α-Ketoglutarate Accelerates the Initial Differentiation of Primed Human Pluripotent Stem Cells

**Highlights**
- hPSCs produce TCA cycle metabolites despite low OXPHOS
- α-ketoglutarate (αKG) promotes early differentiation of hPSCs
- Accumulation of succinate or depletion of αKG delays differentiation of hPSCs
- αKG/succinate alters histone methylation corresponding to differentiation kinetics

**In Brief**
α-ketoglutarate (αKG) is an important cofactor for demethylation reactions that helps to maintain naive pluripotent stem cells. TeSlaa et al. show that at later stages of pluripotency, αKG can promote early differentiation, highlighting that the cellular context and potentially the stage of cellular maturity can alter the effect of αKG.

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**SUMMARY**

Pluripotent stem cells (PSCs) can self-renew or differentiate from naive or more differentiated, primed, pluripotent states established by specific culture conditions. Increased intracellular \( \alpha \)-ketoglutarate (\( \alpha \)KG) was shown to favor self-renewal in naive mouse embryonic stem cells (mESCs). The effect of \( \alpha \)KG or \( \alpha \)KG/succinate levels on differentiation from primed human PSCs (hPSCs) or mouse epiblast stem cells (EpiSCs) remains unknown. We examined primed hPSCs and EpiSCs and show that increased \( \alpha \)KG or \( \alpha \)KG-to-succinate ratios accelerate, and elevated succinate levels delay, primed PSC differentiation. \( \alpha \)KG has been shown to inhibit the mitochondrial ATP synthase and to regulate epigenome-modifying dioxygenase enzymes. Mitochondrial uncoupling did not impede \( \alpha \)KG-accelerated primed PSC differentiation. Instead, \( \alpha \)KG induced, and succinate impaired, global histone and DNA demethylation in primed PSCs. The data support \( \alpha \)KG promotion of self-renewal or differentiation depending on the pluripotent state.

**INTRODUCTION**

Human pluripotent stem cells (hPSCs) may self-renew or differentiate into all three germ layers (Takahashi et al., 2007; Thomson et al., 1998), but use in regenerative medicine is limited by generally inefficient differentiation strategies (Glanpain et al., 2012). During in vitro differentiation, hPSCs undergo a metabolic shift that increases respiration (oxidative phosphorylation [OXPHOS]) and reduces glycolysis, with inhibition of this transition impeding differentiation (Moussaieff et al., 2015; Zhang et al., 2011; Zhou et al., 2012). Despite the importance of this metabolic shift, differentiation protocols have focused on manipulating key signaling pathways and have overlooked metabolic contributions.

\( \alpha \)-ketoglutarate (\( \alpha \)KG), a tricarboxylic acid (TCA) cycle metabolite, is a cofactor for \( \alpha \)KG-dependent dioxygenase enzymes, which include JmjC-domain containing histone demethylases (JHDMS) and ten-eleven translocation (TET) enzymes (Kaelin and McKnight, 2013). \( \alpha \)KG can also bind and block the mitochondrial ATP synthase and inhibit mechanistic target of rapamycin (mTOR) signaling (Chin et al., 2014). Addition of cell-permeable dimethyl-\( \alpha \)KG (dm-\( \alpha \)KG) to culture media enhances self-renewal and inhibits the differentiation of naive-state mouse embryonic stem cells (mESCs) likely by promoting histone and DNA demethylation (Carey et al., 2015). The hPSCs grown in standard conditions are in a primed, or more developmentally mature, pluripotent state, similar to post-implantation mouse epiblast stem cells (EpiSCs) (Greber et al., 2010; James et al., 2005; Tesar et al., 2007). A role for \( \alpha \)KG in primed mouse or hPSCs has not been explored.

Naive and primed pluripotent stem cells (PSCs) show many molecular differences including self-renewing conditions, epigenetic states, and metabolism (Greber et al., 2010; Leitch et al., 2013; Marks et al., 2012; Ware et al., 2014; Zhou et al., 2012). A consensus naive state for hPSCs, however, remains somewhat elusive. Culture conditions that establish naive-like hPSCs yield slightly different transcriptional profiles (Huang et al., 2014). Uncertainty about whether naive hPSCs offer differentiation advantages over traditional primed hPSCs (Pera, 2014)
emphasizes the remaining importance of primed hPSCs as options for potential clinical applications.

Metabolites other than αKG have been shown to play a role in PSC self-renewal and differentiation. Removal of methionine, which provides methyl groups for DNA and histone methylation, potentiates PSC differentiation (Shiraki et al., 2014). Increased acetyl-coenzyme A delays PSC differentiation and histone acetylation and maintains expression of OCT4 (Moussiaff et al., 2015). Oxygen levels can enhance reprogramming to pluripotency or differentiation of hPSCs, depending on environmental context (Mathieu et al., 2014; Xie et al., 2014). The mESCs are dependent on threonine catabolism for histone and DNA methylation (Shyh-Chang et al., 2013; Wang et al., 2009). Here, we investigate the role for αKG during primed PSC differentiation.

RESULTS

TCA Cycle Metabolite Production in hPSCs

Respiration is reduced in hPSCs compared to their differentiated counterparts, suggesting that TCA cycle metabolite production could be low (Zhang et al., 2011; Zhou et al., 2012). To examine the TCA cycle, stable isotope labeling experiments were performed in Essential 8 (E8) media promoting self-renewal or Essential 6 (E6) media encouraging differentiation (Figures S1A and S1B). Using the E8 or E6 system, culture media differ by only two factors that are excluded from the E6 media, basic fibroblast growth factor (bFGF) and transforming growth factor β (TGF-β), ensuring differences in metabolism are due to different cell states. A shift in the oxygen consumption rate (OCR)-to-extracellular acidification rate (ECAR) ratio confirmed a shift toward OXPHOS with E6 media differentiation (Figure 1A). Furthermore, glutamine withdrawal reduced oxygen consumption, implicating glutamine as a TCA cycle fuel in hPSCs (Figure 1B).

Despite a low OCR-to-ECAR ratio, hPSCs showed a robust contribution of [U-13C] glucose into TCA cycle metabolites (Figure 1C). Glutamine withdrawal increased the glucose contribution to TCA cycle metabolites αKG, succinate, and malate in E8, but not in E6, culture conditions (Figure 1C). The mass isotopologue distribution (MID) of citrate indicates the contribution of [U-13C] glucose to the TCA cycle, with m+2 and m+3 isotopologues indicating 13C glucose carbons that have cycled through one or two turns (Figure 1D). In glutamine-sufficient conditions, no differences in the citrate MID in E8 and E6 cultures were detected (Figure 1E), but glutamine withdrawal for 18 hr resulted in an increase in m+4 and m+6 citrate isotopologues in undifferentiated human embryonic stem cells (hESCs) (E8) (Figure 1F). Thus, glucose-derived carbons are retained through one or two turns of the TCA cycle in the absence of glutamine in self-renewing hPSCs. Consistent with this result, glutamine withdrawal led to a decrease in unlabeled αKG (m+0) in E8 conditions (Figure S1C).

The MID of citrate with [U-13C] glutamine indicates the amount of citrate derived from glutamine after one turn (m+4) or two turns (m+2) of the TCA cycle (Figure 1G). Increased m+4 citrate was detected in E8 compared to E6 conditions, suggesting a lower contribution of glutamine to the TCA cycle in E6 differentiated hPSCs (Figure 1H). However, 40% of glutamate, which is generated directly from glutamine, was unlabeled in E6 conditions (Figure S1D). Measurement of extracellular glutamate levels revealed net uptake of glutamate by cells cultured in E6 medium (Figure 1I). Detection of [U-13C] glutamate uptake confirmed these results (Figure 1J), with increased conversion of glutamate into αKG occurring in differentiated cells (E6) only in the absence of glutamine (Figure 1K). The data suggest that glucose and glutamine are the major contributors to the TCA cycle in hPSCs; other metabolites, such as glutamate, fuel the TCA cycle in early differentiated hPSCs.

In proliferative cells, αKG-producing transaminases (TAs), which transfer amine groups from glutamate to α-keto acids to form amino acids (Figure S1E), have high activity (Coloff et al., 2016). Glutamine inclusion in hPSC culture medium, which provides glutamate for TAs, increases αKG, alanine, and aspartate levels, products of these TAs (Figures S1F and S1G). To further study TA activity, [15N2] glutamine was used to quantify the transfer of 15N from glutamate to amino acids (Figure S1E), and 15N was detected in alanine, aspartate, isoleucine, serine, and methionine (Figure 1L). Expression of glutamic-pyruvic transaminases (GPTs) and glutamic-oxaloacetic transaminases (GOTs) was confirmed in hPSCs, verifying their contribution to αKG production (Figure S1H). Robust αKG production in primed hPSCs prompted studies into a role for αKG in PSC differentiation.

αKG Accelerates Multi-lineage Primed PSC Differentiation

Neuroectoderm (NE) differentiation was induced in primed H1, H9, UCLAl1, and HIPS2 hPSCs by dual SMAD inhibition (Chambers et al., 2009). The dm-αKG significantly increased the percentage of PAX6, an essential transcription factor for NE specification in humans (Zhang et al., 2010), expressing cells by day 4 of differentiation (Figures 2A–2C). MAP2C- and NESTIN-positive cells modestly increased with αKG treatment (Figures 2C and S2A). To determine whether this effect was lineage specific, endoderm differentiation was induced by high-concentration activin A exposure (D’Amour et al., 2009). On day 2, αKG significantly increased the percentage of H9 cells expressing SOX17, a definitive endoderm transcription factor (Figures S2B and S2C). Combined, the data support that αKG accelerates the early differentiation of multiple hPSC germlineages.

NE differentiation was examined with added αKG, which unlike dm-αKG, is dependent on membrane transporters for uptake, and resulted in an increase in PAX6-positive cells (Figures 2D and 2F). In contrast to dm-αKG, which shows a dose-dependent increase in cell differentiation, αKG levels beyond 4 mM did not further stimulate NE differentiation (Figures 2D–2G). Added αKG and dm-αKG each increased intracellular αKG and TCA cycle metabolite levels, although only 12 mM dm-αKG reached significance (Figures 2H and 2I).

Consistent with prior results showing that dm-αKG supports naive mESC self-renewal (Carey et al., 2015), an increase in alkaline phosphatase staining was detected in naive mESCs incubated with dm-αKG during 48 hr of leukemia inhibitory factor (LIF) withdrawal (Figure S2D). Because dm-αKG accelerates primed hPSC differentiation, we examined the role for αKG in primed mouse PSCs, or EpiSCs. Addition of dm-αKG to EpiSCs induced to differentiate by withdrawal of bFGF and activin A
Figure 1. Production of TCA Cycle Metabolites in hPSCs

(A) Ratio of OCR to ECAR in H9 hESCs cultured in medium containing (E8) or lacking (E6) bFGF and TGF-β.
(B) OCR quantified in H9 hESCs grown in media containing glutamine or with glutamine removed for 1 or 18 hr.
(C) Fractional contribution of [13C]-labeled metabolites from [U-13C] glucose after 18 hr, quantified by UHPLC-MS.
(D) Schematic illustrating how the MID of citrate from [U-13C] glucose reveals the contribution of glucose-labeled metabolites through multiple turns of the TCA cycle.
(E) MID of citrate in H9 hESCs from [U-13C] glucose in conditions containing glutamine (E) or lacking glutamine (F).
(G) Schematic of [U-13C] glutamine labeling of the TCA cycle. The m+5 isotopologue of citrate can be produced by reductive carboxylation of glutamine (blue). The m+4 and m+2 isotopologues contain carbons derived from glutamine after one and two turns of the TCA cycle, respectively.
(H) MID of citrate from [U-13C] glutamine in H9 hESCs cultured in E8 or E6 medium.
(I) Measurement of glutamate uptake from culture medium in H9 hESCs maintained in E8 medium or differentiated in E8 medium. The m+5 isotopologue of glutamate in H9 hESCs, indicating increased uptake of [U-13C] glutamate from the culture medium in differentiated cells (E6) grown with or without glutamine.
(J) The m+5 isotopologue of αKG in H9 hESCs grown in [U-13C] glutamate.
(K) The m+1 isotopologue of listed amino acids in H9 hESCs grown with [15N2] glutamine, reflecting the activities of multiple αKG-producing TAs.

Data represent mean ± SD of at least three biological replicates. *p ≤ 0.05; **p ≤ 0.01; ***p ≤ 0.001; ****p ≤ 0.0001. The p values were determined by an unpaired two-tailed Student’s t test (A and I) or by one-way ANOVA (B) or two-way ANOVA (C, E, F, H, J, and K) with correction for multiple comparisons.
Figure 2. αKG Accelerates Differentiation of Primed PSCs

(A) Flow cytometry analysis of PAX6 transcription factor expression in UCLA1 hESCs encouraged to differentiate into NE at 4 or 6 days. The figure shows 12 mM dm-αKG incubated traces (blue) and control traces (black).

(B) Flow cytometry quantification of the percentage of PAX6-expressing cells at 4 days of NE differentiation for H9, UCLA1, and H1 hESCs and HIPS2 human-induced pluripotent stem cells (hiPSCs). H9, UCLA1, and H1 cells were incubated with (blue) or without (black) 12 mM dm-αKG, and HIPS2 hPSCs were incubated with (blue) or without (black) 6 mM dm-αKG. Lines connect pairs of independent biological replicates.

(C) Immunoblot of ectoderm markers PAX6 and MAP2C after 2, 4, and 6 days of NE differentiation of H9 hESCs.

(D and E) Flow cytometry of PAX6 expression in H9 hESCs on day 4 of NE differentiation, with indicated amounts of αKG (D) or dm-αKG (E) added to the culture medium.

(F and G) Percentage of H9 hESC cells positive for PAX6 expression on day 4 of differentiation plotted against the concentration of added αKG (F) or dm-αKG (G). Lines connect independent biological replicates.

(H and I) UHPLC-MS quantification of fold change of αKG levels (H) and other TCA cycle metabolite levels (I) in H9 hESCs incubated with the listed concentrations of αKG or dm-αKG. Error bars represent SEM of three biological replicates.

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(Greber et al., 2010) accelerated the rate of Oct4 inactivation (Figures 2J, 2K, and S2E–S2G). Therefore, αKG can accelerate differentiation of both mouse and human primed PSCs.

αKG has been shown to bind to and inhibit the mitochondrial ATP synthase subunit β1, leading to mTOR inhibition (Chin et al., 2014). To determine whether ATP synthase inhibition contributes to αKG-accelerated primed PSC differentiation, an OXPHOS uncoupling agent, carbonyl cyanide 4-(trifluoromethoxy)-phenylhydrazone (FCCP), was used during NE differentiation. FCCP inhibits ATP production as a mitochondrial inner membrane protonophore that dissipates the H+ ion electrochemical gradient that runs the ATP synthase. That dm-αKG accelerated NE differentiation in H9 hESCs incubated with FCCP (Figures S2H and S2I) suggested an alternative mechanism. Furthermore, hPSCs treated with an inhibitor of ATP synthase, oligomycin, during NE differentiation showed almost no PAX6 expression after 4 days, despite the addition of pyruvate and uridine to promote cell survival (Figures S2J and S2K) (Birsoy et al., 2015; Sullivan et al., 2015). Because ATP synthase inhibition delays or inhibits NE differentiation, which opposes the accelerating effect of dm-αKG, we conclude that αKG does not accelerate differentiation of primed PSCs through inhibition of ATP synthase.

### Succinate Accumulation Delays hPSC Differentiation

A second potential mechanism for αKG-accelerated hPSC differentiation is stimulation of epigenome-modifying dioxygenases. In this event, an αKG-dependent dioxygenase competitive inhibitor, such as succinate (Xiao et al., 2012), would impair differentiation. Cell-permeable succinate, dimethyl succinate (dms), resulted in a decreased percentage of PAX6-expressing hPSCs during NE differentiation compared to hPSCs incubated with dm-αKG alone (Figures S2L and S2M). Inhibition of succinate dehydrogenase A (SDHA), which converts succinate to fumarate either with a chemical inhibitor, 3-nitropironic acid (NPA), or with small hairpin RNA (shRNA), should also cause succinate accumulation (Figure 3A). NPA treatment decreased the αKG-to-succinate ratio 34-fold, with a 14-fold mean increase in succinate and 2.4-fold mean decrease in αKG (Figures 3B and S3A). NPA delayed NE-specifying PAX6 expression (Figures 3C and S3B–S3D), and the loss of pluripotency marker SSEA3 in hPSCs differentiated into NE (Figure S3E). Knockdown of SDHA, confirmed by immunoblot and reduced OCR (Figures S3F–S3H), also delayed PAX6 and MAP2B expression during NE differentiation, which was rescued by dm-αKG (Figures 3D, 3E, and S3I).

We further assessed the role of succinate in differentiation by embryoid body (EB) formation, which contrasts with lineage-directed differentiation by removal of bFGF rather than by addition of supplements or inhibitors. OCT4 was almost eliminated in EBs expressing a scrambled shRNA, whereas shRNA targeting SDHA maintained elevated OCT4 expression (Figures S3J and S3K). Validating these results, inhibition of SDHA with NPA significantly impaired OCT4 repression in EB differentiation compared to control (Figure S3L).

### TA Inhibition Delays hPSC Differentiation

To decrease αKG levels, chemical inhibitors of αKG-producing TAs were used (Figure 3A). Aminoacycatic acid (AOA), a pan-TA inhibitor, reduced 15N transfer from [15N] glutamine to alanine, aspartate, isoleucine, serine, and methionine in hPSCs (Figure S3M). L-cycloserine (cyclo), a GPT inhibitor, decreased 15N transfer to alanine in hPSCs treated with cyclo (Figure S3M). Both inhibitors caused a significant decrease in αKG levels and other TCA cycle metabolite levels, but they had no effect on basal respiration (Figures 3F, S3N, and S3O). AOA impaired PAX6 and MAP2B activation during NE differentiation (Figures 3G, 3H, and S3P). A mixture of non-essential amino acids (NEAAs), including alanine and aspartate, had no effect on AOA treatment, whereas supplementation with dm-αKG rescued the block in differentiation caused by AOA (Figures 3I and 3J). Supplementation with dm-αKG restored MAP2B levels in cyclo-treated cells (Figures 3I and 3J). Therefore, decreased αKG levels cause a delay or inhibition of directed differentiation.

To determine whether TCA cycle flux affects differentiation, dichloroacetate (DCA), an inhibitor of pyruvate dehydrogenase kinase (PDK), was used. DPK inhibits pyruvate dehydrogenase (PDH) activity. Therefore, DCA increases PDH activity and glucose flux into the TCA cycle, which elevates electron transport chain (ETC) activity (Figure S4A). Low levels of DCA increased PAX6-positive cells on day 4 of NE differentiation, whereas higher DCA levels had no effect (Figures S4B and S4C). Furthermore, dm-αKG enhanced differentiation at 0 and 3 mM DCA but not at 1 mM (Figures S4B and S4C). These results do not support changes in TCA cycle flux as a mechanism for αKG in accelerated hPSC differentiation. Rather, the data indicate that succinate delays and αKG promotes the initial differentiation of primed PSCs, most likely through actions on αKG-dependent dioxygenases.

### αKG/Succinate Regulates the Epigenome of Differentiating hPSCs

To evaluate the role of αKG-dependent dioxygenases during hPSC differentiation, an inhibitory αKG mimetic, dimethyloxalylglycine (dmog) was used. Exposure to dmog inhibited PAX6 expression at concentrations that did not affect cell number (Figures 4A, 4B, and S4D). To assess the effect of the αKG on TET enzymes, dot blots were performed and levels of 5-hydroxymethylcytosine (5hmC) and 5-methylcytosine (5mc) in DNA were measured from H9 and UCLA1 hESCs after 4 days of NE differentiation. Although dm-αKG exposure caused a 2-fold increase in the 5hmC-to-5mc ratio, NPA caused a significant decrease in this ratio (Figures 4C, 4D, and S4E). The data suggest a significant role for TET enzymes in αKG-accelerated differentiation of primed hPSCs. To evaluate αKG regulation of JHDMs, histone
Figure 3. Decrease in the αKG-to-Succinate Ratio Delays hPSC Differentiation

(A) Inhibition or depletion of SDHA causes succinate accumulation. Inhibition of TAs, such as GPT or GOT, results in a decrease in αKG levels. Both of these manipulations cause a decrease in the αKG-to-succinate ratio, which is predicted to inhibit αKG-dependent dioxygenases.

(B) The αKG-to-succinate ratio in control (black) or 10 μM NPA incubated (red) H9 hESCs for 18 hr, quantified by UHPLC-MS.

(C) Immunoblot of OCT4 and PAX6 at indicated time points of NE differentiation with or without 50 μM NPA for H9 hESCs. β-TUB, β-tubulin.

(D and E) Flow cytometry analysis of PAX6 expression in H9 hESCs expressing shRNA targeting SDHA or scramble control shRNA treated with dm-αKG where indicated at day 4 of NE differentiation.

(F) UHPLC-MS quantification of fold change in αKG levels with TA inhibitors AOA and cyclo compared to drug carrier controls.

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lysine 4 trimethylation (H3K4me3) and histone lysine 27 trimethylation (H3K27me3) were assessed by intracellular flow cytometry. In general, NPA treatment during NE differentiation in UCLA1 hESCs led to an increase in histone marks, whereas dm-αKG led to a decrease (Figures 4E–4H and S4F–S4J). Analysis of an array of histone post-translational modifications by immunoblot revealed an overall repressive effect of dm-αKG on global lysine trimethylation but little effect on global monomethylation or acetylation marks (Figure 4I).

Overall, the data suggest that an increased αKG-to-succinate ratio accelerates, and a decreased αKG-to-succinate ratio retards, initial primed PSC differentiation by titered cofactor/inhibitor activities on epigenome remodeling enzymes, including TET enzymes and JHDMs. This interpretation is consistent with the proposed αKG/succinate mechanism for maintained naive pluripotency in mESCs through epigenetic regulation (Carey et al., 2015), only with an inverse cell fate outcome in the primed pluripotent state with induced differentiation.

**DISCUSSION**

This study reveals an unanticipated differentiation-promoting role for αKG in primed PSCs. Results have shown that αKG supports self-renewal of naive mESCs, potentially by promoting the demethylation of histones and DNA (Carey et al., 2015). Our data support a similar mechanism with an opposite outcome in the context of primed PSCs induced to differentiate. Consistent with a context-specific role for αKG, TET enzymes and JHDMs have dual roles in the self-renewal and differentiation of mESCs. TET1 promotes reprogramming to naive pluripotency, whereas triple knockout of the TET enzymes impedes differentiation (Costa et al., 2013; Dawlaty et al., 2014). JHDMs are a large class of enzymes that have functions in both naive pluripotency and differentiation. For example, JMJD3 and UTX promote naive pluripotency (Carey et al., 2015), whereas Jarid1b is involved in neurodevelopment (Mathieu et al., 2014; Xie et al., 2014). Both DNA and histone methylation levels are lower in naive compared to primed PSCs (Hackett and Surani, 2014; Leitch et al., 2013). These epigenome differences, along with metabolome differences, support a similar mechanism with an opposite outcome in the state-dependent effects of αKG in primed PSCs (Hackett and Surani, 2014; Leitch et al., 2013). These epigenome differences, along with metabolome differences.
between naive and primed pluripotent states, could support a differential role for αKG identified in this study. Our results suggest a model for αKG promotion of induced differentiation by primed PSCs from demethylation reactions that help silence pluripotency genes and activate lineage-specific genes to accelerate induced multi-lineage differentiation. A key goal of hPSC research is to develop mature and functional cells for regenerative medicine. Increased αKG could be useful for improving primed-state PSC differentiation.

**EXPERIMENTAL PROCEDURES**

Standard procedures were followed for immunoblotting, confocal microscopy, and siRNA knockdown, as described in the Supplemental Experimental Procedures. Maintenance of university compliance, including ESCRO, IRB, and Biological Safety, was overseen by K.H. and M.A.T.

**Cell Culture**

Primed hPSCs were passaged onto feeder-free matrigel (Fisher Scientific) in mTeSR1 medium with Gentle Cell Dissociation Reagent (STEMCELL) for most experiments. For experiments performed in EB or e6 media (STEMCELL), hPSCs were switched from mTeSR1 to appropriate media at passage. EpiSCs were grown in feeder-free conditions in medium containing bFGF and activin A on fibronectin. Further details are provided in the Supplemental Experimental Procedures.

**Seahorse Measurements**

OCR and ECAR assays were performed as previously described (Zhang et al., 2012). The hPSCs were plated onto an XF24 microplate (Seahorse Bioscience) at 10^4 or 10^5 cells/well with 10 μM Y-27632 (BioPioneer). The next day, 1 hr before the assay, the medium was changed to XF Media (Seahorse Biosciences) supplemented with 17.5 mM glucose. Cell metabolic rates were measured using an XF24 Extracellular Flux Analyzer (Seahorse Biosciences). Basal respiration was determined by quantifying OCR before and after the addition of 1 μM rotenone and 1 μM antymycin A (Sigma).

**Metabolite Extraction and Analysis**

Cellular metabolites were extracted with 80% ice-cold methanol, and ultra-high performance liquid chromatography-mass spectrometry (UHPLC-MS) measurements of metabolite levels were performed and analyzed as previously described (Thai et al., 2014). Details are provided in the Supplemental Experimental Procedures.

**Glutamate Uptake**

Levels of glutamate in culture media were measured using a BioProfile Basic Analyzer (Nova Biomedical). Details are provided in the Supplemental Experimental Procedures.

**NE Differentiation**

NE differentiation was performed as previously reported (Chambers et al., 2009; Shirak et al., 2014). Details are provided in the Supplemental Experimental Procedures.

**Flow Cytometry Analysis**

Cells were collected with Gentle Cell Dissociation Reagent and processed using the Cytofix/Cytoperm Kit (BD Biosciences). Cells were analyzed with either LSRII or LSRFortessa (BD Biosciences).

**Dot Blot Analysis and Quantification**

DNA was collected with the DNeasy Blood and Tissue Kit (QIAGEN) and quantified using a NanoDrop 1000 spectrophotometer (Thermo Fisher Scientific). DNA was denatured at 99°C for 5 min, put on ice, and neutralized by adding ammonium acetate to a final concentration of 0.66 M. Then, 400 ng of each sample were spotted on Amersham Hybond-N+ (Fisher) nylon membranes and baked at 80°C for 2 hr. Membranes were blocked with 5% skim milk for 3 hr and incubated with primary antibody overnight. The immunoblot procedure was followed. Blots were imaged with an Odyssey Fc and quantified with Image Studio v.5.2.5 (LI-COR Biosciences).

**Statistical Analysis**

Values are presented as mean ± SD or mean ± SEM. Data were analyzed with Prism (GraphPad). Pairwise comparisons were analyzed using two-tailed Student’s t test. Other data were analyzed using one-way or two-way ANOVA with correction for multiple comparisons. In all cases, p < 0.05 was considered significant.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures and four figures and can be found with this article online at http://dx.doi.org/10.1016/j.cmet.2016.07.002.

**AUTHOR CONTRIBUTIONS**

T.T., A.C.C., S.L.E., and D.B. performed experiments. I.L. and K.H. provided key reagents and expert guidance. T.T., J.H., T.G.G., D.B., and M.A.T. participated at differing levels in designing the study and analyzing data. T.T. and M.A.T. wrote the paper with help from T.G.G. and D.B. Maintenance of university compliance, including ESCRO, IRB, and Biological Safety, was overseen by K.H. and M.A.T.

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**REFERENCES**


