1	Kaposi's sarcoma-associated herpesvirus fine-tunes the temporal expression of		
2	late genes by manipulating a host RNA quality control pathway		
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9	Running title: PPD dampens KSHV late gene expression		
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24 Abstract

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26 Kaposi's sarcoma-associated herpesvirus (KSHV) is a human oncogenic nuclear DNA 27 virus that expresses its genes using the host cell transcription and RNA processing 28 machinery. As a result, KSHV transcripts are subject to degradation by at least two host-29 mediated nuclear RNA decay pathways, PABPN1 and PAPa/y-mediated RNA decay (PPD) and an ARS2-dependent decay pathway. Here, we present global analyses of viral 30 31 transcript levels to further understand the roles of these decay pathways in KSHV gene 32 expression. Consistent with our recent report that the KSHV ORF57 protein increases 33 viral transcript stability by impeding ARS2-dependent decay, ARS2 knockdown has little effect on viral gene expression 24 hours after lytic reactivation of wild-type virus. In 34 35 contrast, inactivation of PPD results in premature accumulation of late transcripts. The up-regulation of late transcripts does not require the primary late gene-specific viral 36 37 transactivation factor, suggesting that cryptic transcription produces the transcripts that then succumb to PPD. Remarkably, PPD inactivation has no effect on late transcripts at 38 39 their proper time of expression. We show that this time-dependent PPD evasion by late 40 transcripts requires the host factor NRDE2, which has previously been reported to protect 41 cellular RNAs by sequestering decay factors. From these studies, we conclude that KSHV 42 uses PPD to fine-tune the temporal expression of its genes by preventing their premature 43 accumulation.

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47 Importance

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49 Kaposi's sarcoma-associated herpesvirus (KSHV) is an oncogenic gammaherpesvirus that causes Kaposi's sarcoma and other lymphoproliferative disorders. Nuclear 50 51 expression of KSHV genes results in exposure to at least two host-mediated nuclear RNA 52 decay pathways, PABPN1 and PAPa/y-mediated RNA decay (PPD) and an ARS2-53 mediated decay pathway. Perhaps unsurprisingly, we previously found that KSHV uses 54 specific mechanisms to protect its transcripts from ARS2-mediated decay. In contrast, 55 here we show that PPD is required to dampen the expression of viral late transcripts that 56 are prematurely transcribed, presumably due to cryptic transcription early in infection. At 57 the proper time for their expression, KSHV late transcripts evade PPD through the activity of the host factor NRDE2. We conclude that KSHV fine-tunes the temporal expression of 58 59 its genes by modulating PPD activity. Thus, the virus both protects from and exploits the 60 host nuclear RNA decay machinery for proper expression of its genes. 61 62 63 64 65 66

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70 Introduction

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72 Kaposi's sarcoma-associated herpesvirus (KSHV; also known as human herpesvirus 8; 73 HHV-8), is an enveloped, double-stranded DNA virus. It is the etiological agent of 74 Kaposi's sarcoma and of the lymphoproliferative disorders primary effusion lymphoma 75 (PEL) and multicentric Castleman's disease (MCD) (1-4). Like other herpesvirus, the KSHV life cycle consists of a latent phase and a lytic phase. During latency, the viral 76 genome resides in the host nucleus as a non-integrated, circular episome, and no virions 77 78 are produced. Upon reactivation, the virus undergoes a well-regulated cascade of gene 79 expression initiated by the viral transactivator ORF50 (Rta) that ultimately results in the production of infectious virus (5-7). KSHV transcription and genome replication occur in 80 81 the nucleus where the virus takes control of the host machinery needed for these processes. Consequently, similar to host RNAs, viral transcripts are subject to host-82 83 mediated RNA guality control (QC) pathways (8-12).

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85 RNA QC pathways play an essential role during RNA biogenesis (8-12). In addition to 86 eliminating misprocessed transcripts, RNA QC pathways prevent the accumulation of 87 unstable, non-coding RNAs such as promoter-upstream transcripts (PROMPTs, also 88 called uaRNAs) (13-15). PROMPTs are polyadenylated, non-coding RNAs with no or few 89 introns that are transcribed from bidirectional promoters antisense to protein-coding 90 genes (16-18). Accumulation of PROMPTs has deleterious effects for the cells as they 91 compete with coding transcripts for the translational machinery (15). In eukaryotes, at 92 least two nuclear RNA decay pathways prevent the accumulation of PROMPTs (13-15,

19, 20). Primarily, they are degraded through the CBCN complex that is recruited to the
RNA via its 5' cap. CBCN consists of the cap-binding complex (CBC), the ARS2 protein,
and the nuclear exosome targeting (NEXT) complex (13, 21). The NEXT complex subunit
MTR4 recruits the RNA exosome to degrade the transcript (13, 22).

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98 PROMPTs and other long polyadenylated nuclear RNAs are also degraded by the PABPN1 and PAPα/y-mediated RNA decay (PPD) pathway in which decay factors are 99 recruited through 3' poly(A) tail (14, 15, 19, 20, 23, 24). In this pathway, the nuclear poly 100 101 (A)-binding protein (PABPN1) promotes poly(A) tail extension of target transcripts by 102 stimulating the function of the poly (A) polymerases (PAP α or PAPy; abbreviated $PAP\alpha/\gamma$). The targeted RNAs are subsequently degraded by the nuclear RNA exosome. 103 104 Recruitment of the exosome to polyadenylated RNAs is mediated by the zinc finger protein ZFC3H1, which links the exosome cofactor MTR4 to PABPN1. This link was 105 106 coined the poly(A) tail exosome targeting (PAXT) connection (also called the polysome 107 protector complex, PPC) (15, 20). Overall, both PPD and the CBCN complex survey the 108 integrity of RNAs to eliminate transcriptional noise, misprocessed and other potentially 109 detrimental RNAs.

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ARS2 directly interacts with the CBC to form a hub that allows the assembly of mutually exclusive complexes that dictate the fate of a transcript (13, 21, 24, 25). For instance, ARS2 interacts with PHAX or ALYREF to promote the nuclear export of properly processed snRNAs and mRNAs, respectively (21, 24-26). Alternatively, ARS2 recruits NEXT or PAXT to target RNAs for exosome-mediated degradation (13, 20, 27). In some

cases, ARS2 targets transcripts for degradation independently of NEXT or PPD/PAXT
(28, 29). These complex interconnections challenge efforts to uncover the degree of
independence or redundancy between nuclear RNA decay pathways. Nonetheless,
ARS2 clearly plays a central role in promoting the decay of a number of nuclear
transcripts.

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122 Like their host counterparts, KSHV mRNAs are capped and polyadenylated, but most 123 KSHV genes are short and intronless (30, 31). Consequently, cellular RNA QC pathways 124 may degrade KSHV RNAs due to their similarity to PROMPTs. The essential 125 multifunctional KSHV ORF57 protein promotes viral transcript accumulation by increasing nuclear RNA stability (32-45). We recently reported that viral transcripts are subject to 126 127 degradation by both PPD and an ARS2-dependent but NEXT-independent decay pathway upon lytic reactivation of virus lacking ORF57 (29). Using pulse-chase assays 128 129 with an unstable form of the KSHV nuclear non-coding PAN RNA (PAN Δ ENE), we further 130 showed that ORF57 preferentially protects viral transcripts from the ARS2-dependent 131 decay pathway (29). Interestingly, although viral transcripts succumb to PPD, ORF57 132 protection of PANΔENE from PPD was modest, suggesting the possibility that a subset 133 of viral transcripts undergo PPD-dependent degradation in the presence of ORF57 (29). 134 However, the role of PPD during viral infection with ORF57 expressing virus has not been 135 fully explored.

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Here, we used RNAi to inactivate PPD and/or ARS2-dependent decay and monitored
their contributions to KSHV gene expression in the presence of ORF57 by RNA-seq at

24 hours after lytic reactivation. ARS2 depletion resulted in few changes in viral gene 139 expression. However, PPD inactivation resulted in increased expression levels of several 140 141 viral genes. Interestingly, the most upregulated transcripts were late transcripts that are 142 otherwise expressed at ~48 hours after lytic reactivation. Our data suggest that PPD prevents the premature accumulation of late transcripts which presumably arise as a 143 144 consequence of cryptic transcription. Notably, at their proper time of expression, PPD 145 inactivation has no effect on viral late transcripts, and the host factor NRDE2 is needed for evasion of PPD. We conclude that KSHV exploits PPD to fine-tune the temporal 146 expression of viral genes by dampening steady-state levels of prematurely transcribed 147 late genes. 148

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- 150 Results
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152 PPD inactivation results in aberrant temporal expression of KSHV late genes

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154 Our previous studies showed that viral RNAs were subject to both PPD and ARS2-155 mediated decay in the absence of ORF57 (29). ORF57 more potently protected viral 156 transcripts from ARS2-mediated decay than from PPD, suggesting that viral transcripts 157 succumb to PPD even in the presence of ORF57. To further characterize the role of 158 these decay pathways during KSHV infection, we performed an RNA-seg experiment to 159 monitor the levels of viral transcripts after siRNA depletion of ARS2, the PPD component 160 PAPa/y, or both simultaneously (dKD) in iSLK cells latently infected with the KSHV 161 infectious clone BAC16 (iSLK WT) (46, 47)(Fig 1A). Efficiency of knockdown was

validated by western blot, qRT-PCR and/or loss of function assays (Fig 1B). Lytic 162 163 reactivation was induced using doxycycline (dox) to promote expression of the dox-164 inducible RTA integrated into the iSLK host cell chromosomes and by the histone 165 deacetylase inhibitor sodium butyrate (NaB). We prepared libraries from RNA harvested 166 24 hours post induction (hpi), and the samples were subjected to high-throughput 167 sequencing (Fig 1A). Expression of several KSHV genes significantly increased in 168 samples depleted of PAP α /y and in the dKD compared to samples treated with a control siRNA. However, we observed minimal alterations in gene expression in the ARS2 169 170 knockdown samples, consistent with the idea that ARS2-mediated decay is inhibited by 171 ORF57 (Fig 1C and Table S1). Surprisingly, the most upregulated genes (> 4-fold 172 change) upon PAP α /y depletion and in the dKD were late genes, 76% and 70% 173 respectively (Fig 1D). Typically, these KSHV late genes are not expressed at ~24 hpi in 174 these cells. These data suggest that KSHV exploits $PAP\alpha/y$ to temporally control the 175 expression of late genes.

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177 **PPD/PAXT targets KSHV late genes for degradation**

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Our RNA-seq data show that $PAP\alpha/\gamma$ depletion results in increased expression of late genes at 24 hpi and suggest that PPD controls the expression late genes that prematurely arise due to cryptic transcription. To validate these observations and extend the findings to additional PPD factors, we focused our attention on three KSHV late genes, ORF52, ORF75, and K8.1. These genes were selected because they have different structural features. ORF52 is a short transcript (395 base pairs (bp)) while ORF75 is a long

transcript (3890 bp). Both ORF52 and ORF75 are intronless, but K8.1 contains one intron. 185 186 In spite of these structural differences, the expression of each gene increased upon 187 PAPα/y depletion at 24 hpi (Fig 2B, D and F). None were affected by ARS2 depletion 188 alone. Similar results were obtained when the mRNA levels of these genes were 189 monitored by qRT-PCR (Fig 2C, E and G). In principle, PAP α /y knockdown may affect 190 gene expression independent of PPD due to its function in 3'-end formation. To confirm the role of PPD, we tested whether depletion of PPD factors other than PAP α/γ caused a 191 192 similar phenotype. We depleted cells of the unique PPD/PAXT factor ZFC3H1 or the 193 exosome co-factor MTR4. Efficiency of knockdown was monitored by western blot, gRT-194 PCR and loss of function assays (Fig 2A). Consistent with PPD inactivation, depletion of ZFC3H1 or MTR4 also increased ORF75, ORF52 and K8.1 levels (Fig 2C, E and G). 195 These data support the conclusion that PPD/PAXT suppresses KSHV late gene 196 197 expression during the early stages of the virus lytic phase.

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PPD regulates virus late gene expression independently of the viral transactivation
 factor ORF24

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To elucidate the mechanism of late gene expression upregulation in the context of PPD inactivation, we first focused our attention on the requirements for late gene expression. In KSHV, transcription of late genes takes place after initiation of viral DNA replication, and their expression requires the action of several viral transactivation factors (vTF) (48-53). The viral TATA box-binding protein homolog, ORF24, is a KSHV transactivation factor that plays a critical role when the virus progresses from DNA replication to

expression of late genes. ORF24 binds to late gene promoters and recruits RNA 208 209 polymerase II (pol II) and other vTFs to the promoter region of late genes to induce their 210 expression. Because ORF24 mRNA levels were increased in cells depleted of PAP α/γ 211 and in the dKD at 24 hours post lytic reactivation (Fig 3A and 3B), it is possible that PPD 212 inactivation results in ORF24 upregulation which drives the increased expression of late 213 genes. In this case, the effects of PPD on other late transcripts would result from a 214 secondary effect of ORF24 upregulation. To determine whether the increased expression 215 of late genes at 24 hpi is due to an upregulation of ORF24 upon PPD inactivation, we 216 used iSLK cells transfected with a bacmid encoding the viral genome in which ORF24 contains a point mutation, R328A, that renders it inactive (51). ORF24^{R328A} maintains the 217 218 interaction with pol II but is unable to interact with other vTAs resulting in strongly impaired 219 expression of late genes (Fig 3C). If the up-regulation of late genes by PPD inactivation 220 is due to secondary effects of ORF24 upregulation, then the up-regulation will be 221 abrogated in the mutant virus. In contrast to this prediction, depletion of PPD components 222 PAPα/y, MTR4 or ZFC3H1 resulted in increased mRNA levels of ORF52, ORF75 and K8.1 in ORF24^{R328A} reactivated cells (Fig 3D, E and F). Thus, the role of PPD in late gene 223 224 expression at 24 hpi is independent of ORF24 transactivation. Moreover, these data are 225 consistent with a proposed role for PPD in the posttranscriptional inhibition of premature 226 late transcript accumulation.

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PPD control of late transcripts is restricted to early phases of reactivation but does
 not affect genome replication or virus production in iSLK cells

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KSHV late genes are expressed after the initiation of viral genome replication (48-52), 231 232 which occurs prior to 48 hpi in our iSLK WT cells. As PPD inactivation increases late gene 233 expression at 24 hpi, we tested whether the expression of late genes at 48 and 72 hpi is 234 also affected by PPD inactivation. Interestingly, at 48- or 72-hpi, PAP α /y depletion had 235 no effect on the expression levels of the late genes tested (Fig 4A, B and C). These data 236 suggest that at the proper time of expression, KSHV late transcripts are able to avoid 237 PPD-mediated degradation. They further support the model that PPD dampens 238 premature expression of KSHV late genes.

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240 Because of the aberrant timing of viral late gene expression, we investigated whether 241 PPD inactivation affected virus fitness. To test genome replication, iSLK WT cells were 242 depleted of PAP α/γ or ZFC3H1, and ORF57 DNA levels were monitored over time by gRT-PCR. Depletion of PAPa/y or ZFC3H1 had no significant effect on the levels of viral 243 244 genome replication at 48 hpi (Fig 4D). We next investigated whether PPD inactivation 245 affects production of infectious virions. To test this, we collected media from iSLK WT 246 cells at 0, 8, 12, 24, 48 and 72 hours after lytic reactivation and used it to infect HEK293 247 cells. Two days later, viral infection was analyzed by flow cytometry to detect the GFP 248 expressed by BAC16. Most of the HEK293 cells infected with media collected from iSLK 249 WT cells at 48 and 72 hpi were GFP positive (Fig 4E). Importantly, depletion of PAP α/γ 250 or ZFC3H1 had no effect in the production of infectious virus as the percentage of GFP 251 positive cells was similar to that of cells treated with a control siRNA (Fig 4E). These data 252 suggest that the premature expression of late genes observed upon PPD inactivation 253 does not dramatically perturb the virus life cycle in cultured cells.

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NRDE2 is needed for proper expression of late genes at 48 hours post lytic reactivation

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258 To determine how late transcripts avoid degradation at their proper time of expression, 259 we centered our attention on the human nuclear RNAi-defective 2 (NRDE2) protein. 260 NRDE2 localizes to nuclear speckles where it forms a 1:1 complex with MTR4 to inhibit 261 its recruitment and RNA degradation (54). Given this protective activity, we hypothesized 262 that KSHV uses NRDE2 to protect late transcripts from degradation. Therefore, we 263 depleted cells of PAP α /y, NRDE2 (Fig 5A) or both simultaneously and measured 264 expression levels of late transcripts by gRT-PCR at 24 hpi (Fig 5B) or 48 hpi (Fig 5C). As 265 expected. PAP α /y depletion resulted in increased expression levels of late genes at 24 266 hpi but no effect at 48 hpi (Fig 5B and C) (green bars). In contrast, NRDE2 depletion 267 caused a reduction in expression levels of all late transcripts at 48 hpi but had no effect 268 at 24 hpi (Fig 5B and C) (orange bars) suggesting that NRDE2 protects KSHV late transcripts from PPD-mediated degradation at 48 hpi. Importantly, co-depletion of PAP α/ν 269 270 and NRDE2 restored late transcripts levels to that of control siRNA treated cells (Fig 5C) 271 (blue bars). We conclude that after viral genome replication, the host NRDE2 protects 272 KSHV RNAs from PPD.

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277 Discussion

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279 KSHV transcripts are subject to degradation by at least two host-mediated nuclear RNA 280 decay pathways, PPD and an ARS2-dependent decay pathway (29). KSHV ORF57 281 increases viral transcript stability by protecting RNAs from ARS2-dependent decay (29). 282 Our work here suggests that KSHV uses PPD to post-transcriptionally control the 283 premature accumulation of late transcripts during the early stages of the viral lytic phase. 284 In the context of PPD inactivation, late transcripts aberrantly accumulate at 24 hpi (Fig 1 285 and 2), but the premature production of late transcripts does not require functional 286 ORF24. Therefore, the transcripts do not accumulate as a secondary consequence of the 287 up-regulation of the late gene inducer ORF24 (Fig 3). Presumably, the open chromatin 288 and high transcription of the viral genome during early lytic phase allows low-level cryptic 289 transcription of late genes (Fig 6A). The transcripts are eliminated by PPD, so no proteins 290 are produced. At their proper time of expression, late transcripts evade PPD by an 291 NRDE2-dependent mechanism (Fig 5). Thus, we propose that KSHV fine-tunes temporal 292 expression of its genes using a specific host RNA quality control pathway.

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Increasing evidence shows that RNA decay pathways play a critical role in controlling viral infection (29, 55-57). The nuclear RNA decay factors, MTR4 and ZCCHC7, translocate to the cytoplasm where they promote exosome-mediated degradation of viral transcripts of multiple RNA virus (55). In the case of KSHV, the ORF57 protein protects viral transcript from ARS2-dependent decay by preventing MTR4 recruitment (29). In these examples, the viruses must circumvent the RNA QC machinery to properly express

its genes. Here we propose that the virus hijacks PPD activity to fine-tune the temporal 300 301 expression of its genes. That is, KSHV allows PPD to degrade late transcripts that arise as a consequence of cryptic transcription at 24 hpi (Fig 2 and 6A). However, at the 302 303 appropriate time of expression, KSHV blocks PPD so late genes are expressed (Fig 4A-304 C). The long co-evolution of herpesviruses with their specific hosts has selected for 305 sophisticated host-pathogen interactions. These contrasting interactions with the host 306 RNA QC pathways represent intriguing examples of virus-host co-evolution that ensures 307 KSHV expresses its genes in a precise temporal manner.

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309 Our observations also contribute to the understanding of the distinctions between host 310 PPD, PAXT and ARS2-mediated decay processes. While it seems likely that PPD and 311 PAXT represent the same pathway, it has been reported that ARS2 is involved in PAXT 312 recruitment to unstable transcripts (20). However, in the context of KSHV infection, PPD 313 and an ARS2-mediated decay pathway appear to be independent pathways. For 314 example, simultaneous depletion of PAP α /y and ARS2 resulted in greater stabilization of 315 viral transcripts than depletion of either alone during iSLK- Δ ORF57 reactivation (29). 316 Furthermore, we show that depletion of $PAP\alpha/\gamma$ enhances specific viral genes, while ARS2 depletion has little effect (Figs 1 and 2). Importantly, we show that depletion of 317 318 ZFC3H1, a PAXT component, mimics PAP α/γ depletion as the same group of viral genes 319 are upregulated (Fig 2). Overall, these data suggest that PPD and PAXT represent the 320 same process, but ARS2 is not absolutely required for PPD/PAXT-mediated decay. 321 However, more work is needed to completely define the overlap and independence of 322 these RNA QC pathways.

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PPD inactivation results in the aberrant temporal expression of KSHV late genes. 324 325 However, this atypical expression of late genes does not affect viral genome replication 326 or production of infectious virions in our iSLK cells (Fig 4D & E). Several reasons may 327 explain this unexpected result. Even though PAP knockdown increases late gene 328 expression levels relative to cells treated with a control siRNA, the expression levels 329 reached may not be high enough to affect viral fitness. Indeed, expression of late genes 330 at 48 or 72 hpi is considerably higher than at 24 hpi (Fig 4A-C). Consequently, the levels 331 of late genes reached at 24 hpi may not be sufficient to disrupt viral physiology in iSLK 332 cells. Another possibility is that the virus uses PPD to keep expression levels of late genes 333 low during early phases of lytic infection as these may elicit an immune response in an 334 infected organism. Indeed, circulating anti-K8.1 antibodies detected in Kaposi's sarcoma 335 patients support that this PPD-restricted gene elicits an immune response (58-60). If the 336 role of PPD is to keep potentially immunogenic genes low, we would miss this phenotype 337 in a cell culture system.

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339 Late transcripts evade PPD during late phases of replication, but the mechanism of PPD 340 evasion is not completely understood. Our data show that the host factor NRDE2 is 341 required for evasion of PPD during late infection (Fig 5). NRDE2 function is linked to its 342 residence in nuclear speckles, where it interacts with MTR4 to inhibit its activity thereby 343 protecting speckle-associated mRNAs (54). As expression of KSHV late genes occurs at 344 the onset of viral genome replication, we speculate that NRDE2 re-localizes to replication 345 compartments (Fig 6B). When this occurs, NRDE2 interacts with MTR4 preventing the 346 recruitment of the nuclear exosome to viral transcripts. Consequently, late transcripts

evade PPD and are properly expressed. Supporting this idea, HSV1 replication
compartments coalesce with nuclear speckles (61), but whether this occurs in KSHV has
yet to be tested.

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351 Additional factors may contribute to protection of late transcripts from PPD during late 352 phases of KSHV reactivation. For example, expression of KSHV late genes requires the 353 action of several viral transactivation factors (48-52). ORF24 is essential to recruit RNA pol II and other viral transactivation factors to late gene promoters (51). In principle, 354 355 ORF24-induced transcription may promote the co-transcriptional recruitment of factors 356 that protect transcripts from degradation. Evasion and exploitation of nuclear RNA decay 357 pathways by KSHV is only beginning to be understood. Further experimentation is 358 needed to substantiate the role of NRDE2 and identify other factors involved in these 359 processes.

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362 Materials and Methods

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364 **RNA-seq: library preparation**

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iSLK WT cells were transfected with a non-targeting control siRNA or a two-siRNA pool targeting PAP α and PAP γ (PAP α/γ), ARS2 or both PAP α/γ and ARS2 combined (dKD) using the concentrations specified in the siRNA transfection section. Total RNA was harvested three days after siRNA transfection and 24 hours post lytic reactivation. One

µg of intact total RNA per condition was used to make stranded mRNA-seq libraries with
the Stranded mRNA-Seq kit (KAPA Biosystems) as per manufacturer's protocol. The
strand-specific single-end RNA-sequencing was performed using Illumina HiSeq2500.

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374 **RNA-seq analysis**

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376 The qualities of sequencing reads were evaluated using NGS QC Toolkit (v2.3.3) (62) 377 and high-quality reads were extracted. The human reference genome sequence and gene 378 downloaded Illumina annotation data, hg19, were from iGenomes 379 (https://support.illumina.com/sequencing/sequencing_software/igenome.html). The viral NCBI 380 genome was downloaded from GenBank 381 (https://www.ncbi.nlm.nih.gov/nuccore/GQ994935.1). The qualities of RNA-sequencing libraries were estimated by mapping the reads onto human transcript and ribosomal RNA 382 383 sequences (Ensembl release 89) using Bowtie (v2.2.9) (63). STAR (v2.5.2b) (53) was 384 employed to align the reads onto the human and viral genomes, Picard (v1.140) 385 (https://broadinstitute.github.io/picard/) was employed to sort the alignments, and HTSeq 386 Python package (64) was employed to count reverse-stranded reads per gene. DESeq2 387 R Bioconductor package (65) was used to normalize read counts and identify differentially 388 expressed (DE) genes. The resulting gene expression analyses are given in 389 Supplementary Tables S1 for viral genes exclusively and S2 for human plus viral genes. 390 The enrichment of DE genes to pathways and GOs were calculated by Fisher's exact test 391 in R statistical package. Genome coverages were calculated using SAMtools (v0.1.19) 392 (66), **BEDTools** (v2.26) (67), bedGraphToBigWig and

(https://genome.ucsc.edu/index.html). The heatmap was generated using Morpheus from 393 394 the Broad Institute (https://software.broadinstitute.org/morpheus/). 395 The data discussed in this publication have been deposited in NCBI's Gene Expression 396 Omnibus (68) and are accessible through GEO Series accession number GSE144747 397 (https://www.ncbi.nlm.nih.gov/geo/guery/acc.cgi?acc=GSE144747). 398 **Cell Culture** 399 400 401 iSLK cells were grown at 37°C with 5% CO₂ in DMEM (Sigma) supplemented with 402 10% Tet-Free fetal bovine serum (FBS, Atlanta Biologicals), 1x penicillin-streptomycin 403 (Sigma), and 2 mM L-glutamine (Fisher). iSLK WT cells were grown in the presence of 404 0.1 mg/mL G418 (Fisher), 1 µg/mL puromycin (Sigma) and 50 µg/mL hygromycin. iSLK-ORF24R328A cells (gift from Dr. Britt Glaunsinger, University of California Berkeley) were 405 406 grown under the same conditions, except 200 µg/mL of hygromycin was used.

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408 siRNA Transfections

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iSLK cells were transfected with 20 or 40 nM siRNA (Silencer Select, ThermoFisher)
using RNAiMAX transfection reagent (Invitrogen) per manufacturer's instruction.
Specifically, we used final concentrations of 40 nM siRNAs for ZFC3H1, MTR4 and
NRDE2 and 20 nM siRNAs ARS2. For PAPα/γ, we used 20 nM each of siRNAs that target
PAPα or PAPγ for a total of 40 nM siRNA. Twenty-four hours after siRNA transfection,
cells were split into new plates and allowed to grow for another 24 hours, after which

doxycycline and NaB was added to induce lytic reactivation. Thus, total RNA was
harvested 72 hours post siRNA transfection and 24 hours post lytic reactivation.
Nontargeting control, PAPα/γ, ZC3H1 and MTR4 siRNAs are the same as previously
used (29). NRDE2 siRNAs are: 5' GGUGUUGUUUGAUGAUAUUtt 3' (s30063) and 5'
GUUUAGUACCUUUUCGAUAtt 3' (s30064).

421

422 Quantitative RT-PCR

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RNA was harvested using TRI reagent (Molecular Research Center, Inc.) according to
the manufacturer's protocol. Following extraction, RNA was treated with RQ1 DNase
(Promega). Oligo dT₂₀ was used to prime cDNA synthesis with MuLV reverse
transcriptase (New England Biolabs). Real-time reactions used iTaq Universal SYBR
Green Supermix (Biorad). Primers are listed in Table S3.

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430 KSHV Reactivation and Infection

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Lytic reactivation of iSLK derived cells was achieved by adding doxycycline (1 μ g/ml) and NaB (1mM). Tissue culture supernatants from iSLK WT cells were collected at 0, 8, 12, 24, 48 and 72 hpi, centrifuged for 5 min at 1000 x g and passed through a 0.45 um filter. Polybrene was added (8 μ g/mL final concentration), and 300 μ L were applied to HEK293 cells grown in a 12-well plate. Cells were centrifuged for 45 min at 30°C and then incubated in 5% CO₂ at 37°C for 2 hours. After this, media was replaced and cells were analyzed by flow cytometry 24 hours later.

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440 Western Blotting

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Cells were lysed in buffer containing 100 mM NaCl, 50 mM Tris-HCl pH 7.4, 1% Triton X-442 100, 1X Protease Inhibitor cocktail (PIC) (Calbiochem) and 250 µM PMSF. Proteins were 443 444 resolved by SDS-PAGE and analyzed by western blot using standard procedures. Antibodies used are rabbit polyclonal anti-ARS2 (Abcam, ab192999), rabbit polyclonal 445 anti-MTR4 (Abcam, Ab70551), rabbit polyclonal anti-NRDE2 (Proteintech, 24968) and 446 447 mouse monoclonal anti-Actin (Abcam, ab6276). Quantitative westerns were performed using infrared detection with an Odyssey Fc and guantification was performed using 448 ImageStudio software (LI-COR Biosciences). 449

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462 Figure Legends

463

464 Fig 1. PPD inactivation affects the temporal expression of KSHV late genes. (A) 465 Diagram of the RNA-seg experiment. iSLK WT cells were transfected with a non-targeting 466 control siRNA or a two-siRNA pool targeting PAPa/y, ARS2, or both PAPa/y and ARS2 467 combined (dKD). Total RNA was harvested three days after siRNA transfection and 24 468 hours after lytic reactivation. Stranded mRNA-seq libraries were prepared and sequenced. (B) Efficiency of knockdown of PAPa, PAPy, and ARS2 in iSLK WT cells. 469 470 Due to the lack of robust antibodies, PAP α and PAPy knockdown efficiency was 471 determined by gRT-PCR. Bar graphs show PAPα and PAPγ mRNA levels in iSLK WT cells treated with siRNAs targeting PAPa and PAPy. Because RNA knockdown does not 472 473 necessarily correlate with protein loss, we assayed for loss of functional activity. To do 474 so, we measured the RNA levels of a known PPD target, NEAT1, under the same conditions used for RNA-seq. Bar graph shows NEAT1 levels determined by gRT-PCR 475 476 in iSLK WT cells depleted of PAP α /y. Values are displayed relative to siCtrl after 477 normalization to the 18S rRNA level. All values are averages, and the error bars are 478 standard deviations (n = 3). ARS2 knockdown efficiency was determined by quantitative 479 western blot. Actin serves as loading control. (C) Heatmap showing the log2 fold change 480 (FC) relative to siCtrl for all KSHV genes in samples depleted of PAP α/γ , ARS2 or dKD. 481 Genes are arranged in increasing log2 FC order based on PAP α/γ , where red represents 482 maximum fold change and blue represents Log2 FC <1.5. (D) Pie charts showing the 483 distribution of upregulated (>4-fold) KSHV genes in PAP α/γ (top) and dKD (bottom) 484 according to their phase of expression (30).

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Fig 2. KSHV late transcripts are degraded by PPD/PAXT. (A) Efficiency of knockdown 486 487 of ZFC3H1 and MTR4 in iSLK WT cells. Bar graphs show ZFC3H1 and NEAT1 mRNA levels in iSLK WT cells treated with ZFC3H1 siRNAs. Increased levels of NEAT1 indicate 488 489 that PPD was effectively inactivated upon ZFC3H1 depletion. MTR4 knockdown 490 efficiency was determined by quantitative western blot. Actin serves as loading control. (*) nonspecific band. (B, D and F) Integrative genome viewer (IGV) browser screenshots 491 showing ORF75 (B), ORF52 (D) and K8.1 (F) reads in samples depleted of PAPα/y, ARS2 492 493 or dKD. Each condition is depicted at the same scale. (C, E and G) Bar graphs showing 494 relative ORF75 (C), ORF52 (E) and K8.1 (G) mRNA levels in iSLK WT cells depleted of PAPa/y (green), ARS2 (orange), dKD (red), ZFC3H1 (purple) and MTR4 (light blue). Total 495 RNA was harvested 24 hpi and analyzed by qRT-PCR. Values are displayed relative to 496 497 siCtrl after normalization to the 18S rRNA level. All values are averages, and the error 498 bars are standard deviations (n = 3). P values were determined by two-tailed unpaired Student's *t* test: * < 0.05; ** < 0.01; *** < 0.001. 499

500

Fig 3. PPD upregulation of late genes is independent of ORF24. (A) IGV browser
screenshot showing ORF24 reads in samples depleted of PAPα/γ, ARS2 or dKD. (B) Bar
graphs showing relative ORF24 mRNA levels in iSLK cells depleted of PAPα/γ (green),
ARS2 (orange) and dKD (red). (C) Bar graphs showing relative ORF52, ORF75 and K8.1
mRNA level in iSLK WT (gray) and iSLK ORF24^{R328A} (white) cells at 48 hpi. (D, E and F)
Bar graphs showing relative ORF52 (D), ORF75 (E) and K8.1 (F) mRNA levels in iSLK
ORF24^{R328A} cells depleted of PAPα/γ (green), ARS2 (orange), dKD (red), ZFC3H1

508 (purple) and MTR4 (light blue). Total RNA was harvested 24 hpi and analyzed by qRT-509 PCR. Values are displayed relative to siCtrl after normalization to the 18S rRNA level. All 510 values are averages, and the error bars are standard deviations (n = 3). *P* values were 511 determined by two-tailed unpaired Student's *t* test: * < 0.05; ** < 0.01; *** < 0.001.

512

513 Fig 4. KSHV late transcripts evade PPD degradation at their proper time of expression (A, B and C) Bar graphs showing relative ORF75 (A), ORF52 (B) and K8.1 514 (C) mRNA levels in iSLK WT cells treated with siRNAs targeting PAP α/γ (green) or a 515 516 control siRNA (gray). Total RNA was harvested at 24, 48 and 72 hpi. Values were 517 calculated relative to siCtrl at 24 hpi (gray) and normalized to the 18S rRNA level. Note 518 that the data are plotted on a log scale due to the strong up-regulation of late genes after 519 48 hpi. (D) Bar graphs showing relative ORF57 DNA levels in iSLK WT cells depleted of 520 PAPa/y (green) or ZFC3H1 (purple). DNA was harvested at 0, 12, 24 and 48 hpi. Values 521 were calculated relative to siCtrl (gray) at 0 hpi. (E) Bar graph of flow cytometry analysis 522 showing percentage of GFP-positive HEK293 cells infected with supernatants collected 523 from iSLK WT cells at 0, 8, 12, 24, 48 and 72 hpi. All values are averages, and the error 524 bars are standard deviations (n = 3). P values were determined by Student's t test: * < 525 0.05; ** < 0.01; *** < 0.0001.

526

Fig 5. NRDE2 protects viral late transcripts from degradation. (A) NRDE2 knockdown
efficiency was determined by quantitative western blot. Actin serves as loading control.
(*) nonspecific band. (B and C) Bar graphs showing relative ORF75, ORF52 and K8.1
mRNA levels at 24 (B) and 48 (C) hpi in iSLK WT depleted of PAPα/γ (green), NRDE2

(orange) and PAP α/γ and NRDE2 combined (blue). Total RNA was harvested 24 or 48 hpi and analyzed by qRT-PCR. Values are displayed relative to siCtrl after normalization to the 18S rRNA level. All values are averages, and the error bars are standard deviations (n = 3). *P* values were determined by two-tailed unpaired Student's *t* test: * < 0.05; ** < 0.01; *** < 0.001.

536

Fig 6. Model of PPD regulation of KSHV late genes. (A) KSHV late genes are cryptically transcribed at 24 hpi, but the transcripts do not accumulate due to PPD. (B) At 48 hpi, KSHV late transcripts evade PPD and accumulate at high levels. We speculate that KSHV replication compartments coalesce with nuclear speckles where NRDE2 protects viral transcripts from PPD degradation by sequestering MTR4.

542

Table S1. Differential expression of KSHV genes. This spreadsheet contains the expression levels of all KSHV genes in samples depleted of $PAP\alpha/\gamma$, ARS2 and dKD relative to siCtrl.

546

Table S2. Differential expression of human and KSHV genes. This spreadsheet
contains the expression levels of human and KSHV genes in samples depleted of
PAPα/γ, ARS2 and dKD relative to siCtrl.

550

Table S3. Primers used in this study. Target, sequence, and primer number (ID) forall PCR primers used herein.

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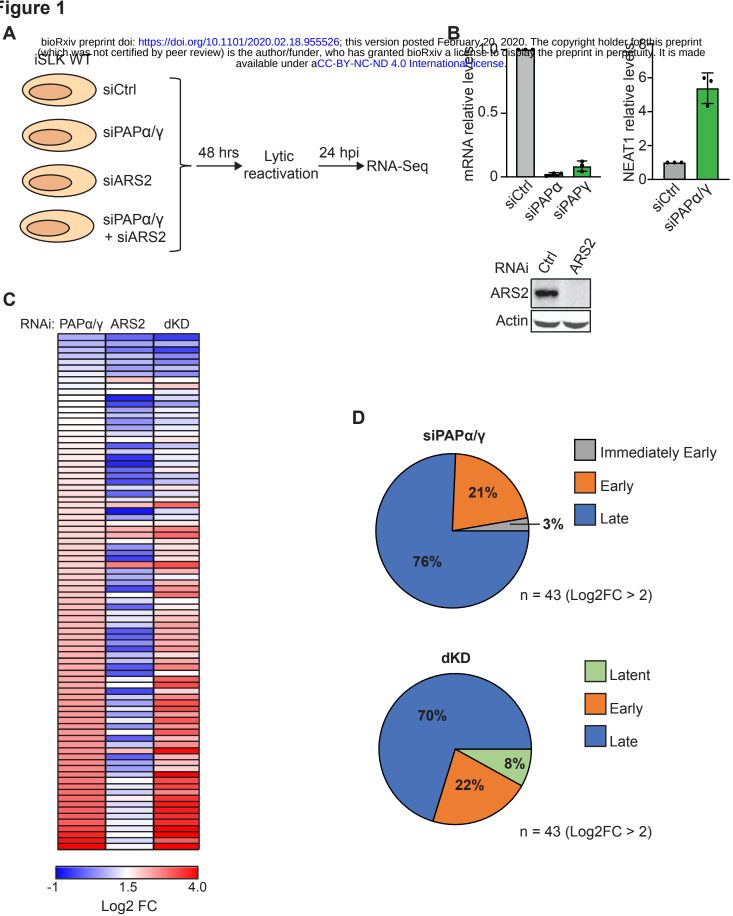
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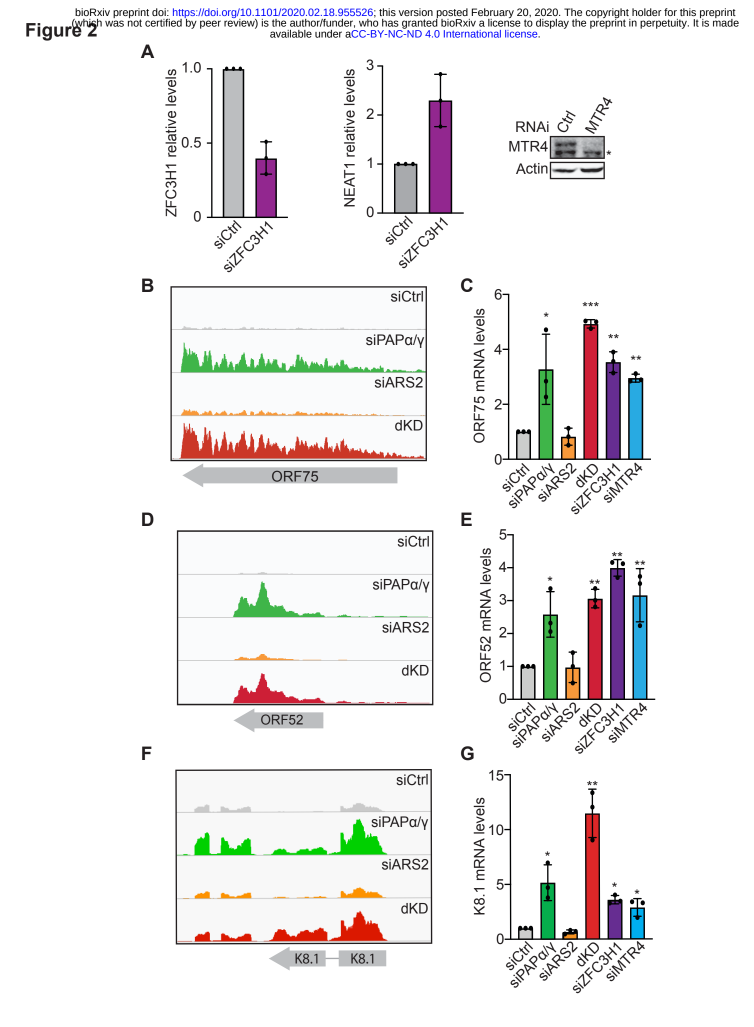
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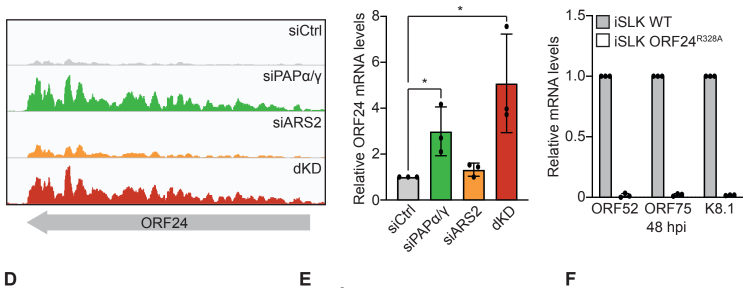
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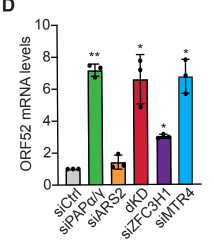
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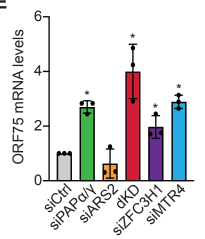
Figure 1

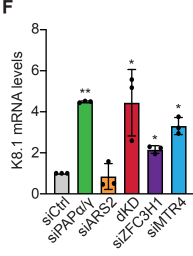


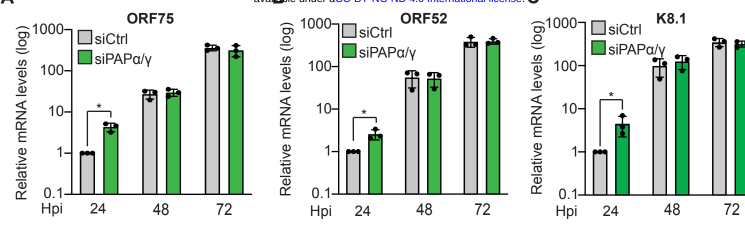


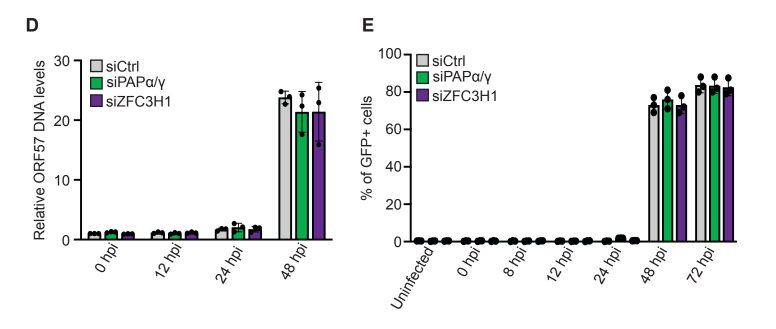












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Figure 5 bioRxiv preprint doi: https://doi.org/10.1101/2020.02.18.955526; this version posted February 20, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license. ORF75 - 24 hpi ORF52 - 24 hpi K8.1 - 24hpi WOR 3 6-5 **Relative mRNA levels Relative mRNA levels** Relative mRNA levels Å **RNAi** 4 2 4 NRDE2 3 2 Actin 2 1 • 1 silvedez sipapati ett Silved Ez sip_{*N} sipapat a Sib_×N SiCtry silvedez 🕨 SiCtry sipana, 0 0 Sib XN sictry 🕨 0 С ORF75 - 48 hpi ORF52 - 48 hpi K8.1 - 48hpi 1.5 1.5 Relative mRNA levels 1.5 Relative mRNA levels Relative mRNA levels 1

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Figure 6

