Neogenin Facilitates the Induction of Hepcidin Expression by Hemojuvelin in the Liver

Received for publication, February 10, 2016, and in revised form, April 12, 2016. Published, JBC Papers in Press, April 12, 2016, DOI 10.1074/jbc.M116.721191

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Hemojuvelin (HJV) regulates iron homeostasis by direct interaction with bone morphogenetic protein (BMP) ligands to induce hepcidin expression through the BMP signaling pathway in the liver. Crystallography studies indicate that HJV can simultaneously bind to both BMP2 and the ubiquitously expressed cell surface receptor neogenin. However, the role of the neogenin–HJV interaction in the function of HJV is unknown. Here we identify a mutation in HJV that specifically lowers its interaction with neogenin. Expression of this mutant Hjv in the liver of Hjv−/− mice dramatically attenuated its induction of BMP signaling and hepcidin mRNA, suggesting that interaction with neogenin is critical for the iron regulatory function of HJV. Further studies revealed that neogenin co-immunoprecipitated with ALK3, an essential type-I BMP receptor for hepatic hepcidin expression. Neogenin has also been shown to facilitate the cleavage of HJV by furin in transfected cells. Surprisingly, although cleavage of HJV by furin has been implicated in the regulation of HJV function in cell culture models and furin-cleaved soluble Hjv is detectable in the serum of mice, mutating the furin cleavage site did not alter the stimulation of hepcidin expression by Hjv in mice. In vivo studies validated the important role of HJV–BMP interaction for Hjv stimulation of BMP signaling and hepcidin expression. Together these data support a model in which neogenin acts as a scaffold to facilitate assembly of the HJV-BMP-BMP receptor complex to induce hepcidin expression.

Hemojuvelin (HJV for humans, and Hjv for mouse),2 also called repulsive guidance molecule c (RGMc), is a glycosylphosphatidylinositol-linked membrane protein. It belongs to the highly conserved RGM family that also contains RGMa and RGMb. Although RGMa and RGMb are predominantly expressed in the developing and adult central nervous system and are essential for neural development (1), HJV is expressed mainly in the liver, skeletal muscle, and heart and is essential for iron homeostasis (2). Mutations in the HJV gene in humans markedly reduce hepcidin expression in the liver and result in juvenile hemochromatosis (JH), the most severe form of iron overload disorder (2). Hepcidin is a 25-amino acid peptide hormone that inhibits the iron efflux from enterocytes and macrophages into circulation by binding to and targeting ferroportin, the only known iron exporter, for degradation (3). It is synthesized in hepatocytes as an 84-amino acid prepropeptide that contains an N-terminal 24-amino acid signal sequence, a 35-amino acid proregion, and a C-terminal 25-amino acid bioactive peptide. After post-translational processing, the bioactive C-terminal 25-amino acid peptide is secreted into the circulation as a mature form to regulate iron homeostasis (4). Consistently, low hepatic hepcidin expression and a marked iron overload were also observed in Hjv knock-out (Hjv−/−) mice (5, 6). Further studies in mice with tissue-specific Hjv knockout demonstrate that only the hepatic Hjv is indispensable for hepcidin expression and iron homeostasis (7, 8).

HJV, in the liver, acts as a co-receptor for BMP6 to stimulate hepcidin expression through the BMP signaling pathway (9–11). BMP signaling is initiated upon the binding of BMP ligands to type-I and type-II BMP receptors on the cell surface. Upon BMP binding, the type-II receptors phosphorylate the type-I receptors, leading to the phosphorylation of SMAD1/5/8 in the cytoplasm. The phosphorylated SMADs form heteromeric complexes with SMAD4 and then translocate to the nucleus where they induce the transcription of target genes. HJV most likely uses two type-I BMP receptors, ALK2 and ALK3, to induce hepcidin expression, because liver-specific deletion of either Alk3 or (to a lesser extent) Alk2 causes iron overload in mice (12).

Structural studies of the HJV ectodomain demonstrate that it can simultaneously bind BMP2 and neogenin with nanomolar affinities through its N-terminal portion (amino acids 1–145) and C-terminal portion (amino acids 146–401), respectively, and identify the key residues in these molecules that are responsible for these interactions (13, 14). Neogenin is a ubiquitously expressed type-I transmembrane protein that contains four immunoglobulin (Ig)-like domains and six fibronectin III (FNIII) domains in its large extracellular region. HJV specifically binds to the FNIII 5–6 subdomains (15). However, the precise role of neogenin in HJV induction of hepcidin expression is still unclear, largely because of lack of an appropriate animal model. In a hepatoma cell line that expresses HJV, deprivation of neogenin abolishes BMP4 induction of hepcidin expression (16). In humans, the most common JH-causing mutation in HJV, G320V, disrupts its interaction with neogenin.
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(17). In mice, neogenin deficiency results in low hepcidin expression and severe iron overload that are indistinguishable from Hjv<sup>−/−</sup> mice. But interpretation of these results is complicated by other developmental defects and early death, because neogenin is also a receptor for netrins, RGMa, and RGMb (18).

Neogenin is required for the cleavage of HJV by the ubiquitously expressed furin proprotein convertases as well as matr intrinsic-2 (MT2) in transfected cells (19, 20), and cleaved HJV is readily detectable in the conditioned medium (21–23). Furin-cleaved soluble HJV suppresses hepcidin expression in HepG2 cells (24), likely by acting as a decoy to compete with membrane HJV for BMP6. MT2 is a type II transmembrane serine protease that is predominantly expressed in hepatocytes and is a key suppressor of hepcidin expression likely by cleaving membrane HIV into inactive fragments (22, 25, 26). Although soluble HIV has been detected in the serum of humans and rats (27, 28), the role of furin-cleaved Hjv in hepatic hepcidin expression and the necessity of neogenin for the cleavage of HIV in vivo are still unknown.

HJV also interacts with hemochromatosis protein (HFE) and transferrin receptor-2 (TfR2) (29), which are highly expressed in hepatocytes. In humans, mutations in either HFE or TfR2 decrease hepcidin expression and cause hereditary hemochromatosis. Although the mechanisms by which HFE or TfR2 up-regulate hepcidin expression has not been fully defined, a recent study indicates that HJV, HFE, and TfR2 operate in the same pathway (30). In the present study, we systematically examined the role of neogenin in Hjv-mediated induction of hepcidin expression in the liver of mice. Results demonstrate that an efficient induction of hepcidin expression by Hjv requires its interaction with neogenin.

Experimental Procedures

cDNA Constructs—We generated mouse Hjv ORF in pGEM-T vector (Hjv-pGEM-T) in our previous study (31). Hjv with a glycine to valine substitution at amino acid 92 (G92V-Hjv; Table 1) was generated by site-directed mutagenesis using the QuikChange kit (Stratagene). After verification by sequencing, both Hjv and G92V-Hjv constructs were subcloned into an AA8V construct containing human HJV coding sequence with a deletion of Ig 1–4 domains but still retaining an intact signal sequence (neogenin/ΔIg) by PCR using site-directed mutagenesis and the following primers: 5′-gggccccgccccgccgctcagctctccggaggtg-3′ (forward) and 5′-acatccggagttcgaagcccgccgccccgccgccc-3′ (reverse). The sequence was verified by DNA sequencing. We obtained human ALK2 ORF with an HA tag in pcDNA3.1 (pcDNA3-neogenin) were obtained from Dr. Eric R. Fearon at the University of Michigan Medical School, Ann Arbor, MI. We generated a truncated neogenin construct with a deletion of Ig 1–4 domains but still retaining an intact signal sequence (neogenin/ΔIg) by PCR using site-directed mutagenesis and the following primers: 5′-gggccccgccccgccgctcagctctccggaggtg-3′ (forward) and 5′-acatccggagttcgaagcccgccgccccgccgccc-3′ (reverse). The sequence was verified by DNA sequencing. We obtained human ALK2 ORF with an HA tag in pcDNA3.1 (pcDNA3-ALK2/HA) from Dr. Peter ten Dijke (Leiden University, The Netherlands).

Animal Analysis—Hjv<sup>−/−</sup> mice on the 129/SvEvTac (129/S) background were obtained from Dr. Nancy Andrews (Duke University). Both Hjv<sup>−/−</sup> and wild-type 129/S mice were bred and maintained in the Oregon Health and Science University DCM. We performed three separate sets of studies with at least 5 animals for each group.

For the first study, 8-week-old male Hjv<sup>−/−</sup> mice were intraperitoneally injected with PBS, AA8V-Hjv, A183R-fHjv, or G313V-fHjv at ~5 × 10<sup>11</sup> and/or ~10<sup>13</sup> genome-particles per mouse. Three weeks later, mice were euthanized for analysis. Blood was collected by cardiac puncture for serum iron and Hjv analysis. Livers were rapidly removed, snap-frozen in liquid nitrogen, and then stored at −80°C for qRT-PCR, Western blotting, immunoprecipitation, and tissue nonheme iron assays. Age and gender-matched wild-type and Hjv<sup>−/−</sup> mice were included as additional controls.

For the second study, male Hjv<sup>−/−</sup> mice were fed an iron-deficient (ID) diet (less than 1 ppm iron, TestDiet, Richmond, IN; low iron group) starting from 5 weeks old until euthanization to deplete bodily iron stores. Animals at the age of 8 weeks old were intraperitoneally injected with PBS, ~5 × 10<sup>11</sup> AA8V-fHjv, R281A-fHjv, R325A-fHjv, or GFP genome-particles per mouse. Three weeks later, mice were euthanized for analysis as described above. At 4, 16, and 24 h before euthanization, mice in AA8V-fHjv, R281A-fHjv, and R325A-fHjv groups were intraperitoneally injected with aprotinin (G Biosciences) at ~80 mg/kg body weight in PBS or with the vehicle only.

The third study for G92V-Hjv was performed in the same setting as in our previous report for Hjv (31). Briefly, 8-week-old male Hjv<sup>−/−</sup> mice were injected with ~5 × 10<sup>11</sup> AA8V-Hjv or G92V-Hjv vector particles per mouse via the tail vein. Two
weeks later, mice were euthanized for analysis. All procedures for animal use were approved by the Oregon Health and Science University DCM. The viral vector stocks were handled according to Biohazard Safety Level 2 guidelines published by the National Institutes of Health.

Quantitative Real-time RT-PCR (qRT-PCR)—qRT-PCR analysis of \( \beta \)-actin, Activin-B, Bmp6, hepcidin, Hjv, Id1, and IL-6 transcripts in the liver were conducted as previously described (31). The mouse primers used for qRT-PCR are listed in supplemental Table S1. All primers were verified for linearity of amplification. The results are expressed as the amount relative to that of \( \beta \)-actin for each sample.

Serum Iron and the Liver Nonheme Iron Assays—Serum iron concentrations were measured using a serum iron/TfR2 Reagent Set (Teco Diagnostics, Anaheim, CA). Nonheme iron concentrations in the liver tissues were determined as described in our previous studies (31).

Immunodetection of fHjv and Smad Proteins in Mice—Liver extracts were separated using SDS-PAGE, probed with mouse anti-FLAG M2 IgG (1:10,000; Sigma), rabbit anti-phosphorylated Smad1/5/8 (pSmad, 1:1,000; Cell Signaling Technology), rabbit anti-Smad1/5/8 (Smad, 1:500; Santa Cruz), and mouse anti-\( \beta \)-actin (1:10,000; Sigma), and immunodetected by using an Alexa Fluor 680 goat anti-\( \beta \)-actin secondary antibody or an Alexa Fluor 800 goat anti-mouse secondary antibody. Images were visualized and quantified using an Odyssey Infrared Imaging System (Licor). Additionally, fHjv in the liver extracts was also detected directly by using an HRP-coupled mouse anti-FLAG M2 IgG (1:10,000; Sigma) and chemiluminescence (Super Signal, Pierce) to reduce the nonspecific background. For immunodetection of serum fHjv, fHjv in \( \sim 60 \mu l \) of serum was pulled down with mouse anti-FLAG M2 beads (Sigma), followed by elution using 1\( \times \) Laemmli buffer, separation using SDS-PAGE, and immunodetection using HRP-coupled mouse anti-FLAG M2 IgG and chemiluminescence. The intensities of fHjv bands in x-ray films were quantified by using ImageJ software.

Immunoprecipitation—We used HeLa cells that stably express both human TfR2 and HFE with a C-terminal FLAG tag (HeLa-TfR2/\( \beta \)FHE) for co-immunoprecipitation analysis of human HJV and mutants with TfR2 and HFE. Briefly, HeLa-TfR2/\( \beta \)FHE cells were transiently transfected by pcDNA3 empty vector, pcDNA3-HJV, and mutants using TransIT-2020 transfection reagent (Mirus). After 48 h of transfection, cell lysates were prepared. Immunoprecipitation was performed using protein A-agarose beads and rabbit anti-neogenin FNIII 1–6 as described above. Immunoprecipitated proteins were probed with mouse anti-HJV, goat anti-neogenin (C20; Santa Cruz), or mouse anti-\( \beta \)-actin antibody, followed by immunodetection using an Alexa Fluor 800 goat anti-mouse secondary antibody or an Alexa Fluor 680 donkey anti-\( \beta \)-actin antibody and visualized using a Licor.

We used HEK293 cells that stably express ALK2/\( \beta \)A (HEK293-ALK2/\( \beta \)A) for the immunoprecipitation analysis with neogenin and neogenin/\( \Delta \)lg. Briefly, HEK293-ALK2/\( \beta \)A and ALK3/\( \beta \)A cells were transiently transfected by pcDNA3 empty vector, pcDNA3-neogenin, and pcDNA3-neogenin/\( \Delta \)lg plasmid DNA using TransIT-2020 transfection reagent. After 48 h of transfection, cell lysates were prepared. Immunoprecipitation was performed using protein A-agarose beads and rabbit anti-neogenin FNII 1–6 as described above. Immunoprecipitated proteins were probed with mouse anti-\( \beta \)-actin (12CA5; Abcam), goat anti-\( \beta \)-actin (C20; Santa Cruz), or mouse anti-\( \beta \)-actin antibody, followed by immunodetection using an Alexa Fluor 800 goat anti-mouse secondary antibody or an Alexa Fluor 680 donkey anti-\( \beta \)-actin antibody and visualized using a Licor.

We used liver extracts from \( Hjv^{-/-} \) mice injected with AA/8-fHjv, A183R-fHjv, and G313V-fHjv vectors at \( \sim 5 \times 10^{11} \) genome particles per mouse for the co-immunoprecipitation analysis with neogenin in vivo. Briefly, about 3 mg of extract protein was immunoprecipitated by using anti-FLAG M2 beads for fHjv as described above. The immunoprecipitated proteins and about 250 \( \mu g \) of extract protein (input) were subjected to SDS-PAGE, followed by immunodetection using a HRP-coupled mouse anti-FLAG M2 IgG and chemiluminescence for fHjv and using a rabbit anti-neogenin FNIII 1–6, an Alexa Fluor 800 goat anti-rabbit secondary antibody, and a Licor for neogenin.

Biotinylation of Cell Surface Proteins—HEK293 cells were plated in the polylysine-coated 6-well plate and transfected with pcDNA3 empty vector, pcDNA3-fHjv, A183R-fHjv, G313V-fHjv, neogenin and neogenin/\( \Delta \)lg plasmid DNA using Lipofectamine 2000 (Invitrogen). After 48 h of transfection, cell surface proteins were biotinylated with 0.25 mg/ml of Sulfo-NHS-Biotin (Thermo Fisher Scientific) at 4 \( ^\circ \)C for 30 min. After the reaction was terminated, cells were immediately solubilized in NET-Triton, 1\( \times \) protease inhibitors mixture (Roche Applied Science). Biotinylated proteins were isolated using streptavidin-agarose beads (Thermo Fisher Scientific). Bound proteins were eluted with NET-Triton, 1\( \times \) Laemmli buffer, subjected to SDS-PAGE, probed with mouse anti-HJV, mouse anti-\( \beta \)-actin antibody, and immunodetected using an
Alexa Fluor 800 goat anti-mouse secondary antibody or an Alexa Fluor 680 goat anti-rabbit secondary antibody and visualized using a Licor.

**Analysis of HJV Secretion**—HepG2 cells that stably expressed HJV (HepG2-HJV) were plated in 12-well plates. After 2 days of incubation, fresh medium (MEM, 10% FCS) containing various concentrations of aprotinin (0, 0.005, 0.01, 0.05, 1, 5, and 10 μM) was replaced, and cell lysate and conditioned medium (CM) cells were collected after an additional 16 h of incubation. About 50% of cell lysate and 10% of CM were subjected to SDS-PAGE and immunodetection using rabbit anti-HJV antibody, HRP-conjugated goat anti-rabbit secondary antibody, and chemiluminescence.

**Statistical Analysis**—One-way analysis of variance and Tukey’s post test were used to compare three or more sets of data. Only the results for the groups of interest were presented.

**Results**

Identification of a Mutation in HJV That Disrupts Its Interaction with Neogenin, but Not TfR2 or HFE—To determine the role of the HJV-neogenin complex in the induction of hepcidin expression in vivo we generated a new mutant form of HJV that fails to interact with neogenin. G320V-HJV (corresponding to G313V-Hjv in mouse; Table 1) that does not interact with neogenin (17) is not ideal because of the contradictory results of its cell surface localization in previous studies (32, 33). Based on the crystallographic and binding studies, Ala-186 is a key highly conserved residue involved in the interaction of the closely related RGMb with neogenin (14). In human HJV the equivalent residue is Ala-190 and in murine Hjv it is Ala-183. We tested whether mutations in this site in human HJV (A190R-HJV; Fig. 1A) and in mouse Hjv (A183R-fHjv, Fig. 1B) could disrupt interaction with neogenin. For mouse Hjv a construct

### Table 1

**Mutations in HJV used in this study**

| Human residue | Eq. murine residue | Disrupted interaction/function | References |
|---------------|-------------------|--------------------------------|------------|
| G99V          | G92V              | BMP2                          | 33         |
| A190R         | A183R             | Neogenin                      | This study |
| R288A         | R281A             | Neogenin                      | This study |
| G320V         | G313V             | Neogenin                      | 17         |
| R332A         | R325A             | Furin cleavage                | 24         |

**Figure 1.** Interaction of mutant hemajuvelin with neogenin and cell surface localization of Hjv. A. A190R-HJV did not co-immunoprecipitate with neogenin. HEK293-neogenin cells were transfected by pcDNA3 (C), pcDNA3-HJV or A190R-HJV, immunoprecipitated (IP) by using anti-HJV antibody generated against residues 1–401 of HJV, and immunodetected for neogenin (Neog) and HJV, and β-actin. B. co-immunoprecipitation of WT and mutant fHjv with neogenin. HEK293 cells were co-transfected by pcDNA3-neogenin with pCMV9-fHjv, A183R-fHjv, or G313V-fHjv, immunoprecipitated by using anti-FLAG M2 beads, and immunodetected for neogenin (Neog) and fHjv. HEK293 cells transfected with pCMV9 alone (C) were included as a control. The entire images are illustrated in supplemental Fig. S1A. C. co-immunoprecipitation of WT and mutant HJV with neogenin. HEK293-neogenin (HEK-Neog) cells were transfected by pcDNA3 (C), pcDNA3-HJV, and G320V-HJV, immunoprecipitated by using anti-HJV antibody, and immunodetected for neogenin (Neog) and HJV. HEK293 cells (HEK) were included as a control. The entire images are illustrated in supplemental Fig. S1B. D. co-immunoprecipitation of WT and A190R-HJV with HFE. HeLa-TfR2/HFEC cells were transfected by pcDNA3 (C), pcDNA3-HJV, and A190R-HJV, immunoprecipitated by using anti-HJV antibody, and immunodetected for TR2, HJV, and HFE. E. biotinylation of cell surface Hjv. HEK293 cells were transfected with pCMV9 (C), pCMV9-fHjv, A183R-fHjv, and G313V-fHjv, followed by biotinylation of cell surface proteins at 4 °C. Hjv, Na+/K+ ATPase, and β-actin in the biotinylated proteins (Cell surface) and cell lysate input (Input) were immunodetected. F. co-immunoprecipitation of WT and G320V-HJV with HFE and TR2. HeLa-TfR2/HFEC cells were transfected by pcDNA3 (C), pcDNA3-HJV, and G320V-HJV, immunoprecipitated by using anti-HJV antibody, and immunodetected for TR2, HJV, and HFE. The entire images are illustrated in supplemental Fig. S1C. All experiments were repeated at least three times with consistent results.
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with a 3× FLAG sequence at the site immediately after the signal peptide was used because the FLAG tag was necessary for detection of Hjv in subsequent in vivo experiments. A190R-HJv, A183R-fHjv, and wild-type counterparts were transfected into HEK293 cells that express endogenous and transfected neogenin. Both A190R-HJv and A183R-fHjv had reduced interaction with human neogenin as judged by co-immunoprecipitation using anti-HJV or anti-FLAG antibody, respectively (Fig. 1, A and B, and supplemental Fig. S1A). As positive controls, both Hjv and fHjv were able to pulldown neogenin (Fig. 1, A and B). As negative controls, both G320V-HJv and G313V-fHjv had decreased interaction with neogenin by co-immunoprecipitation (Fig. 1, B and C, supplemental Fig. S1, A and B). These results confirmed Ala-190 in HJv or Ala-183 in Hjv as key residues for a high affinity interaction with neogenin, mutation of which could serve as a new tool for interrogating the importance of the HJv-neogenin interaction in vivo.

We next examined the effect of the A190R mutation in HJv on the interaction with TfR2 and HFE. To avoid the potential effects of sequence difference between species, only human proteins were used. HeLa cells that express TfR2 and FLAG-tagged HFE (HeLa-TfR2/HFE) were transfected with HJv or mutants. Similar to wild-type HJv, A190R-HJv was co-immunoprecipitated with TfR2 and fHFE using anti-HJV antibody (Fig. 1D). These results indicate that disruption of the HJv-neogenin interaction does not affect HJv interaction with TfR2 or HFE. Additionally, the A183R mutation did not reduce fHjv localization on the plasma membrane in HEK293 cells by biotinylation of cell surface proteins (Fig. 1E). These results indicate that disruption of the HJv-neogenin complex does not affect either HJv interaction with TfR2 and HFE or its cell surface localization, making A183R-Hjv a good tool for assessing the importance of the HJv-neogenin interaction in vivo. Interestingly, similar results were also obtained for G320V-HJv and G313V-fHjv (Fig. 1, E and F, and supplemental Fig. S1C). Thus G313V-fHjv is also a good candidate for in vivo studies.

Disruption of HJv-Neogenin Complex Attenuates HJv Induction of Hepcidin mRNA in the Liver of HJv−/− Mice—To determine the in vivo role of the HJv-neogenin interaction, we established an animal model to express fHjv in the liver of HJv−/− mice using an AAV8 vector. Our previous studies indicate that AA8 with a liver-specific promoter limits HJv expression specifically to the liver and does not induce inflammation that could complicate interpretation of the results (31). Addition of a 3xFLAG tag to Hjv was used to detect Hjv in mouse tissues and serum, because in our hands there was no available antibody to detect Hjv in vivo. Hjv, A183R-fHjv, or G313V-fHjv in AAV8 (∼5 × 1011 genome particles per mouse) were intraperitoneally administered into 8-week-old male HJv−/− mice fed a regular diet. Animals were euthanized for analysis 3 weeks after injection. The mRNA levels of fHjv, A183R-fHjv, and G313V-fHjv in the liver were comparable with that of wild-type mice (Fig. 2A). Consistently, Western blotting analysis using anti-FLAG antibody detected similar levels of fHjv and A183R-fHjv, and about 50% higher level of G313V-fHjv, which migrated at ∼55 kDa and corresponded to the expected full-length Hjv (Fig. 2, B, top panels, lanes 5–8, 11, and 12, and C). As negative controls, no corresponding band was detected either in wild-type mice or in Hjv−/− mice (Fig. 2, B and C). In line with our previous studies using Hjv with no tag (31), expression of fHjv was able to fully correct the low hepcidin status in HJv−/− mice, and the level of hepcidin mRNA was even greater than that of wild-type mice (Fig. 2D). The levels of hepatic hepcidin mRNA were used as an indicator for the levels of serum hepcidin throughout the animal studies, because recent studies demonstrated a strong correlation between the levels of hepatic hepcidin mRNA and serum hepcidin in mice (34–36). The increased hepcidin mRNA was associated with an increase in phosphorylated Smad1/5/8 (pSmad) and the mRNA level of Id1 (Fig. 2, B and E, and supplemental Fig. S2A), consistent with the idea that fHjv induces hepcidin expression by activating the BMP signaling pathway. The phosphorylation of Smad1/5/8 is a direct indicator of the activation of BMP signaling, whereas Id1 is a direct target of BMP signaling. Both IL-6 and Activin-B are inflammatory factors that can induce hepcidin expression through Stat3 and the BMP signaling pathway, respectively, independent of Hjv (37). There was no significant change in either IL-6 or Activin-B mRNA levels in the livers (supplemental Fig. S2, B and C). Thus expression of fHjv in the liver was able to specifically induce hepcidin expression, which was correlated with stimulation of the BMP signaling. Consistent with the role of hepcidin in the suppression of iron efflux from enterocytes and macrophages, induction of hepcidin expression by fHjv significantly decreased the levels of both serum iron and the liver non-heme iron (Fig. 2, F and G). The latter is an indicator of bodily iron stores.

Expression of a similar level of A183R-fHjv in mouse livers to that of fHjv (Fig. 2B, lanes 7 and 8 versus 5 and 6, and C) was only able to mildly increase hepcidin mRNA (Fig. 2D). As a negative control, expression of a comparable level of G313V-fHjv, a mutant form of Hjv that causes JH, also exhibited a similarly attenuated induction of hepcidin expression (Fig. 2, A–D). In contrast to fHjv, neither A183R-fHjv nor G313V-fHjv was able to significantly increase the levels of pSmad1/5/8 (Fig. 2B, lanes 7/8 and 11/12 versus 5/6, supplemental Fig. S2A) and Id1 mRNA (Fig. 2E). Both Smad1/5/8 and β-actin were included as loading controls (Fig. 2B). These results indicate that the decreased hepcidin induction by A183R or G313V-fHjv resulted from lack of stimulation of BMP signaling in the liver. In agreement with low hepcidin mRNA, expression of neither A183R-Hjv nor G313V-fHjv was able to lower the levels of serum iron and liver non-heme iron (Fig. 2, F and G). Consistent with the findings in transfected cells, both A183R-fHjv and G313V-fHjv exhibited reduced interaction with neogenin in the liver extracts by co-immunoprecipitation using anti-FLAG antibody, when compared with fHjv (Fig. 2H). These results indicate that disruption of HJv-neogenin interaction markedly reduces Hjv induction of hepcidin expression in the liver.

To test whether increased expression of fHjv mutants could correct the status of low hepcidin expression, HJv−/− mice were intraperitoneally administered a 20-fold higher dose of AA8-V183R-fHjv and G313V-fHjv. The levels of hepatic Hjv mRNA in A183R-fHjv and G313V-fHjv groups reached about 25- and 16.5-fold higher than that of fHjv group, respectively (Fig. 2A). The protein levels of A183R and G313V-fHjv in the liver were similarly increased (Fig. 2, B, lanes 9/10 and 13/14, and C).
Interestingly, overexpression of these mutants was able to increase hepatic hepcidin mRNA to a level similar to the mice with wild-type fHJV (Fig. 2D). Consistently, elevated hepcidin mRNA was correlated with increases in pSmad (Fig. 2, B, lanes 9/10 and 13/14, supplemental Fig. S2A) and Id1 mRNA (Fig. 2E). Importantly, injection of a large amount of AAV8 did not induce any evident inflammation as measured by a lack of increase in IL-6 and Activin-B mRNA in the liver (supplemental
As expected, increased hepcidin mRNA led to a significant reduction of serum iron and the liver non-heme iron (Fig. 2, F and G). These results indicate that both A183R and G313V-fHjv are partially functional and in excess can induce hepcidin expression through the BMP signaling pathway. At physiological levels, however, the inability to interact with neogenin prevents these Hjv mutants from properly controlling iron levels through regulation of hepcidin expression.

Interaction with Neogenin Is Not Required for Furin Cleavage of HJV in Vivo and Abolishment of Furin Cleavage in HJV Does Not Affect the Induction of Hepatic Hepcidin mRNA—The cleavage of HJV by furin is promoted by neogenin in cell culture models, and furin-cleaved soluble HJV is proposed to suppress the function of membrane HJV. The essential furin cleavage site in HJV was mapped to Ala-332 (corresponding to Ala-325 for mouse Hjv) at the conserved polybasic RNRR motif, and the R332A mutation in HJV abolishes furin cleavage in cultured cells (21, 23, 24). To determine whether neogenin regulates the cleavage in vivo and the role of furin-cleaved Hjv in hepatic hepcidin expression, both fHjv and R325A-fHjv were expressed in the livers of Hjv<sup>−/−</sup> mice. Expression of AAV8-GFP was included as a negative control. Mice were fed an ID diet starting from 5 weeks of age to deplete excess liver iron. In cellular studies, furin cleavage of HJV is negatively regulated by iron (21, 23, 28).

Hjv<sup>−/−</sup> mice fed an ID diet for 6 weeks led to decreased liver non-heme iron by about 50%, serum iron by 10%, and hepatic hepcidin mRNA by about 100-fold compared with Hjv<sup>/+</sup> mice fed a regular iron diet (Fig. 3, A–C). Consistent with the finding that BMP6 mRNA levels are positively regulated by iron in the liver (38), hepatic BMP6 mRNA levels decreased by about 50% (supplemental Fig. 3A). Hjv mRNA levels for both fHjv and G325A-fHjv groups were comparable with those of wild-type mice (Fig. 3D). Western blotting analysis revealed similar levels of fHjv and G325A-fHjv in the liver extracts (Fig. 3E, top panel, and lanes 3/4 versus 7/8, supplemental Fig. S3B). To detect the furin-cleaved fHjv, −60
A single distinct fHjv band migrating at ~43 kDa was observed in the serum of fHjv mice but not in GFP mice (Fig. 3E, lower panel, lanes 3 and 4). The molecular weight corresponds to that of the predicted size of the furin-cleaved product. In contrast, no serum fHjv was detected in the serum of R325A-fHjv mice (Fig. 3E, lower panel, lanes 7 and 8). These findings indicate that furin uses the same site to cleave Hjv in the liver as in transfected cells.

To test the role of neogenin in furin cleavage of Hjv, we compared the relative levels of serum fHjv in Hjv+/- mice expressing fHjv, to the A183R, and G313V mutants that have reduced interaction with neogenin. The sizes of serum Hjv from all three groups corresponded to the furin-cleaved products (Fig. 4, A and B). When comparable levels of fHjv protein were expressed in the liver, we detected similar levels of serum fHjv between fHjv and A183R-fHjv groups, and a relatively low level in the G313V-fHjv group (Fig. 4A, lower panel). Interestingly, when the levels of A183R and G313V-fHjv proteins were increased in the liver, serum fHjv levels from both groups were proportionally increased (Figs. 2, B and C, and 4, B and C), indicating that mutated proteins were able to be processed and secreted. These results suggest that interaction with neogenin is not required for furin cleavage of Hjv in the liver.

Because furin-mediated cleavage of HJV negatively regulates HJV activity in the human hepatoma cell line HepG2 cells (24), we compared hepcidin expression in fHjv mice to that of R325A-fHjv mice to determine whether soluble Hjv inhibits hepcidin expression in vivo. Expression of fHjv in the liver of Hjv+/- mice was able to robustly increase hepcidin mRNA by about 1000-fold, reaching a level well above that of wild-type mice (Fig. 3C). As a negative control, expression of GFP displayed no significant effect indicating that expression of an exogenous protein was unable to perturb hepcidin expression (Fig. 3C). Surprisingly, R325A-fHjv exhibited a similar extent of induction of hepcidin expression as fHjv (Fig. 3C). This is in contrast to our initial prediction that abolishment of furin-cleaved Hjv in the serum would enhance hepcidin expression by increasing BMP6 signaling in the liver. In comparison with GFP, expression of both fHjv and R325A-fHjv increased hepatic mRNA of Id1 to a similar extent (Fig. 3F). Neither induced the expression of the inflammation factors, IL-6 and Activin-B, in the liver (supplemental Fig. S3, C and D). These results provided further evidence to support that fHjv induces hepcidin expression specifically through the BMP signaling pathway. As expected, the increased hepcidin mRNA observed with fHjv and R325A-fHjv reduced serum iron to a similar extent (Fig. 3B). No further decrease in the liver non-heme iron was detected because these animals were fed an ID diet (Fig. 3A). Together, these observations indicate that serum Hjv derived from the liver does not repress the ability of membrane-anchored Hjv to induce hepcidin expression in vivo.
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Disruption of the Predominant MT2 Cleavage Site in Hjv Abolishes Hjv-Neogenin Interaction and Reduces the Induction of Hepcidin mRNA in Vivo—Neogenin also interacts with MT2 to facilitate MT2 cleavage of HJV in cultured cells (20). MT2 cleaves HJV at distinct sites from furin (24). The major MT2 cleavage site was mapped to Arg-288 in HJV (corresponding to Arg-281 for murine Hjv), and the R288A mutation in HJV abolishes MT2 cleavage in transfected cells (24). Because mutations of the MT2 gene in humans or deletion of the catalytic domain of MT2 in mice result in inappropriately high hepcidin expression and iron deficiency (25, 39), we hypothesized that disruption of the MT2 cleavage site in Hjv would increase its induction of hepcidin expression in vivo. To test this hypothesis, we first examined whether MT2 cleaves Hjv in the liver. Surprisingly, no predicted MT2-cleaved fHjv band was detected in the serum of Hjv−/− mice expressing fHjv, A183R-fHjv, G313V-fHjv, or R325A-fHjv (Figs. 3E, and 4, A and B). When compared with fHjv, expression of R281A-fHjv that is expected to be resistant to MT2 cleavage in the liver of Hjv−/− mice was expected to be lower band furin convertase inhibitor in our previous studies (24), and the R281A-fHjv mice were significantly higher than that of fHjv mice (Fig. 3B). This unexpected finding is in agreement with a clinical study, which shows that R288W mutation in human HJV (corresponding to R281W in murine Hjv) causes JH (40). R288A-HJV traffics to the plasma membrane (24), indicating that it is folded correctly. To explore the potential mechanism, we determined whether this mutation affects its interaction with other important proteins. Similar to wild-type HJV, R288A-HJV was co-immunoprecipitated with both Tfr2 and Hfe (Fig. 5A). Intriguingly, both R281A-fHjv and R288A-HJV had reduced interaction with neogenin in transfected HEK293 cells by immunoprecipitation (Fig. 5, B and C). These observations indicate that R288A mutation in HJV disrupts its association with neogenin, but not with Tfr2 or Hfe, and suggest that the attenuated induction of hepcidin expression by R281A-fHjv in Hjv−/− mice results from a lack of interaction with neogenin.

These findings provided another line of evidence supporting the critical role of neogenin in the induction of hepcidin by HJV.

Neogenin Interacts with ALK3 through Its Ig Domains—Because the interaction of HJV with neogenin was critical for the induction of hepcidin expression through the BMP signaling pathway in vivo, we wanted to investigate whether neogenin itself had any direct interactions with the BMP receptors, and whether these interactions occurred through domains other than FNIII 5–6 in neogenin (Fig. 6A). FNIII 5–6 subdomains are responsible for interaction with HJV (15). Human liver expresses two type-I BMP receptors, ALK2 and ALK3 (41). Both ALK3 and (to a lesser extent) ALK2 are essential for hepcidin expression in mice (12). We examined the interaction of neogenin with ALK2 and ALK3 by immunoprecipitation. HEK293 cells that stably express ALK2 with a HA tag (ALK2/HA) or ALK3 with a HA tag (ALK3/HA) were transfected with neogenin. Interestingly, rabbit anti-neogenin antibody was able to pull down ALK3/HA, but not ALK2/HA (Fig. 6, B and C). These observations indicate that neogenin interacts with ALK3.

To map the domains in neogenin responsible for this interaction, a truncated construct without all four Ig subdomains but with an intact signal peptide sequence (neogenin/ΔIg; Fig. 6A) was generated. Deletion of four Ig subdomains did not affect its trafficking to the plasma membrane in HEK293 cells (Fig. 6D). An anti-neogenin antibody was able to pull down neogenin/ΔIg. But no detectable ALK3/HA was co-immunoprecipitated with neogenin/ΔIg (Fig. 6C). These observations suggest that neogenin interacts with ALK3 through its Ig domains, which is a different region than what binds HJV, implying that these interactions may not be mutually exclusive. These findings also provide a potential mechanism for the involvement of neogenin in the BMP signaling pathway through binding to ALK3.

The induction of hepcidin mRNA by R281A-fHjv was next examined in the liver of Hjv−/− mice. In contrast to our prediction that expression of R281A-fHjv that is resistant to MT2 cleavage would increase hepcidin expression, qRT-PCR analysis revealed an attenuated induction when compared with fHjv (Fig. 3C and supplemental Fig. S3F). This reduced hepcidin mRNA was correlated with a compromised induction of BMP signaling as manifested by significantly lower levels of id1 mRNA than the fHjv group in the liver (Fig. 3F). Consistent with reduced hepcidin expression, serum iron levels in R281A-fHjv mice were significantly higher than that of fHjv mice (Fig. 3B).
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Disruption of Hjv interaction with BMP ligand, but not Neogenin, also reduces the induction of hepcidin mRNA in the liver—Because HJV can simultaneously bind BMP2 and neogenin (13) and the above studies showed that the Hjv-neogenin interaction is essential for hepcidin expression, we next wanted to validate the importance of BMP-HJV interactions in HJV induction of hepcidin expression in vivo. The crystallographic studies indicate that the G99V mutation in HJV causes JH in humans (40). The Gly-99 residue in HJV (corresponding to Gly-92 in mouse Hjv) is located at the interface with BMP2 (13). In transfected cells, G92V mutation in Hjv abolishes the binding to BMP2 (33). Immunoprecipitation analysis indicated that the G99V mutation in HJV did not affect its interaction with neogenin, TR2, or HFE (Fig. 7, A and B). Thus the major defect of G99V-HJV or G92V-Hjv is a lack of interaction with BMP.

G92V-Hjv with no tag was expressed in the liver of Hjv−/− mice by using AAV8. Expression of wild-type Hjv mRNA at ∼15% of the levels seen in the wild-type mice was able to restore hepcidin expression in Hjv−/− mice (Fig. 7, C and D) (31). This suggests that the levels of Hjv in the liver is not a limiting factor for the induction of hepcidin expression (31). Interestingly, expression of a similar level of G92V-Hjv mRNA was only able to induce hepcidin mRNA to 20% of that induced by wild-type Hjv, with no evident increase in pSmad in the liver (Fig. 7, C–F). In this study, we did not detect the levels of expressed Hjv protein because of lack of an appropriate antibody for immunodetection of the liver Hjv. These results indicate that the attenuated induction of hepcidin mRNA by G92V-Hjv resulted from a lack of activation of the BMP signaling. Consistent with low hepcidin mRNA, no significant reduction of serum iron or liver non-heme iron was detected in the G92V-Hjv group (Fig. 7, G and H). These results recapitulated the low hepcidin status in JH patients with G99V-HJv and substantiated the pivotal role of HJV-BMP interactions for HJV induction of hepcidin mRNA in the liver.

Discussion

This study investigated the role of the Hjv-neogenin interaction in the liver of Hjv−/− mice. Our results showed that this interaction is essential for the efficient induction of hepcidin mRNA by Hjv through the BMP signaling pathway. This process likely involves the association of neogenin with ALK3. In contrast to the observations in cultured cells, additional studies revealed that the Hjv-neogenin interaction was not required for furin cleavage of Hjv in the liver, and that the furin-cleaved soluble Hjv from the liver did not affect hepcidin expression. Further studies also substantiated the pivotal role of Hjv-BMP interaction in hepcidin expression. These findings support the key role of neogenin in the assembly of an HJv-BMP-BMP receptor complex to induce BMP signaling and hepcidin expression in the liver.

We established a mouse model to study the function of neogenin in Hjv induction of hepcidin expression in the liver by using AAV8 with a liver-specific promoter, FLAG-tagged Hjv, and adult Hjv−/− mice. This AAV8 vector was able to specifically deliver Hjv cDNA into the liver of mice (31) with no activation of inflammation at a wide range of dosage. Addition of a FLAG tag to Hjv did not affect Hjv induction of the BMP sig-
naling and hepcidin mRNA. Rather it allowed detection of the fHjv protein in the liver and furin-cleaved fHjv in the serum. The use of adult mice circumvented the confounding issues of developmental defects encountered in neogenin mutant mice (18). In the studies of A183R and G313V-fHjv, results revealed a positive correlation between the administered AAV8 particles and the expressed Hjv at both mRNA and protein levels. Thus this model allows an easy control of the expression levels of Hjv simply by adjusting the dosage of AAV8.

The first line of evidence to support the critical role of HJV-neogenin interaction in Hjv induction of hepcidin expression was derived from the studies of A183R-fHjv that did not interact with neogenin. This mutant was generated based on co-crystal studies of RGMb and neogenin FNIII 5–6 domains (14). In the liver of Hjv/Hjv mice, expression of A183R-fHjv exhibited an attenuated BMP signaling and a reduced hepcidin mRNA. Thus the major defect is that these mutations disrupt the Hjv-neogenin interaction. These observations indicate that the interaction with neogenin is essential for Hjv-mediated induction of hepcidin expression through the BMP signaling in the liver. The lack of complete abolishment of hepcidin expression by these mutations might be due to the possibility that they still interact with neogenin with low affinity.

The second line of evidence supporting the role of the neo- genin-HJV interaction in the regulation of iron homeostasis was obtained from studying MT2 cleavage of Hjv. Arg-281 in Hjv is expected to be the major MT2 cleavage site from the studies in transfected cells (24). We initially predicted that disruption of MT2 cleavage of Hjv by the R281A mutation would lead to a greater induction of hepcidin expression than wild-type fHjv, because mutations of the MT2 gene in humans or deletion of the catalytic domain of MT2 in mice result in an inappropriately high hepcidin expression (25, 39). Unexpectedly, expression of R281A-fHjv in the liver of Hjv/Hjv mice revealed a greatly reduced induction of hepcidin expression when compared with wild-type fHjv. The extent of reduction is comparable with A183R- and G313V-fHjv. This unanticipated

**FIGURE 6. Neogenin is co-immunoprecipitated with ALK3 through its Ig domains.** A, diagrams of neogenin and truncated neogenin lacking Ig domains (neogenin ΔIg). TM, transmembrane domain; CD, cytoplasmic domain. B, anti-neogenin FNIII 1–6 antibody pulled down neogenin with ALK3/HA, but not ALK2/HA. HEK293-ALK2/HA or ALK3/HA cells were transfected with pcDNA3 (−) or pcDNA3-neogenin plasmid DNA (+), immunoprecipitated (IP) by using rabbit anti-neogenin FNIII 1–6 antibody, and immunoblotted (IB) with goat anti-neogenin (Neo), mouse anti-HA, or mouse anti-β actin antibodies in lysate input (Input) and IP eluate. C, neogenin ΔIg was not co-immunoprecipitated with ALK3/HA by anti-neogenin FNIII 1–6 antibody. HEK293-ALK3/HA cells were transfected with pcDNA3 (C), pcDNA3-neogenin, or neogenin ΔIg plasmid DNA, immunoprecipitated by using rabbit anti-neogenin FNIII 1–6 antibody, and immunodetected for neogenin, ALK3/HA, and β-actin in lysate input (Input) and IP eluate. D, biotinylation of cell surface neogenin. HEK293 cells were transfected with pcDNA3 (C), pcDNA3-neogenin, or neogenin ΔIg plasmid DNA, followed by biotinylation of cell surface proteins at 4 °C and immunodetection of neogenin, Na+/K+-ATPase, and β-actin in the biotinylated proteins (Cell surface) and cell lysate input. All experiments were repeated at least three times with consistent results.
finding was supported by a clinical study, which shows that the R288W mutation in human HJV (corresponding to R281W in murine Hjv) causes JH (40). This implies that Arg-281 in Hjv acts as a key residue for its function. Further studies indicate that the R281A mutation in Hjv disrupted its interaction with neogenin, but not with TfR2 or HFE, which is similar to A183R- and G313V-fHjv. These findings strengthen the idea that the HJV-neogenin interaction is essential for hepcidin expression. Additionally, we also obtained indirect evidence showing that MT2 did cleave Hjv in the liver and that Arg-281 might not be the only MT2 cleavage site in Hjv.

Previous studies indicate that neogenin is also involved in furin cleavage of HJV. In HepG2 cells, secretion of furin-cleaved HJV is blocked by siRNA knockdown of endogenous neogenin (19, 20). In HEK293 cells and cultured primary mouse muscles, expression of neogenin inhibits HJV secretion (18). In this...
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study, we confirmed that furin cleavage of Hjv in the liver used the same site as in transfected cells (21, 23, 24). Furin-cleaved fHjv in the serum was detectable by Western blot, which indicates that the liver is a source of serum Hjv. Interestingly, when compared with fHjv, there was no evident decrease of furin-cleaved fHjv in the serum of mice expressing A183R-fHjv or R281A-fHjv that did not interact with neogenin. These data indicate that HJV-neogenin interaction may not be required for furin cleavage of HJV in the liver, which is contrary to the previous observations in transfected cells. However, this does not rule out the possibility that neogenin is involved in furin cleavage of HJV in other tissues.

Furin-cleaved-soluble HJV is proposed to be a negative regulator of hepatic hepcidin expression by acting as a decoy that competes with membrane HJv for binding to BMP6. In this study, abolishment of furin cleavage of Hjv in the liver (by expressing R325A-fHjv) resulted in no detectable fHjv in the serum, but had no evident effect on hepcidin mRNA. Thus soluble Hjv does not appear to be in sufficient amounts to compete with membrane-bound Hjv for binding to BMPs. It is likely that furin cleavage of Hjv represents a process of membrane Hjv turnover in vivo. In HepG2 cells, membrane HJv has a half-life of about 1 h, and HJV release from the cells constitutes the major pathway of HJV turnover (42).

Mechanistic studies indicate that neogenin interacts with ALK3 through its Ig subdomains, which provides a potential linkage of neogenin to the BMP signaling. ALK3 is an important type-I BMP receptor in the liver for BMP signaling and hepcidin expression in mice (12). Thus neogenin can simultaneously bind both HJV and ALK3 via its FNIII 5–6 subdomains that are immediately adjacent to the transmembrane domain and its N-terminal Ig domains, respectively. This study also validated the essential role of Hjv-BMP interaction for induction of hepcidin expression by using G92V-Hjv that interacts with neogenin but does not bind to BMP2 (33). Structural analysis indicates that the N-terminal portion of Hjv shares an overlapping BMP2-binding surface with the ectodomain of ALK3 (13).

On the basis of the observations in this and previous studies, we propose a model in which neogenin acts as a scaffold by a direct association with both HJV and ALK3 to facilitate the assembly of BMP ligand-ALK3-type-II BMP receptor complex in the liver, which subsequently initiates the BMP signaling cascade to induce the transcription of the hepcidin gene (Fig. 7I). Consistent with this model, mutations in HJV that disrupt its interaction with either neogenin or BMP cause JH. In summary, this study provided evidence in support of the idea that HJV induction of hepcidin expression in the liver requires its association with both neogenin and BMP.

Author Contributions—N. Z., J. E. M., C. A. E., and A. Z. designed the experiments, analyzed the data, and wrote the paper. N. Z., J. E. M., R. H. Z., M. W., and A. Z. performed the experiments.

Acknowledgments—We thank Dr. Jodie Babitt and Dr. Herbert Lin for HJV-pCMV9, Dr. Eric R. Fearon for Neogenin-pcDNA3.1, Dr. Peter ten Dijke for pcDNA3-ALK2-HA and ALK3-HA, and Dr. Nancy Andrews for Hjv mice.

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