Complexities in the role of acetylation dynamics in modifying inducible gene activation parameters

Samantha Carrera, Amanda O’Donnell, Yaoyong Li, Karol Nowicki-Osuch*, Shen-Hsi Yang, Syed Murtuza Baker, David Spiller and Andrew D. Sharrocks*+

Faculty of Biology, Medicine and Health, University of Manchester, Michael Smith Building, Oxford Road, Manchester M13 9PT, UK

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ABSTRACT

High levels of histone acetylation are associated with the regulatory elements of active genes, suggesting a link between acetylation and gene activation. We revisited this model, in the context of EGF-inducible gene expression and found that rather than a simple unifying model, there are two broad classes of genes; one in which high lysine acetylation activity is required for efficient gene activation, and a second group where the opposite occurs and high acetylation activity is inhibitory. We examined the latter class in more detail using EGR2 as a model gene and found that lysine acetylation levels are critical for several activation parameters, including the timing of expression onset, and overall amplitudes of the transcriptional response. In contrast, DUSP1 responds in the canonical manner and its transcriptional activity is promoted by acetylation. Single cell approaches demonstrate heterogenous activation kinetics of a given gene in response to EGF stimulation. Acetylation levels modify these heterogenous patterns and influence both allele activation frequencies and overall expression profile parameters. Our data therefore point to a complex interplay between acetylation equilibria and target gene induction where acetylation level thresholds are an important determinant of transcriptional induction dynamics that are sensed in a gene-specific manner.

INTRODUCTION

Histone modifications play an important role in controlling the levels of gene transcription. High levels of histone acetylation are associated with transcriptionally active genes where the modifications decorate the nucleosomes surrounding both proximal and distal regulatory elements (1–3). It is generally assumed that high levels of histone acetylation are associated with active transcription and reciprocally, low levels or hypoacetylation are associated with repressed or inactive genes. However, evidence for a more dynamic model has been gathered, chiefly through the use of histone acetylation/deacetylation inhibitors, whereby histone acetylation dynamics rather than overall levels are critical in gene activation (4–6). It is important to note that in addition to histones, other transcriptional regulatory proteins can be targeted by acetylation to influence their activity as exemplified by studies on MMTV promoter activation (7). Thus, acetylation dynamics may have an impact beyond simply through modifying histones associated with regulatory elements.

Lysine acetylation levels are governed by the combined actions of lysine acetyl transferases (KATs) and lysine deacetylases (KDACs) found at gene regulatory elements. By using acute administration of KAT (5) and KDAC (4) inhibitors, both types of enzymatic activity have been

*To whom correspondence should be addressed. Tel: +44 161 275 5979; Fax: +44 161 275 5082; Email: a.d.sharrocks@manchester.ac.uk

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implicated as important for the inducible activation of genes such as *FOS* and *JUN*, providing weight to the model that acetylation dynamics rather than overall levels are critical for inducible transcription to take place. Furthermore, genome-wide ChIP-seq studies have also shown a role for both KDACs and KATs at active genes providing further support for a role for dynamic histone acetylation in gene activation (8). This has led to the broader recognition that KDACs and their deacetylation activity can have activating roles in addition to their previously established role in transcriptional repression (reviewed in 9). More recent work has implicated KDACs in both gene activation and repression (10). Furthermore, class I KDACs are required for the inducible activation of glucocorticoid receptor target genes (11). As both KATs and KDACs are considered as therapeutic targets in a range of human diseases (reviewed in 12,13), it is therefore important to further understand their mechanisms of action.

Many of the insights provided to date have focused on a relatively small number of genes and, in particular, the growth factor regulated *FOS* and *JUN*. We therefore wished to more broadly assess the role of acetylation dynamics in modifying the activation of growth factor inducible genes. Rather than a simple unifying model, we identified two broad classes of genes; one in which high acetylation activity is required for efficient gene activation, and a second group where the opposite occurs and high acetylation is inhibitory. This points to a more complex model where acetylation levels thresholds are an important determinant of transcriptional induction dynamics that are sensed in a gene-specific manner.

**MATERIALS AND METHODS**

**Cell line creation and culture**

Parental MCF10A cells and cell lines derived from these were grown in DMEM/F12 (Gibco, 11320-033) containing 5% horse serum (Biosera, DH291), 20 ng/ml EGF (Sigma, E1257), 10 μg/ml insulin (Sigma, I0516), 100 ng/ml cholera toxin (Sigma, C9903) and 0.5 μg/ml hydrocortisone (Sigma, H0396) (complete medium). When required cells were seeded or changed into starvation media: DMEM/F12 containing 0.5% horse serum, 10 μg/ml insulin, 100 ng/ml cholera toxin and 0.5 μg/ml hydrocortisone. Cells were left for 48 h in starving media and then treated with EGF where required.

To create MCF10A-EGR2-Luc cells, we used the pX335 CRISPR/Cas9 plasmid (14; obtained via Addgene) and the *EGR2* 3′-end-targetting guide RNA sequences; target 1 (ADS5212, ADS5213) and target 2 (ADS5214, ADS5215). These guides were inserted into plasmid pX335 BbsI cloning site as described previously (15) to create plasmids pX335-EGR2-target1 (pAS4883) and pX335-EGR2-target2 (pAS4884). For donor plasmid EGR2-2A-LucP (pAS4885), the full-length nanoLuciferasePEST gene (pNL1.2 Promega) was PCR-amplified along with a 63 bp 5′-end viral 2A sequence using primers ADS5225/ADS6756 then prepared for Gibson assembly using primers ADS5227/ADS5228. The luciferase gene was then cloned into the BamHI site in the pUC19 backbone vector using the Gibson assembly technique, along with the PCR-amplified 782 bp left homology arm (primers ADS5220/ADS5226 incorporating a single base mutation to remove PAM) and 876 bp right homology arm (primers ADS5229/ADS5230), from genomic DNA for specific integration by homologous recombination. Low passage MCF10A cells were transfected at 80% confluency in a 25 cm² flask using 25 μl FugeneHD (Promega) with 6 μg donor plasmid (pAS4885), 1 μg each CAS9 guide RNA plasmid (pAS4883 and 4884) and 200 ng linear hygromycin resistance gene (Clontech) were used. Transfected cells were selected for by adding hygromycin (50 μg/ml) for 72 h before plating surviving cells onto 10 cm dishes without antibiotic selection. Single cells were seeded in 96-well plates using serial dilutions, inserts were checked using primers ADS5337 and ADS5338.

To create MCF10A-DUSP1-Luc cells, we used AIO-GFP as a source of Cas9-D10A (16; obtained via Addgene) and the *DUSP1* targeting guide RNA sequences (ADS6055, ADS6056, ADS6057, ADS6058) were inserted into the AIO-GFP plasmid to create pAS4863. A HDR template containing the full-length nanoLuciferasePEST gene (pNL1.2 Promega) and *DUSP1* homology arms was synthesised as a double-stranded DNA GBlock (Integrated DNA Technologies). Low passage MCF10A cells were transfected at 80% confluency in a 25 cm² flask using 25 μl FugeneHD (Promega) with 6 μg of pAS4863 and 200 ng of GBlock donor template. Transfected cells were grown for 72 h and then cell sorted into a GFP positive population. GFP positive cells were grown for a further 5 days before seeding single cells in 96-well plates using serial dilutions, inserts were checked using primers ADS6061 and ADS6141. Sequencing of the genomic locus confirmed that a single *DUSP1* allele had been tagged and the other two alleles had deletion of the final 2 and 27 amino acids.

All of the PCR primers used are detailed in Supplementary Table S1.

**RNA extraction and qRT-PCR**

RNA was extracted using the RNeasy plus kit (Qiagen) following the manufacturer’s protocol. Subsequently, RNA samples were quantified using a Nanodrop 2000 (Thermo Scientific) and concentrations were normalized to 20 ng/μl. 40 ng of each sample was used per reverse transcription-quantitative polymerase chain reaction (RT-qPCR) reaction using the QuantiTect SYBR® Green RT-PCR Kit (Qiagen, 204243) on Rotor-Gene Q (Qiagen) real-time PCR machine. When indicated, the nanolitre volume RT-qPCR was performed using the Fluidigm Biomark HD system using EvaGreen chemistry following the manufacturer’s protocol. Six housekeeping genes were included in the analysis (*GLUD1*, *C3*, *HADHB*, *SRSF6*, *GAPDH* and *HPRT1*). The geometric mean of the housekeeping genes was calculated for each sample. Samples were normalized to the mean of the housekeeping genes before calculating DeltaCT. The PCR primers are detailed in Supplementary Table S1. The output data were processed following the default quality protocol. Data points with more than one peak in the melt analysis were discarded.
RNA-seq library preparation

The nuclear RNA-seq data were generated from the EGF-induced MCF10A cells at two time points, 0 min and 30 min as previously described (E-MTAB-5370; 17). Additionally, RNA-seq was performed on whole cell RNA extracts using longer EGF time course (0, 30, 90, 180 min) as previously described (17). Briefly, MCF10A cells were seeded in medium without EGF and with 0.5% horse serum 48 h prior to EGF stimulation. EGF (final concentration 20 ng/ml) was added for indicated times and RNA was extracted using an RNeasy plus kit (Qiagen, 74134) with DNase treatment according to the manufacturer’s protocol. Samples with RNA integrity number (RIN) >9 were used for sequencing library construction with the TruSeq Stranded mRNA sample preparation protocol (Illumina).

RNA-seq analysis

Cufflinks was used to identify differential expression (DE) genes between the two time points 0 min and 30 min (18). The R package edgeR was also used to identify DE genes between 0 min and 30 min (19). In order to obtain the EGF inducible genes (i.e. those genes with significantly higher transcription level at 30 min in comparison with that at 0 min), the same criteria, namely fold change >1.5 and adjusted P-value <0.05, was applied separately on the DE genes obtained from Cufflinks and edgeR, resulting in 168 genes from Cufflinks and 160 genes from edgeR. We merged together the two sets of genes and obtained 212 unique EGF inducible genes (Supplementary Table S2). Among the 212 genes in the list, 169 genes were selected whose mean FPKM value was between 0.5 and adjusted >1.5 and ad-

ChIP-seq data analysis

The raw read data of the three ChIP-seq data in MCF10A cells (H3K27ac, H3K79me2, H2BK120ub) were downloaded from the GEO web site (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE85158). The reads were aligned to the human genome hg19 by Bowtie2. The aligned reads from multiple replicates of one type of ChIP-seq data were pooled. The gene annotations of ENSEMBL version 75 were used to locate the TSSs of genes. In-house R scripts were used to calculate the normalised read count (RPKM) within the promoter region (defined as ±500 bp from the TSS) of the selected genes. R functions were used to draw the box plots in Figure 4.

Chromatin Immunoprecipitation

MCF10A cells (4 × 10^6) were seeded in starvation media 48 hours before induction with EGF. Cells were induced with 20 ng/ml EGF for the desired time. Cells were cross linked for 10 min in 1% Formaldehyde before quenching with 0.125 M glycine for 5 min. Cells were harvested in ice-cold PBS with complete protease inhibitor (Roche) and washed sequentially with ChIP Lysis Buffer I (50 mM HEPES–KOH pH 7.5, 140 mM NaCl, 1 mM EDTA, 10% glycerol, 0.5% Igepal, 0.25% Triton X-100) and ChIP Lysis Buffer II (10 mM Tris–HCl pH 8, 200 mM NaCl, 1 mM EDTA pH 8, 0.5 mM EGTA pH 8.0). The resulting nuclei were re-suspended in ChIP Lysis Buffer III (10 mM Tris–HCl pH 8.0, 100 mM NaCl, 1 mM EDTA pH 8.0, 0.5 mM EGTA pH 8.0, 0.1% Na-Deoxycholate, 0.5% N-lauroylsarcosine). Lysates were sonicated on ice to yield 200–600 bp DNA fragments using a Bioruptor (Diagenode).

Magnetic protein A Dynabeads (Invitrogen) were incubated with 0.5 μg of anti-H3K27ac (Abcam-ab4729), NELF-C/D (Cell signaling Technology; TH1L(D5G6W)-12265) antibodies or non-specific IgG overnight at 4°C. Antibody/beads were washed and incubated with nuclear extracts overnight at 4°C. Immunoprecipitates were washed 5 times with RIPA buffer (50 mM HEPES–KOH pH 7.6, 500 mM LiCl, 1 mM EDTA pH 8.0, 1% Igepal, 0.7% Na-Deoxycholate) and once with TE-NaCl (10 mM Tris–HCl pH 8.0, 1 mM EDTA, 50 mM NaCl). ChIP DNA was eluted from Protein A Dynabeads by adding 150 μl elution buffer (50 mM Tris–HCl pH 8.0, 10 mM EDTA pH 8.0, 1% SDS) and incubating at 65°C. Cross-links were reversed by heating to 65°C overnight, then treating with proteinase K for 1 h at 45°C. Chromatin was cleaned using QiaQuick PCR cleanup columns (Qiagen). Nanolitre volume qPCR was performed using the Fluidigm Biomark HD system using EvaGreen chemistry following manufacturer’s protocol and the PCR primers detailed in Supplementary Figure S2. The output data were processed following the default quality protocol. Data points with more than one peak in the melt analysis were discarded.

Bioluminescence detection in live cells

For analyzing cell population timecourses, MCF10A-EGR2-Luc or MCF10A-DUSP1-Luc cells were seeded into 96-well plates in starvation media 48 h before the experiment. Where required, cells were treated with 1.2 nM 4SC202 (Selleckchem), 100 nM A485 (Selleckchem), 100 nM GNE-781 (20; kindly supplied by Karen Gascoigne, Genentech) or vehicle (DMSO) for one hour prior to induction with EGF. NanoGlo Endurazine substrate (Promega) was added to each well 30 min before EGF induction. Cells were induced with 20 ng/ml EGF (Sigma) and luminescence was measured for 5 s every 5 min in FLUOSTar Omega microplate reader.

For single cell analysis, MCF10A-EGR2-Luc or MCF10A-DUSP1-Luc cells were seeded into a 35/10 MM glass bottom dish (Greiner Bio-One) in starvation media 48 hours before the experiment. NanoGlo Endurazine substrate (Promega) was added to each well 30 min before EGF induction. Where required, cells were treated with 1.2 nM 4SC202 or vehicle (DMSO) for one hour prior to induction with EGF. Images were collected on a Zeiss Axiol Observer A.1 microscope using a 10×/0.5 Fluor objective and an iXon Ultra EMCCD Camera (Andor) camera through Micromanager software v1.4.15. Images were then processed and analysed using Fiji ImageJ (http://imagej.net/Fiji/Downloads).

To identify similar patterns in single cell profiles, principal component analysis (PCA) was conducted on normalised counts of luminescent signals from DUSP1- and EGR2-luciferase transgenes across the seven time-points.
Figure 1. Establishment of a reporter gene assay to study EGF-mediated gene activation kinetics. (A) RNA-seq analysis of EGF inducible genes in MCF10A cells at the indicated time points following EGF addition. Data are shown for genes with FPKM > 0.1 as row z-normalised values. (B) Diagrammatic illustration of the luciferase reporter constructs where nanoluciferase was introduced into the 3' end of genomic EGR2 or DUSP1 loci in MCF10A cells. There is a 2A sequence inserted after their coding regions to allow expression of non-fused nanoluciferase. A PEST sequence is incorporated to allow for rapid degradation. (C) Activation kinetics of EGR2- and DUSP1-luciferase reporters after induction with EGF. Cells were treated with vehicle or the MEK inhibitor trametinib 1 hr prior to EGF addition. Data are the average of three biological replicates (n = 3), shaded area represents ± SEM.

UMAP visualization was then generated by taking the first 10 principal components for DUSP1 and 15 principal components for EGR2.

smFISH

smFISH probes for EGR2 and DUSP1 were designed and ordered using the Stellaris Probe Designer (Biosearch Technologies). MCF10A cells were seeded into glass coverslips in starving media and left to grow for 48 h before inducing them with EGF and treated with inhibitors where required. Media was removed and cells were fixed with formaldehyde solution for 15 min at room temperature, washed and incubated in 70% ethanol for 2 h. Samples were hybridized using 100 nM Stellaris probes for EGR2, or DUSP1 overnight. Coverslips were mounted using Pro-Long Gold antifade Mountant with DAPI (ThermoFisher). Images were acquired on an Olympus IX83 inverted microscope using Blue, Red and Green-Yellow Lumencor LED excitation, a 60×/1.42 Plan Apo objective and the Sedat filter set (Chroma 89000). The images were collected using a R6 (Qimaging) CCD camera with a Z optical spacing of [0.2 μm]. Raw images were then deconvolved using the Huygens Pro software (SVI) and maximum intensity projections of these deconvolved images are shown in the results. Mature mRNA transcripts and transcription site quantification was performed using FISHquant (21).

Luminometer data presented is the average ± SEM of three biological replicates with at least three technical repeats each.

Single cell luciferase data and smFISH data are the combination of at least two biological replicates, statistical significance was calculated using unpaired t test GraphPad PRISM v8.

RESULTS

EGF-inducible gene expression profiles in MCF10A cells

To establish the repertoire of EGF-inducible genes in MCF10A cells, we treated cells with EGF and used RNA-seq to profile mRNA expression. We first generated RNA-seq data at two time points, namely 0 and 30 min to identify genes showing rapid induction. A total of 212 genes exhibited enhanced expression at 30 min (1.5 fold; adjusted P-value < 0.05 Supplementary Table S2) (17). To further interrogate the expression profiles of these genes, we generated another RNA-seq dataset at 4 time points over a 3 h time period (Figure 1A). There are several clearly distinct expression profiles, with a group of genes exhibiting rapid and transient responses while others show more delayed and/or extended responses to EGF stimulation (Figure 1A). To study the mechanisms of induction further we decided to focus on two rapidly induced genes, EGR2 and DUSP1. EGR2, shows a robust transient response to EGF stimulation and DUSP1 shows lower amplitude response (see Figure 2C). To allow real-time continuous monitoring of gene induction kinetics, we used CRISPR-Cas9 editing to create MCF10A reporter lines which contain a fusion of the nanoluciferase gene (22) to one allele of either EGR2 or DUSP1 (Figure 1B; Supplementary Figure S1A

Statistical analysis

Data for qRT-PCR and ChIP-qPCR are presented as means of a minimum of three biological replicates. Two-way ANOVA was performed using GraphPad PRISM v8, statistical significance was determined using the Sidak multiple comparisons test.
Figure 2. Differential effects of KDAC inhibitors on EGF-mediated gene activation. (A) Model showing the point of action of the class I KDAC inhibitor 4SC202 in the acetylation-deacetylation equilibrium. (B) Activation kinetics of EGR2- and DUSP1-luciferase reporters after induction with EGF. Cells were treated with vehicle or 4SC202 1 h prior to EGF addition. Data are the average of three biological replicates (n = 3), shaded area represents ± SEM. (C and D) RT-qPCR analysis of relative mRNA expression profiles of the indicated genes after induction with EGF for the indicated times. Cells were treated with vehicle (Control) or 4SC202 for 1 h before induction. Detailed profiles for EGR2 and DUSP1 are shown in (C). Data represents mean of three biological replicates ± SD; * P-value < 0.05, *** P-value < 0.001. The heatmap in (D) is from RT-qPCR analysis using the Fluidigm Biomark system and shows log₂ fold changes of gene expression caused by 4SC202 administration after EGF induction at the indicated times. Data are the average of three biological replicates (n = 3). Samples are grouped according to similar response profiles to KDAC inhibition. (E) H3K27ac levels at the EGR2 or DUSP1 promoters after treatment with 4SC202 for 1 h prior to adding EGF for 30 min. Data represents the mean of three biological replicates ± SD; * P-value < 0.05.

and B). The nanoluciferase has been engineered by incorporating PEST sequences to create a short half-life of 10–30 min thus enabling its use as a readout for monitoring transient induction kinetics. Importantly, the induction kinetics of the EGR2-luciferase and DUSP1-luciferase fused alleles closely mimicked that of the endogenous transcript (Supplementary Figure S1C and D). Indeed, in keeping with the RNA-seq analysis, the EGR2 reporter exhibited the expected high amplitude and transient luciferase activity kinetics whereas the DUSP1 reporter showed reduced activity following treatment with 4SC202 (23) to block deacetylation activity (Figure 2A), and administered this shortly before (60 min) EGF induction to monitor acute responses. The DUSP1 reporter showed increased activity following treatment with 4SC202 as would be expected by inhibiting deacetylases, which are thought to be repressive in nature (Figure 2B, bottom). However, in contrast, EGR2 reporter activity was severely curtailed by the deacetylase inhibitor (Figure 2B, top). This differential response to deacetylase inhibition was unexpected but this was verified at the mRNA level for both genes by RT-qPCR which more directly detects transcriptional changes albeit with differing kinetics due to the time delay between transcription and protein translation of the luciferase reporter (Figure 2C). Increases in DUSP1 expression were also observed after 4SC202 treatment at the pre-mRNA levels, which peak earlier and are more reflective of nascent transcription (Supplementary Figure S2A). This effect of KDAC inhibition appears specific to EGF inducible genes as acute administration of deacetylase inhibitors has little effect on the expression of non-inducible HADHB control gene (Supplementary Figure S2B). Furthermore, an alter-
native deacetylase inhibitor, sodium butyrate, caused similar effects on EGF-inducible gene expression, extinguishing induction of the *EGR2* reporter, but enhancing *DUSP1* reporter activity (Supplementary Figure S2C). RT-qPCR analysis confirmed that sodium butyrate treatment also reduced *EGR2* mRNA levels (Supplementary Figure S2D).

Given these divergent responses, we widened our analysis to a panel of EGF inducible genes and performed RT-qPCR analysis across the time course consistent with transcriptional derepression profiles, and this yielded a final list of 36 genes after excluding ineffective primer combinations. Three different responses to deacetylation inhibition were observed; group 1 (including *DUSP1*) generally had increased expression across the time course consistent with transcriptional derepression; group 2 showed reduced peak expression, whereas group 3 (including *EGR2*) showed broadly reduced expression across the time course (Figure 2D; Supplementary Figure S3). Similar effects were seen at both the mature RNA and pre-mRNA levels as exemplified by *SOCS3*, *CXCL2* and *EREG*, demonstrating an effect on nascent transcription levels, although in the case of *SOCS3* and *CXCL2* mature mRNAs clear trends were observed but statistical significance was not obtained (Supplementary Figure S3).

To explore the potential effects of deacetylase inhibition on the histone acetylation levels found at gene regulatory elements, we monitored H3K27 acetylation levels at the *EGR2* and *DUSP1* promoters. As expected, significant increases were observed to basal acetylation levels at the *DUSP1* promoter following KDAC inhibition (Figure 2E). EGF stimulation enhanced acetylation at this promoter but no further increase was elicited after KDAC inhibition (Figure 2E). However, deacetylation inhibition has no effect on the steady state acetylation levels at the *EGR2* promoter (Figure 2E). Only the molecular changes at *DUSP1* promoter are therefore consistent with the effects of KDAC inhibition on its transcriptional output. However, for *EGR2*, a simple direct cause and effect relationship on histone acetylation levels at its promoter is hard to establish.

Collectively these data demonstrate that there is a heterogeneous effect of inhibition of class I deacetylases on EGF-mediated gene expression. Individual genes show either enhanced or reduced expression following deacetylase inhibition, rather than just the general increases expected from inhibiting deacetylase-mediated transcriptional repression.

### Acetylation inhibitors cause reciprocal responses to EGF-mediated gene induction profiles

Given the unexpected complexities in response to deacetylation inhibition, we asked whether acute acetylase inhibition causes reciprocal effects on the EGF target gene induction profiles. We focussed on two inhibitors of P300/CBP (KAT3B/KAT3A), A485 which is thought to act broadly on histone acetylation levels at promoter and enhancer elements (24), and GNE-781 which is thought to predominantly act at enhancer elements (Figure 3A; 20, 25). The *DUSP1* reporter activity profile was dampened down after treatment with the A485 inhibitor and also by administration of the GNE-781 inhibitor at the early stages of induction (Figure 3B). These effects on *DUSP1* transcription are consistent with the known association between high levels of histone acetylation and gene activation. In contrast however, the *EGR2* reporter activity is elevated following treatment with either acetylation inhibitor (Figure 3B). These effects on reporter activity were broadly recapitulated by directly measuring *EGR2* and *DUSP1* mRNA levels (Figure 3C). Thus, reciprocal effects are seen on both *EGR2* and *DUSP1* expression kinetics following inhibition of acetylation and deacetylation, with high acetylation activity being associated with increased activation of *DUSP1* but decreased activation of *EGR2*. Importantly, acute administration of acetylase inhibitors has little effect on the expression of non-inducible control genes such as *GLUD1* and *HADHB* (Supplementary Figure S4A) which is consistent with what we observed with deacetylase inhibitors.

We next examined whether this reciprocity is observed more generally across the EGF-activated programme, and found two broad categories of genes; group B which is generally upregulated by inhibiting acetylation (including *EGR2*) and a larger group (A) whose expression is reduced (including *DUSP1*) (Figure 3D). There is generally a high level of consistency in the response to the two inhibitors. Similar effects were seen at both the mature RNA and pre-mRNA levels as exemplified by *SOCS3*, *CXCL2* and *EREG*, demonstrating an effect on nascent transcription levels (Supplementary Figure S4B and C). Reduced acetylation levels would not be expected to result in enhanced transcriptional output, and we therefore asked whether these are the same genes identified as decreasing in activity following deacetylation inhibition. All of the genes showing enhanced expression following acetylation inhibition (group B) are in the two groups (2 and 3) which show either decreased expression at all, or a subset, of time points following deacetylase inhibition (compare Figure 2D and Figure 3D; Supplementary Figure S5A). This reinforces the view that decreased acetylation levels lead to the enhanced expression of a cohort of genes (group B) in response to EGF induction. This differential response to disturbing the acetylation equilibrium, suggests that group A genes may share different properties to group B genes, and examination of their expression profiles demonstrates that group B genes generally show earlier peak induction and a faster decrease back towards basal activity (Supplementary Figure S5B).

Next, we asked whether changes in H3K27 acetylation could be found at the *EGR2* and *DUSP1* regulatory elements which could explain the transcriptional regulatory effects we observed. No changes could be observed in H3K27 acetylation at the *EGR2* promoter following acetylation inhibition (Figure 3E). In contrast, we observe attenuation of the EGF-mediated increases in *DUSP1* promoter acetylation following treatment with either acetylation inhibitor (Figure 3E), which is fully consistent with the reduced transcription levels caused by the same treatment. Taken together with the deacetylase inhibitor results, changes to *DUSP1* promoter acetylation occur as predicted from disturbing the acetylation equilibrium and are sufficient to explain the changes in *DUSP1* expression according to established paradigms linking promoter acetylation to gene
Figure 3. Differential effects of KAT inhibitors on EGF-mediated gene activation. (A) Model showing the point of action of the KAT3A/B inhibitors A485 and GNE-781 in the acetylation-deacetylation equilibrium. (B) Activation kinetics of EGR2- and DUSP1-luciferase reporters after induction with EGF. Cells were pre-treated with vehicle (Control), A485 or GNE-781 prior to EGF addition. Data are the average of three biological replicates ($n = 3$), shaded area represents ± SEM. (C and D). RT-qPCR analysis of relative mRNA expression profiles of the indicated genes after induction with EGF for the indicated times. Cells were treated with vehicle (Control) or GNE-781 1 h prior to EGF addition. Detailed profiles for EGR2 and DUSP1 are shown in (C). Data represents mean of three biological replicates ± SD; * $P$-value < 0.05, *** $P$-value < 0.001. The heatmap in (D) is from RT-qPCR analysis using the Fluidigm Biomark system and shows log2 fold changes of gene expression caused by KAT inhibitor administration prior to EGF induction at the indicated times. Data are the average of three biological replicates ($n = 3$). Samples are grouped according to similar response profiles to KAT inhibition. (E) ChIP-qPCR analysis of H3K27ac levels at the EGR2 or DUSP1 promoters after induction with EGF for 30 min after prior treatment with vehicle (Control), A485 or GNE-781. Data represents mean of three biological replicates ± SD; * $P$-value < 0.05.

In summary, we observe a broad degree of reciprocity in effects of manipulating the acetylation equilibrium on gene expression levels, although two different categories of inducible genes can be identified; those that show the expected decrease in expression when acetylation activity is increased and those that show the opposite behaviour to manipulating the acetylation-deacetylation equilibrium.

Gene regulatory features of differentially sensitive inducible gene categories

Having identified two classes of EGF inducible genes that are differentially sensitive to changes in the acetylation equilibrium, we asked whether they had any particular regulatory features which characterised each group. We therefore first examined the basal transcription levels of group 1 (activated by KDAC inhibition) and group 3 (repressed by KDAC inhibition) genes. Group 1 genes had significantly higher basal transcriptional levels prior to EGF stimulation than group 3 genes (Figure 4A) suggesting that group...
3 genes had lower intrinsic activity. We therefore examined the histone marks surrounding the promoter regions of the two gene categories by interrogating published ChIP-seq data from MCF10A cells (26). Higher H3K27ac was observed in group 1 genes (narrowly below statistical significance) and significantly more H3K79me2 and H2BK120ub was also observed in this group of genes (Figure 4B). Example loci for group 1 genes (DUSP1, IER5, KLF6) and group 3 genes (EGR2, NR4A1 and KLF2) clearly illustrate the differential presence of these marks at the two different gene categories (Figure 4C; Supplementary Figure S6A). All of these marks are associated with transcriptional activation and extend into the coding region where the presence of H3K79me2 and H2BK120ub is consistent with ongoing transcriptional elongation (27,28). Given these associations with transcriptional elongation and the previous links between KDAC inhibition and the binding of the transcriptional elongation regulatory factor NELF to promoters (29) we asked whether KDAC inhibition affected NELF association with EGF regulated gene promoters. Reduced binding under basal conditions was observed at the EGR2 promoter but not at the DUSP1 promoter following KDAC inhibition (Supplementary Figure S6B). However, reciprocal effects were not observed following KAT inhibition and again decreased NELF binding was observed at the EGR2 promoter, indicating that changes to NELF binding are not the underlying molecular causes of the differential susceptibility to KDAC/HAT inhibitors.

Differential inducible gene activation susceptibility to altering the lysine acetylation equilibrium is therefore associated with basal activity status, both at the transcriptional level and the chromatin state surrounding the promoter. More active basal expression is associated with reduced induction levels following reductions in acetylation activity.

The impact of acetylation activity levels on transcriptional induction parameters

Having established the impact of acetylation levels on bulk cell populations we next turned to single cell analysis using our EGR2-luc and DUSP1-luc reporter lines to further elucidate the mechanism of action. At the single cell level, the response to EGF stimulation is highly heterogeneous (Figure 5A; Supplementary Figure S7A; Supplementary movies 1 and 2). In the case of EGR2, there is negligible baseline reporter expression and very few cells exhibit high amplitude induction (>10-fold) and the majority fall below this threshold, with some registering barely detectable luciferase activity. In contrast, baseline expression of DUSP1 is detectable in most cells, and their magnitude of induction show a narrower range (ranging between 2.2- and 10-fold) (Supplementary Figure S7A). To further investigate the gene induction profiles we normalised the data to remove fluctuations in gene expression amplitude and used principal component analysis (PCA) to group the different expression profiles (Supplementary Figure S7B). This identified two characteristic profiles for EGR2 and three different profiles for DUSP1 (Figure 3B; Supplementary Figure S7C). For both genes, the timing of their peak expression is a key defining parameter, indicating a non-uniform response of cells to EGF induction.
Figure 5. Single cell analysis of KDAC inhibition effects reveals complex changes in gene induction parameters. (A) Examples of bioluminescence microscopy images of EGR2- (left) and DUSP1- (right) reporters in unstimulated cells (basal level) or in cells stimulated with EGF for 90 min. Bright field (BF) images show cells prior to induction. (B) Expression kinetics of EGR2- (left) and DUSP1- (right) reporters for the different profiles defined by PCA analysis (groups 1–3; see Supplementary Figure S6B and C). The data includes a combination of all single cells measured (EGR2 n = 109; DUSP1 n = 100). Lines represent average Z scores of profiles from single cells ± SEM (shaded area). (C) Kinetics of expression of EGR2- and DUSP1-reporters after either treatment with vehicle or 4SC202 for 1h prior to EGF addition. The bioluminescence of single cells was measured and the lines represent the average Z score of profiles from single cells. The shaded area represents ± SEM. (D) Ratio of cells exhibiting EGF-mediated EGR2- or DUSP1-reporter expression profiles belonging to the groups identified by PCA analysis (groups 1–3; see Supplementary Figure S6B and C) after treatment with vehicle (Control) or 4SC202. (E) The schematic shows different parameters of EGR2-reporter activity measured in single cells following EGF addition. These parameters are plotted with (n = 60) and without (n = 49) 4SC202 addition. In the graphs, each dot represents one cell. * P-value < 0.05, *** P-value < 0.001.

Next, we inhibited deacetylation and asked whether this impacted on various activation parameters. To study activation kinetics more carefully, we normalised for differences in the amplitude of activation, and found that the expression profile of EGR2 is shifted, indicating earlier initiation (Figure 5C). For DUSP1, the changes appear more complex but the initial phase of activation follows the same trajectory in the presence or absence of inhibitor (Figure 5C). To investigate these changes more closely we returned to the different single cell profiles identified by PCA analysis. For EGR2, we still see two distinct profiles but there is a significant shift in the numbers of cells exhibiting the group 1 pattern to group 2, thereby favouring an earlier response to EGF (Figure 5D). For DUSP1, we see more complex changes to the expression profiles with a loss of group 1, compensated for by a gain in group 2 and 3 profiles, although this changed profile does not reach statistical significance. We further examined various parameters associated with the EGR2 expression profile, including peak time, end point, duration and peak intensity and found statisti-
cally significant changes in all parameters at the single cell level when treated with deacetylase inhibitor (Figure 5E). The peak time and the end point are shifted earlier, consistent with the normalised data (Figure 5C). There is also a decrease in the duration of response and the overall peak intensity following deacetylase inhibition, emphasising a multifaceted response to manipulating acetylation activity.

To provide further insights into EGF-inducible gene activation kinetics, we turned to smFISH to examine the effects on allelic activation frequencies following deacetylase inhibition. First, we analysed the distribution of total number of mRNA molecules across different cells. The results were broadly in agreement with the luciferase reporter assays with EGR2 exhibiting a broader distribution of mRNA molecules per cell upon induction in comparison to the tighter distribution associated with DUSP1 (Supplementary Figure S8). Treatment with the deacetylase inhibitor 4SC202 caused a general reduction in the numbers of EGR2 transcripts per cell whereas the opposite was observed for DUSP1 (Supplementary Figure S8A), consistent with what we observed using reporter alleles or bulk mRNA from cell populations. To study allelic activation frequencies, we focussed on DUSP1. In the absence of EGF stimulation very few cells contained active transcriptional loci but after 15 mins stimulation, over 60% of cells showed activation of two or three alleles (MCF10A cells contain an extra copy of the chromosomal segment harbouring the DUSP1 locus) (Figure 6A and B). Following that, there is a general decline to one or no active alleles by 60 min. However, in the presence of the deacetylase inhibitor 4SC202, a large number of cells already show activation of one or more alleles under basal conditions and there is a clear increase in the proportion of cells containing two or three active alleles after 15 min EGF treatment which is sustained at 30 min. In contrast when cells were treated with an acetylation inhibitor, only a small proportion of cells show activation of two or three alleles at any timepoint after EGF stimulation. Thus, acetylation-mediated control over the number of active alleles likely plays a major role in determining the outcome of transcription at the DUSP1 locus. We were unable to design probes to detect active transcriptional loci for EGR2 and instead focussed on FOS which responded in a similar manner and was activated following KAT inhibition (Figure 3D; Supplementary Figure S4A). In contrast to DUSP1 where fewer loci were active following KAT inhibition with GNE-781, more FOS loci per cell were active at all time points following EGF stimulation (Supplementary Figure S8B). Treatment with 4SC202 was less clear cut with minimal changes, reflecting the more limited changes seen to FOS expression at the mRNA level and its placement in a distinct category of KDAC inhibitor responsive genes from EGR2 (Figure 2D).

Collectively, these single cell approaches reveal a complex series of gene activation parameters that are altered by changing acetylation dynamics that underly the expression profiles derived from cell populations.

DISCUSSION

High levels of histone acetylation are typically associated with transcriptionally active genes both at promoter and enhancer elements and this has led to the assumption that increased acetylation leads to high level gene transcription as exemplified by the use of KAT inhibitors on a genome-wide scale (25,30). Nevertheless, reciprocal effects are observed at some loci such as genes encoding core regulatory transcription factors where KDAC inhibitors suppress transcription (31). This indicates that high level acetylation does not always lead to enhanced transcriptional activity. Here, we examined the effect of disturbing the acetylation equilibrium catalyzed by class I KDACs and the KATs P300/CBP on growth factor inducible gene activation. We found that EGF activated genes fall into two broad categories (Figure 6C), those which show the canonical activation response to increased acetylation levels (group I e.g. DUSP1) and those which show the opposite effect and whose expression is dampened when acetylation levels are increased (group I e.g. EGR2). The latter group are characterized by low basal level transcription and low levels of histone marks associated with high transcriptional activity. Thus, rather than a unified response, different genes are tuned to respond differently to changes in acetylation levels, even when responding to the same signaling events.

Using single cell approaches, we gained further insights into the gene activation dynamics at DUSP1 and EGR2 and the mechanistic changes induced by disturbing the acetylation equilibrium. Rather than a uniform response to EGF addition, there is a large variation in the magnitude of peak EGR2 transcription, but there are two basic expression profiles which differ according to their timing onset. In contrast, the DUSP1 expression profile is much more complex, with a more uniform induction level across single cells but three different underlying kinetic profiles. KDAC inhibition led to a switch in the timing of EGR2 expression to the earlier of the two kinetic profiles and a concomitant drop in overall peak expression levels and the duration of activation. The DUSP1 activation profiles were also altered by KDAC inhibition leading to a shift towards later expression onset and peak expression. However, due to the more complex expression profiles for DUSP1, we were unable to further disentangle the changes elicited by KDAC inhibition. Instead we examined allelic activation frequencies and found that KDAC inhibition increased the numbers of active alleles per cell whereas KAT inhibition had the opposite effect. Perturbing the acetylation dynamics therefore affects the allele activation frequency. This is reminiscent of recent studies where KAT inhibition reduced Fos transcriptional burst frequencies in neurons (32) and, reciprocally, high KAT activity and high histone acetylation levels augmented burst frequencies of Bmal1 and hence transcriptional output (33).

The effects of disturbing the acetylation equilibrium on transcriptional levels can be explained largely by the changes to H3K27ac at the DUSP1 promoter, and a recent study demonstrated that a subset of genes respond to acetylation changes at their promoters (34). However, the latter study made the surprising finding that H3K27ac acetylation levels at most promoters are refractory to P300/CBP inhibition and here we show that EGR2 falls into this category. Similarly, an earlier study indicated that acetylation at the majority of promoters is refractory to KDAC inhibition (10). The acetylation event(s) that are altered at the
Figure 6. DUSP1 allele activation frequency in response to disturbing the acetylation-deacetylation equilibrium. (A and B) smFISH of DUSP1 mRNA production in serum starved (basal) and EGF stimulated MCF10A cells in the presence of prior addition of vehicle, 4SC202 or GNE-781. Representative images from cells following 15 mins induction are shown in (A). (B) Quantification of the distribution of cells with 0, 1, 2 or 3 active transcription sites at each time point and condition. Numbers at the top of each column indicates number of cells (n). (C) Model showing the effects of altering the acetylation (Ac) equilibrium on the activation of the EGF inducible genes DUSP1 and EGR2, examples of larger groups of genes that respond in a similar manner. The transcriptional start site (TSS) and putative enhancer regions (En) are indicated.

EGR2 locus therefore remain enigmatic. In many ways, our findings that the effects of KDAC inhibitors on EGR2 activation are independent of obvious changes to promoter histone H3K27 acetylation are reminiscent of findings on MMTV transcription where the effects of KDAC inhibitors are attributed to changes to the basal transcription machinery rather than to histones (7). This suggests a broader effect of KDACs in gene activation beyond their role in histone modifications. However, it should be noted that there are numerous different histone modifications in addition to H3K27ac that we tested here, and it is possible that KDACs function on one or more of these in the context of enhancing EGF-mediated genes activation. Others have questioned the relevance of H3K27ac inactivation of regulatory elements and their associated genes and found that this mark is generally not needed for enhancer activity in mouse ESCs (35). It is possible that different acetylation marks are more relevant, including others such as H3K18ac, H3K56ac, H3K64ac, and H3K122ac that are deposited by P300/CBP (36–38). Moreover, H4K5/8/12ac have been shown to be important at promoters for signal-dependent transcriptional activation (39) and their levels may be influenced by KDAC inhibition. It also remains possible that histone acetylation at enhancer regions is modulated in response to KDAC/KAT inhibition, at least at a subset of genes and/or we have sampled the wrong time point in our analyses. Alternatively, there may be other non-histone proteins involved whose activity is influenced by acetylation levels (30). Further work is needed to probe these possibilities. It is also important to note that although we can detect increased nascent transcription and allelic expression of group II genes like DUSP1 upon increasing lysine acetylation levels, we were unable to assess the reciprocal effects on group I genes due to technical limitations. Instead, we were able to analyze FOS expression (which only partially resembles group I genes) and demonstrated that inhibition of KATs triggered increased mRNA production, which was reflected in an increase in the number of active loci per cell. Thus, KAT inhibition can have opposite effects on the number of active loci per cell depending on the EGF target gene. However, equally this leaves open the possibility that Group I genes like EGR2 may be affected at the post-transcriptional level through decreased stability following KDAC inhibition and/or that there may be a combination of transcriptional and post-transcriptional events at play.

Overall, we identified a group of genes (group II: exemplified by DUSP1) whose activity is affected as expected by disturbing acetylation levels, with high acetylation causing increased levels of gene transcription but another set of genes (group I: exemplified by EGR2) behaves in the opposite manner. Interestingly, at this second group of genes, acetylation inhibitors show an effect earlier in the induction profile than deacetylation inhibitors (see Supplementary Figure S5). This suggests a model whereby initial high level basal acetylation caused by KDAC inhibition is refractory for the onset of gene activation. Conversely, the
inducible acetylation changes that occur following growth factor stimulation are more affected by the acetylation inhibitors where the dampening down of acetylation levels leads to a transcriptional overshoot. This implies that high acetylation levels normally limit the overall activation levels of expression at this set of genes.

While the growth factor inducible genes fall into these two categories based on KDAC inhibition profiles, they respond subtly differently to KDAC inhibition, with group B genes (activated by KAT inhibition) being split between two subgroups, one of which shows general dampening across the profiles and another where only the peak expression is reduced. Thus, although there are commonalities, there are further subtle differences in how individual genes respond, likely caused by their unique regulatory setups and different induction profiles. Indeed, previous work on inducible acetylation changes that occur following growth factor stimulation are more affected by the acetylation inhibitors where the dampening down of acetylation levels leads to a transcriptional overshoot. This implies that high acetylation levels normally limit the overall activation levels of expression at this set of genes.

While the growth factor inducible genes fall into these two categories based on KAT inhibition profiles, they respond subtly differently to KDAC inhibition, with group B genes (activated by KAT inhibition) being split between two subgroups, one of which shows general dampening across the profiles and another where only the peak expression is reduced. Thus, although there are commonalities, there are further subtle differences in how individual genes respond, likely caused by their unique regulatory setups and different induction profiles. Indeed, previous work on FOS induction, demonstrated the importance of the acetylation-deacetylation equilibrium for gene activation, and in that case inhibition of both KDACs and KATs reduced transcriptional levels (4,5). In combination with these earlier case inhibition of both KDACs and KATs reduced transcription, demonstrated the importance of the acetylation-deacetylation equilibrium for gene activation, and in that case inhibition of both KDACs and KATs reduced transcriptional levels (4,5). In combination with these earlier studies, our data therefore indicate that it is important to re-evaluate the assumptions that high steady state acetylation levels mean that acetylation promotes transcriptional activity. Instead, this should be evaluated on a case by case basis, where each gene is tuned to respond differently to dynamically changing acetylation levels catalyzed by the opposing functions of KDACs and KATs.

**DATA AVAILABILITY**

RNAseq data newly generated in this study are deposited in ArrayExpress; accession number E-MTAB-9881.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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