Targeted DNA oxidation by LSD1–SMAD2/3 primes TGF-β1/ EMT genes for activation or repression

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ABSTRACT

The epithelial-to-mesenchymal transition (EMT) is a complex transcriptional program induced by transforming growth factor β1 (TGF-β1). Histone lysine-specific demethylase 1 (LSD1) has been recognized as a key mediator of EMT in cancer cells, but the precise mechanism that underlies the activation and repression of EMT genes still remains elusive. Here, we characterized the early events induced by TGF-β1 during EMT initiation and establishment. TGF-β1 triggered, 30–90 min post-treatment, a nuclear oxidative wave throughout the genome, documented by confocal microscopy and mass spectrometry, mediated by LSD1. LSD1 was recruited with phosphorylated SMAD2/3 to the promoters of prototypic genes activated and repressed by TGF-β1. After 90 min, phospho-SMAD2/3 downregulation reduced the complex and LSD1 was then recruited with the newly synthesized SNAI1 and repressors, NCoR1 and HDAC3, to the promoters of TGF-β1-repressed genes such as the Wnt soluble inhibitor factor 1 gene (WIF1), a change that induced a late oxidative burst. However, TGF-β1 early (90 min) repression of transcription also required synchronous signaling by reactive oxygen species and the stress-activated kinase c-Jun N-terminal kinase. These data elucidate the early events elicited by TGF-β1 and the priming role of DNA oxidation that marks TGF-β1-induced and -repressed genes involved in the EMT.

INTRODUCTION

The epithelial-to-mesenchymal transition (EMT) is the product of a complex transcriptional program that induces mesenchymal characteristics in polarized epithelial cells. These features include reduced cell–cell contacts, enhanced migratory and invasive capacity, and resistance to apoptosis (1). EMT occurs during tumor progression to generate metastatic cells that are endowed with a more motile and invasive phenotype (2). Several transcription factors are implicated in EMT control; particularly, the repressor SNAI1, a member of the zinc finger transcription factor family, is induced by transforming growth factor β1 (TGF-β1) to initiate the EMT transcription program (3). EMT also critically depends on reactive oxygen species (ROS), which support tumor growth by stimulating multiple pathways (4–6). Additionally, high oxidative stress levels in the tumor microenvironment, due to both hypoxia and stromal inflammatory cell recruitment, can promote myofibrob...
last differentiation into activated cancer-associated fibroblasts (CAFs) (7–9). CAFs, in turn, exploit the oxidative stress to produce cytokines and proteases that can elicit EMT in cancer cells (10). Moreover, CAFs induce EMT in prostate carcinoma cells through a redox-dependent mechanism that involves nuclear factor κB (NF-κB), NADPH oxidase (NOX), cyclooxygenase and hypoxia-inducible factor 1 (HIF-1) to sustain tumor progression and metastatic dissemination (11).

ROS and DNA oxidation are also linked to transcription. Demethylation of histone H3 by histone lysine-specific demethylase (LSD1) is stimulated by estrogen and retinoic acid. ROS produced by LSD1 induce oxidation of deoxy-G (oxo-dG), a process that is instrumental for the assembly of a productive transcriptional initiation complex and the formation of transcriptional loops (12,13). In this scenario, the aim of our work was to explore whether a similar mechanism, dependent on nuclear ROS production, could be the driving force behind the transcriptional program leading to EMT in mammary epithelial cells induced by TGF-β1 during the initiation of EMT.

We report here that TGF-β1 triggers an early (30 min) oxidative wave in the nucleus, mediated by LSD1, recruited and targeted by phosphorylated SMAD2/3 to the promoters of activated and repressed TGF-β1/EMT genes. At 90 min, there was a second oxidative wave, detected by the accumulation of oxo-dG glycosylase, OGG1, only at TGF-β-repressed genes. At these promoters, a repressive complex was formed by LSD1 and HDAC3–NCoR1 with the newly synthesized repressor SNAI1. Early (30–90 min) DNA oxidation mediated by LSD1 was necessary for both activation and repression of transcription by TGF-β1. Transcriptional repression by TGF-β1 required an additional signal generated by the ROS-activated kinase c-Jun N-terminal kinase (JNK).

MATERIALS AND METHODS

Materials

PVDF membrane was obtained from Millipore (Bedford, MA, USA); anti-E-cadherin, anti-N-cadherin, anti-SET7/9, anti-OGG1/2, anti-HDAC3, LSD1 shRNA plasmid and JMJD2a shRNA plasmid were obtained from Santa Cruz Biotechnology (Santa Cruz, CA, USA); anti-FLAG and anti-smooth muscle actin antibody and Phalloidin-TRITC were obtained from Sigma-Aldrich (St Louis, MO, USA); DCFDA and Alexa 488 secondary antibody were obtained from Molecular Probes; anti-H3K4me2, anti-H3K4me3, anti-H3K9me2, anti-H3K9me3, anti-H3, anti-LSD1, anti-JMJD2A, anti-SUV39H1, anti-NCoR1, anti-APE1, anti-β-actin, anti-β-tubulin antibodies, anti-IgG mouse and anti-IgG rabbit were obtained from Abcam (Cambridge, UK); anti-pSMAD2/3 (Ser 423/425) and anti-SMAD2/3 antibodies were obtained from Cell Signaling Technology (Danvers, MA, USA); anti-8-oxo-dG antibodies were obtained from Trevigen Inc. (Gaithersburg, MD, USA); and the OxyDNA assay was obtained from Calbiochem (San Diego, CA, USA).

Cell cultures, transfections and drug pretreatments

MCF10A cells were obtained from the American Type Culture Collection. MCF10A cells were cultured in DMEM/Ham’s F12 supplemented with 5% horse serum, 0.5 μg/ml hydrocortisone, 100 ng/ml cholera toxin, 20 ng/ml EGF and 10 μg/ml insulin. Before TGF-β1 stimulation, the cells were starved in serum-free medium containing DMEM/Ham’s F12 supplemented with 0.5 μg/ml hydrocortisone, 100 ng/ml cholera toxin, 20 ng/ml EGF and 10 μg/ml insulin for 24 h. TGF-β1 (10 ng/ml) was added to the cultures at different times (30–60–90 min or 12–24 h). Where necessary, MCF10A cells were pretreated with N-acetylcysteine (NAC, 5 mM), dihydroxyphenidion (DPI, 10 μM), tranylcypromine (TCP, 0.5–1 mM), JIB04 (23 mM), SP600125 (10 μM), SB431542 (10 μM), apocynin (50 μM) and rotenone (5 μM). Several LSD1 inhibitors, SP2509 and CBB1007 (noncatalytic, reversible inhibitors) and OG-L002 and GSK (catalytic MAO inhibitors) (see selleckchem.com/lsd1.html), were tested on MCF10A cells. All of them but TCP reduced the growth at 24–36 h of MCF10A cells at concentrations used (A. Pezone, unpublished data) (14,15).

LSD1–JMJD2A silencing

MCF10A cells were transfected with LSD1 or JMJD2A shRNA/hairpin vectors using Lipofectamine 3000 (Invitrogen, Waltham, MA, USA) according to manufacturer’s instructions or by electroporation. Four single or pooled shRNA/hairpin vectors (Santa Cruz Biotech, LSD1sh sc-60970 and JMJD2A sc-62515) targeting different LSD1 or JMJD2A segments were transfected in MCF10A cells. After 24 h, the cells were exposed to TGF-β1 and the levels of JMJD2A, LSD1, β-actin, GADPH, α-tubulin mRNAs and 18S RNA were measured by qPCR or immunoblot with specific antibodies (except 18S RNA). In some experiments, the wild-type human LSD1 expression vector without the 3′-UTR was included in the transfection to control off-target effects in experiments with single or pooled shRNA/hairpin vectors (Supplementary Figure S2).

Immunoprecipitation and immunoblot analysis

A total of 1 × 10⁶ cells were lysed for 20 min on ice in 500 μl of complete RIPA lysis buffer (50 mM Tris–HCl, pH 7.5, 150 mM NaCl, 1% Nonidet P-40, 2 mM EGTA, 1 mM sodium orthovanadate, 1 mM phenylmethylsulfonyl fluoride, 10 μg/ml aprotinin, 10 μg/ml leupeptin). Lysates were clarified by centrifugation and immunoprecipitated for 4 h at 4°C with 1–2 μg of the specific antibodies. Immune complexes were collected on Protein A/G PLUS agarose, separated by SDS-PAGE and transferred onto nitrocellulose. Immunoblots were incubated in 3% milk, 10 mM Tris–HCl, pH 7.5, 1 mM NaCl and 0.1% Tween 20 for 1 h at room temperature, probed first with specific antibodies and then with secondary antibodies.

Real-time PCR

Total RNA from MCF10A was extracted using TRIzol (Gibco/Invitrogen). mRNA was reverse transcribed for 1
h at 50°C, and the reaction was heat inactivated for 15 min at 70°C. The primers used are shown in Supplementary Table S1. Data were normalized to 18S RNA. Results (mean ± SD) are the mean of three different experiments in triplicate.

Immunocytochemistry

After washing with phosphate-buffered saline (PBS), cells were fixed with 3.7% formaldehyde solution in PBS for 20 min at 4°C. After extensive washing in PBS, cells were permeabilized with 0.1% Triton X-100 in PBS and then stained with anti-E-cadherin antibody overnight and with 50 µg/ml fluorescent phallolidin conjugate, phallolidin-TRITC, in PBS for 1 h at room temperature and then with anti-rabbit Alexa 488 secondary antibodies. After washing with PBS, the cover slides were mounted with glycerol and then observed under a confocal fluorescence microscope (Leica Microsystems).

8-Oxo-G assay

Immunocytochemistry was performed according to the manufacturer's instructions (Trevigen Inc.). The anti-8-oxo-dG antibodies (mouse monoclonal 15A3) were validated by confocal microscopy (16) and chromatin immunoprecipitation (ChiP) (17). The signal was resistant to RNase A and sensitive to DNase I and NAC. Briefly, cells were fixed for 15 min with MeOH at −20°C followed by 15 min at −20°C with acetone. Fixed cells were treated with 0.05 N HCl for 5 min on ice and incubated with 250 µl of 100 µg/ml RNase A in 150 mM NaCl and 15 mM sodium citrate for 1 h at 37°C. After sequentially washing in 1× PBS and 35%, 50% and 75% ethanol (EtOH), DNA was denatured in situ with 250 µl of 0.15 M NaOH in 70% EtOH. A 0.2 µg/ml (250 µl) TO-PRO-3 Iodide (Thermo Fisher Scientific) in 1× PBS was used to stain DNA for 10 min. After several washing with 70% EtOH containing 4% v/v formaldehyde, 50% and 35% EtOH, and 1× PBS, the cells were incubated in 250 µl of 5 µg/ml proteinase K in 20 mM Tris and 1 mM EDTA, pH 7.5 (TE), for 10 min at 37°C. Nonspecific binding was prevented by incubation with 5% normal goat serum in 1× PBS containing 0.05% Tween 20, fixed cells were incubated in 250 µl of fluorescent secondary antibody, goat anti-mouse IgG (Alexa Fluor 488), at 5 µg/ml in 1× PBS containing 1% BSA for 1 h in the dark at room temperature. After extensive washing, cells were mounted with glycerol and observed by a laser scanning confocal microscope.

Colocalization of 8-oxo-G signals in confocal microscopy

Colocalization of confocal images was performed with two specific plugins of ImageJ: (i) Colocalisation (http://www.sussex.ac.uk/gdsc/intranet/pdfs/Colocalisation.pdf) and (ii) Coloc_2 (https://imagej.net/Coloc_2). Briefly, two points colocalized if their ratio of intensity was higher than the setting value (50 was used as default setting value). To normalize to the number of the cells in each field, the values were expressed as the percentage of the area of the cell nuclei in each individual photo. The area of non-colocalized green spots was measured as percentage of the area of the nuclei. For each sample, Pearson's correlation and Manders' split coefficient were calculated.

Intracellular ROS determination

Production of intracellular H₂O₂ was assayed as previously described (18). At 3 min before the end of the incubation time, DCFDA was added to a final concentration of 5 µM. Cells were lysed in 1 ml of RIPA buffer and analyzed immediately by fluorescence analysis using a Perkin Elmer Fluorescence Spectrophotometer 650-10S equipped with a xenon power supply (excitation 488 nm, emission 510 nm).

2D ultra-performance liquid chromatography–tandem mass spectrometry

The 2D ultra-performance liquid chromatography–tandem mass spectrometry (UPLC–MS/MS) analysis was performed following our recently developed method (19). Briefly, DNA hydrolysates were spiked with a mixture of internal standards in a volumetric ratio of 4:1, to concentration of 50 fmol/µl of [D3]-5-hydroxymethyl-5-C (5-hmdC), [13C10, 15N2]-5-formyl-2'-deoxycytidine, [13C10, 15N2]-5-carboxyl-2'-deoxycytidine, [13C10, 15N2]-5-hydroxymethyl-2'-deoxycytidine, [15N2]-8-oxo-dG. Chromatographic separation was performed with a Waters Acquity 2D UPLC system with a photodiode array detector, for the first-dimension chromatography, used for the quantification of unmodified deoxynucleosides and 5-methyl-2'-deoxycytidine, and a Xevo TQ-S tandem quadrupole mass spectrometer for the second-dimension chromatography. An at-column dilution technique was used between first and second dimensions for improving retention at the trap/transfer column. The columns used were a Phenomenex Kinetex C-18 column (150 mm × 2.1 mm, 1.7 µm) at the first dimension, a Waters X-select C18 SH (30 mm × 2.1 mm, 1.7 µm) at the second dimension and Waters X-select C18 SH (30 mm × 2.1 mm, 1.7 µm) as the trap/transfer column. The chromatographic system operated in the heart-cutting mode, indicating that selected parts of effluent from the first dimension were directed to the trap/transfer column via six-port valve switching, which served as ‘injector’ for the second-dimension chromatography system. The flow rate at the first dimension was 0.25 ml/min and the injection volume was 0.5–2 µl. The separation was performed with a gradient elution for 10 min using a mobile phase of 0.1% acetate (A) and acetonitrile (B) (1–5% B for 5 min, column washing with 30% acetonitrile and re-equilibration with 99% A for 3.6 min). The flow rate at the second dimension was 0.35 ml/min. The separation was performed with a gradient elution for 10 min using a mobile phase of 0.01% acetate (A) and methanol (B) (4–50% B for 4 min, isocratic flow of 50% B for 1.5 min and re-equilibration with 96% A up to next injection). All samples were analyzed in three to five technical replicates of which technical mean
was used for further calculation. Mass spectrometric detection was performed using the Waters Xevo TQ-S tandem quadrupole mass spectrometer, equipped with an electrospray ionization source. Collision-induced dissociation was obtained using argon 6.0 at 3 × 10⁻⁶ bar pressure as the collision gas. Transition patterns for all the analyzed compounds as well as specific detector settings were determined using the MassLynx 4.1 IntelliStart feature.

Chromatin immunoprecipitation
MCF10A cells were grown to 95% confluence in DMEM/Ham’s F12. Following the addition of 10 ng/ml TGF-β1 for 30, 60 and 90 min, cells were washed twice with PBS and cross-linked with 1% formaldehyde at room temperature for 10 min. Cells then were washed with ice-cold PBS containing protease inhibitors (1 mM phenylmethylsulfonyl fluoride, 1 μg/ml aprotinin and 1 μg/ml pepstatin A). Cells were then resuspended in 0.2 ml of lysis buffer (1% SDS, 10 mM EDTA, 50 mM Tris–HCl, pH 8.1, 1 mM phenylmethylsulfonyl fluoride, 1 μg/ml aprotinin and 1 μg/ml pepstatin A), sonicated for 17 cycles (12 s followed by 28 s of stop) and centrifuged for 10 min at 10 000 × g at 4°C. Supernatants were collected and diluted in buffer containing 1% Triton X-100, 2 mM EDTA, 150 mM NaCl and 20 mM Tris–HCl, pH 8.1. Immunoprecipitation was performed overnight at 4°C with 2 μg of specific antibodies with 14 μl Protein G Dynabeads (Invitrogen, Waltham, MA, USA). After immunoprecipitation, precipitates were washed sequentially for 10 min each in high-salt solution (0.1% SDS, 1% Triton X-100, 2 mM EDTA, 20 mM Tris–HCl, pH 8.1, 150 mM NaCl), low-salt solution (0.1% SDS, 1% Triton X-100, 2 mM EDTA, 20 mM Tris–HCl, pH 8.1, 500 mM NaCl) and LiCl solution (0.25 M LiCl, 1% NP-40, 1% deoxycholate, 1 mM EDTA, 10 mM Tris–HCl, pH 8.1). Precipitates were then washed three times with TE buffer and extracted two times with 1% SDS and 0.1 M NaHCO₃. Eluates were pooled and heated at 65°C overnight to reverse the formaldehyde cross-linking. DNA fragments were purified with a QIAquick Spin Kit (Qiagen, Valencia, CA, USA). The primers used are shown in Supplementary Table S1.

Statistical analysis
The data we present derive from at least three biological and three technical replicates. Multivariate analysis of variance (MANOVA) or repeated measures analysis of variance (ANOVA) was used to determine differences in multiple dependent variables over time or between treatments. Each figure is associated with specific statistical figures indicated as Figure x-Stat.y. Statistical analysis of the data was performed by Student’s t-test. P-values of ≤0.05 were considered statistically significant.

RESULTS
TGF-β1-induced ROS were generated by multiple sources and were all essential for the EMT program
TGF-β1 evokes a time-dependent increase of ROS measured as DCF fluorescence that peaks at 30–45 min after treatment and thereafter declines, although remain-
Figure 1. TGF-β1-induced ROS are essential for EMT establishment. (A, B) ROS induced by TGF-β1. (A) Cells were serum starved for 24 h before stimulation with 10 ng/ml TGF-β1 for the indicated times. H₂O₂ production was evaluated as described in the ‘Materials and Methods’ section. (B) Cells were pretreated with 5 mM NAC or 1 μg/ml catalase for 15 min before stimulation with 10 ng/ml TGF-β1 for 30 min. The data presented are the mean of at least three experiments in duplicate (n = 6). Comparison between three or more samples was performed with one-way ANOVA test; comparison between each pair was performed with Student’s t-test: *P < 0.01 versus untreated control; **P < 0.01 versus TGF-β1 alone (see Supplementary Figure 1-Stat.1). (C) E–N cadherin switch and α-SMA levels induced by TGF-β1. Cells were pretreated with 5 mM NAC for 15 min and then stimulated with 10 ng/ml TGF-β1. After 72 h, the cells were lysed and subjected to immunoblot analysis for E-cadherin, N-cadherin, α-SMA and β-actin. (D) Photographs of cells exposed to 10 ng/ml TGF-β1 for 72 h were taken with a phase/contrast microscope. (E) Representative images of cells exposed to 10 ng/ml TGF-β1 for 72 h, fixed and examined by immunofluorescence microscopy of phalloidin in red and E-cadherin in green in the presence of NAC. (F, G) MCF10A cells were pretreated with 5 mM NAC or 5 μM DPI for 15 min before stimulation with 10 ng/ml TGF-β1 for 90 min. Total RNA was extracted and analyzed by qRT-PCR with primers corresponding to SNAI1, CDH2, VIM and PAI mRNA (F) and WIF1, cMYC, CDH1 and DSP genes (Supplementary Table S1) (G). The data shown are the mean of at least three experiments performed three times (n ≥ 9). A separate MANOVA was used to examine the association between the mRNA levels after TGF-β1 and NAC/DPI treatment; the interaction was significant (see Supplementary Figure 1-Stat.2 for panel F and Supplementary Figure 1-Stat.5 for panel G). Univariate testing found the effect to be significant for both NAC and DPI (see Supplementary Figure 1-Stat.3 for panel F and Supplementary Figure 1-Stat.6 for panel G). *P < 0.01 versus CTRL + TGF-β1; †P < 0.01 versus basal CTRL (without TGF-β1). Pairwise comparison to evaluate the response to TGF-β1 was performed with Student’s t-test (see Supplementary Figure 1-Stat.4 for panel F and Supplementary Figure 1-Stat.7 for panel G). *P < 0.01 versus each basal (without TGF-β1).
Nuclear G oxidation by LSD1 promoted TGF-β1 activation and repression of transcription

To identify the early (30–90 min) ROS sources induced by TGF-β1 that prime the cells to initiate EMT, we focused our attention on the nucleus. Rather than measuring nuclear ROS, which are extremely unstable and diffuse across membranes, we determined the accumulation of the major DNA oxidation product, 8-oxo-dG, by confocal microscopy using specific antibodies against 8-oxo-dG. Figure 2A shows significant accumulation of nuclear 8-oxo-dG after 30-min TGF-β1 treatment, in strict concomitance with the increased ROS levels (Figure 1A) and EMT engagement (Figure 1C). NAC inhibited the 8-oxo-dG levels induced by TGF-β1. DPI treatment reduced TGF-β1-induced nuclear 8-oxo-dG and increased the 8-oxo-dG signal, mainly in the perinuclear area (Figure 2A, and Supplementary Figures S1B and S1-Stat.2–3).

8-Oxo-dG has been implicated in transcriptional regulation by a variety of signals, including nuclear hormones such as estrogens (12), retinoic acid (13) and c-Myc (31) or hypoxia (32). Histone demethylation is a redox reaction, and under myriad circumstances it induces deoxy-G Oxidation (12,13). The enzymes involved in histone demethylation are mono- or dioxygenases; LSD1 or KDM1 is a monoxygenase that generates hydrogen peroxide during demethylation reactions (33). Since LSD1 has been associated with EMT engagement and SNAI1-mediated repression of epithelial markers (34), we investigated whether the nuclear oxidation wave induced by TGF-β1 was generated by LSD1. To this end, we silenced LSD1 in MCF10A cells by shRNA (Supplementary Figure S1D) interference and measured 8-oxo-dG upon TGF-β1 stimulation. LSD1 silencing was as effective as NAC (compare Figure 2A and B) at inhibiting TGF-β1-induced deoxy-G Oxidation (Figure 2B, and Supplementary Figures S1C and D and S1-Stat.4–5). Additionally, treatment of the cells with a monoamine oxidase inhibitor, namely TCP (15,35), abolished the nuclear 8-oxo-dG signal (Figure 2A and Supplementary Figure S1B). These data showed that LSD1 is the source of nuclear ROS and the cause of G oxidation. As a logical extension of the previous experiment, we analyzed EMT engagement in MCF10A cells upon LSD1 depletion. Confocal analysis of cortical E-cadherin and actin stress fibers confirmed that LSD1 silencing inhibited EMT induced by TGF-β1 (Figure 2C).

The data reported above show that the nuclear TGF-β1-induced ROS were produced by LSD1 (Figure 2B) and that LSD1 depletion inhibited the EMT/TGF-β1 response (Figure 2C). To document the impact of LSD1 on the expression of the genes involved in EMT, we measured SNAI1 and other TGF-β1-induced or -repressed mRNA levels in LSD1-depleted cells. LSD1 knockdown abolished SNAI1 induction as well as the induction of CDH2, VIM and PAI by TGF-β1 (Figure 2D and Supplementary Figure 2-Stat.1–2). Moreover, LSD1 silencing reversed TGF-β1 repression of genes such as WIF1, cMYC, CDH1 and DSP (Figure 2E and Supplementary Figure 2-Stat.3–4). Furthermore, treatment of the cells with TCP, the monoamine oxidase inhibitor (15,35), inhibited the TGF-β1-mediated induction of SNAI1, CDH2, VIM and PAI and prevented the TGF-β1-mediated repression of WIF1, MYC, CDH1 and DSP, replicating LSD1 depletion effects (Figure 2F, Supplementary Figure 2-Stat.5–6, Figure 2G and Supplementary Figure 2-Stat.7–8). The effects of LSD1 shRNAs on SNAI1 and WIF1 induction or inhibition of transcription were prevented by expressing exogenous LSD1 (Supplementary Figure S2).

TGF-β1 induced a genome-wide temporal oxidation wave that modified the DNA

Confocal microscopy revealed that TGF-β1 treatment induced 8-oxo-dG nuclear signal; it was eliminated by LSD1 depletion (Figure 2B). To confirm this finding with a different technique, we subjected the DNA of TGF-β1-exposed cells to MS analysis to identify and quantify the modified DNA bases (see the ‘Materials and Methods’ section). Figure 3A shows that 8-oxo-dG levels were induced by TGF-β1 after 60–90 min (Figure 3A and Supplementary Figure 3-Stat.1A). Induction of 8-oxo-dG by TGF-β1 was prevented by DPI or LSD1 depletion (Figure 3A and Supplementary Figure 3-Stat.3A and 4A). Apocynin, a nonspecific antioxidant, that inhibits peroxides (36) did not modify the 8-oxo-dG levels induced by TGF-β1 (Figure 3A, right panel, and Supplementary Figure 3-Stat.3).

Analysis of the other modified DNA bases revealed that the TGF-β1 effects on DNA also involve deoxy-C metabolites: 5-hmdC and 5-methyl-dC (Figure 3B and C, and Supplementary Figure 3-Stat.1-2B and C). Furthermore, 5-hmdU or deoxy-U levels were reduced by DPI, data that suggest oxidation by NOX enzymes favors C and T deamination (Figure 3D and E, and Supplementary Figure 3-Stat.1–2D and E and 3-Stat.3).

To our knowledge, these data provide the first demonstration that TGF-β1 induces a genome-wide oxidation wave that modifies DNA to initiate the EMT transcriptional program.

Common and differential chromatin changes induced by TGF-β1 at repressed and activated promoters

The above data demonstrate that LSD1 is required for DNA oxidation and for both the activation and repression of TGF-β1-induced transcription, but they do not clarify the mechanism. TGF-β1-activated receptors I and II induce a series of events that culminate in the phosphorylation and nuclear accumulation of the phosphorylated SMAD2/3 complex (37). We analyzed by ChIP the earliest chromatin changes (30–60 min) at the promoter sites of the two prototypic genes induced (SNAI1) or repressed (WIFI1) by TGF-β1. The analyzed genomic regions contain sequences recognized by the complex SMAD2/3 (37,38) (A. Pezone, unpublished data). Upon TGF-β1 challenge, phosphorylated SMAD2/3 (pSMAD2/3) complex was recruited to the SNAI1 and WIFI1 promoters and steadily accumulated at 30 min and disappeared by 90 min (Figure 4A and B). LSD1 was also recruited after 30 min at both promoters, but at 90 min, compared to 0- or 60-min treatment, LSD1 accumulated at WIFI1 promoter (Figure 4A and B, and Supplementary Figure 4-Stat.1). The recruitment at 30 min of
Figure 2. Nuclear G oxidation induced by TGF-β1 through the histone demethylase, LSD1. (A) Immunofluorescence detection of 8-oxo-dG by fluorescein-tagged anti-8-oxo-dG antibody in cells pretreated with 5 mM NAC or 5 μM DPI for 1 h before stimulation with 10 ng/ml TGF-β1 for 30 min. Nuclei
LSD1 and pSMAD2/3 suggests that these proteins form a transient complex at both promoters. The association between LSD1 and pSMAD2/3 was shown by immunoprecipitation with antibodies to LSD1 and immunoblot with anti-pSMAD2/3 (Figure 4D) or immunoprecipitation with antibodies to an exogenous FLAG–LSD1 and western blot with anti-pSMAD2/3 antibodies in extracts derived from transfected cells (Figure 4E). The association was also shown by ChIP and re-ChIP experiments of SNAI1 and WIF promoters with antibodies against LSD1 on pSMAD2/3 (Figure 4G and H, and Supplementary Figure 4-Stat.2). Both assays (immunoprecipitation and ChIP) revealed that non-phosphorylated SMAD2/3 did not bind LSD1. However, the absence of signal with the antibody to total SMAD2/3 suggests that only a fraction of SMAD2/3 binds LSD1 (Figure 4F). The complex that contained pSMAD2/3 was transient (30 min) and rapidly disappeared at the SNAI1 promoter between 60 and 90 min after TGF-β1 treatment (Figure 4A). LSD1 reappeared at the WIF1 promoter 90 min after TGF-β1 treatment, without pSMAD2/3 (Figure 4B and Supplementary Figure 4-Stat.1C and D). The notion that pSMAD2/3 is recruited to promoters repressed by TGF-β1 is not novel, although it has not been connected to a specific mechanism (36). Phosphorylated SMAD2/3 work as pioneer factors that transiently bind DNA sites present in TGF-β1-induced or -repressed promoters in different cell types (38,39). Depending on the concomitant presence of other signals, such as ROS and stress-kinase-induced repressors (40,41), some genes are repressed while others are activated. For example, the repressor SNAI1 was induced at 30 min after TGF-β1 challenge and recruited at 90 min to the WIF1 promoter, where it formed a complex with LSD1 and silenced the expression of the gene (Supplementary Figures S3 and S3-Stat.1A–C). With the same timing, the co-repressors NCoR1 and HDAC3 accumulated at the WIF promoter (Supplementary Figures S4A and B, and S4-Stat.1).

Collectively, these data demonstrate that the pSMAD2/3–LSD1 complex very early upon TGF-β1 challenge targeted the chromatin of TGF-β1-activated and -repressed promoters (Figure 4A and B). Ninety minutes after TGF-β1 challenge, SNAI1 and LSD1 accumulated preferentially at the repressed promoters (Figure 4A and B). Since nuclear G oxidation was the earliest modification we observed upon TGF-β1 challenge, we measured recruitment of OGG1 and the apurinic/apyrimidinic endonuclease 1 (APE1) to the SNAI1 and WIF1 promoters. OGG1 recognizes 8-oxo-dG and APE1 targets, excises and repairs the abasic site generated by OGG1 (42). Since LSD1 demethylates H3K4 and K9m2, we also determined at the same sites the H3K4-K9 methylation status and the recruitment of H3K9 and H3K4 methyltransferases, SUV39 and SET7/9 (43,44).

In summary, OGG1 accumulated at 30 min at the SNAI1 promoter (Figure 5A and Supplementary Figure 5-Stat.1A–C) and at 30 and 90 min at the WIF1 promoter (Figure 5B and Supplementary Figures 5-Stat.1D–F and S3B), a pattern that mirrored the timing of LSD1 recruitment at the same sites (Figure 4A and B). The presence of APE1 at both promoters 90 min after TGF-β1 treatment suggests a prolonged repair step, such as long-patch base excision repair, following dG oxidation (45,46) at both promoters. At WIF1 promoter, the second oxidation peak is marked by OGG1 and overlaps with the rise of 8-oxo-dG levels at 90 min detected by MS (Figure 3A). With the same timing, NCoR1 and HDAC3 accumulate at the WIF1 promoter (Supplementary Figure S4B). Since the TGF-β1-induced dG oxidation is dependent on LSD1 (Figures 2 and 3) and LSD1 demethylates H3K9 and H3K4m2 (12,13), we have asked whether LSD1 contributes to the H3K9 and H3K4 methylation status of SNAI1 and WIF1 chromatin during the early response to TGF-β1. Histone H3 methylation changes induced by TGF-β1 at the SNAI1 and WIF1 promoters are produced by the recruitment of demethylases and methyltransferases (43,44). Following TGF-β1 stimulation, both K9 and K4 methyltransferases, SUV39 and SET7/9, respectively, were recruited to the SNAI1 and WIF1 promoters with similar kinetics (Supplementary Figures S5 and S5-Stat.1A–C).
Figure 3. 2D UPLC–MS/MS analysis of DNA derived from cells stimulated with TGF-β1 for various periods of time (left panels) or for 60 min in the presence of DPI, apocynin, shCTRL or shLSD1 (right panels). The technical and statistical details are indicated in the ‘Materials and Methods’ section and in Supplementary Figure 3-Stat.1–5. Results are the mean of at least three experiments in triplicate (n ≥ 9). Left panels: Multivariate analysis of repeated measures (RM-MANOVA) was used to determine differences in multiple dependent variables over time. Pairwise comparison was performed with Student’s t-test. *P < 0.01 versus time zero. The actual P-values are shown in Supplementary Figure 3-Stat.1–2. Right panels: The effect of TGF-β1 in the various treatment conditions was evaluated with the Wilcoxon signed-rank test (Supplementary Figure 3-Stat.3). Differences between three or more groups were determined by MANOVA (Supplementary Figure 3-Stat.4–5).
Figure 4. TGF-β1-induced phospho-SMAD2/3 interact with LSD1 and form a transient complex at the WIF1 and SNAI1 promoters. Chromatin from MCF10A cells stimulated with 10 ng/ml TGF-β1 for 30, 60 and 90 min was immunoprecipitated with antibodies to SMAD2/3 or phospho-SMAD2/3, anti-LSD1. DNA was extracted and amplified with primers corresponding to SNAI1 (A) or WIF1 (B) promoters or as control the promoter region of human TSH receptor gene, TSHR (C). Normal mouse or rabbit pre-immune IgG was used as a negative control (horizontal lines in each graph). The bar graph shows the qPCR signals in immunoprecipitates, normalized to input DNA from each sample. Multivariate analysis of repeated measures (RM-MANOVA) was used to determine differences in multiple dependent variables over time (Supplementary Figure 4-Stat.1A, C and E). Differences between three or more samples were determined by one-way ANOVA test; pairwise comparison was performed with Student’s t-test (Supplementary Figure 4-Stat.1B, D and F). *P < 0.01, in 90-, 60- or 30-min TGF-β1-stimulated cells versus control. **P < 0.01, in 90-min compared to 60-min TGF-β1-treated cells. The primers used are shown in Supplementary Table S1. (D) Immunoprecipitation of LSD1 and immunoblot with anti-phospho-SMAD2/3, LSD1, SMAD2/3 and LAMININ β1 antibodies in cells exposed for 30 min to TGF-β1. (E) Immunoprecipitation of exogenous FLAG–LSD1 with anti-FLAG antibodies and immunoblot with anti-phospho-SMAD2/3 and SMAD2/3. (F) Immunoprecipitation of SMAD2/3 and immunoblot with LSD1, SMAD2/3 and tubulin antibodies in cells exposed for 30 min to TGF-β1. ChIP of SNAI1 (G), WIF1 (H) and TSHR (I) gene promoters with antibodies versus phospho-SMAD2/3. The DNA immunoprecipitated was subjected to re-ChIP with anti-LSD1 antibody. Data are reported as mean ± SD of three experiments in triplicate (n = 9). Pairwise comparison was performed with Student’s t-test. *P < 0.01, TGF-β1-stimulated cells versus control (Supplementary Figure 4-Stat.2). The primers are shown in Supplementary Table S1.
Figure 5. Chromatin changes induced by TGF-β1 at repressed and activated promoters. Chromatin from MCF10A cells stimulated with 10 ng/ml TGF-β1 for 30, 60 and 90 min was immunoprecipitated with antibodies to OGG1 or APE1 (A–C), anti-H3K4me2/3 (D–F), anti-H3K9me2/3 (G–I) or anti-H3 (J–L). DNA was extracted and amplified with primers corresponding to SNAI1 (left panels) or WIF1 (middle panels) promoters or as control the promoter region of human TSH receptor gene, TSHR (right panels). Normal mouse or rabbit pre-immune IgG was used as a negative control (horizontal lines in each graph). The bar graph shows the qPCR signals in immunoprecipitates, normalized to input DNA from each sample. Data are reported as mean ± SD of three experiments, performed, at least, in triplicate (n ≥ 9). Multivariate analysis of repeated measures (RM-MANOVA) was used to determine differences in multiple dependent variables over time (Supplementary Figure 5-Stat.1–3A, D and G). Differences between three or more samples were determined by one-way ANOVA test (Supplementary Figure 5-Stat.1–3). Pairwise comparison was performed with Student’s t-test. *P < 0.01, in 90-, 60- or 30-min TGF-β1-stimulated cells versus control. **P < 0.01, in 90-min compared to 60-min TGF-β1-treated cells. The primers used are shown in Supplementary Table S1.
ures S4D and E, and S4-Stat.2). In contrast, H3K9me2–3 and H3K4me2–3 levels at the same promoters varied considerably between 30 and 90 min after TGF-β1 treatment (Figure 5D and E). Briefly, the most prominent H3K4 and K9 methylation changes upon TGF-β1 treatment are the following: (i) A time-dependent increase of H3K4me3 at SNAI1 promoter (Figure 5D and E) due to methylation of H3K4me2 (Supplementary Figures S4D and S4-Stat.2). (ii) A progressive reduction of H3K4me2 and H3K4me3 at WIF1 promoter, due to H3K4me2 and H3K4me3 demethylation, notwithstanding the rise at 90 min of the methyltransferase SET7/9 (Figure 5E, and Supplementary Figures S4E and S4-Stat.2). These changes are consistent with the induction of SNAI1 and the inhibition of WIF1 expression by TGF-β1 treatment. (iii) The monoamine oxidase inhibition activity by TCP increased H3K4me2 levels at the WIF1 (Supplementary Figures S5B and S5-Stat.1–2) and the levels of H3K9m2 at the SNAI1 promoters (Supplementary Figures S5D and S5-Stat.3), notwithstanding the recruitment of the methyltransferases SUV39 and SET7/9 (Supplementary Figure S4E). Although TCP inhibits nonspecifically monooxidases, it increases the global levels of H3K4me2, which is the natural LSD1 substrate (Supplementary Figure S5J). The methylation changes induced by TCP are consistent with the inhibition of TGF-β1 induction or repression of SNAI1 and WIF1 transcription as in cells in which LSD1 has been depleted (Figure 2D–G). The methylation changes induced by TCP were associated with a significant depletion of nucleosomes at the WIF1 promoter, as shown by loss of H3 at 30 and 90 min (Supplementary Figures S5H and S5-Stat.4).

Collectively, the levels of H3K9m2 and H3K4m2, the timing of LSD1 recruitment and the effects of TCP on the H3K4m2 and H3K9m2 levels suggest that LSD1 recruitment and activity mediate the induction of SNAI1 and the repression of WIF1 by TGF-β1 (Figure 4A and B).

Also, the H3K9me2 reduction of H3K9me2 levels at the chromatin of the SNAI1 promoter in the first 30 min without a concomitant increase of H3K9me3 (Figure 5G) suggests that H3K9me2 is the main LSD1 substrate at the SNAI1 promoter, as previously shown at the chromatin of genes induced by nuclear receptors (12,13). Finally, the steady H3K4me3 increase and the concomitant reduction in H3K4me2 after 60 min of TGF-β1 stimulation at the SNAI1 promoter can be explained by the recruitment of the methyltransferase SET7/9 following 30 min of TGF-β1 stimulation (Figure 5D and Supplementary Figure S4D). The H3K9me2 chromatin modification induced by LSD1 (Figure 5G) was consistent with the loss of TGF-β1 induction and repression of target genes upon LSD1 depletion or TCP exposure (Figure 2D–G).

Cooperation between two histone demethylases, LSD1 and JMJD2A, mediated TGF-β1-induced EMT

LSD1 demethylates H3K4me2 and H3K9me2 but not trimethyl K4 or K9 (47). To account for the H3K9me3 changes at the SNAI1 and WIF1 promoters, we determined the effects of TGF-β1 on a different type of histone demethylase, KDM4A (also called JMJD2A or JHDM3A), a dioxygenase that demethylates H3K9me3 and H3K9me2 in an oxidation reaction that requires Fe²⁺ (48). Notably, KDM4A/JMJD2A is exquisitely sensitive to oxygen levels and displays a graded response to oxygen depletion (49). JMJD2A depletion inhibited the nuclear 8-oxo-dG signal and TGF-β1-induced EMT (Supplementary Figure S6A–D). JMJD2A, upon TGF-β1 stimulation, was recruited to the SNAI1 promoter and marginally to the WIF1 promoter 30 min after treatment (Supplementary Figure S6I and J). JMJD2A depletion reduced the expression of SNAI1, CDH2, VIM and PAI1 induced by TGF-β1 (Supplementary Figure S6E and F), but did not change TGF-β1-mediated repression of WIF1, MYC, CDH1 and DSP (Supplementary Figures S6G and H, and S6-Stat.1–7). The same phenotype was observed by inhibiting JMJD2 activity with JIB04 (50,51) (Supplementary Figures S3 and S6-Stat.1–7). We conclude that JMJD2A cooperates with LSD1 to activate TGF-β1-induced EMT genes and it is not required for TGF-β1 silencing of epithelial genes.

Repression of transcription by TGF-β1 is dependent on both nuclear and cytoplasmic ROS

Collectively, the above data demonstrated that LSD1 is involved in both TGF-β1 induction and repression of transcription depending on the histone methylation code and partnership with other factors, such as pSMAD2/3 or SNAI1 recruited at activated or repressed promoters, respectively. ROS are central to both activation and repression of transcription by TGF-β1 (Figure 1F and G), and their source might be relevant for the efficiency of the final transcriptional response.

Early (30–90 min) TGF-β1-mediated transcription induction or repression can be explained by the activation of two major signaling pathways, both of which require LSD1. The first, which leads to transcription activation, includes nuclear ROS, pSMAD2/3, LSD1 and JMJD2A/KDM4A. The second, which represses transcription, is established by the LSD1–SNAI1 complex, and, eventually, maintained by the co-repressors NCoR1–HDAC3 (Figures 1F and G, and 2A, F and G).

To get insight into the other signaling element(s) that cooperate with LSD1 to mediate TGF-β1 transcription repression, we exposed TGF-β1-treated cells to several chemical inhibitors to identify the primary source of the ROS: (i) rotenone (52), an inhibitor of mitochondrial complex I, which has been linked to TGF-β1-induced ROS; (ii) apocynin, which is a generic antioxidant, which does not modify nuclear o xo-dG accumulation by TGF-β, but inhibits peroxides (36,53); and (iii) SP600125, a specific inhibitor of JNK (54), the ROS-activated kinase that mediates WIF1 silencing by ROS and ATM (41,55).

Although the ROS inhibitors we have used were not as specific as the silencing probes, we can extract the following information: (i) Apocynin prevented repression of transcription by TGF-β1 (Supplementary Figure S7D) but did not reduce o xo-dG detected by MS (Figure 3A). (ii) DPI prevented repression of transcription by TGF-β1 (Figure 1G) and reduced nuclear o xo-dG detected by MS (Figure 3A). (iii) Both drugs did not influence the induction of transcription by TGF-β1 at early times, while an ROS scavenger like NAC prevented both induction and repression of tran-
scription by TGF-β1 (Figure 1F and Supplementary Figure S6C). At 24 h, DPI inhibited NOX4 and SNAI1 induction by TGF-β1 (Supplementary Figure S1A and B) (26,28,29). (iv) SP600125, the JNK inhibitor, did not influence TGF-β1-induced gene expression (Supplementary Figure S7A), but prevented TGF-β1 silencing of WIF1, MYC, CDH1 and DSP genes (Supplementary Figures S7B and S7-Stat.1–5), data that replicate the effects of DPI (Figure 1F–G).

(v) Unexpectedly, inhibition of mitochondrial complex I by rotenone did not affect early TGF-β1 repression of EMT genes, data that suggest mitochondrial ROS do not mediate TGF-β1 early inhibition of transcription, notwithstanding the presence of SNAI1 repressor (Supplementary Figure S7C and D). These results imply that at 2 and 24 h there are two pharmacologically different sources of ROS induced by TGF-β1. At 24 h, the major player is NOX4 and this has been demonstrated in many laboratories and we now know that NOX4 is induced at 24 h, and its expression is inhibited by DPI (Supplementary Figure S1A) (25,26). At early times, LSD1 is the main source of nuclear ROS and is essential for both induction and repression of transcription by TGF-β1. However, the data shown in Supplementary Figure S7 suggest that another early source of ROS (JNK) is necessary for the early inhibition of transcription by TGF-β1. Here, we provide an experimental tool to separate the induction and repression of transcription by TGF-β1 at early (hours) and late (days) times. Notably, SNAI1, which is considered essential for TGF-β1/LSD1 repression of epithelial genes (31) in the presence of DPI, did not repress early transcription (Figure 1F and G, and Supplementary Figure S7C and D). We believe that this information is relevant because SNAI1 is considered the master repressor of transcription induced by TGF-β1 (31).

**DISCUSSION**

The role of ROS in the regulation of EMT in several cell types, including tumor models, is the subject of active investigation (5,6). Oxidative stress following the exposure of cells to microenvironmental insults, such as hypoxia or contact with stromal or inflammatory cells (i.e. CAFs or macrophages), has been causally associated with EMT activation. In redox-dependent EMT, there are at least two molecular axes driven by pro-inflammatory/pro-oxidant sensors: one involves HIF-1 and NF-κB and the other involves the Rac-1 small GTPase, NOX and Src kinase (5,6,8).

With our data, we identified another element that plays a relevant role in the EMT transcriptional program: ROS induced by TGF-β1 in the nuclear compartment by the histone demethylase LSD1 (Figure 2B) were essential for the induction, repression of transcription and the initiation of the TGF-β1-mediated EMT program (Figures 1 and 2). LSD1 was targeted to promoter sites activated or repressed by TGF-β1 by phospho-SMAD2/3 (Figure 4A and B).

**Phosphorylated SMAD2/3 and LSD1 targeted TGF-β1-induced and -repressed genes**

Induction of phospho-SMAD2/3 by TGF-β1 is a central event (29,37) that initiates the EMT transcriptional program. Although SMAD2 does not directly bind to DNA, the SMAD2/3 complex recognizes specific, essential sequence motifs present at the transcription start site (TSS) and enhancers of many TGF-β1-induced or -repressed genes (29,37,38). These SMAD2/3 signatures were also present in the TGF-β1-induced or -repressed genes analyzed here. Phosphorylated SMAD2/3 work as a pioneer factor that marks several genomic sites of TGF-β1-induced or -repressed promoters in different cell types (38–40). Depending on the cell context and the presence of other factors, some genes will be induced and others will be inhibited. Our data suggest that the cellular context is represented by different ROS sources.

**TGF-β1 genomic oxidative signature initiated the EMT program**

TGF-β induces nuclear migration of SMAD proteins, which are phosphorylated by several kinases, including stress kinases, JNK and p38 (41). PhosphoSMAD2/3 complex binds and targets LSD1 to the promoters of genes such as SNAI1 and WIF1, which are induced or repressed, respectively, by TGF-β1 (Figure 4A and B). LSD1 induced a peak of 8-oxo-dG, detectable by recruitment of OGG1 and APE1 at SNAI1 and WIF1 promoters (Figure 5A and B). Local dG oxidation at these two promoters reflected a massive oxidation wave across the entire genome induced by TGF-β1, detectable by MS and dependent on LSD1 (Figure 3). G and C are oxidized in vivo by two mechanisms. In the first, G is oxidized by a Fenton reaction catalyzed by LSD1, which generates hydrogen peroxide (33), and partly by KDM4A/JMJD2A (dependent on Fe2+) (48). In the second, C is oxidized by ten–eleven translocation (TET) enzymes that use ketoglutarate as a cofactor and Fe2+ as an electron donor (similar to all JMJ enzymes). Hydroxymethylation of 5mC is the obligatory step to C demethylation and transcription activation (56). Although G oxidation was the major genomic change induced by TGF-β, we notice that the global levels of 5OHmdC were higher in cells exposed for 60 and 90 min to TGF-β (Figure 3B).

Collectively, G and C oxidation marks the global transcription changes induced by TGF-β during the initiation of the EMT (Figure 3).

**DNA oxidation and transcription**

Targeted oxidation of DNA at enhancers, TSSs and 3’ ends of genes is essential for the formation of chromatin transcription loops and the initiation complex (12,13,57). In fact, transcription loops induced by retinoic acid or estrogens do not form when LSD1 or OGG1 is downregulated or oxidation (12,13) and/or repair (12,58) are prevented. The 8-oxo-dG peak induced by 30–90-min TGF-β1 treatment corresponds to the time required to stabilize chromatin loops during initiation of transcription by estrogens or retinoic acid. These loops are RNase H sensitive, recruit LSD1, OGG1 and cohesin RAD21, and connect the enhancer–TSS and polyA sites (12,13,57). The oxidized sites form single-strand nicks that recruit topoisomerase II beta, RNA polymerase II including BER (OGG1–APE1).
and NER (12,13). These events occur in a precise temporal window (12,13,57), and histone demethylation and G oxidation are the first critical steps (12,13). We note that the punctuated oxo-dG signal detected by confocal microscopy (Figure 2A and B) resembles the interchromatin granules punctuated oxo-dG signal detected by confocal microscopy (12,13). The TGF-β signal is continuous and persistent (Figure 5). In contrast, during the PDGF-β induced EMT, the TGF-β signal is continuous and persistent (Figure 5), and the induced proteins disappear after 3 h to 9 days, the EMT phenotype (63); see also the comments in (64).

Initiation, establishment and maintenance of TGF-β1-induced EMT

We wish to stress that our observation window of TGF-β1 effects is narrow (0–90 min) compared to the timing of TGF-β1 action analyzed in the majority of published studies, which show that NOXs, and specifically NOX4, are essential for stable EMT (24–48 h to 15 days) (28,29). Our results complement and do not contradict the published data because the changes we document are transient and reversible. This is shown by the activation of the canonical TGF-β1/SMAD pathway, which initiates a transient wave of SMAD2/3 phosphorylation (Figure 4) (60,62). In fact, if TGF-β1 is removed after 3 h to 9 days, the EMT phenotype and the induced proteins disappear. Higher doses and continuous exposure to TGF-β1 stabilize irreversibly the EMT phenotype (63); see also the comments in (64).

Our data show that the early events following TGF-β1 exposure are relevant for the initiation of the EMT program and suggest that the early (1–2 h) and late (24 h) events triggered by TGF-β1 to induce EMT are mediated by a combination of different players.

We propose that the TGF-β1 EMT transcriptional program is carried out in two distinct temporal phases: the first (initiation), between 30 and 90 min, initiates the program by modifying DNA at multiple sites. On a smaller scale, the complex pSMAD2/3–LSD1 primes genomic sites that eventually will be stably activated or repressed only if the TGF-β1 signal is continuous and persistent (Figure 5) (60,63). The second phase (maintenance), between 1 and 14

Repression of EMT genes requires synchronous signaling from JNK and LSD1

LSD1 is neutral with regard to activation or repression, but it is a coincidence sensor at the WIF1 promoter because its presence reflects a steady ROS signal leading to JNK activation and a nuclear oxidation signal, both necessary to repress transcription. In the absence of the ROS–JNK signal, LSD1 binds pSMAD2/3, activates SNAI1 transcription and marks transiently WIF1 promoter 30 min after TGF-β1 treatment. In the presence of a persistent ROS signal (90-min TGF-β1 treatment), LSD1 binds to SNAI1–NCoR1–HDAC3 and induces a new demethylation-oxidative wave by removing activation marks from chromatin (Figure 3E). In fact, while LSD1 depletion inhibited both genes that are positively and negatively regulated by TGF-β1, transcription repression was achieved only in the presence of a concomitant and strong DPI/JNK–ROS signal. The early repressor induced by TGF-β1, SNAI1, does not repress transcription in the absence of this signal (Figure 1G and Supplementary Figure S7D). Instead, nuclear oxidation by LSD1 in the absence of JNK–ROS was sufficient to activate
days, consolidates and stabilizes TGF-β1 transcriptional repression or induction (28,29). Although we did not identify precisely the source of the DPI-sensitive ROS in the initiation phase (90-min TGF-β1 treatment), we suggest that JNK–NCoR1 and HDAC3 are essential for the stabilization and maintenance of early TGF-β1 repression of transcription (55).

Figure 6 shows a simplified scheme of the early TGF-β1 signals. It describes the induction or repression of transcription of EMT genes. Specifically, the repressor complex nucleated by LSD1 at the WIF1 promoter inhibits transcription only when two temporally close and spatially distributed signals are present, i.e. nuclear (LSD1) and cytoplasmic (NOX–JNK) ROS. This repressor complex is an ROS sensor and is linked to the redox state of the cell (55).

We speculate that the source of cytoplasmic ROS is linked to the redox state of the cell (55). We suggest that the source of cytoplasmic ROS is linked to the repressor complex, which substantially modifies the response to TGF-β1 in a cell-specific fashion.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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