Effects of Adding Fat on Rheological Properties of Fish Meat Gel

Yoshinori Mochizuki,*1 Takahide Saito,*2 Naomichi Iso,*2 Haruo Mizuno,*2 Akira Aochi,*2 and Masato Noda*2

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The changes of the rheological properties of fish meat gel which contained various quantities of fat were examined on the bases of the results of differential scanning calorimetry (DSC). Samples were made from the frozen surimi of Alaska pollack and lard. The thermal transition of surimi, i.e., gelation, was irreversible, while that of lard was reversible. The thermal transitions of surimi and lard seemed to proceed independently in the mixture. The instantaneous elastic modulus $E_0$ of the mixtures decreased with the fat content above the melting point of lard (ca. 29°C). The decrease corresponded to the increase of sensory “softness”. The number of chains per unit volume, $\nu$-value, was given by the temperature dependence of $E_0$ of samples according to the theory of rubber elasticity. The $\nu$-values were not changed above 29°C, but increased with the increase of the fat content at the measuring temperature below 29°C. The increase of $\nu$-value below 29°C may come from the fact that the crystals of fat behaved like crosslinks. On the other hand, the $\nu$-values did not change above 29°C, since the fat melted.

In the processing “kamaboko” which is a kind of fish meat gel, ingredients and a method of heating were important factors. Many studies have been published on the rheological properties of fish meat, and also fish meat gel (kamaboko). In addition, many reports on the thermal analysis using differential scanning calorimetry (DSC) of fish meat, meat proteins, and kamaboko have been published. However, only a few studies on both rheological and thermal properties of surimi was reported.

In this work, we examined rheological properties of fish meat gel having various content of fat, by the thermal analysis and the stress-strain experiment.

Materials and Methods

Materials

Samples were made from the frozen surimi of Alaska pollack (an extra fine quality) and lard, both of which were purchased at a market. The frozen surimi was ground for 7 min with water (21%) and sodium chloride (2.5%). Then, the mixture was ground for 7 min with lard (0, 5, 10, 15, 20, 25, 30, 40%) and become paste. Thus, obtained the paste was inserted in a poly vinylidene chloride casing (55 mm × 220 mm) and heated for 1 h in hot water (85°C). The paste containing lard, (25, 30, 40%) could not form kamaboko gel by heating.

Differential Scanning Calorimetry

A portion (ca. 60 μg) of sample was transferred to an aluminium hermetic pan and weighed to within 5 μg. A High Sensitive Differential Scanning Calorimeter (Model SSC-560U, DAINI SEIKOSHA) was used to scan the sample from 10°C to 85°C at 1.0°C/min.

Measuring Compression Instantaneous Elastic Modulus ($E_0$)

This experiments were carried out by using a Tensipresser (Model TTP-50BX, TAKETOMO DENKI) at 0°C, 10°C, 20°C, 30°C, 40°C, 50°C, and 60°C. The plunger was a cylinder of 18 mm in diameter. A constant strain of 0.25 was selected. Five measurements were taken with the samples heated at the same temperature.

Analysis of $\nu$-values

According to the theory of rubber elasticity the

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*1 Toyoko Gakuen Women's Junior College, Todoroki, Setagaya, Tokyo 158, Japan (変月義範: 東京女子短期大学).  
*2 Department of Food Science and Technology, Tokyo University of Fisheries, Konan, Minato, Tokyo 108, Japan (壱藤隆英, 畜産, 水産, 食品, 東京水産大学).
stress-tension strain curve of ideal rubber in expressed as follows:

\[ f = vRT \left( \frac{L}{Lu} \right)^2 - \left( \frac{Lu}{L} \right) \]  

(1)

where \( R \) is the gas constant, \( v \) the number of chains per unit volume (mol/cm\(^3\)), and \( T \) the absolute temperature. The \( Lu \) is the undeformed sample length, and \( L \) is the deformed one. We have assumed that the theory of rubber elasticity can be applied to the compression-strain. On the assumption, eq. (1) must be replaced to the following equation,

\[ F = vRT \left( \frac{L}{Lu} \right)^2 \]  

(1')

Since the compression elastic modulus, \( E \), is defined as

\[ E = \left( \frac{\partial F}{\partial L} \right)_T \]  

(2)

the combination of eqs. (1') and (2) gives the eq. (3),

\[ E = vRT \left( \frac{L}{Lu} + 2 \left( \frac{Lu}{L} \right)^2 \right) \]  

(3)

In this study, the compression ratio is 0.25, i.e. \( L/Lu = 0.75 \), then we get following relation,

\[ E = -4.3vRT \]  

(4)

Results and Discussion

Fig. 1 shows the DSC thermograms of unheated samples. The thermogram of surimi only (0% fat content in Fig. 1) has two endothermic peaks. The peak in the lower temperature region, peak A, is mainly due to the thermal transition of myosin or actomyosin and the peak in the higher one, peak B, is due to that of actin, according to Saito et al.\(^7\) and Akahane et al.\(^8\) The thermogram of lard only (100% fat content in Fig. 1), has one major endothermic peak at temperature of 29°C, peak C, and some minor ones, peak D, E, F, and G. This thermogram is similar to the thermogram which was reported for lard by Imamura et al.\(^9\) and Cornily and Meste.\(^10\) The temperature of 29°C is corresponded with the melting point of lard. The thermograms of the mixture samples of surimi and lard were similar to the synthetic curve of the thermogram of surimi only and that of lard only.

The total enthalpy change, \( \Delta H_t \), was estimated by area between a thermogram and the baseline at temperature range from 10°C to 85°C. Table 1 shows the enthalpy changes of unheated samples. Different enthalpy changes \( \Delta H_o \), were the differences between the total enthalpy change of surimi without lard, and those of surimi with lard. The \( \Delta H_o \) for each of mixture samples was proportional to the fat content, 5%, 10%, 15%, and 20%, respectively. It is suggested that the thermograms of the mixture could be synthesized quantitatively, from those of surimi and lard.

Fig. 2 and Table 2 show the DSC thermograms and the total enthalpy changes of heated samples, respectively. The thermogram of surimi only (0% fat content), has found no peak, but the thermograms of surimi with lard and that of lard only have a major endothermic peak at temperature 29°C, peak H, and some minor peaks, peak I and J. Moreover, each thermogram of surimi with lard was the similar shape as that of lard only. It is suggested that the thermal transition of surimi was irreversible, while that of fat was reversible. The total enthalpy changes of the mixtures, \( \Delta H_t \), were proportional to the fat contents, as well as those of unheated samples. It may be said from the result that the difference of the rheological properties of heated samples related to the fat content because the thermal transition of lard was reversible.
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Table 2. Enthalpy changes of heated samples

<table>
<thead>
<tr>
<th>Fat content (%)</th>
<th>$\Delta H_t^*$ (cal/g)</th>
</tr>
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<tbody>
<tr>
<td>0</td>
<td>0.00</td>
</tr>
<tr>
<td>5</td>
<td>0.31</td>
</tr>
<tr>
<td>10</td>
<td>0.64</td>
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<tr>
<td>15</td>
<td>0.93</td>
</tr>
<tr>
<td>20</td>
<td>1.37</td>
</tr>
<tr>
<td>100</td>
<td>6.27</td>
</tr>
</tbody>
</table>

* $\Delta H_t$: Total enthalpy change.

Fig. 3. Dependence of the instantaneous elastic modulus ($E_0$) on the measuring temperature.

As shown in Fig. 3, the compression instantaneous elastic modulus, $E_0$, decreased with the increase of measuring temperature. In addition, the $E_0$ values decreased with increasing fat content above 29°C. The decrease of $E_0$ may be corresponded to the increase of "softness". The measuring temperature dependence of $E_0$ on the mixtures were different between above 29°C and below 29°C.

The $\eta$-values were shown as a function of fat content in Fig. 4. The increase of $\eta$-values below 29°C were dependent of the increase of the fat content. However, the $\eta$-values above 29°C were independent of the fat content. The temperature of 29°C was the melting point of lard. Thus, the $\eta$-values increased apparently with the fat content below the melting point. According to Sherman, an unworked fat consists of an interlocking network of fat crystals. Therefore, we assume that the arrangement of fat crystals have the same response as the crosslinks of proteins, when the stress is given to kamaboko with lard below 29°C. In other words, it may be said that the fat crystals in the mixtures behave like the crosslinks of actomyosin chains in gel. On the other hand, since the fat crystals in the mixtures melted, the $\eta$-values were not changed above the melting point.

Fig. 4. The relationship between the $\eta$-value and fat content.

References