Loss of Hspa9b in zebrafish recapitulates the ineffective hematopoiesis of the myelodysplastic syndrome

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Myelodysplastic syndrome (MDS) comprises a heterogeneous group of clonal blood disorders characterized by ineffective hematopoiesis arising from dysplasia and accelerated apoptotic death of multipotential hematopoietic progenitors and their progeny. These syndromes display significant clinical variability, and whereas some patients present solely with anemia and erythroid dysplasia, many develop fatal multilineage peripheral cytopenia or even progress to acute myeloid leukemia (AML). Treatment options are limited, and the majority of patients with MDS die within 3 to 4 years. Thus, the identification of genes whose loss or altered function produces defective hematopoiesis is critical for diagnosis, improved therapies, and prevention of neoplastic transformation.

Numerous chromosomal abnormalities have been linked to MDS, and although karyotypes are usually quite complex, interstitial deletions in the long arm of chromosome 5, del(5q), are prevalent in both de novo and therapy-induced cases. A distinct subset of MDS, the 5q− syndrome, is characterized by good prognosis and deletions at chromosome 5q32 as the sole karyotypic abnormality. In contrast, deletions at 5q31 are associated with more aggressive forms of MDS, often progressing rapidly to leukemia. A critical deleted region (CDR) at chromosome 5q31 has been described containing 9 candidate genes, including anemia, dysplasia, increased blood cell apoptosis, and multilineage cytopenia. By positional cloning, rescue, and morpholino knockdown experiments, we demonstrate that crs encodes a conserved mitochondrial matrix chaperone HSPA9B containing a glycine-to-glutamate substitution within the substrate-binding domain. This mutation compromises mitochondrial function, producing oxidative stress and apoptosis distinctly in blood cells. Thus, we identify an essential role for Hspa9b in hematopoiesis and implicate both loss of HSPA9B specifically and mitochondrial dysfunction generally in the pathogenesis of the MDS.

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Introduction

Myelodysplastic syndrome (MDS) comprises a heterogeneous group of clonal blood disorders characterized by ineffective hematopoiesis, arising from dysplasia and accelerated apoptotic death of multipotential hematopoietic progenitors and their progeny. These syndromes display significant clinical variability, and whereas some patients present solely with anemia and erythroid dysplasia, many develop fatal multilineage peripheral cytopenia or even progress to acute myeloid leukemia (AML). Treatment options are limited, and the majority of patients with MDS die within 3 to 4 years. Thus, the identification of genes whose loss or altered function produces defective hematopoiesis is critical for diagnosis, improved therapies, and prevention of neoplastic transformation.

Numerous chromosomal abnormalities have been linked to MDS, and although karyotypes are usually quite complex, interstitial deletions in the long arm of chromosome 5, del(5q), are prevalent in both de novo and therapy-induced cases. A distinct subset of MDS, the 5q− syndrome, is characterized by good prognosis and deletions at chromosome 5q32 as the sole karyotypic abnormality. In contrast, deletions at 5q31 are associated with more aggressive forms of MDS, often progressing rapidly to leukemia. A critical deleted region (CDR) at chromosome 5q31 has been defined containing 9 candidate genes, including HSPA9B, but causative genes have yet to be identified.

Compromised mitochondrial function is implicated in the pathogenesis of a wide variety of diseases, including MDS. Structural abnormalities and iron accumulation are frequently found in the mitochondria of bone marrow cells from patients with MDS, and these cells may display a proapoptotic profile, including activation of the mitochondrial death machinery and spontaneous release of cytochrome c. Furthermore, mutations in the mitochondrial genome have been reported. However, the exact role mitochondrial dysfunction plays in the ineffective hematopoiesis of MDS remains unclear.

Hematopoiesis is well characterized in the zebrafish Danio rerio and demonstrates a high degree of conservation with mammalian counterparts. Large-scale forward genetic screens have isolated zebrafish mutants at various stages in blood development and several mutants model human hematopoietic diseases. Here we present a characterization of the zebrafish developmental blood mutant crimsonless (crs). crs mutants are anemic at the onset of circulation and display defective blood cell differentiation followed by apoptosis and a reduction in erythrocytes, granulocytes, and hematopoietic progenitors. Furthermore, crs haplodeficiency increased apoptosis specifically in adult hematopoietic progenitors. Thus, the crs phenotype closely resembles the symptoms of MDS, and crs animals provide an additional valuable vertebrate model of these disorders.

Using positional cloning, RNA rescue, and morpholino knockdown, we identified the mutated gene in crs as Hspa9b (mortalin/mthsp70/GRP75), a highly conserved, ubiquitously expressed, mitochondrial matrix chaperone. Mutant HSPA9B produces an increase in reactive oxygen species (ROS) and oxidative stress distinctly in blood cells. Thus our results demonstrate that Hspa9b is essential for proper hematopoiesis, and its loss and subsequent mitochondrial dysfunction in blood cells are sufficient to recapitulate characteristic symptoms of MDS.

Materials and methods

Fish stocks and maintenance

Fish were maintained and staged as described. Embryos raised beyond 24 hours postfertilization (hpf) were treated with phenylthiourea (PTU; 0.003% wt/vol; Sigma, St Louis, MO) to prevent melanization.

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Mapping and cloning

Fish carrying the crs mutation were out-crossed to WIK for positional cloning. Polymerase chain reaction (PCR) amplification of simple sequence length polymorphic (SSLP) genetic markers from gDNA of mutant versus heterozygous and wild-type (WT) crs embryos was used to establish linkage to linkage group 14 (LG14) between the markers z43267 and z26376. A chromosome walk was initiated from the closest marker z43267 (0.36 cm; 8 of 2239 recombinants) and single-stranded conformation polymorphism (SSCP) analysis was used to orient PI-derived artificial chromosomes (PAC) clones. Two PACs that displayed zero recombinants (0 of 2239) were identified. The PACs were sequenced, and both contained the complete coding region of the Hspa9b (GenBank accession no. 201326). A mutation in Hspa9b in crs embryos was demonstrated both by sequencing exons from gDNA and by reverse transcription-PCR (RT-PCR) of total RNA from mutant versus wild-type embryos. Full-length zebrafish Hspa9b was cloned by RT-PCR of total RNA from wild-type embryos using the TA cloning vector pCRII (Invitrogen, Carlsbad, CA).

Whole-mount in situ hybridization, o-dianisidine, acridine orange, and dihydrorhodamine 123 staining

Digoxigenin-labeled antisense RNA probes were hybridized at 68 to 70°C and developed with nitroblue tetrazolium (NBT) as described previously.31 Probes for gata1 and α and β-globins were generously provided by Dr Len Zon (Children’s Hospital, Howard Hughes Medical Institute, Chevy Chase, MD). Probes for alas2 and heme oxygenase-1 were isolated by RT-PCR and were TA cloned into pCRII (Invitrogen). Staining with o-dianisidine and labeling with acridine orange (AO) were performed as described.24 Labeling with Mitotracker 580 and dihydrorhodamine 123 (Molecular Probes, Carlsbad, CA) was performed according to the manufacturer’s protocol.

Histologic and TUNEL staining of embryonic blood cells and adult organs

Embryonic blood was isolated by cardiac puncture. Kidneys or liver from adult heterozygous and wild-type animals were dissected out and smeared onto glass slides. Wright-Giemsa staining was performed as previously described25 and TdT-mediated dUTP nick-end labeling (TUNEL) analysis was performed using DeadEnd Fluorometric TUNEL System (Promega, Madison, WI) according to the manufacturer’s protocol.

Whole-mount immunofluorescence

Embryos were dechorionated and fixed in 4% paraformaldehyde for 1 hour at room temperature or 4°C overnight. Embryos were equilibrated with 4 to 5 washes in −20°C methanol and stored overnight (minimum 1 hour) at −20°C, rehydrated through 50:50 methanol/phosphate-buffered saline with 0.1% Tween (PBST), washed once with dH2O, treated with −20°C acetone for 7 minutes, and washed with PBST. Embryos were blocked for 2 hours at room temperature in PBST, 5% normal goat serum (NGS), 2 mg/mL bovine serum albumin (BSA), 0.5% dimethyl sulfoxide (DMSO), and then incubated with 1st antibody (rabbit anti–active caspase-3, PharMingen [San Diego, CA], catalog no. 559565; rabbit antiphospho-histone H3, Upstate Biotechnology [Lake Placid, NY], catalog no. 06-570) in block overnight at 4°C. Embryos were washed in PBST and incubated 2 hours at room temperature with 2nd cyanine 3 (Cy3) antibody in block and washed finally in PBST before mounting and imaging.

RNA and morpholino injections

Capped sense and antisense RNA was synthesized using the mMESSAGE mCARTINE kit (Ambion, Austin, TX) according to the manufacturer’s protocol from linearized pCS2+ plasmids containing full-length Hspa9b. The sequence of the morpholino-modified antisense oligonucleotide (Gene Tools, Philomath, OR) targeting the zebrafish Hspa9b gene is as follows: 5’-TTGCCCGTTCTCGACACAGACAACAT. RNA (~250 pg) or morpholinos (5-10 ng) was microinjected at the 1- to 4-cell stage. At 33 to 36 hpf, injected embryos were processed for o-dianisidine staining.

RT-PCR

RNA was isolated from wild-type versus mutant embryos (Qiagen, Valencia, CA). Either 300 ng was used for Qiagen one-step RT-PCR according to the manufacturer’s protocol and amplification products gel analyzed, or 25 ng for real-time RT-PCR (Applied Biosystems, Foster City, CA) using TdTAMFAM-labeled probe/primer sets for c-mpl with GAPDH as a comparison and α,β-globin as a positive control.

Image acquisition

Images were taken on either (1) a Leica dissecting microscope at 10 to 40× with a Plan Apo objective (Leica Microsystems, Bannockburn, IL) in aqueous solution (Figures 1, 3D-E, and 5A-B); (2) a Nikon TE300 inverted microscope (Nikon, Kanagawa, Japan) with a × 60 oil objective with embryos mounted in Flurromount G (Figure 2A-D), or Plan Fluro × 200.45 air objective (Figures 2G-J and 4C-E), or Plan Fluro × 10/0.30 air objective (Figure 4A-B) in aqueous solution; or (3) a Nikon E800 upright microscope with Plan Apo × 40/0.95 air objective (Figures 2E-F, 4F, and 7), or Plan Apo × 200.75 air objective (Figures 4G-H, 5C-D, and 6) with embryos mounted in Flurmount G. All images were taken at room temperature with an RT Slider Spot camera (Diagnostic Instruments, Sterling Heights, MI) and acquired with Spot 3.52, version software for Windows. Subsequent manipulations were made using Adobe Photoshop (Adobe, San Jose, CA).

Results

crs mutants display a distinctive defect in blood development.

A single autosomal recessive allele of crimsonless (crs) was identified in a chemical mutagenesis screen for developmental mutants. crs embryos appeared visibly normal until approximately 33 hpf when light microscopy revealed hypochromic blood despite a normal number of cells in circulation. Analysis of erythroid hemoglobin content by whole-embryo staining with o-dianisidine provided evidence of hemoglobin in mutant animals compared to wild-type at both 33 hpf (Figure 1A-B) and 48 hpf (Figure 1F). This late generation of hemoglobin was associated with an up-regulation of e1-globin as a positive control. Delayed blood cell development was complemented by the failure of hemoglobin production at 33 hpf as revealed by whole-mount in situ hybridization for scl, gata2, and gata1 (Figure 1G-H and data not shown). Furthermore, mutant animals displayed normal expression of both α,β-globin and β,γ-globin (Figure 1K-L and data not shown), and intravenous injection of iron-dextran failed to rescue hemoglobin production (data not shown). Thus, neither loss of blood formation, loss of globin chain expression, nor inadequate levels of circulatory iron account for the hypochromia of crs mutant embryos.

Accompanying the anemia, mutant blood cells failed to properly differentiate into mature erythrocytes. Hematoxylin and eosin staining at 33 hpf significantly delayed development of red blood cells in crs mutant fish with large nuclei and basophilic cytoplasm characteristic of immature erythroblasts (Figure 2A-B). At 48 hpf mutant blood cells remained arrested at the erythroblast stage compared to the small, ellipsoid-shaped erythrocytes of wild-type animals (Figure 2C-D). Delayed blood cell development in mutant animals was also seen by Wright-Giemsa staining of isolated blood cells at 36 hpf (Figure 2E-F); however Prussian blue staining failed to detect the presence of ringed sideroblasts from altered iron utilization (data not shown). Further signs of dysplastic blood cells in mutant animals included the presence of binucleated cells (arrows in Figure 2B) and the failure to down-regulate expression of the early blood cell marker gata1 (Figure II-J). Yet despite these defects, a small number of crs blood cells did produce hemoglobin by 48 hpf (Figure 1F). This late generation of hemoglobin was associated with an up-regulation of alas2, the
mutant animals (bracket in Fi) are further magnified and indicated by an arrowhead
higher magnification (E-Fi). A few o-dianisidine–positive cells over the yolk sac in
flow in the tail vessels. Blood flow over the yolk sac is bracketed (C-D) and shown
hpf (G,H,K,L) and 48 hpf (I-J) in wild-type (G,I-K) and mutant (H,J,L) embryos.

In contrast to the profound hematopoietic defect at 33 to 46 hpf,
all other organ systems developed normally up to 48 hpf as
revealed by molecular markers for vasculature, gut, and nervous
system (data not shown). One exception was the head and eyes,
which appeared stunted as early as 38 hpf. However, after 48 hpf all
development appeared to halt, including further maturation of the
musculature, fins, and internal organs, and most animals failed to
survive past 72 hpf when the heart slowed and eventually stopped
beating (data not shown).

**Mutant Hspa9b is sufficient to account for the crs phenotype**

To identify the gene responsible for defective hematopoiesis in crs
embryos, we took a positional cloning strategy. Using polymorphic
microsatellite markers, we mapped the crs mutation to LG14
between markers z43267 and z26376 (Figure 3A). A chromosome
walk initiated from the closer marker z43267 (0.36 cM; 8 of 2239
recombinants) identified 2 partially overlapping PAC clones that
displayed zero recombinants (0 of 2239 recombinants) and were
sufficient to rescue the anemia in crs embryos (data not shown).
The PACs were sequenced, and both contained the complete coding
region of the ubiquitously expressed, mitochondrial matrix chaperone
protein HSPA9B (Mortalin/mthsp70/GRP75). Sequence analysis
of Hspa9b in crs mutants uncovered a point mutation leading to
a glycine-to-glutamate substitution at position 492 (G492E) within
the substrate-binding domain of HSPA9B (Figure 3B-C). This
rate-limiting enzyme in heme production (data not shown), suggest-
ing that compensatory gene regulation allowed for the production
of some functional hemoglobin in mutants.

In addition to anemia and dysplasia, the number of blood cells
in circulation was also affected in mutant animals. Although
normal at the beginning of development, the number of circulating
blood cells began to decrease starting at around 36 hpf and by 38 to
42 hpf their numbers were on average one half to one fourth that of
wild-type (Figure 2G-H). There was a reduction of cells flowing
over the yolk sac and through the vessels including the caudal
artery (Figure 2G-H and data not shown) with no signs of pooling
or a decrease in the rate of blood flow. Subsequently, the number of
blood cells in circulation either remained constant or in some
individuals decreased further until death at approximately 72 hpf.

**Figure 1. crs embryos are anemic despite normal blood specification.** (A-F)
Whole-mount o-dianisidine stain for hemoglobin in wild-type (WT) (A,C,E) and mutant
(−/−) (B,D,F) embryos at 33 hpf (A-B) and 48 hpf (C-F). Arrows (C-D) indicate blood
flow in the tail vessels. Blood flow over the yolk sac is bracketed (C-D) and shown
higher magnification (E-Fi). A few o-dianisidine–positive cells over the yolk sac in
mutant animals (bracket in Fi) are further magnified and indicated by an arrowhead
(Fii). (G-L) Whole-mount in situ hybridization for gata1 (G-J) and β-globin (L) at 33
hpf (G,H,K,L) and 48 hpf (I-J) in wild-type (G,J,K,L) and mutant (H,J,L) embryos.

**Figure 2. crs red blood cells fail to differentiate and are reduced in number.**
(A-D) Hematoxylin and eosin staining of blood cells from whole-mount wild-type (A,C)
and mutant (B,D) embryos at 33 hpf (A-B) and 48 hpf (C-D). Binucleated cells were
often observed (B, arrows). (E-F) Wright-Giemsa staining of isolated blood cells from
wild-type (E) and mutant (F) embryos at 36 hpf. (G-H) Bright-field microscopy of blood
cells in wild-type (G) and mutant (H) embryos at 42 hpf in the caudal artery with
spaces between the reduced cell numbers in mutant animals bracketed (H).

**Figure 3. crs encodes a conserved mitochondrial matrix chaperone, HSPA9B.**
(A) The crs mutation is on linkage group 14 (LG14) between the polymorphic
microsatellite markers z43267 and z26376. The distance between crs and each
polymorphic marker is shown in centimorgans on the left, and the number of
recombinants of 2239 meioses is on the right. No recombinants were found on 2
PACs that contain the entire zebrafish hspa9b gene. (B) ABI automated sequencer-
produced chromatographs from wild-type (top) and mutant (bottom) mRNA. A G>A
point mutation produces a glycine-to-glutamate conversion at amino acid 492 in
zebrafish HSPA9B. (C) Domain structure of HSPA9B. HSPA9B has an N-terminal
adenosine triphosphatase domain (amino acids 55-434) immediately followed by a
propeptide-binding domain (amino acids 435-587) within which the G492E mutation
lies (red asterisk). Underneath the domain map is an alignment of the propeptide-
binding domains of zebrafish and human HSPA9B and E. coli DnaK chaperones with
conserved residues in gray and the conserved glycine mutated in crs zebrafish
marked with a red asterisk. (D-E) Whole mount o-dianisidine staining of embryos
injected with control (D) or hspa9b (E) morpholino-modified antisense oligonucleo-
tides (MO) at 36 hpf.
mutation was confirmed by RT-PCR of total RNA from mutant versus wild-type embryos. Zebrafish HSPA9B shares 84.8% identity and 89.4% similarity with human HSPA9B and 53.5% identity and 63.3% similarity with the *Escherichia coli* DnaK chaperone. A near-identical mutation in the conserved glycine at position 443 in DnaK completely abolishes propeptide binding, rendering the chaperone functionless.33 Thus, *crs* likely represents a null allele of *Hspa9b*.

To verify that the mutation in *Hspa9b* is sufficient to cause the *crs* phenotype, we used both rescue and antisense morpholino knockdown experiments. Injection of capped RNA encoding wild-type HSPA9B rescued approximately 95% (53 of 56) injected mutant embryos. The blood of rescued animals was no longer hypochromic with o-dianisidine staining close to wild-type levels, and cytopenia did not occur up to 48 hpf (data not shown). No overexpression phenotype was observed in *Hspa9b*-injected embryos, and embryos injected with antisense RNA as a control were not rescued (0 of 44 injected; data not shown). Conversely, we inactivated zebrafish *Hspa9b* using antisense morpholino-modified oligonucleotides that targeted the 5′ untranslated region (UTR) and initiation methionine. Morpholino-injected embryos recapitulated the anemic phenotype observed in *crs* mutants, and approximately 80% of the injected animals showed decreased o-dianisidine staining compared to wild-type at 36 hpf (Figure 3D-E). Taken together these results provide strong support that the G492E mutation in HSPA9B is the cause of the *crs* phenotype.

**Loss of *Hspa9b* recapitulates MDS**

Human HSPA9B is located on the long arm of chromosome 5 within a CDR at 5q31 associated with MDS10,11 and is thus a candidate for playing an important role in a subset of these blood disorders. Given that ineffective hematopoiesis including dysplasia, anemia, and a decline in circulating blood cells characterize both patients with MDS and *Hspa9b* mutant zebrafish, we further analyzed the blood defect of *Hspa9b* mutants to validate its relevancy to the human disease.

Although anemia is most typical of patients with MDS, it often occurs in conjunction with neutropenia or thrombocytopenia or both.2 We therefore determined if multilineage cytopenia is present in *Hspa9b* mutant fish by examining myeloperoxidase-positive (mpo+ cells) of the neutrophil lineage, L-plastin-positive monocytes/macrophages, and CD41+ thrombocytes.34,35 *L*-plastin-positive myeloid cells, which initially originate outside the intermediate cell mass (ICM) over the anterior yolk sac,34 were unchanged in number (data not shown), and similarly, despite failing to detect CD41 or the thrombopoietin receptor (*c-mpl*) by in situ, real-time RT-PCR revealed equivalent levels of *c-mpl* expression at 46 hpf in wild-type and mutant animals (data not shown). Yet in striking contrast, mpo+ granulocytes, though unaffected at 18 to 24 hpf in the ICM (data not shown), are clearly reduced in number by 38 to 42 hpf in *crs* mutant embryos compared to wild-type (Figure 4).

Thus our results differ from those of other zebrafish blood mutants with exclusive erythroid defects25,26,34 and instead link the loss of *Hspa9b* not only with anemia but also with multilineage cytopenia.

The ineffective hematopoiesis of MDS is associated with increased apoptosis of hematopoietic stem cells and their progeny.1,13 Accordingly, we analyzed whether blood cell apoptosis preceded cytopenia in *Hspa9b* mutant fish. An hour of incubation of live embryos with AO at 36 hpf labeled few dying cells in wild-type animals, and no blood cells were AO+. By contrast, in mutant animals AO labeled circulating blood cells as well as immobile cells in the posterior ICM of the ventral tail (Figure 5B-E). Furthermore, an identical population of mutant cells was immunoreactive for activated caspase-3 (Figure 5F). Because the posterior ICM is a site where both hematopoietic progenitors36 and phagocytic cells have been described,37 these stationary cells may include both blood cells cleared from circulation as well as dying progenitors. Consistent with the latter, at 42 hpf *c-myb*+ progenitors in the dorsal aorta were significantly reduced in mutant animals as compared to wild-type (Figure 5G-H). In contrast to the dramatic increase in cell death, the number of cells immunoreactive for phospho-histone H3, a marker of cell proliferation, was unchanged (data not shown). Cells in the eyes of mutant animals were the only other population that were positive for AO and activated caspase-3 (Figure 5B), and no additional cell populations labeled at 48 hpf or 72 hpf, indicating a lack of generalized cell death in mutant animals despite the lethal phenotype (data not shown).

In contrast to the hematopoiesis defect and lethality of homozygous *crs* embryos, a single wild-type copy of *Hspa9b* was sufficient to sustain embryonic development. Yet MDS manifest primarily later in life, so to determine whether haploid insufficiency affected adult definitive hematopoiesis, we analyzed kidneys from animals at 1 year of age for histologic abnormalities and apoptosis. Whereas Wright-Giemsa staining of kidney smears showed blood cells at all stages of development in approximately similar proportions in heterozygous versus wild-type animals (Figure 6A-B), TUNEL analysis demonstrated rates of apoptosis in heterozygous kidneys 3 to 5 times those seen in kidneys from wild-type clutchmates (Figure 6C-D). By contrast, the number of TUNEL+ cells in smears from adult liver was similar regardless of genotype (data not shown). Thus *Hspa9b* haploid insufficiency specifically increased the susceptibility of developing blood cells to apoptosis.

**Loss of *Hspa9b* produces oxidative stress preceding blood cell apoptosis**

HSPA9B has been implicated in a number of cellular functions but most prominently as a chaperone for the transport, folding, and assembly of mitochondrial matrix proteins.38–40 We thus looked for signs of mitochondrial dysfunction in blood cells from *Hspa9b* mutant animals. Normal mitochondrial respiration generates ROS, but when mitochondrial function is disrupted, levels of ROS can overwhelm the antioxidant capacity of cells, triggering oxidative stress and apoptosis.41–43 To look for elevated ROS production, we incubated live animals with dihydorhodamine 123 for approximately 1 hour. Starting at 34 hpf, dihydorhodamine 123 specifically labeled blood cells in mutant embryos with maximal labeling.
Figure 5. Apoptosis of crs blood cells. (A-E) Live wild-type (A) and mutant (B-E) embryos incubated with acridine orange at 36 to 38 hpf. The area bracketed in ventral tail (B) is shown at higher magnification (C) with far right arrow indicating the same labeled blood cell in panels B and C. Areas over the yolk sac (B) and in the ventral tail (C) are shown at higher magnification in panels D and E, respectively. Arrowheads indicate the same labeled blood cells in panels C and E. Fluorescent images overlay bright-field images (D-E) and labeled cells are surrounded by unlabeled blood cells over the yolk sac (D) and in the tail vessels; CA indicates caudal artery and CV caudal vein (E). (F) Whole-mount immunofluorescence on 36 hpf mutant embryos with an antibody against activated caspase-3 overlies a bright-field image in the ventral tail. The yolk extension (yolk ext) and anus are marked for orientation. (G-H) Whole-mount in situ hybridization for heme oxygenase-1 (ho-1) at 36 to 38 hpf in wild-type (C) and mutant (D) embryos. A single cell in the tail vessels of the wild-type embryos is marked by an arrow in panel C. The region bracketed in panel D is shown at higher magnification (inset) with a ho-1+ cell (arrow) next to an unlabeled cell (arrowhead) in the caudal artery (CA).

Figure 6. Increased blood apoptosis in adult crs heterozygotes. Kidney smears from wild-type (A,C) and heterozygous (B,D) adult crs animals showing hematopoietic progenitors and mature blood cells by Wright-Giemsa staining (A-B) or TUNEL (C-D) with nuclei shown by DAPI (4,6 diamidino-2-phenylindole; blue) and TUNEL (green) labeling dying cells.

Discussion

The zebrafish mutant crs displays a distinct hematopoietic defect including anemia, erythroid dysplasia, apoptosis of blood cells, and multilineage cytopenia. This phenotype closely resembles the characteristic ineffective hematopoiesis of the human blood disorders of MDS. Positional cloning, rescue, and gene knockdown analysis linked the crs phenotype to a point mutation in Hspa9b, a candidate causative gene in a subset of MDS patients, and that encodes a ubiquitous mitochondrial matrix chaperone. Thus we provide strong evidence that Hspa9b plays an essential role in blood cell development and independently implicate loss of HSPA9B and mitochondrial dysfunction in MDS pathogenesis.

HSPA9B is a member of the heat shock protein 70 family of chaperones. These chaperones are composed of 3 domains including a highly conserved C-terminal substrate-binding domain. The crystallized substrate-binding domain of E coli DnaK revealed a β-sandwich containing the peptide-binding channel followed by 5 α helices. Mutation of glycine 443 to aspartic acid was predicted to disrupt this structure and produced a complete loss of substrate binding, rendering the chaperone functionless. The presence of the homologous G492E substitution in crs mutants strongly suggests that crs represents a null allele of Hspa9b.

HSPA9B is involved in the transport, folding, and assembly of proteins in the mitochondrial matrix including those involved in oxidative phosphorylation, metabolite biosynthesis, and protection from ROS. HSPA9B also negatively regulates proteins involved in mitochondrial cell death including p53 and p66Shc, and its overexpression can immortalize cells and increase the lifespan in Caenorhabditis elegans. Conversely, in yeast, hspa9b (ssc1) loss-of-function produces abnormal mitochondrial morphology, protein aggregation, and death. Mitochondrial function in Hspa9b mutant zebrafish appeared grossly normal as dyes that accumulate in functional mitochondria labeled mutant cells (S.E.C., unpublished results, June 2004). Rather, we find that Hspa9b mutant blood cells increased ROS production and induced expression of ho-1 starting at 34 hpf. This delayed onset is common to
mitochondrial diseases, and maternal contribution of wild-type HSPA9B may contribute to the delay. Oxidative stress was almost entirely restricted to blood cells and was accompanied by programmed cell death as indicated by labeling with AO and immunoreactivity for activated caspase-3. Furthermore, neither oxidative stress nor apoptosis became more generalized at 48 hpf, when development arrested, or at death around 72 hpf. Taken together, our results suggest that mitochondrial dysfunction in the absence of Hspa9b produces a mechanistically distinct defect in hematopoiesis that precedes general developmental arrest and thus provides insight into a human somatic blood cell disorder.

Many previously characterized zebrafish blood-specific mutants have defects restricted to the erythroid lineage. And although anemia and erythroid dysplasia alone describe 2 of the 8 World Health Organization categories for MDS, the remaining disease classifications are characterized by bicytopenias or pancytopenias. Here we demonstrate that mutant Hspa9b embryos have a reduction not only in red blood cells but also in mpo +3 myeloid cells. Thus, we link the loss of Hspa9b not just with anemia but also with multilineage cytopenia. In addition, we find that c-myb hematopoietic progenitors were lost, suggesting that stem cell populations were affected by apoptosis as well. Definitive progenitors are observed in the posterior ICM and in the dorsal aorta as early as 24 hpf, but not just with anemia but also with multilineage cytopenia. In addition, we find that c-myb hematopoietic progenitors were lost, suggesting that stem cell populations were affected by apoptosis as well. Definitive progenitors are observed in the posterior ICM and in the dorsal aorta as early as 24 hpf, but not just with anemia but also with multilineage cytopenia. However, 1-year-old heterozygous embryonic phenotype; however, haploid reduction not only in red blood cells but also in populations were affected by apoptosis as well. Definitive progenitors did not manifest obvious anemia or other characteristic symptoms of MDS, suggesting that further aging or alternatively loss of heterozygosity or other genetic lesions may be required.

Conversion of normal hematopoietic stem cells to an MDS state and ultimately a leukemic state is believed to be a multistep process requiring the accumulation of multiple genetic lesions. Initial damage to the DNA of hematopoietic stem cells can arise de novo or as a result of exposure to toxic agents, including chemotherapeutic drugs or radiation. This primary event may precede the induction of well-recognized MDS-associated cytogenetic abnormalities and may act to increase both stem cell proliferation, contributing to clonal hematopoiesis, and susceptibility of the cell to further injury. Clonal selection may be then be assisted by alterations in the bone marrow microenvironment or immune dysregulation. Subsequently, secondary genetic events such as a deletion at chromosome 5q and loss of HSPA9B could provide one route toward clonal evolution and the development of characteristic symptoms of MDS. And finally, further DNA damage may eventually lead to disruptions in the apoptotic response and leukemic transformation. Deletions at 5q31 often result in rapid progression to leukemia. Our finding that loss of functional HSPA9B results in high levels of ROS that can directly damage DNA and lead to further genetic lesions may help explain this poor prognosis.

Mutations in mitochondrial proteins and the mitochondrial genome are the primary defects in a number of diseases affecting organs with high energy demands including the nervous system and musculature. Blood cells are less often affected; however, large deletions of mitochondrial DNA are associated with a rare congenital disorder, Pearson syndrome, which results in pediatric pancytopenia. Also, transplantation of Sod2-deficient hematopoietic stem cells into wild-type mice resulted in a blood defect resembling sideroblastic anemia with oxidative damage and reduced survival of erythroid cells. Our analysis of the ccs zebrafish mutant indicates that a primary defect in a mitochondrial chaperone is sufficient to produce characteristic symptoms of the MDS blood disorders and implicates mitochondrial dysfunction in the pathogenesis of these syndromes.

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