Comprehensive Mapping of Pluripotent Stem Cell Metabolism Using Dynamic Genome-Scale Network Modeling

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Comprehensive Mapping of Pluripotent Stem Cell Metabolism Using Dynamic Genome-Scale Network Modeling

**Graphical Abstract**

**Highlights**

- Systems approach infers metabolic network state using time-course metabolomics
- Characterized the metabolic state of naive and primed murine embryonic stem cells
- Tracked metabolic reprogramming by the pluripotency factor Lin28 in iPS cells
- Simulated and validated the influence of metabolic pathways on histone methylation

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**In Brief**

Chandrasekaran et al. use computational modeling, metabolomics, and metabolic inhibitors to discover metabolic differences between various pluripotent stem cell states and infer their impact on stem cell fate decisions.
Comprehensive Mapping of Pluripotent Stem Cell Metabolism Using Dynamic Genome-Scale Network Modeling

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SUMMARY

Metabolism is an emerging stem cell hallmark tied to cell fate, pluripotency, and self-renewal, yet systems-level understanding of stem cell metabolism has been limited by the lack of genome-scale network models. Here, we develop a systems approach to integrate time-course metabolomics data with a computational model of metabolism to analyze the metabolic state of naive and primed murine pluripotent stem cells. Using this approach, we find that one-carbon metabolism involving phosphoglycerate dehydrogenase, folate synthesis, and nucleotide synthesis is a key pathway that differs between the two states, resulting in differential sensitivity to anti-folates. The model also predicts that the pluripotency factor Lin28 regulates this one-carbon metabolic pathway, which we validate using metabolomics data from Lin28-deficient cells. Moreover, we identify and validate metabolic reactions related to S-adenosyl-methionine production that can differentially impact histone methylation in naive and primed cells. Our network-based approach provides a framework for characterizing metabolic changes influencing pluripotency and cell fate.

INTRODUCTION

Pluripotent stem cells are able to self-renew and have the capacity to regenerate all tissues in the human body. Recently, there has been a resurgence of interest in stem cell metabolism due to its role in affecting signaling pathways and epigenetic processes, apart from synthesizing precursors and generating energy for stem cell renewal. Understanding the metabolism of pluripotent stem cells holds promise for understanding early development as well as for regenerative medicine.

Two distinct states of pluripotent stem cells that differ in their developmental potential have been recently described. Naive murine embryonic stem cells (ESCs) that closely mirror the inner cell mass of the pre-implantation embryo are maintained in culture with ERK/MEK inhibitor and GSK3 inhibitor (2i) (Nichols and Smith, 2009), whereas primed ESCs derived from the post-implantation embryo are maintained in the presence of fibroblast growth factor 2 (FGF2) and Activin (Brons et al., 2007; Tesar et al., 2007). These states also show distinct patterns of metabolism. Naive cells predominantly employ mitochondrial oxidative metabolism (Huang et al., 2014; Takashima et al., 2014; Zhou et al., 2012) and utilize more glucose and less glutamine to make α-ketogluturate, which in turn influences histone and DNA methylation (Carey et al., 2015). Human naive-like cells derived with different protocols (Gafni et al., 2013; Takashima et al., 2014; Theunissen et al., 2014; Ware et al., 2014) consistently exhibit high oxidative metabolism, as well as low S-adenosyl-methionine (SAM) levels due to high consumption that presumably leads to histone hypomethylation (Sperber et al., 2015), a characteristic of the naive state (Marks et al., 2012). It has been shown that mouse ESCs in LIF/serum media (a metastable state between naive and primed) utilize threonine to feed the SAM pool for H3K4 methylation (Shyh-Chang et al., 2013), and human ESCs require methionine to maintain pluripotency (Shiraki et al., 2014). However, the role and activity of enzymes involved in one-carbon metabolism, which impacts nucleotide biosynthesis, redox homeostasis, and methylation reactions, in two clearly defined naive and primed
Figure 1. Overview of Our Approach to Integrate Time-Course Metabolomics (Intracellular or Extracellular) with Genome-Scale Metabolic Models to Reconstruct the Metabolic Network of Different Cell Fates

(A) A simple metabolic network consisting of two parallel pathways to produce the growth-associated metabolite C is shown, along with its corresponding stoichiometric matrix representation. Deleting reactions 2 or 4 should impact the biomass production equally in this model.

(B) Time-course metabolomics data from two different conditions are then overlaid onto the metabolic network model and dynamic metabolites are identified. The accumulation or depletion of metabolites resulting from increased or decreased flux through the reactions involving those metabolites are represented in silico.

(C) Reaction deletion and flux variability analysis are performed to understand the metabolic flux changes.

(D) Metabolic model with 3744 reactions, 2771 metabolite species, and 1487 genes is used for FBA with Flux Activity Coefficients for deviation from steady state.

(E) Genome-scale metabolic model + Time-Course Metabolomics is used to predict differentially sensitive reactions (red nodes).
pluripotency states of mouse ESCs has not been explored from a systems perspective. The enormous size and complexity of the mammalian metabolic networks has so far limited systems-level understanding of stem cell metabolism. Expression analysis of metabolic genes or metabolomics can provide a snapshot of the metabolic status, but is limited because gene or metabolite changes often do not reflect the flux of the metabolic reactions. Isotope tracing analysis can determine flux but only focuses on specific metabolic pathways and does not give a systems perspective of the entire network. Pathway enrichment analysis is another commonly used approach to characterize metabolic changes at the systems level (Subramanian et al., 2005; Xia et al., 2015). However, given the highly inter-connected nature of the metabolic network, the underlying assumption behind pathway enrichment analysis that each pathway is independent of each other does not hold for metabolism (Shlomi et al., 2008). Adjacent pathways on the network can influence each other’s activity, and altered flux activity can arise due to perturbations upstream of a given pathway. Furthermore, individual enzymes in a pathway do not change coherently as a whole and can have different levels of activity. A systems-level model is thus needed to understand metabolic differences between different cell states at both the individual reaction level and the network level.

Here, we use genome-scale computational modeling to comprehensively characterize the metabolism of different pluripotent stem cell states. Genome-scale metabolic network models are manually curated and represent the mechanistic relationships among thousands of genes, proteins, and metabolites within a biological system (Bordbar et al., 2014). Recently, genome-scale metabolic models have been integrated with transcriptomics data to identify a subset of metabolic genes that are active in a system. Such transcriptomics-constrained models have been applied successfully to predict metabolic behaviors of human tissues and cancer cells at steady state (Frezza et al., 2011; Uhlen et al., 2015).

Because stem cell metabolic rewiring is dynamic (i.e., not at steady state) and has been known to be regulated at the post-transcriptional level (Zhang et al., 2016), we developed a genome-scale modeling approach to directly infer metabolic states based on time-course metabolomics data. Similar to pathway enrichment analysis where increased levels of metabolites in a pathway suggests increased pathway activity in a condition, we assume that the accumulation or depletion of a metabolite over time is evidence for increased or decreased overall flux activity of the set of reactions involving the metabolite. By integrating data from several metabolites into a unified genome-scale model, we can identify reactions whose differential activity will most likely explain the observed pattern of metabolic changes.

We first validate this approach by identifying metabolic vulnerabilities of the NCI-60 cancer cell lines. This approach is then applied to characterize murine naive and primed ESC metabolic states, and track the metabolic effect of Lin28, a pluripotency factor (Viswanathan and Daley, 2010) associated with the primed state (Zhang et al., 2016). We confirm a transition of mitochondrial metabolism and uncovered an elevation of one-carbon metabolism pathways from naive to primed state. Using Lin28 knockout pluripotent stem cells as a surrogate of the naive state, we demonstrate a pivotal role of Lin28 in conferring the unique one-carbon and nucleotide metabolism associated with primed pluripotency.

**RESULTS**

**Construction of a Dynamic Genome-Scale Metabolic Model**

Our proposed approach integrates time-course metabolomics data with a genome-scale metabolic network model and makes use of the flux balance analysis (FBA) framework to infer the metabolic state of a condition (Orth et al., 2010). The FBA approach identifies the optimal flux through each reaction in the metabolic network that maximizes a specific objective such as the growth rate, while satisfying stoichiometric and mass-balance constraints.

The integration of metabolite levels with genome-scale metabolic models has been a significant challenge due to the lack of thermodynamic and kinetic parameters (Cotten and Reed, 2013; Yizhak et al., 2015). In addition, integrating metabolomics measurements from mammalian systems poses additional challenges due to noise in the data, the undefined nature of serum components in the media, and compartment-specific metabolism. In microbial dynamic flux-balance models, time-course metabolomics measurements are used to explicitly determine metabolite accumulation/depletion rates (Kleessen et al., 2015; Schmidt et al., 2013). In contrast, given the complexity associated with eukaryotic metabolic measurements due to cellular heterogeneity and compartmentalization, our approach uses time-course metabolomics measurements as cues for likelihood of altered flux activity around a metabolite. Our approach tries to identify a metabolic state that best fits the metabolomics data while allowing for deviations from observed pattern of accumulation or depletion of metabolites. Network integration is then used to combine these cues into a global, consistent metabolic state (Experimental Procedures; Figure 1).

Using our approach, we can infer the impact of the observed differential metabolite levels on the corresponding reaction, the encompassing metabolic pathway, and the entire network of thousands of metabolic reactions. Furthermore, the input data can be either intracellular or extracellular. In the metabolic model, metabolites in each compartment (i.e., extracellular, cytosol, mitochondria, nucleus, or other organelles) are distinct from each other. Transport reactions are used to connect metabolites in different compartments. Because the network is unified, the
impact of changes in metabolites in any compartment can be predicted on the network. For example, changes in extracellular metabolite levels will impact the uptake or secretion flux of these metabolites, which in turn will impact reactions downstream of these transport reactions. Hence, data from extracellular measurements can be directly utilized to constrain the model using the same mathematical framework used for intracellular metabolites. This approach goes beyond traditional pathway enrichment analysis by creating a genome-scale model of the metabolic state of a system. The in silico model can be subsequently used to simulate deletion of metabolic genes or inhibition of enzymes, in addition to identifying differentially active reactions and pathways.

As a validation of our method in complex mammalian systems, we applied it to predict the metabolic vulnerabilities of the NCI-60 cancer cell lines. The metabolic properties of these cell lines have been characterized using metabolomics collected at two different time points (Jain et al., 2012). Using our approach, we integrated these time-course metabolomics data with a genome-scale model of human metabolism (Duarte et al., 2007) consisting of 1,487 genes and 3,744 metabolic reactions to map the metabolic state of each of the NCI-60 cell lines.

We simulated deletion of all metabolic genes in the model and predicted the effect of deletion on the growth of the cell. Some genes affected the growth of all the cell lines, whereas others were more selective for specific cell and tissue type (Figure 2A). We first identified common metabolic vulnerabilities across all 60 cell lines. This was done by identifying genes that when deleted impact growth by at least 5% of the wild-type (Figure 2A; Supplemental Experimental Procedures). Comparing the predicted metabolic vulnerabilities with data from small interfering RNA (siRNA) and CRISPR-Cas9 gene knockdown screens (Cheung et al., 2011; Marcotte et al., 2012; Aguirre et al., 2016) revealed that cell lines were significantly more sensitive to siRNAs targeting these genes compared to other metabolic genes, suggesting that the genes identified by our approach were more likely to be essential to the growth of these cell lines (p value = 8 × 10^{-3}, 2 × 10^{-15}, and 3 × 10^{-20}, respectively, for siRNA [Cheung et al., 2011; Marcotte et al., 2012] and CRISPR data [Aguirre et al., 2016], one-sided Kolmogorov–Smirnov test [KS] test). As a negative control, prediction using the metabolic model without metabolomics constraint resulted in no significant enrichment.

We next used the integrated model to determine metabolic differences between the cell lines. We identified metabolic genes that were differentially vulnerable in a cell line relative to the remaining cell lines (Figure 2A). These genes were also more likely to be predicted as selectively vulnerable in these cell lines based on both siRNA and CRISPR-Cas9 screens (p value = 3 × 10^{-2}, 1 × 10^{-15}, and 1 × 10^{-20}, respectively, for siRNA [Cheung et al., 2011; Marcotte et al., 2012] and CRISPR data [Aguirre et al., 2016], one-sided KS-test) (Figures 2B–2D). These results are notable given the variable nature of high-throughput siRNA screens and metabolomics data across different studies. These findings suggest that our approach can accurately capture metabolic differences between cell states using time-course metabolomics data.

**Modeling Murine Naive and Primed ESC Metabolism**

To characterize the metabolic state of mouse ESCs in naive and primed states, we measured levels for over 200 metabolites at three time points (Figure 3A). The metabolomics data revealed changes in several metabolites, but these were not specifically related to any one pathway. Pathway enrichment analysis of metabolomics data mirrored this complexity and only two pathways—purine metabolism and urea cycle—were predicted to be differentially active at false discovery rate (FDR) < 0.05 (Tables S1A and S1B). Enrichment analysis of gene expression data of these two states also did not reveal any significant differential activity for a metabolic pathway over the background (FDR < 0.1). Changes in metabolic pathways were masked by strong changes in other pathways such as signaling pathways (Table S1E).

Given the limited insight from enrichment analysis, we applied our dynamic genome-scale modeling approach to tease out the underlying metabolic differences between the two cell fates. The metabolomics data were then used to create a dynamic genome-scale metabolic model of the naive and primed state (Figure 3A). We performed deletion analysis of all the metabolic reactions to discover differences between the two metabolic networks (Figure 3B). In addition to reaction deletion analysis, which identified the dominant metabolic differences between the two states, we used flux variability analysis (FVA) to determine the range of feasible fluxes for each reaction based on thermodynamic, stoichiometric, and metabolomics constraints (Mahadevan and Schilling, 2003). FVA can capture flux differences in essential reactions, which cannot be inferred through deletion analysis as they are lethal to both states. The combination of reaction deletion analysis and FVA was used to prioritize reactions that showed differential sensitivity and flux activity.

The majority of the 3,744 reactions (96%) did not show selectivity between the two states, i.e., their knockout affected both states to the same extent. Among the reactions that were selective, the primed state network was more sensitive to knockout of reactions in folate/SAM/one-carbon metabolism, cysteine-methionine synthesis, and purine synthesis, whereas the naive state was more sensitive to reactions in oxidative phosphorylation, tricarboxylic acid (TCA) cycle, and pyrimidine synthesis (Figures 3B and 3C). We also observed that reactions in mitochondria were more sensitive in the naive state. For example, reactions involving folate in the cytoplasm were more active in the primed state, whereas folate reactions in mitochondria were more active in the naive state (Tables S2A and S3A).

**Validating Predicted Metabolic Differences between Naive and Primed Mouse ESCs**

Consistent with the in silico predictions, we found using ^13^C-glucose and ^13^C-glutamine tracing that, in the naive state, glucose had increased incorporation into the mitochondrial TCA cycle metabolites such as citrate, α-ketoglutarate/glutamate, fumarate, and malate, and decreased incorporation into one-carbon metabolism metabolites such as serine, 3-phosphoserine, and methionine; glutamine had decreased incorporation into nucleotides (Figures 3D and 3E; Figures S1A and S1B). Overall, metabolites associated with differentially sensitive reactions were also more likely to be differentially labeled between the two states (p value = 0.005, hypergeometric test; Figure S1F), confirming the strong concordance between two datasets.

To further assess the importance of folate and one-carbon metabolism predicted by the model, we tested our predictions...
by using the anti-folate compound methotrexate. Methotrexate is a well-established inhibitor of nucleotide synthesis and folate synthesis that has been used in many previous studies (Singh et al., 2006). It inhibits dihydrofolate reductase (DHFR), which participates in tetrahydrofolate synthesis and thus impacts de novo synthesis of purines and thymidine. As predicted by the

Figure 2. Validation of Our Approach to Infer Metabolic States from Time-Course Metabolomics Data by Applying It to Predict the Metabolic Behavior of NCI-60 Cancer Cell Lines

(A) Table of cell-line-specific metabolic vulnerabilities predicted by our approach. The predicted growth (measured by ATP flux) of 48 differentially sensitive gene knockouts in the NCI-60 cell lines is shown relative to wild-type. Highlighted genes affect ATP flux across all cell lines.

(B) The set of cell-line-specific metabolic gene vulnerabilities predicted by our approach were consistent with siRNA screens. The GARP (gene activity rank profile) scores of genes that were sensitive (vulnerable) in a cell line were found to be lower than those that were not predicted to be sensitive in a given cell line. The lower the GARP score for a given cell line, the more essential is the gene in that particular cell line (data from Marcotte et al., 2012). The distribution of GARP scores for the sensitive and non-sensitive associations are shown. The distributions were compared using the KS test (p value = 10^-5).

(C) Predictions were also consistent with siRNA knockout screen (p value [KS test] = 0.0034; Cheung et al., 2011). Distribution of ATARIS (analytic technique for assessment of RNAi by similarity) z-scores are shown for sensitive and non-sensitive gene knockouts across all cell lines.

(D) Predictions were also consistent with a CRISPR/Cas9-mediated knockout screen (p value [KS test] = 0.0012; Aguirre et al., 2016). Distribution of ATARIS z-scores are shown for sensitive and non-sensitive gene knockouts across all cell lines.

(E) Sensitivity analysis of the dynamic modeling approach. Increasing the weight parameter (kappa) results in increasing weight for the metabolomics data over growth. Correlation with the default settings (kappa = 1, i.e., equal weights for growth and metabolomics) and after changing kappa is shown for the predicted difference in growth between naive and primed states after deletion of all the 3,744 metabolic reactions in the model.

(F) The approach is robust to noise in the metabolomics data. 10%, 25%, and 33% of the metabolites in the metabolomics data were randomly removed, and the impact on the predictions between naive and primed state was analyzed. Correlation from predictions using this randomly sampled data with the predictions using the entire dataset is shown. Predictions made by using just the direction of change (accumulation or depletion) without the magnitude also gave qualitatively similar predictions as using the entire data (shown as yellow bar).
Figure 3. Mapping the Metabolic Differences between Naive and Primed Stem Cells Using the Computational Model of Metabolism

(A) Overview of our approach to measure time-course metabolomics data and identify differentially active reactions.

(B) To identify differentially active metabolic reactions, we performed reaction knockout analysis on the naive and primed metabolic networks. The histogram shows the distribution of growth rates after in silico deletion of all the metabolic reactions. The horizontal x axis shows the relative selectivity of each knockout for naive state versus the primed state. The stronger the magnitude of selectivity, the greater the difference between the growth rate of the reaction knockout between the two states. The majority (96%) of knockouts do not have a differential effect on naive or primed metabolism. The y axis shows the total number of reactions in each bin. Metabolic reactions that showed the greatest differences in growth between the two states are highlighted. The full list of reactions and their abbreviations are provided in Table S2A. The suffix "m" at the end of each reaction name indicates a reaction occurring in mitochondria; isoforms are represented with numerical suffixes (orange: more sensitive to deletion in naive; purple: more sensitive to deletion in primed).

(C) Schematic diagram of the metabolic pathways that are predicted to differ between the two states (naive, orange labels; primed, pink labels) as identified by the model. The thickness of the reaction name label border is proportional to the extent of the differential activity of the reactions between the two states. Reactions predicted to impact SAM flux in primed state (MAT, AHC, and METS) are also highlighted (Table S4).

(D) 13C tracing of glucose revealed that metabolites in the one-carbon and nucleotide metabolism were differentially labeled between the two conditions suggesting re-routing of metabolic flux. The heatmap shows the total isotopomers labeled at different time points for each metabolite after z-transformation (Supplemental Experimental Procedures).
model, primed cells were more sensitive to inhibition by methotrexate (Figure 3F). In contrast, naive cells were more sensitive to mitochondrial folate metabolism inhibition by the drug MTH1459 that blocks MTHFD2 activity, also confirming the in silico predictions (Figure 3F).

Decreased one-carbon metabolism in the naive state might not be caused by reduced glycolysis because $^{13}$C incorporation to lactate was not lower in the naive state (Figure S2D). Furthermore, even though $^{13}$C incorporation from glucose to the ribose moiety of nucleotide and nucleoside was lower in naive cells (Figure 3D; Figure S2C), it was not caused by the pentose phosphate pathway, as metabolites in this pathway were not differentially measured in either state (Figure S2D). On the other hand, $^{13}$C incorporation from serine to nucleobase moieties was lower in naive cells (Figure S2E), indicating one-carbon metabolism through N$^{10}$-formyl-tetrahydrofolate (f-THF) accounted for reduced nucleotide biosynthesis. $^{13}$C incorporation from glutamine to nucleobase moieties through aspartate was also lower in naive cells (Figure 3E), which could be attributed to lower glutamine incorporation into the TCA cycle (Figure S2B). In summary, these data suggest that the one-carbon metabolism pathway branching from glycolysis is more active in the primed state.

In addition, mRNA transcripts of genes associated with the reactions predicted by the model to be differentially sensitive were also significantly more likely to be differentially expressed ($p = 0.003$, t test; Figure S1G). Although pathway enrichment of transcriptomics data did not identify differential activity of these pathways, analysis of specific transcripts prioritized by the model revealed lower expression of folate pathway genes Mtfhd1, Mthfd2, Mthfd2l, Mthfs, Mthfr, and Dhdr, and nucleotide biosynthesis pathway genes Adsl, Atic, and Gart, in the naive state cells (Figure S1E). Further assessment of in vivo embryonic day 4.5 (E4.5) pre-implantation epiblast (naive) versus E5.5 post-implantation epiblast (primed) also showed lower expression of Phgdh, Psat, Shmt1/2 (serine metabolism), Mtfhd1l, Mthfs, Mthfd2, Dhdr (folate pathway), Matl2a, Matl1, Cbs (methionine/cysteine metabolism), and Adsl, Gart, Tym, Atic, Paic (nucleotide biosynthesis) (Figure S1H). A dormant state of diapause characteristic of the embryonic epiblast showed the lowest expression of most genes listed above (Figure S1H) (Boroviak et al., 2015). Taken together, these data suggest that mouse naive cell pluripotency is associated with lower one-carbon metabolism and glutamine metabolism required for nucleotide biosynthesis.

**Tracking Metabolic Network Rewiring by Lin28**

We next examined the metabolic state of naive induced pluripotent stem cells (iPSCs) cultured in LIF/2i condition depleted of Lin28. Lin28 knockout cells show elevated naive state features compared with wild-type cells; they have defects in priming and do not grow well in the primed condition (FGF2/Activin). Indeed, our prior analysis has indicated that Lin28-deficient cells represent a surrogate for the naive-state metabotype (Zhang et al., 2016).

With the same approach using time-course metabolomics data and reaction deletion analysis, we discovered similar metabolic differences in one-carbon, oxidative phosphorylation and nucleotide metabolism between Lin28 knockout and wild-type cells (Tables S2B and S3B). Lin28 knockout cells were more sensitive to deletion of reactions in the mitochondria, and less sensitive to deletion of one-carbon metabolism reactions, reminiscent of the naive state compared with the primed state (Figures 4A–4C). The comparison between Lin28 knockout cells and wild-type cells (both in 2i) differed from the comparison between naive cells and primed cells in the activity of the pentose phosphate pathway (Figure 4D). This analysis also revealed the importance of the nucleotide salvage pathway in Lin28 knockout cells, with reactions in purine salvage and purine cycle pathways such as Hgprt, predicted to have higher flux in Lin28 knockout cells (Figures 4A and 4B). This could explain a previous report that these cells are deficient in de novo nucleotide biosynthesis and addition of nucleotides in the media rescues this phenotype (Zhang et al., 2016). Time-course analysis of $^{13}$C-glucose incorporation also showed lower flux to nucleotides in Lin28 knockout cells (Figure 4C; Figure S1C). In addition, $^{13}$C-glutamine tracing revealed knockout cells had lower flux to nucleotide, similar to the 2i naive state (Figure 4C; Figure S1D).

We next quantified the overall similarity of the 3,744 reaction sensitivities between naive, primed, lin28 knockout, and wild-type cells using unsupervised clustering and multi-dimensional scaling (MDS) analysis (Figures 4D and 4E). Hierarchical clustering analysis revealed that most reactions that show differential sensitivity between cell states share the same direction of change in both naive versus primed cells and ∆Lin28 versus wild-type cells, suggesting broad similarities between the two scenarios (Figure 4D). A small subset of reactions showed the opposite direction of change in ∆Lin28 versus wild-type compared to naive versus primed cells, suggesting that these two scenarios are similar, but not identical (Figure 4D). MDS analysis, which visualizes the entire set of variation in a data in three dimensions, confirmed that knockout of Lin28 moves the cellular metabolic network of iPS cells closer to the naive ESC state (Figure 4E).

**Simulating Impact of Metabolic Pathways on Histone Methylation**

The one-carbon metabolism pathway is required for histone methylation in ESCs through modulating the SAM/S-adenosyl-L-homocysteine (SAH) ratio (Shyh-Chang et al., 2013) (Figure 3C). To assess the impact of the naive and primed metabolic network states on histone methylation, we simulated the effect of metabolic reaction deletions on SAM flux and subsequently histone methylation under the assumption that SAM is a limiting factor.
Figure 4. Mapping the Metabolic Differences of Lin28 Knockout versus Wild-Type PSCs Using the Computational Model of Metabolism

(A) The histogram shows the distribution of growth rates after in silico deletion of all the metabolic reactions. The horizontal x axis shows the relative effect of each gene knockout on the growth rate of Lin28 knockout versus wild-type cells (selectivity). The y axis shows the total number of reactions in each bin. Top pathways with multiple differential reactions are highlighted; see Table S2 for the full list of reactions (naive, orange; primed, purple).

(B) Schematic diagram of the metabolic pathways that are predicted to differ between the two states (wild-type, orange labels; D Lin28, pink labels) as identified by the model. The thickness of the reaction label border is proportional to the extent of the differential activity of the reactions between the two states.

(C) (Left) 13C tracing of glucose revealed that metabolites downstream of the PHGDH, MTR, MTHFR, UMPS, and DHODH pathways were differentially labeled between the two conditions, suggesting re-wiring of the metabolic network of these two states. (Right) 13C tracing of glutamine revealed that glutamine flux is routed toward nucleotide metabolism in wild-type cells. Metabolites that are differentially labeled (p < 0.05) are displayed in both heatmaps. n = 3 for each time point. The heatmaps show the total isotopomers labeled at different time points for each metabolite after z transformation (Supplemental Experimental Procedures).

(D) Unsupervised hierarchical clustering analysis of the entire set of 3,744 reactions and their differential activity between naive and primed cells, and lin28 knockout versus wild-type cells. Reactions are clustered based on extent of similarity between the two comparisons (naive versus primed and D Lin28 versus WT). The majority of reactions do not have differential impact in any state (yellow color in both rows). The bottom panel shows the cluster of 220 reactions that exhibit differential sensitivity between different cell states. Overall, even reactions that show differential sensitivity have the same direction of change in both naive versus primed cells and D Lin28 versus WT cells, suggesting broad similarities between the different states (96.6% similarity). A small subset of reactions (N = 33) shows opposite differences in D Lin28 versus WT compared to naive versus primed cells; reactions with the strongest differences are highlighted.

(E) MDS analysis captures the overall metabolic differences between cell states and confirmed that knockout of Lin28 moves the cellular metabolic network closer to naive cell metabolism.
Figure 5. Prediction and Validation of Intracellular SAM Levels and Global Histone Methylation

(A) Using the metabolic model, we predicted metabolic changes that can impact SAM flux and subsequently histone methylation. The histogram shows the distribution of SAM flux toward histone methylation after deletion of all the metabolic reactions. The horizontal x axis shows the relative effect of each knockout on SAM flux in the naive state versus the primed state (selectivity). The y axis shows the total number of reactions in each bin. Metabolic reactions that showed the...
metabolite for methylation. Deletion of reactions in one-carbon metabolism and nucleotide metabolism preferentially affected SAM production in the primed state over the naive state (Figure 5A; Table S4). We experimentally validated this prediction by treating cells with 3-deazaadenosine (3-DZA), a SAH hydrolase inhibitor that decreases the SAM/SAH ratio, and found that it more profoundly affected H3K4, H3K9, and H3K27 trimethylation in the primed state relative to the naive state (Figure 5B). Furthermore, treatment with 3-DZA affected wild-type H3K9 trimethylation more than Lin28 knockout, consistent with the role of Lin28 in the primed state (Figure 5C). FBA also revealed higher flux through SAM synthesis reactions in the primed state, suggesting that the primed state relies more on one-carbon metabolism for production of SAM (Table S4). Consistent with the in silico prediction, the primed metabolic state has higher 13C incorporation from serine toward SAM than the naive state based on tracing 13C-labeled serine (Figure 5E). Furthermore, histone methyltransferases and demethylases are not differentially expressed in these two states; hence altered SAM metabolism can explain the observed differential histone methylation (Figure 5D).

**DISCUSSION**

Metabolism influences gene-regulatory networks that govern cell fate choices (Ryall et al., 2015; Zhang et al., 2012), in both pluripotent stem cells (Carey et al., 2015) and various types of adult progenitor/stem cells (Peng et al., 2016; Yang et al., 2016). Given the development potential of naive ESCs compared to primed ESCs, it is critical to understand the metabolic differences between these pluripotent cell fates. Analysis using our dynamic metabolic modeling framework revealed three insights on pluripotent stem cell metabolism. First, previous studies showed higher mitochondria function in mouse ESCs, in comparison to EpiSC or human ESCs (Zhou et al., 2012). Furthermore, human naive-like cells have higher nicotinamide N-methyltransferase (NNMT) activity that consume SAM (Sperber et al., 2015). Here, we comprehensively characterized the metabolic network of two bona fide pluripotent states in mouse ESCs and discovered that, during priming, the entire one-carbon metabolic pathway is activated, including upstream serine production from glucose and downstream SAM and nucleotide production through folate cycle and methionine cycle. This makes mouse primed ESCs more vulnerable to folate metabolism and SAM/SAH repression. Interestingly, this effect is compartment specific, with folate inhibition in mitochondria more sensitive in naive cells, whereas cytoplasmic inhibition is more sensitive in primed cells. Our systems approach has thus uncovered a transition of one-carbon metabolism between two clearly defined naive and primed states.

Second, a previous report found that deleting Lin28 changes mouse ESCs to a naive-like state. At the network level, broad similarities between 2i naive and Lin28 knockout cells were observed. However, unique differences in the pentose phosphate pathway identified by our analysis suggest that these states are not identical (Figure 4D). Third, we found that increased flux through one-carbon pathway leads to higher flux toward SAM and histone methylation in primed state, and inhibiting these pathways reduces methylation disproportionately in primed state. A previous report connected NNMT activity to SAM consumption (Sperber et al., 2015). Our approach revealed that serine flux and the one-carbon metabolic pathway contribute to SAM production and histone methylation. Consistent with a low SAM pool, naive mouse cells have hypomethylated DNA and histone, and are tolerant to loss of epigenetic regulators such as DNMT1 and polycomb protein EED, in contrast to primed cells (Weinberger et al., 2016). Because recent studies also implied that similar metabolic changes in one-carbon and oxidative metabolism may occur in converted human naive-like and primed pluripotent stem cells (PSCs) (Sperber et al., 2015; Takashima et al., 2014), whether the converted naive-like cells become more tolerant to DNMT1 depletion and/or to SAM/SAH repression, merits careful investigation to substantiate the identity of those cells.

Pervasive histone methylation is a key feature of primed cells and our findings directly tie into the biology of primed pluripotency. Our model also suggests that hypomethylation in naive cells is likely due to the lack of one-carbon donor, SAM. Differentiation potential (governed by epigenetic regulations) and proliferation are the two most important cellular phenotypes that distinguish naive and primed pluripotent states. The mechanism of regulation of histone methylation so far is unknown because histone methyltransferases and demethylases are not differentially expressed in these two states (Figure 5D). Our study provides evidence that altered one-carbon and SAM metabolism account for the differential histone methylation in these two states, which in turn determines expression of various pluripotent and developmental genes.

Our work suggests an interplay between mitochondrial respiratory metabolism and one-carbon metabolism during ESC pluripotent state transitions, pointing to an unappreciated role of elevated mitochondrial function in naive state pluripotency. Similar coupling between one-carbon and respiratory metabolism has been recently reported in cancer cell lines (Bao et al., 2016). Furthermore, many of the enzymes predicted to be differentially active between naive and primed ESCs, such...
as PHGDH, MTHFR, and other one-carbon metabolism pathway constituents, have also been associated with tumor initiation, progression and metastases (Locasale, 2013; Piskounova et al., 2013). Comparison of metabolic genes that are sensitive in NCI-60 cancer cell lines (Figure 2) with corresponding sensitivity in naive and primed cells (Figure 3) revealed that the metabolic state of many NCI-60 cell lines is surprisingly similar to naive cell metabolism but not to primed cell metabolism. This intriguing preliminary observation needs to be explored further in a future study. Our findings may provide a better understanding of metabolic rewiring during tumorigenesis as well.

Our systems biology approach overcomes a significant challenge in genome-scale modeling by inferring the metabolic state of mammalian cells directly from time-course metabolomics data. The power of our approach is that it can infer the activity of thousands of reactions based on the measurement of a few hundred metabolites. Interpreting metabolomics data is quite challenging due to the highly interconnected nature of metabolic networks. The metabolites in our time-course data were involved in 834 metabolic reactions in the model, even after excluding ubiquitous metabolites such as ATP and NADH. Hence, it is difficult to manually infer the differentially active reactions from the metabolomics data. By overlaying these data onto the model, we identified a small, prioritized set of reactions that are differentially active between these two states.

Using our approach, we identified compartment-specific differences in folate metabolism, which we validated using cytosolic and mitochondrial folate inhibitors (Figure 3F). This is significant given that information on cellular compartments is lost during metabolomics measurement. Furthermore, the model correctly identified differences in oxidative phosphorylation pathway despite there being no direct measurement of this pathway’s activity. These results highlight the power of our systems approach to infer metabolic changes based on network topology that goes beyond traditional analysis of metabolomics data.

Our computational approach still has some limitations despite these numerous advances highlighted earlier. First, the algorithm requires the levels of numerous metabolites (~100) to infer the activity of reactions in the metabolic model. The accuracy of the approach dropped significantly with data from fewer than 65 metabolites (i.e., 33% of the data removed; Figure 2F). Future studies should reveal which metabolites provide the most predictive power for these models. Second, existing genome-scale metabolic models lack detailed mapping of atom transfer from substrates to products, thus preventing the incorporation of isotopomer labeling distribution from 13C tracing data. 13C flux tracing studies currently use significantly smaller metabolic models with atomic mapping to predict fluxes. Our systems-scale approach can complement such traditional 13C analysis approaches by making coarse-grained predictions for a larger set of reactions. Third, our approach is restricted to predicting the impact of complete enzyme inhibition or gene knockouts on the metabolic network. As enzyme kinetic parameters become available, it should be possible to simulate the systems-level impact of nuanced changes in enzyme levels.

Looking forward, a systems-level understanding of stem cell metabolism could allow us to rationally manipulate specific metabolic modules to facilitate differentiation and aid cell-fate engineering. Our approach could be readily extended to understand metabolic mechanisms underlying other cell-fate changes and could open new avenues for computational identification of metabolic vulnerabilities of cancer cells and other complex metabolic disorders.

**EXPERIMENTAL PROCEDURES**

**Genome-Scale Metabolic Network Modeling**

We used the Duarte et al. (2007) model of human metabolism for all the metabolic modeling in this study as it has been widely applied in literature for understanding cellular metabolism of various normal and disease states. We also used the human Recon 2.0 model and the genome-scale model of mouse metabolism as further validation. The mouse model was derived based on homology with the Duarte et al. human model. Using these models leads to the identification of similar set of genes and pathways that were differentially active between naive and primed states (Table S2C).

We used FBA to determine the optimal metabolic state that satisfies the growth objective and the metabolomics constraints. Mathematically, in FBA, we solve the optimization problem to identify a metabolic state (v, a vector of reaction fluxes) that maximizes a specific cellular objective, such as the biomass production or ATP synthesis rate, while satisfying thermodynamic and mass balance constraints. FBA is formulated as follows:

\[
\text{Minimize } S \cdot v = b, \\
\text{subject to } \begin{cases} 
S \cdot v + \alpha \cdot \beta = \epsilon \end{cases}
\]

where \( S \) is the stoichiometric matrix with \( S_{ij} \) representing the stoichiometric coefficient of metabolite \( i \) in reaction \( j \), \( v \) is the flux vector, and \( b \) represents the rate of change of metabolites \((dM/dt)\). In traditional FBA, the value of R.H.S vector \( b \) is assumed to be zero to represent steady state. We set this value proportional to the measured rate of change from time-course metabolomics data. For each dynamic metabolite, we include a flux activity coefficient, whose value and sign determines whether there is increased or decreased flux activity involving the metabolite.

\[
S \cdot v = \epsilon
\]

where \( \epsilon \) is a vector of flux activity coefficient values (\( \neq 0 \) for dynamic metabolites).

The value of the flux activity coefficients is directly determined based on the rate of change of metabolite levels over time. Because subcellular compartment-specific information is lost during metabolomics measurement, we assumed that the measured metabolites represent the sum total of those in the cytoplasm, nucleus, and the mitochondrial compartments in the model. In addition, some metabolite changes might represent noise in the data, and these constraints would be predicted to be infeasible by the model. To account for these issues, we implemented an approach, defined below, that identifies a metabolic state that best fits the entire set of metabolomics constraints simultaneously:

\[
S \cdot v + \alpha \cdot \beta = \epsilon
\]

\[
\text{Minimize } (\alpha + \beta)
\]

Alpha and beta are positive vectors that represent deviation from the measured experimental data (Supplemental Experimental Procedures). The entire sequence of steps in the dynamic modeling approach is described in the pseudo-code provided in the supplement (Supplemental Experimental Procedures).

The approach is robust to relative weights for metabolomics data and growth objective (as defined by the kappa parameter), and also to noise in metabolomics data (Figure 1E). Using a simple model that takes as input only the
direction of change of metabolites (i.e., accumulating or depleting) without the magnitude also led to qualitatively similar set of predictions for naive and primed states (Figure 1F).

In addition to FBA, we used FVA to determine the range of feasible fluxes for each reaction. Reactions that had either higher minimal flux or both higher minimal and maximal flux compared to the second condition (naive or primed state) were considered to have higher activity in that condition.

Software and Data Availability

The MATLAB implementation of the algorithm and associated datasets are publicly available at the Synapse bioinformatics software repository (https://www.synapse.org/; synapse ID: syn7253624). Examples and instructions for running the analysis are also provided. The entire metabolomics data for naive, primed, and Δlin28 cells are also provided as a supplemental dataset (Tables S5 and S6). The optimization problem was solved using the Gurobi mathematical programming solver.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, two figures, and six tables and can be found with this article online at https://doi.org/10.1016/j.celrep.2017.07.048.

AUTHOR CONTRIBUTIONS

J.Z. and S.C. conceived, designed, and performed research and wrote the manuscript with input from G.Q.D. and J.J.C. S.C. performed all the computational analysis, and J.Z. conducted all the experimental assays with help from Z.S., L.Z., and Y.-C.H. G.Q.D. and J.J.C. contributed to study design and supervised the study. J.M.A. performed metabolomics, and C.A.R. and H.L. performed transcriptomics analysis.

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