

Of mice and men: the iron age

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Commentary

Recently, mutations causing juvenile hemochromatosis have been identified in a novel gene, *hemojuvelin* (*HJV*), located on chromosome 1. Mouse models of this disease have now been developed by 2 groups, Huang et al. and Niederkofler et al., through targeted disruption of the *Hjv* gene (see the related articles beginning on pages 2180 and 2187). These mutant mice will allow further investigation into the role of HJV in the regulation of iron homeostasis, a role that to date remains elusive.

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destination where newly generated peptide pools are made available for binding to them. Absent from the discussion is the obvious consideration of protease specificity: if lysosomal proteases that generate the class II MHC ligands were to display preferred cleavage sites, then the pools of peptides made available to class II molecules would be biased. However, this has never been a popular notion in view of the redundancy, in terms of substrate specificity, of lysosomal proteases. There are examples to the contrary, however: asparagine-specific endoprotease is indeed remarkably selective in the peptide bonds it cleaves, and the enzyme has been implicated in the generation, in a highly selective fashion, of class II MHC-restricted epitopes (4). Nonetheless, there are very few examples of antigens for which the processing pathway

has been characterized comprehensively in terms of intermediates and the proteases that generate them. It may well be that the convergence in peptide binding, observed for DQ8 and I-A*^{g7}, produced by the identical types of antigen-presenting cells, owes at least a component of this repertoire to shared enzymatic machinery upstream. As Suri et al. state (1), expression levels of the source antigen and the possibilities of encounter with the entire processing and peptide loading apparatus are certainly going to affect the final outcome, in terms of the MHC-peptide complexes produced. The best map of a landscape is the landscape itself: seen from this perspective, the comprehensive data set presented here by Suri and coworkers will allow others to navigate the territory of autoimmunity with an accurate map in hand.

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Of mice and men: the iron age

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Recently, mutations causing juvenile hemochromatosis have been identified in a novel gene, *hemojuvelin* (*HJV*), located on chromosome 1. Mouse models of this disease have now been developed by 2 groups, Huang et al. and Niederkofler et al., through targeted disruption of the *Hju* gene (see the related articles beginning on pages 2180 and 2187). These mutant mice will allow further investigation into the role of *HJV* in the regulation of iron homeostasis, a role that to date remains elusive.

The following scenario is classic with respect to genetic disorders: the disease is clinically recognized, the related gene is identified, and a couple of years later, a mouse model of the genetic disease is established. The genes involved in iron metabolism have not been an exception to this rule (in particular those related to hereditary hemochromatosis, a prevalent heterogeneous iron-overload disorder), and the development of mouse models have greatly facilitated our understanding of the pathogenesis of iron overload. The *hemojuvelin* gene (*HJV*, also known as *HFE2* and *RGMc*), which is mutated in a

form of juvenile hemochromatosis linked to chromosome 1, was identified in 2003 (1). Researchers in the field have therefore been waiting for the development of a relevant mouse model of this disease. This feat has now been accomplished by Huang et al. and Niederkofler et al., who in this issue of the *JCI* describe the phenotype of their mutant mice (2, 3). Their findings add further support to the emerging attractive hypothesis that hepcidin, a recently described iron-regulatory peptide, and ferroportin, its target, are common pathogenic denominators in all forms of iron overload.

In mammals, iron homeostasis is complex and depends on regulated dietary iron absorption by mature enterocytes of the duodenum and iron recycling by macrophages, which supply most of the serum iron through recovery of the metal from senescent erythrocytes (4). These 2 fundamental processes are deregulated in

hereditary hemochromatosis; over time, this deregulation leads to iron deposition in parenchymal cells and transferrin saturation (5). As the disease progresses, patients develop iron-induced tissue damage, which results in serious illnesses including cirrhosis, hepatomas, diabetes, cardiomyopathy, arthritis, and endocrinopathies. Unless recognized and treated, this disease is fatal.

While hereditary hemochromatosis has been clinically recognized since 1889, it is only over the 10 past years that the genetic basis for the disease has been elucidated. Hereditary hemochromatosis is now recognized to be genetically heterogeneous and its nomenclature is under debate. Depending on distinct clinical and genetic entities, hereditary hemochromatosis can be divided into 3 classes: classical hemochromatosis, juvenile hemochromatosis, and ferroportin disease.

Classical hemochromatosis is associated with the historical and most prevalent form of hereditary hemochromatosis and is almost always caused by mutations in *HFE*, a gene which encodes a protein of the major histocompatibility complex class I (6). Rarely, another gene can be involved, namely *transferrin receptor 2* (*TFR2*) (7), which encodes a homolog of *TFR1* that mediates

Nonstandard abbreviations used: HAMP, hepcidin antimicrobial peptide; *HJV*, *hemojuvelin*; *TFR2*, transferrin receptor 2.

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stream target of hepcidin, hepcidin level is not expected to be decreased in this type of hemochromatosis (29).

Juvenile hemochromatosis in *Hju*^{-/-} mice results from impaired hepcidin synthesis and consequent overexpression of ferroportin

In this issue of the *JCI*, Huang et al. (2) and Niederkofler et al. (3) report the establishment of murine models of HJV deficiency. Although the data presented do not provide any new information on the functional role of HJV, they effectively describe the increased iron deposition in the liver, heart, and pancreas and the decreased iron levels in tissue macrophages of the mutant mouse. The authors note that liver iron accumulation begins earlier and progresses more quickly in *Hju*^{-/-} mice than in *Hfe*^{-/-} mice. The heterozygous *Hju*^{+/-} mice are indistinguishable from control mice (2). These results definitively confirm that loss-of-function mutations in the *Hju* gene lead to severe hemochromatosis and are consistent with the fact that parents of juvenile hemochromatosis patients are unaffected. Interestingly, both groups report that *HAMP* mRNA is virtually undetectable in the livers of mutant mice, which suggests that HJV acts by depressing *HAMP* gene activity. It is likely that HJV, like HFE, is needed as a sensor or mediator of activating signals of *HAMP* expression, including iron. In fact, the hepcidin gene's decreased sensitivity to iron has been reported in *Hfe*^{-/-} mice (24). Niederkofler et al. show that the *Hju*^{-/-} mice fail to express *HAMP* in response to dietary or injected iron, which demonstrates that the transmission of the iron signal in the liver is completely blocked in the absence of HJV. However, the authors report that *Hju*^{-/-} mice retain their ability to upregulate *HAMP* expression in response to inflammation and emphasize that the 2 inducers of *HAMP* expression, iron and inflammation, act through different pathways, the latter one independent of HJV.

Having established the impairment of hepcidin synthesis in mutant mice, Huang et al. (2) further show that the ferroportin level is considerably enhanced in both intestinal epithelial cells and macrophages of *Hju*^{-/-} mice, most likely reflecting the profound hepcidin deficiency. Accordingly, ferroportin has been shown to be overexpressed in the intestine and macro-

phages of hepcidin-deficient mice (30). It should be noted that in addition to hepcidin deficiency-induced ferroportin hyperactivity, Drakesmith et al. recently reported that ferroportin mutations resulting in insensitivity to hepcidin could also lead to ferroportin hyperactivity and subsequent hemochromatosis (31). Together, these results confirm and extend the key role of the hepcidin/ferroportin axis in regulating iron absorption by the intestine and iron efflux from macrophages.

While HJV mouse models of juvenile hemochromatosis are elegantly described in these important studies (2, 3), some key questions are still awaiting answers: (a) What is the functional role of HJV, in particular its role — if any — in the skeletal and cardiac muscles that express *Hju* at a high level? Will older knockout animals develop specific muscular damage in addition to iron overload?; (b) Does hepcidin gene expression in organs other than the liver (e.g., in the heart, pancreas, and kidney) play any role? Does HJV regulate *HAMP* expression in a similar manner in all tissues?; and (c) How do iron-sensing and inflammatory pathways converge on *HAMP* expression, and at what level is the *HAMP*/iron-activating signal disturbed in the absence of HJV? Could *HAMP* be activated in patients and mice suffering from this type of juvenile hemochromatosis by drugs that, for instance, mimic inflammatory cues but are devoid of their detrimental effects?

There is no doubt that the discoveries of the hormonal iron-regulating role of hepcidin (16, 17) followed by the elucidation of its mechanism of action — to regulate ferroportin degradation in the intestine, macrophages, and probably placenta (18) — herald exciting times in the field of iron metabolism and related disorders. The hepcidin/ferroportin axis occupies a central position in regulating iron homeostasis and explaining the physiopathology of iron-overload syndromes and likely — at the other end of the spectrum of iron disorders — anemia of inflammation (23, 32). Although considerable efforts are still required in order to solve remaining questions, the studies by Huang et al. and Niederkofler et al. have paved the way for rapid progress in the development of inducers or drugs capable of enhancing endogenous *HAMP* expression in spite of HFE or HJV deficiency. This should lead to breakthrough treatments for hereditary hemochromatosis, secondary

hemosiderosis, and chronic inflammatory anemias (thanks to hepcidin antagonists or inhibitors of *HAMP* gene expression) — pathologies that affect millions of patients around the world.

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