Evaluation of In Silico Prediction Possibility of Epitope Sequences Using Experimental Data Concerning Allergenic Food Proteins Summarised in BIOPEP Database

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INTRODUCTION

Allergy is one of the greatest challenges for the contemporary food science and medicine [Skripak & Sampson, 2008; Jędrychowski et al., 2008; Cianferoni & Spergel, 2009; Cummings et al., 2010]. Allergy treatment involves the elimination of allergens from the diet or desensitization therapy [Skripak & Sampson, 2008; Kim & Sicherer, 2010; Prescott et al., 2010]. For elimination diets and desensitization treatment to be effective, allergenic food proteins and protein fragments that are epitopes have to be determined. The relevant research is supported by bioinformatics tools [Tong & Ren, 2009; Mari et al., 2009; Salimi et al., 2010; Tomar & De, 2010]. The bioinformatics tools applied in immunology and allergology include databases containing allergenic protein sequences [Gendel, 2009; Mari et al., 2009; Salimi et al., 2010; Tomar & De, 2010; Darewicz et al., 2011]. Those databases are compatible with applications comparing a given protein sequence with the sequences listed in the database. Most of such applications rely on BLAST [Altschul et al., 1997] and FASTA [Pearson et al., 1991; Pearson, 2000] algorithms. The existing databases support the evaluation of a protein’s potential allergenicity based on a set of bioinformatics criteria recommended by the World Health Organization (WHO), such as the presence of protein sequence fragments containing a minimum of 6–8 amino acid residues which are identical to the fragments of known allergens or fragments containing a minimum of 80 amino acid residues showing at least 35% similarity with the known allergen sequence [Goodman, 2006]. The search for new solutions is still necessary to further advancement of the existing bioinformatics methods and tools [Gowthaman & Agrewala, 2009].

The aim of the study was to evaluate the possibility of predicting potential epitope sequences and location in allergenic proteins from food using EVALLER program by comparison with experimental epitopes summarised in the BIOPEP database of allergenic proteins. Sequences of experimental epitopes from food allergens, present in the BIOPEP database of allergenic proteins were used in the study. Sequences of potential epitopes were found using EVALLER program. The Positive Predictive Value (PPV) has been used as a measure of prediction quality. The potential epitopes fully or partially overlapping with the experimental ones were considered as true positive results whereas these unrelated to the experimental ones as false positive results. The PPV for entire dataset containing 310 potential epitopes was 80.6%. The PPV varied significantly among particular allergen families defined according to the AllFam database. Caseins revealed PPV = 100% (with one exception), proteins from tropomyosin family and proteins from papain-like cystein protease family – exceeding 50%. The last two families possess also relatively low frequency of epitope occurrence. The predictive potential was poor (less than 50%) for plant allergens from cupin superfamily. Families such as lipocalins from milk and EF-hand family (parvalbumins) revealed high variability within family. The EVALLER program may be used as a tool for the prediction of epitope location although its potential varies considerably among allergen families. High PPV is associated with a high number of known experimental epitopes (such as in caseins) and/or a high degree of sequence conservation within family (caseins, tropomyosins).

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insertion and deletion computed using the FASTA algorithm [Pearson, 1991; 2000], and the Smith-Waterman score [Smith & Waterman, 1981]. The algorithm applied to develop the EVALLER program was compared with other algorithms investigating the allergic character of the examined proteins [Soeria-Atmadja et al., 2006]. In reference to the official bioinformatics criteria recommended by the WHO [Goodman, 2006], the discussed program produces fewer false-positive results, i.e. cases in which a non-allergic protein is found to be an allergen [Soeria-Atmadja et al., 2006]. According to a suggestion presented by Björklund et al. [2005], peptides characteristic for allergens could overlap with experimental epitopes determined by mapping, i.e. based on interactions between the peptides corresponding to fragments of protein sequences and the antibodies of persons allergic to the analysed protein [Bohle, 2006; Steckelbroeck et al., 2008]. Comparison of the epitope determination using experimental mapping and prediction using EVALLER is presented in Figure 1.

The aim of the present study was to evaluate the possibility of predicting potential epitope sequences and location in allergic proteins from food using EVALLER program by comparison with experimental epitopes summarised in the BIOPEP database of allergenic proteins.

**METHODS**

Information about bioinformatics tools mentioned in this article, including web addresses and references (if available) is summarised in Table 1. All databases and programs were accessed before 30.04.2011.

Sequences of food allergens and experimental epitopes present in the BIOPEP database of allergenic proteins were used in the study.

The BIOPEP database of allergenic proteins and their epitopes contains the following information: allergen name; sequence, sequences of experimental epitopes; sequences of predicted epitopes; reference describing proteins sequence (bookmark “reference”); reference concerning allergenicity, information about sequences of experimental and theoretical epitopes; ID of particular experimental epitopes in the Immune Epitope Database (bookmark “additional information”); information about AllFam allergen family and epitopes occurring in more than one protein in the BIOPEP database (bookmark “homology”) and information about annotation in general protein database (mainly UniProt), WHO-IUIS (World Health Organization – International Union of Immunological Societies) and Allergome or other allergen databases (bookmark “database reference”). The BIOPEP database of allergenic protein and their epitopes contains also a search engine that enables finding fragments identical with epitopes in protein sequences (e.g. provided by user).

Sequences of potential epitopes were found using EVALLER program with default parameters. Number of sequences displayed in an output covered all sequences revealing 100% identity with fragments of known allergens (as recommended

**TABLE 1. Bioinformatics tools mentioned in this publication.**

| Tool               | Website                                                                 | Reference                  |
|--------------------|-------------------------------------------------------------------------|----------------------------|
| Allergome          | http://www.allergome.org/                                               | Mari et al. [2006, 2009]   |
| AllFam             | http://www.meduniwien.ac.at/allergens/allfam/                          | Radauer et al. [2008]      |
| BIOPEP database of allergenic proteins and their epitopes | http://www.uwm.edu.pl/biochemia                                       | Minkiewicz et al. [2011]  |
| EVALLER            | http://bioinformatics.bmc.uu.se/evaller.html;                         | Martinez Barrio et al. [2007] |
|                    | http://www.slv.se/en-gb/Group1/Food-Safety/e-Testing-of-protein-allergenicity/e-Test-allergenicity/ |                           |
| Immune Epitope Database (IEDB-AR) | http://www.immuneepitope.org/                                       | Vita et al. [2010]         |
| Pfam               | http://pfam.sanger.ac.uk/                                              | Sammut et al. [2008]       |
| UniProt            | http://www.expasy.org                                                  | Jain et al., 2009; The UniProt Consortium [2011] |
| WHO-IUIS allergen database | http://www.allergen.org/                                               |                            |
by Minkiewicz et al. (2011). All such fragments are annotated in the BIOPEP database. Sequences with a lower degree of identity were not taken into account.

The frequency of occurrence of sequential epitopes (A) has been calculated automatically during introduction of protein data into the BIOPEP database according to equation 1 [Dziuba et al., 2003]. Both experimental and potential epitopes are included during calculation.

\[ A = \frac{n}{N} \]

where: \( n \) – number of epitopes, and \( N \) - number of amino acid residues.

Positive predictive value (PPV) has been used as a parameter estimating the quality of prediction of a potential epitope. The PPV may be considered as likelihood that potential epitope will fully or partially overlap with at least one experimental epitope. The PPV value has been calculated using equation 2 [Pulido et al., 2003].

\[ PPV = \frac{tp}{tp + fp} \]

where: \( tp \) – true positive results, and \( fp \) – false positive values.

Definitions of true and false positive values are presented in Figure 1.

The total PPV value has been calculated for all proteins containing both potential and experimental epitopes. The PPV value for an individual allergen has been calculated only if the allergenic protein contains more than one potential epitope.

**RESULTS AND DISCUSSION**

Sixty proteins out of the 135 ones present in the BIOPEP database contain both potential and theoretical epitopes, and 43 of them contain at least two potential epitopes determined by EVALLER program. The data cover 310 allergen-representative FLAPs, of which 187 (60.3%) completely overlapped with the experimental epitopes, 63 (20.3%) partially overlapped with the experimental epitopes, and 60 (19.4%) were unrelated to the experimental epitopes. In line with the adopted sequence coverage criterion, a single FLAP has to be found in a single experimental epitope or a single experimental epitope has to be determined in a single FLAP to be considered as fully overlapping. Such a rule will be maintained in the further discussion. The Positive Predictive Value calculated for the entire dataset (total PPV) is 80.6%.

A group of experimental epitopes forming a continuous protein sequence fragment could overlap with a theoretically-predicted allergenic peptide or a group of theoretically-predicted peptides (FLAPs) forming a continuous protein sequence fragment could overlap with an experimental epitope. An example of the above is an experimental epitope sequence containing residues 131–151 in the precursor of allergen Ara h 3.0101 (BIOPEP ID 47) which overlaps with a group of potential epitopes containing residues 108–135; 124–145; 126–147; 139–163; 144–165 and 149–184, respectively. None of the above FLAPs contains or is a part of an experimental epitope. Based on the above, the discussed FLAPs have been classified as partially overlapping with experimental epitope. All six FLAPs create a continuous fragment of a protein sequence containing 100% experimental epitope amino acid residues. If similar epitope groups were to be classified as completely overlapping, the number of completely overlapping sub-sets would increase to 192 (63.4% of all theoretically-predicted allergenic peptides), and the number of FLAPs that partially overlap with experimental epitopes would be reduced to 51 (16.8% of all theoretically-predicted allergenic peptides). The percentage of unrelated potential epitopes would be thus 19.8%.

Proteins containing at least 2 potential epitopes have been divided into subgroups according to A and PPV value as shown in Table 2. For 27 out of 43 proteins mentioned in this table PPV=100%. It means that all potential epitopes found using EVALLER program in the sequences of proteins mentioned above, fully or partially cover known experimental epitopes. Proteins with A value lower than 0.2 are characterised by PPV from 0 to 100%. The PPV value of proteins possessing A values between 0.2 and 0.6 is between 50 and 100%. All proteins with the frequency of occurrence of sequential epitopes larger than 0.6 are characterised by a positive predictive value of 100%. It means that such proteins do not contain potential

**TABLE 2. Classification of allergenic proteins from the BIOPEP database according to PPV concerning overlapping of potential and experimental epitopes and A, and PPV are defined according to equations 1 and 2 respectively (see Methods section). Allergen names are given according to WHO-IUIS and Allergome database. ID number according to BIOPEP database of allergenic proteins are given in parentheses.**

| PPV (%) | A < 0.2 | 0.2 < A < 0.6 | A > 0.6 |
|---------|---------|---------------|----------|
| 100     | Bos d 6 (15); Cha f 1 (62); Cor a 1.0404 (70); Cru g 1 (3); Equ as ALA (116); Gad c 1 (17); Gal d 2 (4); Hom a 1.0102 (63); Ses i 2.0101 (52) | Budd casein alpha s1 (109); Budd casein alpha s2 (79); Budd casein beta (110); Gal d 1.0101 (5); Met e 1 (73); Ovi a casein alpha s1 (103) | Bos d 5 (14); Bos d 8 alpha s1 (9); Bos d 8 alpha s2 (10); Bos d 8 beta (11); Bos d 8 kappa (12); Bos g casein kappa (111); Bos i casein beta (113); Bos i casein kappa (114); Budd b BLG (108); Cap h BLG (100); Ovi a BLG (77); Ran i BLG (128) |
| 50–99.9 | Ara h 3.0101 (47); Ara h arachin 6 (51); Gyl m Bd 30 KI (42); Gyl m Bd 30 K2 (71); Hom a 1.0101 (72) | Cap h casein alpha s1 (78); Pan s 1 (64) | - |
| 0.1–49.9 | Ara h 4 (98); Ara h 4.0101 (48); Equ c BLG (122); Fag e 1 (40); Ran e 2.0101 (94); Ses i 3.0101 (53) | - | - |
| 0       | Equ as BLG (117); Gad m 1.0201 (92); Gad m 1.0201 (16) | - | - |
epitope unrelated to the experimental ones. Proteins characterised by high A values contain a high number of epitopes found using experimental mapping strategy. Thus, it is easy to find experimental epitopes overlapping with the potential ones. Low PPV (below 50%) values correspond with low A values. This fact may be explained by the incomplete knowledge about epitopes. In addition to sequential epitopes, allergenic proteins also contain conformational epitopes [Pomés, 2010; Ponomarenko et al., 2011]. They are not mentioned in the BIOPEP database, but may also overlap with FLAPs (potential epitopes) pointed out by EVALLER program.

Ranges of A and PPV values of allergenic proteins of various origin are presented in Table 3. The discussed group includes the highest-risk allergens [Jędrzychowski et al., 2008]. Milk proteins form a dominant group among the allergens characterised by high values of A indicating a high number of known experimental epitopes. They were subjected to extensive studies aimed at epitope determination as compared with allergenic proteins from other sources [Vaughan et al., 2010]. Moreover sequences of milk proteins are highly conserved and contain common experimental epitopes as pointed out by Minkiewicz et al. [2011]. The experimental epitopes were attributed to bovine milk proteins [Monaci et al., 2006; Vaughan et al., 2010; IEDB-AR database]. The current list of bovine milk epitopes has been markedly enriched as compared with that presented in the above-mentioned review. On the other hand, calculations restricted to the epitopes mentioned in the review of Monaci and coworkers [2006] give overall results (PPV and frequency of sequential epitope occurrence (A) for allergenic proteins belonging to various homology-based families. ID numbers and names of particular families are taken from AllFam database. A and PPV are defined according to equations 1 and 2 respectively (see Methods section).

### Table 4. Ranges of positive predictive value concerning overlapping of potential and experimental epitopes (PPV) and frequency of sequential epitope occurrence (A) for allergenic proteins belonging to various homology-based families. ID numbers and names of particular families are taken from AllFam database. A and PPV are defined according to equations 1 and 2 respectively (see Methods section).

| PPV (%) | A < 0.2 | 0.2 < A < 0.6 | A > 0.6 |
|---------|---------|---------------|---------|
| 100     | milk (2 proteins); crustacea and mollusca (3 proteins); plants (2 proteins); fishes and amphibians (1 protein); eggs (1 protein) | milk (4 proteins); eggs (1 protein); crustacea and mollusca (1 protein) | milk (12 proteins) |
| 50 -99.9| plants (4 proteins); crustacea and mollusca (3 proteins) | milk (1 protein); crustacea and mollusca (1 protein) | - |
| 0.1 -49.9| plants (4 proteins); milk (1 protein); fishes and amphibians (1 protein) | - | - |
| 0       | milk (1 protein); fishes and amphibians (2 proteins) | - | - |

Ranges of A and PPV values of allergenic proteins belonging to different families are presented in Table 4. Classification
of allergens in the AllFam database is based on the presence of domains summarised in the Pfam database [Radauer et al., 2008]. The presence of characteristic domains is a common basis of protein function annotation [Dessailly et al., 2009], although the alternative solutions of this task are recently proposed [Petrey & Honig, 2009]. Analysis of bioactive peptide profiles of protein containing various domains has been published by Iwaniak & Dziuba [2009]. Information about affiliation of individual allergens to families is available (apart from AllFam database) in Allergome and BIOPEP databases.

### TABLE 5. Allergens containing potential epitopes predicted using EVALLER program, but not containing known, experimentally found sequential epitopes (till the end of April 2011).

| Allergen1 | ID2 | Number of epitopes | Location of epitopes | Origin of protein | Family name and ID according to AllFam database |
|-----------|-----|--------------------|----------------------|------------------|-----------------------------------------------|
| Api g 1.0101 | 57 | 2 | [1–41], [79–135] | plant | Bet v 1-related proteins family (AF069) |
| Api g 1.0201 | 58 | 1 | [38–135] | plant | Bet v 1-related proteins family (AF069) |
| Api g 4 | 60 | 1 | [61–116] | plant | Proline family (AF051) |
| Gad m 1.0101 | 130 | 1 | [1–41] | fish | EF-hand family (AF007) |
| Gad m 1.0102 | 131 | 1 | [1–41] | fish | EF-hand family (AF007) |
| Gal d 3 | 6 | 18 | [5–34], [29–51], [33–54], [40–63], [144–177], [266–288], [270–300], [293–316], [302–325], [339–365], [351–375], [427–465], [521–542], [606–631], [616–639], [637–653], [662–689], [673–700] | egg | Transferrin family (AF068) |
| Gal d 5 | 8 | 13 | [14–35], [90–111], [92–113], [125–149], [131–160], [191–238], [226–247], [334–357], [339–425], [406–427], [413–437], [440–464], [481–503], [530–568] | egg | Serum albumin family (AF056) |
| Gly m 3.0101 | 43 | 1 | [5–45] | plant | Proline family (AF051) |
| Gly m 4.0101 | 44 | 2 | [1–39], [21–70] | plant | Proline family (AF051) |
| Gly m Bd 60K | 41 | 5 | [179–235], [286–372], [216–274], [358–412], [488–583] | plant | Lecithin family (AF045) |
| Hor v 15.0101 | 25 | 6 | [24–86], [68–119], [103–126], [109–130], [119–140], [124–146] | plant | Prolamin superfamily (AF050) |
| Ory s 1.0101 | 37 | 4 | [137–187], [57–99], [168–206], [205–236] | plant | Expansin, C-terminal domain family (AF093) |
| Ory s 12.0101 | 38 | 2 | [9–79], [163–113] | plant | Prolamin family (AF051) |
| Ran e 1.0101 | 33 | 3 | [1–22], [32–53], [64–85] | amphibian | EF-hand family (AF007) |
| Sal s 1.0101 | 22 | 3 | [9–30], [13–34], [16–41] | fish | EF-hand family (AF007) |
| Sal s 1.0201 | 23 | 1 | [2–25] | fish | EF-hand family (AF007) |
| Ses i 6.0101 | 56 | 10 | [25–62], [51–94], [86–128], [112–144], [162–211], [207–243], [231–276], [265–292], [274–323], [393–442] | plant | Lecithin family (AF045) |
| Tri a 3 | 36 | 4 | [1–51], [66–102], [32–67], [92–118] | plant | Expansin, C-terminal domain family (AF093) |
| Tri a TPI | 29 | 2 | [12–47], [32–64] | plant | Triosephosphate isomerase family (AF032) |
| Zea m 14.0101 | 39 | 3 | [7–51], [30–98], [84–120] | plant | Prolamin superfamily (AF050) |

**Total number of potential epitopes:** 83

1. Allergen names according to WHO-IUIS and Allergome database.
2. ID in the BIOPEP Database of allergenic proteins.

(β-lactoglobulins) strongly depends on A value. It indicates that for some proteins belonging to this family knowledge concerning experimental epitopes may be incomplete. Bovine β-lactoglobulin (BIOPEP ID 14) was extensively studied to find fragments interacting with the immunological system. Other proteins from this family contain epitopes identical with these from bovine protein. Lipocalins with a low number of such epitopes are also characterised by a low PPV value. Prediction of epitope location using EVALLER program appears to be promising for proteins revealing low A value and high PPV value, such as proteins from tropomyosin family and papain-like cystein protease family. The group with
the lowest A (A<0.2) and PPV=100% contains also single proteins from four other families (Table 4). Among proteins with a low number of known sequential epitopes prediction quality is rather poor for cupin superfamily. This family does not contain proteins with PPV=100%. Four of the six proteins from this family reveal PPV below 50%. The last family is EF-hand family (fish and amphibian parvalbumins). Among these proteins one has revealed PPV=100%, but two – PPV=0. Parvalbumins are characterised by high sequence variability. Their predicted secondary structure may also vary between species as shown on the example of proteins from Cyprinus carpio and Salmo salar [Iwaniak & Dziuba, 2011]. It is difficult to take any conclusions about predictability of epitope occurrence in proteins form the EF-hand family.

Allergens from the BIOPEP database, containing potential epitopes found by EVALLER program but not containing known experimental sequential epitopes are summarised in Table 5. There is also no data concerning discontinuous epitopes or data concerning these allergens in IEDB or IEDB-3D [Ponomarenko et al., 2011], although we cannot exclude finding such epitopes in the future. Most of allergens indicated in Table 5 are of plant origin (cereals or leguminous plants). Some of allergen AllFam families indicated in Table 5 are represented also among the ones containing both potential and experimental epitopes (Table 4). Proteins from the serum albumin, prolamine or Bet v 1-related family were characterised by a high positive predictive value with a relatively low frequency of epitope occurrence. Results concerning these families look promising in contrast with the proteins belonging to the Cupin and EF-hand families. On the other hand, we cannot exclude that these families contain proteins characterised by poor predictive values such as members of the EF-hand family with PPV=0. The BIOPEP database does not contain proteins from the Profilin, Transferrin and Expansin C-terminal Domain families containing both experimental and potential epitopes.

A comparison with the experimental results is the only way to evaluate in silico prediction. Fragments typical of allergenic proteins, generated by program EVALLER reveal usually good (ca. 80%) likelihood of coverage with experimental epitopes. The suggestion that they cover such epitopes [Björklund et al., 2005] is thus generally confirmed for the used dataset of food allergenic proteins and their epitopes. The positive predictive value may, however, vary among allergen families. The possibility of continuous update is a characteristic property of bioinformatic tools and is considered as major advantage [Wren & Bateman, 2008]. Further enrichment of both databases: BIOPEP and database used by EVALLER program, may lead to the improvement of the Positive Predictive Value concerning overlapping between potential and experimental epitopes.

CONCLUSIONS

Program EVALLER generates fragments characteristic for allergenic proteins (so called Filtered Length-Adjusted Allergenic Peptides – FLAPs). Most of such peptides present in the BIOPEP database of allergenic proteins (ca. 80%) fully or partially overlap with the known experimental ones. Program EVALLER may thus be used as a tool for predicting epitope location although its potential varies considerably among allergen families. The predictive potential was good for milk proteins (α/β-caseins family and κ-caseins family), although these proteins possess numerous experimental epitopes. The predictive potential appears good also for invertebrate tropomyosins (tropomyosins family) and poor for plant allergens from the Cupin superfamily.

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