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Allergenic potency of kiwi fruit during fruit development

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Abstract
Food allergies, including kiwi fruit allergy, have been the subject of extensive research in the last few years. The aim of this study was to examine a possible relationship between the developmental stage of kiwi fruit and its allergenic potency. The protein and allergen patterns of kiwi fruit extracts in September, October, November and December fruit in the period from 2000–2002 were analysed. One of the factors that may contribute to the difficulties in proposing well-defined and standardized fruit extracts should also be the time of fruit harvesting. In this particular case, when the kiwi fruit was edible throughout November and December, we showed discrepancies in allergen content and potencies both in qualitative and quantitative terms. Two major allergens of kiwi fruit, Act c 1 and Act c 2, mainly accounted for the highest allergenic potential of November kiwi extract in vivo and in vitro. Not only the content of major allergens, but also the ratio of different proteins and even isoforms of the same allergen (Act c 2) change with fruit ripening. These findings should be taken into account during preparation of extracts for allergy diagnosis.

Keywords: Kiwi fruit, Actinidia deliciosa, food allergy, allergen, actinidin, thaumatin-like protein, development, ripening.

Abbreviations: Oral-allergy syndrome (OAS), skin prick testing (SPT), sodium dodecylsulfate-polyacrylamide gel electrophoresis (SDS-PAGE), bovine serum albumine (BSA), Tris-buffered saline (TBS), phosphate-buffered saline (PBS), thaumatin-like protein (TLP), polyvinylpoly-pyrrolidone (PVPP), Coomassie Brilliant Blue (CBB), simulated gastric fluid (SGF), Rocket immunoelectrophoresis (RIE).

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Introduction

Food allergy is an important health problem nowadays. Clinical reactions to food, including cutaneous, gastrointestinal or respiratory disorders or systemic anaphylactic reactions (Sampson 1999) are demonstrated by 8% of children and 2% of adults (Helm & Burks 2000). It is likely that both the incidence and prevalence of food allergies are increasing in line with other forms of allergic diseases (Kimber & Dearman 2001). Food allergy can develop as an isolated reaction to food (fruit, fish, milk) or a secondary reaction after sensitization to pollen (oral-allergy syndrome, OAS) or latex (latex-fruit syndrome) when food allergens cross-react with allergens present in pollen or latex. It is generally believed that food allergens are proteins resistant to digestion because only intact proteins (or their larger fragments) are required for processing by antigen-presenting cells located in gut mucosa. Although stability under digestion may not be the defining characteristic of food allergens, it is widely accepted that resistance to digestion would increase the probability of stimulating allergic reactions presumably by retaining protein integrity (Fu 2002). Thus, digestibility tests have been widely accepted as an appropriate method for evaluating the potential allergenicity of newly-introduced proteins in genetically modified plants (Metcalfe et al. 1996, Fu et al. 2002).

Kiwi fruit is very popular throughout the world because of its taste and high vitamin C content. The kiwi fruit harvest season is in late autumn in the Mediterranean area. September and October fruits are not edible because they are hard, sour and lacking aroma. The fruit is fully ripened in November and December.

Accompanying the growing popularity of kiwi fruit in the regular diet, there has been an increasing number of reports of adverse and allergic reactions to kiwi (Garcia et al. 1989, Pastorello et al. 1996, Pastorello et al. 1998, Rudescho et al. 1998, Gavrovic-Jankulovic et al. 2002b, Bublin et al. 2004), which is often pollen associated. According to earlier studies, at least 12 IgE binding proteins (allergens) could be detected in kiwi extract with molecular weights ranging from 64–12 kDa (Pastorello et al. 1996). Two major allergens of kiwi fruit have been isolated and characterized. Act c 1, a protein of molecular mass about 30 kDa, commonly known as actinidin (Pastorello et al. 1998) is a cysteine protease related to papain (Varughese et al. 1992). Act c 2 is a thaumatin-like protein, the plant defense protein with proven antifungal activity (Gavrovic-Jankulovic et al. 2002b, Wang et al. 2002) that, in kiwi extract, exists in two isoforms differing slightly in pI value (9.4 and 9.5) with molecular mass of about 24 kDa (Gavrovic-Jankulovic et al. 2002b).

Allergenicity of fruit during development depends on the expression of IgE-binding proteins. Vieths et al. (1994) demonstrated that the severity of symptoms in patients allergic to apples was highly correlated with the appearance of 18 kDa apple allergen during storage and suggested that this was caused by ripening. Additionally, Paschke et al. (2001) have correlated the number and intensity of bands in immunoblots with ripening stages of mango fruit and observed no difference caused by ripening.

Allergenic extracts, made by different manufacturers, for the use in diagnosis and therapy of allergic diseases usually show variations in allergenic potency. Especially allergen extracts from fruits, vegetables and other plant foods often lack sufficient biological activity due to the presence of proteolytic enzymes, carbohydrates, and phenol components (Vieths et al. 2001). That is the consequence of applying different extraction methods and source materials. In the last few years several in vivo (such as skin prick testing, food challenge, etc.) and in vitro (measurement of the content of the
major allergen) methods for standardization of allergenic extracts have been proposed (Yunginger 1991, Esch 1997, Duffort et al. 2002).

The purpose of this study was to examine a possible relationship between the developmental stage (from September to December) of kiwi fruit and its allergenic potency. We monitored the protein and allergen content during development, digestibility of allergen samples in simulated gastric fluid and allergenic potential in vivo by skin prick testing (SPT). In the purpose of standardization, we correlated three different methods for the quantification of the major allergen Act c 1.

Materials and methods

Fruit collection and protein extract preparation

Kiwi fruits (Actinidia deliciosa, Monti cultivar) were collected monthly from September to December in the period from 2000 to 2002, from the same tree in Bar, Montenegro. The fruits were stored at −20°C without specific treatment until use. The extracts were made according to previously published protocol (Gavrovic-Jankulovic et al. 2002b). Briefly, the fruits were homogenized 1: 2 (w/v) in 100 mM sodium bicarbonate buffer, pH 9.3 containing 2% polyvinylpoly-pyrrolidone (PVPP) and 0.02% NaN₃ in a blender for 1 min. After extraction and centrifugation, ammonium sulfate was dissolved in the supernatants to achieve 90% saturation. After overnight standing at 4°C the solutions were centrifuged for 20 min at 10000 × g. The obtained pellets were dissolved in a minimal volume of a starting buffer and dialyzed extensively against the same buffer. Protein concentration in the extract was determined by the Bradford assay (Bradford 1976).

Sodium dodecylsulfate-polyacrylamide gel electrophoresis (SDS-PAGE). SDS-PAGE was performed on 4% stacking gel and 10% or 12% resolving gel according to the method of Laemmli (Laemmli 1970) under reducing and non-reducing conditions. Twelve micrograms of proteins per lane were applied on the gel for Coomassie Brilliant Blue (CBB) staining or semidry transfer (0.8mAcm⁻²) to a nitrocellulose membrane (Serva) for further examination. The content of major allergen Act c 1 in kiwi fruit extracts was determined by densitometric analysis after SDS-PAGE under reducing conditions.

Two-dimensional polyacrylamide gel electrophoresis. Two-dimensional PAGE isoelectric focusing was performed in a model 2117 Multiphor cell (LKB Pharmacia), according to the manufacturer’s instructions. The proteins (35 µg per lane) were applied to isoelectric focusing gel and further separated by SDS-PAGE under the conditions previously described (Gavrovic-Jankulovic et al. 2000).

Immunoblotting. The separated proteins on the gel were transferred onto a nitrocellulose membrane, as described by Towbin et al. (1979). The quality of the transfer was checked by staining the nitrocellulose with 0.1% Ponceau S in 5% acetic acid. The nitrocellulose membranes were blocked with 1% bovine serum albumine (BSA) in Tris-buffered saline (TBS) containing 0.05% Tween 20, pH 7.8 for 1 hour at room temperature. After blocking, the membranes were incubated with the allergic patients’ serum diluted 1: 5 in 0.1% BSA in TBS containing 0.05% Tween 20, pH 7.8 for 5 hours. The blots were washed extensively with TBS containing 0.05%
Tween 20, pH 7.8 and incubated for 2h with 1: 1000 diluted monoclonal anti-human IgE antibody labeled with alkaline phosphatase (Abcam Ltd., Cambridge Science Park) and, after washing, were visualized using nitro blue tetrazolium and 5-bromo-4-chloro-3-indolyl phosphate (Sigma Chemical Co., St Louis, MO, USA) as substrates as described (Gavrovic-Jankulovic et al. 2002a). A serum pool of non-allergic persons was used as the negative control.

Digestion with simulated gastric fluid. The digestibility of kiwi proteins in simulated gastric fluid (SGF) was examined according to the method of Yagami et al. (2000). Briefly, 140 μg of kiwi extract proteins was dissolved in 40 μL of prewarmed SGF (US Pharmacopoeia) containing 0.32% w/v of pepsin A (Sigma Chemical Co.). Digestion proceeded at 37°C with continuous shaking and an aliquot (8 μL) of the digest was periodically withdrawn (at 8 and 30 minutes). The digestion was stopped with 2.4 μL 0.2 M Na₂CO₃, and samples were mixed with a sample buffer for SDS-PAGE analysis.

Determination of proteolytic activity of actinidin. Proteolytic activity of actinidin in different kiwi fruit samples towards 0.5 mM Nα-Benzoyl-DL-Arginine 4-nitroanilide (Sigma) in 0.1 M potassium phosphate buffer, pH 6.5, containing 1mM EDTA, 13 mM L-cysteine and 5 mM dithiothreitol at RT was evaluated by spectral analysis at 410 nm. Specific proteolytic activity was calculated using excintion coefficient of 4-nitroaniline and protein concentrations data. Relative proteolytic activity was calculated as percent of November extract proteolytic activity (set up as 100%). Effects of the fruit extract pigments were minimized by preparing probes for each sample without addition of the substrate. Absorbance was read after 1 h of incubation.

A zymogram was run according to Grobe et al. (2002) with some modifications. Briefly, 12 μg of November kiwi extract proteins previously resolved by isoelectricfocusing were applied to 12% resolving gel copolymerized with 0.1% gelatin for 2D-PAGE analysis. After electrophoresis, the gel was incubated in a buffer containing 0.1 M glycine, 10 mM Ca²⁺, 5 mM dithiothreitol and 10 mM cysteine, pH 3.6, for 16 h, followed by staining with CBB.

Antisera production. Antibodies against the December kiwi extract were raised in rabbits according to Harboe and Ingild (1983). The animals were injected with 0.5 mL of a 1:1 emulsion of ripe kiwi extract (1 mgmL⁻¹) in complete Freund’s adjuvant. Bleeding was performed 50 days after the first immunization and every two weeks thereafter. The presence of antibodies to kiwi proteins was detected by immunodiffusion. The serum was partially purified by ammonium sulfate fractionation (50% saturation).

Rocket immunoelectrophoresis (RIE). The kiwi extract proteins (5.5 μg) were applied to 1% agarose gel containing 13% rabbit antibodies on a glass plate (12 × 7 cm). Electrophoresis was carried out in a buffer containing 5 mM barbital, 25 mM Na-barbital, pH 8.6, for 12 h at 2 V cm⁻¹. In order to compare the amount of the major allergen Act c 1 in different kiwi fruit samples, we applied 2.75 μg Act c 1 isolated according to Carne and Moore (1978) without derivatization with S-sulphenyl thiosulphate on the same agarose gel.
Patient sera. Five sera from patients allergic to kiwi fruit were used in this study (1, 2, 3, 4 and 5). All the patients showed OAS and one of them had severe contact dermatitis and anaphylaxis when touching the fruit (case No. 4 in Table I).

Skin tests. The extracts for skin prick testing (SPT) were prepared in phosphate-buffered saline (PBS) (10 mM phosphate buffer, 0.9% NaCl, pH 7.2) 1:1 (w/v), centrifuged for 5 min at 10000 ×g, filtered and neutralized with NaHCO₃ yielding a final protein concentration of 0.07 mgmL⁻¹. SPT as performed on the volar surface of the forearm with the kiwi extracts prepared as previously described (Malling 1993). Histamine (0.010 mgL⁻¹) was used as the positive control and saline (0.9%) as the negative control. The results of SPT were read after 20 min. Wheal diameters were expressed relative to the histamine response.

Results

Kiwi fruit extracts

Kiwi fruit extracts (gathered in 2000, 2001 and 2002) were compared using SDS-PAGE, immunoblot and rocket immunoelectrophoresis. The same protein/antigen/allergen patterns were observed in all three years investigated. The concentration of proteins in kiwi fruit extracts increased with fruit development. The results presented here were all obtained using samples from the year 2001.

Protein patterns and actinidin activity in the zymogram. At least 10 protein bands with molecular weights between 67 and 10 kDa were detected by CBB staining in extracts of kiwi fruit from September to December. SDS-PAGE patterns under reducing (see Figure 1a) and non-reducing (Figure 1b) conditions were quite different. The most noticeable difference was in the mobility of the 24 and 30 kDa proteins in reducing conditions (most likely a thaumatin like protein and actinidin, respectively) compared to non-reducing conditions, when they exhibited 20 and 27 kDa molecular weights, respectively. The protein of 29 kDa was constitutively expressed. We also noticed three protein bands of 10, 12 and 14 kDa with similar appearance and in similar amounts from September to December. Several more protein bands of about 42, 56 and 62 kDa were also present in the extracts of fruits from September to December. Protein bands of 17, 22 and 39 kDa appeared in September and reached the highest intensity.

Table I. Clinical data and skin test results of five patients allergic to kiwi.

<table>
<thead>
<tr>
<th>Patient no.</th>
<th>Diagnosis</th>
<th>Kiwi: September SPT</th>
<th>Kiwi: October SPT</th>
<th>Kiwi: November SPT</th>
<th>Kiwi: December SPT</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>RC/ OAS</td>
<td>++</td>
<td>+</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>2</td>
<td>BA/ OAS</td>
<td>+++</td>
<td>+++</td>
<td>+++</td>
<td>+++</td>
</tr>
<tr>
<td>3</td>
<td>RC/OAS</td>
<td>+++</td>
<td>+++</td>
<td>+++</td>
<td>+++</td>
</tr>
<tr>
<td>4</td>
<td>CD/A</td>
<td>++</td>
<td>++</td>
<td>+++</td>
<td>+++</td>
</tr>
<tr>
<td>5</td>
<td>RC/AE</td>
<td>+++</td>
<td>+++</td>
<td>+++</td>
<td>+++</td>
</tr>
</tbody>
</table>

OAS, oral-allergy syndrome; RC, rhinoconjunctivitis; BA, bronchial asthma; AE, angioedema; A, systemic anaphylaxis; CD, contact dermatitis. The protein concentration in the extracts for the experiment of SPT has adjusted to 0.07 mgmL⁻¹.
in December. Other protein bands of 26 and 67 kDa appeared in September and became most abundant in November.

Two-dimensional PAGE (Figure 2) showed fragmentation of the 30 kDa protein (actinidin), pI 3.6 into a 26 kDa protein with a pI value of 3.2. Both the protein and
protein fragment showed proteolytic activity in zymogram (Figure 3). We observed two more proteins of 39 and 17 kDa with pI values of 5.0 and 4.8, respectively.

The major kiwi fruit allergen Act c 2 was present in all extracts as a mixture of isoforms with isoelectric points of 9.4 and 9.5 and a molecular weight of 24 kDa (Gavrovic-Jankulovic et al. 2002b). Its content increased with fruit development. The amount of the 26 kDa protein, pI 8.8, was the highest in the November extract. Also, a protein of 17 kDa with pI 9.0 reached the highest band intensity in the December kiwi fruit extract.

**Human IgE binding.** In immunoblots developed with individual sera (1, 2, 3, 4) or pool of the patients sera (1, 2, 3, and 4) (see Figure 4a) a significant number of IgE binding proteins were detected. The most intensive bands found in all four extracts, were 30 and 24 kDa proteins (corresponding to actinidin and thaumatin-like protein). An allergen of 29 kDa was also present in all extracts. Other allergens clearly showed a dependence on the developmental stage. IgE-binding proteins of 17, 26, 39, 42, 56 and 62 kDa were more abundant in later months. Allergens of 42 and 68 kDa were present in the October and November samples. The allergen of 22 kDa was most pronounced in the November sample.

Another immunoblot developed after incubation with serum from patient No. 5 (Figure 4b), showed no difference between the extracts of kiwi fruits in different stages of development. Each strip had only one band corresponding to a 24 kDa protein. However, according to the SPT results (case No. 5 in Table I), a quantitative difference between the extracts tested for the same patient did exist. An explanation for the phenomenon that *in vivo* testing appeared much more sensitive than *in vitro*
techniques, could be that not all of the mainly conformational IgE-binding epitopes have been renaturated after transfer to the NC membrane for immunoblot analysis, as demonstrated for some other allergens of different sources (Nilsen et al. 1991).

Quantification of actinidin. The quantity of actinidin in our samples (see Table II) was measured using three different methods: rocket immunoelectrophoresis, densitometry and determination of proteolytic activity. Rocket immunoelectrophoresis was run according to Hudson and Hay (1989). According to all three methods used in this study, the quantity of Act c 1 was the most pronounced in November kiwi fruit extract (Table II). The most noticeable difference in Act c 1 content between the November kiwi fruit extract and the other three kiwi fruit extract was obtained by determination of proteolytic activity.

Digestibility of kiwi proteins. All kiwi proteins in September, October, November and December kiwi fruit extracts were decomposed completely within 30 minutes of exposure to simulated gastric fluid (see Figure 5), but the 24 kDa protein (presumably

Table II. Quantity of major allergen Act c 1 determined by three different methods.

<table>
<thead>
<tr>
<th>Method used</th>
<th>S#</th>
<th>S*</th>
<th>O#</th>
<th>O*</th>
<th>N#</th>
<th>N*</th>
<th>D#</th>
<th>D*</th>
</tr>
</thead>
<tbody>
<tr>
<td>RIE</td>
<td>43.6</td>
<td>81.3</td>
<td>43.6</td>
<td>81.3</td>
<td>53.6</td>
<td>100</td>
<td>43.6</td>
<td>81.3</td>
</tr>
<tr>
<td>Densitometry</td>
<td>36.9</td>
<td>93.9</td>
<td>37.6</td>
<td>95.7</td>
<td>39.3</td>
<td>100</td>
<td>35.7</td>
<td>90.8</td>
</tr>
<tr>
<td>Relative proteolytic activity of Act c 1^Δ</td>
<td>51.1</td>
<td>65.7</td>
<td>100</td>
<td>70.6</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

S, September; O, October; N, November; D, December; RIE, Rocket immunoelectrophoresis; \#% of total protein quantity; *% of November actinidin quantity; ^Δ% of November actinidin activity.
a thaumatin-like protein (TLP), Act c 2) appeared to be more resistant to digestion by gut enzymes in unripe fruit extracts.

Skin prick testing. The results of SPT and patients’ clinical characteristics are shown in Table I. All patients exhibited the most pronounced reaction to the November kiwi fruit extract. The intensity of the reaction to November kiwi fruit extract was approximately twice as strong as the response to the other kiwi fruit extracts (September, October and December).

Discussion

The aim of this study was to correlate the allergen profile and content in kiwi fruit during ripening and to answer the question whether the allergenicity and the digestibility of the fruit depend on its developmental stage or not.

We analysed kiwi fruit extracts from three consecutive years (2000, 2001 and 2002) by SDS-PAGE, 2D-PAGE, immunoblot and rocket immunoelectrophoresis. These extracts exhibited the same allergen and protein profile, ruling out the possibility that our results may depend on specific conditions of the year selected.

The allergenic potential in vivo of the different developmental stages of kiwi fruit (September, October, November and December extracts) investigated by skin prick tests was the highest in extract of fruit collected in November. The amount of Act c 1 (actinidin) was the highest in the same sample. The content of this major allergen of kiwi fruit was measured by rocket immunoelectrophoresis, densitometry and proteolytic activity. Proteolytic activity increased in September to November kiwi fruit extracts and decreased in the December extract, while the actinidin contents determined by densitometry and RIE for September, October and December samples were comparable. These results led us to conclude that actinidin might not be fully active in unripe kiwi fruit and that the amount and activity decreased with overripening. According to our results, determination of proteolytic activity cannot be an appropriate method for quantification of actinidin in different kiwi fruit samples.
Moreover, the difference in the content of Act c 1 (measured by RIE and densitometry) was not sufficiently pronounced to fully explain the extent of patients’ reactions. It seems likely that other allergens, whose content also changes during development, contribute to the intensity of skin reactions. In the December extract the intensity of a more basic isoform of Act c 2 on 2D PAGE was higher when compared to the September, October and November extracts. Interestingly, it seems that this isoform, with a presumably higher allergenic potential, appeared to be more prone to digestion in simulated gastric conditions (see Figure 5). Additionally, we noticed that the expression of some until now uncharacterized allergens, started in September and intensified to reach the highest value in November and December (17, 22, 26, 39 and 67 kDa molecular weights). These allergens may also contribute to higher allergenicity of the November extract. Also, it would be of interest to examine the role that these proteins may play in the process of kiwi fruit development.

A previous study by Paschke et al. (2001) showed no difference between allergenic potency of mango fruit during ripening but cannot be directly compared with our results due to the different time-scale of food collection (different fruit developmental stage samples). Kiwi fruits used in our study were harvested in September and October as unripe, while November and December kiwi fruits were ripe and edible. The fruit extracts were made immediately after harvesting. Paschke et al. (2001) performed their experiments only on extracts of consumable and fully developed mango fruit made five to 40 days after harvesting.

According to our results, one of the factors that contribute to the difficulties in proposing well-defined and standardized fruit extracts should also be time (developmental stage) of fruit harvesting. In this particular case, when the kiwi fruit was edible throughout November and December, we clearly showed strong discrepancies in allergen content and potencies both in qualitative and quantitative terms. It seems that in the case of kiwi fruit, the fully ripened fruit contains less allergens and was able to induce less pronounced allergic reaction in the SPT in the group of tested patients. However, when considering the allergen extract preparation, it should be kept in mind that not only the content of major allergens, but also the ratio of different proteins and even isoforms of the same allergen change with fruit ripening and development. For an optimized extract preparation, a well defined harvesting time should be taken into account. For the improvement of in vitro allergy diagnosis preparations based on optimized allergen extracts as well as recombinant allergens of natural fruit allergens (Vieths et al. 2001) may be a promising solution for this very complex problem.

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References


