Abstract

Background: The ERK MAPK pathway plays a pivotal role in regulating numerous cellular processes during normal development and in the adult but is often deregulated in disease scenarios. One of its key nuclear targets is the transcription factor ELK1, which has been shown to play an important role in controlling gene expression in human embryonic stem cells (hESCs). ELK1 is known to act as a transcriptional activator in response to ERK pathway activation but repressive roles have also been uncovered, including a putative interaction with the PRC2 complex.

Methods: Here we probe the activity of ELK1 in hESCs by using a combination of gene expression analysis in hESCs and during differentiation following ELK1 depletion and also analysis of chromatin occupancy of transcriptional regulators and histone mark deposition that accompany changes in gene expression.

Results: We find that ELK1 can exert its canonical activating activity downstream from the ERK pathway but also possesses additional repressive activities. Despite its co-binding to PRC2 occupied regions, we could not detect any ELK1-mediated repression at these regions. Instead, we find that ELK1 has a repressive role at a subset of co-occupied SRF binding regions.

Conclusions: ELK1 should therefore be viewed as a dichotomous transcriptional regulator that can act through SRF to generate both activating and repressing properties at different genomic loci.

Keywords
ELK1, repression, embryonic stem cells
Corresponding author: Andrew D. Sharrocks (andrew.d.sharrocks@manchester.ac.uk)

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Introduction

In vitro studies on human embryonic stem cells are an important step in understanding the molecular basis to human development. Cultured human embryonic stem cells (hESCs) require FGF2-mediated signalling through the ERK pathway to maintain their pluripotent state (Lanner & Rossant, 2010). More recent studies indicate that an earlier ERK pathway-independent state can be achieved whereby ERK pathway suppression is a key event in driving this transition (Theunissen et al., 2014). This earlier state is equivalent to the mouse ESC naïve ground state that is thought to represent the pre-implantation epiblast.

Nevertheless, understanding the role of the ERK pathway in hESCs remains an important goal. Some of the best characterised targets of the ERK MAPK signalling pathway are the E-twenty six (ETS) proteins, which are nuclear transcription factors and as such can directly convert ERK pathway signalling to alternative interpretation and we have now done this. In response to reviewer 1, we have now mentioned that due to the lack of binding motif, ELK1 is likely indirectly recruited to chromatin characterised by repressive features in hESCs. In addition we now show that ELK1 expression is unchanged following RA treatment (Figure 2B). Further discussion has been added concerning the potential role of ELK1 in repression of mesoderm-specific genes and we have linked the results to our data showing an interplay between direct ELK1 and SRF binding to the FOXC1 locus.

In response to reviewer 2, due to the inconclusive nature concerning ELK1 opposing MRTF signalling, we have removed mention of this from the abstract and qualified the discussion to indicate that we cannot completely rule out a role for ELK1 in opposing MRTF recruitment in mesoderm cells. We chose to retain the data we have so far though on this subject to inform future studies in this area.

Further explanation and discussion of the different culture conditions has been added to the text and legend describing Figure 6. We have also clarified that the target genes are bound by ELK1 and hence likely direct targets but leave open the possibility that indirect effects might occur. We have also indicated that it remains possible that ELK1 and PRC2 might function together in repression at a subset of genes we have not tested or under different conditions. An additional reference that is relevant to the study (Lee et al., 2010) is now cited in the revised paper.

Here we extended the analysis of ELK1 function in hESCs, first exploring the relationship between ELK1 and the PRC2 complex, and then its activity through its known binding partner SRF. Despite ELK1 and PRC2 co-occupying a large number of genomic regions, we were unable to uncover evidence to support a repressive role of ELK1 in this context. However, unexpectedly, we were able to uncover a repressive role for ELK1 in the context of a subset of SRF-bound regulatory regions. This repressive role appeared distinct from a simple competition model for SRF binding by the co-activator MRTFA. ELK1 therefore possesses both activating and repressive functions in hESCs, directed through its regulatory partner protein SRF.

Results

Functional interplay between ELK1 and PRC2

Previous studies demonstrated that ELK1 occupies two distinct sets of genomic loci in H1-hESCs (Göke et al., 2013). One set was associated with co-binding with SRF, a configuration which is usually associated with transcriptional activation. However, the second set of loci exhibited co-localisation with members of the repressive PRC2 complex, hence suggesting a role in transcriptional repression. As these conclusions were based on the analysis of promoter-proximal ELK1 binding sites, we re-analysed the chromatin immunoprecipitation sequencing (ChIP-seq) data to establish whether these patterns could be observed in a genome-wide manner. Initially we focussed on the co-association with the PRC2 complex, and segregated ELK1 regions according to whether co-binding of the PRC2 complex subunit SUZ12 could be identified (ELK1+SUZ12) or not (ELK1-SUZ12). A significant overlap in ELK1 and SUZ12 binding regions was observed (108 regions; hypergeometric
p-value = 7.4×10^(-12), although the majority of ELK1 binding peaks showed no overlap (Figure 1A). Next, we asked whether the two classes of ELK1 binding regions showed differences in co-association with the ELK1 binding partner SRF and a variety of histone marks that are characteristic of transcriptional repression or activation. The ELK1+SUZ12 binding loci were enriched for SUZ12 and EZH2 binding and for the H3K27me3 histone modification, suggesting repressed regions of chromatin (Figure 1B). These regions showed no enrichment of the ELK1 DNA binding motif as observed previously for a smaller window (Göke et al., 2013), suggesting an indirect mechanism for recruitment of ELK1 to chromatin at these loci. In contrast, the ELK1-SUZ12 regions showed little co-association with the repressive features and instead high enrichment of SRF binding was observed and co-association with histone marks suggestive of active transcription, H3K9ac, K3K27ac and H3K4me3 (Figure 1C). These “active” sites also showed strong occupancy by its known binding partner, SRF, consistent with a large overlap between ELK1 binding regions when analysed at the individual binding peak level (Figure 1A). These data point to the existence of an active ELK1+SRF module that lacks PRC2 binding, and a distinct repressive ELK1+SUZ12 module (Figure 1D, E) and are broadly consistent with the models previously proposed using a subset of this data (Göke et al., 2013).

Next, we focussed on the potential repressive role of ELK1 in the context of the ELK1+SUZ12 co-bound regions and sought to establish whether the genes associated with these peaks had potential biological relevance. To address this, we first assigned biological function gene ontology terms of genes associated with ELK1 binding loci either overlapping with SUZ12 binding loci (D) or not overlapping with SUZ12 binding loci (E).

Figure 1. ELK1 has binding modules either enriched for PRC2 or enriched for active histone marks. (A) Venn diagram showing the intersection of binding regions from genome-wide chromatin immunoprecipitation-sequencing datasets for ELK1, SUZ12 and SRF in H1-hESCs. (B and C) Transcription factor and histone modification tag density profiles in H1-hESCs in a region 2,500 bp to either side of the centre of the ELK1 binding regions. ELK1 binding regions are portioned according to (A) overlapping with SUZ12 binding regions (+SUZ12) or (B) not overlapping with SUZ12 binding regions (-SUZ12). (D and E) diagrammatic illustration of the ELK1 binding regions associated with co-binding of the repressive PRC2 complex (D) or associated with its binding partner SRF and active regions of chromatin (E). (F and G) Biological function gene ontology terms of genes associated with ELK1 binding loci either overlapping with SUZ12 binding loci (D) or not overlapping with SUZ12 binding loci (E).
genes to binding loci with HOMER (Heinz et al., 2010), using the nearest TSS model. We next analysed the biological process gene ontologies (GO) of genes linked to partitioned ELK1 binding loci. ELK1 binding loci overlapping with SUZ12 binding loci (ELK1+SUZ12) were enriched for terms relating to development (Figure 1F). Terms are provided on figshare (Prise, 2019a). This result is consistent with the known function of PRCs in functionally repressing key developmental genes in hESCs (Lee et al., 2006; Yu et al., 2017). However, an ELK1-mediated mechanism of PRC repression would be novel. These loci are also enriched for the H3K27me3 modification (Figure 1B), a functional indication of PRC2-mediated repression, suggesting that an ELK1+SUZ12 module might be involved developmental gene repression. In contrast, we found that ELK1-binding loci not overlapping with SUZ12 binding loci were not enriched for developmental processes and were instead correlated with gene expression and metabolic processes (Figure 1G). These latter observations are consistent with the role for ELK1 previously identified in HeLa cells (Boros et al., 2009).

Having shown that ELK1+SUZ12 peaks were enriched in development genes, we then identified a set of associated target genes for further study that are induced upon differentiation. We hypothesised that the regulatory regions of these genes would switch from repressed (and PRC2-bound) to active during differentiation. We chose to use retinoic acid (RA), a potent initiator of hESC differentiation. Using the nearest TSS-association model in HOMER, we identified a set of ELK1+SUZ12-bound genes whose expression increased upon 48–96 hours of treatment with RA and focussed on six of these that showed robust induction (Figure 2A). ELK1 expression is unaffected under these conditions (Figure 2B). To understand the role of ELK1 in the context of the repressive modules associated with these genes, we depleted ELK1 using shRNA (Figure 2C, D) and examined the effect on PRC2 occupancy and gene expression in H1-hESCs. ELK1 depletion led to the expected decrease in ELK1 binding in these regions (Figure 2E). However, this was not accompanied with a decrease in SUZ12 binding (Figure 2F), nor did an ELK1 knockdown result in a substantive increase in nearby gene expression (Figure 2G). Thus, although ELK1 co-occupies a set of genomic regions with the PRC2 complex, these results indicate that ELK1 does not appear to have a role in maintaining PRC2 occupancy or in maintaining transcriptional repression through these regions in H1-hESCs. Raw RT-qPCR (Prise, 2019b) and ChIP-qPCR (Prise, 2019c) data are available on figshare.

Functional interplay between ELK1 and SRF

Having shown that ELK1 plays no clear role in PRC2-mediated gene repression, we next returned to the ELK1 module, which preferentially shows enrichment for co-binding of SRF. Of the 710 ELK1-bound regions not found in the ELK1-SUZ12 dataset, 282 (38%) also show SRF co-occupancy (Figure 1A; hypergeometric p-value = 3.2 × 10^-209). Notably, there are very few regions co-bound by ELK1, SRF and SUZ12, indicating a clear distinction between regions bound by ELK1 and either SRF or SUZ12. We decided to focus on the role of ELK1 in differentiation to mesoderm as SRF has previously been implicated in this developmental process (Arsenian et al., 1998). An additional advantage of the mesoderm differentiation protocol is that it produces a decrease in pluripotency factors and high expression of the marker gene T after a short 3-day treatment time (Figure 3A–C), and is therefore compatible with a siRNA depletion approach. ELK1 knockdown revealed no general change in expression of pluripotency factors in H1-hESCs (Figure 3D, E). ELK1 protein expression increased in mesoderm cells (Figure 3C) but there was little change in the expression of two mesoderm-marker genes MSX2 and PITX2 following ELK1 depletion, indicating the absence of a general role in mesoderm differentiation (Figure 3F). In contrast two other marker genes (FOXC1 and HAND2) exhibited increased expression following differentiation to mesoderm, suggesting a potential repressive function in this context (Figure 3F).

To determine possible functional interactions between ELK1 and its known cofactor, SRF, we first identified a set of genes which are located close to potential regulatory regions that are co-bound by ELK1 and SRF. Next, we treated H1-hESCs with either siSRF or siELK1 and then either maintained the hESCs in their pluripotent state or differentiated them to mesoderm cells. First we analysed SRF and ELK1 binding to chromatin and performed ChIP-quantitative PCR (qPCR) on regions which were bound by both ELK1 and SRF. Binding of both factors was specifically detectable on the known target genes EGR1 and EGR2 (Figure 4). Generally, we saw a decrease in SRF binding and a concomitant decrease in ELK1 binding in the same regions following SRF depletion (Figure 4A, C and Figure 5A, C). This pattern was detected in 19/19 of the regions tested H1-hESC (Figure 5A) and 18/22 of the regions tested in mesoderm (Figure 5C). This is consistent with the existing models, whereby SRF acts as a platform to aid ELK1 recruitment to chromatin (Gille et al., 1995; Janknecht & Nordheim, 1992; Latinkič et al., 1996; Treisman et al., 1992). Conversely, when we depleted ELK1 and performed ChIP-qPCR on regions which were bound only by ELK1 (Figure 5B, D) and 16 of the 22 regions tested in mesoderm (Figure 5D). These results confirm a widespread role for SRF in stabilising ELK1 occupancy on chromatin but suggest an unexpected role for ELK1 in apparently reducing SRF occupancy on chromatin. Raw Fluidigm data are available on figshare (Prise, 2019d).

To investigate whether these changes in transcription factor binding correlated with gene expression, we again performed a knockdown of ELK1 in H1-hESCs or during mesoderm differentiation and tested the expression of a number of target genes. First, we assessed whether we could detect the known activating role of ELK1 at the immediate-early genes EGR1, EGR2 and...
Figure 2. ELK1 is not involved in PRC2-mediated repression. (A and B) Reverse transcription-quantitative PCR (RT-qPCR) analysis of ELK1+SUZ12-bound genes (A) and ELK1 (B) upon 48 and 96 hours of retinoic acid (RA) treatment. Data are shown relative to DMSO-treated cells (taken as 1) and are the average of at least three independent experiments. Asterisks represent p-value <0.05. (C) RT-qPCR of GAPDH and ELK1 expression after 96 hours of shELK1 knockdown in H1-hESCs. Two different shRNA vectors were tested (plasmid containing shRNA_2 was subsequently used in all experiments). Data are shown relative to control empty plasmid treated cells (taken as 1) and are the average of three independent experiments. Asterisks represent p-value <0.05. (D) Western blot analysis of ELK1 and HDAC1 expression in H1-hESC after 96 hours of treatment with an empty plasmid or a plasmid containing shELK1. (E and F) Chromatin immunoprecipitation-qPCR of ELK1-binding regions for ELK1 (E) or SUZ12 (F) occupancy after 96 hours of treatment with either control empty vector or shELK1. (G) RT-qPCR of ELK1+SUZ12-bound genes after 96 hours of treatment with either control empty vector or shELK1. Data are normalized to GAPDH expression and “control plasmid” (taken as “1”) and are the average of 5 independent experiments, with the exception of the data for ONECUT2 and POU3F2 which are the average of 4 independent experiments.
Figure 3. The role of ELK1 in gene expression during mesoderm differentiation. (A and B) Reverse transcription-quantitative PCR (RT-qPCR) measuring the expression of pluripotency markers (A) or the mesoderm marker (T/Brachyury) after 72 h of growth in MIM. (C) Western blots for ELK1, the indicated pluripotency markers, T/Brachyury, and the control TUBB protein in H1-hESC cells and H1-hESCs grown in MIM for 72 h (mesoderm cells). (D) Western blots for ELK1, OCT4, SOX2, and TUB expression in H1-hESC cells treated with non-targeting (NT) siRNA or siRNAs targeting ELK1. (E and F) RT-qPCR measuring the expression of ELK1 and the indicated pluripotency markers (E) or the indicated ELK1-bound genes in H1-hESCs and H1-hESCs after 98 h of growth in MIM (mesoderm cells) (F) after treatment with non-targeting (NT) siRNA or siRNAs targeting ELK1. RT-qPCR data are normalised to H1-hESCs in the presence of siNT and are the average of 3 independent experiments. *p-value < 0.05.
Figure 4. Reciprocal effects of ELK1 and SRF depletion on each other’s chromatin association. (A–D) Chromatin immunoprecipitation-quantitative PCR showing ELK1 and SRF binding (indicated above each graph) after treatment of H1-hESCs (A and B) or mesoderm cells (C and D) with non-targeting (NT) siRNA or siRNAs against SRF (A and C) or siELK1 (B and D). Binding to the promoter regions of EGR1 and EGR2 are shown and a negative control region located 2 kb upstream from the EGR1 locus. Data are the average of 3 independent experiments. *p-value < 0.05.
EGR3 in these cells. As these genes are inducibly activated by ERK pathway signalling, we activated the ERK pathway by culturing mesoderm cells (derived from H1-hESCs) in DMEM/F12 media followed by serum starvation for 24 hrs and then switched the media to MIM for 15 mins. This treatment caused increased levels of active, phosphorylated ERK (Figure 6A) and the activation of EGR1, EGR2 and EGR3 expression (Figure 6B). ELK1 was efficiently depleted by siRNA treatment (Figure 6A) and this depletion caused a significant decrease in the expression of EGR1, EGR2 and EGR3 under stimulating conditions (Figure 6B). Having established the known activating role of ELK1 in our system, we switched to investigating whether ELK1 plays an activating role at different genes which are not expected to be activated by the ERK pathway as exemplified by SPARCL1 (Figure 6C). In both H1-hESC and mesoderm cells grown under steady state levels, ELK1 depletion caused an increase rather than a decrease in the expression of a panel of its target genes bound by the ELK1-SRF complex (Figure 6D, E). This indicates that ELK1 is necessary for the repression rather than the activation of these target genes. ELK1 is also responsible for EGR2 repression in basal and steady state levels (Figure 6B) but switches to an activating role following acute stimulation (Figure 6E) which is consistent with previous data showing that it can both activate and repress transcription (Marais et al., 1993; Yang et al., 2001; Lee et al., 2010).

Figure 5. Inter-relationship between ELK1 and SRF binding to chromatin across co-bound loci. (A–D) Heatmaps of chromatin immunoprecipitation-quantitative PCR showing fold change in ELK1 and SRF binding after treatment of either H1-hESCs with siSRF (A) and siELK1 (B) or mesoderm cells after treatment of either H1-hESCs with siSRF (C) and siELK1 (D). Data are the average of three independent experiments. *p-value < 0.05.

ELK1 status does not affect the response to changes in actin dynamics

A plausible model is that SRF is an activator at these loci, and ELK1 would then act in a repressive manner to keep SRF activity in check. This model was previously proposed for the role of ELK1 in opposing the recruitment of the MRTF family co-activator myocardin in smooth muscle cells (Wang et al., 2004). Indeed, ELK1 and MRTFs occupy the same binding surface on SRF, meaning that binding is mutually antagonistic (Zaromytidou et al., 2006). If ELK1 was opposing the actions of MRTF family members, depletion of ELK1 would be predicted to hypersensitise target gene expression to activators of this pathway. To address this possibility, we tested the effect of ELK1 depletion in the context of cytochalasin D stimulation, which acts through inhibiting actin polymerisation and has been shown to activate SRF-target genes through the MRTFs (Esnault et al., 2014; Posern & Treisman, 2006).

We stimulated serum-starved mesoderm cells with cytochalasin D, concurrently with ELK1 knockdown, and tested the expression of four target genes for the ELK1-SRF complex, EGR2, CDK91A, FOSL1 and SPARCL1. Neither FOSL1 nor SPARCL1 were responsive to cytochalasin D treatment under serum-starved conditions, suggesting a lack of involvement of MRTFs (Figure 7). As observed previously, we saw an increase in the expression of both genes upon ELK1 knockdown in mesoderm...
cells under both normal culture conditions and also when treated with cytochalasin D (Figure 7). However, these genes did not become responsive to MRTF pathway activation when the putative MRTF-binding inhibition via ELK1 was removed by ELK1 depletion (Figure 7). In contrast, EGR2, a gene usually activated by ELK1, is activated by cytochalasin D and this effect is potentiated by depletion of ELK1 (Figure 7). Thus, ELK1 appears to interact differently with the MRTF pathway at...
different loci, but in regions where it acts as a repressor such as the SPARCL locus, it does so without influencing the response to this pathway. This suggests that binding by another co-activator or the intrinsic activity of SRF alone may be responsible for the increased gene activation we observe in the absence of ELK1.

**Discussion**

Previous work studying the role of ERK signalling in H1-hESC led to a focus on ELK1 as a potential regulator of pluripotency (Göke et al., 2013). In this context, ELK1 was proposed to act in combination with the PRC2 complex to repress the expression of genes involved in hESC differentiation. In line with this previous analysis on promoter proximal events, we demonstrated that ELK1 binding occupies two distinct modules throughout the genome, one enriched for the repressive PRC2 complex and one enriched for the presence of active histone marks and binding of the known ELK1 partner protein SRF. However, we were unable to demonstrate a repressive role for ELK1 through these elements indicating that ELK1 does not act in the context of PRC2-mediated repression in hESCs. Although we have surveyed a panel of genes, it remains possible that other genes may be controlled through this complex and/or the repressive function is only revealed under particular conditions. Based on our analysis though, the widespread binding of ELK1 and the PRC2 complex appears to be coincidental rather than functionally linked (Figure 8A).

This led us to question whether ELK1 acts merely as a transcriptional activator rather than a repressor in hESCs, consistent with its known role as a mediator of ERK pathway-mediated gene activation (Gille et al., 1995; Janknecht et al., 1993; Marais et al., 1993). We confirmed that ELK1 acts as an activator of canonical target genes for the ELK1-SRF complex such as EGR1 (Figure 8C). However, we found that ELK1 loss broadly resulted in increased SRF ChIP signal at a panel of target genes for this complex, suggesting that it might act as a repressor in this context, perhaps by destabilising SRF binding to DNA. Indeed, we showed that ELK1 acts in a repressive manner at a subset of target genes for this complex, as exemplified by FOSL1 (Figure 8B). However, both ELK1 and SRF binding is detectable at this and other loci by ChIP-seq analysis, making it unlikely that ELK1 destabilises SRF binding. Moreover, previous studies have shown that ELK1 can stabilise SRF binding to DNA rather than inhibiting its binding (Ling et al., 1998). Others have previously shown that depletion of ELK1 along with other TCF subfamily proteins, results in increased ChIP signal for SRF at co-bound genes in murine cells, although they provided no molecular explanation for this phenomenon (Gualdrini et al., 2016). Instead, an alternative technical explanation might be increased epitope exposure on SRF after ELK1 knockdown, either through loss of steric hindrance or due to conformational changes in the DNA-bound SRF.

In theory, loss of ELK1 binding to SRF might reveal a binding surface for another co-activator protein as well as potentially allowing access to antibodies used in the ChIP procedure. Such a scenario would be consistent with the previously defined role for ELK1 in opposing binding by the MRTF family of co-activator proteins (Wang et al., 2004; Zaromytidou et al., 2006). However, loss of ELK1 did not generally make its target genes more responsive to MRTF pathway activation. It remains possible that basal MRTF signalling may be operative in these cells and that ELK1 presence opposes this and future additional studies are needed to fully rule out a role for MRTFs. Whilst our current data do not support a role for MRTFs as an important co-activator in this context, it may be that an as-yet-unknown factor may play a role in SRF-mediated gene activation. Future studies are needed to address this possibility. Alternatively, ELK1 may itself impart repressive properties on an SRF-bound regulatory region, such as through recruitment of the SIN3A complex (Yang et al., 2001) or through SUMO-mediated recruitment of histone deacetylases (Yang & Sharrocks, 2004) as previously shown in other cell types. However, it is unclear why ELK1 should be repressive at only a subset of its binding regions and activating at others. Finally, it remains possible that ELK1 may repress its

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**Figure 8. Model for the role of ELK1 at different genomic loci.** (A) ELK1 binds to the same genomic regions as SUZ12 and EZH2 of the PRC2 complex. In this scenario, repression is mediated at genes like PAX7 by the PRC2 complex but not ELK1. (B) ELK1 binds to regions alongside its partner protein SRF. In this scenario, repression is mediated at genes like FOSL1 by ELK1, potentially by competing for another co-activator protein. (C) ELK1 binds to regions alongside its partner protein SRF and is activated by the ERK pathway. Here ELK1 acts as a canonical transcriptional activator protein at genes like EGR1.
targets through an indirect mechanism, although it appears more likely that it would act directly through its binding to the regulatory regions of its target genes.

In summary, our work has identified a role for ELK1 in acting in its traditional role as a transcriptional activator downstream from the ERK pathway in hESCs. In addition, it also plays a repressive role in hESCs through SRF-bound regulatory regions. However, we were unable to find evidence to support a role in transcriptional repression in conjunction with the PRC2 complex as previously proposed (Göke et al., 2013). ELK1 therefore acts as dichotomous transcriptional regulator in hESCs, through imparting both activating and repressive activities to SRF-bound target genes.

**Methods**

**Cell Culture**

H1-hESC cells (Wicell) were routinely cultured in mTeSR1™ (StemCell Technologies). Plates were coated with Matrigel (Corning) at 37°C for 1 hour before passage. To passage the cells, the cells were coated with a thin layer of ReLeSR™ (StemCell Technologies) and incubated at 37°C for 5 minutes.

For shELK1 and RA treatment, conditioned H1 media was used, containing DMEM/F12 (Invitrogen), 20% (v/v) knockout serum replacement (Thermo Fisher Scientific), 1 mM L-glutamine (Gibco), 1% (v/v) nonessential amino acids (Gibco), 0.1 mM 2-mercaptoethanol (Gibco), and 4 ng/ml basic fibroblast growth factor (Invitrogen). This media was conditioned with CF1 mouse fibroblasts (MTI-GlobalStem) for 24 hr prior to adding to the H1-hESCs. Media was then vacuum filtered (0.22 µM), and an additional 8 ng/ml of basic fibroblast growth factor (Invitrogen) was supplemented to conditioned medium before usage. To passage the cells, the cells were coated with a thin coat of dispase (StemCell Technologies) and incubated at 37°C for 5 minutes.

For differentiating hESCs to mesoderm, cells were first treated with 10 µM Y-27632 (ROCK inhibitor) for 1 hour before passage and dissociated with TrypLE™ Express (StemCell Technologies). Cells were initially seeded at a density of 5×10^4 per cm^2 in mTeSR1™ supplemented with 10 µM Y-27632. After 24 h, cells were grown in STEMdiff™ mesoderm Induction medium (StemCell Technologies) for an additional 3 days.

To activate the ERK signalling pathway, H1-hESCs were first allowed to differentiate into mesoderm cells, maintained in DMEM/F12 media lacking serum for 24 h and then switched to DMEM/F12 containing 2 µM cytochalasin D for 60 minutes to study gene expression. DMEM/F12 media containing DMSO at a final concentration of 1:10,000 was used as a control for cytochalasin D stimulation.

For cytochalasin D stimulation, H1-hESCs were first allowed to differentiate into mesoderm cells, maintained in DMEM/F12 media lacking serum for 24 h and then switched to DMEM/F12 containing 2 µM cytochalasin D for 60 minutes to study gene expression. DMEM/F12 media containing DMSO at a final concentration of 1:10,000 was used as a control for cytochalasin D stimulation.

**shRNA and siRNA treatment regimes**

For shRNA treatment, cells were first treated with ROCK inhibitor and dissociated with TrypLE™ Express. Next, 5×10^5 cells were treated with 7.5 µl of TransIT®-LT1 (Mirus) plus 2.5 µg of shRNA plasmid prepared in 250 µl of OptiMEM. shELK1 plasmid was a pSuper-derived plasmid containing the shRNA hairpin for ELK1: 5’-GCCAGAAGTTCGTCTACAA-3’ (Göke et al., 2013). An empty pSuper plasmid was used as a control for shELK1 treatments.

For siRNA transfection cells were prepared as above and after dissociation seeded at a concentration of 5×10^5 per cm^2 in mTeSR1™ supplemented with 10 µM Y-27632. Each 5×10^5 cell sample was treated with 7.5 µl Lipofectamine RNAiMAX reagent (Thermo Fisher Scientific) and 2.5 µl of siRNA (20 µM stock concentration), prepared in 150 µl Opti-MEM (Gibco). A non-targeting siRNA (siNT) was used as a control for siELK1 and siSRF treatments.

**ChIP assays**

Cells were incubated at room temperature with 1% (v/v) formaldehyde (Sigma), for 10 minutes (3×10^6 cells were seeded per immunoprecipitation (IP)). The crosslinking reaction was then quenched with 0.125 M glycine for 5 minutes. Cells were washed with ice cold 1x PBS. Next, 3×10^6 cells were harvested in FA cell lysis buffer (10 mM Tris-HCl, pH 8.0, 0.25% (v/v) Triton-X100, 10 mM EDTA, 0.1M NaCl) rotated for 10 minutes at 4°C, the nuclei pelleted at 13.1 krpm at 4°C for 5 minutes and the supernatant discarded. Cells were resuspended in FA Cell Lysis Buffer, rotated for 10 minutes at 4°C, pelleted at 13.1 krpm at 4°C for 5 minutes and the supernatant discarded. Nuclei were then resuspended in 1% SDS solution (50 mM HEPES-KOH, pH 7.5, 150 mM NaCl, 2 mM EDTA, 1% (v/v) Triton-X100, 0.1% (v/v) Na-DOC, 1% (w/v) SDS), rotated for 10 minutes at 4°C and the chromatin pelleted at 13.1 krpm at 4°C for 5 minutes and the supernatant discarded. Chromatin was then suspended in 0.1% SDS solution (50 mM HEPES-KOH, pH 7.5, 150 mM NaCl, 2 mM EDTA, 1% (v/v) Triton-X100, 0.1% (v/v) Na-DOC, 0.1% (w/v) SDS), rotated for 10 minutes at 4°C, pelleted at 13.1 krpm at 4°C for 5 minutes and the supernatant discarded. Chromatin was then resuspended in 0.1% SDS solution and sonicated to produce chromatin fragments of 100–500 bp.

For 3×10^6 cells, 12.5 µl of Dynabeads® Protein G (Thermo Fisher Scientific) and 1.25 µg of antibodies were conjugated at room temperature for 2 hours, after which, conjugated beads were washed with 0.1% SDS buffer. The lysate was then rotated with the conjugated beads overnight at 4°C. The next day, beads were washed sequentially with 0.1% SDS solution high salt wash.

For retinoic acid (RA) treatment, cells were dissociated with TrypLE™ Express and seeded at a density of 5×10^4 per cm^2 in mTeSR1™ supplemented with 10 µM Y-27632. After 24 h, cells were grown in mTeSR1™ supplemented with 5 µM RA (Sigma) for up to 96 h. DMSO was used at a final concentration of 1:10,000 as a control for RA treatment.
(50 mM HEPES-KOH, pH 7.5, 500 mM NaCl, 2 mM EDTA, 1% (v/v) Triton-X100, 0.1% (v/v) Na-DOC, 0.1% (w/v) SDS), NP40/LiCl wash (10 mM Tris-HCl, pH 8.0, 250 mM LiCl, 1 mM EDTA, 0.5% (v/v) NP-40, 0.1% (w/v) Na-DOC) and TE (1 0M Tris-HCl pH 8.0, 1 mM EDTA). The beads were then resuspended in ChIP elution buffer (50 mM Tris-HCl, pH 7.4, 10 M EDTA, 1% (w/v) SDS) and shakened at 65°C at 1000 rpm for 1 hour. This supernatant was then transferred to a new tube, treated with 1:50 Proteinase K (20 mg/ml) (Roche) and shakened at 55°C at 600 rpm for 1 hour. For siELK1 and MIM treatment, DNA was then further purified using the QIAquick PCR purification kit (Qiagen). For shELK1 and RA treatment, the eluted DNA was mixed with an equal volume of phenol-chloroform (Thermo Fisher Scientific). The aqueous layer was isolated following centrifugation, for 10 minutes at 13.1 krpm at 4°C, and mixed 1:1 with isopropanol and frozen at −80°C for 30 minutes. The solution was then spun at 13.1 krpm at 4°C for 20 minutes and the supernatant discarded. The pellet was then washed with 70% ethanol and spun two more times. It was then air-dried for 24 h and 100 µl of water was added.

A PCR reaction was then run with the following settings: 50°C for 30 min (only for RT-PCR), followed by 95°C for 20 min then [95°C for 20 s, 55°C for 30 s, 72°C for 30 s] for 40 cycles, melt curve 72–95°C. ChIP-qPCR samples in Figure 3 and RT-qPCR in Figure 4 were analysed with the BioMark HD System (Fluidigm), used as per the manufacturer’s instructions. The 14-cycle Specific Target Amplification was used for pre-amplification of the ChIP product and Exonuclease I treatment was used to remove unincorporated primers. BioMark Data Analysis (Fluidigm) was used for data analysis. PCR primers are shown in Extended data, Supplementary Table S1 (Sharrocks, 2019).

### RT-qPCR assays
For MIM stimulation and siELK1 experiments, RNA was purified with an RNaseasy Kit (Qiagen) using the manufacturer’s protocol. For RA and shELK1 experiments, cells were collected into 350 µl of RNAzol (Sigma) homogenised with a Gilson pipette, and spun at 13.1 krpm at 4°C for 20 minutes. The aqueous layer was then mixed 1:1 with isopropanol and RNA precipitated and dried as described for ChIP-isolated DNA above. cDNA was then prepared using SuperScript2 (Thermo Fisher Scientific) according to the manufacturer’s instructions. PCR was then carried out using Power SybrGreen (Thermo Fisher Scientific) with an annealing temperature of 55°C, according to the manufacturer’s instructions. Data was collected with the ViiA 7 (Thermo Fisher Scientific) and analysed with ViiA7 V1.2 software (Thermo Fisher Scientific). The PCR primers are shown in Extended data, Supplementary Table S1 (Sharrocks, 2019).

### Western blotting
For Western blot analysis, cells were harvested in RIPA buffer by scraping on ice. Cell lysates were then centrifuged at 13.1 krpm at 4°C for 2 minutes. The supernatant was then measured using a Bradford Protein Assay. Next, 1 µl of sample was added to 1 ml of Coomassie Brilliant Blue (ThermoFisher Scientific) and measured against BSA standards ranging from 0.2 mg/ml to 2 mg/ml. Approximately 20 µg of protein was used for each well. 1x SDS loading buffer was added to the lysate, which was then boiled at 99°C for 10 minutes. Proteins were resolved on the 12% gel in 1x SDS running buffer, and transferred to a nitrocellulose membrane using transfer buffer. Finally, cells were incubated with primary and secondary antibodies (Extended data, Supplementary Table S2 (Sharrocks, 2019)), diluted 1:500–2000 and 1:10000, respectively, in Licor Odyssey Buffer and imaged on the Odyssey Imaging System (Licor biosciences).

### Bioinformatics analysis
For ChIP-seq analysis from published datasets from human H1-hESCs (in Extended data, Supplementary Table S3 (Sharrocks, 2019)), reads were mapped to the genome using Bowtie2 (v2.2.9) with default settings (Langmead & Salzberg, 2012). Bowtie2 output was then sorted, compressed and unaligned reads removed using samtools (v0.1.18), using the settings -shu -F4, which removed unmapped reads (Li et al., 2009). Finally, peaks were called with MACS2, using the default settings (Zhang et al., 2008). To identify intersecting peaks from two datasets, after MACS2 peak calling, narrowPeak files were intersected with the intersectBed tool in bedtools (v2.21)(Quinlan & Hall, 2010), using the -f 0.1 and -r settings, creating a reciprocal overlap of 10%.

To create tag density graphs, the mapped, sorted and compressed ChIP-seq files were converted to BED files using the bamtobed tool in bedtools. BED files were converted to tag directories using the makeTagDirectory.pl tool in HOMER (v4.8.3) (Heinz et al., 2010). Finally, ChIP-peaks were annotated using annotatePeaks.pl tool in HOMER, using the settings -size -2500, 2500 -hist 25 -norm 0, which created a tag density profile with a 25 bp bin, averaged to tag count in all tag directories, 2500 bp on either side of the peak centre.

To associate peaks to potential target genes we used the nearest gene model in HOMER (with default settings; Heinz et al., 2010). Enriched functional or biological processes associated with these genes were identified from lists of gene ontology (GO) terms using DAVID v6.7 (Huang et al., 2008; Huang et al., 2009).

### Statistical analysis
Pairwise student’s t-tests were performed in GraphPad v7. Statistical significance determined using the Holm-Sidak method, with significance set to p < 0.05. Hypergeometric p-values were calculated using the phyper function in R v3.4.1.

### Data availability
**Underlying data**
Raw data underlying the findings of this study are available from figshare. These include raw GO data (Figure 1F, G; GO terms), RT-qPCR data (Figure 2A: RT-qPCR data for RA stimulation; Figure 2B, D–F: RT-qPCR data plus shELK1; Figure 3A: RT-qPCR data for pluripotency genes in hESCs and mesoderm cells; Figure 3E: RT-qPCR data for pluripotency genes plus siELK1; Figure 3F: RT-qPCR data for differentiation factor
genes plus siELK1; Figure 6D: RT-qPCR data in mesoderm and hESCs plus siELK1; Figure 6B and C: RT-qPCR data for EGR1-3 plus/minus MIM stimulation; and Figure 7: RT-qPCR data with CytoD treatment). ChIP-qPCR data (Figure 2D: ChIP-qPCR data for ELK1 plus siELK1; Figure 2E: ChIP-qPCR data for SUZ12 plus siELK1), Fluidigm data (Figure 4 and Figure 5: ChIP-qPCR data plus siELK1 or siSRF), and uncropped western blots (Figure 2C, Figure 3C, D and Figure 6).

DOIs: GO, https://doi.org/10.6084/m9.figshare.7657958 (Prise, 2019a); RT-qPCR, https://doi.org/10.6084/m9.figshare.7667099 (Prise, 2019b); ChIP-qPCR, https://doi.org/10.6084/m9.figshare.7657961 (Prise, 2019c); Fluidigm, https://doi.org/10.6084/m9.figshare.7657931 (Prise, 2019d); western blots, https://doi.org/10.6084/m9.figshare.7657682 (Prise, 2019e).

Extended data

Extended data are available from figshare.

Supplemental Table S1. Human H1-hESCs ChIP-seq data set accession numbers used in this study.

Supplemental Table S2. Antibodies used in the study.

Supplemental Table S3. Human H1-hESCs ChiP-seq data set accession numbers used in this study.

DOI: https://doi.org/10.6084/m9.figshare.7695617 (Sharrocks, 2019).

Data are available under the terms of the Creative Commons Attribution 4.0 International license (CC-BY 4.0).

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The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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Reviewer Report 08 July 2019
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Ron Prywes
Department of Biological Sciences, Columbia University, New York, NY, 10027, USA

The authors have satisfactorily responded to my comments. While they have not ruled out a role for Elk1 repression through blocking MRTF-A/B binding to SRF, they have qualified their conclusions. (This still appears plausible for the Egr2 gene, which is activated by Cytochalasin D in Figure 7, and further activated by Elk1 knockdown.) In general this is an interesting study showing Elk1 repression of target genes co-bound by SRF. The mechanism of Elk1 repression remains to be determined.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Cell signaling.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 08 July 2019
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Alexis Verger
UMR 8576, Unité de Glycobiologie Structurale et Fonctionnelle (UGSF), CNRS, Université de Lille, Lille, France

The authors have addressed appropriately all my concerns.

Competing Interests: No competing interests were disclosed.
Reviewer Expertise: ETS transcription factor family, regulation of gene expression.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

In this paper, the authors evaluate the dual activating and repressive role of the ETS transcription factor ELK-1 in human embryonic stem cells. The paper builds upon a previous study (Göke et al. Molecular Cell 2013) in which ERK genome-wide chromatin interactions in human embryonic stem cells was characterized. In the present paper, the authors observed a significant overlap in ELK-1 and SUZ12 (PRC2 complex) binding regions but they don’t find any evidence that ELK-1 does have a role in mediating transcriptional repression through these regions. Interestingly they find that ELK-1 has a repressive role during mesoderm differentiation that appears not to be exerted through competition with MRTF co-activators of RhoA-mediated gene expression. The paper represents an interesting follow up of the earlier work and is written in a clear and logical fashion. The purpose of the study is well defined and technically sound. However some data and aspects of the manuscript are worth clarifying.

My understanding is that ELK-1 binding motif is only enriched in the subset of ELK-1 binding sites that are co-occupied by ERK2 whereas promoters bound by ELK-1 without ERK2 (and thus with SUZ12 and EZH2) do not have this ELK-1 binding motif, at least for a 500bp window. Is this also the case with your 2500bp window? If true, do the authors have any explanation of how ELK-1 is recruited?

The RT-qPCR experiments after retinoic acid (RA) treatment appear somewhat highly variable especially for PAX7 and POUF3F2. Do the authors identify more than the six ELK-1+SUZ12 bound genes whose expression increased upon RA treatment? If yes, it should be worth to include them. Is the expression of ELK-1 affected by RA?

The role of ELK-1 during mesoderm expression should be discussed in more detail. The authors nicely show that ELK-1 depletion caused an increase in the expression of FOXC1 and HAND2 (Figure 3F) but in the meantime the Western-blots (Figure 3C) suggests that ELK-1 protein expression is induced during mesoderm differentiation. Is this possible that these genes are ELK-1/PRC2-mediated repression dependent?
Figure 5. FOXC1 (in relation with RT-qPCR Figure 3F) is included in the heatmaps of chromatin Ip but not HAND2. If the ChiP-qPCR has been done for HAND2, it could be nice to add it.

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Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Partly

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: ETS transcription factor family, regulation of gene expression

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Reviewer Report 12 March 2019
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Ron Prywes
Department of Biological Sciences, Columbia University, New York, NY, 10027, USA

The authors nicely evaluate the role of Elk1 in regulating gene expression in human embryonic stem cells. Elk1 co-binds to DNA with SRF, but can also bind DNA without SRF. They don’t find evidence that the sites bound without SRF are functional here. Interestingly, they find that Elk1 can function as an activator
or a repressor depending upon the exact gene binding site they examine.

One conclusion that I feel is less well proven is that Elk1 repression is independent of its blocking another SRF co-factor MRTF-A and -B. Their evidence is that activation of MRTF-A/B by cytochalasin D is not stimulated after Elk1 knockdown. However, the activation of specific genes in these cells could be due to basal MRTF-A/B activation (or is the related myocardin gene expressed in these cells?). If so, perhaps there is something about these cells that limits cytochalasin activation of MRTF-A/B at these genes. A more thorough way to test this question is to knockdown the members of the MRTF family along with Elk1 knockdown to see whether the derepression is affected.

Some more minor points:
1. In Figure 6, different cell treatments are used in 6B and 6D, E. The differences are not entirely clear. In 6B, there is a brief treatment of MIM media. What is done in the other figures? This is important as in 6B, Elk1 activates expression and in 6E it represses.
2. It’s concluded for Fig. 6, on page 6, that Elk1 represses specific genes. This implies that it is doing so directly by binding to them. What rules out an indirect effect? This should at least be discussed.
3. In Figure 2, the authors examine retinoic acid induction of the cells and conclude that Elk1 binding to PRC2 is not important. Later in the paper they use media to induce the cells to differentiate to mesodermal cells. Is it possible that Elk1/PRC2 target genes are important for this type of induction of these cells?
4. We previously found that Elk1 could act as a repressor of MRTF target genes in NIH3T3 cells particularly when grown continuously and found a redundancy for immediate early activation of some genes by Elk1 and MRTF. Are the embryonic stem cells different than other cell types for Elk1 activity and does the growth state of the cells affect Elk1 activity?

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Is the work clearly and accurately presented and does it cite the current literature? Yes

Is the study design appropriate and is the work technically sound? Yes

Are sufficient details of methods and analysis provided to allow replication by others? Yes

If applicable, is the statistical analysis and its interpretation appropriate? Yes

Are all the source data underlying the results available to ensure full reproducibility? Yes

Are the conclusions drawn adequately supported by the results? Partly

Competing Interests: No competing interests were disclosed.
Reviewer Expertise: Cell signaling

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.