E3L in the transfected cells (Fig. 6E). While this enhanced expression of E3L-DM could contribute to its retention of PKR inhibitory activity, these data together with the yeast (Fig. 5A) and in vitro (Fig. 6C) data demonstrate that the PKR inhibitory activity of E3L requires the NTD but is independent of Z-DNA binding.

Identification of E3L ZBD mutations that impair PKR inhibition but not Z-DNA binding

Having found that the E3L NTD, but not its Z-DNA binding activity, is required for inhibition of PKR, we subjected the E3L NTD to random mutagenesis and screened for mutants that lost the ability to inhibit PKR in yeast. Error-prone PCR was used to generate a library of E3L mutants containing mutations in the NTD. The mutant library was introduced into the yeast strain expressing PKR under the control of the galactose-inducible promoter. Approximately 2000 transformants were patched on SD medium where PKR and E3L are not expressed and then replica-printed to galactose medium to induce PKR and E3L expression. The E3L plasmid was isolated from the colonies growing on SD medium that failed to grow on galactose medium. Following retesting and sequencing, only two of the mutants were found to contain single amino acid changes. These two mutants, E3L-L36R and E3L-L50P, were selected for further characterization. As shown in Figure 7A, E3L-L36R and E3L-L50P, unlike wild-type E3L, were unable to restore the growth of yeast coexpressing PKR. The mutant proteins were expressed at levels comparable to wild-type E3L in cells lacking PKR (Fig. 7B). The expression of E3L-L36R and E3L-L50P was slightly lower than E3L in cells expressing PKR, consistent with the notion that induction of PKR inhib-
PKR inhibition (Fig. 7D, lanes 3, 5, 6; Romano et al. 1998). Finally, the E3L-L50P mutant was tested in the mammalian cell transfection assay. In contrast to WT E3L, E3L-L50P was unable to lower eIF2α phosphorylation (Fig. 6E, lane 5) and restore luciferase expression in cells transfected with the PKR expression vector and dsRNA (Fig. 6D). Thus, the L50P mutation impairs E3L inhibition of PKR in both yeast and mammalian cell assays. The finding that the E3L-L36R and E3L-L50P mutants retain Z-DNA binding activity, yet fail to inhibit PKR, further demonstrates that the PKR inhibitory function of the E3L NTD is distinct from its nucleic acid–binding activity.

DISCUSSION

Successful viral replication is dependent on subversion of host cell defense mechanisms. It was previously shown that vaccinia virus expresses an inhibitor of the interferon-induced host cell kinase PKR, and this activity was eventually linked to the viral E3L protein (Rice and Kerr 1984; Whitaker-Dowling and Youngner 1984; Chang et al. 1992). As the E3L inhibition of PKR could be overcome by adding excess dsRNA (Davies et al. 1993; Jagus and Gray 1994), and as E3L was found to contain a dsRBD at its C terminus (Chang et al. 1992), it was concluded that E3L inhibits PKR by sequestering dsRNA activators of the kinase (Davies et al. 1993). Consistent with this idea, point mutations that blocked dsRNA binding to E3L likewise interfered with the ability of E3L to inhibit PKR (Shors et al. 1997). At odds with this simple model for E3L inhibition of PKR, truncation mutations that removed the NTD of E3L (E3L–Δ80), yet left the dsRBD intact, blocked the ability of E3L to suppress PKR toxicity in yeast (Romano et al. 1998). The inability of this truncated E3L mutant, which retains dsRNA-binding activity (Fig. 3B), to inhibit PKR indicates that sequestration of dsRNA activators is not sufficient for E3L inhibition of PKR. In accord with this latter idea, it was shown that E3L could bind directly to PKR, leading to the hypothesis that E3L inhibits PKR both by sequestering dsRNA activators of the kinase and by forming inactive heterodimers with PKR (Romano et al. 1998; Sharp et al. 1998). In this study, we further explored the role of the E3L NTD and found that the ZBD, but not its nucleic acid–binding function, is important for PKR inhibition. It is noteworthy that the E3L ZBD, but not its Z-DNA binding activity, is likewise required to block the induction of the innate immune response and cytokine production in poxvirus-infected plasmacytoid dendritic cells (Dai et al. 2011; Cao et al. 2012). In contrast, the Z-DNA binding activity of E3L was previously shown to be critical for viral pathogenicity (Kim et al. 2003). Taken together, we propose that the E3L ZBD performs a nucleic acid–binding independent function to inhibit PKR and perhaps other components of the innate immune response, whereas the Z-DNA binding activity of this domain is critical for E3L subversion of other cellular anti-viral defenses.

Our studies provide several insights into the mechanism of E3L inhibition of PKR. We show that the abundance of PKR activators determines the requirements for PKR inhibition. Whereas full-length E3L impairs the phosphorylation of eIF2α and the inhibition of reporter gene expression by PKR in either untreated mammalian cells or in cells treated with dsRNA (Fig. 4), the inhibition of PKR in mammalian cells by E3L–Δ80 (Fig. 4) or the Z-DNA binding mutant E3L-DM (Fig. 6) was reversed by increasing the abundance of dsRNA. We propose that under native conditions, where dsRNA activators are not abundant, the E3L RBD is sufficient to sequester the dsRNA activators and prevent PKR activation. However, under conditions where dsRNA activators are plentiful, E3L inhibition of PKR is dependent on both the RBD and the ZBD. The requirement for the E3L ZBD observed in mammalian cells treated with dsRNA mimics the E3L requirements for PKR inhibition in yeast, supporting the notion that PKR activators are abundant in yeast. While it could have been proposed that the ZBD enhances the nucleic acid–binding properties of E3L and thus enables the protein to effectively titrate the activators, the fact that the Z-DNA binding defective mutant E3L-DM effectively inhibits PKR both in yeast and in mammalian cells treated with excess dsRNA (Figs. 5, 6) suggests that the ZBD contributes to PKR inhibition in a manner independent of nucleic acid binding. Further supporting this notion, we identified the E3L-L36R and E3L-L50P mutants that readily bind dsRNA as well as Z-DNA but are unable to inhibit PKR in yeast or in mammalian cells treated with dsRNA. Taken together, our molecular dissection of E3L has revealed a PKR inhibitory function in the NTD that is distinct from its nucleic acid–binding activity.

We reasoned that the NTD could contribute to PKR inhibition by promoting heterodimer formation with PKR. Previously, the N-terminal half of vaccinia virus E3L was shown to interact with the PKR KD in yeast two-hybrid assays (Romano et al. 1998; Sharp et al. 1998). We found that full-length variola E3L interacted with full-length PKR in yeast two-hybrid assays (data not shown). Moreover, this interaction was not disrupted by the N44D and Y48A mutations that impair Z-DNA binding. However, as both full-length E3L and PKR contain dsRBDs, the two-hybrid interaction we detected could be mediated solely by the dsRBD in E3L. Unfortunately, we were unable to test the effect of the ZBD mutations on binding of the isolated E3L NTD to PKR. While the variola E3L ZBD interacted with full-length PKR in a yeast two-hybrid assay, thus eliminating the concern about dsRNA-mediated interactions, the ZBD mutations caused the DNA-binding domain fusion to promiscuously activate the two-hybrid reporter in the absence of a partner activation domain (data not shown). Moreover, and consistent with previous reports (Romano et al. 1998; Sharp et al. 1998), issues with nonspecific binding prevented the use of pull-down assays to monitor an interaction between the E3L ZBD and PKR. Thus, while we were unable to directly
show that disruption of its Z-DNA binding activity does not affect the ability of the E3L NTD to bind PKR, our data are consistent with the notion that the E3L NTD contributes to PKR inhibition in a manner that is independent of its nucleic acid–binding function and is presumably mediated by direct protein–protein interactions with PKR.

The E3L gene is critical for vaccinia virus replication in mammalian cells (Chang et al. 1995) and for viral pathogenicity in mice (Brandt and Jacobs 2001). As knockdown of PKR expression in HeLa cells complemented the defect in virus yield and virus-induced apoptosis by E3L-deleted vaccinia virus (Zhang et al. 2008), it is clear that PKR is an important target of E3L. Consistently, the attenuated phenotype of E3L-deleted virus following intratracheal infection of wild-type mice was partially reversed upon infection of mice lacking PKR (Rice et al. 2011). The fact that deletion of PKR does not fully rescue the pathogenicity of the E3L-deleted virus (Rice and Kerr 1984; Xiang et al. 2002) indicates that E3L likely targets other components of the cellular anti-viral defense, such as the mitochondrial anti-viral signaling (MAVS) pathway for interferon induction (Deng et al. 2008). In support of this idea, the E3L-deleted virus was pathogenic in mice lacking both PKR and the anti-viral endonuclease RNAse L (Rice et al. 2011). However, the PKR and RNase L pathways are likely not the sole targets of E3L, as E3L-deleted virus failed to replicate to high titers in mice triply lacking PKR, RNase L, and the interferon-induced Mx1 protein (Xiang et al. 2002). Our results are also consistent with the notion that E3L inhibits anti-viral targets other than PKR. While our data show that the Z-DNA binding domain of E3L is not required for PKR inhibition in transfected mammalian cells (not treated with excess dsRNA) and that the Z-DNA binding activity is dispensable for PKR inhibition in yeast, in vitro, and in mammalian cells treated with dsRNA, mutations impairing the Z-DNA binding function of E3L were previously shown to reduce the pathogenicity of vaccinia virus in mice (Kim et al. 2003). Thus, while the E3L ZBD has a nucleic acid–binding independent function to inhibit PKR, the nucleic binding property of this domain is important for viral pathogenicity. Taken together, these results lead us to suggest that the Z-DNA binding properties of E3L are necessary for the inhibition of other components of the interferon-induced anti-viral defense, consistent with the recent report that the pathogenicity of ΔE3L vaccinia virus is fully rescued by knockout of the type I interferon receptor in mice (White and Jacobs 2012).

**MATERIALS AND METHODS**

**Plasmids**

Plasmids used in this study are listed in Table 1. Construction of the plasmid pEMBLyex4–E3L (p2245), which expresses hemagglutinin (HA) epitope-tagged vaccinia E3L under the control of a yeast GAL–CYC1 hybrid promoter, was described previously (Romano et al. 1998). A Sacl–BamH1 fragment carrying HA-E3L was subcloned in pUC19 to generate pC2490. Site-directed mutagenesis was performed at 10 residues to convert vaccinia E3L to variola E3L (N11D, T21N, I24L, A27V, A30V, D78M, A79T, G115N V121L, and R130K), generating pUC19-varE3L (pC2491). The Sacl/BamH1 fragment from pC2491 was subcloned in pEMBLyex4 to generate pEMBLyex4-varE3L (pC2492). Site-directed mutagenesis of pC2492 was performed to generate E3L single (K167A, W66A, N44D, Y48A, P63A, L36R, and L50P), double (N44D/Y48A; Y48A/P63A), and triple (N44D/Y48A/P63A) point mutants.

| Table 1. Plasmids used in this study |
|-------------------------------------|
| **Plasmid** # | **Description** | **Source** |
| pC2490 | vacc. E3L in pUC19 | Romano et al. (1998) |
| pC2491 | E3L in pUC19 | This study |
| pC2492 | E3L in pEMBLyex4 | This study |
| pC4419 | E3L-ΔBO in pEMBLyex4 | This study |
| pC4420 | E3L-RR in pEMBLyex4 | This study |
| pC4462 | PKR-ΔK in pEMBLyex4 | This study |
| pC4421 | E3L-W66A in pEMBLyex4 | This study |
| pC4422 | E3L-K167A in pEMBLyex4 | This study |
| pC4423 | E3L-N44D in pEMBLyex4 | This study |
| pC4424 | E3L-Y48A in pEMBLyex4 | This study |
| pC4426 | E3L-N44D,Y48A in pEMBLyex4 | This study |
| pC4427 | E3L-Y48A,P63A in pEMBLyex4 | This study |
| pC4428 | E3L-N44D,Y48A,P63A in pEMBLyex4 | This study |
| pC4429 | E3L-L36R in pEMBLyex4 | This study |
| pC4430 | E3L-50P in pEMBLyex4 | This study |
| pC2812 | pGEX-E3L | This study |
| pC4432 | pGEX-E3L-ΔBO | This study |
| pC4433 | pGEX-E3L-DM | This study |
| pC4434 | pGEX-E3L-K167A | This study |
| pC4435 | pGEX-E3L-L36R | This study |
| pC4436 | pGEX-E3L-L50P | This study |
| pC4437 | pGEX-E3L-ZBD | This study |
| pC4438 | pGEX-E3L-W66A-ZBD | This study |
| pC4439 | pGEX-E3L-N44D,Y48A-ZBD | This study |
| pC4440 | pGEX-E3L-L36R-ZBD | This study |
| pC4441 | pGEX-E3L-L50P-ZBD | This study |
| pC2252 | His6-eIF2α | Dey et al. (2003b) |
| pC1685 |FLAG-His6-PKR | Dey et al. (2003b) |
| pC4460 | Knockdown-resistant human PKR in pSG5 | Rothenburg et al. (2009) |
| pC4442 | E3L in pSG5 | This study |
| pC4443 | E3L-ΔBO in pSG5 | This study |
| pC4444 | E3L-N44D in pSG5 | This study |
| pC4445 | E3L-Y48A in pSG5 | This study |
| pC4446 | E3L-P63A in pSG5 | This study |
| pC4447 | E3L-N44D,Y48A in pSG5 | This study |
| pC4448 | E3L-N44D,Y48A,P63A in pSG5 | This study |
| pC4449 | E3L-W66A in pSG5 | This study |
| pC4450 | E3L-K167A in pSG5 | This study |
| pC4451 | E3L-L36R in pSG5 | This study |
| pC4452 | E3L-L50P in pSG5 | This study |

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by inserting the relevant PCR products between the BamHI and EcoRI sites of the vector pGEX-5x-3. Point mutations in GST-E3L were generated by site-directed mutagenesis of pC2812.

Construction of knockdown-resistant human PKR (pSG5-PKR) was described previously (Rothenburg et al. 2009), pSG5-E3L-HA (pC4442) and pSG5-E3L-Δ80-HA (pC4443) were generated by PCR inserting a C-terminal HA-epitope and cloning between the EcoRI and BamHI sites of pSG5. Point mutants pSG5-E3L-N44D (pC4446), pSG5-E3L-N44D,Y48A (pC4445), pSG5-E3L-P63A (pC4447), pSG5-E3L-N44D,Y48A,P63A (pC4448), pSG5-E3L-W66A (pC4449), and pSG5-E3L-K167A (pC4450) were generated by site-directed mutagenesis using pC4442 as a template.

Yeast strains

Construction of yeast strains H2544 (MATa ura3-52 leu2-3 112 trp1-Δ63 gen2a GAL1-CYC1-PKR, LEU2+) and J110 (MATa ura3-52 leu2-3 112 trp1-Δ63 gen2a LEU2+) has been described previously (Romano et al. 1995; Seo et al. 2008). Strain H2766 (MATa ura3-52 leu2-3 112 trp1-Δ63 gen2a LEU2+) was used for expression and purification of Flag- and His6-tagged PKR (Krishnamoorthy et al. 2001).

Immunoblot analysis assay

Yeast transformants were grown in SC-Ura medium (synthetic minimal medium with all amino acids) overnight at 30°C to saturation, diluted in fresh medium to OD600 ≈ 0.1, and grown to 30°C until OD600 ≈ 0.6. Cells were harvested and transferred to SCGal-Ura medium (SC-Ura except 10% galactose) for 2 h at 30°C to induce PKR and E3L expression. Cells were harvested, and whole cell extracts (WCEs) were prepared as described previously (Dey et al. 2005a). Five micrograms of WCEs was resolved by SDS-PAGE and subjected to immunoblot analysis using rabbit phosphospecific antibodies directed against phosphorylated Ser51 of eIF2α (BioSource International), rabbit polyclonal antiserum that recognizes yeast eIF2α irrespective of Ser51 phosphorylation, mouse monoclonal anti-HA (Roche), or mouse monoclonal anti-PKR antibodies (R&D Systems). Immune complexes were detected by enhanced chemiluminescence.

dsRNA-binding assay

Polyinosinic–polyribocytidylic acid bound to agarose [Poly(I:C) agarose, Pharmacia] was equilibrated in Buffer A (50 mM Tris–HCl at pH 8.0, 150 mM NaCl) and incubated with 2 µg of purified GST-E3L variants in 200 µL of Buffer A for 1 h at 4°C. Agarose beads were collected, washed five times with Buffer B (50 mM Tris–HCl at pH 8.0, 150 mM NaCl, 0.1% Triton X-100), resuspended in 30 µL of SDS sample buffer, and boiled for 5 min. Proteins were resolved by 10% SDS-PAGE, transferred to nitrocellulose membranes, and immunoblotted with anti-GST antibody (Santa Cruz Biotechnology). Immune complexes were detected by enhanced chemiluminescence.

Recombinant protein expression and purification

Flag- and His6-tagged PKR was expressed in yeast strain H2766. Yeast transformants were grown to saturation in 5 mL of SC-Ura medium at 30°C. Cells were harvested, transferred to 50 mL of SGR medium (SC medium containing 10% galactose and 2% raffinose), and incubated for 48 h at 30°C. Cells were then harvested, washed with ice-cold water, and resuspended in Flag Binding Buffer (20 mM Tris–HCl at pH 7.5, 500 mM NaCl, 0.1% Triton X-100, 1 mM phenylmethylsulfonyl fluoride [PMSF], 10 mM NaF, 50 mM β-glycerophosphate, 125 µM Na2VO4, and complete protease inhibitor mixture [Roche]). Cells were broken by high speed mixing with glass beads on a vortex. WCEs were mixed with 100 µL of anti-Flag-M2 affinity gel (50% slurry; Sigma-Aldrich) pre-equilibrated with Flag Binding Buffer, and incubated with gentle rocking for 1 h at 4°C. Immune complexes were collected by centrifugation and washed three times with Flag Binding Buffer and once with 1× Kinase Buffer (20 mM Tris–HCl at pH 8.0, 50 mM KCl, 25 mM MgCl2, and 1 mM PMSF). Bound proteins were eluted with 25 µL Flag-peptide (500 µg/mL), and individual aliquots were used for immunoblot analysis and for protein kinase assays with recombinant yeast eIF2α as described previously (Dey et al. 2005a).

GST fusion proteins and His6-eIF2α1–200 were expressed in E. coli strain BL21(DE3) and purified using glutathione Sepharose 4B and Ni2+–NTA resin, respectively, according to the manufacturer’s protocol.

In vitro kinase assay

Purified Flag-His6-PKR, GST-E3L fusion proteins, and His6-eIF2α were incubated with 0.5 mM ATP for 10 min at room temperature in Kinase Buffer (20 mM Tris–HCl at pH 8.0, 50 mM KCl, 25 mM MgCl2, and 1 mM PMSF). Reaction products were separated by SDS-PAGE, and eIF2α phosphorylation was monitored by immunoblot analysis using phospho-specific and total eIF2α antibodies.

Mammalian cell luciferase assay and immunoblot analysis

HeLa PKRkd cells (Zhang and Samuel 2007) were grown in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% (v/v) fetal bovine serum and 1 µg/mL puromycin (Sigma-Aldrich) at 37°C in 5% CO2. For luciferase assays, 3 × 105 cells were seeded in a 24-well plate, incubated for 24 h, and then transfected at 70%–80% confluence with 2 µL of GenJet (SigmaGen Laboratories), 0.05 µg of pGL3 Promoter luciferase plasmid (Promega), 0.1 µg of PKR expression vector, and varying amounts of variola E3L plasmids using Lipofectamine 2000 reagent (Invitrogen) according to the manufacturer’s instructions. Transfections were performed in triplicate, and cells were harvested 48 h post-transfection using passive lysis buffer (Promega). Luciferase activity was determined using the Luciferase Assay System (Promega) and a luminometer (Centro XS Microplate Reader LB 960).

For Western blot analysis, 1 or 5 × 105 cells were seeded into 12-well or six-well plates, respectively, a day before transfection. Cells were cotransfected with GenJet reagent (SigmaGen Laboratories), PKR expression vector, and E3L or mutant expression vectors. At 24 h post-transfection, WCEs were obtained by lysis in 50 mM Tris (pH 8.0), 150 mM NaCl, 1% Triton X-100, 100 mM sodium fluoride, 50 mM β-glycerophosphate, and 10 mM sodium pyrophosphate. Six micrograms of WCEs were subjected to SDS-PAGE, and immunoblot analysis was performed using monoclonal anti-HA.
(Roche) or anti-PKR (R&D Systems) antibodies, or with polyclonal antibodies against total eIF2α (Santa Cruz Biotechnology) or rabbit phosphospecific antibodies directed against Ser51 on eIF2α. Immune complexes were detected by enhanced chemiluminescence.

Z-DNA binding assay

The synthetic oligomer (5′-dCG[5Br-dC]GGCGG[5Br-dC]GGCGG[5Br-dC]GGCGG[5Br-dC]GGCGG[5Br-dC]GGCGG[5Br-dC]GGCGG[5Br-dC]G), which is stable in the left-handed Z-DNA conformation (Schwartz et al. 1999; Kim et al. 2006), was annealed (95°C for 2 min, ramp-cooled to 25°C over a period of 45 min) and 5′-end-labeled using poly(adenylate kinase and [γ-32P]ATP). The labeled DNA was incubated with GST-E3L proteins in Binding Buffer (10 mM Tris–HCl at pH 7.5, 10 mM MgCl2, 20 mM NaCl, 5 mM DTT, 0.1 mg/mL BSA, and 0.5 μg/mL calf thymus DNA) for 10 min in a final volume of 15 μL. The mixture was separated on a 6% TBE–Polyacrylamide Gel at 10 V/cm using 0.5× TBE as running buffer, and protein–DNA complexes were detected by autoradiography. Synthetic annealed B-DNA (TCCCTCTGTAGTTGGTCGCCC GGTTCATCCTCGGAT) was used as a competitor.

E3L random mutagenesis

An SmaI restriction site was engineered between amino acid residues 80 and 81 of the yeast E3L expression vector pC2492. Error-prone PCR (GeneMorph II Random Mutagenesis Kit, Agilent Technologies, Inc.) was used to amplify the SacI–SmaI fragment encoding E3L residues 1–80 under low-stringency conditions (one or two mutations per cycle), and the products were subcloned to pC2492. Sequencing of randomly selected clones from the mutagenesis library revealed that 70% of the plasmids contained mutations. The mutant library was transformed in the PKR strain H2544, and about 2000 independent yeast transformants were patched on glucose medium and then replica-printed to galactose medium to induce E3L and PKR expression. The E3L plasmid was isolated from about 2000 independent yeast transformants, and then E3L single mutants were generated by site-directed mutagenesis of pC2492 and tested in H2544.

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REFERENCES

Barber GN, Wambach M, Wong ML, Dever TE, Hinnebusch AG, Katze MG. 1993. Translational regulation by the interferon-induced double-stranded-RNA-activated 68-kDa protein kinase. Proc Natl Acad Sci 90: 4621–4625.

Beattie E, Denzler KL, Taggart J, Perkus ME, Paolotti E, Jacobs BL. 1995. Reversal of the interferon-sensitive phenotype of a vaccinia virus lacking E3L by expression of the reovirus S4 gene. J Virol 69: 499–505.

Bierle CJ, Semmens KM, Geballe AP. 2013. Double-stranded RNA binding by the human cytomegalovirus PKR antagonist TRS1. Virology 442: 28–37.

Brandt TA, Jacobs BL. 2001. Both carboxy- and amino-terminal domains of the vaccinia virus interferon resistance gene, E3L, are required for pathogenesis in a mouse model. J Virol 75: 850–856.

Bratke KA, McLysaght A, Rothenburg S. 2013. A survey of host range genes in poxvirus genomes. Infect Genet Evol 14: 406–425.

Cao H, Dai P, Wang W, Li H, Yuan J, Wang F, Fang CM, Pitha PM, Liu J, Condit RC, et al. 2012. Innate immune response of human plasma- cytoid dendritic cells to poxvirus infection is subverted by vaccinia E3 via its Z-DNA/RNA binding domain. PLoS ONE 7: e63823.

Carroll K, Elroy-Stein O, Moss B, Jagus R. 1993. Recombinant vaccinia virus K3L gene product prevents activation of double-stranded RNA-dependent, initiation factor 2α-specific protein kinase. J Biol Chem 268: 12837–12842.

Chang HW, Jacobs BL. 1993. Identification of a conserved motif that is necessary for binding of the vaccinia virus E3L gene products to double-stranded RNA. Virology 194: 537–547.

Chang HW, Watson JC, Jacobs BL. 1992. The E3L gene of vaccinia virus encodes an inhibitor of the interferon-induced, double-stranded RNA-dependent protein kinase. Proc Natl Acad Sci 89: 4825–4829.

Chang HW, Uribe LH, Jacobs BL. 1995. Rescue of vaccinia virus lacking the E3L gene by mutants of E3L. J Virol 69: 6605–6608.

Chang KL, Feng L, Schappert K, Meurs E, Donahue TF, Friesen JD, Hovanessian AG, Williams BR. 1992. Human p68 kinase exhibits growth suppression in yeast and homology to the translational regulator GCN2. EMBO J 11: 1553–1562.

Dai P, Cao H, Merghoub T, Avogadi F, Wang W, Parikh T, Fang CM, Pitha PM, Fitzgerald KA, Rahman MM, et al. 2011. Myxoma virus induces type I interferon production in murine plasmacytoid dendritic cells via a TLR9/MyD88, IRF3/IRF7–, and IFNAR-dependent pathway. J Virol 85: 10814–10825.

Dar AC, Sicheri F. 2002. X-ray crystal structure and functional analysis of vaccinia virus K3L reveals molecular determinants for PKR subversion and substrate recognition. Mol Cell 10: 295–305.

Dar AC, Dever TE, Sicheri F. 2005. Higher-order substrate recognition of eIF2α by the RNA-dependent protein kinase PKR. Cell 122: 887–900.

Davies MV, Elroy-Stein O, Jagus R, Moss B, Kaufman RJ. 1992. The vaccinia virus K3L gene product potentiates translation by inhibiting double-stranded-RNA-activated protein kinase and phosphorylation of the α subunit of eukaryotic initiation factor 2. J Virol 66: 1943–1950.

Davies MV, Chang HW, Jacobs BL, Kaufman RJ. 1993. The E3L and K3L vaccinia virus gene products stimulate translation through inhibition of the double-stranded RNA-dependent protein kinase by different mechanisms. J Virol 67: 1688–1692.

de Rosa M, Zacarias S, Athanasiadis A. 2013. Structural basis for Z-DNA binding and stabilization by the zebrafish Z-DNA-dependent protein kinase PKZ. Nucleic Acids Res 41: 9924–9933.

Deng L, Dai P, Parikh T, Cao H, Bhoj V, Sun Q, Chen Z, Merghoub T, Houghton A, Shuman S. 2008. Vaccinia virus subverts a mitochondrial antiviral signaling protein-dependent innate immune response in keratinocytes through its double-stranded RNA binding protein, E3, J Virol 82: 10735–10746.

Dever TE, Chen JJ, Barber GN, Cigan AM, Feng L, Donahue TF, London IM, Katze MG, Hinnebusch AG. 1993. Mammalian eukaryotic initiation factor 2α substrate recognition. Mol Cell Biol 13: 2501–2507.

Dey M, Cao C, Dar AC, Tamura T, Ozato K, Sicheri F, Dever TE. 2005a. Mechanistic link between PKR dimerization, autophosphorylation, and eIF2α substrate recognition. Cell 122: 901–913.

Dey M, Trieselmann B, Locke EG, Lu J, Cao C, Dar AC, Krishnamoorthy T, Dong J, Sicheri F, Dever TE. 2005b. PKR and GCN2 kinases and guanine nucleotide exchange factor eukaryotic translation initiation factor 2B (eIF2B) recognize overlapping surfaces on eIF2a. Mol Cell Biol 25: 3063–3075.
Pei J, Kim BH, Grishin NV. 2008. PROMALS3D: A tool for multiple protein sequence and structure alignments. *Nucleic Acids Res* 36: 2295–2300.

Rice AP, Kerr IM. 1984. Interferon-mediated, double-stranded RNA-dependent protein kinase is inhibited in extracts from vaccinia virus-infected cells. *J Virol* 50: 229–236.

Rice AD, Turner PC, Embury JE, Moldawer LL, Baker HV, Moyer RW. 2011. Roles of vaccinia virus genes E3L and K3L and host genes PKR and RNase L during intratracheal infection of C57BL/6 mice. *J Virol* 85: 550–567.

Romano PR, Green SR, Barber GN, Mathews MB, Hinnebusch AG. 1995. Structural requirements for double-stranded RNA binding, dimerization, and activation of the human eIF2α-kine DAI in *Saccharomyces cerevisiae*. *Mol Cell Biol* 15: 365–378.

Romano PR, Zhang F, Tan SL, Garcia-Barrio MT, Katze MG, Dever TE, Hinnebusch AG. 1998. Inhibition of double-stranded RNA-dependent protein kinase PKR by vaccinia virus E3: Role of complex formation and the E3 N-terminal domain. *Mol Cell Biol* 18: 7304–7316.

Rothenburg S, Deigendesch N, Dittmar K, Koch-Nolte F, Haag F, Lowenhaupt K, Rich A. 2005. A PKR-like eukaryotic initiation factor 2α kinase from zebrafish contains Z-DNA binding domains instead of dsRNA binding domains. *Proc Natl Acad Sci* 102: 1602–1607.

Rothenburg S, Seo EJ, Gibbs JS, Dever TE, Dittmar K. 2009. Rapid evolution of protein kinase PKR alters sensitivity to viral inhibitors. *Nat Struct Mol Biol* 16: 63–70.

Rothenburg S, Chinchar VG, Dever TE. 2011. Characterization of a ranavirus inhibitor of the antiviral protein kinase PKR. *BMCMicrobiol* 11: S6.

Ryter JM, Schultz SC. 1998. Molecular basis of double-stranded RNA-protein interactions: Structure of a dsRNA-binding domain complexed with dsRNA. *EMBO J* 17: 7505–7513.

Saunders LR, Barber GN. 2003. The dsRNA binding protein family: Critical roles, diverse cellular functions. *FASEB J* 17: 961–983.

Schade M, Turner CJ, Lowenhaupt K, Rich A, Herbert A. 1999. Structure–function analysis of the Z-DNA-binding domain Zα of dsRNA adenosine deaminase type I reveals similarity to the (α+β) family of helix–turn–helix proteins. *EMBO J* 18: 470–479.

Schwartz T, Rouš MA, Lowenhaupt K, Herbert A, Rich A. 1999. Crystal structure of the Zα domain of the human editing enzyme ADAR1 bound to left-handed Z-DNA. *Science* 284: 1841–1845.

Schwartz T, Behlke J, Lowenhaupt K, Heinemann U, Rich A. 2001. Structure of the DLM–1–Z-DNA complex reveals a conserved family of Z-DNA-binding proteins. *Nat Struct Biol* 8: 761–765.

Seo EJ, Liu F, Kawagishi-Kobayashi M, Ung TL, Cao C, Dar AC, Schier E, Dever TE. 2008. Protein kinase PKR mutants resistant to the poxvirus pseudosubstrate K3L protein. *Proc Natl Acad Sci* 105: 16894–16899.

Sharp TV, Schwemmle M, Jeffrey I, Laing K, Mellor H, Proud CG, Hille K, Clemens MI. 1993. Comparative analysis of the regulation of the interferon-inducible protein kinase PKR by Epstein-Barr virus RNAs EBER-1 and EBER-2 and adenosine VAI RNA. *Nucleic Acids Res* 21: 4483–4490.

Sharp TV, Moonan F, Romashko A, Joshi B, Barber GN, Jagus R. 1998. The vaccinia virus E3L gene product interacts with both the regulatory and the substrate binding regions of PKR: Implications for PKR autoregulation. *Virology* 250: 302–315.

Shors T, Jacobs BL. 1997. Complementation of deletion of the vaccinia virus E3L gene by the *Escherichia coli* RNase III gene. *Virology* 227: 77–87.

Shors T, Kibler KV, Perkins KB, Seidler-Wulff R, Banaszak MP, Jacobs BL. 1997. Complementation of vaccinia virus deleted of the E3L gene by mutant of E3L. *Virology* 239: 269–276.

St Johnston D, Brown NH, Gall JG, Jantsch M. 1992. A conserved double-stranded RNA-binding domain. *Proc Natl Acad Sci* 89: 10979–10983.

Tome AR, Kus K, Correia S, Paulo LM, Zacarias S, de Rosa M, Figueiredo D, Parkhouse RM, Athanasiadis A. 2013. Crystal
structure of a poxvirus-like Zα domain from cyprinid herpesvirus 3. 
*J Virol* **87**: 3998–4004.

Vázquez de Aldana CR, Dever TE, Hinnebusch AG. 1993. Mutations in the α subunit of eukaryotic translation initiation factor 2 (eIF-2α) that overcome the inhibitory effect of eIF-2α phosphorylation on translation initiation. *Proc Natl Acad Sci* **90**: 7215–7219.

Watson JC, Chang HW, Jacobs BL. 1991. Characterization of a vaccinia virus-encoded double-stranded RNA-binding protein that may be involved in inhibition of the double-stranded RNA-dependent protein kinase. *Virology* **185**: 206–216.

Whitaker-Dowling P, Youngner JS. 1984. Characterization of a specific kinase inhibitory factor produced by vaccinia virus which inhibits the interferon-induced protein kinase. *Virology* **137**: 171–181.

White SD, Jacobs BL. 2012. The amino terminus of the vaccinia virus E3 protein is necessary to inhibit the interferon response. *J Virol* **86**: 5895–5904.

Xiang Y, Condit RC, Vijaysri S, Jacobs B, Williams BR, Silverman RH. 2002. Blockade of interferon induction and action by the E3L double-stranded RNA binding proteins of vaccinia virus. *J Virol* **76**: 5251–5259.

Yue Z, Shatkin AJ. 1997. Double-stranded RNA-dependent protein kinase (PKR) is regulated by reovirus structural proteins. *Virology* **234**: 364–371.

Yuwen H,cox JH, Yewedell JW, Bennink JR, Moss B. 1993. Nuclear localization of a double-stranded RNA-binding protein encoded by the vaccinia virus E3L gene. *Virology* **195**: 732–744.

Zhang P, Samuel CE. 2007. Protein kinase PKR plays a stimulus- and virus-dependent role in apoptotic death and virus multiplication in human cells. *J Virol* **81**: 8192–8200.

Zhang S, Lockshin C, Herbert A, Winter E, Rich A. 1992. Zuotin, a putative Z-DNA binding protein in *Saccharomyces cerevisiae*. *EMBO J* **11**: 3787–3796.

Zhang F, Romano PR, Nagamura-Inoue T, Tian B, Dever TE, Mathews MB, Ozato K, Hinnebusch AG. 2001. Binding of double-stranded RNA to protein kinase PKR is required for dimerization and promotes critical autophosphorylation events in the activation loop. *J Biol Chem* **276**: 24946–24958.

Zhang P, Jacobs BL, Samuel CE. 2008. Loss of protein kinase PKR expression in human HeLa cells complements the vaccinia virus E3L deletion mutant phenotype by restoration of viral protein synthesis. *J Virol* **82**: 840–848.