The protein structure determines the sensitizing capacity of Brazil nut 2S albumin (*Ber e1*) in a rat food allergy model

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**Abstract**

It is not exactly known why certain food proteins are more likely to sensitize. One of the characteristics of most food allergens is that they are stable to the acidic and proteolytic conditions in the digestive tract. This property is thought to be a risk factor in allergic sensitization. The purpose of the present study was to investigate the contribution of the protein structure of 2S albumin (*Ber e1*), a major allergen from Brazil nut, on the sensitizing capacity in vivo using an oral Brown Norway rat food allergy model. Disulphide bridges of 2S albumin were reduced and alkylated resulting in loss of protein structure and an increased pepsin digestibility in vitro. Both native 2S albumin and reduced/alkylated 2S albumin were administered by daily gavage dosing (0.1 and 1 mg) to Brown Norway rats for 42 days. Intraperitoneal administration was used as a positive control. Sera were analysed by ELISA and passive cutaneous anaphylaxis. Oral exposure to native or reduced/alkylated 2S albumin resulted in specific IgG1 and IgG2a responses whereas only native 2S albumin induced specific IgE in this model, which was confirmed by passive cutaneous anaphylaxis. This study has shown that the disruption of the protein structure of Brazil nut 2S albumin decreased the sensitizing potential in a Brown Norway rat food allergy model, whereas the immunogenicity of 2S albumin remained preserved. This observation may open possibilities for developing immunotherapy for Brazil nut allergy.

**Keywords:** Brazil nut 2S albumin, *Ber e1*, Food allergen, Rats, Reduction, Alkylation, Allergenicity, Immunogenicity

**Introduction**

Humans rather frequently suffer from allergic reactions after consumption of dietary proteins. The prevalence of food allergy is approximately 1-2% in adults and 6-8% in children and most food allergies are mediated by antigen-specific IgE and are characterized as type-I reactions [1]. Cases of severe allergic reactions including anaphylaxis were reported for Brazil nut (*Bertholletia excelsa*) [2] and the storage protein 2S albumin was identified as allergen and classified as *Ber e1*. It has gained particular interest as in the early 1990's it was considered to transfer the gene coding for the Brazil nut 2S albumin by transgenic techniques to soybean in order to improve its nutritional value for animal feed which however never reached the market due to the allergenic nature of the Brazil nut 2S albumin [3].

An important characteristic of food allergens is that they tend to be more stable to the proteolytic and acidic conditions of the digestive tract, which results in an increased probability of reaching the intestinal mucosa, where absorption can occur and allergic sensitization may be induced [4,5]. Many of the major food allergens are comparatively resistant to digestion, and this general characteristic has been considered a risk factor for food allergy induction. Furthermore, *in vitro* digested Brazil nut 2S albumin retains about one quarter of its IgE-binding potency [6], and similar observations were made for 2S albumin-like allergens from peanut [7]. This suggests that even if 2S albumins are digested, their allergenicity in terms of IgE-binding is not completely abolished. Taken together, resistance to digestion does
not provide sufficient information for safety aspects in terms of risk of allergenicity.

The purpose of this study is to evaluate the relation between specific structural aspects of Brazil nut 2S albumin and its sensitizing capacity. Therefore, we prepared two forms of 2S albumin from Brazil nuts, a native- and a reduced/alkylated 2S albumin (2S albumin-R) in which the disulphide bridges are reduced. Both have the same amino acid sequence that provides the same set of potential antigenic linear epitopes, but with different structural organization. Reduction of the disulphide bridges of Brazil nut 2S albumin led to a remarkable decrease in stability toward digestion [8]. Both preparations are here tested in their ability to induce allergic sensitization in a Brown Norway rat model for food allergy.

**Findings**

**Materials and methods**

Disulphide bridges of 2S albumin were reduced and alkylated resulting in unfolding of the protein [8], here referred to as 2S albumin-R. To study the sensitizing capacity of the structurally different forms of 2S albumin [8] in vivo we used an oral Brown Norway rat food allergy model. Young male Brown Norway (BN) rats (3-4 weeks old at arrival) obtained from Charles River (Sulzfeld, Germany) were bred and raised on a commercially available Brazil nut free rodent diet (SDS Special Diet Service, LAD1 (E) SQC, Witham, England). For the oral sensitization animals (n = 10) were exposed by gavage dosing during 6 weeks to either 2S albumin-R or native 2S albumin (0.1 mg or 1 mg protein/ml tap water; 1 ml/animal) without the use of an adjuvant or only water (controls). Blood samples were obtained days 0, 28 and 42 and sera were prepared. Positive control animals (n = 5) were injected intraperitoneally (i.p.) with 0.5 ml of a 0.2 mg/ml RA- or native 2S albumin solution in sterile saline on days 0, 2, 4, 7, 9 and 11. In the positive control animals, the immune response was potentiated at day 0 with 0.2 ml of a 25 mg/ml Al(OH)3 adjuvant suspension in sterile saline mixed with the 0.5 ml of 2S albumin or 2S albumin-R solution. The animals were bled on day 28. The sera were analyzed for anti-native- and 2S albumin-R-specific IgG1 (Thelper-1 mediated), IgG2a and IgE (Thelper-2 mediated) titers by ELISA and by Passive Cutaneous Anaphylaxis (PCA) essentially as described previously [9]. Data were analysed using one-way ANOVA with Bonferroni’s post-hoc test using GraphPad Prism version 5.00 (GraphPad Software, San Diego, CA, USA). The welfare of the animals was maintained in accordance with the general principles governing the use of animals in experiments of the European Communities (Directive 86/209/EEC) and Dutch legislation (The experiments on Animals Act, 1997), which includes approval of the study by TNO’s ethical review committee under DEC number 1732.

**Results and discussion**

From the animals orally exposed to native 2S albumin the majority developed 2S albumin-specific IgG1 and IgG2a responses at days 28 and 42 in a dose-dependent way whereas oral exposure to 2S albumin-R resulted in a lower number of responding animals (Figure 1A + B). At day 28, i.p. exposure of the animals to native 2S albumin resulted in the development of very pronounced 2S albumin-specific IgG1 and IgG2a responses in all animals (Figure 1A). I.p. exposure to 2S albumin-R resulted in 2S albumin-R-specific IgG1 and IgG2a responses in 80% of the animals. Both oral (0.1 and 1 mg protein/rat/day) and i.p. exposure of the animals to 2S albumin-R did not result in the development of 2S albumin-R-specific IgE antibodies. In contrast, oral dosing of the
animals to either 0.1 mg or 1 mg native 2S albumin resulted in 2S albumin-specific IgE responses in 50% of the animals and i.p. exposure lead to 100% responders. These results were confirmed by PCA testing (Figure 2). These results show that reduction of disulfide bonds and concomittal loss of protein structure and an increased sensitivity for digestion of Brazil nut 2S albumin [8], decreases the prominent sensitizing potential of 2S albumin in the oral BN rat food allergy model. Not all animals developed specific antibody responses upon oral exposure to 2S albumin. This phenomenon is also observed with other food allergens like ovalbumin and cow’s milk using the described BN rat food allergy model [9]. While the allergenicity of 2S albumin-R was dramatically decreased, the immunogenicity of the protein still existed since specific IgG1 and IgG2a antibodies against 2S albumin were produced, although at a lower level compared to native 2S albumin.

Evidence for an important role of the structural conformation (the secondary and tertiary structure) of the protein with respect to food protein allergenicity also comes from animal studies. Prefeeding of an endopeptidase inhibitor (aprotinin) to mice results in an inhibition of oral tolerance induction by protein feeding [10] while feeding of protein antigen to mice is known to induce substantial systemic tolerance for specific antibody and cell mediated immune responses under normal circumstances [11]. Recently it was shown that by regulating the gastric pH the dose-dependency of food allergy induction was influenced [12]. The effect of anti-ulcer drugs that increase the gastric pH on the formation of IgE in humans was investigated previously [13]. It was found that pre-existing IgE was boosted to higher levels and that de novo IgE towards digestion-labile food proteins was induced. The implications of regulating the gastric pH on the risks for food allergic consumers were recently reviewed [14]. While the above mentioned studies investigated the effect of decreasing the susceptibility to proteolysis, we investigated the effect of increasing the susceptibility to proteolysis.

Disruption of the structural conformation of protein by alkylation/denaturation has previously been used to generate hypoallergenic variants of Ara h2 [15] and Pru p3 [16], respectively the major peanut and peach allergen. Although no hypoallergenic derivatives have received marketing authorization to date, they are suggested to be promising vaccine candidates for immunotherapy. The reduced IgE-antibody-binding capacity reduces the frequency of adverse reactions during allergen-specific immunotherapy, allowing the application of higher amounts of allergen with a reduced risk of serious adverse effects [17,18]. Moreover, T cell immunogenicity needs to be preserved to maintain the therapeutic potential [19,20]. In line with this, we here show that the reduced/alkylated Brazil nut 2S albumin showed a reduction in biologically active IgE (by PCA), reflecting a reduced allergenic potential, whereas the immunogenicity remained intact.

Interestingly, if the native 2S albumin and 2S albumin-R form were injected i.p. together with alum as an adjuvant, specific IgE antibodies were only observed in animals treated with the native 2S albumin. As upon i.p. exposure degradation by the digestive tract will be absent and moreover the tolerogenic mucosal sites are bypassed, it is suggested that other aspects may also influence the potential allergenicity of a protein. It is known that proteases present in dendritic cells (DC) generate peptides from foreign and self proteins for eventual capture and display to T cells, demonstrating that individual proteolytic enzymes may have a clear contribution to antigen processing [21]. In fact, it was recently shown that the immunogenicity of Bet v1, the major birch pollen allergen, could be influenced by changing the susceptibility towards digestion [22]. Similar observations were made for 2S albums from other allergenic plant foods when injected i.p. [16]. Native 2S albumin compared to 2S albumine-R is relatively stable to digestion [8], an altered degradation by individual proteolytic enzymes and subsequent presentation of both peptide fragments by antigen presenting cells may thus lead to an altered antibody response. This may explain the observed difference in response against 2S albumin and 2S albumin-R upon i.p. administration. In conclusion, the results of this study demonstrate the importance of the protein structure of Brazil nut 2S albumin in inducing food allergy. The observation that modified
Brazil nut 2S albumin is still immunogenic while its allergenicity is reduced opens possibilities for developing immunotherapy for Brazil nut allergy.

**Abbreviations**

2S albumin-R: Reduced and alkylated 2S albumin; BN: Brown Norwegian; DC: Dendritic cells; i.p: Intraperitoneally; PCA: Passive cutaneous anaphylaxis.

**Competing interests**

The authors report no conflicts of interest. SJK is consultant to DBV Technologies, a company involved in developing epicutaneous immunotherapy for peanut allergy.

**Authors’ contributions**

JHMB carried out data analysis. LMJK carried out animal data acquisition and analysis. WPN and HHU prepared the proteins. AHP participated in the coordination of the animal studies. SJK was responsible for overall design of the studies and the qualification of the proteins. All authors were involved in the preparation of the manuscript and gave their final approval of the submitted version.

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**References**

1. Rona RJ, Keil T, Summers C, Gislon D, Zuidmeer L, Sodergren E, Sigurdardottir ST, Lindner T, Goldhahn J, Dahlstrom J, McBride D, Madsen C. The prevalence of food allergy: a meta-analysis. *J Allergy Clin Immunol* 2007, 120(3):38-46.

2. Pastorella EA, Faralli L, Pravettoni V, Ispano M, Conti A, Ansaloni R, Rotondo F, Incorvaia C, Bentgson A, Rivolta F, Tramaboldi C, Previd M, Ortolani C. Sensitization to the major allergen of Brazil nut is correlated with the clinical expression of allergy. *J Allergy Clin Immunol* 1998, 102(6 Pt 1):1021-1027.

3. Nordbre JA, Taylor SL, Townsend JA, Thomas LA, Bush RK. Identification of a Brazil-nut allergen in transgenic soybeans. *N Engl J Med* 1996, 334(11):688-692.

4. Astwood JD, Leach JN, Fuchs RL. Stability of food allergens to digestion in vitro. *Nat Biotechnol* 1996, 14(10):1269-1273.

5. Moreno FJ. Gastrointestinal digestion of food allergens: effect on their allergenicity. *Biomed Pharmacother* 2007, 61(1):50-60.

6. Moreno FJ, Melion FA, Wickham MS, Bottrill AR, Mills EN. Stability of the major allergen Brazil nut 25 albumin (Ber e 1) to physiologically relevant in vitro gastrointestinal digestion. *FEBS J* 2005, 272(2):341-352.

7. Sen M, Kopper R, Pons L, Abraham EC, Burks AW, Bannon GA. Protein structure plays a critical role in peanut allergen stability and may determine immunodominant IgE-binding epitopes. *J Immunol* 2002, 169(2):882-887.

8. Koppelman SJ, Nieuwenhuizen WF, Gaspari M, Knipps LM, Penninks AH, Krol EF, Heffle SL, de Jongh HH. Reversible denaturation of Brazil nut 25 albumin (Ber e1) and implication of structural destabilization on digestion by pepsin. *J Agric Food Chem* 2005, 53(1):123-131.

9. Knipps LM, Penninks AH, Spanhaak S, Houben GF. Oral sensitization to food proteins: a brown Norway rat model.* Clin Exp Allergy* 1998, 28(3):368-375.

10. Hanson DG, Roy MJ, Green GM, Miller SD. Inhibition of orally-induced immune tolerance in mice by prefeeding an endopeptidase inhibitor. *Reg Immunol* 1993, 5(2):76-84.

11. Strobel S, Mowat AW. Immun responses to dietary antigens: oral tolerance. *Immunol Today* 1998, 19(4):173-181.

12. Diesner SC, Knittelfelder R, Krishnamurthy D, Pali-Scholl I, Gajdzik L, Jensen-Jarolim E, Untersmayr E. Dose-dependent food allergy induction against ovalbumin under acid-suppression: a murine food allergy model. *Immunol Lett* 2008, 121(1):45-51.

13. Untersmayr E, Bakos N, Scholl I, Kundi M, Roth-Walter F, Szalai K, Riemer AB, Ankersmit HJ, Scheiner O, Boltz-Nitulescu G, Jensen-Jarolim E. Anti-ulcer drugs promote IgE formation toward dietary antigens in adult patients. *FASEB J* 2005, 19(6):656-658.

14. Untersmayr E, Jensen-Jarolim E. The role of protein digestibility and ant- acids on food allergy outcomes. *J Allergy Clin Immunol* 2008, 121(6):1301-1308.

15. Starkl P, Felix F, Krishnamurthy D, Streminitzer C, Roth-Walter F, Prickett SR, Vockamp AL, Willendraper A, Szalai K, Weichselbaumer M, O’Hehir RE, Jensen-Jarolim E. An unfolded variant of the major peanut allergen Ara h 2 with decreased anaphylactic potential. *Clin Exp Allergy* 2012, 42(12):1801-1812.

16. Toda M, Reese G, Gadenmaier G, Schulten V, Lauer I, Egger M, Briza P, Randow S, Wolfheimer S, Kigongo V, Del Mar San Miguel Moncin M, Fatsch K, Bohle B, Vietts S, Scheurer S. Protein unfolding strongly modulates the allergenicity and immunogenicity of Pru p 3, the major peach allergen. *J Allergy Clin Immunol* 2011, 128(5):1022–1030. e1-7.

17. Linhart B, Varlet A. Molecular design of allergy vaccines. *Curr Opin Immunol* 2005, 17(6):646-655.

18. Vrtala S. From allergen genes to new forms of allergy diagnosis and treatment. *Allergy* 2008, 63(3):299-309.

19. Akdis CA, Akdis M. Mechanisms and treatment of allergic disease in the big picture of regulatory T cells. *J Allergy Clin Immunol* 2009, 123(4):735–746. quiz 747-8.

20. Larche M. Peptide immunotherapy for allergic diseases. *Allergy* 2007, 62(2):325–331.

21. Watts C, Moss CX, Mazzee D, West MA, Matthews SP, Li DN, Manoury B. Creation versus destruction of T cell epitopes in the class II MHC pathway. *Ann N Y Acad Sci* 2003, 987:9–14.

22. Egger M, Jurets A, Wallner M, Briza P, Ruzek S, Haizl S, Pichler U, Kitzmuller C, Bohle B, Huber CG, Ferreira F. Assessing protein immunogenicity with a dendritic cell line-derived endosomosomal degradation. *PloS One* 2011, 6(2):e17278.