

HPV type 16 E6 and NFX1–123 augment JNK signaling to mediate keratinocyte differentiation and L1 expression

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ABSTRACT

The HPV life cycle is differentiation-dependent, with cellular differentiation driving initiation of the late, productive stage of the viral life cycle. Here, we identify a role for the protein NFX1–123 in regulating keratinocyte differentiation and events of the late HPV life cycle. NFX1–123 itself increased with differentiation of epithelial cells. Greater NFX1–123 augmented differentiation marker expression and JNK phosphorylation in differentiating 16E6-expressing human foreskin keratinocytes (16E6 HFks). This was associated with altered expression of MKK4 and MKK7, upstream kinase regulators of JNK phosphorylation. Modulating levels of NFX1–123 in HPV16-positive W12E cells recapitulated the effects on differentiation markers, JNK phosphorylation, and MKK4/7 seen in 16E6 HFks. Crucially, levels of NFX1–123 also correlated with expression of L1, the capsid protein of HPV. Altogether, these studies define a role for NFX1–123 in mediating epithelial differentiation through the JNK signaling pathway, potentially linking expression of cellular genes and HPV genes during differentiation.

1. Introduction

The human papillomaviruses (HPVs) are small, non-enveloped, double-stranded DNA viruses that infect keratinocytes of cutaneous and mucosal stratified squamous epithelium (Doorbar et al., 2012; zur Hausen, 2002). There are over 200 types of HPV, and those that specifically target mucosal epithelium are further categorized as low-risk HPV (LR HPV) or high-risk HPV (HR HPV) based on their epidemiologic association with cancer (Bosch et al., 2002; Muñoz et al., 2003; zur Hausen, 2009). HR HPV cause nearly all cervical cancer, the fourth most common cancer in women, in addition to other anogenital and oropharyngeal cancers (Bosch et al., 2002; Stratton and Culkin, 2016; Kreimer et al., 2005; Chaturvedi, 2010; Forman et al., 2012). Altogether, cancers caused by HR HPV account for approximately 5% of the global burden of cancer (de Martel et al., 2012).

HPV has a unique life cycle that is tied to the differentiation programming of its target cell, the keratinocyte (Doorbar et al., 2015; Lee and Laimins, 2007). The viral life cycle can be divided into two stages: in the first, HPV infects keratinocytes in the basal layer of stratified squamous epithelium and maintains its genome as an episome at a low

copy number (Doorbar et al., 2012, 2015; Lee and Laimins, 2007; Doorbar, 2005; Roden and Stern, 2018). In this stage, the early genes (E1, E2, E4, E5, and oncogenes E6 and E7) are expressed from the early viral promoter (Doorbar, 2005; Graham, 2010; Schiffman et al., 2016). The second stage occurs once an infected cell rises through the epithelial suprabasal layers and begins to differentiate. This triggers the late stage of the viral life cycle, whose events include vegetative genome amplification, expression of the capsid proteins L1 and L2 from a viral late promoter, and ultimately, complete virion formation (Doorbar et al., 2012; Graham, 2010, 2017). As a keratinocyte differentiates and moves up through the layers of stratified squamous epithelium, it expresses a successive series of proteins that commonly serve as differentiation markers. These include Keratins 1 and 10, Involucrin, Loricrin, and Filaggrin (Fuchs, 1990; Watt, 1983). Similarly, HPV expresses its genes in a precisely ordered fashion in association with specific epithelial layers. There is thus a tight link between events dictating the expression of cellular genes during differentiation and those dictating expression of HPV genes. Although it is known that terminal differentiation of epithelial cells is required for completion of the viral life cycle, the cellular factors activated during epithelial

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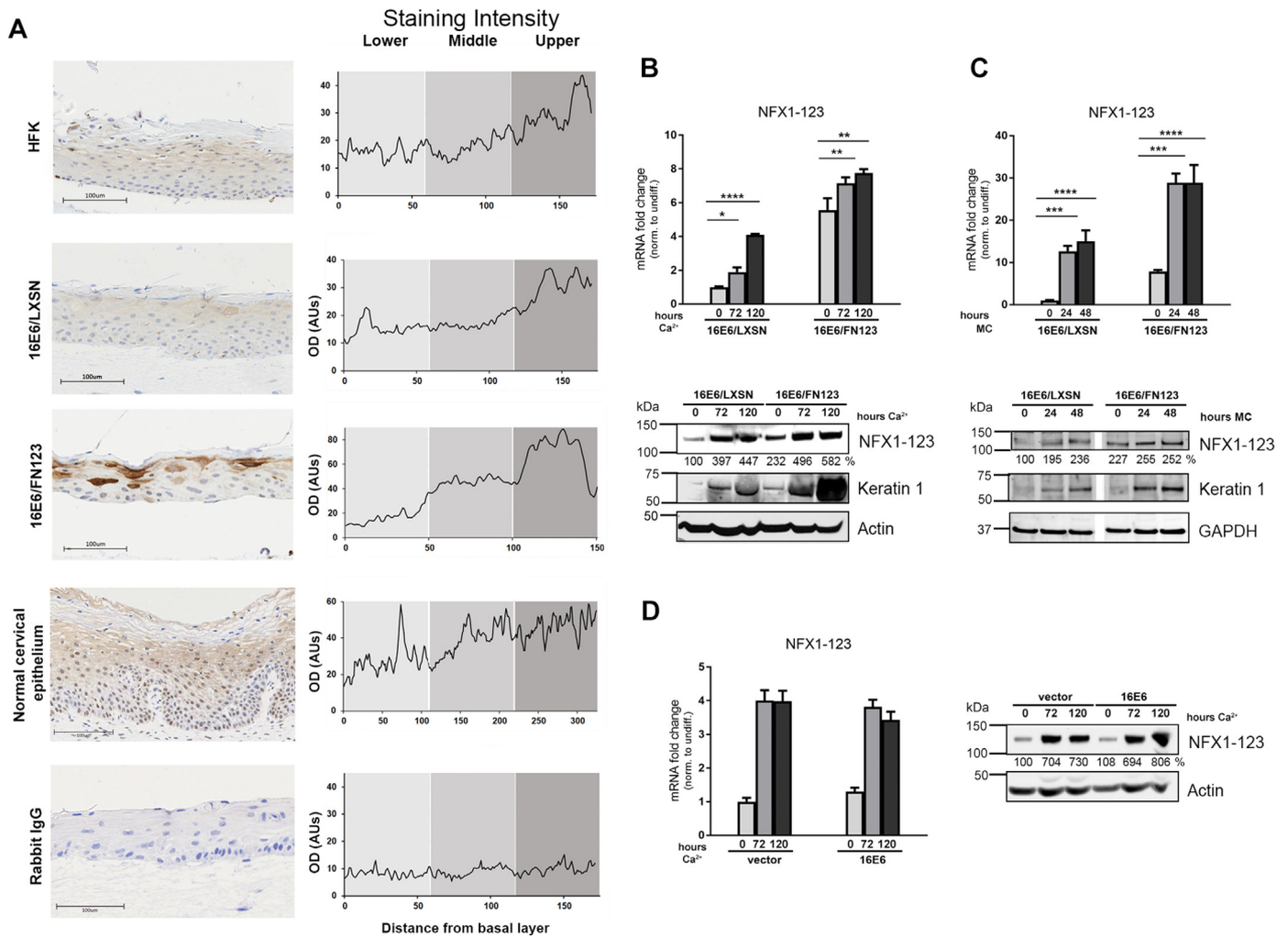


Fig. 1. NFX1–123 expression increases with keratinocyte differentiation. (A) Organotypic raft cultures were grown using HFKs, HFKs serially transduced with 16E6 and vector control (16E6/LXSN), and HFKs transduced with 16E6 and FLAG-tagged NFX1–123 overexpression construct (16E6/FN123). These rafts and normal cervical epithelial biopsies were stained for NFX1–123 protein via immunohistochemistry or stained using a rabbit polyclonal IgG as a control. Staining intensity was measured over a linear distance from the basal layer in ImageJ and the average of four plots over two sections is shown. (B and C) 16E6/LXSN and 16E6/FN123 HFKs were differentiated with 1.8 mM Ca²⁺ treatment for 0, 72, or 120 h (B) or suspension in 1.7% methylcellulose medium (MC) (C) and total mRNA and protein collected. Mean expression of NFX1–123 mRNA was measured by qPCR and compared to 16E6/LXSN 0 h. Protein levels of NFX1–123 were measured by Western blot using actin or GAPDH as a loading control. Induction of the differentiation marker Keratin 1 indicates successful differentiation. White spaces in Western blots indicate removal of empty or irrelevant lanes from original image for clarity. (D) HFKs transduced with empty vector or with 16E6 were differentiated with 1.8 mM Ca²⁺ treatment for 0, 72, or 120 h, and total mRNA and protein collected. Mean expression of NFX1–123 mRNA was measured by qPCR and each sample was compared to HFK/vector 0 h. Protein levels of NFX1–123 were measured by Western blot using actin as a loading control. For B, C, and D, mRNA expression of NFX1–123 for each sample was normalized to the housekeeping gene 36B4, and all error bars represent 95% confidence intervals from replicates. Protein expression of NFX1–123 for each sample was quantified using ImageJ and normalized to the loading control for that sample. One representative experiment is shown from at least three conducted in biologically independent HFK cell lines. Statistical significance was calculated using one-way ANOVA with Bonferroni correction. * p < 0.05, ** p < 0.01, *** p < 0.001, **** p < 0.0001.

differentiation, and how these factors signal to induce the appropriate viral promoters, are not fully defined.

Efforts to understand the regulation of differentiation-dependent HPV gene transcription have implicated a wide variety of transcription factors and cellular processes. These include transcription factors specific to differentiating keratinocytes, CDP and EPOC-1, as well as ubiquitously expressed transcription factors whose levels may change during epithelial cell differentiation such as AP-1 family members, Oct1, C/EBP, YY1, and others (O'Connor et al., 2000; Ai et al., 1999; Yukawa et al., 1996; Chan et al., 1990; Chong et al., 1991; Dong et al., 1994; Hadaschik et al., 2003; Frattini et al., 1997; Gunasekharan et al., 2016; Ruesch et al., 1998). In addition to global changes in levels of transcription factors, mechanisms including shifting ratios of factors (Sp1/Sp3) (Apt et al., 1996), increased binding to HPV promoters during epithelial differentiation (Carson and Khan, 2006), and

activation of cellular signaling pathways (protein kinase C) (Bodily et al., 2006) have been associated with HPV transcription. Furthermore, linker scanning mutational studies have mapped cis regulatory elements upstream of the HPV promoters that may be important during different stages of the life cycle (Sen et al., 2002, 2004).

The understanding of how differentiation-dependent HPV gene transcription is regulated is further complicated by a switch from the early viral promoter to the late viral promoter. The late viral promoter does not become active until the host cell rises to the upper granular and cornified layers of the stratified squamous epithelium. Even after infected cells have begun differentiating, there are additional signals required to activate the viral late promoter. Studies have shown a role for SRp20, a cellular splicing factor, and transcription elongation factors (Jia et al., 2009; Songcock et al., 2017), but many gaps remain in our knowledge of what controls this transition in HPV transcription.

Despite the abundance of transcription factors and complexity of mechanisms shown to be associated with differentiation-dependent HPV transcription, there is still not a complete understanding of the process.

We have previously shown that HR HPV type 16 E6 (16E6) and NFX1–123, a known cellular protein partner of 16E6, collaboratively increase levels of Notch1 (Vliet-Gregg et al., 2013). Notch1 is a master regulator of cellular growth and differentiation (Lefort and Dotto, 2004; Rangarajan et al., 2001). Upon receptor stimulation in these cells, the increased Notch1 expression resulted in an upregulation of its canonical target genes, *Hes1* and *Hes5*, as well as the differentiation markers Keratin 1 and Keratin 10 (Vliet-Gregg et al., 2015). Blocking Notch1 signaling, however, did not abrogate expression of these differentiation markers, suggesting additional Notch1-independent mechanisms by which 16E6 and NFX1–123 are able to affect keratinocyte differentiation.

In the current study, we further explore the role of NFX1–123 in mediating epithelial differentiation. We utilize targeted differentiation stimuli to further delineate these Notch1 independent mechanisms and demonstrate that NFX1–123, a host cellular protein partner of HPV type 16 E6, is able to regulate differentiation marker expression in HFKs and HPV16-positive cells through JNK signaling. Finally, we show that NFX1–123's role in mediating epithelial differentiation is ultimately associated with expression of HPV16 L1, pointing to its potential as a cellular factor connecting epithelial differentiation and the differentiation-dependent HPV life cycle.

2. Results

2.1. Expression levels of NFX1–123 increased with epithelial cell differentiation

NFX1–123 is a known cellular protein partner of HR HPV type 16 E6 (16E6) and is endogenously expressed in epithelial cells (Gewin et al., 2004). Although NFX1–123 is increased in cervical cancer cell lines (Vliet-Gregg et al., 2015), HPV positive cervical precancerous lesions, and HPV-positive cervical cancers (Vliet-Gregg et al., 2019), the overall expression of NFX1–123 in normal epithelial tissue has not been characterized. To do so, we utilized organotypic raft cultures, which approximate the growth and physiology of stratified squamous epithelium, as well as archived normal cervical epithelial biopsies. Raft cultures were grown using human foreskin keratinocytes (HFKs), HFKs expressing 16E6 and endogenous levels of NFX1–123 (16E6/LXSN HFKs), and HFKs expressing 16E6 and overexpressing FLAG-tagged NFX1–123 (16E6/FN123 HFKs). Expression of 16E6 in HFKs was confirmed by immunoblot documenting a reduction of p53 (Supp Figure 1). These rafts and normal cervical epithelial samples were stained for NFX1–123 via immunohistochemistry (Fig. 1A).

Matching previous findings that NFX1–123 has primarily cytoplasmic subcellular localization, there was cytoplasmic staining in cells throughout the raft cultures and cervical epithelium, with some perinuclear staining seen in basal layer cells. This perinuclear staining in basal layer cells was sporadic in the raft cultures, but especially evident in the normal cervical epithelium (Fig. 1A). In normal cervical epithelium, NFX1–123 staining was predominately perinuclear in basal layers and became predominately cytoplasmic in intermediate and upper layers. Expression of 16E6 did not affect expression of NFX1–123 (HFK vs 16E6/LXSN). Rafts grown with 16E6/FN123 cells that overexpressed NFX1–123 had darker staining of NFX1–123 indicating that NFX1–123 overexpression was maintained.

Interestingly, we noted that expression of NFX1–123 rose overall with increasing distance from the basal layer in the raft cultures and normal cervical epithelium. Indeed, when staining intensity was plotted as a function of distance from the basal layer, there was a direct relationship (Fig. 1A, graphs). Because epithelial cell differentiation also progresses with increasing distance from the basal layer, we hypothesized that NFX1–123 expression was associated with cells undergoing

differentiation.

To identify a direct link between cellular differentiation and NFX1–123 expression, we induced differentiation in 16E6 HFKs using high-dose calcium or suspension in methylcellulose and quantified NFX1–123 mRNA and protein levels. 16E6/LXSN and 16E6/FN123 HFKs were cultured in media containing 1.8 mM Ca^{2+} and total mRNA and protein were serially collected. There was a significant induction of the differentiation marker Keratin 1 after calcium treatment, indicating effective differentiation (Fig. 1B, Western blot). mRNA and protein levels of NFX1–123 were higher after 72 h of calcium exposure, compared to undifferentiated cells with no calcium exposure, and rose even further after 120 h (Fig. 1B). Induction of NFX1–123 with differentiation was observed even in 16E6/FN123 cells that began with greater expression. To further test the association between epithelial differentiation and NFX1–123 expression, we induced differentiation in 16E6 HFKs using another well-characterized stimulus, suspension in methylcellulose (Green, 1977; Watt et al., 1988; Rice and Green, 1978). Again, induction of Keratin 1 indicated successful differentiation (Fig. 1C). With methylcellulose suspension, 16E6/LXSN and 16E6/FN123 cells had rapid and robust increases NFX1–123 levels at 24 h; these levels were slightly increased or maintained at 48 h (Fig. 1C, Western blot). NFX1–123 expression increased during suspension-induced differentiation in both 16E6/LXSN and 16E6/FN123 cells. Finally, we compared endogenous NFX1–123 expression in HFKs and 16E6 HFKs induced to differentiate with high-dose calcium. NFX1–123 was increased with differentiation in HFKs, and 16E6 did not have any significant impact on its endogenous expression or induction. Altogether, these data indicated that NFX1–123 itself was increased with keratinocyte differentiation, even in cells that began with overexpressed NFX1–123, and that this upregulation was unaffected by 16E6 expression.

2.2. NFX1–123 mediated keratinocyte differentiation in both Notch1-dependent and Notch1-independent manners

We have demonstrated previously that greater NFX1–123 affected keratinocyte pathways (Vliet-Gregg et al., 2013, 2015; Katzenellenbogen et al., 2009) and increased expression of differentiation markers Keratin 1 and Keratin 10 (Vliet-Gregg et al., 2015). Because these studies had been conducted in non-differentiating conditions, we were interested in interrogating whether NFX1–123 regulates expression of these markers in the context of full keratinocyte differentiation. 16E6/LXSN and 16E6/FN123 raft cultures were grown and stained for differentiation markers Keratin 1 and Loricrin (Fig. 2A). FN123 rafts with greater levels of NFX1–123 had more intense staining of these differentiation markers compared to LXSN rafts. These data confirmed that greater NFX1–123 increased expression of these markers when cells were actively differentiating.

Previous data from our laboratory has shown that NFX1–123 regulated Notch1 expression and consequently affected its canonical pathway targets (Vliet-Gregg et al., 2013, 2015). This effect of NFX1–123 and Notch1 was also linked to keratinocyte differentiation, but regulation of that pathway by NFX1–123 was less direct. Specifically, when Notch1 was activated in 16E6 HFKs overexpressing NFX1–123, canonical and differentiation targets were increased; however, when Notch1 activation was blocked in these same cells, the canonical pathway targets of Notch1 rapidly fell while differentiation pathway targets did not (Vliet-Gregg et al., 2015). To more deeply interrogate the interplay between Notch1, NFX1–123, and keratinocyte differentiation, we selected methods to stimulate differentiation in epithelial cells that would limit or permit Notch1 signaling. In selecting differentiation stimuli, we leveraged the fact that Notch1 pathway activation requires cell-to-cell contact for signaling to occur (Kopan and Ilagan, 2009; Dotto, 2008; Lai, 2004). We additionally wanted to identify how NFX1–123 altered differentiation pathways when 16E6 was co-expressed, to model how NFX1–123 might regulate

16E6 HFKs

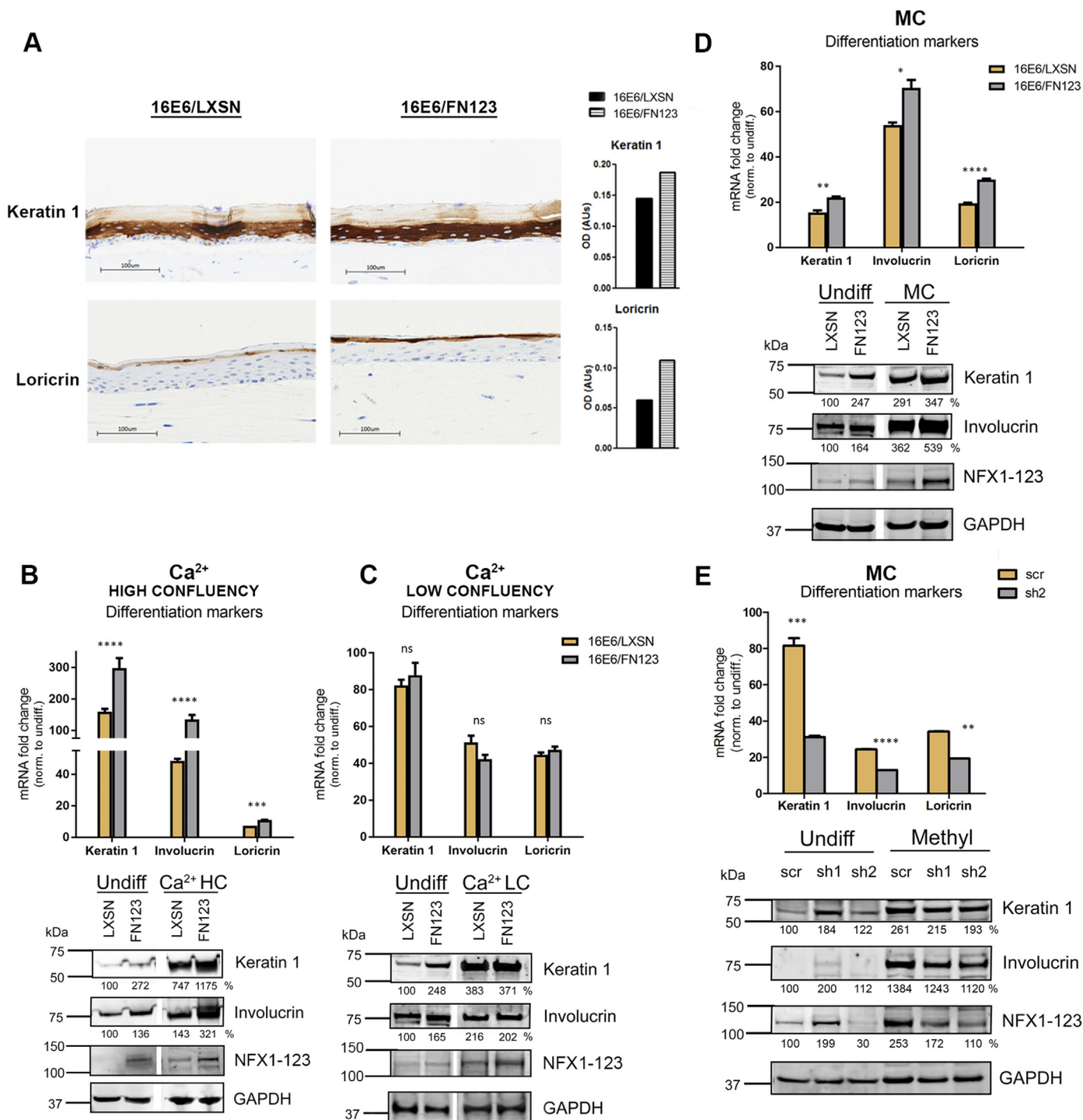


Fig. 2. NFX1-123 regulates expression of differentiation markers in 16E6 HFKs. (A) 16E6/LXSN and 16E6/FN123 raft cultures were stained for protein expression of differentiation markers Keratin 1 and Loricrin via immunohistochemistry. Total staining intensity for the section area was calculated in ImageJ. (B and C) 1×10^6 16E6/LXSN and 16E6/FN123 HFKs (B) or 3×10^5 16E6/LXSN and 16E6/FN123 HFKs (C) were plated and treated with 1.8 mM Ca^{2+} for 0 or 72 h, and total mRNA and protein collected. (D) 1×10^6 16E6/LXSN and 16E6/FN123 HFKs were suspended in methylcellulose (MC) for 0 or 24 h, and total mRNA and protein collected. (B, C, and D) mRNA levels of Keratin 1, Involutrin, and Loricrin were measured by qPCR and compared to 16E6/LXSN 0 h. Protein levels of Keratin 1, Involutrin, and NFX1-123 were assessed by Western blot. GAPDH was used as a loading control. Densitometry analysis was done in ImageJ. Protein levels were normalized to the loading control, and all samples compared to undifferentiated 16E6/LXSN. Statistical significance was calculated using unpaired *t*-tests. White spaces in Western blots indicate removal of empty or irrelevant lanes from original image for clarity. (E) 16E6 HFKs were transduced with short hairpins targeting NFX1-123 (sh1 and sh2) or scramble short hairpin control (scr). 1×10^6 scr, sh1, and sh2 cells were suspended in methylcellulose (MC) for 0 or 24 h, and total mRNA and protein collected. mRNA levels of Keratin 1, Involutrin, and Loricrin were measured by qPCR and compared to scr 0 h. Statistical significance was calculated using one-way ANOVA with Bonferroni correction. Protein levels of Keratin 1, Involutrin, and NFX1-123 were assessed by Western blot. GAPDH was used as a loading control. Densitometry analysis was done in ImageJ. Protein levels were normalized to the loading control, and all samples compared to undifferentiated 16E6/scr. For B, C, D, and E, expression levels of genes of interest were normalized to the housekeeping gene GAPDH, and all error bars represent 95% confidence intervals from replicates. One representative experiment is shown from at least three conducted in biologically independent HFK cell lines. * $p \leq 0.05$, ** $p \leq 0.01$, *** $p \leq 0.001$, ****

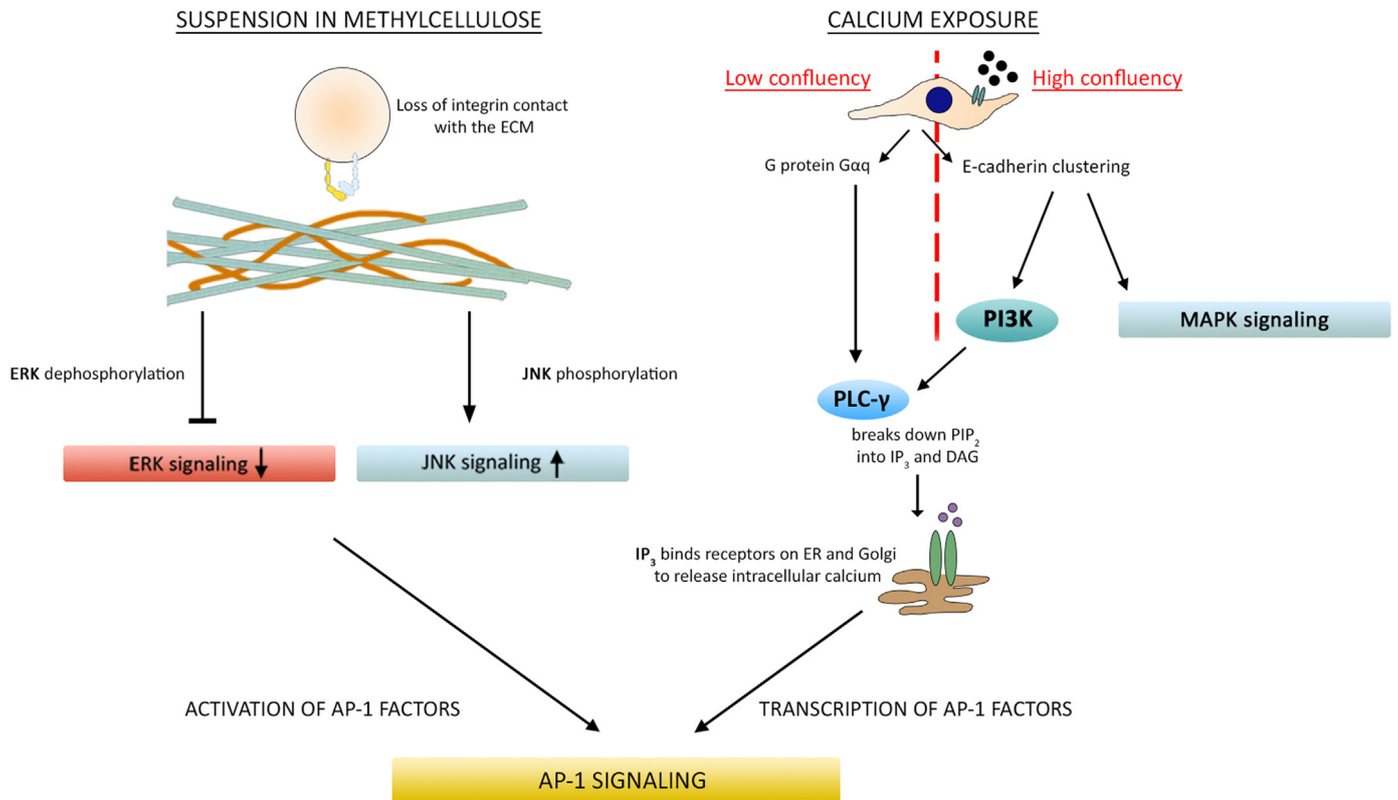


Fig. 3. Diagram of intracellular signaling pathways activated by differentiation stimuli. A representation of the intracellular signaling pathways known to be activated by differentiation stimuli used in Figs. 1 and 2. Following suspension in methylcellulose, phosphorylation and signaling activity of extracellular signal-regulated kinase (ERK) decreases. Phosphorylation and signaling activity of c-Jun N-terminal kinase (JNK) increases. This leads to activation of AP-1 transcription factors and subsequent transcriptional expression of differentiation markers. For cells plated at low confluency, exposure to extracellular calcium activates the G protein Gαq, which in turn activates phospholipase C gamma (PLC-γ). PLC-γ hydrolyzes the second messenger phosphatidylinositol 4,5-bisphosphate (PIP₂) and activates a series of reactions that lead to the transcription of AP-1 subunits, AP-1 signaling and subsequent transcriptional expression of differentiation markers. For cells plated at high confluency, exposure to extracellular calcium leads to E-cadherin clustering, activating mitogen-activated protein kinase (MAPK) and phosphoinositide 3-kinase (PI3K) activity. PI3K activates PLC-γ and here, the pathway converges with that of low confluency cells treated with calcium terminating in AP-1 signaling and transcription of differentiation markers.

differentiation in HPV infected cells. NFX1–123 interacts with E6, and our previous work showed synergy of NFX1–123 with 16E6 in increasing differentiation markers (Vliet-Gregg et al., 2015). Modulating NFX1–123 in the presence of 16E6 allowed us to probe the role of NFX1–123 itself.

First, 16E6/LXSN and 16E6/FN123 HFKs were plated at high confluency and treated with high-dose calcium for 72 h to induce differentiation. The high confluency allows for ample cell-cell contact and is thus permissive for Notch1 activation and signaling (Dotto, 2008; Lai, 2004). This Notch1 signaling was reflected in the high induction of the canonical Notch1 target gene Hes1 (Supp Fig 2) (Kopan and Ilagan, 2009). 16E6/LXSN HFKs had a robust upregulation of differentiation markers Keratin 1, Involucrin, and Loricrin (Fig. 2B, gold bars), and there was an even greater induction in the 16E6/FN123 HFKs (Fig. 2B, gray bars). This was reflected at the protein level as well; in agreement with our previous studies, we saw a greater expression of differentiation markers with more NFX1–123, even in undifferentiated cells (Fig. 2B). Following differentiation, 16E6/FN123 HFKs had greater levels of Keratin 1 and Involucrin compared to 16E6/LXSN (Fig. 2B). This confirmed that NFX1–123 played a role in mediating keratinocyte differentiation when Notch1 signaling could occur.

Next, cells were plated at low confluency (~10%), which limited cell-cell contact and therefore limited Notch1 signaling. Accordingly, there was no change in Hes1 expression in these cells (Supp Fig 2). These low confluency 16E6 HFKs were treated with high dose of calcium to induce differentiation. There was again an upregulation of Keratin 1, Involucrin, and Loricrin, indicating epithelial differentiation

was triggered; there was, however, no difference in the mRNA or protein levels of differentiation markers in 16E6/LXSN cells compared to 16E6/FN123 (Fig. 2C). Without the availability of Notch1 signaling, greater expression of NFX1–123 did not augment differentiation marker induction in 16E6 HFKs stimulated by high-dose calcium alone.

Lastly, 16E6 HFKs were stimulated to differentiate with suspension in semisolid methylcellulose medium. This stimulus virtually eliminates cell-cell contact and Notch1 signaling, similar to low confluency plating, but the intracellular signaling pathways that are activated and lead to differentiation are distinct from those activated by high-dose calcium (Watt et al., 1988; Connelly et al., 2010). Again, confirming restricted Notch1 signaling, there was no induction of Hes1 (Supp Fig 2). Examining levels of differentiation markers, 16E6/FN123 HFKs had a greater induction of these markers at both the mRNA and protein level compared to 16E6/LXSN cells, indicating that NFX1–123 was able to mediate keratinocyte differentiation stimulated by suspension and independent of Notch1 (Fig. 2D). Overexpression of NFX1–123 alone increased induction of differentiation markers even in the absence of 16E6; however, expression of 16E6 augmented this trend (data not shown). To further confirm the involvement of NFX1–123, 16E6 HFKs expressing short hairpin RNAs targeting NFX1–123 (sh1 and sh2) were differentiated via methylcellulose suspension. Knock down of NFX1–123 reduced induction of differentiation markers at the mRNA level relative to control (scr) (Fig. 2E). Although the knock down of NFX1–123 was not evident in undifferentiated sh1 cells, there was still a dose-dependent association between NFX1–123 expression and differentiation marker expression. When NFX1–123 was knocked down,

this did affect the low levels of Keratin 1 and Involucrin protein in undifferentiated cells (scr vs. sh2). After methylcellulose suspension, both sh1 and sh2 cells had decreased levels of NFX1–123 and had reduced induction of Keratin 1 and Involucrin compared to scr (Fig. 2E). These data collectively identified that NFX1–123 itself was increased during differentiation (Fig. 1); that with its increased expression, NFX1–123 augmented differentiation stimulated by methylcellulose suspension (Fig. 2D); and that NFX1–123 could achieve this through Notch1-independent signaling pathways.

2.3. Intracellular signaling and MAP kinase pathways triggered by differentiation stimuli

Extensive work in the field of epidermal differentiation has characterized the intracellular signaling pathways that are activated by the high-dose calcium and methylcellulose suspension, and how these signaling pathways subsequently lead to induction of differentiation markers (Watt et al., 1988; Connelly et al., 2010; Bikle et al., 2012; Charest et al., 2009; Pece and Gutkind, 2000; Müller et al., 2008; Watt, 2016). These complex pathways have been simplified in the schema shown in Fig. 3. In addition to their utility for investigating differentiation with or without Notch1 involvement, the stimuli chosen have unique intracellular signaling pathways (Fig. 3). Identifying where these signaling pathways overlap and differ may indicate a cellular pathway through which NFX1–123 affects differentiation. NFX1–123 regulated keratinocyte differentiation stimulated by suspension and by calcium in high confluency cells, but not low confluency. Examining the intracellular signaling pathways activated by high confluency calcium and suspension in methylcellulose, we saw that they share components of mitogen-activated protein kinase (MAPK) signaling. In the case of methylcellulose suspension, these are specifically the MAPKs ERK and JNK (Connelly et al., 2010; Trappmann et al., 2012). We hypothesized that these two kinases were therefore potential targets through which NFX1–123 mediated epithelial cell differentiation.

2.4. 16E6 and NFX1–123 mediate levels of activated JNK through upstream kinases

To investigate whether 16E6 and NFX1–123 alter JNK and ERK signaling, we returned to 16E6 HFKs with endogenous (LXSN) or overexpressed NFX1–123 (FN123) that were suspended in methylcellulose. Upon differentiation, there were no significant differences seen in total protein levels of JNK or ERK, regardless of the amount of NFX1–123 (Fig. 4A). Because these kinases are activated by phosphorylation, we also examined levels of phospho-JNK (P-JNK) and phospho-ERK (P-ERK). Suspension-induced differentiation resulted in decreased P-ERK; the small amounts were difficult to quantify and thus differences due to NFX1–123 could not be determined (data not shown). However, upon methylcellulose-induced differentiation, cells with more NFX1–123 had greater levels of P-JNK (Fig. 4A). JNK can be phosphorylated by a number of MAP kinase kinases (MKKs), but in particular by MKK4, which is specific for JNK and p38, and MKK7, which is solely JNK-specific (Johnson and Nakamura, 2007; Pearson et al., 2001). Intriguingly, 16E6/FN123 HFKs showed greater amounts of MKK4 and MKK7 compared to 16E6/LXSN HFKs. This greater MKK4 and MKK7 with more NFX1–123 was observed in both undifferentiated and differentiated cells (Fig. 4A).

To confirm the required role of NFX1–123 in JNK phosphorylation and MKK expression, we examined levels of these signaling proteins in the 16E6 HFKs expressing short hairpins against NFX1–123 that had been differentiated in methylcellulose. When NFX1–123 was knocked down by short hairpin RNA, there were again no differences in total levels of ERK or JNK (Fig. 4B). Knockdown of NFX1–123 did, however, result in decreased levels of P-JNK, MKK4, and MKK7 upon differentiation (Fig. 4B). These data indicate that NFX1–123 was required for full MKK4 and MKK7 expression and JNK phosphorylation upon

16E6 HFKs

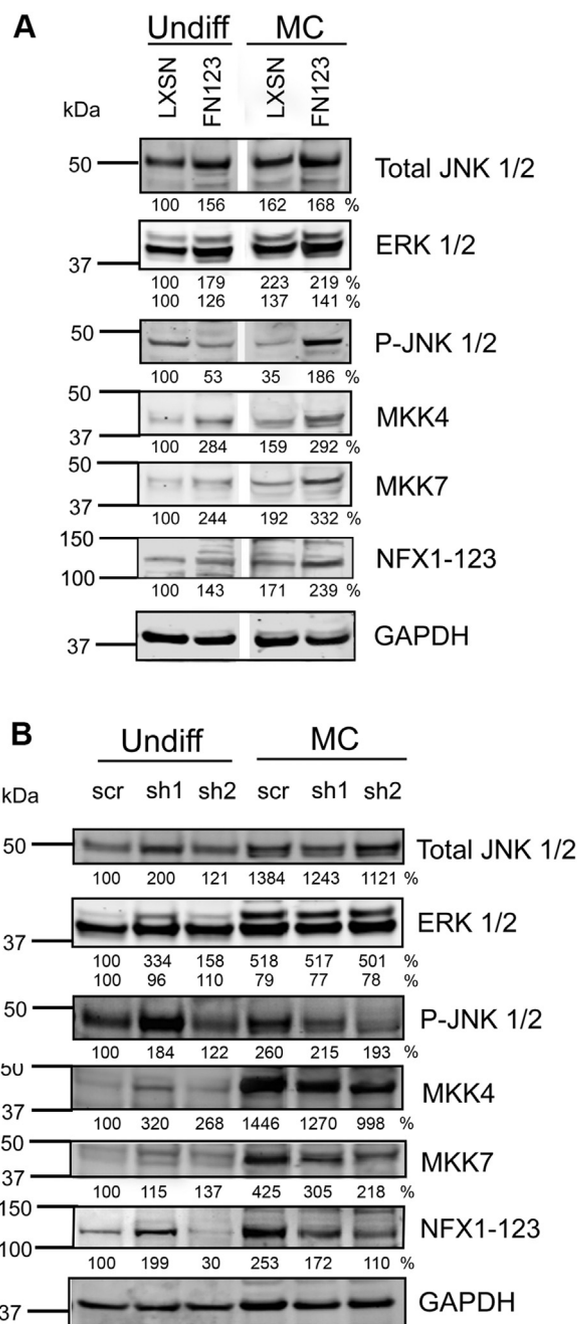
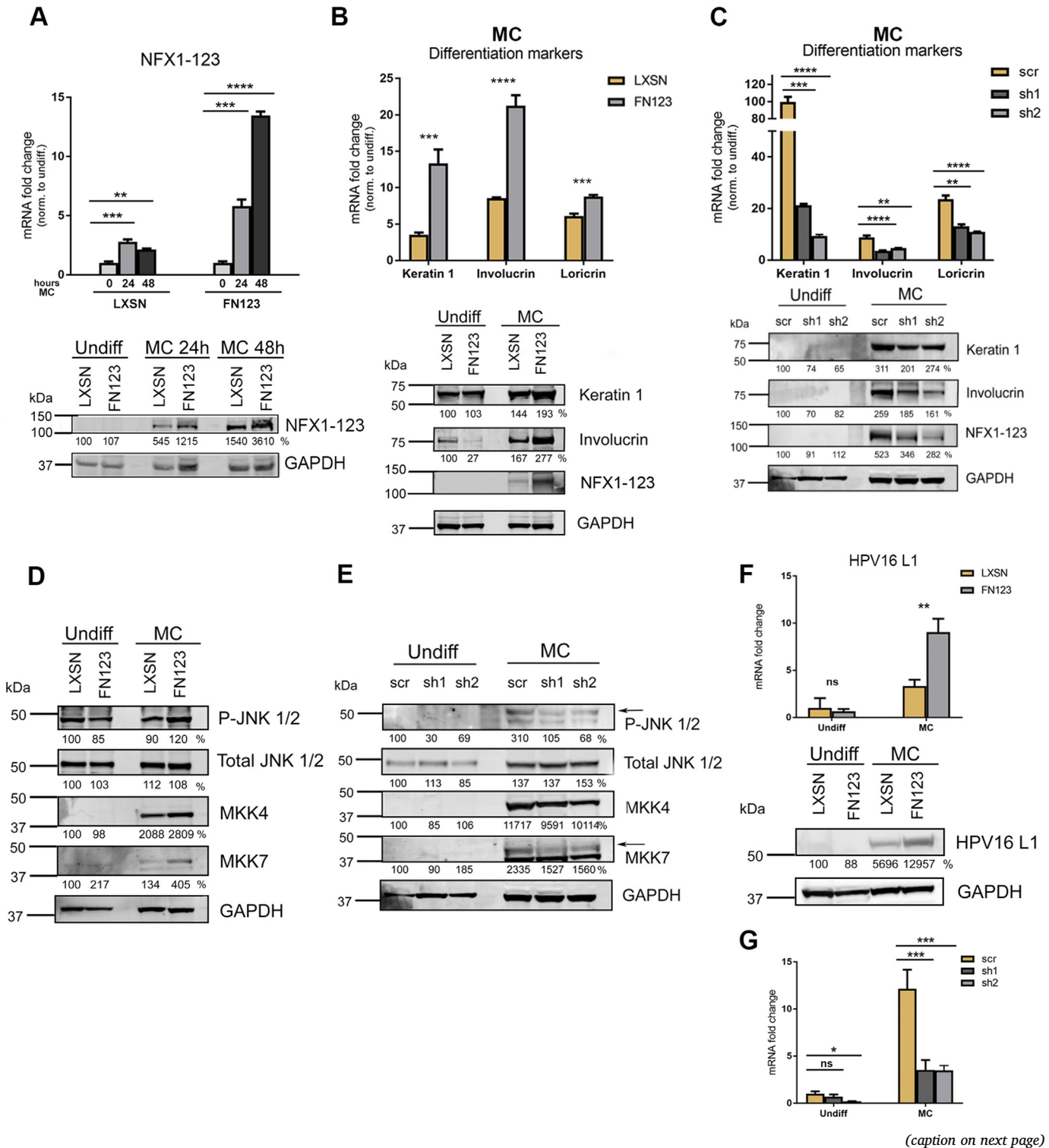


Fig. 4. NFX1–123 regulates phosphorylation of JNK through MKK4 and MKK7. (A) 1×10^6 16E6/LXSN and 16E6/FN123 HFKs were suspended in methylcellulose (MC) for 0 or 24 h, and total protein collected. Protein levels of phosphorylated JNK (P-JNK), total JNK, total ERK, total MKK4, and total MKK7 were assessed by Western blot. GAPDH was used as a loading control. Densitometry analysis was done in ImageJ. Protein levels were normalized to the loading control, and all samples compared to undifferentiated 16E6/LXSN. White spaces in Western blots indicate removal of empty or irrelevant lanes from original image for clarity. (B) 16E6 HFKs were transduced with a short hairpin targeting NFX1–123 (sh1 or sh2) or a scramble short hairpin control (scr). 1×10^6 scr, sh1, and sh2 cells were suspended in methylcellulose for 0 or 24 h, and total protein collected. Protein levels of P-JNK, total JNK, total ERK, total MKK4, and total MKK7 were assessed by Western blot. Densitometry analysis was done in ImageJ. Protein levels were normalized to the loading control, and all samples compared to undifferentiated scr. For A and B, one representative experiment is shown from at least three conducted in biologically independent HFK cell lines.

W12Es



(caption on next page)

suspension-induced differentiation.

2.5. NFX1-123 regulated JNK signaling and differentiation in HPV16-positive W12E cells and modulated HPV16 L1 expression

Given that NFX1-123 regulated keratinocyte differentiation marker expression, that this regulation by NFX1-123 was augmented by 16E6 (Vliet-Gregg et al., 2015), and that HPV late gene transcription is

differentiation-dependent, we wished to assess if NFX1-123 impacted and supported these events of the late HPV life cycle. To explore this, we utilized W12E cells that harbor the full viral genome (Doorbar et al., 1990). First, we wanted to confirm that NFX1-123 was endogenously expressed in these cells, and that its expression responded to differentiation in a similar manner as HFKs. Cells were transduced with vector control (LXSN) or NFX1-123 over-expression construct (FN123) and then suspended in methylcellulose

Fig. 5. NFX1–123 mediates differentiation and L1 expression in HPV16-positive W12E cells. (A) W12E cells were transduced with LXSN control vector (LXSN) or FLAG-tagged NFX1–123 overexpression construct (FN123) and 1×10^6 cells were suspended in methylcellulose (MC) for 0, 24, or 48 h. Mean expression of NFX1–123 mRNA measured by qPCR and compared to LXSN 0 h. Protein levels of NFX1–123 were measured by Western blot using GAPDH as a loading control. (B) 1.5×10^6 LXSN and FN123 W12E cells were suspended in methylcellulose (MC) for 24 h and total mRNA and protein collected. mRNA levels of Keratin 1, Involucrin, and Loricrin were measured by qPCR and compared to W12E/LXSN 0 h. Statistical significance was calculated using unpaired *t*-tests. Protein levels of Keratin 1, Involucrin, and NFX1–123 were assessed by Western blot. GAPDH was used as a loading control. Protein levels were normalized to the loading control, and all samples compared to undifferentiated LXSN. (C) W12E cells were transduced with a short hairpin targeting NFX1–123 (sh1 or sh2) or a scramble short hairpin control (scr), and 1.5×10^6 cells differentiated by suspension in methylcellulose for 0 or 24 h. mRNA expression of Keratin 1, Involucrin, and Loricrin was measured by qPCR and compared to scr 0 h. Statistical significance was calculated using one-way ANOVA with Bonferroni correction and *p* values for the difference in means between scr and sh1 or sh2 are shown. Protein levels of Keratin 1, Involucrin, and NFX1–123 were measured by Western blot using GAPDH as a loading control. (D and E) Protein levels of P-JNK, total JNK, total ERK, total MKK4, and total MKK7 were assessed by Western blot in LXSN and FN123 W12E cells (D) or scr, sh1, and sh2 W12E cells (E). (F) mRNA expression levels of HPV16 L1 were measured by qPCR and compared to LXSN or scr W12E cells. Protein levels of L1 were assessed by Western blot in LXSN and FN123 W12E cells. (G). For all qPCRs, expression levels of the genes of interest were normalized to the housekeeping gene GAPDH, and all error bars represent 95% confidence intervals from replicates. For all Western blots, protein levels were normalized to the loading control GAPDH, and all samples compared to the undifferentiated control (LXSN in 5A, B, D, F; or scr in 5C and E). Densitometry analysis was done in ImageJ. All experiments in Fig. 5 were performed three independent times. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$, **** $p < 0.0001$.

for 24 or 48 h to induce differentiation. Comparable to the results observed in 16E6 HFKs, NFX1–123 mRNA was expressed in W12E cells and increased in both W12E LXSN and W12E FN123 cells with 24 h of methylcellulose suspension (Fig. 5A). These levels were further increased or maintained at 48 h (Fig. 5A). Endogenous protein levels of NFX1–123 in W12E cells appeared to be lower than in HFKs, as NFX1–123 protein could not be detected by immunoblot in either undifferentiated W12E LXSN or W12E FN123 cells. Upon differentiation, however, NFX1–123 protein increased to detectable amounts, and greater protein could be seen in W12E FN123 cells relative to W12E LXSN cells (Fig. 5A).

We next examined whether NFX1–123 also regulated expression of differentiation markers in W12E cells. Overexpression of NFX1–123 in W12E cells recapitulated the results seen in 16E6 HFKs, with greater levels of NFX1–123 resulting in increased induction or differentiation markers at the mRNA and protein level (Fig. 5B). As expected, knock down of NFX1–123 resulted in decreased induction of differentiation markers in response to suspension (Fig. 5C). Analysis of MAPK proteins revealed that modulating levels of NFX1–123 had no effect on total levels of ERK or JNK, but did alter levels of P-JNK, MKK4, and MKK7 (Fig. 5D). P-JNK, MKK4, and MKK7 levels were increased when NFX1–123 expression was increased (Fig. 5D, FN123 versus LXSN) and were decreased with knock down of NFX1–123 (Fig. 5E, sh1 and sh2 versus scr). In total, these results support what was observed in HFKs: NFX1–123 was required for full induction of epithelial differentiation marker induction, augmented phosphorylation of JNK, and increased expression of its upstream kinases.

To explore whether this regulation of cellular differentiation also impacted the transcription of the HPV late genes that are triggered during differentiation, we quantified the expression of HPV16 L1; L1 encodes the HPV major capsid protein and is a common reading frame of all mRNA products expressed from the viral late promoter. As expected, both W12E LXSN and W12E FN123 cells upregulated L1 mRNA following differentiation (Fig. 5F and G Undiff versus MC). However, W12E FN123 cells overexpressing NFX1–123 had an even greater induction of L1 mRNA compared to W12E LXSN cells (Fig. 5F). This upregulation in mRNA translated to increased protein production as well, with W12E FN123 cells containing more L1 protein than W12E LXSN (Fig. 5F). Therefore, increased expression of NFX1–123 affected both cellular differentiation and subsequent differentiation-dependent viral transcription.

To determine if NFX1–123 was required for full induction of L1 during differentiation, W12E were transduced with short hairpin RNAs against NFX1–123 (sh1 or sh2) or a scrambled control (scr) and then differentiated with methylcellulose suspension. W12E scr cells, with endogenous expression levels of NFX1–123, had a robust upregulation of L1 upon differentiation, while W12E cells with NFX1–123 knocked down had a lowered induction of L1 (Fig. 5G). This demonstrated that the ability of NFX1–123 to augment epithelial cell differentiation

pathways in HPV 16-positive cells, and its requirement for full activation of cellular differentiation, subsequently influenced the differentiation-dependent late gene transcription of HPV16.

3. Discussion

This study confirms that the cellular protein NFX1–123 is a regulator of epithelial cell differentiation, identifies an intracellular signaling cascade affected by NFX1–123 expression, and subsequently provides insight into the linkage between epithelial differentiation and the HPV life cycle. The data presented here extend our previous findings, which were conducted in keratinocytes grown in non-differentiating media. We now demonstrate that NFX1–123 is able to regulate induction of epithelial differentiation markers in cells actively undergoing differentiation. Previously, we had shown that NFX1–123 was able to increase expression of differentiation markers when Notch1 was stimulated, but also determined that NFX1–123 employed Notch1-independent mechanisms (Vliet-Gregg et al., 2015). Here, we use targeted differentiation stimuli and modulation of NFX1–123 expression to explore the Notch1-independent mechanisms by which NFX1–123 regulates epithelial differentiation markers. Confirming earlier studies, we show that overexpressed NFX1–123 results in increased expression of differentiation markers when cells are stimulated to differentiate in the context of Notch1 signaling. This is seen both in raft cultures and in high confluency plating with calcium exposure (Fig. 2). When Notch1 signaling is limited, however, NFX1–123 has a more selective role. We show that NFX1–123 mediates differentiation downstream of methylcellulose suspension-induced differentiation, but not in low confluency cells treated with calcium (Fig. 2). This indicates that NFX1–123 does not have a global effect on differentiation marker expression but rather affects a particular intracellular signaling cascade.

Previous studies in our laboratory have focused on the post-transcriptional regulatory role NFX1–123 has on gene expression, both with and without 16E6.

For the case of telomerase, which is activated by 16E6, NFX1–123 post-transcriptionally stabilized the mRNA of hTERT, the catalytic subunit of telomerase (Katzenellenbogen et al., 2009, 2007, 2010). For Notch1, NFX1–123 increased its mRNA and protein expression, and like hTERT, it required its RNA binding and regulatory motifs to do so (Vliet-Gregg et al., 2013). However, in our studies of differentiation marker and pathway regulation, NFX1–123 likely does not regulate target protein expression in the same manner. If post-transcriptional stabilization of differentiation genes were the mechanism by which NFX1–123 increased their expression, we would expect NFX1–123 to affect levels of differentiation markers, irrespective of what had stimulated their expression. We instead identified that NFX1–123 specifically affected the JNK signaling pathway, with greater NFX1–123 expression robustly activating proteins within that cascade.

Further studies are required to elucidate the role of 16E6 in

partnering with NFX1–123 in the context of epithelial differentiation. NFX1–123 appears to play a role in mediating keratinocyte differentiation by itself, and data suggest that 16E6 is able to enhance this while protecting against cellular growth arrest (Vliet-Gregg et al., 2015). This mirrors what we have previously published on the regulation of Notch1 expression; NFX1–123 itself regulated the expression of Notch1, and this role was augmented by 16E6 (Vliet-Gregg et al., 2013). In contrast, the control of hTERT transcription in keratinocytes required 16E6 expression, and NFX1–123 functioned to augment this activation (Katzenellenbogen et al., 2007). These studies collectively demonstrate two mechanisms by which 16E6 functions in keratinocytes. 16E6 augments the typical regulation of pathways and genes by NFX1–123, and 16E6 also co-opts NFX1–123 to control genes and pathways it normally would not. This duality in the partnership between 16E6 and NFX1–123 has been seen with HR HPV E6 and other host protein partners, such as E6AP. E6 binds to and co-opts the functional role of E6AP in cells, targeting E6AP to degrade proteins and activate genes (Scheffner et al., 1990, 1993). These targets are both novel genes and pathways, as well as ones typically controlled by E6AP (Beaudenon and Huibregtse, 2008). More work is required to elucidate the exact contributions of E6 and its partnership with NFX1–123 in the context of differentiation. Although we know that 16E6 and NFX1–123 collaborate to modify cellular signaling pathways, and that the HPV late promoter is dependent on and responsive to cellular signaling, we do not yet know the exact molecular mechanism by which E6 is acting.

Ultimately, the data presented in this study adds to previous work demonstrating that NFX1–123, as an endogenous host protein partner, is fundamental to the cellular dysregulation driven by HPV 16E6. Without adequate expression of NFX1–123, the ability of HR HPV to activate, augment, or accelerate key pathways, such as epithelial differentiation, is muted. Our studies utilized L1 expression as one indicator of the late HPV life cycle events that are activated during differentiation. Furthermore, these studies utilized experimental systems of differentiation in order to discern the cellular networks that were activated and altered. Future experiments will further dissect differentiation-dependent events of the late viral life cycle, such as genome amplification and E1^{E4} expression, in additional systems that promote the full virus life cycle such as organotypic raft cultures.

While keratinocytes differentiating in squamous cell epithelium would have multiple, overlapping, and concurrently active stimuli and pathways, we applied a single stimulus to cells to isolate unique intracellular signaling pathway cascades. MAPK signaling, specifically involving ERK and JNK, is central to triggering suspension-induced epithelial differentiation (Connelly et al., 2010; Watt, 2016). Analysis of ERK and JNK demonstrated that after differentiation, NFX1–123 had a significant effect on levels of phosphorylated JNK (Fig. 4). Examining the kinases upstream of JNK, we identified that greater NFX1–123 resulted in increased expression of MKK4 and MKK7 (Fig. 4). Signal amplification in MAPK cascades can be driven by increased abundance of a kinase within any step of the cascade (Pearson et al., 2001). Cells with more NFX1–123 may have increased baseline levels of MKK4 and MKK7 that, when the cascade is activated, result in increased phosphorylation of JNK. In HPV-positive W12E cells, the regulation of differentiation by NFX1–123 was paralleled by expression of L1, a product controlled by the differentiation-dependent viral late promoter. Although more studies are needed to interrogate whether this regulation of JNK signaling is directly or indirectly controls L1 expression, elevation of P-JNK levels during HPV infection has been documented before (Egawa et al., 2017). In that study, increased P-JNK drove differentiation-dependent genome amplification and transcription of HPV16.

Our study joins others that document HPV's co-opting and usage of MAPK signaling. MAPK signaling is a central axis around which HPV achieves many of the events required for its life cycle (Satsuka et al., 2015): E1 phosphorylation by ERK and JNK promotes its nuclear accumulation, which is critical for viral DNA replication (Bergvall et al., 2013); activated p38 and JNK enhances E6 targeting of PDZ domain-

containing targets (Pim and Banks, 2010; Massimi et al., 2006); and sustained JNK signaling engendered by E7 prolongs the G₂ phase of cells to drive viral DNA replication (Banerjee et al., 2011; Davy et al., 2005). Alteration of MAPK signaling in the context of HPV infection has been shown previously, but this is the first time that involvement of NFX1–123 has been identified.

Another novel finding was that NFX1–123 expression was itself induced during differentiation, with the highest expression in the upper, more differentiated layers of the epithelium. This is of interest considering that even within differentiating cells, HPV genes expression is restricted to certain layers of stratified squamous epithelium. What controls this tight linkage between HPV gene expression and progressive epithelial differentiation is ill-defined. NFX1–123 may be a candidate for regulating this component of the viral life cycle. Intriguingly, we also observed basal cells with intense perinuclear staining for NFX1–123 in raft cultures and normal cervical samples. In upper layers, NFX1–123 became more predominately cytoplasmic. We see in multiple contexts that greater expression of NFX1–123 can significantly affect cellular processes and engender a milieu beneficial to HR HPV infection. These cellular processes include ones are typically thought of as opposing, such as cellular proliferation and differentiation (Vliet-Gregg et al., 2013, 2015, 2019; Katzenellenbogen et al., 2009, 2007). The subcellular localization of NFX1–123 in different epithelial layers may have a correlation with the cellular processes being affected. We have also observed in this study and others that there is a dose-response relationship between NFX1–123 levels and alterations in these cellular processes, especially in the context of 16E6. HPV may prefer to infect these basal cells that have greater than typical levels of NFX1–123. Additionally, endogenous expression of NFX1–123 may vary from person to person across normal cervical samples. It is therefore possible that upon infection with HR HPV, a person who naturally has higher endogenous levels of NFX1–123 will display more acute changes in infected cells and potentially better harbor HPV infection.

The study presented here extends previous work in which greater NFX1–123 indirectly induced differentiation marker expression through increased Notch1 expression. Now, we show that NFX1–123 itself is increase during differentiation and that it is required for and can augment differentiation marker expression in HFKs and HPV16-positive cells. These data also identify NFX1–123 as a cellular protein partner through which HPV increases MAPK signaling and engenders a cellular milieu favorable to the viral life cycle. This work adds to our understanding of the cellular pathways altered by HPV to promote infection and lends insight into the control of the differentiation-dependent HPV life cycle.

4. Materials and methods

4.1. Cell culture and generation of organotypic raft cultures

Primary human foreskin keratinocytes (HFKs) were cultured as described previously (Gewin et al., 2004). Briefly, HFKs were derived from neonatal foreskins and grown in EpiLife medium supplemented with calcium chloride (60 μM), human keratinocyte growth supplement (Life Technologies, Carlsbad, CA), and penicillin-streptomycin. 293 T cells were grown in Dulbecco's modified Eagle's medium (Thermo Fisher Scientific, Grand Island, NY) containing 10% fetal bovine serum and penicillin-streptomycin. W12 clone 20863 cells (W12E), cervical epithelial cells containing full copies of the HPV16 genome, were a gift from the laboratory of Paul Lambert. W12E cells were maintained in E-media supplemented with 5% fetal bovine serum at sub-confluency as published (Doorbar et al., 1990). Organotypic HFK raft cultures were grown in E-media containing 5% fetal bovine serum, as published with 6-well transwells (Corning, Corning, NY) in place of wire mesh, also as published (Shamir and Ewald, 2014; Anacker and Moody, 2012).

4.2. Plasmids

The FLAG-tagged NFX1–123WT (FN123), pBabe-puro 16E6 (16E6), LXS vector control (LXSN), and c-FUGW plasmids have been described previously (Katzenellenbogen et al., 2007). Scrambled short hairpin control (scr) and short hairpin 1 RNA (sh1) have also been described previously (Katzenellenbogen et al., 2007). Short hairpin 2 RNA to NFX1–123 (sh2) was designed specifically to the novel C terminus of NFX1–123. BLAST (Basic Local Alignment Search Tool) was used to confirm no off-target effects of shRNA and scramble sequences. Annealed oligonucleotides were ligated into the C-FUGW vector using BamHI and EcoRI restriction sites.

4.3. Retrovirus production and infection

Retrovirus was produced in 293T cells by a transient vesicular stomatitis virus G-pseudotyped virus (VSV-G) production protocol as previously described (Bartz and Vodicka, 1997). Lentivirus was also produced as previously described (Katzenellenbogen et al., 2007). Briefly, short hairpin NFX1–123 (sh1 or sh2) c-FUGW or scramble c-FUGW (scr) constructs were co-transfected with cytomegalovirus-vesicular stomatitis virus G and Δ8.9 plasmids into 293 T cells using FuGENE6 (Roche, Alameda, CA), and lentiviral retrovirus was serially collected. Retrovirus was concentrated by ultracentrifuge, mixed with Polybrene (8 μg/mL) (EMD Millipore, Billerica, MA), and incubated with 50–60% confluent HFKs or W12E cells. Virus was left on overnight and medium replaced the next day. For VSV-G pseudotyped virus infections (FN123, LXSN vector control, pBabe-puro 16E6) the cells were expanded 24 h post-transduction, and after 48 h the cells were placed under neomycin/G418 selection (50 μg/mL) or puromycin selection (0.5 μg/mL). All lentivirus infections (scramble, sh1 and sh2) were confirmed by green fluorescent protein expression.

4.4. Immunohistochemistry and histologic analysis

Organotypic HFK raft cultures were formalin fixed and embedded in paraffin (FFPE) following standard procedures. 32 normal cervical epithelium specimen FFPE blocks were obtained from the University of Washington HPV Research Group Specimen Repository. They were deidentified and considered to be not human subjects by the Seattle Children's Research Institute IRB. Sections 4 μm thick were stained for NFX1–123 using a rabbit polyclonal anti-NFX1–123 antibody, a gift from Dr. Ann Roman, a rabbit polyclonal anti-Loricrin antibody (BioLegend, San Diego, CA), a rabbit polyclonal anti-cytokeratin 1 (BioLegend, San Diego, CA), or a rabbit polyclonal IgG control at a 1:1000 dilution. Epitope retrieval was done using citrate for 10 min at 100 °C. Slides were scanned in brightfield at 20x magnification using the Hamamatsu NanoZoomer Digital Pathology System.

For semiquantitative analysis of NFX1–123, Keratin 1, and Loricrin staining, ImageJ was used. Color deconvolution for hematoxylin and DAB was performed and the deconvolved image displaying only the DAB channel used for subsequent quantitation. For NFX1–123, plot profiles of intensity were obtained over a set linear distance from the basal layer upwards through the apical surface of the raft culture or tissue specimen. For NFX1–123, eight plots over four sections were averaged per sample and are shown as a function of distance from the basal layer. For Keratin 1 and Loricrin, total intensity of a section were obtained.

4.5. Differentiation of keratinocytes

Prior to differentiation, HFKs were trypsin harvested from monolayer cultures. For calcium differentiation, HFKs were grown in EpiLife medium without human keratinocyte growth supplements, with 1.8 mM calcium chloride and penicillin-streptomycin added. After trypsin harvest, cells were counted using the Bio-Rad TC20 Automated

Cell Counter (Bio-Rad, Hercules, CA); for low confluency calcium differentiation, 3×10^5 HFKs were plated on a 10 cm tissue culture dish; for high confluency calcium differentiation, 1×10^6 HFKs were plated. Cells were trypsin harvested after three days' of daily calcium treatment. For differentiation in semisolid medium, 1×10^6 HFKs were plated into 25 mL of EpiLife containing 1.7% methylcellulose in a non-treated 10 cm dish. Cells were harvested using washing in ice-cold PBS and centrifugation as previously described. Untreated controls for differentiation were cells plated into monolayer in normal EpiLife and collected at the same timepoint as differentiated samples.

4.6. Differentiation of W12E cells

W12E cells were differentiated in a similar manner to HFKs. W12E cells were trypsin harvested from subconfluent monolayer cultures. 1.5×10^6 W12E cells were plated into 25 mL of semisolid medium, E-media containing 5% FBS and 1.7% methylcellulose in a non-treated 10 cm dish. Cells were harvested using washing in ice-cold PBS and centrifugation as above. Untreated controls for differentiation were cells plated into monolayer in E-media containing 5% FBS.

4.7. Quantitative real-time PCR

RNA was isolated with TRIzol reagent (Thermo Fisher Scientific, Grand Island, NY) as previously described (Katzenellenbogen et al., 2007). Total RNA (1 μg) was DNase treated and cDNA generated using random hexamer primers and SuperScript IV reverse transcriptase (Thermo Fisher Scientific, Grand Island, NY). Quantitative real-time PCR (qPCR) was performed using an ABI StepOne Plus system (Applied Biosystems, Foster City, CA). Primer sequences to amplify NFX1–123 and 36B4 were described previously (Katzenellenbogen et al., 2009, 2007). Primers used to amplify HPV16 L1 are as follows: forward primer – 5' GGTGTTGAGGTAGGTCGTGG 3' (nucleotides 5948–5967) and reverse primer – 5' CACACCTGCATTTGCTGCAT 3' (nucleotides 6042–6061) (Bienkowska-Haba et al., 2018). Amplification was carried out in replicative triplicates using Power UP SYBR Green master mix (Thermo Fisher Scientific, Grand Island, NY). Error bars represent 95% confidence intervals. For differentiation markers, amplification was carried out using TaqMan master mix and the following pre-designed Taqman probes: GAPDH (4333764F), KRT1 (Hs00196158_m1), KRT10 (Hs00166289), IVL (Hs00846307_s1), LOR (HS01894962_s1), HES1 (Hs00172878_m1) as published previously (Vliet-Gregg et al., 2015) according to the manufacturer's instructions (Applied Biosystems, Foster City, CA). Analyses were done using the ΔΔCt method, with Pfaffl correction for PCR efficiency. GAPDH was used as a reference gene for normalization of cDNA amounts, and then each experimental sample was compared to the undifferentiated control.

4.8. Western blot

Whole cell lysates were prepared directly in 2X sample buffer (100 mM Tris pH 6.8, 4% SDS, 20% glycerol). Equal amounts of protein lysates were electrophoresed on a 4–12% gradient SDS-polyacrylamide gel (Thermo Fisher Scientific, Grand Island, NY) and transferred to Immobilon-FL membrane (Millipore Sigma, Burlington, MA). Blots were probed with antibodies: Cytokeratin 1 (PA5–26699) (Thermo Fisher Scientific, Grand Island, NY); p53 (OP03) at 1:1000 (Millipore Sigma, Burlington, MA); Involucrin (SY5) at 1:1000; total JNK (D-2) at 1:500, Actin (I-19) at 1:1000, (Santa Cruz Biotechnology, Santa Cruz, CA); P-JNK/SAPK (81E11) at 1:1000, total SEK1/MKK4 (9153) at 1:1000, total MKK7 (4127S) at 1:1000, (Cell Signaling Technology, Danvers, MA); GAPDH (6C5) at 1:100,000 (Abcam, Cambridge, MA); total ERK (W15133B) at 1:1000, (BioLegend, San Diego, CA). The rabbit polyclonal anti-NFX1–123 antibody was used 1:1000 and generously provided by Dr. Ann Roman.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.virol.2019.03.008.

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