Constitutive mTORC1 activation by a herpesvirus Akt surrogate stimulates mRNA translation and viral replication

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All viruses require cellular ribosomes to translate their mRNAs. Viruses producing methyl-7 (m7) GTP-capped mRNAs, like Herpes Simplex Virus-1 (HSV-1), stimulate cap-dependent translation by activating mTORC1 to inhibit the translational repressor 4E-binding protein 1 (4E-BP1). Here, we establish that the HSV-1 kinase Us3 masquerades as Akt to activate mTORC1. Remarkably, Us3 displays no sequence homology with the cellular kinase Akt, yet directly phosphorylates tuberous sclerosis complex 2 (TSC2) on the same sites as Akt. TSC2 depletion rescued Us3-deficient virus replication, establishing that Us3 enhances replication by phosphorylating TSC2 to constitutively activate mTORC1, effectively bypassing S6K-mediated feedback inhibition. Moreover, Us3 stimulated Akt substrate phosphorylation in infected cells, including FOXO1 and GSK3. Thus, HSV-1 encodes an Akt surrogate with overlapping substrate specificity to activate mTORC1, stimulating translation and virus replication. This establishes Us3 as a unique viral kinase with promising drug development potential.

[Keywords: Translational control; herpesvirus replication; viral kinase; mTOR activation; Akt signaling]

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Recruiting 40S ribosome subunits to mRNA is a critical step controlling translation. For eukaryotic mRNAs, the vast majority of which have methyl-7 (m7) GTP caps at their 5’ termini, this involves assembling the multi-subunit, cap-binding translation initiation factor eIF4F (Sonenberg and Hinnebusch 2009). Comprised of the cap recognition subunit eIF4E and an RNA helicase eIF4A bound to the eIF4G assembly platform, eIF4F associates with eIF3-bound 40S subunits and localizes them to the mRNA 5’ end. Interaction of eIF4E with eIF4G represents a pivotal step controlling eIF4F assembly, and is regulated in part by small 4E-binding proteins (4E-BPs) that sequester eIF4E, inhibit the eIF4E–eIF4G association, and thereby repress mRNA translation. Phosphorylation of 4E-BP by the cellular kinase mTORC1 releases eIF4E, a prerequisite for eIF4G binding and forming an active eIF4F complex. Integration of diverse signaling inputs by mTORC1 facilitates translation initiation factor complex remodeling and rapid response to nutrient availability, energy supplies, and a battery of environmental stresses, including viral infection. (Fig. 1A; Mohr et al. 2007; Buchkovich et al. 2008; Ma and Blenis 2009)

Herpesvirus’ lifelong latency within their cellular hosts is punctuated by productive viral replication episodes (Roizman et al. 2007). Successful completion of their replicative cycle during these outbreaks requires conscripting host ribosomes to translate viral mRNAs. This is achieved in part by viral functions that commandeer cellular translational control functions, one of which is ICP6. As a Herpes Simplex Virus-1 [HSV-1]-encoded eIF4G-associated protein that stimulates eIF4G binding to eIF4E, ICP6 mediates eIF4F assembly and eIF4E phosphorylation by the eIF4G-associated kinase Mnk. However, despite stimulating eIF4F assembly, ICP6 did not detectably contribute to the fate of the translational repressor 4E-BP1 in infected cells [Walsh and Mohr 2006]

Productive replication of representative α [HSV-1] and β [HCMV] herpesviruses, together with γ herpesvirus reactivation from latency, inactivate 4E-BP1 (Kudchodkar et al. 2004; Walsh and Mohr 2004; Walsh et al. 2005;
Moorman et al. 2008; Arias et al. 2009). In HSV-1-infected primary human cells, this involves 4E-BP1 hyperphosphorylation, which requires viral gene expression, is sensitive to the mTORC1 inhibitor rapamycin, and is followed by proteasome-mediated degradation [Walsh and Mohr 2004]. Here we show that phosphorylation of the 4E-BP1 translational repressor, an mTORC1 substrate, is required for eIF4F assembly and normal viral protein synthesis levels. Furthermore, we define the mechanism of mTORC1 activation and 4E-BP1 inactivation in HSV-1-infected cells by demonstrating that the virus-encoded Us3 Ser/Thr protein kinase is required. Surprisingly, Us3 catalytic function acts analogously to the cellular kinase Akt to promote mTORC1-mediated 4E-BP1 phosphorylation, enabling Akt-independent mTORC1 activation. By stimulating site-specific phosphorylation of the tuberous sclerosis complex (TSC) TSC2 subunit at S939 and T1462, Us3 ensures constitutive mTORC1 activation in infected cells by antagonizing TSC antiviral activity. Finally, siRNA-mediated TSC2 depletion selectively enhanced Us3-deficient virus replication, providing genetic evidence that TSC2 restricts mTOR activation, which is normally antagonized by Us3 in HSV-1-infected cells.

This establishes that the HSV-1 Us3 protein kinase overcomes host TSC antiviral activity and remodels host translational initiation factor complexes, defining a new strategy whereby protein kinases bearing little or no primary sequence homology with Akt regulate mTORC1. Remarkably, by encoding a constitutively active kinase to mimic Akt and activate mTORC1, HSV-1 bypasses feedback inhibitory circuits, antagonizes TSC2, and ensures that the 4E-BP1 translational repressor remains inactive throughout the productive growth cycle.

Results

Regulation of eIF4F assembly and protein synthesis in HSV-1-infected cells by mTORC1-mediated 4E-BP1 phosphorylation

Although 4E-BP1 phosphorylation in HSV-1-infected primary human cells was sensitive to the mTORC1 inhibitor rapamycin, significant reductions in viral replication or translation rates were not detected [Walsh and Mohr 2004]. Recently developed mTOR active site inhibitors (PP242 and Torin) are superior to rapamycin in repressing translation by preventing 4E-BP1 phosphorylation [Feldman et al. 2009; Thoreen et al. 2009]. Indeed, Torin reduced HSV-1 replication in immortalized mouse cells in a 4E-BP1-dependent manner (Moorman and Shenk 2010), and PP242 reduced HSV-1 replication in primary human fibroblasts (Supplemental Fig. S1). Thus, wild-type levels of HSV-1 replication and protein synthesis in normal,
primary human cells required mTOR catalytic activity. However, the extent to which site-specific 4E-BP1 phosphorylation by mTOR restricted HSV-1 replication by regulating infected cell protein synthesis and, most significantly, the underlying mechanism responsible for activating mTOR in infected cells remained unexplored.

To determine if preventing site-specific 4E-BP1 phosphorylation reduced protein synthesis in HSV-1-infected cells, normal human dermal fibroblasts [NHDFs] stably expressing epitope-tagged [Flag] wild-type 4E-BP1 or a double-mutant derivative (AA) with alanine substituted for each of the critical T37 and T46 phosphoacceptor residues were isolated. Unable to be inactivated by mTORC1, the AA mutant behaved as a constitutively activated translational repressor [Gingras et al. 1999]. Growth-arrested NHDFs expressing wild-type or AA 4E-BP1 were either mock-infected or HSV-1-infected, and metabolically pulse-labeled proteins were analyzed by SDS-PAGE (Fig. 1B). Quantification of \(^{35}\)S amino acid incorporation in six independent experiments by TCA precipitation revealed a modest 10%–15% decrease in uninfected cells expressing AA versus wild-type 4E-BP1 (Fig. 1B, lanes 1,3). Under these conditions, in growth-arrested primary fibroblasts, the introduced wild-type and AA 4E-BP1 were both active repressors that bound to eIF4E in infected NHDFs even though endogenous eIF4E (Fig. 1C). This provides evidence that 4E-BP1 were associated with eIF4E in mock-infected wild-type. Importantly, repairing the Us3 allele markedly decreased 4E-BP1 hyperphosphorylation, correcting the mutant phenotype and proving it was due to the Us3 deficiency (Fig. 2A).

4E-BP1 phosphorylation is followed by proteasome-mediated degradation in HSV-1-infected and uninfected cells [Walsh and Mohr 2004; Elia et al. 2008; Braunstein et al. 2009]. Fractionation identical samples by SDS-PAGE in a 7.5% gel easily monitors steady-state 4E-BP1 abundance, as phosphorylated isoforms are not resolved and 4E-BP1 migrates as a single band. Indeed, overall 4E-BP1 abundance was reduced in wild-type virus-infected compared with mock-infected cells, whereas steady-state control antigen (actin) levels remained unchanged. However, 4E-BP1 levels in ΔUs3-infected cells remained similar to mock-infected cells (Fig. 2B). Thus, Us3-deficient viruses are unable to inactivate 4E-BP1 by promoting its phosphorylation and degradation in infected cells.

To determine if Us3 Ser/Thr kinase activity was required to inactivate 4E-BP1 in infected cells, NHDFs were either mock-infected or infected with ΔUs3, a recombinant virus expressing catalytically inactive Us3 (K220A), the Us3 repair virus, or wild-type HSV-1. Analysis of 4E-BP1 phosphorylation by SDS-PAGE followed by immunoblotting revealed that K220A and ΔUs3 were similarly defective in promoting 4E-BP1 phosphorylation (Fig. 2C). Finally, the ability of Us3 to interfere with eIF4E binding to the 4E-BP1 repressor in HSV-1-infected cells was evaluated by collecting eIF4E-bound proteins in cell-free extracts prepared from mock-infected or HSV-1-infected NHDFs on mGTP-Sepharose. While the amount of 4E-BP1 associated with cap-bound eIF4E dropped significantly in lysates from wild-type or Us3 repair virus-infected cells, amounts of 4E-BP1 associated with eIF4F were either mock-infected or HSV-1-infected, normal human dermal fibroblasts [NHDFs] stably expressing epitope-tagged [Flag] wild-type 4E-BP1 or a double-mutant derivative (AA) with alanine substituted for each of the critical T37 and T46 phosphoacceptor residues were isolated. Unable to be inactivated by mTORC1, the AA mutant behaved as a constitutively activated translational repressor [Gingras et al. 1999]. Growth-arrested NHDFs expressing wild-type or AA 4E-BP1 were either mock-infected or HSV-1-infected, and metabolically pulse-labeled proteins were analyzed by SDS-PAGE (Fig. 1B). Quantification of \(^{35}\)S amino acid incorporation in six independent experiments by TCA precipitation revealed a modest 10%–15% decrease in uninfected cells expressing AA versus wild-type 4E-BP1 (Fig. 1B, lanes 1,3). Under these conditions, in growth-arrested primary fibroblasts, the introduced wild-type and AA 4E-BP1 were both active repressors that bound to eIF4E in infected NHDFs even though endogenous eIF4E (Fig. 1C). This provides evidence that 4E-BP1 were associated with eIF4E in mock-infected wild-type. Importantly, repairing the Us3 allele markedly decreased 4E-BP1 hyperphosphorylation, correcting the mutant phenotype and proving it was due to the Us3 deficiency (Fig. 2A).

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cap-bound eIF4E in cells infected with either ΔUs3 or K220A remained similar to those in mock-infected cells (Fig. 2D). Furthermore, stimulation of eIF4G binding to eIF4E was detected only in cells infected with wild-type or Us3 repair viruses (Fig. 2D). Taken together, these results indicated that Us3 Ser/Thr kinase activity was required to stimulate 4E-BP1 phosphorylation in HSV-1-infected cells, thereby inactivating the translational repressor and disrupting the 4E-BP1-eIF4E complex to release the cap-binding, translation initiation factor eIF4E. Finally, Us3-mediated release of eIF4E from 4E-BP1 was required to recruit eIF4G, a critical eIF4F subunit required to recruit 40S ribosomes and initiate translation.

Us3 acts analogously to Akt to stimulate mTORC1 signaling

HSV-1 infection activated Akt in established, transformed cell lines (Benetti and Roizman 2006) and growth-arrested primary human cells (Fig. 3B). To determine if Akt is required to inactivate 4E-BP1 in HSV-1-infected cells, the extent of 4E-BP1 phosphorylation was evaluated after treatment with Akt inhibitor VIII, a highly selective allosteric Akt activation inhibitor (Fig. 3B; Calleja et al. 2009). Growth-arrested NHDFs were mock-infected or infected with wild-type or ΔUs3 virus and treated with...
Akt inhibitor, and, at 15 hpi, total protein was collected, fractionated by SDS-PAGE, and analyzed by immunoblotting. Significantly, phosphorylated 4E-BP1 and p70 S6K, another mTORC1 substrate, still accumulated in wild-type virus-infected cells treated with the Akt inhibitor [Fig. 3C, lanes 6 vs. 7], indicating that Akt activity is not required for HSV-1 to stimulate mTOR. p70 S6K and 4E-BP1 phosphorylation, however, was only partially impaired in ΔUs3 or K220A Us3-infected cells [Fig. 3C, lanes 3,4] compared with those infected with HSV-1-expressing wild-type Us3 [wild-type or ΔUs3 repair in Fig. 3C, lanes 2,5], suggesting that HSV-1 activates mTOR through both Akt-dependent and Akt-independent mechanisms. The Akt inhibitor did not detectably effect viral thymidine kinase [tk] accumulation, indicating that lack of detectable 4E-BP1 phosphorylation in ΔUs3 or K220A Us3 virus-infected cells did not result from nonspecific inhibition of viral gene expression [Fig. 3C]. Thus, 4E-BP1 and p70 S6K phosphorylation in HSV-1-infected cells treated with an Akt inhibitor was Us3-dependent [Fig. 3C, lanes 8,9], suggesting that Us3 activates mTOR to phosphorylate p70 S6K and 4E-BP1 by phosphorylating a target downstream from and independent of Akt.

Lying downstream from Akt, TSC is a Rheb•GTPase-activating protein [GAP] that negatively regulates mTORC1. TSC is a heterodimer composed of TSC1 and TSC2 subunits. TSC2 phosphorylation [T1462, S939] by Akt inhibits TSC GAP activity and allows Rheb•GTP accumulation and subsequent mTORC1 activation [Fig 3A; Ma and Blenis 2009]. To determine if TSC2 phosphorylation on sites normally targeted by Akt occurred in HSV-1-infected cells, total protein isolated from NHDFs infected with wild-type or Us3 mutant viruses was analyzed by immunoblotting. In DMSO-treated cultures, reduced TSC2 p-T1462 levels accumulated in cells infected with Us3 mutant viruses [ΔUs3 or K220A], and this correlated with increased 4E-BP1 hypophosphorylation [Fig. 3C, lanes 3,4 vs. 2,5]. However, TSC2 p-T1462 and hyperphosphorylated 4E-BP1 were detected only in Akt inhibitor-treated cultures infected with functional Us3-expressing viruses [wild-type or Us3 repair in Fig 3C, lanes 7,10 vs. lanes 8,9]. Similar results were observed using antisera specific for TSC2 p-S939 [data not shown]. This demonstrates that TSC2 Thr1462/Ser939 phosphorylation was Us3 Ser/Thr kinase-dependent in HSV-1-infected cells treated with an Akt inhibitor. Moreover, it suggested that TSC2 may be a substrate for the HSV-1-encoded kinase Us3, and this could be the mechanism underlying mTORC1 activation in infected cells.

Us3 phosphorylates TSC2 in vivo and in vitro

While phospho-TSC2 accumulation in HSV-1-infected cells was Us3-dependent, Us3 is itself regulated by another viral protein kinase [Kato et al. 2006]. In addition, profound physiological changes in infected cells could conceivably contribute to TSC2 phosphorylation. To determine whether Us3 expression was sufficient to promote TSC2 phosphorylation in the absence of a concurrent HSV-1 infection, Us3 was transiently expressed in uninfected cells. Following transfection of U2OS cells with epitope-tagged vector expressing wild-type Us3, K220A catalytically inactive Us3, or empty vector, total protein was collected and fractionated by SDS-PAGE, and TSC2 phosphorylation along with Us3 expression were evaluated by immunoblotting [Fig. 4A]. Despite being expressed to equivalent levels, wild-type Us3 was more effective than the K220A variant in promoting p-T1462 TSC2 accumulation in DMSO-treated cells [Fig. 4A, lanes 1–3]. However, background p-S939 phosphorylation in this transformed, established osteosarcoma cell line was high. To eliminate any contribution of endogenous, constitutively activated Akt, the experiment was performed in parallel cultures treated with the PI3K inhibitor wortmannin. Under these conditions [1] Akt activation measured by S473 phosphorylation was not detected [Fig. 4A, lanes 4–6], and [2] p-TSC2 [T1462/S939] was detected only in cultures expressing wild-type Us3. Thus, Us3 expression was sufficient to promote p-TSC2 accumulation in uninfected cells; moreover, TSC2 phosphorylation in transiently transfected cells required Us3 Ser/Thr kinase catalytic activity.

To refine requirements for TSC2 phosphorylation, Us3-dependent TSC2 phosphorylation was reconstituted in vitro using immunopurified components. Transiently expressed epitope-tagged Us3 and TSC2 were immunopurified using protein G-Sepharose bound to anti-Flag antibody. TSC2-expressing cultures were wortmannin-treated to enrich for unphosphorylated TSC2. Furthermore, to reduce nonspecific protein binding, immune complexes were washed with high-salt buffer prior to equilibrating the beads in kinase reaction buffer. A similar procedure established that TSC2 was a bona fide Akt substrate [Manning et al. 2002]. After incubating at 30°C, reactions were fractionated by SDS-PAGE and TSC2 phosphorylation was evaluated by immunoblotting using phospho-specific antibodies [Fig. 4B]. Phospho-TSC2 accumulation was both Us3- and ATP-dependent, and was not detected in reactions containing catalytically inactive K220A Us3 [Fig. 4B, lanes 4–6]. While Akt was present in the initial cell-free lysate, it was not detected in the immunopurified kinase reactions, demonstrating the effectiveness of the purification procedure in depleting the endogenous cellular TSC2 kinase [Fig. 4B, cf. lanes 1–3 and 4–6]. To further control for specificity, samples programmed with [γ-32P]-ATP were fractionated by SDS-PAGE to visualize all radiolabeled products. Remarkably, only two major [32P]-incorporated products were observed on overexposed films in reactions that contained wild-type but not K220A Us3. The largest product corresponds in size to TSC2, while the smaller one corresponds to the Us3 autophosphorylation product [Fig. 4C; Sagou et al. 2009]. Thus, Us3 stimulated TSC2 phosphorylation in vitro and the reaction required catalytically active Us3 and ATP.

TSC2 is a substrate phosphorylated directly by Us3

While Us3 stimulated TSC2 phosphorylation in vivo and in vitro, other cellular Us3-responsive or copurifying
kinase contaminants might still be responsible. To evaluate if a direct kinase–substrate interaction accounted for Us3-mediated TSC2 phosphorylation, Us3 was engineered to accept a bio-orthogonal ATP analog not used by the remainder of the kinome (Allen et al. 2007). This was achieved by introducing a Leu→Gly substitution at residue 259, and allows the Us3 active site to accept an unusual bulky ATP analog. While wild-type Us3 can use ATPγS to thiophosphorylate substrates, only analog-sensitive Us3 (AS-Us3) can use N6-alkylated ATPγS (A*TPγS) to thiophosphorylate its substrates. Thiophosphorylated proteins are subsequently reacted with a thiol-specific alkylating agent that generates a bio-orthogonal thiophosphate ester. The resulting labeled proteins are readily detected by immunoblotting using a thiophosphate ester-specific antibody (Allen et al. 2007). Following immunopurification from transfected 293 cells, wild-type and AS-Us3 used ATPγS to thiophosphorylate themselves and TSC2 (Fig. 4D, lanes 1,2,4). Both Us3 and TSC2 thiophosphorylation were not detected in reactions containing K220A catalytically inactive Us3 (Fig. 4D, lane 3). Significantly, whereas AS-Us3 accepted ATPγS and A*TPγS to thiophosphorylate both Us3 and TSC2, wild-type Us3 used ATPγS but not A*TPγS (Fig. 5, lanes 5 vs. 6). This demonstrated that A*TPγS is bio-orthogonal, as it was detectably used by only AS-Us3 and not wild-type Us3. Moreover, it established that Us3 directly phosphorylates TSC2, since AS-Us3 is the only kinase in the reaction capable of accepting and using the A*TPγS analog to thiophosphorylate TSC2.

Inactivation of 4E-BP1 requires TSC2 phosphorylation in HSV-1-infected cells

Stimulation of TSC2 T1462/S939 phosphorylation by Us3 suggested that TSC inactivation was required to activate mTORC1 and in turn stimulate 4E-BP1 phosphorylation in HSV-1-infected cells. To test this, epitope-tagged wild-type TSC2 or a double-mutant derivative with alanine substituted at each of the critical S939 and T1462 phospho-acceptor residues (SATA) were transiently expressed in 293 cells. While both ectopically expressed...
wild-type and SATA TSC2 subunits bind TSC1, SATA-TSC2 was unable to be inactivated by Akt, and behaves as a constitutively activated Rheb-GAP that prevents mTORC1 activation (Manning et al. 2002). HSV-1 infection of 293 cells resulted in rapamycin-sensitive 4E-BP1 hyperphosphorylation followed by proteasome-mediated degradation, as reported in NHDFs (Fig. 5A; Walsh and Mohr 2004). Cells transiently transfected with vector alone or TSC2 expression plasmids (wild type or SATA) were infected with wild-type HSV-1, and, at 12 hpi, total protein was isolated, fractionated by SDS-PAGE, and analyzed by immunoblotting. As expected, HSV-1 infection reduced 4E-BP1 steady-state levels in cells transfected with either vector alone or wild-type TSC2 expression plasmid, while control antigen abundance (actin) remained relatively constant [Fig. 5B, cf. lanes 1 and 2, and lanes 3 and 4]. However, 4E-BP1 levels were not detectably reduced in HSV-1-infected cells expressing TSC2 SATA [Fig. 5B, lanes 5 vs. 6]. Both wild-type and SATA proteins accumulated to similar levels, and comparable viral tk levels confirmed the cultures were infected equivalently [Fig. 5B, lanes 2,4,6]. This established that TSC2 S939/T1462 phosphorylation in HSV-1-infected cells was restricted by host TSC2, which limited 4E-BP1 inactivation and replication in the absence of Us3; and [3] Us3 antagonized the antiviral actions of TSC2. To test this hypothesis, 4E-BP1 phosphorylation and viral replication were evaluated in TSC2-depleted NHDFs. Following treatment with either nonsilencing siRNA or a TSC2 siRNA pool, NHDFs were infected with ΔUs3 or wild-type virus. At 15 hpi, total protein was isolated, fractionated by SDS-PAGE, and analyzed by immunoblotting to verify TSC2 depletion and evaluate 4E-BP1 phosphorylation. 4E-BP1 was largely hypophosphorylated in ΔUs3-infected cells treated with nonsilencing siRNA, as evidenced by the relative abundance of fast-migrating, hypophosphorylated forms compared with the slower-migrating hyperphosphorylated forms [Fig. 6A, lane 1]. In contrast, relatively greater amounts of slower-migrating, hyperphosphorylated 4E-BP1 accumulated in ΔUs3-infected cultures, where TSC2 protein levels were effectively depleted by TSC2 siRNA treatment [Fig. 6A, cf. lanes 3 and 1]. Furthermore, hyperphosphorylated 4E-BP1 levels achieved by silencing TSC2 in ΔUs3-infected cells were similar to those in wild-type virus-infected cells treated with nonsilencing siRNA [Fig. 6A, lanes 2 vs. 3]. This established that endogenous host TSC2 restricted 4E-BP1 inactivation by Us3-deficient HSV-1, and provided genetic evidence that the Us3–TSC2 interaction is required to activate mTORC1.

To determine the impact of TSC2 depletion on replication and spread of Us3-deficient virus, NHDFs treated with either nonsilencing or a TSC2 siRNA pool were infected with either ΔUs3 or wild-type HSV-1. After 4 d, the virus produced was quantified by plaque assay in Vero cells. Both wild-type HSV-1 and encephalomyocarditis (EMCV), an unrelated +strand RNA virus, replicated equivalently in the presence of either nonsilencing or TSC2 siRNA, demonstrating that TSC2 depletion did not generally increase viral replication in mammalian cells [Fig. 6B; Supplemental Fig. S2]. In contrast, TSC2 siRNA selectively enhanced ΔUs3 replication by ~20-fold [Fig. 6B; While substantial, TSC2 depletion did not fully restore ΔUs3 replication to wild-type levels. This was not surprising, since Us3 is multifunctional and, as a Ser/Thr kinase, has numerous viral and cellular substrates (Mou et al. 2007). Without Us3, critical virus functions, including those needed for egress, remain impaired and are unaffected by TSC2 depletion. Nevertheless, these results established that endogenous cellular TSC2 restricted Us3-deficient virus replication in NHDFs. Furthermore, they suggest that Us3 stimulates HSV-1 replication in part by antagonizing TSC2 function, which in turn stimulates mTORC1, promotes 4E-BP1 translational repressor inactivation, and facilitates eIF4F assembly.

Figure 5. Inactivation of the 4E-BP1 translational repressor in HSV-1-infected cells requires site-specific TSC2 phosphorylation. (A) 293 cells were mock-infected (−) or infected (+) with HSV1 [MOI = 10] and, after 3 h, were treated with DMSO or rapamycin. Total protein was collected at 12 hpi, fractionated by SDS-PAGE (17.5% gel), and immunoblotted with anti-4E-BP1 antibody. [B] 293 cells were transiently transfected with either empty vector (−), pcDNA3-Flag-TSC2 (wild type [WT]), or pcDNA3-Flag-TSC2SATA (SATA). After 6 h, cells were serum-starved for 18 h and subsequently mock-infected or infected [MOI = 10] with HSV-1. At 12 hpi, total protein was fractionated by SDS-PAGE (7.5% gel) and analyzed by immunoblotting with the indicated antibodies.

Stimulation of Us3-dependent, Akt-substrate phosphorylation in HSV-1-infected cells

Having established Us3 phosphorylated TSC2 on identical residues targeted by the cellular kinase Akt, the possibility that other Akt substrates were phosphorylated...
in a Us3-dependent manner in infected cells was investigated. An Akt-phosphosubstrate (Akt-pSub) antibody used to identify TSC2 as an Akt substrate and that recognizes the RxRxxS/T Akt substrate motif in a growth factor- and PI3K-dependent manner was employed [Manning et al. 2002]. Akt-pSub accumulation was stimulated in HSV-1-infected NHDFs in the presence and absence of Akt inhibitor VIII [Fig. 7A]. Significantly, while similar profiles of Akt-pSub immunoreactive proteins were detected in ΔUs3-infected cells, their accumulation was Akt inhibitor VIII-sensitive, consistent with their being Akt substrates. This suggested that Akt-pSub accumulation in HSV-1-infected cells was Us3-dependent. Indeed, phosphorylation of Akt substrates GSK3 and FOXO1 on Akt target sites was stimulated in HSV-infected cells and fully dependent on Us3 in the presence of the Akt inhibitor [Fig. 7A]. However, unlike Akt, which requires PI3K-dependent activation, GSK3 and 4E-BP1 phosphorylation in HSV-1-infected cells was insensitive to the PI3K inhibitor wortmannin (Supplemental Fig. S3). Finally, wild-type but not K220A Us3 effectively stimulated FOXO1 and GSK3 phosphorylation in uninfected, Akt inhibitor-treated cells (Fig. 7B, lanes 4–6), demonstrating that Us3 was necessary and sufficient to promote phosphorylation of these well-characterized Akt substrates in uninfected cells. Despite stimulating Akt substrate phosphorylation, α herpesvirus Us3-like Ser/Thr kinases occupy a separate branch distinct and equidistant from other cellular kinases. In particular, the Akt/ PKA subfamily is not more closely related to Us3 than any other subfamily [Fig. 7C]. Overall 28% identity between Akt1 and Us3, along with a 3e-15 blast E-value, suggests that, while both are functional Ser/Thr protein kinases, they are at best very distantly related at the phylogenetic level.

Discussion

By preventing eIF4G from associating with the cap-binding protein eIF4E and restricting assembly of a multisubunit initiation factor complex, the translation repressor 4E-BP1 regulates 40S ribosome subunit recruitment to mRNA. Moreover, having 4E-BP repressor activity responsive to mTORC1 allows translational output to respond to changes in intracellular and environmental status. For example, viruses whose mRNAs contain a 5’ m7GTP cap must ensure that translational repression by 4E-BP1 does not limit viral protein production [Mohr et al. 2007]. Here, we establish that the HSV-1 Us3 Ser/Thr kinase is required to inactivate 4E-BP1 in infected cells. Acting analogously to the cellular kinase Akt, Us3 stimulated accumulation of phosphorylated Akt substrates, including FOXO1, GSK3, and TSC2. Direct, site-specific TSC2 phosphorylation by Us3 turned on mTORC1, inactivating the translational repressor 4E-BP1 in infected cells. Finally, TSC2 depletion increased Us3-deficient virus replication in normal human fibroblasts, illustrating that TSC restricts replication of a virus unable to activate mTORC1 and that Us3 is required to antagonize TSC antiviral activity. This demonstrates that the HSV-1 Us3 Ser/Thr protein kinase can remodel host translational initiation factor complexes, and establishes a new strategy whereby protein kinases bearing little or no primary sequence homology with Akt are used to regulate mTORC1.

By encoding a Ser/Thr protein kinase like Us3, HSV-1 capitalizes on protein phosphorylation as a potent cell signaling mechanism. Moreover, by imbuing this kinase with a broad yet specific substrate specificity, this single gene product is able to subvert signaling within multiple, fundamental pathways to favor viral productive replication without occupying substantial space in the viral genome [Munger and Roizman 2001; Ogg et al. 2004; Poon et al. 2006; Leach et al. 2007; Mou et al. 2007]. While not essential for replication, Us3-deficient viruses exhibit cell type-specific replication defects in culture and are severely impaired in mouse pathogenesis models [Meignier et al. 1988, Nishiyama et al. 1992, Ryckman and Roller 2004]. In addition, Us3 bears little resemblance to any specific cellular Ser/Thr kinase obstructing target substrate prediction. Despite phosphorylating some peptide
substrates shared with PKA and PKC in vitro, Us3 exhibited distinct synthetic peptide substrate specificity (Leader et al. 1991). One study suggested Us3 specificity overlapped with PKA, whereas another found this too limiting (Benetti and Roizman 2004; Mou et al. 2007). Nevertheless, recognition motifs are unlikely to provide the specificity required in vivo, since (1) many kinases share in vitro recognition motifs (i.e., Arg residues N-terminal to phospho-acceptor site for p90 RSK, PKA, and Akt), (2) recognition motifs may not be physiologically phosphorylated, (3) peptides may not mimic intact protein phosphorylation kinetics, and (4) not all kinases have a clear consensus motif in peptide substrates (Biondi and Nebreda 2003). Instead, biological specificity is often achieved by interactions outside the active site that dock the kinase to physiological substrates (Pawson and Nash 2000; Remenyi et al. 2006). To further complicate matters, Us3 also targets viral substrates and is itself phosphorylated by a second HSV-1 protein kinase, UL13 (Kato et al. 2006). While UL13 might alter Us3 specificity, Us3 alone is sufficient to promote TSC2, GSK3, and FOXO1 phosphorylation. Us3 not only facilitates nuclear lamina disassembly and egress of newly assembled progeny virions, it has anti-apoptotic activity and stimulated phosphorylation of the Akt substrate BAD in vitro (Munger and Roizman 2001; Kato et al. 2005). Our findings that Us3 stimulates site-specific, phosphorylated Akt substrate accumulation—including FOXO1, GSK3, and TSC2—now raise the possibility that Us3 may in fact be a mimic for this critical cellular regulator. Importantly, although Us3 targets Akt substrates and constitutively activates Akt signaling, it is unrelated to Akt at the primary sequence level and therefore is unresponsive to endogenous regulatory pathways that limit Akt activity.

By restricting mTORC1 activation, TSC controls an Akt-regulated antiviral checkpoint limiting access to the...
host cap-dependent translation machinery. Indeed, co-opting control of Akt signaling by inactivating TSC2 figures prominently in the biology of different representative α, β, and γ herpesvirus subfamily members. However, distinct mechanisms of targeting the TSC2 subunit by Akt. Notably, only the herpesvirus HCMV encodes a TSC2-binding protein [UL38] required for mTORC1 activation (Moorman et al. 2008), whereas the γ herpesvirus KSHV stimulates TSC2 phosphorylation via a virus-encoded GPCR (Sodhi et al. 2006). We now define Us3 as the first viral Ser/Thr kinase capable of directly phosphorylating TSC2 on sites normally targeted by Akt. Notably, only α herpesvirus subfamily members (HSV-1, PRV, and VZV) encode a Us3-related Ser/Thr kinase protein. Conceivably, the diverse mechanisms by which TSC is inactivated by different herpesviruses could reflect the specialized relationship each virus has with the differentiated host cells it colonizes. For example, the Ser/Thr kinase is encoded uniquely by herpesviruses (which colonize neurons), whereas an inhibitory TSC2-binding protein is specified by HCMV (which colonizes monocyte/macrophage precursors), and a GPCR is encoded by KSHV [which targets B cells and endothelial cells].

Regardless of their TSC inactivation mechanism, the fact that representative herpesvirus family members all target TSC signaling to manipulate mTORC1 is striking, highlighting the significance of the TSC regulatory intersection in regulating mTORC1 and implying that it is an intrinsic host defense component whose actions must be antagonized by a viral function. Other unrelated DNA viruses that produce caged, polyadenylated mRNAs also target TSC2, as the HPV E6 protein promotes TSC2 degradation [Lu et al. 2004]. Furthermore, physiological dampening of cap-dependent translation to stresses—including hypoxia and energy depletion via AMPK and REDD1, respectively—all proceed through TSC (for review, see Ma and Blenis 2009). Importantly, having a virus-encoded function activate mTORC1 through TSC overrides p70 S6K-mediated cellular feedback controls in place to limit both receptor-mediated activation of the PI3K/Akt/mTORC1 signaling axis (Harrington et al. 2005) and mTORC2-mediated Akt activation (Dibble et al. 2009). Thus, activating mTORC1 via a PI3K-coupled cellular receptor would ultimately only be transient. Indeed, transient Akt activation early in the replication cycle is observed in primary human fibroblasts infected with wild-type HSV-1 or HCMV (Benetti and Roizman 2005); PNBM was from Epitomics (3700-1); and N 6-phenylethyl-ATP (ATCC) cells were propagated in 5% calf serum. F-strain wild-type and ΔUs3 repair (VRR1202repair) HSV1 were described (Ryckman and Roller 2004). UL13-deficient virus [dl 13 lac z] was a gift from S. Rice. Antibodies Akt (#9272), Akt-pSer473 (#9271), Akt-pThr 308 (#9275), TSC2 (#3611), TSC2-pSer939 (#3615), TSC2-pThr1462 (#3611), AktSub (#9611), p70 S6K T389 and total (#9430), FOXO1 (#2488), FOXO1-pSer256 (#9461), and GSK3a/β-pSer9/21 (#9331) were from Cell Signaling; anti-GSK3a/β (#05-903) was from Millipore; anti-thiophosphate ester was from Epitomics [2686-1]; anti-HSV-1 tk was a gift from B. Roizman; and anti-FlagM2 Monoclonal and ATPγS (#AI388) were from Sigma. All other antibodies were described [Walsh and Mohr 2004]. Rapamycin, wortmannin, puromycin, and Akt inhibitor VIII were from Calbiochem; PP242 was described (Feldman et al. 2009); NPNM was from Epitomics [3700-1]; and N'-Phenylethyl-ATPγS was from BioLog (P026).

Western blotting and mGTP-Sepharose chromatography

These procedures were performed as described (Walsh and Mohr 2004; Walsh et al. 2005).

siRNA and plasmids

siGENOME SMARTpool targeting human TSC2 and siGENOME Non-Targeting siRNA #2 were from ThermoScientific. pcDNA3-FlagTSC2 (14129), pcDNA3-FlagTSC2SATA (14131), and pRK7-FlagTSC2 [8996] were from AddGene. pcDNA3-Us3-Flag was described previously (Ogg et al. 2004); pcDNA3-Us3-Flag K220A and L256G were constructed by standard techniques and verified by sequencing (R. Roller and U. Chuluunbaatar, unpubl.). Flag-PAK3-4E-BP1 wild-type and AA mutant cloned into the retroviral vector pBABE-puro were from R. Schneider.
Cell transfection and infection

NHDFs were seeded at $3 \times 10^4$ cells per well in six-well dishes. The next day, cells were washed with PBS, serum-deprived for 72 h as described (Walsh and Mohr 2004), and then infected with HSV-1 (multiplicity of infection [MOI] = 5). HEK293 cells were seeded [4 \times 10^5 cells per well in a 12-well dish], and the following day were transfected using Lipofectamine 2000 [Invitrogen] at 1:2.5 [w/v] ratio for 6 h, serum-deprived for 18 h, and infected with HSV-1 (MOI = 10). U2OS cells were seeded [1.5 \times 10^5 cells per well in a 12-well dish], and the next day were transfected as described above. After 6 h, fresh media was added. At 24 h post-transfection, cells were treated with 200 nM wortmannin, Akt inhibitor VIII, or DMSO for 30 min before collection. For multicyle growth experiments, cultures were infected with 100 plaque-forming units [pfu] per well. After 4 d, cell-free lysates were prepared by freeze–thawing and the virus produced was quantified by plaque assay in Vero cells.

siRNA depletion

NHDFs were seeded at 2.7 \times 10^4 cells per well in 12-well dishes, and the next day were transfected with siRNA using DharmaFECT Duo (ThermoScientific) according to the manufacturer. After 24 h, the transfection procedure was repeated. Forty-eight hours later, cultures were serum-deprived as described (Walsh and Mohr 2004) and then infected with HSV-1 or EMCV.

Immunoprecipitation and in vitro kinase assay

HEK293 cells [60-mm dishes] were transfected with pRK7-FlagTSC2, pcDNA3-Us3Flag, or pcDNA3-K220AUs3Flag. DNA quantity was adjusted to achieve comparable Us3 and K220A expression. Twenty-four hours post-transfection, FlagTSC2-expressing cells were treated with wortmannin [200 nM for 30 min]. Cells were lysed in 50 mM Tris-HCl [pH 7.5], 150 mM NaCl, 1% NP-40, 0.5% deoxycholate, and 0.1% SDS plus Complete Mini protease inhibitor [Roche], and extracts were clarified by centrifugation [12,000 g, for 10 min at 4°C]. Soluble fractions from Flag TSC2 transfected cells were mixed 1:1 with an equivalent fraction from Us3Flag or K220AUs3Flag transfected cells. Protein G-Sepharose conjugated to anti-Flag antibody [10 \mu L of settled bed volume] was added and the mixture was rocked [2 h at 4°C]. Immune complexes were collected by brief centrifugation, resuspended in buffer (10 mM Tris-HCl at pH 8.0, 1 M NaCl, 0.2% NP-40), rocked gently [15 min at 4°C], and then washed three times with 0.5 mL of Us3 kinase buffer supplemented with 10 mM ATP, 0.5 mM DTT, and 10 \mu Ci [gamma-32P]ATP for 30 min at 30°C and stopped by adding equal volume of 2× sample buffer [125 mM Tris-HCl at pH 6.8, 4% SDS, 20% glycerol, 10% beta-mercaptoethanol]. Reaction mixtures were fractionated by SDS-PAGE and processed for immunoblotting or autoradiography. Two-hundred-fifty micromolar ATPyS or NAD-Phenylethyl ATPyS was included in place of ATP where indicated. ATP analog-containing reactions were incubated for 1 h at 30°C, terminated by adding EDTA to 40 mM, and alkylated with 2.5 mM  p-nitrobenzyl mesylate (PNBM) for 2 h at room temperature. After SDS-PAGE and immunoblotting, alkylated proteins were detected using anti-thiophosphate ester antibody.

Retroviral expression

HEK293 Gag-Pol packaging cells [80% confluent on 10-cm dishes] were transfected with 8 \mu g each of pcMV-VSV-G and pBABE-puro vectors using Lipofectamine 2000 [Invitrogen]. Retrovirus-containing supernatants were harvested after 48 h, filtered [0.45 \mu m], and frozen at −80°C until use. Early passage NHDFs were infected with retrovirus-containing media plus polybrene [10 \mu g/mL] for 6 h. Puromycin [2 \mu g/mL] was added 48 hpi and the population was maintained in the presence of drug. Puromycin was removed from cultures before infecting with HSV-1.

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