Short Communication

Cyn d 1 airborne allergen in a Southern Brazilian city

Cinty Thom de Souza1*, Nelson Augusto Rosario Filho1, Ernesto Akio Taketomi2, Juliana Silva Miranda2 and Ricardo Henrique Moreton Godoi3

1Department of Pediatrics, Federal University of Paraná, Brazil
2Laboratory of Allergy and Clinical Immunology, Biomedical Science Institute, Federal University of Uberlândia, Brazil
3Department of Environmental Engineering, Federal University of Paraná, Brazil

By researching the factors related to exposure to indoor and outdoor allergens, such seasons, climate changes and particulate matter, allergists can screen the sensitization profile of individuals according to their exposures and conduct preventive treatment and individualized immunotherapy. Molecular allergology has improved aerobiological screening of allergenic components toward more specific results on allergic exposure, sensitization, and symptoms [1,2]. The Enzyme-Linked Immunosorbent Assay (ELISA) is a colorimetric enzyme immunoassay technique used to quantify soluble substances such as proteins, peptides, antibodies, and hormones. Due to its high sensitivity and specificity, ELISA can quantify substances at low concentrations, such as allergens [3].

In Brazil, pollinosis, also known as hay fever or seasonal allergic rhinitis, is mainly attributed to Poaceae grasses [4]. In the 1970s, the first case reports of the disease appeared in the southern region of Brazil, in individuals with sensitization to pollen extracts and spring symptoms of allergic rhinitis and conjunctivitis [5]. A local air pollen count revealed grass pollen between September and December, the spring season, with a peak on the second half of November [6]. In the temperate climate of southern Brazil, Lolium multiflorum (rye grass) was found to be the main allergenic sensitizing grass. The climate in southern Brazil consists of lower temperatures during winter that precede an exuberant pollen season in spring. Subtropical grass species like Cynodon dactylon (Bermuda grass) and Paspalum notatum (Bahia grass) became increasingly prominent due to their invasiveness and ubiquity [7]. In addition, climate change, global warming, and air pollution are gradually contributing to an intense growth and distribution of allergenic plants, prolonged pollination periods, and an increased pollen allergenicity [8,9].

Group-1 grass pollen allergens are present in all species of Poaceae. However, Group-1 allergens have limited cross-reactivity between subfamilies of temperate and subtropical climate grasses. Cyn d 1 is a major allergen of C. dactylon, followed by Cyn d 4. Phl p 1 and Phl p 5 are the major Pooidae pollen allergens. In light of the high allergenic homology between Phleum pratense and L. multiflorum, the allergenic components of the former may be interpreted to sensitization to the latter [10,11]. Through a series of Immunoglobulin E (IgE) microarray analyses, Araujo, et al. [12] demonstrated sensitization to several of allergens in children with asthma and allergic rhinitis in Curitiba, southern Brazil; 16.8% of those children were sensitized to Cyn d 1, while 14.8% were sensitized to Phl p 1 and 12.9% to Phl p 4. Six out of 17 children were only sensitized to Cyn d 1 and Phl p 4, suggesting true sensitization to C. dactylon.

This study was prompted by the broad distribution of Bermuda grass in subtropical and tropical areas, and the growing sensitization rates to Cynodon dactylon in some areas of Brazil. The present study, the first of its kind in Brazil, was undertaken to determine airborne Cyn d 1 in Maringá, a large city in southern Brazil. To this end, we obtained air samples using a total particulate impactor placed outdoors, 90 cm from the ground, with a flow rate of 1,13 m³ of air/hour. The collection point was an urban area, at latitude 23.41 S,
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longitude 51.97 W, 537 m above sea level. The samples were collected from March 2017 to March 2018. Seventy filters were collected during the 12 months with a median interval of 5 days (ranging from 1 to 14 days). To elute the filters, small pieces (3 cm²) were incubated with f NH4HCO3, 0.1% BSA (bovine serum albumin) and 0.1% Tween 20® (polysorbate 20) in an orbital shaker overnight at 4 °C. The solution was then centrifuged twice at 4000 rpm for 5 minutes and the supernatants extracted and stored at -80 °C. Finally, the samples were analyzed for Cyn d 1 (ALK-Abelló®) via ELISA, a method with highly specificity and a detection limit of 0.24 ng/mL [13]. The test was performed with samples, controls and blank filter, all in duplicate.

We also collected daily meteorological data from the city of Maringá during the study at the weather station (OMM:83767, latitude 23,4 S; longitude 51,91 W; 542 m above the sea). Temperature, relative humidity, rain precipitation, insolation, wind speed and direction were available for analysis.

From the total of filters, 10 were excluded from the analysis due to technical issues during sampling. Therefore, we analyzed 28 filters collected during a 24-hour period and 32 filters collected during a 48-hour period. The 60 samples were tested for Cyn d 1 using the ELISA method. None of the samples showed any detectable levels.

To this day, the ELISA method used for Cyn d 1 in air samples has been an unprecedented technique. To ensure that the material used in the ELISA was working and that the elution method employed were reliable for extracting Cyn d 1, we carried out other assays. We tested the pure allergenic extracts (Immunotech®) of Cynodon dactylon and Paspalum notatum as positive and negative controls, respectively. Cyn d 1 levels of 50 ng/mL were found in the Cynodon dactylon extract, and 2.28 ng/mL in the Paspalum notatum extract. In order to clarify whether the elution method was effective for Cyn d 1, the diluted Cynodon dactylon extract (1:1,000) was applied to the blank filter, and after drying, it was used the elution protocols. Cyn d 1 was detectable in concentrations ranging from 1.3 to 3.2 ng/mL, which proves the reliability of the method.

The lack of Cyn d 1 allergen in any of the sample must be considered. The use of only Cynodon dactylon allergen for analysis introduces a bias in the study since Cyn d 4, which is also considered a major allergen of this species, was not analyzed. Another aspect to discuss is the number of sampled days. There were 60 samples collected over a one-year period from a single site in the city. These samples could not be representative of an entire year and the allergen could have been detected in other days. However, this hypothesis is unlikely if we consider that the meteorological data found during the 60 days of sample collection were equivalent to those found throughout the year (p > 0.05, CI 95%, t - student test). Thus, one site of air sample collection is enough to capture pollens of a region, since they can travel hundreds of kilometers, regardless of the distance of their source [14].

Given that studies have found an increasing sensitization to Cynodon dactylon in the Brazilian population [12,15], the allergic sensitization to Cynodon dactylon found in the region of Maringá is incoherent with the undetected Cyn d 1 levels. We suggested two hypotheses to justify these results. The first one is that sensitization could occur even in the presence of low concentrations of this allergen. In aeropallinological studies conducted in Caxias do Sul, another southern Brazilian city, the presence of pollen grains was detected throughout the year [16]. Recently, in Curitiba, grass pollen dispersion was observed from August to April [9]. This pattern of pollen dispersion is most evident in subtropical and tropical areas, given that there are no well-defined seasons between the Tropic of Capricorn and equator [17].

The second hypothesis for this result was that the presence of Cyn d 1 serum-specific IgE antibodies may occur at low levels in individuals with high positivity to Phl p 1 due to cross-reactivity to group 1, without representing proper sensitization to Cynodon dactylon [18]. In addition, the allergenic component used in this test is a natural allergen (nCyn d 1) glycosylated in the amino-terminal portion, which may imply a specific IgE response to nCyn d 1 in populations not exposed to Cynodon dactylon. The glycosylation present in nCyn d 1 is responsible for the production of IgE to cross-reactive carbohydrates determinants (CCD), epitopes in several plant species and insect venoms that determine a sensitization to clinically irrelevant antigens [19].

Some limitations to the study are that the samples came from a single location in the city and reflect only that point. The sampled days were chosen based on convenience. Excessively rainy days were automatically ruled out since the device could not be exposed to heavy rain. The 48 –hour samples in particular may have shown protein losses due to the long sampling period in higher-than-average temperatures. The absence of detectable Cyn d 1 in the air should be viewed with caution, considering that the allergenic grass is present in the region and people are sensitized to this type of grass.

Despite the undetectable levels of Cyn d 1 in the present study, this was the first attempt at using the ELISA method to identify this Cynodon dactylon allergen in the environment. Environmental studies of aeroallergens are essential to the practice of allergists because it enables us to understand the patient’s environmental exposure, identify risks, introduce preventive measures, and draw up personalized treatment. We recommend that this study be reproduced with different components to highlight the diversity and degree of allergenic exposure of individuals in the environment that they live.

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Author contributions

CCTS and JSM contributed with data collection, interpretation and analysis of data, as well writing; NARF contributed with analysis of data, revision and content of this manuscript. EAT and RHMG contributed with critical revision and content of this manuscript. All authors have read and agreed to the published version of the manuscript.

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