

## Article

# Structural Basis for the IgE-Binding Cross-Reacting Epitopic Peptides of Cup s 3, a PR-5 Thaumatin-Like Protein Allergen from Common Cypress (*Cupressus sempervirens*) Pollen

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**Abstract:** The present work was aimed at identifying the IgE-binding epitopic regions on the surface of the Cup s 3 allergen from the common cypress *Cupressus sempervirens*, that are possibly involved in the IgE-binding cross-reactivity reported between Cupressaceae species. Three main IgE-binding epitopic regions were mapped on the molecular surface of Cup s 3, the PR-5 thaumatin-like allergen of common cypress *Cupressus sempervirens*. They correspond to exposed areas containing either electropositive (R, K) or electronegative (D, E) residues. A coalescence occurs between epitopes #1 and #2, that creates an extended IgE-binding regions on the surface of the allergen. Epitope #3 contains a putative N-glycosylation site which is actually glycosylated and could therefore comprise a glycotope. However, most of the allergenic potency of Cup s 3 depends on non-glycosylated epitopic peptides. The corresponding regions of thaumatin-like allergens from other closely related Cupressaceae (*Cryptomeria*, *Juniperus*, *Thuja*) exhibit a very similar conformation that should account for the IgE-binding cross-reactivity observed among the Cupressaceae allergens.

**Keywords:** pollen allergen; thaumatin-like protein; Cup s 3; Cupressaceae; allergenicity; IgE-binding epitope; IgE-binding cross-reactivity



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## 1. Introduction

Allergy to common cypress (*Cupressus sempervirens*) pollen is widely distributed over Mediterranean countries, especially in Italy [1], Israel [2], Greece [3], Spain [4], and the south of France [5]. Pollens from other closely related Cupressaceae species are responsible for seasonal rhinitis in Japan (Japanese cypress *Chamaecyparis obtusa*, Japanese cedar *Cryptomeria japonica*) [6], Australia (*Callitris glaucophylla*) [7], USA (Mountain cedar *Juniperus ashei*, Arizona cypress *Hesperocyparis arizonica*) [8], France (Northern whitecedar *Thuja occidentalis*) [9], and Mediterranean countries (prickly juniper *Juniperus oxycedrus*) [10]. The exposure to Cupressaceae pollen allergens and consequently allergy levels have increased these last 50 years and is still increasing for several reasons. Firstly, besides the classical geographical distribution, Cupressaceae trees are spreading over many other regions due to their increasing use as ornamental trees, especially in urban and suburban areas [11]. Secondly, because of global warming, the pollination period of cypresses and other Cupressaceae species lasts longer, which promotes the currently observed spreading

of their associated allergies, especially in northern Europe [12,13]. Thirdly, the expression of a group 3 allergen of Cupressaceae that belongs to the thaumatin-like protein family was reported to increase under an increasingly polluted environment [14].

Different allergens have been characterized in the pollen of *Chamaecyparis obtusa* (Cha o 1, Cha o 2, Cha o 3), *Cryptomeria japonica* (Cry j 1, Cry j 2, Cry j 7), *Cupressus arizonica* (Cup a 1), *C. sempervirens* (Cup s 1, Cup s 2, Cup s 3, Cup s 7), *Juniperus ashei* (Jun a 1, Jun a 2, Jun a 3, Jun a 7), *J. oxycedrus* (Jun o 1, Jun o 4), *J. sabinoide*s (Jun s 1), and *J. virginiana* (Jun v 1, Jun v 3) (WHO/IUIS Allergen Nomenclature Sub-Committee, [www.allergen.org](http://www.allergen.org), assessed on 15 October 2022). They essentially correspond to ubiquitous proteins such as pectate lyase (group 1 allergens), polygalacturonase (group 2 allergens), PR-5 thaumatin-like protein (group 3 allergens), and Ca<sup>2+</sup> binding protein (calmodulin, group 4 allergen). Besides the WHO/IUIS certified allergens, about 20 additional allergens have been described in Cupressaceae pollen [13]. Recently, three additional allergens belonging to the family of Gibberellin-Regulated Proteins (GRP), Cup s 7 from *Cupressus sempervirens* [15], Cry j 7 from *Cryptomeria japonica* [16], and Jun a 7 from *Juniperus ashei* [17], have completed the list of pollen allergens from the Cupressaceae. Interestingly, these pollen GRP allergens display some IgE-binding cross-reactivity with the corresponding GRPs from fruits and vegetable [18,19]. According to the ubiquitous character of the major Cupressaceae allergens, IgE-binding cross-reactivity has been reported to occur between closely related Cupressaceae pollens [20]. PR-5 thaumatin-like proteins of Cupressaceae pollen offer an example of conserved allergens able to trigger IgE-binding cross-reactions and thus constitute an appropriate model to decipher the structural basis of this IgE-binding cross-reactivity [14,21–25].

Here, we report on the characterization of IgE-binding epitopic peptides of the PR-5 thaumatin-like allergen Cup s 3 [26] and their structural features responsible for the reported IgE-binding cross-reactivity among different Cupressaceae species. In addition, the identification of IgE-binding epitopic regions of pollen allergens is an exciting goal since synthetic peptides mimicking the IgE-binding epitopic stretches identified on the molecular surface should be used instead of recombinant allergens for the purpose of diagnosis, e.g., in immobilized epitope micro-arrays.

## 2. Materials and Methods

### 2.1. Cypress Pollen Extract

A crude protein extract was prepared from common cypress (*Cupressus sempervirens*) pollen (Allergon, Angelholm, Denmark) by suspending 1 vol. of pollen in 10 vol. of 50 mM NH<sub>4</sub>HCO<sub>3</sub> (pH 6.5). After stirring for 3 h at 4 °C, the slurry was centrifuged at 20,000× *g* for 30 min and the supernatant was collected and kept frozen at −20 °C until used. The protein content of the protein extract was evaluated using the bicinchoninic acid kit reagent (Pierce) [27], with the banana TLP Mus a 4 [28] as a standard. The occurrence of both Cup s 1 (pectate lyase) and Cup s 3 in the protein extract was checked by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) in 15% polyacrylamide gels using Tris-glycine as trailing ion [29] and staining with Coomassie blue or silver nitrate.

### 2.2. Sera from Allergic Patients

Blood samples were collected after informed consent of patients selected according to symptoms as rhino-conjunctivitis and/or asthma to Cupressaceae pollen and individual sera were used in SPOT experiments and ELISA. All the used cypress pollen allergic patient (CPAP) sera interacted with Cupressaceae pollen extracts (ImmunoCAP<sup>®</sup> t23, Thermo Fisher Scientific, Phadia 67400 Illkirch, France) (Table 1). Sera from this list are the same as previously reported in [30].

**Table 1.** List of sera from patients allergic to cypress pollen, used in SPOT and ELISA inhibition experiments.

Subjects	Sex/Age	Allergic History	Specific IgE (kU/mL)
1	F/5	* CY,DERF,PAR	0.19
2	M/4	CY,DERP,PAR,DAC,PN,SIN,SHR	0.44
3	F/29	CY,DERP/F,PAR	0.70
4	M/11	CY,DERP/F,CAT,DOG	1.10
5	M/5	CY,OLI,PAR	1.27
6	F/42	CY,DERP/F,CAT	1.39
7	F/43	CY,PAR,DAC	1.98
8	F/16	CY,PAR	4.49
9	F/46	CY,DERP,PAR,DAC	4.92
10	M/33	CY,DERP,PAR,DAC,ALT	6.70

\* CY: cypress, DERF: *Dermatophagoides farinae*, DERP: *D. pteronyssinus*, PAR: *Parietaria*, DAC: *Dactylis*, PN: peanut (*Arachis hypogaea*), SIN: *Sinapis*, SHR: shrimp, OLI: olive tree, ALT: *Alternaria*.

### 2.3. IgE-Binding Epitope Mapping

Overlapping 15-mer peptides, frame shifted by three residues, corresponding to the entire amino acid sequences of Cup s 3 (Accession number Uniprot. Q69CS2), were synthesized using the SPOT technique [31]. For these experiments we used the Multiprep automatic SPOT synthesizer (Intavis Bioanalytical instruments, Cologne, Germany). After different treatments to bind the peptides as described previously [32] the membranes were soaked overnight in tris-buffered saline (TBS) containing 2 mL blocking buffer (Roche-Diagnostics, Meylan, France) and 1 g sucrose (pH 7.0). Then, membranes were incubated (2 h) with CPAP serum (1:10 *v/v*) in the presence of an anti-protease cocktail (Roche) in a moist chamber. Membranes were then soaked in a 1:4000 dilution of mAb anti human-IgE coupled to Alkaline Phosphatase (AP) (Sigma-Aldrich, St Louis, MO, USA) for 1 h. After, the interacting peptide spots were stained for 30 min by adding the AP substrate, 5-Bromo-4-Chloro-3-Indolyl Phosphate (BCIP, Promega corporation, Madison, WI, USA). Then the membranes were washed three times with deionized water and dried for scanning. Negative controls were obtained by using either TBS without serum or a serum from non-allergic patients to Cupressaceae pollen. Between each incubation step, 3 washes with TBS containing 0.1% (*v/v*) tween 20 (TBS-Tw) were performed. Membranes can be used repeatedly after regeneration steps consisting of 1–3 washes in dimethylformamide for 10 min each, 3 washes in deionized water, 3 washes in 8M urea containing 1% (*w/v*) SDS and 1% (*w/v*)  $\beta$ -mercaptoethanol and finally 3 washes in a mixture of ethanol–acetic acid–H<sub>2</sub>O (50:10:40, *v/v/v*).

### 2.4. ELISA Inhibition Experiments

Three 9-mer peptides that broadly correspond to the predicted epitope #1 (peptide #1: 9PGGGKRLDQ27), epitope #2 (peptide #2: 45RFGWRTGIT53), and epitope #3 (peptide #3: 161TNYSKIFKN169) of Cup s 3, were synthesized using the Fmoc (Fluorenylmethyloxycarbonyl) chemistry (JPT Peptide Technologies GmbH, Berlin, Germany). The capacity of synthetic peptides to inhibit the IgE-cypress extract interaction was checked by ELISA (Enzyme-Linked Inhibition Sorbent Assay). Briefly, the wells were coated with cypress pollen extract in phosphate buffer saline (PBS, pH 7.5), containing 1  $\mu\text{g}\cdot\text{mL}^{-1}$  protein, and after overnight incubation (4 °C), the wells were washed 3 times with PBS and then incubated (2 h, room temperature) with PBS containing 0.1% tween 20 (*v/v*) and 1% BSA (Bovine serum albumin) (*w/v*) (PBSTB). Then, 50  $\mu\text{L}$  of 1:30 diluted CPAP serum in PBSTB, previously incubated with 1 mM or 2 mM of synthetic peptide, were added and the plates were incubated for 1 h 30, at room temperature, under constant stirring. Then, mAb anti human-IgE-AP (Sigma-Aldrich, St. Louis, MO, USA) was added (diluted 1:500) for 1 h incubation followed by BCIP (45 min in the dark). Between each incubation step, 3 washes with PBST were performed. The absorbance at 405 nm was recorded on a Titertek

Multiscan spectrophotometer (Labsystems, Thermo Fisher Scientific, Villebon-sur-Yvette, France). Each value is the mean of three separate experiments and appropriate controls were performed under the same conditions.

### 2.5. Bioinformatics

Multiple amino acid sequence alignment of Cupressaceae PR-5 thaumatin-like proteins (TLP) was carried out with CLUSTAL-X [33]. An unrooted phylogenetic tree was built from the multiple amino acid sequence alignment of PR-5 TLP, using the neighbor-joining method.

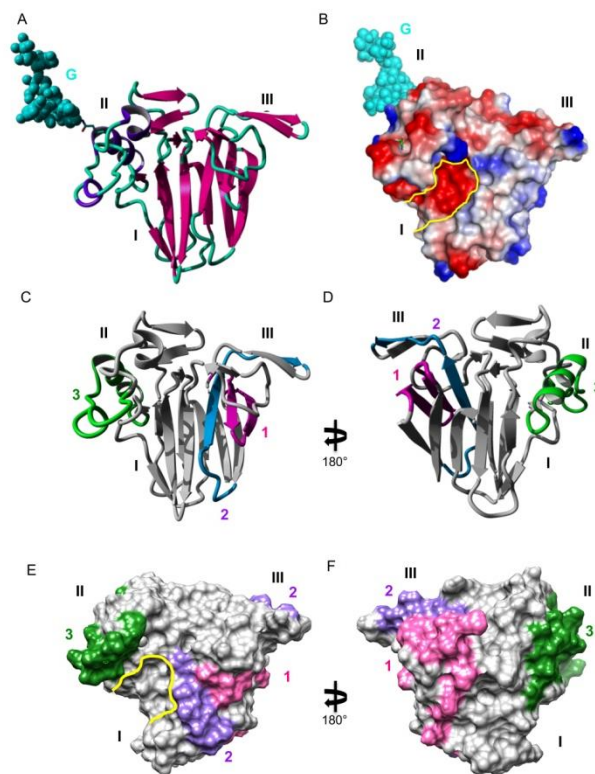
Homology modeling of Cup s 3 from *Cupressus sempervirens* was performed with the YASARA Structure program [34]. Different models of Cup s 3 were built from the X-ray coordinates of the banana (*Musa acuminata*) thaumatin-like protein allergen Mus a 4 (PDB code 1Z3Q) [35], the kiwi (*Actinidia deliciosa*) fruit allergen Act d 2 (PDB code 4BCT), the grape (*Vitis vinifera*) thaumatin-like protein (PDB code 4JRU) [36], the Sodom apple (*Calotropis procera*) laticifer osmotin (PDB code 4L2J), and the antifungal protein zeamatin of maize (*Zea mays*) (PDB code 1DU5) [37], used as templates. Finally, a hybrid model of Cup s 3 was built up from the different previous models. PROCHECK [38], ANOLEA [39], and the calculated qualitative model energy analysis (QMEAN) scores [40,41], were used to assess the geometric and thermodynamic qualities of the three-dimensional model. Only three residues (Asn101, Cys118, Gln173) over 199, occurred in the non-allowed regions in the Ramachandran plot. Using ANOLEA to evaluate the model, only 6 residues (over 199) of the Cup s 3 model exhibited an energy over the threshold value. Both residues are mainly located in the loop regions connecting the  $\beta$ -sheets to the  $\alpha$ -helices in the model. The calculated QMEAN score of the model gave a value of 0.664. Other thaumatin-like proteins from other Cupressaceae species including TLP-Ca from *Hesperocyparis/Cupressus arizonica*, TLP-Cj from *Cryptomeria japonica*, Jun a 3 from *Juniperus ashei*, TLP-Jr from *Juniperus rigida*, TLP-To from *Thuja occidentalis* and TLP-Ss from the closely related Taxodiaceae species *Sequoia sempervirens*, and TLP-Txm from the Taxaceae species *Taxus  $\times$  media*, were similarly modeled using the same PDB code proteins as templates. Reliable values of 0.749 (TLP-Ca), 0.768 (TLP-Cj), 0.642 (Jun a 3), 0.699 (TLP-Jr), 0.717 (TLP-To), 0.731 (TLP-Ss) and 0.704 (TLP-Txm), were obtained for the QMEAN score of the different modeled allergens, respectively.

The surface electrostatic potentials were calculated and rendered with YASARA using the Amber96 forcefield with dielectric constants applied to the protein and the solvent fixed at 4.0 and 80.0, respectively. Electrostatic potentials were displayed on the molecular surface as red (electro-negatively charged) and blue (electro-positively charged) patches. Neutral surfaces are white. Assuming the putative N-glycosylation sites at Asn/N residues of Cup s 3 is glycosylated by a high-mannose bi-antennary glycan chain with a tri-mannoside core (Man)<sub>5</sub>-(GlcNAc)<sub>2</sub>, the glycan chains was built on the allergen using the GlyProt server (<http://www.glycosciences.de/modeling/glyprot/php/main.php>, accessed on 4 September 2022) and further represented in CPK on the molecular surface. Molecular cartoons were drawn with YASARA. The surface occupied by the SPOT identified sequential IgE-binding epitopic stretches of the modeled Cup s 3 and other modeled allergens were differently colored and displayed on the molecular surface of the PR-5 TLP allergen models with Chimera [42]. The root-mean-square deviation of atomic positions (rmsd, in Å) between the superposable C $\alpha$  of pairwise superposed PR-5 TLP allergens was calculated at the SuperPose web server (<http://www.wishart.biology.ualberta.ca>, accessed on 4 September 2022).

### 3. Results

The modeled Cup s 3 displayed the canonical three-domain structure of PR-5 thaumatin-like proteins, made of a  $\beta$ -sandwich central domain (I) flanked on both sides by an  $\alpha$ -helical domain (II) and a short  $\beta$ -hairpin domain (III) (Figure 1A,B). Both domains I and II delineate a strongly electronegative central groove responsible for the endo-1,3- $\beta$ -glucanase activity reported for PR-5 thaumatin-like proteins [43] (Figure 1C,D). The face opposite to the groove displays a more pronounced electropositive character. A very similar three-

dimensional conformation was observed for other modeled PR-5 allergens of Arizona cypress (TLP-Ca), Japanese cedar (TLP-Cj), mountain cedar (Jun a 3), and Northern white cedar pollen (TLP-To) (results not shown). All these allergens share a high degree of both amino acid sequence identity (from 74 to 98% with reference to Cup s 3) and similarity as shown on multiple amino acid sequence alignment (Table 2).



**Figure 1.** (A). Ribbon diagram of the Cup s 3 model showing the overall organization of the allergen in three distinct domains labeled I, II, and III, and the N- glycan chain (G) attached to residue Asn162 in domain II, represented in CPK colored cyan. (B). Molecular surface of Cup s 3 showing the electrostatic potentials patches. The electronegatively charged groove is outlined in yellow. (C,D). Mapping of the sequential IgE-binding epitopes 1 (colored pink), 2 (colored blue), and 3 (colored green) on the front face (C) and the back face (D) of the ribbon diagram of Cup s 3. (E,F). Mapping of the sequential IgE-binding epitopes 1 (colored pink), #2 (colored blue) and 3 (colored green) on the front face (E) and the back face (F) of the molecular surface of Cup s 3. Epitope 2 consists of two separate patches on the molecular surface of the TLP allergen. Note the coalescence of epitope 1 with both patches of epitope 2.

Three main IgE-binding epitopic stretches, that strongly interacted with all the tested patient's sera, were identified using the SPOT technique (Figure 2A,B). They correspond to IgE-binding epitopic regions 1, 2, and 3, respectively. Two additional IgE-binding spots constantly interacted with all of the IgE-containing CPAP sera but the interaction was weaker and, therefore, they were not retained as major IgE-binding epitopes for Cup s 3. A few other spots, especially at the C-terminal end of the sequence, were more scarcely and faintly stained by a restricted number of patient sera and were not considered as relevant epitopic regions.



**Table 2.** Multiple amino-acid sequence alignments of seven thaumatin-like proteins from Cupressaceae pollen. The sequence of Jun v 3 is incomplete in the NCBI databank. Grey boxes correspond to the absence of insertion which are only present in TLP-Cj and TLP-To. Cystein residues are in red and red dotted lines indicate the eight disulfide bridges.

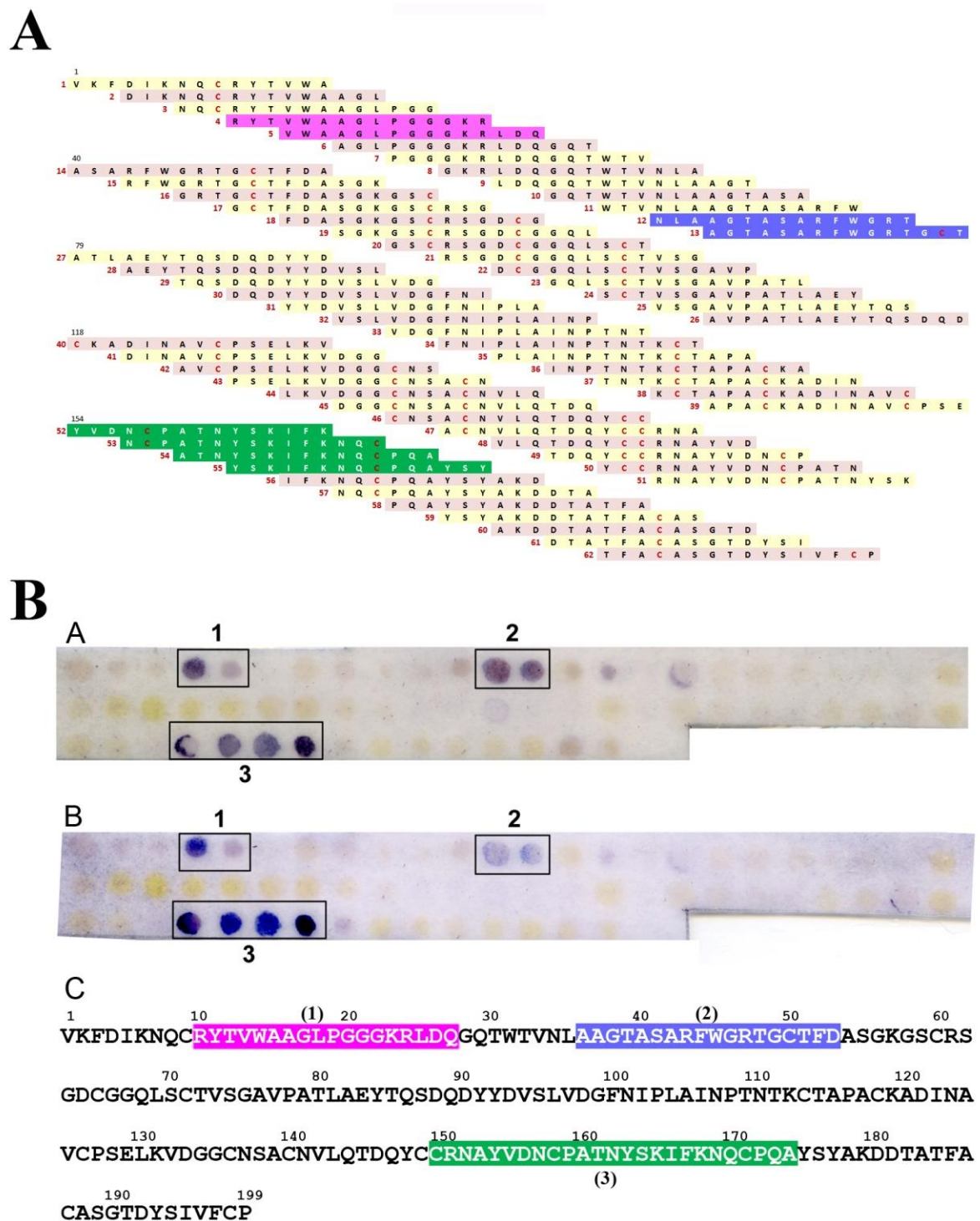
SPECIES	Accession Number	Allergen	1	5	10	15	20	25	30	35	40	45
Common cypress	<i>Cupressus sempervirens</i>	Q69CS2	Cup s 3	VKFDIKNQ	CRYTVWAAGLPGGGKRLD	QGQTWTNLAAGTASARFWG						
Arizona cypress	<i>Hesperocypariz arizonica</i>	Q9FY35	TLP-Ca	-----G	-----EF	-----						
Mountain cedar	<i>Juniperus ashei</i>	P81295	Jun a 3	-----G	-----	-----						
Temple juniper	<i>Juniperus rigida</i>	Q69CS6	TLP-Jr	-----G	-----L	-----						
Eastern red cedar	<i>Juniperus virginiana</i>	Q9LD79	Jun v 3	-----G	-----	-----						
Japanese cedar	<i>Cryptomeria japonica</i>	Q4W6C7	TLP-Cj	-EL	-E	-QQ	-P	-EVP	-KG			
Northern whitecedar	<i>Thuja occidentalis</i>	Q5RZ93	TLP-To	-N	-R	-G	-Q	-S	-DV	-KG		
RTGCTFDASGKGS CRS GD C G G Q L S C T V S G A V P A T L A E Y T Q S D Q D Y Y D V S L V D G F N I P L A I N P T N T K C T A P A C K A D I N A V												
-----QT-----AQ-----												
-----KT-----AQ-----												
-----QT-----R-----												
-----S-----R-----T-----KT-----NS-----Q-----G-----T-----LNG-----GNK-----F-----V-----S-----SQ-----F-----V-----A												
-----S-----G-----R-----QT-----L-----SLNGNDNK-----F-----V-----S-----G-----T-----V-----A												
CPSELKVDGG CNSAC NVLQTDQYCCRNAYVDNCPATNYSKIFKNQCPQAYS YAKDDTA TFACASG TDYSIVFCP												
-----N-----												
-----FK-----												
-----E-----F-----												
-----AQ-----T-----F-----E-----GTH-----SPSS-----M-----SSS-----T-----P-----T-----												
-----A-----NS-----F-----G-----N-----M-----SS-----T-----P-----T-----												
												identity (%)*
												100
												98
												96
												96
												96
												75
												81

\*: Percentage of identity calculated with reference to Cup s 3. -: Identical amino-acids with Cup s 3.

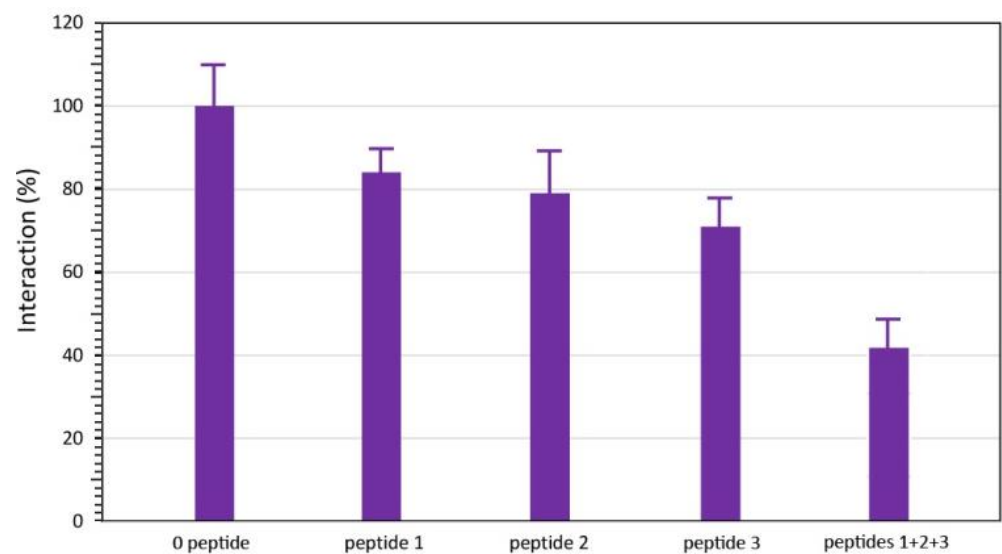
Mapping of the major IgE-binding amino-acid stretches previously identified by the SPOT technique on the ribbon diagram (Figure 1C,D) and the molecular surface of Cup s 3 (Figure 1E,F), allowed us to accurately define those amino acid residues sufficiently exposed on the molecular surface of Cup s 3 to readily interact with the corresponding IgE. Exposed residues in each of the three IgE-binding stretches were thus considered as forming the major IgE-binding epitopic regions of Cup s 3 (Figure 1E,F). They correspond to more restricted areas on the surface of Cup s 3 than the whole IgE-binding peptide because some amino-acids are buried in the molecule and not accessible to antibodies. Epitope 1 (stretch 10RYTVWAAGLPGGGKRLDQ27) and part of epitope 2 (stretch 34NLAAGTASAR43) are distributed on domain I whereas the other part of epitope 2 (stretch 49RTGCTFD53) occurs only in the hairpin-containing domain III.

Epitope 3 (stretch 150CRNAYVDNCPATNYSKIFKNQCPQA174) coincides with the  $\alpha$ -helix-containing domain II. These major IgE-binding epitopes are arrayed on both faces of Cup s 3 but do not interfere with the catalytic groove located in the front face of the molecule. Some of these epitopic regions, epitopes 1 and 2, are coalescent and generate an extended epitopic region on the molecular surface of Cup s 3 that could correspond to discontinuous epitopes. Epitope 3 contains a putative N-glycosylation site 162NYS that is predicted to be glycosylated (Figure 1B,D), and should therefore consists of a glycotope undetectable in the SPOT technique since all the assayed peptides are not glycosylated. This N-glycosylation site is conserved in other thaumatin-like allergens (TLP-Jr, TLP-Ca, Jun a 3, and TLP-To) (except TLC-Cj) from the Cupressaceae species (Table 2).

Synthetic peptides corresponding to the three main epitopic regions of Cup s 3 (epitopes 1, 2, and 3) were tested for ELISA inhibition of the interaction of IgE-containing sera from CPAPs with a total protein from cypress pollen extract. An inhibition in the range between 16–28% of the interaction occurred with all of the tested peptides, and up to 58% inhibition was obtained with an equimolar mixture of the three synthetic peptides (Figure 3). The synthetic peptide corresponding to epitope 3 was the most potent inhibitor. These results suggest that Cup s 3 participates in the allergenicity of Cupressaceae pollen.



**Figure 2.** (A). Amino acid sequences of the 62 overlapping peptides used in the SPOT technique. Epitope 1 (pink background), 2 (purple background), and 3 (green background) are indicated. Cysteine residues are in red. Peptide numbering in red. (BA, BB). Examples of SPOT results obtained with two different sera from cypress allergic patients used as a probe. (BC). Sequential IgE-binding epitopic stretches 1, 2, and 3 deduced from the stained spots are indicated in colored boxed white letters along the amino acid sequence of Cup s 3.

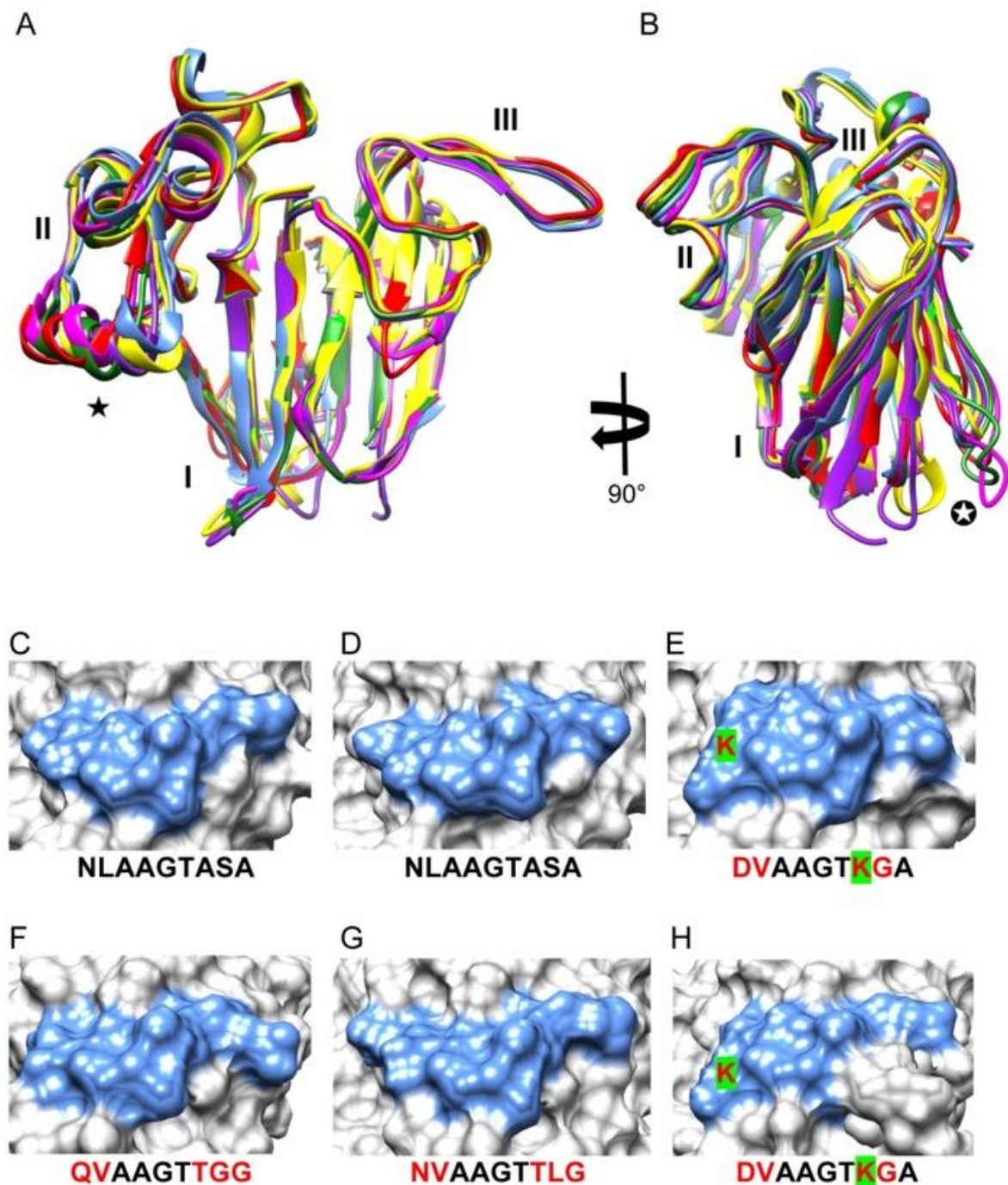


**Figure 3.** ELISA inhibition of the interaction between cypress pollen extract and IgE from CPAPs, in the presence of 2 mM of synthetic peptides corresponding to epitope 1 (peptide 1: 9PGGGKRLDQ27), epitope 2 (peptide 2: 45RFGWGRTGIT53), epitope 3 (peptide 3: 161TNYSKIFKN169), and an equimolar mixture of the three peptides 1, 2, and 3 (peptides 1 + 2 + 3). All values are the mean  $\pm$  s.d. of three separate measurements performed in the same experimental conditions.

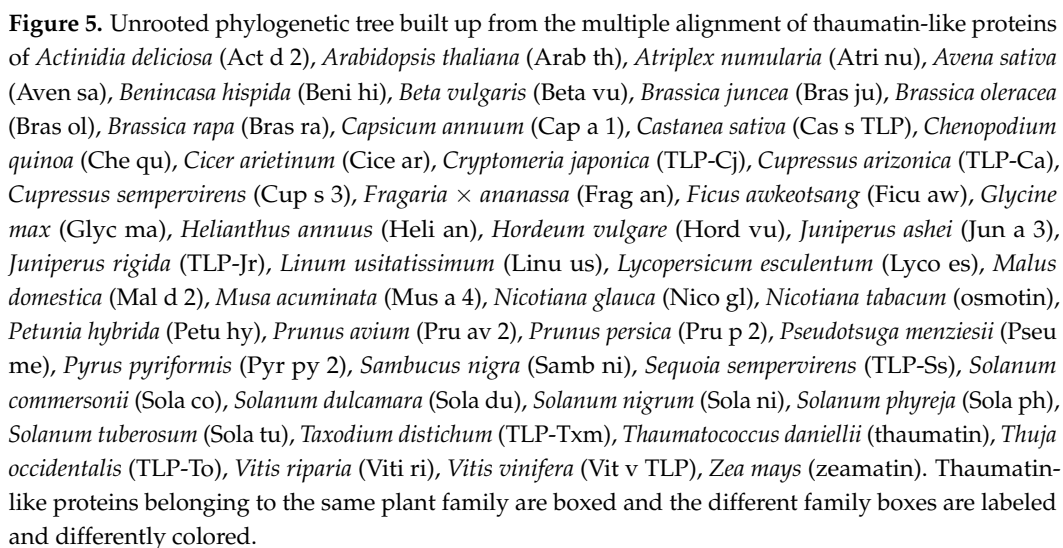
All of the three-dimensional models built up for other TLP pollen allergens consist of exhibit a very similar three-domain organization and are readily superimposed with a mean rms  $< 1.0\text{-}\text{\AA}$  (Figure 4A,B). Accordingly, all the IgE-binding epitopic regions identified on the molecular surface of Cup s 3 consist of conserved residues and exhibit a high degree of conformational similarity (similar shape and extent) with the corresponding regions of TLP allergens from other Cupressaceae pollens. Despite amino acid changes that have occurred along the amino acid sequence stretch corresponding to epitope #2, the overall conformation of this epitopic region looks very similar in other Cupressaceae allergens (Figure 4C–H). However, the introduction of a bulky lysine residue in the sequence stretch TLP-Jr from Temple juniper and TLP-To from Northern whitecedar, induces some distortion into the overall conformation of epitope #2 (Figure 4E,H).

These close conformational similarities most probably account for the currently reported IgE-binding cross-reactivity between Cupressaceae pollen species. In this respect, the unrooted phylogenetic tree built from the multiple alignment of the PR-5 TLP belonging to different species of higher plants, shows that pollen allergens from Cupressaceae, Taxodiaceae and Taxaceae, consist of a homogeneous clade of extremely closely related species, poorly related to other higher plant family clades (Figure 5).





**Figure 4.** Front view (A) and lateral view (B) of the superimposed ribbon diagrams of Cup s 3 (red), TLP-Ca (blue), TLP-Jr (yellow), TLP-Cj (magenta), TLP-Ss (green) and TLP-To (purple). The three domains I, II and III are labeled. Stars ★ and ☆ indicate the poorly superimposed regions in the ribbon diagrams of the TLP allergens, respectively. Overall conformation of epitope 2 at the molecular surface of Cup s 3 (C), TLP-Ca (D), TLP-Jr (E), TLP-Cj (F), TLP-Ss (G) and TLP-To 3 (H). The amino acid sequence stretches corresponding to epitope 2 are indicated and amino acid changes are shown in red letters. The K residue change responsible for the distortion of the overall conformation of epitope 2 observed in TLP-Jr (E) and TLP-To (H), is shown in green highlighted red letter.



As a typical PR-5 thaumatin-like protein, the modeled Cup s 3 allergen exhibits the canonical three-dimensional conformation made of three domains delineating a central electronegative catalytic groove responsible for the  $\beta$ -glucanase activity (Figure 1A,B). Three distinct sequential IgE-binding epitopic peptides distributed in four epitopic regions were identified on the molecular surface of Cup s 3. Epitope #2, named according to the results of SPOT techniques, splits into two exposed epitopic regions because one part is buried in the molecule most likely due to the eight disulfide bridges. In this respect, although the SPOT technique used herein clearly identified three epitopic regions, the existence of undetected additional discontinuous epitopes cannot be ruled out because of the very folded nature of these allergens. Epitopes are essentially distributed on the face opposite to that containing the catalytic groove. Two of these epitopes (1 and the larger part of 2) occur in domain I and epitope 3 in domain II. Epitope 3 possesses a *N*-glycosylation site, that could possibly correspond to a glycotope that does not prevent the induction of

IgE specific to the underlying peptide. Alternatively, the TLP allergen might consist of two isoforms, glycosylated and not glycosylated. As could be expected, the protruding hairpin domain III is an epitope (smaller part of epitope 2). The identified peptides in the SPOT techniques were confirmed to be accessible on the intact allergen since they were able to significantly inhibit the binding of the specific serum IgE to the TLP allergen in its native form in pollen extract.

Conformational analysis of the IgE-binding peptides identified on Cup s 3 revealed striking similarities with the corresponding regions of the modeled regions from other closely related Cupressaceae TLP, from *Cryptomeria japonica* and from *Thuja occidentalis* (Figure 4). These conformational mimics most probably account for the IgE-binding cross-reactivity reported between different Cupressaceae allergens [26,30,44,45]. In agreement with these conformational similarities, some sequence similarity was observed among the epitopic regions of the thaumatin-like allergens from different Cupressaceae. In this respect, four positively IgE-reacting peptides corresponding to sequence stretches 120–131, 132–145, 155–145, and 169–179 have been previously identified on the molecular surface of Jun a 3 from the mountain cedar pollen [46]. Epitope 3 (sequence stretch 156–173) of Cup s 3 overlaps with two of the major IgE-binding epitopic regions (sequence stretches 155–165 and 169–173) characterized on Jun a 3. The Jun a 3 epitope is located in the  $\alpha$  helix-containing domain II [46]. Cross-reactivity of Cup s 3 and Jun a 3 was demonstrated by ELISA inhibition assays in which the binding of IgE to Cup s 3 was inhibited by a preincubation in the presence of purified Jun a 3 [22]. Cross reactivities between TLPs from various plant species were also studied in TLP sensitized patients from restricted areas in Spain [24]. However, sensitization to Cup a 3 was barely represented and poorly significant “co-sensitization links” were observed between Cup a 3 and Pru av 2 or Mald 2, TLPs from cherry and apple, respectively. A similar value of “co-sensitization link” was also reported with Ana c 2, a cysteine protease from pineapple. Whether these “co-sensitization links” might be attributed to cross-reactive oligosaccharide moieties present on TLPs and/or cysteine protease was not studied.

The possible role of *N*-glycans with respect to the allergenicity and IgE-binding cross-reactivity of Cupressaceae allergens was deeply investigated by Italian authors using different allergens as *N*-glycosylated probes [47–50]. Their results indicated that a significant proportion of IgE from CPAP specifically recognizes the *N*-glycan moieties of the different glyco-allergens, pectate lyase (group 1), and polygalacturonase (group 2) [51]. However, the cross-reactivities observed for group 1 and group 2 allergens of Cupressaceae pollen do not rely only on oligosaccharides moieties since AA sequences are closely related, generating similar peptidic epitopic regions. This is particularly exemplified in the Cup s 3 results reported herein where epitopic region 3 of Cup s 3, including a glycosylation site, exhibited the most intense IgE-binding reactivity.

In agreement with the close structural similarities of their epitopic regions, PR-5 TLP allergens from the Cupressaceae, Taxaceae, and Taxodiaceae form a very homogeneous clade distinct from other PR-5 TLP family clades, as shown in the phylogenetic tree built up for the plant PR5 TLP (Figure 5). The lack of close relationship with other families of PR-5 TLP within the phylogenetic tree suggests that peptidic cross-reactivities between Cupressaceae TLPs and TLPs from other plant species are unlikely to occur.

## 5. Conclusions

The identification of putative IgE-binding cross-reactive epitopes of Cup s 3 and other closely related PR-5 TLP allergens from the Cupressaceae, Taxaceae, and Taxodiaceae pollen has been completed. The position in the phylogenetic tree of these pollen allergens, which form a distinct clade separated from other PR-5 TLP family clades, suggests that their IgE-binding cross-reactivity (1) is limited to the Cupressaceae and the closely related Taxaceae and Taxodiaceae families, irrespective of their sequential and structural similarities, and (2) should exclude cross-reactivity to plant-derived foods and food products.



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