NCOR1 modulates erythroid disorders caused by mutations of thyroid hormone receptor α1

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Thyroid hormone receptor α (THRA) gene mutations, via dominant negative mode, cause erythroid abnormalities in patients. Using mice expressing a dominant negative TRα1 mutant (TRα1PV; Thra1PV/mice), we showed that TRα1PV acted directly to suppress the expression of key erythroid genes, causing erythroid defects. The nuclear receptor corepressor 1 (NCOR1) was reported to mediate the dominant negative effects of mutated TRα1. However, how NCOR1 could regulate TRα1 mutants in erythroid defects in vivo is not known. In the present study, we crossed Thra1PV/mice with mice expressing a mutant Ncor1 allele (NCOR1ΔID; Ncor1ΔID mice). TRα1PV mutant cannot bind to NCOR1ΔID. The expression of NCOR1ΔID ameliorated abnormalities in the peripheral blood indices, and corrected the defective differentiation potential of progenitors in the erythroid lineage. The defective terminal erythropoiesis of lineage-negative bone marrow cells of Thra1PV/mice was rescued by the expression of NCOR1ΔID. De-repression of key erythroid genes in Thra1PV/Ncor1ΔIDΔID mice led to partial rescue of terminal erythroid differentiation. These results indicate that the inability of TRα1PV to recruit NCOR1ΔID to form a repressor complex relieved the deleterious actions of TRα1 mutants in vivo. NCOR1 is a critical novel regulator underpinning the pathogenesis of erythroid abnormalities caused by TRα1 mutants.

Thyroid hormone receptors (TRs) mediate the nuclear actions of thyroid hormone (T3) in growth, development, differentiation, and maintaining metabolic homeostasis. There are two TR genes, THRA and THRB, that encode three major T3 binding receptors, TRα1, TRβ1, and TRβ2. These receptors share high sequence homology in the DNA binding and T3 binding domains, but differ in the amino terminal A/B domain. The transcriptional activity of TR is dictated the type of thyroid hormone response elements on the promoter of T3 target genes, and is modulated by T3-dependent interaction with nuclear coregulatory proteins, e.g., nuclear corepressors and coactivators. The classical bimodal switch model of TR action is that in the absence of T3, TR complexed with the retinoid acid receptor (RXR) recruits corepressors to repress gene transcription. Binding of T3 releases corepressors to allow the liganded-TR-RXR to recruit coactivators to activate gene transcription. The best-studied nuclear corepressors are the nuclear receptor corepressor 1 (NCOR1) and silencing mediator of retinoid and thyroid hormone receptors (SMRT; NCOR2). These two corepressors share about 50% identity in the amino acid sequences, but have similar functional domains. However, the three receptor interaction domains located in the C-terminal region of these two corepressor proteins show some preferential avidity to associate with TRs. It is possible that these two corepressors could have non-overlapping functions to regulate TR actions in target tissues.

Because TR isoforms share high sequence homology in the functional DNA and T3 binding domains, the question of whether TR isoforms have redundant or isoform-specific roles has been intensively studied. Mice deficient in Thra, Thrb, or both genes showed that TR isoforms have a redundant role as well as overlapping functions. However, in mice expressing an identical knock-in dominant negative mutation (hereafter referred to as PV) in the Thrb or Thra genes, the phenotypic expression is distinct. The ThrbPV mice exhibit the hallmark of resistance to thyroid hormone (RTH) with dysregulation of the pituitary-thyroid axis, hyperglycemia, and enlarged fatty livers. In contrast, Thra1PV mice have nearly normal thyroid functions tests, but exhibit growth retardation, delayed bone development, and reduced fat mass and liver size. These observations indicated that TR mutant isoforms exhibit distinct biological functions in vivo and predicted that mutations of TR subtypes could lead to different human diseases. While autosomal dominant resistance was first recognized in 1967 and
mutations of the *THRB* gene were identified to cause the disease (RTHβ) in 1989, three patients with mutations of the *THRA* gene were not discovered until 2012. Indeed, similar to *Thra1PV* mice, in spite of nearly normal thyroid function tests, these patients exhibit classical hypothyroidism with growth retardation and delayed bone development, indicating resistance of target tissues to thyroid hormones (RTHα). The discovery of RTHα patients displaying symptoms distinct from those of RTHβ patients unambiguously shows that the in vivo molecular actions of TR mutant isoforms are distinct.

Interestingly, the mutated C-terminal sequences in TRα1PV share the identical truncated sequence in two RTHα patients. Through use of *Thra1PV* mice, much has been learned about how mutated TRα1 led to bone abnormalities at the molecular levels. Moreover, the *Thra1PV* mouse has been used as a preclinical model to test whether long-term treatment of T4 could be beneficial to patients with mutations of the *THRA* gene. One notable pathological manifestation in patients with RTHα is erythroid disorders (e.g., anemia) that were not observed in RTHβ patients. Recently, we have shown that *Thra1PV* mice, similar to RTHα patients, also exhibited erythroid abnormalities. We further elucidated that TRα1PV, via dominant negative action, impaired erythropoiesis by suppressing the expression of the key erythroid genes, *Gata1*, *Klf1*, and their several downstream target genes in the bone marrow of *Thra1PV* mice. These findings prompted us further to ascertain how the dominant negative actions TRα1PV is regulated in mediating the erythroid disorders in *vivo*. NCO1 has been shown to modulate the in vivo dominant negative action of TRα1PV in the adipocytes. Accordingly, we adopted the loss of function approach by crossing *Thra1PV* mice with mice expressing a mutant Ncor1 allele (*Ncor1ΔID; Ncor1ΔID* mice) that cannot recruit TRα1PV mutant. Remarkably, we found that the disruption of the interaction of NCO1 to complex with TRα1PV ameliorated the deleterious actions of TRα1PV on erythropoiesis. Thus, aberrant interaction of TRα1 mutants underpinning the pathogenesis of erythroid disorders. Importantly, the present studies uncovered NCO1 as an important regulator in TRα1 signaling in erythropoiesis.

**Results**

**Expression of NCO1 ΔID reverts abnormal erythropoietic parameters and ameliorates defective progenitor differentiation capacity of *Thra1PV* mice.** Previously, we have shown that peripheral erythropoietic indices were lower in *Thra1PV* mice than in wild-type (WT) mice. Consistent with those findings, we found that the red blood indices were reduced 16.1% (red blood cell count), 11.2% (hemoglobin content), 9.2% (hematocrit) and 27% (platelets) as compared with WT mice (Fig. 1A, bars 3 versus bars 1 in panels a,b,c and d). Remarkably, the expression of NCO1ΔID in *Thra1PV* mice nearly completely corrected the decreased blood indices (bars 4 in all panels). These data indicated that the abnormal red blood cell indices of *Thra1PV* mice could be reverted by the expression of NCO1ΔID.

It is known that anemia stress induces the expression of erythropoietin (EPO) in *vivo*. Accordingly, we determined EPO levels in mice with four genotypes. *Thra1PV* mice which are anemic, exhibited elevated EPO (bar 3, Fig. 1A-e). The reversal of anemic phenotypes by the expression of NCO1ΔID in *Thra1PV* *Ncor1ΔIDΔID* mice (bars 4 in Fig. 1A, panels a–d) led to the lowering of EPO (bar 4, Fig. 1A-e). These EPO data further support that the expression of NCO1ΔID in *Thra1PV* *Ncor1ΔIDΔID* mice ameliorated the erythroid disorders in *Thra1PV* mice.

Figure 1B shows that the expression of NCO1ΔID partially corrected the decreased total bone marrow cells from a reduction of 58.0% in *Thra1PV* mice (bar 3 versus bar 1, Fig. 1B) to 38.2% in *Thra1PV* *Ncor1ΔIDΔID* mice (bar 4 versus bar 1). There were no significant differences in the total bone marrow cells between WT mice and *Ncor1ΔIDΔID* mice (bar 1 versus bar 2). That the expression of NCO1ΔID could partially correct the deficiency in the total bone marrow cells of *Thra1PV* mice prompted us to ascertain the effect of the expression of NCO1ΔID on the ability of colony forming units of the progenitors derived from colony forming unit (CFU) granulocyte-erythroid-monocyte-megakaryocyte (CFU-GEMM) downstream of hematopoietic stem cells (HSC; see Fig. 1C). The number of CFU-GEMM colonies was decreased 71.2% in *Thra1PV* mice compared with WT mice (Fig. 1C-b, bar 3), but was corrected to only 17% reduction in *Thra1PV* *Ncor1ΔIDΔID* mice (Fig. 1C-b, bar 4). The number of burst-forming unit erythroid (BFU-E) and CFU erythroid (CFU-E) was also decreased 81.5% and 68.0%, respectively, in *Thra1PV* mice (bars 3 in panels c and d, Fig. 1C), but was corrected to the reduction of 54.9% and total recovery, respectively, in *Thra1PV* *Ncor1ΔIDΔID* mice (bars 4 in panels c and d). The number of CFU-granulocyte (CFU-GM) and CFU-megakaryocyte (CFU-MK) was decreased 70.8% and 78.8%, respectively, in *Thra1PV* mice (bars 3 in panels e and f, Fig. 1C), but was corrected to only reduction of 48.3% and 61.8%, respectively, in *Thra1PV* *Ncor1ΔIDΔID* mice (bars 4 in panels e and f). These results indicated that the expression of NCO1ΔID in *Thra1PV* mice could ameliorate the impaired capacity of progenitor cells to differentiate from GEMM to the mature erythrocytes and megakaryocytes in *Thra1PV* mice.

**Expression of NCO1ΔID rescues the terminal erythropoiesis in Lin negative (Lin-) bone marrow cells.** Because patients with mutations of the *THRA* gene exhibit anemia, we focused our studies on the erythroid lineage. To further confirm that the effect of NCO1ΔID on the maturation of erythrocytes in *Thra1PV* mice, we used an *in vitro* terminal differentiation system. Using an equal number of total bone marrow cells from *Thra1PV* mice and *Thra1PV* *Ncor1ΔIDΔID* mice (Fig. 2A-a and e, respectively; the mature erythrocyte population shown in the gated boxes identified by Ter119+ with low FSC population), we isolated lineage depleted bone marrow cells (Lin-BM) as shown in Fig. 2A-b and f, for *Thra1PV* mice and *Thra1PV* *Ncor1ΔIDΔID* mice, respectively. After induction of terminal differentiation, we found 14% and 17%, respectively, of Ter119+ with low FSC population (gated in red boxes). The quantitative comparison shows that the expression of NCO1ΔID led to a 18% increase in matured erythrocytes in *Thra1PV* *Ncor1ΔIDΔID* mice as compared with *Thra1PV* mice (bar 2 versus bar 1, Fig. 2B). These findings indicated that the decreased number of mature erythrocytes in *Thra1PV* mice is markedly increased by the expression of NCO1ΔID.
TRα1PV-mediated repression of erythropoietic genes is de-repressed by the expression of NCOR1ΔID in the bone marrow of Thra1PV/+ mice. To understand how the expression of NCOR1ΔID ameliorated the erythroid disorders in Thra1PV/+ mice, we analyzed the expression of key erythroid regulators in Thra1PV/+Ncor1ΔID/ΔID mice. The GATA1 (erythroid transcription factor; GATA-binding factor 1) is essential for erythroid development by regulating a large ensemble of genes that mediate both the development and function of red blood cells. We have recently shown that the Gata1 gene is directly regulated by TRα1 and T3, and that TRα1PV acted to repress its expression in the bone marrow of Thra1PV/+ mice. Interestingly, the TRα1PV-mediated repression of the Gata1 gene was totally de-repressed by the expression of NCOR1ΔID (bar 4 versus bar 3, Fig. 3A). The expression of Gata1 mRNA in the bone marrow of Ncor1ΔID/ΔID mice was similar to that in WT mice (bar 2 in Fig. 3A). We further showed that GATA1 protein abundance was detected by co-immunoprecipitation assay in WT mice and Ncor1ΔID/ΔID mice (lanes 7 & 8, Fig. B-I), but was too low to be detected in the bone marrow of Thra1PV/+ mice under identical experimental conditions (lane 9, Fig. 3B-I). Remarkably, a similar level of GATA1 protein as in WT mice was found in the bone marrow of Thra1PV/+Ncor1ΔID/ΔID mice.

Figure 1. The expression of NCOR1ΔID partially corrects the abnormal blood indices, bone marrow cell number and colony forming ability of progenitors in Thra1PV/+ mice. (A a–d) Peripheral blood indices as marked among adult mice (3–5 months old) with indicated genotypes (n = 13–58). P values are indicated. (A e) Serum EPO levels from mice with 4 genotypes as marked (n = 4–6) were determined as described in Methods, p values are indicated, NS, not significant. (B) Total bone marrow cells in among adult mice with indicated genotypes (n = 4–12). (C-a) Schematic representation of the erythroid lineage. The CFU-GEMM (panel b), BFU-E (panel c), CFU-E (panel d), CFU-GM (panel e) and CFU-MK (panel f) colonies from total bone marrow cells in mice with the indicated genotypes. The p values are indicated (duplicates in each assay; n = 5–7).
ID in expression of NCOR1Δ1PV to interact with NCOR1ΔID mice. (A) Total bone marrow profile from Thra1PV/+ mice (A-a) and Thra1PV/+Ncor1ΔID/ΔID mice (A-e). (Ter119+ FSClow) population is boxed in red. Population of Lin-BM cells from Thra1PV+ (A-b) and Thra1PV+ Ncor1ΔID/ΔID (A-f) mice. Terminal induced differentiated Ter119+/FSClow population is boxed in red (A-c for Thra1PV/+ mice and A-g for Thra1PV+/+Ncor1ΔID/ΔID mice). (B) Quantitative analysis shows the fold changes of erythrocytes after terminal erythrocyte differentiation of Lin-BM cells of Thra1PV/+ mice and Thra1PV+/+Ncor1ΔID/ΔID mice. P-values are indicated (mean ± SEM; n = 3).

The inability of TRα1PV to interact with NCOR1ΔID leads to the reversal in the expression of the Gata1 gene in Thra1PV+ΔID/ΔID mice. Next we sought to understand the molecular basis by which the expression of NCOR1ΔID rescued the erythroid abnormalities caused by TRα1 mutants in Thra1PV+ΔID/ΔID mice. Previously, we have elucidated that the Gata1 gene is directly and positively regulated by TRα1 via binding to one positive thyroid hormone response element (denoted as TRE2) on the promoter of the Gata1 gene18. Using specific antibody against TRα1 (designated as C4) in ChIP analysis, we found a strong binding of TRα1 to TRE2 in the bone marrow of erythroid WT mice (bar 2 versus bar 1, Fig. 4A). In mice expressing NCOR1ΔID, similar binding of TRα1 to TRE2 as in WT mice was found in the bone marrow of Ncor1ΔID/ΔID mice (bar 4 versus bar 3). As expected, a decreased binding of TRα1 to TRE2 was detected in the bone marrow of Thra1PV+ΔID/ΔID mice (bars 6 and 8 versus bar 2, Fig. 4A) because anti-TRα1 antibody C4 cannot recognize TRα1PV. However, significant binding of TRα1 to TRE2 was detected (compare bars 6 to 5, Fig. 4A). To demonstrate the binding of TRα1PV to TRE2, we used anti-TRα1PV specific antibodies, T1, in the ChIP assays. As
shown in Fig. 4B, T1 did not recognize TRα1 in the WT mice (bar 2, Fig. 4B), nor in Ncor1ΔID/ΔID mice (bar 4). In contrast, specific binding of TRα1PV to TRE2 was detected (compare bar 6 with bar 5), indicating that TRα1PV was bound to the promoter of the Gata1 gene. A low but not significant binding of TRα1PV was detected in Thra1PV/+ mice (bar 8, Fig. 4B). Using anti-NCOR1 antibody in ChIP analysis, we detected a significantly higher recruitment of NCOR1 by TRE2-bound TRα1PV to the promoter of the Gata1 gene in Thra1PV/+ mice (bar 6, Fig. 4C) than in Thra1PV/+ Ncor1ΔID/ΔID mice (bar 8, Fig. 4C). Very low binding of NCOR1 to TRE2-bound TRα1 was detected in euthyroid WT mice. Virtually no NCOR1 binding to TRE2-bound TRα1

Figure 3. TRα1PV-mediated repression of key erythroid genes is de-repressed by the expression of NCOR1ΔID in the bone marrow of Thra1PV/+ Ncor1ΔID/ΔID mice. (A) Relative Gata1 mRNA levels. (B) GATA1 protein levels determined by co-immunoprecipitation (B–I) (Full gel/blot is shown in the Supplemental Figure I) and quantitative data (B–II). (C) Klf1 mRNA with indicated genotypes (P values indicated; mean ± SEM; n = 3 mice per group; each triplicates). (D–I) Western blot analysis of KLF1 protein abundance (Full gel/blot is shown in the Supplemental Figure II), and (D–II) quantification of the band intensities in (D–I) (n = 3 mice per group). Relative mRNA levels of β-globin (E), Bzrp (F), Ahsp (G), and Dematin (H) in total bone marrow cells of mice with indicated genotypes were determined by quantitative real-time PCR. Values are mean ± SEM (n = 2–3 mice per group; each triplicates).
gene to suppress its expression, thereby impairing erythropoiesis 18. On the basis of Gata1 the promoter of the critical roles in erythroid development. Moreover, the findings from these two studies are complementary in that erythroid defects in the bone marrow of adult mice. While the findings of these two studies were derived from antibody in ChIP analysis, we found that only TRE2 bound-TRα1PV-NCOR1 complex recruited HDAC3 to α gene in α ID led to reversal in expression of the of TRα1PV with NCOR1 ΔID leads to the reversal in the expression of the TRα1PV act as a dominant negative suppressor to induce erythroid disorders. We took advantage of a mutant mouse that expresses the NCOR1ΔID mutant to test the hypothesis that the loss of the interaction of TRα1 mutants with NCOR1ΔID could reverse the deleterious effects of TRα1 mutants in erythropoiesis. Indeed, we found that the expression of NCOR1ΔID in Thra1PV mice led to partial reversal in the erythroid blood indices, corrected differential potential of progenitors in the erythroid lineage, increased the capacity of the terminal differentiation, and the reversal of the TRα1PV-mediated repression of key erythroid regulatory genes. These results indicated that aberrant association of NCOR1 with TRα1 mutants in vivo underlies the pathogenesis of erythroid disorders caused by TRα1 mutations.

The important role of NCOR1 in erythropoiesis has been documented in a mouse model deficient in NCOR1 (Ncor1ΔID−/− mice)36. Ncor1ΔID−/− mice exhibit anemia at E13.5, and the severity of the anemia increases with age, resulting in eventual death. Phenotypic analysis during embryoid development showed that NCOR1 deficiency leads to defects in definitive erythropoiesis. The fetal liver size of Ncor1ΔID mice was about half that of Ncor1+/+ litter mates. Further, the BFU-E forming capacity was reduced in embryos at E13.5-E14.5 of Ncor1ΔID−/− mice. These observations clearly demonstrated that NCOR1 regulates erythroid development.

In line with these findings from Ncor1ΔID−/− mice, the present studies highlighted the critical regulatory role of NCOR1 in erythropoiesis using Thra1PV+/+ mice expressing NCOR1ΔID. Our studies focused on dissecting the erythroid defects in the bone marrow of adult mice. While the findings of these two studies were derived from two different mutant mice, the collective evidence allowed us to reach the same conclusion that NCOR1 plays critical roles in erythroid development. Moreover, the findings from these two studies are complementary in that the deficiency in functional NCOR1 caused defective erythropoiesis during development as shown in Ncor1ΔID−/− mice and that the defects could persist into adulthood as shown in Thra1PV+/+ Ncor1ΔID−/− mice. These two studies jointly indicate the need of NCOR1 in normal erythropoiesis not only during development, but also in the maintenance of normal erythropoiesis in adults. However, how NCOR1 was involved in regulating definitive erythropoiesis was not elucidated in Ncor1ΔID−/− fetal livers 36. By using Thra1PV+/+ Ncor1ΔID−/− mice, we found that one mechanism by which NCOR1 could impact erythropoiesis was via its aberrant association with TRα1PV. NCOR1 was recruited by TRE-bound TRα1PV on the promoter of the Gata1 gene to suppress its expression, thereby impairing erythropoiesis 18. On the basis of these findings, it is reasonable to postulate that TRα1 could be involved in the NCOR1 functions in definitive erythropoiesis during development. The association of NCOR1-TRα1 with certain erythroid regulatory genes to suppress their expression could be critical in definitive erythropoiesis. The loss of the suppression function

Figure 4. The inability of TRα1PV to interact with NCOR1ΔID leads to the reversal in the expression of the Gata1 gene in Thra1PV+/+ Ncor1ΔID−/− mice. (A) ChIP assay was carried out using normal mouse IgG (solid bar) or anti-WT TRα1 (C4, n = 6–8) antibody (open bar), and (B) anti-TRα1PV-specific antibodies, T1, or normal rabbit IgG (solid bar) or (C) anti-NCOR1 antibody (open bar), (D) normal mouse IgG (solid bar) or anti-HDAC3 antibody (open bar), from total bone marrow cells of mice with indicated genotypes as described in Materials and Methods (n = 3–4 mice for each group).
of erythroid regulatory genes in Ncor1ΔID mice would result in defective erythropoiesis during development. Identification of these genes in the future would help us understand the functions of not only TRα, but also NCOR1 in erythropoiesis.

Previously we have shown that expression of NCOR1ΔID in Thra1PV mice ameliorated the abnormalities in the pituitary–thyroid axis and partially reverted infertility, growth retardation, impaired bone development, and lipid abnormalities. The present studies showing that TRα1 mutants caused erythroid disorders further expanded the scope of the RTHα-resistant target tissues regulated by NCOR1 and further strengthened the conclusion that aberrant recruitment of NCOR1 by TRα1 mutants leads to clinical hypothyroidism in patients. However, it is noteworthy that the extent of the correction of abnormalities in the Thra1PV mice by the expression of NCOR1ΔID varies across target tissues. As shown previously, the mildly elevated serum total T3 and TSH levels were totally corrected to the basal levels of WT mice by the expression of NCOR1ΔID. However, similar to those observed in growth, bone length, and white adipose tissues, the correction in the erythroid defects was partial as shown in the incomplete recovery of total bone marrow cells and the colony forming capacity of progenitors in the erythroid lineage (see Fig. 1). The incomplete recovery would suggest that the dominant actions of TRα1 mutants could also be regulated by other nuclear co-repressors such as NCOR2/SMRT. Erythropoiesis is a complex biological process and is modulated by large networks of regulators. NCOR1 could be affecting only a subset of erythropoietic genes. Still, the finding of partial corrections of the erythroid abnormalities by NCOR1 is a step forward in understanding how mutations of the THRA gene leads to erythroid defects in patients.

Materials and Methods

Mice and treatment. All animal studies were performed according to the approved protocols of the National Cancer Institute Animal Care and Use Committee. Mice harboring the mutated Thra1PV gene (Thra1PV mice) were prepared and genotyped by PCR as described earlier. Ncor1ΔID mice were prepared as described previously. The Thra1PV mice were crossed with Ncor1ΔID mice to obtain different genotypes for studies. These mice were intercrossed several generations, and littersmates with a similar genetic background were used in all experiments.

Cells. Bone marrow cells were isolated from femurs and tibiae of mice with different genotypes (Thra1PV+/−, Ncor1ΔID+/−, Thra1PV+/−Ncor1ΔIDΔID, Thra1PV+/−Ncor1ΔID+, Thra1PV+/−Ncor1ΔID−ΔID, age: 3–5 months). Single cell suspensions were prepared by passing bone marrow through a 70 µM cell strainer.

Peripheral blood profile analysis. For analysis of complete blood counts, peripheral blood was collected in a heparinized microtube and analyzed by hematology analyzer (HEMAVED HV950FS, Drew Scientific, Miami Lakes, FL).

Determination of serum mouse erythropoietin (EPO). Erythropoietin levels were analyzed in mice (3–5 months old) with 4 genotypes. Collected blood were allowed to clot for 2 hours at room temperature before centrifuging for 20 minutes at 2000 X g. Serum EPO levels were measured by Quantikine mouse erythropoietin kit (Cat.no: MEP00B, R&D Systems, Minneapolis, MN, USA). Serum erythropoietin levels were quantified using a microplate reader set to 450 nm.

Colony assays. Bone marrow cells were isolated from femurs and tibiae of mice with different genotypes at age of 3–5 months. To detect burst-forming units-erythroid (BFU-E), granulocyte/macrophage progenitor (CFU-GM), and multi-potential progenitor cells (CFU-GEMM) colonies, 4×10⁴ bone marrow cells were mixed with semisolid medium (Methocult GF M3434) by vortexing. The colony forming units-erythroid (CFU-E) colonies, 3.2×10⁴ bone marrow cells were mixed with semisolid medium (Methocult M3334). The colony forming units-megakaryocytes (CFU-MK) colonies, 8×10⁴ bone marrow cells were mixed with semisolid medium (MethocultGF M04974, supplemented with 10 ng/mL Interleukin (IL)-3, 20 ng/mL Interleukin (IL)-6, 50 ng/mL thrombopoietin (TPO). All reagents and Methocults were purchased from STEMCELL Technologies, Vancouver, BC. Bone marrow cells in medium were seeded in duplicates on 6- well plates and cultured for 8 days for BFU-E, 10⁴ bone marrow cells were mixed with semisolid medium (Methocult M3334). The colony forming units-erythroid (CFU-E) colonies, 3.2×10⁴ bone marrow cells were mixed with semisolid medium (Methocult M3334). The colony forming units-megakaryocytes (CFU-MK) colonies, 8×10⁴ bone marrow cells were mixed with semisolid medium (MethocultGF M04974, supplemented with 10 ng/mL Interleukin (IL)-3, 20 ng/mL Interleukin (IL)-6, 50 ng/mL thrombopoietin (TPO). All reagents and Methocults were purchased from STEMCELL Technologies, Vancouver, BC. Bone marrow cells in medium were seeded in duplicates on 6- well plates and cultured for 8 days for BFU-E, CFU-GM, and CFU-GEMM, 2 days for CFU-E, or 6 days for CFU-MK for scoring.

RNA extraction and quantitative RT-PCR. Total RNA was isolated from bone marrow cells using Trizol (Thermo Fisher Scientific, Waltham, MA). RT-qPCR was performed with one step SYBR Green RT-qPCR Master Mix (Qiagen, Valencia, CA). The mRNA level of each gene was normalized to the GAPDH (glyceraldehyde-3-phosphate dehydrogenase) mRNA level. The primer sequences used in RT-qPCR are listed in Supplemental Table 1.

Western blot analysis and co-immunoprecipitation. Cell lysates from bone marrow were prepared similarly as described previously. The detection of KLF1 in the bone marrow of WT and mutant mice by western blot analysis was performed as described previously. For the detection of GATA1 proteins in the bone marrow of WT and mutant mice, bone marrow lysates (600 µg each) were first immunoprecipitated with rat anti-GATA1 antibody (4 µg) or mouse IgG (4 µg: negative controls) followed by pulling down the enriched GATA1-anti-GATA1 antibody-complex with protein G-agarose beads. GATA1 proteins were subsequently detected by western blot analysis as described above using rabbit anti-GATA1 antibody. The antibodies used are listed in Supplemental Table 1.

Chromatin immunoprecipitation assays (ChIP). ChIP assay with bone marrow cells was performed as described previously. Briefly, mouse bone marrow cells isolated from WT and mutant mice fixed in 1% of formaldehyde, washed, and sheared, followed by immunoprecipitation overnight at 4°C with IgG (control), anti-TRα1 monoclonal antibody (C4), Anti-nuclear receptor corepressor 1 (NcoR1) antibody (ChIP grade; ab24552). Quantitative PCR was performed to detect the upstream fragment in Gata1 genes with primer pairs (Supplemental Table 1). DNA binding was calculated as percentage of the input.
**In vitro terminal erythropoiesis assay.** For lineage depleted bone marrow cell preparation, lineage marker positive cells were depleted using the biotin based selection kit (cat # 19856, STEMCELL Technologies, Vancouver, BC) according to the manufacturer’s instructions. Lin-BM cells were seeded in fibronectin-coated wells (Corning Inc, Corning, NY). To induce erythropoiesis, Lin-BM cells were cultured as described. On the first day, Lin-BM cells were cultured in Iscove’s Modified Dulbecco’s Medium (IMDM medium) supplemented with 15% FBS, 1% detoxified bovine serum albumin (BSA; Sigma-Aldrich, St. Louis, MO), 200 µg/mL holo-transferrin (Sigma-Aldrich, St. Louis, MO), 10 µg/mL recombinant human insulin (Sigma-Aldrich, St. Louis, MO), 2 mM L-glutamine, 10^−4 M β-mercaptoethanol, 50 units/ml penicillin G, 50 µg/mL streptomycin (Thermo Fisher Scientific, Waltham, MA) and 2 U/mL Epo (STEMCELL Technologies, Vancouver, BC). The medium was replaced with IMDM with 20% FBS, 2 mM L-glutamine and 10^−4 M β-mercaptoethanol for erythropoiesis differentiation for the second and third day.

**Flow cytometry analysis.** All antibodies used in flow cytometry were from eBiosciences (Thermo Fisher Scientific, Waltham, MA). To exclude dead cells from analysis, 7-aminactinomycin D (7-AAD) was used. Antibodies were chosen to determine terminal erythropoiesis using Lin-bone marrow cells as follows: anti-Ter119 (TER-119, APC-eFluor® 780) and anti-CD44 (IM7, eFluor® 450). The flow cytometry analyses were performed on a BD LSR II flow cytometer (BD Bioscience, San Jose, CA) and analyzed with FloJo, LLC (Tree Star Inc, Ashland, OR).

**Statistical analysis.** All statistical analyses and the graphs were performed and generated using GraphPad Prism version 6.0 (GraphPad Software, La Jolla, CA). Student's t test was used to examine whether differences between groups are statistically different from each other. P < 0.05 is considered statistically significant. All data are expressed as mean ± SEM.

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Author Contributions
C.R. Han: Design experiments, Data curation, Methodology, Validation, and writing-original draft. S. Park: Data curation, Methodology, Validation. S.Y. Cheng: Conceptualization, Formal analysis, Funding acquisition, Project administration, Resources, Supervision, Validation, and Writing – original draft, Writing – review & editing.

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