Retrospective Microbiological Study of Atypical Recurrent Pharyngitis in Patients Presenting the White-line Clinical Sign

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ABSTRACT

Background: Chronic atypical oropharyngeal disease in adults, accompanied with chronic cough, can occur at any age manifesting itself with different grade evolutive diseases. Often their pathogenesis is attributed to gastroesophageal reflux, to virosis or to unspecified immune deficiencies but some clinical aspects, such as the simultaneous presence of urinary disorders, the temporal scanning of the recurrence and the reduced response to antibiotic therapy, suggests a different or a superimposed pathology.

Methods: The present study was carried out to assess retrospectively biopsy and biological materials from a population afflicted by atypical recurrent pharyngitis, presenting a “white line” clinical sign into the context of respiratory difficulties, manifesting chronic choking cough (CCC), laryngopharyngeal (LPR) and gastroesophageal (GERD) reflux diseases. This population, already clinically, endoscopically and histologically characterized, was newly studied following the microbial approach by cultural and molecular procedures.

Results: We analyzed 14 biopsy, 60 biological pooled materials from lingual, pharyngeal, post nasal drip mucoid secretions and sputum (here initialled: LPNS) and 60 lingual cell and salivary secretions (LCSS) resulted positive to Chlamydiaceae [(Chlamydia pneumoniae (Cp) and Chlamydia trachomatis (Ct)], to urogenital Mycoplasmas [Mycoplasma hominis (Mh) and Ureaplasma urealyticum (Uu)], to Helicobacter pylori (Hp) into the context of a changeable overlapping with other typical bacteria, belonging to Corynebacteria, Enterobacteria, Streptococci and Staphylococci groups.

Conclusions: Our data indicated that atypical infections [C. trachomatis and urogenital Mycoplasmas (Mh and Uu)], together with Cp, were the underhand pathogens of an initial chronic oropharyngeal scenery until now unrecognized, triggering, after decades, the respiratory problems in middle and old subjects genetically susceptible. The presence of white line clinical sign, endoscopically observed, together with an altered pH salivary secretion, into the scenario of CCC, LPR and GERD reflux manifestations, refractory to non-specific medical therapy, represents a pathognomonic triad to include routinely these valuations into the diagnostic protocol of an atypical recrudescence pharyngitis.

Keywords: Whiteline reflux; Chronic choking cough, Laryngopharyngeal and gastroesophageal reflux diseases; Chlamydiaceae; Mycoplasmataceae; Helycobacter Pilory; Molecular age

INTRODUCTION

Oral cavity, rhinopharynx and larynx, are anatomical structures of access to respiratory and digestive tracts [1] and their mucosae are continuously stressed by the chemical and microorganism agents. The healthy human microbiome is the consequence of a durable complex interaction developing between microbe distributions and different human modulating mechanisms, regulated (and/or harmonized) by the coexistence of several factors, beginning from conception to death [2,3]. Physiologically, the laryngeal reflex (LR) represents an involuntary and necessary protective response to stimuli in the larynx to regulate the three principle
functions: airway protection, respiration and voice production. The LR entity is valued by ENT specialists to predict dysphagia or to portend clinical phenotypes of chronic cough, vocal cord dysfunction or paediatric apnoea [4]. In adult, the LR has been observed frequently in population showing gastroesophageal reflux, obstructive sleep apnoea, chronic obstructive pulmonary disease, neurodegenerative disorders and advanced age [4,5]. An important defence mechanism against external noxious stimuli is operated by mucociliary transport which carrying out the mucous released by goblet cells, protect their mucosae from chemical agent and microorganism [6]. In spite of different typological aetiology, the evaluation and the management of chronic cough is still a much debated question [7]. In neonatal age, a cough is considered chronic if present for more than four weeks, while in most childhood cases it is caused by chronic bronchitis, asthma and/or upper airway cough syndrome. Initially, the evaluation of these exacerbating scenarios should be focused on bacterial aetiologies, targeting at treatment and monitoring for resolution, but unfortunately still nowadays we are witnessing a persistence of an old vision of these infective diseases [8,9], without the inclusion of atypical sexual bacteria, like Chlamydia trachomatis and urogenital Mycoplasmas. On the contrary, in adults, several clinical chronic oropharyngeal scenarios, accompanied with chronic cough, can occur at any age manifesting itself with different grade evolutive diseases [10-12]. Further, chronic choking cough may be produced by a number of different disorders in distinct anatomic sites and still represent a frustrating and common problematic condition resulting in significant psychological and physical sequelae as well as enormous financial costs in terms of expensive health care and loss of working days [13]. Further, chronic choking cough (CCC) can be caused by other worsening symptomatologies as gastroesophageal reflux disease (GERD), laryngopharyngeal reflux disease (LPR) and no asthmatic eosinophilic bronchitis overlapping themselves with other bacterial aetiologies assignable to different principal conditions. So, recently it is also reported a possible H. pylori association with chronic tonsillitis and laryngopharyngeal reflux in the tonsillar biopsy samples [14]. Although from human proteomic studies on the salivary protein characterizations and them genetic polymorphisms [15] emerged the important role of saliva as a non-invasive approach for diagnostic and prognostic purposes, only recently has been clarified the different role of atypical pathogen [M. pneumoniae (Mp) or C. pneumoniae (Cp)] in terms of ethnogenesis [16], in terms of polymorphic genetic susceptibility [17] and that of common pathogens [M. catarrhalis (Mc), H. influenzae (Hi), or S. pneumoniae (Sp)] in terms of recrudescent childhood asthmatic episodes [18]. At this purpose, the oral and nasal cavity, rhinopharyngeal, laryngeal and upper airway mucosae represent perfect niches for common and atypical microbial adhesion and biofilm formation [19]. From long time, Chlamydia trachomatis, an endocellular obligate parasite prevalent in genital tract, had been defined “Trojan horse” [20], presenting different sequelae and representing a significant source of morbidity. At the same time, urogenital Mycoplasmas, esoparasites of mucosae membrane surfaces, living their inside, were frequently found together with Ct, when they were researched in any human chronically inflamed source investigated by appropriate cultural procedures [10-12, 21-23]. On the basis of these considerations, we have investigated the biopsy, LPNS and LCSS sources to evaluate the presence of atypical and common bacteria in patients affected by chronic pharyngeal inflammatory process presenting CCC, GERD and LPR reflux diseases, showing the white-line clinical sign [24]. Virus and micete were excluded for their well typified clinical manifestations and resolutive medical therapy.

MATERIALS AND METHODS

Cohort patients

Biopsy (14/60), LPNS and LCSS (60/60) samples, from outpatients of an initial cohort of 200 (30%) of Caucasian subjects, examined from June 2011 to 2011 and already clinically selected for rhino-pharyngeal globe (RPG) sensation, manifesting CCC, LPR and GERD reflux symptomatologies into the chronic pharyngitis scenery, presenting a white line sign, were reassessed hypothesizing a causative microbial involvement.

Ethics statement

The informed consent was already obtained from each outpatient during the enrolment in the prior study [24], while attending at the first visit in the ambulatory of the ENT Clinic Unit, SS. Annunziata Hospital (Chieti). A waiver of informed consent was not necessary included, because this study excludes a direct contact with a population already studied and all patient identifiers were removed from the dataset on initial collection. All patients already had adhered to the Declaration of Helsinki and to the ICH-GCP, GU 184/2003. The methodologies of this study were commercial products and others were conform to conventional procedure standard of literature.

Sample preparations

Biopsy material from rhinopharynx wall, below the white line and from the reticular tissue, above the white line, was chopped finely by little bistoury and immediately placed into an Eppendorf tube, stored at -30°C. This material was thawed before DNA extraction and recovered immediately with 1,0 mL of sterile saline solution, vigorously stirred to disperse cellular material and immediately processed to detect the Cp-, Ct-, Hp- DNA by PCR and to culture for common and atypical urogenital bacteria.

LPNS sample of the pooled biological materials, from epithelial lingual and pharyngeal living cells, added to mucoid post nasal drip and sputum, collected by scraping and expectorant modalities respectively, were pooled for each patient with 2.0 mL sterile saline solution and stored at -30°C. Before use, this suspension was thawed, vigorously stirred to disperse cellular material and treated with 8.0 mL of 2.0 mM cysteine saline solution, incubated at 37°C to solubilize mucoid material; afterwards it was centrifuged at 5000 rpm at 4°C for 10 min. The pellet was collected with 2.0 mL of sterile saline solution and used for molecular and cultural researches.

LCSS material from lingual surface was sampled by sterile wood spatula scraping, dispersed with 1.0 mL sterile saline solution and stored at -30°C. An aliquot of each of three sources was used to measure the pH value immediately before of freezing. It was thawed and immediately used as above reported for molecular and cultural analyses.

Molecular section

Each of biopsy, LPNS and LCSS aliquots (300 µL) for Ct-DNA analysis were centrifuged at 12,000 rpm for 5’ at 4°C; each pellet was immediately scattered with 180 µL of 0.025 M buffer phosphate pH=7.0 plus 1.0 mM EDTA, added with 25 µL of protease K, incubated at 56°C overnight; 210 µL of absolute ethanol was
added to precipitate the DNA extracted, filtered and eluted from column with 60 µL of elution buffer. 50 µL of this solution was amplified into a Thermocycler of Applied Biosystem (Clemens GmbH), following the manufacturer’s instructions [BioAesis srl, Jesi (AN), Italy (Line-20 Chlamydia trachomatis code 04L120). Cp-DNA sample (300 µL) was carried out following the procedure already reported [25,26] and used for our previously studies [8-10]. Hp-DNA sample (300 µL) was processed for cagA and ureC templates, using QIAamp DNA mini kit (Qiagen GmbH, Hilden, Germany) following the manufacturer’s procedure [27].

**Cultural section**

Aliquot of 75 µL from biopsy, LPNS and LCSS was dispersed separately and cultured by Mycoplasma IST 2, following the BioMerieux Manual procedures. 1.0 µL of this remaining suspension was plated onto chromID CPS agar and Columbia CAN agar. Total Microbial Charge (TMC), expressed as the total number of colony-forming units per milliliter (CFU/mL), was calculated and the relative presence of each species was reported as percentage of TMC (considered as 100%), and as previously reported [10]. Monoculture was considered significant at ≥10^8 CFU/mL. API Coryne galleries (code 20 900) were identified by an API automated system, the confidence ranged between 94.7% and 99.9%, indicating a high level of identification. Cultural section and relative colony characterization were carried out using products purchased from BioMerieux Italia. Streptococcus pneumoniae (Spn), grown on Columbia CNA, underwent PCR analysis to confirm the eventual misidentification of Streptococcus pseudopneumoniae as true S. pneumoniae [26].

**RESULTS**

All biopsies, LPNS and LCSS sources were positive for atypical pathogens (Table 1). In detail, C. pneumoniae (Cp-DNA) was positive into 3/14 biopsies (21.43%), while it was found also into 13/60 LPNS (21.67%), but only 6/60 into LCSS (10.00%). C. trachomatis (Ct-DNA) was positive into 12/14 biopsies (85.71%), while it was present into 53/60 (88.33%) of both LPNS and LCSS sources, overlapping itself between them (Cp and Ct) with twelve cases. The mycoplasmal positiveness (Mh and Uu) entered into the biopsy was of the 100%. Particularly, the Mh positivity was of the 64.28% (n=9/14), while the Uu positivity was of the 78.57% (n=11/14), spreading themselves completely in all samples and overlapping into 42.86% (n=6/14) with a very low infectant charge (10^3 UCC/mL). The Mh positiveness for LPNS and LCSS sources was higher than biopsy, resulting in 80.00% (48/60) and 85.00% (51/60) with a UCC/mL of 10^6 and 10^7 respectively, while the Uu positiveness for the same sources were both of the 88.33% (53/60), but with a major infectant charge 10^4 and 10^5 UCC/mL. At the same time, each of two different patients resulted positive respectively for H. pylori-DNA and for S. pneumoniae (Sp-DNA) in all sources together with atypical sexual pathogens. Corynemycetes, Enterobacteria, Streptococci and Staphylococci species were random distributed in the same samples together atypical pathogens at a variable percentage, ranging from 5.0 x 10^3 to 5.0 x 10^4 UFC/mL of the TMC. Median pH value of the three sources were pH=6.7 ± 0.2, pH=7.7 ± 0.4 and pH=8.4 ± 1.4 for biopsy, LPNS and LCSS respectively.

**DISCUSSION AND CONCLUSION**

It is universally accepted that these sexual atypical bacteria are asymptotically and symptomatically present into 70-80% and into 20-30% respectively of the whole population with different grade and evolutive sequelae, depending on the individual host genetic susceptibility [28]. Recent immunological advances demonstrated the pivotal roles of different individual host genetics in directing the innate immune response to Cp infection [29] and in modulating Ct pathogenesis [30]. Considering the initial Chlamydia trachomatis and urogenital Mycoplasmas contaminations an event always possible during sexual activity, these atypical infections, impairing the trophoblast function [31], could inflict serious injuries on different cellular structures of foetal tissue in active accretion, colonizing the humankind from early stage of conception [32]. At this purpose, it has been noted from several times, but recently updated [33], that chlamydial and mycoplasmal infections in early asymptomatic pregnant women [34] are linked to increased risk of spontaneous abortions, preterm birth and/or stillbirth [35]. Although the atypical bacteria, like C. pneumoniae and M. pneumoniae, were frequently considered in the acute and/or chronic manifestations of the different oropharyngeal and upper respiratory diseases of the pediatric age [8,9], only very poor studies had been carried out on neonates including both those atypical sexual-derived and evaluating them into mother’s cervical-vaginal ecosystem [23]. Actually, although recent acquisitions have clarified the human microbiome compositions and its development during three crucial developmental stages, like pregnancy, birth, and infancy [2], and new research strategies have been established [3], new approaches on the maintenance of human health and microbial perturbations into the contest of human different pathophysiology of autoimmune disorders will be necessary. In spite of these recent acquisitions on human microbiome and on mechanisms of chlamydial and mycoplasmatic diseases, the clinical studies on ill hospitalized population are carried out still on the knowledge of old microbiological protocols following outdated laboratory procedures, excluding them from routine investigations. Thus, in this retrospective preliminary remark, we found a full dispersion and overlapping of sexual atypical bacteria into the biopsy, LPNS and LCSS of patients affected by chronic pharyngitis evolting, decades later, in CCC, LPR and GERD manifestations. These results constitute a new indication in directing further the methodological approaches for studying

Table 1: Atypical bacteria results from biopsy, LPNS and LCSS sources from oropharyngeal cavity and upper respiratory tract.

<table>
<thead>
<tr>
<th>Biopsy (n=14)</th>
<th>LPNS (n=60)</th>
<th>LCSS (n=60)</th>
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<tbody>
<tr>
<td>Cp-DNA: positive n=3/14 (21.43%)</td>
<td>Cp-DNA: positive n=13/60 (21.67%)</td>
<td>Cp-DNA: positive n=6/60 (10.00%)</td>
</tr>
<tr>
<td>Ct-DNA: positive n=12/14 (85.71%)</td>
<td>Ct-DNA: positive n=53/60 (88.33%)</td>
<td>Ct-DNA: positive n=53/60 (88.33%)</td>
</tr>
<tr>
<td>Mh: 10^3 UCC/mL n=9/14 (64.28%)</td>
<td>Mh: 10^3 UCC/mL n=48/60 (80.00%)</td>
<td>Mh: 10^3 UCC/mL n=51/60 (85.00%)</td>
</tr>
<tr>
<td>Uu: 10^3 UCC/mL n=12/14 (85.71%)</td>
<td>Uu: 10^3 UCC/mL n=53/60 (88.33%)</td>
<td>Uu: 10^3 UCC/mL n=53/60 (88.33%)</td>
</tr>
<tr>
<td>Hp-DNA: positive n=1/14</td>
<td>Hp-DNA: positive n=1/60</td>
<td>Hp-DNA: positive n=1/60</td>
</tr>
<tr>
<td>Sp-DNA n=1/14</td>
<td>Sp-DNA n=1/60</td>
<td>Sp-DNA n=1/60</td>
</tr>
<tr>
<td>pH = 6.7 ± 0.2</td>
<td>pH = 7.7 ± 0.4</td>
<td>pH = 8.4 ± 1.4</td>
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(*) molecular detected.

(•) culturally isolated and molecularly confirmed.
a population of patients presenting a white line clinical sign into the contest of CCC, LPR and GERD reflux diseases, as recently detailed in a new clinical case [36]. Further, the lack of a careful clinical evaluation of these atypical unrecognized scenarios (Figures 1 and 2), of a careful timing choice of the molecular procedures [Ct-DNA stability [36-38] against Ct-RNA increased clearance in spontaneous pharyngeal chronic scenery [39], and of a sampling procedural adequacy (swabbing in “acute” [23]) against (scraping in “chronic” moment of the inflammatory state [12,12,21,23,36-38]) make the choice of molecular procedure a technique yet to be well defined. The results for chlamydial and mycoplasmal presences into biopsy, LPNS and LCSS demonstrate once again the temporal and procedural correctness of our molecular choice in detecting the nucleic acids presence for these atypical bacteria. No additional oropharyngeal alterations were observed for *H. pylori* (Figures 1 and 2, panel B and B') and for *S. pneumoniae* (Figures 1 and 2, panel C and C') positive patients into the contest of to Chlamydiaceae and urogenital Mycoplasmas. An exhaustive description of the biopsy histological pictures, performed above and below the white line, previously reported [24], revealed a changeable morphology of the inflammatory state, varying from chronic flogistic infiltration to intraepithelial apoptosis, changing through the different evolutive phases of chronic inflammatory setting, probably determined by genetic susceptibility and by a temporary spontaneous pharyngeal chlamydia trachomatis RNA clearance [39], but decisively assignable to sexual atypical pathogens. To confirm this, there were the progressive quantification of atypical bacteria culturally detected into LPNS and LCSS sources respect to biopsy. Further, we can affirm that LPNS and LCSS biological materials from oral cavity represent the best sources to evaluate these pathogens during a precocious contamination. Thus, the highest mean value of pH of LCSS (pH=8.4 ± 1.4) compared to LPNS (pH=7.7 ± 0.4), directly confirmed the high significance of this parameter in signaling in advance a silent mycoplasmal infection in two different, but contiguous localizations of the oral cavity. At this purpose, it had already been reported that the high pH value represent the first step towards the unbalance of hypothiocyanite ion (OSCN-) human saliva formation [39], one of the physiological antimicrobial products of the salivary peroxidase system of human microbiome [41,42]. The physical salivary valuation (“serous” against “mucous” change), together with the high pH value, would represent an
additional precocious marker of a complex protein mixture, indicative of an oral altered microbiota [10] and of a source material easily, quickly and noninvasively obtainable to evaluate human microbiome in different phenotype diseases, as already reported for ocular microbiota [23]. Thus, an increase of pH value from physiological range would represent the first parameter accurately measurable and easily obtainable to orient their researches. In conclusion, identification of sexual atypical infections in oral cavity, collecting an accurate anamnesis on the clinical signs and symptomatology of oral cavity sexually involved, introducing two simple chemical and physic parameters, paying utmost attention to the sampling modality, swabbing vs scraping, for chronic chlamydial infections [23,36,38], introducing routinely their researches, we ascertained their involvement in oropharyngeal and respiratory diseases, that sometimes could trigger reactive arthritis also in younger patient genetically susceptible [12,21] and/or in other clinical case still to reveal. Nowadays, it remains hard for us to comprise the motivation of their persistent exclusion in clinical studies on pathogens causing upper respiratory tract infections in outpatients, considering the chronicity of these infections and relative diseases, the necessity of health economy reduction and of loss working hours, providing a better medical assistance, attending the development of effective and durable vaccine [43]. Lastly, considering their dangerous effects on embryonal development during any phase of foetal growth [28], depending on their capacity to escape from mother’s innate immune system, manifesting their pathogenicity at the birth [23] and/or on their capacity to modify the estrogen-progesterone hormonal levels [31], they could prompt a major tissue alterations causing a variety of other embryonal and genetic anomalies. Further studies will be necessary to comprehend truly complexity of the human microbiome considering their asymptomatic presence into the major part of human population.

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