

Original Research

Lysyl hydroxylase 2-induced collagen cross-link switching promotes metastasis in head and neck squamous cell carcinomas[☆]



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Abstract

Head and neck squamous cell carcinoma (HNSCC) is the 6th most common cancer worldwide and incidence rates are continuing to rise globally. HNSCC patient prognosis is closely related to the occurrence of tumor metastases, and collagen within the tumor microenvironment (TME) plays a key role in this process. Lysyl hydroxylase 2 (LH2), encoded by the Procollagen-Lysine,2-Oxoglutarate 5-Dioxygenase 2 (*PLOD2*) gene, catalyzes hydroxylation of telopeptidyl lysine (Lys) residues of fibrillar collagens which then undergo subsequent modifications to form stable intermolecular cross-links that change the biomechanical properties (i.e. quality) of the TME. While LH2-catalyzed collagen modification has been implicated in driving tumor progression and metastasis in diverse cancers, little is known about its role in HNSCC progression. Thus, using gain- and loss-of-function studies, we examined the effects of LH2 expression levels on collagen cross-linking and cell behavior *in vitro* and *in vivo* using a tractable bioluminescent imaging-based orthotopic xenograft model. We found that LH2 overexpression dramatically increases HNSCC cell migratory and invasive abilities *in vitro* and that LH2-driven changes in collagen cross-linking robustly induces metastasis *in vivo*. Specifically, the amount of LH2-mediated collagen cross-links increased significantly with *PLOD2* overexpression, without affecting the total quantity of collagen cross-links. Conversely, LH2 knockdown significantly blunted HNSCC cells invasive capacity *in vitro* and metastatic potential *in vivo*. Thus, regardless of the total “quantity” of collagen crosslinks, it is the “quality” of these cross-links that is the key driver of HNSCC tumor metastatic dissemination. These data implicate LH2 as a key regulator of HNSCC tumor invasion and metastasis by

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☆ Abbreviations: ECM, extracellular matrix; HNSCC, head and neck squamous cell carcinoma; LOX, lysyl oxidase; PLOD2, Procollagen-Lysine,2-Oxoglutarate 5-Dioxygenase 2.

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modulating collagen cross-link quality and suggest that therapeutic strategies targeting LH2-mediated collagen cross-linking in the TME may be effective in controlling tumor progression and improving disease outcomes.

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Introduction

Head and neck squamous cell carcinomas (HNSCC) are the sixth leading cancer by incidence worldwide with an estimated 600,000 new cases per year [4, 10, 24]. In the United States, HNSCC accounts for ~3% of all cancers with an estimated 54,000 new cases and 11,000 deaths per year [52]. More than half of these new cases involve patients diagnosed with advanced stage disease that is associated with poor prognosis and low 5-y survival rates [51]. The frequent emergence of local disease recurrence and tumor metastasis contributes to the high morbidity and mortality associated with this cancer [46]. Notably, HNSCCs often exhibit desmoplastic features characterized by excessive accumulation of fibrillar collagen within the stroma that forms a stiff extracellular matrix (ECM) [26]. The progressive linearization and thickening of collagen is known to augment cell growth and survival, via cellular mechanosignaling [29, 39], and lead to increased cell migration and metastatic progression [28, 37].

Fibrillar type I collagen is the major component of the tumor stroma in solid cancers [16, 27, 55]. Type I collagen is a heterotrimeric molecule composed of 2 $\alpha 1$ chains and one $\alpha 2$ chain, which consist of 3 structural domains: Amino-terminal nonhelical telopeptide (N-telo), central triple helix (helical), and carboxy-terminal nonhelical telopeptide (C-telo) domains [63]. After procollagen molecules are synthesized and post-translationally modified within cells, they are secreted into the extracellular space and undergo additional enzymatic processing before self-assembling into a fibril, which is then stabilized by covalent intermolecular cross-linking [62]. Cross-linking is initiated by the conversion of telopeptidyl lysine (Lys) and hydroxylysine (Hyl) to the respective aldehyde, Lys^{ald} and Hyl^{ald}, via the action of lysyl oxidase (LOX) and LOX-like enzymes (LOXL1-4). These aldehydes then undergo a series of condensation reactions involving the juxtaposed Lys, Hyl and histidine (His) residues on the neighboring collagen molecules. The critical feature that determines the stability of cross-links is the state of Lys hydroxylation in telopeptides; Hyl^{ald}-derived cross-links are more stable in comparison to Lys^{ald}-derived cross-links [8, 20, 61]. The Lys hydroxylation reactions are catalyzed by lysyl hydroxylases 1-3 (LH1-3). However, only LH2, encoded by the Procollagen-Lysine,2-Oxoglutarate 5-Dioxygenase 2 (*PLOD2*) gene (LH2 hereafter), catalyzes Lys hydroxylation specifically within the N- and C-telopeptides [45, 58, 60].

Recent studies have shown that aberrant collagen cross-linking plays a significant role in controlling tumor ECM stiffness, which promotes cancer progression and metastasis [9, 28, 48, 62]. Thus, much attention has been paid to the role that collagen modifying enzymes such as LOX family members and LH2 play in determining the quantity and quality of collagen cross-linking, respectively. It has been reported that LOX and LOXL2 are widely up-regulated in tumors [1, 11, 32], suggesting that the overall quantity of collagen cross-links is increased across many cancers. More recent reports have shown that LH2 is specifically up-regulated in late-stage, metastatic cancers [9, 14, 19, 48], strongly suggesting that a ‘switch’ in tumor-ECM quality resulting from LH2-mediated stable Hyl^{ald}-derived cross-links drives aggressive tumor behaviors. Taken together, it is now abundantly clear that the

biomechanical nature of fibrillar collagens plays a critical role in modulating cancer cell behaviors associated with migration and metastasis [25, 41, 62]. However, little is known about the functional role of LH2 in the pathobiology of HNSCC maintenance and/or progression.

In this study, we mechanistically examined the effects of LH2 expression levels on collagen cross-linking, and specifically how these modifications affect tumor cell behavior, both *in vitro* and *in vivo*, using tractable HNSCC models. Using both gain- and loss-of-function studies we report that LH2 is a key regulator of HNSCC cell migration, invasion and lymph node metastasis and that stable LH2-modified collagen cross-links are both necessary and sufficient to augment these invasive behaviors and promote metastatic dissemination.

Materials and methods

Cell culture

All cell lines used in this study are of human origin. “Normal” immortalized keratinocyte cell lines OKF4-TERT and OKF6-TERT cells were kindly provided by Drs. Jim Rheinwald and Matthew Ramsey at Harvard University [13] and cultured in Keratinocyte Serum Free media (KSFM media; Gibco, cat#: 17-005-042) supplemented with 25 ug/mL of Bovine pituitary extract (Sigma-Aldrich, cat#: P1167-5MG) + 1x Penicillin-Streptomycin-Glutamine (PSG; Thermo fisher, cat#: 10378-016) + 0.2 ng/mL Epithelial Growth Factor (Sigma-Aldrich, cat#: E9644) + 0.3 mM Calcium Chloride. The GMSM-K normal immortalized keratinocyte cell line was kindly provided by Dr. Valerie Murrah at the University of North Carolina-Chapel Hill [18] and cultured in KSFM supplemented with 50 ug/mL of Bovine pituitary extract + 1x Penicillin-Streptomycin-Glutamine + 50 ng/mL Epithelial Growth Factor. The UM-SCC-5, UM-SCC-11A, UM-SCC-14A, and UM-SCC-74A cells were kindly provided by Dr. Thomas Carey at the University of Michigan-Ann Arbor [5] and cultured in Dulbecco’s modified Eagle’s medium (Gibco, cat#: 11965-118) supplemented with 10% Heat-inactivated Fetal Bovine Serum (Atlanta Biologicals, cat#: S11550) + 1x Glutamax (Gibco, cat#: 35050061) + 1x Nonessential amino acids (Gibco, cat#: 11140050) + 1x PSG. The FBS was heat inactivated by a 30-min incubation at 55°C. The UM-SCC-15 and UM-SCC-25 cells, also obtained from Dr. Thomas Carey at the University of Michigan-Ann Arbor [5], were cultured in DMEM/F12 medium (Gibco, cat#: 11320033) supplemented with 10% FBS + 15 mM HEPES (Corning, cat#: 25-060-CI) + 0.5 mM Sodium Pyruvate (LifeTech, cat#: 11360070) + 400 ng/mL Hydrocortisone (Sigma-Aldrich, cat#: H0888). The UM-SCC-47, UM-SCC-2, SCC-090 and 93-VU-147T cells were kindly provided by Dr. Randall Kimple at the University of Wisconsin and cultured in DMEM supplemented with 10% Fetal Bovine Serum + 1x Glutamax + 1x PSG. STR profiling (LabCorp DNA Identity) were performed on the OKF6-TERT, UM-SCC-5, and UM-SCC-74A cell lines to confirm their authenticity.

qPCR

RNA was isolated using the NucleoSpin RNA kit (Macherey-Nagel, #740955) following the manufacturer's protocol. RNA was eluted in RNase-free water and quantified using a Cytation 5 plate multimode reader spectrophotometer. cDNA was synthesized using the iScript cDNA Synthesis Kit (BioRad cat, #: 170-8891). Quantitative RT-PCR was performed using FastStart Universal SYBR Green Master (Roche cat, #: 04913850001), using 2 μ L of cDNA (10 ng/ μ L) per reaction. qPCR primers used were as follows:

LH1:

sense: 5'-GAGCAGCATCCCTCGTTTCT-3'

antisense: 5'-AACTGAGCTGAGCGCTTGAA-3'

endogenous LH 2:

sense: 5'-CTCCCAAAGCTAAGTG CAGG-3'

antisense: 5'-TCAGCGCGGACGCTATTT-3'

exogenous by overexpressed LH 2:

sense: 5'-GACCCGAGGAAGTGCTGAAG-3'

antisense: 5'-CTGTTCGCCAGTCTCTTGT -3'

LH3:

sense: 5'-ATACGCTGACCGGGAGGATA-3'

antisense: 5'-CTGCCACTCTGGACGAACCT-3'

LOX:

sense: 5'-CCTACTACATCCAGGCGTCC-3'

antisense: 5'-GCCCTGTATGCTGTACTGGC-3'

LOXL2:

sense: 5'-CACCTGCGAGAATGGGCTAC-3'

antisense: 5'-GGGGTTGCTCTGGCTTGTA-3'

RPL23:

sense: 5'-TGATGGCCACAGTCAAGAAA-3'

antisense: 5'-ACACGCCATCTTTTCTACGG-3'

LH2 isoform-specific PCR

mRNA was extracted from GSMK, UM-SCC-5 and UM-SCC-74A cells using the NucleoSpin RNA kit (Macherey-Nagel, #740955) following the manufacturer's protocol. RNA was eluted in RNase-free water and quantified using a Cytation 5 plate multimode reader spectrophotometer. cDNA was synthesized using the SuperScript IV Synthesis Kit (BioRad cat, #: 170-8891) using oligo-dT primers. LH2 was amplified via PCR using previously validated and published isoform-specific primers and quantified as described [21].

LH 2 a versus LH2b isoform:

sense: 5'-CGATCAGAGATGAATGAAAGG-3'

antisense: 5'-GCAGTGGATAATAGCCTTCC-3'

Western blotting

Cells were lysed (250 mM NaCl, 50 mM Tris pH7.4, 50 mM NaF, 0.1 mM NaVO₄, 5 mM EDTA, 0.1% Triton-X, and protease and phosphatase inhibitors) and total protein concentration was quantified by BCA assay (Thermo Scientific, cat#: 23225). SDS-PAGE electrophoresis was performed using ~20 μ g protein. Blots were incubated with primary antibodies diluted in 5% skim milk/TBST overnight at 4°C. Horseradish peroxidase-conjugated secondary antibody (#7074, Cell Signaling Technology, USA) was diluted 1:10,000 in 5% skim milk/TBST and incubated for 1 h at room temperature. Protein bands were visualized using Clarity Western ECL Substrate (BioRad, USA) and analyzed with ImageJ 4.7v software (National Institutes of Health, Bethesda, MD, USA). anti-LH2 polyclonal antibody (1:100 dilution, Proteintech Group, USA), β -actin (1:10000 dilution, Sigma-Aldrich, USA).

Wound healing assay

Cells were seeded in a 6 well plate in media with or without 1 μ g/mL doxycycline and cultured for 3 d (UM-SCC-5) or 7 d (UM-SCC-74A) to ensure maximal overexpression/knockdown, respectively. Wells were washed and fresh media/Dox added daily. Cells were then trypsinized, counted and seeded for the wound healing assay. Ibidi 2 well culture inserts (Ibidi, cat#: 81176) were placed in a 24 well dish and cells were seeded (UM-SCC-5= 100,000 per side and UM-SCC-74A= 30,000 per side) in a 100 μ L volume on each side of the insert. 24 h postseeding, once cells had formed a 100% confluent monolayer, the Ibidi culture inserts were gently and carefully removed from the wells. Wells were washed once with 1xDPBS and 1 mL of fresh media with/without doxycycline was added. Wound healing/scratch closing was imaged once every hour over a period of 24 h using a Cytation 5 plate imager (BioTek Instruments, Inc). Three independent fields of view were imaged per well for each biologic replicate. Wound size was quantified manually using the area measurement function in ImageJ software.

Transwell Matrigel Invasion Assay (Boyden chamber)

Cells were seeded in a 6 well plate in media with or without 1 μ g/mL doxycycline. Cells were cultured for 2 d (UM-SCC-14A), 3 d (UM-SCC-5) or 7 d (UM-SCC-74A) to ensure maximal overexpression/knockdown, respectively. Wells were washed and fresh media/Dox added daily. Cells were then trypsinized, washed, counted, and resuspended in low-serum media (regular culture media without FBS, supplemented with 0.1% BSA). Prior to seeding cells onto the transwell chamber (Falcon permeable-transwell chambers- 8 μ m pore membrane, cat#: 353097), the transwell membrane was first coated with 5 % Matrigel (Corning, cat#: 356234). Briefly, 100 μ L of 5 % Matrigel solution was added to the transwell chamber, which was then incubated for 1 h in a humidified 37C, 5% CO₂ cell culture incubator to ensure polymerization. Postpolymerization, the transwell chamber was placed in a 24 well plate well containing 750 μ L of complete UM-SCC cell culture media (chemoattractant). Cells were seeded (UM-SCC-5 = 500,000/transwell, UM-SCC-14A = 125,000/transwell, or UM-SCC-74A = 100,000/transwell) into the transwell chamber in 1 mL of low-serum media and allowed to invade through the Matrigel and onto the bottom side of the transwell membrane for 24 h. Twenty-four hours postseeding, cells were fixed using 10% formalin, 5 min at room temperature. Cells were visualized using 0.05% crystal violet, 30 min incubation at room temperature followed by 5 washes using distilled water. Noninvaded cells still adhered to the 'top-side' of the transwell chamber were removed using cotton swabs. Membranes were dried overnight, and invaded cells were imaged in brightfield using a Cytation 5 plate imager. In order to accurately quantify the number of fully invaded cells, we sought to quantify the number of nuclei, instead of performing the quantification based solely on the crystal violet staining. Cells were de-stained by incubating in 1% Triton-X100 for 45 min at room temperature. Nuclei were stained with DAPI (0.5 μ M in 1xDPBS, for 15 min) and imaged using a Cytation 5 plate imager. Number of nuclei was counted across at least 5 random fields of view using the Cell Counter plugin in ImageJ software.

Plasmids

The pRRL-rtTA3 lentiviral vector was kindly provided by J. Zuber, Research Institute of Molecular Pathology, Vienna, Austria, TRE-KRAB-dCas9-IRES-BFP was a gift from Eric Lander (Addgene plasmid # 85449), pgRNA-CKB vector for guide RNA expression was a gift from Bruce Conklin (Addgene plasmid # 73501), and Hs-LH2 plasmid was a gift from Gavin Wright (Addgene plasmid # 51756).

Doxycycline-inducible LH2 retrovirus cloning

Generating the RetroX-TRE3G::MCS_PGK::GpNLuc vector

First, the 2nd generation pTight promoter from the RetroX-pTight::MCS_PGK::GpNLuc (Addgene plasmid # 70185) was replaced with a 3rd generation doxycycline response promoter; TRE3G. Briefly, the pTight promoter was excised using BamHI/XhoI restriction enzymes (New England Biosciences, cat#; R3136 and R0146, respectively). The BamHI digested vector was blunted using T4 polymerase (NEB, cat#; M0203) prior to the XhoI digestion in order to generate a blunt end/sticky end vector backbone. The dually digested plasmid was run in a 1% agarose gel and the digested backbone fragment (~7150 bp) was excised and purified (Macherey-Nagel PCR clean up kit, cat. #: 740609). Next, the TRE3G promoter was excised from a previously published plasmid (TRE-KRAB-dCas9-IRES-BFP, Addgene plasmid # 85449) using SmaI/XhoI restriction enzymes (NEB, cat#; R0141). The digested vector backbone and insert, 1:6 ratio, were ligated overnight using T4 DNA ligase (NEB, cat#; M202) and transformed into homemade (Zymo Research, cat#; T3001) chemically competent E.coli Stable3 cells (NEB, cat#; C304). Multiple clones were minipreped (Macherey-Nagel NucleoSpin kit, cat. #: 740588) and sequence verified (Eton Biosciences, Inc). All enzyme incubations were performed in accordance with the manufacturer's recommendations.

Generating the RetroX-TRE3G::HsLH2-T2A-Hygromycin_PGK::GpNLuc vector

The newly generated and sequence verified RetroX-TRE3G::MCS_PGK::GpNLuc vector was digested and dephosphorylated using BamHI/EcoRI (NEB, cat#; R3136 and R3101, respectively) and recombinant Shrimp Alkaline Phosphatase (NEB, cat#; M0371). The digested/dephosphorylated plasmid was column purified (Macherey-Nagel PCR clean up kit, cat. #: 740609). The LH2 cDNA insert was PCR amplified from a previously published vector (Addgene plasmid # 51756) [53] using Pfu Ultra DNA polymerase (Agilent, cat. #: 600380), and simultaneously, restriction enzyme sites 5'-BamHI-LH2-XbaI-3' were added into the amplicon using the PCR primers (*sense* = 5'-ctctcGGATCCggcgccaccatgg-3' + *antisense* = 5'-ctctcTCTAGAcgcccagggtcgaag-3'). The T2A-Hygromycin insert was PCR amplified from a previously published vector (Addgene plasmid # 154263) [35], and simultaneously, restriction enzyme sites 5'-XbaI-T2A-Hygromycin-EcoRI-3' were added into the amplicons using the PCR primers (*sense* = 5'-CTCCTCctagaATGGACGAGCTGTACAAGGGCT-3' + *antisense* = 5'-CTCCTCgaattcCTATTCCTTTGCCCTCGGACGAG-3'). The PCR amplicons were run in a 1% agarose gel and the appropriate bands (LH2 = 2220 bp and T2A-Hygromycin = 1120 bp) were excised and purified. The PCR amplicons were digested with BamHI/XbaI or XbaI/EcoRI as indicated (NEB, XbaI-cat#; R0145), and the digestion reaction was column purified. The digested/dephosphorylated backbone and digested PCR amplicon inserts were ligated, 1:3:3 ratio, using T4 DNA ligase overnight and transformed into homemade chemically competent E.coli Stable3 cells. Multiple clones were minipreped and sequence verified (Eton Biosciences, Inc). All enzyme incubations were performed in accordance with the manufacturer's recommendations. These vectors have been deposited with Addgene (https://www.addgene.org/Antonio_Amelio/).

Design and cloning of PLOD2 gene promoter-targeting guide RNAs

PLOD2 guide RNA design

sgRNA for knocking down PLOD2 using CRISPR-interference (dCas9-KRAB) were designed using the CRISPR-ERA resource (<http://crispr-era.stanford.edu/>). The top 3 guide RNA sequences were cloned (see below) and validated for PLOD2 knockdown efficiency by qPCR and Western blotting.

PLOD2 targeting sgRNA 3

sense: 5'-TCAGCCGGCGGCCAATAGCC-3'

PLOD2 targeting sgRNA 4

sense: 5'-TGAGCAAACAGTCCAGACGT-3'

PLOD2 targeting sgRNA 5

sense: 5'-CTGAGCAAACAGTCCAGACG-3'

PLOD2 targeting guide RNA cloning

The pgRNA-CKB vector (Addgene plasmid # 73501) was used to constitutively express the guide RNAs. Guide RNAs were cloned into the pgRNA-CKB vector using previously published methods [30]. Briefly, pgRNA-CKB backbone was digested with Esp3I (NEB, cat#; R0734) and dephosphorylated using recombinant Shrimp Alkaline Phosphatase. The required guide RNAs were synthesized (Eton Biosciences, Inc) as oligonucleotides with Esp3I compatible overhangs (forward oligo overhang = 5'-TTGG—sgRNA, reverse oligo overhang = 5'-AAAC—sgRNA). Oligonucleotides (50 μ L reaction, 2 μ M Forward/Reverse oligo final) were phosphorylated using T4 Poly Nucleotide Kinase (NEB, cat#; M0201) and annealed. Annealing was performed by adding 50 μ M sodium chloride to the phosphorylated oligo reaction, denaturing the oligo mix via boiling (95°C for 10 min) followed by an extended, slow cooling step (~0.1°C/sec). The annealed and phosphorylated sgRNAs were diluted to a final concentration of 0.1 μ M. 100 ng of the digested/dephosphorylated pgRNA-CKB backbone was ligated with 1 μ L of the 0.1 μ M phosphorylated/annealed guide RNA oligos using T4 DNA ligase overnight and transformed into homemade chemically competent E. coli Stable3 cells. Multiple clones were minipreped and sequence verified (Eton Biosciences, Inc) using a 5'-gagatccagtttggttagtaccggg-3' sequencing primer. All enzyme incubations were performed in accordance with the manufacturer's recommendations. The various pgRNA-CKB-PLOD2 guide RNA vectors described in this study have been deposited with Addgene (https://www.addgene.org/Antonio_Amelio/).

Viral transductions (spinflection)

All virus transductions were performed using established spinflection protocols. Briefly, cells were seeded in 6 well/24 well plates. Twenty-four hours postseeding, wells were washed and fresh media was added (1.5 mL/500 μ L respectively). The required virus was diluted into fresh media (500 μ L/250 μ L respectively) and polybrene was added to a final concentration of 4 μ g/mL. Virus/polybrene mixture was gently mixed and added dropwise onto the cells. Plates were spininfected by centrifugation at 1200 g for 1.5 h at 30°C (Sorvall LYNX 4000, Thermo Scientific, cat#; 75006580). Plates were returned to a humidified 5% CO₂, 37°C cell culture incubator post spinflection. Cells were supplemented with 1 mL of fresh media 48 h postspinflection. Cells were split 72 h postspinflection and expanded for downstream applications (selection/FACS).

Generation and validation of doxycycline inducible LH2 overexpression cells

UM-SCC-5 and UM-SCC-14A cells were first transduced with a lentivirus encoding the doxycycline-responsive reverse tetracycline transactivator 3 protein (rtTA³). rtTA³ is constitutively expressed from this vector under the control of the EF1 α promoter. A positively transduced polyclonal population was obtained by applying a puromycin antibiotic selection and rtTA3 mRNA expression was confirmed via real-time PCR. Next, these cells were transduced via spinflection with the retrovirus containing the doxycycline inducible LH2 and constitutively expressed GpNLuc LumiFluor cassette (see above for TRE3G:LH2 plasmid cloning information). A positively transduced polyclonal population was obtained by FACS selection. Untransduced cells were used as a GFP-negative gating controls.

The doxycycline induced *LH2* overexpression was validated at both mRNA and protein levels. Cells were doxed (125 ng/mL-2 mg/mL doxycycline) over a period of 72 h, with fresh doxycycline added daily. Cells were harvested for mRNA (NucleoSpin RNA kit (Macherey-Nagel, cat#; 740955)) and protein (RIPA buffer- [150 mM sodium chloride, 1.0% NP-40, 1.0% sodium deoxycholate, 0.1% sodium dodecyl sulfate, 25 mM Tris, pH 7.6, 20 mM NaF, 1 MM EDTA, supplemented with protease inhibitors (cOmplete, EDTA-free Protease Inhibitor Cocktail - Roche, cat. #; 04693132001) and phosphatase inhibitors (phosphoSTOP, - Roche, cat. #; 10917400)]). Unmodified 'parental' UM-SCC-74A and UM-SCC-5 cells were used as positive/negative controls (high/low expressor, respectively).

Generation and validation of doxycycline inducible LH2 knockdown cells

Stable CRISPRi (dCas9-KRAB) UM-SCC-74A cells were generated by first transducing cells with a previously published lentivirus encoding constitutively expressed GpNLuc Lumifluor and puromycin selection cassettes (Addgene plasmid # 135935). A positively transduced polyclonal population was obtained by applying puromycin to the cells for 14 d. Next, these GpNLuc positive cells were simultaneously transduced with 2 additional lentiviruses, (1) encoding the doxycycline-responsive reverse tetracycline transactivator 3 protein ($rtTA^3$), constitutively expressed via an EF1a promoter, and (2) encoding the doxycycline induced dCas9-KRAB-IRES-BFP (Addgene plasmid # 85449) cassettes. Since the BFP protein is under the control of the doxycycline-responsive TRE3G promoter, only cells dually transduced with both $rtTA^3$ and dCas9-KRAB-IRES-BFP viruses will express BFP upon doxycycline treatment. Thus, transduced triple positive cells were selected by applying doxycycline treatment for 72 h, followed by FACS selection for the GFP+ve/BFP+ve cell population, and single GpNLuc stable cells were used as gating controls. Lastly, these triple positive UM-SCC-74A cells (GpNLuc/ $rtTA^3$ /dCas9-KRAB) were transduced with lentiviruses encoding the *LH2* targeting sgRNAs of interest. These *LH2* targeting sgRNAs expression vectors were generated from a previously published guide RNA vector (Addgene plasmid # 73501), which also contains a constitutively expressed 2xNLS-mKate2 fluorescent reporter and blasticidin antibiotic resistance cassettes. Positively transduced cells were selected via blasticidin antibiotic selection. The quadruple positive stable cells (GpNLuc/ $rtTA^3$ /dCas9-KRAB/guide RNA) were confirmed by fluorescent microscopy as being positive for GFP, BFP (doxycycline induced) and nuclear mKate2.

The doxycycline induced *LH2* knockdown was validated at both mRNA and protein levels. Cells were doxed for 7 d with 1 μ g/mL doxycycline, with fresh doxycycline added daily. Cells were harvested for mRNA (NucleoSpin RNA kit, Macherey-Nagel, cat#; 740955) and protein (RIPA buffer [150 mM sodium chloride, 1.0% NP-40, 1.0% sodium deoxycholate, 0.1% sodium dodecyl sulfate, 25 mM Tris, pH 7.6, 20 mM NaF, 1 MM EDTA, supplemented with protease inhibitors (cOmplete, EDTA-free Protease Inhibitor Cocktail, Roche; cat.# 04693132001) and phosphatase inhibitors (phosphoSTOP, Roche, cat#; 10917400)]). Unmodified UM-SCC-74A and UM-SCC-5 cells were used as positive/negative controls (high/low expressor, respectively).

In vivo orthotopic xenograft

Female nude mice (6–10 wk old) were injected with 10^5 trypsin-dissociated single cells in a total volume of 30 μ L (15 μ L Hanks Balanced Salt Solution + 15 μ L Matrigel) into the left lateral tongue. Mice were kept anesthetized during the procedure (2–2.5% isoflurane gas). Doxycycline was administered via chow (1000 ppm Doxy Diet, Envigo, cat# TD.180096). Animals in the Dox cohort were prefed Dox chow for 3 d prior to orthotopic implantation and subsequently maintained on the Dox chow diet for the course of this experiment. Endpoint was determined by a

more than 20% loss of their maximum body weight or if the animals had become moribund. To determine tumor volume by external caliper, the greatest longitudinal diameter (length) and the greatest transverse diameter (width) were determined. We defined tumor volume as $1/2(\text{length} \times \text{width}^2)$. Consequently, lymph nodes, submaxillary gland, lung and liver were harvested and placed into 12 or 24 well plate to perform *ex vivo* analysis. This research was approved by The University of North Carolina at Chapel Hill Institutional Animal Care and Use Committee (Protocol No.17-202).

In vivo longitudinal bioluminescent imaging

Bioluminescent-fluorescent BRET signal was quantified noninvasively as previously described [49] with slight modifications. Briefly, animals were i.p. injected with 250 μ M (1:20 dilution, \sim 500 μ g/kg) Nano-Glo Luciferase Assay Substrate (Promega, cat. #N1120) in sterile DPBS. Isoflurane-anesthetized (3%) animals were imaged using the AMI Optical Imager (Spectral Instruments Imaging, Inc.) 5 min after injection. Mice were imaged using the following camera settings: Exposure time= 5 min, Binning= Medium, *F*-stop= 1.2, Field of view= 25. Data were analyzed using the Aura software package (<https://spectralin vivo.com/software/>).

Collagen biochemical analysis

After harvesting tongue tumors from each mouse, tissues were pulverized in liquid nitrogen, washed with cold PBS and distilled water, and lyophilized. Dried samples (\sim 3.0 mg each) were reduced with standardized NaB^3H_4 , hydrolyzed with 6N HCl and subjected to amino acid and cross-link analyses as previously reported [64]. Collagen composition (% of collagen in total proteins) was determined based on the value of 100 hydroxyproline (Hyp) residues per 1000 total amino acids in collagen. The level of Hyl per collagen was determined by Hyl/HypX300. The reducible cross-links, i.e. dehydrodihydroxylysine/its ketoamine (deH-DHLNL), dehydrohydroxylysine/its ketoamine (deH-HLNL) and dehydrohistidinohydroxymerodesmosine (deH-HHMD) were analyzed as their reduced forms, i.e. DHLNL, HLNL, and HHMD, respectively. The nonreducible, stable cross-links, pyridinoline (Pyr) and deoxypyridinoline (d-Pyr), were simultaneously analyzed by their specific fluorescence. The total number of aldehydes was calculated as $\text{DHLNL} + \text{HLNL} + 2 \times \text{Pyr} + 2 \times \text{d-Pyr} + 2 \times \text{HHMD}$ and the ratio of Hyl^{ald} - derived collagen cross-links (HLCCs) to Lys^{ald} derived collagen cross-links (LCCs) was calculated as $(\text{DHLNL} + \text{Pyr} + \text{d-Pyr})/\text{HHMD}$ as previously reported [9].

Histological analysis

Harvested tissues were fixed in 10% buffered formalin solution for at least 72 hrs. Following fixation, tissues were processed on an ASP6025 automated tissue processor (Leica Biosystems). Paraffin embedded tissue blocks were sectioned at 5 μ m, mounted on glass slides, and deparaffinized prior to Hematoxylin and Eosin (H&E) staining. Immunohistochemistry was performed on the Discovery Ultra (Ventana Medical Systems) using manufacturer's reagents. For *LH2* immunohistochemistry, anti-*LH2* (Proteintech #21214-1-AP) was prepared using Discovery PSS Diluent (cat. #: 760-212). Antigen retrieval was performed using Ventana's CC1 (pH 8.5) for 64 min at 90°C and slides were given a hydrogen peroxide block for 8 min at room temperature. Slides were then incubated in the primary antibody diluent (1:50) for 1 h at room temperature, followed by anti-Rabbit HRP secondary antibody for 32 min at room temperature.

Picrosirius red staining was performed to visualize collagen fibers as previously described [33, 48]. Briefly, FFPE sections were stained with Hematoxylin (Sigma-Aldrich, cat#; MHS32-1L) to visualize cell nuclei. Then, 0.1% Sirius red stain (Electron Microscopy Sciences, cat#; 26357-02), dissolved in saturated picric acid, was applied to FFPE sections for 1

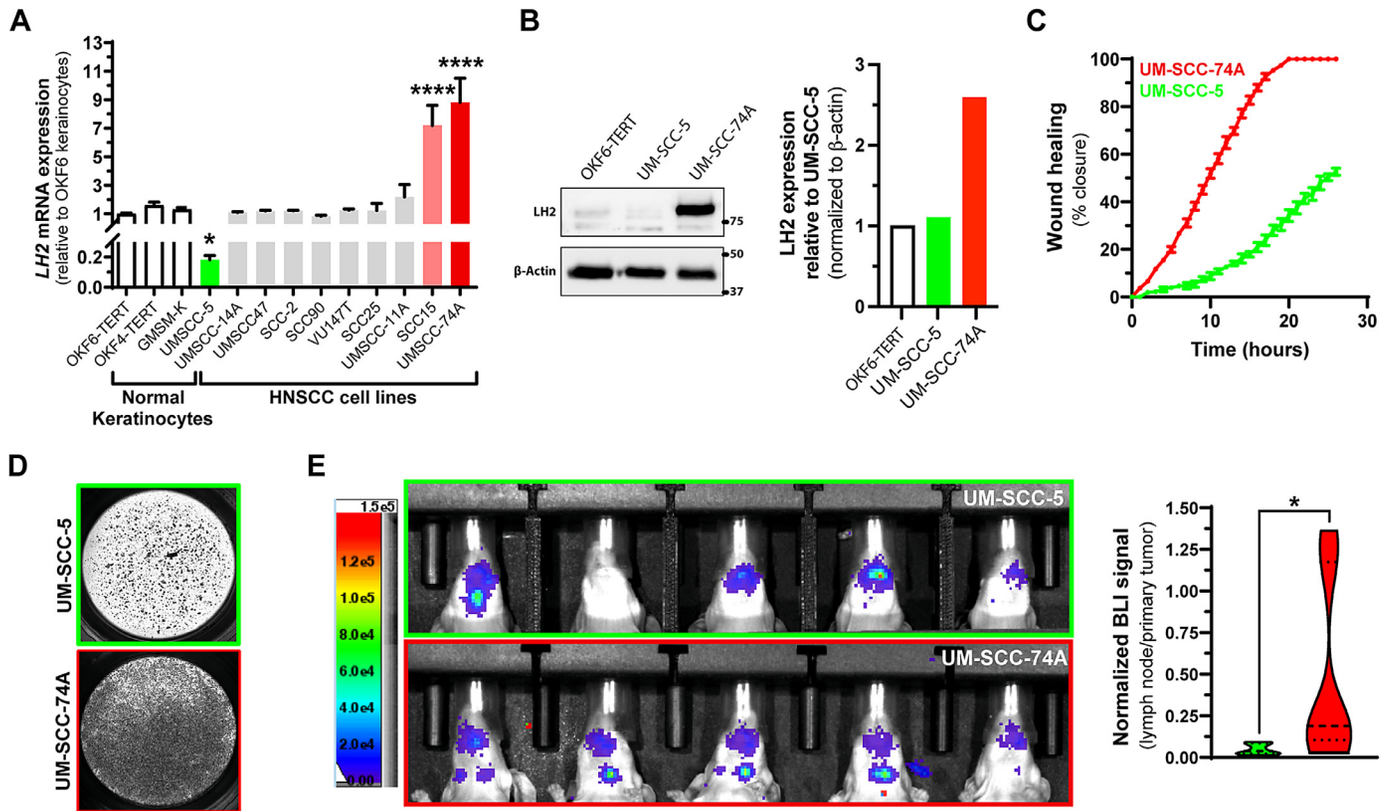


Fig. 1. LH2 expression levels in HNSCC cancer cell lines correlate with migratory and invasive potential. (A) Quantitative real-time PCR analysis of *LH2* mRNA levels in immortalized oral keratinocytes and HNSCC cell lines. *LH2* expression was normalized to *RPL23* mRNA levels and fold expression was calculated relative to the normal oral keratinocytes; OKF6-TERT cells. Data are presented as the mean \pm SEM ($n = 3$ biologic replicates; One-way ANOVA test, * $P < 0.05$, **** $P < 0.0001$). (B) *Left*, western blot showing LH2 protein expression levels in OKF6-TERT oral keratinocytes compared to UM-SCC-5 and UM-SCC-74A cells. This blot is representative of independent biologic replicates. *Right*, quantification of LH2 protein levels. Band intensities were normalized to β -actin and are shown as the fold change relative to LH2 expression in OKF6-TERT oral keratinocytes. (C) Wound healing (scratch) assay performed with LH2-high UM-SCC-74A versus LH2-low UM-SCC-5 cells. Images were captured every hour for 24 h to quantify rates of wound closure. Data are presented as the mean \pm SEM ($n = 3$ biological replicates, with 3 independent FOV per replicate). (D) Transwell invasion assays were performed for LH2-high UM-SCC-74A versus LH2-low UM-SCC-5 cells using Boyden chambers precoated with Matrigel. After 24 h, cells were stained with 0.05% crystal violet and DAPI, imaged, and counted to quantify rates of invasion. Representative data of independent biologic replicates are presented as the mean \pm SD (>5 independent FOV were quantified per biologic replicate). (E) *Left*, *in vivo* bioluminescent imaging (BLI) of LumiFluor-labeled LH2-high UM-SCC-74A versus LH2-low UM-SCC-5 cells 21 d postorthotopic tongue transplantation. *Right*, BLI signals were quantified by region of interest (ROI) analysis of images obtained at endpoint. Also see Supplementary Figure S1 for ROI definition.

h to stain the collagen fibers. Slides were washed with 0.01 N hydrochloric acid, dehydrated in 100% ethanol and xylene and mounted in resinous medium (DPX Mounting Medium, Electron Microscopy Sciences, cat# 13510). Collagen fibers were observed under polarized light microscopy. Immunohistochemistry was performed as previously described [6].

Statistical analysis

Statistical analyses were performed with GraphPad Prism (version 9) using student's *t*-test, one-way ANOVA or 2-way ANOVA as applicable. Data are presented as mean \pm SD or mean \pm SEM as indicated in the figure legends.

Results

Elevated LH2 correlates with enhanced HNSCC cell motility and metastasis

LH2 is amplified and transcriptionally upregulated in late-stage human primary HNSCCs and this elevated expression significantly correlates with

regional lymph node metastasis (RLNM) and poor patient outcome [48]. To determine if *LH2* levels are associated with invasion and metastasis *in vivo*, *LH2* expression in ten independent cell lines derived from oral and oropharyngeal sites of different patients, representative of both HPV(-) and HPV(+) subtypes, (Table S1) was compared to 3 independent normal control oral cell lines (nontransformed, immortalized oral keratinocytes). We found that *LH2* levels across a majority of HNSCC cell lines were overall comparable to oral keratinocytes (Fig. 1A). However, UM-SCC-5 cells were found to have significantly lower expression (~ 5 -fold decrease, $P < 0.05$) while SCC-15 and UM-SCC-74A cells displayed significantly higher mRNA expression (~ 8 -fold increase, $P < 0.0001$) relative to oral keratinocytes, a finding also consistent at the protein level in the UM-SCC-74A cells (Fig. 1A-B). Notably, isoform-specific analysis of *LH2* expression revealed that both UM-SCC-5 and UM-SCC-74A cells primarily express the LH2b (long) isoform (Figure S1C). In contrast to *LH2*, the closely related *LH1* and *LH3* family members displayed near-equivalent levels of expression between the UM-SCC-5 and UM-SCC-74A cells, and the expression of 2 additional collagen-modifying enzymes responsible for

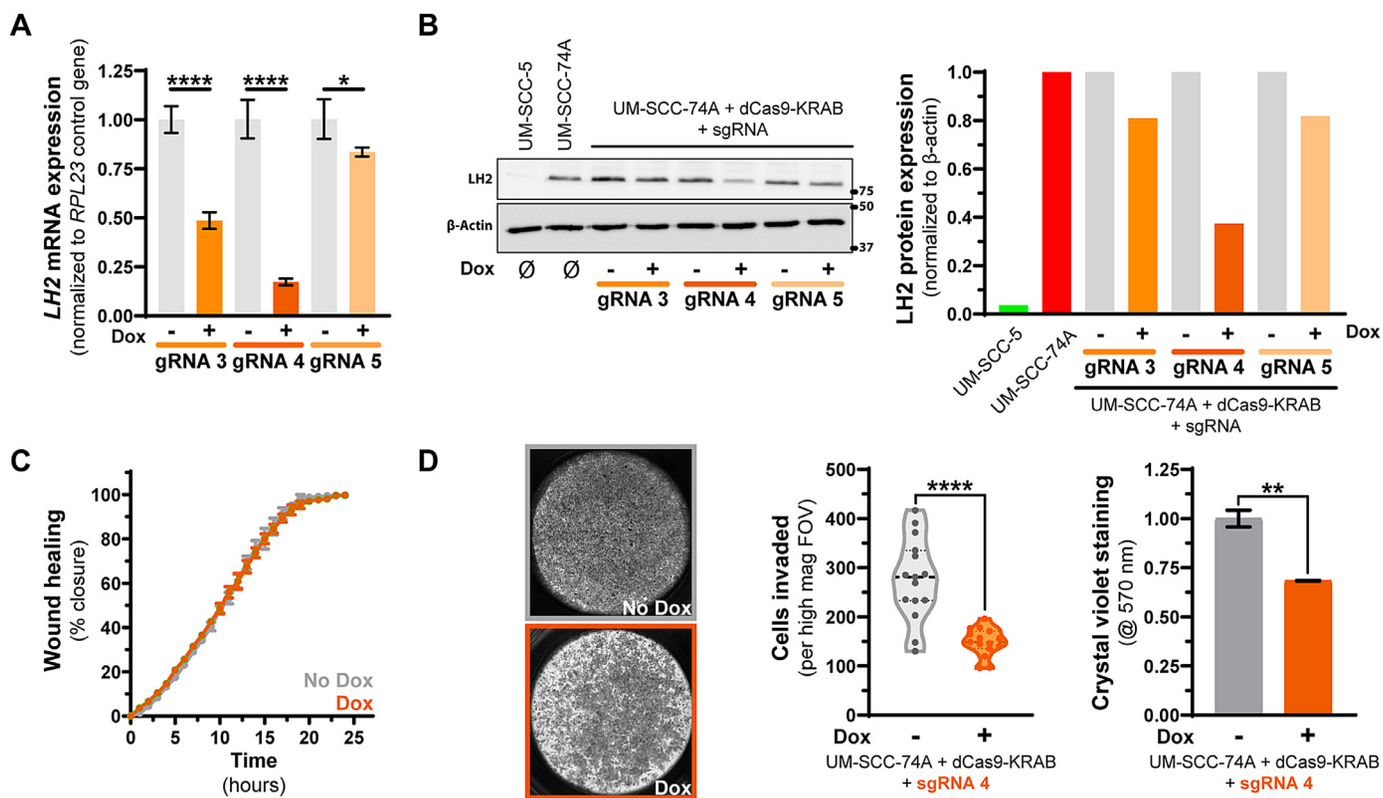


Fig. 2. Repressing *LH2* expression blunts HNSCC cell migration and invasion. (A) Evaluation of the CRISPRi system for inducible silencing of *LH2* expression. To validate 5 sgRNAs targeting the *PLOD2* gene promoter region, LH2-high UM-SCC-74A cells stably expressing doxycycline-inducible dCas9-KRAB were transduced with each sgRNA, cells were treated +/- 1 µg/mL doxycycline for 7 d, and quantitative real-time PCR was used to assess *LH2* mRNA levels. Representative data of independent biologic replicates are presented as the mean ± SD (4 technical replicates per biologic replicate; Student's *t* test, **P* < 0.05, *****P* < 0.0001). Also see Supplementary Figure 2A. (B) *Left*, UM-SCC-74A_TRE3G^{dCas9-KRAB}U6^{sgRNA} cells were treated +/- 1 µg/mL doxycycline for 7 d, total protein extracted, and western blot run to validate repression at the protein level. This blot is representative of independent biological replicates. *Right*, quantification of LH2 protein levels. Band intensities were normalized to β-actin and are shown as the fold change relative to the no doxycycline condition. (C) Wound healing (scratch) assay performed with the stable UM-SCC-74A_TRE3G^{dCas9-KRAB}U6^{sgRNA4} cells treated +/- 1 µg/mL doxycycline for 7 d. Images were captured every hour for 24 h to quantify rates of wound closure. Representative data of independent biologic replicates are presented as the mean ± SD (3 independent FOV were captured and quantified per biologic replicate). Also see Supplementary Figure 2B. (D) *Left*, transwell invasion assays were performed for UM-SCC-74A_TRE3G^{dCas9-KRAB}U6^{sgRNA4} cells using Boyden chambers precoated with Matrigel. After 24 h, cells were stained with 0.05% crystal violet, imaged, and cell invasion quantified. *Right*, quantification of invasion presented as either the number of cells invaded per field of view (FOV), or by extracting the crystal violet and analyzing absorbance at 570 nm. Representative data of independent biologic replicates are presented as the mean ± SD (>5 independent FOV were quantified per biologic replicate).

initiating collagen cross-linking, *LOX* and *LOXL2*, were also significantly increased (>10-fold, *P* < 0.001) but only in the UM-SCC-11A and UM-SCC-74A cells (Figure S1A-B). Since elevated *LH2* has been correlated with aggressive tumor behavior, we next examined the association between expression levels and cell behavior *in vitro* and *in vivo*. Analysis of 2D cell migration in wound healing assays confirmed that the higher *LH2* expression characteristic of UM-SCC-74A cells correlated with enhanced rates of wound closure (~75% increase) compared to UM-SCC-5 cells (Fig. 1C). Moreover, higher *LH2* levels also correlated with a dramatic increase in 3D cell invasion through Matrigel in transwell assays for UM-SCC-74A cells compared to UM-SCC-5 cells (Fig. 1D). We next investigated the ability of these cell lines to metastasize to regional lymph nodes using a previously established orthotopic tongue transplant model of HNSCC [23, 36, 40]. Quantitative bioluminescent imaging (BLI) of each cell line stably expressing our LumiFluor GpNLuc optical reporter [49] revealed that UM-SCC-74A cells rapidly and consistently develop RLNs compared to UM-SCC-5 cells *in vivo* (Fig. 1E and S1D).

LH2 is necessary for HNSCC invasion and metastatic dissemination

To directly test the functional role of *LH2* toward driving tumor cell migration, invasion and HNSCC disease dissemination, we next performed loss- and gain-of-function studies using the LH2-high (UMSCC-74A) and LH2-low (UMSCC-5) cell lines identified above, respectively. We engineered UM-SCC-74A cells (UM-SCC-74A_TRE3G^{dCas9-KRAB}U6^{sgRNA}) with an inducible CRISPRi repression system (dCas9-KRAB) [17] which allows for the controlled downregulation of *LH2* gene expression (Figure S2A). A panel of sgRNAs targeting the *LH2* promoter were designed and validated for efficiency of *LH2* repression at both the RNA and protein levels in stably transduced cell lines. In particular, sgRNA4 produced the most robust decrease (~80% and 60% decrease, respectively) in LH2 levels (Fig. 2A-B). Upon Dox-mediated CRISPRi repression, the UM-SCC-74 sgRNA4 stable cells displayed a significant decrease in invasive ability (*P* < 0.0001), however 2D cell migration was not affected by suppressing *LH2* (Figs. 2C-D and S2B). Beyond its role in collagen-I modification, *LH2* has also been shown to

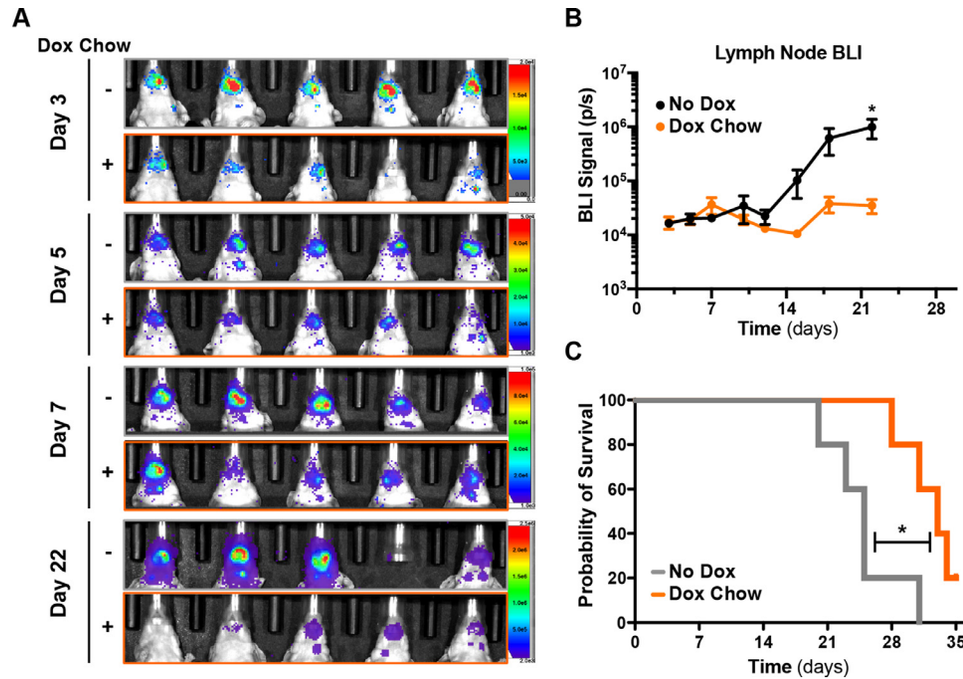


Fig. 3. Repressing *LH2* expression inhibits metastasis and improves overall survival. (A) Representative *in vivo* BLI images of LumiFluor-labeled UM-SCC-74A_TRE3G^{dCas9-KRAB}U6^{sgRNA4} cells at various times postorthotopic tongue xenograft transplantation in animals administered either normal or doxycycline chow. (B) Longitudinal quantification of metastatic lymph node BLI signals by ROI analysis of images obtained at various times. Detection of regional lymph node metastases was significantly reduced ($P < 0.05$) in *LH2* knockdown mice fed doxycycline compared to control chow animals. Also see Supplementary Figure S1 for ROI definition. (C) Kaplan-Meier analysis of animals administered either normal or doxycycline chow and monitored at various times postorthotopic tongue transplantation with UM-SCC-74A_TRE3G^{dCas9-KRAB}U6^{sgRNA4} cells. Tumor progression was significantly reduced ($P < 0.05$) in *LH2* knockdown mice fed doxycycline (median overall survival = 32 d) compared to control chow animals (median overall survival = 25 d).

play a critical role in integrin- $\beta 1$ activation in HNSCC cells [57], which may explain why *LH2* knockdown robustly retards tumor cell invasion but does not significantly affect 2D cell migration. Based on these findings, we next examined whether repressing *LH2* levels affects the efficiency of metastasis *in vivo* using the orthotopic tongue transplant model. Orthotopic xenografts were established with UM-SCC-74A_TRE3G^{dCas9-KRAB}U6^{sgRNA4} cells and mice were maintained on either normal or Dox chow to induce CRISPR-mediated repression of *LH2*. Knockdown of *LH2* *in vivo* led to a significant reduction in RLNM burden compared to the control group. (Figs. 3A-B). Moreover, *LH2* downregulation significantly prolonged disease-specific overall survival ($P < 0.05$) and decreased the onset of tumor-induced weight loss, which could be driven in part by a slight reduction in overall tumor volumes at endpoint (Fig. 3C and S3A-B).

LH2 overexpression is sufficient to enhance HNSCC cell migration, invasion and promote metastasis in vivo

Our loss-of-function findings indicated that *LH2* is necessary for promoting aggressive, invasive cell behavior *in vitro* and metastasis *in vivo*. Next, to determine if *LH2* expression is sufficient to regulate these processes, we engineered gain-of-function UM-SCC-5 cells (UM-SCC-5_PGK^{TetOn3G}TRE3G^{LH2}) with an inducible Tet-on 3G system (TRE3G; rtTA³) which enables precise, reversible, and efficient inducible control of ectopic *LH2* gene expression (Figure S4A). Dox dose-titration experiments with this cell line demonstrated robust >500-fold induction of *LH2* at both the RNA and protein levels in stably transduced cell lines using as little as 125 ng/mL Dox (Fig. 4A-B). Notably, the induced *LH2* protein levels were

comparable to levels observed in the highly metastatic UMSCC-74A cell line. *LH2* overexpression led to significantly faster wound closure ($\sim 2x$ faster, $P < 0.001$) and a significant increase ($P < 0.0001$) in 3D invasion in the dox-induced UM-SCC-5_PGK^{TetOn3G}TRE3G^{LH2} cells compared to control (Fig. 4C-D and S4B). Consequently, orthotopic xenografts established with these cells gave rise to robust RLNM (cervical lymph nodes) by 21 d post-transplant in animals administered Dox chow compared to animals maintained on a control diet (Fig. 5A). Remarkably, *ex vivo* BLI of surgically resected tissues confirmed the presence of local metastases to the cervical lymph nodes as well as distant metastases to the liver (Fig. 5B-C and S5B). Additionally, pan-cytokeratin staining confirmed the presence of lymph node metastasis only in the *LH2*-overexpressing SCC-5 tumors (Fig. 5C, right) Importantly, analysis of intraoral tumor volumes throughout the experiment and tumor volumes at endpoint demonstrate that *LH2* overexpression does not affect primary tumor growth rate or size (Fig. 5D and S5C). Collectively, these data indicate that, while *LH2* does not appear to play a major role in driving primary tumor growth and survival, its expression is necessary and sufficient to augment tumor cell invasive behaviors and promote metastatic dissemination *in vivo*.

LH2 upregulation changes collagen cross-link quality in the tumor ECM without affecting overall cross-link quantity

As described above, collagen cross-linking is initiated by the formation of Lys^{ald} or Hyl^{ald} in collagen telopeptides via the action of LOX and LOXL1-4 (Figure S6). Depending on the state of Lys hydroxylation in collagen telopeptides, 2 major cross-linking pathways are evolved: Lys^{ald}- and Hyl^{ald}-

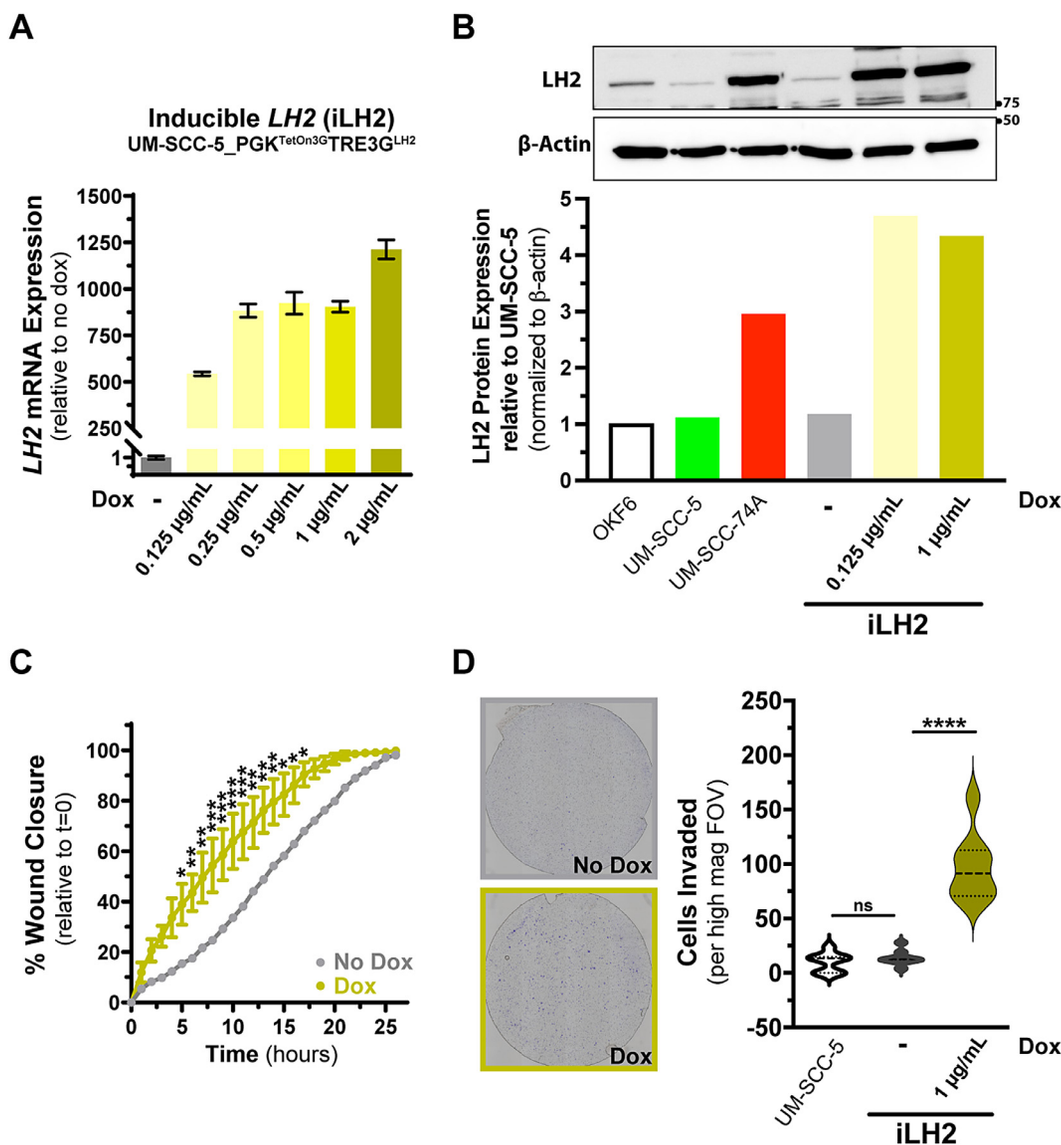


Fig. 4. Ectopic *LH2* overexpression in *LH2*-low cells augments HNSCC migration and invasion. (A) Evaluation of the doxycycline inducible Tet-On 3G system for inducing *LH2* expression. Quantitative real-time PCR analysis of *LH2* mRNA levels in UM-SCC-5_PGK^{TetOn3G}TRE3G^{LH2} cells treated with increasing concentrations of doxycycline for 72 hrs. Representative data of independent biologic replicates are presented as the mean ± SD (4 technical replicates per biologic replicate). (B) *Top*, UM-SCC-5_PGK^{TetOn3G}TRE3G^{LH2} cells were treated with the indicated concentrations of doxycycline for 72 h, total protein extracted, and western blot run to validate induced expression at the protein level. This blot is representative of independent biologic replicates. *Bottom*, quantification of *LH2* protein levels. Band intensities were normalized to β-actin and are shown as the fold change relative to the no doxycycline condition. (C) Wound healing (scratch) assay performed with UM-SCC-5_PGK^{TetOn3G}TRE3G^{LH2} cells treated +/- 1 μg/mL doxycycline for 72 h. Images were captured every hour for 24 h to quantify rates of wound closure. Representative data of independent biologic replicates are presented as the mean ± SD (3 independent FOV were captured and quantified per biologic replicate; Two-way ANOVA test with Sidak correction, **P* < 0.05, ***P* < 0.01, ****P* < 0.001). (D) Transwell invasion assays were performed for UM-SCC-5_PGK^{TetOn3G}TRE3G^{LH2} cells treated +/- doxycycline using Boyden chambers precoated with Matrigel. Cells were stained with 0.05% crystal violet after 24 h, imaged, and cell invasion quantified. Representative data of independent biologic replicates are presented as the mean ± SD (>5 independent FOV was quantified per biologic replicate; Student's *t* test, ns = not significant, *****P* < 0.0001).

derived cross-links [63], with the latter displaying enhanced stability over the former. Since *LH2* is responsible for Lys hydroxylation in telopeptides, its activity is key to determining the stability, that is, “quality”, of collagen cross-linking [8, 20, 61]. While *LH2* hyperactivity has been implicated in fibrosis for decades [59, 60], mounting evidence suggests that *LH2* upregulation can also drive proinvasive and prometastatic tumor behaviors across many different cancers.

To characterize the effect of *LH2* levels on the collagen cross-link profiles in our orthotopic HNSCC model, we analyzed primary tumors isolated from UM-SCC-5_PGK^{TetOn3G}TRE3G^{LH2}-transplanted tongues of animals administered either normal or Dox chow and performed histology and immunohistochemistry (Fig. 6A). Analysis of Picrosirius red staining under polarized light revealed that the amount of fibrillar collagens in animals administered Dox chow was significantly higher and more mature

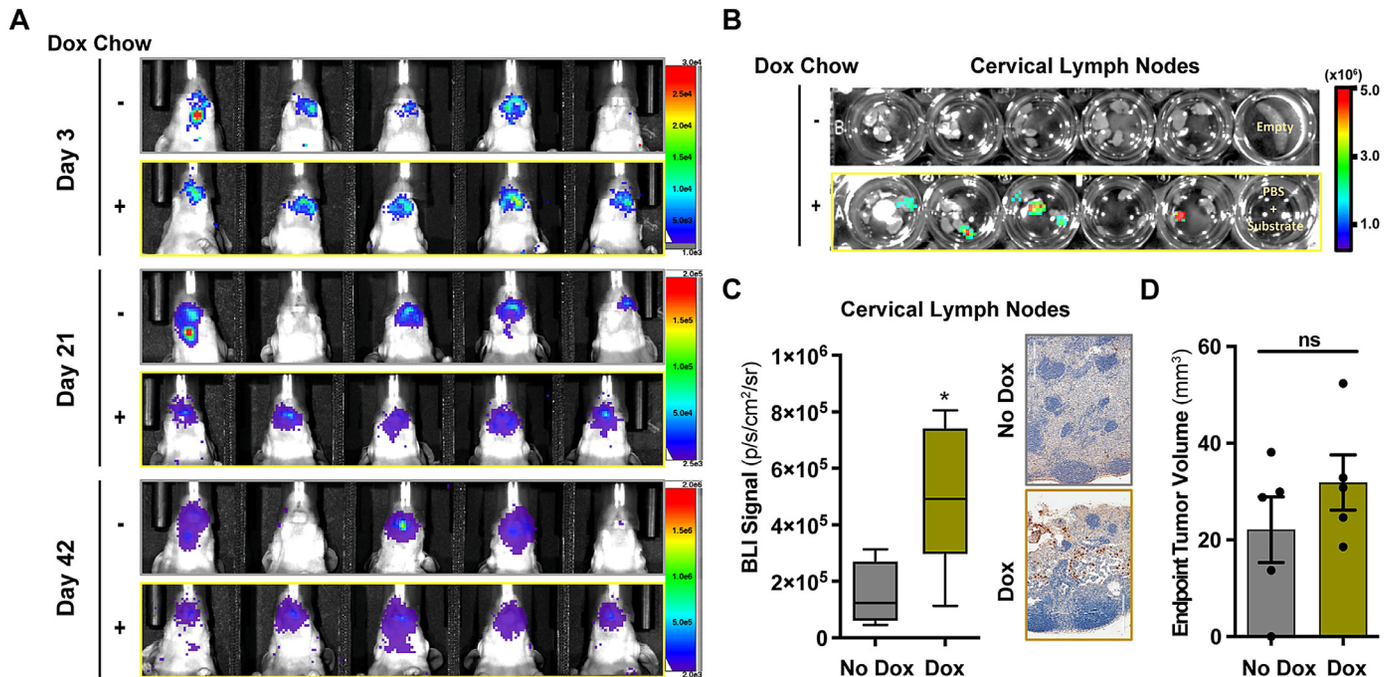


Fig. 5. Increased *LH2* expression is sufficient to drive HNSCC tumor metastasis.
(A) Representative *in vivo* BLI images of LumiFluor-labeled UM-SCC-5_PGK^{TetOn3G}TRE3G^{LH2} cells at various times postorthotopic tongue transplantation in animals administered either normal or doxycycline chow.
(B) Representative *ex vivo* BLI images of cervical lymph nodes resected from control- and doxycycline-chow fed animals at study endpoint.
(C) Quantification of metastatic lymph node *ex vivo* BLI signals by ROI analysis of images obtained at endpoint. Detection of regional lymph node metastases was significantly increased in induced *LH2* overexpression mice fed doxycycline compared to control chow animals. ($n = 5$ mice per group, mean \pm SEM; Student's t test, $P < 0.05$).
(D) Primary orthotopic tumor volumes are similar in control- and doxycycline-chow fed animals at study endpoint. ($n = 5$ mice per group, mean \pm SEM; Student's t test, ns = not significant).

as indicated by orange-bright red color compared to control (No Dox) diet animals (greenish yellow-orange) or nontransplanted normal tongues. Amino acid analysis revealed that collagen composition (collagen/total proteins) was identical (20–25%) across the normal, No Dox and Dox groups (Fig. 6B). The Hyl level per collagen was slightly but significantly higher in the No Dox group when compared to normal, which could be due to slight differences in tissue sampling (i.e., geographic location of the tumor tissue piece that was processed for analysis). Importantly, though, there was no difference between the normal and Dox, or the No Dox and Dox groups. However, quantitative collagen cross-link analysis revealed that the tumor tissue in the *LH2* overexpression group contained high *LH2*-mediated Hyl^{ald}-derived collagen cross-links (HLCCs) compared to control animals (Fig. 6C). Furthermore, the ratio of HLCC (DHLNL and HLNL cross-links) to Lys^{ald}-derived collagen cross-links (LCCs, HHMD) was significantly increased with *LH2* overexpression. HLNL was not included in the ratio since it can be derived from Lys^{ald}- or Hyl^{ald} [63]. However, considering that *LH2* levels are high in tumor tissues (i.e., telopeptidyl Lys hydroxylation), a significant amount of the HLNL in tumors is likely derived from the Hyl^{ald}. In this case, the ratio of HLCCs to LCC in tumors is expected to be even higher. Collectively, these findings confirm that the stable collagen cross-links mediated by increased *LH2* activity are responsible for generating a modified tumor ECM that promotes tumor metastasis.

Discussion

Invasive tumor phenotypes are responsible for the poor overall survival observed in patients with advanced HNSCC. These aggressive, metastatic

cases are often associated with a desmoplastic ECM characterized by excessive accumulation of fibrillar collagens [44, 62]. Utilizing a well-established orthotopic HNSCC model, we identified a functional role for *LH2*-mediated collagen modification in tumor progression to metastasis. Our findings corroborate previous studies indicating a role for collagen cross-linking in HNSCC disease severity [48] and mechanistically elaborate upon previous reports suggesting that *LH2*-mediated HLCC collagen cross-links are associated with metastatic dissemination. A recent report from Ueki and colleagues [57] found that elevated *LH2* expression activates Integrin β -1 driven cell migration, invasiveness and tumor dissemination of HNSCC tumors, further implicating *LH2* in regulating HNSCC metastasis and disease outcome. Thus, *LH2* levels may have prognostic value in predicting disease progression, while also serving as an actionable therapeutic target for prophylactic treatments aimed at preventing HNSCC tumor progression and adverse disease outcomes.

Accumulating evidence indicates that increased lysyl hydroxylase (*LH*) activity derived from both tumor intrinsic and extrinsic sources (e.g. cancer associated fibroblasts) is associated with tumor progression in many cancers [9, 38, 62]. While all *LH*s appear to be able to hydroxylate Lys residues in the helical domain of collagen, only *LH2* is capable of catalyzing hydroxylation of telopeptidyl Lys residues, a prerequisite to form Hyl^{ald}-derived stable collagen cross-links. Thus, there is significant interest in therapeutic strategies capable of inhibiting *LH2* activity and blunting tumor progression. Notably, *LH2* is part of a superfamily of oxygenases that require α -ketoglutarate (α KG) for their enzymatic activity. A recent high-throughput small molecule drug screen identified KD-1 (4,4,4-Trifluoro-1-(pyridine-3-yl)-1,3-butanedione) using a novel luminescence-based assay that measures *LH2* activity based on

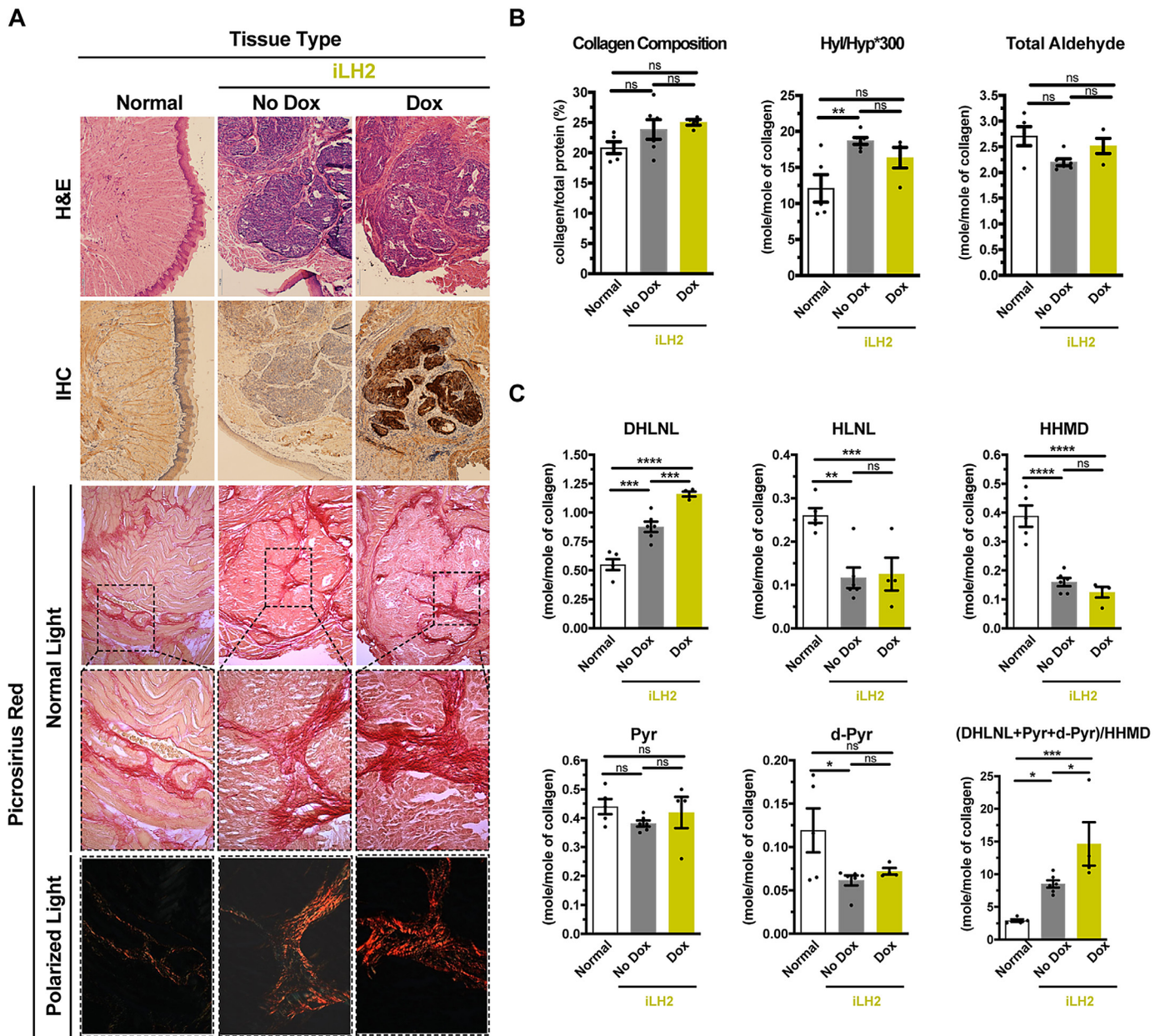


Fig. 6. LH2 upregulation in HNSCC cells remodels collagen cross-link quality but not total cross-link quantity in the tumor microenvironment. (A) Histologic and immunohistochemical analysis of LH2 levels and LH2-induced fibrillar collagen tracks. Representative coronal sections from resected tongues of animals transplanted with UM-SCC-5_PGK^{TetOn3G}TRE3G^{LH2} cells and administered either normal or doxycycline chow. Tissues were formalin fixed, embedded, and stained images of normal tongue versus SCC5 +/- LH2 overexpression primary tumors were captured for H&E, IHC staining for LH2, and Picrosirius Red staining for collagen fibrils. (B) Total collagen cross-link quantities in naïve normal tissues were compared to animals transplanted with UM-SCC-5_PGK^{TetOn3G}TRE3G^{LH2} cells and administered either normal or doxycycline chow. Quantification of total collagen cross-links, aldehyde levels, and the ratio of hydroxylysine (Hyl) to hydroxyproline content (Hyp/Hyp x 300). Data are presented as the mean ± SEM (*n* = 4-6 mice per group; One-way ANOVA with Fischer's LSD test, ns = not significant, ***P* < 0.01). (C) Analysis of collagen cross-link quality in naïve normal tissues were compared to animals transplanted with UM-SCC-5_PGK^{TetOn3G}TRE3G^{LH2} cells and administered either normal or doxycycline chow. Quantification of DHLNL, HLNL, HHMD, Pyr, d-Pyr and the ratios of DHLNL-to-HLNL, Pyr-to-dPyr, or HLCC-to-LCC [(DHLNL+Pyr+d-Pyr)/HHMD] were calculated. Data are presented as the mean ± SEM (*n* = 4-6 mice per group; One-way ANOVA with Fischer's LSD test, ns = not significant, **P* < 0.05, ***P* < 0.01, ****P* < 0.001, *****P* < 0.0001).

the amount of succinate produced from α KG. Since α KG is a rate-limiting metabolic intermediate of the TCA cycle, these results suggest that treatment of LH2-high tumors with KD-1 may lead to an accumulation of intratumoral α KG levels [12, 22]. Interestingly, recent work has demonstrated that reduced α KG levels are associated with the premalignant to malignant transition and interventions that lead to elevated α KG promote tumor cell differentiation and halt tumor progression [34]. Therefore, our findings have clear translational implications since strategies to inhibit LH2 may not only disrupt collagen cross-linking and tumor metastasis but may also lead to increased α KG levels that promote tumor cell differentiation and a less aggressive phenotype. Future studies aimed at investigating these exciting possibilities are warranted.

The TME is largely composed of ECM, whose composition, biochemical, and biophysical properties are known to influence the differentiation, proliferation, survival, migration, and invasion of tumor cells [25]. Fibrillar collagens such as types I and III, especially type I collagen [55], are the major constituents of solid tumor ECM. These collagens provide stability and tensile strength to tissues and therefore contribute to the stiffness observed in desmoplastic cancers such as HNSCCs. The tensile strength of collagen fibrils is achieved by formation of covalent intermolecular cross-linking initiated by LOX/LOXL1-4 [56]. However, the stability of collagen fibrils also depends on the quality/type of cross-links formed, not necessarily the quantity of cross-links alone. Collagens containing LH2-mediated stable cross-links are more resistant to degradation than those not modified by LH2 [8, 20, 61]. Moreover, such a stiffened and highly cross-linked tumor ECM may diminish accessibility of immune cells to the cancer cells within the TME. Immune exclusion from tumors can occur by several mechanisms including suppression of antigen processing machinery, impaired immune cell trafficking, impaired infiltration into tumors, poor T cell activation, and/or establishment of a PD-1/PD-L1 checkpoint [2, 3, 7]. Notably, tumor-associated hypoxia induces TGF- β 1, which not only promotes exclusion of T lymphocytes from tumors [31, 43, 54] but also functions as a key driver of LH2 and LOX/LOXL2 gene expression [15, 47], thus creating a direct association between tumor hypoxia, ECM stiffness, mechano-sensation [57], and immunosuppression [41].

Conclusions

The present study sheds new insight into the role that the tumor ECM – established by LH2-mediated cross-links – plays in promoting HNSCC metastasis. Several studies have suggested that increased total collagen cross-links resulting from upregulation of LOX family members are associated with poor prognosis in patients with breast, lung, and oral cancers [28, 42, 48, 50]. However, here we show that knockdown of LH2 alone is sufficient to blunt invasion and metastasis despite robust cellular expression of LOX and LOXL2. This indicates that the quality/type of collagen cross-links (i.e. stability), and not simply the total quantity of cross-links, is a necessary and crucial driver of HNSCC metastatic dissemination. Lastly, a limitation of our study is the number of HNSCC cell line models tested. However, given the striking nature of our findings and their potential disease relevance toward HNSCC patient outcomes, it is both prudent and urgent that these studies be further validated in additional independent HNSCC model systems.

Author contributions

Conception and design: K.P.-S., M.Y., and A.L.A.

Development of methodology: K.S., K.P.-S., M.T., and A.L.A.

Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): K.S., K.P.-S., A.M.M., M.T., and R.M.M.

Interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): K.S., K.P.-S., M.T., M.Y. and A.L.A.

Writing of the manuscript: K.S., K.P.-S., M.Y., and A.L.A.

Review and revision of the manuscript: K.S., K.P.-S., A.M.M., M.T., M.Y., and A.L.A.

Administrative, technical, or material support (i.e., reporting or organizing data, constructing databases): M.R.R., H.H., M.Y., and A.L.A.

Study supervision: M.Y. and A.L.A.

Acquisition of funding: A.L.A.

Declaration of competing interest

The authors declare no competing interests.

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Supplementary materials

Supplementary material associated with this article can be found, in the online version, at [doi:10.1016/j.neo.2021.05.014](https://doi.org/10.1016/j.neo.2021.05.014).

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