**Reticular dysgenesis–associated AK2 protects hematopoietic stem and progenitor cell development from oxidative stress**

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Reticular dysgenesis–associated AK2 protects hematopoietic stem and progenitor cell development from oxidative stress

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Adenylate kinases (AKs) are phosphotransferases that regulate the cellular adenine nucleotide composition and play a critical role in the energy homeostasis of all tissues. The AK2 isoenzyme is expressed in the mitochondrial intermembrane space and is mutated in reticular dysgenesis (RD), a rare form of severe combined immunodeficiency (SCID) in humans. RD is characterized by a maturation arrest in the myeloid and lymphoid lineages, leading to early onset, recurrent, and overwhelming infections. To gain insight into the pathophysiology of RD, we studied the effects of AK2 deficiency using the zebrafish model and induced pluripotent stem cells (iPSCs) derived from fibroblasts of an RD patient. In zebrafish, Ak2 deficiency affected hematopoietic stem and progenitor cell (HSPC) development with increased oxidative stress and apoptosis. AK2-deficient iPSCs recapitulated the characteristic myeloid maturation arrest at the promyelocyte stage and demonstrated an increased AMP/ADP ratio, indicative of an energy-depleted adenine nucleotide profile. Antioxidant treatment rescued the hematopoietic phenotypes in vivo in ak2 mutant zebrafish and restored differentiation of AK2-deficient iPSCs into mature granulocytes. Our results link hematopoietic cell fate in AK2 deficiency to cellular energy depletion and increased oxidative stress. This points to the potential use of antioxidants as a supportive therapeutic modality for patients with RD.

SCID includes a heterogeneous group of immunological diseases characterized by severe defects of T-lymphocyte number and function, variably associated with numerical and functional abnormalities of B and NK cells (Fischer et al., 2005; Aloj et al., 2012). Reticular dysgenesis (RD) is a

* A. Rissone and K.G. Weinacht contributed equally to this paper.
** L.D. Notarangelo and F. Candotti contributed equally to this paper.

Abbreviations used: AK, adenylate kinase; AP, alkaline phosphatase; ATRA, all-trans-retinoic acid; CHT, caudal hematopoietic tissue; DIG, digoxigenin; dpf, day postfertilization; dsRed, Discosoma sp. RFP; EB, embryoid body; EGFP, enhanced GFP; GSH, glutathione; hpf, bryoid body; EGFP, enhanced green fluorescent protein; HSPC, hematopoietic stem and progenitor cell; HSCT, hematopoietic stem cell transplantation; iPSC, induced pluripotent stem cell; ISH, in situ hybridization; M3, morpholino oligonucleotide; NAC, N-acetyl-L-cysteine; PFA, paraformaldehyde; RD, reticular dysgenesis; ROS, reactive oxygen species; RQ, relative quantification; WISH, whole mount ISH; ZFN, zinc-finger nuclease.
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AKs are phosphotransferases that play an important role in cellular and mitochondrial energy homeostasis; moreover, they contribute to modulating the AMP-mediated response to stress signals (Dzeja and Terzic, 2003, 2009; Dzeja et al., 2007). Different AKs set the adenine nucleotide ratio in different intracellular compartments. In particular, AK2 is expressed in the mitochondrial intermembrane space where it regulates

particularly severe, autosomal-recessive form of SCID and is characterized by virtual absence of circulating lymphocytes and neutrophils, as well as sensory neural hearing loss (Lagresle-Peyrou et al., 2009; Pannicke et al., 2009). The lack of innate and acquired immune function in patients with RD leads to extreme susceptibility to infections and early death, unless hematopoietic and immune reconstitution is achieved by allogeneic hematopoietic stem cell transplantation (HSCT). In 2009, two groups have independently reported that RD is caused by mutations in adenylate kinase 2 (AK2; Lagresle-Peyrou et al., 2009; Pannicke et al., 2009).

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Figure 1. Zebrafish AK2 alternative splicing isoforms are expressed in hematopoietic regions during embryo development. (A) Schematic representation of human (hAK2) and zebrafish (zak2) conserved gene structure and alternative splicing mechanism. STOP indicates the position of the STOP codons. Neither exons nor introns are drawn to scale. (B) RT-PCR analysis on cDNA of 4 dpf WT embryos. The reverse primer AK2-ISO B R2 (Table S1) was used in the PCR reaction to bind to isoforms A and B. MW, molecular weight size marker; C−, negative control. Blue arrowheads point to the two different alternative splicing isoforms. The results are representative of two independent experiments. (C) Multiprotein sequence alignment of human (hAK2) and zebrafish (zAk2) AK2 splicing isoforms (indicated as ISOA and ISOB). Features of human AK2 structure are depicted above the amino acid sequences. Different colors indicate different physicochemical properties. (D) mRNA quantitative real-time PCR analysis showing the relative expression of AK2 splicing isoforms during embryo development using specific probes for isoforms A or B (ak2-A and ak2-B) or a common probe for A and B isoforms (ak2-AB). Expression levels were normalized to β-actin 2 (β-act 2) and elongation factor 1α (ef1α), and mRNA from 2–5-somite stage embryos was used as reference. Error bars indicate the calculated maximum (RQ_max) and minimum (RQ_min) expression levels that represent the standard error of the mean expression level (RQ value). Data are pooled from at least three independent experiments. (E) Spatiotemporal analysis by stereomicroscopy of AK2 expression by WISH using a common probe for A and B isoforms at the indicated times. For each stage, at least 25 embryos have been analyzed. Data show one representative experiment out of two independently performed. ICM, intermediate cell mass; CHT, caudal hematopoietic tissue. Bars, 100 µm.
concentration of mono- and diphosphorylated adenine nucleotides (Dzeja and Terzic, 2003, 2009; Dzeja et al., 2007). Through their distinct subcellular localization, AKs create a shuttle system to serve the needs of different cellular compartments for energetic and metabolic signaling; therefore, they represent a sensitive reporter of the cellular energy state (Dzeja and Terzic, 2003, 2009; Dzeja et al., 2007). Knockout studies have shown that the lack of a single AK enzyme is usually compensated at different levels through remodeling of the whole energetic network (Janssen et al., 2000; Pucar et al., 2000, 2002). For example, in Ak1−/− mice, mitochondrial AK2 and AK3 can compensate at least in part for the lack of AK1 (Pucar et al., 2000). However, within the myeloid and lymphoid lineages, as well as in the stria vasularis of the inner ear, AK1 expression is absent or too low to compensate for the lack of AK2 activity in patients with RD (Le Guyader et al., 2008; Pannicke et al., 2009).

Since its first description (De Vaal and Seynhaeve, 1959), RD was assumed to be a disorder of primitive multipotent “reticular” progenitor cells primarily failing to develop into granulocytic and lymphoid precursors. However, the pathophysiology underlying the concurrent myeloid and lymphoid defects in RD has long remained unresolved.

To explore this topic, we studied the effects of AK2 deficiency in zebrafish embryos and human induced pluripotent stem cells (iPSCs) derived from an RD patient. Our results showed severe impairment of hematopoietic stem and progenitor cell (HSPC) development associated with increased oxidative stress and apoptosis in AK2-deficient zebrafish. Directed myeloid differentiation of AK2-deficient iPSCs resulted in the characteristic promyelocyte maturation arrest observed in RD patients and was associated with an energy-depleted adenine nucleotide profile. Notably, we found that treatment with antioxidants rescued the hematopoietic abnormalities in AK2 mutant zebrafish and restored granulocytic differentiation of AK2-deficient iPSCs. Our results support the hypothesis that AK2 deficiency results in energy depletion and abnormal redox state and suggest that antioxidants have potential as a supportive therapeutic modality in RD.

RESULTS
AK2 deficiency induces a wide array of hematopoietic defects in zebrafish
In humans, alternative splicing of the AK2 gene can produce two different protein isoforms (A and B); isoform B is encoded by seven exons instead of six and has a different termination codon (Lee et al., 1998; Pannicke et al., 2009). One major characteristic of the zebrafish AK2 gene (ak2) is the lack of intron 6, resulting in the juxtaposition of the exons 6 and 7 coding sequences (Fig. 1 A). Bioinformatic and RT-PCR analyses revealed a previously unreported isoform in zebrafish, corresponding to the human AK2 isoform B. Although the zebrafish genomic sequence corresponding to the human AK2 exon 7 represents part of the 3′-UTR region of isoform A, RT-PCR and sequence alignment analyses indicated that alternative splicing mechanisms are conserved (Fig. 1, A–C).

To assess the characteristics of physiological ak2 expression during zebrafish development, we performed quantitative real-time PCR and whole mount in situ hybridization (ISH [WISH]) experiments. The quantitative real-time PCR analysis showed that zebrafish embryos inherit both maternal A and B transcripts; moreover, relative quantification (RQ) using the 2−ΔΔCT method as reference indicated a significant decline in transcription levels after the beginning of zygotic transcription (Fig. 1 D). WISH experiments at different stages of development using a common ISH probe for isoforms A and B (Fig. 1 E) localized ak2 expression to hematopoietic regions such as the intermediate cell mass and the caudal hematopoietic tissue (CHT) around 24 h postfertilization (hpf) and 5 d postfertilization (dpf), respectively.

No mouse model for AK2 deficiency has been reported in the literature to date, and mouse lines carrying homozygous ak2-inactivating retroviral insertions are embryonically lethal (unpublished data). To study the role of AK2 in hematopoietic development, we set out to generate a zebrafish model of AK2 deficiency. Because RD patients present with a variety of different mutations (including large intragenic deletions, single nucleotide substitution, and frameshift mutations; Lagresle-Peyrou et al., 2009, 2011; Pannicke et al., 2009; Al-Zahrani et al., 2013; Henderson et al., 2013), we decided to generate various ak2 zebrafish mutants. Using zinc-finger nuclease (ZFN) technology, we introduced frameshift mutations in the first exon of the ak2 gene (a 2-bp deletion and a 4-bp insertion, indicated as ak2del2 and ak2ins4, respectively; Fig. 2 A; Sood et al., 2013). In addition, we screened a zebrafish DNA library of ENU-induced mutations (Sood et al., 2006) and uncovered a mutant fish line carrying a T371C/L124P missense mutation within exon 4 of the AK2 gene. The mutated amino acid in this mutant (Fig. 2 B, red arrows) falls in a highly conserved region of AK2 (Liu et al., 2009) located upstream of the β sheet connecting the CORE domain to the LID domain. The L124P mutation has a PROVEAN score (Choi et al., 2012) equal to −6.9 and was predicted to affect protein stability and enzymatic activity. Genotyping of in-crossed heterozygous embryos confirmed the presence of these recessive mutations in a normal Mendelian ratio (Fig. 2 B, right). Contrary to the ZFN-induced frameshift mutations that resulted in abrogation of ak2 mRNA, the L124P substitution did not affect production of the mRNA transcript (Fig. 2 C).

A previous study has shown that morpholino oligonucleotide (MO)–mediated knockdown of ak2 expression induced defects in zebrafish leukocyte development (Pannicke et al., 2009). We confirmed and refined these findings in our zebrafish ak2 mutant lines that showed profound impairment of lymphoid and myeloid development, as indicated by lack of ikaros and rag1 expression in the thymic anlage and defective mpx expression at 5 dpf (Fig. 2 D). Because RD is characterized by agranulocytosis, we used Sudan black, a lipid stain which labels the granules of granulocytes (Sheehan and Storey, 1947; Le Guyader et al., 2008), to visualize mature granulocytes in zebrafish larvae. When comparing Sudan black–stained AK2 mutants to control embryos, we found a progressive
Figure 2. Zebrafish AK2 mutants present a wide array of hematopoietic defects. (A) Alignment of protein sequences encoded by the WT and ZFN-induced mutant alleles (del2 and ins4) shows the predicted frameshift after the Arg 15 codon targeted by the ZFN cutting site. Premature stop codons are also indicated (*). (B) 3D structure of human AK2 (left). (bottom) Partial sequence multi-alignment of WT (zAK2) and missense mutant zebrafish Ak2 (zAK2L124P/L124P) and human AK2 (hAK2). The mutated amino acid position is highlighted in both panels (red arrows). (right) Sequence chromatograms showing the nucleotide change for L124P missense mutation in heterozygous (ak2L124P/+) and homozygous (ak2L124P/L124P) embryos compared with the WT sequence. (C) Stereomicroscope images of WISH analysis on 24 hpf WT and mutant embryos using an antisense probe against ak2 mRNA (A and B transcripts). For each sample, at least 25 embryos have been analyzed. (D) Analysis of definitive hematopoietic phenotypes in ak2L124P/del2 and ak2L124P/del2 mutants by WISH and stereomicroscopy. Red and black dashed circles indicate thymus and kidney regions, respectively. (left) Lateral views of embryos hybridized with rag1, mpx, and hbae1 and ventral views of ikaros hybridization at 5 dpf. (right) Ventral views of rag1 and lateral views of mpx and hbae1 hybridized embryos at 5 dpf and lateral views of embryos stained using Sudan black solution at different stages of development. For each sample, at least 25 embryos have been analyzed. Data in C and D show one representative experiment out of two independently performed. Bars, 100 µm.
Figure 3. ak2 knockdown by MO injection phenocopies hematopoietic defects observed in ak2 zebrafish mutants. (A) Schematic representation of the two different MOs used in the study: ak2 MO1 blocking the mRNA translation and ak2 MO2 targeting the intron 3–exon 4 splice junction of the ak2 gene. Neither exons nor introns are drawn to scale. Numbers below the introns (straight lines) and exons (white boxes) indicate their size in bases. (B) Schematic representation of the predicted AK2 transcripts in MO-treated embryos. Black arrowheads in the bottom left panel indicate noncanonical STOP codons introduced by ak2 MO2–induced skipping of exon 4. Horizontal black arrows indicate the pair of primers used in RT-PCR (right panels) to test the effects of different doses of ak2 MO2 during development as indicated. MW, molecular weight size marker. (C) mRNA quantitative real-time PCR analysis of splicing MO activity (three different doses) at 28 hpf and 3 dpf. In both cases, ST-CTRL MO–injected embryos serve as reference. The error bars indicate the calculated maximum (RQ_{max}) and minimum (RQ_{min}) expression levels that represent the standard error of the mean expression levels.
reduction in Sudan black–positive cells throughout the larva bodies (Fig. 2 D). Interestingly, we also observed markedly impaired expression of the erythroid marker hbae1 (Fig. 2 D). These results were confirmed by injecting translation-blocking (ak2 MO1) or aberrant splicing-inducing (ak2 MO2) MOs (Pannicke et al., 2009), either individually or in combination (Fig. 3). The expected effects of the splice-blocking MO injection (ak2-MO2, targeting intron 3–exon 4 of AK2 pre-mRNAs) included the skipping of exon 4 and a frameshift resulting in the introduction of multiple STOP codons (Fig. 3, A and B). RT–PCR and quantitative real-time PCR analyses confirmed the skipping of exon 4 and indicated that the splice-blocking MO was still active at 3–dpf stage, although with reduced activity (Fig. 3, B and C). The findings of impaired lymphoid and myeloid development in both AK2 mutant and ak2 MO-injected (morphant) zebrafish embryos (Fig. 3, D–F) strongly indicate that the observed hematopoietic phenotype represents a direct consequence of AK2 deficiency.

AK2 deficiency affects zebrafish HSPC development

Because AK2 deficiency in zebrafish resulted in developmental defects of all hematopoietic lineages, we hypothesized that HSPC development might also be affected. Analysis of c-myb and runx1 expression in hematopoietic tissues revealed only minor differences between WT and AK2 mutants or morphants during early phases of HSPC development (i.e., 30–36 hpf; Fig. 4, A, B, and D). Moreover, WISH analysis with a pan-endothelial vascular marker (dh5) and confocal analysis on embryos from incross of ak2del1/del1 mutants in the Tg(fl1a:EGFP) f1; Tg(gata1a:dsRed) d42 background indicated a correct vascular development (marked by enhanced GFP [EGFP] expression) and the presence of active circulation (marked by Discosoma sp. RFP [dsRed] expression) in mutant embryos from 24 hpf to 2 dpf (Fig. 4 E). Interestingly, however, both AK2 mutants and morphants showed a strong reduction of c-myb expression at 2 dpf through 5 dpf (Fig. 3, G and H; and Fig. 4, A–C).

To obtain additional evidence that HSPC development was in fact impaired, we injected Tg(db41:GFP) embryos with the ak2 MOs, either separately or in combination at low dose (Fig. 4, F and G). In this specific transgenic line, HSPCs and thrombocytes are marked by GFP expression (Lin et al., 2005; Bertrand et al., 2008; Kissa et al., 2008; Renaud et al., 2011). At 4.5 dpf, control embryos displayed GFP+ cells (HSPCs and thrombocytes) in the CHT region and in the circulation. However, consistent with the aforementioned described findings, Tg(db41:GFP)/AK2 morphant embryos showed strong reduction of GFP+ cells in the CHT region (Fig. 4, F and G). Collectively, these data support an important role for AK2 in HSPC development during zebrafish definitive hematopoiesis.

AK2 deficiency increases oxidative stress and apoptosis in zebrafish hematopoietic tissues

Under physiological conditions, the amount of oxidative stress produced in the mitochondria is counteracted by an intricate defense system that scavenges and detoxifies reactive oxygen species (ROS). Mitochondrial dysfunction can lead to excess ROS production that exhausts the cellular antioxidant capacity. Unopposed oxidative stress leads to apoptosis (Craven et al., 2005). Attempts to compensate for increased levels of ROS include activation of the expression of different genes such as the inducible isoform of heme oxygenase (HMOX1; Ryter and Choi, 2002). Therefore, HMOX1 expression serves as a marker of increased levels of cellular oxidative stress. We reasoned that the reduced ADP recycling caused by AK2 deficiency would result in decreased ATP production, mitochondrial dysfunction, and increased ROS production. Indeed, CHT cells from ak2 MO1 and ak2 MO2 (2.4 ng) embryos showed evidence of oxidative stress as detected by the MitoSOX and CellROX indicators (Fig. 5 A and not depicted). In addition, FACS analysis of WT and ak2 MO–derived cells stained with MitoSOX showed significantly increased levels of superoxide in mitochondria of null mutant cells at 4 and 5 dpf, with similar levels of apoptotic and dead cells as indicated by Annexin V–APC and 7–AAD staining, respectively (Fig. 5, B and C). CellROX staining at 4 dpf confirmed the presence of higher levels of oxidative stress in null mutant embryo cells (Fig. 5 D), and hmx1a expression in the CHT of both ak2 MO1 and ak2 MO2 mutant embryos showed dramatic induction starting at 2 dpf, in contrast to WT control embryos that showed very few or no hmx1a-positive cells in the same region (Fig. 5, E and F). Analogous to what is observed in human fibroblasts derived from individuals with RD (Pannicke et al., 2009), these results demonstrate that, in zebrafish, AK2 deficiency induces oxidative stress in hematopoietic tissues.

Previous reports have linked decreased AK2 activity to induction of apoptosis (Pannicke et al., 2009; Burkart et al., 2005).
These data suggest that the progressive reduction of HSPCs and committed hematopoietic precursors in AK2-deficient embryos can be attributed to increased levels of ROS, resulting in cell death by apoptosis, which is in line with observations made in AK2-deficient in vitro model systems (Lagresle-Peyrou et al., 2009; Pannicke et al., 2009).

To test this hypothesis, we assayed for cell death using acridine orange staining and TUNEL assay. We detected apoptotic cells in the CHT of ak2 morphants at 3 dpf (Fig. 5 G) and confirmed the increased number of apoptotic cells by TUNEL assay in CHT regions of ak2<sup>del2/del2</sup> and ak2<sup>L124P/L124P</sup> mutant embryos (Fig. 5, H and I). These data suggest that the progressive reduction of HSPCs and committed hematopoietic precursors in AK2-deficient embryos can be attributed to increased levels of ROS, resulting in cell death by apoptosis, which is in line with observations made in AK2-deficient in vitro model systems (Lagresle-Peyrou et al., 2009; Pannicke et al., 2009).
Figure 5. AK2 mutant zebrafish demonstrate increased levels of cellular oxidative stress and apoptosis in hematopoietic tissues. (A) Confocal microscopy assessment of oxidative stress in the CHT region of WT and ak2<sup>ins4/ins4</sup> mutant embryos probed with the MitoSOX or CellROX indicators at the indicated stages. Yellow arrowheads in the dashed insets indicate MitoSOX- or CellROX-positive cells in CHT regions. (B) Representative quantitative analysis by flow cytometry of oxidative stress, apoptosis, and cell death on 5 dpf ak2<sup>del2/del2</sup> and control embryos (60 embryos each) using MitoSOX Red and Annexin V + 7-AAD, respectively. (C and D) Photometric analysis of MitoSOX Red (C) and CellROX Green (D) staining at 4 and 5 dpf in ak2<sup>del2/del2</sup> mutants and WT siblings (60 embryos each). (E) WISH analysis of hmoxA expression in ak2<sup>del2/del2</sup> mutants and their WT siblings at different stages of development. (F) WISH analysis of hmoxA expression in ak2<sup>L124P/L124P</sup> mutants and their WT siblings at different stages of development. (E and F) Open arrowheads indicate hmoxA-positive cells. (G) Stereomicroscopy analysis of acridine orange staining of AK2 morphants at 3 dpf. Dashed insets show CHT regions. (H) Confocal analysis of fluorescent TUNEL staining in CHT regions of WT siblings and ak2<sup>L124P/L124P</sup> and ak2<sup>L124F/L124F</sup> mutant embryos from 2 to 5 dpf. Each panel represents a crop of the CHT region from a 14x magnification image. The green signal indicates TUNEL-positive cells and the blue signal...
Antioxidant treatment rescues hematopoietic phenotypes in AK2-deficient embryos

To further corroborate our findings, we investigated the effects of different antioxidant compounds on oxidative stress and hematopoietic abnormalities noted in ak2 mutant zebrafish. We exposed an unselected population of embryos derived from intercross of ak2<sup>2/124⁸</sup> heterozygous missense mutants to different concentrations of N-acetyl-L-cysteine (NAC). This treatment resulted in decreased cellular oxidative stress, as indicated by a 50% reduction of hmox1a induction in NAC-treated ak2<sup>L124P/L124P</sup> missense mutant zebrafish versus untreated mutants (Fig. 6 A). Moreover, antioxidant treatment rescued the defective expression of c-myb, hba1, and rag1 in ~50% of the embryos expected to show an abnormal phenotype (Fig. 6 A, left).

Importantly, similar results were observed for c-myb and rag1 expression after treatment of ak2<sup>6/6</sup>–null mutant embryos with different concentrations of NAC, glutathione (GSH), and GSH reduced ethyl ester (GSH-MEE), a membrane/lipid-permeable derivative of GSH (Fig. 6, B–E). Notably, we did not observe macroscopic defects, nor enhanced expression of hematopoietic markers in WT embryos treated with antioxidants. Overall, these data indicate that antioxidant treatment ameliorates the hematopoietic phenotypes observed in zebrafish ak2 mutants and suggest a crucial role of oxidative stress in the pathogenesis of RD.

Patient-derived iPSCs as an in vitro model for RD

To assess whether our results in zebrafish may be translated into an in vitro model for RD, we developed a disease model based on in vitro hematopoietic differentiation of iPSCs. Dermal skin fibroblasts were derived from a patient who presented with agranulocytosis, severe T cell lymphopenia, and absent brain stem auditory-evoked response at birth, the characteristic triad that clinically defines RD. The patient was found to carry a homozygous point mutation (c.524G>A) in exon 6 of the AK2 gene, leading to an arginine to glutamine substitution (p.R175Q) in the LID domain of the AK2 protein (Fig. 7, A and B; Henderson et al., 2013). Western blot analysis from patient fibroblasts and CD3<sup>+</sup> T cells demonstrated reduced protein expression compared with control (Henderson et al., 2013), and a PROVEAN score of −3.871 predicted this mutation to be deleterious. The cultured fibroblasts were reprogrammed to pluripotency with an excisable, self-inactivating, codon-optimized lentiviral vector carrying OCT4, SOX2, KLF4, cMYC, and the reporter dTomato in a single cassette (Warlich et al., 2011). Immunofluorescence imaging and quantitative real-time PCR analysis confirmed the stemness of the generated iPSC line by expression of the characteristic pluripotency markers TRA-1–81, TRA-1–60, NANOG, OCT-4, SSEA-3, and SSEA-4 (Fig. 7, C and D). Karyotype and G-banding analyses demonstrated cytogenetic integrity, and gDNA sequencing showed genotypic identity with the parental fibroblast line (Fig. 7, E and F). Of note, the reprogramming efficiency of 0.0096% in AK2-deficient fibroblasts (mean over eight experiments) was ~500-fold lower than the reprogramming efficiency of 4.75% seen in the WT control (mean over three experiments).

AK2 deficiency affects in vitro granulopoiesis and erythropoiesis

To model the hematopoietic phenotype of RD in vitro, patient-derived iPSCs were grown in suspension culture to form embryoid bodies (EBs), dissociated into single cells, and plated in a CFU assay in the presence of SCF, FLT3, BMP4, G-CSF, IL3, and IL6. AK2<sup>R175Q/R175Q</sup> iPSCs recapitulated the characteristic maturation arrest of the myeloid lineage at the promyelocyte stage that is observed in the bone marrow of RD patients (Lagresle-Peyrou et al., 2009; Pannicke et al., 2009), whereas the control demonstrated differentiation into mature neutrophils with distinctive nuclear segmentation (Fig. 8 A). For unequivocal delineation of the differentiation stage, we performed electron microscopy on AK2<sup>R175Q/R175Q</sup> iPSC-derived myeloid precursors and control cells. Cells morphologically identified as promyelocytes by light microscopy displayed abundant electron-dense primary granules, validating their promyelocyte nature (Fig. 8 B). In addition to the maturation arrest, the quantitative potential of AK2-deficient cells to form mixed myeloid lineage colonies was significantly decreased (Fig. 8 C). Interestingly, we also noted abnormalities in the erythroid differentiation of RD iPSCs. Red cell precursors derived from AK2<sup>R175Q/R175Q</sup> iPSCs displayed obvious dysmorphic features such as bilobate nuclei and inconsistent nucleus-to-cytoplasm ratio (Fig. 8 D), and their potential to give rise to CFU-erythroid (CFU-E) and burst-forming units–erythroid (BFU-E) was severely compromised (Fig. 8 E). The impact of AK2 deficiency on more than one hematopoietic cell lineage in the iPSC model supports our findings in the zebrafish, suggesting that RD affects HSPC development and differentiation.

AK2 deficiency affects the cellular adenine nucleotide profile

AK2 is considered the primary mitochondrial ADP generator (Dzeja and Terzic, 2009). We therefore reasoned that decreased AK2 activity would alter the cellular balance between AMP and ADP toward a relative increase in AMP. Using tandem mass spectrometry, we quantified the intracellular nucleotide content. In AK2<sup>R175Q/R175Q</sup>–derived myeloid cells, the AMP/ADP ratio was markedly skewed toward AMP whereas intracellular ADP levels were decreased (Fig. 8 F). These data show that AK2 deficiency results in decreased cellular ADP supply, which may limit substrate availability for the ATP synthase and compromise cellular respiration.
As a proof of concept that our in vivo findings in zebrafish can be translated into human cells, we tested the effect of antioxidant treatment in the iPSC model. GSH treatment led to a highly significant improvement in myeloid maturation in a dose-dependent manner. Maximal response was observed at a concentration of 3 mM GSH (Fig. 9 A) and led to an increase.

Antioxidant treatment rescues myeloid differentiation of AK2R175Q/R175Q iPSCs

Our experience in zebrafish has suggested that endogenous ROS production is increased in RD and has demonstrated efficacy of exogenous antioxidant treatment in rescuing hematopoietic developmental defects.
Impairment of AK2 activity in humans leads to the severe lymphocyte and granulocyte differentiation defects and sensorineural deafness characteristic for RD (Lagresle-Peyrou et al., 2009; Pannicke et al., 2009). Our studies in zebrafish and iPSCs show that AK2 deficiency is associated with an even broader hematological phenotype than originally anticipated, which affects the development of the erythroid lineage, in addition to the known abnormalities in granulocyte and lymphocyte differentiation. Although anemia has been reported in several RD patients (Lagresle-Peyrou et al., 2009; Pannicke et al., 2009; Poliani et al., 2009), these observations have been mainly attributed to infections. Our findings in zebrafish and iPSC models strongly suggest that reduced RBC production may be part of the clinical phenotype of the disease.

During substrate binding, AKs undergo a conformational change in the LID and NMP domains from “open” to “closed” (Dzeja and Terzic, 2009; Daily et al., 2010). Once the catalytic process has been completed, the enzyme resumes its original configuration. This transition is important as it limits the catalytic turnover rate (Shapiro et al., 2000; Whitford et al., 2007; Daily et al., 2010). Notably, both missense mutations carried by human AK2R175Q/R175Q iPSCs and ak2L124P/L124P zebrafish mutants are located within the catalytic region of AK2 (inside the LID domain and in a conserved region upstream of the 3D structure of the human AK2 protein. The amino acid position mutated in patient-derived AK2R175Q/R175Q mutant fibroblasts and iPSC lines is highlighted in violet (yellow arrowhead). The amino acid position mutated in ak2L124P/L124P mutant zebrafish is marked in light blue (red arrowhead). Yellow arrowhead marks the amino acid position mutated in patient-derived cell lines. (C–F) iPSCs were generated from AK2-deficient dermal and control foreskin fibroblasts. (C) Representative confocal microscopy images showing the expression of human pluripotency markers (TRA-1-81 and SSEA3 [magenta], Oct4 and NANOG [red], and SSEA4 and TRA-1-60 [green]) using immunofluorescently labeled antibodies in AK2R175Q/R175Q iPSCs; cellular content is highlighted by nuclear staining with Hoechst 33342 (blue). Bars, 100 µm. (D) mRNA analysis using quantitative real-time PCR of the indicated pluripotency-associated genes in AK2R175Q/R175Q iPSCs; cellular content is highlighted by nuclear staining with Hoechst 33342 (blue). Bars, 100 µm. (E) Karyotype and G-banding analysis of AK2R175Q/R175Q iPSCs. (F) Sequencing of the genomic region surrounding the mutation in the control and patient-derived AK2-mutated iPSC line. Data (C–F) are representative of at least two independent experiments.

DISCUSSION
RD is a rare and life-threatening disease affecting infants in the immediate newborn period. Although previous studies have shown that mutations in the AK2 gene are responsible for RD, the rarity of this condition and the lack of suitable animal and cellular models have represented significant obstacles to defining its molecular and cellular pathophysiology. Here we describe ak2 mutant zebrafish and patient-derived iPSCs as platforms to investigate the effects of AK2 mutations on hematopoiesis and to explore novel therapeutic approaches.

Manipulations of AK activity in different organisms result in impaired adenine nucleotide homeostasis, block of ATP export from mitochondria, and reduced cell growth and metabolism, as well as increased susceptibility to metabolic stress (Bandlow et al., 1988; Janssen et al., 2000; Pucar et al., 2000, 2002). Impairment of AK2 activity in humans leads to the severe lymphocyte and granulocyte differentiation defects and sensorineural deafness characteristic for RD (Lagresle-Peyrou et al., 2009; Pannicke et al., 2009). Our studies in zebrafish and iPSCs show that AK2 deficiency is associated with an even broader hematological phenotype than originally anticipated, which affects the development of the erythroid lineage, in addition to the known abnormalities in granulocyte and lymphocyte differentiation. Although anemia has been reported in several RD patients (Lagresle-Peyrou et al., 2009; Pannicke et al., 2009; Poliani et al., 2009), these observations have been mainly attributed to infections. Our findings in zebrafish and iPSC models strongly suggest that reduced RBC production may be part of the clinical phenotype of the disease.

During substrate binding, AKs undergo a conformational change in the LID and NMP domains from “open” to “closed” (Dzeja and Terzic, 2009; Daily et al., 2010). Once the catalytic process has been completed, the enzyme resumes its original configuration. This transition is important as it limits the catalytic turnover rate (Shapiro et al., 2000; Whitford et al., 2007; Daily et al., 2010). Notably, both missense mutations carried by human AK2R175Q/R175Q iPSCs and ak2L124P/L124P zebrafish mutants are located within the catalytic region of AK2 (inside the LID domain and in a conserved region upstream of the
β-sheet connecting the AK2 CORE domain to the LID domain, respectively), which is expected to result in a perturbation of AK2 enzymatic activity and explains the severe phenotype observed in both models.

Our findings in zebrafish null mutants (ak2ΔΔ and ak2Δ+4) and the experiments with ak2 MOs in transgenic Tg(cd41:GFP) reporter zebrafish strongly suggest that, although specification of HSPC is preserved in the absence of AK2, maturation and maintenance of this cell population in zebrafish are dependent on AK2 expression. Although further experiments are necessary to prove a direct effect of AK2 deficiency on human HSPC development, the extended phenotypic spectrum of myeloid and erythroid lineage abnormalities observed in human iPSCs further supports this theory.

In adult bone marrow, HSPCs rely heavily on anaerobic glycolysis for energy production, keeping the generation of ROS and therefore oxidative damage to DNA to a minimum (Jang and Sharkis, 2007; Takubo et al., 2010). Although AK2 deficiency is not expected to have major bearing on glycolysis, it is expected to have a greater impact on cells relying mostly on oxidative phosphorylation to meet their metabolic needs. This could explain how the phenotypic abnormalities observed in RD become more apparent as the cell transitions from predominantly anaerobic to oxidative means of ATP production.
production, which corresponds to the differentiation of HSPCs into lineage-specific mature cells. Alternatively, this might suggest a more subtle effect of AK2 deficiency on the stem cell compartment in humans that has remained unnoticed until now.

Previous studies have linked AK2 function to cell survival and proliferation. In particular, AK2 deficiency in *Drosophila melanogaster* is embryonically lethal (Fujisawa et al., 2009; Chen et al., 2012). Moreover, it has been suggested that AK2 deficiency may induce the unfolded protein response (Burkart et al., 2011) and apoptosis (Single et al., 1998; Köhler et al., 1999; Lee et al., 2007). The intracellular localization of AK2 is strictly limited to the mitochondrial intermembrane space (Dzeja et al., 2007) where it is required to generate ADP. This, in turn, serves as substrate for the ATP synthase and fuels the mitochondrial ATP production critical to meet the energy needs of the cell (Hatefi, 1985). Our observations in zebrafish and iPSCs point to the disturbance of cellular energy homeostasis and increased oxidative stress as key mechanisms responsible for the hematopoietic abnormalities in RD. The decrease in cellular ADP observed in AK2-mutated iPSCs is expected to have a major impact on the activity of the ATP synthase. When substrate unavailability becomes the rate-limiting factor for ATP synthesis and ATP production declines, the cell enters “state 4 respiration,” leading to a transient rise in membrane potential followed by an escalation in the formation of ROS (Liesa and Shirihai, 2013). Whether this promotes mitochondrial uncoupling, resulting in a decline of the mitochondrial membrane potential (Liesa and Shirihai, 2013), or whether ROS acts as second messenger, leading to cell cycle arrest, senescence, and apoptosis (Sauer et al., 2001), remains to be investigated.

Overall, our data suggest that oxidative stress could represent a promising target for novel therapeutic approaches to this syndrome. Accordingly, our models represent a unique opportunity to study the effect of different antioxidant compounds in AK2 deficiency.

The current standard of care in RD is allogeneic HSCT (Antoine et al., 2003; Lagresle-Peyrou et al., 2009). However, the severe neutropenia and lymphopenia in this disease often lead to death before transplantation can be attempted. Furthermore, at variance with other forms of SCID, patients with RD require intensive conditioning to allow for durable donor cell engraftment (Bertrand et al., 2002), further contributing to increased morbidity and mortality. Therefore, addressing the increased oxidative stress by using antioxidant compounds may represent a previously unexploited and easy-to-implement strategy to treat patients with RD and may have important implications for the development of clinical
trials aimed at alleviating the disease phenotype, while preparing for HSCT.

MATERIALS AND METHODS
Zebrafish lines and maintenance
Zebrafish were maintained and used following protocols approved by the National Human Genome Research Institute Animal Care and Use Committee. Zebrafish handling, breeding, and staging were performed as previously described (Kimmel et al., 1995; Westerfield, 1995). Beginning from 24 hpf, embryos were cultured in fish water containing 0.003% 1-phenyl-2-thiourea (PTU) to prevent pigmentation. The following stages were used: WT EK (Etkwill) and Tg(y1a41:GFP) (Lin et al., 2005) and Tg(y1a1a: EGFP)y1;Tg(gata1a:dsRed)y12 (Lawson and Weinstein, 2002; Traver et al., 2003).

Databases and bioinformatic analysis of data
The following genome assemblies were searched on the Ensembl database (Flicek et al., 2013; Homo sapiens (GRC37.p12, Feb 2009) and Danio rerio (Zv9, Apr 2010)). Multi-alignment of human and zebrafish AK2 was performed with Clustal Omega (Goujon et al., 2010). In each multi-alignment, an asterisk indicates positions that have a single, conserved residue, a colon indicates conservation between groups of strongly similar properties, and a period indicates conservation between groups of weakly similar properties. Structural features of amino acid sequence of human AK2 were defined on 3D structure of human protein (MMDDB ID: 37222). In all of the 3D structures, α-helix structures are depicted in green, β-strands are depicted in yellow, and Bio(Adenoine)-5'-Triphosphate and 1,2-Ethanediol are shown as ball and stick models. PROVEAN score was used to predict whether a protein sequence variation affects protein function (Choi et al., 2012).

Accession number
The GenBank accession number of zebrafish ak2 isoform B is KJ651951.

RT-PCR
Total RNAs were prepared from different zebrafish embryos at different developmental stages using the RNeasy Mini kit (QiAGEN), treated with DNase I RNA-free (Roche) to avoid possible contamination from genomic DNA, and then reverse transcribed using Superscript III (Life Technologies) and random primers. The cDNAs were then subjected to PCR amplification using specific primers and Platinum Taq DNA Polymerase or Platinum Taq DNA Polymerase High Fidelity (Life Technologies) as previously described (Risone et al., 2012). To the extent possible, all primer pairs were designed on different exons to avoid the amplification of DNA isoforms. PCR products were cloned into the pCR4-TOPO vector (Life Technologies), and DIG RNA Labeling kit (Roche); lba1, lplastin, mpx, irata, rqt1, and c-myb. For hmx1a isoform antisense probes, specimen-specific fragments were amplified by RT-PCR on suitable templates. The primers (lhx1a ISHI FOR, 5'-TTGGACTCTCACACAAAAAAGAGC-3' and lhx1a ISHI REV, 5'-CCTTCC-TCTTGTATGCTACGAC-3') were designed to cover the coding sequence of the gene of interest. PCR products were cloned into the pCR-4-TOPO vector (Life Technologies), and DIG RNA Labeling kit (SP6/T7; Roche) was used to synthesize antisense or sense RNA probes. All probes were purified using Spin Purification Clean-Up Columns (Sigma-Aldrich) before use. All embryos used for WISH were fixed overnight in 4% paraformaldehyde (PFA)/PBS, rinsed with PBS-Tween, dehydrated in 100% methanol, and stored at −20°C until being processed for WISH. WISHs were performed as described by Thiise and Thiise (2008). Hybridization was performed at 70°C. Hybridized probes were then detected by using an anti-DIG antibody conjugated to alkaline phosphatase (AP; Roche) at a 1:5,000 dilution. Nitro blue tetrazolium/5-bromo-4-chloro-3-indolyl phosphate (NBT/BCIP; Roche) or INT/BCIP Stock Solution (Roche) was used as a substrate for AP. Stained embryos were stored in 4% PFA until imaging.

Sudan black and acridine orange stainings
For Sudan black staining, embryos were fixed with 4% PFA (Polysciences) in PBS for 2 h at room temperature, rinsed in PBS, and incubated in Sudan black solution (Sigma-Aldrich) as described previously (Sheehan and Storey, 1947; Le Guyader et al., 2008). For acridine orange staining, zebrafish larvae were anesthetized with Tricane (Sigma-Aldrich), incubated in a solution of 3 μg/ml acridine orange (Sigma-Aldrich) in E3 medium with Tricane (Sigma-Aldrich) for 30 min, washed twice in E3 with Tricane, and then analyzed under a fluorescent stereomicroscope.

TUNEL assay
TUNEL assay was performed as previously described (Finckbeiner et al., 2011) with minor modifications. After incubation in STOP/Wash solution followed by 3 × 10 min PBT washes, the embryos were incubated in anti-DIG Fluorescein-conjugated antibody (EMD Millipore) or in anti-DIG AP-conjugated antibody (Roche) using NBT/BCIP (Roche) as substrates for AP.

Cell sorting and oxidative stress staining
Whole mount staining with the Mitotracker and CellROX oxidative stress probes was performed as previously described (Mugoni et al., 2014) on living zebrafish.
zebrafish embryos at different stages of development. Embryos from incross of ak2+/- and ak2+/- were subjected to tail resorption 30 mm before being stained with 5 μM MitoSOX or CellROX solutions for 30 min, followed by fluorescence analysis using confocal microscopy. At the end of the analysis, embryos were recovered and genotyped. Quantitative analysis of the oxidative stress using flow cytometry was performed as previously described (Maugoni et al., 2014) with the following modifications. Embryos from incross of ak2+/- homozygous fish were grown until the selected stage (4 or 5 dpf), and then null mutants and their siblings (~60 embryos each) were separated and collected in different tubes. After euthanasia, embryos were washed twice with 1× PBS and then dissociated with a mix of collagenase P and Trypsin-EDTA in 1× PBS at 28°C for 20 min. Homogenization of the samples was obtained by extensive periodical pipetting. Then the cells were maintained in DMEM 10% FCS at 28°C until staining. For MitoSOX analysis, cells were resuspended in HBSS containing calcium and magnesium and then stained with 5–2.5 μM MitoSOX Red (Life Technologies) for 15 min. For CellROX Green (Life Technologies) staining, the probe was added to the DMEM 10% FCS medium at a final concentration of 2.5–1 μM for 30 min. For Annexin V–APC and 7-AAD (BD) double staining, cells were washed and resuspended in 1× binding buffer and then incubated with the probes according to the manufacturer’s instruction. In all cases, the cells were incubated at 28°C in the dark and then washed twice to remove excess probe. Flow cytometry data were collected on a FACSAna II equipped with FACSiva software using the blue 488-nm and red 640-nm lasers (BD). Further data analysis was performed using FlowJo software (FlowJo LLC). Unstained cells were used to set the gates in the FlowJo program.

Imaging

The embryos obtained by WISH, Sudan black staining, acridine orange, and TUNEL assay were observed with an MZ16F stereomicroscope (Leica), and pictures were taken with a DC500 camera using a FireCam (version 1.7.1; Leica). Confocal images were acquired at room temperature using an LSM 510 NLO Meta system mounted on an Axiovert 200M microscope (Carl Zeiss) with a Plan-Apochromat 10×/0.45 or a Plan-Apochromat 20×/0.75 objective lens. Excitation wavelengths of 488, 561, and 770 nm were used for detection of EGFP, dRed, and DAPI, respectively. A range of z-slices was used depending on the zebrafish orientation to capture all desired structures using optical slices of ~4 μm. All confocal images were of frame size 512 pixels by 512 pixels, scan zoom of 0.7×, and line averaged four times. Images were collected, and maximum projections were processed using the ZEN 2009 V.5.3 SP2 software package (Carl Zeiss).

Morpholino injection

Antisense morpholinos targeting the ak2 ATG region (ak2 MO1, 5′-CATGG-GCTACAGCTTCTTCTTAATCT-3′) or the splice acceptor site of ak2 intron 3 (ak2 MO2; Pannicke et al., 2009) and the standard control MO (ST-CTRL MO, 5′-CCCTTACCTCAGTTACATTATATA-3′) were manufactured by Gene Tools. The morpholinos were used according to the manufacturer’s instructions, and the indicated amount of each morpholino was injected into the yolk of one-cell-stage WT embryos. To confirm that AK2 pre-mRNA was specifically targeted, RT-PCR was performed on ST-CTRL MO–injected embryos was injected into the yolk of one-cell-stage WT embryos. To confirm that AK2 pre-mRNA was specifically targeted, RT-PCR was performed on ST-CTRL MO–injected embryos. RT-PCR on ST-CTRL MO–injected embryos was performed as described above. The starting cell count of 3 × 10⁴ fibroblasts was maintained in hESC medium containing DMEM, 20% KOSR, 10 mg/ml bFGF1 mmol/liter l-Glutamine, 100 mmol/liter nonvesseous amino acids, 100 mmol/liter 2-mercaptoethanol, and 100 mg/liter Primocin as previously described (Park et al., 2008b). Karyotype analysis was performed by Cell Line Genetcs. Genomic DNA was isolated using the QIAamp DNA kit (Qiagen), and the genomic regions flanking the reported mutations were PCR amplified using primers AK2 M13 F (5′-ACAGCATTGGGCGCGGGAAGATT-3′) and AK2 M13 R (5′-ATTCCTCACCCTATGGGCGCTAC-3′) and sequenced in a commercial core facility (Eon Bioscience).

Immunohistochemistry

iPSCs were selected based on morphology, cloned for five to seven passages, and then manually picked as single-colony cells and seeded into a 96-well plate wells containing 8.5 × 10⁴ irradiated mouse embryonic fibroblasts per well in standard hESC medium. Cells were washed with PBS, fixed at room temperature in 4% PFA/PBS for 15 min, washed with PBS, and permeabilized/blocked for 1 h at room temperature with 0.1% Triton X-100 in animal-free blocker (Vector Laboratories) followed by three washes with PBS. Cells were subsequently incubated with primary antibodies (OCT4: clone 40/0ct-3, Alexa Fluor 555, 1:50 dilution; NANOG: N31-355, PE, 1:30; SSEA4: MC813-70, Alexa Fluor 488, 1:200; SSEA3: MC-631, Alexa Fluor 647, 1:30; TRA-1-81: Alexa Fluor 647, 1:30; TRA-1-60: Alexa Fluor 488, 1:50; all from BD) and 2 µg/ml Hoechst 33342 in low-binding buffer. After 1 h, cells were washed twice with PBS, fixed at room temperature in 4% PFA/PBS for 15 min, and then incubated with the secondary antibodies (anti-Alexa Fluor 555, Alexa Fluor 488, and Alexa Fluor 647, 1:500). Images were acquired using a DMi8 microscope (Leica) using a 10× lens. Background images (empty well images) were used depending on the imaging parameters using the ImageJ software (National Institutes of Health). Immunofluorescence intensity was compared with a control hES cell line as internal standard to yield semiquantitative results.

Quantitative real-time PCR. RNA was extracted from 10⁵ cells using the mirVana RNA isolation kit (Ambion) and reverse transcribed to cDNA with qScript cDNA supermix (Quanta BioScience) according to the manufacturer’s instructions. Expression of the OCT4, KLF4, MYC, SOX2, NANOG, GDF3, and DNMT3B genes was quantified by real-time PCR with Power SYBR Green PCR Master Mix (Applied Biosystems) on a 7500 Real Time PCR analyzer (Applied Biosystems) and normalized to the human B-actin gene expression (hACTB) using primers previously reported (Park et al., 2008b). Relative expression compared with parental fibroblasts was calculated using the ∆∆Ct (ddCt) method.

Directed myeloid differentiation

EBs were generated in suspension culture from iPSCs using a bFGF-free hESC cell culture medium containing 300 ng/ml SCE, 300 ng/ml FTLD, 50 ng/ml G-CSF, 10 ng/ml IL3, 10 ng/ml IL6, and 50 ng/ml BMIP4 on low-adhesion plates as previously described (Park et al., 2008a). After 14 d, EBs were dissociated using 0.5 mg/ml Collagenase B, nonenzymatic cell dissociation buffer (Gibco) and mechanical sheering. EB–derived single cells were plated on methylcellulose (MethoCult; STEMCELL Technologies) and cultured for 14 further days according to the manufacturer’s instructions in 21% O₂, 5% CO₂. For samples grown in the presence of GSH, a 100 mM stock solution was freshly prepared from l-Glutathione (reduced; Sigma-Aldrich) and added to the methylcellulose to yield a final concentration of 1–3 mM. For samples grown

Reprogramming.

Human dermal fibroblasts were collected from a patient with homozygous c.524G>A mutation in the human AK2 gene (Henderson et al., 2013) by punch biopsy. All human protocols were approved by Boston Children’s Hospital Institutional Review Board. Fibroblasts were expanded in hFIB medium containing DMEM, 20% FBS, 1 mmol/liter l-Glutamine, 100 mmol/liter nonvesseous amino acids, and 1% PenStrep. Fibroblasts were reprogrammed using an excisable, self-inactivating, codon-optimized lentiviral vector carrying OCT4, SOX2, KLF4, MYC, and dTomato CDNAs as previously described (Warlich et al., 2011). Transduction was performed with a starting cell count of 3 × 10⁴ fibroblasts using a multiplicity of infection (MOI) of 1. iPSC colonies emerged after 17–21 d of culture. To generate control iPSCs, human neonatal foreskin fibroblast “BJ” (ATCC CRL-2522) was used, and reprogramming was performed as described above.

iPSC culture. iPSCs were maintained on irradiated mouse embryonic fibroblasts (Global Stem) in hESC medium containing DMEM, 20% KOSR, 10 mg/ml bFGF1 mmol/liter l-Glutamine, 100 mmol/liter nonvesseous amino acids, 100 mmol/liter 2-mercaptoethanol, and 100 mg/liter Primocin as previously described (Park et al., 2008b). Karyotype analysis was performed by Cell Line Genetcs. Genomic DNA was isolated using the QIAamp DNA kit (Qiagen), and the genomic regions flanking the reported mutations were PCR amplified using primers AK2 M13 F (5′-ACAGCATTGGGCGCGGGAAGATT-3′) and AK2 M13 R (5′-ATTCCTCACCCTATGGGCGCTAC-3′) and sequenced in a commercial core facility (Eon Bioscience).
in the presence of ATRA and G-CSF retinoic acid powder (Sigma-Aldrich) was reconstituted in DMSO to make a 1 mM stock solution and added to methylcellulose to a final concentration of 1 µM. G-CSF (Neupogen) powder was reconstituted in sterile PBS with 0.1% serum albumin and added to methylcellulose to a final concentration of 25 ng/ml.

Electron microscopy
One million iPSC-derived myeloid cells were pelleted and resuspended in 200 µl of 25% glutaraldehyde and 200 µl PFA-picric acid. Sections were cut and grids prepared by the Harvard Medical School Electron Microscopy Core Facility. Images were obtained using a Tecnai G2 Spirit BioTWIN microscope (FEI).

Tandem mass spectrometry
Shock-frozen pellets from 1–2 × 10^7 iPSC-derived myeloid cells were reconstituted in 300 µl water and sonicated on ice for 10 s. Lysate volumes of equal total cell numbers were precipitated by adding 250-µl volume of acetone-titrated and centrifuged at 16,200 rpm for 2 mm. Supernatant was transferred to a new vial, dried under nitrogen flow, and resuspended in 50 µl water. A calibration curve, containing mixtures of ATP, ADP, and AMP, ranging from 0 to 10 µmol/liter (0.01, 0.1, 1, and 10 µmol/liter) was used to quantify nucleotides. Nucleotide quantification was performed using a Triple-Quad Mass Spectrometer (8040; Shimadzu) equipped with a turbo ion spray source and coupled to a Nexera UPLC (Shimadzu).

Statistical analysis
Statistical analyses were performed using Excel software (Microsoft). Asterisks refer to p-values. P-values and statistical tests applied are listed in the figure legends.

Online supplemental material
Table S1 shows sequences of primers and quantitative PCR probes used in this study. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.201412566/D1C1.

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REFERENCES


