A Novel and Simple Method of Insolubilization of Ovomucoid in Cookies Prepared from Batter Containing Egg White

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Heat stable ovomucoid (OM) is a major allergenic protein of chicken eggs, which are widely used in wheat flour processed food. PBS soluble OM remained in the cookies prepared by the usual method. A procedure for insolubilizing OM in batter was designed for cookies supplemented with egg white. Immunoblotting and ELISA competitive inhibition using rabbit anti-OM serum were used to analyze soluble OM in the cookies. PBS soluble OM remained both in the batter and the dough as observed in control egg white solution. Soluble OM in the cookies made from batter baked at 180˚C for 8 min decreased to about one-tenth of the batter, while it decreased to almost nothing in bread made from dough baked at 180˚C for 8 min. When the batter was mixed for 15 min before baking at 180˚C for 12 min, PBS soluble OM disappeared in the cookies as in bread. OM in the cookies was soluble with SDS+2-ME solution, but with neither the PBS nor SDS solution. This suggests that a disulfide exchange reaction between OM and wheat components is induced in the cookies. Both mixing and baking time of batter played an important role in the elimination of PBS soluble OM in the cookies. This simple method is useful for insolubilization of OM in processed foods made from batter supplemented with egg white.

Keywords: insolubilization, ovomucoid, egg white, wheat flour, batter, dough

Allergens in foods are mostly proteins, and major allergenic proteins have been isolated from milk, egg, soybean, wheat, etc. Many food allergens identified in water and salt soluble fractions are relatively low molecular weight proteins or glycoproteins and are often resistant to heat denaturation and proteolytic digestion (Taylor, 1980; Matsuda et al., 1982). Ovomucoid (OM) is a major allergenic protein in chicken egg white, (Langeland, 1982a, b; Hoffman, 1983) and is even soluble and antigenic in shelled eggs boiled for 1 h (Gu et al., 1986), indicating that OM is not rendered insoluble by heating in the presence of other egg white proteins.

Chicken eggs are widely used in processed foods, especially wheat flour foods such as bread and cookies, but have been little examined for their soluble antigenic proteins in these foods. We reported a remarkable decrease in soluble antigenic OM in bread supplemented with egg white during the process of breadmaking (Kato et al., 1997). The effect of wheat gluten on the heat-induced insolubilization of OM was compared with that of soybean protein and milk casein. It was found that OM was easily rendered insoluble by heating in the presence of gluten but not of casein (Kato et al., 2000). This result suggested that wheat gluten might lead to the disappearance of soluble OM through disulfide exchange in the process of baking.

In this report, cookies made from a batter of soft wheat flour, which includes egg white, are used as a model for wheat flour processed food. Our interest is that OM is insolubilized in the process of creating these cookies through the interaction between OM and gluten. Consequently, the presence of PBS soluble OM was compared in cookies and bread baked from batter or dough of soft flour supplemented with egg white, respectively.

Materials and Methods

Preparation of cookies or bread, and their soluble protein fractions Soft wheat flour used for cookies or bread was a product of Nisshin Mills Co., Ltd. (Tokyo). The other ingredients were sugar, butter and salt.

For cookie making, fresh egg white (110 g, 28% to total ingredients) and soft flour (120 g) were added to a mixture of butter (90 g) and sugar (70 g) and were thoroughly mixed. The batter sample was divided into small and flat pieces (7 g) and baked at various temperatures and times in an oven range.

For bread making, soft flour, fresh egg white (25% of total ingredients) and other ingredients (instant dry yeast, sugar, skim milk powder, salt and water) were mixed at a ratio of 140 : 63 : 49, and kneaded for 35 min at 450 rpm using an electric kneader (Taisho Denki Co. Ltd., Kusatsu). The resulting dough was fermented for 150 min in a chamber (Yamato Scientific Co., Ltd., Model IG-43H.M., Tokyo) maintained at 30˚C and 75% RH (relative humidity), then punched and divided into small pieces (50 g), followed by a second fermentation at 38˚C and 85% RH until the loaf volume reached 2.5 times the original bulk. The fermented dough was baked at 180˚C for 8 min unless otherwise noted.

Each sample (0.364 g) was suspended in 1 ml of phosphate buffered saline of pH 7.4 (PBS), and homogenized with a vortex mixer for 1 min at the maximum speed (Vortex Genie Scientific Industries, INC, Bohemia, NY) and subsequently ultrasonicated for 15 min (100 W, 39 kHz). Each suspension was kept overnight at 4˚C and then centrifuged at 16,000×g for 20 min to remove the insoluble materials. The supernatant was subjected to analy-
proteins. Proteins were extracted with 4% sodium dodecyl sulfate (SDS), and 4% SDS containing 10% 2-mercaptoethanol (SDS + 2-ME) from cookies baked at 180°C for 12 min (0.346 g). These SDS or SDS + 2-ME extracts were heated for 5 min in boiled water, then directly used for the SDS-polyacrylamide gel electrophoresis (SDS-PAGE) and immunoblotting analyses.

**SDS-PAGE and immunoblotting** Proteins were first separated by SDS-PAGE in 12.5% acrylamide gel according to the method of Laemmli (1970). One sheet of the gel was stained with Coomassie Brilliant Blue (CBB) R-250, and a second gel was used for immunoblotting. The proteins separated by SDS-PAGE were transferred electrophoretically onto a nitrocellulose sheet (0.45 mm, Advantec Toyo, Tokyo) by the method of Towbin et al. (1979). The nitrocellulose sheet was incubated overnight at 4°C with 3% bovine serum albumin (BSA) in tris buffered saline of pH 7.5 containing 0.02% Tween-20 (TBST). After being washed with TBST, the sheet was incubated at 37°C for 2 h in a mixture of 5 ml of 1% BSA in TBST and 5 µl of a rabbit anti-OM serum. It was then incubated at 37°C for 1 h with a 10⁻⁴ fold diluted peroxidase-conjugated goat anti-rabbit IgG antibody (Cappel Laboratories, Aurora, OH) with 1% BSA/TBST. The protein bands reactive to the specific antiserum were thereafter activity-stained for peroxidase using 4-chloro-1-naphthol (Bio Rad Laboratories, Hercules, CA).

**Enzyme-linked immunosorbent assay (ELISA) competitive inhibition** The antigenic activity of OM in samples was evaluated by the competitive inhibition analyses of ELISA (Engvall & Perlman, 1971; Matsuda et al. 1983) using rabbit anti-OM serum. One hundred microliters of OM solution (1 µg/ml) was added to each well of flat-bottomed microtiter plates, and incubated for 3 h at 37°C, then washed with PBS-T (phosphate buffer saline containing 0.05% of Tween 20). One percent of BSA/PBS-T solution was added to each well and incubated for 30 min at 37°C. Fifteen microliters of competitors, PBS extract from sample, was gradually diluted with 135 µl of rabbit anti-OM serum in 1% of BSA/PBS-T solution (10⁻⁴), applied to the wells after washing with PBS-T twice, and incubated for 3 h at 37°C. Each well was washed with PBS-T four times. One hundred microliters of peroxidase conjugated goat anti-rabbit IgG (DAKO Co., Ltd., Glostrup, Denmark) in 1% of BSA/PBS-T solution (10⁻⁴) was added to each well and incubated for 1 h at 37°C. After washing each well with PBS-T three times, 100 µl of o-phenylenediamine in citrate-phosphate buffer (0.4 mg/ml) containing 0.03% of H₂O₂ was added to each well and the reaction was stopped with of 50 µl of H₂SO₄ solution (2.5 mol/l) after 20 min at RT. The color development of the well was photometrically measured at 492 nm.

**Results and Discussion**

Pieces of batter of similar size were baked at 100°C or 120°C for 2 to 30 min and at 150°C or 180°C for 2 to 12 min. The OM in soluble fractions from cookie samples was examined by immunoblotting using rabbit anti-OM antibody, PBS-extractable OM in cookies baked below 150°C was clearly detected on the blotting sheet (data not shown). The extraction efficiencies of soluble OM from cookies baked at 180°C for 2 to 12 min were compared by ELISA competitive inhibition analysis (Fig.1-A),
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and expressed as the amounts required to attain 50% inhibition of the reaction between OM and anti OM serum (Fig.1-B). The soluble OM concentrations in cookie samples baked for 2 min and 6 min were almost the same as in the batter, which faithfully reflected the amount of OM occurring in egg white. The amount of soluble OM in the cookies was reduced to below one-tenth by baking at 180˚C for 8 min or more.

Bread was baked from dough so as to make comparisons between cookie and bread. The PBS soluble fractions were prepared by buffer extraction from both dough and bread baked at 180˚C for 8 min. The soluble OM in bread was about 1/10000 as much as that in dough, indicating that soluble OM almost disappeared in the bread (Fig. 2). These results with reference to our previous report show that disulfide exchange occurred between OM and wheat compounds (Kato et al., 1997). The disappearance of OM in bread may be attributed to the dough formed by kneading.

In cookie making, the watery batter was fully mixed instead of kneading as an additional treatment before baking. This resulted in a disulfide exchange reaction between wheat protein and OM, leading to the loss of soluble OM. The effects of mixing for from zero to 15 min using a hand mixer (650 rpm/min) were examined in detail before baking at 180˚C for 8 min. Zero time of mixing was defined as the point at which all materials were well mixed. The soluble OM in baked cookies was analyzed by ELISA competitive inhibition using anti OM antibody (Fig. 3).

Soluble OM in cookie samples decreased along with mixing time. Moreover, the content of soluble OM in the cookies mixed for 15 min was greater about 400 times than that in batter to attain 50% inhibition of the reaction between OM and anti OM serum. The amount of soluble OM in the 15 min-mixed cookies

Fig. 3. Effect of mixing time on antigenic OM in the PBS extracts of cookies analyzed with ELISA competitive inhibition. Batter was mixed for zero (●), 1 (○), 5 (△), 10 (●), or 15 min (□) using a hand mixer, followed by baking at 180˚C for 8 min (A). The $I_{50}$ value of each sample was compared to the value of batter (=0) (B).

Fig. 4. ELISA competitive inhibition for antigenic OM in the PBS extracts of cookies baked for different times. Batter (●) after mixing for 15 min was divided into pieces, which were baked at 180˚C for 2 (○), 6 (△), 8 (●), or 12 min (□) (A). The $I_{50}$ value of each sample was compared to the value of batter (=0) (B). The assay procedures for soluble OM are the same as shown in the legend of Fig. 1.

Fig. 5. Separation by SDS-PAGE (A) and immunoblotting (B) of proteins extracted with PBS, SDS or SDS+2-ME from baked cookies. Proteins were extracted from cookies baked at 180˚C for 12 min after 15-min mixing using PBS, SDS and SDS+2-ME solutions. Each extract (10 μl) or a 10 μl solution of authentic OM (1 μg/μl) was subjected to SDS-PAGE as usual. The gel sheet and its blotted membrane were stained with CBB and a rabbit anti-OM serum, respectively. SDS-PAGE was carried out using a 12.5% separating gel and a 4% stacking gel.
corresponded to 1/40 of that in the cookies prepared without mixing.

The relationship between baking time at 180˚C and PBS soluble OM remaining in the cookies baked after 15-min mixing of the batter is shown in Fig. 4. The soluble OM decreased with prolonged baking time, although remaining appreciably in cookies baked from the usual batter without mixing. Meanwhile, the soluble OM in the cookies disappeared when the batter was mixed beforehand for 15 min instead of kneading in breadmaking, and then baked at 180˚C for 12 min.

Soluble proteins extractable with PBS, SDS or SDS+2-ME from the cookies baked at 180˚C for 12 min were analyzed by means of SDS-PAGE and immunoblotting (Fig. 5). Several proteins capable of being stained with CBB were found in the SDS extract, but not in the PBS extract, and also proteins with higher molecular weights were also observed in the SDS+2-ME extract. OM occurring in these extracts was identified by immunoblotting using anti OM serum. OM was undetectable in both PBS- and SDS-extracts as shown in Fig. 5. On the other hand, multiple protein bands with higher and lower molecular weights present in the SDS+2-ME-extract were positive to immunoblotting. Higher molecular bands might be the polymers interacting with wheat proteins, and the immunoblotted lower molecular band was presumed to be a degradation product of the complex of wheat protein and OM by mixing and heat-treatment. This result can be explained by disulfide exchange reactions between wheat gluten and OM leading to insolubilization of an irreversible denaturation of antigenic OM as described previously (Kato et al., 1997; 2000).

Watery batter does not change into dough by mixing for 15 min, although the time is relatively long. Processed wheat flour foods are produced from either dough or batter. The treatment of additional mixing and slightly longer baking is an alternative procedure for making cookies; in this case, however, the taste of the cookies is still almost the same as in the usual way. This procedure can be applied to processed wheat flour foods such as doughnuts and fritters made from the batter with supplemental egg. A study to estimate the metabolic modulation of insoluble OM in such processed foods as cookies is in progress.

Urisu et al. (1997) recently reported that oral challenge of egg-sensitive patients to thermo-coagulated egg white did not induce any allergic response in 21(55%) of 38 persons who were hypersensitive to raw egg white and that egg white coagulation deprived of soluble protein is much more hypoallergenic than heat-denaturated egg white. These results suggest that heated and insolubilized egg white allergens are clinically hypoallergenic compared with the native and soluble ones. It is very interesting to evaluate the allergenicity of the cookies containing insolubilized OM by clinical tests such as skin prick and oral challenge. If the insolubilized OM has little or no clinical allergenicity, the insolubilization procedure proposed in the present study surely serves as a convenient and useful means of converting hyperallergenic foods to hypoallergenic ones.

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References


