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ABSTRACT

Low levels of hepcidin are responsible for the development of iron overload in p.Cys282Tyr HFE related hemochromatosis. Every genetic factor lowering the hepcidin gene expression could contribute to a more severe phenotype in HFE hemochromatosis. Based on this hypothesis, we identified a heterozygous nc.-153 C>T mutation in the hepcidin gene promoter sequence in a patient homozygous for the p.Cys282Tyr HFE mutation who presented massive iron overload, resisting to well conducted iron depletive treatment. Our results demonstrate that the nc.-153 C>T mutation, located within a BMP-RE (Bone Morphogenetic Protein-Responsive Element): i) decreases the transcriptional activity of the hepcidin promoter, ii) alters its IL-6 (Interleukin-6) total responsiveness, and iii) prevents the binding of the SMAD protein complex (1/5/8 and 4) to the BMP-RE. In conclusion, our results suggest that a mutation in the BMP-RE of hepcidin promoter may impact on human iron metabolism.

Key words: hepcidin, BMP, hemochromatosis, iron, gene expression.


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Introduction

Hepcidin, synthesized mainly by hepatocytes and secreted in plasma, plays a role in the control of iron metabolism by interacting with ferroportin, the major iron exporter expressed especially on macrophages and on basolateral membranes of enterocytes. Hepcidin induces degradation of ferroportin and thus iron sequestration within these cells. Therefore, hepcidin may modulate plasma iron concentration and transferrin saturation.

The homozygous p.Cys282Tyr mutation of HFE gene is the most frequent etiology of genetic hemochromatosis (GH) linked to an inaccurate low level of hepcidin regarding iron status. The molecular mechanisms linking the p.Cys282Tyr mutation and low hepcidin levels are not fully characterized. Mutations in other genes, including transferrin receptor 2 (TFR2), hemojuvelin (HJV) and ferroportin (FPN), as well as in coding and non-coding regions of hepcidin (HAMP), are known to induce phenotypic presentations similar to HFE related GH. HFE related GH is a disease with incomplete penetrance. This suggests that modifier genes modulate the expressivity of HFE hemochromatosis. Mutations in hepcidin (HAMP), TFR2 or HJV genes have been demonstrated to favor the development of iron overload in p.Cys282Tyr heterozygote patients and to increase iron burden in p.Cys282Tyr HFE homozygotes.

In HFE-GH it can be hypothesized that every genetic factor lowering hepcidin gene expression could lead to a more severe phenotype. Therefore, we searched for mutations in iron metabolism regulatory genes in one patient with homozygous p.Cys282Tyr mutation presenting severe iron overload. We identified a heterozygous mutation located within the recently reported BMP-RE which could influence hepcidin gene expression. In this report, we demonstrate that the nc.-153 C>T hepcidin promoter mutation decreases its transcriptional activity thus underlining the potential impact of mutation in BMP-RE of hepcidin promoter on iron metabolism in humans.
**Design and Methods**

**Patient and genetic studies**

In 1994, the diagnosis of genetic hemochromatosis was established in a 37 year old man with massive hepatic iron overload (450 µmol of iron/g dry liver weight, n<36) involving mainly hepatocytes and associated to liver fibrosis stage 2-3 in Metavir classification. Iron depletive treatment based on weekly 400 mL phlebotomies was initiated. In 1997, the patient was demonstrated to be homozygous for the p.Cys282Tyr mutation. In 2004 (47 years), 468 venesections had been performed, corresponding to the removal of 85g of iron. However, the patient was always exhibiting abnormal iron parameters with 40 µmol/L for serum iron, 100% for transferrin saturation, and 2,066 µg/L for serum ferritin. In addition, hepatic iron concentration remained very high (close to 300 µmol/g dry liver weight) at MRL. Serum hepcidin level quantified by Elisa 19 was low (24 ng/L; 29<N<254 for men) despite major iron overload (10 µg) were preincubated with 250 ng of poly(dI-dC),poly(dI-dC), used as non-specific competitor DNA, in a binding buffer (10 mM Tris-HCl pH 7.5, 50mM NaCl, 1mM DTT, 1 mM EDTA, 5% glycerol) for 10 min on ice. This mixture was then added to the 32P-end-labeled probe either with or without specific competitor oligonucleotides, and the incubation was carried out for a further 30 min at room temperature. Gel shift assay with purified polyclonal rabbit IgG anti-SMAD4 or anti-SMAD1/5/8 (Santa Cruz) or non-relevant antibody were performed under the same conditions except that the antibodies were preincubated with nuclear extracts for one hour on ice. Complexes were resolved by electrophoresis on prerun Tris-Glycine 5% native polyacrylamide gels. The following chemically synthesized double-stranded oligonucleotides were used as probes and as specific competitors (nucleotide mutation is underlined): BMPRE-wt GCCTTTTCCGCGCCACCACC, BMPRE-mut GCCTTTTCCGCTGCCACCCACC.

**Cell culture**

Human hepatoma HepG2 cells were grown in Minimum Essential Medium α (invitrogen) supplemented with 10% fetal bovine serum (invitrogen), 100 UI/mL penicillin, 100 mg/mL streptomycin and 2 mM L-glutamine.

**Plasmid constructions**

The -1024 bp from the translation start site of the human hepcidin promoter were amplified from HuH7 genomic DNA and subcloned into pGL4.17 (Promega) to generate -1024/BMP-RE wt Hep/Luc plasmid construct. The internal mutations in the BMP responsive element or in the STAT3 DNA binding site were made using -1024 Hep/Luc as template and the internal following primers: GCCCTTTTCGGT GCCACCACC; antisense GGTGGTGGCAGCCAAAAGGC for BMP-RE or CACCTCTTGCCGCGTGAAGC, antisense GTCTCAGGCGCAAGAGGTC for STAT3 binding site. The nucleotide mutations are underlined. The identity of the constructs was confirmed by DNA sequencing.

**Transfection, luciferase assay**

Cells were transfected, using transfectine (Bio-Rad) with 100 ng of pGL4-hepcidin promoter vector with a normalization plasmid (pRL-TK, Promega). After 48 h of transfection, luciferase activities were measured in cell lysates using the Dual-Luciferase-Reporter assay system (Promega) and a Centro LB 960 luminometer (Berthold Technologies). For treatments, either 10 ng/mL of human BMP9 or 20 ng/mL of human BMP4 or 20 ng/mL IL-6 (R&D systems) were added to the cells for 48 h.

**Electrophoretic mobility shift assay**

HepG2 cells were incubated with either 10 ng/mL of human BMP9 or 20 ng/mL of human BMP4 for one hour. Then crude nuclear extracts were prepared as described. For gel retardation assays, nuclear extracts (10 µg) were preincubated with 250 ng of poly(dI-dC),poly(dI-dC), used as non-specific competitor DNA, in a binding buffer (10 mM Tris-HCl pH 7.5, 50mM NaCl, 1mM DTT, 1 mM EDTA, 5% glycerol) for 10 min on ice. This mixture was then added to the 32P-end-labeled probe either with or without specific competitor oligonucleotides, and the incubation was carried out for a further 30 min at room temperature. Gel shift assay with purified polyclonal rabbit IgG anti-SMAD4 or anti-SMAD1/5/8 (Santa Cruz) or non-relevant antibody were performed under the same conditions except that the antibodies were preincubated with nuclear extracts for one hour on ice. Complexes were resolved by electrophoresis on prerun Tris-Glycine 5% native polyacrylamide gels. The following chemically synthesized double-stranded oligonucleotides were used as probes and as specific competitors (nucleotide mutation is underlined): BMPRE-wt GCCTTTTCCGCGCCACCACC, BMPRE-mut GCCTTTTCCGCTGCCACCCACC.

**Results and Discussion**

**The heterozygous nc -153 C>T was found in hepcidin promoter**

In genetic hemochromatosis, iron loading results from abnormally low levels of hepcidin linked to mutations in the hepcidin gene itself or in genes involved in hepcidin expression control, such as the HFE, HJV and TFR2 genes.10,11 This prompted us to search for additional mutations in iron regulatory genes in a p.Cys282Tyr homozygous patient presenting a massive iron overload. The sequencing of the HFE gene confirmed the homozygous p.Cys282Tyr mutation and excluded another mutation. In addition, sequencing of HJV, TFR2 and FPN genes were normal. Finally, sequencing of the hepcidin gene led to the identification of a punctual mutation in position -153 from the transcription start site. This mutation was localized within the BMP-RE as recently proposed,17 and was found neither in the web databases nor in 200 chromosomes issued from 100 healthy volunteers.
The heterozygous nc.-153 C>T mutation decreases hepcidin promoter transcriptional activity

As described previously, the BMP-RE controls human hepcidin promoter transcriptional activity under steady state-condition. In order to assess the effect of the heterozygous nc.-153 C>T mutation on basal hepcidin gene expression, plasmid constructs containing the -1024 bp of the human hepcidin promoter with the wild type or the mutated BMP-RE inserted among the luciferase gene (-1024/BMP-RE wt Hep/Luc and -1024/BMP-RE mut Hep/Luc) were transfected in HepG2 cells. Luciferase activity from the -1024/BMP-RE mut Hep/Luc construct was about 2.4 fold reduced compared to the luciferase activity from the plasmid construct containing the wild type BMP-RE (Figure 1, lanes 1-2), demonstrating that this mutation decreased hepcidin transcriptional activity in basal condition. As this BMP-RE mediates the induction of hepcidin gene expression in response to BMP, HepG2 cells were transfected with the -1024/BMP-RE wt Hep/Luc and -1024/BMP-RE mut Hep/Luc plasmid construct, which was arbitrarily set at 100%. Data represent means of at least three independent experiments (± SEM). *p<0.001 compared to the -1024/BMP-RE wt Hep/Luc construct and #p<0.001 compared to the -1024/BMP-RE mut determined by 1-way ANOVA and the Student-Newman-Keuls comparisons test.

The heterozygous nc.-153 C>T mutation impairs the binding of SMAD1/5/8/4 to the BMP-RE

Figure 2. The -1024 BMPRE-wt Hep/Luc but not the mutated constructs responds in a dose-dependent manner to BMP4. HepG2 cells were transfected with the -1024 BMPRE wt or mutated and were treated with increasing amount of BMP4.

These results demonstrate that the heterozygous nc.-153 C>T mutation affects the basal hepcidin gene expression and impairs its BMP4/9 response.

Hepcidin gene responsiveness to IL-6 has been recorded decreasing when the BMP-RE or STAT3 binding sites are mutated or in SMAD4-/- mice. Accordingly, we found (Figure 1, lanes 8-9), by comparing the impact of IL-6 on wild type and mutated promoter, that the nc.153C>T was critical for IL-6 total responsiveness. However, the mutation did not affect the ability to respond in terms of fold induction, by comparing the ratio between basal and IL-6 stimulated mutated construct (Figure1 lanes 2,9) to the ratio between basal and IL-6 stimulated wild type construct. (Figure 1, lanes 1,8).

To test the severity of the mutation we performed a dose dependent assessment of the BMP4 effect (Figure 2). Whereas we found a dose dependent induction of Luciferase activity from the -1024/BMP-RE wt Hep/Luc construct, the BMP9 treatment increased luciferase activity by about 2.7 fold and BMP4 treatment by 1.8 fold (Figure 1, lanes 1,4,6). By contrast, with the -1024/BMP-RE mut Hep/Luc construct, after BMP9 or BMP4 treatment, there was no change observed for luciferase activities compared to the basal condition (Figure 1, lanes 2,5,7). These results suggest that the heterozygous nc.-153 C>T mutation affects the basal hepcidin gene expression and impairs its response to BMP4/9 and the IL-6 total responsiveness.
for SMAD1/5/8 transcription factors, which are phosphorylated upon BMP binding to their receptors and migrate to the nucleus associated to SMAD4, to activate transcription of target genes.\(^1\) In order to determine if the heterozygous nc.-153 C>T mutation impaired the formation of a nucleoprotein complex of SMAD1/5/8 with the BMP-RE, an EMSA was performed using nuclear extracts from HepG2 cells treated or not with BMP4 or BMP9 and probes containing the wild type or mutated BMP-RE. A complex was observed using the probe containing the wild type BMP-RE (Figure 3A, lanes 1,3,5; complex I). This complex I is present to a lower extent in nuclear extracts obtained from basal condition compared to those from BMP stimulated cells. The complex I formation was inhibited by using a probe which contained the mutated BMP-RE (Figure 3A, lanes 2,4,6). The specificity of formation of this complex with the probe containing the wild type BMP-RE was demonstrated using unlabeled oligonucleotides containing wild type or mutated BMP-RE in excess. Complex I binding was competed with the wild type BMP-RE used as cold competitor (Figure 3B, lanes 1-4) whereas competitor with the mutated BMP-RE had no effect (Figure 3B, lanes 5-8), thus demonstrating its binding specificity. The lower band was not affected by the wt competitor, demonstrating its unspecificity.

To identify the protein content of complex I, HepG2 nuclear extracts were incubated with antibodies against a non-relevant protein as control, SMAD4 or SMAD1/5/8. Supershifts were observed with anti-SMAD4 antibody and anti-SMAD1/5/8 antibody specifically blocked the nucleoprotein complex I formation, thus inducing its disappearance. These results suggest that in HepG2 cells, a nucleoprotein complex containing the SMAD1/5/8/4 proteins is observed with the wild type BMP-RE DNA binding site and that this nucleoprotein complex formation is abrogated by the nc.-153 C>T mutation. Thus, the impairment of SMAD1/5/8/4 binding to the nc.-153 C>T mutated BMP-RE could reduce hepcidin gene expression.

Taken together, our data supports the deleterious impact of the nc.-153 C>T hepcidin promoter mutation. The iron depletive treatment (85g of iron subtracted, including 18g potentially absorbed iron during the whole venesection period, according to Olsson et al.\(^2\)) was only partially successful as compared with what can be expected in classical HFE hemochromatosis as well as in juvenil hemochromatosis related to either hepcidin or hemojuvelin gene mutations. These data suggest that: i) the nc.-153 C>T mutation may have an additive effect to the homozygous p.Cys282Tyr mutation, which remains to be characterized and, ii) that the addition of an iron chelator treatment could be relevant both to potentiate iron elimination and to limit venesecion-related iron absorption.

Our overall results suggest that mutations in BMP-RE of hepcidin promoter may alter iron homeostasis, contributing to the increase in the iron burden in HFE related hemochromatosis.

**Authorship and Disclosures**

MLI designed and performed experiments, analyzed data and wrote the manuscript. PB diagnosed the patient and participated in writing the manuscript. AMJ and AM performed sequencing and analysed data. VD and YD coordinated the study in healthy volunteers and participated in writing the manuscript. OL initiated the study, analyzed data and wrote the paper.

The authors reported no potential conflicts of interest.
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