DNA and Chromatin Modification Networks Distinguish Stem Cell Pluripotent Ground States*

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Pluripotent stem cells are capable of differentiating into all cell types of the body and therefore hold tremendous promise for regenerative medicine. Despite their widespread use in laboratories across the world, a detailed understanding of the molecular mechanisms that regulate the pluripotent state is currently lacking. Mouse embryonic (mESC) and epiblast (mEpiSC) stem cells are two closely related classes of pluripotent stem cells, derived from distinct embryonic tissues. Although both mESC and mEpiSC are pluripotent, these cell types show important differences in their properties suggesting distinct pluripotent ground states. To understand the molecular basis of pluripotency, we analyzed the nuclear proteomes of mESCs and mEpiSCs to identify protein networks that regulate their respective pluripotent states. Our study used label-free LC-MS/MS to identify and quantify 1597 proteins in embryonic and epiblast stem cell nuclei. Immunoblotting of a selected protein subset was used to confirm that key components of chromatin regulatory networks are differentially expressed in mESCs and mEpiSCs. Specifically, we identify differential expression of DNA methylation, ATP-dependent chromatin remodeling and nucleosome remodeling networks in mESC and mEpiSC nuclei. This study is the first comparative study of protein networks in cells representing the two distinct, pluripotent ground states, and points to the importance of DNA and chromatin modification processes in regulating pluripotency. In addition, by integrating our data with existing pluripotency networks, we provide detailed maps of protein networks that regulate pluripotency as well as efforts to reliably control the differentiation of these cells into functional cell fates.

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Pluripotency, the ability to give rise to all cell types within the body, is a unique cellular property restricted to very early embryogenesis. Pluripotent stem cells have been derived from early embryos and represent a powerful system for understanding mammalian development as well as providing a scalable source of cells for regenerative medicine (1–7). Significant effort has been directed to understanding how the pluripotent state is specified in stem cells, and transcriptional regulatory mechanisms are now known to be central to maintenance of the pluripotent state. Embryonic stem cells are able to self-renew and yet remain in a poised state, ready to differentiate as required (8). In addition, pluripotent stem cells may be derived from different embryonic tissues, such as the epiblast (6), with distinct properties and responses to cellular signals suggesting that there are different pluripotent ground states.

Uncovering the molecular basis of stem cell pluripotency has been the principal goal of many ‘omics studies. Protein expression analysis has been used to globally define proteins specific to embryonic stem cells (9, 10). In addition, targeted proteomics using affinity-purification methods has been used to define protein complexes important for maintenance of pluripotency (11, 12). Genome-wide profiling of DNA-binding sites and gene-expression analyses have been used to study stem cells and their differentiated derivatives and to uncover the molecular networks that regulate the pluripotent state (13). Meta-analyses of multiple data sets or aggregation of large-scale data has enabled the definition of pluripotency networks (14, 15). The latter study integrated gene-expression profiles from multiple different embryonic stem cell lines to construct a core pluripotency gene-network named PluriNet (15). A key conclusion from this and other studies is the existence of a defined network common to multiple different embryonic stem cells lines that differentiates these cells from nonpluripotent cells and functions to regulate the pluripotent state.

Processes that regulate chromatin assembly and conformation are major determinants of the required activation and repression of pluripotency gene-expression. Dynamic changes to chromatin occur as part of many developmental transitions in mammals, and also appear to regulate the maintenance of pluripotency. Processes such as DNA methylation and ATP-dependent chromatin-remodeling facilitate these developmental transitions through regulation and assembly of...
The SWI/SNF complexes (also known as Brg associated factors, BAF), first described in yeast, have been shown to be important chromatin remodeling components in many other eukaryotes (16). Detailed maps of BAF complexes have shown that there are both core components and complex configurations that occur in specific cell types. In embryonic stem cells, the esBAF complex has been defined and shown to differ in its subunit composition from differentiated derivatives (17). The important role of BAF components such as Smarca4/Brg1 and Smarcc1/Baf155 in pluripotency has been shown by their role in direct repression of key pluripotency-regulating genes during differentiation of embryonic stem cells (18, 19).

Although many studies have focused on understanding the pluripotent state in mouse embryonic stem cells, it is now appreciated that distinct pluripotent ground states exist (mESC-like and mEpiSC-like) and it is important to decipher how pluripotency is regulated in these distinct states. These comparisons are especially pertinent because human embryonic stem cells share molecular properties and defining features with the epiblast state (6, 20, 21). Our study used label-free proteomics to quantitatively compare the nuclear proteomes of mouse embryonic stem cells (mESCs) and mouse epiblast stem cells (mEpiSCs). We show that proteomics approaches add significantly to the growing understanding of regulatory networks underlying pluripotency. We identify specific differences in expression of components of DNA methylation and chromatin remodeling protein networks, indicating that these processes differ between mESCs and mEpiSCs and may dictate the different pluripotent states of mESCs and mEpiSCs. This study provides the first global comparison of the mESC and mEpiSC proteomes, and will provide a foundation for manipulation and future functional studies of mESC and mEpiSC pluripotency.

EXPERIMENTAL PROCEDURES

Cell Culture, Cell Lysis, and Nuclear Extraction—mESCs and mEpiSCs were grown as described previously (6, 22). Whole colonies were lifted from the plates using 1.5 mg/ml (w/v) collagenase IV (Invitrogen). Colonies were pelleted at 200g for 5 min and washed twice with ice-cold PBS buffer (137 mM NaCl, 2.7 mM KCl, 100 mM Na2HPO4 and 2 mM KH2PO4) and either processed immediately or frozen at −80 °C until use. Cell lysis buffer I (50 mM Tris-HCl, pH 7.5, 1 mM EDTA, 150 mM NaCl, 1% Triton-100, Protease inhibitor mixture) was added to the cell pellets to suspend the cells. The cell suspension was kept on ice for 30 min before low speed centrifugation at 500g for 5 min in 4 °C. The pellets were resuspended in 100 µl 0.1% formic acid and further cleaned up by reverse phase chromatography using C18 column (Harvard, Southborough, MA). The final volume was reduced to 10 µl by vacuum centrifugation and adding 0.1% formic acid.

Tryptic peptides were separated by on-line reverse phase nanoscale capillary liquid chromatography (nano-LC, Dionex Ultimate 3000 series HPLC system) coupled to electrospray injection (ESI) tandem mass spectrometer (MS-MS) with octopole collision cell (Thermo Fisher Scientific LTQ Orbitrap XL). Loaded peptides were trapped in a C18 trap column and eluted on nano-LC with 90-min gradients ranging from 6 to 66% Acetonitrile in 0.1% formic acid with a flow rate 300 nl/min. Data dependent acquisition was performed on the LTQ-Orbitrap using Xcalibur software (version 2.0.6, Thermo Scientific) in the positive ion mode with a full scan MS1 at resolution of 60,000 in the m/z range 325.0 to 1800.0 followed by collision-induced dissociation (CID) fragmentation of the top five precursor ions using 35% normalized collision energy. Appropriate dynamic exclusion settings were based on a nominal LC peak width of 45 s. Adequate LC re-equilibration time was factored into the LC-MS/MS method.

In-solution Based Approach and LC-MS/MS—Nuclear extracts (three replicates for each cell type) were added with cold acetone for protein precipitation overnight such that 40 µg of total protein was isolated from each sample. Solubilization of all the proteins was performed with the addition of 4% SDS in Tris 50 mM buffer, pH 8.0. Processing of the solubilized samples for in-solution digest was done using the FASP approach (24). Briefly, a 3K cutoff filter (Millipore Inc, Billerica, MA) was used as a proteomic reactor for detergent cleaning, cysteine reduction and alkylation prior to tryptic digestion. The digests prepared above were analyzed by LC-MS/MS system using a Waters nano acquity UPLC systems (Waters Inc, MA) that was interfaced to a LTQ Velos-Orbitrap mass spectrometer (Thermo-Finnigan, Bremen, Germany). The platform was operated in the nano-LC mode using the standard nano-ESI API stack fitted with a picotip emitter (uncoated fitting, 10 µm spray orifice, New Objective, Inc., Woburn, MA). The solvent flowrate through the column was maintained at 300 nl/min using the split-free acquisition system. The protein digests (5 µl) were injected into a reversed-phase column C18 trapping column (0.18 × 20 mm, 5 µm particle size, Waters Inc.) equilibrated with 0.1% formic acid/2% acetonitrile (v/v) and washed for 5 min with the equilibration solvent at a flow rate of 15 µl/min, using the sample trap mode of UPLC. After the washing step, the trapping column was switched in-line with a reversed-phase bridged ethyl hybrid (BEH) C18 nanoacquity UPLC column (0.075 × 250 mm, Waters Inc.) and the peptides were chromatographed using a linear gradient of acetonitrile from 5% to 50% in aqueous 0.1% formic acid over a period of 210 min at the above-mentioned flow rate such that the eluate was directly introduced to the mass spectrometer. A 100% acetonitrile elution step was subsequently performed for 15 min prior to resetting the analytical column to the initial equilibration conditions for 15 more
minutes at the end of the chromatographic run, accounting for a total of 240 min/h of LC-MS/MS time. The mass spectrometer was operated in a data-dependent MS to MS/MS switching mode, with the 12 most intense ions in each MS scan subjected to MS/MS analysis. The full scan was performed at 60000 resolution in the Orbitrap detector and the MS/MS fragmentation scans were performed in the Velos dual ion trap detector (IT) CID mode such that the total scan cycle frequency was ~1.5 s. The threshold intensity for the MS/MS trigger was always set at 1000 and the fragmentation was carried out using the CID mode using a normalized collision energy (NCE) of 35. The data was entirely collected in the profile mode for the full scan and centroid mode for the MS/MS scans. Dynamic exclusion function for previously selected precursor ions was enabled during the analysis such that the following parameters were applied: repeat count of 2, repeat duration of 45 s, exclusion duration of 60 s and exclusion size list of 450. Xcalibur software (version 2.0.7), Thermo-Finnigan Inc., San Jose, CA was used for instrument control, data acquisition, and data processing.

Chromatographic reproducibility and full scan MS (also termed MS 1) intensity robustness are critical to the success of this label-free method. To monitor this efficiently and exhaustively, spiked external peptides from yeast enolase digest (400 fmols on column-load) was used to keep a track of the chromatographic performance of the LC-MS/MS system.

Data Processing—Raw MS data were processed using Mascot search engine (version 2.2.0; Matrix Science, London, UK). The raw data were searched against Mouse International Protein Index (IPI) database (released on August 10, 2009 and containing 56,733 protein sequences) with decoy database search enabled. The searches were executed with fixed modification carbamidomethyl cysteine and variable modification oxidized methionine. Peptide tolerance and MS/MS tolerance were set at 15 ppm and 0.8 Da, respectively. In addition, peptide charges of +2 and +3 were selected and maximal missed cleavage of one was allowed. Scaffold (Proteome Software Inc., Portland, OR, USA; version 3.0.0.04) was used for spectral count analysis and to validate LC-MS/MS-based peptide and protein identifications (25). Peptide identifications were accepted if they could be established at greater than 95.0% probability as specified by the Peptide Prophet algorithm (26). Protein identifications were accepted if they could be established at greater than 99.0% probability and contained at least two identified peptides (27). With these stringent parameters of Peptide Prophet and Protein Prophet, the false discovery rate was zero (27). Proteins that contained similar peptides and could not be differentiated based on MS/MS analysis alone were grouped to satisfy the principles of parsimony.

Data Analysis—For the gel-based fractionation, spectral counts and protein identifications from each gel slice were merged within each lane using Scaffold. Data normalization of spectral counts across experiments was performed using the Scaffold software. Pseudocounts were added to each spectral count value to account for missing values (28). Log2 fold-change values were computed for mESC versus mESIC and p values computed using Student’s t test. The normalized gene expression data of mouse mESC and mEpiSC in three biological replicates using Agilent whole-genome microarrays was obtained from a previous study (6). From this data, the mean log2 fold change and t test were used to calculate statistical significance p value of each probe for a given gene. For those genes with multiple probes, the value of the probe with the most significant p value was used to represent the corresponding gene. The p values from the Student’s T-tests were corrected using multiple hypothesis testing, and q-values were calculated using the bootstrap method (29). The set of 299 PluriNet genes were downloaded from http://www.stemcellmatix.org (15).

**Immuno blot Analyses and Quantification**—Equal amount (20 μg) of proteins from different samples were loaded on precast 4–12% Bis-Tris gel (NP-0335, Invitrogen, Carlsbad, CA) and subjected to electrophoresis. Gels were either stained with Coomassie Brilliant Blue or transferred to nitrocellulose membrane (10402594, Whatman, Dassel, Germany). Immunoblotting was used to detect the protein with super signal ELISA Pico chemiluminescent substrate (37070, Pierce, Rockford, IL). Primary antibodies anti-DNMT1 (5119, Cell Signaling Technology, Inc., Danvers, MA), anti-DNMT3L (ab3493, Abcam, Cambridge, MA), anti-Brg1 (3508, Cell Signaling Technology, Inc., Danvers, MA), anti-BAF53 (sc-271226, Santa Cruz Biotechnology, Inc.), anti-BAF155 (sc-10756, Santa Cruz Biotechnology, Inc., Santa Cruz, CA), anti-BAF60a (sc-135843, Santa Cruz Biotechnology, Inc.) and anti-α-tubulin (2144, Cell Signaling Technology, Inc.) as loading control were applied at 1:1000, 1:1000, 1:1000, 1:1000, 1:1000 and 1:2000, respectively and secondary antibody HRP-conjugated anti-Rabbit (NB730-H, Novus Biologicals, Littleton, CO) were added at 1:20000. Chemiluminescence detection using SuperSignal® ELISA Pico Chemiluminescent Substrate (PI-37070, Thermo Scientific, Rockford, IL) was applied to all immunoblots. The bands shown in blots were then quantified by ImageJ (http://rsbweb.nih.gov/ij/) (32, 33) and the mean values were plotted as well as the standard deviation for the two replicates of each protein.

**RESULTS**

**Analysis of mESC and mEpiSC Data Sets**—The goal of this study is to identify proteins and their associated networks that determine the distinct pluripotent states of mouse embryonic (mESC) and epiblast (mEpiSC) stem cells. An overview of the work flows used is shown in Fig. 1. Because pluripotency is largely regulated through control of gene-expression programs, we focused our proteomics experiments on analysis of the nuclear mESC and mEpiSC proteomes. In preliminary experiments, we analyzed both nuclear and cytoplasmic fractions of mESC and mEpiSCs and observed significant enrichment of known nuclear proteins and significant overlap with two previously described networks of pluripotency genes, PluriNet (15) and PluriNetWork (30) (Fig. 3). We also ascertained that spectral counts from nuclear and cytosolic fractions were similar for “housekeeping” proteins for mESCs and mEpiSCs showing that subcellular fractionation was similarly effective for each cell type (supplemental Table S7). In our extended studies we analyzed nuclear proteomes for both cell types (Fig. 1) and subsequent data analyses focused on these nuclear proteomes. To maximize coverage of the mESC and mEpiSC nuclear proteomes, two separate analytical strategies were applied. In the first, proteomes were separated by 10-fold 1D-SDS-PAGE fractionation (hereafter referred to as...
the in-gel fractionation experiment) (supplemental Fig. S2). In
the second study, an extended fractionation-free 4 h long
chromatographic gradient was used (hereafter referred to as
the in-solution experiment). In each study, 3 replicate mESC
and 3 replicate mEpiSC samples were analyzed (a total of 6 ×
10 = 60 MS/MS runs in the 1D SDS experiment and 6 MS/MS
runs in the in-solution fractionation free long gradient exper-
iment). These studies were performed using the same biolog-
ic samples (3 mESC and 3 mEpiSC samples) after subcel-
ular fractionation to enrich for nuclear associated proteins.
In addition, we also previously acquired gene-expression mi-
croarray data (6) from these two cell types, enabling analysis
of comparative gene-expression profiles from mESCs and
mEpiSCs.

Global comparison of the in-solution and in-gel fraction-
ation experiments showed substantial overlap of the proteins
identified. In total, the in-solution experiment identified 1462
proteins and the in-gel fractionation experiment identified 955
proteins according to the peptide and protein selection crite-
ria used (≥ two peptides/protein, protein probability ≥ 0.99).
Proteins identified in the in-gel fractionation experiment were
largely a subset of those identified using the in-solution ap-
proach (85.1% of in-gel proteins were also identified in the
in-solution experiment) (supplemental Fig. S3). To identify
quantitative differences in protein expression between mESC
and mEpiSC, spectral count analyses (Scaffold; Proteome
Software Inc.) were performed. mESC/mEpiSC ratios, p val-
ues and q-values (to account for multiple testing) were then
computed for each protein in each data set (in-solution or

Fig. 1. Experimental workflow. Sample preparation and compar-
ative microarray and proteomics analysis of mouse embryonic stem
cells (mESC) and mouse epiblast stem cells (mEpiSC) by LC-MS/MS.

Fig. 2. Biological trends in mESC and mEpiSC nuclear pro-
teomes. A, Top functional categories represented in combined mESC
and mEpiSC data sets (using all identified proteins) showing signifi-
cant representation of proteins involved in embryogenesis and develop-
ment. B, Proteins with significantly increased expression (p < 0.1;
Student’s t test) in either mESC and mEpiSC show significant differ-
ences in representation of several functional categories. C, Analysis
of mutant phenotypes of genes corresponding to mESC and mEpiSC
proteins (proteins differentially expressed in mESCs or mEpiSCs with
p < 0.1; Student’s t test). mESC proteins represent genes with
embryonic lethal phenotypes at earlier developmental stages than
mEpiSC proteins. Phenotypes are arranged according to develop-
mental stage: (1) Prenatal lethality, (2) embryonic lethality before
implantation (E0–E4.5), (3) embryonic lethality before somite formation
(E4.5–E8), (4) embryonic lethality before turning of embryo (E8–E9), (5)
embryonic lethality during organogenesis (E19–E14), (6) postnatal
lethality, (7) absent somites, (8) open neural tube.
in-gel). **supplemental Tables S1 and S2 provide complete sets**

We then assessed the concordance between the in-gel and in-solution experiments by analyzing the agreement of expression trends (mESC/mEpiSC) for proteins found in both. The trends between in-gel and in-solution experiments were highly concordant when proteins with \( p < 0.05 \) or \( p < 0.1 \) were compared (98.04% of proteins show same expression change trend; **supplemental Fig. S1**). In addition, we compared the in-solution and in-gel experiments to the microarray data sets and for those features in common, found similar concordance (**supplemental Fig. S4**). Two data sets were initially created for subsequent analyses. In the first, we considered all proteins with \( p < 0.05 \) (in in-gel or in-solution experiments; maximal q-value = 0.08) (466 total proteins) and in the second, all proteins with \( p < 0.1 \) (in in-gel or in-solution experiments; maximal q-value = 0.16) (646 total proteins). These two data sets showed similar concordance in expression trends between proteins identified in in-solution and in-gel experiments. In addition, we noted that the extra proteins included by considering proteins with \( p < 0.1 \) largely fell in to similar biological process and pathways as proteins with \( p < 0.05 \). For subsequent global analyses therefore, a union set of proteins was created consisting of all proteins in the in-solution or in-gel experiments with expression differences significant at \( p < 0.1 \) in one or both experiments. This union data set consists of 646 proteins, 336 showing increased expression in mESC and 310 showing increased expression in mEpiSC (**supplemental Table S5**). For the focused analyses of protein networks in Figs. 3, 4 and 5, we primarily focused on proteins with \( p < 0.05 \) and included selected proteins with \( p < 0.1 \) where necessary.

**Differing Functional Properties of the mESC and mEpiSC Nuclear Proteomes**—In analyzing functional annotations of the sets of mESC and mEpiSC proteins, we noted very significant representation of proteins involved in developmental processes. As shown in Fig. 2A, the most significant func-

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**FIG. 3. Intersecting proteomics studies with existing pluripotency networks.** A. Overall overlap of all identified proteins in union set of proteins identified from gel-free and gel-based proteomics studies (“union set all” - 2 peptides; Scaffold probability ≥ 99%) with PluriNet and PluriNetWork. B. The intersection between proteomics data and PluriNet (analyzed using Ingenuity Pathways Analysis) - 125 proteins. C. The intersection between proteomics data and PluriNetWork - 94 proteins. Each node in (B) and (C) represents a protein identified in mESC or mEpiSC samples (red shaded nodes indicate mESC>mEpiSC protein expression with \( p < 0.1 \) (Student’s t test), green shaded indicate mEpiSC>mESC protein expression with \( p < 0.1 \) (Student’s t test) and gray shaded nodes indicate identified protein in either mESC or mEpiSC but without statistically significant quantitative difference.

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tional categories pertain to developmental processes such as embryonic and tissue development.

To compare active cellular processes in mESCs and mEpiSCs, the sets of proteins either enriched \( p < 0.1 \) in mESC cells (336 proteins) or enriched \( p < 0.1 \) in mEpiSC (310 proteins) were analyzed. The most significant functional categories for each of these two sets of proteins are shown in Fig. 2B. Notably, mESCs are enriched in proteins encoding fundamental cellular processes such as DNA replication, recombination and repair \( (p = 4.46E-8) \). A greater enrichment of proteins involved in embryonic development is found in the mESC than in mEpiSC samples \( (p \) values equal to 6.69E-06 and 3.66E-04 respectively). Of the 56 proteins from mEpiSC included in this category, several pertain directly to epiblast related functions such as survival of embryonic stem cells (DNMT1), hatching of blastocyst (SMARCA4) and the formation or morphology of the visceral endoderm (MYH10). These global analyses show that proteins with pertinent biological functions are identified in the proteomics screens and identify functional categories that significantly differ between mESCs and mEpiSCs.

To further understand the differences in developmental functions represented by mESC and mEpiSC proteins, we analyzed mouse phenotypes of corresponding mutations. All mESC and mEpiSC proteins from the union set were mapped to embryonic phenotypes using the Mammalian Phenotype resource (31). Embryonic lethal phenotypes and their associated numbers of genes represented in the mESC and mEpiSC sets are shown in Fig. 2C. The overall numbers of proteins that could be assigned to a phenotype using the Mouse Genome Informatics database (31) were approximately similar for mESC and mEpiSC proteins (140 and 147 respectively), representing \( \sim50\% \) of the mESC and mEpiSC sets overall. However, the phenotypes differ markedly between mESC and mEpiSC. We focused on embryonic lethality phenotypes as shown in Fig. 1C, and found that mESC proteins were more highly associated with very early embryonic lethality (pre-implantation failure or lethality prior to somite formation) than mEpiSC proteins, which showed greater representation of genes with lethality occurring during organogenesis and post-natally. Thus, aside from the differences in functional categorizations (Fig. 1B), the mESC and mEpiSC proteins represent at least in part, proteins expressed as part of the ordered process of embryonic development. In addition, the association of significant proportions of the identified proteins with embryonic mutant phenotypes suggests that many of the identified proteins play key, nondispensable roles in development.

We also observed many known markers of embryonic stem cells in our proteomics data sets. Table I shows these markers with their respective fold-differences between mESC and mEpiSCs. Notably, most of the known markers show increased expression in mESC samples. We observed 4 distinct peptides corresponding to Nanog protein (uniquely in mESC samples). Although the Nanog protein is a known Oct4 binding partner, Nanog is not consistently detected using Oct4 immuno-purification (12, 34, 58), possibly because of difficulty in digesting the Nanog protein to tryptic peptides (34). Thus proteomics profiling approaches, such as our own, are complementary to focused immuno-purification of pluripotency associated proteins.

Integrated Pluripotency Networks—To understand more fully what our proteomics data reveal about pluripotency, we integrated our data with two previously defined pluripotency networks. A previous study used gene-expression profiling to provide a large-scale classification of diverse stem cell lines (15). By integrating these transcriptional profiles with protein-protein interaction networks, a core pluripotency network, named PluriNet (15), was defined. A second study, using an alternative approach of curating literature, created a pluripotency network, PluriNetwork, focused on molecules and interactions underlying pluripotency mechanisms in mouse stem cells (30). We first compared the complete set of proteins identified in our study to those proteins represented in PluriNet and PluriNetwork. As shown in Fig. 3A, there are significant overlaps of our identified proteins with PluriNet and PluriNetwork, showing that the mESC and mEpiSC nuclear

### Table I

| Gene symbol (mESC/mEpiSC)* | Protein name | mESC/mEpiSC protein fold change | p value (T-test) | References |
|---------------------------|--------------|--------------------------------|-----------------|-----------|
| Oct4/Pou5f1                | POU class 5 homeobox 1 | 1.59 | 0.0005 | 51 |
| Sox2                      | SRY (sex determining region Y)-box 2 | 2.00 | NA | 51 |
| Nanog                     | Nanog homeobox | 0.96 | 0.098 | 51 |
| Utf1                      | Undifferentiated embryonic cell transcription factor 1 | 0.57 | 0.008 | 52 |
| Dppa4                     | Developmental pluripotency associated 4 | 1.56 | 0.002 | 53 |
| Trim28                    | Tripartite motif containing 28 | 3.53 | 0.002 | 54 |
| Dnmt3a                    | DNA (cytosine-5')-methyltransferase 3 alpha | 4.39 | NA | 45, 57 |
| Dnmt3b                    | DNA (cytosine-5')-methyltransferase 3 beta | −1.00 | 0.02 | 57 |

\*Log2 (mESC/mEpiSC) values (positive values indicate higher expression in mESCs and negative values indicate higher expression in mEpiSCs).
proteomes are highly enriched for proteins with roles in pluripotency. We detected 42% of the total proteins in PluriNet and 35% of the total proteins represented in PluriNetwork. In addition, we noted that there is relatively little overlap between PluriNet and PluriNetwork; indeed only 31 genes are represented in both PluriNet and PluriNetwork (~10% of each network). This likely reflects the different methods used to derive PluriNet (unbiased clustering of gene-expression profiles and integration with high-throughput protein-protein interaction data) and PluriNetwork (curation of pluripotency-related publications). It also suggests that each resource most likely represents an incomplete network of pluripotency-related genes/proteins. This finding prompted us to further integrate our data with these pluripotency networks to see whether the networks might be extended using our proteomics data.

Proteomics data was integrated with PluriNet and PluriNetwork as shown in Figs. 3B and 3C respectively. In each case, all proteins present in both our proteomics data and PluriNet (Fig. 3B) or our proteomics data and PluriNetwork (Fig. 3C) were analyzed using Ingenuity Pathways Analysis. Proteins shown in gray were detected in our study but not significantly different between mESC and mEpiSC, whereas proteins in red are those proteins significantly (p < 0.1) more abundant in mESC than mEpiSC and proteins in green those proteins significantly more abundant in mEpiSC than mESC proteomes. Considering only proteins with statistically significant differences between mESC and mEpiSC (p < 0.1) in the union data set, we observed a greater overlap between mESC and PluriNet (44 proteins) than between mEpiSC and PluriNet (15 proteins) (Fig. 3B). Similarly, we observed a greater overlap between mESC and PluriNetWork (38 proteins) than between mEpiSC and PluriNetWork (19 proteins) (Fig. 3C). The greater representation of proteins enriched in mESC as compared with mEpiSC suggests that existing pluripotency networks represent the mESC pluripotency state better than the mEpiSC pluripotency state. In addition, there is greater representation of embryonic stem cell samples in PluriNet than epiblast cell samples (15).

We also noted differences in the properties of the PluriNet/proteomics intersecting network (Fig. 3B) and the PluriNetwork/proteomics intersecting network (Fig. 3C). The PluriNet network (Fig. 3B) is noticeably less dense in terms of connectivity than the PluriNetwork (Fig. 3C), presumably attributable to the different sources of information for PluriNet and PluriNetwork. In addition, proteins represented in Figs. 3B and 3C differ in their functional categories. Table II lists the most significant functional categories. Interestingly, although some of these functional categories are the same, the degree of enrichment of the PluriNetwork associated categories is greater than that for PluriNet. For example, Embryonic development (p value = 3.41E-09 for PluriNetwork versus p value = 5.53E-05 for PluriNet) and Tissue development (p value = 1.85E-08 for PluriNetwork versus p value = 1.68E-07 for PluriNet).

Integrating the proteomics data with PluriNet and PluriNetWork also suggested that known pluripotency networks might be extended. The relatively small overlap between PluriNet and PluriNetWork (31 proteins representing ~10% of the total from each network) and the different properties of the networks as noted above suggest that the underlying pluripotency network is incomplete. By considering PluriNet, PluriNetWork and the proteomics data set, we observed several protein complexes that are incompletely represented in the known pluripotency networks. For example, although PluriNet includes components of the chromatin remodeling BAF (Brg1 associated factor) complexes, such as SMARCD1 and SMARCA1, and PluriNetWork includes SMARCA4 and SMARCA1, our data suggest that additional BAF complex components SMARCB1, SMARCD2, and SMARCC2 are also intrinsic to the pluripotency network. Similarly, a tightly clustered homeobox subnetwork of PBX1, PBX2, MEIS1, MEIS2,
and HOXA5 (Fig. 4) suggests that these may also be included in the core pluripotency network along with PBX1. In summary, our data add many key proteins to the clustered functional protein complexes in PluriNet and PluriNetWork. 

Nuclear Proteomics Differentiates mESC and mEpiSC Protein Networks—The principal objective of this study is to identify proteins and networks that determine the pluripotent ground states of mESCs and mEpiSCs. We therefore analyzed the complete set of proteins in the union set (p < 0.1) using Ingenuity Pathways Analysis to identify subnetworks connecting proteins with significant differential expression in either the mESC or mEpiSC proteome. We noted in particular, that networks regulating gene-expression through chromatin modification and DNA methylation were highly represented with significant differential expression in the mESC or mEpiSC proteomes. Fig. 4 shows several selected protein networks showing coherent differential expression in mESC and mEpiSC proteomes. In Fig. 4A, several DNA and chromatin-modification associated protein networks showed significant enrichment in either mESCs or mEpiSCs. The Oct4-Nanog subnetwork and associated proteins is more highly expressed in the mESC proteome, along with Polycomb complex components. In contrast, several DNA methyltransferases and components of SWI/SNF nucleosome remodeling complexes (also known as BRG associated factors, BAF) were found to be enriched in the mEpiSC proteome. Other notable subnetworks are shown in Figs. 4B–4D. C-terminal binding protein 1 (CTBP1) is a transcriptional co-repressor with known roles in transcriptional repression during development and in oncogenesis (35). As shown in Fig. 3C, several known interacting partners of CTBP1 show significant increased expression in the mEpiSC proteome. ZEB1 binds to CTBP1 and functions in concert with CTBP1 as a transcriptional repressor and regulator of the epithelial to mesenchymal transition (EMT) (36). The zinc-finger protein, WIZ, was shown to physically link the EHMT1 histone methyltransferase to the CTBP repressor machinery in mESCs (37). Thus, coherent increased expression of this complex suggests increased activity in mEpiSCs. Fig. 4D shows a subnetwork of interacting homeobox transcription factors also significantly enriched in mEpiSCs. PBX, MEIS, and HOX classes of homeobox are known to form trimeric complexes (38). PBX and MEIS homeoproteins act as cofactors for HOX domain transcription factors and play important roles during early mammalian development (39).

We were particularly intrigued by the differential expression of DNA methyltransferases, the polycomb complexes and the SWI/SNF-related Baf components that we observed between mESCs and mEpiSC proteomes. To further qualify these observations, we cross-referenced our proteomics data set to our previous microarray-based comparison of mESCs and mEpiSC gene-expression (Gene Expression Omnibus accession GSE26814; supplemental Table S4). By integrating the gene-expression microarray and proteomics data at the gene level, we observed that the global trends between the data sets show significant similarity (supplemental Fig. S4). We analyzed in more detail subsets of genes represented in our protein networks. Supplemental Table S4 contains mESC/mEpiSC ratios from the stem cell proteomics and gene-expression data sets for genes represented in the subnetworks in Fig. 4.

We first analyzed the expression of several SWI/SNF-related Baf complex components. Two Baf complex components, Smarcd1 and Smarca4 were observed to be more abundant in mEpiSCs than in mESCs at the protein and gene-expression levels. To further validate the findings from proteomics and gene-expression microarray data sets, several proteins were selected for immunoblot analysis. Fig. 5 shows immunoblot analysis (Fig. 5A) and quantification of expression (Fig. 5B) of selected components from the chromatin and DNA modification subnetwork in Fig. 4A. Proteins were selected based upon their observed differences in expression according to the spectral count analyses as well as the availability of a suitable antibody. Two independent immunoblots were performed for each protein and the results used to quantify and measure ratios of expression (Fig. 5B). We were also interested to further explore differences in expression of DNA methyltransferases. In particular, in our preliminary proteomics comparisons of mESCs and mEpiSCs, we observed dramatically increased expression of Dnmt3l (DNA methyltransferase-3 like) protein in mESCs as compared with mEpiSCs (supplemental Table S6). Since we could not confirm this finding with subsequent experiments (peptides corresponding to Dnmt3l protein were not observed in our union data set), we used anti-Dnmt3l antibody in immunoblots to compare expression in mESCs and mEpiSCs. As shown in Fig. 5, Dnmt3l protein shows markedly increased abundance in mESCs as compared with mEpiSCs. In contrast, a binding partner of Dnmt3l, Dnmt1 was more abundant in mEpiSC proteomes than in mESCs (Fig. 4A). Thus, these results combined with the proteomics data show that specific perturbations of DNA and chromatin modification networks are detectable using nuclear proteomics of mESCs and mEpiSCs and that these perturbations may determine the pluripotency ground states of mESCs and mEpiSCs.

DISCUSSION

Understanding the molecular basis of pluripotency in stem cells is imperative to being able to manipulate stem cells for applications in regenerative medicine. In addition, it is clear that there exist multiple types of stem cell with potentially different therapeutic applications. The mechanisms that maintain pluripotency in different classes of embryonic stem cells, exemplified by mESCs and mEpiSCs, may differ and it is important to understand these differences for subsequent efforts to manipulate these different cell types. In this study, we have undertaken an in depth comparison of the nuclear proteomes of mESCs and mEpiSCs. Our proteomics data...
sets are highly enriched in proteins important during embryogenesis and overlap and extend known networks of pluripotency associated proteins, providing a framework for future manipulation of the pluripotency state. In addition, our analysis of mutation phenotypes associated with mESCs or mEpiSCs proteins showed that there is an association of proteins expressed in each cell with known mutations at different stages of embryogenesis (Fig. 2C). mESC proteins are...
are more likely to correspond to genes that when mutated show embryonic lethality prior to somite formation, whereas mEpiSC proteins correspond to genes associated to lethality during organogenesis and post-natal lethality. This is significant because it shows that cultured epiblast stem cells recapitulate developmentally dependent patterns of expression, thus reinforcing their value as a model for early mammalian development. Finally, qualitative and quantitative comparisons of the mESC and mEpiSC proteomes reveal key differences in particular in the expression of protein networks associated with DNA and chromatin-modification, pointing to the importance of these processes in specifying mESC and mEpiSC pluripotency states.

This study represents the first in-depth proteomics study of epiblast stem cell pluripotency, significant because of the apparent similarity between mEpiSC state and human embryonic stem cells. The quantitative and qualitative differences between the mESC and mEpiSC proteomes have the potential to shed significant light on the different pluripotency states and properties of these two stem cell types. We observed many proteins, biological processes and protein networks that were more abundant in mEpiSCs than mESCs (Fig. 4). In particular, we observed networks of transcription factors such as the Meis-Pbx-Hox complex (Fig. 4D) that suggest the epiblast stem cells are primed for the expression of downstream developmental programs. Increased expression of Cbpt and associated proteins (Fig. 4C) in the epiblast proteome and their role in transcriptional repression and the epithelial to mesenchymal transition, also suggest that whereas the epiblast stem cells maintain pluripotency, they are also primed for key developmental transitions.

Our results emphasize the importance of chromatin remodelling protein complexes and networks in regulating the pluripotent state. Together with other studies indicating the specific role of BAF complex components (11, 17), our results suggest that different pluripotent states may in part be determined by the balance (or reconfiguration) of BAF complex components that both operate to activate and repress gene-expression. Specifically, we identified several components of ATP-dependent chromatin remodeling complexes that exhibit differential expression between mESCs and mEpiSCs. Expression of Smarcd1/Baf60a and Smarca4/Brg1 were both significantly higher in mEpiSCs than mESCs. Interestingly, Smarcd1/Baf60a mediates the interaction between several signaling pathways and BAF complexes (40). In human cells, a direct interaction between p53 with SMARCD1/BAF60A (but not between p53 and SMARCA4/BRG1) has been reported (41). Uncoupling of the interaction between SMARCD1/BAF60A and p53 resulted in repression of p53-dependent cellular functions. SMARCD1/BAF60A also mediates the interaction between nuclear hormone receptors and BAF complexes (42), governing the efficacy of chromatin remodeling in response to glucocorticoid receptor hormone signaling.
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mouse lymphosarcoma cells, Dnm3A was shown to co-immunoprecipitate with Smarca4/Brg1 (49). Given the roles of DNA and chromatin modification networks in transcriptional regulation, future studies that identify the DNA target sites of these proteins in mESCs and mEpiSCs should be highly informative. Genome binding sites of core SWI/SNF components and their protein-protein interactions, have revealed a complex picture of diverse activities and both repression and activation of gene-expression by SWI/SNF complexes (50). The potentially different and significant patterns of DNA-binding of SWI/SNF components in mESCs and mEpiSCs and their significance remain to be discovered.

Integration of mESC and mEpiSC nuclear proteomics data with existing pluripotency networks, PluriNet and Pluri-Net, shows that protein expression profiling may significantly augment existing pluripotency networks. The pluripotency network serves as a framework for studies such as ours, but as evident from our study, expression differences between components of pluripotency networks may be key to understanding the differences between pluripotent stem cell types and their pluripotent ground states. In this study, we have identified specific protein expression differences between mESCs and mEpiSCs, in protein complexes with known roles in maintaining the pluripotent state. This study supports the notion that mESCs and mEpiSCs maintain distinct pluripotent ground states. Future work will focus on understanding the functional significance of findings in this study with a view to using the protein networks as indicators of pluripotent state and as a foundation for identification of factors that can be used to manipulate the different pluripotent states.

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