The N-terminus of hepcidin is essential for its interaction with ferroportin: structure-function study

Elizabeta Nemeth, Gloria C. Preza, Chun-Ling Jung, Jerry Kaplan, Alan J. Waring, and Tomas Ganz

Hepcidin is the principal iron-regulatory hormone. It acts by binding to the iron exporter ferroportin, inducing its internalization and degradation, thereby blocking cellular iron efflux. The bioactive 25 amino acid (aa) peptide has a hairpin structure stabilized by 4 disulfide bonds. We synthesized a series of hepcidin derivatives and determined their bioactivity in a cell line expressing ferroportin-GFP fusion protein, by measuring the degradation of ferroportin-GFP and the accumulation of ferritin after peptide treatment. Bioactivity was also assayed in mice by the induction of hypoferremia. Serial deletion of N-terminal amino acids caused progressive decrease in activity which was completely lost when 5 N-terminal aa's were deleted. Synthetic 3-aa and 6-aa N-terminal peptides alone, however, did not internalize ferroportin and did not interfere with ferroportin internalization by native hepcidin. Deletion of 2 C-terminal aa's did not affect peptide activity. Removal of individual disulfide bonds by pairwise substitution of cysteines with alanines also did not affect peptide activity in vitro. However, these peptides were less active in vivo, likely because of their decreased stability in circulation. G71D and K83R, substitutions previously described in humans, did not affect hepcidin activity. Apart from the essential nature of the N-terminus, hepcidin structure appears permissive for mutations. (Blood. 2006;107:328-333)

Introduction

Hepcidin, a peptide hormone produced by the liver, is the principal regulator of iron homeostasis. Hepcidin production increases with iron loading and inflammation and decreases with anemia and hypoxia.1,2 Hepcidin acts by binding to ferroportin, the sole cellular iron exporter, and inducing its internalization and degradation.3 Because ferroportin is expressed in the duodenum, spleen, liver, and placenta,4 hepcidin increase, and the subsequent decrease of ferroportin, results in inhibition of duodenal iron absorption, release of recycled iron from macrophages, mobilization of iron stores from the liver, and iron transfer across placenta. Excessive hepcidin production resulting from increased expression in liver tumors5 or transgene expression,6 or acute administration of synthetic hepcidin,7 leads to either acute or chronic hypoferremia. Alternatively, hepcidin deficiency resulting from mutations in the hepcidin gene itself, or the genes encoding HFE, transferrin receptor 2, or hemojuvelin,8-11 appears to be the cause of most types of hereditary hemochromatosis.

The hepcidin gene encodes an 84-aa prepropeptide with a typical signal sequence and a consensus cleavage site for a prohormone convertase that generates the mature bioactive 25-aa form found in plasma and urine.12,13 The 25-aa peptide (hepc25) forms a hairpin loop stabilized by 4 disulfide bonds, one of which is an unusual vicinal bond between adjacent cysteines at the hairpin turn.14 Three-dimensional nuclear magnetic resonance (3D-NMR) structure studies also showed that hepcidin is an amphipathic peptide similar to most antimicrobial peptides. However, unlike antimicrobial peptides that display great species-to-species variation, hepcidin sequence is conserved across species, ranging from fish to mammals, with human sequence showing 60% similarity to the zebra fish hepcidin-1.15

Here, we report on an investigation of hepcidin structure in relation to its function and show that the N-terminus of hepcidin is critical for its iron-regulatory activity.

Materials and methods

Peptide synthesis

The peptides were synthesized on an ABI 431A peptide synthesizer (PE Biosystems, Applied Biosystems, Foster City, CA) using fmoc amino acids, Wang resin (AnaSpec, San Jose, CA), and double coupling for all residues. After cleavage, 30 mg crude peptides was reduced with 1000-fold molar excess of dithiothreitol (DTT) in 0.5 M Tris buffer (pH 8.2), 6 M guanidine hydrochloride, and 20 mM EDTA at 52°C for 2 hours. Fresh DTT...
(500-molar excess) was added and incubated for an additional hour at 52°C. The reduced peptides were purified on the 10-g C18 Sep-Pak cartridges (Waters, Milford, MA) equilibrated in 0.1% TFA and eluted with 50% acetonitrile. The eluates were lyophilized and resuspended in 0.1% acetic acid. The reduced peptides were further purified by reversed-phase high-performance liquid chromatography (RP-HPLC) on Vydar C18 column (218TP910; Waters) equilibrated with 0.1% trifluoroacetic acid and eluted with an acetonitrile gradient. The eluates were lyophilized, dissolved in 0.1% acetic acid, 20% DMSO, to the approximate concentration of 0.1 mg/mL (pH 8), and air oxidized by stirring for 18 hours at room temperature. The refolded peptides were also purified sequentially on the 10-g C18 Sep-Pak cartridge and on the RP-HPLC Vydar C18 column using an acetonitrile gradient. The eluates were lyophilized and resuspended in 0.016% HCl. The conformation of refolded synthetic hepcidin derivatives was verified by electrophoresis in 12.5% acid-urea polyacrylamide gel electrophoresis (PAGE), and peptide masses were determined by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF-MS; UCLA Mass Spectrometry Facility, Los Angeles, CA). Of note, the C11A/C19A peptide was modified at the C-terminus by tert-butyl because of incomplete removal of the protective group following the crude peptide synthesis. All of the other peptides had masses that matched predictions.

Circular dichroism spectroscopy

Circular dichroism spectra (185-260 nm) of the hepcidin peptides were recorded on an AVIV 62DS spectropolarimeter (AVIV Associates, Lake- wood, NJ). The instrument was calibrated for wavelength and optical rotation using d-10-camphorsulphonic acid.16 Samples were scanned at a temperature of 25°C using 0.01-cm path length cells at a rate of 10 nm per minute and a sample interval of 0.2 nm. Sample concentration was determined by quantitative amino acid analysis (UC Davis Core Facility, Davis, CA), and the sample concentration of the peptides was 200 μM in a solution containing 25 mM HEPES buffer, pH 7.5 with 150 mM NaCl. Before analysis, spectra were baseline corrected by subtracting spectra of peptide-free buffer solution from the peptide-containing sample and expressed as the Mean Residue Ellipticity $\theta_{BRE}$. Quantitative estimates of the secondary structural contributions were made with SELCON 317 using spectral basis set 8 for soluble proteins implemented in the Olis Global Works software package (Olis, Bogart, GA).

Flow cytometry

ECR293-Fpn, a cell line stably transfected with the mouse ferroportin-GFP construct under the control of ponasterone-inducible promoter, was prepared and maintained as previously described.3 The cells were plated on poly-D-lysine–coated plates in the presence of 20 μM FAC, with or without 10 μM ponasterone. After 24 hours, ponasterone was washed off, and cells were treated with peptides for 24 hours. Cells were then trypsinized and resuspended at $1 \times 10^6$ cells/mL, and the intensity of green fluorescence was analyzed by flow cytometry at the UCLA Jonsson Comprehensive Cancer Center and Center for AIDS Research Flow Cytometry Core Facility (supported by NIH awards CA-16 042 and AI-28 697, Jonsson Cancer Center, the UCLA AIDS Institute, and the UCLA School of Medicine). Flow cytometry was performed on FACSscan (fluorescence activated cell scanner) Analytic Flow Cytometer (Becton Dickinson, San Jose, CA) with CellQuest version 3.3 software (Becton Dickinson). Cells not induced with ponasterone to express Fpn-GFP were used to establish a control under the same conditions. The effect of each peptide was assessed by comparing the intensity of green fluorescence in the presence of the peptide to the control. The relative intensity of green fluorescence was calculated as $I = (F_{\text{uninduced}} - F_{\text{ponasterone}}) / (F_{\text{uninduced}} - F_{\text{naposta}})$, where F was the mean of the gated green fluorescence and x was the peptide.

Ferritin measurements

ECR293-Fpn cells were incubated with 20 μM FAC with or without 10 μM ponasterone. After 24 hours, ponasterone was washed off, and hepcidin derivatives were added for 24 hours in the presence of 20 μM FAC. Cellular protein was extracted with 150 mM NaCl, 10 mM EDTA, 10 mM Tris (pH 7.4), 1% Triton X-100, and a protease inhibitor cocktail (Sigma-Aldrich, St Louis, MO). Ferritin levels were determined by an enzyme-linked immunosorbent assay (ELISA) assay (Ramco Laboratories, Stafford, TX, or Biotech Diagnostic, Laguna Nigel, CA) according to the manufacturer’s instructions and were normalized for the total protein concentration in each sample, as determined by the bicinchoninic acid (BCA) assay (Pierce, Rockford, IL). The results were expressed as the relative change in ferritin, according to the formula $(F_{\text{Funtreated}} - F_{\text{hep25}})/(F_{\text{Funtreated}} - F_{\text{hep25}},$ where F was the ferritin concentration (ng/mg protein) and x was the peptide.

Mouse serum iron measurements

Animal studies were approved by the Animal Research Committee at UCLA. C57BL/6 mice were obtained from The Jackson Laboratory (Bar Harbor, ME) and were maintained on NIH 31 rodent diet (iron content 336 mg/kg; Harlan Teklad, Indianapolis, IA). Two weeks before the experiment, the mice were switched to an iron-deficient diet containing 2 to 4 ppm iron (Harlan Teklad) to suppress endogenous hepcidin. Mice were injected intraperitoneally either with 100 μL PBS (control) or with 50 μg peptide in 100 μL PBS. Mice were killed 2 hours later, blood was collected by cardiac puncture, and serum was separated using Microtainer tubes (Becton Dickinson, Franklin Lakes, NJ). Serum iron was determined by using a colorimetric assay (Diagnostic Chemicals, Oxford, CT), which was modified for the microplate format18 so that 50 μL serum was used per measurement. The results were expressed as the percentage of decrease in serum iron when compared with the average value of serum iron levels in PBS-injected mice.

Statistics

We used Sigma Stat version 3.0 for statistical analyses (Systat Software, Point Richmond, CA).

Results

We synthesized a series of hepcidin peptides with alterations targeting the N-terminus, C-terminus, and the disulfide bonding pattern, or with substitutions found in patients with iron overload (Figure 1). After the peptides were folded and purified, their structure was analyzed by circular dichroism (CD) spectroscopy. All spectra were typical of disulfide-stabilized β sheet peptides connected by turn-loop segments.13,19,20 Analysis of the CD spectra indicated that the general conformational characteristics were preserved in all the structures (Table 1). Even CD spectra of disulfide-modified hepcidins (Figure 2), including the shortened peptide lacking 3 of the 4 disulfide bonds, indicated little effect on conformation.

We next analyzed the effect of structural alterations on the peptide bioactivity. We previously used a cell line (ECR293) stably transfected with the mouse ferroportin-GFP fusion protein1 to show that hepcidin causes internalization and degradation of iron export channel ferroportin. In this study, we used the same ECR293 in vitro model but accurately measured the degradation of ferroportin-GFP by flow cytometry after treatment with different peptides (Supplemental Figure S1, available at the Blood website; see the Supplemental Figure link at the top of the online article). Table 2 shows a summary of the peptide activities and their statistical analysis. The serial deletion of the N-terminal amino acids caused a progressive loss of activity, with almost complete loss when all 5 residues were deleted (del1-5(DTHFP), also known as hep20) (Figure 3A). To determine whether the N-terminus by itself could serve as hepcidin agonist or antagonist, we synthesized the 3-amino acid (DTH) and 6-amino acid (DTHFP) N-terminal peptides. The
peptides did not internalize ferroportin and, even at 20-fold higher concentrations, did not interfere with ferroportin degradation induced by hep25 (Table 2). The N-truncated inactive peptide Hep20 (5 μM) also did not interfere with the full-length peptide (1 μM) when the 2 were simultaneously added to the cells (Table 2).

The peptide extended at the N-terminus by the addition of Ala (hep26(+A)) had activity equivalent to hep25 (Figure 3A). When amino acid residues from the C-terminus were deleted (del24-25(KT)), the peptide activity was only slightly affected (90% activity; Figure 3A).

We also altered the disulfide-bonding pattern by replacing pairs of cysteines with alanines so that C7A/C23A lacked the vicinal bond (bond 4). The 3 peptides exhibited nearly full activity (85% ± 3%, 92% ± 6%, 92% ± 12%, mean ± SD, respectively; Figure 3A). CD spectroscopy indicated that these substitutions did not significantly change the molecule’s hairpin structure (Figure 2). The peptide with a significantly shortened hairpin and lacking 3 of 4 disulfide bonds (del19-22(FCCG), del19-22(CGMC), C13A/C14A; abbreviated as 1SS) still showed 54% ± 5% activity.

**Table 1. Secondary structures of hepcidin variants**

<table>
<thead>
<tr>
<th>Sample</th>
<th>β sheet, %</th>
<th>Turn, %</th>
<th>Disordered, %</th>
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<tbody>
<tr>
<td>hep25</td>
<td>30.0</td>
<td>28.0</td>
<td>42.0</td>
</tr>
<tr>
<td>hep20</td>
<td>32.0</td>
<td>29.0</td>
<td>39.0</td>
</tr>
<tr>
<td>delKT</td>
<td>30.0</td>
<td>30.0</td>
<td>40.0</td>
</tr>
<tr>
<td>K83R</td>
<td>29.0</td>
<td>27.0</td>
<td>44.0</td>
</tr>
<tr>
<td>G71D</td>
<td>29.0</td>
<td>29.0</td>
<td>42.0</td>
</tr>
<tr>
<td>1SS</td>
<td>31.0</td>
<td>25.0</td>
<td>44.0</td>
</tr>
<tr>
<td>C7A/C23A</td>
<td>26.0</td>
<td>26.0</td>
<td>49.0</td>
</tr>
<tr>
<td>C11A/C19A</td>
<td>25.0</td>
<td>26.0</td>
<td>49.0</td>
</tr>
<tr>
<td>C13A/C14A</td>
<td>30.0</td>
<td>26.0</td>
<td>44.0</td>
</tr>
</tbody>
</table>

*The percentage of helix for all samples was 0.

Figure 2. Representative CD spectra of parent hep25 peptide and disulfide bond variants. Hep25 (solid lines) was compared to hepcidin C13A/C14A (A), hepcidin C7A/C23A (B), hepcidin C11A/C19A (C), and hepcidin 1SS (del9-12[FCCG], del19-22[CGMC], C13A/C14A; abbreviated as 1SS) (D).

Table 2. Summary of hepcidin derivative bioactivity

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Hepcidin-25 activity, %</th>
<th>P</th>
<th>Ferritin ELISA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hep24</td>
<td>95 ± 4</td>
<td>.018</td>
<td>77 ± 19</td>
</tr>
<tr>
<td>Hep23</td>
<td>88 ± 4</td>
<td>&lt;.001</td>
<td>47 ± 28</td>
</tr>
<tr>
<td>Hep22</td>
<td>46 ± 6</td>
<td>&lt;.001</td>
<td>13 ± 16</td>
</tr>
<tr>
<td>Hep21</td>
<td>22 ± 9</td>
<td>&lt;.001</td>
<td>3 ± 6</td>
</tr>
<tr>
<td>Hep20</td>
<td>12 ± 8</td>
<td>&lt;.001</td>
<td>&lt;2 ± 10</td>
</tr>
<tr>
<td>Hep3</td>
<td>6 ± 10</td>
<td>&lt;.001</td>
<td>5 ± 5</td>
</tr>
<tr>
<td>Hep6</td>
<td>1 ± 3</td>
<td>&lt;.001</td>
<td>—</td>
</tr>
<tr>
<td>Hep26</td>
<td>100 ± 1</td>
<td>NS</td>
<td>—</td>
</tr>
<tr>
<td>delKT</td>
<td>90 ± 7</td>
<td>.01</td>
<td>87 ± 18</td>
</tr>
<tr>
<td>C7A/C23A</td>
<td>85 ± 3</td>
<td>&lt;.001</td>
<td>107 ± 12</td>
</tr>
<tr>
<td>C11A/C19A</td>
<td>92 ± 6</td>
<td>.014</td>
<td>101 ± 9</td>
</tr>
<tr>
<td>C13A/C14A</td>
<td>92 ± 12</td>
<td>NS</td>
<td>84 ± 21</td>
</tr>
<tr>
<td>1SS</td>
<td>54 ± 5</td>
<td>&lt;.001</td>
<td>36 ± 29</td>
</tr>
<tr>
<td>G71D</td>
<td>108 ± 1</td>
<td>&lt;.001</td>
<td>91 ± 11</td>
</tr>
<tr>
<td>K83R</td>
<td>101 ± 3</td>
<td>NS</td>
<td>107 ± 17</td>
</tr>
<tr>
<td>zebrafish</td>
<td>106 ± 10</td>
<td>NS</td>
<td>103 ± 8</td>
</tr>
<tr>
<td>hep25 + 20; 5 μM</td>
<td>102 ± 2</td>
<td>.05</td>
<td>—</td>
</tr>
<tr>
<td>hep3; 20 μM</td>
<td>−10 ± 7</td>
<td>&lt;.001</td>
<td>—</td>
</tr>
<tr>
<td>hep25 + 3; 20 μM</td>
<td>102 ± 5</td>
<td>NS</td>
<td>—</td>
</tr>
<tr>
<td>hep6; 20 μM</td>
<td>6 ± 4</td>
<td>&lt;.001</td>
<td>—</td>
</tr>
<tr>
<td>hep25 + 6; 20 μM</td>
<td>96 ± 4</td>
<td>.047</td>
<td>—</td>
</tr>
</tbody>
</table>

The results, expressed as percentage of the hep25 activity, are the mean ± SD of 3 to 6 independent experiments. The values obtained by flow cytometry represent the ability of peptide to cause degradation of ferroportin-GFP, and the values obtained by ferritin ELISA represent the peptides’ ability to cause cellular iron retention. Each peptide was added at the concentration of 1 μM, unless indicated otherwise. Paired t test was used for statistical analysis for each peptide in comparison to hep25.

NS indicates not significant (> .05); —, not done.
The unmodified, naturally occurring active forms of hepcidin are shown as black bars. Del19-22(CGMC), C13A/C14A; zebra fish (87% C11A/C19A, and C13A/C14A) showed nearly full activity. N-terminus deletion resulted in the loss of activity, C-terminus results obtained from flow cytometry (Figure 3B; Table 2).

We also synthesized peptides with substitutions found in patients with iron overload disorders. Heterozygous G71D substitution was described in patients with HFE-related hemochromatosis,21,22 and heterozygous K83R substitution was found in a patient with porphyria cutanea tarda and iron overload (J.K. and John Phillips, University of Utah; unpublished observations, December 2004). G71D and K83R peptides, however, had activity similar to wild-type hep25 (Figure 3A).

Zebra fish hepcidin-1, which has only 52% identity with human hepcidin but preserves the disulfide structure and the general amphipathic character of the molecule, also caused ferroportin degradation comparable to human hep25.

Hepcidin-induced ferroportin internalization and degradation cause cellular iron retention.3 We measured accumulation of the iron storage protein ferritin after treatment with hepcidin derivatives as an additional assessment of peptide bioactivity. The activity of peptides as determined by ferritin ELISA paralleled the results obtained from flow cytometry (Figure 3B; Table 2). N-terminus deletion resulted in the loss of activity, C-terminus deletion (delKT), or disulfide bonds replacements (C7A/C23A, C11A/C19A, and C13A/C14A) showed nearly full activity (87% ± 18%, 107% ± 12%, 100% ± 9%, and 84% ± 21%, mean ± SD, respectively), while the short hairpin lacking 3 disulfide bonds (1SS) had 36% ± 29% activity when compared with hep25. The N-terminal 3-aa or 6-aa peptides failed to cause ferritin accumulation, whereas zebrafish hepcidin-1 was fully active (103% ± 8% of the human hep25 activity).

Injection of synthetic human hep25 causes rapid hypoferremia in mice.23 We therefore measured serum iron in mice 2 hours after the injection of 50 μg selected hepcidin derivatives (Figure 4). Hep25, G71D, and K83R peptides caused a similar and significant drop in serum iron compared with control mice injected with diluent. C13A/C14A peptide lacking the vicinal disulfide bond and delKT lacking the 2 C-terminal amino acids also significantly suppressed serum iron. Serum iron levels did not change significantly in mice injected with hep20, the short hairpin lacking 3 disulfide bonds (1SS), or the 2 peptides lacking disulfide bonds across the hairpin (C7A/C23A, C11A/C19A).

**Discussion**

Hepcidin regulates extracellular and plasma iron by controlling iron flows from the duodenum, macrophages, and liver.25 This is achieved by hepcidin binding to the cellular iron exporter ferroportin and causing its internalization and degradation. Bioactive hepcidin is a 25–amino acid peptide; a simple hairpin stabilized by 4 disulfide bonds, one of which is an unusual vicinal bond.13,25 Hepcidin structure is conserved across species from fish to mammals, but some species (mouse, zebra fish) contain an additional hepcidin gene of unknown functional significance. Zebra fish hepcidin-1 shows only 52% identity with human hepcidin, but all 8 cysteines are in the same position as in human hepcidin. The substitutions in the important N-terminal are conservative, as indicated by similarity matrices derived from analyses of amino acid substitutions in evolutionarily-related proteins. For example, in the BLOSUM80 matrix,27 the first 4 amino acids in the zebrafish sequence QSHL represent respectively the second, second, first, and third most conservative substitution for the amino acids of the human sequence DTHF. In our study, synthetic zebrafish hepcidin-1 was equivalent to human hepcidin in its ability to induce degradation of mouse ferroportin-GFP and cause cellular iron retention.

We next introduced a series of structural alterations into human hep25 to assess the functional importance of the N-terminal and C-terminal amino acids, as well as the disulfide-bonding pattern. The peptide bioactivity was assessed by quantitating ferroportin-GFP degradation and by measuring ferritin accumulation as an indicator of the inhibition of iron efflux. Some peptides were also tested for their ability to cause hypoferremia in mice. Surprisingly, hepcidin structure was quite permissive to changes. The only major effect on activity was seen when the 5 N-terminal amino acids were serially deleted, resulting in a gradual loss of activity, with hep20
showing an almost complete loss of the activity both in vitro and when injected in mice. Addition of 1 amino acid to the N-terminus, however, had no effect on the peptide’s ability to induce ferroportin degradation. Deletion of 2 C-terminal amino acids also had no significant functional consequences.

Considering the essential nature of the N-terminus, we synthesized the 3- and 6-amino acid N-terminal peptides to assess their potential use as agonists or antagonists. The 2 peptides, however, did not mimic or interfere with the activity of hep25. The only other inactive peptide, N-terminally truncated hep20, also did not interfere with ferroportin degradation induced by full-length hep25.

Of interest, pairwise replacement of cysteines with alanines, which removed disulfide bonds 1, 3, or 4 one at a time, did not cause major changes in the peptide structure when analyzed by circular dichroism spectroscopy. Accordingly, the peptide bioactivity in vitro was not significantly affected. However, in mice, the 2 peptides lacking disulfide bonds across the hairpin (C7A/C23A, C11A/C19A) were inactive at the given dose, whereas C13A/C14A peptide lacking the vicinal disulfide bond was somewhat less effective than wild-type hep25. The high degree of disulfide bonding in small peptides likely confers greater stability and resistance to proteolytic degradation, and peptides made to lack one or more disulfide bonds may have a shorter half-life in circulation.

We generated a shortened hairpin peptide with only one disulfide bond, by removing 8 residues [del9-12(FCCG), del19-22(CGMC)] in addition to replacing vicinal cysteines with alanines. Despite the significant structural alteration, the peptide still exhibited 50% activity in vitro. In vivo, however, injection of the single dose of shortened peptide in mice failed to cause a significant change in serum iron. The discrepancy between in vitro and in vivo activity is consistent with our expectations of decreased stability and shortened half-life of the mutant peptide.

Substitutions affecting cysteine residues that form disulfide bonds in the mature hepaticin have been identified in patients with juvenile hemochromatosis (C70R and C78T, both affecting bond 3) and in a patient with adult hemochromatosis (compound heterozygous hepaticin mutations R59G and C82Y, the latter affecting bond 1). The substitution of a single cysteine residue, however, would lead to the appearance of an unpaired cysteine and would likely initiate the formation of incorrect disulfide bonds and degradation of the aberrant peptides inside the cell, resulting in reduced production of the bioactive peptide. Hepcidin concentrations in these patients, however, have not yet been assessed.

Hepcidin disease-causing mutations described so far include the previously mentioned substitutions of single cysteines, as well as the premature stop codon, out-of-frame start codon, and frameshift, all expected to produce no mature peptide. The only other missense mutations in the region coding for mature peptide were generally found in the heterozygous state and have an as yet uncertain phenotypic effect. G71D was found as a possible modifier in patients with hemochromatosis carrying HFE C282Y mutations, and K83R was found in a patient with porphyria cutanea tarda and iron overload (J.K. and John Phillips, University of Utah; unpublished observations, December 2004). Synthetic G71D and K83R peptides were fully functional in in vitro assays as well as in the mouse hypoferrernia assay. These findings would suggest that neither of these hepaticin mutations is likely to be important modifiers in iron overload diseases.

It may appear counterintuitive that the hepaticin-ferroportin interaction is so tolerant of hepaticin mutations. However, the estimated affinity of the hepaticin-ferroportin interaction is relatively low both in vitro and in vivo (~500 nM), so a maximal molecular fit between the 2 structures is not necessary. In vivo, the low affinity of the hepaticin-ferroportin interaction would require large concentration of hepaticin in the target tissues, achievable because of the very large size of the hepaticin-producing organ (the liver) and relatively high rates of hepaticin synthesis, at least as estimated from mRNA abundance in gene array experiments.

In conclusion, our study revealed the N-terminus of hepaticin is essential for its bioactivity, but the structure was otherwise surprisingly permissive for changes. Further studies of hepaticin structure in relation to its function are vital for the design of hepaticin antagonists and agonists, which could be used for treatment of iron disorders.

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References

10. Papanikolaou G, Samuels ME, Ludwig EH, et al. Mutations in HFE cause major changes in the peptide structure when analyzed by circular dichroism spectroscopy. Accordingly, the peptide bioactivity in vitro was not significantly affected. However, in mice, the 2 peptides lacking disulfide bonds across the hairpin (C7A/C23A, C11A/C19A) were inactive at the given dose, whereas C13A/C14A peptide lacking the vicinal disulfide bond was somewhat less effective than wild-type hep25. The high degree of disulfide bonding in small peptides likely confers greater stability and resistance to proteolytic degradation, and peptides made to lack one or more disulfide bonds may have a shorter half-life in circulation.

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