The SMAD2/3 interactome reveals that TGFβ controls m^6^A mRNA methylation in pluripotency

Alessandro Bertero1†*, Stephanie Brown1†*, Pedro Madrigal1,2, Anna Osnato1, Daniel Ortmann1, Loukia Yiangou1, Juned Kadiwala1, Nina C. Hubner1, Igor Ruiz de los Mozos4, Christoph Sadée4, An-Sofie Lenaerts1, Shota Nakanoh1, Rodrigo Grandy1, Edward Farnell5, Jernej Ule4, Hendrik G. Stunnenberg3, Sasha Mendjan1† & Ludovic Vallier1,2

The TGFβ pathway has essential roles in embryonic development, organ homeostasis, tissue repair and disease. These diverse effects are mediated through the intracellular effectors SMAD2 and SMAD3 (hereafter SMAD2/3), whose canonical function is to control the activity of target genes by interacting with transcriptional regulators. Therefore, a complete description of the factors that interact with SMAD2/3 in a given cell type would have broad implications for many areas of cell biology. Here we describe the interactome of SMAD2/3 in human pluripotent stem cells. This analysis reveals that SMAD2/3 is involved in multiple molecular processes in addition to its role in transcription. In particular, we identify a functional interaction with the METTL3–METTL14–WTAP complex, which mediates the conversion of adenosine to N^6^-methyladenosine (m^6^A) on RNA. We show that SMAD2/3 promotes binding of the m^6^A methyltransferase complex to a subset of transcripts involved in early cell fate decisions. This mechanism destabilizes specific SMAD2/3 transcriptional targets, including the pluripotency factor gene NANOG, priming them for rapid downregulation upon differentiation to enable timely exit from pluripotency. Collectively, these findings reveal the mechanism by which extracellular signalling can induce rapid cellular responses through regulation of the epitranscriptome. These aspects of TGFβ signalling could have far-reaching implications in many other cell types and in diseases such as cancer.

Activin and NODAL, two members of the TGFβ superfamily, play essential roles in cell fate decision in human pluripotent stem cells (hPSCs). Activin–NODAL signalling is necessary to maintain pluripotency, and inhibition of this pathway drives differentiation towards the neuroectoderm lineage. Activin–NODAL signalling also cooperates with BMP and WNT pathways to drive mesendoderm specification. Therefore, we used the differentiation of hPSCs into definitive endoderm as a model system to investigate the SMAD2/3 interactome during a dynamic cellular process. To allow a comprehensive and unbiased examination of the proteins that interact with SMAD2/3, we developed an optimized SMAD2/3 co-immunoprecipitation protocol that is compatible with mass-spectrometry analyses (Extended Data Fig. 1a, b and Supplementary Discussion). By examining human embryonic stem cells (hESCs) and hESC-like cells that have been induced to differentiate to endoderm (Fig. 1a), we identified 89 interacting partners of SMAD2/3 (Fig. 1b, Extended Data Fig. 1c, d and Supplementary Table 1). Eleven of these proteins interacted with SMAD2/3 either in hESCs or the differentiating cells but not in both (Extended Data Fig. 1e), suggesting that the SMAD2/3 interactome is largely conserved across these two cell types (Supplementary Discussion). Notably, this list includes known SMAD2/3 transcriptional and epigenetic cofactors (including FOXH1, SMAD4, SNON, SKI, EP300, SETDB1 and CREBBP). We also performed functional experiments on FOXH1, EP300, CREBBP and SETDB1, which uncovered essential functions of these SMAD2/3 transcriptional and epigenetic cofactors in hPSC fate decisions (Extended Data Figs 2, 3 and Supplementary Discussion).

These proteomic experiments also show that SMAD2/3 interacts with complexes involved in functions that have, to our knowledge, not previously been associated with TGFβ signalling (Fig. 1b and Extended Data Fig. 1f), such as ERCC1–XPF and DAPK3–PAWR, which are involved in DNA repair and apoptosis, respectively. We also identified several factors involved in mRNA processing, modification and degradation (Fig. 1b), such as the METTL3–METTL14–WTAP complex (involved in deposition of N^6^-methyladenosine (m^6^A)), the PABP-dependent poly(A) nuclelease complex (PAN, involved in mRNA decay), the cleavage factor complex CFIm (involved in pre-mRNA 3’ end processing) and the NONO–SFPQ–PSPC1 factors (involved in RNA splicing and nuclear retention of defective RNAs). Overall, these results suggest that SMAD2/3 could be involved in a large number of biological processes in hPSCs, including not only transcriptional and epigenetic regulation, but also non-canonical functions of SMAD2/3.

To further investigate these functions, we examined the function of activin–NODAL signalling in m^6^A deposition. m^6^A is the most common RNA modification, and it regulates multiple aspects of mRNA biology including decay and translation. However, it is not known whether this is a dynamic event that can be modulated by extracellular cues. Furthermore, whereas m^6^A is known to regulate haematopoietic stem cells and the transition between the naive and primed pluripotency states, its function in hPSCs and during germ layer specification is unknown. We first validated the interaction of SMAD2/3 with METTL3–METTL14–WTAP by co-immunoprecipitation followed by western blotting with both hESCs and human induced pluripotent stem cells (hiPSCs; Fig. 2a and Extended Data Fig. 4a, b). Inhibition of SMAD2/3 phosphorylation blocked this interaction (Fig. 2b and Extended Data Fig. 4c). Proximity ligation assays (PLA) also demonstrated that the interaction occurs in the nucleus (Fig. 2c, d). These observations suggest that SMAD2/3 and the m^6^A methyltransferase complex interact and that this interaction depends on activin–NODAL signalling.

To investigate the functional relevance of this interaction, we assessed the transcriptome-wide effects of inhibition of activin–NODAL signalling on the deposition of m^6^A by performing nuclear-enriched m^6^A-methylated-RNA immunoprecipitation followed by deep sequencing (NeMeRIP-seq; Extended Data Fig. 5a–d, and Supplementary Discussion). Consistent with previous reports, deposition of m^6^A onto exons was enriched around stop codons and transcription start sites, and occurred at a motif corresponding to the m^6^A-consensus sequence. Assessment of differential m^6^A deposition revealed that inhibition of activin–NODAL
signalling predominantly resulted in decreased m^6^A levels in selected transcripts (Supplementary Table 2; mean absolute log_2^ fold-change of 0.56 and 0.35 for m^6^A decrease and increase, respectively). Decreases in m^6^A deposition were mostly observed on peaks located near stop codons (Extended Data Fig. 5h), where m^6^A deposition has been reported to decrease the stability of mRNAs\(^{16,24,25}\). Transcripts with reduced m^6^A levels after inhibition of activin–NODAL signalling largely and significantly overlapped with genes bound by SMAD2/3 (\(P = 2.88 \times 10^{-18}\); Extended Data Fig. 5i), including well-known transcriptional targets such as NANOG, NODAL, LEFTY1 and SMAD7 (Fig. 2e and Extended Data Fig. 5j). Accordingly, activin–NODAL-sensitive m^6^A deposition was largely associated with transcripts that rapidly decreased in abundance during the exit from pluripotency triggered by inhibition of activin–NODAL signalling (Extended Data Fig. 6a). Transcripts that behaved in this fashion were enriched in pluripotency regulators and factors involved in the activin–NODAL signalling pathway (Supplementary Table 3). On the other hand, the expression of a large number of developmental regulators associated with activin–NODAL-sensitive m^6^A deposition remained unchanged following inhibition of activin–NODAL signalling (Extended Data Fig. 6a–c and Supplementary Table 3). Considered together, these findings show that activin–NODAL signalling can regulate m^6^A deposition on a number of specific transcripts.

We then examined the molecular mechanisms that underlie the regulation of m^6^A deposition by activin–NODAL signalling. RNA immunoprecipitation experiments on nuclear RNAs showed that inhibition of activin–NODAL signalling impaired binding of WTAP to several m^6^A-labelled transcripts including NANOG and LEFTY1 (Fig. 2f and Extended Data Fig. 4d, e), whereas SMAD2/3 itself interacted with these transcripts in the presence of activin–NODAL signalling (Fig. 2g and Extended Data Fig. 4e). Thus, SMAD2/3 appears to promote the recruitment of the m^6^A methyltransferase complex to nuclear RNAs. Notably, recent reports have established that m^6^A deposition occurs co–transcriptionally and involves nascent pre-RNAs\(^{16,20,26}\). Considering the broad overlap between SMAD2/3 transcriptional targets and transcripts showing activin–NODAL-sensitive m^6^A deposition (Extended Data Fig. 5i), we hypothesized that SMAD2/3 could facilitate co–transcriptional recruitment of the m^6^A methyltransferase complex onto nascent transcripts. Consistent with this notion, inhibition of activin–NODAL signalling mainly resulted in downregulation of m^6^A, not only on exons, but also on pre-mRNA-specific features such as introns and exon–intron junctions (Extended Data Fig. 6d–i and Supplementary Table 2). Moreover, we observed a correlation in activin–NODAL sensitivity across m^6^A peaks within the same transcript (Extended Data Fig. 6j), suggesting that SMAD2/3 regulates m^6^A deposition at the level of the genomic locus rather than on a specific mRNA peak. Nevertheless, we did not detect stable and direct binding of the m^6^A methyltransferase complex to DNA (Extended Data Fig. 4f). Thus, co–transcriptional recruitment might rely on indirect and dynamic interactions with chromatin. Considering all these results, we propose a model in which activin–NODAL signalling promotes co–transcriptional m^6^A deposition by facilitating the recruitment of the m^6^A methyltransferase complex onto nascent mRNAs (Fig. 2h).

To understand the functional relevance of this regulation in the context of hPSC cell–fate decisions, we performed inducible knockdown of the subunits of the m^6^A methyltransferase complex\(^{27}\) (Extended Data Fig. 4g, h).
comparisons. Two-way analysis of variance (ANOVA) with post hoc Holm–Sidak controls as they do not contain m6A. In Fig. 7a, b, METTL3, METTL14 and WTAP decreased m6A deposition (Extended Data Fig. 7c, d); however, prolonged knockdown did not affect pluripotency (Extended Data Fig. 7e, f). We also found that expression of m6A methyltransferase complex subunits was necessary for neuroectoderm differentiation induced by the inhibition of activin–NODAL signalling without being necessary for activin-driven endoderm specification (Fig. 3a and Extended Data Fig. 8a–c). Notably, Activin–NODAL is known to block neuroectoderm induction by promoting NANOG expression, whereas NANOG is required for the early stages of endoderm specification. Accordingly, we found that NANOG transcript and protein were upregulated, and the stability of NANOG mRNA increased when m6A methyltransferase activity was impaired (Fig. 3b and Extended Data Fig. 9a–c). These results show that m6A deposition decreases the stability of NANOG mRNA, facilitating its downregulation upon loss of activin–NODAL signalling, and thereby facilitating exit from pluripotency and neuroectoderm specification (Extended Data Fig. 9d). Additional transcriptomic analyses showed that WTAP knockdown resulted in global upregulation of genes that were transcriptionally activated by SMAD2/3 in hESCs and impaired the upregulation of genes induced by inhibition of activin–NODAL signalling during neuroectoderm differentiation (Fig. 3b, Extended Data Fig. 10a–e).
defective neuroectoderm differentiation (Fig. 3d and Extended Data Fig. 8e, f). Together, these results show that the interaction of SMAD2/3 with METTL3–METTL14–WTAP can promote m^6^A deposition on a subset of transcripts, including a number of pluripotency regulators that are also transcriptionally activated by activin–NODAL signalling. The resulting negative feedback destabilizes these mRNAs and causes their rapid degradation following inhibition of activin–NODAL signalling. This mechanism allows timely exit from pluripotency and induction of neuroectoderm differentiation (Extended Data Fig. 9d).

In conclusion, this analysis of the SMAD2/3 interactome reveals interactions between TGF^3^ signalling and a wide variety of cellular processes. Our results suggest that SMAD2/3 could act as a hub, coordinating several proteins known to have a role in mRNA processing and modification, apoptosis, DNA repair and transcriptional regulation. This function is illustrated by our results that show activin–NODAL-sensitive regulation of m^6^A. Activin–NODAL signalling connects transcriptional and epitranscriptional regulation through the interaction between SMAD2/3 and the METTL3–METTL14–WTAP complex, and primes its transcriptional targets for rapid degradation upon withdrawal of signalling (Extended Data Fig. 9d). This avoids overlaps between the pluripotency and neuroectoderm transcriptional programs, thereby facilitating changes in cell identity. We anticipate that further studies will clarify the other non-canonical functions of SMAD2/3, and will dissect how they are related to epigenetic, transcriptional and epitranscriptional regulation of gene expression.

Our findings also clarify and broaden our understanding of the function of m^6^A in cell-fate decisions. They establish that depletion of m^6^A in hPSCs does not lead to differentiation, contrary to predictions from studies in mouse-epiblast stem cells. This could imply that there are important functional differences in epitranscriptional regulations between human and mouse pluripotent states. Moreover, widening the conclusions from previous reports, we demonstrate that deposition of m^6^A is specifically necessary for neuroectoderm induction, but not for definitive endoderm differentiation. This can be explained by the fact that in contrast to its strong inhibitory effect on the neuroectoderm lineage expression of NANO^G^ is necessary for the early stages of mesendoderm specification. Finally, our results establish that m^6^A modification of RNA is a dynamic event that is directly modulated by extracellular cues such as TGF^3^. Considering the many functions of TGF^3^ signalling, the regulation we describe here may have an essential function in many cellular contexts that require a rapid response or change in cell state, such as the inflammatory response or cellular proliferation.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Author Contributions A.B. conceived the study, performed or contributed to most of the experiments, analysed data and wrote the manuscript with input from the other authors. S.B. contributed to study conception, performed immunoprecipitation, RNA-IP and selection experiments, analyzed data. P.M., I.R.D.I.M. and C.S. analysed NeMeRIP–seq. A.O. performed PLA and co-immunoprecipitations and analysed RNA-seq data. D.O., L.Y. and J.K. assisted with hPSC gene editing and differentiation; N.C.H. performed quantitative proteomics and data analysis. A.-S.L., S.N. and R.G. assisted with hPSC culture. E.F. optimized NeMeRIP–seq libraries. J.U. contributed to study conception and supervision. H.G.S. supervised quantitative proteomics. S.M. contributed to study conception and supervision, and assisted with SMAD2/3 co-immunoprecipitation. L.V. conceived, supervised and supported the study, and wrote and provided final approval of the manuscript.

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METHODS

hPSC culture and differentiation. Feeder- and serum-free culture of hESCs (H9/WA09 line; WiCell) and hiPSCs (A1ATR/R; ref. 31) have been previously described. In brief, cells were plated on gelatin and MEF medium-coated plates, and cultured in chemically defined medium (CDM) containing bovine serum albumin (BSA). CDM was supplemented with 10 ng/ml activin-A and 12 ng/ml FGF2 (both from M. Hyvonen, Dept. of Biochemistry, University of Cambridge). Cells were passaged every 5–6 days with collagenase IV and plated as clumps of 50–100 cells. Once cells reached an density of 100–150 cells per cm², passaging was initiated in adherent hESC cultures 48 h after passages. Deriving clonal endoderm specification was induced for three days (unless stated otherwise) by culturing cells in CDM (without insulin) with 20 ng/ml FGF2, 10 μM L2Y294002 (PI3K inhibitor; Promega), 100 ng/ml activin-A and 10 ng/ml BMP4 (R&D), as previously described. Neuroectoderm was induced for three days (unless stated otherwise) in CDM-BSA with 12 ng/ml FGF2 and 10 μM SB431542 (activin–NODAL–TGF3 signalling inhibitor; Tocris), as previously described. These same culture conditions were used for activin–NODAL signalling inhibition experiments. hiPSCs were routinely monitored for absence of karyotypic abnormalities and mycoplasma infection. As hESCs were obtained from a commercial supplier, cell line identification was not performed. hiPSCs were previously generated in house and genotyped by Sanger sequencing.

Molecular cloning. Plasmids carrying inducible shRNAs were generated by cloning annealed oligonucleotides into the pAAV-Puro_JKD or pAAV-Puro_SKD vectors as previously described. All shRNA sequences were obtained from the RNAi Consortium TRC library (https://www.broadinstitute.org/rnaa/public/). Of the shRNAs that had been validated, the most powerful ones were chosen (the sequences are reported in Supplementary Table 5). Generation of a vector containing shRNAs against METTL3, METTL14, and WTAP and matched controls expressing a scrambled shRNA was performed by Sanger sequencing.

A targeting vector for the AAVSI locus carrying constitutively-expressed NANOG was generated starting from pAAV_TRE-eGFP. First, the TRE-eGFP cassette was removed using PspXI and EcoRI, and substituted with the CAG promoter (cut from pR26-CAG-eGFP30 using SpeI and BamHI) by ligating blunt-ended fragments. The resulting vector (pAAV_Puro_CAG) was then used to clone the full-length NANOG transcript, which includes its full 5′ and 3′ UTRs. The full-length NANOG transcript was constructed from three DNA fragments. The 5′ (bases 1–301) and 3′ (bases 1878–2105) ends were synthesized (IDT) with 40 bp overlaps corresponding to pGemZ vector linearized with SmaI. The middle fragment was amplified from cDNA of H9 hESCs obtained by retrotranscription with poly-T primer using primers 5′-TTGGTCTTGGAGACACAGCTGG-3′ and 5′-AAAAACCGTAAAGACATCATAA-3′. The three fragments and the linearized vector were assembled using a Gibson reaction (NEB). The full-length NANOG transcript was then cloned into pAAV_Puro_CAG vector following KpnI and HincII digestion. The resulting vector was named pAAV-Puro_MsiKD-M3M14W. Generation of the matched control vector containing three copies of the scrambled shRNA sequence (pAAV-Puro_MsiKD-SCR3+) has been described previously.

A targeting vector for the AAVSI locus carrying constitutively-expressed NANOG was generated starting from pAAV_TRE-eGFP. First, the TRE-eGFP cassette was removed using PspXI and EcoRI, and substituted with the CAG promoter (cut from pR26-CAG-eGFP30 using SpeI and BamHI) by ligating blunt-ended fragments. The resulting vector (pAAV_Puro_CAG) was then used to clone the full-length NANOG transcript, which includes its full 5′ and 3′ UTRs. The full-length NANOG transcript was constructed from three DNA fragments. The 5′ (bases 1–301) and 3′ (bases 1878–2105) ends were synthesized (IDT) with 40 bp overlaps corresponding to pGemZ vector linearized with SmaI. The middle fragment was amplified from cDNA of H9 hESCs obtained by retrotranscription with poly-T primer using primers 5′-TTGGTCTTGGAGACACAGCTGG-3′ and 5′-AAAAACCGTAAAGACATCATAA-3′. The three fragments and the linearized vector were assembled using a Gibson reaction (NEB) and the sequence of the construct was confirmed by Sanger sequencing. The full-length NANOG transcript was then cloned into pAAV_Puro_CAG vector following KpnI and HincII digestion. The resulting vector was named pAAV-Puro_CAG-NANOG.

Inducible gene knockdown. Clonal inducible knockdown hESCs for METTL3, METTL14, WTAP or matched controls expressing a scrambled shRNA were generated using pAAV-Puro_iKD vectors27 in hESCs expressing a randomly integrated wild-type tetracycline resistance gene. Two wells were transfected for each individual shRNA. This same approach was followed to generate multiple inducible knockdown hESCs for METTL3, METTL14 and WTAP (plasmid pAAV-Puro_MsiKD-M3M14W) or matched controls expressing three copies of the SCR shRNA (plasmid pAAV-Puro_MsiKD-SCR3) in hESCs expressing the metabolically active OCT4-SV1 reporter and MsiKD-SCR3. pAAV-Puro_JKD containing exon 1–3 of OCT4 was linearized with KpnI, digested with HindIII and purified. The resulting vector was named pAAV-Puro_MsiKD-M3M14W. Expression of the full length OCT4 transcript was confirmed by Sanger sequencing. The full length OCT4 transcript was confirmed by Sanger sequencing. The full length OCT4 transcript was confirmed by Sanger sequencing.
Peptide spectra were compared against the human database (Uniprot) using the integrated Andromeda search engine, and peptides were identified with FDR < 0.01, determined by false matches against a reverse decoy database. Peptides were assembled into protein groups with an FDR < 0.01. Protein quantification was performed using the MaxQuant label-free quantification algorithm requiring at least two ratio counts, in order to obtain label-free quantification (LFQ) intensities. Collectively, the MS/MS spectra were matched to ~20,000 known peptides, leading to the identification of 3,635 proteins in at least one of the conditions analysed. Statistical protein analysis was performed using Perseus (MaxQuant). First, common contaminants and reverse hits were removed, and only proteins identified by at least two peptides (one of those being unique to the respective protein group) were considered as high-confidence identifications. Proteins were then filtered for those identified in all replicates of at least one condition. LFQ intensities were converted to their log2 values, and missing intensity values were imputed by representative noise values. One-tailed t-tests were then performed to determine the specific interactors in each condition by comparing the immunoprecipitations with the SMAD2/3 antibody to those with the IgG negative controls. Statistical significance was set with a permutation-based FDR < 0.05 (250 permutations). Fold-enrichment over IgG controls was calculated from LFQ intensities.

This same pipeline was used to analyse mass-spectrometry data based on dimethyl labelling, with the following two exceptions. First, an additional mass of 28.03 Da (light) or 32.06 Da (heavy) was specified as ‘labels’ at the N terminus and at lysines. Second, during statistical analysis of mass-spectrometry data, the outlier significance was calculated based on protein intensity (significance B6), and was required to be below 0.05 for both the forward and the reverse experiments.

**Biological interpretation of mass-spectrometry data.** The SMAD2/3 protein–protein interaction network was generated using Cytoscape v.2.8.3. First, all the annotated and curated interactions involving the SMAD2/3-binding proteins were inferred by interrogating protein–protein interaction databases through the PSICQUIC Universal Web Service Client. IMEx-complying interactions were retained and merged by union. Then, a subnetwork involving only the SMAD2/3 interactors was isolated. Finally, duplicate nodes and self-loops were removed to simplify visualization. Note that based on our results all the proteins shown would be connected to SMAD2/3, but such links were omitted to simplify visualization and highlight those interactions with SMAD2/3 that were already known. Proteins lacking any link and small complexes of less than three factors were not shown, in order to improve presentation clarity. Note that since the nodes representing SMAD2 and SMAD3 shared the same links, they were fused into a single node (SMAD2/3).

Functional enrichment analysis was performed using Fisher’s exact test implemented in Enrichr44, and only enriched terms with a Benjamini–Hochberg adjusted P value < 0.05 were considered. For Gene Ontology (GO) enrichment analysis, the 2015 GO annotation was used. For mouse phenotype enrichment analysis, level 3 of the Mouse Genomic Informatics (MGI) annotation was used. To compare protein abundance in different conditions, a cut-off of absolute LFQ intensity log2, fold-change larger than 2 was chosen, as label-free mass spectrometry is currently not sensitive enough to detect smaller changes with confidence.

**Proximity ligation assay.** Proximity ligation assay (PLA) was performed using the Duolink in situ Red Starter Kit Goat/Rabbit (Sigma–Aldrich). Cells were cultured on glass coverslips and prepared by fixation in 4% paraformaldehyde (PFA) in PBS for 10 min at room temperature, followed by two gentle washes in PBS. All subsequent incubations were performed at room temperature unless otherwise stated. Samples were permeabilized in PBS containing 0.25% Triton X-100 for 20 min, blocked in PBS with 0.5% BSA for 30 min, and incubated with the two primary antibodies of interest (diluted in PBS with 0.5% BSA; see Supplementary Table 6) for 1 h at 37 °C in a humid chamber. The Duolink in situ PLA probes (anti-rabbit minus and anti-goat plus) were mixed and diluted 1:5 in PBS with 0.5% BSA and incubated for 20 min at room temperature. The coverslips were incubated with the PLA probe solution for 1 h at 37 °C in a humidified chamber. Single-antibody and probes-only negative controls were performed for each antibody tested to confirm assay specificity. Coverslips were washed twice in wash buffer A for 5 min under gentle agitation, and incubated with 1 × ligation solution supplemented with DNA ligase 1:40 dilution for 30 min at 37 °C in a humidified chamber. After two more washes in wash buffer B for 2 min with gentle agitation, coverslips were incubated with 1 × amplification solution supplemented with DNA polymerase (1:80 dilution) for 1 h at 40 min at 37 °C in a humid chamber. Samples were protected from light from this step onwards. Following two washes in wash buffer B for 10 min, the coverslips were dried overnight, and finally mounted on a microscope slide using Duolink in situ Mounting Medium with DAPI. Images of random fields of view were acquired using a LSM 700 confocal microscope (Leica) using a Plan-Apochromat 40×/1.3 OIL DIC M27 objective, performing z-stack with optimal spacing (~0.36 μm). Images were analysed automatically using ImageJ. For this, nuclear (DAPI) and PLA z-stacks were first individually flattened (max intensity projection) and thresholded to remove background noise. Nuclear images were further segmented using the watershed function. Total nuclei and PLA spots were quantified using the ‘analyse particle’ function of ImageJ, and nuclear PLA spots were quantified using the ‘spleckle inspector’ function of the ImageJ plugin BioVoxxel.

**RNA immunoprecipitation.** Approximately 2 × 106 cells were used for each RNA immunoprecipitation (RIP). Unless stated otherwise, all biochemical steps were performed on ice or at 4 °C, and ice-cold buffers were supplemented with 10 mM Tris–HCl pH 7.5, 3 mM CaCl2, 2 mM MgCl2, 0.32 M sucrose) for 12 min to induce cell swelling. Then, Triton X-100 was added to a final concentration of 0.3%, and cells were incubated for 6 min to lyse the plasma membranes. Nuclei were pelleted at 600g for 5 min, washed once with ten volumes of IBL, and finally resuspended in two volumes of nuclear lysis buffer (NLB; 50 mM Tris–HCl pH 7.5, 100 mM NaCl, 50 mM KC1, 3 mM MgCl2, 1 mM EDTA, 10% glycerol, 0.1% Tween) supplemented with 800 U/ml RNasin Ribonuclease Plus Inhibitor (Promega) and 1 μM DTT. The nuclear suspension was transferred to a Dounce homogenizer (Jencons Scientific) and homogenized by performing 70 strokes with a tight pestle. The nuclear lysate was incubated with rotation for 30 min, homogenized again by performing 30 additional strokes with the tight pestle, and incubated in rotation for 15 min at room temperature after addition of 12.5 μl/ml of DNase I (Sigma).

The protein concentration was assessed, and approximately 1 mg of protein was used for overnight immunoprecipitation with rotation with the primary antibody of interest (Supplementary Table 6), or with equal amounts of non-immune species-matched IgG. Ten per cent of the protein lysate used for immunoprecipitation was saved as pre-immunoprecipitation input and stored at ~80 °C for subsequent RNA extraction. Immunoprecipitation reactions were then incubated for 1 h with 30 μl of protein–G agarose, then washed twice with 1 ml of Lici wash buffer (50 mM Tris–HCl pH 7.5, 250 mM LiCl, 0.1% Triton X-100, 1 mM DTT) and twice with 1 ml of NLB. Beads were resuspended in 90 μl of 30 mM Tris–HCl pH 9.0, and DNase-digested using the RNase-free DNase kit (QIAGEN) by adding 10 μl of RDD buffer and 2.5 μl of DNase. The pre-immunoprecipitation input samples were similarly treated in parallel, and samples were incubated for 10 min at room temperature. The reaction was stopped by adding 2 mM EDTA and by heating at 70 °C for 5 min. Proteins were digested by adding 2 μl of proteinase K (20 mg/ml; Sigma–Aldrich) and by incubating at 37 °C for 30 min. Finally, RNA was extracted with 1 ml of TRIReagent (Sigma–Aldrich) according to the supplier’s instructions.

The RNA was resuspended in nuclease-free water, and half of the sample was used in a reverse-transcription reaction using SuperScript II (ThermoFisher) using the manufacturer’s protocol. The other half was used in a control reaction with no reverse transcriptase to confirm successful removal of DNA contaminants. Samples were quantified by quantitative real-time PCR (qPCR), and normalized first to the housekeeping gene GAPDH and then to the IgG control using the ΔΔCt approach (see below). Supplementary Table 5 shows the primers used.

**Chromatin immunoprecipitation.** Approximately 2 × 106 cells were used for each chromatin immunoprecipitation (ChiP), and cells were fed with fresh medium 2 h before collection. ChiP was performed using a previously described protocol10,30. Briefly, cells were crosslinked on plates, first with protein–protein crosslinkers (10 mM dimethyl 3,3′-dithiopropionimidate dihydrochloride and 2.5 mM 3,3′-dithiodipropionic acid di-N-hydroxy succinimide ester; Sigma–Aldrich) for 15 min at room temperature, then with 1% formaldehyde for 15 min. Crosslinking was quenched with glycine, after which cells were collected, subjected to nuclear lysis, and centrifuged for 5 min at 1000 g. The supernatant was incubated overnight with the antibodies of interest (Supplementary Table 6) or non-immune IgG. ChiP was completed by incubation with protein–G agarose beads followed by subsequent washes with high salt and LiCl-containing buffers (all exactly as previously described10,30). Crosslinking was reversed, first by adding DTT (for disulfide bridge-containing protein–protein crosslinkers), then by incubating in high salt at high temperature. DNA was finally purified by sequential phenol–chloroform and chloroform extractions. Samples were analysed by qPCR using the ΔΔCt approach (see Supplementary Table 5 for primer sequences). First, a region in the last exon of SMAD7 was used as internal control to normalize for the enrichment of transcription input and then to the enrichment observed in non-immune IgG ChiP controls. m^A dot blot. m^A dot blots were performed as described with minor modifications23. Poly-A RNA was purified from total cellular RNA using the Dynabeads mRNA Purification Kit (ThermoFisher), diluted in 50 μl of RNA loading buffer (RLB; 2.2 M formic acid, 50% formamide, 0.5 × MOPS buffer containing 0.6 M NaCl, 0.25% SDS).
(20 mM MOPS, 12.5 mM CH₃COONa, 1.25 mM EDTA, pH 7.0), incubated at 55°C for 15 min, and snap-cooled on ice. An Amersham Hyper-CL XL column was equilibrated in water for 3 min, then in 10X saline-sodium citrate buffer (SSC; 1.5 M NaCl, 150 mM Na₂HPO₄, pH 7.0) for 10 min, and finally 'sandwiched' in a 96-well dot blot hybridization manifold (ThermoFisher Scientific). Following two washes of the wells with 150 μl of 10X SSC, the RNA was spotted onto the membrane. After UV cross-linking for 2 min at 254 nm using a Stratalinker 1800 (Stratagene), the membrane was washed once with Tris-buffered saline TWEEN buffer (TBST; 10 mM Tris·HCl pH 7.4, 150 mM NaCl, 0.05% Tween-20). Each wash was performed by incubating the beads with 500 μl of TBST with 0.1% SDS, 10 mM DTT, 0.1% Tween-20. Each wash was performed by incubating the beads with 500 μl of TBST for 10 min to induce cell swelling. Then, Triton X-100 was added to a final concentration of 0.3% and cells were incubated for 6 min to lyse the plasma membranes. Nuclei were pelleted at 600 g for 5 min and washed once with ten volumes of lysis buffer (IB). RNA was extracted from the nuclear pellet using the Neasy® kit (QIAGEN) according to the manufacturer’s instructions. Residual contaminating DNA was digested in solution using the RNase-free DNase set from QIAGEN, and RNA was re-purified by sequential acid phenol–chloroform and chloroform extractions followed by ethanol precipitation. At this stage, complete removal of DNA contamination was confirmed by qPCR of the resulting RNA without a reverse-transcription step. RNA was then chemically fragmented in 20 μl reactions each containing 20 μg of RNA in 20 mM ZnCl₂, 10 mM Tris·HCl pH 7.0. Such reactions were incubated at 95°C for 5 min, followed by inactivation with 50 mM EDTA and storage on ice. The fragmented RNA was then cleaned up by ethanol precipitation. In preparation for MeRIP, 15 μg of anti-m6A antibody (Synaptic Systems, 202-003) or equivalent amount of rabbit non-immune IgG was crosslinked to 0.5 μg of magnetic beads using the Dynabeads Antibody Coupling Kit (ThermoFisher Scientific) according to the manufacturer’s instructions. Following equilibration of the magnetic beads by washing with 500 μl of binding buffer (30 mM Tris·HCl pH 7.5, 150 mM NaCl, 1% NP-40, 1 mM EDTA), MeRIP reactions were assembled with 300 μg of the fragmented RNA in 3 ml of binding buffer supplemented with 3,000 U of RNasin RNase inhibitor (Promega), 1 mM DTT for 10 min to induce cell swelling. Then, RNA was extracted from the nuclear pellet using the Neasy® kit (QIAGEN) according to the manufacturer’s instructions. Residual contaminating DNA was digested in solution using the RNase-free DNase set from QIAGEN, and RNA was re-purified by sequential acid phenol–chloroform and chloroform extractions followed by ethanol precipitation. At this stage, complete removal of DNA contamination was confirmed by qPCR of the resulting RNA without a reverse-transcription step. RNA was then chemically fragmented in 20 μl reactions each containing 20 μg of RNA in 20 mM ZnCl₂, 10 mM Tris·HCl pH 7.0. Such reactions were incubated at 95°C for 5 min, followed by inactivation with 50 mM EDTA and storage on ice. The fragmented RNA was then cleaned up by ethanol precipitation. In preparation for MeRIP, 15 μg of anti-m6A antibody (Synaptic Systems, 202-003) or equivalent amount of rabbit non-immune IgG was crosslinked to 0.5 μg of magnetic beads using the Dynabeads Antibody Coupling Kit (ThermoFisher Scientific) according to the manufacturer’s instructions. Following equilibration of the magnetic beads by washing with 500 μl of binding buffer (30 mM Tris·HCl pH 7.5, 150 mM NaCl, 1% NP-40, 1 mM EDTA), MeRIP reactions were assembled with 300 μg of the fragmented RNA in 3 ml of binding buffer supplemented with 3,000 U of RNasin RNase inhibitor. Samples were incubated with rotation at 7 r.p.m. for 1 h at room temperature. Fragmented RNA (5 μg; 10% of the amount used for MeRIP) was first treated with 0.1 μl of Turbo DNase (Ambion) for 15 min at 37°C, followed by 1 μl of RNasin RNase inhibitor blocked for 1 h at room temperature with TBST supplemented with 4% non-fat dry milk. Incubations with the anti-m6A primary antibody (Synaptic Systems, 202-111; used at 1 μg/ml) and the mouse-β-Actin antibody (Supplementary Table 6) were each performed in TBST with 4% milk for 1 h at room temperature, and were followed by three 10-min washes at room temperature in TBST. Finally, the membrane was incubated with ECL2 Western Blotting Substrate (Pierce), and exposed to X-Ray Super RX film.

m6A nuclear-enriched methylated RNA immunoprecipitation. m6A MeRIP of nuclear-enriched RNA for analysis by deep sequencing (MeRIP–seq) was performed using modifications of previously described methods.24,25 For each sample, 7.5 x 10⁷ hESCs were used, and three biological replicates were performed per condition. Cells were fed with fresh medium for 2 h before washing with PBS, scraping in cell dissociation buffer (CDB, Gibco), and pelleting at 250 x g for 5 min. The cell pellet was incubated in five volumes of isotypic lysis buffer (ILB; 10 mM Tris·HCL pH 7.5, 3 mM CaCl₂, 2 mM MgCl₂, 0.32 M sucrose, 1,000 U/ml RNasin RNase inhibitor (Promega), 1 mM DTT) for 10 min to induce cell swelling. Then, Triton X-100 was added to a final concentration of 0.3% and cells were incubated for 6 min to lyse the plasma membranes. Nuclei were pelleted at 600 g for 5 min and washed once with ten volumes of ILB. RNA was extracted from the nuclear pellet using the Neasy® kit (QIAGEN) according to the manufacturer’s instructions. Residual contaminating DNA was digested in solution using the RNase-free DNase set from QIAGEN, and RNA was re-purified by sequential acid phenol–chloroform and chloroform extractions followed by ethanol precipitation. At this stage, complete removal of DNA contamination was confirmed by qPCR of the resulting RNA without a reverse-transcription step. RNA was then chemically fragmented in 20 μl reactions each containing 20 μg of RNA in fragmentation buffer (10 mM ZnCl₂, 10 mM Tris·HCl pH 7.0). Such reactions were incubated at 95°C for 5 min, followed by inactivation with 50 mM EDTA and storage on ice. The fragmented RNA was then cleaned up by ethanol precipitation. In preparation for MeRIP, 15 μg of anti-m6A antibody (Synaptic Systems, 202-003) or equivalent amount of rabbit non-immune IgG was crosslinked to 0.5 μg of magnetic beads using the Dynabeads Antibody Coupling Kit (ThermoFisher Scientific) according to the manufacturer’s instructions. Following equilibration of the magnetic beads by washing with 500 μl of binding buffer (30 mM Tris·HCl pH 7.5, 150 mM NaCl, 1% NP-40, 1 mM EDTA), MeRIP reactions were assembled with 300 μg of the fragmented RNA in 3 ml of binding buffer supplemented with 3,000 U of RNasin RNase inhibitor. Samples were incubated with rotation at 7 r.p.m. for 1 h at room temperature. Fragmented RNA (5 μg; 10% of the amount used for MeRIP) was first treated with 0.1 μl of Turbo DNase (Ambion) for 15 min at 37°C, followed by 1 μl of RNasin RNase inhibitor blocked for 1 h at room temperature with TBST supplemented with 4% non-fat dry milk. Incubations with the anti-m6A primary antibody (Synaptic Systems, 202-111; used at 1 μg/ml) and the mouse-β-Actin antibody (Supplementary Table 6) were each performed in TBST with 4% milk for 1 h at room temperature, and were followed by three 10-min washes at room temperature in TBST. Finally, the membrane was incubated with ECL2 Western Blotting Substrate (Pierce), and exposed to X-Ray Super RX film.

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and DESeq\textsuperscript{25}. For assessment of reproducibility, regularized log transformation of count data was computed, and biological replicates of input samples of the same condition were clustered together in the PC space\textsuperscript{26}. Estimation of differential m\textsubscript{6}A deposition onto each peak in NeMeRIP samples versus input controls was performed using an analogous approach. Function-enzymatic analysis of m\textsubscript{6}A-marked transcripts was performed using Enrichr\textsuperscript{44}, as described above for mass-spectrometry data. The coordinates of SMAD2/3 ChIP-seq peaks in hESCs\textsuperscript{36} were transferred from their original mappings on hg19 to hg38 using liftOver. Overlap of differential expression intervals with m\textsubscript{6}A peaks significantly downregulated after 2 h of SB431542 treatment was determined using CAT\textsuperscript{46} with default parameters. SMAD2/3-binding sites were assigned to the nearest gene using the annotatePeaks.pl function from the HOMER suite\textsuperscript{61} with standard parameters. The significance in the overlap between the resulting gene list and that of genes encoding for transcripts with m\textsubscript{6}A peaks that are significantly downregulated after 2 h of SB431542 treatment was calculated by a hypergeometric test where the population size corresponded to the number of genes in the standard Ensembl annotation (GRCh.38.83). m\textsubscript{6}A peaks on introns were identified in three steps (Extended Data Fig. 6d). First, MetaDiff was used to simultaneously perform peak calling and differential methylation analysis. Since MetaDiff only accepts a transcriptome GTF annotation as an input to determine the genomic space onto which it identifies m\textsubscript{6}A peaks, in order to determine peaks on introns, we followed the strategy recommended by the package developers of running the software using a custom transcriptome annotation that includes introns\textsuperscript{46,62}. This ‘extended’ transcriptome annotation was built using Cufflinks\textsuperscript{2.2.1}\textsuperscript{63} with parameters ‘--library-type = fr-firststrand -m 100 -s 50’ and guided by the Ensembl annotation (GRCh.38.83). This was assembled using all available pre-NeMeRIP input reads. The result was an extended transcriptome annotation including all of the transcribed genome that could be detected and reconstructed from our nuclear-enriched input RNA samples, thus including most expressed intron sequences. Then, MetaDiff was run using this extended annotation as input for GENE\_ANNO\_GTF, pooled inputs for each condition, WINDOW\_WIDTH = 40, SLIDING\_STEP = 20, FRAGMENT\_LENGTH = 250, PEAK\_CUTOFF\_PVVALUE = 1E-03, FOLD\_ENRICHMENT = 2, MINIMAL\_MAPQ = 20, and all other parameters as default. In a second step, the peaks identified by MetaDiff were filtered for robustness by requiring that they overlapped with MACS2 peak calls, exactly as for exome-focused MetaDiff peak calls (Extended Data Fig. 5d). Finally, only peaks that strictly did not overlap with any exon based on the Human Gencode annotation V.27 were retained to ensure specificity of mapping to introns (Supplementary Table 2; ‘intron m6A’). MetaDiff scores for the resulting peak list were used to assess differential m\textsubscript{6}A deposition based on the cutoff of FDR < 0.05.

m\textsubscript{6}A exons spanning splice sites were selected from those identified by both the MetaDiff analysis on the transcribed genome that was just described and by MACS2. Among these peaks, those presenting sequencing reads that overlapped both an exon and an upstream or downstream intron were further selected (Supplementary Table 2; ‘splice-site spanning m6A’). Peaks accomplishing MetaDiff-calculated FDR < 0.05 and absolute fold-change > 1.5 (log, fold-change < −0.85) were used to create densities of RPKM-normalized reads inside exons and in the ±500 bp surrounding introns. Biological replicates were merged and depicted on 10 bp-binned heatmaps for visualization purposes. To study the composition of m\textsubscript{6}A peaks inside each transcriptional unit, the exonic peak with the greatest downregulated MetaDiff fold-change was compared to the mean fold-change of the rest of the m\textsubscript{6}A peaks found within the gene (both on exons and on introns). The resulting correlation was significant (\(r < 2 \times 10^{-16}\), adjusted \(r^2 = 0.2221\))

**RNA sequencing.** Poly-A purified opposing-strand-specific miRNA libraries were prepared from 200 ng of total RNA using the TruSeq Stranded mRNA HT sample preparation kit (illumina). Samples were individually indexed for pooling using a dual-index strategy. Libraries were quantified both with a Qubit (ThermoFisher) and using the Qubit RNA Quantitation Kit (Invitrogen). Libraries were then normalized and pooled. Pooled libraries were diluted and denatured for sequencing on the NextSeq 500 (illumina) according to the manufacturer’s instructions. Samples were pooled so as to obtain >30 million unique clusters per sample. (18 samples were split in two runs and multiplexed across four lanes per run). The phiX control library was diluted using the MB.2D function with all parameters set in each figure or figure legend. In cases where multiple control samples were included in each experiment (no actinomycin D treatment). The data were then fitted to a one-cycle threshold (\(\Delta\Delta C_t\)) approach\textsuperscript{69} using RPLPO as housekeeping gene. The reference sample used as control to calculate the relative gene expression is indicated in each figure or figure legend. In cases where multiple control samples were used as reference, the average \(\Delta C_t\) from all controls was used when calculating the \(\Delta\Delta C_t\). All primers were designed using PrimerBlaster (http://www.ncbi.nlm.nih.gov/tools/primer-blast/), and were validated to have a qPCR efficiency > 98% and to produce a single PCR product.

**mRNA-stability measurements.** RNA stability was measured by collecting RNA samples at different time points following transcriptional inhibition with 1 μg/ml actinomycin D (Sigma-Aldrich). Following qPCR analyses using equal amounts of mRNA, gene expression was expressed as relative to the beginning of the experiment (no actinomycin D treatment). The data were then fitted to a one-phase decay-regression model\textsuperscript{57}, and statistical differences in mRNA half-life were evaluated by comparing the model fits by extra sum-of-squares test.

**Western blots.** Samples were prepared by adding Laemmli buffer (final concentrations: 30 mM Tris–HCl pH 6.8, 6% glycerol, 2% SDS, 0.02% bromophenol blue and 0.25% β-mercaptoethanol), and were denatured at 95 °C for 5 min. Proteins were loaded and run on 4–12% NuPAGE Bis-Tris Precast Gels (Invitrogen), then transferred to polyvinylidene fluoride (PVDF) membranes by liquid transfer using NuPAGE Transfer buffer (Invitrogen). Membranes were blocked for 1 h at room temperature in PBS, 0.05% Tween-20 (PBST) supplemented with 4% non-fat dried milk, and incubated overnight at 4 °C with primary antibody diluted in the same blocking buffer (Supplementary Table 6). After three washes in PBST, membranes were incubated for 1 h at room temperature with horseradish peroxidase (HRP)-conjugated secondary antibodies diluted in blocking buffer (Supplementary Table 6), then washed a further three times with PBST before incubation with Pierce ECL2 Western Blotting Substrate (Thermo) and exposure with X-Ray Super RX Films (Fujifilm).

**Immunofluorescence.** Cells were fixed for 20 min at 4 °C in PBS with 4% PFA, rinsed three times with PBS, and blocked and permeabilized for 30 min at room temperature using PBS with 10% donkey serum (Biorad) and 0.1% Triton X-100. Primary antibodies (Supplementary Table 6) were diluted in PBS with 1% donkey serum and 0.1% Triton X-100 and incubated overnight at 4 °C. This was followed by three washes with PBS and further incubation with AlexaFluor secondary antibodies (Supplementary Table 6) for 1 h at room temperature away from light. Cells were finally washed three times with PBS, and DAPI (Sigma-Aldrich) was added to the first wash to stain nuclei. Images were acquired using a LSM 700 confocal microscope (Leica).

**Flow cytometry.** Single-cell suspensions were prepared by incubation in cell–lysing buffer (Invitrogen) for 10 min at 37 °C followed by extensive pipetting. Cells were washed twice with PBS and fixed for 20 min at 4 °C with PBS, 4% PFA. After three washes with PBS, cells were first permeabilized for 20 min at room temperature with PBS, 0.1% Triton X-100, then blocked for 30 min at room temperature with PBS containing 10% donkey serum. Primary and secondary antibody incubations (Supplementary Table 6) were performed for 1 h each at...
room temperature in PBS, 1% donkey serum, 0.1% Triton X-100, and cells were washed three times with this same buffer after each incubation. Flow cytometry was performed using a Cyan ADP flow cytometer, and at least 10,000 events were recorded. Data analysis was performed using FlowJo X.

**Statistics and reproducibility.** Unless described otherwise in a specific section of the Methods, standard statistical analyses were performed using GraphPad Prism 7 using default parameters. The type and number of replicates, the statistical test used, and the test results are described in the figure legends. The level of significance in all graphs is represented as follows: *P* < 0.05, **P** < 0.01 and ***P*** < 0.001. Test assumptions (for example, normal distribution) were confirmed where appropriate. For analyses with n < 10, individual data points are shown, and the mean ± s.e.m. is reported for all analyses with n > 2. The mean is reported when n = 2, and no other statistics were calculated for these experiments owing to the small sample size. No experimental samples were excluded from the statistical analyses. Sample size was not pre-determined through power calculations, and no randomization or investigator blinding approaches were implemented during the experiments and data analyses. When representative results are presented, the experiments were reproduced in at least two independent cultures, and the exact number of such replications is detailed in the figure legend.

**Code availability.** Custom bioinformatics scripts used to analyse the data presented in the study have been deposited in GitHub (http://github.com/pmb59/neMeRIP-seq).

**Data availability.** The mass-spectrometry proteomics data that support the findings of this study have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the identifier PXD005285. Nucleotide sequencing data that support the findings of this study have been deposited to Array Express with identifiers E-MTAB-5229 and E-MTAB-5230. Source Data for the graphical representations found in all figures and Extended Data figures are provided in the Supplementary Information of this manuscript. Electrophoretic gel Source Data (uncropped scans with size marker indications) are presented in Supplementary Fig. 1. Supplementary Tables 1 to 4 provide the results of the proteomics mass-spectrometry experiments analyzed in this study. The mass-spectrometry proteomics data that support the findings of this study have been deposited in the ProteomeXchange Consortium (identifiers PXD005285). The mass-spectrometry proteomics data that support the findings of this study have been deposited to the ProteomeXchange Consortium (identifiers PXD005285). The mass-spectrometry proteomics data that support the findings of this study have been deposited to the ProteomeXchange Consortium (identifiers PXD005285).

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Extended Data Figure 1 | See next page for caption.
Extended Data Figure 1 | Optimized co-immunoprecipitation protocol to define the SMAD2/3 interactome in hPSCs and early endoderm cells.

a. Western blots of SMAD2/3 or control (IgG) immunoprecipitations from nuclear extracts of hESCs following the co-IP1 or co-IP2 protocols. Input is 5% of the material used for immunoprecipitations. Results are representative of two independent experiments. For gel Source Data, see Supplementary Fig. 1.

b. Scatter plots of the log2 ratios of label-free quantification (LFQ) intensities for proteins identified by quantitative mass spectrometry in SMAD2/3 co-immunoprecipitations compared with IgG negative control co-immunoprecipitations. The experiments were performed from nuclear extracts of hESCs. The SMAD2/3 and IgG negative control co-immunoprecipitations were differentially labelled after immunoprecipitation using the dimethyl method, followed by a combined run of the two samples in order to compare the abundance of specific peptides and identify enriched peptides. The values for technical dye-swap duplicates are plotted on different axes, and proteins whose enrichment was significant (significance $B < 0.01$) are shown in black and named. As a result of this comparison between the two co-immunoprecipitation protocols, co-IP2 was selected for further experiments (see Supplementary Discussion).

c. Volcano plots of statistical significance against fold-change for proteins identified by label-free quantitative mass spectrometry in SMAD2/3 or IgG negative control immunoprecipitations in pluripotent hESCs or early endoderm (see Fig. 1a). The black lines indicate the threshold used to determine specific SMAD2/3 interactors, which are located to the right ($n = 3$ co-immunoprecipitations; one-tailed $t$-test: permutation-based FDR < 0.05).

d. Selected results of the analysis described in c for SMAD2, SMAD3 and selected known bona fide SMAD2/3-binding partners (full results can be found in Supplementary Table 1).

e. Mean label-free quantification (LFQ) intensity log2 ratios in endoderm (endo) and pluripotent cells (pluri) for all SMAD2/3 interactors. Differentially enriched proteins are shown as green and blue bars.

f. Selected results from gene ontology (GO) enrichment analysis, and enrichment analysis for mouse phenotypes annotated in the MGI database. All putative SMAD2/3-interacting proteins were considered for this analysis ($n = 89$ proteins; Fisher’s exact test followed by Benjamini-Hochberg correction for multiple comparisons). For each term, its rank in the analysis, the adjusted $P$ value, and the number of associated genes are reported.
Extended Data Figure 2 | See next page for caption.
Extended Data Figure 2 | Functional characterization of transcriptional and epigenetic cofactors of SMAD2/3 in hPSCs. a, Western blots of SMAD2/3 or control (IgG) immunoprecipitations from nuclear extracts of pluripotent hESCs (pluri) or hESCs differentiated into endoderm for 36 h (endo). Input is 5% of the material used for immunoprecipitations. Results are representative of two independent experiments. b, Schematic of the experimental approach for the generation of iKD hESC lines for SMAD2/3 cofactors. c, qPCR screening of iKD hESCs cultured in the absence (CTR) or presence (TET) of tetracycline for three days. Three distinct shRNAs were tested for each gene. Expression is normalized to the mean level in hESCs carrying negative control shRNAs (scrambled (SCR) or against B2M) and cultured in the absence of tetracycline. The mean is indicated, n = 2 independent clonal pools. Note than for the B2M shRNA only the scrambled shRNA was used as negative control. shRNAs selected for further experiments are circled. d, Phase-contrast images of iKD hESCs expressing the indicated shRNAs (sh) and cultured in the presence of tetracycline for six days to induce knockdown. Scale bars, 400 μm. Results are representative of two independent experiments. e, Immunofluorescence for the pluripotency factor NANOG in iKD hESCs for the indicated genes cultured in the absence (CTR) or presence of tetracycline (TET) for six days. Scale bars, 400 μm. Results are representative of two independent experiments. f, Heat map summarizing qPCR analyses of iKD hESCs cultured as in e. log2 fold-changes (FC) are compared to scrambled control (n = 2 clonal pools). Germ-layer markers are grouped in boxes: green, endoderm; red, mesoderm; blue, neuroectoderm.
Extended Data Figure 3 | Functional characterization of transcriptional and epigenetic cofactors of SMAD2/3 during endoderm differentiation.

a, qPCR validation of iKD hESCs in pluripotent cells (PLURI) or following endoderm differentiation (ENDO). Pluripotent cells were cultured in the absence (CTR) or presence (TET) of tetracycline for six days. For endoderm differentiation, tetracycline treatment was initiated in undifferentiated hESCs for three days in order to ensure gene knockdown at the start of endoderm specification, and was then maintained during differentiation (three days). For each gene, the shRNA resulting in the strongest level of knockdown in hPSCs was selected (refer to Extended Data Fig. 2). Expression is normalized to the mean level in pluripotent hESCs carrying scrambled control shRNA and cultured in the absence of tetracycline. The mean is indicated, \( n = 2 \) independent clonal pools.

b, Immunofluorescence of the endoderm marker SOX17 following endoderm differentiation of iKD hESCs expressing the indicated shRNAs and cultured as described in a. Scale bars, 400 \( \mu \)m. Results are representative of two independent experiments.

c, qPCR following endoderm differentiation of iKD hESCs. The mean is indicated, \( n = 2 \) independent clonal pools.

d, Table summarizing the phenotypic results presented in Extended Data Fig. 2 and in this figure. E, endoderm; N, neuroectoderm; M, mesoderm.
Extended Data Figure 4 | Mechanistic insights into the functional interaction between SMAD2/3 and the m^6^A methyltransferase complex.

a–c, Western blots of SMAD2/3, METTL3, METTL14 or control (IgG) immunoprecipitations from nuclear extracts of hESCs (a, c) or hiPSCs (b). Input is 5% of the material used for immunoprecipitations. In c, immunoprecipitations were performed from hPSCs maintained in the presence of activin or treated for 1 h with the activin-NODAL signalling inhibitor SB431542. Results are representative of three (a) or two (b, c) independent experiments.

d, qPCR validation of hESCs constitutively overexpressing NANOG (NANOG OE) following gene targeting of the AAVS1 locus with pAAV-Puro_CAG-NANOG. Parental wild-type H9 hESCs (H9) were analysed as negative controls. Cells were cultured in the presence of activin or treated with SB431542 for the indicated times. The mean is shown, n = 2 cultures. NANOG-overexpressing cells are resistant to downregulation of NANOG following inhibition of activin–NODAL signalling.

e, RNA immunoprecipitation experiments for WTAP, SMAD2/3 or control (IgG) in NANOG-overexpressing hESCs maintained in the presence of activin or treated for 2 h with SB431542. Enrichment of the indicated transcripts was measured by qPCR and expressed relative to background levels observed in control IgG RNA immunoprecipitations in the presence of activin. RPLP0 was tested as a negative control transcript. Mean ± s.e.m., n = 3 cultures. Significance of differences from activin (left) or IgG (right) RIP was tested by two-way ANOVA with post hoc Holm–Sidak comparisons.

f, ChIP–qPCR in hESCs for ChIP against the indicated proteins or the negative control ChIP (IgG). qPCR was performed for validated genomic SMAD2/3-binding sites associated with the indicated genes. hESCs were cultured in the presence of activin or treated for 2 h with SB431542. Enrichment is normalized against background binding observed with IgG ChIP. The mean is shown, n = 2 technical replicates. Results are representative of three independent experiments.

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Extended Data Figure 5 | See next page for caption.
Extended Data Figure 5 | Monitoring changes in m\textsuperscript{6}A deposition that are rapidly induced by inhibition of activin–NODAL signalling.

\textit{a, b,} MeRIP–qPCR results from purified mRNA, total cellular RNA or cellular RNA species separated by nuclear and cytoplasmic subcellular fractionation. hESCs were cultured in pluripotency-maintaining conditions containing activin or in conditions in which activin–NODAL signalling was inhibited for 2 h with SB431542. IgG MeRIP experiments were performed as negative controls. The mean is indicated, \( n = 2 \) technical replicates. Differences between activin- and SB431542-treated cells were observed only in the nuclear-enriched fraction. Therefore, the nuclear-enriched MeRIP protocol (NeMeRIP) was used for subsequent experiments (refer to the Supplementary Discussion). Results are representative of two independent experiments. 

\textit{c,} Overlap with the indicated genomic features of m\textsuperscript{6}A peaks identified by NeMeRIP–seq using two different bioinformatics pipelines in which peak calling was performed using MetDiff or MACS2. For each pipeline, the analyses were performed on the union of peaks identified from data obtained in hESCs cultured in the presence of activin or with inhibition of activin–NODAL signalling for 2 h with SB431542 (\( n = 3 \) cultures). Note that the sum of the percentages within each graph is not 100% because some m\textsuperscript{6}A peaks overlap several feature types. MetDiff is an exome peak caller, and, accordingly, 100% of peaks map to exons. MACS2 identifies peaks throughout the genome. 

\textit{d,} Venn diagrams showing the overlap of peaks identified by the two pipelines. Only MetDiff peaks that were also identified by MACS2 were considered for subsequent analyses focused on m\textsuperscript{6}A peaks on exons. 

\textit{e,} Top sequence motifs identified de novo on all m\textsuperscript{6}A exon peaks, or on those that showed significant downregulation following inhibition of activin–NODAL signalling (activin–NODAL-sensitive m\textsuperscript{6}A peaks; Supplementary Table 2). The position of the methylated adenosine is indicated by a box. 

\textit{f,} Coverage profiles for all m\textsuperscript{6}A exon peaks across the length of different genomic features. Each feature type is expressed as 100 bins of equal length with 5’ to 3’ directionality. 

\textit{g, h,} Overlap of m\textsuperscript{6}A exon peaks and transcription start sites (TSS) or transcription end sites (TES). In \( g \), the analysis was performed for all m\textsuperscript{6}A peaks. In \( h \), only activin–NODAL-sensitive peaks were considered. 

\textit{i,} Left, activin–NODAL-sensitive m\textsuperscript{6}A exon peaks were evaluated for direct overlap with SMAD2/3-binding sites as indicated by ChIP–seq\textsuperscript{50}. \( n = 482 \) peaks; FDR = 0.41 as calculated by the permutation test implemented by the GAT python package; N.S., not significant based on 95% confidence interval. Right, overlap was calculated after the same features were mapped to their corresponding transcripts or genes, respectively. \( n = 372 \) genes; hypergeometric test \( P = 2.88 \times 10^{-18} \), significant based on 95% confidence interval. 

\textit{j,} m\textsuperscript{6}A NeMeRIP–seq results for selected transcripts (\( n = 3 \) cultures; replicates combined for visualization). Coverage tracks represent read enrichments normalized by million mapped reads and size of the library. Blue, sequencing results of m\textsuperscript{6}A NeMeRIP; orange, sequencing results of pre-NeMeRIP input RNA (negative control). GENCODE gene annotations are shown (red, protein coding exons; white, untranslated exons; all potential exons are shown and overlaid). The location of SMAD2/3 ChIP–seq-binding sites is also shown. Compared to the other genes shown, m\textsuperscript{6}A levels on SOX2 were unaffected by inhibition of activin–NODAL signalling, showing specificity of action. 

POU5F1 (also known as OCT4) is used as a negative control since it is known to not have a m\textsuperscript{6}A site\textsuperscript{23}, as confirmed by the lack of m\textsuperscript{6}A enrichment compared to the input.
Extended Data Figure 6 | See next page for caption.
Extended Data Figure 6 | Features of activin–NODAL-sensitive differential m\(^6\)A deposition. a, Scatter plot of the average log\(_2\) fold-change in SB431542 versus activin-treated hESCs for m\(^6\)A NeMeRIP–seq and pre-NeMeRIP input RNA \((n = 3\) cultures). The analysis was performed for all m\(^6\)A exon peaks (left), or for those peaks that were significantly downregulated following inhibition of activin–NODAL signalling (right). Data are colour coded according to the square of the difference between the two values (square diff.). b, c, As Extended Data Fig. 5j, but for representative transcripts that are stably expressed following inhibition of activin–NODAL signalling for 2 h \((n = 3\) cultures; replicates combined for visualization). The m\(^6\)A NeMeRIP and input tracks were separated and are shown at different scales to facilitate comparison between the conditions. The m\(^6\)A peaks and those significantly downregulated after 2 h of SB431542 treatment are indicated. d, Venn diagram illustrating the strategy for identification of m\(^6\)A peaks on introns. Peaks mapping to the transcribed genome were obtained by running MetDiff using an extended transcriptome annotation based on the pre-NeMeRIP input RNA, which has a high abundance of introns. The resulting peaks were first filtered by overlap with genome-wide MACS2-identified peaks, and then by lack of overlap with annotated exons. e, Results of MetDiff differential methylation analysis in activin versus 2 h SB431542 treatment for m\(^6\)A peaks on introns. \(n = 3\) cultures; \(P\) value calculated by likelihood ratio test implemented in the MetDiff R package, and adjusted to FDR by Benjamini–Hochberg correction. See Supplementary Table 2 for the FDR of individual peaks. f, As Extended Data Fig. 5j, but for a representative transcript that shows activin–NODAL-sensitive m\(^6\)A deposition in introns \((n = 3\) cultures; replicates combined for visualization). The m\(^6\)A peaks on exons, introns, and those significantly downregulated after SB431542 treatment within each subset are indicated. g, Plots of RPKM-normalized mean m\(^6\)A coverage for m\(^6\)A exon peaks significantly downregulated after SB431542 treatment (absolute fold-change > 1.5). Data for all such peaks is in blue, whereas green lines report coverage for only those peaks characterized by next generation sequencing reads that span exon–intron junctions. Exons were scaled proportionally, and the positions of the 3′ and 5′ splice sites (SS) are indicated. A window of 500 bp on either side of the splice sites is shown. m\(^6\)A, signal from m\(^6\)A NeMeRIP–seq; input, signal from pre-NeMeRIP input RNA. The results show that coverage of activin–NODAL-sensitive m\(^6\)A peaks often spans across splice sites (highlighted by the dotted lines). h, Heat map representing in an extended form the data shown in g for all activin–NODAL-sensitive m\(^6\)A exon peaks in hESCs cultured in the presence of activin. There are multiple regions in which sequencing coverage extends across exon–intron junctions (see Supplementary Table 2). i, Example of an activin–NODAL-sensitive peak located in the proximity of a 3′ splice site \((n = 3\) cultures; replicates combined for visualization). This peak is shown within its genomic context in c, where it is indicated by a dotted box. Top, m\(^6\)A NeMeRIP–seq coverage; bottom, individual next generation sequencing reads. Multiple reads span the exon–intron junction (indicated by the dashed line). j, Relationship between the decrease of m\(^6\)A on the most affected exonic peak located on a transcript \((y\) axis) and the mean change of all other peaks mapping to the same transcript \((x\) axis). The analysis considered transcripts with multiple m\(^6\)A peaks and with at least one peak significantly decreasing after inhibition of activin–NODAL signalling with SB431542 (absolute fold-change > 1.5). Sensitivity of m\(^6\)A deposition to activin–NODAL signalling across these transcripts is correlated.
Extended Data Figure 7 | Generation and functional characterization of hPSCs following iKD of the subunits of the m^6^A methyltransferase complex. a, qPCR validation of iKD hESCs cultured in the presence of tetracycline for five days (TET) to drive gene knockdown. Two distinct shRNAs and multiple clonal sublines (cl) were tested for each gene. Expression is normalized to the mean level in hESCs carrying a negative control scrambled (SCR) shRNA. For each gene, sh1 cl1 was selected for further analyses. The mean is indicated, n = 2 cultures. b, Western blot validation of selected iKD hESCs for the indicated genes. TUBA4A (α-tubulin), loading control. Results are representative of three independent experiments. c, MeRIP–qPCR in iKD hESCs cultured for ten days in the absence (CTR) or presence of tetracycline (TET). m^6^A abundance is shown relative to control conditions in the same hESC line. The mean is shown, n = 2 technical replicates. Results are representative of two independent experiments. d, m^6^A dot blot in WTAP or scramble shRNA control iKD hESCs treated as described in c. Decreasing amounts of mRNA were spotted to allow semiquantitative comparisons, as indicated. Results are representative of two independent experiments. e, Immunofluorescence of the pluripotency markers NANOG and OCT4 in iKD hESCs cultured for three passages (15 days) in the absence (CTR) or presence of tetracycline (TET). Scale bars, 100 μm. Results are representative of two independent experiments. f, Flow cytometry showing NANOG expression in cells treated as in e. The percentage and median fluorescence intensity (MFI) of NANOG-positive cells (NANOG^+) are shown. The gates used for the analysis are indicated, and were determined on the basis of a secondary-antibody-only negative staining (NEG). Results are representative of two independent experiments.
Extended Data Figure 8 | Function of the m^6^A methyltransferase complex during germ-layer specification. a, qPCR analysis following neuroectoderm or endoderm differentiation of iKD hESCs cultured in absence (CTR) or presence of tetracycline (TET). Tetracycline treatment was initiated in undifferentiated hESCs for ten days and was maintained during differentiation (three days). Expression was normalized against the mean level in undifferentiated hESCs. Mean ± s.e.m., n = 3 cultures. Significant differences versus the same iKD line in control conditions were calculated by two-way ANOVA with post hoc Holm–Sidak comparisons.
b, Flow cytometry quantification of the percentage of SOX1-positive cells (SOX1^+^) in cells treated as in a. Mean is shown, n = 2 cultures.
c, Immunofluorescence of the lineage marker SOX17 in endoderm-differentiated hESCs treated as in a. Scale bars, 100 μm. Results are representative of two independent experiments. d, qPCR validation of multiple inducible knockdown (MiKD) hESCs simultaneously expressing shRNAs against WTAP, METTL3 and METTL14. Cells expressing three copies of the scrambled shRNA (SCR3×) were used as negative control. Cells were cultured in the presence of tetracycline (TET) for five days to drive gene knockdown. Mean ± s.e.m., n = 3 cultures. Significant differences versus SCR3× hESCs in control conditions were calculated by two-way ANOVA with post hoc Holm–Sidak comparisons.
e, f, qPCR analysis following endoderm differentiation of WTAP, METTL3 and METTL14-MiKD hESCs treated as described in a. Mean ± s.e.m., n = 3 cultures. Significant differences versus control conditions were calculated by two tailed t-test (e) or two-way ANOVA with post hoc Holm–Sidak comparisons (f).
Extended Data Figure 9 | Function of the m<sup>6</sup>A methyltransferase complex during exit from pluripotency induced by inhibition of activin–NODAL signalling. a, qPCR analyses in iKD hESCs cultured in absence (CTR) or presence (TET) of tetracycline for ten days, then subjected to inhibition of activin–NODAL signalling with SB431542 (SB) for the indicated time (see Extended Data Fig. 10a). Activin, cells maintained in standard pluripotency-promoting culture conditions containing activin and collected at the beginning of the experiment. Mean ± s.e.m., n = 3 cultures. Significant differences versus same iKD line in control conditions were calculated by two-way ANOVA with post hoc Holm–Sidak comparisons. b, Western blots of cells treated as described in a. TUBA4A, loading control. Results are representative of two independent experiments. c, Measurement of mRNA stability in WTAP iKD hESCs cultured in absence (CTR) or presence (TET) of tetracycline for ten days. Samples were collected following transcriptional inhibition using actinomycin D (ActD) for the indicated time. The statistical significance of differences between the mRNA half lives in tetracycline versus control is shown (n = 3 cultures, comparison of fits to one-phase decay model by extra sum-of-squares F-test). The difference was significant for NANO7 but not for SOX2 (95% confidence interval). d, Model showing the interplays between activin–NODAL signalling and m<sup>6</sup>A deposition in hPSCs (left), and the phenotype induced by impairment of the m<sup>6</sup>A methyltransferase complex (right).
Extended Data Figure 10 | See next page for caption.
Extended Data Figure 10 | Genome-wide analysis of the relationship between WTAP and activin–NODAL signalling. 

a, Schematic of the experimental approach to investigate the transcriptional changes induced by the knockdown of the m^6^A methyltransferase complex subunits during neuroectoderm specification of hESCs. b, qPCR analyses of WTAP iKD hESCs subjected to the experiment outlined in a (n = 3 cultures). Activin, cells maintained in standard pluripotency-promoting culture conditions containing activin and collected at the beginning of the experiment. Z-scores indicate differential expression measured in number of standard deviations from the mean across all time points. c, RNA-seq analysis at selected time points from the samples shown in panel b (n = 3 cultures). The heat map shows Z-scores for the top 5% differentially expressed genes (1789 genes as ranked by the Hotelling T^2^ statistic). Genes and samples were clustered based on their Euclidean distance, and the four major gene clusters are indicated (see Supplementary Discussion). d, Expression profiles of genes belonging to the clusters indicated in c. Selected results of gene-enrichment analysis and representative genes for each cluster are shown (cluster 1: n = 456 genes; cluster 2: n = 471 genes; cluster 3: n = 442 genes; cluster 4: n = 392 genes; Fisher’s exact test followed by Benjamini–Hochberg correction for multiple comparisons). e, Principal component analysis of RNA-seq results in c (n = 3 cultures). The top 5% differentially expressed genes were considered for this analysis. For each of the two main principal components (PC1 and PC2), the fraction of inter-sample variance that they explain and their proposed biological meaning are reported. f, Proportion of transcripts marked by at least one high-confidence m^6^A peak^23^ in transcripts significantly up- or downregulated following WTAP iKD in hESCs maintained in the presence of activin (left), or following inhibition of activin–NODAL signalling for 2 h with SB431542 in control cells (right). Differential gene expression was calculated in three cultures using the negative binomial test implemented in DEseq2 with a cutoff of P < 0.05 and abs.FC > 2. The number of genes in each group and the hypergeometric probabilities of the observed overlaps with m^6^A-marked transcripts are reported (n.s.: not significant at 95% confidence interval).
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1. Sample size
   Describe how sample size was determined.
   No power calculations were performed. Sample size was determined depending on the experiment type based on what is standard practice in the field of pluripotent stem cell biology to statically examine a large effect within an in vitro system which experiences only limited biological variability (n=2-4, see Figure Legends).

2. Data exclusions
   Describe any data exclusions.
   No data points were excluded from any of the analyses presented.

3. Replication
   Describe whether the experimental findings were reliably reproduced.
   All of the presented experiments were successfully reproduced.

4. Randomization
   Describe how samples/organisms/participants were allocated into experimental groups.
   No randomization was performed since this was not relevant to the study: all treatment/control experiments were performed on the same starting cell population (no covariates).

5. Blinding
   Describe whether the investigators were blinded to group allocation during data collection and/or analysis.
   No blinding was performed. This was deemed unnecessary since none of the analyses reported involved procedures that could be influenced by investigator bias (such as manual counting/measuring and/or morphological assessments). Indeed, all analyses presented involved automated processing of data through experimental instrumentation and/or computing procedures (including counting of PLA signals presented in Fig. 2d, see the Methods).

Note: all studies involving animals and/or human research participants must disclose whether blinding and randomization were used.
6. Statistical parameters

For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or in the Methods section if additional space is needed).

- **n/a**
  - For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or in the Methods section if additional space is needed).
  - The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement (animals, litters, cultures, etc.).
  - A description of how samples were collected, noting whether measurements were taken from distinct samples or whether the same sample was measured repeatedly.
  - A statement indicating how many times each experiment was replicated.
  - The statistical test(s) used and whether they are one- or two-sided (note: only common tests should be described solely by name; more complex techniques should be described in the Methods section).
  - A description of any assumptions or corrections, such as an adjustment for multiple comparisons.
  - The test results (e.g., P values) given as exact values whenever possible and with confidence intervals noted.
  - A clear description of statistics including central tendency (e.g., median, mean) and variation (e.g., standard deviation, interquartile range).
  - Clearly defined error bars.

*See the web collection on statistics for biologists for further resources and guidance.*

7. Software

**Policy information about availability of computer code**

Describe the software used to analyze the data in this study.

All of the software used for the analyses presented is described in detail in the relevant Methods, which also describes the relevant parameters used when these were not the default. The software used was: MaxQuant, Perseus, Cytoscape, Enrichr, ImageJ, R/Bioconductor, Trimomatic, TopHat 2.0.13, MetDiff, FCCAC v1.0.0, RSeqC-2.6, RCAS, DREME, DESeq2, Cufflinks, GraphPad Prism 6, Sickle, Samtools view, SeqMonk, timecourse (Bioconductor). All bioinformatic scripts have been deposited to GitHub (http://github.com/pmb59/neMeRIP-seq).

For manuscripts utilizing custom algorithms or software that are central to the paper but not yet described in the published literature, software must be made available to editors and reviewers upon request. We strongly encourage code deposition in a community repository (e.g., GitHub). *Nature Methods* guidance for providing algorithms and software for publication provides further information on this topic.

8. Materials availability

**Policy information about availability of materials**

Indicate whether there are restrictions on availability of unique materials or if these materials are only available for distribution by a for-profit company.

All unique materials (cell lines and plasmids) are readily available from the authors.

9. Antibodies

Describe the antibodies used and how they were validated for use in the system under study (i.e., assay and species).

All of the antibodies used are detailed in Supplementary Table 5, which also reports the application for which they were used and the amount/dilution. The list includes: anti-CREBBP (Cell Signalling 7389); anti-EP300 (Santa Cruz sc-584); anti-FOXH1 (R&D BAF4248); anti-METTL14 (Sigma-Aldrich HPA38002); anti-METTL3 (Proteintech 15073-1-AP, and Bethyl Lab A301-567A); anti-NANOG (R&D AF1997, and Abcam ab21624); anti-OCX4/POUSIF1 (Santa Cruz sc-5279); anti-SETDB1 (Cell Signalling 2196); anti-SMAD2/3 (R&D AF3797, and Cell Signalling 12470S); anti-SNON/SKIL (Santa Cruz sc-9595x); anti-SOX1 (R&D AF3369); anti-SOX17 (R&D AF1294); anti-TUB4A4 (Sigma-Aldrich T6199); anti-WTAP (Bethyl Lab A301-436A). All of the antibodies were validated to recognize the relevant human protein, and most of them were specifically validated for the relevant application, as specified on the relevant catalog pages on the suppliers’ websites. In the few cases in which the antibody was not previously validated for a specific application, extensive testing in house with the appropriate negative controls was performed to confirm the specificity (see Methods and Figure legends). Such applications were: anti-METTL14 (IP and ChIP); anti-METTL3 (ChIP); anti-NANOG (flow cytometry and PLA); anti-SMAD2/3 (RIP and PLA); anti-SOX1 (flow-cytometry); anti-WTAP (PLA, RIP, and ChIP).
10. Eukaryotic cell lines
   a. State the source of each eukaryotic cell line used. The H9 hESC line was obtained from WiCell (Madison, Wisconsin). The A1ATR/R hiPSC line was obtained in house and previously described in Yusa et al 2011. Yusa, K. et al. Targeted gene correction of alpha1-antitrypsin deficiency in induced pluripotent stem cells. Nature 478, 391-4 (2011).
   b. Describe the method of cell line authentication used. No authentication was performed on the H9 hESCs as they were used directly from the commercial supplier. The A1ATR/R was genotyped in house to confirm the presence of A1AT R/R allele. Both cell lines were routinely karyotyped by standard G-banding to confirm euploidy.
   c. Report whether the cell lines were tested for mycoplasma contamination. Yes, mycoplasma screening was performed every month.
   d. If any of the cell lines used are listed in the database of commonly misidentified cell lines maintained by iCLAC, provide a scientific rationale for their use. No commonly misidentified cell lines were used.

Animals and human research participants

Policy information about studies involving animals; when reporting animal research, follow the ARRIVE guidelines

11. Description of research animals
    Provide details on animals and/or animal-derived materials used in the study. 
    No animals were used

Policy information about studies involving human research participants

12. Description of human research participants
    Describe the covariate-relevant population characteristics of the human research participants.
    The study did not involve human research participants.
Flow Cytometry Reporting Summary

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▶ Data presentation

For all flow cytometry data, confirm that:

1. The axis labels state the marker and fluorochrome used (e.g. CD4-FITC).
2. The axis scales are clearly visible. Include numbers along axes only for bottom left plot of group (a 'group' is an analysis of identical markers).
3. All plots are contour plots with outliers or pseudocolor plots.
4. A numerical value for number of cells or percentage (with statistics) is provided.

▶ Methodological details

5. Describe the sample preparation. Biological samples: H9 hESCs (Extended Data Figure 7f), or hESC-derived neuroectoderm (Extended Data Figure 8b). Single cell suspensions were prepared by incubation in cell dissociation buffer (CDB; Gibco) for 10’ at 37° followed by extensive pipetting. Cells were washed twice with PBS and fixed for 20’ at 4°C with PBS 4% PFA. After three washes with PBS, cells were first permeabilized for 20’ at RT with PBS 0.1% Triton X-100, then blocked for 30’ at RT with PBS 10% donkey serum. Primary and secondary antibodies incubations (Supplementary Table 5) were performed for 1h each at RT in PBS 1% donkey serum 0.1% Triton X-100, and cells were washed three times with this same buffer after each incubation.

6. Identify the instrument used for data collection. Cyan ADP flow cytometer (Beckman Coulter)

7. Describe the software used to collect and analyze the flow cytometry data. Data collection: Summit software (Beckman Coulter). Data analysis: FlowJo X

8. Describe the abundance of the relevant cell populations within post-sort fractions. No cell sorting was performed in the study

9. Describe the gating strategy used. Cells were first gated on the basis of forward and side scatter properties, after which singlets were isolated on the basis of relationship between side scatter area peak area and width. A secondary-only negative control was used to determine the background fluorescence, and positive cells were quantified by setting a boundary so that less than 1% of the secondary-only control cells would be considered positive.

Tick this box to confirm that a figure exemplifying the gating strategy is provided in the Supplementary Information. ☑