The homeoprotein DLX3 and tumor suppressor p53 co-regulate cell cycle progression and squamous tumor growth

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Abstract

Epidermal homeostasis depends on the coordinated control of keratinocyte cell cycle. Differentiation and the alteration of this balance can result in neoplastic development. Here we report on a novel DLX3-dependent network that constrains epidermal hyperplasia and squamous tumorigenesis. By integrating genetic and transcriptomic approaches, we demonstrate that DLX3 operates through a p53-regulated network. DLX3 and p53 physically interact on the p21 promoter to enhance p21 expression. Elevating DLX3 in keratinocytes produces a G1-S blockade associated with p53 signature transcriptional profiles. In contrast, DLX3 loss promotes a mitogenic phenotype associated with constitutive activation of ERK. DLX3 expression is lost in human skin cancers and is extinguished during progression of experimentally induced mouse squamous cell carcinoma (SCC). Reinstatement of DLX3 function is sufficient to attenuate the migration of SCC cells, leading to decreased wound closure. Our data establish the DLX3-p53 interplay as a major regulatory axis in epidermal differentiation and suggest that DLX3 is a modulator of skin carcinogenesis.

Keywords

keratinocytes; cell cycle; differentiation; p53; p63; DLX3; SCC

INTRODUCTION

Skin cancer is the most common form of all cancers, with cutaneous squamous cell carcinoma (SCC) comprising approximately 20% of skin malignancies. A wide variety of skin cancers such as basal cell carcinomas (BCC), SCC and melanomas harbor mutations...
in the tumor suppressor gene p5322,32. Frequently acquired mutations in RAS or p53 lead to altered responses to growth factors, perturbing the balance between keratinocyte proliferation and differentiation that is crucial to prevent neoplastic transformation 41, 45.

During epidermal differentiation, keratinocytes acquire a specific gene expression profile which includes cell cycle inhibitors and tumor suppressor genes 43. The expression of the cyclin-dependent kinase inhibitor p21 during growth arrest is controlled by the tumor suppressor p53 and the activation of p53 itself, within its role as a caretaker gene in regulating cell cycle progression 36, 57.

An important p53 family member with essential roles in epidermal homeostasis is the transcription factor p63 3. The p63 gene (TP63) encodes for multiple isoforms products of alternate promoters (ΔN and TA) and carboxy-terminal ends (α, β, δ, ε, γ, ζ) 3, 38, 56. Due to the complexity of p63 isoforms, it has been challenging to determine their exact roles in terms of enhancing or blocking cell proliferation. While rarely deleted or mutated, TP63 is frequently deregulated in human cancers 12, 28. In cutaneous SCC, high levels of p63 is used as a diagnostic marker 15 and recent characterization of isoform-specific deletions has highlighted the tumor suppressive functions or oncogenic function of the TA versus the ΔNp63 isoforms 47, 54–56.

Homeobox transcription factors play critical roles in gene regulatory networks that control developmental homeostasis 17, with their expression being also dysregulated in cancer 2. It has been shown that homeoproteins can act as drivers of tumor initiation and progression through regulation of proliferation, migration and survival pathways 39. The DLX3 homeodomain regulator is expressed during calcium (Ca++)-dependent epidermal differentiation process 37, 40 and epidermal-specific deletion of DLX3 leads to epidermal hyperplasia, accompanied by barrier disruption and associated development of an inflammatory response 24. DLX3 mutations have been associated with Tricho Dento Osseous (TDO), an ectodermal dysplasia (ED) 38 characterized by abnormalities in hair, teeth and bone 42. DLX3 is a target of p63 during ectodermal development and is involved in a regulatory feedback loop with p63 which is crucial for the maintenance of the stratified epithelia 14, 44. Mutations in p63 are also associated with human hereditary syndromes 60.

The functional interplay between p53, p63 and transcription factors in the regulation of keratinocyte differentiation has been recently highlighted for Runx1 35. Here we show that by co-regulation with p53, DLX3 affects p53 downstream targets to modulate cell cycle exit in the skin and acts as a proliferative brake. On the other hand, loss of DLX3 is conducive to a pre-neoplastic state. Consistent with this model, DLX3 is lost in human and experimentally induced murine SCCs, supporting a function of DLX3 in the context of cutaneous tumorigenesis.

RESULTS

DLX3 promotes cell cycle arrest

We assessed the impact of DLX3 transcriptional function by transducing proliferative human epidermal keratinocytes with a retroviral vector expressing DLX3 (pHAN-DLX3/Flag)
DLX3 expression induced morphological changes characteristic of keratinocyte differentiation (Figure 1a). Gene ontology analysis of differentially regulated transcripts showed that exogenous DLX3 expression promoted the dysregulation of genes involved in DNA replication, cell cycle, G1 to S cell cycle control, as well as p53 network and TGFβ signaling pathways (Figure 1b and Supplementary Table 1). Of particular relevance were the increased levels of TP53 and cyclin-dependent kinase inhibitors (CKIs) p21 and p57 transcript, as well as those of proteins involved in the formation of the G1/S transition complexes (Figure 1c left and right panel; Supplementary Table 2 and 3). The CKIs p21 and p57 share a wide range of cyclin/cdk targets to modulate G1 phase progression. Studies of p21 null mouse keratinocytes support that p21 is required for initial commitment of keratinocytes to differentiate. DLX3 transduced keratinocytes showed high levels of BAX and GADD45 which are critical factors for cell cycle arrest in differentiating cells (Figure 1c right panel; Supplementary Table 3). We also found significant upregulation of MYC, a well established effector of epidermal differentiation, in contrast with specific downregulation of proliferation markers such as proliferating cell nuclear antigen (PCNA) (Figure 1c, right panel). Alteration in multiple networks were filtered by focusing on factors of known function in cell cycle control. The downregulation of proteins involved in DNA replication are also consequences of DLX3-regulated functions (Supplementary Table 2 and 3). Significant DLX3-dependent upregulation of TP53, p21, p57 and TAp63 was verified by qPCR (Figure 1d). ΔN- and TAp63 have essential roles in epidermal differentiation, being associated with proliferation or cell cycle break/apoptosis, respectively. A role for TAp63 in the transcriptional control of keratinocyte differentiation has been recently reported. While the expression of ΔNp63 was not significantly modulated in DLX3-overexpressing human keratinocytes, we found significant modulation of a p63 target, NOXA. Our findings support a model in which DLX3 function is critical for cell cycle regulation and triggering terminal differentiation.

Thus, our results indicate that DLX3 expression in proliferating keratinocytes globally drives cell cycle block and a switch to a suprabasal profile (Supplementary Table 4), and promotes the upregulation of p53 target genes. These data were corroborated by results from an RT-PCR profiler array (Supplementary Figure 1). Cytofluorimetric analysis confirmed the increased accumulation of cells in G0/G1 and a 50% decrease in cells in S-phase (Figure 1e).

**DLX3 and p53 co-regulate p53 target, p21**

Given the DLX3-dependent modulation of p53-regulated networks in skin, we investigated the transcriptional regulation and expression of the well-known p53 targets utilizing RNA interference experiments. Reduced expression of p53 resulted in a significantly decreased upregulation of p21 and epidermal differentiation markers such as Krt10 (Figure 2a). Interestingly, the broader impact of p53 silencing was also reflected in the significantly reduced DLX3, p21 and p57 expression in DLX3-transduced cells. These results were corroborated using p53 knockout HCT116 cells, where p21 expression and endogenous DLX3 transcription were not induced by DLX3 in the absence of p53 function (Figure 2b).

The role of p21 in the epidermal differentiation process and its transcriptional regulation by p53 has been reported. Analysis of the p21 promoter region determined a potential
binding site for DLX3 in the promoter region of p21 (Figure 2c, top panel) (−2475 to −2457 in mouse and −2261 to −2242 in human). Pull down assays with anti-H3K4me3 or anti-
H3K27me3 antibodies confirmed that this region was bound by DLX3 and was
transcriptionally active in differentiating keratinocytes (Figure 2c, bottom panel). Luciferase
assays using the p21 promoter containing a WT, deleted, or mutated DLX3 binding site
(Figure 2d) as well as electromobility shift assays (EMSA) with anti-DLX3 antibody
(Supplementary Figure 2) supported the functional relevance of DLX3 binding to the p21
promoter during Ca\(^{++}\)-induced differentiation. This region of the p21 promoter also
contained well-established conserved p53-binding sites (Figure 2c)\(^49\), raising the possibility
of the formation of a transcriptional complex necessary to regulate p21 transcription. We
confirmed a specific, physical interaction between DLX3 and phosphorylated p53 on the p21
promoter that is enhanced in differentiating media (Figure 2e). These data highlight the role
of DLX3 in the regulation of p21 gene transcription during differentiation. Furthermore, the
association of DLX3 with p53 supports an essential transcriptional modulation activity
within p53 network pathways in keratinocytes.

**DLX3 expression is extinguished in skin cancer models**

We next investigated a possible contribution of DLX3 to cutaneous tumorigenesis. The
expression levels of DLX3 were monitored during tumor progression in the two-stage mouse
model of skin cancer\(^13\). We show the extinction of DLX3 expression at both mRNA and
protein level as skin papillomas progress from low risk to high risk for conversion to
squamous cell carcinomas (Figure 3). Selective ablation is noted in tumor areas expressing
KRT13, a marker of progression from benign to malignant stage\(^13\) (Figure 3c).

The strong association between DLX3 extinction and skin tumor progression prompted us to
test the impact of DLX3 loss in keratinocytes transduced with v-ras\(^{Ha}\) as an experimental
cell system. Transduction of normal keratinocytes with an oncogenic RAS allele is sufficient
to initiate skin carcinogenesis\(^48\). We found that the absence of DLX3 function correlates
with a significantly lower expression of p21, p57 and p53 transcripts when compared to WT
cells, independent of v-ras\(^{Ha}\) (Figure 4a). Interestingly, DLX3cKO cells had higher levels of
\(\Delta\)Np63, supporting a function for DLX3 in the regulation of \(\Delta\)Np63alpha levels, a
mechanism that may contribute to reduce the abundance of \(\Delta\)Np63alpha during
differentiation of stratified epithelia\(^14\). The lower levels of \(\Delta\)Np63 and the higher level of
the TA63 target p57 in v-ras keratinocytes may potentially be due to the oncogene-induce
replicative senescence\(^21,59\).

While v-ras\(^{Ha}\) transduction reduced Krt10 transcription and abolished protein expression in
keratinocytes (Supplementary Figure 3a and Figure 4b), DLX3cKO cells display a profile
characterized by lack of keratin 10 expression in the absence of v-ras transduction. Analysis
of the RAS-MAPK pathway in DLX3cKO keratinocytes showed that the loss of DLX3 in
cultured keratinocytes was associated with activation of ERK, even in the absence of v-ras\(^{Ha}\)
(Figure 4b). Proliferation was markedly increased, and growth arrest in response to Ca\(^{++}\)
was impeded in v-ras\(^{Ha}\) transduced DLX3cKO keratinocytes (Figure 4c). These results also
support the inability of v-ras\(^{Ha}\) keratinocytes to differentiate in the absence of DLX3.
Exogenous DLX3 significantly increased p57 expression in ras-keratinocytes (Figure 4d). On the other hand, while exogenous DLX3 did not restore the expression of Krt10 (Supplementary Figure S3c) it significantly modified p21, p53 or ΔNp63 expression in ras-keratinocytes, indicating a selective influence of DLX3 on ras-mediated transformation (Figure 4d). This indicates that DLX3 may induce cell cycle exit in ras-transformed keratinocytes through a p57-dependent pathway.

Interestingly, the expression of Cxcl1, a CXCR2 ligand essential for RAS mediated tumor formation was significantly higher in DLX3cKO v-ras Ha keratinocytes when compared to WT v-ras Ha keratinocytes (Supplementary Figure 3a). In contrast, exogenous expression of DLX3 in v-ras Ha transduced keratinocytes reduced to non-transformed levels the release of CXCL1 and IL1α that is critical for transformation of keratinocytes by v-ras Ha (Supplementary Figure 3b). Altogether, these results support that the loss of DLX3 leads to a mitogenic phenotype conducive to a proneoplastic state.

### DLX3 expression is lost in human SCC as a function of RAS mutation

Our data highlight the correlation of DLX3 function and skin cancer progression, and that keratinocytes without DLX3 have characteristics of initiated cells. We further assessed the expression of DLX3 on immunohistochemical analysis of human skin carcinoma tissue microarrays containing sections of SCC and metastatic adenocarcinoma lesions. DLX3 expression was lost in all tumor stages, from well or moderately differentiated SCC (grade I and II) to metastatic poorly differentiated adenocarcinoma (grade III) and undifferentiated carcinoma samples (grade IV) (Biomax Tissue Array; SK208) (Figure 5a). All of these skin lesions strongly express the malignancy marker KRT13 (Supplementary Figure 4).

Examination of published human transcriptome data sets (Oncomine) showed that DLX3 mRNA levels were downregulated in a cohort of HNSCC lesions from oral mucosa (Fold-Change: −2.310; p-value: 5.53E-11; Figure 5b). These lesions display decreased expression of TP53 and KRT1, and increased expression of TP63 (Fold-Change: −1.892; p-value= 2.10E-11; Fold-Change: −5.895; p-value= 1.78E-16; Fold-Change: 1.238; p-value= 2.68E-4 respectively; Figure 5b). In the same data sets, a significant negative fold-change for DLX3 was found in melanoma (Fold-Change: −10.453; p-value: 3.24E-6), colorectal (Fold-Change: −3.249; p-value: 4.61E-16) and breast cancer (Fold-Change: −2.524; p-value: 9.756E-10) (Figure 5c). Thus, comparative analysis of expression data highlights the association between loss of DLX3 expression and the progression of squamous cell carcinomas.

### Reinstatement of DLX3 in SCC cell lines alters cell cycle and differentiation

Since DLX3 expression is undetectable in human SCCs, we examined the levels of DLX3 expression in SCC lines and confirmed that it is abolished in SCC13 and SCC9 (Figure 6a). We hypothesized that the reinstatement of DLX3 into SCC cell lines would impact cell cycle. To analyze the effect of DLX3 function in SCC9 and SCC13 lines, we utilized a scratch assay to study migration upon infection with Ad-GFP or Ad-DLX3. We determined that wound closure occurred at a delayed rate upon addition of Ad-DLX3 in both SCC13
and SCC9 (Figure 6b and c), and could be morphologically observed by 18 hours post infection.

To investigate the transcriptional programs underlying the reduced wound closure upon DLX3 restoration, we performed transcriptomic profiling of SCC13 infected with Ad-GFP or Ad-DLX3 at 18, 24, 36, or 48 hours post infection. An Ingenuity Pathway Analysis of the genes at 48 hours (fold change >2 and ≤−2) revealed a negative activation Z-score for genes related to cell movement, cancer, growth, and cell cycle (Figure 6d and Supplementary Table 5). At 48 hours post infection we show that, while TP53 was unaffected, specific transcriptional targets related to the G1/S phase were differentially regulated (Supplementary Table 6). Isoforms of the CDKN2a gene, p16INK4A and p14ARF were upregulated (6.5 and 8.8 fold, respectively), with downstream targets regulated to promote inhibition of the G1 to S transition (Figure 6e).

We then sought to understand the global transcriptional changes of the differentially regulated genes at 18, 24, 36, and 48 hours post-infection. A heat map of hierarchically clustered fold changes reveals a trend toward upregulation of genes involved in negative regulation of the cell cycle, signaling and cell adhesion in clusters 1 and 2. In clusters 3 to 6, we observe downregulation of apoptosis regulation, cell cycle, and cell chemotaxis with the most pronounced differences at 48 hours (Figure 6f). This suggests that DLX3 function may lead to a metabolically active cell cycle arrest in SCC cells through the upregulation of genes involved in cell cycle arrest and a downregulation of genes involved in the pro-apoptosis pathway (Supplementary Table 6 and 7). Thus, our results indicate that the reinstatement of DLX3 is sufficient to attenuate SCC cell migration and affect transcriptional changes involved in signaling, adhesion, cell cycle, and cell movement. Collectively, our results support the emerging concept that DLX3 acts as a key regulator of cell cycle progression and squamous tumor growth.

**DISCUSSION**

This work establishes the functional association of the DLX3 within p53-regulated networks in skin keratinocytes and in cutaneous tumorigenesis. Functionally, we show that DLX3 expression is necessary for cell cycle block and differentiation. An integrated transcriptome analysis identified a DLX3-dependent gene signature characterized by downregulation of the DNA replication signaling pathway and upregulation of the p53-dependent cell cycle related genes. The p53 network signaling, which includes p63 isoforms (ΔN- and TAp63) and p21, is determinant in promoting keratinocyte maturation.

Notably, ΔNp63 isoform overexpression is characteristic of SCC and other epithelial cancers and it correlates with decreased differentiation and increased oncogenic activity. Recent work by Chakravarti and colleagues provides a mechanistic understanding on the role of Np63 in epidermal differentiation by demonstrating that ΔNp63 transcriptionally regulates the miRNA processing cofactor DGCR8 to reprogram adult somatic cells into multipotent stem cells. In contrast, TA isoforms are associated with cell cycle arrest by the transactivation of p53 target genes, such as p21, BAX or PERP and assume the role of tumor suppressors. TAp63 was recently reported to suppress a wide
variety of tumors in vivo including cuSCC and to transcriptionally regulate p57 to maintain stem cells in the dermis in quiescence using in vivo models. Recent work establishes a significant p63-p53 interplay in the differential expression of the transcription factor Runx1 in proliferative or differentiated human keratinocytes. Interestingly, as for ΔNp63, RUNX1 expression was strongly expressed in BCC and, in parallel to p53, was downregulated in SCC.

It has been previously reported that exogenous expression of the homeobox transcription factor DLX3 decreases Np63 protein levels through proteasomal degradation. Here we report that loss of DLX3 increases ΔNp63 expression and that p63 transcriptional targets, such as NoxA, decrease upon the exogenous expression of DLX3 in keratinocytes. Therefore, DLX3 may be responsible for the reduction of ΔNp63 expression during the differentiation of the stratified epithelia.

We also found that exogenous DLX3 increases the expression of TAp63 and its transcriptional target p57, thus regulating p63 signalling at multiple levels. Furthermore, exogenous DLX3 expression significantly increased p57 levels in ras-keratinocytes and SCC13 cells. Therefore, DLX3 function in the context of transformed cells could induce cell cycle exit through the TAp63-p57 axis.

Here we present evidence that loss of DLX3 removes a proliferative brake from keratinocytes conducive to a preneoplastic state with the development of a molecular signature characteristic of tumorigenic SCC cells. In the case of the p53 target p21, the association between p53 and DLX3 within the p21 promoter is directly required to p21 expression. In a broader sense, our results establish DLX3 and p53 signaling networks as interconnected circuits that regulate epidermal differentiation in normal epidermal keratinocytes.

We demonstrate that DLX3 is lost in human and experimentally induced murine SCCs corroborating a potential tumor suppression function of DLX3 in the context of cutaneous tumorigenesis. Mechanistically, we show that DLX3 expression itself may modulate ERK activation, and the loss of DLX3 is potentially linked to an activated mitogenic phenotype required for the development of neoplastic lesions.

Collectively, our data support the correlation between DLX3 and p53-regulated network in skin tumorigenesis, identifying DLX3 as an essential modulator in the regulation of cell cycle exit and revealing its tumor suppressor activity in epidermal cells. We show that addition of DLX3 function in SCC lines is sufficient to cause altered migration in a scratch wound assay resulting in decreased wound closure. Molecularly, we demonstrate that DLX3 alone impacts cell cycle exit. We propose that reinstatement of DLX3 function in squamous tumors could potentially be used to reduce tumor growth by restoring responses to a differentiation signal in the skin.
MATERIALS AND METHODS

Generation and analysis of mice

Mouse experiments were approved by NIAMS Animal Care and Use Committee. DLX3cKO mice were generated and genotyped as previously reported. The two-stage model of skin carcinogenesis was performed as previously described. For this experiment all animals, male or female, received treatments while untreated skin was used as control. No blind studies were performed.

Cell culture and transduction

Normal human epidermal keratinocytes (NHEK) were obtained from surgically resected human foreskins from the NIH Clinical Center, Dermatology Branch, after provision of written consent. NHEK were isolated as previously described and maintained in KGM (Lonza), with Ca++ concentration of 0.05 mM (proliferating media). To induce human keratinocyte differentiation, a final concentration of 1.8mM Ca++ was used. We transduced NHEK with retroviral vectors containing or not DLX3 cDNA (pHan-DLX3/flag and pHan-EV, respectively) in the presence of 4μg/ml polybrene (Sigma). Retroviral particle were generated in Phoenix cells using Promega’s ProFection® Mammalian Transfection Systems kit. We monitored morphological changes of transduced cells in culture using a Carl Zeiss AxioVert S100 microscope, AxioCam HRm Carl Zeiss digital camera and visualized using AxioVision Rel. 4.4 software. Primary murine keratinocytes (PMK) were isolated from DLX3cKO and WT mice as previously described. For each experimental replicate, a minimum of 10 newborn mice were used. PMK were cultured in S-MEM (Gibco) with 8% Fetal Bovine Serum (FBS, Atlanta Bio), 1% antibiotic (PSA, Sigma), with Ca++ concentration of 0.05 mM. To induce keratinocyte differentiation, a final concentration of 0.12 mM Ca++ was used. HCT116 cells were cultured in McCoy’s 5A media (Gibco) supplemented with 10% FBS (Atlanta Bio) and 1% antibiotic (Pen-strep, Sigma). SCC13 and SCC9 cells were cultured in DMEM/F12 + GlutaMAX + sodium pyruvate (Gibco) supplemented with 10% FBS (Atlanta Bio) and 1% antibiotic (Pen-step, Sigma).

Cells were transfected with scramble siRNA or p53 siRNA (100nM, Thermo Scientific), or scramble siRNA; transduced with GFP or DLX3 adenoviruses (50 MOI; Vector Biolab); control or v-rasHa retrovirus (1 MOI) in the presence of 4 μg/ml polybrene (Sigma). Retroviral particles were generated as previously described.

Site-Directed mutagenesis and luciferase reporter assay

The p21 promoter region (−2475/+1) was ligated into pGL3 basic vector (Promega), and confirmed by restriction digestion and sequencing. Using Quickchange® (Stratagene), we introduced mutations on the Dlx3 binding site within the p21−2475/+1 WT promoter construct. Mutagenesis was confirmed by sequencing. The following primers were used to mutate or delete the Dlx3 binding sites: p21WT_Forward: TTTTTTTCTCGAGTGTGGAGGTGACTTCTTCTGAAAATCTGA; p21WT_Reverse: TTTTTTAAGCTTTGACCAACTGTGGAGGACTAATCGT; p21Del_Forward: TTTTTTCTCGAGGAGGAGGTGTCACACGACTACAGCTTCAGTTT; p21Del_Reverse: TTTTTTAAGCTTTGACCAACTGTGGAGGACTAATCGT;
p21Mut_Forward: GACCCCAGATGGCCAAGGATGTCCCACTTTGC; p21Mut_Reverse: GCAAAGTGGGACATCCTTGGCCATCTGGGGT. Using Dual-Luciferase® Reporter Assay System (Promega), lysates from transfected keratinocytes were analyzed using a 20/20® Luminometer (Turner Systems) 48 hours after transfection.

**Electrophoresis Mobility Shift Assay (EMSA)**

EMSA was performed as previously described. A probe containing the Dlx3 consensus-binding site (GGGGGATAAAATTGCTGG) and a probe containing the Dlx3 binding site motif of the p21 promoter (GGGAGATAATTAAGGA) were used to perform the assay in presence of the recombinant protein DLX3. For the competition assays, mutated (GGGGGATAAATTGCTGG) and WT unlabeled probes were used. To perform the super-shift, the reaction was incubated with 3μg of anti-Dlx3 (Abnova; H00001747).

**Thymidine incorporation proliferation assay**

Primary keratinocytes (NHEK and PMK) were plated in 24-well tissue culture plates and RAS-transduced on day 3. Cells were pulsed for 4 h with 1 μCi [3H]thymidine (GE Healthcare). Cells were fixed using methanol and acetic acid (in a 3:1 ratio) and then solubilized in 5 N NaOH. The incorporated counts were measured using a scintillation counter.

**RNA extraction and qRT-PCR**

Total RNA was extracted using RNeasy kit (Qiagen) according to the manufacturer’s instructions. cDNA was prepared using the ImProm-II Reverse Transcription System (Promega). For each gene, qRT-PCR analysis was done in triplicate using the iQSYBR Green Supermix (Bio-rad). The following genes were analyzed: DLX3, p53, p21, p57, ΔNp63, TAp63, KRT14, KRT10 and Cxcl1. Primer sequences used in qRT-PCR as indicated (Supplementary Table 8). Relative expression was normalized against the housekeeping gene RPLPO or β-actin. Fold changes were calculated by the Ct method as described previously.

**Western blot**

Lysates of cells and tissues were prepared in SDS sample buffer (60 mM Tris-HCl at pH 6.8, 10% glycerol, 2% SDS, and 5% 2-mercaptoethanol). Equal amounts proteins (25 μg) were separated on 4% to 12% SDS-PAGE gels, transferred to PVDF membranes (Invitrogen), and incubated with primary antibodies specific to ERK (Cell Signaling; 4695), P-ERK (Cell Signaling; 4370S), H-RAS (Santa Cruz; sc-520), Krt10 (Covance; PRB-159P), Vinculin (Sigma; V4139). Blots were rinsed in TBST and incubated in peroxidase-conjugated anti-mouse or anti-rabbit secondary antibodies, respectively. Proteins were visualized using an ECL detection system (Pierce Biotech) and exposed to film. All experiments were repeated in triplicate.

**ChIP**

NHEK and PMK maintained in proliferative or differentiating media, were fixed with formaldehyde and processed using the ChIP Assay Kit (Millipore) as per manufacturer’s...
instructions. The following antibodies were used: anti-IgG (Abcam; ab46540); anti-DLX3 (Abcam; ab66390); anti-H3K4me3 (Diagenode; C15410003) and anti-H3K27me3 (Diagenode; C15410195). The following primers were used to amplify p21 promoter region: for mouse: p21Forward CATGTTCCAGCCTTGAATTGAAGAG, p21Reverse: ATACAGTACCTGACACATACACC; for human: p21Forward: CATTCCCTACCCCCATGCTGCTC; p21Reverse: CTTCAGGGCTGAAAGGGTTT.

**Flow Cytometry**

Cells were prepared in 70% ethanol and stained with Propidium Iodide (20mg/ml). Data were acquired on BD Biosciences FACS Canto and analysis was performed using Flowjo Software.

**Histology and immunohistochemistry**

For histological (Hematoxylin and Eosin staining – H&E) and immunohistochemical analysis, tissues were fixed with 4% PFA and embedded in paraffin. 5 μm sections were prepared for specific staining. Each section was incubated overnight at 4°C with the primary antibodies which included: DLX3 (Epitomics; 8434-1), Krt5 (Acris; BP5006), Krt13 (Abcam; ab92551). The Alexa Fluor 488 and 555 –conjugated were used as secondary antibodies.

**mRNA Expression Profiling and RNA sequencing**

For microarray analysis, the Affymetrix GeneChip scanner 3000 running Affymetrix (GeneChip Operating Software) was used. Significantly affected genes (p<0.05 and fold change>1.5) were determined on ANOVA. RNA from SCC13 cells infected with Ad-GFP or Ad-DLX3 was extracted with Trizol (Life Technologies). RNA-seq was performed as previously described. mRNA expression profiling were performed in the NIAMS Genome Core Facility at the NIH. Reads were mapped to the human genome using TopHat software. Expression values (rpkm) and fold changes were calculated and analyzed with the Partek Genomics Suite and ANOVA statistics. Genes with robust expression (rpkm > 1) were considered for further analysis. The row clustering for the heat map was clustered using Pearson correlation across all samples. The heat map values represent the Log2 fold change (>1=2 fold up and <=−1=−2 fold down). Analysis of clusters was done using DAVID Bioinformatics Functional Annotation Tool (http://david.abcc.ncifcrf.gov/summary.jsp). Completed microarray and RNA-seq data have been deposited in the Gene Expression Omnibus (GSE63049) site.

**Human skin SCC**

Human skin and SCC samples were from Biomax SCC tissue array (US Biomax; Cat#: sk-208), that contained 60 cases of skin cancer (squamous cell carcinoma and metastatic adenocarcinoma) plus 9 cases of normal tissue from autopsy. Squamous tumor and adenocarcinoma are ranked from grade I to grade 4 as pathological diagnosis, which is equivalent to well differentiated, moderately differentiated, poorly differentiated or undifferentiated, respectively. Pathological grade was confirmed for all cases.
Scratch Wound Assay

$2 \times 10^4$ cells of SCC13 and SCC9 cells were seeded in 96-well ImageLock plates (ESSEN BioScience) and grown to 90 – 95% confluency. Subsequently, a scratch was placed in the middle of the wells with the WoundMaker (ESSEN BioScience). After washing with phosphate buffered saline, the respective virus (Ad-GFP or Ad-DLX-GFP at a titre of $10^{10}$ PFU) was added to each well at an MOI of 50. The cells were grown in DMEM/F-12 + GlutaMAX + sodium pyruvate (Gibco) supplemented with 5% BSA. Pictures were taken every two hours for 48 hours using the IncuCyte ZOOM 40008 instrument and analyzed with the IncuCyte Zoom 2013A software (ESSEN BioScience).

Computational Analysis

Microarray expression fold-changes were calculated using Partek Genomics Suite (http://www.partek.com). Pathway enrichment was calculated using the hypergeometric distribution and WikiPathways 26. All heatmaps were generated with the R programming environment using Wards method with Euclidean distance (http://www.r-project.org/). Cytoscape was used for network visualization 53.

Statistical Analysis

For all experiments in vitro technical and biological triplicates were performed. The data were analyzed by Prism Software (GraphPad Software) and significance values were assessed by the Student’s t-test. For all experiments with error bars, standard deviations were calculated to indicate variations within each experiment and data, and values represent mean ± standard deviation of mean. P<0.05, P<0.01, P<0.001 were considered significant.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. DLX3-dependent p53 signaling network activation drives cell cycle exit and differentiation

(a) Upper panel: Normal Human Epidermal Keratinocytes (NHEK) transduced with Empty Vector (EV) or pHan-DLX3/Flag (DLX3/Flag) expressing vectors in proliferating media (0.05mM Ca^{++}). Cell morphology was assessed 24 hours post infection. Lower panel: Western Blot analysis with anti-Flag or β-actin antibodies. (b) Top significant biological pathways in microarray data of DLX3-transduced keratinocytes ranked by fold-change average and determined by Wikipathways analysis. (c) Left panel: A subset of the G1 to S cell cycle control pathway determined by Wikipathways database overlaid with DLX3-transduced keratinocyte expression profile. Red and green boxes represent up and down-regulated genes, respectively. Right panel: p53 target gene fold change (FC) in DLX3 transduced keratinocytes (Fold-change > ±2; P< 0.05). (d) Bar graphs showing relative expression level of TP53, p21, p57, ΔNp63, TAp63, and NOXA in DLX3 transduced keratinocytes. The results are shown as mean ± SD of three independent experiments. *P<0.05, **P<.001, and *** P<0.01. (e) PI staining of EV and DLX3-transduced keratinocytes (mean ± SD, ** P<0.05).
Figure 2. DLX3 promotes p53-dependent p21 transcription

(a) Relative expression levels of TP53, DLX3, p21, p57 and KRT10 in keratinocytes transfected with scramble or p53 siRNA and transduced with Ad-GFP or Ad-GFP-DLX3 (mean ± SD; *P<0.05, **P<0.01, ***P<0.001), (b) Relative expression levels of TP53, DLX3, and p21 in TP53 null HCT116 cells and TP53 wild type HCT116 cells transduced with Ad-GFP or Ad-GFP-DLX3 (mean ± SD; *P<0.05 and **P<.001). (c) Top panel: p53 and DLX3 consensus sites on the p21 promoter. Bottom panel: ChIP assay with DLX3, H3K4me3 and H3K27me3 antibodies showing specific binding on p21 promoter. IgG

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antibody was used as control. (d) Bar graph showing p21 promoter activity induced by DLX3 measured by luciferase assay. Normal human keratinocytes were co-transfected with p21 promoter constructs (Wild type (WT), deleted (Del) or mutated (Mut) DLX3 binding site), and pHan-DLX3/Flag or pHan-EV. (e) Western blot showing P-P53 pull-down by DLX3 antibody of primary keratinocyte cell lysates in proliferating and differentiating media.
Figure 3. DLX3 expression is lost in mouse skin carcinogenesis model
(a) DLX3 expression level by microarray data analysis of normal, low- and high-risk papilloma and SCC lesions and (b) Hematoxylin and Eosin and immunohistochemical staining with antibodies against KRT5, DLX3 and KRT13 of, 1–2) Low risk papilloma, 3–4) high risk papilloma, 5 – 6) SCC. Scale bar, 20 μm.
Figure 4. DLX3 loss promotes a mitogenic phenotype associated with constitutive activation of ERK

(a) Upper panel: Relative expression levels of Tp53, p21, p57, and ΔNp63 in WT and DLX3cKO keratinocytes transduced with v-ras\(^{Ha}\). (b) Western blot analysis utilizing H-RAS, KRT10, ERK, phospho-ERK and Vinculin antibodies in proliferative (0.05mM Ca\(^{++}\)) and differentiating (0.12mM Ca\(^{++}\)) media. Total-ERK vs Phospho-ERK Ratio was normalized to Vinculin expression

(c) Cell proliferation assessed by 3H-thymidine in proliferating or differentiating media. (d) Relative expression levels of Tp53, p21, p57, and ΔNp63 from Ras-transformed keratinocytes infected with Ad-GFP or Ad-DLX3. (mean ± SD; *P<0.05, ** P<0.01, *** P<0.001),
Figure 5. DLX3 expression is lost in human squamous carcinoma
(a) Immunohistochemical staining of normal skin, squamous cell carcinoma of grade I and II, metastatic adenocarcinoma and undifferentiated carcinoma sections with antibodies against K5 and DLX3. Scale bars, 20 μm. (b) Box-plots showing DLX3, TP53, KRT1 and TP63 expression levels in oral SCC and normal samples by Oncomine Database. (c) Box-plots showing DLX3 expression levels in HNSCC samples determined by TCGA Database. G1 to G4 indicate increased severity in tumor stages.
Figure 6. DLX3 attenuates the migration of SCC cells

(a) Relative mRNA expression levels of DLX3 in Normal Human Epidermal Keratinocytes, SCC13, and SCC9 (mean ± SD; ***P<0.001). Scratch wound assay of (b) SCC13 cells and (c) SCC9 cells infected with Ad-GFP or Ad-DLX3 with GFP fluorescent images taken 18, 24, 36, and 48 hours after scratch and infection. Right panel: Relative Wound Closure percentage of cells measured every two hours for 48 hours after scratch and infection. (d) Top significant biological function annotations identified by Ingenuity Pathway Analysis of differentially regulated SCC13 mRNA fold changes comparing Ad-DLX3 versus Ad-GFP samples 48 hours after infection. (Fold change>2 or <−2). (e) G1 to S cell cycle control
pathway determined by Wikipathways database overlaid with RNA-seq of SCC13 cells infected with Ad-DLX3 compared to Ad-GFP after 48 hours of infection. Red and green boxes represent up and down regulated genes, respectively. Right panel: fold changes of differentially expressed mRNA at 48 hours after infection. (f) Heat map of hierarchically clustered fold changes of SCC13 mRNA infected with DLX3 compared to GFP control at 18, 24, 36, and 48 hours post-infection. Clusters were analyzed by DAVID Bioinformatics Functional Analysis Tool with the top five significant functional annotations listed for each cluster.