Meis1 regulates the metabolic phenotype and oxidant defense of hematopoietic stem cells

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The role of Meis1 in leukemia is well established, but its role in hematopoietic stem cells (HSCs) remains poorly understood. Previously, we showed that HSCs use glycolytic metabolism to meet their energy demands. However, the mechanism of regulation of HSC metabolism, and the importance of maintaining this distinct metabolic phenotype on HSC function has not been determined. More importantly, the primary function of Meis1 in HSCs remains unknown. Here, we examined the effect of loss of Meis1 on HSC function and metabolism. Inducible Meis1 deletion in adult mouse HSCs resulted in loss of HSC quiescence, and failure of bone marrow repopulation after transplantation. While we previously showed that Meis1 regulates Hif-1α transcription in vitro, we demonstrate here that loss of Meis1 results in down-regulation of both Hif-1α and Hif-2α in HSCs. This resulted in a shift to mitochondrial metabolism, increased reactive oxygen species production, and apoptosis of HSCs. Finally, we demonstrate that the effect of Meis1 knockout on HSCs is entirely mediated through reactive oxygen species where treatment of the Meis1 knockout mice with the scavenger N-acetylcysteine restored HSC quiescence and rescued HSC function. These results uncover an important transcriptional network that regulates metabolism, oxidant defense, and maintenance of HSCs. (Blood. 2012; 120(25):4963-4972)

Introduction

Hematopoietic stem cells (HSCs) are defined by their abilities to self-renew and to differentiate into all blood cell types.1,2 Much of the advancement in HSC therapy is credited to decades of pioneering work that led to the development of HSC enrichment techniques based on staining of cell-surface antigens or vital dyes followed by fluorescence-activated cell sorting (FACS).3,5 However, little is known about metabolic characteristics of HSCs, its regulation, or how the metabolic phenotype may influence HSC function.

In 1978, the concept of the special microenvironment, or niche, of HSCs was introduced.6 Since then, it has become clear that the niche plays a crucial role in self-renewal and differentiation of HSCs.7,8 One of the hallmarks of the HSC niche is its low oxygen tension, hence the term “hypoxic niche.”9 Numerous studies indicate that this low oxygen environment is not only tolerated by HSCs, but is also essential for their function.10 We recently demonstrated that HSCs rely on glycolysis and have lower rates of oxygen consumption,11 which may be crucial for survival of HSCs within hypoxic bone marrow niches.

In the mitochondria, oxygen is used as the terminal electron acceptor for the respiratory chain, and in the absence of oxygen the proton gradient generated by the respiratory chain collapses and mitochondrial ATP production declines. Under these hypoxic or anoxic conditions, energy production is derived from cytoplasmic glycolysis through the fermentation of glucose, and in the final step of anaerobic glycolysis, pyruvate is converted to lactate to replenish NAD+. Anaerobic glycolysis produces 18 times less ATP than mitochondrial oxidative phosphorylation,12 which may be well suited for quiescent cells, but certainly cannot sustain cells with high-energy demands.

The energy advantage of mitochondrial oxidative phosphorylation over glycolysis is, unfortunately, not without deleterious consequences, as the mitochondrion is considered a major source of reactive oxygen species (ROS) production.13,14 ROS are believed to be important mediators of aging, and of numerous degenerative diseases, including HSC dysfunction and senescence.15 In fact, within the HSC compartment, the repopulation capacity is localized to only those HSCs with low levels of free radicals.16 Therefore, the glycolytic metabolic phenotype of HSCs may not only protect them against hypoxic insults, but may also serve to minimize oxidant damage that result from mitochondrial oxidative phosphorylation.

Hypoxia-inducible factor-1α (Hif-1α) is a major transcriptional regulator of hypoxia response. Hif-1α mediates the metabolic switch from aerobic mitochondrial metabolism, to anaerobic cytoplasmic glycolysis17-19 by increasing both the expression,20 and kinetic rate21 of key glycolysis enzymes. Moreover, Hif-1α inhibits the use of pyruvate by the mitochondria,22,23 and inhibits mitochondrial biogenesis.24 Takubo and colleagues recently demonstrated that Hif-1α is enriched in HSCs, and that loss of Hif-1α in HSCs results in HSC dysfunction,25 while our group recently showed that Meis1 is required for optimum transcriptional activation of Hif-1α in HSCs ex vivo.11


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Meis1, which is a 3-amino-acid loop extension homeodomain protein, plays an important role in leukemogenesis as well as normal hematopoiesis. Meis1 was first identified as a common viral integration site in myeloid leukemic cells of BXX-2 mice, and it is also frequently up-regulated in human primary acute myeloid leukemia (AML) and acute lymphoblastic leukemia (ALL) samples. Moreover, overexpression of Meis1 accelerates the initiation of AML in murine models. In normal hematopoiesis, Meis1 is expressed in the most primitive hematopoietic populations and is down-regulated on differentiation. Targeted Meis1 knockout causes lethality by embryonic day 14.5 with multiple hematopoietic and vascular defects. Moreover, Pbx-1, a cofactor of Meis1, has been shown to regulate self-renewal of HSCs by maintaining their quiescence. However, the role of Meis1 regulating the function and metabolism in HSCs remains poorly understood.

In the current report, we show that Meis1 regulates both HSC metabolism and oxidant stress response, through transcriptional regulation of Hif-1α and Hif-2α, respectively.

Methods

Mouse breeding and genotyping

Meis1 knockout (KO) mice were genotyped with Meis1 F0r1: 5’-CCAAAGTACGACCAATATCATGA-3’ and Meis1 Rev: 5’-AGCGTGACTTGGAAAAGCAGATT-3’ primers. Wild-type (WT) allele is determined by a 332-bp-long PCR product and mutant allele determined by a 440-bp long PCR product on 1.2% agarose gel. HSC-specific deletion of Meis1 was achieved by following crosses of Meis1WT with Scl-Cre-ERT primers. Scl-Cre mice were genotyped using Scl-Cre-ER primer 1: 5’-GAACTGGAAGATGTTCGCGAT-3 and Scl-Cre-ER primer 2: 5’-ACCTGTAATGCAGTGGATAC-3. To generate Meis1−/− mice, Meis1f/f;Scl-Cre-ERT+ mice were injected intraperitoneally with tamoxifen (40 mg/kg, T5648-1G; Sigma-Aldrich) daily for 14 days. We used age-matched tamoxifen-injected Meis1+/-;Scl-Cre-ER(T+) or Meis1f/f;Scl-Cre-ER(T−) mice as controls (Meis1+/- mice). Genotyping of Cre-deleted Meis1 locus (Meis1 exon8 deleted) was performed using Meis1 F0r2: 5’-CATTGACCTTAATGTATGGTGTC-3’ and Meis1 Rev: 5’-AGCGTCACTTTGGAAAAGCAGATT-3’ primers. Cre-deleted Meis1 locus gives rise to a 261-bp product while wt and mutant (nondeleted) shows no amplicon.

Dr Joseph A. Garcia (UT Southwestern) kindly provided the Hif-1α floxed mice. HSC-specific deletion of Hif-1α was achieved by following crosses of Hif-1αLox/Lox with Scl-Cre-ERT mice and tamoxifen injections (40 mg/kg for 14 days). We used age-matched tamoxifen-injected Hif-1α+/-;Scl-Cre-ER(T+) or Hif-1αLox/Scl-Cre-ER(T−) mice as controls (Hif-1α+/- mice).

Flow cytometry

Donor bone marrow (BM) cells were isolated from 8- to 12-week-old Meis1−/− or Meis1+/- mice (or Hif-1α−/− and Hif-1α+/− mice). Lin+Sca1+Kit+Flk2−CD34+ cells (long-term HSCs [LT-HSCs]) were isolated by staining with a biotinylated lineage cocktail (anti-CD3, anti-CD5, anti-B220, anti–Mac-1, anti-Gr-1, anti-Thy1.2; StemCell Technologies) followed by streptavidin-PE/Cy5.5, anti–Sca-1–FITC, anti–Kit–allophycocyanin (APC), anti–Flk-2–PE, and anti-CD34–PE. For analyzing repopulation of mouse HSCs, peripheral blood cells of recipient CD45.1 mice were collected by retro-orbital bleeding, followed by lysis of red blood cells and staining with anti-CD45.2–FITC, anti-CD45.1–PE, anti–Thy1.2–PE (for T-lymphoid lineage), anti-B220–PE (for B-lymphoid lineage), anti–Mac-1–PE, or anti–Gr-1–PE (cells costaining with anti–Mac-1 and anti–Gr-1 were deemed to be of the myeloid lineage) monoclonal antibodies (BD Pharmingen). The “percent repopulation” shown in all figures was based on the staining results of anti-CD45.2–FITC and anti-CD45.1–PE. In all cases, FACS analysis of the above-listed lineages was also performed to confirm multilineage reconstitution as previously described.

For analyzing LT-HSCs in the peripheral blood by flow cytometry, peripheral blood of Meis1−/− or Meis1−/− mice was collected by retro-orbital bleeding and stained with LT-HSCs markers as described in the previous paragraph.

The cell-cycle analysis with Hoechst 33342 and pyronin Y staining was performed as we described. Samples were immediately analyzed by flow cytometry (FACSAria; BD Biosciences). To examine the apoptosis, Lin−Kit−Sca-1− cells were stained with PE-conjugated anti–annexin V and 7AAD according to the manufacturer’s manual (BD Pharmingen). To study the apoptosis in LT-HSCs, bone marrow cells were stained for HSC markers Sca-1-PE/Cy5.5, C-Kit-APC, CD34-PE, and Flk2-PE after lineage depletion and stained with FITC-conjugated anti–annexin V according to the manufacturer’s manual (eBioscience).

Competitive reconstitution analysis

The indicated numbers of CD45.2 donor cells from Meis1−/− or Meis1−/− mice were mixed with 1 × 106 freshly isolated CD45.1 competitor BM cells and the mixture was injected intravenously via the retro-orbital route into each of a group of 6- to 8-week-old CD45.1 mice previously irradiated with a total dose of 10 Gy. To measure reconstitution of transplanted mice, peripheral blood was collected at the indicated time points posttransplantation, and the presence of CD45.1+ and CD45.2+ cells in lymphoid and myeloid compartments was measured as described.

Homing

BM Lin− cells were labeled with 5- (and -6) carboxyfluorescein succinimidyl ester (CFSE), and 3 × 106 cells were transplanted into indicated strains of lethally irradiated mice. After 16 hours, the total number of CFSE+ cells in the BM, spleen, or liver was determined by flow cytometry. When CFSE+ LSKitCFSE− Sca-1+ (HSCs) were analyzed, the BM cells were stained with a biotinylated lineage cocktail followed by streptavidin-PE/Cy5.5, anti–Sca-1–PE, and anti–Kit–APC before analysis, as described.

 Colony-forming assays

Normal BM cells were diluted to the indicated concentration in IMDM with 2% FBS, and were then seeded into methylcellulose medium M3434 (StemCell Technologies), for CFU-GM and BFU-E colony formation according to the manufacturer’s instructions.

Real-time PCR

Total RNA was isolated from Meis1−/− and Meis1−/− HSCs using the RNeasy Mini Kit (QIAGEN) according to the manufacturer’s instructions. cDNA was synthesized using SuperScript II RT (Invitrogen). Predesigned primers (Table 1) from the National Institutes of Health (NIH) mouse primer depot (http://mouseprimerdepot.nci.nih.gov/) were ordered from Integrated DNA Technologies. Real-time PCR was performed with SyberGreen (Applied Biosystems) on an ABI Prism 7700 Sequence Detector (Applied Biosystems). β-actin was used as control to normalize results.

PCR array

The Mouse Hypoxia Signaling Pathway RT2 Profiler PCR Array was performed as described previously. A 488 bp of RNA was extracted from Meis1−/− and Meis1−/− LT-HSCs (Lin−Sca1+Kit+Flk2−CD34+ cells) using the RNeasy Mini Kit (QIAGEN) according to manufacturer’s instructions. cDNA was retrotranscribed by using SuperScript II RT (Invitrogen). Mouse Real-Time Syber Green PCR Master Mix (SuperArray) and Cell Cycle primer sets (SA Biosciences) used for real-time PCR on an ABI Prism 7700 Sequence Detector (Applied Biosystems). The data were analyzed using the ΔΔCT method. Fold change was calculated as difference in gene expression between Meis1−/− and Meis1−/− LT-HSC samples.
Table 1. List of primers used for real-time PCR

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward primer</th>
<th>Reverse primer</th>
</tr>
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<tbody>
<tr>
<td>β-actin</td>
<td>GAACCTTAGCCACAACGGTGAAGAT</td>
<td>ACGCCTGGTCCAAATAGTGAT</td>
</tr>
<tr>
<td>p16</td>
<td>GGTTCGGCACAAGCCCGGA</td>
<td>TGGCAACACCCACGGTCC</td>
</tr>
<tr>
<td>p19</td>
<td>GTTTCTTGGAAGTCTGTCG</td>
<td>TCTAATCTGGTCAGGATTTC</td>
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<tr>
<td>Meis1</td>
<td>GTGTGCGGATCACCTCCTT</td>
<td>ATCCTACTGGTCAGGAGAA</td>
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<td>Hif-1α</td>
<td>GCGGAGAAAGACGGAAGA</td>
<td>AAAATTTGACACTTTTCGTCC</td>
</tr>
<tr>
<td>Hif-2α</td>
<td>ATACGCGATTGTTCTCTCCTC</td>
<td>GGTAAAGGAACCCAGGTGCT</td>
</tr>
<tr>
<td>Hif-3α</td>
<td>TGTGAACTCTAGTCACGACG</td>
<td>GCAATGCGGTTGCTTATCT</td>
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Generation of luciferase reporter vectors

Conserved Meis1 motifs in Hif-2α gene were determined using a genome browser (http://genome.ucsc.edu). A 779-bp-long DNA fragment containing conserved Meis1 sites (located next to start codon sequence, human chr2:46 502:46 525 064) from the Hif-2α promoter was amplified by PCR from mouse genomic DNA with the following primers: conserved Meis1 site: pHif-2α-F, 5’-GGGCTAAGCGCTCAGG-3’ and pHif-2α-R, 5’-CATAGGAAACGCTCAGGAAAGC-3’. PCR fragments were subcloned into the pCR2.1-TOPO vector (Invitrogen) according to the manufacturer’s instructions. pHif2α-TOPO and E1b-pGL2 vectors were digested with XhoI and KpnI. Then, the PCR fragments containing conserved Meis1 sites were cloned into E1b-pGL2 to generate the pHif2α-pGL2 luciferase reporter vector. To test Meis1 site specificity, Meis1-binding sites (TGAC) at the TGCAGCTGACAA (Meis1 binding site is indicated in bold) sequence were mutated to TGGCCGGCCGCAA (Nori site insertion) using lProof High-Fidelity DNA Polymerase (Bio-Rad) from the Hif-2α-pGL2 vector with the following primers: Mut-Hif2α-F, 5’-AAgGccgGccCAAGGAAAAAAAGTTAAGCGGGG-3’ and Mut-Hif2α-R, 5’-AAgGgGcGCTTACGCGTCGCGCTC-3’. The Hif-2α-Mut-pGL2 reporter was generated after Nori digestion and ligation of the PCR product. Transcriptional activation of Hif-2α by Meis1 was evaluated using a luciferase reporter system (Promega) as described previously.11

Metabolic assays

Metabolic assays are carried out as described previously11 with some modifications.

Oxygen consumption assays

Meis1+/+ and Meis1−/− HSCs were separated flowcytometrically as described in “Flow cytometry.” Equal numbers of cells (5-10 x 10^6 cells/well) were incubated for 6 hours in the provided 384-well plate (BD Oxygen Biosensor System) and sealed to prevent air exchange before measurement. Culture media lacking cells was used as a negative control and sodium sulfite (100 mM) was used as a positive control. Oxygen consumption is presented as relative units.

ATP assays

Meis1+/+ and Meis1−/− HSCs were sorted as described and centrifuged at 1200g for 10 minutes. At least 50,000 cells were used for each single ATP measurement. Fifty microliters of ATP standards (10^-9.5-10^-12M) and 50 μL of cell lysates were quantified using the ATP Bioluminescence Assay Kit CLS II (Roche) using Fluostar Optima plate reader (BMG Labtech). Finally, data were normalized to cell count and protein content.

Glycolytic flux assay

13C-lactate production, end product of glycolysis, was measured as described previously11 using glycolytic flux medium supplemented with 10mM D-[1-6-13C]-glucose (Cambridge Isotope Labs) to allow up to all of the glucose-derived lactate pool to be labeled on C-3. A minimum of 50,000 cells were cultured in 40 μL of flux medium overnight. Then, the cells were pelleted and supernatant collected and prepared for gas chromatography–mass spectrometry. Lactate abundance was determined by monitoring m/z at 117 (unenriched), 118 (lactate containing 13C from glucose), and 119 (internal standard) as described previously.11

Measurement of ROS

Bone marrow cells from Meis1+/+ and Meis1−/− mice were isolated as described in “Flow cytometry.” After lineage depletion (leucine depletion kit; BD Biosciences), cells were incubated with 1μM 5- (and-6)-carboxy-2’,7’-dichlorofluorescein diacetate (carboxy-DCFDA; Invitrogen) for 30 minutes in a 37°C water bath in the dark. Then, cells were stained for HSC markers Sca-1-PE/Cy5.5, C-Kit-APC, CD34-PE, and Flk2-PE and assayed with a flow cytometer.

NAC administration in vivo

Six-week-old Meis110/Scl-Cre-ERT+ mice were injected intra-peritoneally with tamoxifen (40 mg/kg) daily for 14 days. After tamoxifen injections, animals were treated daily for 2 weeks with N-acetyl-L-cysteine (NAC; 100 mg/kg; Sigma-Aldrich) by subcutaneous administration or provided in drinking water (500 mg x kg^-1 x d^-1) in drinking water as described previously39,40 and were subsequently analyzed by flow cytometry. We used age-matched tamoxifen-injected Meis1+/+,Scl-Cre-ER(T+) or Meis110/Scl-Cre-ER(T−) mice as controls (Meis1+/− mice).

Chromatin immunoprecipitation assay

Chromatin immunoprecipitation assay (ChIP) assays were performed to evaluate the in vivo binding of Meis1 to its consensus sequence in the Hif-2α gene. The assays were done in Kasumi-1 cells, a hematopoietic progenitor cell line, using the ChIP kit (Upstate) as described previously.11 Meis1 antibody (Santa Cruz Biotechnology) and normal goat IgG (Santa Cruz Biotechnology) were used. The DNA isolated from immunoprecipitated fragments and from the precipitated chromatin fragments by anti-Meis1 antibody or control IgG was subjected to real-time PCR using primers flanking the consensus Meis1-binding sites on Hif-2α promoter: 5’-GGTTCCTCCAGCTACATTTCTCC-3’ and 5’-CTCACCAACCACCTCGCTC-3’.

Measurement of blood cell counts (CBC/differential)

Approximately 50 μL of peripheral blood collected by retro-orbital bleeding from Meis1+/+ and Meis1−/− mice post 1 month and 3 months of tamoxifen injections in K+EDTA tubes. Samples are submitted to University of Texas Southwestern Medical Center Animal Resources Center (UTSW ARC) Diagnostic Laboratory and analyzed with the HemaVet 950FS analyzer. The following parameters are reported for the sample submitted: white blood count (WBC), neutrophil count, lymphocyte count, monocyte count, eosinophil count, basophil count, red blood cell count, hemoglobin, HCT, MCV, MCH, MCHC, platelet count, and MPV.

Generation of the Meis110/Scl-Cre+; rtTA; TRE-Hif-1α mice and rescue analysis by viral Hif-1α and Hif-2α overexpression

To determine the role of Hif-1α in Meis1 KO HSCs, inducible HSC-specific, Hif-1α transgenic mice were generated. The transgenic construct was made by cloning Hif-1α cDNA (Open Biosystems) into the pTRE-Tight vector (Clontech), which makes it responsive to rtTA regulatory proteins in the Tet-On system. We crossed TRE-Hif-1α mice with rtTA mice which has a stop-codon flanked by loxp sequence. Cre-mediated removal of the stop region by deletion of the loxp flanked sequence allows expression of rtTA as described.41 Therefore, sequential administration of tamoxifen will result in Meis1 deletion, and concomitant deletion of the stop-codon,
which allows for Hif-1α overexpression on administration of doxycycline. We generated Meis1<sup>fl/ft</sup>;Scl-Cre<sup>+</sup>;rtTA; TRE-Hif-1α mice by a series of crosses to overexpress in Hif-1α in HSC-specific manner after 14 days of tamoxifen (1 mg/d/mice) and providing doxycycline in drinking water (500 mg/L). HSCs from these mice were either used for in vivo repopulation studies, or for in vitro colony forming assay.

We also used a viral strategy to overexpress Hif-1α or Hif-2α in HSCs (generated by cloning into XZ201 vector) and lentivirus-expressing Hif-2α (kindly provided by Dr Joseph Garcia, UT Southwestern). Fetal liver cells were harvested from Meis1<sup>fl/ft</sup>;Scl-Cre<sup>+</sup> pups at P1 and infected with either lentivirus-expressing Hif-2α or retrovirus-expressing Hif-1α with IRES-GFP. Then, infected cells were transplanted into CD45.1<sup>+</sup> recipient host mice. The repopulation was measured 2 months posttransplantation before induction of deletion of Meis1 with tamoxifen injections. The repopulation analysis performed at indicated time points post-tamoxifen treatments either measuring GFP<sup>+</sup> cells or CD45.2<sup>+</sup> donor cells (n = 3-5).

Finally, in a separate set of experiments, we used cobalt chloride (CoCl<sub>2</sub> 100μM), which is a known stabilizer of both Hif-1α and Hif-2α, in methocult culture after deletion of Meis1 to examine the effects of stabilization of Hif-1α and Hif-2α on colony formation assay after Meis1 deletion. After 14 days of tamoxifen injection into aged matched Meis1<sup>+/+</sup>; Scl-Cre<sup>+</sup> (Control) and Meis1<sup>fl/ft</sup>;Scl-Cre<sup>+</sup> (Meis1 KO) mice, we performed colony-forming assays using Methocult medium supplemented with 100μM CoCl<sub>2</sub>. Same number of bone marrow cells (20 x 10<sup>5</sup> cells/plate) was plated as recommended by manufacturer and colonies were quantified after 12 days of culture.

Statistical analysis
Results are expressed as mean ± SEM, and a 2-tailed Student t test was used to determine the level of significance. P < .05 was considered statistically different.

Results
Deletion of Meis1 in LT-HSCs
Because the global loss of Meis1 is embryonic lethal,<sup>33,34</sup> we sought to pursue an inducible deletion of Meis1 to study the role of Meis1 in adult HSCs. Meis1<sup>fl/ft</sup> mice with loxp flanking exon 8 were crossed with transgenic mice expressing the tamoxifen-inducible Cre recombinase under the control of stem cell leukemia (Scl) HSC enhancer, which drives deletion in HSCs.<sup>4</sup> Upon tamoxifen treatment, exon 8 is deleted which results in the loss of Meis1 expression in HSCs (supplemental Figure 1A, available on the Blood Web site; see the Supplemental Materials link at the top of the online article). To verify deletion of Meis1, we performed genotyping analysis and quantitative RT-PCR in peripheral blood cells and phenotypic LT-HSCs (Lin<sup>-</sup> Sca1<sup>-</sup> Kit<sup>-</sup> Flik2<sup>-</sup> CD34<sup>-</sup>) in the bone marrow after tamoxifen treatment. Meis1 in peripheral blood cells was deleted 14 days after tamoxifen treatment, and Meis1 mRNA level in LT-HSCs was markedly decreased to approximately 10% of control values (supplemental Figure 1B).

Meis1 is required for the maintenance of LT-HSCs
To explore the role of Meis1 in LT-HSCs, we first examined the frequencies of phenotypic LT-HSCs in control (Meis1<sup>+/+</sup>) and Meis1 conditional KO (Meis1<sup>-/-</sup>) mice. As shown in Figure 1A, 7 days after tamoxifen treatment, there was a 4.5-fold increase of HSC frequency in Meis1<sup>-/-</sup> mice compared with Meis1<sup>+/+</sup> controls.
The increase of HSC frequency may result from cell-autonomously accelerated proliferation or the compensatory effect of increased apoptosis or mobilization in Meis1−/− HSCs. To address this concern, we first examined the cell cycle of HSCs by using Hoechst 33342 and pyronin Y staining, and we found that the percentage of CFSE− cells in different tissues showed no difference between Meis1−/− and Meis1+/− mice (n = 5). (F) Quantification of repopulation of lineages for Meis1−/− using whole bone marrow from first BMT mice demonstrates no defects in lineage repopulation (n = 5); *P < .05, **P < .01.

To further evaluate the function of Meis1 in HSCs in vivo, we performed competitive bone marrow transplantation with Meis1+/− and Meis1−/− HSCs. Two weeks after tamoxifen treatment, 150 Lin−/−Sca1+Kit−/−Flk2−/−CD34− Meis1+/− or Meis1−/− CD45.2 HSCs, together with 1 × 10⁶ CD45.1 competitors, were administered into CD45.1 recipients through retro-orbital injection. Repopulation was examined at 4, 10, 16 weeks after transplantation. Strikingly, as shown in Figure 2A, we could not detect any engraftment with Meis1−/− HSC donors, indicating a severe impairment of HSC repopulation ability after loss of Meis1. A repeat transplantation experiment with total BM cells showed similar results (Figure 2B). The impaired repopulation of Meis1-deficient LT-HSCs (Figure 2A) and progressive decline in repopulation of Meis1-deficient cells in primary transplant recipients (Figure 2B) suggested a defect in self-renewal. Therefore, we performed secondary transplantation experiments to confirm the reduction of function Meis1-deficient LT-HSCs in the bone marrow of primary transplant recipients. Eight weeks after primary transplantation, total bone marrow from primary recipients from whole transplanted mice were harvested and transplanted into secondary recipients. Meis1-deficient cells from primary recipients (0.5-1 × 10⁶ cells) were unable to repopulate secondary recipients (Figure 2C). The decreased repopulation may result from defects in HSC maintenance, or homing, or increased apoptosis. To exclude the possibility that the decreased engraftment is caused by defect of homing in Meis1−/− HSCs, we labeled Meis1−/− or Meis1+/− Lin−/− cells with CFSE and injected into lethally irradiated recipients. Sixteen hours later, we examined CFSE− or LSK CFSE− cells in spleen, liver, and BM. No significant difference in homed total CFSE− or LSK CFSE− cells was detected between Meis1−/− and
Meis1−/− donors (Figure 2D-E). In addition, quantification of lineage repopulation of Meis1+/+ and Meis1−/− donors demonstrates no lineage-specific defects (Figure 2F). Taken together, these data provide strong evidence that Meis1 plays a crucial role in the maintenance of HSCs and prevention of apoptosis in HSCs.

**Role of Meis1 in regulating Hif-1α for glycolytic metabolism of LT-HSCs**

We have shown previously that the Meis1 gene acts upstream of Hif-1α by regulating its transcriptional activity by an enhancer located in the first intron of Hif-1α.17 We further examined whether hypoxia-inducible factors (Hif-1α, Hif-2α, and Hif-3α) are regulated by Meis1 in LT-HSCs. Quantitative real-time PCR analysis demonstrated that mRNA levels of both Hif-1α and Hif-2α were down-regulated but no change in Hif-3α levels was observed in Meis1−/− LT-HSCs (Figure 3A). Moreover, we found that several downstream targets of Hif-1α and/or Hif-2α are down-regulated in Meis1−/− LT-HSCs as determined by quantitative PCR (qPCR) array (supplemental Figure 2). We used the same inducible Cre system to delete Hif-1α specifically in HSCs (supplemental Figure 3A). Hif-1αfl/fl mice with loxp flanking exon 2 were crossed...
with transgenic mice expressing the tamoxifen-inducible Cre recombinase under the control of stem cell leukemia (Scl) HSC enhancer. Upon tamoxifen treatment, exon 2 is deleted resulting in the loss of Hif-1α expression in HSCs (supplemental Figure 3A). To verify deletion of Hif-1α, we performed quantitative RT-PCR in LT-HSCs (Lin− Sca1+ Kit+ Flk2− CD34−) in bone marrow after tamoxifen treatment. Hif-1α mRNA level in LT-HSCs was dramatically decreased, approximately 400-fold lower than that of control (Hif-1+/+ ) after 14 days of tamoxifen treatment (supplemental Figure 3B). We showed previously that LT-HSCs primarily rely on cytoplasmic glycolysis rather than mitochondrial oxidative phosphorylation. However, the role of Hif-1α in HSC metabolism was not examined before. Here, we demonstrate that HSC-specific deletion of Hif-1α results in increased mitochondrial respiration as shown by increased oxygen consumption (supplemental Figure 3C), and decreased glycolytic flux measured by the rate of glucose-derived 13C-lactate production (supplemental Figure 3D).

To further examine the role of Hif-1α in HSCs, we examined expression of Hif-2α and Hif-3α in Hif-1α−/− LT-HSCs. We found that depletion of Hif-1α in LT-HSCs results in a profound increase in Hif-2α mRNA levels (>120-fold), with no significant change in the levels of Hif-3α mRNA (supplemental Figure 3E). This marked compensatory up-regulation of Hif-2α in Hif-1α−/− LT-HSCs is in stark contrast to the down-regulation of Hif-2α in the Meis1−/− LT-HSCs.

HSC-specific deletion of Hif-1α also demonstrated hematopoietic defects similar (supplemental Figure 4) to global deletion (Mx-1-Cre) in the bone marrow shown by Takubo and colleagues, including the increased frequency of HSCs (supplemental Figure 4A), decreased myeloid progenitors (supplemental Figure 4B), loss of quiescence (supplemental Figure 4C), and similar multilineage defects (supplemental Figure 4F-G).

Meis1 is a transcriptional activator of Hif-2α

Down-regulation of Hif-1α and Hif-2α in Meis1−/− LT-HSCs (Figure 3A) led us to examine their metabolic profile and ROS levels. We found that Meis1−/− LT-HSCs have increased rates of oxygen consumption (Figure 3B) and decreased glycolytic flux (Figure 3C). In addition, Meis1−/− LT-HSCs showed significantly higher ROS levels compared with control Meis1+/+ LT-HSCs (Figure 3D). This increased ROS was associated with increased expression of p16INK4a and p19ARF in Meis1−/− LT-HSCs, which are known to induce HSC senescence and apoptosis, respectively (Figure 3E). Therefore, the increased ROS production in Meis1−/− LT-HSCs is compounded by the down-regulation of the master antioxidant gene Hif-2α. Moreover, measurement of peripheral blood counts of Meis1−/− mice post 1 month and 3 months of tamoxifen treatments demonstrates decreased red blood cells, white blood cells, as well as platelets, similar to the phenotype observed in Hif-2α−/− mice (supplemental Figure 5). To determine the mechanism of down-regulation of Hif-2α in Meis1−/− LT-HSCs, we identified 2 Meis1 consensus-binding motifs in the promoter of the Hif-2α gene (Figure 3F). Using a Hif-2α-pGL2 reporter which includes the conserved Meis1-binding sites, we demonstrate a dose-dependent activation of Hif-2α by Meis1 expression vector (CMV-Meis1; Figure 3G). In addition, this activation demonstrates dependence on binding of Meis1 to its consensus-binding sequences in the Hif-2α promoter because mutation of the seed sequences (Hif-2α–Mut-pGL2) completely abolished the activation of Hif-2α by Meis1. Given down-regulation of Hif-2α in Meis1−/− LT-HSCs (Figure 3A), these results indicate that Meis1 is required for optimal transcriptional activation of Hif-2α in LT-HSCs. We also confirmed the in vivo binding of Meis1 to its conserved sequences in the Hif-2α gene by ChIP assays in Kasumi1 cells as determined by real-time PCR after immunoprecipitation with Meis1 antibody (Figure 3H).

Effect of ROS scavenging on the Meis1−/− phenotype

To study effects of ROS observed in Meis1−/− mice, we used NAC, an antioxidant, in an attempt to rescue the Meis1 phenotype by scavenging of ROS in Meis1 HSCs. After 14 days of tamoxifen injection, we administered NAC intraperitoneally for 12 days (Figure 4A). Mice where then harvested and HSCs were isolated for analysis of HSC frequency, HSC cell-cycle status, apoptosis rate, ROS levels, and expression of redox-sensitive cell-cycle regulators P16INK4a and P19ARF. Here, we show that NAC administration rescues the Meis1−/− phenotype. We found that frequency of HSCs in Meis1−/− mice become similar to Meis1+/+ mice (Figure 4B). Flow cytometric analysis of cell cycle of Meis1−/− HSCs (Figure 4C). Quantification of apoptosis in Meis1+/+ and Meis1−/− mice shows an increased apoptosis with Meis1−/− mice; however, this increase in the number of apoptotic cells was not statistically significant (P = .051; Figure 4D). In addition, quantification of ROS after NAC treatments restored ROS to Meis1−/− control levels in Meis1−/− HSCs (Figure 4E). Finally, scavenging ROS restored the transcript levels of P16INK4a and P19ARF in HSC to Meis1+/+ control levels (Figure 4F). While these effects are most likely attributable to the antioxidant effect of NAC, other unknown mechanisms of NAC may also contribute to this rescue.

To further evaluate the effect of ROS on engraftment defect observed after Meis1 deletion, we performed another bone marrow transplantation from Meis1−/− and Meis1−/− mice treated with and without NAC. Two weeks after tamoxifen treatment and NAC administration, 1×103 BM cells from Meis1−/− or Meis1−/− CD45−2, together with 1×105 CD45.1 competitors, were injected into CD45.1 recipients through the retro-orbital complex (Figure 4G). After bone marrow transplantation, NAC is provided in drinking water for 2 weeks and injected daily for 2 more weeks. Repopulation was examined at 4 weeks posttransplantation. As shown in Figure 4H, scavenging of ROS by NAC administration restored engraftment defects in Meis1−/− donors. In conclusion, scavenging ROS, through administration of NAC, rescues the Meis1 phenotype in HSC by decreasing levels of ROS, thereby normalizing P16INK4a and P19ARF expression and restoring HSC quiescence and repopulation defects.

Effect of HIFs on stem cell function in Meis1−/− phenotype

Meis1 KO BM cells treated with 100μM CoCl2 resulted in restoration of total number of colonies to the WT levels (supplemental Figure 6A). Moreover, we found that Hif-1α overexpression using the Meis1F/F;Scl-Cre−; rtTA; TRE-Hif-1α mouse strategy, which resulted in marked up-regulation of Hif-1α in HSCs, resulted in restoration of the BFU-E and total number of colonies to the WT levels, as well as partial restoration of number of mixed colonies (CFU-GEMM) which is a measure of most undifferentiated hematopoietic progenitors (supplemental Figure 6C). In addition, we attempted to rescue repopulation defect in Meis1−/− HSCs using viruses that express Hif-1α and Hif-2α (supplemental Figure 6D) or using Meis1F/F;Scl-Cre−; rtTA; TRE-Hif-1α (supplemental Figure 6E); however, these strategies failed to rescue the in vivo reconstitution capacity up to 4 months after Meis1 deletion. In
In retrospect, these results are not entirely surprising in light of recent results demonstrating a narrow beneficial dose range of Hif-1α on HSC function, where overstabilization of Hif-1α results in worsening HSC function which might explain worsening of the HSC repopulation defect in Meis1 KO HSCs in vivo (supplemental Figure 6D-E).

In summary, we demonstrate that Meis1 deletion results in a shift in HSC metabolism toward oxygen consumption, with the resultant increase in ROS production. This phenotype is compounded by down-regulation of the oxidant stress response gene Hif-2α, and as a result Meis1 deletion results in HSC dysfunction and apoptosis.

**Discussion**

In the current report, we demonstrate that Meis1 functions upstream of a transcriptional network that regulates HSC metabolism and oxidant defense. The Meis1 deletion-induced metabolic shift and oxidant injury is compounded by the down-regulation of
Hif-2α, which is in stark contrast to the marked up-regulation of Hif-1α in HSC after Meis1 deletion. This phenotype is associated with up-regulation of p16Ink4a and p19Arf, loss of quiescence, increased apoptosis, and marked HSC dysfunction.

Endothelial PAS domain protein 1 (EPAS1), also known as Hif-2α,4,44 is closely related to Hif-1α in structure and is likewise activated during hypoxia.29 While Hif-1α is a master regulator of metabolism, Hif-2α is a master regulator of oxidant stress response45 and is induced by ROS.46 It is involved in regulation of numerous antioxidant genes that minimize the oxidant damage that results from mitochondrial respiration.43 Hif-2α cells were pancreatic,42,47 and have high levels of oxidative stress,45 which suggests that Hif-2α is required for normal hematopoiesis. Our results indicate that Hif-2α is markedly up-regulated after Hif-1α deletion, which is an indication of a robust antioxidant response to Hif-1α deletion, likely secondary to a shift toward oxidative metabolism, with the subsequent increase in ROS.46 In stark contrast, we show that Hif-2α is down-regulated after Meis1 KO in HSCs, which may partially explain the severity of the Meis1−/− phenotype compared with the Hif-1α−/− phenotype.

Delicate control of ROS levels in HSCs is crucial for HSC maintenance, where elevated ROS levels in HSCs is associated with defects in HSC self-renewal and increased apoptosis. Regulation of ROS in HSCs is a highly complex process that involves regulation of the metabolic phenotype of HSCs, as well as regulation of antioxidant defense mechanisms. The contribution of oxidative metabolism to ROS production has been extensively studied.48 It is estimated that 2% of all electrons flowing through the mitochondrial respiratory chain result in the formation of oxyradical.49 Cells leaking from the respiratory chain interact with oxygen, partially reducing it to superoxide anion (O2−).48 Even though O2− itself is not a strong oxidant, it is the precursor of most other ROS.14 ROS overwhelm the natural antioxidant defense mechanisms over time, and result in widespread cellular damage.48 ROS can also induce the cell-cycle regulators p16Ink4a and p19Arf, which cause loss of quiescence and apoptosis of HSCs.49 Our results indicate that Meis1 regulates ROS production in HSC through regulation of both the metabolic phenotype (through Hif-1α) and oxidant defense mechanisms (through Hif-2α). Finally, although our ROS scavenging studies indicate that the effect of loss of Meis1 on HSC cell cycle and engraftment defect are entirely mediated through an increase in ROS production, rescue studies using Hif-1α and/or Hif-2α overexpression or stabilization failed to fully rescue the phenotype. These results support previous reports which indicate that the precise dose of Hif-1α is necessary for optimal HSC function.25 Although our results implicate Hif-1α and Hif-2α in the HSC phenotype after Meis1 deletion, there certainly may be additional ROS modulators that are targeted by Meis1.

In the current report, we highlight the role of Meis1 in a transcriptional network that regulates HSC metabolism and antioxidant defense. These results implicate Meis1 as an important regulator of the redox state of HSCs, which may be echoed by its role in leukemogenesis. Therefore, it would be important for future studies to determine the role of Meis1 in regulation of leukemia stem cell metabolism, survival, and self-renewal. Our findings suggest that HSCs are endowed with redundant mechanisms for regulation of their metabolic phenotype, rather than being solely dependent on environmental signals, such as the hypoxic microenvironment. Deciphering the role of these transcriptional networks in regulating HSC fate and function may provide valuable clues for understanding HSC disease and malignancies.

Acknowledgments

The authors thank Dr Keith Humphries, Michelle Miller, and Patty Rosen for sharing some initial data on characterization of the KO line genotyping, Dr Joachim R. Goethert at Universitätsklinikum Essen for providing the Scl-Cre-ER T2 mice, and Dr Joseph A. Garcia for Hif-1α floxed mice.

This work is supported by grants from the American Heart Association (AHA; GIA12060240 [H.A.S.]), the Gilead Research Scholars Program in Cardiovascular Disease (H.A.S.), National Institutes of Health (NIH) grant R01HL115275 (H.A.S.), NIH grant K01 CA120099 (C.Z.), and Cancer Prevention and Research Institute of Texas (CPRIT) RP100402 (C.Z.).

Authorship

Contribution: F.K. designed and performed the research, analyzed data, and wrote the manuscript; J.Z., S.T., N.G.C., N.A.J., and R.J.D. performed research and analyzed data; and C.Z. and H.A.S. designed and supervised research and wrote the manuscript.

Conflict-of-interest disclosure: The authors declare no competing financial interests.

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Meis1 regulates the metabolic phenotype and oxidant defense of hematopoietic stem cells

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