Role of duodenal iron transporters and hepcidin in patients with alcoholic liver disease

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Abstract

Patients with alcoholic liver disease (ALD) often display disturbed iron indices. Hepcidin, a key regulator of iron metabolism, has been shown to be down-regulated by alcohol in cell lines and animal models. This down-regulation led to increased duodenal iron transport and absorption in animals. In this study, we investigated gene expression of duodenal iron transport molecules and hepcidin in three groups of patients with ALD (with anaemia, with iron overload and without iron overload) and controls. Expression of DMT1, FPN1, DCYTB, HEPH, HFE and TFR1 was measured in duodenal biopsies by using real-time PCR and Western blot. Serum hepcidin levels were measured by using ELISA. Serum hepcidin was decreased in patients with ALD. At the mRNA level, expressions of DMT1, FPN1 and TFR1 genes were significantly increased in ALD. This pattern was even more pronounced in the subgroups of patients without iron overload and with anaemia. Protein expression of FPN1 paralleled the increase at the mRNA level in the group of patients with ALD. Serum ferritin was negatively correlated with DMT1 mRNA. The down-regulation of hepcidin expression leading to up-regulation of iron transporters expression in the duodenum seems to explain iron metabolism disturbances in ALD. Alcohol consumption very probably causes suppression of hepcidin expression in patients with ALD.

Keywords: DMT1  FPN1  DCYTB  HEPH  TFR1  HFE  hepcidin  alcoholic liver disease  iron  gene expression

Introduction

Iron metabolism disturbances are common findings in patients with chronic alcohol consumption. It ranges from anaemia to iron overload [1–3]. The pathogenesis of anaemia in alcoholic liver disease (ALD) is complex; it includes hypersplenism with splenic pooling and increased destruction of erythrocytes, blood loss because of gastrointestinal bleeding as a complication of alcoholic cirrhosis, which can lead to anaemia with iron deficiency [4–8]. Inadequate diet with nutrient deficits, in alcoholic patients, can result in anaemia with megaloblastic and sideroblastic features [9]. Additionally, anaemia of chronic disease can occur in patients with chronic ALD [10, 11]. Conversely, some alcoholics develop iron overload [1, 3, 12]. It has been documented that chronic alcohol consumption in moderate to excess amounts leads to elevated serum ferritin concentration and transferrin saturation, and can result in increased hepatic iron stores [13]. Additionally, increased intestinal iron absorption has been observed in patients with chronic alcoholic disease [14]. Both iron and ethanol individually cause oxidative stress and lipid peroxidation and the cumulative effects of ethanol and iron on liver cell damage, in patients with ALD, exacerbate liver injury [15–17].

As there is no physiological way of eliminating excess iron, iron homoeostasis is regulated primarily by iron absorption. The process of intestinal non-haeme iron uptake starts with the reduction in ferric iron to the ferrous form by, the brush border enzyme, ferrireductase DCYTB (CYBRD1, duodenal cytochrome b reductase 1) [18]. Iron is then transported across the apical membrane of duodenal enterocytes

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by means of the divalent metal ion transporter 1 (DMT1, SLC11A2, NRAMP2, DCT1) [19, 20]. From the enterocyte, iron is exported across the basolateral membrane by ferroportin (FPN1, SLC40A1, IREG1, MTP1) [21–23]; a ferroxidase haemochromatin (HEPH) oxidizes iron to its soluble and non-reactive ferric state, which is then delivered to transferrin [24]. HFE, responsible for hereditary haemochromatosis, is another regulatory protein associated with iron metabolism [25]. HFE affects the interaction of transferrin-bound iron with the transferrin receptor (TFR1, TFRC). Binding of HFE to TFR1 lowers its affinity for iron-transferrin, resulting in a reduction of cellular iron uptake [26, 27]. HFE forms complexes also with transferrin receptor 2 (TFR2) [28]. The HFE:TFR2 complex is thought to serve as an iron sensor that regulates hepcidin expression [29]. Hepcidin, a key hormone in the regulation of iron metabolism [30], is produced mainly in the liver. Hepcidin appears to act via binding to and internalization of ferroportin and its subsequent degradation [31, 32]; although, evidence that hepcidin inhibits apical uptake via DMT1 is also available [33]. Hepcidin was found to be up-regulated by iron overload and down-regulated by iron deficiency anaemia and hypoxia [30, 34–36]. It has been documented that hepcidin is decreased in patients with haemochromatosis [37–39]. In addition to its response to iron homeostasis, hepcidin is induced by inflammation [30]. Recently, with the use of animal models, ethanol was shown to down-regulate the expression of hepcidin in the liver which resulted in elevated expression of the iron transporters DMT1 and ferroportin in the duodenum [40, 41]. Deregulation of hepcidin synthesis may be one of the causes of iron disturbances during chronic alcohol consumption.

So far, the effect of ethanol on iron uptake via duodenal iron transporters and its relation to hepcidin have only been analysed using cell lines and animal models [40–43]. Therefore, the aim of this study was to evaluate the expression of duodenal iron transporters both on mRNA and protein levels and their relation to hepcidin in alcoholic patients either with anaemia, iron overload or normal iron stores.

Materials and methods

Patients

A total of 54 individuals (35 male, 19 female), mean age of 57.4 years, ranging from 25 to 82 years were enrolled in the study. The diagnosis of ALD (N = 24) was based on patients’ history of consumption of more than 30 g alcohol per day, presence of elevated serum AST (aspartate aminotransferase, EC 2.6.1.1) or ALT (alanine aminotransferase, EC 2.6.1.2) and GGT (gamma-glutamyltransferase, EC 2.3.2.2) activity and sonographically observed fatty changes in the liver (liver steatosis). According to laboratory parameters, ALD patients were categorized into three subgroups: ALD with anaemia (N = 8), ALD with iron overload (N = 6) and ALD without overload (N = 10). First, patients were divided according to the presence of anaemia (haemoglobin levels <11 g/dl). These patients had minor decreases in serum iron parameters, however, they did not meet criteria for iron deficiency anaemia (serum ferritin <20 μg/l, haemoglobin <11.0 g/dl and transferrin saturation <16%). Patients without anaemia were then divided according to the presence of iron overload, defined as elevated ferritin levels (cut-off: 200 μg/l for women and 250 μg/l for men) or increased transferrin saturation (cut-off = 45%). The control group (N = 30) had an upper GI (gastrointestinal) endoscopy to evaluate dyspeptic symptoms and their iron parameters were in normal ranges (serum iron 11–26 μmol/l, serum ferritin male 30–250 μg/l, female 30–200 μg/l, transferrin saturation 20–45%). Controls were participants of another study by our group concerning the gene expression in haemochromatosis [44]. To analyse the effect of HFE gene mutations, the genotyping for C282Y, H63D and S65C mutations of HFE gene was performed using the PCR-RFLP method, as described previously [45]. DNA for HFE genotyping was available from 23 ALD patients and from 10 controls. Patients were recruited at our outpatient department between 2005 and 2009. Informed consent was obtained from all patients and the study was approved by the Ethics Committee of the Third Faculty of Medicine, Charles University and conducted in accordance with the Helsinki Convention.

Sample collection

Duodenal biopsy samples were obtained from 54 individuals during GI endoscopy. For RNA analysis, samples were stored at −20°C in RNAlater solution (Sigma-Aldrich, St. Louis, MO, USA) prior to RNA isolation and for protein analysis at −80°C prior to protein isolation.

RNA isolation and real-time quantitative polymerase chain reaction

Total RNA was isolated from RNAlater-stored duodenal biopsies by using an RNAeasy MiniKit (Qiagen, Hilden, Germany), and included DNase digestion according to manufacturer’s instructions. After estimation of RNA integrity by using gel electrophoresis and determination of each sample concentration, one sample was found to be inadequate for further analysis. The following analyses were carried out as previously described in detail [44]. Briefly, cDNA synthesis was performed using a reverse transcription kit TaqMan Reverse Transcription Reagents (Applied Biosystems, Foster City, CA, USA) with random primers according to the manufacturer’s instructions. Real-time quantitative PCR was performed using an ABI Prism 7000 Sequence Detection System (Applied Biosystems) and commercially available kits: Taq Man Universal PCR Master Mix and Sybr green PCR Master Mix (Applied Biosystems). For the amplification of FPN1, TFR1, HFE and GAPDH cDNA, Applied Biosystems pre-designed gene expression assays were used: FPN1 – Hs00205888_m1, TFR1 – Hs00174609_m1, HFE – Hs00373474_m1 and GAPDH – Hs99999905_m1. For amplification of DMT1, DCYTB and HEPH cDNA, previously described analyses were used [46]. DMT1 analysis was performed on the DMT1(IRE) variants because these are the isoforms that are regulated by iron status in cell [47]. All data were normalized to the amount of GAPDH cDNA in the sample. The 2−ΔΔCt method was used to calculate relative changes in gene expression.

Western blot analysis

Because of the limited amount of biological material, Western blot analyses were performed only on duodenal iron transporters and coupled oxido-reductases (DMT1, ferroportin, DCYTB and hephaestin). After protein extraction from duodenal biopsy by using the RIPA buffer (Sigma-Aldrich) and determination of concentration, seven samples were found to be inadequate for further analysis with Western blot. A
small amount of tissue, obtained by biopsy, was detected in two samples; therefore, the Western blot analysis could only be performed for hephaestin and DMT1 detection in these samples. Western blot analyses of DMT1 (IRE variants), DCYTB, ferroportin, hephaestin and actin (loading control) levels were performed using goat polyclonal antibodies NRAMP 2 and Hphaestin against human DMT1 and hephaestin (Santa Cruz Biotechnology, Santa Cruz, CA, USA), goat polyclonal anti-Cytochrome b reductase 1 antibody against human DCYTB (Everest Biotech, Upper Heyford, UK), rabbit polyclonal antibody MTP11-A against human ferroportin (Alpha Diagnostic International, San Antonio, TX, USA) and mouse monoclonal antibody against human actin (Sigma-Aldrich). Analysis was carried out as previously described [44]. Briefly, proteins separated using SDS-PAGE were blotted onto a 0.2 μm nitrocellulose membrane for 2 hrs at 0.25 A, by using a Mini-Protein II blotting apparatus (Bio-Rad, Hercules, CA, USA). The membrane for DMT1, ferroportin, DCYTB and hephaestin was blocked with 5% BSA in TBS (100 mM Tris-HCl, 150 mM NaCl, pH 7.5), whereas, for actin, 5% non-fat milk in TBS was used. The washed membrane was incubated with the relevant primary antibody. After incubation (overnight, 4°C), the washed membrane was incubated for 2 hrs with the corresponding horseradish peroxidase-conjugated secondary antibody, which was then detected by using enhanced chemiluminescence (overnight, 4°C). Bands intensities were quantified by densitometry and the data were analysed by using ImageJ software (version 1.42q; NIH, USA, available on http://rsb.info.nih.gov/ij/).

**ELISA**

To evaluate the role of hepcidin, a subgroup of 24 individuals, whose serum was available for hepcidin analysis, was created. It represented samples from eight controls and 16 ALD patients, which were further divided into the following subgroups: ALD with anaemia (N = 8), without overload ALD (N = 6), and ALD with iron overload (N = 2). Mature bioactive hepcidin was measured in serum samples by using a commercial ELISA (EIA-4705) kit (DRG International Inc., Springfield, NJ, USA) according to the manufacturer’s instructions. In both patients and controls, blood sample was drawn between 7:30 and 8:30 a.m. after overnight fasting.

**Statistical analysis**

Data are expressed as mean ± SEM. Because of non-normality of the measured variables, the non-parametric Mann–Whitney test or the Kruskal–Wallis ANOVA with multiple comparison tests were used as needed. Correlations were assessed using the Spearman rank method. P value less than 0.05 was considered significant. Statistical analyses were performed using the Statistica program (Version 9; StatSoft, Tulsa, OK, USA) and the GraphPad Prism program (Version 5.00; GraphPad Software, Inc., San Diego, CA, USA).

**Results**

**Clinical and laboratory characteristics of patients**

Laboratory parameters of controls and ALD group are shown in Table 1. DNA genotyping showed that among ALD patients, one was found to be a C282Y/H63D compound heterozygote, one was a H63D homozygote, three were H63D heterozygotes and one was a S65C heterozygote. Tested controls were wild-type for all three tested mutations. In further analyses, gene expression values of individuals with HFE gene mutations were not outliers or extremes and were within the non-outlier range of values measured in their respective subgroups.

**RNA expression**

Gene expression at the mRNA level was measured for DMT1, FPN1, DCYTB, HEPH, HFE and TFR1. A significant increase in the ALD group was found when DMT1, FPN1 and TFR1 were examined (2.51-fold, \( P = 0.0147 \), 1.54-fold, \( P = 0.0342 \), and 2.02-fold, \( P = 0.0011 \), respectively). Gene expression of the other tested molecules (DCYTB, HEPH and HFE) was not significantly different in ALD patients compared to controls (Fig. 1A).

To study the ALD group in more detail, we divided the group into three subgroups: (i) patients with the signs of iron overload, (ii) patients without iron overload and (iii) patients with anaemia according to parameters defined in Materials and Methods. A significant increase in DMT1 and FPN1 mRNA was found in the ALD subgroup without iron overload compared to ALD with iron overload.

| Variable                        | ALD (N = 24) | Controls (N = 30) | \( P \)-value |
|---------------------------------|--------------|-------------------|--------------|
| **Age (years)**                 | 57.38 ± 1.57 | 57.40 ± 2.85       | 0.4431       |
| **Gender**                      | 17M/7F       | 18M/12F           | 0.5673       |
| **Serum iron (μmol/l)**         | 18.03 ± 1.94 | 17.55 ± 0.67       | 0.6240       |
| **Serum ferritin (μg/l)**       | 296.22 ± 96.11 | 151.0 ± 13.83   | 0.9640       |
| **Transferrin saturation (%)**  | 32.22 ± 4.88 | 31.22 ± 0.96       | 0.4455       |
| **Hb (g/dl)**                   | 12.66 ± 0.44 | 14.00 ± 0.31       | 0.0138       |
| **Ht (%)**                      | 37.16 ± 1.16 | 41.48 ± 0.93       | 0.0058       |
| **ALT (μkat/l)**                | 0.62 ± 0.05  | 0.54 ± 0.05        | 0.0478       |
| **AST (μkat/l)**                | 0.81 ± 0.09  | 0.47 ± 0.03        | 0.0003       |

Data are presented as arithmetic mean ± SEM. Hb: haemoglobin; Ht: haematocrit; ALT: alanine aminotransferase; AST: aspartate aminotransferase. Normal ranges: serum iron (11–26 μmol/l), serum ferritin (male 30–250 μg/l, female 30–200 μg/l), transferrin saturation 20–45%, Hb (male 13.0–18.0 g/dl, female 11.5–16.0 g/dl), Ht (male 38–54%, female 35–47%), ALT (0.1–0.75 μkat/l) and AST (0.1–0.75 μkat/l).
overload (3.02 versus 0.34, \(P = 0.0312\) and 1.72 versus 0.70, \(P = 0.0420\), respectively). Moreover, a significant elevation in DMT1 and TFR1 mRNA expression was observed in ALD patients without iron overload compared to controls (3.02-fold, \(P = 0.0479\), and 1.91-fold, \(P = 0.0066\), respectively). Although FPN1 gene expression was also increased compared to controls, the increase did not reach statistical significance (1.72-fold, \(P = 0.0599\)). When the ALD subgroup with anaemia was examined, a significant increase in DMT1, FPN1, DCYTB and HEPH mRNA levels was observed compared with controls (3.50-fold, \(P = 0.0018\), 1.93-fold, \(P = 0.0115\) and 2.84-fold, \(P = 0.0075\), respectively). The same applies to comparison of the ALD anaemia subgroup to the ALD iron overload patients when DMT1 and FPN1 mRNA were analysed (3.50 versus 0.34, \(P = 0.0007\) and 1.93 versus 0.70, \(P = 0.0080\), respectively). Additionally, the HFE mRNA level was elevated in ALD patients with iron overload compared with controls (1.40-fold, \(P = 0.0376\)) (Fig. 2A–F).

### Protein expression

To investigate whether the changes in mRNA expression correlated with changes in protein expression, we examined DMT1, ferroportin, DCYTB and hephaestin levels by using Western blotting.

There was a significant increase in ferroportin protein expression in ALD patients compared with controls (1.76-fold, \(P = 0.0495\)). Unchanged DCYTB and HEPH mRNA expression was paralleled by unchanged respective protein expression (1.20-fold, \(P = 0.8292\) and 0.96-fold, \(P = 0.8898\), respectively). DMT1 protein expression was not different in the ALD group compared with controls (1.03-fold, \(P = 0.3879\)) (Fig. 1B). Further analysis of the ALD group, divided based on iron overload or anaemia, revealed no significant differences in protein expression. The ferroportin protein expression had the same pattern as seen at the mRNA level; however, the significance found in the ALD group was lost...
Fig. 2 Gene expression of the analysed molecules in controls, patients with alcoholic liver disease without iron overload (ALD without overload), with iron overload (ALD overloaded), and anaemia (ALD anaemia). (A) mRNA expression of DMT1, (B) FPN1, (C) DCYTB, (D) HEPH, (E) TFR1, (F) HFE. (G) Protein expression of DMT1, (H) FPN1, (I) DCYTB, (J) HEPH. Results are depicted as means ± SEM. Statistically significant differences are indicated by *P < 0.05, **P < 0.01 and ***P < 0.001.
and the increase in ferroportin protein in ALD patients without iron overload and anaemia did not reach statistical significance compared with controls (1.90-fold, \( P = 0.0746 \) and 2.08-fold, \( P = 0.1685 \), respectively; Fig. 2G–J).

**Relationships among gene expression of tested molecules**

Relationships among DMT1, FPN1, DCYTB, HEPH, TFR1 and HFE expression were analysed by using the Spearman rank correlation. When all individuals regardless of underlying disease were evaluated, there was a positive association among mRNA expression of the tested molecules. The strongest correlation was found between DMT1 and FPN1 \( (r = 0.778, \ P < 0.0001) \), between DCYTB and HEPH \( (r = 0.687, \ P < 0.0001) \) and between DMT1 and TFR1 \( (r = 0.660, \ P < 0.0001; \) Table 2).

When the gene expression, at the protein level, in the cohort of all individuals was investigated, a positive relationship between the following proteins was found: DCYTB and ferroportin \( (r = 0.632, \ P < 0.0001) \), DMT1 and hephaestin \( (r = 0.580, \ P < 0.0001) \), ferroportin and hephaestin \( (r = 0.482, \ P = 0.0008) \), DCYTB and hephaestin \( (r = 0.391, \ P = 0.0080) \) and DMT1 and DCYTB \( (r = 0.314, \ P = 0.0358; \) Table 3).

When all individuals were analysed together, a correlation between gene expression at the mRNA level and the protein level was found only for hephaestin \( (r = 0.338, \ P = 0.0215) \).

When correlations in ALD and controls were investigated separately, positive relationships were found, and are summarized in Tables 2 and 3.

### Table 2 Associations among analysed molecules—mRNA level

|       | HFE    | TFR1   | HEPH   | DCYTB  | FPN1   |
|-------|--------|--------|--------|--------|--------|
| **DMT1** |        |        |        |        |        |
| ALD   | ns     | 0.613**| ns     | ns     | 0.863***|
| Controls | ns   | 0.634***| ns     | ns     | 0.683***|
| All   | ns     | 0.660***| ns     | ns     | 0.778***|
| **FPN1** |        |        |        |        |        |
| ALD   | ns     | 0.593**| ns     | ns     |        |
| Controls | ns   | 0.523**| 0.593***| 0.364**|
| All   | ns     | 0.591***| 0.447***| 0.322* |
| **DCYTB** |        |        |        |        |        |
| ALD   | 0.600**| ns     | 0.570**|        |        |
| Controls | ns   | ns     | 0.790***|        |        |
| All   | 0.468***| ns     | 0.687***|        |        |
| **HEPH** |        |        |        |        |        |
| ALD   | ns     | ns     |        |        |        |
| Controls | 0.414*| ns     |        |        |        |
| All   | 0.399**| 0.315*|        |        |        |
| **TFR1** |        |        |        |        |        |
| ALD   | ns     |        |        |        |        |
| Controls | 0.604***|        |        |        |        |
| All   | 0.452***|        |        |        |        |

Spearman rank correlation, only statistically significant data are shown, statistically significant differences are indicated by * \( P < 0.05 \), ** \( P < 0.01 \) and *** \( P < 0.001 \), ns - not significant.
Spearman rank correlation, only statistically significant data are shown. Significance levels indicated by * P < 0.05, ** P < 0.01 and *** P < 0.001, ns - not significant.

### Table 3: Associations among analysed molecules—protein level

| Proteins   | DMT1 | DCYTB | FPN1 | Hepcidin |
|------------|------|-------|------|----------|
| HEPH       |      |       |      |          |
| ALD        | 0.792*** | 0.471* | ns   |          |
| Controls   | 0.490*  | ns    | ns   |          |
| All        | 0.580*** | 0.314* | ns   |          |
| Hepcidin   |      |       |      |          |
| ALD        | ns    | ns    | ns   |          |
| Controls   | ns    | ns    | ns   |          |
| All        | ns    | ns    | ns   |          |
| FPN1       |      |       |      |          |
| ALD        | 0.428*  | 0.719*** |      |          |
| Controls   | 0.580** | 0.601** |      |          |
| All        | 0.482*** | 0.632*** |      |          |
| DCYTB      |      |       |      |          |
| ALD        | 0.465*  |      |      |          |
| Controls   | ns     |      |      |          |
| All        | 0.391** |      |      |          |

### Discussion

In this study, iron metabolism in alcoholic patients with normal iron indices, iron overload and anaemia was examined. Gene expression of molecules participating in iron absorption in duodenum (DMT1, ferroportin, DCYTB and hephaestin) was analysed both at mRNA and protein levels. Additionally, TFR1 and HFE mRNA expression was investigated. Serum hepcidin, a key regulator of iron metabolism, was evaluated as well. This study is, to our knowledge, the first to investigate duodenal gene expression of iron transport molecules both at the mRNA and protein level, together with the serum hepcidin level in ALD patients.

The effect of alcohol consumption on hepcidin expression has been previously demonstrated [40, 41]. It has been shown, by using animal models, that alcohol down-regulates hepcidin expression,
which affects ferroportin and DMT1 and leads to increased iron absorption in the duodenum. It has been documented that ethanol down-regulates hepcidin promoter activity and DNA binding activity of transcription factor C/EBPα [40, 41]. In the present study, hepcidin levels were decreased in ALD patients compared with controls, supporting the abovementioned mechanism of alcohol on hepcidin expression in humans. We found a significant elevation of FPN1 at both the mRNA and protein level and an increase in DMT1 and TFR1 mRNA in ALD patients. However, the elevation of DMT1 mRNA was not paralleled by the elevation of DMT1 protein. It could be speculated that some post-transcriptional mechanisms are involved e.g. IRE/IRP system [48], Ndfips/WWP2 system responsible for ubiquitination and degradation of DMT1 [49], or PAP7 which inhibition was shown to cause a reduction in the expression of DMT1[IRE] protein but not mRNA [50]. The regulation by miRNA can also be considered but the only miRNA documented to affect DMT1 acts on the non-IRE variant [51]. Also, the heterogeneity of patients’ genetic background or the small sample size can be the reason of this discrepancy. Further analysis of the ALD subgroups showed that at the mRNA level, there was a significant increase in DMT1 and FPN1 in ALD patients without iron overload compared with ALD with iron overload and a significant increase in DMT1 and TFR1 mRNA when compared with controls.

Additionally, hepcidin levels in these subgroups were significantly lower compared with controls. Thus, it can be hypothesized that in ALD patients, chronic alcohol consumption decreases hepcidin levels, which results in up-regulation of DMT1 and FPN1 mRNA in patients with yet normal iron indices. It could be expected that when iron stores become increased, the iron burden is sensed to be inappropriately high by the ‘iron status’ hepcidin regulatory pathway (probably HJV/BMP and/or HFE/TFR2 pathway) and decreased hepcidin levels tend to normalize to compensate iron absorption. This would lead to suppression of increased DMT1 and FPN1 mRNA levels in ALD patients with iron overload as seen in our study. However, we did not detect this normalization of hepcidin levels in the ALD iron overload subgroup, probably because of limited number of serum samples; thus we can only speculate at this point. It can be argued that these two factors—alcohol, which tends to decrease hepcidin expression and physiological regulation, which tends to compensate for high iron stores by increasing hepcidin expression, act on various regulatory pathways and this interaction finally establishes some sort of balance; an analogous situation has been documented in treated and untreated haemochromatosis patients [52–54]. This is in agreement with another study where the effect of alcohol together with iron was investigated [42]. In the study, animal models were used and the results showed that iron elevated and alcohol decreased liver hepcidin expression. Alcohol was shown to suppress up-regulation of hepcidin mRNA in iron-overloaded rats to levels similar to those in control animals. The duodenal ferroportin expression was elevated in alcohol-treated mice. When both factors were investigated together duodenal ferroportin expression was increased and reached levels between alcohol-treated and iron-treated mice. In addition, the iron-induced increases in the DNA binding activity of C/EBPα were diminished by alcohol to levels found in controls. Similar observations using HFE knockout mice were documented in another study [43], which also suggested that alcohol decreases hepcidin expression independently of the HFE pathway, possibly by alcohol-induced hypoxia.

We also investigated ALD patients with anaemia. The pathogenesis of anaemia in ALD is complex: it may include splenic pooling and haemolytic anaemia caused by hypersplenism; chronic bleeding into the gastrointestinal tract resulting in iron deficiency; secondary malnutrition, leading to anaemia with folic acid deficiency; anaemia as a consequence of the direct toxic effect of alcohol on erythrocyte precursors in bone marrow [10]. These symptoms lead to hypoxia, anaemia and iron deficiency, which all inhibit hepcidin synthesis via several pathways and corresponding mediators: hypoxia inducible factor α, erythropoietin (EPO), growth differentiation factor 15 and twisted gastrulation protein homologue 1. In iron deficiency, the regulation pathway for iron status includes activity of the BMP/HJV and/or HFE/TFR2 pathway [55, 56]. On the other hand, anaemia of chronic disease can occur in patients with chronic ALD [10], when hepcidin expression is induced by inflammatory stimuli (e.g. IL-6) [55, 56]. Finally, with chronic alcohol consumption, the effect of ethanol on hepcidin promoter activity and the DNA binding activity of transcription factor C/EBPα must also be considered [41]. All these factors may have played a role in our ALD patients with anaemia and the final level of hepcidin represented the combined effect of all these various pathways. We also found decreased hepcidin level in our ALD patients...
with anaemia, which was consistent with the abovementioned facts. The effect of increased hepcidin synthesis because of anaemia of chronic disease seems to be minor, especially when the erythroid demand for iron is thought to be a more powerful regulator of hepcidin expression than inflammation [57]. Consequently, these patients displayed increased mRNA expression of DMT1 and FPN1 in the duodenum, demonstrating the efforts of the organism to increase iron absorption in response to iron needs associated with enhanced erythropoiesis. However, these elevations were not detected at the protein level of DMT1 and FPN1. Although the expression of FPN1 protein was increased, it did not reach statistical significance. Surprisingly, serum hepcidin levels were similar in ALD patients with anaemia (when the effect of anaemia plus ethanol affects hepcidin synthesis) and without anaemia (when only ethanol is implicated). We were not able to distinguish between the effect of erythropoietic stimulation, iron deficiency and ethanol when analysing the combined effect of all factors playing a role in the pathogenesis of anaemia in our ALD patients. It seems that the effect of various mediators on hepcidin expression pathway is not simply additive, and crosstalk among the different pathways, can be hypothesized, to influence different signal transduction intensities. On the other hand, EPO has been proposed to act directly to repress hepcidin through EPOR-mediated regulation of C/EBPα [58], which is also affected by ethanol. It could be possible that this pathway can suppress hepcidin synthesis, but only to some limit; and once the C/EBPα is influenced by a mediator, others are unable to exert their full influence. However, the small sample size of our associated ALD subgroup did not permit definite conclusions in this respect.

DMT1 mRNA was increased in ALD patients with anaemia and without iron overload. The increase of DMT1 in ALD patients without overload cannot be explained by iron deficiency and anaemia as in the ALD anaemia subgroup [59], thus the effect of ethanol needs to be considered. It has been shown that ethanol exposure can increase the expression of DMT1 in hepatocytes [60, 61]. The increase in DMT1 expression is partially because of the increased activity of iron regulatory proteins (IRPs) linked to the oxidative stress of ethanol metabolism [60]. It can be speculated that a similar mechanism is implicated in duodenal cells; however, this mechanism has yet to be elucidated. In addition, if ethanol can affect DMT1 mRNA expression in duodenal cells it may also play a role in the expression of other iron transport molecules.

We detected a strong positive correlation between DMT1 and FPN1; DCYTB and HEPH; and DMT1 and TFR1 at the mRNA level, which suggests a coordinated regulation of these genes. We also investigated the association between serum iron parameters and duodenal iron transporters. Significant inverse correlations between DMT1 mRNA and serum ferritin, in controls and ALD patients, were detected confirming that body iron stores play a role in the regulation of duodenal expression of this transporter. With regard to controls, our observations are in agreement with several studies [54, 62]; however, there were other studies that did not find any correlation at all [52, 53].

In conclusion, our results demonstrate that serum hepcidin levels are decreased by alcohol consumption resulting in increased expression of iron transporter ferroportin at the mRNA and protein level and DMT1 at the mRNA level in duodenum of ALD patients. Detailed analyses revealed that these changes were observed in ALD patients without iron overload and ALD patients with anaemia. The increase in duodenal TFR1 mRNA expression is a consequence of anaemia and probably also an effect of ethanol on duodenal cells. Positive correlations among DCYTB, HEPH, DMT1, FPN1 and TFR1 mRNA indicate coordinated regulation of these genes. Further research is required to elucidate the complex pathogenesis of ALD and the effect of ethanol and oxidative stress derived from its metabolism in duodenal cells deserves further investigation.

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Conflicts of interest

The authors confirm that there are no conflicts of interest.

Author contribution

Dostalikova-Cimburova M., Kratka K., Kovar J., and Horak J. designed the research study; Dostalikova-Cimburova M., Balusikova K., Chmelikova J., and Neubauerova J. performed the research (performed molecular analyses); Kratka K., Hejda V., Hnanicek J. performed the research (examined patients, collected the clinical data and biological material); Dostalikova-Cimburova M. and Vranova J. analysed the data; Dostalikova-Cimburova M. wrote the paper; Kratka K., Balusikova K., Chmelikova J., Hejda V., Hnanicek J., Neubauerova J., Vranova J., Kovar J., and Horak J. critically revised the manuscript.

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