HELICOBACTER PYLORI GENOTYPES AMONG BELARUS PATIENTS WITH GASTRODUODENAL DISORDERS AND THEIR ASSOCIATION WITH CLINICAL OUTCOME

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The aim of this study was to evaluate the prevalence of Helicobacter pylori genotypes (vacA and cagPAI) directly in gastric biopsy specimens in patients with gastric diseases in Belarus. Gastric biopsies were collected from 461 patients with different gastrointestinal disorders: superficial gastritis (287 subjects), atrophy gastritis (59 subjects), erosive gastritis (47 subjects), duodenal ulcer disease (54 subjects), and stomach ulcer (14 subjects). PCR-based genotyping was used to detect s1a, s1b, s2, m1a, m1b, m2, cagM, cagA, and cagT genes. Overall prevalence of vacA s1a allele was 60.5% followed by m2 (47.1%) and m1a (37.5%). The analysis of data showed that genotype s1a/m1a was significantly more prevalent in patients with duodenal ulcer (21.4% vs. 45.1%, OR = 3.0, 95% CI = 1.5–6.1). The cagA gene was found with a high incidence in most patients with inflammatory diseases of stomach and duodenum. There was a significant increase in the frequency of cagT in patients with duodenal ulcer as compared to superficial gastritis. A high cagM prevalence was found in patients with atrophy gastritis and duodenal ulcer disease. All three island genes of pathogenicity of cagPAI are more often detected in patients with duodenal ulcer, which increases the risk of developing duodenal ulcer by 4.5 times.

Keywords: Helicobacter pylori genotypes, clinical outcome, cagPAI

Introduction

Helicobacter pylori (HP) is possibly the most widespread human pathogen that causes of gastritis, peptic ulceration, and gastric cancer. According to World Health Organization statistics, HP is reported as a class 1 carcinogen in gastric adenocarcinoma [1].

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However, only a small part of infected people develop disease, and the disease outcome is influenced by a combination of host, bacterial, and environmental factors [2]. It is important to determine the most prevalent genotypes in a specific region of human population to find a correlation between putative virulence genes and clinical outcome. In areas of high HP infection prevalence, this information may potentially help us to predict the patient’s disease course.

HP is a slow-growing Gram-negative curved or S-shaped rod with a single tuft of multiple-polar flagella and characteristic terminal bulb. The organism does not form spores but, under adverse conditions, undergoes transformation from spiral to coccal morphology often accompanied by a loss of culturability on conventional media. The organism is fastidious and cultivation in vitro requires a microaerophilic environment and complex media.

The core and pan genomes of HP were studied by Cao et al. [3] that compared 75 HP genomes and found an open and diverse core genome with 1,173 conserved protein families and 673 conserved protein domains in the Helicobacter genus.

Approximately 89% of the predicted open-reading frames are common to different strains, confirming HP as a single species. A region containing 45% of HP strain-specific open-reading frames, termed the plasticity zone, is present on the chromosomes, verifying that some strain variability exists. The plasticity zone in HP exhibits great variability [4].

According to the literature, HP studies have focused on two groups of bacterial virulence factors, such as the cag pathogenicity island (cagPAI) and the vacuolating cytotoxin vacA. The vacA cytotoxin is a well-known HP virulence factor with multiple cellular effects. Bacterial strains have revealed that the vacA gene differs in its signal sequence (s1a, s1b, s1c, or s2) and in its mid-region sequence (m1 or m2). Its virulence function has been implicated in the pathogenesis of peptic ulceration and a number of gastroduodenal disorders [5–7].

The cagPAI of HP is a ~40 kb region of chromosomal DNA that contains ~27 Cag proteins, whose part of them encodes components of a bacterial type-IV secretion system (T4SS). One of the most studied putative virulence factors is the CagA protein, encoded by the cagA gene. Some HP strains contain only parts of the cagPAI and not the whole set of 27 genes [8]. The combination of cagPAI genes can also have an effect on the HP virulence and the clinical outcome.

The aim of this study was to evaluate the prevalence of HP genotypes (vacA and cagPAI) directly in gastric biopsy specimens in patients with gastric diseases in Belarus.
Material and Methods

Gastric biopsies were collected from 461 patients (average age of 42.9 years, 269 women and 192 men) with different gastrointestinal disorders in the medical center “Nordin” (Minsk, Belarus) from 2013 to 2015. All patients signed an informed consent form.

The specimens were examined for the presence of HP by polymerase chain reaction (PCR). Based on endoscopic and histological findings, 287 patients presented superficial gastritis (SG), 59 had atrophy gastritis (AG), 47 had erosive gastritis (EG), 54 had duodenal ulcer disease (DU), and 14 patients were diagnosed to have stomach ulcer (SU).

Total DNA was extracted from frozen gastric biopsy specimens by DNA-sorbC (Apmlisens, Russia).

HP genotyping

A 267-bp fragment of the 23S rRNA gene of HP was amplified using primers HP-F and HP-R [9].

Gene amplification by PCR was prepared in a volume of 25 µl containing 1 x PCR buffer, 3 mM of MgCl₂, 0.2 mM of deoxynucleoside triphosphate, 0.5U of Taq polymerase, 10 pmol of each primer, 1.5 µl of DNA, and sterile distilled water. PCR amplifications were performed in an automated thermal cycler (Thercyc, Russia). All runs included a negative DNA control consisting of PCR grade water.

PCR-based genotyping was used to detect s1a, s1b, s2, m1a, m1b, m2, cagM, cagA, and cagT genes (Table I) using previously published primers [10, 11].

Subsequently, DNA fragments were separated by electrophoresis on 1.2% agarose gel, stained with ethidium bromide, and visualized under UV transilluminator. Standards of 50 or 100 bp (Life Technologies, Lithuania) were used as molecular size markers.

Statistical analysis

The $\chi^2$ test was applied to estimate the statistical differences between disease and various genotypes. The $p$ value $<0.05$ was considered as significant statistical differences.
Results

Studied population

A total of 461 patients were studied [192 men (41.6%) and 296 women (58.4%)], aged between 22 and 84 years with mean age of 43.0 ± 0.7 years. Analysis showed that 64.9% of patients were HP-infected (41.8% of men vs. 58.2% of women).

The histological data exhibited a significantly higher degree of inflammation (78%) and neutrophil activity (50.3%) in HP-positive patients ($p < 0.05$).

The clinical examination shows (Table II) that, among HP-positive, 11% patients have AG, 57.8% have SG, 10.4% have EG, 17% have DU, and 3.7% have SU. The analysis shows a significant increase in the number of people with DU among HP-positive patients; the risk of the disease increases by 10.8 times ($p < 0.05$, 95% CI = −3.2 to 44.5).

Figure 1 shows the comparison of the frequency of HP infection between groups depending on age and clinical outcome.

Table I. Primers used for HP genotyping by PCR

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer sequence</th>
<th>Size of the amplified product (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HP</td>
<td>F: 5′-AGGTATAGGGTGTCCTCAGTC-3′ &lt;br&gt;R: 5′-CGCATTATTTCCATTAGCAG-3′</td>
<td>267</td>
</tr>
<tr>
<td>s1a</td>
<td>F: 5′-CTGTTCTAATGCGGCAAC-3′ &lt;br&gt;R: 5′-CTGCGTATCAGGCAAC-3′</td>
<td>190</td>
</tr>
<tr>
<td>s1b</td>
<td>F: 5′-AGCGCATTACCGGCAAC-3′ &lt;br&gt;R: 5′-CTGCTTGAATGCGGCAAC-3′</td>
<td>187</td>
</tr>
<tr>
<td>s2</td>
<td>F: 5′-GCTAACAGCGCAATGATCC-3′ &lt;br&gt;R: 5′-CTGCTTGAATGCGGCAAC-3′</td>
<td>199</td>
</tr>
<tr>
<td>m1a</td>
<td>F: 5′-GGCTCAAATGCGGTCATGG-3′ &lt;br&gt;R: 5′-CTGTTAGTGCCGCCAGAAC-3′</td>
<td>290</td>
</tr>
<tr>
<td>m1b</td>
<td>F: 5′-CGTCTCTTGCTAGCAG-3′ &lt;br&gt;R: 5′-GCTGTTAGTGCCCTAACAGCAT-3′</td>
<td>295</td>
</tr>
<tr>
<td>m2</td>
<td>F: 5′-AGAGCCACGAGAACATTTG-3′ &lt;br&gt;R: 5′-CTAATACTACCTTTCAGC-3′</td>
<td>352</td>
</tr>
<tr>
<td>cagA</td>
<td>F: 5′-AACACCGCACTTACAGC-3′ &lt;br&gt;R: 5′-TTTGTTGGCTTTCAGC-3′</td>
<td>400</td>
</tr>
<tr>
<td>cagM</td>
<td>F: 5′-AATACTACAAAAGAGGAAGG-3′ &lt;br&gt;R: 5′-ATTATTTCACAAAGTTAGAAGGAGGGCC-3′</td>
<td>586</td>
</tr>
<tr>
<td>cagT</td>
<td>F: 5′-CCATGTTTATACCGCTGT-3′ &lt;br&gt;R: 5′-CATACCACACCCCTTTTGT-3′</td>
<td>301</td>
</tr>
</tbody>
</table>

Note: HP: Helicobacter pylori; PCR: polymerase chain reaction; F: forward; R: reverse.
Among the patients less than 50 years of age, SG dominated \((p < 0.01)\) regardless of \(HP\) status (significantly higher than patients >50 years of age, \(p < 0.01\)).

In the group of \(HP\)-negative patients with AG and EG, an increase in the number of people over the age of 50 years was found. In group with AG in the

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**Table II.** The frequency of \(HP\) infection between groups depending upon age and clinical diagnosis

<table>
<thead>
<tr>
<th>Groups of patients</th>
<th>(HP) detection</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Negative (%)</td>
</tr>
<tr>
<td>Men</td>
<td>41.4</td>
</tr>
<tr>
<td>Women</td>
<td>58.6</td>
</tr>
<tr>
<td>Age</td>
<td></td>
</tr>
<tr>
<td>&lt;50 years</td>
<td>59.9</td>
</tr>
<tr>
<td>&gt;50 years</td>
<td>40.1</td>
</tr>
</tbody>
</table>

**Morphological diagnosis**

<table>
<thead>
<tr>
<th></th>
<th>(HP)-positive patients in different clinical outcome groups depending upon age. SG: superficial gastritis; AG: atrophy gastritis; EG: erosive gastritis; DU: duodenal ulcer; SU: stomach ulcer. *Significant in comparison with (HP)-negative group, (p &lt; 0.05).</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SG</strong></td>
<td><strong>AG</strong></td>
</tr>
<tr>
<td>HP -</td>
<td></td>
</tr>
<tr>
<td>&lt;50 year</td>
<td></td>
</tr>
<tr>
<td>&gt;50 year</td>
<td></td>
</tr>
</tbody>
</table>

*Significant in comparison with \(HP\)-negative group, \(p < 0.05\).
presence of HP infection, an increase in the number of patients less than 50 years of age is observed.

Thus, the presence of HP is a risk factor for developing AG in younger age people. Among HP-positive patients, a significant increase in the percentage of people with EG and DU in a group less than 50 years is observed. In the group of HP-negative patients with gastric ulcer, an increase in the number of individuals less than 50 years was not detected. In the detection of HP, the incidence of gastric ulcer in this group was 45.5%.

Pathogenicity genes detection frequency in HP-positive patients’ gastric biopsies

The presence of the pathogenicity genes was examined in all 299 HP-positive biopsies from patients with gastrointestinal diseases (Figure 2).

The predominant vacA genotype in the biopsies was the s1a (60.5%), followed by m2 (47.1%) and m1a (37.5%). The HP vacA s1a/m1a genotype was predominantly detected in our samples (28%). Genotypes s1a/m2 and s2m2 were almost equally present, i.e., 24% and 21%, respectively. In this study, the rare s2m1a and s2/m1b genotypes were detected in 4% and 1.3%, respectively.

The cagA gene was observed in 202 of 299 (67.6%) biopsies, and cagT and cagM detected with frequencies of 47.5% (142/299) and 35.8% (107/299), respectively. A significant correlation between the simultaneous presence of cagA, cagM, and cagT genes was found ($p < 0.0001$).

![Figure 2](image_url)

Figure 2. Distribution of genotypes for all HP-positive samples

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The most of cagA-positive samples possessed the s1a form of vacA gene (73.3%), whereas among cagA-negative samples, only 14.8% contained s1a vacA. The analysis showed that combination of three cagPAI genes (cagA, cagM, and cag T) was found in 30.4% samples.

**Relationship between HP genotypes and clinical outcomes**

The genotypes of HP found in patients with different gastrointestinal diseases are presented in Table III.

Statistical comparative analysis of the frequency of each allele in the different form of disease revealed an association between allele s1a and the presence of AG (p < 0.05, OR = 3.7, 95% CI = 1.4–10.0), EG (p < 0.05, OR = 2.4, 95% CI = 1.0–6.1), and DU (p < 0.02, OR = 3.2, 95% CI = 1.5–7.1). The vacA s1b genotype was detected only in 12% of 299 patients: 9.1% with AG, 17.3% with SG, 3.2% with EG, and 7.8% with DU.

The association of middle vacA regions with clinical outcomes shows that m1a allele was more frequently identified in patients with DU (53%). There were no differences in m1b allele between all clinical groups, whereas m2 was the most predominant genotype in patients with SG and EG.

**Table III. Association of HP genotypes with clinical diagnosis**

<table>
<thead>
<tr>
<th>Allele/gene</th>
<th>Clinical diagnosis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>AG [n (%)]</td>
</tr>
<tr>
<td>s1a</td>
<td>26 (78.8)*</td>
</tr>
<tr>
<td>s1b</td>
<td>3 (9.1)</td>
</tr>
<tr>
<td>s2</td>
<td>4 (12.1)</td>
</tr>
<tr>
<td>m1a</td>
<td>13 (39.4)</td>
</tr>
<tr>
<td>m1b</td>
<td>5 (15.1)</td>
</tr>
<tr>
<td>m2</td>
<td>15 (45.5)</td>
</tr>
<tr>
<td>s1a/m1a</td>
<td>12 (36.4)</td>
</tr>
<tr>
<td>s2/m2</td>
<td>3 (9.1)</td>
</tr>
<tr>
<td>s1a/m2</td>
<td>12 (36.4)</td>
</tr>
<tr>
<td>s2/m1a</td>
<td>0</td>
</tr>
<tr>
<td>cagM</td>
<td>17 (51.5)*</td>
</tr>
<tr>
<td>cagA</td>
<td>25 (75.8)*</td>
</tr>
<tr>
<td>cagT</td>
<td>17 (51.5)</td>
</tr>
<tr>
<td>Gene combination</td>
<td></td>
</tr>
<tr>
<td>cagM/cagA/cagT</td>
<td>13 (39.4)</td>
</tr>
<tr>
<td>s1a/cagM/cagA/cagT</td>
<td>12 (36.4)</td>
</tr>
</tbody>
</table>

*Note: SG: superficial gastritis; AG: atrophy gastritis; EG: erosive gastritis; DU: duodenal ulcer; SU: stomach ulcer.

*Significant difference in comparison with SG group, p < 0.05.
Genotype $s1a/m1a$ was significantly found predominantly in patients with DU ($p = 0.002$, OR $= 3.0$, 95% CI $= 1.5$–6.1). Although in this study, $s1a/m1a$ alleles were also more prevalent in $HP$ samples in patients with AG (36.4%), but the results showed no statistical significance.

More than 20% of the SG, EG, and SU patients were infected with $s2m2$ genotype bacteria. However, no significant correlation was observed between this virulence marker and disease status.

In this study, the $cagA$ gene was found with a high incidence in most patients with inflammatory diseases of the stomach and duodenum: 75.8% in cases of AG, 83.9% – EG, 90.2% – DU, and 90.1% in cases of gastric ulcer. Simultaneously, there was a significant increase in the frequency of $cagT$ in patients with DU as compared to SG ($p < 0.001$, OR $= 7.3$, 95% CI $= 3.2$–16.8). A high $cagM$ prevalence was also found in patients with AG and DU diseases.

The analysis of the data showed that in the biopsy samples obtained from patients with DU, all three island genes of pathogenicity of $cagPAI$ are reliably more often detected in comparison with SG patients – 56.9% vs. 22.5%, which increases the risk of developing DU by 4.5 times (OR $= 4.5$, 95% CI $= 2.2$–9.2).

The combination $s1a/cagM/cagA/cagT$ was detected in 49% ($p = 0.001$, OR $= 4.2$, 95% CI $= 2.1$–8.7) and 36.4% ($p = 0.05$, OR $= 2.5$, 95% CI $= 1.0$–6.1) of patients with PU and AG, respectively.

**Discussion**

Gastric and systemic immune responses develop within a short period of time after $HP$ infection. The gastric inflammatory response induced by $HP$ consists of neutrophils, lymphocytes, plasma cells, dendritic cells, and macrophages, along with varying degrees of epithelial cell degeneration and injury [12]. Contact between $HP$ and gastric epithelial cells results in activation of NF-$\kappa B$, which is followed by increased interleukin-8 (IL-8) expression [13]. IL-18 is essential for inducing CD4 + CD25 + FOXP3 + regulatory T cell in response to $HP$, which in turn restricts excessive effector T-cell activation and promotes infection persistence [14].

After entering the host mucosa stomach, $HP$ utilizes its urease activity to neutralize the hostile acidic condition at the onset of infection. Then, $HP$ use flagella-mediated motility to move toward host gastric epithelium cells. Finally, $HP$ releases several effector proteins/toxins causing host tissue damage [7].

Differences in the prevalence of $HP$ genotypes could play a role in the incidence diseases. This study analyzed the association between gastric mucosa pathology and $HP$ genotypes.
VacA is a key toxin for HP pathogenesis. VacA is bound to cell surface and initiates a pro-inflammatory response. It modulates T- and B-cell activation and proliferation, and if the toxin is transported to the mitochondria, it induces apoptosis [15]. The vacA gene is present in all HP strains and comprises variable regions (signal – s, middle – m, and intermediate – i).

The vacA signal region encodes the N terminus of the toxin and may be active (type s1) or encode an N-terminal extension, which blocks activity (type s2). The vacA mid-region may bind to a wide range of cells, causing toxicity (type m1) or to a smaller range (type m2) [7]. Genotypic combination of vacA alleles results in different pathogenicity level as follows: s1m1 produces high amount of toxin and consequently the most virulent, s1m2 – produces moderate vacuolating toxins [11].

Conclusions about any association between the presence of HP genotypes and clinical outcome may not be true because there is a geographic variability of HP genotypes [16, 17]. It is important to determine the most prevalent genotypes in a specific region and to confirm whether there is a correlation between putative virulence genes and diseases. According to the literature, the vacA s1a allele is found most predominantly in Northern and Eastern European strains, whereas s1b is prevalent in Latin America [18–20].

It is known that patients with the vacA s1a gene had a significantly more strong degree of chronic inflammation, whereas those with vacA s2 had significantly less activity inflammation [21, 22].

In agreement with other studies, we confirmed that the most determined vacA allele is s1a [22–24] and less common is allele m1a. Our findings demonstrate an association between the presence of the s1a allele and the presence of AG, EG, and DU. The explanation for these results may be stated that HP strains with the s1a genotype produce more vacuolating toxin and might contribute to the more effective development of inflammation.

Atherton et al. [25] demonstrated that m1a alleles are associated with gastric epithelial injury. Our results also demonstrate an association between the presence of the m1a allele and presence of DU.

We found that the frequency of the genotype s1a/m1a vacA gene in patients with chronic SG was 21.4%; it is significantly lower compared with the group of patients with DU – 45.1% \((p < 0.05, OR = 3.0, 95\% CI = 1.5–6.1)\).

The cagPAI is one of the important virulence determinants of HP. The cagPAI encodes 27–31 proteins and comprises two regions – cagI and cagII. The cagPAI genes encode a T4SS that forms a structure to translocate the cagA protein into the gastric epithelial cells. In addition, the cagPAI stimulates the release of pro-inflammatory cytokines in gastric mucosa, including IL-8 [8].
In this investigation, the *cagA* and *cagM* were selected to represent *cagI*, and *cagT* was selected to represent *cagII*.

The primary virulence determinant of *HP* is CagA oncoprotein, which is associated with development of severe outcomes of the infection [26]. When CagA protein is introduced into a host cell, toxic change leads to abnormal proliferation, motility, and cytoskeletal change in gastric epithelial cells [27].

In European studies of *HP* strains, the frequency of *cagA* has shown variations (72%–81%) and has been higher in southeast European countries (Bulgaria, Greece, and Turkey) [18, 28, 29]. A high *cagA* gene frequency (86%–87%) is observed in East Asia [30, 31]. The results of this study also showed that the prevalence of *cagA* is significantly higher among different disease groups in comparison with SG group (*p* < 0.05). The highest percentage is observed in patients with ulcer (90.2%).

The *cagT* is located at the 3′ end of the *cagII* region. Studies showed that strains lacking the *cagT* gene had a defective “molecular syringe” that is encoded by the PAI [32], thus reflecting on inability of the type-IV system to eject out the *cagA* protein. Our results show that the prevalence of *cagT* was significantly different in patients with DUs compared with those with non-ulcer dyspepsia and it is a better marker than the *cagA* status for ulcer clinical outcome. Other studies conducted elsewhere also support our findings; in England, Kauser et al. [19] found an association between *cagT* and ulcer development.

We found a significant correlation between the simultaneous presence of *cagPAI* genes in the samples studied. It was found that combination *cagM/cagA/cagT* is detected more frequently in patients with peptic ulcer than in patients with chronic gastritis. The data show the significant roles of CagT and CagM in the T4SS effectivity in *HP*–host interaction that results in an enhanced ability of the bacterium to colonize the antral gastric niche.

This study analyzed the relationship between *HP* genotypes and clinical data of gastrointestinal diseases. The results presented in this study show that the *HP* samples from patients suffering DU had a high expression of s1a/m1a genotype (45.1%). We conclude that genotyping of *HP* (*cagPAI*) may be useful in clinical practice for identifying patients with a high risk of developing gastric diseases.

We think that the prevalence of virulence-associated *HP* genotypes has clinical implications for our understanding of the pathogenesis of infection and related gastric disorders. In future studies, it would be of interest to evaluate the influence of other pathogenicity genes in helicobacteriosis clinical outcome.
Conflict of Interest

The authors declare no conflict of interest.

References


