**Helicobacter pylori** colonization of the adenotonsillar tissue: Fact or fiction?

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**ABSTRACT**

Objective: The transmission of the gastric pathogen **Helicobacter pylori** involves the oral route. Molecular techniques have allowed the detection of **H. pylori** DNA in samples of the oral cavity, although culture of **H. pylori** from these type of samples has been sporadic. Studies have tried to demonstrate the presence of **H. pylori** in adenotonsillar tissue, with contradictory results. Our aim was to clarify whether the adenotonsillar tissue may constitute an extra gastric reservoir for **H. pylori**.

Methods: Sixty-two children proposed for adenoidectomy or tonsillectomy were enrolled. A total of 101 surgical specimens, 55 adenoid and 46 tonsils, were obtained. Patients were characterized for the presence of anti-**H. pylori** antibodies by serology. On each surgical sample rapid urease test, immunohistochemistry, fluorescence in situ hybridization (FISH) with a peptide nucleic acid probe for **H. pylori**, and polymerase chain reaction–DNA hybridization assay (PCR–DEIA) directed to the vacA gene of **H. pylori** were performed.

Results: Thirty-nine percent of the individuals had anti-**H. pylori** antibodies. Rapid urease test was positive in samples of three patients, all with positive serology. Immunohistochemistry was positive in samples of two patients, all with negative serology. All rapid urease test or immunohistochemistry positive cases were negative by FISH. All samples tested were negative when PCR–DEIA for **H. pylori** detection was used directly in adenotonsillar specimens.

Conclusions: The adenotonsillar tissue does not constitute an extra gastric reservoir for **H. pylori** infection, at least a permanent one, in this population of children. Moreover, techniques currently used for detecting gastric **H. pylori** colonization are not adequate to evaluate infection of the adenotonsillar tissues.

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1. Introduction

**Helicobacter pylori** is considered the most frequent chronic infection of humans, with more than half of the world’s population being infected [1]. **H. pylori** is a microaerophilic, gram negative bacterium that persistently colonizes the human gastric mucosa. The infection is usually acquired early in childhood and can persist for the lifetime of the host if not treated [1]. **H. pylori** infection occurs worldwide, but its prevalence is significantly different between countries [2]. Studies of risk factors for **H. pylori** infection have associated the prevalence of infection with the socioeconomic indicators, with higher prevalence in developing regions and lower prevalence in developed regions [2].

The transmission of **H. pylori** is still poorly understood, but the faecal–oral and the oral–oral are the most consensual routes of transmission [1,2]. The presence of **H. pylori** in the oral cavity has been detected in samples from supragingival plaque, dental plaque and saliva by polymerase chain reaction [2–4], but culture of the bacteria from this type of samples has been sporadic. More recently, it has also been suggested that adenotonsillar tissues may be a reservoir for **H. pylori**, and several studies evaluated the presence of **H. pylori** in adenoid and tonsil specimens [5–7]. However, results have been contradictory and, most importantly, techniques used to detect **H. pylori** in these tissues were not appropriate. Therefore, our aim was to clarify whether the adenotonsillar tissue may constitute an extra gastric reservoir for **H. pylori**. We have studied 62 children proposed for further studies.
adenoidectomy and/or tonsillectomy, and a total of 101 samples (55 adenoid and 46 tonsils) were characterized for the presence of *H. pylori* using different techniques.

2. Materials and methods

2.1. Patients and surgical procedures

Patients were consecutively selected from the surgical waitlist for adenoidectomy and/or tonsillectomy, with or without treatment of middle ear disease, and that had not been treated with antibiotics nor with proton pump inhibitors on the 4 weeks prior to surgery. Sixty-two patients (mean age = 7.9 ± 5.5 years and male/female ratio of 1:1.2) fulfilled these criteria and were recruited for this study. Thirty-nine patients (62.9%) underwent adenotonsillectomy, 16 patients (25.8%) underwent adenotonsillectomy and seven patients (11.3%) underwent tonsillectomy at the Hospital São Marcos, Braga, Portugal. Routine surgical indications for adenotonsillectomy were followed. All patients were operated under general anaesthesia. Blockage of gastro-pharyngeal reflux was obtained with gauze placed on the hypopharynx after oro-tracheal intubation and before trendelenburg positioning. Protocols followed in the study were in accordance with the institutional ethical standards. Informed consent was obtained from the patients' parents. Samples were delinked and unidentified from their donors.

2.2. Serology

Serum samples were used for detection of IgG antibodies against *H. pylori*, using the Pyloriset EIA-A-G III Kit (Orion Diagnostics, Espoo, Finland). The assay was performed following the manufacturer's instructions.

2.3. Rapid urease test

A 1 mm³ sample from each surgical specimen was immersed in urea broth containing phenol red (Fluka, Buchs, Switzerland) and incubated for 24 h at 37 °C. At the end of that time period, colour changes in medium were registered.

2.4. Histology and immunohistochemistry

Surgical specimens were formalin fixed, paraffin embedded, and sections of 3 μm were cut. Slides were deparaffinised in xylol and hydrated with grade ethanol concentrations, and stained with haematoxylin and eosin, modified Giemsa and with a rabbit anti-human polyclonal antibody against *H. pylori* (Cell Marque, CA, USA). Briefly, antigen retrieval was performed by microwave pre-treatment in 10 mmol/L citrate buffer (pH 6.0) for 15 min. Endogenous peroxidase activity was blocked with hydrogen peroxide block solution (Labvision, UK) for 10 min. After washing in phosphate buffered saline (PBS), sections were incubated for 30 min with primary antibody (1:500). Staining was achieved using Dako RealTM EnvisionTM/HRP rabbit/mouse polymer, for 30 min at room temperature. Binding was detected using diaminobenzidine (DAB), which upon oxidation forms a stable brown end-product. Nuclei were slightly counterstained with Mayer's haematoxylin. Gastric biopsies positive for *H. pylori* infection were used as positive controls. Slides were visualized by two experienced pathologists that were blind to the patients' clinical data.

2.5. PNA–FISH

*H. pylori* was detected in the adenotonsillar samples using the Hpy769 probe, a peptide nucleic acid probe (PNA) that is highly specific and sensitive for the detection of *H. pylori* [8]. The surgical specimens were deparaffinised and rehydrated in xylol and ethanol. Samples were immersed twice in xylol for 15 min, and in decreasing concentrations of ethanol for 5 min, and were finally washed with distilled water for 10 min. Samples were allowed to air dry and the fluorescence in situ hybridization (FISH) procedure was performed as described in Guimaraes et al. [8].

2.6. DNA isolation

DNA was isolated of 8 mm³ fragments from all surgical specimens using the DNA Invisorb Spin Tissue Mini Kit (Invitek, Berlin, Germany). The extraction was performed following the manufacturer's instructions. In a subset of cases, the whole surgical specimen was fully homogenized in Tris–EDTA buffer (10 mM Tris pH 7.5, 1 mM EDTA) using an ultraturax homogenizer and DNA was isolated.

2.7. PCR and DEIA

Detection of the vacA gene of *H. pylori* was performed using primers VA1F and VA1XR, described previously [9]. PCR reaction mixtures were performed in a volume of 25 μL, 1× PCR Buffer, 2.5 mmol/L of MgCl₂, 0.25 mmol/L dNTPs, 0.25 U AmpliTag Gold, and 25 pmol of each primer. PCR was performed with 9 min pre-denaturation at 95 °C, followed by 40 cycles of 30 s at 95 °C, 45 s at 50 °C, and 45 s at 72 °C. Final extension was performed for 10 min at 72 °C. Amplimers carrying a biotin moiety at the 5¢ terminus of the reverse primer were detected in a microtiter well-based DNA hybridization assay (DEIA) using a specific general probe for the vacA gene [10,11], DNA isolated from *H. pylori* Tx30a (ATCC 51932, American Type Tissue Culture) and 26695 (ATCC 700392) strains was used as reference.

2.8. Statistical analyses

Age distribution between *H. pylori* seropositives and seronegatives was analysed by the Student's t-test. Association between gender and *H. pylori* serology was assessed by the Pearson Chi-Square test. Logistic regression models were computed with Statview for Windows software (version 5.0; SAS Institute Inc., Cary, NC). Differences in data values were considered significant at *p* values lower than 0.05.

3. Results

The study population was constituted by 62 children living in the North of Portugal, which were indicated to surgery due to adenotonsillar hypertrophy (47 cases), infection (5 cases) or both (10 cases). Sixty-two blood samples and 101 surgical samples (55 adenoids, 23 left tonsils and 23 right tonsils) were obtained.

The presence of anti-*H. pylori* antibodies in serum was detected in 24 (39%) cases. Infected patients were older than uninfected patients (*p* = 0.013) and were more frequently females (*p* = 0.044) (Table 1). In a logistic regression analysis, including both gender and age in the model, only age remained associated with *H. pylori* seropositivity (*p* = 0.039).

To evaluate the presence of *H. pylori* in the adenotonsillar tissues, we have used the rapid urease test, immunohistochemistry with a polyclonal anti-*H. pylori* antibody, FISH with a specific *H. pylori* PNA probe, and PCR–DEIA for *H. pylori* vacA gene (Table 2).

The rapid urease test was positive in two (3.6%) adenoid and in two (4.3%) tonsil specimens from three patients. All urease test positive samples were from patients infected with *H. pylori* as evaluated by serology.
Immunohistochemistry was positive in three tonsil specimens from two patients (Fig. 1), and these patients were H. pylori serology negative. FISH using a specific PNA probe for H. pylori was negative in paraffin cuts consecutive to those used for immunohistochemistry (Fig. 2). PNA–FISH was negative in all studied specimens.

The presence of H. pylori in the adenotonsillar samples was also assessed by PCR, directed to the vacA gene, which is present in all strains of H. pylori. In order to improve the sensitivity of the PCR, amplimers were hybridized with a vacA specific probe in DEIA assays. All samples were negative for the presence of H. pylori. To exclude sampling bias, in a subset of cases corresponding to those that were ELISA positive, DNA isolation was performed after fully homogenizing the whole surgical specimen. All cases were negative for H. pylori.

4. Discussion

Data concerning the presence of H. pylori in adenoids and/or tonsils are inconsistent. In this study we have initially characterized all patients for the presence of serum anti-H. pylori IgG antibodies. In agreement with previously published data on the prevalence of H. pylori infection in the Portuguese population in paediatric age, 39% of the patients were H. pylori positive [12]. The presence of the infection was associated with older patients and, in keeping with previous studies, gender did not constitute a risk factor for H. pylori infection [13].

To clarify whether H. pylori is present in the adenotonsillar tissues of these children, we have used a variety of approaches.
directly aimed at the tissue samples. Therefore, each adenoidal and tonsillar specimen was submitted to rapid urease test, immunohistochemistry, PNA–FISH and PCR–DEIA. We detected urease-positive microorganisms in four tissue specimens of three patients with positive *H. pylori* serology. However, we also demonstrated by immunohistochemistry, PNA–FISH and PCR–DEIA that these urease-producing microorganisms were not *H. pylori*. Though immunohistochemistry was positive in three tonsil specimens from two patients, these specimens were *H. pylori*-negative by PNA–FISH and PCR–DEIA, and the patients did not present antibodies anti-*H. pylori*.

All adenotonsillar specimens included in this study were negative when PNA–FISH or PCR–DEIA were used. The PNA–FISH method has been previously tested in different types of samples and with different microorganisms, and has shown a high specificity for *H. pylori* [8]. PCR is recognized as having high specificity if the target gene and the primers are carefully chosen. In this study, we have used as amplification target *H. pylori* vacA gene that, besides having no homology to other bacterial species, is present in all *H. pylori* strains. The primers used are well established and their high sensitivity and specificity has been previously demonstrated [10,11]. PCR sensitivity was increased by hybridization of amplified products with a specific vacA probe [11].

Considering that PNA–FISH and PCR–DEIA are the most specific and sensitive methods and that all tissue samples analyzed were negative, results obtained by rapid urease test and immunohistochemistry are false positives. Several studies have reported the presence of *H. pylori* in adenoid and/or tonsillar tissues based on the solely use of rapid urease test [6,7]. However, we and others showed that, in cases positive for the rapid urease test, the use of additional detection methods reveals that those are likely false-positive results [14,15]. One very important aspect concerns the specificity of the rapid urease test and of immunohistochemistry. These are related with the probability of the presence in the sample of microorganisms, other than *H. pylori*, that could origin cross reactivity leading to false-positive results. So, although these techniques have a high specificity when used in gastric samples, the specificity may be lower in samples from polymicrobial environments such as the adenotonsillar tissues [14]. The rapid urease test is not in fact a specific test for *H. pylori*, but for urease-producing microorganisms. Therefore, the positive results that we have obtained probably reflect the presence in adenotonsillar tissues of other urease-producing bacteria. Likewise, and although immunohistochemistry allows higher reproducibility of *H. pylori* detection in gastric samples when compared with conventional histological techniques [16], in polymicrobial extra gastric samples antibodies can recognize bacterial epitopes other than *H. pylori*, leading to false-positive results.

The negative results we have obtained by PNA–FISH and PCR–DEIA are also in accordance with previously published data using PCR and/or culture [14,15,17–20]. The use of PCR for detection of *H. pylori* in adenotonsillar tissues has sporadically given positive results [21]. However, primer choice needs to be viewed with caution, since certain primer sets used for detection of *H. pylori* produce a very high rate of false-positive results [22].

The main limitation of this study is related with the fact that the gastric infection status of the studied population was only evaluated by IgG serology, which is not specific for active gastric *H. pylori* infection. Therefore, we cannot exclude the possibility that the adenoids and tonsils are infected only in cases of active or recurrent disease. Furthermore, and because *H. pylori* infection is less common in the paediatric population than in adults, the lack of adenotonsillar *H. pylori* infection in children does not exclude this possibility in the adult population.

5. Conclusion

The adenotonsillar tissue does not constitute an extra gastric reservoir for *H. pylori* infection, at least a permanent one, in this population of children. Furthermore, techniques currently used for detecting gastric *H. pylori* colonization are not adequate to evaluate infection of the adenotonsillar tissues.

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