In vivo and in vitro assessment and proteomic analysis of the effectiveness of physical treatments in reducing allergenicity of hazelnut proteins

Elisabetta De Angelis¹, Danilo Di Bona², Rosa Pilolli¹, Roberto Loiodice², Anna Luparelli¹, Lucia Giliberti², Maria Pia Rossi², Luigi Macchia² and Linda Monaci*¹

Abstract: Hazelnut is a widespread nut species, especially in Europe, that can be consumed raw or roasted, due to its pleasant taste and nutritional properties. Despite the renowned beneficial properties, hazelnuts contain several proteins capable of inducing food allergy in sensitized individuals such as Cor a 2 (profilin), Cor a 8 (lipid transfer protein), Cor a 9 (11S seed storage globulin, legumin-like) and Cor a 11 (7S seed storage globulin, vicilin-like). In the present paper we investigated the effectiveness of autoclave based treatments in decreasing the allergic potential of hazelnut as also assessed by submitting the treated material to in vivo skin prick test and to in vitro immunoblot analysis with sera of allergic individuals exposed to the treated food material. This preliminary analysis shows that autoclave treatment, preceded by hydration and/or followed by drying, seems to be a promising approach and appears to be effective in reducing the allergenicity of hazelnut in most patients, probably due to the denaturation of most major and minor allergenic proteins. This work will open to the opportunity to produce, in perspective, hypoallergenic hazelnut derivatives tolerated by the allergic subjects.

Keywords: food allergy, allergens, allergenicity reduction, skin prick test, proteomic analysis, physical treatments

1. Introduction

Hazelnut is one of the foods most frequently responsible for allergic reactions with an estimated prevalence in Europe of approximately 9.3% of hazelnut specific IgE detected in adults (20 - 54 years)[¹]. Several allergens have been identified in hazelnuts and included in the WHO-IUIS list of allergens [²], but the most studied proteins in relation to food allergy are Cor a 2 (profilin), Cor a 8 (lipid transfer protein), Cor a 9 (11S seed storage globulin, legumin-like) and Cor a 11 (7S seed storage globulin, vicilin-like). Sensitization to Cor a 2 (thermo- and gastro-labile) is associated to oral allergic syndrome, whereas sensitization to Cor a 8, 9 or 11 (thermo- and gastro-resistant) is generally associated to systemic reactions, even severe and life-threatening. Hazelnut consumption is widespread all over the world, especially in Europe and in the Mediterranean diet, due to its pleasant taste and nutritional properties. It can be consumed fresh or roasted, but it is mostly used as ingredient in several food products or preparations (spreads, bakery, pastry, chocolates, and confectionary products). Although some common preparation processes can modify food proteins, they do not necessarily reduce the allergenic potential of the containing food, especially in case of thermo-resistant proteins; this is just the case of nut proteins belonging to the class of seed storage globulins, which maintain their allergenicity even after roasting. The European regulation 1169/2011 on the labeling of food products mandates the obligatory and clear labeling for 14 allergic ingredients, including...
On the other hand, hazelnut might contaminate food products as "hidden allergen", as a result of cross-contamination occurring during manufacturing and this can inadvertently expose allergic consumers to its ingestion. The identification of physical treatments capable of reducing the potential of hazelnut allergens to trigger reactions in sensitized individuals can pave the way for the production of hypoallergenic hazelnut derivatives tolerated by the allergic subjects; this kind of investigation has already been carried out by our team for other nuts, as the case of almonds with an evaluation done in vitro [4]. In the present study we investigated the effects on hazelnut subjected to autoclaving under different schemes, preceded or not by hydration and eventually followed by drying; after obtaining treated hazelnut extracts, we assessed their protein composition, performing gel electrophoresis, western blot and mass spectrometry, and also their skin reactivity, performing a skin prick testing in patients with hazelnut allergy.

2. Materials and Methods

2.1 Patients enrollment for the study

Twenty-two patients with presumed hazelnut allergy were selected at the University Hospital of Bari (Unit of Allergology). Presumed hazelnut allergy (no oral food challenge was performed to confirm diagnosis) was defined based on allergic symptoms following hazelnut ingestion (such as urticaria, angioedema, allergic oral syndrome, asthmatic symptoms, anaphylaxis) and a positive skin prick test and hazelnut specific serum IgE test [5].

Skin prick tests (SPTs) were carried out on the volar surface of the forearm by puncturing the skin through a droplet of a 100 µg/mL allergen with an 1 mm point length standardized needle (ALK-Abellò, Milan, Italy). Histamine 10 mg/mL was used as the positive control. SPTs were performed by placing a drop of extract on the skin of the forearm and subsequently pricking the skin underneath this drop. The test responses were considered positive if the wheal produced had a mean diameter of at least 3 mm. Skin reactivity was also expressed by measuring as the area of the wheals (mm²). All the SPT for any allergen were performed in duplicate.

Patients underwent skin testing with 3 different commercial extracts of hazelnut (Lo-farma®, 2% w/v, ALK-Abello®, 1:20 w/v, and Stallergenes® (Alyostal®), 1 IC/mL) and the specifically prepared raw hazelnut seed extracts (Corylus avellana, var. Italiana) at concentration of 2 mg/mL, left untreated (unmodified, denominated A1) or subjected to differential physical treatments (marked as A2, A3 and A4, detailed in the following paragraph).

Saline was used as the negative control of commercial extracts, whereas an urea buffer solution used as diluent for all the hazelnut extracts was used as negative controls for hazelnut extracts.

2.2 Chemicals

The study was performed on raw hazelnut seeds (Corylus avellana, var. Italiana) provided by Besana S.p.A. (San Gennaro Vesuviano, Napoli, Italia). Trizma-base, urea, sodium chloride, ammonium bicarbonate (AMBIC), iodoacetamide (IAA), dithiothreitol (DTT), and all chemicals for electrophoresis, namely sodium dodecyl sulfate-SDS, glycine, glycerol, Coomassie brilliant blue-G 250, were provided by Sigma Aldrich (Milan, Italy). Acetonitrile (Gold HPLC ultragradient), Methanol (HPLC grade), trifluoroacetic acid (TFA) and Bromophenol blue were purchased from Carlo Erba Reagents (Cornaredo, Milan, Italia). Ultrapure water was produced by a Millipore Milli-Q system (Millipore, Bedford, MA, USA) while formic acid (MS grade) was purchased from Fluka (Milan, Italy). Polytetrafluoroethylene (PTFE) filters (0.45µm) were obtained from Sartorius (Göttingen, Germany) while syringe filters in cellulose acetate (CA) 1.2 µm were purchased from Labochem Science S.r.l. (Catania, Italy). As for in gel protein digestion, trypsin (proteomic grade) was from Promega (Milan, Italy).

2.3 Hazelnut autoclave treatments
Autoclaving treatments were accomplished on a total of 3 raw hazelnut kernels (corresponding to approximately 15 g) placed into a centrifuge tube. Specifically, three different processing schemes were taken into consideration, namely i) autoclaving, ii) sample pre-hydration followed by autoclaving and iii) sample pre-hydration followed by autoclaving and finally drying in a stove overnight at 60°C. For the hydration step, 50 mL of ultrapure water were added to raw hazelnut kernels followed by 2 hours of shaking at room temperature in an orbital shaker (KS 4000 i-control shaker, IKA Works GmbH & Co. KG, Staufen, Germany). Water was discarded before autoclaving. Autoclaving was accomplished by setting the equipment as following: temperature 134 °C, pressure of 2 atm, cycle time 10 min. The system took about 40 min to reach the final temperature of 134 °C. The three different schemes investigated are hereafter summarized:

a) Hazelnut autoclaved for 10 min (A2),
b) Hazelnut prehydrated + autoclaved (A3),
d) Hazelnut prehydrated + autoclaved + drying (A4).

As positive control, raw hazelnut not undergoing any treatment was also included in the study (A1).

The workflow of the proteomic analysis carried out in this work is reported in Figure 1.

2.4 Protein extraction and quantification

Before analysis, raw and processed hazelnut seeds were milled by using an electric miller (Mulinex, Milan, Italy) and proteins were extracted by following the protocol described by Bavaro et al., 2018 [6]. Briefly, 10 mL of TBS buffer (50 mM Tris-HCl, 150 mM NaCl, pH 8 + 1M Urea) were added to 0.4 g of flour and left shaking for 2h at room temperature in an orbital shaker (KS 4000 i-control shaker, IKA Works GmbH & Co. KG, Staufen, Germany). The samples were then centrifuged for 15 min at 1734 x g at 4°C and the supernatant carefully collected while the upper phase discarded. The samples were filtered through 1.2 µm CA syringe filters before successive analysis.

The total protein content of raw and autoclaved hazelnut was calculated by Bradford assay (Quick Start™ Bradford Protein Assay, Bio-Rad Laboratories s.r.l., Segrate MI, Italy) that was carried out according to the manufacturer’s instruction. As reference standard the protein bovine serum albumin (BSA, 0.125-1 mg/ml) was used. Samples were stored at -20°C until their use and filtered through 0.45 µm PTFE filters just before electrophoretic analysis.

2.5 Electrophoretic analysis of hazelnut proteins

Proteins extracted from raw and processed hazelnut samples were then profiled by means of sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE). Specifically, ten micrograms of proteins were separated under reducing condition on a 8-16% polyacrylamide pre-cast gel (13.3 cm x 8.7 cm x 1.0 mm), using a Mini-Protean Tetra Cell equipment (Bio-rad Laboratories, Segrate, MI, Italy). Before electrophoresis analysis, samples were denatured with Laemmli buffer (62.5mM TrisHCl, pH 6.8, 25%
glycerol, 2% SDS, 0.01% Bromophenol Blue, 100 mM DTT) (1:1 ratio) for 5 min at 100 °C. Electrophoretic separation was performed at 60V for the first 20 min and then at 100V until the end of the run by using a TGS (25 mM Tris, 192 mM glycine, 0.1% SDS) solution as running buffer. Gels were finally stained with Coomassie Brilliant Blue G-250 solution and the protein profiles detected on a ChemiDoc™ Imaging System (Bio-Rad Laboratories, Segrate, MI, Italy). Precision Plus Protein™ all blue standards (10-250 kDa, Bio-Rad Laboratories, Segrate, MI, Italy) was used as protein reference for molecular weight.

2.6 In-gel tryptic digestion and LC-MS analysis

The most relevant protein bands detected along the electrophoretic gel of raw and treated hazelnut samples were then submitted to in gel digestion for identification purposes. Briefly, the selected bands were excised from the polyacrylamide gels and in-gel trypsin digested according to the protocol described by De Angelis et al., 2017 [7]. After drying, each sample was re-suspended in 100 µl of H₂O/ACN 95/5+0.1% formic acid (v/v) and 5 µl were injected into LC/MS apparatus. Peptide mixtures obtained from protein bands in-gel digested referred to samples A1, A2, A3 and A4 were analyzed by untargeted proteomic analysis by HPLC-MS/MS equipment consisting in a Q-Exactive™ Plus Hybrid Quadrupole-Orbitrap™ Mass Spectrometer coupled to a UHPLC pump systems (Thermo Fisher Scientific, Bremen, Germany). For the peptide chromatographic separation, a reversed phase Acclaim™ PepMap100, C18 analytical column (1 mm × 15 cm × 3m, 100˚A porosity, Thermo Fisher Scientific, Bremen, Germany) was used, by setting the flow rate at 60 µl/mL. The elution gradient used for peptide separation was the following: 0-60 min solvent B increased from 10% to 60%, 60-61 min further increase from 60% to 80%, then kept constant for 10 min, 70-90 min at a

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**Figure 1.** Workflow for proteomic investigation, encompassing extraction step, electrophoretic analysis, protein identification by Mass Spectrometry and immunoblot analysis for immunoreactivity assessment.
constant 10% for column conditioning before next injection. Solvent A = H2O+0.1% FA, solvent B= Acetonitrile/H2O (80/20 v/v) + 0.1% FA. Volume injection was set to 5 µL and each sample was injected twice in MS. Spectra were acquired in the mass range of 150-2000 m/z by applying the data dependent (FullMS-dd2) acquisition mode analysis and only positive ions were considered. Up to 10 most intense ions in MS1 were selected for subsequent fragmentation in MS/MS mode. A resolving power of 70,000 full width at half maximum (FWHM), a microscan of 1, an automatic gain control (AGC) target of 1.00 e6 and a maximum injection time (IT) of 40 ms were set to generate precursor spectra (full MS analysis). The settings for MS2 fragmentation experiments were: the following; resolving power 17,500 FWHM, microscan of 1, AGC target 5.00 e5, maximum IT 50 ms, loop count 10, MSX count 1, isolation window of 2.0 m/z, isolation offset 0.4 m/z and normalized collision energy (NCE) at 27 and 30 eV by activating the stepped option; as for dd settings maximum AGC target was set at 5.00 e5 with intensity threshold of 1.0 e3, dynamic exclusion at 20 s, peptide match set to preferred and exclude isotopes enabled. All ions with charge equal to 1 and higher than 4 were excluded. As for HESI source, the following conditions were set: spray voltage 3400, capillary T °C 320, sheat and auxiliary gas 25 and 15 respectively, probe heater T °C 290 and S-Lens 55.

Raw MS data were then processed via the commercial software Proteome Discoverer™ version 2.1.1.21 (Thermo-Fisher-Scientific, Bremen, Germany) and protein identification was obtained by SequestHT search against the hazelnut proteins database extracted by Swiss Prot DB on the base of the taxonomy code of Corylus avellana (ID: 13451, containing about 501 sequences). Trypsin was selected as cleavage enzyme and in all cases mass tolerance on the precursor and fragment ions was set to 5 ppm and 0.05 Da, respectively. Finally, only trustful peptide-spectrum matches were taken into account and a minimum of three peptides was set as threshold for protein identification, after filtering the peptide list to the sequences assigned with at least medium confidence (FDR<5%).

2.7. Immunoblot for IgE-Binding Assay

Proteins of raw and differently treated hazelnuts separated by SDS-PAGE (corresponding to 10 µg of proteins loaded for A1, A2, A3 and A4) were electroblotted onto a 0.2 µm nitrocellulose membrane (Bio-Rad Laboratories, Segrate, MI, Italy) using a Trans-Blot Cell (Bio-Rad Laboratories, Segrate, MI, Italy) for 7 min (1.3 A, 25 V). Immunoblotting tests were accomplished according to the protocol reported by Bavaro et al., 2019 [8]. As primary antibody, the pooled sera of a total of 14 allergic individuals showing different clinical symptoms (urticaria, itching, hands/throat angioedema) previously diluted in TBS-T (pH 7.4, 10 mM Tris, 50 mM NaCl, 0.1% Tween 20) at a 1/50 ratio was used and kept shaking overnight at 4 °C. As a secondary antibody goat anti-human IgG (H + L) horseradish peroxidase (HRP) conjugated (Bio-Rad Laboratories) and anti-human IgE (ε-chain specific)–horseradish peroxidase conjugated (Sigma Aldrich, Milan Italy) each diluted 1/5000 (v/v) in TBS-T were added. Specifically, four different immunoblotting experiments were accomplished: in the first two experiments two different sera pools, each obtained by combining the sera of 7 patients per group, were used and the reactive bands detected by using the goat anti-human IgG (H + L) horseradish peroxidase (HRP) conjugate (vide infra) as secondary antibody. In the third and fourth experiments, the sera of all 14 patients were pooled together and the reactive proteins were identified after incubation with anti-human IgE (ε-chain specific)–peroxidase secondary antibody binding and goat anti-human IgG (H + L) horseradish peroxidase (HRP) (vide infra). Final images were obtained on a ChemiDoc™ MP Imaging System.

2.8 Statistical analysis

T-test was used to compare the average areas of the wheals.

3. Results and Discussion

3.1. Effects of hazelnut thermal/pressure treatment on patient response
Firstly, we sought to assess whether the modified hazelnut extracts were associated with a reduced patients’ response in vivo. To this aim, 22 patients with diagnosed hazelnut allergy (16 females [72.7%]; mean age, 28.1 ± 9.4 years; range: 17-48 years) underwent skin prick testing [9]. Specifically, 19 (86.4%) out of 22 patients reported urticaria/angioedema, 3 patients reported oral allergy syndrome (13.6%) and 1 patient reported both manifestations (4.5%), suggesting that most patients were likely sensitized to hazelnut proteins, like Cor a 8, Cor a 9, Cor a 11, that are known to be responsible for systemic symptoms (Table 1).

Table 1. Hazelnut allergens. Overview of different protein families, MWs and specific related clinical pictures.

| Protein name | Protein family | MW (kDa) | Clinical picture          |
|--------------|----------------|----------|---------------------------|
| Cor a 1.04   | PR-10 (Bet v 1 like) | 17       | OAS                       |
| Cor a 2      | profillin (Bet v 2 like) | 14       | OAS                       |
| Cor a 8      | LTP             | 9        | Systemic reactions        |
| Cor a 9      | SSP (11S globulin - legumin like) | 40       | Systemic reactions        |
| Cor a 11     | SSP (7S globulin - vicilin like) | 48       | Systemic reactions        |
| Cor a 12     | Oleosin         | 17       |                           |
| Cor a 13     | Oleosin         | 14 – 16  |                           |
| Cor a 14     | SSP (2S albumin) | 10       |                           |

The SPT reactivity (defined as the mean wheal area of 2 independent measurements) reported with the hazelnut extract A1 (unmodified hazelnut seed extracts - Corylus avellana, var. Italiana) was 24.6 ± 12.3 mm², comparable to that produced by the 3 commercial extracts (ALK, 26.5 ± 13.2 mm²; Stallergenes, 22.8 ± 12.3 mm²; Lofarma, 18.7 ± 9.0 mm². This suggested that the native extract A1 was effective and suitable to be used for comparisons with modified extracts. Negative controls (either saline or urea solutions) did not elicit any skin reaction (no wheal, no erythema, no pruritus).

Only 6 patients (27.3%) showed reactivity with the A2 extract. This reactivity appeared lower than that of the native extract (mean wheal area of the 6 reactive patients, 15.2 ± 4.1 mm²; P [A1 vs. A2] = 0.005) (Figure 2; Table 2).

Only 2 out of the 6 patients reactive to A2 showed reactivity with the A3 extract. The mean wheal area of the 2 reactive patients was 13.5 mm² (patient #15) e 4.5 mm² (patient #20), respectively (Figure 2; Table 2).

Finally, only 1 patient (patient #15) showed SPT reactivity (wheal area, 13.5 mm²) with the A4 extract.

Collectively, these results suggest that all the 3 different treatments are effective in reducing hazelnut allergenicity in a meaningful proportion of patients. Treatments 3 and 4 appeared to be the most effective ones. Notably, one patient out of 22 did not show a significant reduction of SPT reactivity even with the A4 extract. Further analyses will be necessary to define the specific epitopes this patient is reactive to.
Table 2. SPT reactivity of patients with differentially treated hazelnut extracts.

| Extract | # reactive patients N (%) | Wheal area, mm² (mean ± SD) | P          |
|---------|---------------------------|-----------------------------|------------|
| A1      | 22 (100%)                 | 24.6 ± 13.1                 | Reference  |
| A2      | 6 (27.3)                  | 15.2 ± 4.1²                 | p < 0,01   |
| A3      | 2 (9.1)                   | 13.5 (patient #15)          | n.e.¹      |
|         |                           | 4.5 (patient #20)           |            |
| A4      | 1 (4.5)                   | 13 (patient #15)            | n.e.¹      |

¹n.e = not estimable.
²mean ± SD calculated with data from only the 6 reactive patients; reference means that this group (A1) is used for comparisons with the other groups (A2, A3, A4) for statistical analyses.

3.2 Effect of thermal/pressure treatment on hazelnut protein solubility

Food processing is known to alter the final structure and function of proteins, modifying some crucial properties such as the final solubility. Denaturation, hydrolysis of peptide bonds, restructuring of disulphide bonds and interaction with other components (carbohydrates and lipids) that frequently could occur during treatments are among the causes leading to reduced protein solubility [10]. In the light of this, we first sought to evaluate how autoclaving/pressure processing could alter the final solubility of hazelnut proteins by estimating the total protein content of samples undergoing the different treatments. The Bradford assay was used for this purpose (Figure 3). A progressive reduction in protein recovery was shown in the treated hazelnut samples compared to the untreated counterpart. The total protein content of the autoclaved samples (A2) appeared to be 40% lower than control sample (A1), with a more dramatic reduction observed for pre-hydrated/autoclaved sample (A3) and pre-hydrated/autoclaved/dried samples (A4), which showed a 70% decrease in protein recovery compared to the untreated hazelnut sample (Figure 3).
Protein content, estimated by Bradford protein assay, corresponding to untreated hazelnut (A1) and hazelnut submitted to autoclaving (A2), prehydration/autoclaving (A3) and prehydration/autoclaving followed by drying (A4). These results are in accordance with previous studies, reporting the effects of autoclaving on other nuts species, such as almonds or peanuts [4, 6,11, 12]. In general, it was demonstrated that autoclave based treatments produce a decrease in protein recovery and this trend is more marked when treatment is accomplished at harsher conditions or when it is preceded by a hydration step. This phenomenon could be attributed to the numerous biochemical and structural modifications occurring on protein moiety taking place during the combination of heat and pressure treatments. The latters, in combination with aggregation phenomena due to the formation of intra- or inter-molecular covalent and non-covalent interactions between proteins or protein-food matrix, could promote protein precipitation with a consequent decrease in protein solubility and protein content in the final extract [13]. As for hazelnut, Lopez et al. in 2012 demonstrated that autoclaving could alter the secondary and tertiary structure of this tree nut proteins also inducing a glycosylation reaction [14]. In the current study, the reduced protein content observed in treated hazelnuts confirmed that autoclave-based treatments could modify the protein structure of hazelnut proteins, leading to a reduction of their solubility. This effect appears to be enhanced by preceding autoclaving with water incubation. In addition to a reduced solubility, a fragmentation of the allergen protein is also likely to occur during autoclaving due to the pressure and temperature applied. It is already known that according to some OFC studies [15, 16] new thresholds might be proposed for allergen undergoing heating / baking treatments in food matrices as they can induce modification of protein structure and decrease in its allergenicity. In line with this, it is reasonable that the reduced protein extraction or protein degradation after physical treatments might have influenced the overall SPT reactivity.

3.3 SDS-PAGE profiling of processed hazelnuts and proteins identifications

Raw (A1) and differently processed hazelnuts (A2, A3 and A4) were electrophoretically profiled in order to evaluate possible changes in hazelnut protein content and structure induced by the different treatments. Specifically, 10 µg of untreated (A1) and treated hazelnut proteins (A2, A3 and A4, respectively) were analyzed by SDS-PAGE and peculiar protein profiles were observed for each sample analyzed (Figure 4). In the untreated

![Figure 3. Protein content, estimated by Bradford protein assay, corresponding to untreated hazelnut (A1) and hazelnut submitted to autoclaving (A2), prehydration/autoclaving (A3) and prehydration/autoclaving followed by drying (A4).]
sample (Figure 4A, lane A1), several bands in the region of 30 - 50 kDa and 10 - 22 kDa were shown. As known from the literature, in the absence of reducing agents, Cor a 9 (11S legumins) is organized in hexameric structures made up of six subunits interacting non-covalently and arranged in an open ring conformation with 360 kDa [17].

Figure 4. A) Comparison between SDS-PAGE protein profiles of hazelnut untreated (A1), autoclaved (A2), prehydrated and autoclaved (A3), prehydrated/autoclaved and subsequently dried (A4). M, MW reference standards. B) Protein bands excised from the gel (numbered from 1 to 6) to be submitted to tryptic digestion and analysis by untargeted High Resolution Mass Spectrometry.

Each subunit is composed by an acidic polypeptide (30-40 kDa) linked to a basic polypeptide (around 20 kDa) by a disulphide bond [18]. Under reducing conditions, acid and basic subunits are released and they are clearly visible in the lane of the untreated sample (Figure 4A, lane A1). Other bands were visible over the 37 kDa region and below 20 kDa. They are likely to be attributed to Cor a 11, Cor a 8 and Cor a 14, whose MWs were reported to be approximately 48 Da, 9 kDa and 15-16 kDa, respectively [18]. After autoclaving (Figure 4 A, lane A2), a general decrease in bands intensity was recorded, with a concomitant disappearance of the proteins banding at 50 kDa and below 20 kDa that were putatively attributed to Cor a 11, Cor a 8 and Cor a 14 allergens. As for Cor a 9, a marked reduction of signals corresponding to acid and basic subunits was observed, likely attributable to a reduced content of the allergen following the thermal/pressure treatment applied. On the contrary, protein profiles of hazelnut samples incubated with water before autoclaving (Figure 4 A, lanes A3 and A4) appeared as a smear of peptides, with low MW (10–20 kDa), probably produced by fragmentation occurring during the treatments. It is worthy to be noted that drying process after autoclaving (A4 treatment) did not produce any significant difference in protein profile with respect to the A3 sample.

SDS-PAGE profiles are in accordance with protein assay results (Figure 3), where a progressive reduction in protein content was displayed among the hazelnut samples differently treated, with a more marked decrease observed for pre-hydrated/autoclaved and pre-hydrated/autoclaved/dried samples.
To gain insights on the protein content of the specific bands identified by SDS-PAGE, the most relevant bands, which resulted differently expressed after different treatments (Figure 4B, lanes A2, A3 and A4) were excised from the gel (the excised bands are numbered from 1 to 6), submitted to tryptic digestion and analyzed by untargeted High Resolution Mass Spectrometry. MS spectra were then processed by Proteome Discoverer software for protein identification. Specifically, the Uniprot database referred to *Corylus avellana* (last accession on 26th of November 2021) was interrogated and results are resumed in Table 3.

Table 3. Summary of proteins identified by Proteome Discoverer software referred to SDS-PAGE protein bands excised and in gel digested from protein pattern of autoclaved (A2), prehydrated/autoclaved (A3), prehydrated/autoclaved/dried (A4) hazelnut samples.

| Sample | Band | Accession | Description | Allergen | Coverage (%) | #Peptides (unique) | Score |
|--------|------|-----------|-------------|----------|--------------|--------------------|-------|
| A2     | 1    | A0A0A0P7E3 | Cor a 9 allergen (*Corylus avellana*) | Cor a 9 | 52 | 21 (2) | 124 |
|        |      | Q8W1C2    | 11S globulin-like protein (*Corylus avellana*) | Cor a 9.0101 | 45 | 23 (2) | 112 |
|        | 2    | A0A0A0P7E3 | Cor a 9 allergen (*Corylus avellana*) | Cor a 9 | 44 | 17 (4) | 207 |
|        |      | Q8W1C2    | 11S globulin-like protein (*Corylus avellana*) | Cor a 9.0101 | 37 | 14 (1) | 202 |
|        | A3   | A0A0A0P7E3 | Cor a 9 allergen (*Corylus avellana*) | Cor a 9 | 40 | 16 (3) | 118 |
|        |      | Q8W1C2    | 11S globulin-like protein (*Corylus avellana*) | Cor a 9.0101 | 33 | 14 (1) | 116 |
|        |      | A0A1I9RG40| Ribulose bisphosphate carboxylase large chain (*Corylus avellana*) | Cor a 9 | 19 | 8 (8) | 16 |
|        | 4    | A0A0A0P7E3 | Cor a 9 allergen (*Corylus avellana*) | Cor a 9 | 28 | 11 (11) | 38 |
|        |      | Q8W1C2    | 11S globulin-like protein (*Corylus avellana*) | Cor a 9.0101 | 32 | 18 (2) | 111 |
|        | A4   | A0A0A0P7E3 | Cor a 9 allergen (*Corylus avellana*) | Cor a 9 | 34 | 19 (2) | 109 |
|        |      | D0PWG2    | 2S albumin (*Corylus avellana*) | Cor a 14/Cor a 14.0101 | 44 | 8 (8) | 15 |
|        |      | A0A0A0P7E3 | Cor a 9 allergen (*Corylus avellana*) | Cor a 9 | 30 | 14 (2) | 98 |
|        |      | Q8W1C2    | 11S globulin-like protein (*Corylus avellana*) | Cor a 9.0101 | 28 | 14 (2) | 98 |
|        | 6    | D0PWG2    | 2S albumin (*Corylus avellana*) | Cor a 14/Cor a 14.0101 | 31 | 8 (8) | 4 |
|        |      | Q8S4P9    | Vicilin Cor a 11.0101 (*Corylus avellana*) | Cor a 11/Cor a 11.0101 | 8 | 2 (2) | 2 |

As expected, bands 1 and 2 of the autoclaved sample (Figure 4 B, lane A2) were attributed to Cor a 9 allergen, suggesting the resistance of this allergen to autoclave processing although the reduced intensity of the relative bands leads to supposing that some partial degradation/structural alteration of the proteins likely occurred during the treatment. Cor a 9 allergen was found also in bands 3 and 4 (Figure 4 B, lane A2). The smeared bands visible in protein profiles of pre-hydrated/autoclaved samples followed or not by drying (approximately 10-20 kDa, bands 5 and 6, respectively) (Figure 4 B, lanes A3 and A4) were instead produced by a mix of peptides belonging to Cor a 9, Cor a 11 and Cor a 14 allergens (Table 3).
The impact of autoclaving on protein stability of tree nuts and peanuts was already reported in the literature [4, 6, 14, 19, 20], with some papers exploring the effect of water incubation before thermal/pressure treatment [4, 6]. Similar to what described in the present work, these authors observed that samples pretreated with water before autoclaving showed a SDS-PAGE protein profile more fragmented and degraded with respect the autoclaved counterparts. To explain these phenomena, they suggested that probably water absorbed by seeds during incubation facilitates the propagation of heat in the inner part of the seed promoting protein disaggregation with consequent decrease in band intensity. In addition, our results are in line with what was described by Lopez et al. in 2012, who studied the effect of different conditions of autoclaving and high-pressure treatments on the final allergenicity of hazelnut flour. The authors highlighted the disappearance of the main hazelnut allergens protein bands in SDS-PAGE profile of hazelnuts autoclaved at 131°C for 15 or 20 min, ascribing these results to molecular alterations or post-translational modifications (PTMs, e.g. glycosylation) that likely occur during autoclaving. Indeed, by generating a series of bioinformatics homology-based 3D models, they found that the structure of the main hazelnut allergens (Cor a 8, Cor a 9) resulted altered after autoclaving with a new site of glycosylation found in Cor a 11 allergen [14].

3.4 Immunoblotting experiments

To assess the immunogenicity and the allergenicity of hazelnut seeds subjected to the different autoclaving treatments (A2, A3, A4), immunoblotting experiments were performed. Sera from 14 patients showing different clinical symptoms, both systemic and OAS, were used for immunoblotting experiments (Figure 5).
Figure 5. SDS-PAGE along with Immunoblots of hazelnut untreated (A1) or submitted to autoclave (A2), pre-hydration/autoclave (A3), prehydration/autoclave/drying (A4). M, Precision Plus Protein™ All Blue Standard (10–250 kDa, Bio-Rad Laboratories, Segrate, Milan, Italy). Panel A and B: immunoblotting profiles of 14 patients allergic to hazelnut pooled together and incubated with goat anti-rabbit IgG Ab (Panel A) and anti-human IgE Ab (ε-chain specific) secondary antibody (Panel B). Panel C and D: immunoblotting profiles of two subgroups of 14 allergic patients (7 allergic sera pooled together for each group) both incubated with goat anti-rabbit IgG Ab.

The pooled sera of a total of 14 allergic patients were used as the primary antibody. Then, the membranes were incubated with a goat anti-rabbit IgG Ab (Figure 5 A) or an anti-human IgE Ab (ε-chain specific) (Figure 5 B), as the secondary antibodies. Finally, to increase the assay sensitivity, 2 additional subgroups with 7 pooled sera per group (Figure 5 C-D) were incubated with a goat anti-rabbit IgG Ab (vide infra). Figure 5 A shows in untreated hazelnut (lane A1) one main reactive band with MW of approximately 20 kDa visible, along with weak intensity bands at 50 kDa and in the range of 10 and 15 kDa. Two additional blurred signals between 21-22 kDa were displayed in the A1 sample as well. According to the literature, protein banding at 20 kDa could be putatively attributed to the basic subunit Cor a 9 hazelnut allergens as well as bands in the range of 21-22 kDa. On the contrary, the reactivity displayed at 50 kDa could be ascribable to Cor a 11 allergen, which original MW is 48-50 kDa, while the weak signal in the range of 10-15 kDa could be attributed to Cor a 8 or Cor a 14 allergens with MWs falling in this range. After autoclaving treatment (Figure 5 A, lane A2), the weak reactivity of signals at 50 kDa and below 15 kDa became negligible, while the intense spot at 20 kDa and bands in the range of 21-22 kDa still persisted. Grounding on proteomic investigation accomplished on SDS-PAGE protein profile, these bands were attributed to Cor a 9 and its isoform allergen Cor a 9.0101 (Table 3), confirming that these allergens survived to autoclave processing preserving its allergic
potential. On the contrary, no reactive signal was displayed for the band at 50 kDa, putatively ascribed to Cor a 11, along with the 10-15 kDa hazelnut allergens (likely Cor a 8 and Cor a 14), confirming the susceptibility of these proteins to autoclave processing (Figure 5 A). As for the prehydrated/autoclaved (Figure 5 A, lane A3) and prehydrated/autoclaved/dried (Figure 5 A, lane A4) samples, no clearly identifiable reactive bands were shown, thus confirming the key role of the water imbibition step in the structural and conformational alteration/degradation phenomena induced and enhanced by the thermal/pressure treatment applied on proteins. Cor a 9 was demonstrated to be a highly well-structured protein, enriched with a beta-sheet core and with long unstructured loops. These loop regions were found structurally unstable and were predicted to retain linear epitopes located at the external faces of protein, thus being exposed to solvent [14].

After submitting hazelnut samples to autoclaving (121 °C or 138 °C for 15 or 30 min), Lopez and co-workers obtained a reduction of the Cor a 9 allergenicity, since no bands corresponding to this protein was visible by SDS-PAGE analysis. In the light of this, the authors hypothesized that the allergenicity of Cor a 9 could be predominantly ascribable to structural conformation and not to linear epitopes [14]. Our results seem to support this hypothesis. Indeed, the enhanced action of previous water incubation on autoclaving (A3, A4) seems effective in impairing the Cor a 9 structure with no reactive epitopes surviving after processing, differently from with what was observed for the solely autoclaved sample (Figure 5 A, lane A2). By visual inspection of the SDS-PAGE pictured in figure 4B, despite the visible smear bands below 20 kDa, (produced by a mixture of Cor a 9 and Cor a 14 allergens peptides in A3 sample and Cor a 9, Cor a 14 and Cor a 11 allergens peptides in A4 sample as highlighted by proteomic investigation in Table 3), no reactive signals were displayed in the putatively corresponding lanes of immunoblotting profiles. As already emerging from the electrophoretic analysis, no significant changes in immunoblotting profile and consequently in the final allergenicity of prehydrated/autoclaved hazelnut were found by drying the sample (Figure 5, lane A3).

Similar results were obtained by incubating the A1-A4 samples with anti-human IgE Ab (vide infra) (ε-chain specific) secondary antibody, confirming that specific IgE can bind protein bands detected in untreated and autoclaved samples and, thus, likely trigger an allergic reaction in vivo. On the contrary, no IgE reactivity was displayed for A3-A4 samples pointing out their likely lack of allergic potential.

Considering that sensitization to specific hazelnut proteins could vary among allergic individuals, additional immunoblotting experiments were performed dividing the 14 patient sera in 2 groups of 7 patient sera. Our aim was to unveil possible different pattern of sensitization of different patient subgroups. In particular, Figure 5 C and D shows immunoblotting experiments conducted using 2 different sera pools each of which made of 7 patients’ sera. These immunoblots showed different sensitivity compared to those showed in Figure 5 A and B (using the pool made by all the 14 patients), depending on the differential antibody concentration of each patient’s serum in the subgroups. As a matter of the fact, the reactivity profile varied among the different pools. Figure 5 C showed more intense bands at approximately 50 kDa (Cor a 11), 37 kDa (Cor a 9 acid subunit), 20-22 kDa (Cor a 9 basic subunit/Cor a 1.04) and 10 kDa (Cor a 8/Cor a 14), compared to Figure 5 A, B and D. This was expected, considering that pools represent patients’ average response, which can vary along with the differential pool array. Further analyses on individual patient sera will be necessary to accurately identify the individual reactivity profile, and how this could be modified by specific treatments.

The effect of the autoclaving on the final allergenicity of hazelnuts was already investigated in 2012 by Lopez et al. The authors submitted hazelnut defatted flour to autoclave processing at different conditions (121°C 15 min, 121°C 30 min, 138°C 15 min, 138°C 30 min) and investigated the IgE-reactivity of 15 allergic patients via Western blot experiments. In addition, they studied the changes induced by autoclaving to the conformational structure of hazelnut allergens by generating a series of homology-based-bioinformatics 3D-models for allergens Cor a 1, Cor a 8, Cor a 9 and Cora 11. As result, the authors observed that at harsher conditions (138°C 15 min, 138°C 30 min) autoclaving induced a
severe reduction of hazelnut allergenicity in the patients studied. Indeed, the specific-IgE binding of some immunoreactive hazelnut protein-bands such as Cora 1, Cora 8, Cora 9 and Cora 11 (vide infra) decreases. Moreover, the structural analysis (3D-modelling) of these allergens highlighted that a relevant glycosylation occurred in the protein-allergen Cora 11 (vide infra) after autoclaving, pointing out that the combination of temperature/pressure could promote the interaction of protein-matrix, likely altering the final allergenicity of the protein [14].

Very recently Cuadraro et al. investigated the effect of autoclaving on the final allergenicity of Cor a 9, Cor a 14 and Cor a 8 hazelnut allergens by testing the two different autoclaving temperatures, 121°C and 138°C, for 30 min. Whole hazelnut seeds were processed and, similarly to what described by Lopez et al., they observed in the different immunoblot profiles a marked reduction of Cor a 9, Cor a 14 and Cor a 8 reactivity after autoclaving hazelnut material at 138°C for 30 min [21].

In the present investigation we observed, by autoclaving hazelnut seeds at 134°C for 10 min, the intensity of Cor a 9 band appeared to be reduced, with the disappearance of the 50 kDa and 10 kDa bands, putatively ascribed to Cor a 11 and Cor a 8 allergens, as displayed in the SDS-PAGE picture (Figure 3, lane A2). These results are comparable to what obtained by the mentioned authors by applying similar autoclaving conditions (temperature of 121°C for 30 min) [14, 21].

Furthermore, the immunoblot profiles presented here confirmed the IgG and the IgE reactivity of Cor a 9 basic subunits after the autoclaving of hazelnut material only (Figure 5, panel A-D). On the contrary, by incubating hazelnut seeds with water before autoclaving a full degradation/fragmentation of the proteins was observed, with the disappearance of the main allergenic bands in the corresponding SDS-PAGE (Figure 3, lane A3 and A4) and immunoblotting profile (Figure 5, panel A-D). These results are comparable with what observed by Lopez et al. and Cuadraro et al. by analyzing hazelnut autoclaved at 138°C for 30 min via SDS-PAGE and Western blot experiments [14, 21]. In the light of this, it is reasonable to suppose that water incubation strengthens the alternative phenomena induced by autoclaving on proteins allowing to produce the same effect obtained by autoclaving hazelnut at very harsh conditions, such as 138°C - 30 min.

5. Conclusions

This preliminary study shows that autoclave treatments, also preceded by hydration and/or followed by drying, appears to be effective in reducing the allergenicity of hazelnut proteins in most of the patients herein screened by skin prick test, probably due to denaturation/fragmentation of most major and minor allergenic proteins. The reduction of hazelnut proteins solubility due to the specific treatment applied must be also considered in the interpretation of the reduced allergenicity proved by immunoblot tests. Anyway, taking into account that immunoblotting experiments were accomplished by loading the same amount of proteins for all treated samples, the reduction of reactivity in the observed bands could be reasonably attributed to a reduced allergenicity of those proteins. The appearance of a smear corresponding to low MW proteins (lower than 20 kDa) along with the absence of higher MW bands also suggests a possible protein fragmentation caused by physical treatments, and this could be responsible for both reduced IgG and IgE reactivity. Furthermore, unique peptides corresponding to Cor a 9 and Cor a 11 were detected in the low MW degraded proteins (protein bands 5 and 6 in Figure 4 B and Table 3) confirming the hypothesis of protein degradation. Notably, Cor a 9 and Cor a 11 are two of the three most important and harmful hazelnut allergens causing systemic reactions. The third allergen causing systemic reaction, Cor a 8, was likely present in the band approximately at 10 kDa. However, a possible fragmentation of this protein can only be hypothesized, but not demonstrated, because the fragments resulting from the effects of each treatment would be too small to be identified with our immunoblots as they could have escaped the gel meshes during gel running. Further specific experiments will be performed to confirm this hypothesis. Nonetheless, according to studies based on oral food
challenges, threshold doses below which reactions do not occur exist for allergens. Consequently, it is possible that the reduced protein extraction after physical treatments might have influenced SPT reactivity. Whether the reduced SPT reactivity that we observed in our patients could be correlated to a decreased response to Oral food challenge (OFC) needs to be elucidated and will be investigated in further studies.

In conclusion this study, based on patients’ reactivity in vivo (SPT) and their immunoblotting profiles, showed that specific physical treatments can reduce the allergenicity of hazelnut. These results need to be confirmed by double blind placebo-controlled food challenge with the treated extracts. Finally, in the perspective of using these physical treatments in food production, the effects of them on the organoleptic properties of the preparation (smell, taste) is also worthy to be further investigated and optimized for commercialization purposes.

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