Mitochondrial heme: an exit strategy at last

Mark D. Fleming¹ and Iqbal Hamza²

¹Department of Pathology, Boston Children’s Hospital, Harvard Medical School, Boston, Massachusetts, USA. ²Department of Animal and Avian Sciences, Department of Cell Biology and Molecular Genetics, University of Maryland, College Park, Maryland, USA.

The transport of heme across membranes is critical for iron absorption, the formation of hemoglobin and other hemoproteins, and iron recycling in macrophages. However, the identity of heme transport proteins has been elusive. In this issue of the JCI, Chiabrando et al. reveal that an isoform of the feline leukemia virus subgroup C receptor (FLVCR1) exports heme from the mitochondria and is critical for erythroid differentiation.

Nearly two-thirds of the body’s iron endowment is in the form of hemoglobin in erythrocytes, and each erythrocyte contains more than a billion iron atoms in the form of heme (1). Consequently, it is not surprising that inherited or acquired defects in hemoglobin synthesis, including the thalassemias, hemoglobinopathies, and iron deficiency, are among the most prevalent human diseases. Importantly, the approximately 360 billion erythrocytes produced daily require over 250 mg of heme to assemble into hemoglobin. Heme is synthesized in the mitochondria, but globin is translated in the cytosol, and it is unclear how newly synthesized heme is transported out of the mitochondria for incorporation into hemoglobin (2). Heme transport across membranes is important for dietary iron absorption and crucial for erythrocyte heme iron recycling in the reticuloendothelial system (RES) macrophage. Despite the physiologic importance of these processes, the molecular pathways of transmembrane heme transport have, for the most part, remained obscure, in large part due to technical difficulties in identifying heme-specific transporters in mammalian cells and the inability to translate these findings to whole organisms. In this issue of the JCI, using a combination of siRNA studies and targeted mutations in mice, Chiabrando et al. provide compelling evidence that an isoform of the feline leukemia virus subgroup C receptor (FLVCR1) exports heme from the mitochondria (3).

The requirement for heme transport

For more than 50 years, it has been known that nutritional heme iron is absorbed in the intestine by an active, energy-dependent, and inducible process that requires a heme transporter in enterocytes. This is because elemental iron has limited bioavailability in the intestine due to the presence of natural iron chelators, such as phytates and tannins, as well as its tendency to oxidize (i.e., to rust) and precipitate. In contrast, even though heme-iron constitutes only one-third of total dietary iron, it is more easily absorbed and is the source for two-thirds of body iron in meat-eating individuals (4). This is a consequence of heme’s solubility at intestinal pH and because its uptake is not known to be influenced by other dietary factors.

Heme must also be transported across mitochondrial membranes because the final steps of heme synthesis occur in the mitochondria, but some hemoproteins such as hemoglobin are cytosolic (2). Likewise, heme transport out of phagolysosomes is an essential component of iron recycling by macrophages, as heme oxygenase (Hmox), the enzyme that catalyzes the oxidation of heme to biliverdin, carbon monoxide, and ferric iron, is found largely tethered to the cytosolic surface of the endoplasmic reticulum (5). Iron recycling out of the macrophage is mediated by the ferrous iron exporter ferroportin (FPN1) (6), whose cell-surface expression is tightly controlled by the systemic iron regulatory hormone hepcidin (7). In this manner, iron catalyzed by RES macrophages from heme can be released immediately to the plasma to replenish the Fe2-Tf pool necessary for erythropoiesis or stored as ferritin for subsequent use.

Mammalian heme transporters

The identification and characterization of heme and other porphyrin transporters in mammals has proven to be difficult (2), in part due to a lack of genetic and molecular tools, but also as a consequence of the promiscuity of proteins capable of transporting heme with low affinity. Furthermore, in vivo investigation of transporter proteins identified in vitro or on the basis of their expression pattern has been misleading or has provided ambiguous results. For example, Hep1, a putative apical intestinal heme importer, was eventually proven to be essential for folate transport in the intestine when it was found to be mutated in patients with congenital folate deficiency (8, 9). Whether Hep1 also contributes to the absorption of heme in the intestine is uncertain at this time. Some evidence suggests that the breast cancer

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erythroid progenitors in the bone marrow. That the virus infected all bone marrow cells, but produced an erythroid-specific phenotype, was a clue to the function of the protein. Endogenous production of the viral capsid protein interferes with the cellular expression of FLVCR1, effectively resulting in a knockdown of the protein (17). The singular toxicity to erythroid precursors suggested that FLVCR1 performed a function uniquely essential to bone marrow cells of this lineage. Abkowitz and colleagues eventually demonstrated in vitro that FLVCR1 is a heme exporter and that targeted deletion of FLVCR1 in mice results in a severe macrocytic anemia characterized by a proerythroblast maturation arrest (18). Based upon the cell-surface localization of FLVCR1, these authors hypothesized that the phenotype is a consequence of the requirement of early erythroid precursors to export heme to avoid heme toxicity in the absence of active globin synthesis. While the murine knockout data were entirely supportive of the feline viral infection phenotype, the pathogenesis of the erythroid dysfunction remained an enigma.

A mitochondrial FLVCR1 isoform

In this issue, Emanuela Tolosano’s group describes the identification of an mRNA transcript in EST databases encoding an isoform of FLVCR1 termed FLVCR1b (3). This isoform contains a shortened N terminus as a result of an alternative transcriptional start site in intron 2 of the previously described FLVCR1a mRNA. The protein encoded by the FLVCR1b transcript includes amino acids 277–555 of FLVCR1a and encodes a mitochondrial targeting signal at the new N terminus. Targeted deletion of the exon specific to FLVCR1a resulted in skeletal defects and vascular abnormalities, but not anemia, indicating that loss of the cell-surface FLVCR1a is dispensable for erythropoiesis. In contrast, siRNA-mediated knockdown of the FLVCR1b isoform impaired erythroid differentiation in vitro. While the authors do not directly demonstrate that FLVCR1b transports heme out of mitochondria, they do show that overexpression of the protein increases cytosolic heme and that knockdown of FLVCR1b results in mitochondrial heme retention (3). Taken together, these data suggest that the FLVCR1b-null phenotype may be due to cytosolic heme deficiency rather than heme accumulation, as had been originally proposed (Figure 1).

FLVCR1 and human disease

The association of pure red cell aplasia and skeletal abnormalities has suggested that mutations in FLVCR1 might be found in human patients with the inherited form of pure red cell aplasia termed Diamond-Blackfan anemia (DBA), which is also associated with skeletal abnormalities. FLVCR1 mutations in DBA patients have not been found as of yet (19); however, if they were to exist, one would predict that they would occur in the FLVCR1b isoform. Interestingly, missense mutations in FLVCR1 have recently been described in patients with the rare autosomal recessive disease posterior column ataxia and retinitis pigmentosa (PCARP) (20, 21). Three of the four reported disease-associated FLVCR1 variants, all of which are present in the homozygous state in PCARP patients, occur in exon 1 of the gene—the part of the gene that is omitted in the FLVCR1b transcript and protein. Based on this finding, one might speculate that heme export by the cell-surface FLVCR1a may have a uniquely important function in neuronal cells. It is equally possible, as has been observed with many other proteins capable of transporting heme, that FLVCR1a binds other substrates with higher affinity than heme. It is, however, more problematic to reconcile the fourth patient mutation, which occurs in a transmembrane domain common to the FLVCR1a and FLVCR1b isoforms. Individuals harboring this mutation are not anemic, as the murine model would predict (20), raising the possibility that this genetic lesion has a unique effect on transport of an alternative substrate. If FLVCR1b localizes to the inner membrane of the mitochondria, does heme exit the mitochondrial outer membrane through unconventional means? Furthermore, how do eukaryotes that lack obvious FLVCR homologs, such as yeast, export heme from the mitochondria? Pertinent biochemical questions also remain: FLVCR1a is proposed to transport heme out of the cytoplasm, while FLVCR1b transports heme into the cytoplasm. Because FLVCR1 is an MFS family member of secondary transporters, are the sources of energy to translocate heme and the kinetic mechanisms (uniporter, symporter or antiporter) different between the two isoforms? Only time will reveal the truth in the myriad of theoretical possibilities.

Address correspondence to: Mark D. Fleming, Department of Pathology, Boston Children’s Hospital, S. Burt Wolbach
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The glomeruli of the human kidney filter almost 200 liters of fluid every day, but must retain the albumin and other large proteins within the circulation. The fluid has to cross three layers: the endothelium, the glomerular basement membrane (GBM), and the epithelial cells or podocytes. The GBM (1) and the slit diaphragm (2), a structure that joins adjacent podocytes, are two barriers to protein filtration, but the endothelial cell also likely plays a role. Disruption of these barriers results in proteinuria, a consequence of many kidney diseases.

Podocytes begin life as tall epithelial cells that have typical tight junctions near the apex separating the apical surface from the lateral cell membranes. As development progresses, these junctions migrate downward toward the basal surface of the mature podocytes, becoming the slit diaphragm (3). Unlike other polarized epithelia, the lateral surface is above these specialized tight junctions and is continuous with and identical to the apical surface. The basal surface differentiates into thin interdigitating structures called foot processes that are joined together by the cell-cell junctions of the slit diaphragm. During proteinuric disease, the foot processes lose their fine structure and collapse into a mass of cytoplasm on top of the basement membrane (4). The observation that a great abundance of actin and actin-binding proteins accumulates in these diseased foot processes suggested that actin dysregulation might play a key role in the pathogenic progression (5). The discovery that α-actinin-4 mutations were also associated with heavy proteinuria also helped bring attention to the role of actin (6).

Since then, mutations in humans or animals in other actin-binding proteins—including Nck, CD2AP, and dynamin—were also found to mediate proteinuria (7). Further, the cytoplasmic domains of many slit diaphragm components can bind actin or signal to actin, further suggesting a key role for this cytoskeletal protein (8). But what was the actual cellular biological process in which actin was so central? Was the primary function of actin to maintain the slit diaphragm, or was the actin rearrangement merely a consequence of the permeability change? Here, one might turn to basic cell biology and physiology to find the definitive answers. Unfortunately, the podocyte, when grown in culture, loses its complex shape; it has neither foot processes nor a real slit diaphragm. Given that actin is directly involved in cell shape determination and maintenance, this has placed the field in a difficult position. However, an article in this issue by Soda et al. has provided the first insight into the process of maintenance of the slit diaphragm (9). The authors show that clathrin-mediated endocytosis is required for podocyte function and is likely the effector of the changes in actin that are so often observed.

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