Electrophoretic and Immunochemical Demonstration of the Existence of Four Human Pepsinogens *

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In previous work from this laboratory, extracts of noncancerous human gastric mucosa were studied by immunochemical and electrophoretic means (1). Gastric mucosal extracts obtained by homogenization at pH 8 were subjected to electrophoresis in agar at pH 8.2. Zones of proteolytic activity at pH 2 were developed directly on the agar by the technique of Uriel (2). Four electrophoretically distinct zones showing proteolytic activity maximal at or near pH 2.0 were found in each of 45 stomachs studied. Thirty-six stomachs were from patients with duodenal ulcer, seven from patients with gastric ulcer, and in two cases, were normal stomachs obtained at autopsy from patients dying of heart disease less than 24 hours before autopsy. These constituents were designated proteases (P) I, II, III, and IV in order of decreasing electrophoretic mobility.

Immunoelectrophoretic studies of gastric mucosal extracts, using antiserum to mucosal extracts, showed three of the proteolytic constituents, P II, III, and IV, to be antigenic and to differ immunologically from one another (1). These antigens were found in all noncancerous stomachs studied. Preliminary studies of the noninvolved gastric mucosa of 10 patients with gastric carcinoma suggested that the proteolytic constituents were diminished or absent (3). It was thus of considerable interest to define more fully the nature of these gastric mucosal components.

The activation of pepsinogen involves the splitting off of basic peptides, resulting in an increased electrophoretic mobility for pepsin. This reaction is almost immediate at pH 2.0 and is much slower at pH values greater than 4.0 (4). Once activation has occurred, return to a pH greater than 6 results in denaturation of pepsin with loss of proteolytic activity (5). When tested under appropriate conditions, a strong immunologic cross-reaction between pepsin and pepsinogen can be shown (6).

The assumption was made that the proteolytically active components of gastric mucosal extracts (P I, II, III, and IV) were in fact proenzymes (pepsinogens) and that the proteolytic activity they manifested after electrophoresis resulted from activation in situ in the agar by the acidification attendant upon addition of the substrate at pH 1.9.

The present work demonstrates that: 1) The electrophoretic mobility of P I, II, III, and IV is rapidly increased by acidification at pH 2 and slowly or not at all changed at pH values of 4.4 or greater. 2) Alkalization following acidification results in loss of proteolytic activity of three of these constituents. 3) An immunologic relationship can be shown between the acidified and unacidified forms of the three antigenic proteolytic constituents. 4) After acidification at least five electrophoretically distinct enzymes are detectable. 5) One of these human pepsinogens (P II) is shown to be immunologically related to purified porcine pepsinogen, and another (P IV), after activation, displays several of the characteristics of pork pepsin B (7).

Methods

Extracts of gastric mucosa. Extraction of human gastric mucosa was carried out as follows: Stomachs were obtained at surgery from patients with gastric or duodenal ulcers, and in two cases, at autopsy from patients without either gastric or duodenal disease. The gastric mucosa was separated from the underlying mus-
icularis and connective tissue, cut into small fragments, and washed in 0.033 M phosphate buffer at pH 8.0. The wash solution was centrifuged at 6,000 rpm for 30 minutes, and the precipitate, designated "mucus," lyophilized. The mucosal fragments were homogenized in 0.033 M sodium phosphate buffer at pH 8.0 and centrifuged at 15,000 g for 30 minutes, and the supernatant fluid was lyophilized. This supernatant fluid, referred to as mucosal extract, was employed for these studies. The experiments described in this report were performed with mucosal extracts from patients with duodenal ulcer.

Preparation of antisera. Rabbits were immunized with either mucosal extracts or the "mucus" residue obtained from the wash solution. Immunization was carried out by the injection of these preparations in Freund's adjuvant into the foot pads. Five to eight mg protein in 1 ml was administered. Two weeks later the same preparation in alum was injected subcutaneously and then intravenously on the succeeding 2 days. Five to eight mg protein was given with each injection. Rabbits were test bled 5 days after the last injection and exsanguinated if antibody was present. In the absence of antibody formation, repeated courses of antigen in alum or in Freund's adjuvant were given. Antisera to each preparation were pooled. Antibodies to P II and III were found in antisera prepared to mucosal extracts. One rabbit produced antibodies to P II but not to P III, and this antisera was not pooled with the others. Antibodies to P IV were present in high concentration in antisera to "mucus" and in weak concentration in some antisera to mucosal extract. Reactivity with plasma proteins was removed by absorption with lyophilized normal human plasma.

Immunoelectrophoretic techniques. Immunoelectrophoretic analysis (IEA) was carried out by standard techniques (8) with only slight modification. The technique of Uriel (2) was employed to demonstrate proteolytic activity after electrophoresis in agar (see below). For analysis of chromatographic fractions, both IEA and electrophoresis before development of proteolytic activity were carried out in agar buffered at pH 8.2 with 0.05 M Veronal buffer. The same conditions were employed for IEA of absorption studies. All other electrophoretic studies and IEA were carried out at pH 5.6. Difco Noble agar was made up to 1.5% in 0.02 M citrate buffer, pH 5.6, and electrophoresis was carried out with the same buffer in the electrophoresis trays. Double diffusion in agar (Ouchterlony technique) was carried out with 1.5% agar in citrate buffer at pH 5.6, or in Veronal buffer at pH 8.2.

Characterization of proteolytic activity. To demonstrate proteolytic activity after electrophoresis, 24-× 18-cm glass plates were covered with a 2 mm-thickness of agar. Reservoirs were cut near the cathodic end of the plate and filled with 0.05 ml of a mixture containing equal parts of unbuffered 3% agar and a solution of whole gastric extract containing 5 mg of protein per ml. When purified products were studied, appropriately smaller quantities of protein were used. Protein concentration was estimated by the biuret procedure with human serum albumin as standard. Electrophoresis was carried out for 2 hours at a potential difference of 7 v per cm, the plate being refrigerated by trays of ice. After electrophoresis, the agar plate was immersed for 30 minutes in a solution of 0.2% human serum albumin in 0.2 M glycine-HCl buffer at pH 1.9. This solution was then decanted, and the agar plates (containing the imbibed substrate-buffer solution) were incubated in a moist chamber for 2 hours at 37° C. The agar was fixed overnight in a solution of 5% acetic acid in 50% alcohol, dried, and stained for protein with amido black. The imbibed albumin was stained by the dye, whereas sites of proteolytic activity in the agar remained unstained.

For IEA, the agar was dissolved in citrate buffer, pH 5.6, as described above, and electrophoresis was carried out for 1 to 1½ hours at 6 to 7 v per cm.

Anti-exchange chromatography. Chromatography was carried out at 4° C. The lyophilized gastric extract was dissolved in distilled water at a concentration of 20 to 40 mg protein per ml and dialyzed against 0.01 M KH2PO4 buffer, pH 7.0, for at least 4 hours. Ten to 15 ml was introduced on to a 20-× 2-cm DEAE-cellulose column equilibrated with the same buffer. This starting buffer was passed through the column until the absorption at 280 mμ returned to a stable low level. A gradient was then begun employing a modification of the technique of Fahey, McCoy, and Goulain (9) with 500 ml of starting buffer in the round mixing chamber and 250 ml of 0.4 M phosphate buffer, pH 5.65, in the conical reservoir. A flow rate of 20 ml per hour was utilized. After both flasks were empty, 0.4 M PO4 buffer, pH 5.65, was passed until the curve of absorption at 280 mμ returned to a stable level. NaCl, 0.8 M in 0.4 M phosphate buffer at pH 5.4, was then passed through the column. Fractions containing 10 ml of eluate were collected, and the curves of absorption at 280 and 260 mμ were determined to permit estimation of the relative amount of protein and nucleic acid present. Pools of two or three adjacent tubes were made, dialyzed against distilled water, and lyophilized. Samples were analyzed for the presence of gastric antigens by immunoelectrophoresis and for proteolytic activity at pH 1.9 by characterization after electrophoresis in agar as described above. Five different mucosal extracts were separated in separate chromatographic runs, and comparable results were found for each.

Acidification of proteases before electrophoresis. Whole gastric extracts and products separated by chromatography were acidified by the addition of an equal volume of 0.06 M HCl (with a resultant pH of 2.0) and incubated at 37° C for 10 minutes. The acidified extract was then mixed with an equal volume of melted unbuffered 3% agar and pipetted into the appropriate reservoir in the agar at pH 5.6. In experiments in which time of exposure to a given pH had to be limited, the 3% agar was made up in citrate buffer, pH 5.6.

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Determination of pH optimum of separated activated proenzymes. The method of Anson (10) was slightly modified. Separated and partially purified proteolytic constituents were dissolved in 0.9% NaCl. Five-tenths ml of 0.06 N HCl was added to each of a series of tubes containing 0.5 ml of the enzyme solution to be tested. After 10 minutes at 25°C, 2 ml of 1% bovine hemoglobin* dissolved in 0.2 M glycine-HCl or 0.2 M acetate buffer at the pH to be studied was added to the activated enzyme, and the mixture was incubated for 15 minutes at 25°C. The reaction was stopped and protein precipitated by the addition of 3 ml of 10% trichloroacetic acid. The contents of each tube were filtered, and the absorbation at 280 mμ was read against a blank composed of hemoglobin in the same buffer, to which the acidified enzyme had been added after the addition of trichloroacetic acid. A third tube containing substrate, buffer, and enzyme was not precipitated with trichloroacetic acid and was used to determine the pH of the substrate enzyme mixture.

Diffusion coefficient. The method of Allison and Humphrey was employed for the determination of the diffusion coefficient of P III (11).

Results

Chromatographic separation. Figure 1 shows the results of a chromatographic separation of gastric mucosal extract on DEAE-cellulose. The first proteolytic constituent eluted from the column was P IV. A nonproteolytic gastric antigen and serum albumin were found in the same peak. No further purification of P IV was attempted. P III was eluted shortly after a peak rich in 260 mμ-absorbing material and remained present throughout the rest of the chromatograph in gradually decreasing quantities. The subsequent addition of PO₄ buffer, 0.4 M, pH 5.65, resulted in a protein peak rich in P II, but still containing P III. Finally, passage of 0.8 M NaCl in 0.4 M PO₄ buffer at pH 5.4 eluted a peak rich in 260 mμ-absorbing material. This peak contained P I, previously not eluted from the column, as well as traces of P II and P III.

Passage of the P II rich eluate through Sephadex G 25, G 50, G 100, and G 200 failed to separate the P III present from P II (12).

Change in electrophoretic mobility and immunchemical properties after acidification. Three representative mucosal extracts were acidified before electrophoresis to determine whether the mobility of the proteolytic constituents would be increased by this treatment. Electrophoresis was then carried out in agar buffered at pH 5.6, since both pepsinogen and pepsin are stable at this pH (6). Results were the same for all stomachs studied. Representative findings are shown at the right of Figure 2. After acidification there was an in-

![Fig. 1. DEAE-cellulose chromatography of a representative gastric mucosal extract. From patient with duodenal ulcer. Absorptions at 280 mμ and 260 mμ are indicated by a solid line and dashed line, respectively. The distribution of proteolytic constituents is indicated by horizontal bars above the curve. Diminished concentration is indicated by diminished thickness of the bar.](http://www.jci.org/downloads/pdfs/105072/Figure1.jpg)
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FIG. 2. PROTEOLYTIC ACTIVITY AFTER ELECTROPHORESIS IN AGAR. MUCOSAL extracts from duodenal ulcer patients. The anode is at the top. The starting reservoir is visible at the bottom. The two strips at the right show whole mucosal extract before (0) and after (A) acidification. Four distinct proteolytic constituents are visible before acidification, the fourth being very weak. After acidification, five proteolytic spots of increased mobility are seen. Two of these are labeled 3. The pattern of proteolytic activity before and after acidification of the chromatographic fractions containing proteases (P) III, P II, and P I are also shown. P III gives rise to two proteolytic spots after acidification.

FIG. 3. IMMUNOELECTROPHORETIC PATTERN OF GASTRIC MUCOSAL EXTRACT BEFORE AND AFTER ACIDIFICATION. Duodenal ulcer patient. Migration in agar at pH 5.6 for 1 hour. Well 0, unacidified mucosal extract; well A, acidified mucosal extract; aM, antimucosal antiserum containing antibodies to P II and P III. Both P II and P III migrate more rapidly after acidification.

crease in mobility of the ensemble of proteolytic constituents compared to that seen before acidification. Control studies showed that neither human serum albumin nor ovalbumin increased their electrophoretic mobility after acidification.

Upon IEA with antiserum containing antibodies to P II and P III, the mucosal extract showed two precipitin lines before acidification, and two somewhat weaker lines of more rapid mobility after acidification (Figure 3). When tested by the Ouchterlony technique at pH 5.6, the precipitin lines formed against the acidified product were weaker than those against the original material. A very marked degree of cross-reactivity between the original and activated material was shown, with little if any spur formation (Figure 4). The same experiment performed in agar at pH 8.2 revealed no reaction with the acidified product. Absorption of this antiserum by acidified gastric extract removed activity against P II and P III (Figure 5a).

The effect of acidification of the proteolytic
constituents separated or partially separated by chromatography was studied. A fraction containing P IV as the sole proteolytically active constituent demonstrated increased electrophoretic mobility of the proteolytic spot after acidification. Upon immunoelectrophoresis the mobility of the precipitin line representing this antigen increased by a like amount.

Figure 2 shows a similar experiment performed using other chromatographically separated fractions. When a fraction containing P III but devoid of other proteolytic activity and other gastric antigens was tested, two electrophoretically distinct

![Ouchterlony Plate](image)

**FIG. 4.** Ouchterlony plate showing immunologic relationship between acidified and unacidified gastric mucosal extract. Duodenal ulcer patient. Agar at pH 5.6. Well 0, unacidified extract; well A, acidified extract; aM, antiserum (anti-P II and P III); aM2, antiserum (anti-P II). The reaction of aM with well A shows two lines weaker than those seen with well 0. A sharp angle formed between the lines with little if any spur formation. At the concentration of extract employed, no other gastric antigens are detectable.

sites of proteolytic activity could be detected after acidification, both with a mobility more rapid than the original material. Upon immunoelectrophoresis with antiserum to P II and P III, the precipitin line formed by acidified P III was detected closer to the anode than the line formed against unacidified P III and showed a double curvature (Figure 6) consistent with the two proteolytic sites noted. This double curvature indicated that both of these enzymes are derived from a single precursor. Unacidified and acidified P III were placed in a single reservoir, and IEA was carried out. The precipitin line corresponding to the unacidified antigen fused with the more rapidly migrating line representing the activated enzyme to form a continuous line.

When a fraction containing primarily P II with a trace of P III was acidified, the mobility of the

![Immunelectrophoretic Pattern](image)

**FIG. 5.** Absorption of antiserum by activated and by inactivated extract. Duodenal ulcer patient. Immunoelectrophoretic analysis, migration in agar at pH 8.2 for 1½ hours. The well contains mucosal extract. a. Absorption of antiserum aM with acidified extract (aM-A) abolished detectable activity against the original extract. b. Absorption of antiserum with extract that has been inactivated by acidification and subsequent alkalization (aM-I). The precipitin lines are closer to the aM-I antibody reservoir and are shorter, indicating partial absorption of antibody.

**FIG. 6.** Immunelectrophoretic pattern of P III before and after acidification. Duodenal ulcer patient. Migration in agar at pH 5.6. Well 0, P III unacidified; well A, P III acidified; aM, antiserum (anti-P II, P III). The precipitin line formed against acidified P III presents a double curvature and is more anodally located than that formed with the unacidified P III.
proteolytic P II spot was increased after acidification (Figure 2). Immuno-electrophoresis showed that the precipitin line of acidified P II was weaker and more anodally located. The chromatographic peak containing P I, as well as small amounts of P II and P III, similarly showed that the mobility of P I increased after acidification (Figure 2).

All experiments described in this section were performed a total of three to five times on three different mucosal preparations from patients with duodenal ulcer. Findings were identical in each case.

**pH requirement and speed of activation.** A single gastric extract was acidified at pH 2 for periods of 1, 3, 5, and 10 minutes. At the end of these intervals the extract was brought to pH 5.6 by mixture with an equal volume of 3% agar in citrate buffer and placed in the reservoir. The results shown in the left half of Figure 7 indicate that by 1 minute activation was almost complete and that by 3 minutes it was completed.

The effect of incubating the same extract at pH 2.0, 3.0, 4.4, 5.4, and 6.2 for 10 minutes was studied. Activation was completed within this period of time at pH 2 and 3, but only minimally at pH 4.4 and not at all at higher pH's (Figure 7, right).

**Alkaline inactivation after acidification.** Figure 8 shows that alkalinization of acidified gastric extract destroyed proteolytic activity. An equal quantity of gastric extract was placed in each of four tubes. The first and second tubes were diluted with normal saline, whereas the third and fourth were acidified for 10 minutes. The pH of the second and fourth tubes was then brought to pH 8.6 by the dropwise addition of 0.5 M NaOH and 0.1 M NaOH, whereas the first and third tubes were diluted with a comparable amount of normal saline. After 15 minutes, the contents of each tube were pipetted into reservoirs in the agar, and electrophoresis was performed. Alkalinization following acidification essentially eliminated the proteolytic activity of P I, P II, and P III. Identical findings were noted in each of three such experiments employing different mucosal extracts. Absorption of antiserum to P II and P III by this realkalinized extract in each case partially reduced
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3. Change of Electrophoretic Mobility of Pork Pepsinogen and Pepsinogen C After Acidification. Both the range of mobility and the change of mobility upon acidification seen with these porcine pepsinogens are of the same order as is observed with human mucosal extracts. Pepsinogen C, in this case, is contaminated with pepsinogen. O = before and A = after acidification.

4. Absorption of Antiserum with Pork Pepsinogen. The well contains mucosal extract. Antiserum containing antibodies against P II and P III (aM) was absorbed with porcine pepsinogen (aM-P'gen). Upon immunoelectrophoresis at pH 8.2, the absorbed antiserum did not react with P II.

5. That their mobility increased in a similar manner after acidification.

6. A direct precipitin reaction was not demonstrated between porcine pepsinogen or pepsinogen C and antiserum containing antibodies to human P II and P III. Addition of pork pepsinogen to such antiserum, however, removed detectable reactivity against human P II (Figure 10) in each of two trials.

7. Diffusion coefficient. The diffusion coefficient of P III, determined 4 times on three different samples by the technique of Allison and Humphrey (11) was $7.7 \times 10^{-2}$ cm$^2$ per second, only slightly different from that reported for porcine pepsinogen by Van Vunakis, Lehrer, Allison, and Levine.

8. Both kindly supplied by Dr. A. P. Ryle, the University of Edinburgh Medical School, Edinburgh.
human gastric mucosa contains four distinct pro-
enzymes giving rise to five enzymes with pepsin-
like activity. These four constituents differed
from one another physicochemically, demonstrat-
ing differing electrophoretic mobilities and elution
patterns upon chromatography on DEAE-cellu-
lose. In addition, they differed from one another
immunochemically in that three of them demon-
strated unrelated precipitin arcs on immunoel-
trophoresis, whereas the fourth constituent (P I)
did not appear to be antigenic.

The evidence that these constituents represent
precursors of pepsin-like enzymes may be sum-
marized as follows: 1) Each constituent 
demonstrated proteolytic activity at or near pH 2.0, either
when tested in tubes by the Anson hemoglobin as-
say method or in agar after electrophoresis. 2) The 
components were extracted from gastric mu-
cosa under alkaline conditions. Pepsinogen is
stable under these conditions, whereas all but one
of the known pepsins are inactivated (see below)
(7, 15). 3) After acidification, the electrophoretic
mobility of each proteolytically active constituent
increased. This is consistent with the loss of basic
peptides known to occur upon the conversion of
pepsinogen to pepsin. This change in mobility
occurred under the same conditions of pH and as
rapidly as does the activation of pepsinogen to
pepsin (4). 4) Once activated, return to an alk-
line pH resulted in complete or almost complete
loss of proteolytic activity of all constituents ex-
cept P IV. 5) The range of electrophoretic mo-
bility and the change in mobility following activa-
tion are similar to those of porcine pepsinogen
and pepsinogen C.

The studies described in the present report,
demonstrating that the four proteolytic constitu-
ents of gastric mucosal extracts are proenzymes,
were performed with mucosal extracts of stomachs
from patients with duodenal ulcer. These studies
strongly suggest that the four proteolytic con-
stituents of identical mobility and immunochemi-
ical characteristics also found in normals and in
gastric ulcer patients similarly represent four
distinct pepsinogens.

Preliminary studies of human gastric juice per-
formed with Dr. H. Hirsch-Marie have revealed
proteolytic activity in at least three electropho-
etically separate sites comparable in mobility to
those reported here for the three more rapid pro-

Discussion

Previous studies performed in this laboratory
(1) have shown four electrophoretically distinct
proteolytic constituents active at pH 2 in extracts
of each of 45 noncancerous stomachs studied.
Extraction was carried out at pH 8.0 and electo-
phoresis at pH 8.2, ruling out the possibility that
these four components represented modifications
of a single protein due to acid activation.

In the present work evidence is presented that

![Figure 11. pH Optima of Activated P II and P III. Duodenal ulcer patient. See text for details. P III is at the right. A fraction containing primarily P II as well as a small amount of P III is at the left.](http://www.jci.org)
enzymes after activation. In addition, antiserum to P II and P III reacted with gastric juice on immunoelectrophoresis to give two lines with a mobility and appearance identical to those of activated P II and P III.

In the present work it has been shown that antisera to the three antigenic human proenzymes, P II, III, and IV, are reactive with the activated more rapidly migrating enzymes. After activation each antigen was shown by IEA to migrate more rapidly. Absorption of antiserum to P II and P III by activated mucosal extracts abolished activity against these constituents, whereas absorption by activated and subsequently alkalinized extracts only partially abolished reactivity to these antigens.

Studies employing the Ouchterlony technique revealed a high degree of cross-reactivity between P II and activated P II and between P III and activated P III. Fusion of the lines representing unactivated and activated P II upon immunoelectrophoresis also indicated a close immunologic relationship between these components. No reaction between antiserum to P II and P III and acidified extract could be detected at pH 8.2, consistent with denaturation of these acidified constituents at this pH. In studies of the immunochemical behavior of the pepsinogen-pancreatin system, Van Vunakis and his colleagues (14), and Schlamowitz, Varandani, and Wissler (6) found porcine pepsinogen to be weakly antigenic. Cross-reactivity between pepsinogen and pepsin was demonstrated by both of these groups at pH 5.5 with greatly diminished cross-reactivity at pH 8. Thus, the immunochemical findings described in the present report are in agreement with the known immunochemical behavior of the pork pepsinogen system and lend additional support to the thesis that each of these constituents is indeed a distinct molecular species of human pepsinogen.

Multiple porcine pepsins were demonstrated by Ryle and Porter (7) who separated para-pepsin I and para-pepsin II (since designated pepsin B and pepsin C) from pepsinogen. Pepsin B produces almost no degradation of protein substrates, although it hydrolyzes the synthetic substrate acetyl-DL-phenylalanyl L-diodotyrosine (APD). Its activity is unaffected by alkalinization. Pepsin C digests bovine hemoglobin and bovine plasma albumin, but not APD, and is inactivated by alkalinization. Pepsin itself digests both APD and the protein substrates and is known to be quite unstable in alkaline solution.

In subsequent work Ryle (13) reported the isolation of pepsinogen C, the zymogen of pepsin C, from extracts of porcine gastric mucosa and demonstrated that activation of the proenzyme was extremely rapid at pH 2.0 and much slower at pH values greater than 4.0. Recently, Lee and Ryle (16) have succeeded in isolating a fourth zymogen, pepsinogen D, from porcine gastric mucosa.

Tang, Wolf, Caputto, and Trucco (17) demonstrated a proteolytic enzyme, gastricsin, in human gastric juice with optimal activity at pH 3.0 to 3.2, whereas the pH optimum of human pepsin was 2.0. Tang and Tang (15) subsequently reported the purification from human gastric mucosa of zymogen IA which, upon acidification, was converted to two proteolytically active constituents demonstrating the properties of pepsin and gastricsin, respectively. Both of these latter enzymes were inactivated by alkalinization.

It is difficult to relate human pepsinogen I described in this report to known components of the porcine or human pepsinogen systems. It could not be shown to be antigenic under the experimental conditions employed. In view of the pH at which extraction was carried out it seems unlikely that P I represents a proteolytically active product of autodigestion of the other proenzymes or enzymes. The possibility that this component was split from another pepsinogen during lyophilization cannot be excluded.

Human pepsinogen II, on the other hand, is clearly immunologically related to Ryle’s pork pepsinogen as shown by absorption studies. The pH maximum at pH 2.0 of activated P II was not followed by a clear second maximum at pH 3, even after activation for 1 hour. However, the broad pH range over which P II is active may suggest that it corresponds to human zymogen IA (15), which has been shown to give rise to pepsin and gastricsin with pH maxima at 2 and 3, respectively. The possibility that this proenzyme actually gives rise to two electrophoretically different enzymes after activation cannot be ruled out, since the sample studied contained a trace of P III, which might mask a second proteolytic spot.

Human pepsinogen III apparently gives rise to two enzymes upon activation, as demonstrated by
electrophoresis and confirmed by immunoelectrophoresis. However, the sharp pH optimum at pH 2.0 shown by P III after activation makes it unlikely that gastricsin could be derived from this proenzyme.

Because of its mucosal origin and accelerated mobility following acidification, P IV probably represents a proenzyme as well. However, the possibility that this constituent is the enzyme itself cannot be ruled out, since it is not inactivated by alkalinization and would not be denatured by the extraction procedure. P IV does not appear to be immunologically related to the other human pepsinogens, and it is thus unlikely that it represents an enzymatically active degradation product of the others or that it is a denatured pepsin. The pH optimum shown by P IV in agar establishes it as belonging to the pepsin group. P IV is stable upon alkalinization, is the most basic of the human pepsinogens, and demonstrates only weak activity against protein substrates, all properties analogous to those of swine pepsinogen B. These findings suggest that P IV may be related to this porcinezymogen.

The nomenclature we have employed for the human pepsinogens is based on the electrophoretic mobility of these components. It may lead to problems vis-à-vis the well studied pork pepsinogen-pepsin system, where nomenclature has been based on other criteria. Until more complete characterization of the human pepsinogens can be carried out, it seems wise to retain the nomenclature that we have employed.

The cellular origin of these pepsinogens is unknown. Germane to this problem are experiments performed in this laboratory (18) which demonstrated that the pyloric portion of the stomach contains P III and P IV, whereas all four pepsinogens are present in the body and fundus. The results are consistent with the findings reviewed by Taylor of peptic activity in mammalian pyloric mucosa (19).

Summary

Four electrophoretically distinct constituents present in alkaline extracts of human gastric mucosa display peptic activity. Three of these (P I, P II, and P III) have been shown to behave like pepsinogen by an augmentation of electrophoretic mobility following acidification, by the speed and pH requirements of the augmentation, by loss of proteolytic activity following realkalization, and, in the case of two of these constituents (P II and P III), by the demonstration of an immunologic relationship between the acidified and the original antigen. The fourth component (P IV) also migrated more rapidly after acidification, but was not inactivated by subsequent alkalinization. Antiserum to P IV reacted with acidified P IV upon immunoelectrophoretic analysis to give a precipitin reaction more anodally located than unacidified P IV.

These human pepsinogens could be separated from one another partially or completely by anion exchange chromatography. P II was shown to be immunologically related to purified pork pepsinogen. P III showed a diffusion coefficient approximately the same as that of pork pepsinogen and gave rise to two electrophoretically distinct proteolytic components upon activation. Activated P IV displayed several characteristics in common with pork pepsin B.6

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References


6 During revision of this manuscript, Seijffers, Segal, and Miller reported the separation of three pepsinogens (I, II, and III) and four pepsins from human gastric mucosa by chromatographic means (20, 21). These pepsinogens apparently correspond respectively to pepsinogens III, II, and I described in the present report.


