Blomia tropicalis Blo t 5 and Blo t 21 recombinant allergens might confer higher specificity to serodiagnostic assays than whole mite extract

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Abstract

Background: Blomia tropicalis is a dust mite and an important source of allergens in tropical regions. Up to now, the assays to diagnose atopy to this mite use whole body extract as antigens. However, anti-\(B.\) tropicalis IgE antibodies cross-react with Ascaris lumbricoides antigens, hindering the diagnosis of allergy to this mite. In this study, \(B.\) tropicalis recombinant allergens were evaluated with the purpose of developing an immunodiagnostic assay for allergy to this mite with greater specificity than those commercially available.

Methods: Two \(B.\) tropicalis allergens (Blo t 5 and Blo t 21) were cloned into a plasmidial expression vector, expressed in Escherichia coli and purified by affinity chromatography. Sixty-three sera containing anti-\(B.\) tropicalis extract (BtE) IgE antibodies were used to investigate IgE reactivity to the recombinant Blo t 5 and 21 allergens. Inhibition assays with 20 sera pre-adsorbed with \(A.\) lumbricoides extract were performed using rBlo t 5, rBlo t 21, and BtE as antigens. All the assays were carried using indirect ELISA.

Results: Eighty-two point nine percent and 80.0% of the sera with anti-BtE antibodies from 35 children reacted with rBlo t 5 and rBlo t 21, respectively, whereas 92.8% and 89.3% of the 28 sera with anti-BtE antibodies from adult asthma patients reacted with the same allergens, and 96.4% of these sera reacted with a mixture of rBlo t 5 and rBlo t 21. In an inhibition ELISA, the absorption of sera by \(A.\) lumbricoides extract affected less the reaction with rBlo t 5 and rBlo t 21 than with BtE.

Conclusions: The rBlo t 5 and rBlo t 21 allergens contain important epitopes recognized by IgE antibodies of individuals allergic to \(B.\) tropicalis antigens. Moreover, the assays using the recombinant allergens had lower IgE cross-reactivity with \(A.\) lumbricoides antigens, a fact which would confers higher specificity to serodiagnostic assays than the crude mite extract. However, additional recombinant allergens should be evaluated in order to reach the same sensitivity of the commercially available assays based on mite extract.

Keywords: Blomia tropicalis, Recombinant allergens, Immunodiagnosis, Cross-reactivity, Ascaris lumbricoides, Sensitivity, Specificity

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Background

Allergic respiratory diseases, like asthma and rhinitis, are worldwide spread and their prevalences have increased over the past decades, reaching epidemic proportions [1]. Mite-allergen sensitization is a well-documented risk factor for asthma and allergic diseases in atopic individuals. In tropical regions of the world, Blomia tropicalis and Dermatophagoides pteronyssinus are predominant in house dust and are commonly found together [2,3].

B. tropicalis mite is the major source of indoor allergens in the northeastern cities of Brazil [4,5]. Clinical history, skin prick test (SPT) results and specific IgE detection are the triad for the diagnosis of allergic diseases. The majority of the methods for detecting anti-allergen IgE antibodies is based on the use of natural allergenic extracts, a fact that may compromise the interpretation of their results. Natural allergen extracts are highly complex molecular mixtures, containing many proteins, of which only some have allergenic properties, and with their relative contents varying greatly from one extract to another [6,7]. Caraballo and collaborators [2] described at least 25 IgE-binding protein bands in B. tropicalis crude extract. Around 12 recombinant allergens have been molecularly and immunologically characterized and deposited into the allergen database, according to International Union of Immunological Societies (IUIS) Allergen Nomenclature (www.allergen.org). Among these allergens, it includes Blo t 1 (cysteine protease, 26 kDa), Blo t 3 (trypsin protease, 25 kDa), Blo t 5 (unknown molecular function, 14 kDa), Blo t 11 (paramyosin, 110 kDa), Blo t 12 (unknown molecular function, 14 kDa), and Blo t 21 (unknown molecular function, 13.2 kDa), which are present in 50% or more of the sera from B. tropicalis-sensitized individuals [3,4]. Blo t 5 and Blo t 21 represent the major allergens in B. tropicalis mite. These allergens share some structural similarities determined by NMR consisting of three anti-parallel α-helices, assembled in a helical bundle. Despite the fact that some putative IgE epitope residues are conserved in both Blo t 5 and Blo t 21 three-dimensional structures, these allergens present a low to moderate cross-reactivity [5,8].

Nevertheless, the IgE-binding properties to these allergens may vary depending on several factors, such as genetic background of the study population, frequency of mite sensitization, and intrinsic molecular characteristics of the allergens, such as isoform expression profile [3]. Furthermore, in tropical regions of the world, where the high prevalence of helminth infections is a public health problem, the serodiagnosis of allergic diseases may be compromised by cross-reactive antibody to epitopes shared by mite and helminths [9,10].

In this study we produced B. tropicalis recombinant allergens (rBlo t 5 and rBlo t 21) in a prokaryotic system, based on sequences obtained in the GenBank database. A total of 63 sera with IgE antibodies reacting with B. tropicalis extract (BE), from asthmatic patients and from atopic children from Salvador, a large urban center in northeastern Brazil, were used to investigate IgE reactivities to the recombinant Blo t 5 and Blo t 21 allergens by indirect ELISA. The cross-reactivities of IgE antibodies between B. tropicalis (crude extract and recombinant allergens) and A. lumbricoides (crude extract) were also evaluated.

Methods

B. tropicalis and A. lumbricoides extracts

An ether-treated extract of B. tropicalis mite, obtained from house dust samples in Salvador, was prepared according to Baqueiro and collaborators [11]. Briefly, B. tropicalis was collected from bed dust in Salvador, Brazil, cultured with a fish food medium, at 25°C and 75% humidity and cloned. The mites were purified from the medium by flotation on a 5 M sodium chloride solution, followed by several washings by filtration in a 100 μm-pore polystyrene sieve with endotoxin-free distilled water. The washings were carried out until no food residues were seen under microscopy. The mites were lysed in 0.15 M phosphate-buffered saline, pH 7.4 (PBS), in an electric blender (Waring Commercial, Torrington, CN, USA). Lipids from the lysate were extracted by five or six ether extractions and discarded. The protein content of the B. tropicalis aqueous extract was determined by the Folin reagent method [12]. The extract was standardized using a commercially available capture ELISA for detection of Blo t 5 allergen. (INDOOR Biotechnologies, Charlottesville, VI, USA), with the Blo t 5 content in the extract established at 200 ng per mg of total protein, and subsequently aliquoted and cryopreserved until use. The A. lumbricoides aqueous extract was prepared from adult worms obtained from infected albendazol and bisacodyl-treated children in Salvador, Brazil, according to Ponte and collaborators [9].

B. tropicalis recombinant allergens

Sequences encoding B. tropicalis allergens (Blo t 5 and Blo t 21) were obtained from their deposits in the GenBank database. The selection criterion was based on published analysis of the frequency of allergic patients’ sera with IgE antibody reactivity to the recombinant allergens (50% or more reactivity with the tested sera) [3,4]. Pairs of primers (5′-3′) were designed based on the published full length nucleotide sequence of Blo t 5 and Blo t 21 (GenBank: U59102 and GenBank: AY800348, respectively), which allowed their amplification by polymerase chain reaction (PCR) flanked by restriction enzyme sites for BamHI/Xhol (Blo t 5, forward – CCGATCCATGAAAGTTCCGCA TCGTTCC; reverse - GGGCTCGAGTATTGGTTTGG AATATC) or BamHI/HindIII (Blo t 21, forward - CC CGATCCATGAAATTATCATCGCATTT; reverse –
GGGAAGCTTCTATTCGGAATCTTGGAC). As a B. tropicalis cDNA library had already been constructed in our laboratory, an aliquot of it was used as template for the PCR. The Blo t 5 and Blo t 21 amplicons were ligated to the pRSETA expression vector and used for transforming DH-10 B-strain Escherichia coli. The bacteria were cultivated and plasmid DNAs were obtained and sequenced. All sequences were aligned and homology analysis was performed using the BLAST network service provided by NCBI. Constructs containing Blo t 5 and Blo t 21 were then used to transform BL21(DE3) E. coli BL21 (DE3)pLysS, generating His (histidine)-tagged proteins (His-rBlo t 5 and His-rBlo t 21) after induction with 0.1 mM isopropyl β-D-thiogalactoside (IPTG). The recombinant proteins were subsequently purified by affinity chromatography using the Ni-NTA agarose resin (QIAGEN Biotecnologia Brasil Ltda; São Paulo, SP, Brazil), according to the manufacturer’s protocol.

**Polyacrylamide gel electrophoresis and IgE immunoblot assays**

His-rBlo t 5 and His-rBlo t 21 recombinant proteins were subjected to polyacrylamide gel electrophoresis in the presence of sodium dodecyl sulphate (SDS-PAGE; Merck & Co., Inc., White House Station, NJ, USA), according to Laemmli [13], using a Mini-PROTEAN III Electrophoresis Cell (Bio-Rad Laboratories, Hercules, CA, USA), followed by electroblotting onto nitrocellulose membrane (Sigma, St Louis, MO, USA). Immunoblotting was then carried out to determine the reactivities of anti-allergen serum IgE antibodies to these fusion proteins. Briefly, the blots were blocked with 0.15 M phosphate-buffered saline, pH 7.2 (PBS) containing 0.05% Tween (PBS/T) and 10% (w/v) dry non-fat milk for at least two hours at room temperature, followed by overnight incubation with continuous shaking at 4°C with human sera diluted 1:10 in PBS/T containing 5% dry non-fat milk. The blots were incubated with mouse anti-human IgE:HRP monoclonal antibody (ABCAM, Cambridge, UK) at room temperature for one hour. The reaction was developed using H₂O₂ and diaminobenzidine (DAB) as substrate and chromogen, respectively (Sigma, St Louis, MO, USA). Between all steps, the blots were washed three times with PBS/T followed by two washes with PBS alone.

**Serum samples**

Thirty five of the 63 studied sera were obtained from atopic children living in Salvador, Bahia, Brazil. Atopy in these children was characterized by a positive result in the ImmunoCAP assay (Phadia Diagnostics AB, Uppsala, Sweden) for at least one of four tested allergens (from B. tropicalis, D. pteronyssinus, Blattella germanica and Periplaneta americana). The children, aged 4 to 11 years, were enrolled in the SCAALA (Social Change in Asthma and Allergy in Latin America) program, a study performed in 2005 to study risk factors for asthma and allergies in children [14]. The 28 remaining sera were from adult patients with moderate to severe asthma attending the outpatient facility of the Centro de Estudos em Alergias Respiratórias (CEAR), Salvador, Brazil. The cross-reactivity between B. tropicalis recombinant allergens and A. lumbricoides antigens was assessed in 20 adult sera from the CEAR patients. Informed consent was obtained from the SCAALA project children’s parents or guardians and ethical approval was granted by the Instituto de Saúde Coletiva of the Universidade Federal da Bahia, Salvador, Brazil, and by the National Commission on Ethics in Research (CONEP), Brazil. Written consent was obtained from all adult asthma patients and ethical approval was provided by the Ethical Committee of the Maternidade Clínico de Oliveira, Universidade Federal da Bahia, Salvador, Brazil (protocol no. 044/2010).

**ELISA for human IgE antibody reactivity with B. tropicalis recombinant proteins**

ELISA was performed in microassay plate (Costar, Bethesda, MD, USA) wells, which were incubated with 100 μg of B. tropicalis extract or 5 μg of each fusion proteins (His-rBlo t 5 or His-rBlo t 21) when tested individually or 2.5 μg of each when used mixed together in the same solid phase (rBlo t 5 plus rBlo t 21), per mL of sodium carbonate-bicarbonate buffer, pH 9.6, overnight at 4°C. The reaction was developed as described previously [9] and the IgE cut-off for the BtE, rBlo t 5 or rBlo t 21 assays defined according to the mean plus two standard deviations of the results obtained using sera from 10 individuals without history of allergy and with negative SPT reaction to the tested allergens [9].

**Detection of anti-B. tropicalis IgE antibodies cross-reactive with A. lumbricoides antigens**

Inhibition ELISA was performed to investigate the presence of anti-B. tropicalis extract, anti-rBlo t 5, and anti-rBlo t 21 IgE antibodies that cross-reacted with Ascaris lumbricoides antigens. The inhibition was accomplished by pre-incubating the sera with 0.3 or 3 μg/mL of A. lumbricoides extract. These concentrations were chosen based on previous work [9] and are within the concentration range that led to the greatest differences between non-absorbed and absorbed sera. As control for the specificity of the inhibition of the binding of anti-B. tropicalis IgE antibodies, total IgE was determined in the same sera, before and after absorption with Ascaris extract. The assay was carried out as described elsewhere [9]. Results were expressed in percentage of reduction and specific sIgE levels in sera non-adsorbed and adsorved with Ascaris extract (OD 450 nm). The percentage of reduction in anti-BtE anti-rBlo t 5 and anti-rBlo t 21 IgE
antibodies or total IgE level = \[1 - (\text{mean OD obtained in duplicates of serum that had been pre-incubated with } A.\ lumbricoides \text{ antigen/mean OD obtained in duplicates of untreated serum}) \] \times 100.

**Statistical analysis**

The D’Agostino and Pearson test was used for verification of the data normality. The significance of the differences among the different allergens was assessed by the Wilcoxon signed rank test (when the data distribution was found to be non-Gaussian). Differences with a P value equal to or less than 0.05 were considered statistically significant.

**Results**

Expression and validation of recombinant fusion proteins

*B. tropicalis* recombinant allergens were expressed and purified as full length fusion proteins containing N-terminal His-tags (6 His). Once the fusion proteins were purified, their tags (6 His) were not removed. The achieved yields of rBlo t 5 and rBlo t 21 were 3.1 to 4.6 mg and 2.9 to 4.5 mg per liter of bacterial culture, respectively. Each His-tagged (rBlo t 5 and rBlo t 21) proteins produced an expected band in SDS-PAGE (Figure 1A and B). Validation of recombinant fusion proteins was performed by IgE immunoblotting, in which antibodies from atopic and asthmatic individuals, but not negative control sera, were found to react with rBlo t 5, rBlo t 21.

**Antigenicity of B. tropicalis recombinant proteins**

Indirect ELISA was used to evaluate the reactivity of IgE antibodies with the fusion His-rBlo t 5 and His-rBlo t 21 fusion proteins. The IgE reactivity profile of the tested sera with the recombinant proteins is summarized in Table 1. The panel of 35 sera from children with IgE antibodies against BtE had IgE antibody reactivity percentages of 82.9%
(29 sera out of 35) for rBlo t 5 and 80% (28 out of 35) for rBlo t 21. The IgE reactivity percentages for the 28 sera from adult asthma patients with IgE antibodies against BtE were 92.8% (26 out of 28) for rBlo t 5 and 89.3% (25 out of 28) for rBlo t 21. When the indirect ELISA assay was performed using a mixture of rBlo t 5 and rBlo t 21 allergens, a positive reaction was observed in 96.4% (27 out of 28) of the sera (Table 1). Despite the fact that some sera reacted with BtE but not with the recombinant allergens, the levels (expressed by OD450 nm) of IgE antibodies against rBlo t 5 in the children group, and of IgE antibodies in the adult asthmatic patient group as detected in assays using rBlo t 5 or a mixture of rBlo t 5 plus rBlo t 21, were significantly higher than in the assay using BtE (p < 0.0067; p < 0.0314 and p < 0.0034, Wilcoxon signed rank test; Figure 2, A and B). Although the percentages of IgE antibody reactivity to the recombinant antigens were higher in the adults’ serum group than in the children’s serum group, there were no statistically significant differences between these two populations (p > 0.05, Mann-Whitney’s test, data not shown). Of the total sera panel, 66.7% (42 out of 63 children’s and adults’ sera) reacted with both rBlo t 5 and rBlo t 21; 92.1% of these sera (58 out of 63) reacted with at least one of the recombinant allergens and 7.9% (5 out of 63) recognized only the B. tropicalis crude extract (data not shown).

### Cross-reactivity of IgE antibodies against B. tropicalis extract and recombinant allergens with A. lumbricoides antigens

There was a higher reduction in the levels of anti-B. tropicalis crude extract IgE antibodies (median inhibition of 39.2%) than in the levels of anti-rBlo t 5 (median inhibition of 3.6%) and anti-rBlo t 21 (median inhibition of 0.1%) IgE antibodies, when they were incubated with A. lumbricoides extract at both concentrations (Figure 3A). However, the difference between the recombinant protein and the BtE assays was statistically significant only with the concentration of 3 μg/mL of A. lumbricoides extract (p < 0.0052 for rBlo t 5 and p < 0.0020 for rBlo t 21; Wilcoxon signed rank test). As expected, the anti-B. tropicalis crude extract IgE antibody levels were lower in sera pre-adsorbed with A. lumbricoides extract than in the non-adsorbed sera (Figure 3B). This difference between the non-adsorbed and adsorbed sera was statistically significant for both concentrations, 0.3 μg/mL and 3 μg/mL of A. lumbricoides extract (p < 0.0217 and p < 0.0012; Wilcoxon signed rank test). Furthermore, there was also statistically significant difference between sera adsorbed with 0.3 μg/mL and 3 μg/mL of A. lumbricoides extract (p < 0.0011; Wilcoxon signed rank test). The proportion of total IgE that was reduced by incubation with A. lumbricoides natural extract (median inhibition of 0.1%) was lower than the proportion of anti-B. tropicalis extract IgE antibodies that was reduced by the same treatment (p < 0.0008, Wilcoxon signed rank test; Figure 3).

### Discussion

The B. tropicalis mite has been the target of several studies highlighting its role as being one of the major asthma sensitizing agent in tropical areas of the world [15-17]. These studies have demonstrated that serum IgE antibodies in 42% to 98% of B. tropicalis-sensitized individuals react with Blo t 5 and Blo t 21 [2,14]. In the present work it is shown that a large proportion of serum IgE antibodies from BtE-sensitized children and adults with asthma reacted with rBlo t 5 and rBlo t 21 obtained as described herein. The rBlo t 5 in acidic pH solutions, like under purification denaturing conditions, tends to form molecular aggregates [8]. Likely, the rBlo t 21 behave like the rBlo t 5, forming aggregates under the same conditions, since both allergens share structural similarities [5]. This could explain the nonspecific band recognition by antibodies from asthmatic patient sera that were used in immunoblotting assay of this work. B. tropicalis is found more frequently (71.8%) in bed dust in Salvador, a major city in Northeast of Brazil [18]. In environments where B. tropicalis predominates in the mite fauna, it is common to find a high rate of co-sensitization to Blo t 5 and Blo t 21, as highlighted in our study (66.7% of 63 sera from B. tropicalis sensitized individuals) [4,19]. Although several

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### Table 1 Reactivity of anti-B. tropicalis IgE antibodies, assessed by ELISA, using recombinant proteins (rBlo t 5 and rBlo t 21) and crude extract as antigens, in sera containing anti-B. tropicalis extract IgE antibodies

| Antigens                  | N°. of sera with positive results (%) | N°. of sera with negative results (%) |
|---------------------------|--------------------------------------|--------------------------------------|
| rBlo t 5                  | 29 (82.9)                            | 6 (17.1)                             |
| rBlo t 21                 | 28 (80.0)                            | 7 (20.0)                             |
| rBlo t 5                  | 26 (92.8)                            | 2 (7.2)                              |
| rBlo t 21                 | 25 (89.3)                            | 3 (10.7)                             |
| rBlo t 5 + rBlo t 21      | 27 (96.4)                            | 1 (3.6)                              |

Sera from 35 children with anti-B. tropicalis extract IgE antibodies

Sera from 28 adult asthma patients with anti-B. tropicalis extract IgE antibodies

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individuals had a higher IgE antibody reactivity to rBlo t 5 and rBlo t 21 than to BtE (i.e., their sera led to higher optical densities when tested in ELISA for IgE antibodies against these recombinant allergens than against the extract; Figure 2), there is a small percentage of sera (7.9%, corresponding to 5 out of 63 sera) that does not recognize any of the recombinant allergens, a fact that would limit the sensitivity of assays using these allergens. When the two recombinant allergens were used together in the same immunoassay, the IgE reactivity increased so that 96.4% of the sera from asthmatic patients had anti-BtE antibodies. Some studies indicate that mite allergy could be detected using a mixture of two or more major recombinant allergens (component-resolved diagnostics), replacing the natural extract [20]. Furthermore, the use of multiple recombinant allergens to detect sensitization profiles could be very important to guide and improve immunotherapy for mite allergies [19-21].

This study, despite not comparing the reactivity of IgE antibodies to rBlo t 5 and rBlo t 21 with the reactivity to their native counterparts, suggests that recombinant allergens produced as described in this paper can replace natural allergen extracts in the diagnosis of allergies, confirming previously published data showing that rBlo t 5 expressed in E. coli and native rBlo t 5 have comparable IgE reactivity in terms of percentage of sera with antibodies [22]. However, our data also indicate that it is desirable to introduce other recombinant antigen(s), in addition to rBlo t 5 and rBlo t 21, in an assay for IgE antibodies to diagnosis hypersensitivity to *B. tropicalis* in order to increase its sensitivity. Blo t 7, which has been shown to be as reactive as Blot t 5 with IgE antibodies in the sera from allergic children in a study also carried out in a tropical environment [19], may be a good candidate to be included to a pool of recombinant antigens to increase the sensitivity of an immunoassay. Meanwhile, Blo t 10 and Blo t 11, which shown to cross-react with *A. lumbricoides* [23], would not be good candidates to be included in an immunoassay with multiple recombinant antigens.

The relationship between house dust mite sensitization and triggering of asthma and other atopic diseases is well documented [24,25]. Given the difficulty of completely eliminating the sensitizing agents of the affected individuals’ homes, immunotherapy has been playing a key role in alleviating the clinical aspects of allergic diseases. Thus, a more specific mite allergy diagnosis is necessary and it is also a key point to develop appropriate, more specific and individualized therapies [26]. The antigenic extracts used in the diagnosis and immunotherapy of allergic diseases are obtained from natural sources. This fact brings many disadvantages, such as the presence of large amounts of non-antigenic proteins; contamination by other potentially immunostimulating compounds, like endotoxin; allergen variability in the sample composition (depending on the season of the year, the mite life cycle and differences in protein extraction protocols). In addition, the more sensitizing antigenic proteins in the mite extract can be found in low concentrations, requiring the use of higher doses of total extract, which is not always convenient in clinical practice [20,25,27,28].

In addition to the problems described above with the use of whole allergenic extracts in the immunodiagnosis of allergic diseases, there is another factor that should be considered by the companies that commercialize these products: the common association of *B. tropicalis* sensitization with helminth infection in tropical and underdeveloped regions of the world. It is in fact believed that
about 1.5 billion people worldwide are infected with *A. lumbricoides* [10,29]. Salvador city, where the donors of the sera used in the present work live, is one helminth parasite endemic areas in Brazil, having high prevalence of *A. lumbricoides*, *Trichuris trichiura* and *Toxocara* spp (*Toxocara canis* and *Toxocara cati*) infections. In fact, most of the sera studied in the present work had anti-*Ascaris* IgG antibodies, indicating present and/or past *A. lumbricoides* infection. In this area, an overlap of parasitic endemicity and high prevalence of allergic diseases is observed [30-32]. There is often dissociation between positive skin prick test results and detectable serum IgE antibodies in allergic individuals who are also co-infected with *A. lumbricoides* [9]. This finding could be explained by the existence of IgE antibodies, raised in response to the helminth infection, that cross-react with allergens but are unable to lead to degranulation of mast cells, consequently increasing anti-allergen IgE antibody levels but not SPT positivity [9,33]. The increase of knowledge on cross-reactive IgE antibodies to common
environmental respiratory allergens and helminth antigens will certainly improve allergy diagnosis and perhaps even immunotherapy. In the present study, pre-incubation of sera with *A. lumbricoides* extract led to varying degrees of inhibition of the binding of IgE to mite allergens. It would also be useful to further demonstrate this cross-reaction by inhibiting the binding of IgE antibodies to *A. lumbricoides* extract by incubation of the sera with *B. tropicalis* antigens. As the high levels of IgG antibodies raised in the immune response to this helmint hinder the detection of IgE anti-Ascaris antibodies, a possible way to carry out the experiment would be to use an assay to detect IgE antibodies more sensitive than the one used in the present work.

The inhibition by adsorption with *A. lumbricoides* extract was higher for the binding of antibodies to *B. tropicalis* crude extract than to rBlo t 5 and rBlo t 21, although in a few sera there was also a high level of IgE cross-reaction between the helmint extract and the recombinant allergens. Cross-reactivity to both recombinant allergens was observed in the same sera (data not shown). This finding could be explained if the anti-Ascarias antibody response in these sera donors had been more polyclonal than the immune response of the other serum donors, so that antibodies against additional cross-reactive epitopes would be produced. Indeed, it is known that the repertoire of the antigens that are recognized in complex antigenic mixtures may vary greatly in different individuals of the same species [34].

The observed difference in the degree of adsorption by *A. lumbricoides* extract of anti-rBlo t 5 and anti-rBlo t 21 in relation to anti-BtE antibodies cannot be explained by the presence of lower levels of anti-recombinant allergen antibodies, since the amounts of anti-rBlo t 5 IgE antibody activity, on the contrary, were found to be higher than those of anti-BtE IgE antibody activity. Due to this less intense cross-reactivity, assays using a mixture of rBlo t 5 and rBlo t 21 as antigen should be more specific than assays using the crude extract.

IgE antibodies raised by helminths can cross-react with several epitopes of mite antigenic extracts often used in diagnosis of allergic diseases possibly affecting the correct diagnosis. It becomes increasingly relevant, therefore, to obtain more specific allergens to make an accurate diagnosis of allergy and, consequently, improving immunotherapy for mite allergy. However, the price of assays using bacteria-produced recombinant antigens, as compared to those of assays using crude extracts obtained from mite cultures, is a factor that should be taken into consideration.

**Conclusions**

We demonstrated that rBlo t 5 and rBlo t 21 were antigenic for the *B. tropicalis*-sensitized population evaluated in this work. In addition, we showed that the IgE reactive to these allergens had less cross-reactivity with *A. lumbricoides* extract than anti-*B. tropicalis* extract IgE, as assessed by an IgE-binding inhibition assay. Thus, rBlo t 5 and rBlo t 21 expressed in *E. coli* may be potential candidates to be used in a pool of different recombinant allergens for improving serodiagnosis assays of allergy to *B. tropicalis*. More studies are needed to obtain other recombinant allergens in order to develop a highly sensitive and specific assay for the diagnosis of allergy induced by *B. tropicalis* allergens.

**Abbreviations**

rBlo t: Blomia tropicalis recombinant antigen; cDNA: Complementary DNA; pRSETA: Plasmidial vector pUC-derived expression vectors that designed for high-level protein expression and purification from cloned genes in *Escherichia coli;* BtE: *B. tropicalis* crude extract; ELISA: Enzyme-linked immunosorbent assay; SPT: Skin prick test; CEAR: Centro de Estudos em Alergias Respiratórias; SCAALA: Social Change in Asthma and Allergy in Latin America; mRNA: Messenger RNA; PCR: Polymerase chain reaction; BLAST: Basic Local Alignment Search Tool; IPTG: Isopropyl β-D-1-thiogalactopyranoside; SDS-PAGE: Sodium dodecyl sulfate polyacrylamide gel electrophoresis; PVDF: Immobilon-P polyvinylidene difluoride; PBS: Phosphate-buffered saline; PBS/T: Phosphate-buffered saline containing 0.05% Tween; NFM: Non-fat milk; OD: Optical density; CONEP: National Commission on Ethics in Research.

**Competing interests**

All authors report no conflict of interest financial or otherwise, with the findings of this study.

**Authors’ contributions**

KAC carried out the laboratory assays and wrote the first draft of the manuscript; OPMN: helped in the study design, supervised the molecular biology work and helped in the manuscript revision; FBM: helped with the molecular biology assays; JCMP: helped in the inhibitory assays and revised the manuscript; MCAS: has performed the statistical analysis; LCP: helped in the study design and reviewed the manuscript; NMAN: conceived the work, the study design and supervised the immunological assays; CSP: helped in obtaining the Blo t 5 and Blo t 21. All authors read and approved the final manuscript.

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