Evaluation of atopy in patients with COPD*

Avaliação de atopia em portadores de DPOC

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Abstract

Objective: To determine the prevalence of atopy and to evaluate clinical, laboratory, and radiological profiles in patients with COPD. Methods: This was a cross-sectional study involving outpatients with stable COPD (defined by the clinical history and a post-bronchodilator FEV₁/FVC < 70% of the predicted value). The patients completed a questionnaire regarding clinical characteristics and atopy, after which they underwent nasal lavage cytology, skin prick testing, chest X-rays, arterial blood gas analyses, and determination of total serum IgE. Results: Of the 149 subjects studied, 53 (35.6%), 49 (32.8%), and 88 (59.1%) presented with nasal eosinophilia, a positive skin prick test result, and symptoms of allergic rhinitis, respectively. Correspondence analysis confirmed these findings, showing two distinct patterns of disease expression: atopy in patients with COPD that was less severe; and no evidence of atopy in those with COPD that was more severe (reduced FEV₁ and hyperinflation). There was a statistically significant association between nasal eosinophilia and a positive bronchodilator response. Conclusions: Using simple and reproducible methods, we were able to show that there is a high frequency of atopy in patients with COPD. Monitoring inflammation in the upper airways can be a useful tool for evaluating respiratory diseases in the elderly and in those with concomitant asthma and COPD, a clinical entity not yet fully understood.

Keywords: Pulmonary disease, chronic obstructive; Allergy and immunology; Nasal lavage fluid; Asthma; Rhinitis, allergic, perennial.

Resumo

Objetivo: Determinar a prevalência de atopia e avaliar o perfil clínico, laboratorial e radiológico de pacientes com DPOC. Métodos: Estudo de corte transversal com pacientes ambulatoriais portadores de DPOC estável (definida pela história clínica e relação VEF₁/CVF < 70% do previsto após broncodilatador). Os pacientes responderam um questionário clínico e de atopia e foram submetidos a citologia de lavado nasal, teste cutâneo de alergia, radiografia de tórax, hemogasometria arterial e dosagem de IgE total. Resultados: Dos 149 indivíduos avaliados, 53 (35,6%), 49 (32,8%) e 88 (59,1%), respectivamente, apresentavam eosinofilia no lavado nasal, teste cutâneo positivo e sintomas de rinite alérgica. A análise de correspondência confirmou esses achados, evidenciando dois perfis distintos de doença: a presença de atopia em pacientes com estágios mais leves de DPOC, e a ausência de características de atopia em pacientes com aspectos de doença mais grave (VEF₁ reduzido e hiperinsuflação). Houve uma associação estatisticamente significante entre eosinofilia no lavado nasal, teste cutâneo positivo e sintomas de rinite alérgica. A análise de correspondência confirmou esses achados, evidenciando dois perfis distintos de doença: a presença de atopia em pacientes com estágios mais leves de DPOC, e a ausência de características de atopia em pacientes com aspectos de doença mais grave (VEF₁ reduzido e hiperinsuflação). Houve uma associação estatisticamente significante entre eosinofilia no lavado nasal e prova farmacodinâmica positiva. Conclusões: Este estudo identificou uma alta frequência de atopia em pacientes com DPOC, utilizando ferramentas simples e reproduzíveis. A monitorização inflamatória de vias aéreas parece ser uma ferramenta útil para avaliar as doenças respiratórias em idosos, assim como em pacientes com sobreposição de asma e DPOC, entidade clínica ainda pouco compreendida.

Descritores: Doença pulmonar obstrutiva crônica; Alergia e imunologia; Líquido da lavagem nasal; Asma; Rinite alérgica perene.
Introduction

A progressive inflammatory disease of the lower airways, COPD is characterized by airflow limitation that is not fully reversible.\(^1,2\) The symptoms result from an abnormal inflammatory response of the lungs to the inhalation of noxious particles or gases and are mainly caused by smoking. It is estimated that 10% of the world population over 40 years of age have COPD, with a major impact on the quality of life of patients and on health care systems.\(^1,2\) Despite being preventable and treatable, COPD is the fifth leading cause of death worldwide (2.7 million deaths/year)\(^3,4\) and will become the third one by 2020, according to World Health Organization estimates.\(^1\)

A heterogeneous and multifactorial disease, COPD results from genetic and environmental factors.\(^1,4\) The classical phenotypes are emphysema and chronic bronchitis, which is often associated with bronchial hyperreactivity (BHR). Inflammation of the bronchial wall is characterized by the presence of neutrophils and macrophages and by increased concentrations of IL-8 and Th1 cytokines, as well as by a proteinase/antiproteinase imbalance, and this could explain the poorer response to inhaled corticosteroids (ICs) seen in COPD.\(^5-7\)

The association of COPD with asthma and allergic rhinitis (AR), which are diseases characterized by atopy, with Th2 inflammatory response, eosinophilia, and increased IL-4 levels, has been discussed. This association is more common in the elderly with late-onset asthma and risk factors for COPD (a post-bronchodilator FE\(_{V1}\) < 70% of the predicted value, hyperinflation, and smoking).\(^6\) Patients with this profile can present with a positive skin prick test (SPT) result, increased serum IgE levels, bronchial remodeling, and eosinophilic inflammation, similarly to those with asthma.\(^8-11\) Common to allergic respiratory diseases, BHR has been described as a risk factor for the development of COPD, even without reported asthma.\(^12,13\)

The association of COPD and atopy, despite current evidence, has been discussed since 1960, with the advent of the “Dutch hypothesis”, and has been corroborated by recent studies demonstrating the importance of sputum eosinophilia in patients with COPD and the potential benefits of the use of ICs in such patients. The existence of a subgroup of patients with COPD who have a greater bronchodilator response (a positive bronchodilator response), according to the Dutch hypothesis, is due to the fact that asthma, chronic bronchitis, and emphysema have the same genetic basis, are modulated by the environment, and have varying phenotypic expressions (chronic non-specific lung disease).\(^14\) This hypothesis is supported by the occurrence of COPD in only 10-15% of smokers, supposedly more genetically predisposed to developing COPD.\(^4\)

The relationship between asthma and COPD appears to be bidirectional and complex. Asthma patients who smoke are known to have more frequent exacerbations, a poorer response to ICs, and early functional impairment, and sometimes it is difficult to distinguish between COPD and severe asthma, the inflammation of which can mimic the neutrophilic inflammation of COPD.\(^15\) Paradoxically, a high frequency of eosinophilic bronchitis has been described in patients with stable COPD.\(^9\) There is evidence of a subgroup of COPD patients who have a greater response to high doses of ICs, a positive bronchodilator response, frequent exacerbations, and an FE\(_{V1}\) < 50% of the predicted value. However, it is necessary to establish the profile of these patients more accurately, reducing the side effects and costs related to the use of high doses of ICs in nonresponding patients.\(^16\)

Inflammatory response markers that are useful as indicators for therapy with ICs in patients with COPD have yet to be established; however, studies have revealed that sputum examination by nasal lavage cytology (NLC) can measure the degree of inflammation in asthma, in COPD, and in other lung diseases both qualitatively and quantitatively, with the advantage of being a simple, inexpensive, and reproducible test.\(^17,18\) There is evidence of eosinophilic bronchitis in up to 17% of patients with COPD, although the authors of that study did not conclude whether it was associated with a greater response to ICs.\(^9\) There is an association between eosinophilia on NLC and asthma severity in adults,\(^19\) but it is unclear whether this relationship occurs in COPD. This seems important since recent studies have described the overlap between COPD and asthma (overlap syndrome),\(^9,11,20\) although this association has yet to be further investigated, as well as requiring a clinical and laboratory approach, in order to establish the profile of COPD patients with evidence of atopy.
The Dutch hypothesis suggests that atopy and BHR, which are important markers of asthma, can be involved in the pathogenesis of COPD, although there is no clear evidence regarding the frequency of atopy (including asthma), AR, eczema, or increased IgE levels in patients with COPD, because most studies have involved small samples and limited evaluation of atopy, usually using only SPTs and not evaluating clinical parameters. In addition, the profile of disease severity in these COPD patients with evidence of atopy is unknown. The objective of the present study was to determine the prevalence of atopy in patients with COPD, as well as to establish their clinical, laboratory, and radiological profiles.

Methods

This was a cross-sectional study aimed at estimating the frequency of atopy in outpatients with COPD who were followed in the Department of Pulmonology of the Federal University of Bahia Professor Edgard Santos University Hospital, located in the city of Salvador, Brazil. Patients who were admitted to the outpatient clinic with a diagnosis of stable COPD (any stage) between November of 2008 and March of 2011 and who gave written informed consent were consecutively included in the study. The diagnosis of COPD was defined by clinical history (dyspnea, chronic cough, sputum, pulmonary exposure to noxious particles or gases) and by spirometry (a post-bronchodilator FEV₁/FVC < 70% of the predicted value). We excluded patients who had respiratory tract infection in the previous month, who used nasal or systemic corticosteroids in that period, or who used antihistamines in the previous week. We also excluded patients with a diagnosis of stable COPD (any stage) who were admitted to the outpatient clinic with a diagnosis of stable COPD (any stage) who used nasal or systemic corticosteroids in that period, or who used antihistamines in the previous week. We also excluded patients with COPD who were followed in the Department of Pulmonology of the Federal University of Bahia Professor Edgard Santos University Hospital, located in the city of Salvador, Brazil. Patients who were admitted to the outpatient clinic with a diagnosis of stable COPD (any stage) between November of 2008 and March of 2011 and who gave written informed consent were consecutively included in the study. The diagnosis of COPD was defined by clinical history (dyspnea, chronic cough, sputum, pulmonary exposure to noxious particles or gases) and by spirometry (a post-bronchodilator FEV₁/FVC < 70% of the predicted value). We excluded patients who had respiratory tract infection in the previous month, who used nasal or systemic corticosteroids in that period, or who used antihistamines in the previous week. We also excluded patients with chest X-ray findings of parenchymal lesions not consistent with COPD. The study was approved by the local research ethics committee.

The patients underwent thorough history taking and a complete physical examination, with an emphasis on the clinical parameters of COPD and atopy and on the history of comorbidities. We used the quality of life questionnaire known as Airway Questionnaire (AQ20), which has been validated for use in Brazil. A case definition of AR was established by the presence of nasal symptoms triggered by aeroallergens (at least two of the following: rhinorhea; sneezing fits; congestion, or nasal itching). The patients underwent SPTs, spirometry, determination of serum IgE, arterial blood gas analyses, chest X-rays, and stool examination for parasites. All tests were required during a single visit and were performed at the aforementioned hospital by trained staff within two months after the initial evaluation of each patient.

All SPTs were performed with material from the US Food and Drug Administration, using antigens from Aspergillus fumigatus, Blomia tropicalis, Dermatophagoides pteronyssinus, cat and dog dander, house dust, and histamine. A positive test result was defined as the presence of papules > 3 mm in diameter to at least one antigen. For NLC, eosinophilia was defined as an eosinophil count ≥ 5%, whereas neutrophilia was defined as a neutrophil count ≥ 5%. Infectious vasomotor rhinitis was defined as a cell count > 1,000,000 cells. In addition, the presence of bacteria and fungi was assessed.

The parameters measured by spirometry included FEV₁, the FEV₁/FVC ratio, and response to a short-acting bronchodilator (albuterol challenge), expressed as the difference between pre- and post-bronchodilator FEV₁ divided by the predicted value for FEV₁. A positive challenge response was defined as a ratio ≥ 7%. Arterial blood samples were collected for blood gas analysis, with the patients breathing spontaneously on room air. Serum IgE levels were measured by ELISA. Posteroanterior and lateral chest X-rays were requested for evaluation of hyperinflation, defined as an increased anteroposterior chest diameter, increased retrosternal air space, a reduced/compressed cardiac silhouette, the presence of bullae, vascular attenuation, and flattening of the hemidiaphragms. Three stool samples were requested from each patient for stool examination for parasites, including testing for Strongyloides stercoralis by the Baermann method, in order to rule out parasitic eosinophilia.

The patients were divided into two groups: atopic group (those with a positive SPT result and/or AR symptoms accompanied by eosinophilia on NLC) and non-atopic group (the remaining patients, who did not meet any of the previous criteria).

We used the Statistical Package for the Social Sciences, version 11.0 (SPSS Inc., Chicago, IL, USA). Proportions were compared with the chi-square test or Fisher’s exact test. The Mann-Whitney test for independent samples was used to compare the distribution of quantitative variables between
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the two groups studied. Multivariate logistic regression analysis was performed to obtain adjusted ORs (the Z approximation), and 95% CIs were calculated. For all analyses, the level of significance was set at 5%.

Multiple correspondence analysis (classification technique) was performed to obtain the final model, which was configured on a Cartesian grid with four quadrants, each representing a profile of patients. Each point in the model represents one category of variables, and a closer proximity between points in the same quadrant translates into a higher affinity between the corresponding variables in the sample.

Results

We evaluated 149 patients. The general characteristics of the sample are presented in Table 1. Eight patients were receiving supplemental oxygen via nasal cannula, at a flow of 1–2 L/min, at the time of arterial blood gas analysis and were therefore excluded from it.

Regarding the division of patients into groups, 62 (41.6%) showed evidence of atopy, whereas 87 (58.4%) did not have a positive SPT result or AR symptoms accompanied by eosinophilia on NLC. Among the patients in the atopic group, 49 (79.0%) had a positive SPT result, 38 (61.3%) had AR symptoms with eosinophilia on NLC, and 25 (40.3%) presented with both parameters. The differences between the groups studied are shown in Table 2.

When eosinophilia on NLC and the other study variables were compared, some of the associations found were statistically significant, as shown in Table 3.

Table 4 presents the result of the logistic model with multiple variables, showing adjusted OR values. There were risk relationships between several independent variables and eosinophilia on NLC (dependent variable of the model). Eosinophilia on NLC was found to be positively associated with a positive SPT result and vasomotor rhinitis, the association being statistically significant. The variables Medical Research Council (MRC) dyspnea scale score and smoking history correlated negatively with eosinophilia, but these associations did not achieve statistical significance. Gender and hyperinflation showed a weak association with the dependent variable of the model (OR values close to 1).

Figure 1 shows the multiple correspondence analysis results. The characteristics are found to be distributed into two distinct profiles of patients—one group of patients with variables indicating less severe COPD and with evidence of atopy, such as eosinophilia on NLC, a positive SPT result, and AR symptoms (left lower quadrant of the graph), and one group without evidence of atopy but with indicators of more severe COPD (i.e., MRC scale scores ≥ 4, reduced FEV1, hyperinflation, hypercapnia, longer hospital stays, and a long smoking history) and with a higher proportion of female patients (upper right quadrant).

Discussion

The results of the present study show a high frequency of patients with evidence of atopy, considering SPT results and AR symptoms accompanied by nasal eosinophilia: 41.6% of the patients studied, a value that is higher than that reported in a previous study, which found

| Variable                  | Result       |
|--------------------------|--------------|
| Male gender              | 104 (70.0)   |
| Age, years               | 70.3 ± 8.5   |
| COPD stage               |              |
| I                        | 1 (0.7)      |
| II                       | 37 (24.8)    |
| III                      | 100 (67.1)   |
| IV                       | 11 (7.4)     |
| MRC dyspnea scale        |              |
| 1                        | 8 (5.4)      |
| 2                        | 23 (15.4)    |
| 3                        | 42 (28.2)    |
| 4                        | 34 (22.8)    |
| 5                        | 42 (28.2)    |
| Hyperinflation           | 130 (87.2)   |
| Positive skin prick test | 49 (32.9)    |
| Childhood asthma         | 6 (4.0)      |
| Heart disease            | 30 (20.1)    |
| Diabetes mellitus        | 26 (17.5)    |
| Helminthiasis            | 12 (8.1)     |
| FEV1, L/min              | 1.25 ± 0.6 (0.36–3.56) |
| FEV1, % of predicted     | 50.6 ± 20.7 (16.0–110.0) |
| SaO2, %b                 | 93.5 ± 3.7 (74.0–99.0) |
| PaO2, mmHg               | 76.8 ± 10.9 (46.0–100.0) |
| PaCO2, mmHg              | 42.3 ± 6.2 (28.0–63.0) |
| Total serum IgE, b       | 523.6 ± 728.2 (1.5–4392.0) |

MRC: Medical Research Council. *Values expressed as n (%), except where otherwise indicated. †Values expressed as mean ± SD or as mean ± SD (range).
Table 2 – Clinical, laboratory, and radiological characteristics of the COPD patients studied, by group (atopic vs. non-atopic).a

| Variable                  | Atopic Group (n = 62) | Non-atopic Group (n = 87) | p    |
|---------------------------|-----------------------|---------------------------|------|
| Age, years                | 65.7 ± 8.5            | 69.7 ± 8.4                | < 0.01|
| Male genderb              | 48 (77.4)             | 56 (64.4)                 | 0.09  |
| Height, m                 | 1.63 ± 0.1            | 1.62 ± 0.1                | 0.25  |
| Weight, kg                | 67.5 ± 14.5           | 61.6 ± 13.2               | 0.01  |
| FEV₁, L/min               | 1.4 ± 0.6             | 1.2 ± 0.6                 | 0.03  |
| FEV₁, % of predicted      | 52.8 ± 21.7           | 49.6 ± 20.1               | 0.48  |
| COPD stageb               |                       |                           | 0.65  |
| I and II                  | 17 (27.4)             | 21 (24.1)                 |      |
| III and IV                | 45 (72.6)             | 66 (75.9)                 |      |
| Exacerbations/year        | 4.7 ± 3.5             | 5.8 ± 4.9                 | 0.17  |
| Hospital admissions/year  | 1.5 ± 1.6             | 1.8 ± 1.5                 | 0.17  |
| Length of hospital stay, days | 10.7 ± 14.0         | 17.5 ± 26.7               | 0.20  |
| AQ20 questionnaire, %     | 57.1 ± 24.8           | 60.3 ± 24.2               | 0.44  |
| Smoking history, pack-years | 44.1 ± 18.1         | 47.3 ± 18.6               | 0.24  |
| Positive bronchodilator responseb | 26 (42.6)        | 22 (25.3)                 | 0.03  |
| Hyperinflationb           | 51 (82.3)             | 79 (90.8)                 | 0.12  |
| SaO₂, %                   | 93.7 ± 3.8            | 93.4 ± 3.6                | 0.59  |
| PaO₂, mmHg                | 78.7 ± 10.0           | 75.5 ± 11.4               | 0.19  |
| PaCO₂, mmHg               | 41.7 ± 6.3            | 42.8 ± 6.2                | 0.53  |
| Asthmab                   | 4 (6.5)               | 2 (2.3)                   | 0.20  |
| Nasal lavage fluid characteristicsb |               |                           |      |
| Eosinophilia              | 47 (75.8)             | 6 (6.9)                   | < 0.001|
| Neutrophilia              | 22 (35.5)             | 28 (32.2)                 | 0.67  |
| Bacteria                  | 18 (29.0)             | 22 (25.3)                 | 0.61  |
| Vasomotor rhinitis        | 16 (25.8)             | 22 (25.3)                 | 0.94  |
| Fungi                     | 7 (11.3)              | 4 (4.6)                   | 0.12  |

AQ20: Airway Questionnaire 20. aValues expressed as mean ± SD, except where otherwise indicated. bValues expressed as n (%).

Table 3 – Variables showing statistically significant differences (p < 0.05) between the patients with and without eosinophilia on nasal lavage fluid cytology.a

| Variable                  | Eosinophilia on NLC | p    |
|---------------------------|---------------------|------|
|                          | Yes (n = 53)        | No (n = 96) |
| FEV₁, L/min              | 1.5 ± 0.5           | 1.1 ± 0.6 | < 0.01 |
| FEV₁, % of predicted     | 55.3 ± 22.2         | 48.0 ± 19.5 | 0.04  |
| Positive bronchodilator responseb | 24 (45.3)        | 24 (25.3) | 0.01  |
| Symptoms of allergic rhinitisb | 38 (71.7)        | 50 (52.1) | 0.02  |
| Bacteria on NLCb         | 20 (37.7)           | 20 (20.8) | 0.03  |
| Vasomotor rhinitis on NLCb | 19 (35.8)          | 4 (19.8) | 0.03  |
| Fungi on NLCb            | 7 (13.2)            | 4 (4.2) | 0.04  |

NLC: nasal lavage cytology. aValues expressed as mean ± SD, except where otherwise indicated. bValues expressed as n (%).

17% of cases of atopy among patients with COPD.[9] Although studies have estimated that the worldwide prevalence of AR in adults is 10%,[22] in specific subgroups of patients, such as patients with COPD, the rate has yet to be determined. In the present study, 88 patients (59.1% of the sample) had AR symptoms triggered by allergens, a frequency that is much higher than the world average.[22] However, the fact that the diagnosis of AR is determined on the basis of information...
Table 4 - Logistic model with multiple independent variables in relation to the presence of eosinophilia on nasal lavage fluid cytology.

| Independent variable                      | OR (95% CI)   |
|-------------------------------------------|---------------|
| Positive skin prick test result           | 11.2 (4.72-28.72) |
| Male gender                               | 1.1 (0.43-2.98) |
| Hyperinflation                            | 1.2 (0.33-4.77) |
| Vasomotor rhinitis                        | 4.1 (1.59-11.39) |
| Symptoms of allergic rhinitis             | 2.4 (0.97-6.17) |
| MRC scale score > 3                       | 0.5 (0.18-1.45) |
| Smoking history ≥ 20 pack-year            | 0.6 (0.14-2.23) |
| Length of hospital stay > 1 day           | 0.6 (0.26-1.57) |

MRC: Medical Research Council. Area under the curve = 158; pseudo $R^2$ = 49.9%.

Figure 1 - Correspondence analysis of multiple variables. gen0: female gender; gen1: male gender; fev0: FEV$_1$ ≥ 1 L/min; fev1: FEV$_1$ < 1 L/min; pac0: PaCO$_2$ > 46 mmHg; pac1: PaCO$_2$ ≤ 46 mmHg; sp20: SpO$_2$ > 90%; sp21: SpO$_2$ ≤ 90%; mrc1: Medical Research Council (MRC) dyspnea scale = 1; mrc2: MRC dyspnea scale = 2; mrc3: MRC dyspnea scale = 3; mrc4: MRC dyspnea scale = 4; mrc5: MRC dyspnea scale = 5; hy0: no hyperinflation; hy1: hyperinflation; sh0: smoking history < 20 pack-years; sh1: smoking history ≥ 20 pack-years; spt0: negative skin prick test (SPT) result; spt1: positive SPT result; rs0: no allergic rhinitis (AR) symptoms; rs1: AR symptoms; eo0: no eosinophilia on nasal lavage cytology (NLC); eo1: eosinophilia on NLC; atp0: eosinophilia on NLC and no symptoms of AR; atp1: eosinophilia on NLC and/or symptoms of AR; vr0: no vasomotor rhinitis on NLC; vr1: vasomotor rhinitis on NLC; fn0: no fungi on NLC; fn1: fungi on NLC; ht0: length of hospital stay < 1 day; ht1: length of hospital stay > 1 day; be2: blood eosinophilia; be3: blood eosinophilia.

One group of authors concluded that only half of the cases of rhinitis have an allergic etiology, the remaining cases being classified as nonallergic rhinitis, without systemic manifestations of atopy provided by the patient, although this is the recommendation in major consensus guidelines for rhinitis$^{[22]}$ represents a limitation of the study. In addition, one cannot state that it is in fact rhinitis of allergic etiology. Therefore, we sought to associate this finding from history taking with the laboratory finding of eosinophilia on NLC in order to increase the specificity of the criterion for defining the disease (AR).
(as determined by SPTs or testing for serum IgE specific to environmental allergens). In contrast, other authors have described an entity designated local AR, which seems to account for 25% of the total number of patients with rhinitis in allergy clinics and 40% of patients previously diagnosed with nonallergic rhinitis. Local AR is characterized by an inflammatory response limited to the nasal mucosa, with local production of specific IgE, in the absence of systemic atopy. However, locally, there is inflammation, involving eosinophil degranulation, positive responses to nasal challenge with allergens, and a local Th2 immune response. This entity has been found to be associated with ocular symptoms, childhood onset of symptoms (in approximately 40% of cases), and bronchial asthma. Therefore, these data support the fact that, in our study, subjects with allergen-induced nasal symptoms and eosinophilia on NLC, even without a positive SPT result or increased specific IgE levels, were included in the group of patients with evidence of atopy and those with symptoms but no nasal eosinophilia were removed from this group. The resulting number of patients was still much larger than the populations statistics (38 patients had symptoms accompanied by nasal eosinophilia, and this is equivalent to 25.5% of our sample). These data underscore the association between COPD and atopy, which has been described since 1960.

Atopy, a condition that has an increasing prevalence worldwide and remains little studies in many countries, is associated with worsening of quality of life. In Brazil, data have been collected for the International Study of Asthma and Allergies in Childhood, a population survey assessing the prevalence of allergy in children and adolescents. Disease prevalence was found to be 25.7% in Brazilian schoolchildren and 29.6% in Brazilian adolescents, on the basis of the clinical criterion (symptoms) of disease status. However, the clinical impact of AR and atopy on patients with COPD, especially in terms of quality of life, has yet to be clearly defined. In the present study, there was no significant difference in quality of life, as assessed by the AQ20 questionnaire, between subjects with and without atopy.

Bivariate analysis comparing the groups revealed that atopic patients showed evidence of more severe COPD (less reduction in absolute FEV₁) and greater reversibility of airflow obstruction, as measured by bronchodilator response (p = 0.03). Although they were not statistically significant, differences were found in the frequency of exacerbations, length of hospital stay, hyperinflation, and PaO₂, suggesting less disease severity among atopic individuals, which is in contrast to previous findings. It is possible that these observed trends could have reached statistical significance if the sample size (and, consequently, the power of the study) were larger.

The difference found between absolute FEV₁ values (L/min), which were significantly higher in the atopic group, and the absence of this difference when percentage values were compared can be explained, at least in part, by the fact that this group comprised patients who were younger (a 4-year difference, on average, in relation to non-atopic patients) and heavier (approximately a 6-kg difference, on average). These data could affect the absolute values of this spirometric parameter but not its relationship with the predicted values.

In order to evaluate the influence of upper airway inflammation on the clinical, spirometric, and laboratory characteristics of COPD, we analyzed the variable eosinophilia on NLC and its relationship with the other study variables. Eosinophilia was found to be significantly associated both with higher absolute and higher percentage FEV₁ values. These results suggest that this inflammatory marker is more strongly associated with less severe COPD (less impaired pulmonary function) than is the SPT.

In addition, there was a higher frequency of AR symptoms and other NLC findings in patients with nasal eosinophilia, which indicates agreement between symptoms and laboratory findings in nasal lavage fluid. The fact that patients with nasal eosinophilia had greater bronchodilator response can be explained by the type of airway inflammation and its association with reversibility of airflow obstruction. This finding is consistent with those of previous studies.

Logistic regression analysis confirmed the inverse relationship between atopy and severity of COPD in the study sample—there was an inverse relationship of eosinophilia on NLC with higher MRC scale scores, smoking history, and length of hospital stay. In contrast, a positive SPT result and vasomotor rhinitis correlated positively with nasal eosinophilia, which shows the agreement between these two atopy parameters studied.
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Correspondence analysis confirmed the results of the previous analyses by graphically defining the profile of patients with evidence of atopy and less severe COPD and of those with no evidence of atopy and more severe COPD.

Unpublished data from a multicenter study involving Latin American cities have shown that 13.2% of patients with COPD have a combined pattern of asthma and COPD (overlap). This finding can contribute to the development of strategies for the therapeutic management of COPD. However, only 4% of the patients studied reported a concomitant diagnosis. This can suggest a local characteristic of the patients in the present study, although the sample was similar to that described in the literature in terms of gender, age, severity, pulmonary function, hyperinflation, and smoking history.

Many authors have evaluated markers to identify patients with COPD and atopy (atopic phenotype of COPD), who seem to show a better response to ICs. One group of authors suggested that a definition of COPD on the basis of phenotypic characteristics, instead of the current definition, which is based on severity, be used to define treatment strategies. In addition, there have been reports of side effects of ICs in elderly patients, who are often taking several medications, as well as of the high costs of large-scale use of this treatment approach. Major consensus guidelines for COPD recommend the use of ICs in patients with an FEV₁ < 50% of predicted or with frequent exacerbations of COPD.

Preliminary analyses at our facility revealed that more than 80% of patients used ICs (data not shown). The use of these drugs was not based on bronchodilator response or on evidence of atopy but rather on pulmonary function deterioration (FRV₁ < 50% of predicted) and on the occurrence of frequent exacerbations. There seems to be no difference between the study groups in terms of the use of ICs; therefore, it is unlikely that these drugs could have affected, to some extent, the results obtained. The use of ICs by nearly all patients puts them on a similar footing for evaluation. In addition, the systemic effects associated with ICs are small, compared with those of systemic corticosteroids (excluded from the study), and the influence of these effects on NLC have not been described. Furthermore, the influence of the use of ICs on the study population would be that it would underestimate the finding of an increased prevalence of evidence of atopy, and not the other way around.

The presence of fungi on NLC in 11 patients (7.4%) who were not taking concomitant ICs is relevant because these are uncommon agents, the progression of which has yet to be fully elucidated. There is evidence of fungal sinusitis increases due to previous antibiotic therapy, use of ICs and systemic corticosteroids, and immunosuppression, these microorganisms being associated with greater severity and the need for surgical treatment. With the exception of the use of ICs, none of these risk factors were found in the study population. This underscores the importance of monitoring the upper airways in COPD, even in immunocompetent subjects who are not taking risky medications.

In the present study, we chose NLC over induced sputum cytology, which is associated with complications, especially in the presence of impaired pulmonary function. The NLC technique, which is inexpensive and easy to perform, was used by one group of authors who described the association between asthma severity and the degree of upper airway inflammation. Inhaled nitric oxide, which is a marker of airway eosinophilia, was not used because of its cost and its lower reproducibility. The fact that total IgE varied widely in the present study, although the mean total IgE was much higher than normal, and that it has low specificity in diagnosing atopy explained the exclusion of this variable from the study analyses. Determination of specific IgE was not used because of its high cost.

Compared with previous studies evaluating atopy and COPD, the present study differs in sample size and in that it used several variables to establish evidence of atopy, previously determined only on the basis of positive SPT results and increased serum IgE levels. Our results, together with those of future studies, may inform individualized treatment with the use of ICs in the subgroup of COPD patients with evidence of atopy.

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