Hereditary hemochromatosis: study of laboratory alterations related to polymorphism

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INTRODUCTION

Hereditary hemochromatosis (HH) is a genetic, autosomal recessive disorder characterized by the overload of iron in tissues. The HH gene (HFE) is located on the short arm of chromosome 6, a major histocompatibility complex (MHC). The protein expressed by this gene is related to the control of intestinal iron absorption: interacts with the transferrin receptor (TIR) detecting its degree of saturation, thus providing signaling to the enterocyte whether there is a greater or lesser need for intestinal iron uptake.

The HH classification is performed according to the genetic alteration found and the gene associated with iron overload. Classical HH is associated with HFE gene mutations (homozygosis for C282Y or compound heterozygosis for C282Y/H63D). Mutations in this gene, for example, have an inadequate mechanism of hepcidin production (systemic iron-regulatory hormone) in response to iron overload.

Three allelic variants of the HFE gene were associated with HH: C282Y, significantly associated with HH; H63D and S65C. In the first case, there is a single mutation at position 282, in which cysteine is replaced by tyrosine; in the second, the mutation was identified at position 63, where histidine is replaced by aspartate; in the third, the mutation results from the substitution of cysteine for serine at position 65.

The frequency of C282Y and H63D mutants is still the subject of studies in the Brazilian population, but it is known that they are present in 2/3 of the Brazilian patients with HH, probably suggesting other mutations in the HFE gene or that other genes may be involved in the regulation of iron metabolism.
those of the northern European studies, where the mutations are less frequent\(^{(2,5,7-9)}\).

The southern region of the country, with Caucasian origin, should present different frequencies for the C282Y mutation compared to the population of the Northeast region, for example, of predominantly African origin\(^{(10)}\). Therefore, the present study aims at clarifying the possible link between HH-associated polymorphisms and laboratory tests, identifying the \textit{HFE} gene mutations in the study population, relating them to the disease, as well as evaluating the association between C282Y, H63D and S65C polymorphisms and different biochemical and hematological markers in patients attended at a laboratory in Caxias do Sul, Rio Grande do Sul, Brazil.

**METHODS**

We studied the medical records of 283 patients who had their blood tests collected in a laboratory in the city of Caxias do Sul (RS), from January 2010 to January 2013. Patients who performed the genotyping for HH-associated genes, among other conventional laboratory tests, were selected, thus indicating that these patients were under specific investigation for this pathology.

The inclusion criteria for the study were: to present a molecular biology test for HH, to be older than 18 years of age and not having positive serologic markers for hepatitis B and C. The sample was selected by the annual report filed in the laboratory information system. Patients with negative HH genotyping were considered as controls. From these 283 patients, 11 were excluded from the results due to the presence of hepatitis B and C markers reactive to the diseases, which could lead to a false result in the relationship between HH and biochemical tests.

The analysis of the reports was performed through the system used by the data grantor. Results of the following tests were collected:

- molecular biology — polymerase chain reaction (PCR) for detection of mutations\(^{(11,12)}\) H63D, C282Y and S65C;
- biochemical tests — lactate dehydrogenase (LDH), iron (Ir), gamma-glutamyl transferase (GGT), glucose (GLU), aspartate aminotransferase (AST) and alanine aminotransferase (ALT) performed by Cobas Integra 400 plus device, that uses absorption photometry technique;
- immune exams — hepatitis B surface antigen (HbsAg), antibodies to hepatitis C virus (HCV) and ferritin (FER) performed by Cobas 6000 device, that uses electrochemiluminescence method. Transferrin saturation index (TSI) was performed by the supporting laboratory using the immunoturbidimetry method;
- hematometric exams: red blood cells (RBC), hemoglobin (Hb), hematocrit (Hct), mean corpuscular volume (MCV), mean corpuscular hemoglobin concentration (MCHC) and total leukocytes (LEU) performed by flow cytometry method in the Sysmex-XE 2100 device.

After collection, patients with complete laboratory data were selected and the information was stored in a database constructed by the researcher. Statistical analysis was performed in the SPSS 20.0 program. The frequency of genotypes present in the population and the average of the biochemical parameters were evaluated and the Chi-square test was used to verify the significance of the relationship between the biochemical and hematological markers and the genetic mutations. Values of \(p < 0.05\) were considered significant.

The present study follows the ethical requirements for research and was approved by the Research Ethics Committee of the Associação Cultural e Científica Virvi Ramos, on 18 August 2015, under the number of opinion 1.181.585.

**RESULTS**

From the 272 records analyzed, 19.1% (\(n = 52\)) were female patients and 80.9% (\(n = 220\)), were male patients. The mean age was 55.74 years, with minimum age of 23 years and a maximum of 85 years.

The Table shows the frequency of genotypes in the study records, and the highest percentage is 54.4% (\(n = 148\)) of normal patients, that is, without the presence of genotype mutations for HH, followed by H63D/WT 31% (\(n = 84\)) and C282Y/WT 6.2% (\(n = 17\)) genotype. Only medical records with complete laboratory data were analyzed, and FER levels could be significantly increased (\(p < 0.05\)) throughout the study population – all with values above 300 ng/ml [reference values (RV): men (M): 12-300 ng/ml and women (W): 12-150 ng/ml] —, even in patients without mutation in the \textit{HFE} gene.

The group of homozygote for C282Y mutation was the only one that presented TSI values compatible with the diagnosis for HH, where TSI ≥ 55% (RV: 20%-45%) and serum FER concentration ≥ 500 ng/ml (RV M: 12-300 ng/ml and W: 12-150 ng/ml). In this group it is also possible to verify the increase in the other biochemical tests, compatible with the present mutation, but this data was not statistically significant due to the reduced number of patients with this mutation (1.1% – three patients).
The hematimetric indexes were evaluated in all study groups, and we observed an increase in MCV levels in the C282Y/C282Y mutation carriers (mean: 99.17 fl/RV: 82-98 fl) and the mean corpuscular hemoglobin (MCH) in patients with H63D/C282Y genotype (mean: 37.79 pg/RV: 28-33 pg), but these data were not statistically significant. The other hematimetric parameters were within the range of normality.

In the Figure, the data demonstrate the elevated FER levels according to the study groups.

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In the Figure, the data demonstrate the elevated FER levels according to the study groups.

<table>
<thead>
<tr>
<th>Genotypes</th>
<th>Incidence</th>
<th>%</th>
<th>GGT</th>
<th>GLU</th>
<th>TSI</th>
<th>AST</th>
<th>ALT</th>
<th>LDH</th>
<th>Ir</th>
<th>FER</th>
<th>RBC</th>
<th>Hb</th>
<th>Hct</th>
<th>MCV</th>
<th>MCH</th>
<th>MCHC</th>
<th>LEU</th>
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<tr>
<td>Normal</td>
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<td>54.4</td>
<td>64.39</td>
<td>97.98</td>
<td>39.08</td>
<td>30.04</td>
<td>45.89</td>
<td>174</td>
<td>134</td>
<td>731</td>
<td>4.9</td>
<td>15.07</td>
<td>44.34</td>
<td>90.58</td>
<td>30.69</td>
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<td>102.26</td>
<td>50.64</td>
<td>26.73</td>
<td>37.6</td>
<td>165.7</td>
<td>122.4</td>
<td>813.81</td>
<td>4.85</td>
<td>14.9</td>
<td>43.64</td>
<td>90.02</td>
<td>30.63</td>
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<td>42.32</td>
<td>29.67</td>
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<td>166.37</td>
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<td>39.6</td>
<td>73.4</td>
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<td>-</td>
<td>-</td>
<td>152.5</td>
<td>296.8</td>
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<td>45.2</td>
<td>94</td>
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<td>25</td>
<td>153</td>
<td>155.5</td>
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<td>33.33</td>
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</tbody>
</table>

Reference values: GGT: M 10-50 U/l, W 7-32 U/l; GLU: 70-99 mg/dl; TSI: 20%-45%; AST: < 40 U/l; ALT: < 40 U/l; LDH: 100-190 E4/V18; Ir: 30-160 micrograms/dl; FER: M 12-300 ng/ml, W 12-150 ng/ml; RBC: M 5.3 ± 3 g/dl, W 13.6 ± 2.5 g/dl; Hb: 12-15 g/dl; Hct: M 47 ± 7, W 42 ± 6; MCV: M/W 82-98 fl; MCH: M/W 31-36 g/dl; MCHC: M/W 30.5-35 g/dl; LEU: 5,000-10,000.

GGT: gamma-glutamyl transferase; GLU: glucose; TSI: transferrin saturation index; AST: aspartate aminotransferase; ALT: alanine aminotransferase; LDH: lactate dehydrogenase; Ir: iron; FER: ferritin; RBC: red blood cells; Hb: hemoglobin; Hct: hematocrit; MCV: mean corpuscular volume; MCH: mean corpuscular hemoglobin; MCHC: mean corpuscular hemoglobin concentration; LEU: total leukocytes; M: man; W: woman.

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Records from a laboratory in the city of Caxias do Sul (RS), surveyed for the diagnosis of HH, show a high number of male patients in relation to the number of women. Women have an advantage over men regarding the overload of iron, since they have the monthly loss of this element through menstruation and also in the gestational period. Although women benefit from the physiological loss of iron through menstruation, both men and women are affected by the disease, the ratio is 4 to 10:1(1).

Most cases of HH that have northern European origin are associated with HFE(13). Caxias do Sul is a city that officially began with European colonization, with Italian immigrants and people from other European countries. Even so, HH not associated with HFE, although rare, is found in several parts of the world, regardless of race.

The hematimetric indexes found here are expected and, because of the FER levels in these patients, they are unlikely to have iron-deficiency anemia, as well as the increased MCV and MCH values in the C282Y/C282Y and H63D/C282Y mutation carriers, as also demonstrated by other biochemical markers.

Patients without mutation for the study genotypes were the majority (54.4%), but even in those patients that did not have HH mutations, FER values were elevated, with mean value of 731 ng/ml. Elevated FER may be linked to several factors, such as liver diseases, acute and chronic infection, B12 and folic acid deficiency, and excessive alcohol consumption(14, 15). Thus, the HFE gene mutation seems to be a necessary condition, but not a requirement for the development of the disease, evidencing the theory of Beutler (2003)(16).

Cançado et al. (2007)(9) found a frequency of the HFE gene mutation in patients with iron overload of 76%, higher than the frequency found in the present study. The most prevalent mutation
in the present study was H63D/WT (30.9%), corroborating other Brazilian studies that also showed a higher frequency of this mutation\(^{(17, 18)}\). These patients are at low risk of developing the disease; they have the heterozygous gene and may present an increase in FER and iron levels, as observed in this study. The other markers such as AST and ALT, which indicate the level of iron toxicity caused by HH, are not altered.

Patients with C282Y/C282Y genotype, admitted in studies as the most critical for the development of the disease, presented biochemical markers values altered, however with data not statistically significant. The presence of other genotypes should be considered for HH if iron overload is also diagnosed. In patients who presented these mutations, there was increase of serum iron (not compatible with HH), but they may develop the disease in the future according to their living habits.

C282Y mutation of HFE gene presents a wide variation, which will depend fundamentally on the study population and the criteria used in the selection of each sample and in the diagnosis of iron overload. The frequency of mutation in patients with histological diagnosis of iron overload in the studies of Altes et al. (2003)\(^{(19)}\) and Brandhagen et al. (2000)\(^{(20)}\) was 16% and 85%, respectively. In the present study, it was the genotype that presented higher concentrations of serum FER and iron, besides saturation of the major transferrin, values compatible with the diagnosis for HH. These results confirm the correlation between C282Y/C282Y genotype and increased risk of iron overload\(^{(19)}\).

When the selection of the sample occurs by the histological diagnosis of iron overload, the frequency of patients with C282Y/C282Y genotype is higher when the sample is assessed on the biochemical parameters. The estimated percentage of C282Y/C282Y patients who will develop laboratory evidence of iron overload ranges from 40% to 70%\(^{(16, 21)}\), thus evidencing the patient’s predisposition to develop HH when the genetic mutation is present\(^{(16)}\).

Among studies performed in Brazil, Bittencourt et al. (2002)\(^{(22)}\) showed that C282Y and H63D mutations are present in about two thirds of Brazilian individuals with HH, in the sample we observed 53% of patients with C282Y/C282Y genotype and 13.5% with C282Y/WT or H63D/WT. On the other hand, Bueno et al. (2006)\(^{(22)}\) reported that 62% of patients with HH diagnosis had abnormal HFE, 37.5% homozygous for C282Y, 12.5% compound heterozygous (C282Y/H63D) and 12.5% heterozygous for C282Y/WT mutation. The results differ from those found in the present study, thus showing that there is also a wide variation in the frequency of mutations among the samples evaluated in the country.

Individuals homozygous for H63D mutation (H63D/H63D) and heterozygotes (C282Y/H63D) have a low phenotypic expression rate\(^{(21, 22)}\). The S65C mutation implies disease usually when compound with C282Y (C282Y/S65C)\(^{(16, 22)}\), and, likewise the mutations above mentioned, has a low rate of phenotypic expression, which corroborates the results found in the present study.

It was not possible to evaluate the association between biochemical, immunological and hematimetric markers with all genotypes, since the alterations were not statistically significant, most likely because the stage of the disease has not yet led to these laboratory alterations. On the other hand, it is clear the presence of genetic mutations in the study population, and early investigation should be performed in any and all evidence of iron overload in these patients to prevent the disease itself from occurring, often leading to irreversible damage to the body.

**CONCLUSION**

Data from the present study suggest that elevated FER values are associated with HH genotypes, indicating the importance of analyzing this marker in asymptomatic patients. It was also observed that the gene indicated as the most severe for the involvement of the disease, C282Y/C282Y, had a low frequency in the study population.

The correlation of the genotypes with the other markers was not statistically significant. Probably the laboratory changes are not yet noticeable due to the present stage of the disease. It is possible to observe a significant amount of HH-related genotypes in the research sample for the disease, thus showing a high frequency of this pathology, and the need of laboratory follow-up of these patients, as well as that of their relatives, for the purpose of preventing consequences of iron overload that may be triggered by external factors.

**ACKNOWLEDGEMENTS**

Thanks to Doctor Cleber Cremonese for his contribution in this work, and in particular to the Unimed Nordeste (RS) laboratory for making the company’s database available.
RESUMO

Introdução: A hemocromatose hereditária (HH) é uma doença hereditária associada ao armazenamento em excesso de ferro. Sua forte presença é subestimada por ser assintomática nos pacientes. Na cidade de Caxias do Sul (RS), por conta da maior frequência da doença, é necessário realizar um estudo que relaciona os genótipos HH com as análises laboratoriais. Objetivo: Avaliar a associação dos polimorfismos C282Y, H63D e S65C e dos indicadores laboratoriais dos casos estudados. Métodos: O estudo foi realizado com base na análise de registros médicos de 283 pacientes atendidos no período de janeiro de 2010 a janeiro de 2013, maiores de 18 anos, os quais realizaram o protocolo de genotipados para exames HH e laboratoriais. A análise de indicadores entre o genótipo e os exames laboratoriais foi realizada pelo software SPSS 20.0. Resultados: As alterações genéticas mais frequentes foram H63D/WT 84 (31%) e C282Y/WT 17 (6,2%). O genótipo que apresentou alteração mais compatível com a HH foi o C282Y/C282Y, mesmo ele não tendo demonstrado resultados significativos. O aumento da ferritina foi ligeiramente maior (p < 0,05) nas grupos estudados, estando todos acima do valor de referência. Observou-se nos genótipos C282Y/C282Y e H63D/C282Y aumento insignificante no volume corpuscular médio (VCM) e na hemoglobina corpuscular média (HCM). Conclusão: De acordo com o estudo, foi possível observar um aumento significativo da ferritina em todos os genótipos HH.

Unitermos: hemocromatose; polimorfismo genético; genótipo.

REFERENCES

