STUDIES ON THE PHYSIOLOGICAL CHEMISTRY
OF THE SALIVARY GLANDS XXXVIII*
PROTEOLYTIC ACTIVITY OF HUMAN SALIVA

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Recently it was reported that submaxillary gland homogenate of rat degraded 
$^{131}$I-labeled Parotin, growth hormone, and casein with remarkable rapidity as 
compared with kidney, liver, spleen, pancreas, testis, muscle, and parotid gland 
homogenates. This rapid degradation of the labeled proteins by rat submaxillary 
homogenate was found to be proteolytic in nature, by quantitative paper-
chromatographic analysis (Kim, 1958).

Concerning to the proteolytic activity in human saliva, no study has been 
found except the early work by Voss (1931).

This paper deals with the occurrence and some characteristics of the protease 
in human saliva, using $^{131}$I-labeled casein as substrate.

MATERIALS AND METHODS

Preparation of $^{131}$I-labeled casein The techniques employed in trace labeling of casein (Kasein 
nach Hammarsten, E. Merk) were almost similar to those previously reported (Tanaka and Ito, 
1958). A hundred and fifty mg of casein were dissolved in 30 ml of M/10 phosphate-borate 
buffer, pH 8.8. Separately, 10 ml of 0.002 N KI labeled with ca. 10 mc of $^{131}$I were oxidized 
with nitrous acid (1.0 ml of 0.15 N NaNO$_2$ and 2.0 ml of 0.18 N H$_2$SO$_4$), the excess nitrous 
acid being destroyed with 1.0 ml of 1N ammonium sulfamate. This iodine solution was made 
slightly alkaline with 0.2 N NaOH and mixed with the above casein solution, and was kept 
at 26°-27°C for 2.5 hrs. After the addition of 1.5 ml of 0.02 N Na$_2$S$_2$O$_3$, this reaction mixture 
was dialized against ice-cold distilled water adjusted to pH 8 for a day, adding, at intervals, 
about 1 ml of 0.2 N KI as carrier into the inside phase. Then the labeled casein, contained 
in the inside fluid, was precipitated twice at the isoelectric point, and the dialysis against 
ice-cold distilled water was repeated once. After lyophilization of the solution, purified $^{131}$I-
labeled casein was obtained.

The purity of the labeled casein as expressed by the per cent of the total radioactivity 
found in trichloroacetic acid (TCA) precipitable fraction was 98-99%. About 30% of the 
initially used $^{131}$I was found to be combined with the casein.

Collection of saliva Human mixed saliva, collected directly into an ice-cold cylinder, was 
used without centrifugation. The midmorning saliva was used throughout all experiments.

Method of estimation of the proteolytic activity of human saliva $^{131}$I-labeled casein was dissolved 
in distilled water adjusted to pH 8 (200 μg/ml, 400 μg/ml). One half ml of this solution
was mixed with 0.5 ml of M/10 phosphate buffer, pH 7.2 and 1.0 ml of saliva in a small test tube, and aerobically incubated at 37°C for the designated periods. When the incubation was over, tubes were cooled in ice-cold water for several mins., and 0.5 ml of casein solution (TCA-carrier casein: usually 30 mg/ml) and 2.5 ml of 10% TCA were added to each tube. After centrifugation, the supernatant fluid was transferred to another test tube (specially prepared for the gamma-counting experiments, i.e. having three marks at 2, 4, and 6 cm from the bottom), neutralized with sodium hydroxide solution, and the radioactivity was assayed with a well-type scintillation counter in relation to the total I\(^{131}\)-casein added to each test tube.

Quantitative radio-paperchromatography Saliva (1/2 final conc.) and I\(^{131}\)-casein (500 \(\mu\)g/ml final conc.) were incubated at 37°C and pH 7.2 for 3 hrs. The incubation mixture was centrifuged at low temperature without the addition of TCA solution, and the supernatant fluid was spotted on a filter paper in 2 cm width and developed in 95% ethanol and 2 N ammonia (90:10) for about 12 hrs. After being dried, each paper was cut into 21 fragments along the Rf value (Fig. 1). Radioactivity in each fragment was assayed separately or in groups with a well-type scintillation counter, as per cent of the total radioactivity of 21 fragments.

RESULTS AND DISCUSSION

Upon incubation of I\(^{131}\)-labeled casein with human saliva at pH 7.2, there was a rapid rise in the radioactivity in the TCA supernatant fluid. This is demonstrated in Figure 2. The increased supernatant radioactivity was thought to

![Fig. 1. Quantitative radio-paperchromatography](image1)

Radioactivity of each fragment was assayed separately or in groups with a well-type scintillation counter avoiding the geometrical error.

![Fig. 2. Degradation of I\(^{131}\)-labeled casein by human saliva](image2)

Incubation of I\(^{131}\)-casein (100 \(\mu\)g/ml final conc.) and saliva (1/2 final conc.) at 37°C and pH 7.2 for the designated periods; TCA-carrier casein (6 mg/ml final conc.)
be either the result of proteolysis or of direct deiodination of the labeled casein. By the quantitative radio-paper chromatography, the former mechanism, i.e., proteolytic degradation, was demonstrated as shown in Figure 3. In the case of non-incubated $^{131}$-casein, almost all radioactivity was found around the original line (Rf 0.025–0.025), while iodide ion labeled with $^{131}$ appeared in the highest Rf area (Rf 0.875–1.025). In the case of $^{131}$-casein incubated with human saliva at $37^\circ$C and pH 7.2 for 3 hrs., approximately 70% of the total radioactivity was found around the original line and considerable amounts of radioactivity were observed in the higher Rf areas. But in the highest Rf area (Rf 0.875–1.025) where radioactive iodide

Fig. 3. Quantitative radio-paper chromatographic analysis of the mode of degradation of $^{131}$-casein by human saliva
Abscissas: Per cent radioactivity in each designated Rf area
a: $^{131}$-casein non-incubated (control)
b: $^{131}$-casein mixed at low temperature with similarly preincubated saliva (control)
c: Iodide ion labeled with $^{131}$
d: $^{131}$-casein incubated with saliva; incubation of $^{131}$-casein (500 μg/ml final concent.) and saliva 1/2 final concent.) at $37^\circ$C and pH 7.2 for 3 hrs.
pH

Fig. 4. Effect of pH on the proteolytic activity of human saliva
Incubation of I\(^{131}\)-casein (50 μg/ml final conc.) and saliva (1/2 final conc.) at 37°C and the designated pHs for 2 hrs.; TCA-carrier casein (6 mg/ml final conc.)

ion was developed, scarcely any radioactivity was found. Therefore, it was concluded that the degradation of the labeled casein by human saliva was not deiodinative but proteolytic in nature. In addition, these results of paperchromatographic study seemed to be in good agreement with the finding that about 23% of the total radioactivity was observed in the TCA supernatant fluid, following the simultaneous incubation of the labeled casein with saliva under the same conditions.

The effect of pH on the proteolytic activity of human saliva, as measured by the per cent radioactivity in TCA supernatant fluid, was determined at fourteen different pHs, ranging from 3.8 to 9.0 (Fig. 4). The activity was found maximal

Fig. 5. Effect of concentration of saliva on the degradation of I\(^{131}\)-casein
Incubation of I\(^{131}\)-casein (50 μg/ml final conc.) and saliva at 37°C and pH 7.2 for 1 hr. (a) or 3 hrs. (b); TCA-carrier casein (6 mg/ml final conc.)

Fig. 6. Effect of substrate concentration on the degradation of I\(^{131}\)-casein by human saliva
Incubation of I\(^{131}\)-casein (50 μg/ml final conc.), casein (0, 50, 100, 200, 400, and 800 μg/ml final conc.) and saliva (1/2 final conc.) at 37°C and pH 7.2 for 1 hr. (a) or 3 hrs. (b); TCA-carrier casein (6 mg/ml final conc.)
at pHs between 6 and 8, rapidly dropping on either side of the maximum. Consequently pH 7.2 was adopted for the usual assay of the proteolytic activity of human saliva.

That the degradation of $^{131}$-casein was related to the quantity of the enzyme present in the incubation mixture was revealed when the same amount of the labeled casein was incubated with variously diluted saliva at $37^\circ$C and pH 7.2 for 1 or 3 hrs. (Fig. 5).

The effect of substrate concentration on the degradation of $^{131}$-casein by saliva was studied by incubating the same amount of saliva plus labeled casein with different amounts of casein as shown in Figure 6. It was found that not only the degradation of the labeled casein as expressed by the per cent radioactivity in TCA supernatant, but $^{131}$-casein and casein were similarly degraded by saliva. And the latter also seemed to support the conclusion that the labeled casein was degraded proteolytically by human saliva.

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**Fig. 7.** Effect of preincubation of saliva on its proteolytic activity

Preincubation was carried out by incubating the saliva at $37^\circ$C (control: $0^\circ$C) and pH 7.2 for the designated periods. Proteolytic activity was measured by incubating the labeled casein (100 $\mu$g/ml final conc.) and the preincubated saliva (1/2 final conc.) at $37^\circ$C and pH 7.2 for 1 hr. Final conc. of TCA-carrier casein was 6 mg/ml.

**Fig. 8.** Thermal inactivation of proteolytic activity of human saliva

A: Untreated saliva (control)
B: Saliva heated in boiling water for 30 mins.

Proteolytic activity was measured by incubating the saliva (untreated or heat-treated, 1/2 final conc.) at $37^\circ$C and pH 7.2 for 1 hr. (hashed: incubated for 0 min.)
The effect of preincubation of saliva at 37°C and pH 7.2, on its proteolytic activity was expressed in Figure 7. Although a gradual decrease in the activity was observed, the protease in human saliva was found to be fairly stable at pH 7.2.

The proteolytic activity completely disappeared after heating the saliva in boiling water for 30 mins. (Fig. 8).

In assaying the protease activity of human saliva, as measured by the per cent radioactivity in TCA supernatant, considerably large amounts of TCA-carrier casein (i.e. final concent. of 6 mg per ml) were used throughout all experiments. When smaller amounts of TCA-carrier casein (e.g. final concent. of 1 mg per ml)
were used, remarkably large amount of radioactivity was observed in the TCA supernatant fluid of the nonincubated mixture of $^{131}$-casein (100 $\mu$g/ml final concent.), TCA-carrier casein (1 mg/ml final concent.), and saliva (1/2 final concent.). This TCA supernatant radioactivity was increased even further when boiled saliva was used in place of untreated one. This unusual increase of the radioactivity in the TCA supernatant (“anti-TCA activity” of saliva) should not be considered as the result of the degradation of the labeled casein by saliva, but another cause must be sought for. Figure 9 shows the effect of concentration of untreated and boiled saliva on their anti-TCA activity.

Not only the boiled saliva exhibited higher anti-TCA activity than the untreated one, but remarkably rapid dropping of the activity was observed as the saliva was diluted.

The anti-TCA principle in untreated or boiled saliva was not precipitated at pH 4 to 5, but was almost insoluble in 80% acetone (pH 5). From these results, together with the finding that the anti-TCA activity was not destroyed by boiling, salivary mucoid or its related substance was presumed to be responsible for the anti-TCA activity of saliva.

It was shown by Lineweaver and Murray (1947) that ovomucoid was resistant to precipitation by TCA. Jansen et al. (1952) found that cholinesterase prevented the precipitation of relatively small amounts of easily precipitable protein (e.g., $\beta$-lactoglobulin) by TCA, and that relatively large amounts of a carrier protein resulted in the complete precipitation of all the proteins including cholinesterase. The latter phenomenon was also reported by Michel and Krop (1951) using horse serum as carrier proteins.

Figure 10 illustrates the effect of various concentrations of TCA-carrier casein on the anti-TCA activity of untreated and boiled saliva. Remarkably rapid dropping of the anti-TCA activity was observed as the concentration of the carrier casein increased. These results have made it possible to measure the real proteolytic activity of saliva, distinguished from the anti-TCA activity, using the sufficient amounts of TCA-carrier casein. Throughout all experiments, proteolytic activity was measured using TCA-carrier casein.
casein in the final concentration of 6 mg per ml, and it was always confirmed, in every experiment, that the radioactivity in the TCA supernatant fluid of the non-incubated mixture did not increase significantly.

Figure 11 shows the proteolytic and anti-TCA activities of saliva of twelve healthy persons, indicating large variations from person to person. As expressed in Figure 12, considerably large variations were also observed from day to day.

**Figure 12. Variations in proteolytic and anti-TCA activities of saliva of four individuals over a four day period**

- - - : Proteolytic activity

- - - - - : Anti-TCA activity

Proteolytic and anti-TCA activities were measured by the methods described in the legend of Figure 11.

**SUMMARY**

Using I$^{131}$-labeled casein as substrate, the presence of proteolytic activity in human saliva was demonstrated, by measuring the increase of the radioactivity in the non-protein fraction of the incubation mixture and also by employing the
so-called quantitative radio-paperchromatography.

This proteolytic activity of saliva had its optimal pH between 6 and 8. It was fairly stable at pH 7.2 and 37°C, but was completely inactivated by heating the saliva in boiling water for 30 mins.

Besides the proteolytic activity, human saliva was found to possess the activity to prevent the precipitation of relatively small amounts of easily precipitable protein by TCA ("anti-TCA activity"), for which salivary mucoid or its related substanc was presumed to be responsible.

Considerably large variations in proteolytic and anti-TCA activities of saliva were observed from person to person as well as from day to day.

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