Mutant KLF1 in Adult Anemic Nan Mice Leads to Profound Transcriptome Changes and Disordered Erythropoiesis

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Anemic Nan mice carry a mutation (E339D) in the second zinc finger of erythroid transcription factor KLF1. Nan-KLF1 fails to bind a subset of normal KLF1 targets and ectopically binds a large set of genes not normally engaged by KLF1, resulting in a corrupted fetal liver transcriptome. Here, we performed RNAseq using flow cytometric-sorted spleen erythroid precursors from adult Nan and WT littermates rendered anemic by phlebotomy to identify global transcriptome changes specific to the Nan Klf1 mutation as opposed to anemia generally. Mutant Nan-KLF1 leads to extensive and progressive transcriptome corruption in adult spleen erythroid precursors such that stress erythropoiesis is severely compromised. Terminal erythroid differentiation is defective in the bone marrow as well. Principle component analysis reveals two major patterns of differential gene expression predicting that defects in basic cellular processes including translation, cell cycle, and DNA repair could contribute to disordered erythropoiesis and anemia in Nan. Significant erythroid precursor stage specific changes were identified in some of these processes in Nan. Remarkably, however, despite expression changes in large numbers of associated genes, most basic cellular processes were intact in Nan indicating that developing red cells display significant physiological resiliency and establish new homeostatic set points in vivo.

In the inbred semi-dominant mouse model, Nan (neonatal anemia), heterozygotes suffer lifelong anemia due to a missense mutation (E339D) in KLF1 (Krüppel-like factor 1)1,2. Homozygotes die in utero (E10-11). Klf1 expression is restricted to megakaryocyte-erythroid progenitors and erythroid lineage cells where it plays a global role in lineage determination and initiating and maintaining the erythroid-specific transcriptome including expression of globin, membrane skeleton, heme biosynthetic, iron regulating, and cell cycle genes3–10. Indeed, characteristics of both hereditary spherocytosis and thalassemia are evident in adult Nan/+ (hereafter Nan) mice with membrane skeleton defects, decreased expression of adult globin, and increased expression of embryonic globin1,2. KLF1 is a C2H2 transcription factor with three tandem zinc fingers (ZFs), each recognizing a nucleotide triplet in its 9bp cognate DNA binding site, 5′-CCM-CRC-CCN3,11,12. Glutamic acid at position 5 in the middle triplet interacts with the middle triplet. Despite the conservative amino acid change to aspartic acid, the effect on DNA recognition is profound. Previous work1 showed that Nan-KLF1 binds normally to target genes containing guanine at position 5 in the middle triplet but fails to bind, or binds with reduced affinity, to genes containing adenine including β-globin (Hbb), the cell cycle regulator E2F2 (E2f2), and the membrane skeleton protein dematin (Dmtin). Notably, the normal Klf1 allele is expressed in Nan erythroid cells. Despite this, expression of genes that mutant Nan-KLF1 does not bind is somehow disrupted through an unknown mechanism.

Nan-KLF1 also ectopically binds sites not normally engaged by KLF113,14. In fetal liver, only 18% of up-regulated and 52% of down-regulated genes in Nan overlapped known KLF1 targets. ChiPseq confirmed...

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ectopic Nan–KLF1 binding to an altered consensus sequence, CCM-NGC-CCN, with the result that >60% of Nan-KLF1 occupied sites do not overlap wild type (WT) KLF1 sites. Ectopic binding contributes to anemia in Nan through extrinsic mechanisms15. For example, hepcidin (Hamp) and interferon regulatory factor 7 (Irf7), normally expressed in adult liver13 and macrophages13, respectively, show dramatically increased expression in Nan fetal liver13 and adult WT spleen and bone marrow13 leading to increased serum hepcidin and interferon beta. Increased hepcidin with markedly decreased erythroferrone, which is not bound by Nan-KLF115, limits iron availability18,19. Interferons inhibit erythropoiesis at the BFU- and CFU-E stages20.

Nan is relevant to human erythroid-related disease. Human KLF1 mutations lead to benign defects21,22 and anemia, sometimes severe23. For example, a different substitution (E325K) at the position corresponding to Nan causes congenital dyserythropoietic anemia (CDA) type IV22. Prior transcriptome studies in Nan focused on fetal liver erythroid cells1,3,15. Here, we performed RNAseq in adult spleen erythroid precursors, comparing Nan to littermates made anemic by phlebotomy (WT-PHB). The use of anemic WT controls allowed identification of expression changes occurring primarily in response to the Nan KLF1 defect. We show that differential expression in Nan differs in fetal liver and spleen erythroid cells, that expression variation is driven primarily by precursor cell type with mutation effects most prominent in late erythroblasts, and that progression of erythropoiesis is significantly impaired. PCA reveals two major patterns of differential gene expression and predicts that defects in basic cellular processes (e.g., translation, cell cycle) contribute to anemia in Nan. Indeed, erythroid precursor stage specific alterations in some of these processes were identified. Overall, however, despite extensive differential expression, the basic cellular machinery appears largely intact in Nan, suggesting substantial tolerance to transcriptional changes and physiological resiliency in developing red blood cells (RBCs) in vivo.

Results

Distortion of the Nan transcriptome increases during differentiation. Both Klf1 alleles, E339 and D339, are expressed in Nan spleen erythroid precursors (Fig. 1a), as previously demonstrated in fetal liver1. We performed RNAseq analysis on sorted erythroid precursors (pro- and basophilic erythroblasts) from adult male Nan and WT-PHB spleens to identify global transcriptome changes specific to Nan-KLF1. Peripheral blood studies confirm that WT-PHB mice develop anemia with significantly decreased RBC count, hemoglobin, and hematocrit; and increased RBC and hemoglobin distribution width, reticulocyte percentage, and spleen weight (Supplementary Table S1). Cell morphology is similar in Nan and WT-PHB with significant anisocytosis and polychromasia (Fig. 1b). Poikilocytosis including fragmented cells is rare. Circulating erythroblasts are not seen. Total bilirubin and LDH, indicators of hemolysis, did not differ significantly between Nan and WT-PHB (Supplementary Table S1); bilirubin, but not LDH, was increased over WT in both.

We obtained ~36 million mapped reads per sample (3 male mice per genotype with 4 erythroid precursor populations each for a total of 24 samples with 3 technical replicates each) and analyzed for differential expression. The number of differentially expressed genes (DEGs) increases dramatically during differentiation (Table 1), likely reflecting progressively increasing Klf1 expression (Fig. 1c; expression profiles can be generated at http://nanexpression.mdibl.org/). In total, 6,965 unique genes were differentially expressed across the combined precursor populations. Surprisingly, these overlapped with only 61% of fetal liver DEGs. Analysis of gene ontology (GO) terms using DAVID23 by precursor cell type revealed no significant functional annotations (Benjamini corrected p-values) associated with pro- and basophilic erythroblasts and just three associated with polychromatophilic erythroblasts. A great many, however, emerge in orthochromatophilic erythroblasts including processes that could potentially contribute to anemia (translation, RNA processing, cytoskeletal organization, apoptosis, oxidation-reduction, cell cycle) (Fig. 1d,e). Indeed, cytoskeletal (membrane skeleton) defects were documented previously in Nan3,13. The most significant differential genes in orthochromatophilic erythroblasts, suggesting that failure to engage normal DNA targets, or engaging with altered targets, is relevant to human erythroid-related disease. Human KLF1 mutations lead to benign defects21,22 and anemia, sometimes severe23. For example, a different substitution (E325K) at the position corresponding to Nan causes congenital dyserythropoietic anemia (CDA) type IV22. Prior transcriptome studies in Nan focused on fetal liver erythroid cells1,3,15. Here, we performed RNAseq in adult spleen erythroid precursors, comparing Nan to littermates made anemic by phlebotomy (WT-PHB). The use of anemic WT controls allowed identification of expression changes occurring primarily in response to the Nan KLF1 defect. We show that differential expression in Nan differs in fetal liver and spleen erythroid cells, that expression variation is driven primarily by precursor cell type with mutation effects most prominent in late erythroblasts, and that progression of erythropoiesis is significantly impaired. PCA reveals two major patterns of differential gene expression and predicts that defects in basic cellular processes (e.g., translation, cell cycle) contribute to anemia in Nan. Indeed, erythroid precursor stage specific alterations in some of these processes were identified. Overall, however, despite extensive differential expression, the basic cellular machinery appears largely intact in Nan, suggesting substantial tolerance to transcriptional changes and physiological resiliency in developing red blood cells (RBCs) in vivo.

Nan-specific expression differences in fetal liver and spleen. We showed previously that expression of Hamp and Irf7 is significantly up-regulated in Nan whole fetal liver and adult whole spleen vs. WT fetal liver and spleen15, and confirmed upregulation in fetal liver here (Supplementary Table S2). Importantly, Hamp and Irf7 upregulation was confirmed by qRT-PCR in fetal liver cells depleted of adherent cells and expanded in culture with erythropoietin in previous studies15. These data indicate that Hamp and Irf7 are ectopically activated by Nan-KLF1 in fetal liver, as neither is normally expressed in erythroid cells17,18. Here, however, we find no Hamp expression in Nan or WT-PHB spleen erythroid precursors (http://nanexpression.mdibl.org/). Thus, Hamp expression driven by mutant Nan-KLF1 differs in fetal liver and anemic adult spleen erythroid cells even though both are stress environments. On the other hand, Irf7 is expressed at equally high levels in both Nan and WT-PHB spleen (Fig. 2a), indicating that Irf7 can also be upregulated as a general consequence of stress erythropoiesis in the spleen and is not specific to Nan.

Niche-specific differential expression of Nan β-like globin genes is also seen. Embryonic βh1 (Hbb-bh1) is significantly upregulated in Nan vs. WT fetal liver and non-anemic (hereafter WT) spleen15,13. However here, no differential upregulation is seen in adult Nan vs. WT-PHB spleen precursors (Fig. 2b), indicating that increased expression in the spleen is due to stress erythropoiesis in response to anemia, as documented previously13,15. βh1 is expressed at low levels in proerythroblasts and basophilic erythroblasts where the embryonic globin repressor Bcl11a is highly expressed in both Nan and WT-PHB (Fig. 2c). Bcl11a expression falls in poly- and orthochromatophilic erythroblasts; the decrease is greater in Nan than WT-PHB, but their βh1 expression is equivalent in these precursors (Fig. 2b). We confirmed βh1 and Bcl11a expression patterns by qRT-PCR (Fig. 2d). Two
Figure 1. Aberrant erythroid transcriptome in adult anemic Nan mice. (a) cDNA sequence chromatograms showing transcription of Klf1 alleles in wild type control (WT, +/+) and sorted Nan spleen erythroid cells. Both the normal and mutant alleles are equally expressed in Nan throughout terminal differentiation. (b) Peripheral blood smears from untreated (non-anemic) WT and phlebotomized WT control (WT-PHB) and Nan adult mice. Bars, 10 μM. (c) Expression level (log2 counts) of Klf1 in WT-PHB and Nan spleen erythroid precursors. Differences in Klf1 expression do not meet filtering criterion (fold change ≥2, false discovery rate ≤0.05) for differentially expressed genes in any of the precursor populations. The graph shows the three technical replicates for each of the 3 biological replicates, giving 9 points per genotype per cell type. Tick marks, mean ± SD. (d,e) Top functional annotations associated with up- (red) and down- (green) regulated gene expression in Nan spleen poly- and orthochromatophilic erythroblasts suggest multiple mechanisms could contribute to anemia in Nan. Note: In panel e terms broadly associated with translation and RNA processing are shown separately (middle) in order to display other highly significant functional annotations (bottom).
matophilic erythroblasts in WT-PHB precursors, and how expression diverges from WT-PHB in Nan of gene expression emerge in terms of how expression changes during differentiation from pro- to orthochromatophilic differentiation proceeds with Nan-KLF1 driving increasingly aberrant transcription in later stages of differentiation.

The PCA confirms the role of cellular differentiation in driving expression variation as terminal differentiation proceeds with Nan-KLF1 driving increasingly aberrant transcription in later stages of differentiation.

Patterns of gene expression in Nan. We generated pairwise correlations of all samples using a reduced set (~8,600) of high variance genes. Hierarchical clustering based on these correlations shows that expression differences are much more severe in late (poly- and orthochromatophilic) than in early (pro- and basophilic) erythroblasts (Fig. 3a), confirming that progression of differentiation contributes a major source of expression variation. Indeed, PCA of the spleen expression matrix reveals that the first principle component (PC1) is dominated by cell type, whereas PC2 separates the samples within each cell type by genotype with modest effect of cell type at the pro- and basophilic stages but significantly more dramatic effects at the poly- and, particularly, the orthochromatophilic stages (Fig. 3b). PC1 and PC2 account for 59.5% and 13.9% of expression variation, respectively. The PCA confirms the role of cellular differentiation in driving expression variation as terminal differentiation proceeds with Nan-KLF1 driving increasingly aberrant transcription in later stages of differentiation.

When each PC is plotted separately as a function of genotype and precursor cell type, two major patterns of gene expression emerge in terms of how expression changes during differentiation from pro- to orthochromatophilic erythroblasts in WT-PHB precursors, and how expression diverges from WT-PHB in Nan precursors. Because PCA groups genes by their correlation with the components and represents both correlated and anti-correlated patterns, complimentary patterns of expression (meaning both increasing and decreasing) are included within each PC. Thus, the PC1 expression pattern is one of gradual, progressive change in expression in WT-PHB (either increasing or decreasing steadily) during differentiation, while expression in Nan shows a parallel overall course but progressively diverges from WT-PHB during differentiation such that expression differences are most prominent in poly- and orthochromatophilic erythroblasts (Fig. 3c, top panel). Examples of PC1 positively correlated genes (expression decreases during differentiation) include Sf3b3 (splicing factor 3b, subunit 3) (Fig. 3d) and Steap3 (STEAP family member 3) (Fig. 3e). Epor (erythropoietin receptor) exemplifies a PC1 negatively correlated gene (expression increases during differentiation) (Fig. 3f).

In the second expression pattern, that for PC2, expression in WT-PHB trends consistently up or down as differentiation proceeds (Fig. 3c, bottom panel). In Nan, expression is consistently higher or lower than WT-PHB at all stages and follows the same overall trend (either increasing or decreasing with differentiation) until the orthochromatophilic erythroblast stage, where a sharp, often dramatic reversal occurs in Nan (Fig. 3c, bottom panel). Moreover, while the direction of the differential expression in Nan is constant, the magnitude generally increases during differentiation. Thus, PC2-correlated genes show more severe dysregulation throughout differentiation in Nan. Proxl3 (peroxiredoxin 3) is positively correlated with PC2 and its expression increases with differentiation; Srx4 (sorting nexin 4) and Clmb1 (beta catenin) are negatively correlated (Fig. 3g–i) and expression decreases. The complete list of genes correlated with PC1 and PC2 is provided in Supplementary Table S9.

We predict that the overall functional output of the altered Nan transcriptome will be most influenced by genes highly correlated with PC2, where Nan expression diverges the greatest throughout terminal differentiation, and to a lesser degree, PC1. For genes with the most significant negative correlations with PC1 (expression increases during differentiation), the top GO functional annotations include ubiquitination, autophagy/mitophagy, cell cycle, transcription, and erythroid differentiation (Fig. 4a). Conversely, the top annotations for the most significant PC1 positively correlated genes (expression decreases) are dominated by terms related to translation and mRNA processing. As PC1 is driven largely by cell type with little mutation effect, these up- and down changes in gene expression are consistent with major processes occurring during normal terminal differentiation. For PC2, where mutation effects dominate, protein transport, transcription and translation are the most significant GO functional annotations for positively correlated genes (expression increases); for negatively correlated genes (expression decreases) cell cycle, cell division, oxidation-reduction, and DNA repair dominate (Fig. 4b).

Erythropoiesis in Nan. Previous studies revealed membrane skeleton defects and impaired globin expression in Nan. Here, based upon striking differential expression and predictions arising from PCA, we tested the hypothesis that additional processes go awry and contribute to anemia in Nan. We first examined erythropoiesis in Nan compared to both anemic and non-anemic WT spleen and bone marrow to achieve a full picture of erythroid status in Nan. Significantly, unlike KLF1-null mice, expression of cell surface markers used to assess erythropoiesis by flow cytometry (Ter119, CD71, CD44) is normal or near normal in Nan (Supplemental Fig. 2).

### Table 1. Differentially expressed genes in Nan spleen erythroid precursors.* 

| DEGs | Pro | Baso | Poly | Ortho |
|------|-----|------|------|-------|
| Up-regulated | 505 | 860 | 1438 | 2209 |
| Down-regulated | 535 | 799 | 1241 | 2916 |

*FC ≥ 2, FDR = 0.05.
Figure 2. Nan- and erythroid niche-specific expression changes. Expression of (a) Irf7 and (b) Hbb-bh1 is not differentially regulated in Nan vs. WT-PHB spleen erythroid precursor cells while both are markedly upregulated in Nan fetal liver (fold change in Nan fetal liver from Supplemental Table 2 in blue). (c) Expression of Bcl11a. Fold change vs. WT-PHB in blue. (d) Confirmation of Hbb-bh1 (top) and Bcl11a (bottom) expression by quantitative RT-PCR. AU, arbitrary units, no significant differences. n = 3 for Nan and WT-PHB. (e) Hbb-bh2 is progressively upregulated during erythropoiesis in Nan, reaching a 534-fold increase at the orthochromatophilic stage. (f) Confirmation of dramatically up-regulated Hbb-bh2 expression in Nan by quantitative RT-PCR. Average values for WT-PHB are shown in blue to facilitate comparison to Nan. AU, arbitrary units; *p < 0.01 and **p < 0.001 Nan vs. WT-PHB. n = 5 (Nan) and 6 (WT-PHB) except Ortho (n = 2 and 3, respectively). (g) Adult β-globin, Hbb-bs, is downregulated in Nan pro- and basophilic erythroblasts. The duplicated adult globin locus, Hbb-bt, shows the same pattern (http://nanexpression.mdibl.org/). Neither Hbb-bh2 nor Hbb-bt are differentially expressed in Nan whole fetal liver. Tick marks, mean ± SD. Note: All expression values are log2-transformed, normalized counts; therefore, values below 1 are considered background (no expression).
Figure 3. Principle component analysis (PCA). (a) Hierarchical clustering showing that expression differences are greatest in later stages of development (poly- and orthochromatophilic erythroblasts). (b) The first PC is dominated by cell type and accounts for 59.5% of the variation in expression while PC2 (13.9%) separates the samples within each cell type by genotype. (c) PC1 (top) and PC2 (bottom) plotted by genotype as differentiation proceeds from pro- to orthochromatophilic erythroblasts reveal two major patterns of differential expression in Nan. (d) Sf3b3 (splicing factor 3b, subunit 3) and (e) Steap3 (STEAP family member 3) are positively, and (f) Epor (erythropoietin receptor) negatively correlated with PC1 where the most prominent expression differences are late in differentiation at the orthochromatophilic stage. (g) Prdx3 (Peroxiredoxin 3) is negatively correlated with PC2 while (h) Snx4 (Sorting nexin-4) and (i) Ctnnb1 (β-catenin) are positively correlated with PC2. Genes correlated with PC2 show severe dysregulation throughout differentiation in Nan with a striking reversal at the orthochromatophilic stage. Tick marks, X ± SD.
Spleen total cell counts (Fig. 5a) are significantly higher in Nan and WT-PHB vs. WT, as expected due to anemia. Non-erythroid (CD45+ cells) are significantly increased in WT-PHB but not Nan compared to WT. Erythroid cell (CD45−) counts are increased approximately two-fold over WT in Nan and WT-PHB, but do not reach statistical significance. Nan erythroid cell counts showed more variability than WT and WT-PHB, overlapping WT in many cases (Supplemental Fig. 3), suggesting that stress erythropoiesis is inefficient to varying degrees in Nan mice. Bone marrow counts do not differ among the three groups (Fig. 5a).

Flow-cytometric analysis of differentiation29 revealed significantly delayed progression of erythropoiesis in Nan spleen, with cells accumulating at the proerythroblast stage compared to WT and at the baso- and polychromatophilic stages compared to both WT and WT-PHB (Fig. 5b). Orthochromatophilic erythroblasts are significantly increased in Nan compared to WT, but strikingly depleted compared to WT-PHB (Fig. 5b), as is also evident upon inspection of flow cytograms (Supplemental Fig. 4). The accumulation of earlier precursors with striking depletion of late orthochromatophilic erythroblasts in Nan vs. WT-PHB, where a progressive and dramatic increase in the percentages of each successive cell type is seen throughout differentiation, again suggests suboptimal stress erythropoiesis in Nan. A significant delay in erythropoiesis also occurs at the baso- and polychromatophilic stages in Nan bone marrow with depletion of orthochromatophilic erythroblasts compared to WT-PHB mice (Fig. 5b). These results confirm highly disordered basal and stress erythropoiesis in Nan.

Cell cycle. We asked if cell cycle defects contribute to accumulation of precursors and delayed progression of maturation in Nan. Both Nan and WT-PHB spleen proerythroblasts, which do not differ from each other, show evidence of accelerated cell cycling compared to WT; the percentage of proerythroblasts in G0/G1 is significantly decreased and the percentage in S phase significantly increased (Fig. 6a). This pattern was not maintained, however, in subsequent precursors. In WT-PHB, no further differences except a small but significant increase in orthochromatophilic erythroblasts in S phase is seen. As expected, nearly all orthochromatophilic erythroblasts have exited the cell cycle. The percentage of Nan bone marrow polychromatophilic erythroblasts in G2M is significantly higher than WT-PHB but does not differ from WT (Fig. 6a). Thus, with the possible exception of spleen
proerythroblasts, delayed differentiation with accumulation of precursors in Nan (Fig. 5) cannot be explained by cell cycle changes.

**Apoptosis.** The percentage of apoptotic proerythroblasts in both WT and Nan spleen is dramatically and significantly higher than WT-PHB (Fig. 6b). Apoptotic basophilic erythroblasts are also higher in WT and Nan vs. WT-PHB, but do not reach statistical significance in Nan. Notably, apoptosis in Nan does not differ from WT in any spleen precursor population. Apoptosis in WT-PHB spleen is maintained at a low and consistent level throughout terminal differentiation in the spleen, while early precursors in Nan undergo significant apoptosis. Thus, a high percentage of pro- and basophilic erythroblasts are lost in the Nan spleen when increased RBC production is needed.

**Figure 5.** Erythropoiesis is delayed in Nan spleen. (a) Total, CD45 positive and CD45 negative cell counts in WT, Nan, and WT-PHB spleen (n = 3, 10, and 4, respectively) and bone marrow (n = 4/group). (b) Quantitation of the different spleen populations demonstrates an accumulation of pro-, baso and- polychromatophilic erythroblasts in Nan spleen (top panel) and baso- and- polychromatophilic erythroblasts in Nan bone marrow (bottom). As a result, production of orthochromatophilic erythroblasts markedly lags that of WT-PHB. n = 4/group. *P < 0.05; **p < 0.01; ***p < 0.001.
In bone marrow, apoptotic Nan and WT-PHB pro- and basophilic erythroblasts do not differ from each other but are significantly less than WT (Fig. 6b). Apoptosis in Nan poly- and orthochromatophilic erythroblasts is decreased relative WT.

**ROS generation, mitophagy.** To measure ROS, we utilized peroxide- and superoxide-sensitive fluorescent dyes, DCF and DHE, respectively. Preliminary studies suggested increased ROS generation in Nan whole blood; subsequent analyses were done on CD71 and Ter119 stained whole blood to obtain quantitative data on
enriched reticulocyte (CD71hi) and mature erythrocyte (CD71lo) fractions (Supplemental Fig. 5). Superoxide and peroxide generation is increased in Nan mature erythrocytes compared to WT and WT-PHB (Fig. 7A). In Nan reticulocytes peroxide generation did not differ significantly from WT but was significantly less than WT-PHB. Reticulocyte superoxide generation in Nan and WT-PHB did not differ but both were increased compared to WT (Fig. 7A).

Increased ROS production can result from increased mitochondrial content due to defective mitophagy. Hence, we examined RBC mitochondria and ribosome content using Mitotracker Red and Thiazole Orange (Fig. 7B). A small increase in retained mitochondria is seen in Nan compared to WT (1.5 ± 0.8 vs. 0.2 ± 0.2, X ± SD, n = 3) although the difference was not statistically significant. Thus, increased ROS production without significant mitochondria retention contributes to hemolysis in Nan.

DNA damage. We examined DNA damage using a flow-cytometric peripheral blood micronucleus (MN) assay, which provides a quantitative measure of both spontaneous and radiation induced chromosome damage30,31. A small, but significant, increase in Nan reticulocyte MN is seen 48 hours post-irradiation (Fig. 7C,D), returning to baseline (time 0) at 72 hours. A similar pattern, although exaggerated compared to Nan, is seen in WT-PHB. The exaggerated response in WT-PHB may be due to acute induction of anemia in this group. Overall, these data indicate that WT-PHB and Nan are more susceptible to induced MN formation vs. WT; however, the DNA damage response is intact, as reticulocyte MN returned to baseline at 72 hours in both. RBC MN, indicative of spontaneous damage, did not change pre-and post-irradiation, as expected (not shown).

Discussion

Prior studies revealed that Nan-KLF1 acts as both a hypomorphic and neomorphic allele, failing to bind a subset of normal KLF1 target genes while inappropriately engaging non-KLF1 targets32. The result is a corrupted transcriptome that produces anemia by both cell intrinsic and extrinsic mechanisms33. Nan is relevant to human disease where a mutation at the corresponding position causes CDA type IV; the different mouse-human anemia phenotype may reflect the different amino acid changes.

Here, we compared adult Nan and WT-PHB sorted spleen erythroid precursors as a way to discern effects on gene expression specific to Nan-KLF1 induced anemia. We chose phlebotomy-induced anemia over phenyldrazine treatment to avoid confounding factors induced by phenylhydrazine such as widespread oxidative damage, severe hemolysis, and near 100% reticulocytosis. RNAseq revealed progressively increasing transcriptome changes in Nan during erythrocyth differentiation. PCA revealed that PC1 was dominated by cell type, contributing 60% of the expression variation. PC2, dominated by genotype effects, contributed 14% with strongest effects in poly- and orthochromatophilic cells reflecting increasing mutant Klf1 expression during differentiation. In total, 6,965 unique genes were differentially expressed with only a subset overlapping those identified in fetal liver.

We assessed overall functional effects of the extensive global transcriptome changes in the context of the whole animal. We first examined erythropoiesis by flow cytometry and found evidence of suboptimal stress erythropoiesis in Nan. Increased percentages of all erythroid precursors are seen in WT-PHB compared to WT spleen, as expected during stress erythropoiesis in response to anemia. Nan, however, shows a significant accumulation of pro-, baso-, and polychromatophilic erythroblasts compared to WT, and a significant accumulation of baso- and polychromatophilic erythroblasts in conjunction with a severe decrement of orthochromatophilic erythroblasts compared to WT-PHB. These data suggest that progression though the final stages of differentiation fails in a significant portion of Nan precursors, results concordant with analyses of Klf1-null fetal liver erythroblasts32. Strikingly, erythropoiesis in the Nan bone marrow shows the same overall pattern where a significant accumulation of polychromatophilic over orthochromatophilic cells is also evident. Despite the decrement in orthochromatophilic erythroblasts in Nan, reticulocytosis is significantly elevated, possibly due to slower maturation or faster kinetics in reaching the peripheral circulation over WT, or both. Whether the latter occurs at the level of enucleation and/or extrusion from the bone marrow and spleen remains to be elucidated. Nonetheless, total circulating RBCs (reticulocytes plus mature erythrocytes) are significantly decreased in Nan.

Two major patterns of expression divergence between WT-PHB and Nan during differentiation emerged when each PC was plotted individually as a function of cell type and genotype (Fig. 3c). Compared to PC1, PC2 show relatively greater divergence of expression in Nan precursors, particularly late precursors. Using top functional annotation GO terms associated with PC2 as a starting point, we identified processes including cell cycle, DNA damage, and apoptosis that could contribute to anemia with delayed progression of erythropoiesis and/or suboptimal stress erythropoiesis in Nan for functional screening. In terms of the cell cycle, significant changes indicative of increased cell cycle progression were detected in Nan and WT-PHB proerythroblasts compared to WT but were not maintained in later precursors. Overall, cell cycle changes fail to account for disordered erythropoiesis in Nan. Likewise, although Nan appears more susceptible to induced DNA damage, the DNA damage response is intact.

Interestingly, differences in apoptosis could at least partially underlie suboptimal stress erythropoiesis in Nan. Previous studies34 demonstrated that stress-responsive pro- and basophilic erythroblasts undergo a high rate of apoptosis in non-anemic WT mice, and that this response is suppressed in anemic mice under the influence of erythropoietin (Epo). Here, greater than 60% of WT spleen proerythroblasts and nearly 30% of basophilic erythroblasts are apoptotic; spleen apoptosis in early WT-PHB precursors, most strikingly proerythroblasts, is markedly decreased compared to WT (Fig. 6b). In Nan, despite anemia accompanied by high serum erythropoietin (Epo) levels15 and normal Epor expression in stress-responsive precursors (Fig. 3i; nanexpression.mdibl.org), apoptosis is inappropriately high, not differing significantly from WT. Epo-responsive mechanisms mediating suppression of apoptosis in anemia and facilitating erythroid expansion during stress in WT mice include downregulation of cell-surface expression of apoptosis-promoting Fas and FasL, and induction of anti-apoptotic Bcl-xL35-37. Bcl2l1 encoding Bcl-xL is also a well described direct KLF1 target gene46. KLF1 and STAT5 likely direct expression of Bcl2l1 via independent intrinsic enhancers45. Neither Fas nor FasL are expressed in WT-PHB or...
Figure 7. ROS generation, mitophagy, and DNA damage in Nan. (A) Quantitation of ROS production in Ter119 positive, CD71 low (mature red blood cells) and Ter119 positive CD71 high (reticulocytes) populations. As expected, DCF and DHE production in mature red cells is lower than in reticulocytes, n = 3 per group. (B) Flow cytometry of MitoTracker Red (MTR) and Thiazole Orange (TO) stained WT and Nan red blood cells to assess mitochondria and ribosome content three days post-phlebotomy. Early reticulocytes are ribosome-positive, mitochondria-positive (upper right quadrant), late reticulocytes ribosome-positive, mitochondria-negative (lower right quadrant), and retained mitochondria ribosome-negative, mitochondria-positive (upper left quadrant). Cytograms representative of 3 samples/group. (C) Representative micronucleus assay results in non-irradiated mice. (D) Quantitation of induced micronuclei 48 and 72 hours following irradiation. n = 4 per group; ***p < 0.001 (within group comparison at 0 and 48 hours). PI, propidium iodide.
**Methods**

**Animals.** All mice were maintained at The Jackson Laboratory in climate- and light cycle-controlled rooms. Acidified water and 5K52 chow (PMI LabDiet) were provided *ad libitum*. Nan is a fully inbred strain. The original mutant Nan mouse was a female offspring of an ENU–mutagenized C3H/101 F1 male and a “PT” female at the Medical Research Council, Harwell, U.K. The PT stock was a mixed-background mutation-testing stock. The anemic female was mated to an untreated C3H/101 F1 male, and the mutation was propagated by intercrossing offspring. Nan was imported to the Jackson Laboratory in 1994 and maintained in the research colony of Dr. Luanne Peters by brother-sister matings for more than 20 generations. Hence, according to mouse nomenclature rules (http://www.informatics.jax.org/mgihome/nomen/), the original Harwell Nan stock used in this study is a fully inbred subline designated Nan/Llp. The official designations for homozygous, heterozygotes and WT littermates are Nan/Llp Kfl+/Kfl+/Kfl+, Nan/Llp Kfl+/Kfl−, and Nan/Llp Kfl−/Kfl−, respectively. All experiments were carried out in accordance with National Institutes of Health Laboratory Animal Care Guidelines and were approved by the Animal Care and Use Committee (ACUC) of the Jackson Laboratory.

**Phlebotomy.** Adult WT littermates were phlebotomized via daily retro-orbital collection of ~350 μL whole blood followed immediately by intraperitoneal injection of sterile normal saline (37 °C) for four days.
Peripheral blood studies. Complete blood counts were determined using a Siemens Advia 120 analyzer. Blood smears were stained with Wright-Giemsa (Sigma-Aldrich). Serum lactate dehydrogenase (LDH) and bilirubin levels were determined using a Beckman AU680 analyzer.

Sorting of erythroid precursors. Single cell suspensions of adult (6–8 weeks) whole spleens were depleted of CD45+ cells using mouse CD45 MicroBeads (Miltenyi Biotec), stained (CD45-FITC, Ter119-PE and CD44-APC), and sorted on a FACSaria II flow cytometer (BD Biosciences) as described.

RNAseq. Total RNA was extracted with the PureLink RNA Mini Kit (Invitrogen) and libraries prepared using the KAPA mRNA HyperPrep Kit (KAPA Biosystems), according to the manufacturer’s instructions. Briefly, the protocol entails isolation of polyA containing mRNA using oligo-dT magnetic beads, RNA fragmentation, first and second strand cDNA synthesis, ligation of Illumina-specific adapters containing a unique barcode sequence for each library, and PCR amplification. Libraries were checked for quality and concentration using the D5000 ScreenTape assay (Agilent Technologies) and quantitative PCR (KAPA Biosystems), according to the manufacturer’s instructions. Paired end sequencing was performed on the HiSeq2500 System using TruSeq SBS Kit v4 reagents (Illumina). Reads were aligned to mouse genome GRCm38 with STAR Aligner. Quality control was performed with RSeQC and Picard (http://broadinstitute.github.io/picard/). Quantification to the mouse transcriptome ENSEMBL v85 was implemented using Gencode version M10. Count matrix normalization (rlog transformation) and differential expression analysis was performed with DESeq2. Differential expression analysis was performed on unnormalized counts assigned to each gene rather than FPKM (Fragments Per Kilobase of gene per Million reads), as recommended for DESeq2. Differentially expressed genes were extracted from DESeq2 output files using threshold values of ≥ 2 absolute fold change and ≤ 0.05 adjusted p-value.

Previously obtained fetal liver RNAseq data (SRA SRP061646, GEO GSE71396) was re-analyzed as above to facilitate valid comparisons between datasets.

To examine global patterns, further analysis was performed on the rlog regularized matrix. The regularized matrix was further Z-transformed by the function Z := (exp avg − avg stdev, where avg and stdev are the average and sample standard deviation of the regularized expression level of each gene across all samples. To focus the analysis on genes with appreciable expression and/or variation, the matrix was reduced from all 45,232 annotated genes to 8,668 with a 0.05 adjusted p-value.

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Sanger sequencing. cDNA was synthesized using the Ambion MessageSensor RT kit and Invitrogen random decamer primers (Invitrogen). PCR was performed using Invitrogen Platinum SuperFi PCR Master Mix (Invitrogen) with an annealing temperature of 65 °C. Kl/l forward (5′-AGACGCCAGGTGTGCCGGGACT) and reverse (5′-GGAGATGCTACTGAGGCTGAGCCT) primers were in exons 2 and 3, respectively, and spanned the Nan mutation. PCR products are purified using MagBio’s HighPrep PCR magnetic beads. Sequencing was performed using Applied Biosystems BigDye Terminator ready reaction kit Version 3.1, run on an Applied Biosystems 3730xl, and analyzed using Applied Biosystems Sequencing Analysis software version 5.2.

Quantitative real-time RT-PCR (qRT-PCR). cDNA was prepared as above. qRT-PCR reactions consisted of 4.5 μl of cDNA, 5 μl of TaqMan Gene Expression Master Mix and 0.5 μl of primers and probes (TaqMan® gene expression assay, assay ID: Hbb-bh1, Mm00433932_g1; Hbb-bh2, Mm01273444_g1; Bcl11a, Mm00479358_m1) (Applied Biosystems). Samples were prepared in triplicate. qRT-PCR was performed using a ViiA7 Real-Time PCR System (Applied Biosystems) (40 cycles at 95 °C for 15 sec and 60 °C for 1 min). The relative quantification of each gene was determined by the 2−∆∆CT method using control CT values and Gapdh (TaqMan® gene expression assay ID: Mm99999915_g1) expression for normalization.

Erythropoiesis, cell cycle and apoptosis. Terminal erythroid differentiation was monitored using CD44, Ter119, and forward scatter as described and analyzed using a LSRFortessa and FlowJo v9.9.4 software. Cell cycle and apoptosis were analyzed using BD Bioscience’s APC BrdU Flow Kit and Annexin V™, respectively, per manufacturer’s instructions.

ROS production, mitophagy and DNA damage. Intracellular ROS accumulation was determined using chloromethyl, dichlorodihydrofluorescein diacetate (M-H2DCFDA) and dihydroethidium (DHE) (ThermoFisher) as described. Samples were analyzed on a FACSymphony A5 flow cytometer (BD Biosciences) and FlowJo v9.9.4. Mitophagy was assessed using Mitotracker Red CMXRos (MTR; Molecular Probes) and thiazole orange (TO, BD Biosciences) as described. DNA damage was assessed by micronucleus assay at baseline and following irradiation as previously described. Micronuclei (MN) arise from chromosomes or chromosome fragments that are not incorporated into the nucleus during cell division and remain in peripheral blood cells. As reticulocytes arise following the last mitosis, radiation induced chromosome damage is detected as an increase in reticulocyte MN. Intrinsic or spontaneous damage is seen as an increase in MN in mature red cells, which do not contain a nucleus at the time of irradiation.

Data Acquisition and Imaging. Images of peripheral blood were acquired using a Nikon Eclipse E600 microscope equipped with a Nikon DS-Fi3 color camera and NIS-Elements Documentation imaging software (Nikon). Smears were examined with a 100/0.5–1.3 oil-immersion objective.

Statistics. Significant differences were identified using Tukey honestly significant differences (HSD) or 2-tailed Student t test using JMP Statistical Software v11.2.1.
Data Deposition. RNAseq data were submitted to Sequence Read Archive (Bioproject PRJNA432205). Expression profiles are available at http://nanexpression.mdibl.org/.

References
1. Siatecka, M. et al. Severe anemia in the Nan mutant mouse caused by sequence-selective disruption of erythroid Kruppel-like factor. Proc Natl Acad Sci USA 107, 15151–15156, https://doi.org/10.1073/pnas.1004996107 (2010).
2. Heruth, D. P. et al. Mutation in erythroid specific transcription factor KLF1 causes Hereditary Spherocytosis in the Nan hemolytic anemia mouse model. Genomics 96, 303–307, https://doi.org/10.1016/j.ygeno.2010.07.009 (2010).
3. Miller, I. J. & Bicker, J. I. A novel, erythroid cell-specific murine transcription factor that binds to the CACCC element and is related to the Kruppel family of nuclear proteins. Mol Cell Biol 13, 2776–2786 (1993).
4. Drissen, R. et al. The erythroid phenotype of KLF1-null mice: defects in hemoglobin metabolism and membrane stability. Mol Cell Biol 25, 5205–5214, https://doi.org/10.1128/MCB.25.12.5205-5214.2005 (2005).
5. Hodge, D. et al. A global role for EKLF in definitive and primitive erythropoiesis. Blood 107, 3359–3370, https://doi.org/10.1182/blood-2005-07-2888 (2006).
6. Frontelo, P. et al. Novel role for EKLF in megakaryocyte lineage commitment. Blood 110, 3871–3880, https://doi.org/10.1182/blood-2007-03-082065 (2007).
7. Pilon, A. M. et al. Failure of terminal erythroid differentiation in EKLF-deficient mice is associated with cell cycle perturbation and reduced expression of E2F2. Mol Cell Biol 28, 7394–7401, https://doi.org/10.1128/MCB.01087-08 (2008).
8. Siatecka, M. & Bicker, J. J. The multifunctional role of EKLF/KLF1 during erythropoiesis. Blood 118, 2044–2054, https://doi.org/10.1182/blood-2011-03-331371 (2011).
9. Perkins, A. C., Sharpe, A. H. & Orkin, S. H. Lethal beta-thalassemia in mice lacking the erythroid CACCC-transcription factor EKLF. Nature 375, 318–322, https://doi.org/10.1038/375318a0 (1995).
10. Perkins, A. C., Gaensler, K. M. & Orkin, S. H. Silencing of human fetal globin expression is impaired in the absence of the adult beta-globin gene activator protein EKLF. Proc Natl Acad Sci USA 93, 12267–12271 (1996).
11. Feng, W. C., Southwood, C. M. & Bieker, J. J. Analyses of beta-thalassemia mutant DNA interactions with erythroid Kruppel-like factor (EKLF), an erythroid cell-specific transcription factor. J Biol Chem 269, 1493–1500 (1994).
12. Gregory, R. C. et al. Functional interaction of GATA1 with erythroid Kruppel-like factor and Sp1 at defined erythroid promoters. Blood 87, 1793–1801 (1996).
13. Gillinder, K. R. et al. Promiscuous DNA-binding of a mutant zinc finger protein corrupts the transcriptome and diminishes cell viability. Nucleic Acids Res 45, 1130–1143, https://doi.org/10.1093/nar/gkw1014 (2017).
14. Tallack, M. R. et al. Novel roles for KLF1 in erythropoiesis revealed by mRNA-seq. Genome Res 22, 2385–2398, https://doi.org/10.1101/gr.135707.112 (2012).
15. Planutis, A. et al. Neomorphic effects of the neonatal anemia (Nan-Eklf) mutation contribute to deficits through development. Development 144, 430–440, https://doi.org/10.1242/dev.145656 (2017).
16. Ganz, T. Hepcidin and iron regulation, 10 years later. Blood 117, 4425–4433, https://doi.org/10.1182/blood-2011-01-258467 (2011).
17. Honda, K. et al. IRE-7 is the master regulator of type-I interferon-dependent immune responses. Nature 434, 772–777, https://doi.org/10.1038/nature03464 (2005).
18. Ganz, T. & Nemeth, E. Hepcidin and iron homeostasis. Biochim Biophys Acta 1823, 1434–1443, https://doi.org/10.1016/j.bbamcr.2012.01.014 (2012).
19. Kautz, L. et al. Identification of erythроferrone as an erythroid regulator of iron metabolism. Nat Genet 46, 678–684, https://doi.org/10.1038/ng.2996 (2014).
20. Hom, J. et al. Characterization of zinc finger proteins that interact with angiopoietin-1. J Biol Chem 278, 2798–2806, https://doi.org/10.1074/jbc.M309531200 (2003).
21. Paulson, R. F. et al. Stress erythropoiesis: new signals and new stress progenitor cells. Curr Opin Hematol 18, 139–145, https://doi.org/10.1097/MOH.0b013e32835421c8 (2011).
22. Xiang, J., Wu, D. C., Chen, Y. & Paulson, R. F. In vitro culture of stress erythroid progenitors identifies distinct progenitor populations and analogous human progenitors. Blood 125, 1803–1812, https://doi.org/10.1182/blood-2014-07-591453 (2015).
23. Masuda, T. et al. Transcription factors LFPR and BCL11A independently repress expression of fetal hemoglobin. Science 351, 285–289, https://doi.org/10.1126/science.aad3312 (2016).
24. Nandakumar, S. K., Ulirsch, J. C. & Sankaran, V. G. Advances in understanding erythropoiesis: evolving perspectives. Br J Haematol 173, 206–218, https://doi.org/10.1111/bjh.13938 (2016).
25. Liu, J. et al. Quantitative analysis of murine terminal erythroid differentiation in vivo: novel method to study normal and disordered erythropoiesis. Blood 121, e43–49, https://doi.org/10.1182/blood-2012-09-456079 (2013).
26. Balmain, A. et al. A high-throughput in vivo microincule assay for genome instability screening in mice. Nat Protoc 10, 205–215, https://doi.org/10.1038/nprot.2015.010 (2015).
27. Shima, N. et al. Phenotype-based identification of mouse chromosome instability mutants. Genetics 163, 1031–1040 (2003).
28. Gnanapragasam, M. N. et al. EKLF/KLF1-regulated cell cycle exit is essential for erythroblast enucleation. Blood 128, 1631–1641 (2016).
29. Liu, Y. et al. Suppression of Fas-FasL coexpression by erythropoietin mediates erythroblast expansion during the erythropoietic stress response in vivo. Blood 108, 123–133, https://doi.org/10.1182/blood-2005-11-4458 (2006).
30. Koulins, M. et al. Contrasting dynamic responses in vivo of the BCL-xL and Bim erythropoietic survival pathways. Blood 119, 1228–1239, https://doi.org/10.1182/blood-2011-07-365346 (2012).
31. Gillinder, K. R. et al. Direct targets of pSTAT5 signalling in erythropoiesis. PLoS One 12, e0180922, https://doi.org/10.1371/journal. pone.0180922 (2017).
32. Tallack, M. R. et al. A global role for KLF1 in erythropoiesis revealed by ChIP-seq in primary erythroid cells. Genome Res 20, 1052–1063, https://doi.org/10.1101/gr.110655.7.110 (2010).
33. Thomas, L. W., Lam, C. & Edwards, S. W. Mc1-l; the molecular regulation of protein function. FERS Letters 584, 2981–2989 (2010).
34. Wensveen, F. M. et al. BH3-only protein Noxa contributes to apoptotic control of stress-erythropoiesis. Apoptosis 18, 1306–1318, https://doi.org/10.1007/s10495-013-0980-y (2013).
35. Wu, H., Medeiros, L. J. & Young, K. H. Apoptosis signaling and BCL-2 pathways provide opportunities for novel targeted therapeutic strategies in hematologic malignancies. Blood Rev 32, 8–28, https://doi.org/10.1016/j.blre.2017.08.004 (2018).
36. Rûfkind, J. M. & Nagababu, E. Hemoglobin redox reactions and red blood cell aging. Antioxid Redox Signal 18, 2274–2283, https://doi.org/10.1089/ars.2012.4867 (2013).
41. Nagababu, E., Mohanty, J. G., Friedman, J. S. & Rifkind, J. M. Role of peroxiredoxin-2 in protecting RBCs from hydrogen peroxide-induced oxidative stress. Free Radic Res, 164–171 (2013).
42. Neumann, C. A. et al. Essential role for the peroxiredoxin Prdx1 in erythrocyte antioxidant defence and tumour suppression. Nature 424, 561–565, https://doi.org/10.1038/nature01819 (2003).
43. Friedman, J. S. et al. SOD2-deficiency anemia: protein oxidation and altered protein expression reveal targets of damage, stress response, and antioxidant responsiveness. Blood, 104, 2565–2573 (2004).
44. Friedman, J. S. et al. Absence of mitochondrial superoxide dismutase results in a murine hemolytic anemia responsive to therapy with a catalytic antioxidant. J Exp Med, 193, 925–934 (2001).
45. Huh, J. Y. et al. Peroxiredoxin 3 is a key molecule regulating adipocyte oxidative stress, mitochondrial biogenesis, and adipokine expression. Antioxid Redox Signal 16, 229–243, https://doi.org/10.1089/ars.2011.395210.1089/ars.2010.3766 (2012).
46. Iuchi, Y. et al. Peroxiredoxin 4 knockout results in elevated spermatogenic cell death via oxidative stress. Biochem J 419, 149–158, https://doi.org/10.1042/BJ20081526 (2009).
47. Da Costa, L. et al. Molecular approaches to diagnose Diamond-Blackfan anemia: The EuroDBA experience. Eur J Med Genet, https://doi.org/10.1016/j.ejmg.2017.10.017 (2017).
48. Adam, S. et al. Concise Review: Getting to the Core of Inherited Bone Marrow Failures. Stem Cells, 35, 284–298, https://doi.org/10.1002/stem.2543 (2017).
49. Ruggiero, D. & Shimamura, A. Marrow failure: a window into ribosome biology. Blood, 124, 2784–2792, https://doi.org/10.1182/blood-2014-04-526301 (2014).
50. Vlachos, A., Blanc, L. & Lipton, J. M. Diamond Blackfan anemia: a model for the translational approach to understanding human disease. Expert Rev Hematol 7, 359–372, https://doi.org/10.1586/17474086.2014.897923 (2014).
51. Dobin, A. et al. STAR: ultrafast universal RNA-seq aligner. Bioinformatics, 15, 21, https://doi.org/10.1093/bioinformatics/bts635 (2013).
52. Wang, L. et al. Measure transcript integrity using RNA-seq data. BMC Bioinformatics, 17, 58, https://doi.org/10.1186/s12859-016-0922-z (2016).
53. Wang, L., Wang, S. & Li, W. RSeQC: quality control of RNA-seq experiments. Bioinformatics, 28, 2184–2185, https://doi.org/10.1093/bioinformatics/bts356 (2012).
54. Mudge, J. M. & Harrow, J. Creating reference gene annotation for the mouse C57BL6/J genome assembly. Mamm Genome, 26, 366–378, https://doi.org/10.1007/s00335-015-9583-x (2015).
55. Love, M. I., Huber, W. & Anders, S. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol, 15, 550, https://doi.org/10.1186/s13059-014-0550-8 (2014).
56. Vermes, I., Haanen, C., Steffens-Nakken, H. & Reutelingsperger, C. A novel assay for apoptosis. Flow cytometric detection of phosphatidylserine expression on early apoptotic cells using fluorescein labelled Annexin V. J Immunol Methods, 193, 35–43 (1995).
57. Zhang, J. et al. Mitochondrial clearance is regulated by Atg7-dependent and -independent mechanisms during reticulocyte maturation. Blood, 114, 157–164, https://doi.org/10.1182/blood-2008-04-151639 (2009).

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Author Contributions
D.N., S.L.C., R.F.R., J.P. and E.H performed experiments. A.C.P., J.J.B., K.R.C. and L.B. provided data and analysis; J.H.G. analyzed RNAseq data; D.N. and L.L.P. performed all other statistical analyses; L.L.P. and D.N. directed experiments and wrote the paper with input from all authors.

Additional Information
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