Comparison of Acoustic and Stroboscopic Findings and Voice Handicap Index between Allergic Rhinitis Patients and Controls

Eltaf Ayça Özbal Koç1, Bülent Koç2, Selim Erbek1

1Department of Otorhinolaryngology, Başkent University Faculty of Medicine, Ankara, Turkey
2Department of Otorhinolaryngology, Pendik Hospital, İstanbul, Turkey

Background: In our experience Allergic Rhinitis (AR) patients suffer from voice problems more than health subjects.

Aims: To investigate the acoustic analysis of voice, stroboscopic findings of larynx and Voice Handicap Index scores in allergic rhinitis patients compared with healthy controls.

Study Design: Case-control study.

Methods: Thirty adult patients diagnosed with perennial allergic rhinitis were compared with 30 age- and sex-matched healthy controls without allergy. All assessments were performed in the speech physiology laboratory and the testing sequence was as follows: 1. Voice Handicap Index (VHI) questionnaire, 2. Laryngovideostroboscopy, 3. Acoustic analyses.

Results: No difference was observed between the allergic rhinitis and control groups regarding mean Maximum Phonation Time (MPT) values, Fo values, and stroboscopic assessment (p>0.05). On the other hand, mean VHI score (p=0.001) and s/z ratio (p=0.011) were significantly higher in the allergic rhinitis group than in controls.

Conclusion: Our findings suggest that the presence of allergies could have effects on laryngeal dysfunction and voice-related quality of life.

Key Words: Allergic rhinitis, voice handicap index, dysphonia

INTRODUCTION

Allergic rhinitis (AR) is a worldwide health issue affecting 10%-25% of the population (1). AR has been intensely related with many chronic diseases such as asthma, rhinosinusitis, and ear infections (2). AR affects everyday activities, and is not a fatal disease so its morbidity is often underestimated. The problems caused by AR are often worsened by the variety of complaints involved (3). Voice disorders have been commonly associated with respiratory allergy (3). In a population of 80 AR cases, Baker et al. (4) found voice disorders in 44.75% of the patients.

The nose and other tissues of the supraglottic vocal tract are responsible for vocal quality and perceived characters of speech sounds (5,6). Niedzielska et al. (7) studied the acoustic estimation of voice and stated that in patients with AR, the mean Jitter and Shimmer values were higher than in the control group; however, the difference did not reach statistical significance. Simberg et al. (5) studied the relationship between vocal symptoms and allergy, and stated that vocal fold injuries, such as haemorrhage and mucosal tears, may significantly accompany concurrent reflux, asthma or allergy.

Although there is a lot of literature dealing with the nasal symptoms of AR, comparatively little information has been published on the effects of AR on voice and speech. The aim of the present study was to investigate the acoustic and stroboscopic findings of the larynx and Voice Handicap Index (VHI) scores in patients with AR and to compare them with healthy controls.

MATERIAL AND METHODS

Study design

The present study was approved by the Başkent Institutional Review Board (KA12/223). Written informed consent was obtained from all participants. Thirty adult patients diagnosed with perennial AR between February 2013 and November
2013 were compared with 30 age- and sex-matched healthy control subjects without allergy. To qualify for enrolment, patients with AR needed to have a positive skin prick test result. The healthy controls without rhinitis and allergy symptoms were randomly selected among patients admitted to the dental clinic of our centre for routine dental examination. None of the controls were aware of having any allergy and their skin prick test results were negative.

Exclusion criteria for the study and the control groups were as follows: age below 17 or above 70 years, nasal problems (nasal polyposis, severe nasal deviation etc.), pregnancy, smoking, history of laryngeal pathology or laryngeal surgery, diagnosed reflux or gastroesophageal reflux (GER), patients with asthma, patients who received systemic corticosteroids, systemic antihistamines, decongestants or leukotriene modifiers in the past six months or immunotherapy within the past two years.

**Skin prick test**

All subjects underwent skin prick testing performed on the forearm. Saline solution was used as the negative control and histamine as the positive control. The presence of allergy was assessed through skin-prick test (Alk-Abello S.A., Madrid, Spain). A positive response was judged by a wheal diameter at least 3 mm larger than the wheal diameter of the negative control. Skin-test responsiveness was assured with placement of the positive control.

**Outcome parameters**

All assessments occurred in the speech physiology laboratory in our clinic. The testing sequence was as follows: 1. VHI questionnaire, 2. Laryngovideostroboscopy, 3. Acoustic analyses.

1. **Voice Handicap Index**: The Turkish version of a validated questionnaire, the Voice Handicap Index (VHI), was performed for all patients (8, 9). The VHI consists of 10 questions on emotional, functional and physical aspects of voice. Each question is graded on a 5-point scale (0= Never, 1= Almost Never, 2= Sometimes, 3= Almost Always, 4= Always). The total score ranges from 0 (unaffected) to 40 (severely affected).

2. **Laryngovideostroboscopy**: Larynx examination is made by using a 70° magnifying laryngoscope (Richard Wolf, GmbH, Knittlingen, Germany). Vibratory behaviour was assessed by a Digital Strobe View 5570 stroboscope (Richard Wolf, GmbH, Knittlingen, Germany). Closure level, amplitude, vocal fold edge, supraglottic involvement, mucosal wave, non-vibrating portion and closure phase were observed. The results were marked on the “Stroboscopic Assessment Form” (10). Endolaryngeal secretion, excessive mucus and oedema were also noted.

As for the aerodynamic assessment, maximum phonation time (MPT) was determined by measuring the duration of the /a/ vowel after maximum inspiration. MPT was elicited over three trials, the longest of which was considered definitive. The s/z ratio was calculated by measuring the MPT of single consonants /s/ and /z/ in two separate breaths. The best /s/ and /z/ effort of at least three attempts at each was used to obtain the ratio.

3. **Acoustic analysis**: Multi-Dimensional Voice Program (MDVP, model 5105, version 2.5, Kay Elemetrics Corp, Lincoln Park, NJ, USA) with Multi-Speech software (Model 3700, version 2.4, Kay Elemetrics Corp, Lincoln Park, NJ, USA) was coupled to a microphone (AKG, Model C-1000, Nashville, Tennessee, USA), with a standard sound board (Creative Sound Blaster Audigy 2ZS, Creative Technology Ltd, Singapore). The recordings were performed in a silent room by means of a microphone at a stable mouth-to-microphone distance of 15 cm. The patients were educated to sustain the vowels at a comfortable pitch and level of loudness three times before recording in order to obtain maximum phonation during recording. The acoustic analysis was performed by measuring the pitch levels. The habitual pitch level was determined using the strobe unit as the patients read the words “adanayaaaaa” and prolonged the final vowel /a/ (11).

**Statistical analysis**

Data were analysed using the Statistical Package for Social Sciences 15.0 for Windows (SPSS Inc., Chicago, IL, USA). Parametric tests were applied to data of normal distribution and non-parametric tests were applied to data of questionably normal distribution. The Student-t test, Chi-square test, Fisher’s Exact test and Continuity (yates) correction test were used to analyse the differences between two groups. Data are expressed as mean±SD. All differences associated with a chance probability of 0.05 or less were considered statistically significant. The power analysis of the study for the continuous variables for s/z ratio was 74% (the standard deviation was 0.175, hypothesis was two-sided, and the α value was 0.05). Since more than 20% of the cells in the expected frequencies less than 5 in some of the cross table, the Yates correction was made (number was less than 30).

**RESULTS**

Thirty patients with AR (17 females, 13 males) and 30 sex- and age-matched healthy controls (17 females, 13 males) met the eligibility criteria for the study. The mean age was 32.70±10.68
years for the AR group and 34.87±9.40 years for the control group. All patients and controls completed the study.

The mean VHI score of the AR group was significantly higher than those of the control group (p=0.001). No significant differences were found between the AR group and control group in either mean MPT values (p=0.083) or mean Fo values (p=0.825). The s/z ratio was significantly higher in the AR group than the control group (p=0.011) (Table 1). Habitual pitch and MPT values were within the normal limits for both groups.

As for the stroboscopic assessment, no significant differences were found between the AR group and control group in closure level (p=1.000), vocal fold edge (p=0.770), supraglottic involvement (p=0.293), amplitude (p=0.778), mucosal wave (p=0.320), non-vibrating portion (p=1.000), and closure phase (p=0.492) (Table 2).

### DISCUSSION

In the present study, we investigated the acoustic and stroboscopic findings of the larynx and VHI scores in AR patients and compared with normal healthy subjects. The present study showed that no difference was found between the AR and control groups regarding mean MPT values, Fo values, and stroboscopic assessment. However, the mean VHI score and s/z ratio were significantly higher in the AR group.

In the event of AR, several mechanisms are responsible for dysphonia. First, the hypersecretion of the nasal glands causes postnasal drainage: the consequences are cough, throat clearing, and dysphonia (12). Secondly, the rhino-laryngeal reflexes (sympathetic and parasympathetic fibres demonstrated in the musculus vocalis) secondary to AR may contribute to the presence of dysphonia (13). Thirdly, specific receptors sensitive to negative pressure in the nasal cavity and in the pharynx may increase the muscular activity of the posterior cricoarytenoid muscle (14,15). Fourthly, serous otitis diagnosed in nearly 66% of children affected by AR may induce vocal abuse (4). Finally, Gastroesophageal Reflux (GER) is a frequent finding in laryngeal disorders (16). In the present study, patients with GER were excluded from the study.

In the present study, we revealed that patients with AR had a higher prevalence of dysphonia than controls. This is consistent with the findings of previous studies which showed that singers with voice problems were more likely to have AR and that patients with AR who needed allergen immunotherapy were more likely to have dysphonia (12, 17).

The high prevalence of AR in cases with voice disorders causes difficulties in identifying the particular pathogenetic role of AR as an independent variable. Physical examination findings in patients with AR may be normal or nonspecific, such as oedema, redness, or thick mucus labelled as functional dysphonia (14). In the present study, no significant differences were found between the AR group and control group in stroboscopic findings such as closure level, vocal fold edge, supraglottic involvement, amplitude, mucosal wave, non-vibrating portion, and closure phase. However, apart from these findings, it is well known that excess mucous in the larynx causes vocal symptoms. Duncan et al. (18) examined patients with allergy and stated that nearly 33% exhibited laryngeal symptoms such as throat irritation, soreness, burning, and laryngitis. Also they noted that after medication 94% of these patients experienced improvements in their symptoms. Jackson-Menaldi et al. (19)
described 17 allergic patients who complained of concomitant laryngeal symptoms and revealed that vocal fold edema was a common pathological feature among these patients.

Patients with AR have increased excretion of mucus from the nose. Jackson-Menaldi et al. (19) and Sala et al. (20) stated that patients with allergy frequently had thick secretions, hoarseness, and laryngeal oedema/erythema. Unfortunately, these findings were not specific enough to reveal a direct relationship between allergy and laryngeal findings. In the present study, eight of the 30 patients with AR had thick secretions which were thought to be from postnasal drips.

The Voice Handicap Index is a validated tool measuring the impact of vocal problems on quality of life (21). Millqvist et al. (22) and Krouse et al. (23) studied patients groups with allergy, and both showed the negative effect of allergies on voice-related quality of life. Consistent with these studies, the two groups were significantly different from one another on total VHI scores in our study.

In 1971, Boone, who first described the s/z ratio, also hypothesised that cases with normal vocal folds could be expected to prolong the voiceless /s/ and the voiced /z/ phonemes for approximately the same duration of time as the s/z ratio is estimated to be longer than 1.2-1.4 in laryngeal pathologies. In the present study, the s/z ratio was significantly higher in the AR group than in the control group; however, the mean value is within normal limits according to the previously described values. Our finding suggested that patients with AR had laryngeal pathology, so they had difficulty in prolonging the voiced sound /z/ for the same length of time as voiceless /s/ in many studies (3). Therefore, the s/z ratio is estimated to be longer than 1.2-1.4 in laryngeal pathologies. In the present study, the s/z ratio was significantly higher in the AR group than in the control group; however, the mean value is within normal limits according to the previously described values. Our finding suggested that patients with AR had laryngeal pathology, so they had difficulty in prolonging the voiced sound /z/, although the mean s/z ratio was within the normal limits.

In conclusion, our study demonstrated a significant difference in the mean VHI values and the mean s/z ratio between AR patients and healthy controls, indicating a relationship between allergy and dysphonia. Allergy should be considered a potential cause of nonorganic dysphonia and laryngeal symptoms in the diagnostic process.

Ethics Committee Approval: Ethics committee approval was received for this study from the ethics committee of Başkent University.

Informed Consent: Written informed consent was obtained from patients who participated in this study.

Peer-review: Externally peer-reviewed.


Conflict of Interest: No conflict of interest was declared by the authors.

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