1	E6 proteins from high-risk HPV, low-risk HPV, and animal papillomaviruses activate the
2	Wnt/ β -catenin pathway through E6AP-dependent degradation of NHERF1
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4	Camille M. Drews ¹ , Samuel Case ¹ , Scott B. Vande Pol ^{1*}
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6	¹ Department of Pathology, University of Virginia, Charlottesville, VA, USA
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8	Short title: High and low-risk E6 degrade NHERF1 to activate Wnt signaling
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10	* Corresponding author
11	Email: vandepol@virginia.edu (SVP)
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24 Abstract

High-risk human papillomavirus (HPV) E6 proteins associate with the cellular 25 26 ubiquitin ligase E6-Associated Protein (E6AP), and then recruit both p53 and certain 27 cellular PDZ proteins for ubiguitination and degradation by the proteasome. Low-risk 28 HPV E6 proteins also associate with E6AP, yet fail to recruit p53 or PDZ proteins; their 29 E6AP-dependent targets have so far been uncharacterized. We found a cellular PDZ 30 protein called Na+/H+ Exchanger Regulatory Factor 1 (NHERF1) is targeted for 31 degradation by both high and low-risk HPV E6 proteins as well as E6 proteins from 32 diverse non-primate mammalian species. NHERF1 was degraded by E6 in a manner 33 dependent upon E6AP ubiquitin ligase activity but independent of PDZ interactions. A 34 novel structural domain of E6, independent of the p53 recognition domain, was 35 necessary to associate with and degrade NHERF1, and the NHERF1 EB domain was 36 required for E6-mediated degradation. Degradation of NHERF1 by E6 activated canonical Wnt/ β -catenin signaling, a key pathway that regulates cell growth and 37 38 proliferation. Expression levels of NHERF1 increased with increasing cell confluency. This is the first study in which a cellular protein has been identified that is targeted for 39 40 degradation by both high and low-risk HPV E6 as well as E6 proteins from diverse 41 animal papillomaviruses. This suggests that NHERF1 plays a role in regulating 42 squamous epithelial growth and further suggests that the interaction of E6 proteins with 43 NHERF1 could be a common therapeutic target for multiple papillomavirus types.

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47 Author summary

48	Papillomaviruses cause benign squamous epithelial tumors through the action of virally
49	encoded oncoproteins termed E6 and E7, which are classified as either high or low-risk
50	based upon the propensity of the tumor to evolve into cancer. E6 proteins from both
51	high and low-risk HPVs interact with a cellular ubiquitin ligase called E6AP. High-risk E6
52	proteins hijack E6AP ubiquitin ligase activity to target p53 for degradation. Degradation
53	targets of the low-risk E6 proteins in complex with E6AP have not been described.
54	Here, we describe a protein called NHERF1 that is targeted for degradation by both
55	high and low-risk E6 proteins, as well as E6 proteins from diverse animal species.
56	Degradation of NHERF1 resulted in activation of an oncogenic cellular signaling
57	pathway called Wnt. Identification of NHERF1 as a highly conserved E6 degradation
58	target could inform therapies directed against both low-risk HPVs and cancer-inducing
59	high-risk HPVs.
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70 Introduction

Human papillomaviruses (HPVs) are small DNA tumor viruses that cause 71 72 squamous epithelial papillomas in which the virus replicates. The papillomas are initially 73 benign and the host is usually able to clear the underlying HPV infection over time. 74 However, a subset of HPV infections may result in lesions that persist and grow to 75 harmful size or that have a propensity to evolve into carcinomas [1]. The cancer-causing 76 HPV types are called high-risk and the most commonly occurring high-risk types are 77 HPV16 and HPV18. Worldwide, high-risk HPVs are responsible for 5% of cancers, with cervical cancer being the most common [2]. HPV types that are not associated with 78 79 malignancies are termed low-risk HPV; although low-risk for malignancies, the size and 80 location of the benign papillomas can render these lesions medically serious [3]. 81 Beyond HPVs, papillomaviruses have been isolated from mammalian species 82 including rodents, primates, bats, cetaceans, and ungulates [4], and are clustered into 83 related genera based upon the divergence of the L1 capsid protein nucleotide sequence (both high and low-risk mucosal HPV types discussed in this study belong to the 84 85 primate Alpha genera) [5]. Most non-human papillomaviruses encode E6 proteins that are similar in predicted fold to high-risk HPV16 E6 [6]. When diverse mammalian 86 87 papillomaviruses are clustered based on their E6 sequence similarity, two main groups 88 of papillomaviruses emerge: those that encode E6 proteins that bind to the Notch coactivator Mastermind Like 1 (MAML1), and those that bind to a cellular E3 ubiquitin 89 ligase called E6-Associated Protein (E6AP) [7]. An E6 protein that preferentially binds 90 91 MAML1 suppresses MAML1 transcriptional activation, while an E6 protein that 92 preferentially binds E6AP stimulates E6AP E3 ubiquitin ligase activity to then target

additional cellular proteins recruited by E6 to E6AP for ubiquitination and degradation by
the proteasome [7].

The difference between the propensity of high and low-risk HPVs to cause 95 96 cancer is secondary to differences between their respective E6 and E7 oncoproteins [8]. 97 E6 and E7 from both high and low-risk HPVs bind cellular E3 ubiquitin ligases and 98 hijack their ubiquitin ligase activity to perturb certain cellular proteins that are recruited 99 by E6 or E7 [9]. Both high-risk and low-risk E7 proteins interact with ubiguitin ligases of 100 the cullin and N-end rule families and target the degradation of additional cellular 101 proteins recruited to E7 such as pocket-family proteins (RB, RBL1, and RBL2) and 102 PTPN14 [10, 11]. High and low-risk E7 proteins target certain cellular proteins in 103 common (such as the RBL2 p130 pocket protein) [12]. However only high-risk HPV E7 104 types interact with and target the degradation of the retinoblastoma (RB) pocket protein 105 [13, 14], which has implications for the carcinogenic properties of high-risk E7. High and 106 low-risk Alpha genera HPV E6 proteins interact with the cellular E3 ubiquitin ligase 107 E6AP [15-17], but only cellular proteins targeted for degradation by the high-risk E6 108 protein (such as p53) are well established [18, 19].

Another striking difference between high and low-risk E6 is the presence of a PDZ binding motif (PBM) at the extreme carboxy terminus of high-risk E6 proteins [20-22]}. The high-risk PBM enables E6 to interact with a group of cellular proteins termed PDZ proteins, all of which contain PDZ (PSD-90/Dlg1/ZO-1) homology domains [23]. The targeted degradation of cellular proteins that are recruited through interaction with the high-risk E6 PBM has been controversial, but the E6 PBM functionally promotes retention of the viral DNA plasmid within infected cells [24]; the E6 PBM function can be

rescued by disruption of p53 function [25]. Although low-risk E6 proteins bind E6AP,
they do not have a PBM at the carboxy-terminus [20], do not interact with p53 [17], and
no cellular targets of the low-risk E6+E6AP complex have been described. Such cellular
targets would be presumed to be of exceptional interest since they would be common to
both high and low-risk E6 proteins, just as RBL2 is a common target of the high and
low-risk HPV E7 protein.

122 In this study, we identify the PDZ-adapter protein NHERF1 as degraded by both 123 high and low-risk E6 proteins, in a manner dependent upon the ubiquitin ligase activity 124 of E6AP and the proteasome. Other E6 proteins from diverse species where E6 could bind E6AP were also able to initiate NHERF1 degradation, indicating the conservation 125 126 of this function. Interaction of NHERF1 with E6 required prior association of E6 with 127 E6AP, and we identified a novel interaction domain within 16E6 that is required. Finally, 128 the targeted degradation of NHERF1 by both low and high-risk E6 proteins resulted in 129 the activation of canonical Wnt signaling, connecting the degradation of NHERF1 by E6 130 to the activation of an oncogenic signaling pathway.

131

132 **Results**

133 NHERF1 protein levels are reduced by high and low-risk E6 proteins in the

134 presence of E6AP_WT.

NHERF1 was previously shown to be degraded by HPV16 E6 (16E6) (but not by
HPV18 E6 (18E6) or HPV11 E6 (11E6)) through an interaction requiring the PBM of
16E6 [26]. In our proteomic studies of cellular proteins that associate with the 16E6 and
18E6 PBMs [27], we did not identify NHERF1, but in other experiments observed a

139 reduction of NHERF1 protein levels by 16E6, 18E6, and 11E6. In order to characterize 140 the reduction of NHERF1 by these E6 proteins, we performed transient transfections into E6AP-null 8B9 cells reconstituted with either WT E6AP (E6AP_WT) or a mutant 141 142 E6AP defective in ubiquitin ligase activity (E6AP_Ub⁻). E6APs were co-transfected with 143 plasmids encoding p53, NHERF1, and 16E6, 16E6 deleted of the PBM (16E6\(\Delta PBM)\), 144 11E6, or 18E6. Consistent with published literature, p53 was degraded by high-risk E6 145 proteins (16E6 and 18E6) independently of a PBM [25] and dependent upon E6AP 146 ubiguitin ligase activity [28] (Fig 1). Expression of 11E6 together with E6AP WT 147 resulted in a lack of p53 degradation by low-risk E6 (11E6), corroborating published findings [17, 29]. However, NHERF1 protein levels were reduced by each observed E6 148 149 protein (Fig 1A), in contrast to what has been previously published [26].

150 To ensure the reduction of NHERF1 by either high or low-risk E6 proteins was 151 not due to an overexpression artifact, we performed an E6 titration experiment (Fig 1B). 152 Representative western blots from which the quantification in Fig 1B was derived are 153 shown in S1 Fig. Three different E6 proteins were used: 16E6 WT, 16E6∆PBM, and 154 11E6_WT. We used p53 as a control for 16E6-mediated degradation, as multiple 155 studies have shown low-risk E6 proteins (11E6) do not degrade p53 [17, 29] while high-156 risk 16E6 is able to degrade p53 independent of the presence of a PBM [25]. Observing 157 the degradation of p53 in cells expressing variable amounts of E6 provided a guide for 158 physiologically relevant E6 expression levels. NHERF1 and p53 protein levels were 159 similarly reduced by both 16E6 WT and 16E6 Δ PBM at the various E6 titrations (Fig 160 1B). 11E6 WT was unable to initiate the degradation of p53 but targeted NHERF1 at

161 levels similar to those required by 16E6. Deletion of the 16E6 PBM did not impact the

degradation of p53 or reduction of NHERF1 protein levels by 16E6.

163 NHERF1 protein levels are sensitive to cell confluency.

164 We established that NHERF1 protein levels are reduced by E6 in a transient transfection system. To determine whether low levels of stable 16E6-expression could 165 166 initiate the reduction of NHERF1 protein levels, we retrovirally transduced normal 167 immortalized keratinocytes with either empty vector or 16E6 WT and observed 168 NHERF1 protein levels. Initially, our results were variable. We hypothesized that 169 keratinocyte confluency may affect NHERF1 protein levels. To test this possibility, we 170 seeded vector-transduced and 16E6_WT-transduced keratinocytes at three different cell densities: 5x10³ cells/cm² (very sub-confluent), 1.3x10⁴ cells/cm² (sub-confluent), 171 172 and 2.6x10⁴ cells/cm² (mid-confluent). NHERF1 protein levels increased with an increase in cell density and 16E6 WT consistently reduced NHERF1 (Fig 2A). In order 173 174 to determine if changes in NHERF1 levels with confluency were secondary to changes 175 in NHERF1 RNA levels, we performed qPCR on RNA extracted from keratinocytes 176 retrovirally transduced with vector or 16E6 WT and plated as in Fig 2A. Interestingly, 177 NHERF1 RNA levels did not differ between keratinocytes seeded at different densities 178 expressing either empty vector or 16E6 WT (Fig 2B).

179 **E6-mediated degradation of NHERF1 occurs via the proteasome.**

Because the ability of each E6 protein to reduce NHERF1 protein levels was also dependent upon the ubiquitin ligase activity of E6AP (Fig 1A), we hypothesized that E6 reduction of NHERF1 levels would be secondary to proteasome activity. We seeded retrovirally transduced keratinocytes expressing either empty vector or 16E6_WT at

184 similar confluency and treated with either DMSO, mitomycin C (MMC) to induce p53 185 [30], or the proteasome inhibitor MG132 at differing concentrations for 8 hours. As 186 expected, p53 levels increased in vector keratinocytes treated with MMC compared to 187 untreated cells as well as in 16E6_WT cells exposed to increasing concentrations of 188 MG132 [28] (Fig 3). NHERF1 protein levels increased significantly in a dose dependent 189 manner upon treatment with MG132 in parallel to that seen with p53. (Fig 3, lanes 3-8). 190 This indicated that NHERF1 is degraded through the proteasome by E6 in a manner 191 dependent upon WT E6AP. 192 E6-mediated degradation of NHERF1 is conserved across papillomaviruses from diverse hosts. 193 194 The observation that NHERF1 was targeted by both high and low-risk HPV E6 195 proteins suggested that NHERF1 may also be a target of diverse non-primate E6 196 proteins. We examined the ability of E6 proteins from multiple different genera and 197 different mammalian species to target NHERF1 for degradation (Fig 4). E6 proteins that 198 preferentially bind MAML1 were unable to degrade NHERF1 (Fig 4A and 4B). All of the 199 tested Alpha (primate), Dyodelta (boar), and Dyopi (porpoise) genera E6 proteins that 200 bind E6AP targeted NHERF1. While E6AP-binding was necessary it was not sufficient, 201 as E6 proteins from Omega (polar bear, UmPV1) and Omikron (cetaceans, PphPV1 202 and TtPV5) did not degrade NHERF1 (Fig 4A). Interestingly, E6 proteins that bind E6AP 203 but did not target NHERF1 degradation sequence-clustered separately from E6 proteins 204 that did target NHERF1 degradation, suggesting evolutionary divergence of this function 205 (Fig 4B).

A novel 16E6 substrate interaction domain is required for 16E6 degradation of NHERF1.

Because the ability of E6 to degrade NHERF1 was not dependent upon the 208 209 presence of a PBM (Figs 1 and 4), we attempted to identify which residue(s) of 16E6 210 were required to mediate degradation of NHERF1. The crystal structure of 16E6 211 complexed with the E6-binding peptide from E6AP [31] (Fig 6A) was examined to 212 identify amino acids that were at least 20% exposed, resulting in over eighty candidate 213 residues (S2 Fig). Candidate residues were individually mutated in the context of the 214 16E6 gene and the resulting point mutants were screened for their ability to degrade 215 NHERF1 in the presence of E6AP_WT in transiently transfected C33 cells. To ensure 216 our point mutants were not functionally defective (i.e. could not fold properly or could not 217 interact with E6AP), we also screened the mutants for ability to degrade p53. A 218 selection of mutants and the results of the screen are shown in Fig 5. Four mutants 219 stood out as selectively defective in their ability to degrade NHERF1 (Fig 5B) while still 220 being able to degrade p53 (Fig 5C): F69A, K72A, F69R and a double mutant: 221 F69A/K72A. As evidenced in the crystal structure of 16E6, the side chains of F69 and 222 K72 (Fig 6B) are located along the connecting alpha-helix that links the amino-terminal 223 and carboxy-terminal zinc-structured domains of 16E6. The F69 and K72 side chains 224 are aligned and adjacent on the connecting helix, which is on the opposite side of 16E6 225 from the p53 interaction surface [32] (Fig 6C).

We had identified the 16E6_F69A/K72A mutant in a transient transfection screen. To ensure the identified 16E6_F69A/K72A double mutant was selectively defective for degrading NHERF1 in the context of a stable cell line, keratinocytes

229 retrovirally transduced with empty vector, 16E6 WT, 16E6∆PBM, 16E6 F69A/K72A, or 230 11E6_WT were seeded at equal confluency and lysates prepared. Keratinocytes 231 expressing 16E6 WT, 16E6∆PBM, and 11E6 WT degraded NHERF1 (Fig 7, lanes 2, 3, 232 5). However, keratinocytes expressing 16E6_F69A/K72A were unable to stimulate the 233 degradation of NHERF1 (Fig 7, lane 4), indicating a novel substrate interaction domain 234 important for 16E6-mediated degradation of NHERF1. 235 Degradation of NHERF1 by 16E6 requires the NHERF1 EB domain. 236 Because the PBM of E6 proteins is not required to initiate the degradation of 237 NHERF1 (Figs 1, 4, and 7), we hypothesized that neither of the PDZ domains of 238 NHERF1 would be required for 16E6 to initiate NHERF1 degradation. We truncated 239 NHERF1 and deleted several characterized domains within the protein [33, 34] (Fig 8A). 240 E6AP-null 8B9 cells were co-transfected with 16E6 WT, NHERF1 truncations, HA GFP, and either E6AP Ub⁻ or E6AP WT. NHERF1 protein levels were guantified, 241 and then normalized to the internal transfection control (HA_GFP). The various 242 243 NHERF1 truncations displayed different expression levels. To account for these 244 variations, levels of NHERF1 truncations in the presence of E6AP Ub- were set to 245 100% and the expression level of the corresponding NHERF1 truncation in the 246 presence of E6AP WT was normalized accordingly (Fig 8B and 8C, bar graphs). All 247 NHERF1 truncations containing the EB domain were targeted for degradation by 16E6 248 in the presence of E6AP_WT (highlighted in green in Fig 8A). Truncations of NHERF1 249 that lacked the EB domain were not targeted for degradation by 16E6 (highlighted in red 250 in Fig 8A). In addition, the NHERF1 PBM was not required for 16E6 mediated 251 degradation (Fig 8C, lanes 5 vs. 6 and 9 vs. 10).

252 We identified the NHERF1 EB domain as a requirement for 16E6 mediated 253 degradation and the importance of 16E6 residues F69 and K72. In order to examine the 254 interactions between the 16E6+E6AP+NHERF1 complex, all three proteins were 255 expressed in a yeast three-hybrid system so as to detect the heterotrimeric complex. We fused 16E6_WT and ubiquitin ligase dead E6AP (E6AP_Ub⁻) to the LexA DNA 256 257 binding domain and co-expressed this fusion with either vector, 16E6 WT, or 258 16E6 F69A/K72A in yeast containing a LexA responsive LacZ reporter. These yeast 259 were then mated to yeast expressing native p53 or Gal4 (G4) transactivator fusions to 260 NHERF1 121-358 (containing the EB domain), NHERF1 121-297 (deleted of the EB 261 domain), 16E6_WT, or the tyrosine phosphatase PTPN3 (a PDZ protein) (Fig 9). The 262 LexA 16E6 fusion co-expressed with p53 (in the absence of E6AP) resulted in very 263 weak activation of the LacZ reporter (spot 4B) while co-expression with G4 PTPN3 264 resulted in strong transactivation (spot 6B), but no interaction with NHERF1 (spots 2B 265 and 3B). We then co-expressed 16E6 and E6AP by using a LexA_E6AP_Ub⁻ fusion 266 together with native 16E6. When LexA_E6AP_Ub⁻, untagged 16E6_WT, and p53 were 267 co-expressed, a strong activation of the LacZ reporter was observed (Fig 9, spot 4D), 268 illustrating that while p53 has a weak direct interaction with 16E6, it interacts strongly 269 with 16E6 bound to E6AP. This activation was also seen with 16E6 F69A/K72A in the 270 presence of LexA_E6AP_Ub⁻ and p53 (Fig 9, spot 4E), indicating the preserved ability 271 of the 16E6_F69A/K72A double mutant to bind E6AP and recruit p53. When 272 LexA_E6AP_Ub⁻, 16E6_WT, and G4_NHERF1 121-358 (contains the EB domain) were 273 co-expressed, we observed activation of the LacZ reporter, indicating the recruitment of 274 NHERF1 to E6AP by 16E6_WT (Fig 9, spot 2D). Truncating the EB domain from the

275	G4_NHERF1 (G4_NHERF1 121-297, spot 3D) or the use of the 16E6_F69A/K72A
276	double mutant (spot 2E) ablated the reporter transactivation, indicating the requirement
277	of the EB domain and the importance of 16E6 residues F69 and K72 in the interaction
278	of the 16E6+E6AP+NHERF1 complex.
279	E6AP-dependent NHERF1 degradation by E6 activates the canonical Wnt/ β -
280	catenin pathway.
281	It has been shown that high-risk HPV E6 proteins augment the canonical Wnt/ β -
282	catenin signaling pathway [35-39]. Additionally, it has been shown that NHERF1 inhibits
283	the canonical Wnt/ β -catenin signaling pathway through multiple mechanisms. NHERF1
284	forms a complex with β -catenin [40] and can also bind to the intracellular PBM of certain
285	isoforms of Frizzled [41], a G-protein coupled receptor important in the activation of the
286	canonical Wnt signaling pathway. Therefore, we hypothesized E6 degradation of
287	NHERF1 would activate the Wnt/ β -catenin signaling pathway in cells expressing E6. To
288	test this possibility, we utilized the TOP/FOP luciferase reporter assay. 16E6,
289	16E6 Δ PBM, 11E6, and 18E6 all stimulated the activity of the Wnt/ β -catenin pathway
290	over vector-transfected cells (Fig 10). However, cells transfected with 16E6_F69A/K72A
291	were unable to augment the canonical Wnt pathway over vector levels, indicating that
292	the ability of E6 to degrade NHERF1 is required for E6 activation of the canonical
293	Wnt/β-catenin signaling pathway.
294	The earlier Accardi et al. study proposed that expression of 16E7 sensitized
295	NHERF1 for degradation by the induction of NHERF1 phosphorylation [26]. Our

experiments did not show either E7 induction of slow-migrating NHERF1 296

297 phosphorylated isoforms or an enhancement of E6-NHERF1 degradation upon co298 expression of E7 (S3 and S4 Figs).

299

300 Discussion

301 E6 proteins from papillomaviruses can be separated into distinct groups: those 302 that bind MAML1 and repress Notch signaling, and those that bind E6AP and hijack its 303 ubiquitin ligase activity [7, 42-44]. E6 proteins from papillomaviruses in the Alpha, 304 DyoDelta, Dyopi, Omega, and Omikron genera all behave similarly in that they bind E6AP and activate its ubiquitin ligase activity [7]. Here, we describe the degradation of 305 306 NHERF1 by E6 proteins from both high and low-risk HPVs, as well as from 307 papillomaviruses from multiple divergent mammalian species. The ability of these E6s 308 to degrade NHERF1 is dependent upon E6AP (Fig 1) and the proteasome (Fig 3). In 309 addition, we identify two amino acids in 16E6 (F69 and K72) that are necessary for E6-310 mediated degradation of NHERF1. These two residues are aligned, and adjacent in the 311 outwardly oriented face of the E6 connecting alpha helix, suggesting a novel interaction 312 domain (Fig 6B and 6C). NHERF1 degradation by E6 requires the NHERF1 EB domain, 313 but does not require the PBM at the extreme carboxy terminus of NHERF1 (Fig 8B and 314 8C). The ability of E6 proteins to degrade NHERF1 augments the canonical Wnt/ β -315 catenin signaling (Fig 10), an oncogenic pathway frequently active in cancer. NHERF1 is the product of the SLC9A3R1 gene. SLC9A3R1 mRNA is broadly 316 317 expressed in epithelia, with the highest mRNA expression in kidney, gut, and 318 esophagus. NHERF1 is not developmentally essential, although mice have considerably 319 reduced lifespans [45]. NHERF1-null mice are prone to phosphate wasting, brittle bone

structure, and hydrocephaly [45] due to the mislocalization of proteins with which
NHERF1 normally associates [45-47]. NHERF1 contains two PDZ domains and an EB
domain at the carboxy terminus through which it interacts with ezrin, radixin, and
moesin to link itself, and proteins to which it is bound, to the actin cytoskeleton network
[48]. While the functions of NHERF1 are varied due to its role as a scaffold, multiple
studies indicate it regulates cell growth and differentiation, two key cellular functions that
papillomaviruses disrupt in the process of viral infection.

327 Whether NHERF1 is a tumor suppressor or an oncogene has been debated in 328 the literature. There are numerous papers regarding NHERF1 human cancer 329 phenotypes, but they are collectively inconsistent [49, 50]. NHERF1-null mice do not 330 have a direct cancer phenotype, but have lengthened intestines [51], indicating a growth 331 regulatory function of NHERF1. The diminished life span of NHERF1 null mice could 332 limit observation of cancer traits. However, a recent in vivo study provided strong genetic support for NHERF1 as a tumor suppressor. APC^{Min/+} mice bred as either 333 334 heterozygote or knockout for NHERF1 experience considerably shorter survival than 335 their NHERF1-expressing counterparts due to increased tumor burden, demonstrating a tumor suppressor phenotype for NHERF1 [51]. Additionally, these APC^{Min/+} mice lacking 336 337 NHERF1 have greater activation of Wnt/ β -catenin signaling, suggesting NHERF1 acts 338 as a negative regulator of this oncogenic pathway. NHERF1-associated proteins that plausibly could regulate cell proliferation are numerous and include β -catenin [40], 339 340 Frizzled [41], G-protein coupled receptors (β -adrenergic type 2, [52]), receptor tyrosine 341 kinases (PDGFR, [53]), phosphatases (PTEN, [54]), transcriptional coactivators (YAP1,

[55]), ion channels (Kir1.1 and CFTR, [56]), phospholipase-C [57], and actin anchoring
proteins (ezrin, radixin, and moesin, [48]).

Several studies have indicated that HPV E6 proteins can activate canonical 344 345 Wnt/β-catenin signaling [35-39]. Our work expands and builds upon the scope of 346 these studies. The ability of E6 to degrade NHERF1 and activate Wnt signaling may 347 aid in propagation of papillomaviruses by enhancing the stimulation of cellular 348 proliferation and promoting cell survival. There are numerous cell growth regulatory 349 avenues that E6 could manipulate by degrading NHERF1 and within this study we 350 have explored one possibility: the canonical Wnt/β-catenin signaling pathway (Fig 351 10); other possibilities will be the subject of future studies.

The EB domain of NHERF1 is required for E6-mediated degradation in the 352 353 presence of E6AP (Fig 8B and 8C). This domain is responsible for linking NHERF1 to the actin cytoskeleton network via interaction with ERM proteins [48]. NHERF1 has a 354 355 PBM at its extreme carboxy terminus and when the EB domain is not bound to ERM 356 proteins, the NHERF1 PBM can self-associate with the NHERF1 PDZ2 domain, 357 resulting in a closed NHERF1 conformation [33]. The head-to-tail closed NHERF1 confirmation is not required for E6-mediated degradation, as an NHERF1_APBM mutant 358 359 was still targeted for degradation by E6 in the presence of E6AP WT (Fig 8C). Nor was 360 the 16E6 PBM required for degradation of NHERF1 (Figs 1, 4, and 7), contrary to a 361 prior report [26].

In addition to the requirement of the NHERF1 EB domain, two E6 residues, F69 and K72, are necessary for E6-mediated NHERF1 degradation (Figs 5 and 7). Crucially, the 16E6_F69A/K72A double mutant can still initiate the degradation of p53, indicating it

is still able to bind E6AP, recruit p53 to the complex, and trigger ubiquitination. The F69
and K72 residues are also required to form a tri-molecular complex between E6AP, E6,
and NHERF1 in yeast (Fig 9, spot 2E vs. 2D). Like the association of E6 with p53,
NHERF1 does not interact directly with E6, but requires prior association of E6 with
E6AP, indicating that NHERF1 requires an altered conformation of E6 that is secondary
to E6 binding to E6AP [58].

371 As we were testing the ability of E6 proteins to degrade NHERF1 in stable 372 keratinocyte cell lines, we discovered that NHERF1 protein levels are sensitive to cell 373 confluency (Fig 2A). The relationship between NHERF1 and cell confluency may 374 contribute to the lack of identification of NHERF1 as a degradation target of low-risk E6 375 proteins in the past, as well as differences between our studies and a prior publication 376 [26]. It is likely that this observation underlies disparate findings between different 377 laboratories regarding NHERF1 cancer associated traits [49, 50]. Future studies of 378 NHERF1 must take into account and carefully control cell densities when performing 379 experiments.

380 Binding to E6AP is necessary for E6-induced degradation of NHERF1, but it is 381 not sufficient, as three tested E6 proteins that bind E6AP do not target NHERF1 for 382 degradation: UmPV1 E6 (polar bear), PphPV1 E6 (porpoise), and TtPV5 E6 (bottlenose 383 dolphin) (Fig 4A). Interestingly, the three E6 proteins that do not degrade NHERF1 384 cluster together in phylogenetic relatedness (Fig 4B). We utilized transfected human 385 NHERF1 throughout our study, so it is possible that the inability of these three E6 386 proteins to target NHERF1 for degradation may be due to evolutionary divergence in the 387 NHERF1 homologs. Future studies will explore if the lack of degradation of human

388 NHERF1 by UmPV1, PphPV1, and TtPV5 is due to evolutionary divergence of the 389 respective NHERF1 proteins compared to human NHERF1. It would be of interest to 390 determine if NHERF1 is a "universal" target of E6 proteins that act through association 391 with E6AP. 392 Discovery of NHERF1 as a novel target for not only high and low-risk mucosal 393 and cutaneous HPV E6 as well as a wide range of E6 proteins across divergent host 394 species indicates a significant and previously undescribed role for NHERF1 in 395 papillomavirus biology. That NHERF1 is a conserved target of papillomavirus E6 396 proteins further elevates the importance of NHERF1 as a cell growth regulator. Finally, 397 the identification of this highly conserved E6 degradation target may represent a novel 398 avenue for therapeutic intervention against both low and high-risk HPV. 399 Materials and methods 400 401 Cells and cell culture 402 E6AP-null 8B9 cells (a gift of Dr. Lawrence banks, ICGEB, Italy) [59] and HPV-negative 403 C33A cervical cancer cells (ATCC) were maintained and transfected using 404 polyethylenimine (PEI) as previously described [58]. Normal immortalized keratinocytes 405 (NIKS, obtained from ATCC) are spontaneously immortalized foreskin keratinocytes [60] 406 that were cocultured with mitomycin C-treated 3T3 feeder cells in F medium as 407 described previously [61]. NIKS were retrovirally transduced with replication-defective 408 murine retroviruses based on pLXSN [62] as previously described [25]. Retrovirally 409 transduced NIKS cells were counted and seeded at equal confluency in each 410 experiment.

411 Plasmids

- 412 Epitope tagged E6AP, GFP, E6, and NHERF1 were all transiently expressed from the
- 413 pcDNA3 plasmid. HA-tagged NHERF1 originated from Vijaya Ramesh's laboratory
- 414 (from Addgene, plasmid 11635). 16E6 point mutants were created using QuikChange
- 415 primer design (Agilent Technologies). NHERF1 truncations were PCR generated and

416 sequenced.

417 Antibodies and Western blots

- 418 12 well plates of transfected mammalian cells were lysed in 0.5X IPEGAL as described
- 419 previously [7]. Transduced NIKS were lysed in 1% SDS, 5mM EDTA, and 1 mM sodium
- 420 vanadate and equilibrated for protein content (Biorad assay kit). All lysates were
- 421 resolved by SDS-PAGE electrophoresis and transferred to PVDF membranes.
- 422 Antibodies: anti-HA (Bethyl Laboratories, Inc.), anti-FLAG M2 (Sigma), anti-p53 Ab-8
- 423 (ThermoFisher Scientific), anti-16E6 6G6 (a generous gift from Arbor Vita Corporation),
- 424 anti-SLC9A3R1 (Sigma), anti-GAPDH (Cell Signaling Technology), and anti-MYC 9B11
- 425 (Cell Signaling Technology).

426 **RT-PCR**

- 427 Retrovirally transduced NIKS were plated at different cell densities and harvested
- 428 following a TRIzol RNA harvest protocol (Invitrogen). cDNA was generated using
- random hexamers. Quantitative real-time PCR was performed on the cDNA using iQ[™]
- 430 SYBR® Green Supermix (BioRad #1708880). The primers targeted the SLC9A3R1
- 431 gene (BioRad Assay ID: qHsaCEP0050521) and the GAPDH gene (BioRad Assay ID:
- 432 qHsaCEP0041396). Relative values were analyzed using the $\Delta\Delta C_T$ method (where C_T
- 433 is the threshold cycle) and GAPDH as a control.

434 Wnt/β-catenin luciferase reporter assay

- 435 C33A cells plated at 70% confluency were transiently transfected with DNA of the
- 436 TOPFLASH or control FOPFLASH (containing mutated TCF/β-catenin binding sites; 1
- 437 ug) plasmid, Renilla luciferase (0.005 ug) plasmid (used to evaluate transfection
- 438 efficiency), FLAG_E6AP_WT (0.35 ug) plasmid, and the indicated E6 plasmids (0.3 ug).
- 439 18 hrs post-transfection, media was removed and Wnt3A conditioned media was added
- for 8.5 hours to stimulate the Wnt pathway. Luciferase levels were measured using the
- 441 Dual-Luciferase® Reporter Assay System (Promega) and a Cytation1 Plate Reader
- 442 (software version 3.04.17). FOPFLASH luciferase readings were low, and were
- subtracted from the paired TOPFLASH readouts. 10% fetal bovine serum Wnt3A
- 444 conditioned media was generated using L Wnt-3A murine fibroblasts (ATCC, CRL-
- 445 2647) as previously described [63].

446 **Phylogenetic analysis**

- 447 Multiple protein sequence files were downloaded from the Papillomavirus Episteme [64]
- 448 and aligned using the EMBL-EBI MUSCLE (<u>MU</u>Itiple <u>Sequence</u> <u>Comparison by Log-</u>
- 449 <u>Expectation</u>) program [65]. The phylogenetic tree was generated as a neighbour-joining
- 450 tree without distance corrections within the MUSCLE program [65].

451 Yeast expression

- 452 Modified LexA-based yeast three-hybrid assays were performed as previously
- 453 described [58].

454

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- 458 of the manuscript.
- 459

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- 644

645 Figure legends

Fig 1. NHERF1 protein levels are reduced by both high and low-risk E6 proteins.

- (A) NHERF1 protein levels are reduced in an E6 and E6AP dependent manner.
- Plasmids encoding the indicated FLAG_E6AP (1 ug), HA_NHERF1 (0.5 ug), human
- 649 p53 (0.5 ug), HA_GFP (0.08 ug), and the listed E6 proteins (1 ug) were transiently
- transfected into E6AP-null 8B9 cells and HA-NHERF1 expression was analyzed by
- 651 western blot. Reduction of NHERF1 protein levels by high or low-risk E6 requires ligase
- active E6AP (E6AP_WT) but does not require the E6 PDZ binding motif (PBM).
- 653 FLAG_18E6* is a truncated splice isoform of 18E6. E6AP_Ub⁻ denotes an E6AP
- 654 mutant defective for ubiquitin ligase activity. Quantitation is the result of three

655	independent experiments (N=3) where NHERF1 levels are normalized to co-transfected
656	HA_GFP. Shown is a single representative blot. Vertical black line in blots represents
657	removal of an irrelevant sample. The means of triplicate independent experiments \pm
658	standard error are shown. N=3. *<0.05, **<0.01 by Student's t-test. (B) Reduction of
659	NHERF1 protein is not an overexpression artifact. Titrations of the indicated E6 proteins
660	were co-transfected with FLAG_E6AP_WT (1 ug), HA_GFP (0.02 ug), and either
661	HA_NHERF1 (0.5 ug) or p53 (0.5 ug) in murine 8B9 cells. With increased E6
662	expression, NHERF1 decreased for each E6 protein parallel with p53. As expected, p53
663	degradation was observed for the high-risk 16E6 proteins (both WT and ΔPBM) but not
664	by low-risk 11E6 protein despite reduction of NHERF1 protein levels by 11E6. The
665	means of triplicate independent experiments \pm standard error are shown.
666	
666 667	Fig 2. NHERF1 protein levels increase with increased cell density. (A) Protein
	Fig 2. NHERF1 protein levels increase with increased cell density. (A) Protein levels of endogenous NHERF1 increase with cell confluency. Keratinocytes retrovirally
667	
667 668	levels of endogenous NHERF1 increase with cell confluency. Keratinocytes retrovirally
667 668 669	levels of endogenous NHERF1 increase with cell confluency. Keratinocytes retrovirally transduced with either vector or 16E6_WT were counted and plated at the indicated cell
667 668 669 670	levels of endogenous NHERF1 increase with cell confluency. Keratinocytes retrovirally transduced with either vector or 16E6_WT were counted and plated at the indicated cell densities. As confluency increased, NHERF1 protein levels also increased, though still
667 668 669 670 671	levels of endogenous NHERF1 increase with cell confluency. Keratinocytes retrovirally transduced with either vector or 16E6_WT were counted and plated at the indicated cell densities. As confluency increased, NHERF1 protein levels also increased, though still reduced in the presence of 16E6_WT. The means of triplicate independent experiments
667 668 669 670 671 672	levels of endogenous NHERF1 increase with cell confluency. Keratinocytes retrovirally transduced with either vector or 16E6_WT were counted and plated at the indicated cell densities. As confluency increased, NHERF1 protein levels also increased, though still reduced in the presence of 16E6_WT. The means of triplicate independent experiments \pm standard error are shown. (B) NHERF1 RNA levels are not changed by cell
667 668 669 670 671 672 673	levels of endogenous NHERF1 increase with cell confluency. Keratinocytes retrovirally transduced with either vector or 16E6_WT were counted and plated at the indicated cell densities. As confluency increased, NHERF1 protein levels also increased, though still reduced in the presence of 16E6_WT. The means of triplicate independent experiments \pm standard error are shown. (B) NHERF1 RNA levels are not changed by cell confluency or by the presence of 16E6_WT. Total RNA was extracted from
667 668 670 671 672 673 674	levels of endogenous NHERF1 increase with cell confluency. Keratinocytes retrovirally transduced with either vector or 16E6_WT were counted and plated at the indicated cell densities. As confluency increased, NHERF1 protein levels also increased, though still reduced in the presence of 16E6_WT. The means of triplicate independent experiments \pm standard error are shown. (B) NHERF1 RNA levels are not changed by cell confluency or by the presence of 16E6_WT. Total RNA was extracted from keratinocytes retrovirally transduced with either vector or 16E6_WT and plated at the

678

679	Fig 3. Degradation of NHERF1 by 16E6 requires proteasome function.
680	Keratinocytes retrovirally transduced with either vector or 16E6_WT were seeded at
681	equal confluency. Cells were treated with DMSO, mitomycin C (MMC), or the
682	proteasome inhibitor MG132 at varying concentrations for 8 hours as indicated. MG132
683	significantly rescued NHERF1 protein levels in a dose dependent manner. MMC
684	treatment was used to induce p53 levels, which were observed as a positive control.
685	Quantification was normalized to vector-transduced cells treated with DMSO. The
686	means of triplicate independent experiments \pm standard error are shown. N=3, *<0.05,
687	**<0.01, ***<0.001, n.s. = no significance by Student's t-test for samples compared to
688	untreated 16E6 keratinocytes (lane 3).
689	
689 690	Fig 4. E6 proteins from evolutionarily diverse species target NHERF1. (A) E6
	Fig 4. E6 proteins from evolutionarily diverse species target NHERF1 . (A) E6 proteins from divergent animal species degrade NHERF1 via E6AP. HA_NHERF1 (0.4
690	
690 691	proteins from divergent animal species degrade NHERF1 via E6AP. HA_NHERF1 (0.4
690 691 692	proteins from divergent animal species degrade NHERF1 via E6AP. HA_NHERF1 (0.4 ug), HA_GFP (0.1 ug), FLAG_E6AP_WT (0.35 ug), and the indicated FLAG_E6 (0.3 ug)
690 691 692 693	proteins from divergent animal species degrade NHERF1 via E6AP. HA_NHERF1 (0.4 ug), HA_GFP (0.1 ug), FLAG_E6AP_WT (0.35 ug), and the indicated FLAG_E6 (0.3 ug) plasmids were co-transfected into C33 cells. E6 proteins are classified based on their
690 691 692 693 694	proteins from divergent animal species degrade NHERF1 via E6AP. HA_NHERF1 (0.4 ug), HA_GFP (0.1 ug), FLAG_E6AP_WT (0.35 ug), and the indicated FLAG_E6 (0.3 ug) plasmids were co-transfected into C33 cells. E6 proteins are classified based on their known preference for binding E6AP or MAML as indicated. NHERF1 was degraded by
690 691 692 693 694 695	proteins from divergent animal species degrade NHERF1 via E6AP. HA_NHERF1 (0.4 ug), HA_GFP (0.1 ug), FLAG_E6AP_WT (0.35 ug), and the indicated FLAG_E6 (0.3 ug) plasmids were co-transfected into C33 cells. E6 proteins are classified based on their known preference for binding E6AP or MAML as indicated. NHERF1 was degraded by E6 proteins isolated from numerous different mammalian species. Many, but not all, of
690 691 692 693 694 695 696	proteins from divergent animal species degrade NHERF1 via E6AP. HA_NHERF1 (0.4 ug), HA_GFP (0.1 ug), FLAG_E6AP_WT (0.35 ug), and the indicated FLAG_E6 (0.3 ug) plasmids were co-transfected into C33 cells. E6 proteins are classified based on their known preference for binding E6AP or MAML as indicated. NHERF1 was degraded by E6 proteins isolated from numerous different mammalian species. Many, but not all, of the E6 proteins that bind E6AP targeted NHERF1 for degradation, while E6 proteins

standard error are shown. N=5, **<0.01, ***<0.001 by Student's t-test. (B) E6 proteins

701 that degrade NHERF1 cluster phylogenetically. The E6 proteins from the listed 702 papillomaviruses were subjected to a multiple sequence alignment and then clustered 703 phylogenetically using the program MUSCLE [65]. For E6 physical association, blue 704 denotes MAML1 and light purple denotes E6AP. The preferential association of three 705 E6 proteins is unknown. Ability to degrade NHERF1 is denoted in green and lack of 706 ability to degrade NHERF1 is indicated by red. Interestingly, E6 proteins that can bind 707 E6AP but not degrade NHERF1 cluster differently from other E6 proteins that cannot 708 degrade NHERF1. The genera of each papillomavirus is listed. Western blot indicating 709 NHERF1 expression in the presence of HPV1 E6, HPV8 E6, and SfPV1 E6 is shown in 710 S3 Fig. H = Homo sapiens (human), Mm = Macaca mulata (rhesus monkey), Ss = Sus 711 scrofa (wild boar), Pph = Phocoena phocoena (harbor porpoise), Um = Ursus maritimus 712 (polar bear), Tt = Tursiops truncatus (bottlenose dolphin), Oc = Oryctolagus cuniculus 713 (rabbit), Mc = Mastomys coucha (mouse), Ma = Mesocricetus auratus (golden hamster), 714 Sf = Sylvilagus floridanus (Cottontail rabbit; CRPV1). Caption credit: Brimer N, Drews 715 CM, Vande Pol SB. Association of papillomavirus E6 proteins with either MAML1 or 716 E6AP clusters E6 proteins by structure, function, and evolutionary relatedness. PLoS 717 Pathog. 2017;13(12):e1006781.

718

Fig 5. 16E6 mutagenesis screen identified mutants selectively defective in their
ability to degrade NHERF1. (A) Amino acids F69 and K72 are important for
degradation of NHERF1 by 16E6. Plasmids encoding untagged 16E6_WT or 16E6
mutants (0.3 ug) were co-transfected with FLAG_E6AP (0.35 ug), HA_NHERF1 (0.4
ug), MYC_p53 (0.25 ug), and HA_GFP (0.08 ug) into C33 cells and HA_NHERF1 levels

724 determined by western blot. Multiple 16E6 proteins were identified that were unable to 725 degrade NHERF1 but were still capable of degrading p53. (**B**) HA_NHERF1 and (**C**) 726 p53 protein levels were quantified and normalized to co-transfected HA GFP as an 727 internal transfection control. 728 Fig 6. Amino acid side chains F69 and K72 define a novel substrate interaction 729 730 domain on 16E6. (A) HPV16 E6 structure (PDB file 4GIZ) showing the amino-terminal 731 zinc-structured domain in green, connecting alpha helix in yellow, and the carboxy-732 terminal zinc-structured domain in blue. The E6 protein is complexed with the LXXLL peptide of E6AP (pictured in light pink). (B) The E6 protein depicted in A is rotated 45° 733 734 clockwise (C.W.) and the F69 and K72 residues and their side chains are highlighted in 735 red. (C) A similar view as part B is shown complexed with the core p53 DNA binding 736 domain (grey). The E6 interaction face with p53 is opposite the F69 and K72 residues. 737 738 Fig 7. NHERF1 degradation by E6 proteins from both high and low-risk 739 papillomaviruses in stable keratinocytes. Keratinocytes retrovirally transduced with 740 the indicated E6 proteins were seeded at equal confluency and endogenous NHERF1

protein levels were normalized to GAPDH. 16E6_WT, 16E6 deleted of its PBM (Δ PBM),

and 11E6_WT all degraded NHERF1. The 16E6_F69A/K72A double mutant did not

target NHERF1 for degradation. The means of triplicate independent experiments \pm

standard error and one representative blot are shown. N=3, *<0.05, **<0.01, n.s. = no

significance by Student's t-test.

746

747 Fig 8. NHERF1 truncations identify the EB domain as necessary for NHERF1

748 degradation by 16E6. (A) Schematic of NHERF1 truncations. NHERF1 proteins that

- were successfully degraded by 16E6_WT are depicted in green while truncations that
- were not degraded are depicted in red. (**B** and **C**) NHERF1 truncations containing the
- EB domain were degraded, while those lacking the EB domain were not. The listed
- HA_NHERF1 truncations (shown in A in the order loaded in B and C, 0.8 ug), untagged
- 753 16E6_WT (1 ug), FLAG_GFP (0.08 ug), and either FLAG_E6AP_WT (1.2 ug) or
- FLAG_E6AP_Ub⁻ (1.2 ug, defective for ubiquitin ligase activity) were co-transfected in
- 755 E6AP-null 8B9 cells. HA_NHERF1 levels were quantified and normalized to FLAG_GFP
- as an internal transfection control. The bar graph below the blot represents
- 757 quantification of each listed HA_NHERF1 truncation. In panel C, the WT NHERF1 in
- Ianes 2-4 contains an amino terminal 1X HA tag while the WT NHERF1 in lanes 17 and
- 18 contains an amino terminal 2X HA tag. All of the NHERF1 truncations contain amino
- terminal 2X HA tags. Levels of HA_NHERF1 truncations in the presence of
- 761 FLAG_E6AP_WT were normalized to their corresponding expression in the presence of
- FLAG_E6AP_Ub⁻ to account for the differing expression levels. UT = untransfected.
- 763

Fig 9. The E6-E6AP-NHERF1 complex can be modeled in yeast. Yeast three-hybrid
plasmids expressing the LexA DNA binding domain fused to either 16E6_WT or
E6AP_Ub⁻ were co-expressed in yeast (bait) together with either vector, 16E6_WT, or
16E6_F69A/K72A as indicated. The bait yeast were mated to prey yeast expressing
Gal4 activation domain (G4), or G4 fused to 16E6_WT, PTPN3, truncations of NHERF1,
or native p53 and diploids selected. Positive controls for 16E6 expression included the

770	established interaction of the 16E6 PBM with the PDZ domain of tyrosine phosphatase
771	PTPN3 and 16E6-E6AP complex interaction with p53. 16E6_WT recruited NHERF1,
772	p53, and PTPN3 to LexA_E6AP_Ub [_] . The recruitment of NHERF1 to LexA_E6AP_Ub [_]
773	by 16E6 was specifically lost upon mutation of residues F69 and K72, however, p53 and
774	PTPN3 recruitment were maintained. 16E6_WT recruitment of NHERF1 was not seen
775	with an NHERF1 truncation lacking the EB domain (G4_NHERF1 121-297). Caption
776	credit: Ansari T, Brimer N, Vande Pol SB. Peptide interactions stabilize and restructure
777	human papillomavirus type 16 E6 to interact with p53. J Virol. 2012;86(20):11386-91.
778	
779	Fig 10. Activation of the canonical Wnt/ β -catenin pathway is augmented by E6
780	proteins that can degrade NHERF1. The listed E6 proteins were co-expressed with
781	FLAG_E6AP_WT, the TOPFLASH or FOPFLASH luciferase reporter, and a renilla
782	luciferase internal transfection control plasmid in C33A cells. Transfected cells were
783	treated with Wnt3A conditioned media for 8.5 hours, lysed in 1X passive lysis buffer
784	(Promega), and measured for luciferase and renilla luminescence. Fold activation was
785	determined by normalizing the TOPFLASH luminescence by the FOPFLASH
786	luminescence. Each E6 protein that could degrade NHERF1 (16E6_WT, 16E6_ Δ PBM,
787	11E6_WT, and 18E6_WT) augmented the canonical Wnt pathway. 16E6_F69A/K72A,
788	which cannot degrade NHERF1, failed to increase Wnt pathway activation over vector
789	levels. Statistical significance was determined from three independent experiments by
790	Student's t-test (***<0.001, n.s. = no significance).

793 Supporting information legends

794 S1 Fig. Reduction of NHERF1 protein levels is not an overexpression artifact.

- Titrations of the indicated three different E6 proteins (16E6_WT, 16E6_ Δ PBM, and
- 11E6_WT) were co-transfected with FLAG_E6AP_WT (1 ug), HA_GFP (0.02 ug), and
- reither HA_NHERF1 (0.5 ug) or p53 (0.5 ug) in E6AP-null 8B9 cells. A representative
- blot of the triplicate experiments for each E6 protein is shown. Increased E6 expression
- for 16E6_WT, 16E6∆PBM, and 11E6_WT resulted in decreased NHERF1 protein
- levels. Both 16E6_WT and 16E6 Δ PBM degrade p53 with increasing E6 expression.
- 801 Overexpression of 11E6 _WT (>0.1 ug E6) resulted in degradation of co-expressed
- 802 E6AP_WT.

803

804 S2 Fig. 16E6 point mutants screened to determine amino acid(s) necessary for

805 NHERF1 degradation. The 16E6 crystal structure (PDB file 4GIZ) was examined for

residues that were at least 20% exposed as determined by the Swiss PDB Viewer.

807 Point mutants of these identified amino acids were then screened to identify which

808 residue(s) resulted in an E6 protein that was selectively defective for degrading

809 NHERF1 but retained degradation of p53. Residues of interest are indicated in red.

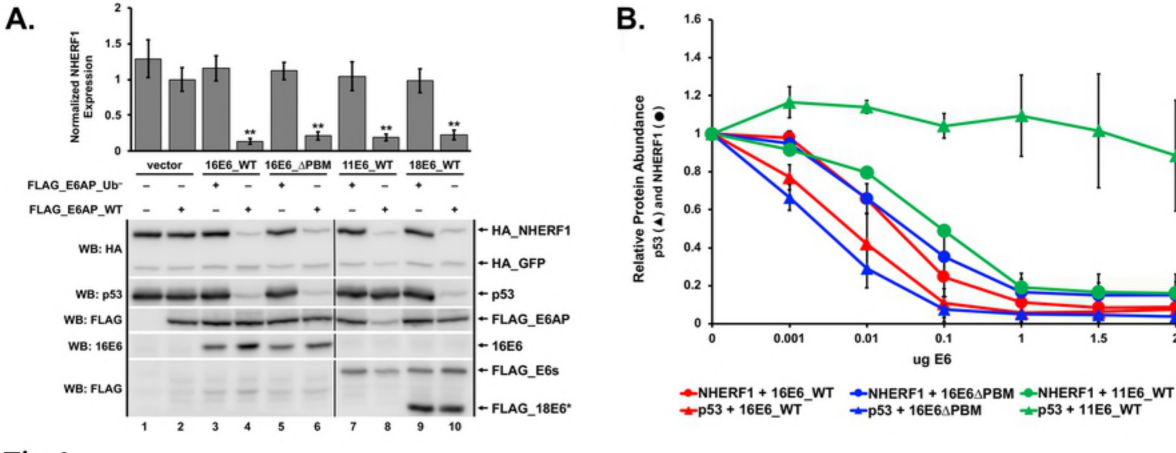
810

811 S3 Fig. The E7 papillomavirus protein does not induce phospho-specific isoforms

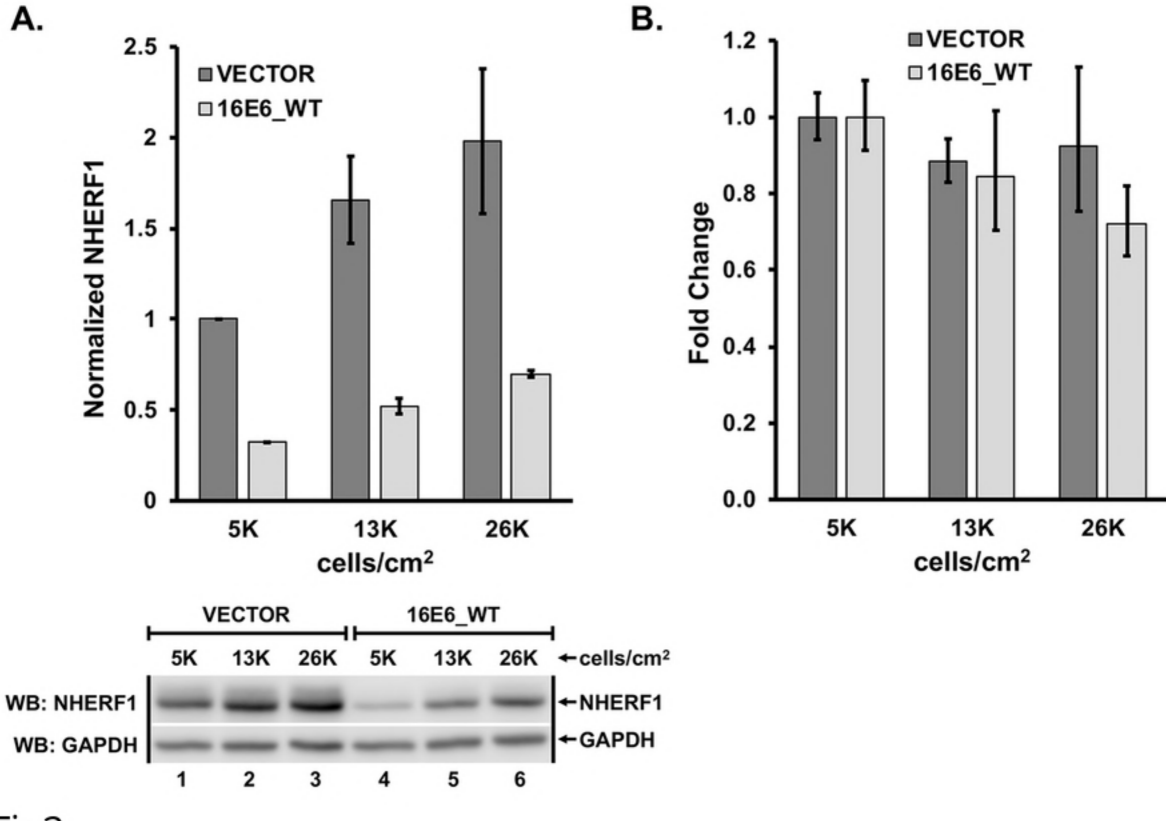
of NHERF1. Keratinocytes retrovirally transduced with vector or the indicated E7 and/or

- E6 proteins were seeded at equal confluency. Levels of endogenous NHERF1 were
- 814 determined by western blot. Levels of phosphorylated NHERF1 (pNHERF1) were
- 815 unchanged in keratinocytes expressing empty vector compared to the various E7

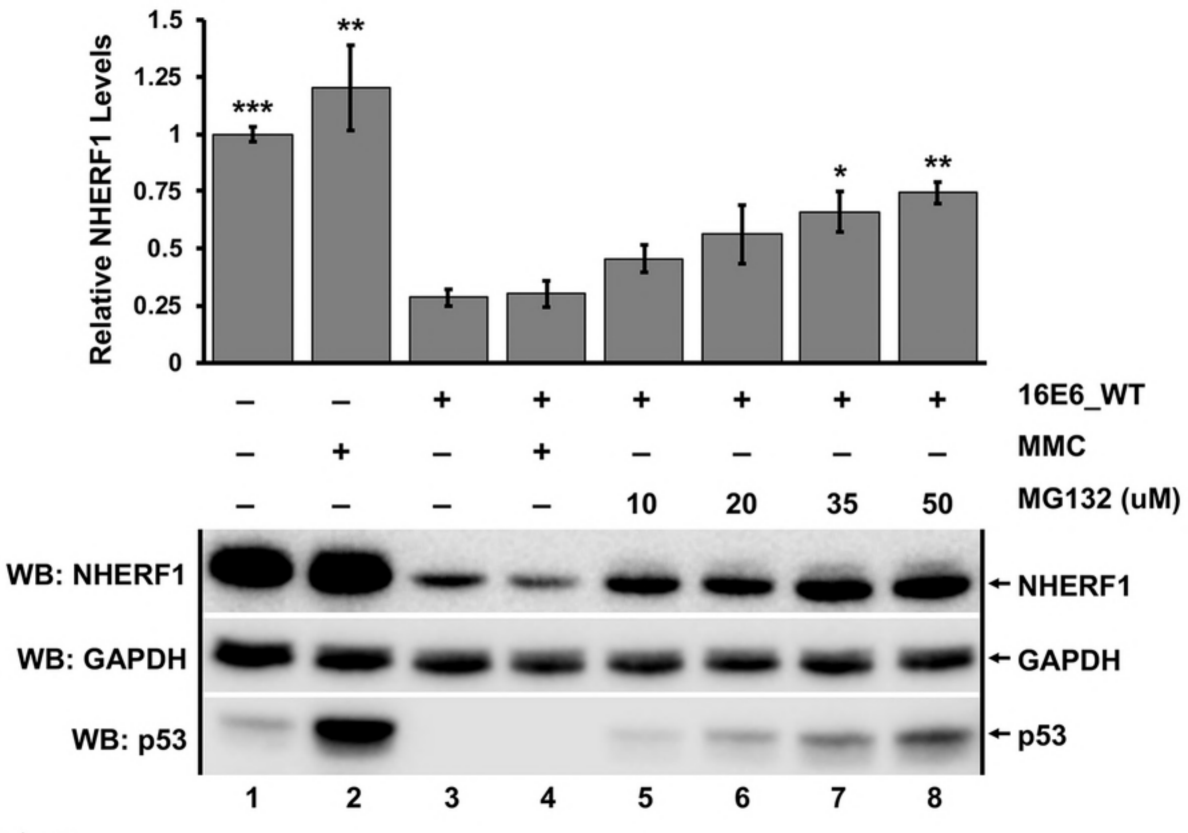
816	proteins. Keratinocytes expressing the E6 protein from high-risk (HPV16) and low-risk
817	(HPV11) degraded NHERF1. H = Homo sapiens (human), Sf = Sylvilagus floridanus
818	(Cottontail rabbit; CRPV1). Caption credit: Accardi R, Rubino R, Scalise M, Gheit T,
819	Shahzad N, Thomas M, et al. E6 and E7 from human papillomavirus type 16 cooperate
820	to target the PDZ protein Na/H exchange regulatory factor 1. J Virol. 2011;85(16):8208-
821	16.
822	
823	S4 Fig. The presence of the E7 oncoprotein does not enhance NHERF1
824	degradation by E6 proteins. C33A cells were co-transfected with the following
825	plasmids: HA_NHERF1 (0.4 ug), FLAG_E6AP_WT (0.35 ug), HA_GFP (0.08 ug), the
826	indicated E6 protein (0.3 ug), and the indicated E7 protein (0.3 ug). HA_NHERF1 levels
827	were determined by western blot. FLAG_18E6* is a truncated splice isoform of 18E6.
828	Quantitation is derived from three experimental replicates. A representative blot and
829	means of triplicate independent experiments \pm standard error are shown. N=3. **<0.01,

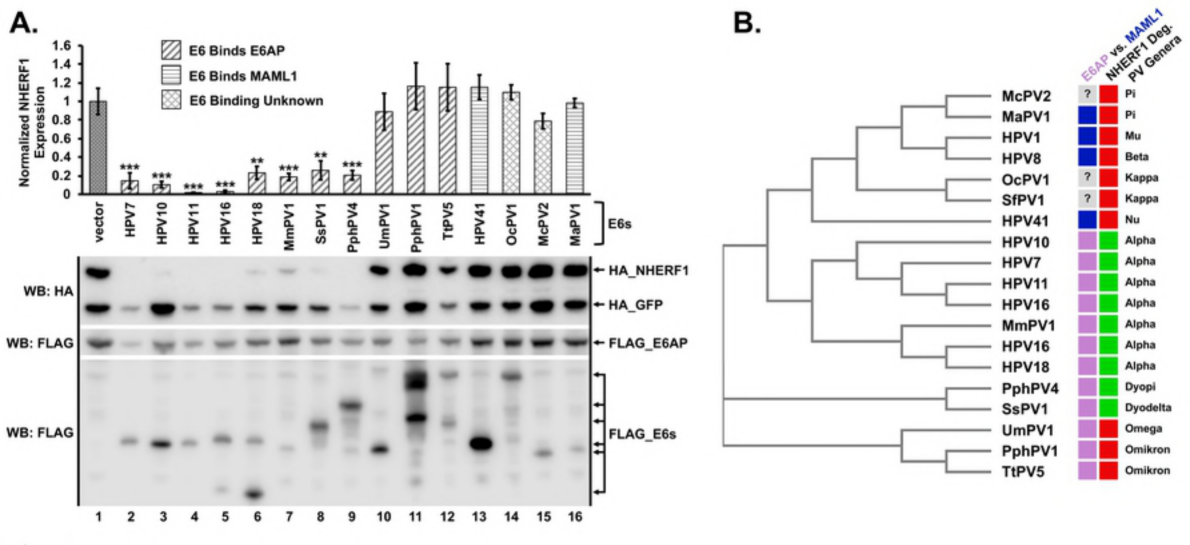


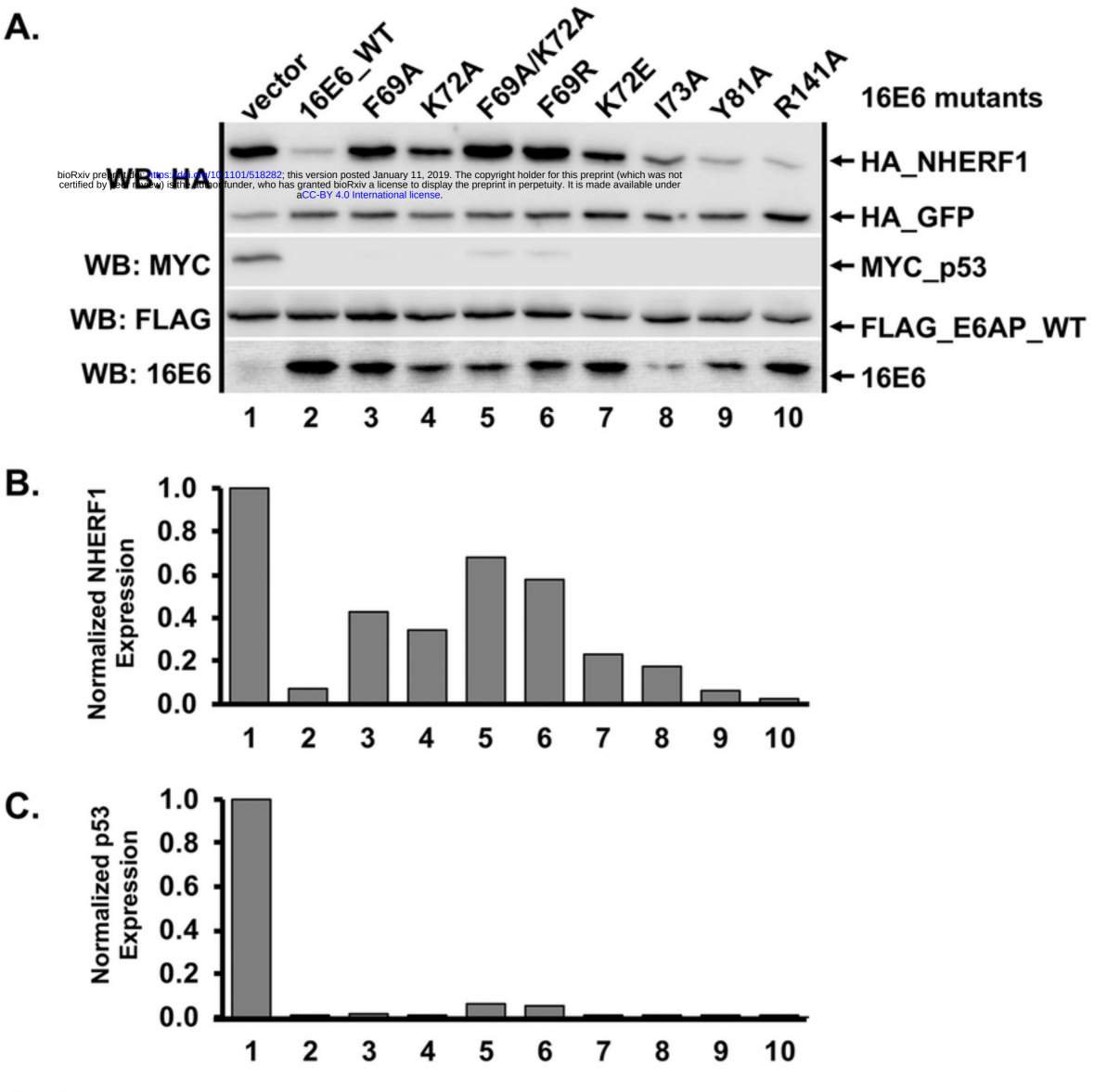
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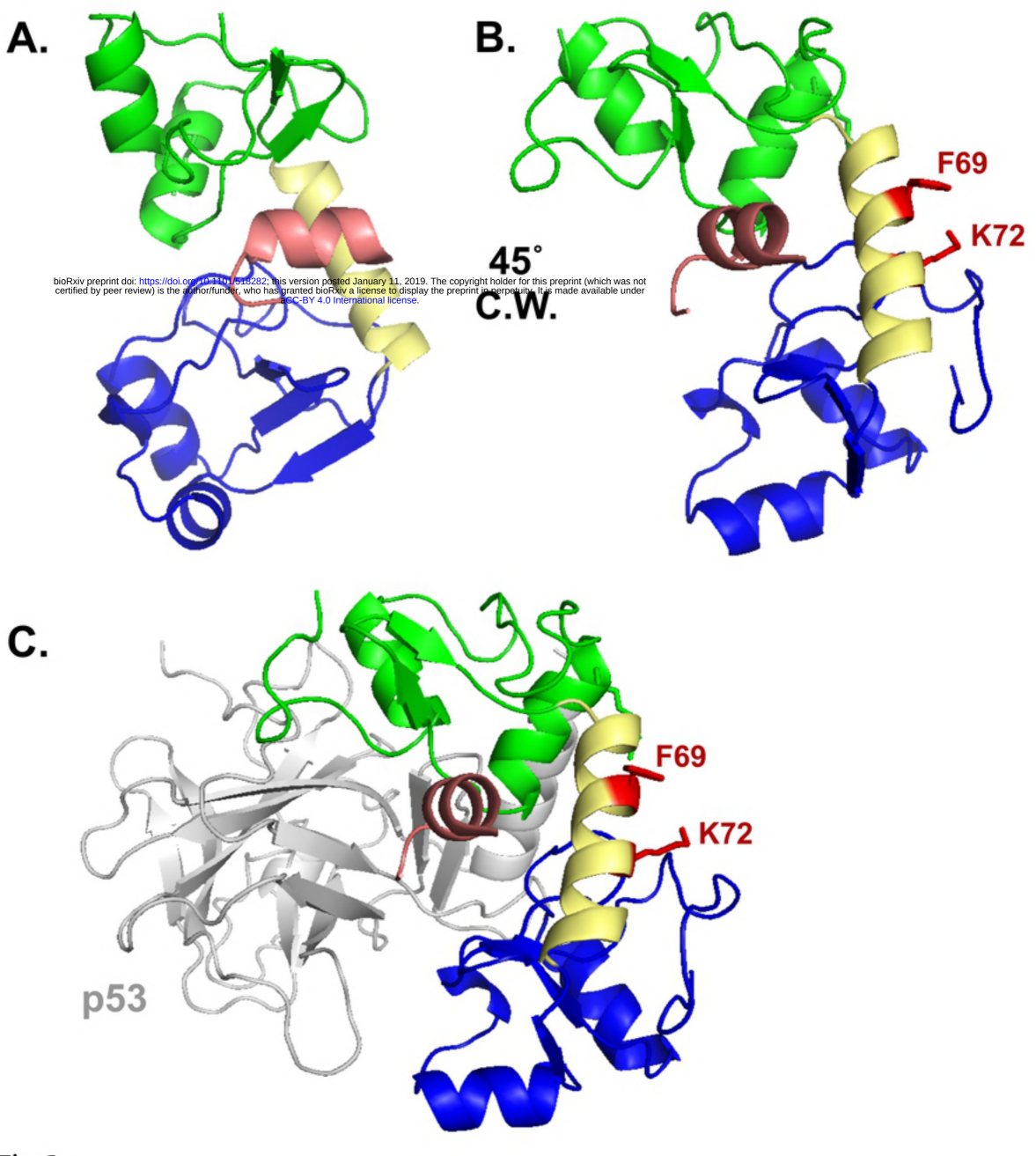


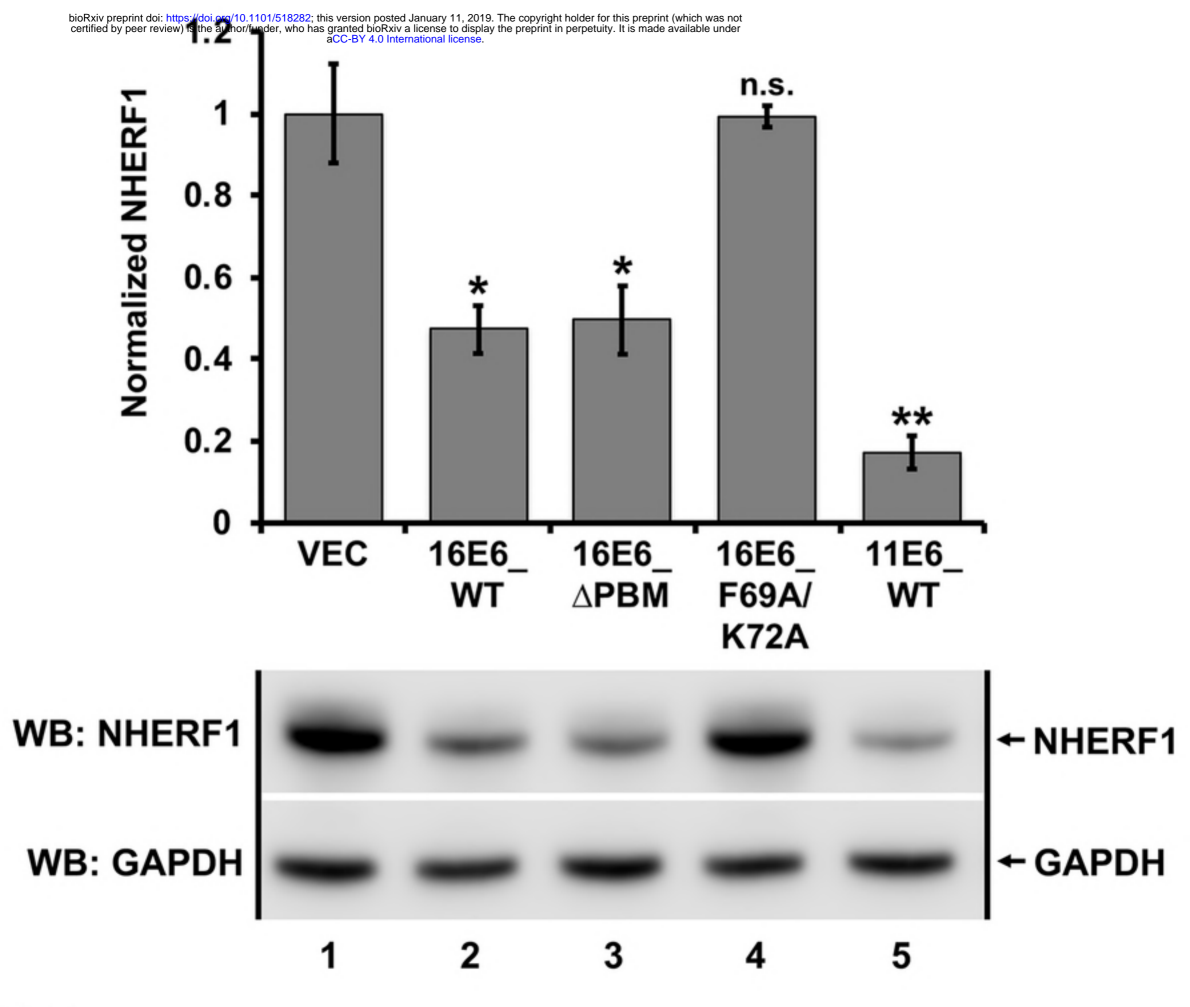
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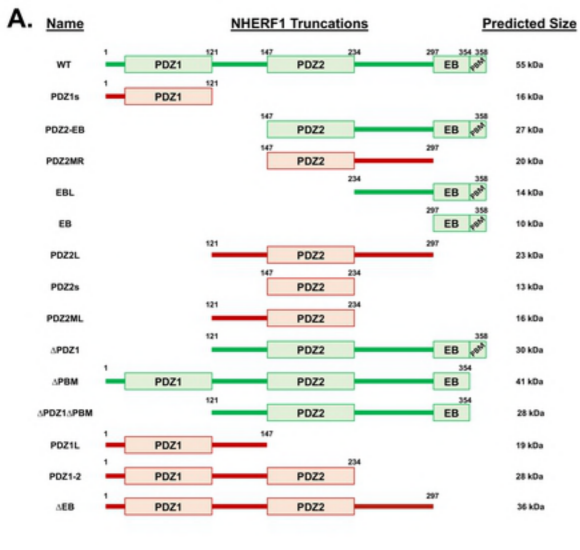


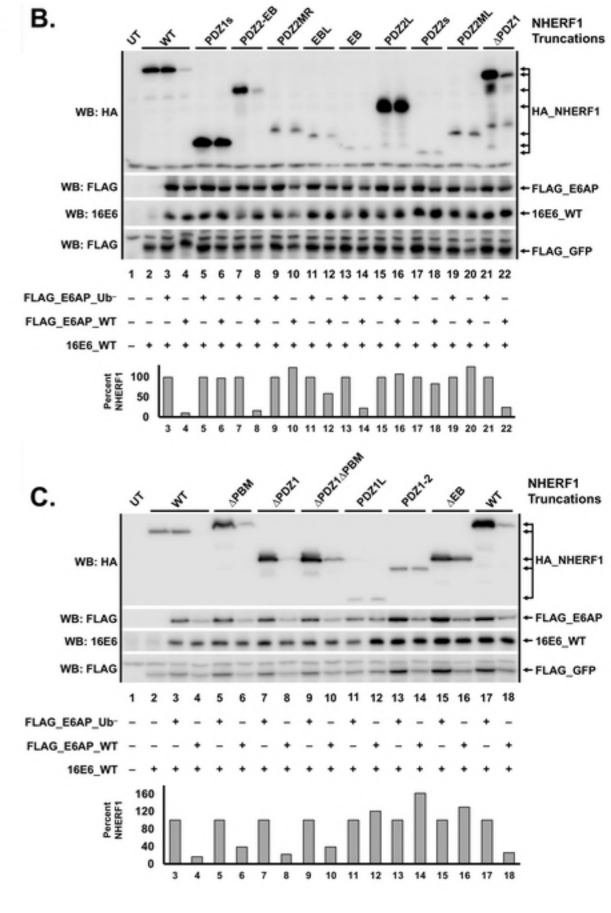


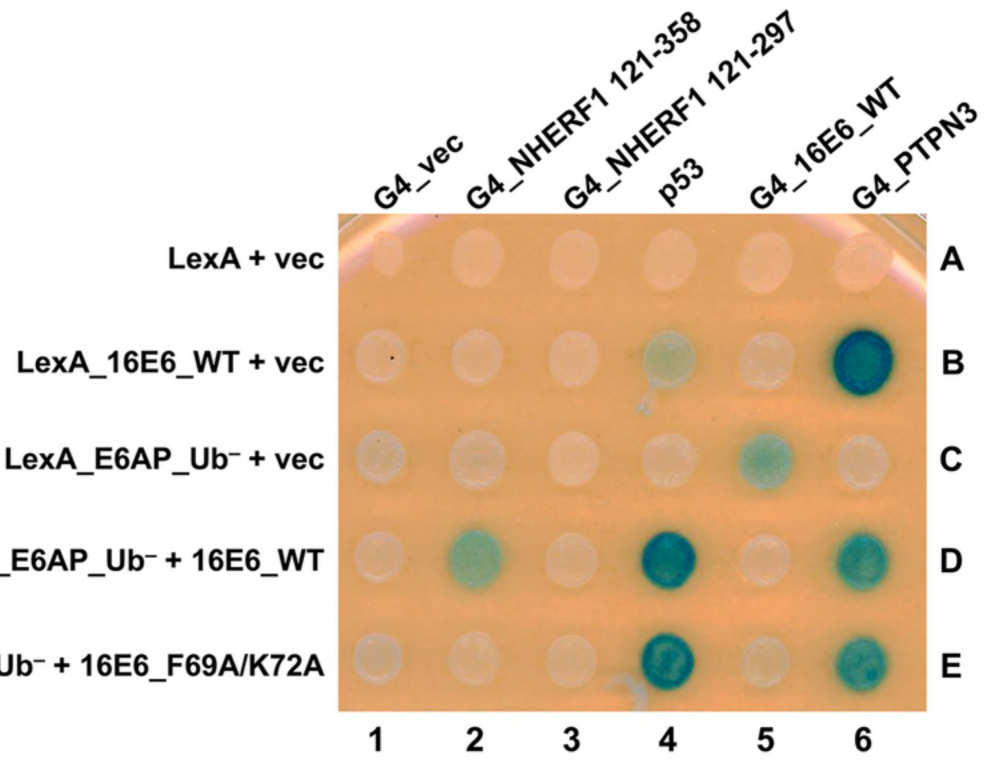












LexA + vec

LexA_16E6_WT + vec

LexA_E6AP_Ub⁻ + 16E6_WT

LexA_E6AP_Ub⁻ + 16E6_F69A/K72A

