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Julien Jarroux, Dominika Foretek, Claire Bertrand, Marc Gabriel, Ugo Szachnowski, et al.. HOTAIR lncRNA promotes epithelial—mesenchymal transition by redistributing LSD1 at regulatory chromatin regions. EMBO Reports, In press, 10.15252/embr.202050193. hal-03222542

HAL Id: hal-03222542 https://hal.sorbonne-universite.fr/hal-03222542

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HOTAIR IncRNA promotes epithelial—mesenchymal transition by redistributing LSD1 at regulatory chromatin regions

Julien Jarroux^{1,†}, Dominika Foretek^{1,†}, Claire Bertrand^{1,†}, Marc Gabriel¹, Ugo Szachnowski¹, Zohra Saci¹, Shuling Guo², Arturo Londoño-Vallejo³, Marina Pinskaya^{1,*} & Antonin Morillon^{1,**}

Abstract

Epithelial-to-mesenchymal transition (EMT) describes the loss of epithelial traits and gain of mesenchymal traits by normal cells during development and by neoplastic cells during cancer metastasis. The long noncoding RNA HOTAIR triggers EMT, in part by serving as a scaffold for PRC2 and thus promoting repressive histone H3K27 methylation. In addition to PRC2, HOTAIR interacts with the LSD1 lysine demethylase, an epigenetic regulator of cell fate during development and differentiation, but little is known about the role of LSD1 in HOTAIR function during EMT. Here, we show that HOTAIR requires its LSD1-interacting domain, but not its PRC2-interacting domain, to promote the migration of epithelial cells. This activity is suppressed by LSD1 overexpression. LSD1-HOTAIR interactions induce partial reprogramming of the epithelial transcriptome altering LSD1 distribution at promoter and enhancer regions. Thus, we uncover an unexpected role of HOTAIR in EMT as an LSD1 decommissioning factor, counteracting its activity in the control of epithelial identity.

Keywords EMT; enhancer; HOTAIR; LSD1; promoter

Subject Categories Cancer; Chromatin, Transcription & Genomics; RNA Biol-

DOI 10.15252/embr.202050193 | Received 11 February 2020 | Revised 2 April 2021 | Accepted 12 April 2021

EMBO Reports (2021) e50193

Introduction

The epithelial-to-mesenchymal transition (EMT) allows normal or neoplastic cells to gradually lose their differentiated epithelial characteristics, including cell adhesion and polarity, and acquire mesenchymal traits, enabling cytoskeleton reorganization and motility (Lamouille et al, 2014). EMT is closely linked to tumor progression: It gradually endows epithelial cells with functions required for invasion and metastasis, as well as stem-like properties that contribute to tumor recurrence and drug resistance (Ye & Weinberg, 2015).

Epithelial-to-mesenchymal transition is a dynamic and reversible process driven by complex changes in signaling circuits and gene expression. Transcription factors such as zinc finger E-box-binding (ZEB1), SNAI2 (Slug), SNAI1 (Snail), and Twist are master regulators of EMT that coordinate the repression of epithelial genes and activation of mesenchymal genes (Lamouille et al, 2014). EMT is also influenced by epigenetic regulators, in particular histone lysine methyltransferases (Polycomb), histone deacetylases (NuRD), and histone demethylases (LSD1, PHF2). Through the control of chromatin modifications, these factors modify the expression of EMT genes, preventing or promoting the transition from an epithelial to a mesenchymal state (Tam & Weinberg, 2013).

Long noncoding (lnc)RNAs are emerging as regulators of EMT and metastasis (Huarte & Marín-Béjar, 2015; Richards et al, 2015; Shi et al, 2015; Liang et al, 2018). They use diverse mechanisms to modify gene expression, including interacting with proteins and DNA, serving as scaffolds, and recruiting chromatin-modifying complexes to specific genomic locations (Morlando et al, 2014; Quinn & Chang, 2015; Hendrickson et al, 2016). One of the bestknown examples is the lncRNA HOTAIR (HOX transcript antisense intergenic RNA), frequently overexpressed in cancer and associated with poor prognosis, metastasis, and stemness (Gupta et al, 2010; Kogo et al, 2011; Tsai et al, 2011; Zhang et al, 2015; Li et al, 2017). HOTAIR was first identified in human fibroblasts as a scaffold RNA, responsible for epigenetic regulation of cell fate during differentiation (Rinn et al, 2007). HOTAIR interacts with Polycomb repressive complex 2 (PRC2) to trigger histone H3 Lys27 (H3K27) methylation of EMT gene promoters (Gupta et al, 2010; Kogo et al, 2011). How it guides PRC2 to specific loci in distinct cell types is still unclear. Mechanisms based on triplex formation between HOTAIR and DNA or on anchoring the transcription factor SNAI1 have been described

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(Kalwa *et al*, 2016; Battistelli *et al*, 2017). However, PRC2 was recently reported to be dispensable for HOTAIR-mediated transcriptional repression in breast cancer cells (Portoso *et al*, 2017). This raises the question of how HOTAIR ensures the control of gene repression and whether PRC2 is required for HOTAIR function in promotion of EMT.

In addition to PRC2, HOTAIR can bind the LSD1/REST/CoREST complex (Tsai et al, 2010; Wu et al, 2013; Somarowthu et al, 2015). LSD1/KDM1A (lysine-specific demethylase-1) demethylates H3K4me2 and thus has been proposed to reinforce HOTAIR/PRC2mediated repression of transcription. However, whether LSD1 contributes to HOTAIR action is incompletely understood. LSD1 is a well-known epigenetic regulator of EMT and cancer. In few cases, LSD1 functions as a tumor suppressor (Wang et al, 2009), but mostly it acts as an oncogene (Lim et al, 2010; Harris et al, 2012; Schenk et al, 2012; Feng et al, 2016; Hino et al, 2016; Sun et al, 2016). This duality of LSD1 reflects its distinct substrates and interacting partners in different biological contexts (Shi et al, 2004; Metzger et al, 2005; McDonald et al, 2011). LSD1 controls the establishment of large organized heterochromatin H3K9 and H3K4 domains (LOCKs) across the genome during EMT of mouse hepatocytes, interacting either with REST/coREST co-repressors in differentiated epithelial cells or with transcriptional co-activators in cells undergoing EMT (McDonald et al, 2011). LSD1 has also been proposed to regulate non-histone targets, such as p53 and DNMT1, and to have non-enzymatic, scaffold roles, particularly in the regulation of enhancer activity in mammals (Wang et al, 2001; Huang et al, 2007; Lan et al, 2007a; Lan et al, 2007b; Scoumanne & Chen, 2007; Wissmann et al, 2007; Roth et al, 2016; Zeng et al, 2016; Maiques-Diaz et al, 2018). However, in the context of HOTAIR, whether and how LSD1 activity is controlled remained unknown.

Here, we investigated the role of the interaction between HOTAIR and LSD1 in EMT. We discovered that HOTAIR-mediated epithelial cell migration required the LSD1-interacting domain but not the PRC2-interacting domain. In addition, overexpression of LSD1 counteracted HOTAIR-induced phenotypic changes, indicating an epistatic crosstalk between both for the regulation of cell migration. Our data suggest that HOTAIR expression in epithelial cells affects LSD1 distribution at regulatory elements, resulting in transcriptome changes that favor mesenchymal traits. These findings further support a role for LSD1 as a guardian of epithelial identity and uncover a PRC2-independent function of HOTAIR in the acquisition of migratory properties by epithelial cells at the very early steps of carcinogenesis.

Results

The LSD1-interacting domain is essential for HOTAIR to promote cell migration

To uncover a potential role for the LSD1-interacting domain within HOTAIR, we generated a series of CMV expression vectors containing the full-length transcript (HOT), a truncation of the first 300 nucleotides (HOT Δ P), disrupting the HOTAIR interaction with PRC2 and a truncation of the last 658 nucleotides, removing the LSD1-interacting domain within HOTAIR (HOT Δ L) (Tsai *et al.*, 2010; Wu *et al.*, 2013) (Fig 1A). These constructs were transduced into an

epithelial cell line to generate stable cell lines through a drug selection of cell populations integrated the constructs constitutively expressing GFP (CTR), HOT, HOT Δ P, and HOT Δ L. Specifically, we used the immortalized human epithelial kidney cell line HA5-early (termed Epi), which has a normal karyotype and epithelial features, such as a rounded cobblestone morphology, low migration, and expression of epithelial markers (Zonula Occludens-1/ZO-1, β -Catenin, Claudin-1; Fig EV1A–C). For comparative studies, we also examined the mesenchymal cell line of the same origin, HA5-late (termed Mes), which expresses key mesenchymal markers and characterizes by high migration (Castro-Vega *et al*, 2013) and has higher endogenous levels of HOTAIR than Epi (Fig EV1A–C).

The levels of HOTAIR in Epi-HOT cells were at least 30 times higher than in Epi-CTR cells (Fig EV1D) and 10 times higher than in Mes cells (Fig EV1B). In addition, transduced Epi cells expressed at least two times higher HOT Δ L compared to the full-length HOTAIR or HOT Δ P (Fig EV1D). RNA immunoprecipitation (RIP) experiments confirmed that truncation of the 5′ or 3′ extremity of HOTAIR resulted in disruption of HOTAIR association with EZH2 and LSD1, respectively (Fig EV1E). We also confirmed that the subcellular distributions of HOTAIR were similar in Mes cells naturally expressing HOTAIR and genetically modified Epi-HOT, -HOT Δ P, and -HOT Δ L cells; in particular, ectopically expressed HOTAIR was associated with chromatin (Fig EV1F). Moreover, the subcellular localization and levels of HOTAIR protein partners EZH2 and LSD1 in total extracts were comparable in all cell lines (Fig EV1G and H, respectively).

Alterations in HOTAIR expression (overexpression or depletion) in epithelial cells are associated with changes in cell migration (Ding et al, 2014; Dong & Hui, 2016; Deng et al, 2017). We used an in vitro wound healing assay to assess a role of HOTAIR in cell migration in our system. As expected, overexpression of HOTAIR promoted a migration of Epi cells (Fig 1B; Appendix Fig S1A), and reversibly, depletion of HOTAIR by antisense oligonucleotides from highly migratory Mes cells reduced cell migration (Fig 1C; Appendix Fig S1B). These results confirmed and reinforced the observations in diverse EMT systems on the role of HOTAIR in promotion of cell migration (Wu et al, 2014; Zheng et al, 2018; Xun et al, 2019). Unexpectedly, deletion of the PRC2-interacting domain within HOTAIR did not abolish the ability of HOTAIR to stimulate migration: Epi-HOTΔP cells migrated as fast as Epi-HOT cells (Fig 1B). In contrast, Epi-HOTΔL migratory properties were similar to the control epithelial cells Epi-CTR, both closing the wound slower than Epi-HOT (Fig 1B). Importantly, the population doubling rates of the cell lines were similar, suggesting that the differences in wound healing efficiencies are not due to cell growth changes (Appendix Fig S1C). The migration changes strongly suggested that Epi-HOTΔL cells maintain their epithelial identity (as Epi-CTR) and this regardless the expression of HOTAIR with the intact PRC2-interacting domain, whereas Epi-HOT and Epi-HOTΔP cells lose some epithelial traits. Both, full-length HOTAIR and HOT Δ P are able to interact with LSD1, suggesting that HOTAIR interaction with LSD1 somehow impedes LSD1 function in regulation of epithelialmesenchymal balance. To challenge this hypothesis, we overexpressed LSD1 in Epi cells in the presence or absence of HOTAIR (Epi-CTR-LSD1-OE and -HOT-LSD1-OE; Appendix Fig S2A and B). It should be noted that LSD1 overexpression resulted in a considerable increase of HOTAIR expression in both Epi-CTR and Epi-HOT backgrounds, that could be explained by direct or indirect effect of LSD1

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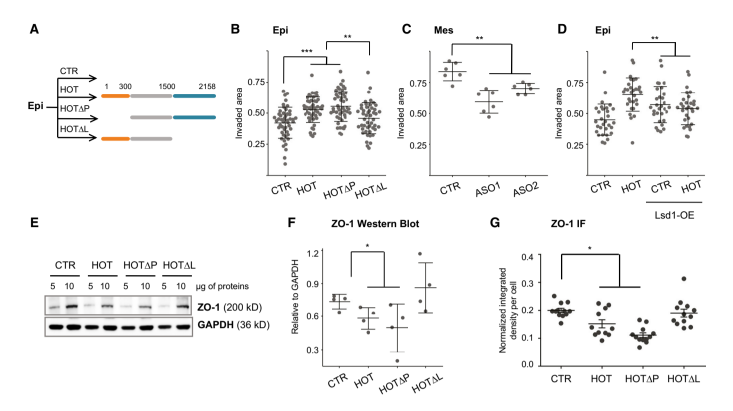


Figure 1. HOTAIR expression in epithelial Epi cells weakens tight junctions and promotes cell migration in LSD1-dependent manner.

- A Stable Epi cell lines ectopically expressing CTR (GFP), full-length HOTAIR (HOT), HOTAIR lacking PRC2- or LSD1-interacting domains, HOTΔP and HOTΔL, respectively.
- B–D Quantification of the wound area invaded in 24 h (gray dots) for Epi cells expressing CTR, HOT, HOTΔP or HOTΔP (B), or Mes cells transfected with ASO CTR (control), ASO1 and ASO2 targeting HOTAIR (C), or Epi cells overexpressing CTR or HOT alone or together with LSD1 (LSD1-OE) (D). Bars correspond to mean values and confidence interval (Cl) of 95%. ***P-value < 10⁻⁴, **P-value < 0.03 (Student *t*-test, *N* = 48 for (B), *N* = 6 for (C), *N* = 30 for (D)).
- E ZO-1 and GAPDH protein levels assessed in Epi cells by Western blot of whole protein extracts.
- F Quantification of ZO-1 levels (gray dots) in four independent Western blot experiments by ImageJ. *P-value < 0.05 (Student's t-test), bars correspond to mean ± standard deviation (N = 4).
- G ZO-1 quantification of IF images performed using the Fiji software and bar-plotted as normalized integrated densities per cell (gray dots) with bars corresponding to mean \pm standard error of the mean (SEM) for at least 11 high-field units of at least 100 cells; *P-value < 0.05 (Student's t-test, N = 11).

on transcription or stabilization of the lncRNA. We further examined the migration capacity of cell lines overexpressing or not LSD1 and full-length HOTAIR. In agreement with other reports, migration of epithelial cells was increased in the presence of high levels of LSD1 (+27%, Epi-CTR-LSD1-OE versus Epi-CTR; Fig 1D) (Boulding et al, 2018; Zhang et al, 2019). Remarkably, in this background, HOTAIR was not able to induce further increase in migration. Interestingly, overexpression of HOTAIR in Epi-HOT-LSD1-OE resulted in 12% loss of wound healing efficiency comparing to the Epi-HOT cell line overexpressing HOTAIR alone (P-value < 0.004, Student's ttest). We concluded that overexpression of LSD1 promoted migration committing epithelial cells to EMT. It can either withdraw, antagonize, or impede HOTAIR function in gain of migratory properties. And reversely, HOTAIR seems to impact negatively LSD1 capacity to stimulate migration. These findings reinforced our hypothesis of the crosstalk between HOTAIR and LSD1 in gene expression regulation involved in EMT.

Cell migration is associated with changes in cell-cell junctions, cytoskeletal organization, and apico-basal polarity of cells (Lamouille

et al, 2014). To investigate whether HOTAIR overexpression in epithelial cells affects the overall cell morphology, we performed phalloidin staining of F-actin fibers. However, we did not detect differences in cell shape or in the formation of cell sheets (Appendix Fig S1D). During EMT, cells acquire migratory properties at least in part due to a reduction in the tight junctions that support cell-to-cell contacts (Tornavaca et al, 2015). Therefore, we examined whether overexpression of HOTAIR affected the tight junction protein ZO-1/TJP1 in Epi cells. We found that ZO-1 protein levels were decreased in Epi-HOT and Epi-HOT ΔP cells compared to the less migratory Epi-CTR and Epi-HOTΔL cells (Fig 1E and F). In addition, ZO-1 displayed a more diffused localization, especially, at cellcell junctions, in Epi-HOT and Ep-HOTΔP compared to Epi-HOTΔL and Epi-CTR (Fig 1G; Appendix Fig S1E). In contrast, the protein levels of other epithelial markers, \u03b3-Catenin and Claudin-1, and mesenchymal markers, SNAI1/2, ZEB1, and Vimentin, were similar in all cell lines (*P*-value > 0.05, Student's *t*-test; Appendix Fig S3). Thus, the key transcription factors that reprogram epithelial cells toward a mesenchymal identity were unchanged in Epi cells

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expressing HOTAIR. Together, these findings suggested that HOTAIR-LSD1 interactions in Epi cells promote migration before all by weakening cell–cell junctions resulting in a mixed epithelial—mesenchymal phenotype.

HOTAIR-induced transcriptome changes depend distinctly on PRC2 and LSD1

To understand molecular mechanisms underlying the altered migratory properties of Epi cells expressing HOTAIR, we studied their transcriptome by RNA sequencing (RNA-seq). First, the transcriptome of Epi-CTR was compared to Epi cells expressing each of HOTAIR variants. We found that the ectopic expression of HOTAIR or its truncated versions in Epi cells induced global changes in mRNA levels, with an overall repressive effect (Fig 2A).

A total of 755 genes showed altered expression in Epi-HOT compared to Epi-CTR, with a fold change (FC) above 2 and adjusted P-value below 0.05 (Fig 2A and B, Dataset EV1). Of them, 505 were altered only by the full-length HOTAIR and hence dependent on both PRC2- and LSD1-interacting domains. We referred to this set of genes as HPL-pos (N=119) and HPL-neg (N=386) for HOTAIR-PRC2-LSD1 up- and downregulated genes (Fig 2B, Dataset EV2). Some of HPL genes were reported to be EMT markers (Gröger $et\ al$, 2012), while others were previously identified as repressed by HOTAIR- and PRC2-mediated histone H3 lysine 27 methylation (Gupta $et\ al$, 2010) (Gröger and Gupta sets, Dataset EV2), including genes involved in proteolysis of extracellular matrix, SERPIN2 and

MMP3, and the protocadherin gene family member *PCDH18* (Gupta *et al*, 2010; Xu *et al*, 2013; Qiu *et al*, 2014). Gene Ontology (GO) analysis revealed that the HPL genes are enriched for several Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways that are tightly linked to EMT, cancer, and metastasis (Fig 2C). The downregulated genes were enriched for extracellular matrix (ECM) receptor interactions, focal adhesion, and Hedgehog signaling, whereas the upregulated genes were enriched for Jak-STAT and bladder cancer pathways (Fig 2C). Notably, expression of EMT drivers and key EMT transcription factors was not stimulated by HOTAIR over-expression in epithelial cells.

On the other hand, 77 and nine protein-coding genes were downand upregulated, respectively, upon expression of any of HOTAIR variants (Fig 2B). These genes were grouped into a core-set of potential HOTAIR targets that do not require neither PRC2 nor LSD1 (Dataset EV2). According to GO analysis, these genes were involved in developmental processes, cell surface receptor signaling pathways, MAPK cascade, and apoptotic processes.

Finally, Epi cells expressing HOT Δ P or HOT Δ L variants showed less transcriptomic perturbations (R^2 of 0.986 and 0.982, respectively) than cells expressing full-length HOTAIR (R^2 of 0.972) (Appendix Fig S4). Only 194 and 353 protein-coding genes were altered in Epi-HOT Δ P and Epi-HOT Δ L, respectively (Fig 2A). Paradoxically, Epi-HOT Δ P cells with the lowest number of differentially expressed genes (N=194) still showed increased migration similar to Epi-HOT cells, while Epi-HOT Δ L cells with 353 differentially expressed genes lost their capacity to migrate (Fig 1C).

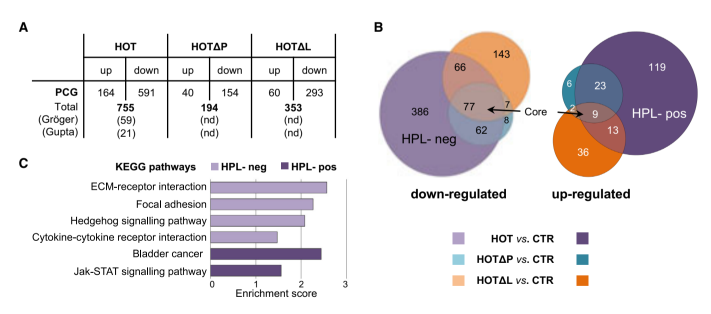


Figure 2. HOTAIR expression in Epi cells induces drastic changes in expression of protein-coding genes, majorly dependent on the presence of both, PRC2- and LSD1-interacting domains.

- A Number of up- and downregulated protein-coding genes (PCG) defined as differentially expressed in Epi-HOT, -HOTΔP, and -HOTΔP cells compared to Epi-CTR by RNA-seq and DESeq analysis (fold change > 2 and adj. *P*-value < 0.05), including those associated with EMT (Gröger set, Dataset EV2) and already identified as HOTAIR/PRC2 targets (Gupta set, Dataset EV2); nd stands for non-determined.
- B Venn diagram of intersection of down- and upregulated protein-coding genes in Epi-HOT, -HOTΔP, and -HOTΔL cells compared to Epi-CTR. HPL-neg and HPL-pos are gene sets down- and upregulated only by the full-length HOTAIR (Dataset EV2).
- C KEGG pathways identified by DAVID as significantly enriched (adj. *P*-value < 0.05) for HPL-neg and HPL-pos sets of protein-coding genes, ranked by enrichment score calculated as $-\log_{10}(P\text{-value})$.

Source data are available online for this figure.

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Together, our protein and global transcriptomic results strongly suggest that HOTAIR is not involved in full reprogramming of epithelial identity during EMT. Instead, HOTAIR modulates the epithelial–mesenchymal plasticity toward acquisition of some mesenchymal traits. Remarkably, PRC2- and LSD1-interacting domains within HOTAIR contribute together but also distinctly to its regulatory function. Specifically, truncation of the LSD1-interacting domain within HOTAIR still allows significant changes in gene expression but these perturbations are insufficient to preserve its capacity in promotion of migration.

The LSD1-interacting domain of HOTAIR shapes the high-migration transcriptome

To define highly expressed genes involved in the maintenance of epithelial properties in low migrating cells (Epi-CTR and -HOT Δ L) and, in contrast, highly expressed genes associated with the

acquisition of migration properties in high migrating cells (Epi-HOT and -HOT Δ P), we performed anew a differential expression analysis, comparing gene expression between these two phenotypically distinct groups. All genes, except those of the core-set, with upregulated expression (FC > 1.5 and adj. *P*-value < 0.05) in low-migrating cells were assigned as a *Low Migration Signature* (LMS), whereas genes with upregulated expression in high-migrating cells were assigned as a *High Migration Signature* (HMS) (Fig 3A and B, Dataset EV3). In total, we defined 131 LMS and 75 HMS genes presenting gene expression patterns specific to each phenotypic group (Dataset EV2).

Gene Ontology analysis showed that the LMS set is enriched for genes involved in pathways linked to cardiac development (as the integrin family), lipid biosynthesis, and cytokine–cytokine receptor interactions, whereas the HMS set includes genes of ECM–receptor interactions and focal adhesion pathways tightly linked to EMT, cancer, and metastasis (Fig 3C). Notably, transcriptome changes

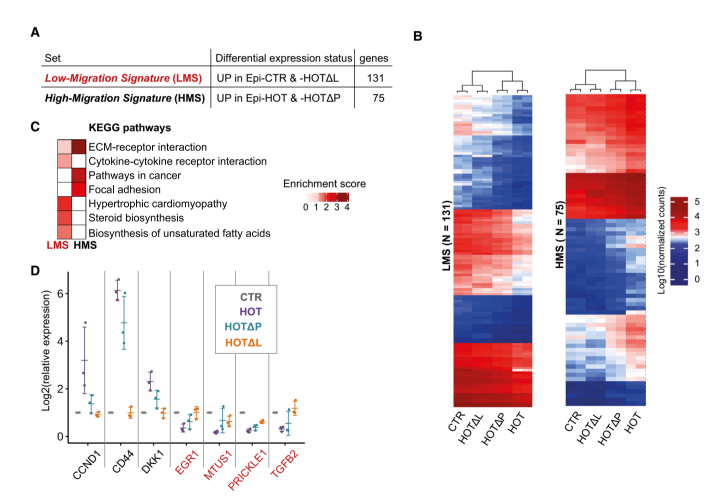


Figure 3. Transcriptome signatures of low- and fast-migrating epithelial cells.

- A Protein-coding gene assignment to Low and High Migration Signature sets according to differential expression features.
- B Heatmap of unsupervised hierarchical clustering based on Low Migration Signature (LMS) and High Migration Signature (HMS) gene expression.
- C Heatmap of enrichment scores (-log₁₀(P-value)) for KEGG pathways significantly enriched for LMS and HMS protein-coding genes (DAVID, P-value < 0.05).
- D LMS (red) and HMS (black) gene expression quantification in Epi cells by random-primed RT–qPCR relative to *RPL11* and represented as log₂ fold change to CTR. Each dot corresponds to an independent replicate; bars represent mean \pm standard deviation values.

Source data are available online for this figure.

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induced by HOTAIR did not result in a complete switch of the EMT program, but rather in a modulation (attenuation or increase) of gene expression (Fig 3B). HMS genes, highly expressed in fastmigrating Epi-HOT and -HOTΔP cells, included the cell cycle regulator CCND1 (Kim & Diehl, 2009) and the CD44 EMT marker (Zhao et al, 2016) (Fig 3D). Among LMS genes, highly expressed in Epi-CTR and -HOTAL cells, were the tumor suppressor MTUS1 (Di Benedetto et al, 2006) and the nuclear receptor PRICKLE1/RILP implicated in the nuclear trafficking of REST/NRSF and REST4 transcription repressors (Shimojo & Hersh, 2006) (Fig 3D). To compare epithelial and mesenchymal traits of Epi cells overexpressing HOTAIR, we also performed the RNA-seq analysis of protein-coding transcriptome of original Epi cells and their mesenchymal compeer-Mes cells. Primary epithelial kidney cells lose epithelial traits and acquire mesenchymal properties with time, through cell division and increase of genomic instability, hence undertaking EMT. When immortalized at different stages before (Epi) and after (Mes) transition, they possess very different transcriptome (Castro-Vega et al, 2013). This was confirmed in our study by RNA-seg and differential expression analysis: 1,274 and 1,423 protein-coding genes were upand downregulated, respectively, in Mes cells, underlying differences in cell phenotypes regarding cell morphology, migration, and expression of EMT markers (Dataset EV4, Appendix Fig S5A and B). However, these alterations were not strictly concurrent with those induced by HOTAIR overexpression in Epi cells reinforcing the idea that HOTAIR is not able to fully reprogram Epi cells to mesenchymal identity. The expression of key transcription factors driving EMT reprogramming, was not induced by HOTAIR in Epi cells, and globally, we did not observe significant enrichments of LMS or HMS groups in down- or upregulated genes from Mes. Only 30 genes downregulated in Mes were part of LMS, and 6 genes upregulated in Mes were found in the HMS group (Appendix Fig S5C), concurrently with low or high migration properties in each system.

In conclusion, the conversion of low- to high-migration phenotype is associated with modulation of epithelial transcriptome altering above all focal adhesion and ECM–receptor interactions. These perturbations depend on the presence of the LSD1-interacting domain within HOTAIR supporting the importance of LSD1 in maintaining epithelial identity and its altered activity during HOTAIR-induced EMT.

The LSD1-interacting domain of HOTAIR alters distribution of LSD1 at promoters

Epithelial-mesenchymal plasticity is determined by transcriptional program. Any change in transcription can cause or, in turn, can

result from epigenetic reconfiguration of chromatin at promoters and distal cis-regulatory elements. Thus, we decided to profile H3K27ac in Epi cells expressing HOTAIR variants. This histone modification is present at promoters of transcribed genes, reflecting transcriptional activity of the locus, but also it is a hallmark of active enhancers and super-enhancers in the genome (Creyghton et al, 2010; Hnisz et al, 2013). We used the recently established CUT&RUN technique (Skene et al, 2018) and applied the HMCan peak calling to profile H3K27ac peaks genome-wide. As expected, H3K27ac tags were specifically enriched in vicinity of transcription start sites (TSS) (Fig EV2A and B). Peaks within the 2-kb window around gene TSS were assigned to promoters. We observed slightly more H3K27ac peaks, and specifically at gene promoters, in lowmigrating Epi-CTR and -HOT L cells (Fig EV2A). Also, low-migrating cells showed more genes with H3K27ac peaks in common than the high-migrating Epi-HOT and -HOTΔP cells (433 and 32, respectively; Fig 4A). In addition, a metagene analysis of H3K27ac distribution around TSS showed global decrease of this mark in highmigrating cells (Fig 4B). Unsupervised hierarchical clustering based on H3K27 mean signal at promoters showed separation of low- from high-migrating cells (Fig 4C). To identify potential pathways which have been reprogrammed between these two phenotypic groups, we divided genes into three classes based on H3K27ac enrichment of their promoters: higher coverage in low-migrating CTR and HOT Δ L conditions (Class 1) or higher in high-migrating HOT and HOTΔP conditions (Class 2), and, finally, all other cases (Class 3; Fig EV2C). Comparisons of a fold change (FC) in H3K27ac density of each peak confirmed pairwise similarity (FC close to 1) between low-migrating Epi-CTR and -HOTAL and between high-migrating Epi-HOT and -HOTΔP cells (Fig EV2D). Class 1 contained far more hyperacetylated genes than Class 2 (2,268 against 136 genes, respectively). The GO term analysis revealed involvement of Class 1 genes in regulation of cell-cell adhesion, cell migration, and actin cytoskeleton whereas Class 2 genes showed poor significance of enrichments in biological processes and the top ranked were not relevant to EMT and cancer (Fig 4D). Globally, very few GO terms were in common between these two classes. This observation was concordant with differential expression and GO term analysis between low- and high-migrating cells (LMS and HMS; Fig 3C). However, comparison between H3K27ac levels at gene promoters and gene expression changes quantified by RNA-seq in Epi cells did not show high concordance (Appendix Fig S6). Notably, Class 1 genes with increased H3K27ac at promoters in low-migrating cells were not specifically ranked as upregulated by RNA-seq comparing to high-migrating cells and reverse. Only 32 genes were shared between LMS (N = 131) and Class 1 (N = 2,268) subgroups defined

Figure 4. HOTAIR expression changes H3K27 acetylation and LSD1 occupancy of gene promoters.

- A Venn diagram representing intersections of gene promoters possessing H3K27ac peaks within the 2-kb window around their TSS in Epi cells.
- B Metagene profiling within the 10-kb window around TSS of H3K27ac, LSD1, and H3K4me2 over IgG in Epi cells expressing none (CTR) or different HOTAIR variants (HOT, HOTΔP, and HOTΔL). bp stands for base pairs.
- C Heatmap of unsupervised hierarchical clustering based on H3K27ac mean values at gene promoters showing separation of two phenotypic groups.
- D Biological process pathways identified by DAVID as enriched for Class 1 or Class 2 genes hyperacetylated in low- or high-migrating cells, respectively (adj. *P*-value < 0.05 for Class 1 and *P*-value < 0.05 for Class 2). Dots represent GO term enrichment with color coding: red for high enrichment, blue for low enrichment, and sizes of the dots representing the significance score of each subset (the bigger the dot, the higher its significance).
- E Visualization of the H3K27ac, Lsd1, and H3K4me2 CUT&RUN and of RNA-seq tracks of the *TGFB2* promoter region in Epi-CTR, -HOT, -HOTΔP, and -HOTΔL cells. Source data are available online for this figure.

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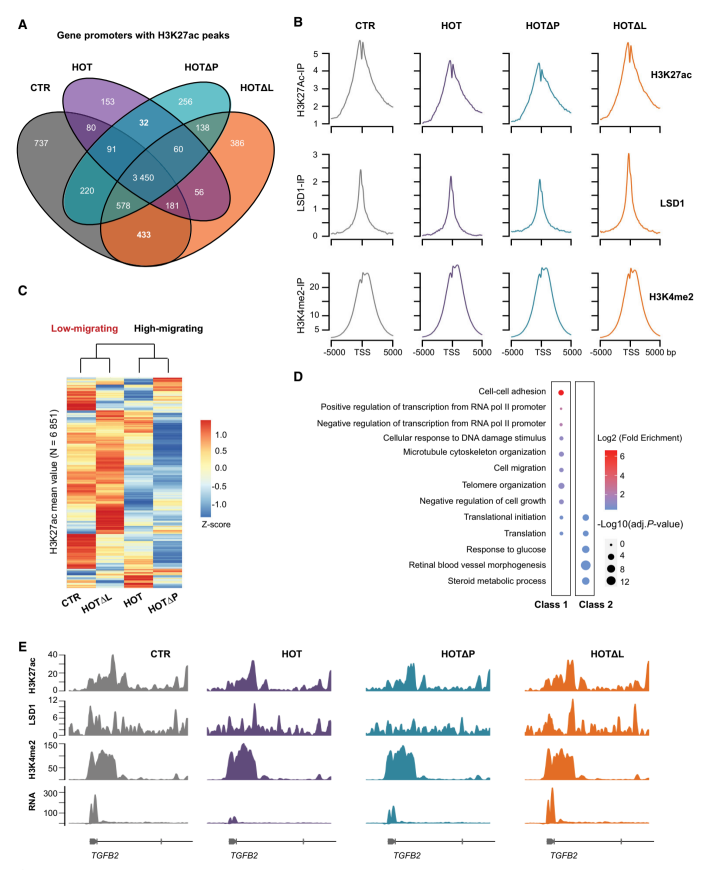


Figure 4.

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by transcriptomic and epigenetic analyses, respectively, including TGFB2 (Fig 4E). And only one NTNG1 gene from HMS (N=75) made part of Class 2 (Dataset EV2). Such low overlap may point to significant differences between steady-state transcriptome and H3K27ac status reflecting differences between chromatin modifications associated with transcriptional activity of the locus and total RNA levels at the equilibrium between RNA synthesis and stability.

Several studies propose LSD1 as a factor involved in regulation of promoter activities (Ferrari-Amorotti et al, 2013; Ambrosio et al, 2017; Carnesecchi et al, 2017; Zhang et al, 2018). To address how LSD1 occupancy of promoters is affected in the presence of HOTAIR, we performed CUT&RUN profiling of LSD1 in our system. Our data revealed that HOTAIR expression rather mildly affected LSD1 overall distribution across the genome (Fig EV3A and B). However, like H3K27ac, low-migrating cells shared more LSD1 peaks than high-migrating cells (Fig EV3C). We observed a positive correlation between LSD1 and H3K27ac levels suggesting cooccurrence of these features at promoters (Fig 4B; Appendix Fig S7). If considering all actively transcribed protein-coding genes, both H3K27ac and LSD1 were lower in high-migrating cells comparing to low-migrating cells (Fig EV2C). At chromatin, LSD1 as a part of several distinct multisubunit complexes can scaffold transcription factors, remodel nucleosomes, and demethylate histones. To determine whether the demethylation of H3K4me2, the prevailing substrate of LSD1 in actively transcribed euchromatic regions, is misregulated, we examined its distribution by CUT&RUN (Fig EV3D-F). Globally, H3K4me2 was preserved across the genome and highly enriched it the vicinity of gene TSS, with no statistically significant changes between low- and high-migrating cells (Fig 4B), nor between Class 1 and Class 2 promoters separately (Fig EV2C).

Overall, our analysis shows that the expression of HOTAIR with its LSD1-interacting domain is associated with changes in H3K27ac and LSD1 occupancy of EMT gene promoters, in striking accordance with the migratory phenotype observed in these cells. Notably, these variations are not concomitant with changes in H3K4me2 levels.

The LSD1-interacting domain of HOTAIR alters distribution of LSD1 at enhancers

Transcription is regulated by enhancers and enhancer clusters, known as super-enhancers (SE). We hypothesized that these distal regulatory elements may also contribute to fine-tune the regulation of an EMT balance in epithelial cells expressing HOTAIR variants. Therefore, we performed the annotation of active enhancers and SEs by the LILY pipeline using H3K27ac and ROSE algorithm (Fig EV4A) (Boeva et al, 2017). Notably, the number of active enhancers in low-migrating Epi-CTR and -HOTΔL was higher than in high-migrating Epi-HOT and -HOTΔP (Fig 5A) whereas the number of SEs was rather constant among conditions (Fig EV4B). Very few SEs were cell line specific, suggesting that, globally, expression of HOTAIR or its variants did not induce a deep reprogramming of cellular identity. Hence, we further focused on enhancers and compared LILY annotations among conditions. A substantial portion of active enhancers was either specific to Epi-CTR or -HOT Δ L (N = 2,096) or shared by both (N = 648), but very

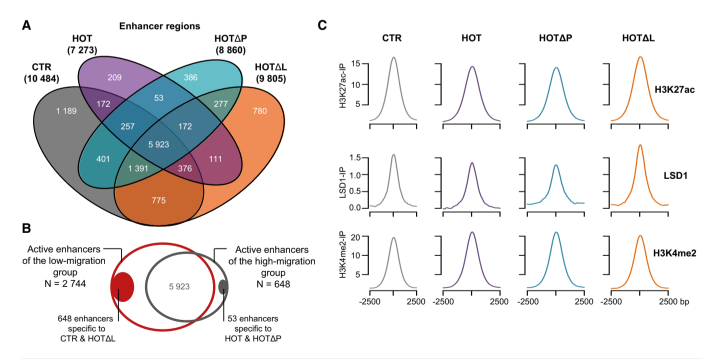


Figure 5. HOTAIR expression perturbs LSD1 activity at enhancers.

- A Venn diagram representing intersections of active enhancer regions in Epi cells.
- B Diagram representing putative active enhancers identified by LILY in each cell line and shared between low- or high-migrating groups, respectively.
- C Metagene profiling of H3K27ac, LSD1, and H3K4me2 over IgG within the 5-kb window of enhancers in Epi cells expressing none (CTR) or different HOTAIR variants (HOT, HOTΔP, and HOTΔL). bp stands for base pairs.

few were inherent to high-migrating Epi-HOT or -HOT Δ P (N = 595) cells or shared by both (N = 53; Fig 5A and B). It suggests that transcriptomic changes and consequent gain of migration by epithelial cells in presence of HOTAIR with intact LSD1-interacting domain may result from changes in enhancer activity. Indeed, consistently with a global decrease in H3K27ac observed in high-migrating cells (Fig 4B), enhancers also showed less acetylation in Epi-HOT and - $HOT\Delta P$ cells. (Fig 5C). CUT&RUN analysis revealed the presence of LSD1 at putative enhancers annotated by LILY (Fig 5C). Remarkably, a metagene profiling showed more LSD1 at enhancers in lowmigrating cells (Epi-CTR and -HOTΔL) in comparison to highmigrating cells (Epi-HOT and -HOT Δ P) (adj. *P*-value < 10^{-12} , Wilcoxon test) (Fig 5C). This was concomitant with slightly lower H3K4me2 levels (adj. P-value < 10^{-10} , Wilcoxon test). While considering separately putative enhancers shared by low- and highmigrating cells (775 and 53, respectively), we also noticed that the enhancers of Epi-CTR and -HOTΔL had higher LSD1 summit than the same enhancers in Epi-HOT and -HOT Δ P (adj. P-value < 0.05, Wilcoxon test) (Fig EV5A and B). At the same time, this pattern was not observed for enhancers of the high-migration group. Moreover, difference in LSD1 occupancy of both types of enhancers was not significantly associated with changes in H3K4me2 levels in all conditions (adj. P-value > 0.05, Wilcoxon test; Fig EV5B and C). One such example of Lsd1-bound regulatory element annotated as super-enhancer by LILY is located downstream the TGFB2 gene and has been reported to control the expression of the gene in the context of fibrosis (Shin et al, 2019) (Appendix Fig S8A). In our system, TGFB2 was expressed to higher extend in low-migrating cell (LMS set) and this was concomitant with higher Lsd1 and H3K27ac levels at its enhancer, measured by CUT&RUN. Considering the overall disbalance in enhancer numbers between these two groups, the latter observation may however suggest that the enhancers of the low-migration group are more perturbed in presence of HOTAIR and, hence, depend on HOTAIR-LSD1 interaction, whereas the enhancers of the high-migration group are rather HOTAIR-independent.

Collectively, these observations supported a model where LSD1 occupies proximal and distal regulatory elements. Expression of HOTAIR with its LSD1-interacting domain in Epi-HOT and -HOΤΔP, perturbs LSD1 distribution. Its action does not seem to be exclusively due to H3K4 demethylation but elicits changes in promoter and enhancer activities. As a consequence, this may imbalance transcription attenuating expression of epithelial genes and promoting expression of mesenchymal genes responsible for cell–cell adhesion and migration (Fig 6). Hence, our results revealed an unexpected role of HOTAIR as a molecular effector of LSD1, which may contribute to EMT at the very early steps of transformation of a normal epithelial cell to a neoplastic one.

Discussion

Epithelial-to-mesenchymal transition is involved in development, tissue homeostasis, and repair, as well as cancer progression. During EMT, epithelial cells acquire mesenchymal traits that promote migration and invasion and thus can underlie a high metastatic potential of tumors. HOTAIR and LSD1 have been independently studied in a variety of cell-based and clinical settings as

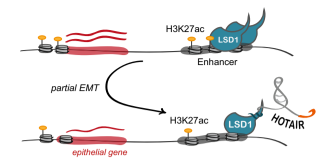


Figure 6. Model of HOTAIR activity.

In epithelial cells, the LSD1-interacting domain of HOTAIR perturbs LSD1 function at enhancers and promoters resulting in alteration of the transcriptional response and change in expression of genes involved in control of cell migration.

factors associated with EMT and metastasis. Although the regulatory function of HOTAIR is unambiguously linked to the acquisition of migration and invasion capacities, the role of the histone demethylase LSD1 is rather context-specific. LSD1 is ubiquitously expressed in both epithelial and mesenchymal cells and as a part of distinct multisubunit complexes can promote transcription repression or activation through its scaffold, demethylase, or chromatin remodeling functions (Shi *et al*, 2004; Wang *et al*, 2007; McDonald *et al*, 2011; Whyte *et al*, 2012; Li *et al*, 2016).

The present work revealed that HOTAIR negatively regulates the function of LSD1 in maintaining epithelial identity. HOTAIR appears to require its interaction with LSD1 but not with PRC2 to promote migration of epithelial cells. Accordingly, upon expression of HOTAIR with intact LSD1-interacting domain, we observed a decrease in level and change in distribution of the ZO-1 tight junction protein in high-migration cells. We found that most of the transcriptome changes induced by HOTAIR require both PRC2- and LSD1-interacting domains, suggesting that these domains have synergic and independent functions as well. Importantly, epithelial cells expressing a HOTAIR truncation that lacks the LSD1-interacting domain (HOTAL) still maintain epithelial identity. In this context, HOTAIR association to PRC2 is not sufficient to induce cell state changes even if the transcriptome of these cells is largely perturbed. We cannot exclude the inhibitory effect of HOTAIR on the PRC2 enzymatic activity at chromatin as proposed by another recent study (Portoso et al, 2017), but we can also consider the fact that in Epi-HOT Δ L cells, LSD1 continues to operate independently of HOTAIR. LSD1 elicits transcriptional programs upon different stimuli and transitions. It has been proposed to control the timing of transcription at enhancers during muscle and embryonic stem cell differentiation or in prostate cancer (Whyte et al, 2012; Cai et al, 2014; Scionti et al, 2017). LSD1 has been shown to regulate negatively or positively transcription from specific promoters (Ferrari-Amorotti et al, 2013; Ambrosio et al, 2017; Carnesecchi et al, 2017; Zhang et al, 2018). In our system, expression of HOTAIR with its LSD1interacting domain results in drastic changes in H3K27ac status, affecting gene expression pattern. Concomitantly, we observed an overall change in LSD1 occupancy of promoters and enhancers and, specifically, at regulatory elements active in low-migrating cells. LSD1 activity at promoters and enhancers by consequence may

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subordinate transcriptional response. At the same time, regulatory elements in high-migrating cells seem to be independent of HOTAIR. The striking correlation between the loss of epithelial traits and changes in the LSD1 genomic landscape supports a pivotal role for LSD1 as a factor that prevents cells from sensing or undergoing EMT, as previously suggested (McDonald et al, 2011). Remarkably, HOTAIR activity in stimulation of cell migration can be counteracted by overexpression of LSD1. Noteworthy, in this context, LSD1 activity seems to be partially inhibited by HOTAIR. Together, these observations highlighted a functional duality of LSD1 in EMT: On one hand, LSD1 is overexpressed in many cancers, it promotes a transition to mesenchymal state and metastasis (Lv et al, 2012; Egolf et al, 2019; Xie et al, 2020) but at the same time LSD1 contributes to the maintenance of epithelial identity (Wang et al, 2009). In the light of our findings, these LSD1 functions can be in part decommissioned by HOTAIR in epithelial cells. Still a lot of questions remain open on how HOTAIR influences LSD1 activity and whether this lncRNA affects its tethering or residence at specific locations or its enzymatic or scaffold activities, at chromatin but also at proteins other than histones. Indeed, even if we could measure only slight variations in H3K4me2, erased by LSD1, they remain rather subtle, indicating that HOTAIR-mediated LSD1 decommissioning from enhancers and/or promoters could affect more its function in chromatin remodeling, scaffolding, or tethering of transcription factors or others.

Intriguingly, several alternative splicing and TSS isoforms of HOTAIR are annotated in the human genome, some of which lack the LSD1- and/or PRC2-binding domains (Mercer *et al*, 2012). Although the clinical relevance of these isoforms has not yet been established, our results suggest that tumors expressing isoforms of HOTAIR that lack the LSD1-binding domain would have a lower metastatic potential and hence better prognosis. Thus, it will be of interest to assess the expression of HOTAIR isoforms in tumors of different grades and prognosis together with the LSD1 level and occupancy but also enhancer activity. Overexpression of the LSD1-interacting domain of HOTAIR in tumor cells with high expression of LSD1 might prove to be a strategy to balance the toxic effect of LSD1 in aggressive cancers.

Materials and Methods

Cell lines and culture

Cells were cultured in a humidified 5% $\rm CO_2$ atmosphere at 37°C in high-glucose DMEM with GlutaMAX for HEK293T (ATCC® CRL-3216TM) and MEM alpha without nucleosides for HA5-late (Mes), HA5-early (Epi), and its derivatives Epi-HOT, -HOT Δ P, and -HOT Δ L. All media were supplemented with 10% fetal bovine serum, essential amino acids, and sodium pyruvate. Cell lines were systematically tested and were negative for mycoplasmas.

Plasmids and oligonucleotides

The CTR construct corresponds to pLenti CMV GFP (659-1) (Addgene, #17445). The HOTAIR full-length and truncated cDNA sequences were PCR amplified from the LZRS-HOTAIR plasmid (Gupta *et al*, 2010), cloned into pENTR and further sub-cloned into pLenti CMV Blast DEST (706-1; Addgene, #17451) (Campeau *et al*, 2009) using the

Gateway system (Thermo Fisher Scientific). LSD1 was PCR amplified from pUNO1-hLSD1a (Isoform NM_015013.3, Invivogen) and sub-cloned into pDEST-ZEO (Addgene, #17294). Oligonucleotide sequences for PCR are available in Appendix Table S1.

Cell transfection and transduction

HEK293T cells at 50–70% of confluence at T25 flasks were co-transfected with 1.3 μg of psPAX2 (Addgene, #12260), 0.8 μg of pVSVG (Addgene, #36399), and 0.8 μg of the lentiviral plasmid bearing cDNA of GFP (CTR), full-length (HOT), truncated HOTAIR (HOΤΔP and HOΤΔL), or 0.4 μg of each of the above constructs and 0.4 μg of LSD1-OE lentiviral plasmid in presence of 5 μl of Lipofectamine® 2000 Transfection Reagent according to manufacturer's protocol (Thermo Fisher Scientific). Virus supernatant was recovered, filtered 48 h post-transfection and added to Epi cells at 50–70% of confluence. After 24 h post-transduction, cells were sub-cultured every 2 days at a 1:4 ratio in 6 μg/ml of blasticidin or 6 μg/ml of blasticidin and 10 μg/ml of zeocin supplemented MEM alpha medium for 1 week and then in MEMalpha medium for additional 2 weeks prior to any experiment.

Antisense oligonucleotides (Appendix Table S1) were added as a supplement to a final concentration of 10 μ M to Mes cells at 50% of confluence for 2 days and again for additional 2–3 days prior to RNA extraction or wound healing assay.

Wound healing assay

Cell suspensions containing 200,000 cells were seeded to each well of the six-well plate and left growing to 95–100% of confluence for 24–36 h. After scratching, each wound was imaged twice at time t=0 and t=24 h post-scratch with a Zeiss inverted microscope with a total of 30–48 high-field units (HFU) per cell line in transduced Epi, six HFU per cell line in ASO treated cells, and 30 HFU per cell line for LSD1-OE. The invaded area was calculated as a ratio of migrated cell surface area to total surface area with a web-based image automated analysis software Tscratch (Gebäck $et\ al.\ 2009$). Results were presented as a mean \pm SEM. P-values were calculated using the Student's t-test.

Subcellular fractionation

This protocol was adapted from Ref. (Gagnon *et al*, 2014). All steps were performed on ice with ice-cold buffers supplemented by 20 U/µl SUPERase-IN (Thermo Fisher Scientific) for RNA extraction and 0.1 mM AEBSF (SIGMA) for protein extraction. RNA extraction from all fractions was performed using TRIzol (Thermo Fisher Scientific) at 65°C for 5 min with regular vortexing and then purified with miRNeasy kit (Qiagen) according to the manufacturer's instructions. For protein extractions, the chromatin fraction was sonicated for 8 min (30 s-on/30 s-off, "high power") on the Bioruptor Plus (Diagenode). Protein content was quantified with the Thermo Fisher Scientific™ BCA Protein Assay Kit (Thermo Fisher Scientific).

Protein extraction and Western blot

Cells at 80% of confluence were lysed in RIPA buffer (Thermo Scientific) supplemented with 0.1~mM AEBSF (SIGMA). A total of 5~mm

and 10 µg of the whole protein extract were separated on a NuPAGE Novex 4–12% Bis-Tris Protein Gel and transferred using Thermo Fisher Scientific iBlot Dry Blotting System (Thermo Fisher Scientific). After blocking, membranes were immunoblotted with the Epithelial-Mesenchymal Transition Western Blot Cocktail (1:2,500 dilution; Abcam, ab157392), anti-GAPDH (Millipore, mab374), EMT Sampler kit (Cell Signaling Technology, 9782), anti-LSD1 (Diagenode, C15410067) overnight at 4°C, then washed and incubated with a corresponding HRP-conjugated secondary antibody (1:5,000 dilution) for 1 h at room temperature and visualized using the SuperSignal West Femto, Dura, or Pico Substrate (Thermo Fisher Scientific). Quantifications of protein amounts were performed relative to GAPDH using the ImageJ software.

Immunofluorescence

A total of 20,000 cells suspension was seeded on a 4-well microscope slide and let growing for 24 h. Following PBS wash, cells were fixed using formaldehyde (3%)–sucrose (2%) solution for 15 min, permeabilized, and blocked with 10% goat serum (Thermo Fisher Scientific). For F-actin staining, cells were incubated with 0.5 μ g/ml DMSO solution of phalloidin-tetramethylrhodamine isothiocyanate (TRITC; Sigma) for 40 min at room temperature. For ZO-1, cells were incubated with the primary antibody (1:100 dilution; Cell Signaling Technology, 8193) for 40 min at 37°C, washed in PBS, and incubated with Alexa Fluor 594 Goat anti-Rabbit secondary antibody (1:400; Thermo Fisher Scientific, A11037) for 30 min at 37°C. Slides were washed and mounted using a ½ dilution of Vectashield Mounting Medium with DAPI (Vector Laboratories).

Fluorescent microscopy and image analysis

For each cell line, three HFUs per well were acquired as stacks composed of twenty slices of 0.5 μm interval over the height of the cells with a 40X objective on a Zeiss AxioVision microscope equipped with an ApoTome module (PiCT-BDD platform, Institut Curie). All images were captured with identical exposure time for DAPI and Alexa Fluor 594 using AxioVision software and then analyzed using the Fiji software (Schindelin $\it et~al,~2012$). Stacks were processed with the "Z-project" tool ("Sum Slices" setting) and an identical minimum threshold value was set for all images. Signal was then quantified using the "Measure" tool and normalized by the number of cells per image. Integrated density per cell was again normalized on overall image background signal to reduce hybridization biases between slides and presented as a mean \pm SEM for at least 11 HFUs.

RNA extraction and reverse transcription

Total RNA extraction was performed directly from cell cultures with miRNeasy kit according to the manufacture' instructions (Qiagen). Only RNAs with the RNA integrity number (RIN) above eight were used for further experiments. Reverse transcription (RT) was performed on 500 ng of RNA with either random and oligo(dT) primers mix (iScript $^{\text{TM}}$ cDNA Synthesis Kit) or specific oligonucleotides (SuperScript II Reverse Transcriptase, Thermo Fisher Scientific; Appendix Table S1). Reactions without RT were included as a negative control for DNA contamination.

RNA-seq library preparation

 $1~\mu g$ of RNA was depleted for ribosomal RNA with the RiboMinusTM Eukaryote Kit for RNA-seq (Thermo Fisher Scientific) and converted into cDNA library using a TruSeq Stranded Total Library Preparation kit (Illumina). cDNA libraries were normalized using an Illumina Duplex-specific Nuclease (DSN) protocol prior to a paired-end sequencing on HiSeqTM 2500 (Illumina).

RNA-seq data analysis

Reads were mapped allowing three mismatches using TopHat 2.0.4 (Trapnell *et al*, 2009) and the human genome version hg19. Uniquely mapped reads were assembled using the BedTools suite (Quinlan & Hall, 2010) and merged in segments if mapped in the same strand to the GENCODE v15 annotation to extract proteincoding genes and annotated noncoding genes including lncRNA, antisense, sense_intronic, sense-overlapped, and pseudogenes.

Differential expression analysis was performed using the DESeq R package (Anders & Huber, 2010). Only genes with an expression fold change (FC) above 2 and adjusted *P*-value below 0.05 were retained as significantly differentially expressed.

CUT&RUN

The experiment was performed in two replicates following the protocol from Henikoff Lab with some modifications (Skene et al, 2018). Briefly, slowly frozen (for histone marks) or fresh (for IgG, LSD1, and EZH2) cells pelleted after scrapping were resuspended in wash buffer at 10⁷ cells/ml. At room temperature, per sample, 10 μl of Concanavalin A beads slurry washed in bead activation buffer was added to 10⁶ of cells suspension in the total volume of 1 ml of wash buffer and incubated for 10 min with rotation at 15-20 rpm. Antibodies against H3K4me2 (Millipore, 07-030), H3K27ac (Millipore, MABE647), H3K9me2 (Active Motif, 39239), LSD1 (Diagenode, C15410067), and EZH2 (Active Motif, 39933) were diluted 100 times and IgG (Invitrogen, 31235)-1,100 times in 50 µl of antibody buffer (wash buffer with 0.1% digitonin, 2 mM EDTA) and added to the nuclei fixed on beads and incubated 10 min for histone marks and 1 h for LSD1, EZH2, and IgG. Afterward, the beads were washed twice with 250 μ l of digitonin buffer (wash buffer with 0.1% digitonin) and resuspended in 50 µl of pAG-MNase (CUTANA EpiCypher, 1:20 dilution) in digitonin buffer, incubated 10 min with agitation. Beads were washed twice in 250 µl of ice-cold digitonin buffer, resuspended in 100 µl of ice-cold digitonin buffer, kept on ice for 5 min before adding 2 µl of 100 mM CaCl₂ and incubating for 30 min on ice. The reaction was stopped by addition of 100 μ l of stop buffer containing 0.1% digitonin and 10 pg/ml spike-in DNA and incubation for 10 min at 37°C. Bead suspension was centrifuged at 16 krcf at 4°C for 5 min, beads recovered for DNA fragments purification with NEB Monarch PCR & DNA purification kit following the protocol enriching for short DNA fragments. DNA was quantified by Qubit HS DNA kit, and 5 ng of each sample was used for DNA library preparation.

CUT&RUN libraries preparation and sequencing

CUT&RUN libraries were prepared using the TruSeq ChIP-Sample Prep Kit-Set A and PCR Box (Illumina) from 5 ng of DNA with 10

PCR cycles for amplification following the manufacture's procedures. Concentration and quality were assessed by Qubit with dsDNA HS Assay Kit (Thermo Fisher Scientific) and High Sensitivity DNA Analysis Chip Kit (Agilent Technologies). Libraries were sequenced on a NovaSeq S1 device in a pair-end 100 bp read length mode. DNA sequencing metrics are present in Appendix Table S2.

CUT&RUN analysis

Reads were first aligned to human genome version GRCh38 using bowtie (version 2.4.1) (Langmead & Salzberg, 2012), with options -I 10 -X 700—no-mixed. Reads with MAPQ value ≥ 30 were kept for further analysis. To equilibrate datasets, the larger samples were randomly down-sampled to approximately 9 million of reads. For histone marks (H3K27ac, H3K4me2) and LSD1, peaks were called relative to IgG control using HMCan (Ashoor et al, 2013) with punctuate parameters according to (github.com/BoevaLab/HMCan/tree/ master/configurations/HMCan.config.punctuate.txt), keeping duplicate reads, and without correction for ploidy. Only peaks showing at least one nucleotide overlap between two replicates were retained and subsequently merged to common peaks. For further analysis, common peaks between conditions were also merged to peak regions (Dataset EV5). The merged peaks were assigned to closest genes, and the shortest distance between each peak and the closest transcript start site (TSS) coordinates were calculated using Gencode v32 annotation. H3K27ac and H3K4me2 peaks within the window of 2 kb around the TSS were assigned to promoters. Regulatory elements (enhancers and super-enhancers) were annotated using the LILY pipeline (Boeva et al, 2017). Peaks overlap with genomic elements was retrieved from genome mapping information of Gencode transcripts using BEDtools. H3K27ac and LSD1 peaks were validated by qPCR for one enhancer regions (see Quantitative PCR section below, Appendix Fig S8B and C).

Gene features analysis

Gene ontology biological process, KEGG pathways, and cellular compartment terms enrichment analyses were performed using DAVID 6.8 or ClusterProfiler (Huang et~al, 2008; Huang et~al, 2009; Yu et~al, 2012). Enrichment score was calculated as $-\log_{10}(P\text{-value})$. Gene sets are present in Dataset EV2.

Quantitative PCR

cDNA was diluted 10–50 times and quantified by real-time PCR with LightCycler® 480 SYBR Green I Master (SIGMA). The *RPL11* gene was used as a reference to quantify a relative abundance of each cDNA (Appendix Table S1). Error bars represent standard deviations in relative quantity of cDNA prepared from at least three independent experiments. To validate CUT&RUN annotated peaks, we performed a qPCR with IP- or IgG-DNA adjected to 0.5 ng/µl, using the *TGFB2* SE_67 region as a region (Appendix Table S1, Appendix Fig S8).

Data availability

The datasets and computer code produced in this study are available in the following links:

- Raw RNA-seq and CUT&RUN datasets: Gene Expression Omnibus GSE106517 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc = GSE106517).
- Computer scripts: GitHub (https://github.com/MorillonLab/HOT AIR_LSD1).

Expanded View for this article is available online.

Acknowledgements

We deeply thank Rocco Cipolla, Elena Battistello, Sandra Carignon, Matthieu Lejars, and Marc Descrimes for technical contributions, Karina Jouravleva, and Luis Castro-Vega for discussions and expertise. The HOTAIR cDNA containing LZRS plasmid was a kind gift of H.Y. Chang. RNA-seq and CUT&RUN sequencing efforts were supported by the grant from the ICGex program of Institut Curie attributed to A.M. and benefited from the facilities and expertise of the NGS platform of Institut Curie, supported by Agence Nationale de la Recherche (ANR-10-EQPX-03, ANR10-INBS-09-08). A.M., M.P., and Z.S. were supported by the European Research Council (ERC-consolidator DARK-616180-ERC-2014) attributed to A.M., J.J. was supported by "La Ligue contre le Cancer."

Author contributions

Project administration and funding acquisition, AM, MP; Conceptualization, supervision and writing, MP; Draft revisions, DF, JJ; Investigation, MP, CB, JJ, DF; Formal analysis, MG, US, ZS; Expertise, AL-V, SG.

Conflict of interest

The authors declare that they have no conflict of interest.

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