Association Between the Phenotypes of Haptoglobin and Tuberculosis in Ivory Coast-Haptoglobin Phenotypes and Tuberculosis

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Abstract
The susceptibility of patients to certain pathologies, such as tuberculosis (TB), is associated to the phenotype of their haptoglobin (Hp). The objective of this study was to investigate the prognostic value of the haptoglobin phenotype in tuberculosis by determining the association of Hp phenotypes with certain epidemiological and clinical characteristics of tuberculosis in Côte d’Ivoire. In a case-control study, 131 tuberculosis and 109 non-tuberculosis as controls, voluntary blood donors were recruited in Abidjan. From venous blood samples, phenotyping of Hp was performed by polyacrylamide gel electrophoresis according to Raymond’s method. Comparisons were made using ch2 test at risk α = 5%. We found three phenotypes: Hp1-1, Hp1-2, Hp2-2 in the respective proportions of 27.5%, 50.5% and 22% in the control population (n = 109) and 36.6%, 54.2% and 9.2% in the tuberculosis population (n = 131). Among the population carrying the Hp1 allele, 58.3% were tuberculosis patients compared to 41.7% in controls subject (p = 0.006). Among TB patients, 33.3% carried Hp2-2 subtype compared to 66.7% in controls (p = 0.011). The Hardy-Weinberg’s equilibrium showed that tuberculosis patients carrying Hp2-2 phenotype died early. Hp phenotype was not associated to TB-HIV co-infection, neither to TB treatment nor to response to anti-tuberculosis treatment. We concluded that there is an association between Hp phenotype and TB infection prognosis. Hp2-2, less antioxidant seemed to be associated to the disease with poor prognosis.

Keywords: haptoglobin, phenotype, tuberculosis, Ivory Coast

1. Introduction
Haptoglobin (Hp) is a glycoprotein with antioxidant and immune modulatory properties in humans (Franck et al., 2001; Mark & David, 2016). Three major phenotypes are found (Hp 1-1, Hp 2-1 and Hp 2-2) (Langlois & Delanghe 1996; Yano, Yamamoto, Miyaishi & Ishuzu, 1998), each of which has a functional specificity that can have a different impact on the prognosis of certain pathologies (Franck et al., 2001; Sadzadeh & Bozorgmehr., 2004; Mark & David, 2016). Hp binds with hemoglobin (Hb) and both form an Hp-Hb complex that plays an important role in the defense of the organism; while iron is necessary for bacterial growth, it’s binding by the Hp-Hb complex reduces its extra-erythrocyte availability and slows down the bacterial growth (Easton, Brandt, Mahoney & Lee., 1992; Barclay, 1985; McDemid & Prentice, 2006; Mark & David, 2016).

A clinical-epidemiological specificity was reported based on the populations (Franck et al., 2001, Sadzadeh &Bozorgmehr, 2004). Few studies reported the causes of these variations, particularly those inherent in ethnogenic factors of the host. Therefore, the susceptibility of patients to certain pathologies including tuberculosis (TB) could be associated to the phenotype of their Hp. Studies have shown that tuberculosis subjects with Hp2-2 phenotype have an increased risk of mortality and increased susceptibility to developing severe renal tuberculosis compared to other phenotypes (Fedoseava, Lusopova, Chukanova & Pospelov, 1993; Kasvosve et al., 2000). The prevalence of tuberculosis in Côte d’Ivoire is around 661 cases / 100 000 inhabitants (Programme national de lutte contre la tuberculose [PNLT], 2012) and also has specific features (PNLT, 2012). In this study, we sought to determine the
association between Hp phenotypes and certain epidemiical and clinical characteristics of TB in Côte d'Ivoire.

2. Material and Methods

2.1 Type of Study
We performed a case-control study comparing the distribution and characteristics of Hp phenotypes in tuberculosis and healthy subjects.

2.2 Population and Study Environment

The study population were TB patients (beginning and ending) receiving treatment at the Anti- Tuberculosis Center of Treichville (ATCT) and voluntary blood donors from the National Blood Transfusion Center (CNTS). For each of the patients included, the survey card containing the variables of the study was filled, followed by a venous blood sampling on a tube containing lithium heparinate. Phenotyping of Hp were carried out at the Center for Diagnosis and Research on AIDS and other infectious diseases (CeDReS). Thus, we recruited 131 TB patients receiving treatment at CATT Abidjan and 109 voluntary blood donors from the CNTS.

2.3 Method of Analysis

The variables in this study were: Hp phenotype, sex, age, smear, clinical status (x-ray of the lungs), response to anti-tuberculosis treatment and presence of HIV / AIDS co-infection. Patient’s data were obtained by history and from the respective medical records. The biological specimen was whole blood obtained through venipuncture on a heparinized tube. The plasma obtained after centrifugation was used for the phenotyping of Hp by electrophoresis of the Hp-Hb complex in 5% polyacrylamide gel (PAGE) (Raiymond, 1962).

2.4 Consent and Ethical Approval

The study was conducted with the authorization from the National Health authorities and the administrative head of the health center. We obtained an ethical clearance from the board of physicians of the center. All authors declare that ‘written informed consent was obtained from the patient for publication of this paper and accompanying images.

2.5 Statistical Analysis

The data collected was entered using SPSS Software v16.0. The proportions of Hp phenotypes obtained were expressed as percentage and compared between the two study populations: tuberculosis patients and healthy subjects. The statistical analysis was carried out using the same software and the comparisons were made by the Chi –Square test. The distribution of alleles in the study populations was evaluated using Hardy Weinberg's equilibrium. The significance threshold was set at 5%.

3. Results

3.1 Demographics Characteristics

In this study, we recruited 240 subjects comprising of 109 non-tuberculosis to serve as controls subjects and 131 tuberculosis patients. The 109 controls included 81.7% male and 18.3% female. Their mean age was 34.4 ± 8.2 years with extremes of 18 and 50 years. The tuberculosis population comprised 75.6% men and 24.4% women, their mean age was 36.10 ± 11.83 years with the extremes of 16 and 74. This predominance of male subjects was statistically significant (p = 0.009).

3.2 Haptoglobin Phenotyping

Three Hp phenotypes were found in the following proportions: Hp1-1 (37.5%), Hp2-1 (50.5%) and Hp2-2 (22%) in controls and Hp1-1 (36.6%), Hp2-1 (54.2%) and Hp2-2 (9.2%) in tuberculosis patients. The Hp1 allele represented 40.7% and 59.3% against 52% and 48% for the Hp2 allele in controls and tuberculosis patients respectively. So, subjects with Hp 2-2 were predominantly observed in the control population whereas those with Hp 1-1 and 2-1 phenotypes were observed among the tuberculosis subjects (p = 0.014) (Table 1). Thus, Hp1 allele was predominantly found in tuberculosis patients (p = 0.005) while Hp2 was more frequent in the control population (p = 0.010) (Table 1). The distribution of these alleles in the population of the tuberculosis subjects was not in the Hardy-Weinberg’s equilibrium (X² = 17.53, p <0.0001); however, in the control population it was in the Hardy-Weinberg’s equilibrium (X² = 0.52, p = 0.48).

3.3 Tuberculosis Infection Location, Treatment and HIV Co-Infection

Concerning the localization of tuberculosis infection, we found 126 pulmonary TB and 05 extra-pulmonary TB. There was no significant association between the Hp phenotype and the smear microscopic analysis result (p = 0.051) (Table 2). High positive smear results (1-3 plus) were found in the Hp1-1 and 2-1 phenotype groups (p = 0.011) (Figure 1).

We did not find any significant link regarding the Hp phenotype either with the treatment lines or with the frequency of TB-HIV co-infections (Table 3).
4. Discussion

This study has enabled us to establish the distribution of Hp phenotypes and Hp 1 and 2 alleles in healthy and tuberculosis populations in Côte d’Ivoire.

Three Hp phenotypes were found. The Hp profiles observed were similar to those reported by Sukaina et al (2015) and Philemon et al (2017); but different from those of Constans, Viau, Gouaillard, and Clerc. (1981), that reported, in addition to these 3 phenotypes, phenotypes Hp0-0 and Hp2-1M in Côte d’Ivoire. Bagat et al. (2015) found Hp0-0 phenotype in many Indians populations. The phenotype Hp0-0 is present in subjects who have anhaptoglobinemia (KoDH et al., 2013). When comparing our results to those of Sutton et al (1959), they agree on the phenotype profile of Hp (3 phenotypes), but according to that team, the Hp1 allele was 70% in Ivorian and Liberian populations, that means a higher frequency than what is observed in our study. These changes in the distribution of Hp phenotypes and alleles in Côte d’Ivoire could be explained, due to several displacements and mingle up of the populations since the first studies performed in 1958. It is therefore necessary to renew the study of Hp profile in the Ivorian population on a larger population sample recruited in several regions.

The subtype Hp 2-2 were predominantly observed in the control population whereas subtypes Hp 1-1 and 2-1 phenotypes were more frequent in the tuberculosis subjects (p = 0.014). The Hp1 allele was predominantly found in tuberculosis patients (p = 0.005) and Hp2 allele was more common in the controls population (P = 0.010). These two results indicated the same trend which was similar to those of Kaminskaia, Abdullaev, Elufimova, Mitinskaia, and Iukhimenko (2004) and Philemon et al. (2017). However, they differed from what reported by many previous studies in which the Hp2-2 phenotype is less protective and therefore more expected to be found in patients than in controls (Langlois & Delanghe, 1996; McDermid & Prentice, 2006; Kasvosve et al., 2000).

The distribution of these alleles in the control population respected the Hardy-Weinberg’s equilibrium (X² = 0.52, p = 0.48), which expresses the random nature of this distribution. Failure to observe this equilibrium in the tuberculosis population (X² = 17.53, p <0.0001) proved the non-random nature of this distribution. This genetic drift in tuberculosis patients could be explained by the increased mortality of tuberculosis among carriers of Hp2-2 phenotype. This result indicated that, Hp2-2 phenotype was less in tuberculosis patients because the prognosis is worse for them and may be they died early than those who have phenotype 1-2 and 1-1. This result could be confirmed by the results of Kaminskaia et al. (2004) who found no carrier of Hp 2-2 phenotype in TB pediatric patients with tuberculous pleurisy. The appearance of pleurisy is a sign of severity of tuberculosis. The carriers of the phenotype Hp 2-2 do not therefore reach this stage; hence their absence in this group.

Concerning the localization of the tuberculosis infection, we found 126 cases of Pulmonary TB and 05 Extra-pulmonary TB. There was no significant association between the phenotype of Hp and the smear microscopic examination result (p = 0.051). The p-value very close to the threshold of statistical significance deserves to be confirmed on a larger sample population or on a population of heterogeneous tuberculosis from several sites in order to favor a random distribution of the Hp alleles. This result disagrees with those of the Kasvosve et al. (2000) study on tuberculosis patients High positive (1 to 3 cross) microscopies were found in groups of Hp1-1 and 2-1 phenotypes but this was not statistically significant.

We did not find any significant link in the Haptoglobin phenotype with neither the treatment lines, nor with the frequency of TB-HIV co-infections. Similar results were reported (Zaccarioto et al., 2006).

Nevertheless, we have not obtained enough studies of less than 5 to 10 years on this theme.

5. Conclusion

Three phenotypes of Hp were found in Côte d’Ivoire: Hp1-1, Hp2-1 and Hp2-2. There was an association between the phenotype of Hp and the frequency of TB infection. Thus, the predominant distribution of the Hp1 allele in tuberculosis subjects was not random and Hp2-2, less antioxidant was associated with TB poor prognosis. The presence of allele Hp² seemed to be associated with a high rate on mortality of TB. The Hp phenotype was not associated with microscopy, HIV-TB co-infection, tuberculosis treatments or response to these treatments.
Table 1. Distribution of Hp phenotypes and alleles in the study population

<table>
<thead>
<tr>
<th>Phenotype of Hp</th>
<th>CASE (n=131)</th>
<th>CONTROL (n=109)</th>
<th>p-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-1</td>
<td>48 (36,6%)</td>
<td>30 (37,5%)</td>
<td></td>
</tr>
<tr>
<td>2-1</td>
<td>71 (54,2%)</td>
<td>55 (50,5%)</td>
<td>0,014</td>
</tr>
<tr>
<td>2-2</td>
<td>12 (9,2%)</td>
<td>24 (22,0%)</td>
<td></td>
</tr>
<tr>
<td>Alleles</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hp1</td>
<td>119 (90,8%)</td>
<td>85 (78,0%)</td>
<td>0,005</td>
</tr>
<tr>
<td>Hp2</td>
<td>12 (9,2%)</td>
<td>24 (22,0%)</td>
<td>0,010</td>
</tr>
</tbody>
</table>

n =population, % = percentage

Table 2. Distribution of microscopic results in the study population

<table>
<thead>
<tr>
<th>Microscopy</th>
<th>Phenotype of Hp</th>
<th>p-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1-1</td>
<td>2-1</td>
</tr>
<tr>
<td>Positive</td>
<td>9 (31,0%)</td>
<td>14 (48,3%)</td>
</tr>
<tr>
<td>Negative</td>
<td>39 (38,2%)</td>
<td>57 (55,9%)</td>
</tr>
</tbody>
</table>

Figure 1. Analysis of diagnostic smear according to Hp phenotypes

BARR: bacilles acido-alcool résistants; CM=Microscopic field; Negative: No BARR/CM; Rare: <1 BARR/CM ;1+: 1 à 9 BARRS/CM; 2+: 10 à 99 BARRS/CM ;3+: ≥100 BARRS/CM

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Table 3. Distribution of clinical-therapeutic characteristics between Hp phenotypes

<table>
<thead>
<tr>
<th>Therapy [n (%)]</th>
<th>Hp phenotypes</th>
<th></th>
<th></th>
<th>p-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1-1</td>
<td>2-1</td>
<td>2-2</td>
<td></td>
</tr>
<tr>
<td>Line 1 (RHZE / RH ; 6 months)</td>
<td>36 (37,9%)</td>
<td>52 (54,7%)</td>
<td>7 (7,4%)</td>
<td>0,502</td>
</tr>
<tr>
<td>Line 2 (SRHZE / RHZE / RH ; 8 months)</td>
<td>12 (33,3%)</td>
<td>19 (52,8%)</td>
<td>5 (13,9%)</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>48 (36,6%)</td>
<td>71 (54,2%)</td>
<td>12 (9,2%)</td>
<td></td>
</tr>
<tr>
<td>FALLEN BACK OR FAILURE [n (%)]</td>
<td>12 (34,3%)</td>
<td>18 (51,4%)</td>
<td>5 (14,3%)</td>
<td>0,470</td>
</tr>
<tr>
<td></td>
<td>36 (37,5%)</td>
<td>53 (55,2%)</td>
<td>7 (7,3%)</td>
<td></td>
</tr>
<tr>
<td>TB-HIV CO-INFECTION [n (%)]</td>
<td>7 (24,1%)</td>
<td>19 (65,5%)</td>
<td>3 (10,3%)</td>
<td>0,283</td>
</tr>
<tr>
<td></td>
<td>41(40,2%)</td>
<td>52 (51,0%)</td>
<td>9 (8,8%)</td>
<td></td>
</tr>
</tbody>
</table>

R: Rifampicine, H: Isoniazide, E: Etambutol, Z: Pyrazinamide, S: Streptomycine

References


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