ISOLATION, PROPERTIES AND THE USE OF A CHYMOSIN-LIKE ENZYME FROM HARP SEAL (PAGOPHILUS GROENLANDICUS)

CENTRE FOR NEWFOUNDLAND STUDIES

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Isolation, properties and the use of a chymosin-like enzyme from harp scal (tagophilus groenlandicus)

A Theafs substited by

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in Partial Puffiment of
the requirements for
the degree of
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ABSTRACT

Four zymogens of acidic processes A, B, C, and C were isolated from the gastric mucosa of harp seal (Pagophtius groenlandicus) by ion exchange chromatography on a DEAR Sephadex A-50 column. Zymogens A and C were further purified by affinity chromatography using carbobengoxy-D-phenyialanine-triethylene-tatromine and gel filtration on a Sephadex G-100 Column. Certain physical and catalytic properties of processes A and C were compared with those of calf chymosin (E.C.3.4.23.4) and porcise pepsin (E.C.3.4.23.1).

Processe A was similar to calf chymosin with respect to several criteria. It had a higher ratio of milk clotting to proteolytic activity than those of seal processe C and porcine pepsin, clotted milk up to pH 7.0, and had a pH optimum of 2.2-3.5 for hemoglobin.

hydrolysis. It did not inactivate ribonuclease, had very low activity on APDT and lost activity in 6N ures. These results indicate that processe A is chymosin-Wike rather than pepsin-like:

Experimental Cinddar cheene made with processe A, cult remet, crude seal gastric protease (SGP) and Fromsac (Moor mideal protease) had comparable yields and proximate compositions. Sensory analytic of experimental cheeses after 30 weeks of aging showed that the product made with crude SGP had higher semsory agrees that those made with the other coagulants. Chemical analyses—showed more extensive degradation of protein in cheeses made with call rennet and Fromsac than in the cheeses made with the seal protesses. The electrophoretic patterns of the proteins from cheeses made with its seal protesses were qualitatively binilar to—those of calf rennet cheese. It was concluded that protesses A and seal gastric processes are promising as rennet substitutes.

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ABBREVIATIONS

APDT N-acetyl-L-phenylalanyl-L-diiodotyrosine

BIS N,N'-methylene-bis-acrylamide

DEAE-Sephadex Diethylaminoethyl-Sephadex

EDTA Ethylenediaminetetraacetic acid

RNA Ribonucleic acid

SDS Sodium dodecylsulphate
TCA Trichloroacetic acid

TEMED N,N,N',N'-tetramethylethylenediamine

Z-D-phe-T-Sepharose Carbobenzoxy-D-phenylalanyl-triethylene-tetramine-

Sepharose

INTRODUCTION

Preliminary studies indicated that a crude extract from the stomachs of harp seal (Pagophilus groenlandiscus) pups had properties similar to those of calf chymosin and different from most pensins. The crude extract had milk clotting activity similar to that of calf chymosin in the pH range 6.0 to 6.7, and different from that of porcine pepsin, had a higher ratio of milk clotting activity to hemoglobin hydrolytic activity than that of pepsin, was more stable than pepsin in casein at pH 6.1 and had a higher pH optimum for hemoglobin hydrolysis than that of pepsin (see Appendices A-D). Cheddar cheese prepared with the crude extract was comparable to that made with calf rennet in yields and sensory qualities (Appendix E) . These observations led to the hypothesis that crude SGP contains a chymosin-like enzyme. Occurrence of a chymosin-like enzyme in rat stomach has been reported by Kotts and Jenness (1976) based on the pH optimum of the gastric juice for hemoglobin hydrolysis, although chymosin is believed to occur only in young ruminant stomachs (Foltmann, 1970). Malpress (1967) demonstrated the absence of any protease that was stable in alkaline buffer in the gastric juice of children and concluded that chymosin does not occur in human stomach. However, Hirsh-Marie et al. (1976) reported the presence of a foetal pepsinogen that did not cross react with antibody against adult human pepsinogen. Therefore, occurrence of chymosin in non-ruminant stomach is still a controversy.

The possible existence of the multiple isoenzyme forms of seal gastric processes was also of interest, in view of the occurrence of

several isomers of gastric proteises in other species including humans (Asato and Rand, 1977; Kageyama and Takahashi, 1976; Meitner and Kassel, 1971; Ryle, 1970; Donta and Yan Vunakis, 1970; Bar-Eli and Merrett, 1970; Etherington and Taylor, 1970). Partial unipo acid sequence of a "pepsinogen" from ring seal (<u>Phoca himpida</u>) has been reported by Klem et al., (1976). However, the enzyme was not characterised to examine whether it is a papain, chymosin, or gastricsin. The possible occurrence of chymosin and of multiple isoenzyme forms was also not addressed. For these reasons gastric proteases from another pinniped species <u>Pagophilus groenlandicus</u> were studied. Harp seal stomach was chosen because it is easily available as a by product from the seal fishery in Newfoundland.

Preparation of Cheddar cheese was also repeated using the crude SGP and protesse A. Cheeses were also prepared with talf remnet as control and Fromase (<u>Mucor mienet</u> protesse), a leading commercial rennet substitute, for comparison. The protectlytic enzyme traditionally used as milk coägulant for cheese making is remnet - the crude preparation from suckling calf stomach containing mainly chymosin (E.C.J.4.23.4). However, during the last twenty years there has been a shortage of calf remnet because of an increasing production of cheese and decrease in the number of young calvess slaughtered (USDA, 1972). This shortage generated considerable interest in a search for new sources of milk coagulants. Protectlytic entrypes from plants, animal organs, and microorganisms have been extansively studied (see Ernstrom, 1974; Green, 1977 for review). A small number of remnet substitutes are now in commercial use. In 1976 calf remnet supplied only 25% of the world demand for milk coagulatis (dekoning, 1978). Milk coagulating enzymes from various sources and their characteristics will be briefly discussed here.

1. Chymosin

Chymosin is an actdic protease secreted by the abomasa of young runfants. It is secreted as a precursor called-prochymosin. Calf prochymosin has a molecular weight of 36,000 daltons. It undergoes autosctivation optimally at pH 2.0. The activation involves removal of a small peptide bond from the aminoterminus of the precursor (Foltmann, 1970). The isoelectric point of calf chymosin is 4.7 (Rispecti et al., 1977). The primary structure of [calf chymosin has been elucidated (Foltmann et al., 1979).

Chymosin possesses some unique characteristics that make it the

highly specific for a particular peptide bond in k-casein (Delfour et al., 1965) under the conditions used in cheese making. As a result of this narrow specificity there is minimum solubilization and loss of curd protein resulting from proteolytic action. The enzyme is relatively stable at the off of the curd so that the residual enzyme carried into the cheese contributes to flavor development by proteolysis during aging. The rate and specificity of proteolysis by chymosin during the ripening stage are conducive to good texture and flavor of the aged cheese. The narrow specificity of chymosin is also evident from its lack of action on ribonuclease (Bang-Jensen et al ... 1964), which is hydrolysed by pepsin (Berger et al., 1959). Moreover, the number of chymosin-susceptible bonds in the B chain of insulin is less than the number of pepsin-susceptible bonds (Bang-Jensen et al., 1964). Chymosin has a higher ratio of milk clotting to proteolytic activities (Green, 1972) and higher pH optimum for protein hydrolysis than does pepsin (Foltmann, 1959; Chiang et al., 1967). Chymosin is also more stable than pepsin under conditions of milk clotting, curd handling and cheese ripening (Green and Foster, 1974). During ripening. of cheese, chymosin preferentially hydrolyses qu-casein (Ledford et al., 1966)

Follmann (1970) separated calf chymosin into three different fractions having minor differences in their properties. Chymosin or crude preparations containing chymosin called rennet from other ruminants have also been studied. Lamb chymosin was chromstographically separated into three different fractions which had different ratios of milk clotting to proteolytic activities (Oruntaeva and

Seltov, 1971). Antiantakis and Green (1980) examined kid and lamb remnet! They found that remnets from these animals have lower ratios of milk clotting to proteolytic activities then calf remnet. againt at al. (1962) isolated "remnet" from adult goats. The yield of "emnet" was low compared to calf remnet. Lamb remnet has been reported to be more efficient in clotting sheep milk than bovine milk (Berfan and Kreal, 1971).

2. Rennet substitutes

a) Pepsins

Pensin (E.C.3.4.23.1) is similar to chymosin in many respects. Both are acidic protesses secreted by the animal stomach and autoactivated in the acidic medium of the organ as a result of the removal of a small peptide from the amino terminus of the corresponding precursors (Foltmann, 1970; Ryle, 1970). Certain sections of the polypeptide chain of chymosin and pepsin have been found to have similar amino acid sequences. However, chymosin was reported to be immunologically distinct from pensin (Rothe et al., 1976). Pensin has been widely studied as a rennet substitute perhaps because of the similarities between chymosin and pepsin discussed above and the fact that commercial rennet used for cheese production always contains some amount of pepsin (Emmons et al., 1978; Emmons et al., 1976). Reports on the suitability of bovine pepsin as a rennet substitute are contradictory. Bottazzi et al. (1974), Corradini et al. (1974) and Phelan (1973) reported successful use of bovine pepsin in cheese production. Fox and Walley (1971) reported no significant difference between the cheeses made with bovine pepsin and those made with calf

rempet either alone or in a 50:50 blend. Nowever, the pR 4.6 soluble to a seed cheene made with 100% bowine pepsin was lower than that of the cheese made with rennet indicating slower rate of protein degradation in cheese made with 100% bowledges alone rate of protein degradation in cheese made with 100% bowledges than in rennet cheese. Fox (1969) showed that bowine pepsin coagulates milk at pH 6.9. This observation suggests that the enzyme is table at this pH and prosumably also at other stages of cheese making and ripening.

Green (1972) found that the ratio of milk-clotting to proteolytic activity of bovine pepsin was slightly higher than that of calf rennet when tested on hemoglobin and slightly lower when based on casein as substrate. However, purified bowine pepsin has a lower ratio of clotting to proteolytic activity than calf chymosin (damer ct al., 1980; deKoning, 1978). The Cheddar cheese made using bovine pepsin was slightly inferior to that made with calf rennet. Emmons et al., (1976), Fhelan (1973) and Stanley and Emmons (1977) have confirmed this observation by reporting that bovine pepsin was associated with slightly reduced yield and lower quality of Cheddar cheese.

Bovine pepsinogen can be separated chromatographically into four components. All these fractions have potential hydrolytic activities on hemoglobin and also on the synthetic substrate N-achtyl-1-phenyl-alanyl-1-difodetyrosine (APDT) according to kassel and Metiner (1970). Foltmann et al., (1979) showed considerable homology in the amino acid sequences of portine pupsinogen and bovine proclymosin.

Use of portine pepsin as a remnet substitute dates back to the earlier part of the twentieth century (See Brnstrom, 1974 for a review). Studies on the use of portine pepsin as a rennet substitute were intensified during the last two decades (Chapman and Burnett, 1968; Malachouris and Tockey, 1964; Emmons et al., 1962; Olson, 1971; Thomasov, 1971; O'Keefe et al., 1977). There are conflicting reports on the quality of cheese prepared with portine pepsin. Davies et al. (1934) reported bitterness in Cheddar cheese made with portine pepsin Maragoudskis et al. (1951) and Malachouris and Tuckey (1964) showed that there was no bitterness in Cheddar cheese made with portine pepsin.

However, there are difficulties in using porcine pepsia's a a remnet substitute. MIL setting takes longer, and results in softer curd and loss of fat (Emmons at al., 1970; Chapman and Surfect, 1969). The longer setting time is attributed to inscrivation of the enzyme above pH 6.0 (Byls, 1970). Green (1972) and Green and Poblers (1973) showed that during the normal Chedder cheese making process all or most of the Forcine pepsin is destroyed. This report is consistent with the observation by Melichouris and Tuckey (1964) that Cheddar wheele made with porcine pepsin riponed slovly;

Emmons et al., (1971) used a 1:12 mixture of call fremmet and porcine pepain in Cheddar cheese manufacture, they found that the cheese made with the mixture and with control femnet were essentially of equal quality. They also suggested that porcine pepain should not be used singly, because slow-milk setting and far loss into whey might present problems. The research committee of the National Cheese Institute (1960) recommended that porcine pepain should not be used alone in cheese manufacture.

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Emmons (1970) observed that porcine pepsin is quickly inactivated by the high pH of hard water at the ambient temperature used in choose plants. Ernstrom (1961) showed that between pH 6.4 and 6.7 porcine papers High much longer milk clotting times than equal milk clotting princey of calf chymosin at pH.6.3. This suggests that porcine pepsinis relatively unstable under these conditions. Mickelsen and Ernstrom (1972) found porcine papers to be unstable above pH 6.0. Cheese made with porcine papers has a bland taste (Sardinas, 1972) because the empire is Anactivated during cheese making (Green, 1972).

Like bovine pepsin; porcine pepsin exists in multiple iscenzyme

forms. Nyle (1956) separated crude porcins pepsin into a major component, A, and minor components B, C and B. They are all believed to be produced of separate pepsinogens. Pepsins B NHS C are stable up to pli 6.9 but clot milk less readily than pepsin A (Ryle and Porter, 1959). Order porcins pepsin contains a gastricain, another gastric proteolytic engine (Ordans et al., 1967). Pepsin and gastric via are differentiated by their substrate specificity. Pepsin can readily hydrolyse the synthetic substrate N-occyl-1-penylalanyl-u-dicodepyrobius (APDT) whereas gastricain cannot (Tang, 1970). The pH optimum for hemoglobin hydrolysis by porcine gastricsin is 1.0, while that of pepsin is 2.0 (Chiang et al., 1967). The pH stability of gastricain is similar to that of pepsin and is probably subject to the same stability problems as porcins pepsin during cheese samufacture (Krmatows, 1974). Rowever, fish gastricains are relatively stable in mikaline sil (Chiang and 960ce, 1981).

Chicken pepsin was used in laboratory scale Khashkhaval and Emmental cheese preparation in Israel (Gordin and Rosenthal, 1978). There was no appreciable difference in taste, flavor, or caxture between the cheeses made with chicken pepsin and those prepared with calf rennet. Neither of the cheeses had any bitterness. While the soluble-nitrogen and free autoo acid contents of the Emmental type cheeses were comparable in both the cases Khashkhaval cheese made with chicken pepain showed much higher soluble nitrogen than rennet cheese suggesting enhanced proteolysis in the Khashkhaval cheese made with chicken pepain. The authors also report the use of chicken pepain for the successful manufacture of many kinds of cheeses in Israel.

Green (1972) reported that a Cheddar cheese trial with chicken pepsin resulted in poor quality cheese with texture and flavor defects, probably due to excessive proteclysis by residual chicken pepsin. Stanley et al., (1980) also reported chicken pepsin to be unsuitable for Cheddar cheese because the cheese made with it resulted in softer body and bitter taste. In addition, there aws an increased loss of nonprotein nitrogen(NPN) into whey from curd made with chicken pepsin, compared to that made with calf rennet. This increased NPN loss suggests a potentially lower yield of cheese than the cheese made with 4 calf rennet. Nonprotein nitrogen in the cheese made with chicken pepsin was approximately twice as much as in the cheese made with calf rennet; Donta and Van Vunakis(1970) characterized chicken pepsin Chicken forestomach 4 different pensingens. Pensingens A and D and their corresponding enzymes were similar in amino acid composition. Pepsinogen C was immunologically different from pepsinogens A and D. Pepsinogen B was active only on the synthetic substrate N-Cbz-L-Glu-L-Tyr. Bohak (1970) showed that chicken pepsin was stable between pH 1.0 and 8.0 at 25°C.

(b) Microbial rennet substitutes

Microorganisms are probably the best means of producing low cost

enzymes having uniform quality. There are thousands of genera to choose from, and it is relatively easy to control production of microbial enzymes. Numerous genera of Microorganisms have been screened for production of suitable remnet substitute (Tendler and Burkholder, 1961; Srinivasan et al., 1964; Schulz et al., 1967; Arina et al., 1967; Ari

Sardhas (1972) reviewed the literature on bacteria reported to produce milk-clotting enzymes including the genera Alkaligenes.

Bacillus, Odrymchacterius, Escherichia, Lactobacillus, Proteus, Pacudononas, Serratia, Staphylococcus, Streptococcus, Streptococcus,

Bacilius cereus processe was patented as a remnet-substitute by Kiles laboratories Inc. (Sardinas, 1969). Cheddar cheese sade with this enzyme was found to be free of bitter flavor (Srinivasan et al., 1962a, b). However, the texture was not typical of Cheddar cheese and maturation was delayed. Melachouris and Tuckey (1968) reported higher proteolytic action of this enzyme than remnet on casein. The enzyme hydrolysed B-casein to a greater extent than did calf rennet.

Parants were issued to Murray and Prince (1970) for making cheese with rennet substitute from <u>Bacilius subtilis</u>. These processes were found to release much more NPM from milk than did calf rennet at the clotting level and to degrade or and B caseins more rapidly and nonspecifically (Pubad, 1969). <u>Macilius subtilis</u> milk coagulant has not been used commercially.

Numerous genera of fungi also have been acreemed for suitable protectytic enzymes, and patents have been granted for some of these enzymes (See Sardinas, 1972 for a review). Whitshill et al. (1960) and Oringer (1960) were granted patents for commarcial preparation of mik clotting enzymes from <u>Confidiobolus</u>, <u>Entomorphora</u>, and <u>Basidiologus</u>. The enzymes were characterized and found to result is bitter flavor in chees.

Proteolytic enzymes from Endothia parastica, Neor pustlus
(Lindt) and Mucor michel are being commercially used: Sardinas (1965,
1967, 1956), Larson and Whitzaker (1970) and Hagemeyer et al. (1965)
characterised the acidic processe from Endothia parasitica. It has a
molecular weight of 34,000 to 39,000, and an isoelectric point of pH
4.6. The milk clotting activity of the enzyme-was less sensitive to
pH change in the range 5.1 to 6.5 than calf remnet:

It was found to have broader substrate specificity than chymosin and pepara. The enzyme has been reported to accelerate the ripening of Cheddar cheese (Showrs and Bavipotto, 1967). Cheddar cheese prepared using a 311 mixture of the enzyme with calf rennet developed poor flavor and texture (Mortis and McKennie, 1970). Endothia parseities protesse was found to be more protectlyfic than calf rennet.

and produced bitterness in Edam, Tilsit and Butter cheeses (Mnomasow et al., 1970). Raper et al., (1969) reported the proparation of good quality Emmental cheese with protease from E. parasitica. Production, of Emmental cheese involves cooking at high temperature. The enzyme is probably inactivated at that temperature before it effects by hydrolysis to produce bitter peptides.

An acidic protease from Nacor pusilus (Lindy) was characterized by Arima et al., (1967) and Iwasaki et al., (1968a, b. 6), and its properties were later reviewed by Arima et al., (1970). The phryme has an isoelectric point of pig 3.5 to 3.8 and is stable in the pil range 4.0 to 6.0. The proteolytic activity of the ensyme is less specific than that of calf rennet. It was found to be more proteolytic than calf remnet, destroying 8-casein to a greater extent than did calf rennet and pepsin (Mtckaleen and Fish, 1970). Mucor pusilus protease e was found to be a matisfactory rennet substitute in the manufacture of Brick, Sutter Camebber, Cheddar, Cottage, Edam, Gooda, come Italian varieties and Tilat cheeses (Arima et al., 1970; Scholz et al., 1967). However, bitter flavor was reported in some bheeses hade with the enzyme (Richardson et al., 1967; Kikuchi and Toyoda, 1970). Richardson et al., reported that MP protease was incompatible with calf remnet, because in solution it destroys the Inter.

Patenta have been granted for manufacture of milk compulant from several strains of M. Mighed. These are M. nichei Cooney ef Berson (Ammstrup, 1968), M. michei NRUL 3420, M. michei ARO 1665.7 (Feldman, 1969), M. michei NRUL 3189 (Bäxter: Laboratories, Inc., 1970). Several companies are selling these milk compulants under various trade names (Rennilase - Novo Industrice). Frommes - G.B. Permentation Industrices, 711.)

Proteolytic enzymes from M. miched Cooney at Emerson was purified and tharacterised by Ottensen and Rickert (1970a, b). Like the other two commercial fungal remnets, M. miched rennet is an acidic processed with an optimum pN for hemoglobin hydrolysis at 4.5. Its specificity on the oxidized B chain of inquiin was reported to be similar to that of calf rennet. Ottensen and Rickert (1970b) observed that the enzyms is quite stable; it retained 90% of its activity in the pN range 3 to 6 during 8 days of incubation at ambient temperature. It has a molecular weight of 38,000 daltons and an isoelectric point of pN 4.2.

Yield and quality of Emental choses manufactured with M. michel

processe were the same as that with calf rennet, and there was no bitter taste (Kiuru, 1969; Mayrg 1971). Thomasow of al. (1971) made similar observations about Edam, Tileit and Butter cheese made with M. michel rennet. Domiati cheese made with M. michel rennet was axisfactory, although it contained more soluble N than calf rennet cheese (Handy, 1970).

(c) Rennet Substitutes from higher plants.

Remnet substitutes from higher plants were sought not only because of remnet shortage but for religious and ethnic reasons as well (Reed, 1975). In India, enzymes from fig and a wild shrub (Strubtus asper) yielded alightly less curd than that made with calf remnet (Veringa, 1961). Both curds were softer than that made with calf remnet. According to Gosthuizen and Scott Blair (1963) the mechanism of casein hydrolysis by ficin is different from that of calf remnet. Properties of ficin, a sulphydryl profeolytic enzyme from fig have been reviewed by Liener and Priedensen (1970). The milk colotting and proteolytic activities of ficin have been reported to be

due to two different factors. Cheese made with ficin had bitter flavor which disappeared with aging, presumably due to degradation of bitter cectides (Krishnaswamy et al., 1961).

Bromelain, a proteolytic enzyme preparation from gineapple was reported to be too proteolytic to be used as a remnet substitute (Poszarne-Eanqla et.al.,1969). Papain from papaya latex was reported to have strong proteolytic activity (Dastur, 1948) but its proteolytic activity was also too high for use as a remnet substitute. Papain had a free sulphydryl group essential for catalytic activity and loses activity reversibly due to oxidation in air in the absence of added cystedne (see review by Arnon, 1970). There is controversy over the catalytic activities of papain, one group of workers supporting the view that proteolytic and milk clotting activities are the same whereas another, group believes those two properties are unrelated (Stelton, 1971; Hinkle and Alford, 1991; Balls and Hoover, 1937).

Kothavalla and Kubchandani (1940) used extract from <u>Michania</u> coagulans for making some Indian varieties of cheese in which no unusual flavor was found. However, there were high fat losses. Extract from cardoon flower is traditionally used by portuguese farmers for Serta cheese from sheep's milk. Studies on the milk clotting properties of the extract showed that the clotting activity was more sensitive to pil of milk than calf rennet (Vieira de Sa and Barbosa, 1970a, b, c). It was more proteolytic than calf rennet when used in Edam cheese which was litter and acid. Emetron (1974) reported extremely bitter flavor in Cheddar cheese made with cardoon extract only after 30 days of ripening. Supta and Eskin (1977) isolated a silk-congulant from ash gourd and used it for Cheddar

chaefs. It was found to be a neutral procease. Cheedar cheese made with the saft gourd procease had a slightly lower sensory score than that of remnet cheese. Fat loss in whey was higher and the yield of cheese was lower.

3. Conservation of rennet and other alternatives

Shortage of remnet may also be circumvented by making changes in the conventional method of cheese production. The amount of computant y required for making cheese may be reduced by (1) concentrating the milk, (2) using CeCl₂, (3) reising the milk temperature, and (4) the congulant may be timobilized for repeated use.

The use of twofold concentrated milk to make Cheddar and Cheshire cheeses resulted in 75-80% saving of remet compared with the conventional method. However, both the cheeses showed less intense flavor than the cheeses made with untreated milk (Chapman et al., 1974). The above-named authors also used milk-concentrated more than twofold to make Cheddar cheese. They found that the cheeses so prepared were of poor quality (Chapman et al., 1979). Green et al. (1981) prepared Cheddar cheeses with 1.7-to 4-fold concentrated milk and found that the fat contents of the cheeses were less than normal. Whreever, with increase in the concentration factor of the milk, the rate of casein breakdown, the intensity of Cheddar flavor and the concentrations of HgS and sethanethfold in the cheeses decreased.

"Addition of Caci; to milk in order to reduce the amount of congulant required to Clot milk is a common practice, particularly with the more proteclytic congulants (Green, 1977). However, the use of CaCl; is reported to reduce the ripening wate of cheese (Eristrom

et al.; 1958) ...

Other conservation methods include the recovery of remet from whey (Green, 1977). This method would have an additional advantage in that the rennet free whey is more suitable for the preparation of concentrated whey protein. Another method is the possible use of fisculated calves to produce femnet continuously. Canguli (1970) reported that such calves produce remest at a satisfactory rate.

The production of microbial remnets by genetically "enhanced" microcranisms has been suggested by Telsey (1981). However, since the commercial rennets of microbial origin are not completely satisfactory in terms of the yield and quality of the cheese they produce (deKoning, 1978) an ideal solution to the rennet shortage would be to mass produce call chymosis by incorporating the calf chymosin genes into appropriate microorganisms. Techniques are how available for the feolation of eukaryotic genes and their incorporation into microorganisms (Dally et al., 1981). The

The use of ismobilised proteases is another possible choice to conserve rennet. Green and Crutchfield(1969) ismobilised calf chymotin to use it repeatedly for clotting milk. However, the enzyme had low activity because of inactivation during the ismobilisation and because of leaching of the enzyme from the support. Successful ismobilisation of chymosin may only partly solve the problem of remnet availability, because the card formed by the ismobilised enzyme would lack the residual chymosin required for the ripening process of the cheese. However, remnet can be added to the turd and there would still be a set saving of rennet. The successful ismobilization of any processe which does not affect the yield of curd should be suitable for this purpose. Objutys at. al., (1979)

imobilized calf rennet and an alkaline protease from <u>Bacillus' subtilia</u> on Dows. Rennet was found to have a very short half-life, whereas the alkaline protease from <u>B. subtilia</u> had a halfilife of 8 days after imobilization. Chedder cheese prepared by using the immobilized alkaline protease was found to ripen at a rate estallar to normal rennet cheese. The organoleptic quality of this cheese was similar to those of normal and immobilized rennet cheeses. However, the ripening rate as judged by the release of NEW for immobilized calf rennet was slower than that of normal calf rennet cheese. These studies cast doubt on the supposition that residual rennet contributes to flavor development. Green (1977) has argued that the interpretation of these studies is dependent on the investigator carefully establishing that no enzymes leach from the reactor into the milk.

4. Chemistry of milk clotting

Enymatic milk cletting occurs in two phases. In the primary or enzymatic phase, k-casein in milk undergoes bytchylysis and thus loses its ability to stabilize the casein micelle. In the secondary or monenzymatic phase the micelles aggregate to form a clot. This takes place in presence of Ca** and the mechanism by which it brings about the aggregation is not fully understood. Alais at al., (1953) has demonstrated this biphasic nature of milk clotting. The asymatic reaction was carried out at 2°C at which no clotting occurred.

Obsgulation took place immediately on warming the milk to shigher temperature. Berridge (1942) made a similar observation. Deligicish (1979) showed that coapplation is not initiated until 88% of the k-casein is hydrolysed. The action of chymosin on K-casein was the

subject of extensive studies (Make, 1959; Garnier, 1958) until Delfour of all, (1965) established that chymosin hydrolyses a peptide band involving phenylaisaine and methionine. As a result a soluble glycomacropeptide and an insoluble peptide are produced. The glycomacropeptide is a heterogeneous glycospetide with a molecular weight of 6,000 to 8,000 daltons (Hill et al., 1979; Nitechmann and Heari, 1959). Besides chymosin other proteases also, e.g. peptin and chymotrypsin (Dennis and Wake, 1965; Green, 1972), E. paramitica processe (Vanderpoorten and Weckx, 1972), N. pustium processe (fu et al., 1965), also attack r-casein at the mane site.

The secondary phase of milk clotting is not clearly understood; because it is not known how the major casein components, namely or, and 8 caseins interact with each other and also with x-casein (which is apparently responsible for micelle stability) and colloidal calcium phosphate to form a stable suspension called casein micelle. Nor is it known how k-casein loses its capacity to stabilize the micelle because of hydrolysis. The function of Catt to bring about aggregation of the micelles is also not clearly understood. There are various models to explain the micellar arrangement of caseins in milk and the mechanism of enzymatic milk coagulations but none of them satisfactorily explains clotting phenomenon (see Ernstrom, 1974 for a review). Most recent model of casein micelle is based on reconstitution of micellerusing gold labelled casein components (Payens, 1981). Electron micrograph of casein micelles reconstituted using gold labelled c-casein showed that all the c-casein is on the surface. Similar experiments using labelled a and & caseins showed that these components are in the core of the reconstituted micelle. However, Payens (1981)

did not mention whether the micelles thus reconstituted behave like the native casein sicelles in mik clotting.

5. Criteria for a good rennet substitute

Evengings and Sandts (1973) recommended that a cosquiant should have the following properties to replace calf rennet: (a) yield of cheese should be equal to that made with calf rennet; (b) cheese should not develop off-flavor or bad texture; (c) the use of the cosquiant should not require any change in standard cheese-making practices; (d) should be free from toxic and antibiotic activity and pathogene; (a) should not have lipolytic or other conteminating entires; (f) should not be too proteolytic; and (g) price should be acceptable.

According to defoning (1978) none of the commercially available remnet substitutes meet all the requirements listed above. For example, pilot-scale trial with portine peps in resulted in lower quality Cheddar cheese. Although the yields of Cheddar cheese made with boving pepsin and engues from Endethia parasitics, Mucor pusition and Motor michel were lower, the cheeses were otherwise satisfactory. All the commercial fungal renners contain anylase.

The most common problem with rennet substitutes is the development of flavor and texture defects in cheese (defening, 1978).—This is attributed to high proteolytic activity of most rennet substitutes relative to their milk electing activity (Green, 1977). Excessive proteolysis also results in located fat in whey (Verings, 1961). To little proteolysis by the residual complaint also has undestable offects on flavor. For example, Cheddar cheeks made with porcine pepsin failed to develop characteristic aged flavor(Green and Foster,

Protectlytic specificity is another important factor. Degradation of *a_-casein is predominant in a normally ripened cheese (Lefford et. al., 1966; §tanley and Emmons. 1977) wherean the cheeses with poor flavor are associated with a high rate of degradation of A_-casein (Fishlari et. al., 1973). The ratio of silk-clotting to protectlytic activities of a rennet subscitute is a generally accepted indication of its narrow specificity. (Green, 1972). However, the best way to determine if a protectlytic engage is against embetture is to prepare cheese on an industrial scale (defoning, 1978).

Lipsse, smootated with microbial remners (Richardson et. al., 1967) is detrimental to the flavor of certain chosens (Green, 1977). Some— Italian chosens require pregastric lipsse present in remnet pasts which cannot be repaired by any other lipsses (Long and Marper, 1956).

6. Flayor, texture and cheese-ripening

The development of flavor in a cheese with aging is a function of the breakdown of carbohydrate, fat and protein by the enzymes from the congulant and the microorganisms growing and declaying in the cheese. The combined actile of these enzymes result in the production of a few classes of compounds. Some of these compounds, e.g. amino acids and carbonyle are qualitatively common to all the cheese-types while others are characteristic of a particular cheese-types while others are characteristic of a particular cheese-types by acute of the cheese compounds broadly into two categories by vacuum distillation. The volatile distillate contributes aroma while the non-volafite register imparts the mensation of taste to the cheese. He showed

that the non-volatile part was composed of lactic acid, andino acids, keto acids and other non-volatile acids and animas and sail. The volatile component is composed of fatty acids, aidehydes, Ketonsa, alcohols, animas, esterm, its and autholian.

An examination of the literature on cheese flavor shows that the relationship between characteristic flavor of cheese and a chemical compound or a group of compounds is based on (a) the presence in the cheese of that compound, (b) increase in concentration of the compound with ripening and increase in flavor intensity of the cheese, and (c) reproduction of the cheese flavor when the compound in question is added to a bland base. However, one has to be cautious in drawing conclusions using such criteria. A compound found to be associated with flavor may actually have been produced in parallel with another compound which is directly responsible for the flavor. Also the flavor compound may be present in chemically undetectable amounts. The fatlure of an added compound to reproduce or to enhance a particular aspect of flavor does not rule out the compound as a flavor component because it is almost impossible to add the compound in its original chemical and physical form along with other compounds which may play supplementary roles in expression of the flavor.

Harper (1959) observed a definite correlation between free amino acids and characteristic Cheddar cheese flavor. Cheese with low free amino acid content lacked flavor and cheese with high antoo acid content had characteristic flavor. Glutamic acid occurred in the highest concentration and appeared to have a direct correlation with flavor. Furthermore, Harper found that free amino acids added to a bland base the concentration found in Cheddar cheese resulted in cheese

fagor lacking aromatic character. Silverman and Kosikowski (1955) made similar observations with Suiss cheese flavor. Others also reported occurrence of free mino acids in cheese (Marper and Susmor, 1997; EstRowski, 1951; kindeviat et al., 1953; Bullock and Irvine, 1956; Malachouris and Tuckey, 1964). O'Kesfe at al., (1976) stated that the full-range of estime acids found in casein also occur in cheese and thefr liberation is accompanied by flavor development. O'Kesfe at al., (1976) observed that coly a limited range of free amino acids, assaly sethioustes, histidine, glycine, series and glutants acid are liberated in measurable amounts in cheese which were chantcally acidulated. Instead of adding starter culture before remeding. This observation led to the conclusion that free amino acids in Cheedear cheese are mainly the result of stcrobial peptidase activity. Sal 31 and Kroper (1981) observed increase in poptidase activity in Cheddan cheese with reparing.

Mabbit (1961) demonstrated that the ratios of the quantities of free axiso acids occurring in Cheddar cheese were in some instances different from those in casein. He added an aqueous solution of a mixture of axiso acids in propertion found in aix-month-old Cheddar cheese to freshly made curd. Sensory evaluation of this "cheese" had a brothy flavor but so Cheddar flavor. Dacre (1953) also made a similar observation. However, Mabbit (1961) concluded from his studies that the free axiso acids contribute towards the background flavor.

There seems to be lack of agreement between workers on the relative contribution of microorganisms and coagulant enzymes towards

protectysts in cheese. Orla-Jensen (1939) showed that remnet along can hydrolyse silk protein but cannot produce free smino acids.

O'Keefe gt al., (1976) concluded from their observations on Cheeder cheese made with and without starter organisms that sic roorganisms when no significant contribution to the formation of large peptides from casein during cheese ripening. Ledford et al. (1966), Reiter et al., (1969) and Oreen and Foster (1974) also reported that remnet is a major proteclytic factor in Cheeder cheese. However, Green and Foster (1974) and Otherya and Sato (1972) suggested that both remnet and microorganisms are responsible for proteclysis in cheese. Zennet was found to accelerate proteclysis by starter bacteria in milk

Using asseptic vat technique Ratter et al., (1969) showed that cheeses prepared by chemical acidulation instead of using starter culture before remeting had failed to develop flavor whereas chesse made with starter alone did develop flavor. These observations considered in the light of the report by Green (1971) that Cheddar cheese prepared using porcine papers (which is completely inactivated during cheese making) is very slow in flavor development ledd to the logical conclusion that remnet brings about the majority of cameia breakdown to form large peptides. These peptides are broken down to sainto acide by sicrobial enzymes in cheese. Emmons at al., (1960;s,b) showed that some strains of starter organisms produce bitter flavor.

Nonstarter microorganisms also affect choose flavor. Res milk with a high bactarial count can give rise to poor-flavored chees(Smith et al., 1856). Pryor (1869) made a comprehensive review of the role of microflora on cheese tipening.

Belative rates of breakdown of come of the peptides formed also determine the cheese flavor. Stanley and Emmons (1977) showed that in a normal Cheldar cheese of casein was degraded to a greater extent than \$6\$ casein. The degradation products had higher electrophoretic mobility than the corresponding—casein components. Hydrolysts of \$6\$ casein was implicated in the development of bitter flavor in Cheddar cheese (Fhelanet al., 1973). However, bitter peptides may originate from hydrolysis of both ag, and \$6\$ caseins (pelissier et.al., 1974). Various components in cheeses are in a state of dynantic equilibrium. The large peptides resulting from casein break, down give riae to smino acides and the latter are then further degradad to saines, sladelyses and ketones; etc. Strecker degradation is an example. In this reaction maino acides and disabonyls interact to produce an aldehyde with one less carbon aton than the smino acid (Schobberg and Moobssher, 1952).

RCO - CO - R + R'CHNE₂COOH + R'CHO + CO₂ + RCHNE₂ - COR
All the necessary reactants occur in cheese (Kenney and Day, 1957).

These authors also demonstrated the Cheédar flavor in test tube by
heating individual amino acid solutions with isatin (Indole-2,3-dions).
Derivatives of each amino acid solutions with isatin (Indole-2,3-dions).
Hethionine derivative had Cheddar cheese flavor: Strecker degradationproducts obtained by steam distillation of an incubation mixture of
casein solution, proteolytic enzymes, isatin, pyruvic and other keto
acids were found to impart Cheddar-like flavor to Cottage cheese curd.

—Although protein and carbohydrate derived compounds are necessary for cheese flavor the ligid component and its derivatives are perhaps much more important (Ernstrom, 1974). Obven and Tuckey (1969) demonstrated that cheese made of skim nik lace typical flavor. Free fatty acids formed from anino acids or intermediates of lactors matabalian (Schormiller, 1988) have been reported in all cheese types. Pattorn of fatty acid seems to be characterizate of a particular type of cheese. Marper (1959) showed that is cheeder cheese accit and butyric acids are the predominant fatty acids whereas propionic acid is the major fatty acid component in a sormal Svisa cheese. In Blue cheese fatty acid with 5 and higher number of carbon atoms occur in higher concentration than the short chain fatty acids, Sutyric acid is predominant in Froviolous cheese. Schormiller (1969) has reviewed the literature on fatty acids of warlow kinds of cheese.

Kristoffersen<u>et al.</u>, (1959) and Bills and Day (1964) determined fatty acids in Cheddar cheese. Notither group found any correlation, between free fatty acids and flavor except in cheese with off flavor which had 10 fold increase in free fatty acids. Patton (1964) considered short chain fatty acids to be indispensable for Cheddar cheese arons.

Carbonyl compounds of warfous chain lengths have been associated with cheese flavor. Acidic carbonyl compounds reported to be present in cheese are exalesuccinic, glyonylic, pruvic, x-ketoglutaric, accesseate and a-ketofscoaproic acids (lasseat and Harper, 1938; Kristoffersen and (ould, 1939; Walker and Harvey, 1959).

Mabbir (1951) reported the following neutral carbonyl comprounds in Cheddar cheene: diacetyl, butyraldehyde, acetaldehyde, acetyl methyl carbinol, acetone, sethyl achyl actone, 2-butypose, 2-sentamons, 2-heptamons, 2-monumons, 2-undecamone, 3-tridecamone, 3-hydraybbirmone, Tormaldehyde, 3-methyl butamol, and proctonaldehyde. Wedamuthu et al.,

(1966) isolated acetaldehyde, propionaldehyde, acetone, pyruvic acid, diacetyl, glyoxal and an a-ketoalkanal from a milk culture inoculated with a lactic starter known to produce normal flavored Cheddar cheese. culture that consistently produced fruity flavor in Cheddar cheese was found to produce more carbonyl compounds than the normal starter under the same conditions. Besides it produced formaldehyde instead of propional dehyde. Beide and Hammond (1979a, b) fractionated Swiss cheese into oil-soluble, water-soluble volatile and water-soluble nonvolatile components. Each of these components were further analysed for individual compounds and their contribution to cheese flavor. Water-soluble volatile fraction was composed of acetic, propionic and butyric acids, diacetyl and ammonia. A synthetic mixture of these compounds reproduced the flavor of water-soluble volatile fraction. The water-soluble nonvolatile fraction contained small peptides and amino acids and these were correlated with the acidity of the fraction, not the lactic acid content or oH. The interaction of calcium and magnesium ions with small peptides and amino acids was attributed to the watersoluble nonvolatile fraction. Free fatty acids were correlated with the nutty flavor of the oil-soluble fraction whereas the neutral portion of the oil-soluble fraction gave undesirable "fermented" . flavor. However, the flavor of the whole cheese could not be predicted from those of its individual fractions.

Sulphur compounds have been reported as important in cheese.

flavor. Singh and Kristoffersen (1970) accelerated flavor development
in Cheddar slurry by treatment with reduced glutathione. Singh and
Kristoffersen (1971) also unde the same observation about Swiss
cheese flavor. Hanning (1978) correlated Cheddar cheese flavor with

the concentrations of 100 and methanethiol. He concluded that reduced sulphur compounds, 9.8. DTT, OSH which are known to accelerate flavor development do so by providing necessary redox potential to form methanethiol from its precursor, namely methionine. Degradation of sulphur-containing amino acids by Strecker degradation has also been considered as important for flavor.

From above discussion it seems that our understanding of cheese flavor is far from clear. Monitoring one or more of the flavor associated compounds gives an indication of the rate and stage of the ripening process but not a complete picture of the cheese quality. "Component balance theory" of Kosikowski and Mocquot (1958) seems to explain flavor development more logically. According to this theory only a small number of compounds are responsible for cheese and other food flavors. Some of these compounds are common to all cheeses and are responsible for background flavor. /Flavor of a particular cheese type results from blending of a number of these compounds in a definite proportion with respect to the background. Intensification of a normal flavor results when concentrations of all these compounds increase, maintaining the balance. For example, in a good Cheddar cheese the amino acids, peptides, carbonyls, fatty acids, salts and residual para-casein may be present in certain proportion to impart background flavor and characteristic Cheddar flavor may be perceived when the volatile sulphur compounds (Manning, 1978) are present in the cheese in a definite proportion with respect to each other and also with respect to the background.

Like flavor, texture is also an important characteristic of cheeses, and the biochemical changes occurring during ripening also influence the texture of cheeses as well. Milk-coagulating engages contribute to cheese texture either by retention of fax during: curding and also through proteolysis (Stanley and Emmons, 1977; Stanley et al., 1980).

MATERIALS AND METHODS

I. Seal Stomachs

Stonachs from harp seals (<u>Pagophilus groenlandicus</u>) were collected by officials of the Department of Fishertes and Oceans, St. John's in 1979 and 1980. The seals were caught off the coast of Newfoundland during the hunting season in March. The stomachs collected in 1979 were from approximately two-week-old pups, These stomachs were sublipped frozen. On arrival in St. John's the stomachs were cleaned and freeze-dried.

The stomachs collected in 1980, were from both 2-week-old and 2-year-old seals, shipped fresh chilled on ice: On arrival the stomachs were cleaned, the nucosal layer separated and freeze-dried. The vet weights of the clean stomachs from two-week-old pups were from 30 to 75 grams and those from the two-year-old seals were from 180 to 250 grams. The freeze-dried samples were stored at -20°C until

II. Chemicals

N-actyl-L-phenylatanyl-L-ddiodotyrosine (APDT), ninhydrin, substrate grade hemoglobin, yeast ribonucleic acid (RNA), dioddum ethylemediamine tetraacetic acid (EDTA), actylemide, N,N'-methyleme-bis-acrylanide (BIS), N,N,N'N'-tetramethylethylemediamine (TEMED), ammonium persulphate, riboflavin, penicillin, strep tomycin, bovine pancreatic ribonuclease (RNAsse E.C.2.7.7.16), twice-crystallized porcine pepsin (E.C.3.4.2.3.1), and calf chymosin (E.C.3.4.2.3.4) were purchased from Sirma Omenical Co., St., Louis, NO. Diethylaninosthyl

Sephadox A-50 were purchased from Pharmacia Time Chemicals (Uppsala, Sweden). Carbobenzoxy-D-phanylalanyl-triethylene-fetranine-Sepharose (Z-D-Pha-T-Sepharose) was purchased from Pierce Chemicala, III.

Hamsen's butterealk culture and calf remnet were purchased from Horan Lally Co., Mississauga, Ontario. <u>Mucor michesi</u> protesse (Fromase 100) was a gift from G.B. Fermentation Industrier-Inc., III.

All other chemicals were of the highest purity available and purchased from various commercial sources.

III. Preparation of crude enzyme

Enyme used in preliminary studies discussed in the introduction was prepared following the method of Qadri, et al., (1962) with a slight modification. Freeze-dried whole stomach (see Section I) was pulverised in a Waring blender at the maximum setting in the cold room at 4°C for a total of 2 minutes with 30-second bursts and a cooling period of 2 minutes in between. The powder was stored at -20°C until used.

For extraction of enzymes 1 gram of the powder was stirred with 20 ml of 10% acetic acid at 4°C for 12 h. The homogenate thus obtained was adjusted to pl 5.3 by adding 1M NaOH and centrifuged in a Sorval NC-5 superspeed refrigerated centrifuge for 30 minutes at 38,000xg. The supernatant was concentrated about 10 fold by ultrafiltration in a 43 mm Amicen ultrafiltration apparatus fitted with a PM₁₀ membrane using nitrogen gas as a propellint. This preparation was used in the preliminary work discussed in the introduction (results in Appendices A-E).

- . Isolation and Purification of zymogens
- (a) Extraction of crude symogens: The freeze-dried samples of gastric pucosa from two week old and two year old seals collected in 1980 (see Section II) were pulverised as in Section III. —One gram of pulverised mucosa was suspended in 20 ml of buffer A (20 mM sodium phosphate pH 7.2 containing 5000 units of penicillin, 50 mg and streptomycin per litro) and stirred gently at 4°C for 12 h. The sturry thus obtained was centrifuged at 48,000xg at 4°C for, 30 minutes. The pellet was resuspended in 10 ml of buffer A and centrifuged at 134,000xg To 60 minutes at 4°C in a Beckman 13-50 ultracentrifuge. The supernatant was used as a crude preparation of symogens. This was converted to enzyme by adjusting the pH to 2.0 with 1N HCl and incubating at 30°C for 30 minutes when needed.

(b) Ion-exchange chromatography: Diethylaminosthyl Sephadex A-50 (DEAE-Sephadex A-50) was packed into a 2.5 x 30 cm column following the manufacturer's instruction. The column was equilibrated with buffer A for 24 h before use.

Forty ml of crude symogen preparation obtained as described above (IVs) were dialysed against 2xZL buffer A at A*C and applied to the ion exchange column. Five ml fractions were eluted with buffer A until the absorbance at 280nm of the fractions and their proteolytic activity approached zero. The column was then eluted by stepwise addition of 0.1%, 0.2M and 0.8M %GLI to buffer A.

(c) Gel filtration: An 88 x 1.6 cm column was packed with Sephadex

O-100 according to manufacturer's instruction and washed with buffer B.

(buffer A containing 1 mM dithiothreitol, 1 mM disodium ethylenediamine retreacetic acid, and 0.5% NaCl).

Fractions eluted from the DEAL-Sephadex column with potential proteolytic activity were pooled, concentrated by ultrafiltration and 4 ml of the concentrate were applied through the bottom of the column and eluted with buffer B in the ascending mode. The column was previously calibrated with a mixture of standard molecular-weight markers consisting of bovine serum albunta (65,000 daltons), chicken egg albunta (43,000 daltons), sheep prolactir (23,800 daltons), cytochrome C (12,700 daltons), and blue dextram (2,000,000 daltons).

(d) Affinity-column chromatography: The method of Fujivara and TBuTu (1977) for purification of pepsin was followed with necessary modifications for symogen. Ten nl of carbobenzoxy-D-phenylalanyl-tricthylene-tetramine-sephanose (Z-D-Phe-T-sephanose) ware packed into a 10-nl disposable plastic syrings with glass wool to support the gel. Two nl of concentrated symogen A from DEAT-Sephandex column were mixed with 4 ml of the starting buffer and applied to the column and eluted quickly with the same buffer (30 mM acctate pH 3.5, containing 0.1M NaCl). When the absorbance of the fractions at 280 mm reached the baseline the column was washed with 1M sodium acctate pH 3.5 containing 1M NaCl. The adsorbed sympogen was then eluted with the starting buffer containing 1M guandine chloride. Aliquots from fractions were incubated with hemoglobin to monitor activity and the fractions eluted with the buffer containing guandine chloride were pooled and mixed with an equal volume of 0.2% sodium phosphate buffer

pH 7.2. The mixture was then dialysed against 2L of water at 4°C for 3 h and concentrated by ultrafiltration using a PMIO membrane.

V. Determination of Enzyme Activity

(a) Measurement of pepsia activity: The method of Schak(1970) was followed to determine the proteolytic activity of anymes using hemoglobin as the substrate with the following modification. A 7.5% (w/v) solution of bovine hemoglobin in 0.06% HGL was dialysed against 50 volumes of 0.06% HGL at 4°C with 2 changes of the acid at 4 h intervals. The dialysed solution was then freeze-dried and stored in screw cap bottles at 4°G with used. Dialysis removed most of the trichloroscetic acid (TCA)-soluble meterials that absorbed at 280 nm.

For assay, 8 ml Volume of a 22 hemoglobin solution in 0.05M HC1 (pH 1.8) was equilibrated at 30°C in a water bath and incubated with 1.0 ml of proteolytic enzyme. At various time intervals, 0.8-ml aliquots were removed, and the reaction stopped by mixing with 1.0 ml of 62 TCA. After cooling on ice for 30 minutes the samples were centrifuged at approximately 3,000xg for 30 minutes. The absorbance of the supernatant at 280 mm was measured in a beckman DU 8 spectrophotometer. For control 0.8 ml solution of hemoglobin was incubated in duplicate for the same length of time and the enzyme added after TCA. One pepein unit is the amount of enzyme that produces an absorbance increase of 0.008% per minute under this condition. Enzyme activity measured with hemoglobin as the substrate has been termed pepsin activity or proteolytic activity in this study.

⁽b) Milk-clotting assay: The milk clotting activity of proteases was

determined following the method of Berridge (1945) except that the volumes of the Berridge substrate (12 g skin milk provder suspended in 100 ml TOIN CaCl2) and enzyme were scaled down to 1.0 ml and 0.1 ml respectively. Instead of clotting milk sample in 100 seconds the concentration of the enzyme was adjusted to klot milk in 600 ± 10 seconds in order to minisize personal error. Milk-clotting unit used in this study was as defined by Berridge (1945). One clotting unit is the amount of enzyme that clots 10 ml of milk in 100 seconds.

(c) AFDT assay: Method of Ryle (1970) was followed to determine the rate of enzymatic hydrolysis of N-acetyl-L-phemylaletyl-L-diodotyrosins (AFDT), except for volumes of substrate and enzyme. To 2.45 ml of 0.01N HG1 was added 0.7 ml solution of 0.002M AFDT in 0.005M AsoBt. After equilibration to 30°C, 0.35 ml of enzyme was added. At intervals 0.4 ml of sample was pipetted out, mixed with 0.2 ml each of the cyanide acetate buffer and ninhydrin solution. After heating for 15 minutes on a boiling water tath, the samples were cooled, and 1 ml of 50% isopropanol solution was added and vigorously shaken. The whosobance of the colour so developed was recorded at 500 mm. The blank incubation mixture contained 0.005M NaON instead of AFDT.

(d) Inactivation of ribonuclease: The method of Bang-Jessen et al., 1964 was used for the enzymatic degradation of ribonuclease. Ribonuclease was dissolved in 0.05M citrate/HGl buffer, pH 2.6 prior to incubation with crude seal gestric processes and the isocharyman A and B. For porcine pepain and seal processes C, ribonuclease was dissolved in

0.01M HCI and for calf chymosin it was dissolved in 0.05M citrate/HCl pH 3.5. The incubation of ribonuclease with proteases was carried out for 2 h instead of 40 minutes and the digestion stopped by adding 0.25 ml of 1.5M sodium phosphate pH 7.0. The residual ribonuclease was diluted by adding 14 ml of water. After 5 minutes 0.1 ml of this diluted mixture containing the residual ribonuclease was added to a reaction mixture consisting of 0.65 ml of 0.1M sodium acetate (pH 5.0). 0.5 ml of 1% RNA in the same buffer. Incubation was carried out at 25°C. After 24 minutes 0.25 ml of 0.75% uranyl acetate in 24% perchloric acid was added to stop the reaction. After cooling on ice for 30 minutes, the samples were centrifuged at 3000xg for 30 minutes at room temperature. Thirty ul of the supernatant were added to 1.0 ml water in a 1 cm quartz cuvette. After thorough mixing the absorbance at 260 nm was noted. To determine any possible ribonuclease in the protease samples the latter were incubated with respective . buffers instead of ribonuclease. To determine A260nm-absorbing materials in RNA which are soluble in uranyl acetate, a blank reaction was set up with 0.75 ml of sodium acetate and 0.5 ml of 1% RNA. After incubation at 25°C for 24 minutes 0.25 ml of uranyl acetate was added and the absorbance at 260 nm of the supernatant was determined as described above.

(e) Determination of the dematuration of proteases by urea: One volume of the respective protease, dissolved in 0.018 MCI was incubated with two volumes of 9M urea in 0.15M sodium accetate, pH 5.66. The final pH of the mixture was 5.40. At intervals 30 µl aliquota were withdrawn in duplicate and added to 0.8 ml of 22 hemoglobin in 0.06M.

VI. Determination of the pH optima for pensin activity

in 30 minutes under the assay conditions.

A 2% solution of hemoglobin in citrate buffer pH 1.2 was used to titrate's all volumes of another 2% hemoglobin solution in sodium acetate buffer pH 5.0 to make hemoglobin solutions of various pH, values between pH 1.2 and 5.0. Both the buffers used in making the stock solutions had the same ionic strength (p=0.05). A 0.8 ml volume of each hemoglobin solution was then incubated with 30 µl of enzyma. containing 0.4 pepsin unit at 30°C for 60 minutes, and the reaction stopped by adding 1.0 ml of 61 TCA. The absorbances at 280 mm of the supernatants were determined as described to Section V(a).

VII. Determination of the influence of pH on the stability of the processes.

Pive hundred microlitres of 10 mM acdium acetata pH 5.3 containing 24-28 pepsin units of the respective processes were mixed with an equal volume of a series of buffers of ionic strength 0.1, one drop of toluene was added to each of the mixtures, the tubes were stoppered and incubated at 25°C. After 24 h 50 µl aliquots of the mixtures were assayed for residual pepsin activity as described above (Section V(a)) except that the 27 hemoglobin solution was prepared in 0.2%

citrate-HC1 buffer pH 1.8.

VIII. Isoelectric focusing

Preparative isoelectric focusing of the crude activated enzyme on Sephadex IEF was done on a Pharmacia flat-bed isoelectric focusing apparatus following the manufacturer's instruction. Pharmalyte 3-i0 was used as pil gradient. The gal was prefocused for 30 minutes at 8 watt setting of the Pharmacia electrophoresis constant power supply model ECPS 3090/150. After applying the sample, the gal was focused for 9 h at 30-watt setting. The gal was cooled by circulating water maintained at 4-8°C. The gal was then divided into 21 equal sections accoped off the flat-bed into test tubes, and the pil of the fractions was measured. The gal fractions were then placed in a hypoderade syringe plugged with glass wool. The protein was then eluted from the gal. Fifty microliter aliquots were then used to detect the proteolytic activity by incubating with 0.8 ml hemoglobin solution, for 60 min. Fractions with prognedlytic activity were pooled into 4 portions to determine QUPU.

Analytical polyacylande gel isociectic focusing of purified symogens A and C was carried out using Pharmalyte 3-10 as the pH gradient. The gel was cast in a 115 x 230 mm glass plate following the manufacturer's suggestion. Samples (65-20 µl)—were applied directly on the surface of the gel. The focusing was carried out for 1 h at 3000V and 20% setting using the power supply described above. After focusing, the pH of the gel was measured using a surface electrode. The protein was stained and destained by following the manufacturer's instruction.

IX. Electrophoresis

Polyacrylamide gel electrophoresis at pH 8.3 was performed following the method of Davis (1964). Sodium dodesyl sulphate (STS) polyacrylamide gel electrophoresis in 7.5% separating gel as and 3% stacking gel was done according to Lesmii (1970). Unless otherwise indicated, the gels were ent off-at the marker-dye after electrophoresis. The gels were stained for 30 minutes (à 0.1% Coomassie blue in 50% methanol, and 10% acetic acid and descated by diffusion in a solution of 5% methanol and 10% acetic acid. Bovine serum albumin (196 65,000 dairons), chicken egg albumin (196 33,000 dairons) assess prelactin (198 23,800 dairons) and cytechrome C (66 12,700 dairons) were used as standard molecular weight markers.

X. Amino acid analysis

Amino acid analysis was performed in a Beckman Model 121MS amino acid analyser using the method described in the Beckman bulletin 121M-TB-013.

XI. Determination of nitrogen and protein estimation

Total and mongrotein nitrogen in whey were determined following the method of Rowland (1938). Total and comprotein Matrogen products of anymatic casein digestion were determined according to the method of Lang (1938).

Protein estimation was carried out following the method of Hartree (1972).

XII. Preparation of Acid Casein

Pastcurfaed skim milk (Sunshine Dairy, St. John's, Nfid.) was used for the isolation of acid casein at pH 4.6 following the method of Fox and Guiney (1972). The product was freeze-dried and stored at 4°C until used.

XIII. Cheddar Cheese

(a) Preparation: Cheddar cheese was prepared using pastsurfased whole milk (Sunshine Dairy, St. John's, Nfild.) by following the method of Kostkowski (1978), adapted for a small volume of milk. Eighteen liters of milk were used for each of the coagulants - calf rennet, activated DEAE peak A, crude SGP, and <u>Mucor michel</u> protease (Fromase 100). DEAE peak A material and crude zymogen from two week old seal were dialysed against 2x21 of 20 mM sodium phosphate pH 7.2 before activation at pH 2.0, in order to remove penicillin and streptomycin used during the extraction of the enzymes (Materials and Nathods, Section IV(a)).

The cheese from an 18L batch was pressed in one-lot using a press constructed with a plastic barrel, 7 inches in dismater following the method of Radke (1974). After pressing for 18 h in this barrel the cheese block was split into a smaller blocks and pressed in perforated square plastic boxes (12 x 12 cm and 5 cm high) lined with cheesecloth. Three bottles each containing 16 gallons of water were used to press these blocks for another 8 h. The cheese blocks (slightly more than a pound each) were vacuum-packed and cured at 870 in a Hotpack temperature-humidity chamber (Hotpack Canada Ltd., Ontario).

- (b) Chemical analyses: Protein, fat and mosture in cheese were determined following the methods of A.O.A.C. (1970). pH of cheese was determined by mixing 5 g of cheese with 5 ml of defonised water using a glass efectrode.
- (c) Sensory evaluation of cheese: Cheeses were evaluated by preference test (Larmond, 1977) at the taste panel room in the Department of Pisheries and Oceane, St. John's. Thirty untrained panelists were provided with number-coded cheese samples (approx. 3x1x2 cm), and general information about Cheddar cheese. Panelists were asked to rate the samples as to overall preference on a numerical scale ranging from 9 (like extremely) to 1 (dislike extremely). The data were evaluated by analysis of variance, (Larmond, 1977).

RESULTS AND DISCUSSION

I. _ Extraction of crude enzyme

Crude seal gastric protease (SGP) used in the preliminary studies (See Appendices A-E) was prepared following the method of Qadri et al., (1962) modified as described in Materials and Methods (Section III). Approximately 4.5 clotting units were obtained pergram of whole stomach powder from a 2-week-old seal pup collected in 1979 (Section I. Materials and Methods). It was later found that extraction with 20 mM phosphate pH 7.2 (buffer A. Materials and Methods, Section III) for 12 h yielded 26.3 clotting units per gram of the same stomach powder. The latter method extracts the zymogens while the method of Qadri et al., (1962) extracts the activated gastric proteases. The lower yield obtained following the method of Oadri et.al.. may be attributed to autodisestion of the protesses. Based on this observation, phosphate buffer was used for extraction in subsequent studies. Also, instead of the whole stomach the mucosal layer was used in order to have higher specific activity of the starting materials. The stomachs used for this purpose were from both twoweek-(approximately) and two-year-old (approximately) harp seals collected in 1980.

The average yield of activated rymagen extracted from mucosal powder was 191.5 clotting units/g for two-week-old pups and 262.5 clotting units/g for two-year-old seals. These averages are based on measurements on two stomachs from sdults and three stomachs from pups. The yield of zymagen from whole stomach collected in 1979 was much less (26.3 clotting units/g whole stomach powder) than from the

stomachs collected in 1980, because (1) only the nucosa was extracted from the stomachs collected in 1980 whereas in the case of the stomachs collected in 1979 the freeze-dried powder of whole stomach was extracted, and (2) the 1980 stomachs were empty but the 1979 stomach was full of curdled milk, indicating that some of the zymogens had already been secreted. The fact that the 1980 stomachs were shipped chilled on ice while the 1979 stomach was shipped frozen and the possible difference between individuals of the same species may also have contributed to the differences observed.

II. Proteolytic activity

Hemoglobin solution (2% in 0.06% HCL, pH 1.8) was used routinely as substrate to monitor the proteclytic activity of the enzymes at various stages of purification. Zymogens were activated by incubating at pH 2.0 for 30 minutes at 25°C (Results and Discussion, Section V). However, this preactivation step was omitted when aliquists of individual fractions from chromategraphic column clustes were assayed to monitor their proteclytic activities. The rate of hemoglobin hydrolysis at pH 1.8 and 30°C was linear with time up to 120 minutes when the enzyme concentration was 1.9 pepsin units per ml of incubation mixture (Figure 1). In a 30 minute incubation under this condition the rate of hydrolysis was also linear as a function of the concentration of the crude enzyme (Figure 1b).

III. Ion exchange chromatography

As shown in Figures 2a and 2b, crude preparations of zymogens from the mucosa of young and adult seal stomachs separated into 4

Figure 1. Proteolytic activity as functions of time and enzyme concentration.

- (4) A 27 hemoglobin solution in 0.06H HCl was incubated with 1.9 pepsinunits of crude extract per all of reaction mixture at 30°C. At intervals indicated duplicate samples of 0.9 ml were withdrawn and the proteolytic activity determined as described in Naterials and Nethods.
- (b) A 2% mesoglobin solution in 0.06M HGl was incubated with the indicated volumes of a dilute solution of crude enzyme so that the final volume of reaction mixture was 0.9 ml. After 30 minutes of incubation at 30°C, the raction was stopped by adding TCA as described in Materials and Methods and extent of proteolysis determined. Absorbances due to control were subtracted for each enzyme concentration.

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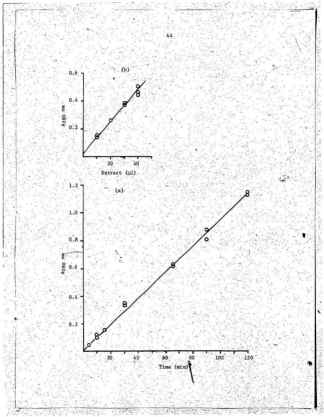
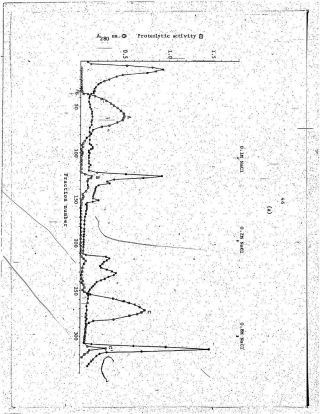
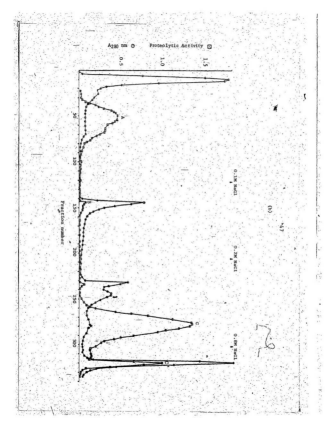


Figure 2. Ion exchange chromatography of extracks from gastric mucosa of harp seal.

Forty ml of homogenate from 2g of dry mucosa were applied to a 2.5x 30 cm DEAE Sephadex A-50 and eluted with buffer A untill—the A280mm and the proteolytic activity approached zero. The column WHE Then eluted stepwise addition of MaCl to buffer A as indicated. Eventy ulaliquots from alternate fractions were incubated with hemoglobin at 30°C for 3h to monitor potential proteolytic activity. Flow rate 20 ml/h, fraction size 5.0 ml. (a) homogenate from a 2-week-old seal,(b) homogenate from a 2-year-old seal. The symogens are activated in the test procedure for proteolytic activity at pH 1.8.

9, Absorbance at 280mm; [D], enzyme activity, absorbance of TCA soluble fraction at 280mm.





pepsingern components, A. B. C and C' of which B and C' were minor and of low specific activity. No further purification of the components B and C' was attempted. The relative proportion of zymogen A as pepsin activity from stomachs of two week-old pups was higher than that from adult stomach. As shown in Table 1 the activity of peak A was 44.5% of the total recovered from one stomach and 51% in the other. The proportion of peak C was 48.5% and 37.8%, respectively. These estimations were based on the areas of the peaks in the activity profiles shown in Figures 2a and 2b. In the adult-stomach miccona peak A activity represented 32.5% in one run and 27% in the other. The proportion of peak C activity in those two samples were 54.8% and 59.1%, respectively. These two samples were from the same antimal. Compared with peaks A and C, the yields of peaks B and C' were highly variable in both the young-and the adult-mucosal gamples.

Nultiple isoeanymagiforms of both chymosin and pepsin have been reported in other species as well. 'Occurrence of 4 caif chymosins from their corresponding symogens has been reported by Asato and Rand (1977). Amer et al., (1980) separated "adult bovine remeet" into two components. The major component was pepsia-like with a CUPPU ratio of 0.032 and the minor component chymosin-like with a CUPPU ratio of 0.639. However, the authors did not define the units of milk-clotting and proteolytic activities. Foltmann at al., (1978) showed that the ratio of milk clotting activity to general proteolytic activity of pigler chymosin was 30 and 70 times as high as those for pepsins from adult funds and pylorous respectively and 4 times as such as that for call chymosin.

49.

Table 1. Relative proportions of various DEAE peaks from mucosal homogenates of the stomachs of young and adult harp seals

Age	CU1/PU2 of Crude	Relativ	e Proportio	ns ³ of Pea	k X
	homogenate	Α	В	c	c'
2 week	4 -0.024	44.5	4.1	48.5	2.9
1	0.028	51.0	5.0	37.8	6.2
2 year	rs ⁵ 0.018	32.5	3.7	54.8	9.0
-10 2	0.016	27.0	1.8	59.1	2.1

- Notes: 1. One C.U. (clotting unit) is the amount of enzyme that will clot 10 ml of 12% skim milk in 100 seconds.
 - One pepsin unit (PU) is the amount of enzyme that produces an increase in the absorbance at 280 nm of TCA-soluble hydrolysis products of 0.0084 per minute under the conditions described in text.
 - 3. Based on pepsin activity (see fig. 2a, b).
 - 4. Stomachs from two different animals.
 - 5. Samples from the same stomach.

- IV. Purifications of zymogens A and C
- (a). Gel filtration: The two major symogens A and C were purified by molecular-sieve chromatography using Sephadox C-100 as described in Materials and Methods (Section IV(c)). Figure 3 shows the protein profile and the pepsin activities of the chromatographic fractions & The column was callbrated with standard molecular weight markers. The peak paptic activity was eluted in a position corresponding to a solecular weight of 26,300 ± 1540° daltons for tymogen A based on 5 trials. The molecular weight of zymogen C was estimated at 37,000 ± 1075 daltons from two trials.
- (b) Purification of zymogen A by affinity chromatography: The zymogen representing peak A on the DEAE-Sephadex column was also purified by chromatography with carbobenzoxy-D-phenylalanyl-triethylenetetramine-Sepharose (Z-D-phe-T-Sepharose), as described in Materials and Methods (Section IV(d)). The major portion of the protein applied to the column was eluted unadsorbed and had very little proteclytic activity (Figure 4). Most of the proteclytic activity was eluted with the buffer containing guanidine-HCl. The active fractions were pooled, mixed with equal volume of 0.2M sodium phosphate (pH 3.2) in order to prevent the conversion of the zymogen to the active enzyme. This mixture was then dislysed against water at 4°C. The dislysate was then concentrated by ultrafiltration to 1.9 ml. The preparation so obtained had some of the zymogen activated as evident from a milkclotting activity of 2.0 clotting units/mg of protein. After activation at pH 2.0 the specific activity increased to 39.4 clotting units/mg (Table 2).

Chromatography of DEAE peaks on Sephadex G-100 column.

Fractions - from DEAE-Sephadex A-50 column with proteolytic activity were pooled, concentrated and 2 ml of the concentrated material were applied to a Sephadex G-100 column (88 x 1.6 cm) equilibrated with buffer B. The column was eluted with the same buffer in the ascending mode. Fraction size 2.7 ml. flow rate 5.0 ml/h. Fifty ul aliquots from alternate fractions were used to monitor potential. proteolytic activity. O, A280 of fractions; , Proteolytic activity, optical density change at 280nm.

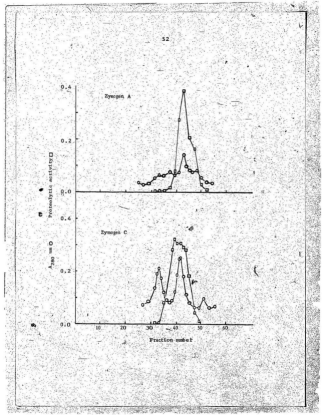


Figure 4. Affinity chromatography of Zymogen A on CSZ-D-phenylalanyl-TETA- Sephatose 4B.

Two ml. of concentrated material from DAE peak A were chromatographed on a column packed with carbobenary; —D-phanylalanyl-tristlylene-tetremine Sepharone (Z-D-Phe-T-Sepharoge) as described in Materials and Methoda. Buffer changes are indicated by arrow. Twenty_lalaquots from alternate fractions were incubated with hemoglobin solution for 60 minutes to monitor potential proteolytic activity. Flow rate 50 ml/h; fraction sizes 2.9 ml.m. Absorbance at 280mm of fractions, a proteolytic activity, change in absorbance at 280mm.

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igure 5. Time-course of activation of zymogens.

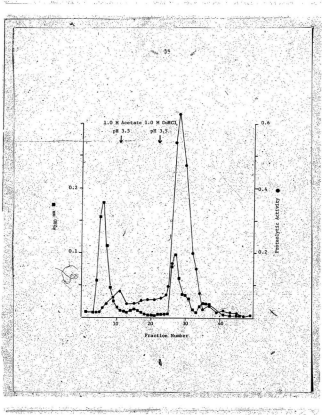
To 1.0 ml of buffer A containing Zymogen A, B, or C, 24 µl of 1M NCl were added to adjust the pH to 2.0 at 25°C. Immediately after adding NCl and at intervals indicated 20 µl aliquots were removed in duplicate to monitor all-clotting activity. A, Zymogen A; • Zymogen B, and A; Zymogen C.

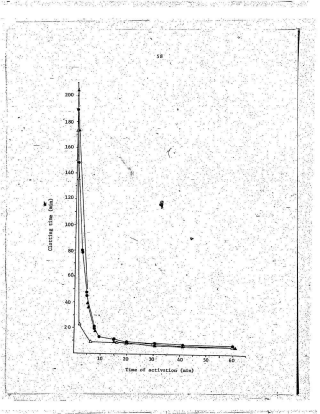
١.	Affinity			Chromatography Sephadex G-100	Peak C'	Peak C	Peak B	Peak A	Chromatography DEAE Sephadex	Crude su	Purifica
•	Affinity chromatography	C	A	graphy on G-100	C	C	B	Α	Chromatography on DEAE Sephadex	Crude supernatant	Purification step
1.9	ly .	7.7	11.4		4.1	6.2	4.4	6.0		46.0	(nl)
18.5		3.8	11.5		1.6	10.9	0.4	35:5	6	9.0	CU
250.2		633.2	153.5		410.0	1457.0	14.1	725.0		589.0	Activity/ml CU PU
35.2		29.3	130.5		6.6	67.6	1.8	213.0		414.0	CU
475.4		4875.6	1750.0		1681.0	9033.4	62.0	4350.0		414.0 27094.0	Total activity CU PU
39.4		5.2	31.8		0.2	2.6	0.1	10.8		1.5	CU PU
532.4		867.5	426.5		48.8	347.0	3.9	220.0		99.8	ty/mg PU
26.3		2.5	21.2		•	1.7		7.2			Purification CU PU
5.3		8.7	4.3		1	3.5	•	2.2		A A	PU
0.075		0.006	0.074		0.004	0.007	0.026	0.049	G	0.015	CU/PU

(c) Summary of purification procedure: Purification steps of the zymogens from the mucosal powder of a two year old harp seal is summarized in Table 2. Zymogen A was purified 21-fold by gel filtration on a Sephadex G-100 column with an increase in its milk clotting activity from 1.5 clotting units/mg of the crude zymogen to 31.5 clotting units/mg. By affinity chromatography of DEAE-peak / a further purification of the milk clotting activity was achieved (39.4 clotting units/mg). Approximately 9 fold purification of zymogen C was achieved by gel filtration; the pepsin activity increased from 99.8 pepsin units/mg of crude extract to 867.5 pepsin units/mg of protein. As a result of purification the ratio of milk clotting activity to proteolytic activity increased in the case of zymogen A compared with this ratio in the crude extract. In the case of zymogen C this ratio decreased by a factor of 4. Under the assay conditions (Sections V(a), V(b)) crystalline calf chymosin had milk clotting and proteclytic activities of 21.4 and 126 units. respectively, per mg of protein based on an average of two determinations.

V. Activation of the 2ymogens

The time course of the activation of symogens A, B and C at 25°C is shown in Figure 5. The pH of an appropriate dilution of the stock solution of each symogen was adjusted to 210 by adding a predeterance delime of IM NCI. The conversion of the symogens to the respective enzymes was monitored by milk clotting assay at pH 0.3 (Materials and Methods, Section V(B)). As shown in the figure the activation of symogen A and B was apparently complete in 15 minutes and that of





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zymogen C in 6 minutes. These observations served as a guideline to activate the zymogens prior to routine assays. Routinely the activation process was carried out for 30 minutes.

VI. Properties of the enzymes

(a) CU/FU ratios: The ratios of milk clotting activity to proteolytic activity of the crude isoenzyms forms A, B, C and C' are shown
in Tables 3 and 4. The ratios for enzyme A were 6 to 9 fold higher
than that of the other major isoenzyme C. Bowever, as shown in
Table 2, the ratio for the purified rymogen A was shout 12 times as
high as that of purified rymogen C. The CU; FU ratios for isoenzyme B
were similar to those of A and the ratios for isoenzyms C' are similar
to those for isoenzyms C irrespective of the age of the similar.

The CU:FU ratios for the total extract from the two-week-old pups were slightly higher than those for the crude enzymes from the adult seals as shown in Table 1. This is consistent with the higher proportion of isoenizyme A in the two-week-old pup stomachs. The CU:FU ratio for protease A purified by gel filtration and affinity chromatography was higher (Table 2) than that of the crude isoenzyme A (Tables 3 and 4). This suggests that the purification process removed some material(s) with low CU:FU ratio. It is also possible that some other momentymatic factors which selectively affect the milk clotting activity of enzyme A were removed by the purification.

(b) Milk clotting as a function of pR: Milk clotting times for active isoenzymes A and C are shown in Figure 6a and that of isoenzyme R is shown in Figure 6b. Clotting times for porcime pepsin and calf

Table 3 Relative rates of milk-clotting and hemoglobin hydrolysis by various DEAE peaks from homogenate of 2-week-old harp seal gastric mucosa

Peak	Pepsin Unit/mg	Milk-Clotting Unit/mg ²	CU PU
Experiment 1 A	278.0	14.2	0.051
В	13.08	0.69	0.053
	223.8	1.83	0.008
c'	1.9	0.02	0.007
		51 July 15	
Experiment 2 ³ A.	104.6	7.48	0.072
В.	6.6	0.36	0.056
c	118.9	1.01	0.009
c'	1.6	0.01	0.006
A Darling to the second		Tall Bereiter 1	

concentrations.

Average of at least two determinations.

The freeze-dried powder was stored in a -20°C freezer for about a year before the extraction. Some activity may have been lost during storage.

Relative rates of milk-clotting and hemoglobin hydrolysis by DEAE peaks from homogenates of 2-year-old harp seal gastric mucosa

Pe	ak Pepsin Unit/mg1	Milk-Clotting Unit/mg ²	CU
Experiment 1 A	219.6 -	10.5 /	0.048
ъ В	3.9	0.12	0.031
	346.9	3.09	0.009
	48.8	1.6	0.004
			4
Experiment 2 A	228.0	12.6	0.055
В	9.70	.46	0.047
c	230.2	1.71	0.007
c c	46.5	0.28	0.006

Pepsin activity was determined at 3 to 4 different concentrations of each sample in duplicate.

Average of at least two determinations.

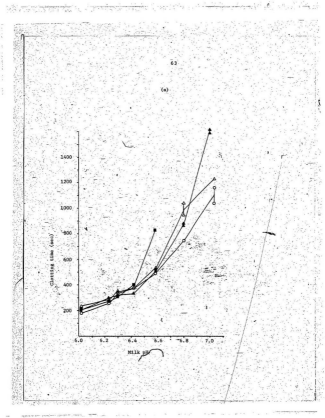
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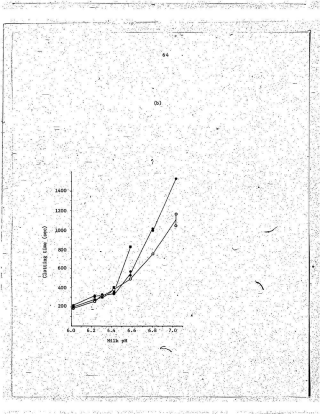
Figure 6. Milk-clotting activity as a function of pH.

A 12% suspension of skin milk in 0.01M CaCl₂ was adjusted to pH 6.0 to 7.02 with 1M BCl or 1M NaOH. The concentration of each enzyme was adjusted so that a 50 μ l volume will clot milk at pH 6.3 in 5 minutes \pm 10 seconds.

(a) Milk-clotting activity of protesses A, and C compared with those of calf chymosin and porcine pepsin.

(b) Protesse 3 compared with porcine pepsin and calf chymosin. O, calf chymosin; m; porcine pepsin; A, protesse A; Δ, protesse C; φ, protesse B.





chymosin are also included for comparison. The three seal gestric proteases exhibited a pH dependency for clotting times similar to that for celf chymosin and different from that of porcine pepsin.

Anifantakis and Green (1980) observed that the milk clotting activities of lamb and kid rennets increased less with decrease in pg than that of calf rennet.

(c) Action of proteases on ribonuclesse: The inactivation of border pancreatic ribonuclesse, by the proteolytic engages was performed as described in Materials and Methods (Section V(d)). Incubation with neal proteases A, B and the crude engage was carried out at pH 2.6 based on the gi optimum of crude sual gastric proteases (Appendix D). Incubation with perfine pepsin and seal processe C was carried out at pH 2.0 and that with calf chymosin was performed at pH 3.5. As shown in Table 5 there was no detectable inactivation of ribonuclesse by the proteases A, B or calf chymosin star. 2 hours of incubation whereas protease C caused an average of 187 inactivation. Forcine pepsin at the potencies of 1.6 and 5.0 pepsin units effected 55% and 84% inactivation, rispectively.

Inactivation of ribonuclease by crede SGP was also examined. Figure 7 shows the percentage losses of ribonuclease activity after 2 h incubation with various concentrations of crude SGF and of porcine pepsin at 30°C. The figure shows that with 76 pepsin units of enzyme, seal gestric proteins inactivated only 22% of the ribbouclease activity whereas porcine pepsin with the same potency inactivated 79% of the Thomsclease.

- Protease	Potency	pH of incubation	7 inactivation
	(Pepsin unit)		
DEAE Peak A	11.8	-2.6 ²	0
Peak B	1.6	2.62	.0
Peak C	6.1	2.0	18
Porcine pepsin	1.6	2.0	. 55
	5.8	2.0	84
Calf chymosin	6.0	3,5	0 1

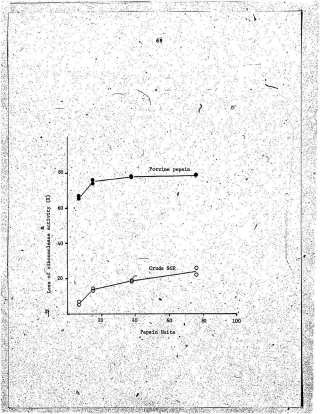
Average of dunlicate measurements

Based on the pH optimum for the pepsin activity of crude SGP.

Figure 7. Inactivation of ribonuclease by crude SGP.

As active proporation of a crude extract from adult-seal nuccess was diluted so that 0.1 hl would contain the popular units indicated in the figure. After 2 hours of incubation with thomoclease the residual ribonuclease activity was determined as described in Materials and Mathematical Seal of the Contained as described in Materials and Mathematical Seal of the 2.0 and that with crude SEP at pH 2.6.

O, crude SGP; , porcine pepsin.



(d) Hydrolysis of APDT: Hydrolysis of the pepsin substrate N-acetyl-L-phenyl-alanyl-L-dijodotyrosine (APDT) by protease A was very low. As shown in Table 6 the activity of protease A expressed as the ratio of APDT unit/pepsin unit of the enzyme was similar to that of calf chymosin, but much lower than those of protease C and porcine pepsin. The activity of protesse B on this substrate was negligible. - Very high concentrations of proteases A and calf chymosin were used in order to obtain detectable increases in absorbance at 570nm. As a result of this, there was high background absorbance and during the initial 2h incubation the increase in A570nm was very low and reproducibility was poor. However, there was appreciable increase in absorbance after 4 h and 13 h of incubation. The relatively low activity of protease A and calf chymosin on APDT shown in this study is consistent with reports that chymosin is much les sensitive to certain synthetic substrates than to K-casein and milk (Martin et al., 1981). These authors also showed that bovine pepsin A was 55 fold as active on the hexapeptide Leu-Ser-Phe(NO2)-Nle-Ala-Leu-OMe than calf chymosin when equal milk clotting units of the two enzymes were used.

(a) Clocking activity as a function of temperature: Clocking activity of calf chymosia and protease A as a function of temperature is shown in Figure 8a. Furified peak A protease had milk-clotking times close to those of calf chymosia in the temperature range 15° to 45°C. At 50°C, the clotking time for protease A was higher than that for calf chymosia.

Arrhenius plots for the milk-clotting activities at various

Table 6. Hydrolysis of N-acetyl-L-phenyl alanyl-L-difodotyrosine

Protease Potency	Incubation	Absorbance (570nm)	
(PU)	time (h) 2 -	4 13	AU ¹ /PUx1
A 11.8	0.06 <u>+</u> 0.06	0.063+0.007 0.222+0.009	2.82
B 1.6	0.055 <u>+</u> 0.055	-0.02 <u>+</u> 0.02 0.08 <u>+</u> 0.08	_3
C 6.1	0.015±0.015	0.172+0.023 0.275+0.028	14.9
Porcine			
pepsin 1.6	0.065 <u>+</u> 0.001	0.106+0.005 0.171+0.001	42.9
Calf			44.40
chymosin 8.8	0.017 <u>+</u> 0.04	0.073+0.011 0.297+0.015	4.4

- One AFDT units (AD) is the amount of enuyme that causes an absorbance increase at 570m of 0.079/min.
 Because of high background absorbance and low activity readings after in were not reliable for A. C. and calf chymosin. Therefore values of Agyons after the were used for calculation of AU in these cases.
- Average of duplicate readings + range after substracting the Asyn of control samples.

Could not ascertain because of low APDT activity.

temperatures for calf chymosin and protesse A are shown in Figure 8b. Activation energies of 13.1 and 15.6 Kcal/mole were calculated for chymosin and protesse A respectively. The calculation was based on the slopes of an Arrhenius plot computed from linear regression of the points in the temperature range 15° to 35°C. Above that range data points were scattered in the case of protesse A. In the case of calf chymosin, there were breaks in the Arrhenius plot, one at 20°C and the other at 35°C. This may relate to the heterogeneity of the chymosin as will be shown later (see Figure 17). Milk clotting activities of chicken pepsin and ealf chymosin as a function of temperature reported by Gordin and Rosenthal (1978) were also plotted flere for comparison. The energies of activation for chicken pepsin and calf remnet calculated from Gordin and Rosenthal's data (1978) are 15.0 and 11.5 Kcal/mole respectively.

It should be noted that the activation energy for the milk clotting activity of an enzyme is a complex property because milk clotting is a two-stage process and the temperature dependence of the nonenzymatic step is different from that of the enzymatic step (Dalgleich, personal communication).

(f) Influence of Ca⁺⁺ on clotting activity: Milk clotting activity of processe A was found equally sensitive to added calcium chloride concentration as calf chymosin (Figure 9). Highest activity was found at 0.02M Ca⁺⁺, for both the enzymes; at higher concentrations activity was inhibited. At 0.IM calcium chloride concentration the clotting times were similar to those without added calcium chloride.

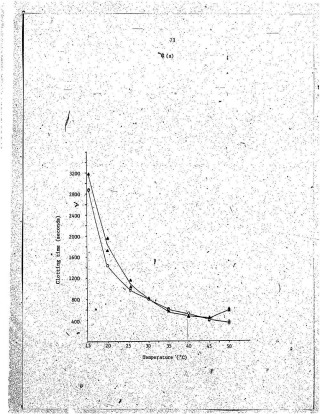
The stimulation of milk clotting activity of coagulants by midded

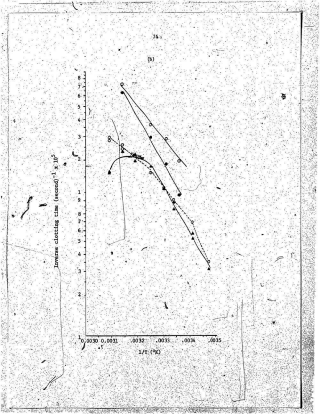
Figure 8. Effect of temperature on clotting activity.

Concentrations of enzymes were so adjusted that both calf chymnein and
Enzyme A had equal milk clotting activity at 30°C in Berridge substrate pH
6.3.

- (a) clotting times as a function of temperature for calf chymosin (O), protesse A (Δ).
- (b) Arthentia plots calculated from this experiment, O—O calf chymosin; A, protease A. Arthentia plots calculated from data reported by Gordin and Rosenthal (1978) are shown as O—O calf chymosin; ——

 chicken pepsin.





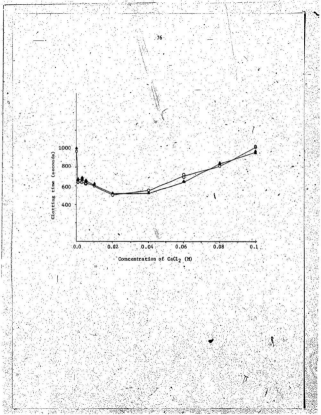
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Figure 9. Influence of calcium chloride concentration on milk-clotting activity.

Suspensions of 12% skin milk were made at various levels of calcius chloride concentrations at p8 6.3. Concentration of each of the enzymes was adjusted so that they had equal clotting time at 0.01% CaCl₂ concentration.

O; Calf.chymosis; A, protease A.

O, call chymosin; A, protease A



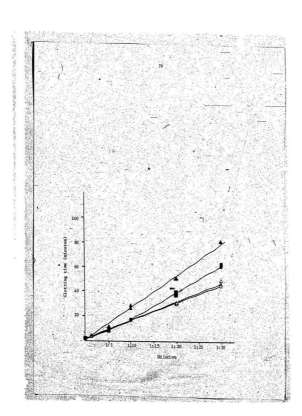
calcium observed in this atudy is in agreement with reports of Gordin and Rosenthal (1978) and of Tsugo and Yamauchi (1959). However these authors observed only small decreases in the clotting activities of calf rennet and other coagulants at high CaCl, concentration compared with the decrease in clotting activities of protease A and calf chymosin in this study. These discrepancies are probably due to differences in the conditions of milk-clotting assay. The drop in milk pH that occurs on the addition of CaCl, was not adjusted in the studies reported by these authors. The pH of reconstituted skim milk in 0.01M CaCl, (Berridge subs(rate) used routinely in this study was 6.3 and its oH dropped to 5.5 on addition of 0.1M CaCl2. The delay. in the clotting times due to the addition of CaClo was probably offset by the lower pH of the milk used by Gordin and Rosenthal (1978). However, it is possible that part of the inhibition of milk-clotting activities observed in this study was due to irreversible change in the casein micelles as a result of pH adjustment as suggested by Gordin and Rosenthal. Amer et al. (1980) reported slight stimulation of milk-clotting activity of boying chymosin and pensin and inhibition of their proteolytic activity by CaCl, and NaCl.

(g) Clotting time as a function of enzyme concentration: Milk-clotting times increased linearly with dilution of both protesse A and calf chysosin as shown. In Figure 10, but the increase was higher in the case of protesse A than the increase in clotting times for calf chysosin at the same dilution. Portice paparin also had a steeper slope than that of calf chysosin whereas protesse C had similar activity with dilution as calf chysosin.

... 78

Figure 10. Effect of enzyme dilution on the clotting activity.

Enzyme concentration was adjusted to clot i al of Serridge substrate in 100 ± 10 seconds. This stock enzyme was then diluted with loam secture at pH 5.3 and immediately 50 µl mixed with l ml of Berridge another the qualification of 30°C, Calf chymosin O; Frofense A&; Frotense CA; and Forcine peptin ...



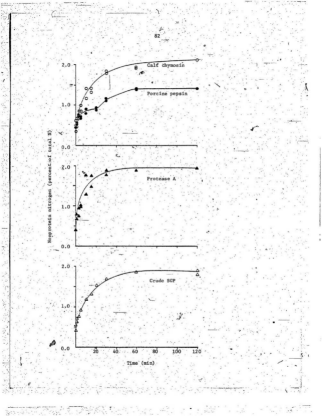
(h) Hydrolysis of casein: The extent of hydrolysis of casein by the coagulants was examined at the enzyme concentrations ordinarily used for cheesemaking. Forty al of a 2% casein solution pH 6.1 was incubated with 2.0 ml of the enzyme at 30°C. The concentrations of the enzymes were standardized so that 2.0 ml volume of the enzymes would clot 40 ml of milk in approximately 20 minutes. At intervals, suples were withdrawn and nonprotein nitrogen determined as described in the legend to Figure 11. As shown in the figure the release of nonprotein trogen due to hydrolysis by protease A and crude SGP with time was similar to that of calf chymosin. In both the cases no further hydrolysis was detectable beyond the 1.7% NPN expressed as percentage of total nitrogen, taking into account the 0.3% NPN already present in the samples at 0 time. With porcine pepsin however, the release of nonprotein nitrogen leveled off at 1%. The release of NPN from casein by chymosin (Alais et al., 1953; Wake, 1959) has been correlated to the action of chymosin specifically on the k-casein fraction which comprises 10-20% of whole casein. A mixture of macropeptides is released as a result (Wake, 1959; Alais and Jolles, 1961) part of which is soluble in 12% TCA. Clotting of milk by bovine, porcine, and chicken pepsins (Green, 1972) and microbial protease (Ohmiya et al., 1979) also involves the same mechanism. However, with non-rennet milk coagulants there is a rapid release of NPN followed by small but gradual increase in NPN indicating nonspecific proteolysis as with chicken pepsin (Gordin and Rosenthal, 1978) and Mucor pulilus protease (Pahkala and Antila, 1980)

⁽i) Influence of pH on hemoglobin hydrolysis: Effect of pH on the

Figure Hydrolysis of casein by proteolytic enzymes.

Forty mi of a 2% solution of casein, pi adjusted to 6.1, was equilibrated at 30°C. To this was added 0.2 pl entrue calculated so clot 40 mi of milk in 10 minutes. At intervals shown in the figure 2 ml samples some withdrawn in duplicate, precipitated by adding 2 ml of 24% TGA.

Contribuged at 3000xg for 10 minutes. One mi aliquet of the superparant was subjected to 8 detention to following the method of Lang (1938). The mosprotein microgen was expressed as percentages of total N. The intital NPH presset in the samples was determined by taking as aliquet before adding the enzymes.



rate of hemoglobin hydrolysis at 30°C by processes A and C arg shown in Figure 12. Peak C pracease had an optimum pil at 2.4 and processe K showed activity over a wide pH range between 2.2 and 3.5. In this respect processe A resembles call chymosin (Berridge, 1945) and gastricsin (Ward et al., 1978).

- h) Influence of pH on enzyme stability: The stability of the active enzymes A and C was compared in buffers of various pH values having an ionic scrength 0.05 after 24 hours of incubation at 25°C. Af pH values below 7.0 procease A lost more bepeatn activity than did the other proteases (Pig. 13). At pH 7.0.4 was much more stable than porcine pepsits and less stable than talf chymosin and protease C. These observations support those summarized fy Figures 6a, 6b. In the acidity pH range all the proceases lost more activity at their pH option for hemoglobin hydrolysis than at any other pH values presumably because of autodigestion (Foltmann, 1959). All the proceases showed minimum loss of activity between pH 5.0 and 6.0.
- (k) Denaturation of proteases by urea: The denaturation of protease: A and C in 6M urea at pH 5.40 and 37°C was compared with that of calf chymosin, and porcine pepsin. As shown in Figure 14, after 90-minute incubation both protease A and calf chymosin were rapidly denatured and lost about 90% of their initial pepsin activity. The initial rate of denaturation of protease A was considerably faster than that of the other proteases. Seal protease C and porcine pepsin lost 62% and 56% of their original activity respectively, in 90 minutes. These results agree with those of Chesseman (1965) who observed only 13% inactivation.

Figure 12.Effect of substrate pH on proteolytic activity.

Hemoglobin solutions of the pH values indicated in the figure were prepared as described in Materials and Methods and incubated at 30°C for 60 minutes with 20 ul of each enzyme containing 0.4 pepsin unit. The ionic strength of the buffers was 0.05.

▲, Protease A; △, protease C.

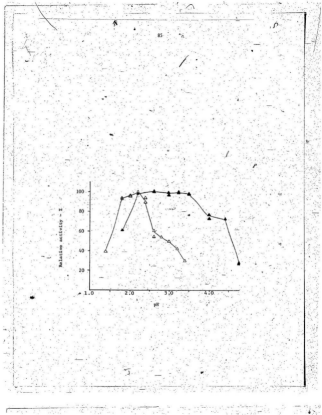


Figure 13. Stability of the proteases with respect to pH.

To 0.5 ml of enzyme solutions 0.5 ml of buffers were added to give the final pil values indicated in the Figure. The mixtures were incubated at 25°C. After 24h residinal proteolytic activities were determined as departed in Naternials and Methods. O. Citrate; O. acetate; and A. phosphate buffers.

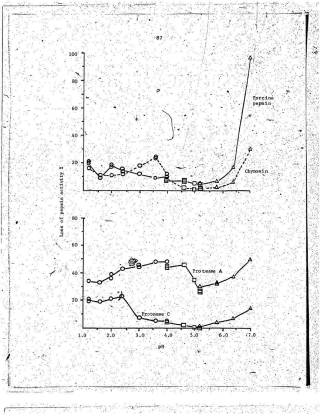
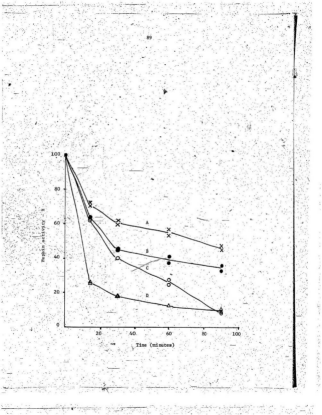


Figure 14. Inactivation of proteases in 6M urea.

To 0.4ml of 9M wrea in 0.15M modium acetate equilibrated at 37°C, 0.2 ml solution of the enzymes was added, so that the final concentration of urea was 6M at pH 5.4. The mixture was then incubated at 37°C and at indicated intervals 30 at aliquots were withdram and the residual hemoglokin hydrolytic activities determined as described in Materials and Methods.

A, protesse C; B, porcine pepsin; C, calf chymosin; and D, protesse A.

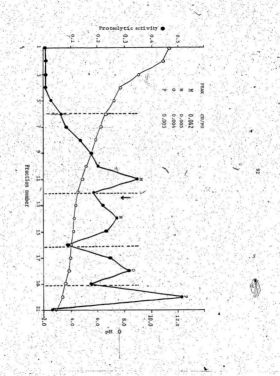


of pepsin whereas purified calf chymosin is lost 90% of its initial activity in 90 minutes. There were not gnough details about the experimental procedures in Cheeseman's report spart from PH, moliarity of urea and incubation temperature. Therefore the difference in the degree of inactivation of pepsin in this study from that found by Cheeseman may be attributed to any possible differencing in the experimental conditions, e.g. type and concentration of buffer and the purity of pepsin used.

(1) Isoelectric points: A crude preparation of proteases from an adult harp seal Stomach resolved into 4 isoenzymes by preparative iscelectric focusing on Sephadex IEF (Figure 15) . The iscelectric. points of the isoenzymes, M, N, O and P were 4.90, 4.31, 3.82 and 3.40 respectively. The result of isoelectric focusing agrees with the separation of four zymogens by ion-exchange chromatography on DEAE-Sephadex column. However, the relative peak sizes of the proteases M, N. O, and P are different from those of the zymogens A, B, C and C separated by DEAE-Sephadex chromatography. It is possible that the recoveries of zymogens B and C' from the DEAE-Sephadex column were less than those of the zymogens" A and C. It was not determined which isoenzyme separated by isoelectric focusing corresponds to which zymogen eluted from the DEAE-Sephadex column. However, isoenzyme M is probably the same as protease A since they had similar CU:PU ratio. The iscelectric point of calf chymosin was reported to be 4.70 ± 0.05 (Righetti et al., 1977) similar to that of the seal iscenzyme M. The iscelectric point of purified zymogen C was 4.02 by analytical polyacrylamide gel isoelectric focusing. No protein band

Figure 15. Preparative isoelectric focusing of crude SGP on sephadex IEF.

Five hundred all of crude protesses from an adult herp seal stocach were subjected to isoelectric focusing as described in Materials and Methods (section VIII). The sample was applied in the position indicated by the arrow. After electrofocusing the protein was cluted from the gel fractions and proteolytic activity determined as described in Nateriala and Methods. Fractions with proteolytic activity were divided as indicated by dotted lines and pooled to determine CU:PU.



was visible when iscelectric focusing was performed on purified symogen A under the same conditions. The protein (9 ug in 20 ul) in the mample probably separated into fractions below detectable level.

(m) Purity of the proteases: Zymogen A purified by gel filtration on a Sephadex G-100 column emerged as a single symmetrical peak by high performance liquid chromatography on a column of Beckman spherogel-TSK3000SW (Figure 16). However, on polyacrylamide gel electrophoresis at pH 8.3 (Davis, 1964) zymogen A appeared as a broad smear covering an area between Rf 0.17 to 0.45 (not shown) and a very faint hazy band between Rf O.46 and O.63, Zymogen C, however, was electrophoretically homogeneous with an Rf 0.713. After activation of zymogen C at pH 2.0 its mobility was lower than that of the zymogen (Rf 0.610), as shown in Figure 17. DEAE-peak A zymogen after purification by affinity chromatography on Z-D-Phe-T-Sepharose column was also electrophoretically heterogeneous (Figure 18). It moved as two closely moving broad bands with Rf's 0.39 (8) and 0.50 (C). There were also two smaller bands with Rf's 0.07 (A) and 0.63 (D). Since the affinity chromatography was done at oH 3.5 it was likely that the electrophoretic heterogeneity was due to slow conversion of zymogen A to protesse A and its possible autodisestion. To examine this possibility zymogen A was incubated at pH 2.0 for 30 minutes for complete activation. On electrophoresis of this protease A it was found that the band A disappeared and two smaller bands a, and as preceded the band B. The absorbance of band B was decreased to 1.1 from 1.3 optical density units on activation. The area under beak B for the zymogen decreased from 53% to 35% after activation. Peak Cincreased in area

Figure 16. Chromatography of Zymogen A on high-pressure liquid schromatography.

A 20 Al volume of symogen A (13 Ag grotein) purified on Sephades C-100 column was subjected to high-performance liquid chromatography on a Beckman model 33 HPLC system with a Bitachi variable wavelength detector and a 7.5 x 600 mm/sepherogel-TSK 3000 SW column (Beckman Instruments). The column was previously calibrated with borine serum albumin (65,000 daltoms), chicken egg albumin (43,000 daltoms), spoglobis (17,000 daltoms), lymosyme (14,500), and cytochrome C (12,700 daltoms). Flowerste, | ml/min.

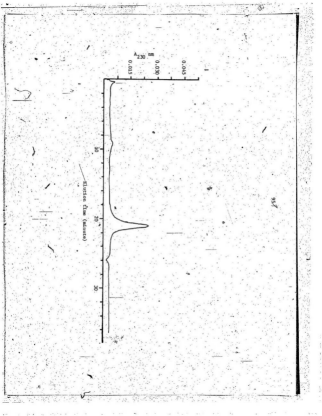


Figure 17. Electrophoresis of zymogens and enzymes:

Polyacrylamide gel electrophoreass was-carried out at pH 8.3 following the gethod of Davis (1964) at 125% constant-voltage setting of a Pharmacia ECFS 2000 power supply for 6 hours with cold-water circulation maintained at 4°C. The gels were cut off at dye front after electrophoreais, stained for 30 minutes in 0.1% Commassic blue and destained by diffusion in solution containing 5% methanol, and 10% acetic acid. 1, zymogen C, 2, protesse C, 3, porcine pepsin, 4, calf chymosin.

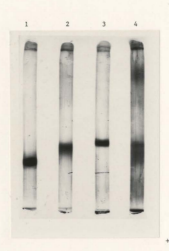
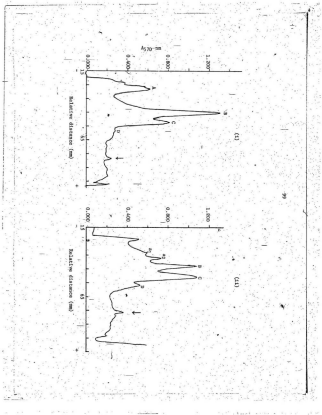


Figure 18. Gel scans of zymogen A and its active form.

Thirty us of symmogen A purified by affinity chromatography and its enzyme form obtained by activating symmogen A at pH 2.0 were subjected to polyacrylamide gel electrophoresis according to the method of Devis (1964). The gels were stained in Commansie blue. After destaining, the gels were scanned at 570 mm. (1) Symmogen A; (11) protesse A. The position of bromophemol blue is indicated by arrow.

The starting point of the scan was set at a point 15 mm from the start of the scan-length scale of the gel holder.



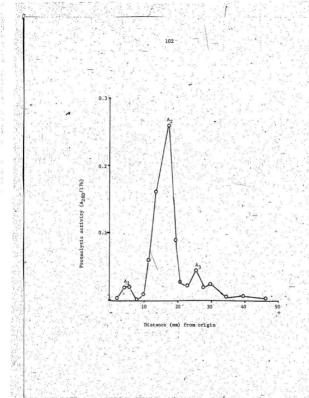
from 29% to 52% after activation. Peak D increased to 2.4% after activation from a faint band in symogen A. The areas under various peaks were estimated by method of triangulation. These results strongly suggested that A and B are symogens which are converted to faster moving bands. Nowever, the activation seems to be partial and complex as the band B did not completely disappear on activation.

To determine which electrophoretic component of zymogen A is proteolytically active 50 ug of zymogen A purified by affinity chromatography was electrophoresed at pH 8.3 as described in Materials and Methods and the gel was sliced in 2 mm thickness. By incubating these slices with hemoglobin solution at pH 1.8 at 30°C for 17 hours highest activity was detected in a position corresponding to Rf 0.37 (peak A2, Figure 19). There were minor activity peaks corresponding to Rf values of 0.10 and 0.55 (peaks A and A3, respectively). However, based on the gel-scanning results shown in Figure 18 peak C is 29% of the total area whereas A3 in Figure 19 represents only about 9% of the total pepsin activity. This may be due to instability of the active enzyme at high pH of the buffer used during electrophoresis. Two of the bands of protesse A had mobilities close to those of two bands in calf chymosin (Rf 0.180 and 0.650). However, the mobility of one of the bands in calf chymosin (Rf 0.590) was also close to that of porcine pepsin which was homogeneous with Rf 0.570 (see Figure 17). The heterogeneity of the crystalline calf chymosin is probably because of the presence of more than one isoenzyme forms (Foltmann, 1960; Asato and Rand, 1977), and products of autodigestion.

⁽n) Molecular weight: Polyacrylamide gel electrophoresis in the

Figure 19. Detection of proteolytic activity on polyacrylamide gel

Pifty Ms. of symogen A purified by affinity chromatography were subjected to-polyacrylamide gel electrophoresis at pi 8.3 following the method of Davis (1964). Two ms thick alices from the gel were incubated with 1.5 ml hemoglobin solution. at pi 1.8 at 30°C for 17 hours, and the product of hydrolysis determined spectrophotometrically as described in Naterials and Methods.



presence of sodifin dodecyl sulphate showed that symogen A purified by affinity chromatography appeared as a single band (Pigure 20) corresponding to a molecular weight of 33,800-1800 daltons based on an average of two estimageges. This Telefipher than the molecular weight estimated by gel filtration (Table 7). Electrophoresis of protesse A under the same conditions showed a broad diffuse band appearing at a position corresponding to molecular weight between 32,000 and 25,000 daltons. Purified tymogen C had a molecular weight of 44,000-2100 daltons and after activation the molecular weight of the main protein band was found to be 38,000 daltons. There was also a havy band in the molecular weight range of 37,000-23,000 daltons

(c) Amino acid compositions: The amino acid composition of the electrophoretic component of zymogan A with the highest proteolytic activity (peak A2 of Figure 19) is shown in Table 8. This grotein was isolated as follows: 480 ye of purified zymogan was electrophoresed on 12 analytical gels at pH 8.3 as described in Materials and Methods. A 3em length from each gel was cut off at a position corresponding to if 0.37. The slices were combined into a test tube, macerated on a Wortex mixer with a spatula dipped into the gel. When the gels were broken to small pieces it was homogenised into a thin slurry with 3.0 ml of 0.1M NaCl using a polytron homogenised (not a thin slurry with 3.0 ml of 0.0M NaCl. The vanishings were combined, dislysed against 2L water, pH adjusted with Ni₂OH to 7.0. The dislysate was freeze-dried and subjected to emino acid analysis after 24 hour acid hydrolysis.

Figure 20. Polyacrylamide gel electrophoresis of the proteases and their symogens in the presence of sodium dodecyl sulphate.

Folyacrylamide gel electrophoresis in the presence of sodium dodecyl sulphate (SDS PACE) was performed as described in Materials and Methods. Zymogens A and C were purified by affinity chromatography and gel filtration respectively. A, symogen A; B, protease A; G, symogen C; D, protease C; and E, standard molecular weight markers as indicated by (a), bovine series albumin (MH 65,000 daltons), (b) chicken egg albumin (MH 65,000 daltons), (c) ovine prolactin (MH 23,800 daltons), and (d), sytochrome C (MH 12,700 daltons).



Table 7. Summary of the estimations of the molecular weights of zymogens A and C.

	*	
Zymogen A	A STATE OF THE STA	Mean + S.D.
Sephadex G-100	0.996	5 26,300 <u>+</u> 1540
HPLC	0.995	26,300
SDS PAGE	0.998	2 33,800+1800

Correlation coefficients

Sephadex C-100 0.996 2 37,100+1075 SDS PAGE - 0.998 2 44,000+2100

lanalysed by the method of least squares.

Zymogen C

As shown in Table 8, there appears to be some similarity in the percentage compositions of aspartic acid, threenine, proline, glutatic acid, lysine, histidine and arginine for zymogen A and calf prochymosin. The percentages of the following amino acids tend to be different in the two symogens: glycine, valine, leucine, tyrosine whenylalmine, serine, two symogens: glycine, valine, leucine, tyrosine whenylalmine, serine, alamine and methionine plus its oxidized product(nethionine sulphone).

Isoleucine in prochymosin appeared to be twice as much as in zymogen A. Cysteine and cysteic acid together seemed higher in zymogen A than in prochymosin. Tryptophan was not determined in zymoggns A and C.

There appeared to be some sinilarity between symogen A and porcine pepsinogen as well. The percentages of serine, proline, valine, tyrosine and phenylalanine seemed similar. There was Biight difference in the percentages of the following: threonine, aspartic acid, alamine, and tyrosine. However, there seemed to be a great difference in the percentages of the following: glutamic acid, glycine, cysteine(together with cysteic acid), methionine(together with its sulphone), isoleucine, leucine, lysine, histidine and arginine. Therefore, there seemed to be more similarity between zymogen A and prochymosin than between pepsinogen and prochymosin.

Zymogens A and C seemed to be similar with respect to aspartic acid, threonine, serine, glutamic acid, alamine, isoleucine, leucine, tyrosine, and phenylalamine. Percentages of proline, cysteine plus cystein acid and valine appeared to be slightly different in the two zymogens. However, these two zymogens seemed greatly dissimilar with respect to the percentages of glycine, histidine, lysine and arginine.

Table 8. Comparison of Amino Acid Compositions

	Zymogen	Α - Δ5 P	rochymosir	² Zymogen (Δ P	Porcine	
Cysteic Acid	0.8		14,13	Tr		, a,	
Methionine sulph	one 0.3			0.0			
Aspartic acid .	9.6	0.7	10.3	11.4	0.7	12.1	1.8
Threonine	6.3	0.1	6.2	6.2	0.1	7.2	1.0
Serine	12.3	2.6	9.7	13.2	3.5	12.7	3.0
Proline"	4.7	0.3	4.4	6.7	2.3	5.2	1.8
Glutamic acid	11.8	0.6	11.2	11.0	0.2	7.7	3.5
Glycine	12.2	3.5	8.7	13.3	4.6	9.6	0.9
Alanine	6.7	2.0	4.7	6.6	1.9	5.2	0.9
Cysteine	2.0	0.1	1.9	1.9	0.0	1.7	0.2
Valine	5.9	1.3	7.2	4.4 5	2.8	6.3	0.9
Methionine	1.2	1.0	2.2	2.7	0.7	1.1	1.1
Isoleucine	3.3	2.6	5.9	3.9	2.0	6.9	1.0
Leucine	7.0	1.1	8.1	7.1	1.0	9.1	1.0
Tyrosine	4.1	1.5	5.6	4.9	0.7	. 4.7:	0.9
Phenylalanine	4.1	0.9	5:0	4.2	0.8	4.1	. 0.9
Lysine	3.6	40.4	4.0	1.0	3.0	2.8	2.2
Histidine	1.9	0.3	1.6	0.5	1.1	0.8	0.8
Tryptophan	N.D.4	100	1.2	N.D.		1.7	1.1.
Arginine	2.3	0.1	2.2	1.1	1.1	1.1	1,1
Mean A		1.18			1.56		1.35

Notes: 1. Expressed as percentage of total.

Calculated from data reported by Ryle (1970).

3: Calculated from data reported by Foltmann (1970).

4. Not determined.

5. Δ = Difference from prochymosin.

VII. Preparation of cheddar cheese

Cheddat cheeses were prepared from 181 batches of milk with commercial calf rennet, <u>Mucor michel</u> protease (Fromase), activated crude homogenate from harp seal gastric mucosa and activated DEAP peak A protease as milk cosgulants. Some aspects of the cheese making process are summarised in Table 9. Although the initial pH of the pasteurised milk was almost the same for all the 4 lots of milk used, the time required for the pH to reach 6.4 after adding starter varied slightly. Although equal clotting units of all the coapulants were used setting time varied slightly, perhaps because of slight temperature difference of the milk. The pH of whey at the end of Cheddaring was between 5.4 and 5:5. There was an initial rapid drop in pH of the cheeses. One week after making the cheeses the pH was between 4.70 to 4.90. The pH of the cheeses were similar after 30 weeks of sging.

The yields of cheeses ware very similar for all coagulants used (Table 10). Although there was slight difference in the total N values in whey, recovery of protein in the chesses was similar (Table 11). There were slight differences in the fat contents of cheeges. It is difficult to judge the significance of these differences from data obtained from one lot of cheeses. However, data from a preliminary cheese-making trial(Appendix E) also appears to show that the yields are/comparable between the cheeses made with ealf remet and SGF.

Table 9. Some parameters of cheese-making process

, /	Protesse A	Crude SGP	Calf rennet	Fromase	
pH of milk	6.60	6.50	6.50	6.50	
pH at renneting	6.40	6.35	6.40	6.40	
Temperature at renneting °C	30	31.	- 31	30	
Time between adding starter					
and coagulant (min)	40	50_	45	50	
Setting time (min)	25	20	20	22	
pH at start of cheddaring	5,40	5.35	5.50	5.40	
pH after one week	4.85	4.70	4.85	4.90	
pH after thirty weeks	5.05	5.20	5.11	5.18	

Table 10. Yields of cheddar cheeses and loss of nitrogen in whey

Enzyme used	Yield of cheese	1 Nitroge	en in Whey?
	, z	Total 2	Non protein
Calf rennet	10.04	0.1091	0.0509
Protease A	10.08	0.1109	0.0515
Crude SGP	9.96	0.1227	0.0517
Fromase /	10.00	0.1316	0.0510

I After pressing and before vacuum packing cheeses, without adjustment for moisture differences.

² Figures represent average of duplicate measurements.

Table 11. Proximate analyses of the Cheddar cheeses

Coagulant	Perc	ent composition 1
16. W. W. 1	Fat Mois	ture Protein ²
Calf rennet	30.79 38	.41 25.19
Protease A	31.49 38.	.22 25.49
Crude SGP	30.58 38	.26 — 25.23
Fromase	31.06 38	.66 25.02

1 Numbers represent average of two determinations:

2 Obtained by multiplying Kjeldahl N values by a factor of 6.38.

Data from one analysis on one lot of cheeses.

VIII. Analyses of the cheeses

The sensory evaluation of the cheeses aged for 4 weeks and 30 weeks was performed by preference test by 30 panelistic. The sensory scores were evaluated by analysis of variance (Larmond, 1977). As shown in Table 12 there was no significant difference in sensory preference accres between the week-old cheeses made with calf remnet and protease A and between the cheeses made with calf remnet and protease A and between the cheeses made with calf remnet and Fromase was significant at 5% level. After 30 weeks of aging the score for the cheese made with crude SGP was significantly higher than that for the cheese made with crude SGP was significantly shown in Table 13. The sensory scores for the cheeses made with Frymase and protease A were not significantly different from that of the cheese made with SGP.

(a) Citrate-MCL extract; Figure 21 shows the absorption spectra of citrate-HCL extracts from 30 week-old cheeses. The vareingths of maximum absorbance of citrate-HCL extract from cheeses were similar. Figure 22 shows the increase in absorbance of the citrate-HCL extracts with aging. It shows that initially the rate of tipening was faster for the cheeses made with crude SGP and Fromage up to 12 weeks when all the cheeses produced about the same-amount of UW absorbing materials. After 30 weeks, however, the citrate-HCL extracts from the cheeses made with calf remmet and Fromase appeared to have higher absorbance than than the extracts from the cheeses made with SGP. This may indicate instability of SGP as aboun by data in figures 10 and 13.

Table 12. Sensory evaluation of 4-week-old Cheddar cheeses

Coagulant	Mean sensory score?
Calf rennet	6.37ª,b
Protease A	6.87 ^à
Crude SGP	5.77a,b,c
Fromase	5.50°

- -1. Evaluated by preference test by 30 panelists on a 9-point scale:
- Standard error of mean = 0.26; least significant difference = 0.95.

Mean values followed by the same superscript are not significantly different at $p \le 0.05$.

Table 13. Sensory evaluation 30-week-old Cheddar cheeses1

Coagulant	Mean sensory score2
Calf rennet	5.67 ^b
Protease A	5.83ª,b
Crude SGP	6.83ª
Fromase	5.90a,b

 Evaluated by preference test by 30 panelists on a 9-point scale.

2.Standard error of mean= 0.29; least significant difference = 1.07.

Mean values followed by the same superscripts are not different at $p \leqslant 0.05$.

Figure 21. Absorption spectrs of citrate-HCl extract from 30 week old cheeses.

Citrate-NCI extracts of the cheeses were prepared by following the method of Vakaletis and Price (1959) and scanned in a Reciman DUS spectrophotometer. a, Fromase; b, calf rennet; c, protesse A; d, crude SGP

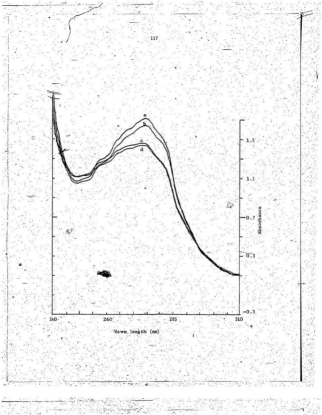
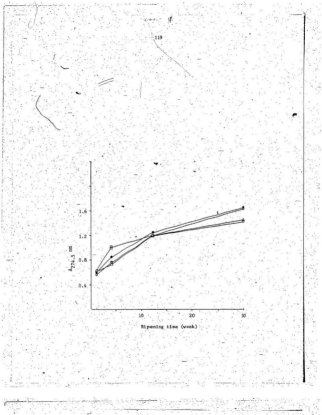


Figure 22. Spectrophotometric monitoring of protein degradation in ripening cheese.

Citrate-HCL extracts of cheeses at various periods of ripening were made following the method of Vakaleris and Price (1959) and absorbance measured at 274.5 mm. D, cheese made with crude SGP; A, cheese made with processe A; O, cheese made with calf rennet; • cheese made with Frommas.

Data from one lot of cheese.



The citrate-HCl extracts from 30 week old cheeses were snalysed by polyacrylamide gel electrophoresis following the method of Davis (1964). The gels were stained and the mobilities of each component was determined using a Beckman DU8 gel scanner at 570 nm. The contribution of each peak to the total was calculated as the percentage of the total area (Table 14; photographs in Appendix L). There are similarities in the number of protein bands present in the extracts from the cheeses made with the SCP, protesse A and calf rennet (bands a to p). The extract from Fromsse cheese showed a lesser number of protein bands than was present in the extracts from the other three cheeses. There was considerable difference in the percentages of the fast-moving peptide designated q (Table 14) in the extracts from the four cheeses. The extract from Fromase cheese had the highest percentage of fast-moving bands whereas that from calf rennet cheese had the lowest percentage. The extract from crude SGP cheese had a percentage of fast-moving bands between these two values. This information may explain the higher sensory preference scores (Table 13) of the SGP cheese after 30 weeks of sging. However, the contribution of the pentides to the sensory quality of the cheeses would not be known until the pentides are isolated and subjected to sensory evaluation.

(b) Electrophoresis of whole cheeses protein: Folyacrylamide gal electrophoresis of the whole cheeses was performed at pH 8.3 favis, 1964) at verious stages of ripening. The protein in whole cheese namples was completely solubilized in citrate buffer containing urea as proviously shown by Stanley and Emmons (1977). Figure 23 shows the sictures of sels superimposed on their scans from 30-week-old

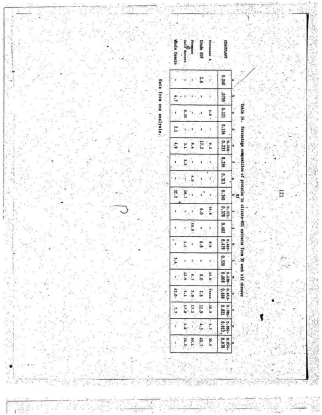
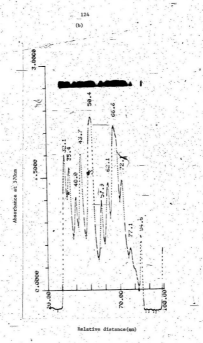
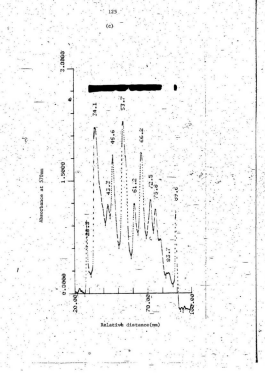
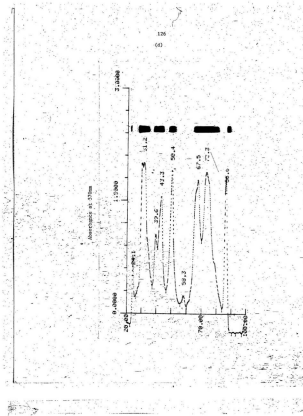


Figure 23. Polyacrylamide gel electrophoresis of whole cheese protein.

Samples From 30-week-old Cheddar cheeses were solved liked following the sathod of Stanley and Emmons (1977) and 10 µl aliquots Electrophoresed according to the method of Davis (1864), stained, destained and scanned as depertured in the legend to Figure 25. Cheese made with (a) protesse A, (b) crude SGF, (c) calf remnet, (d) Fromasé.







cheeses. Whole casein was also electrophoresed as reference (Figure 24) each time the cheese samples were electrophoresed. The reproducibility of the Rf values of the various caseins and of their degradation products was good (0-4.7%) for the fast-moving components (E to M. Table 15) but poor for the relatively slow-moving components (A to D). The components of standard casein were identified on the basis of their relative mobility as shown by Stanley and Emmons (1977). Para K-casein component (Rf=0.07) was not observed in one-week-old cheese samples. However, a band was present at the starting point of the small-pore gel. This may include para-K-casein plus the degradation product of other casein components with overlapping mobility because the percentage of this band was higher than that of K-casein in the whole casein gel. In the 30-week-old cheeses there were two bands in the position of para-K-casein in the cheeses made with protease A and SGP whereas in the case of cheeses made with calf rennet and Fromase the two bands were merged. These two components were estimated together and recorded as para-K-casein. Therefore what has been reported here as para-K-casein is perhaps the sum of para-K-casein and some other peptides. In all the cheeses the percentages of para K-casein. Ts-, and y-casein increased after 30 weeks of aging. However, it should be pointed out here that since changes in percentage composition estimated this way are only relative, an increase in the percentage of one component may also suggest that the component in question has changed slowly or not at all, while the other components have diminished faster (Stanley and Emmons, 1977).

The most remarkable differences are in the major casein components $a_{\rm gl}$ and β . The $a_{\rm gl}$ casein was much more extensively degraded in the

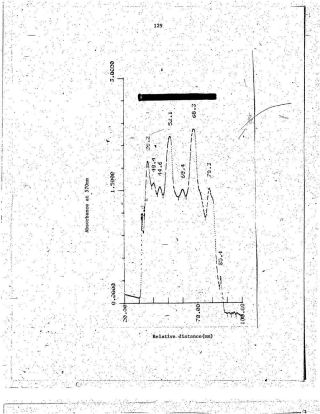
Figure 24. Polyacrylamide gel electrophoresis of whole casein.

Acid casein prepared following the method of Fox and Guiney (1976), Forty mg of freeze-dried casein was dissolved in 10 al of 6H urea containing 0.1%, potassium citrate (Stanley and Emmons, 1977) and 15 HL used for electrophoresis following the method of Davis (1964) Other conditions described in legend to Figure 23.

Identification of casein components

	(mm)
and the state of t	A STATE OF THE STA
K-casein	36.2
Y-casein	40.4
rg-casein	44.6
β-casein.	52.1
Minor as casein	60.4
1-casein	68.3
	A CONTRACTOR OF THE CONTRACTOR
Minor as casein	78.3

Identifying numbers appearing above each peak represents its
distance in millisserer from the start of the scan longth scale
of the gel folder. The starting point of the scan was set at a
point 20 me from the start of the scale.



30-week Ad calf rennet cheese (Table 15, Figure 25s, b) than in the other three cheeses. In the Fromase cheese the degradation of a₉₁ casein was less extensive than in calf rennet cheese but breaseful underwent the highest degradation. In the cheeses made with protease A and SGP, the percentages of residuald₉₁-casein were higher than in calf rennet cheese but the percentages of residual \$\text{P}_{\text{0}}\$-casein were comparable to that in the calf rennet cheese.

Some protein bands with sobilities different from those of standard casein components appeared with aging of the cheeses in positions E, G, H, J, K, L, and M (Table 15). Change in the percentage compositions of most of these proteins in the seal protesse cheeses were similar to those of the calf rennet cheese after 30 weeks of aging (Table 15).

(c) Gel filtration of cheese extracts: Samples' from 30 week old cheeses were dissolved in 0.1M tris citrate buffer pN 8.6 containing 6M ures, 1 mM EDTA and 1 mM DTT according to the method of Foster and Green (1974) and chromatographed on a Sephadex G-100 column to estimate the sizes of proteins and the extent of their degradation. To estimate the percentage compositions of the proteins a triangulation method was followed for calculation of peak areas. Since the peaks were not resolved completely the base-line was connected to the lowest point between the unresolved peaks. Figure 26 shows the clution patterns. The approximate molecular weights of the proteins and their percentage compositions are listed in Table 16. The percentages of low-molecular weight proteins (less than 12,700 daltons) in the cheeses sade with the seal processes were less than that in the Fromase cheese but hither than

(.068-.079)(.156-.18 (.264-.263)(.344-.393)(.484-.593) (.527) (.527) (.527-.528)(.614-.648)(.612-.648)(.741-.763)(.865-.869)(.866-.899) (.988) Figure 25. Degradation of major casein components during ripening of Cheddar cheese.

Samples from 1 week, 4 week and 30 week old Cheddar cheeses were subjected to plyscrylanide gel electrophoresis as described in materials and methods and the gels scanned in a Neckman DUB spectrophotometer equipped with a gel scanner capable of area calculation. The figures show changes in each casein component as a percentage of total casein residues in cheese with eging. (a) and (b) Ks.-casein (——), p-casein (——), for para-R-casein (——), fa-casein (——), for heese made with calf rennet; o, cheese made with Fromser; A, cheese made with crude SGP. The data is for 1 analysis for 1 lot of cheese,

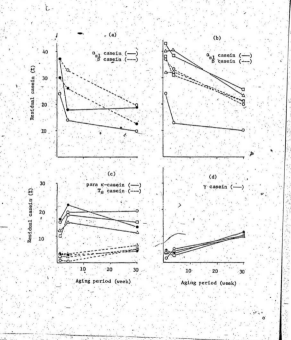


Figure 26. Separation of cheese proteins on Sephadex G-100 column. -

Samples from 30-week-old cheeses were dissolved in buffer as described by Foster and Green (1974) and 5.6 ml solution applied to a 1.6 x 88.5 cm column of sephadex 6-100. Void volume of the column, 32 ml. Flow rate 6ml/h, fraction size, 2.0 ml.

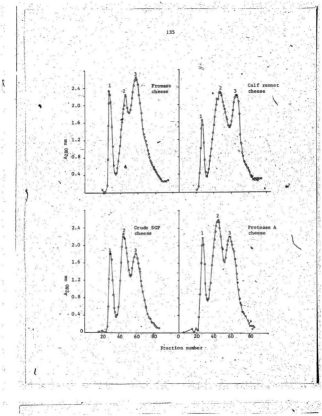


Table 16. Protein degradation patterns in 30 week-old Cheddar cheeses

Coagulant		Protein composi	tion(X)1
	Peak 1 (70,000 daltons)	Peak 2 (30,000 daltons)	Peak: 3 (<12,700 daltons)
Fronase	- 12.6	33.6	53.7
Protease A	14.3	46.4 45.5	39.0
Calf rennet	9.6	58-1	32.3

I Estimated by triangulation method. Since the peaks were not resolved calculation was done by connecting the base line to the lowest point between two unresolved peaks.

that in calf rennet cheese. This observation supports that summarised in Table 14 which shows that the percentages of the fast-moving peptide q in the extract from SGP cheeses were between those of the extracts from Fromase and calf remnet cheese. Although the molecular weights of the casein components are between 18,000 and 23,000 daltons, a considerable amount of protein in the cheeses had molecular weight higher than those of caseins indicating the aggregation of the proteins even in the presence of 6M ures. Foster and Green (1974) also made similar observations: These authors also observed that in the fresh curd the protein peaks of molecular weights between 11,000 and 22,000 daltons were relatively small but increased in size during aging.

(d) Autho actd analysis of cheeses: Citrate-HCI extracts from 1 week and 30-week-old cheeses were analysed for both free and total (i.e. hydrolysate) amino acide. Results are shown in Figure 27 % and B. Autho acid composition of the citrate-HCI extract of old "Forfar" Cheddar cheese - a very popular brand of Cheddar cheese is also included for comparison (Tigure 28).

The free amino acid content of 1-week-old cheeses were about 4 to 6 unoles/g cheese (see Appendix F). The compositions of the total amino acids were also similar for the extracts from all the one-week-old cheeses (Appendix H). After 30 weeks of aging free amino acid levels rose significantly (Appendix F) to 33-39 unoles/g cheese. The levels of free amino acids in the two SOP cheeses were somewhat lower than those of calf remet and much lower than Fromse cheese (Appendix G, J and Figure 27). Cheeses made wish protesse A and crude SOP showed lower levels of observationine and higher levels of cysteck acid and

Figure 27: Amino acid compositions of the Citrate-HCl extracts from Cheedar Cheeses.

Free anino acids were determined on the clear supermatant obtained by treating the Citrate-NCI extracts with 4 volumes of 20% sulphosalicylic acid. The total anino acids were determined by hydrolysing the Citrate-NCI extracts in 6h HCI for Z4h at 110°C. Amino acids have been numbered as follows:

 Cysteic	acid .	

15. Methionine

Glycerophosphoethsnolamine

Cystathionine
 Isoleucine

3. Aspartic scid

18. Leucine

4. Threonine
5. Serine

19. Tyrosine

6. Asparagine

20. Phenylalanine
21. Y-Amino butyric acid

7. Glutamic acid

22. Tryptophan

8. Glutamine 9. Proline

23. Ornithine 24. Lysine

10. Glycine

25. Histidine

12. Citrulline

. - . . /

13. Valine

26. Arginine 27. Taurine

14. Half cystine

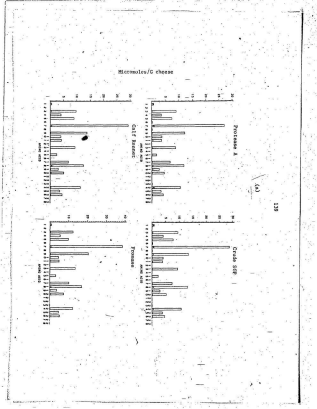
28. \$ -Alanine

Legends: Cross-hatched bar, free amino acid;

Open bar, total amino acids.

. Open par, cocar amino acida

(a) One week old cheeses; (b) Thirty week old cheeses



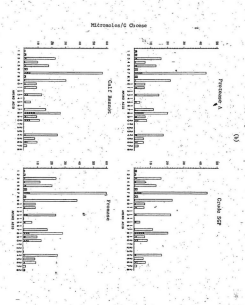
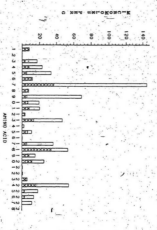


Figure 28. Amino acid composition of the citrate-HCl extract from old

Legend same as in Figure 27.



glutamine than in calf remnet cheese (see Appendix 6). The level of threonine was higher in crude SCP cheeses than in the cheese made with professe. A. Whereas the levels of glycerophosphoethanolamine, serine, and proline were lower. Cheese made with Fromase showed higher levels of threonine, serine, glutamine, valine, methionine and ornithine while the levels of aspartic acid and leucine and argining were lower compared to calf rennet cheese. Except for valine, which had a higher percentage in the extract from 30 wask old crude SCP cheese than in the other three cheeses, the percentage compositions of the total anino acids for all the four cheeses were very similar (see Appendix I).

"Forfar" is considered an outstanding Cheedar cheese, which is manufactured in Ontario. No details were available from the retailer about the cheese except that it was aged for 1-2 years and was prapared with calf rennet with no cheese color or other additives added. This cheese is presumably sanufactured from row milk. None of the cheeses prepared in this study had any added color but there was significant color similarity between the 30-week-old SCF cheeses and the "Forfar" cheese. "Forfar" and the two SCF cheeses had a yellow tinge which appeared to be absent in the Fromase and calf remost cheeses.

Free amino acid levels in the cirrate-MCI extract from "Forfar" theses was much higher than any of the aged cheeses of this study an shown in Figure 28. Percentages of free aspartic acid, arginine and phenylalania are particularly lover and the percentages of free cysteic acid, glutamic acid, citruline, isoleucine, ornithine, lysine and histidine are higher in "Forfar" than in the calf remnet cheese. The percentages of total aspartic acid and arginine are lower and the percentages of total aspartic acid and arginine are ligher in the

"Forfar" than in the calf rennet cheese. High levels of mino acids or differences in relative proportions do not seem to explain the observed sensory quality of the four cheeses made in this laboratory. Likewise, these higher levels of amino acids in "Forfar" cheese than in the calf rennet cheese may not be entirely responsible for its superior quality. "Forfar" cheese is believed to be manufactured with raw milk. Other manufacturing conditions such as the type of starter culture and the type of rennet used in the "Forfar" are not known: These are also very important in the development of flavor (see Gryer, 1969 for a review).

GENERAL DISCUSSION AND CONCLUSION

"Repainogens" were isolated from the stomach success of young and adult harp seals (<u>Fagophilus greenlandicus</u>). The two major pepsinogens were purified, activated and characterised with emphasis on the chymosin-like properties and use as a remnet substitute.

1. Physical properties of SSP and other gastric nuces separated into four active components by ion exchange chromatography on DEAE-Sephadex A-50 column (Figure 2) or isoelectric focusing (Figure 15). Based on the relative peak sizes pepsinogens A and C from ion exchange chromatography comprised the majority of the pepsin activity of the crude extract; components B and Cf occurred in relatively small proportions.

Occurrence of multiple isoenzymes had been reported in stomachs of other aperies as well. Human stomach secretes 7 pepsinogens (Rtherington and Taylor, 1970) and monkey stomach secretes 5 pepsinogens (Rtherington and Taylor, 1970). Chicken stomach secretes 4 (Donta and Van Vanakia, 1970) or 5 pepsinogens (Green and Lewellin, 1973). Stomachs of hog (Fruton, 1971), cow (Meitner and Kassel, 1971) and dogifish (Bar-Rii and Merret, 1970) secrete 4 pepsinogens. Castle and Wheeledck, (1971) showed that call secretes 2 prochymosins. Castricsia, another protoclytic engyme, has been reported to occur in stomachs of several species (Tang, 1970; Waid'si al., 1981; Ryle, 1960; Chiang and Ponce, 1981). However, multiple gastricsins have been reported opy in the figh Medluccium gay' (Chiang and Ponce, 1981).

The electrophoretic mobility of zymogen A determined by the method

of Davis (1964) was much alower than that of zymogen G. The main protein band of protease A also showed much slower mobility than those of proteins G, porcine pepsin and miso those of the major components of calf chymosin. However, both chymosia and protease A had a slow moving band.

It is interesting to note that protesse. A is electrophoretically faster than its precursor and this observation is similar to that of Asato and Rand (1977) who noted that all chymosins from calf stomach were electrophoretically faster than their respective precursors.

On the other hand, chicken pepsins were found to zove slower than the corresponding pepsinogens (Donta and Van Vunakis, 1970), similar to the pepsin/pepsinogens (Donta and Van Vunakis, 1970), similar to the pepsin/pepsinogens (Donta and observed in this study. Therefore the relative electrophoretic mobilities of tymogen and enzyme also suggest this protease A is a chymosin-like enzyme.

Incelectric focusing of the crude enzyme from the gastric nucons of a two year old harp seal revealed that there are 4 isoenzymes N, N, 0, and P as shown in Figure 15. The isoenzyme M which had a higher CU:FU than the other three had an isoelectric point of 4.90, similar to that of calf chymosin with a pl of 4.70 (Righetti et al., 1977). The isoenzymes were 4.31, 3.82, and 3.40. These values are higher than the reported isoelectric points of bovine and porcine pepsine but similar to those of chicken pepsin (Righetti et al., 1977).

The molecular weight of symogen A estimated by gel filtration was 26,300 daltons; a similar molecular weight was estimated by HPLC. This is much lower than those of most pepsinogens and also lower than that of calf prochymosin. However, the molecular weight of symogen A estimated by SDS polyacrylamide gel electrophoresis was close to that of calf prochymonin. Zymogen C had an estimated molecular weight of 37,100 daltons by gel filtration and 44,000 daltons by SDS polyacrylamide gel electrophoresis (Table 7). Similar discrepancies in the molecular weights of porcine pepain estimated by two different methods were reported before (Bies, 1956; Edelhoch, 1957). Chiang and Ponce (1981) reported the molecular weights of gastriceinogens I and II from M. gayl to be 27,000 and 28,000 daltons by gel filtration and 23,300 and 33,300 daltons, respectively, by SDS PAGE. Actual molecular weight of symogen A would be known only after complete electdation of its primary structure.

II. Catalytic properties of SGP and other gastric protesses: All the four isoenzymes of harp seal gastric proteases showed milk-clotting and hemoglobin-hydrolytic activities. However, the milk-clotting activity of the isoenzymes per unit of their peptic activity called CU:PU differed for the iscenzymes. The CU:PU ratio for protease A was higher than that of the other major protease C but was similar to that for chymosins from calf (Green, 1972), kid and lamb (Anifantakis and Green, 1980). Piglet "chymosin" was reported to have a CU:PU higher than that for calf chymosin (Foltmann et al., 1978). Green (1972) observed that a crude extract from adult bowine stomach had a higher CU:PU than calf rennet. However, CU:PU for purified bovine chymosin is much lower than that for purified chymosin (Amer et al., 1980; deKoning, 1978). Gastricsin also has activities on milk and on hemoglobin, but there is little data on its CU:PU ratio. However, Tang (1970) reported that the milk-clotting and proteolytic activities of human gastrics in are less than those of porcine pepsin.

The optimum pil of hemoglobin hydrolysis for each processe A was 3.5 (Figure 12) which is also similar to that for call chymosin (Bertidge, 1945). The pil optimum for processe C was 2.2, which als similar to those for human peppin (Tang, 1970), porcine pepsin (Chiang et al., 1967), and bovies peppin (Amer et al., 1980). An ion-exchange fraction of adult bovine systric processes with CU:FU similar to that of calf chymosin (adult bovine chymosin) had a pH optimum of 4 on homoglobin substrate (Amer et al., 1980). A gastric processe from young rate was called chymosin based on its pH optimum (3.8-4.2) for hydrolysis of homoglobin (Kotts and Jenness, 1976). Piglet chymosin (based on immunological sctivity and CU:FU) also had a pH optimum around 3.5 (Foltmann et al., 1978) Castricsis hydrolyse besoglobin optimally at pH 3.0 (Chiang et al., 1981; Ward et al., 1978; chiang and fonce, 1981; and Tang et al., 1961).

Stability and milk-clotting activity near neutral PH are considered distinguishing characteristics of chymosin (Malpress, 1967; Erestros, 1961; Foltmann, 1959). Although seal gastric protease A was sore unstable than calf chymosia in buffers of various pH values, (Figure 13), it was much more stable at pH 7.0 than portine pepsin. Seal proteases A and C eshibited milk-clotting activity up to pH 7.02 whereas under similar conditions porcise pepsin failed to clot milk above pH 6.5 (Figure 6). More recent reports show however, that other gastric proteases also have these properties. Fox (1969) showed that bovine pepsin clots milk up to pH 6.9. Chicken pepsin (Donts and Van Vunskis, 1970), porcine pepsin B (Myle, 1970), gastricains from fish (Chiang and Ponce, 1981), and toad (Ward et al., 1978) are atable in neutral of slightly sikeline buffers. Protease A and calf chymosin

were senatured by urea to the same extent in 90 minutes although the initial rates of inactivation were different. Protease C and portine pepsia underwent much less inactivation. These observations are consistent with the view that hydrogen bonds may play more important role is maintaining the active configuration of seal protease A and of call chyrogin than of posin (Erastron, 1974).

Calcium chloride added to "reconstituted skin milk influenced the clotting activities of both calf chymosin and processe A similarly (Figure 9). The optimum concentration of added calcium chloride was 0.02M for both the engymes; above that concentration milk cloring was inhibited. This observation was consistent with the reported inhibition of the action of chymosin on x-casein (Kamsnori et al., 1977). However, Cordid and Rosenthal (1978) observed only slight inhibition of milk-cloring activities of chicken pepsia and calf remet at high concentrations of added calcium chlorids. The discrepancy between the results of this study and those reported by Gordin and Rosenthal may be due to differences in the conditions of milk-cloring assay as discussed earlier (Results and Biscussion, Section VI(f)).

III. Conclusion: Tables 17, and 18 compare the properties of calf chymosic and protease A. Although there are some differences between the two enzymes most of the evidences support the hypothesis that protesse A is chymosin-like. One setable difference is the lack of stability of protease A in buffers of various pH willies particularly in pH values near 7 where chymosin is relatively stable (Foltman, 1959). This was considered as a basis for distinction between calf chymosin and protein paps in at least in one instance to conclude that

Table 17. Comparison of the physical-chemical properties of protesses

A and other gastric protesses

Properties	Chymosin-like	Pepsin-like	Gastricsin-like
Molecular weight	similar.	different	similar
Amino acid composition	similar	different	different2
Isoelectric point	similar.	different	-4
Electrophoretic mobility	partly similar	partly simila	r3
Stability in 6M urea	similar	different	-1
Stability in buffers of various pH values	different 4 .	similar	similar 5

By SDS PAGE.

²Mills and Tang (1967).

³ Data not available.

More stable than porcine pepsin at pH 7.

⁵ Fish gastricsin is an exception (Chiang and Ponce, 1981):

Table 18. Comparison of proteases A and other acidic proteases

2		30 T		Ġ
Properties	Chymosin-like	Pepsin-like	Gastricsin-	
			TING	
CU:PU	similar	different	1	
pH optimum for Hb hydrolysis	similar	different	similar	
Time-course of NPN release from casein	similar .	different	_r ·	
'Inactivation of ribonuclease	similar	different	'different	
Activity on APDT	similar i	different	similar	
Sensitivity of milk-clotting activity to Ca ⁺⁺ Clotting activity as a function of milk p ^H	similar similar	different ² .	_1 , _1	
Casein degradation in cheese	partly	different ³	1	
Milk-clotting activity as a function of temperature	partly similar	partly similar	_1	
Activation energy	partly similar	partly similar	_1	
Milk-clotting activity as a function of enzyme dilution	on different	similar	1	

. Data not available

- Gordin and Rosenthal (1978) observed slight difference between calf chymosin and chicken pepsin with respect to their sensitivity to Ca⁺⁺.
- 3.SStanley and Emmons (1977) differences in the electrophoretic patterns of casein in Cheddar cheeses made with calf rennet and bovine pepsin.

human stomach does not secrete chymosin (Malpress, 1967). However, as discussed in the introduction, lack of stability in buffers of certain pH values does not rule out the possibility of existence of chymosin (Hirsch-Marie et al., 1976). The broad pH optimum of protease A for hemoglobin substrate is similar to both chymosins (pH optima 3.5-4.2) and gastricgins (pH optima 3.0). This, coupled with the very low hydrolytic activity of protease A on APDT and the low molecular weight of . zymogen A estimated by HPLC and gel filtration raise the question whether protease A is a chymosin or gastricsin. However, unlike chymosin and seal gastric protease A. gastricsin has been classified as a minor gastric protease (Foltmann and Pedersen, 1977). Besides, gastricsins from pig and human inactivate ribonuclease (Tang. 1970) whereas seal protease A' and chymosins from calf, lamb, kid and piglet do not (Bang-Jensen et al., 1964; O'Leary and Fox, 1975). The last named authors concluded that ribonuclease test is generally applicable to differentiate chymosin from other gastric proteases,

Based on the evidence discussed above, it may be concluded that seal gastric protease A is a chymosin-like enzyme. Boovever, in view of the difficulty in defining an enzyme as chymosin as distinct from the homologous enzymes like pepsin and gastricain, further studies, e.g., immunological relationship between protease A and other homologous gastric proteases (Foltmann, et al., 1981) and the comparison of its primary structure with those of the other gastric proteases (Foltmann and Pedersen, 1977) would confirm whether protease A is a chymosin or not.

IV. Cheese making properties of protease A and SGP: The properties of crude SGP and protease A with respect to their application as milk

cosquiants for making chasse are comparable to those of calf renner (Table 19). Like calf chymosin, the destrable component in-commercial renner, SCP clots milk near pll 7 and has limited specificity. A combination of these properties is not found in other renner substitutes. Curdiing of cheese milk at its physiological pl (near 7) without prior acidification is often practised by cheese industry, particularly to avoid phage contamination (Phelan, 1973). Most pepsians do not clot milk at pll 7; others can clot milk at neutral pll but are too proteolytic and are therefore undestrable for yield, texture and flavor considerations of the aged cheese.

In the actual cheesemaking trials the yield of cheese and fat retention were similar in cheeses made with calf remest and SGP. The sensory preference scores of the cheeses made with crude SGP and procease A were either higher than or comparable to that prepared with calf remnet (Table 20). Protein degradation in the cheeses made with SGP and protease A was low as evident from the low amino acid contents and ripening indices of the citrate-HGl extracts of-the cheeses made with these two congulants. These observations suggest that limited proteolysis is destrable for development of sensory qualities of cheese, and are consistent with the literature reports that excessive proteolysis is associated with poor quality cheese.

In view of the similarity of calf chymnofin and protesse A in enzymatic and physicochemical properties and the results of the Chaddar cheese preparation with respect to yield, chemical and sensory analyses it may be concluded that seel gastric protesse A and the crude SGP containing protesse A may be a good remmet substitute. However, further trials, with larger number of cheese samples on an industrial

Table 19. Some criteria for an ideal rennet.

	Ch	ymosin		SGP	Pepsin	Gastricsin
Ability to clot milk at pH 7	A					9
milk at pH 7		+		+	-1	
Yield of cheese		+		+	-3	2
Fat-retention		+		+	-3 :	2
					- 1	~ .
Specificity		+		+		2
Availability				+4	+ .	-
			-			

- 1. Bovine and chicken pepsins are exceptions
- 2. Data not available
- From reports on bovine (Emmons et. al., 1978) and chicken (Stanley et. al., 1980) pepsins.
- Scal stomach should be easily available as a by-product of seal fishery in Newfoundland.

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. 50				
datio	20.8	21.0	19.2	
8.0	0	2	-	
22				
ation	22.9	3.1	. 4. 6	
Degrad	,			
lex .				
<u>.</u>	09	. 29	38	
gut	0.1	0.167	0.238	0.06.9
Ripen	0,160			
a x				
Total amino activate of the control of Degradation of Degradation trace-HG1 oxt. Degradation of Degradation puple/g Ripening index a _{nl} casein? ß casein ²	220.1	209.4	280.9	0
ettr		-5	,	
Free amino acids pmole/8	32.7	33.0	38.9	
Pi .				
Sensory	6.83	5.83	2.67	00
Coagulant	Crude SGP	Protease A.	Calf rennet 5.67	00 9
		~	-	ľ

sidual

scale will be necessary to conclusively establish the suitability of seal gastric protesses as rennet substitutes.

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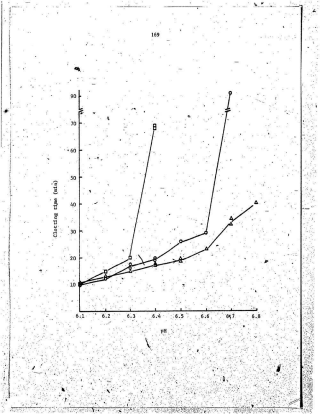
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Appendix A. Influence of pH on milk-clotting times of the enzymes

Enzyme concentrations were adjusted to have equal milk clotting times at pH 6.1. The milk pH was adjusted with 1 M NaOH or HCl. Forcine papsin, C ; seal gastric protease, O; and calf chymosin, A.



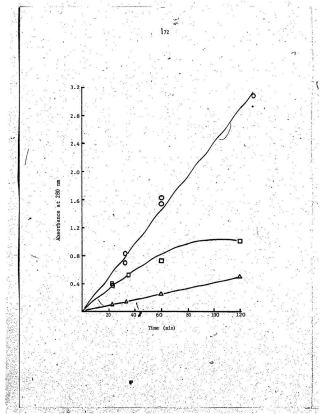
Appendix B. Relative rates of milk-clotting and proteolytic activities of proteases.

	1 18		S	ubstrate	
Protease		1	Hemoglobin CU:PU ¹		Cu:PU
Calf chymosin			0.170		1.61
Porcine pepsin	s ai		0.005		0.39
SGP ·			0.025		0.26

1 Pepsin unit (PU) and clotting unit (CU) are as defined by Anson (1948) and Berridge (1945). The pepsin units are based on initial velocity. Appendix C. Time-course of casein hydrolysis by proteases.

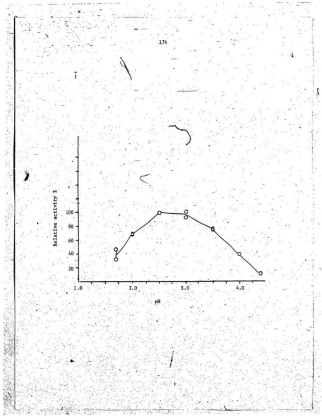
Ten al of a 27 casein solution was incubated with 0.5 ml of each of the three processes (pornine aspain, crude seal gastric protease, and calf dymosin) at 37°C. At intervals 1.0 ml aliquots were withdrawn in duplicate, treated with 1.0 ml of 10% TCA. The mixture was cooled at 4°C for 30 minutes and centrifuged at 3,000xg for 30 minutes. The absorbance at 280m was negaured as described in Materials and Methods.

Porcine pepsin, \Box ; SGP \bigcirc ; calf chymosin, \triangle .



Appendix D. Influence of pH on the hydrolysis of hemoglobin by crude

Three al. of 22 hemoglobin in citrate buffers of various pH values were equilibrated at 30°C. To each sample 0.2 al. of crude SF containing 3.9 papers units of activity was added. After 20 minutes 0.9 al. samples were withdraw, mixed with 1.0 al of 6% TCA, cooled on ice ifor 30 canutes and centrifuged at 3,000 x.g for 30 minutes. The A280 nm of the supernatant was noted. The A280 nm of the coursel samples in which the enzyme was added after the TCA was subtracted from the corresponding teet samples.



apendix E. Analysis of experimental Cheddar cheeses

Coagular	it Y	Leld		0.00			100	
	(8/10	00 ml milk)		141				
100	100			CF 8 Q	A STATE OF THE REAL PROPERTY.			
	Fresh	Dry		%	Citrate-HC1		Sensory_	
2.0	Weight	Weight.	29	moisture	Soluble N1		quality2	
	of a second		- 2		A274 - 5nm	* 10.	1.0	
14			0.0				* 2 ×	
SGP	9.35	6.60		29.3	1.73		Typical	
1917	-,6 9.33				- 1 - C.		Cheddar	
		2.0						
Calf ren	net: 8.64	6.25	100	27.6	1.48		Typical	Ĺ,
	110			,50			Cheddar	

¹ Average of duplicate measurements following the method of Vakaleris and Price (1959).

² Based on-tasting by 5 individuals in the laboratory after II weeks of aging. The cheeses showed mold growth when 17 weeks old and therefore were not suitable for a routine taste panel.

Appendix F. Free amino acid composition of citrate-HCl extracts from one-week old Cheddar cheeses

Amino Acid	Calf	Rennet	Crude	SGP	. Protes	se A	From	ase -	
	µmole/	g z	µmole/g	z	µmole/g		mole/g	x.	
Cysteic acid	- 0.60	13.6	0.79	15.3	0,:73	12.8	0.72	15.6	
Glycerophospho-					4				
ethanolamine	0.00	0.0	0.00	. 0.0	0.00	0.0	0.00	0.0	
Aspartic acid	0.40		0.33	6.4	0.15	2.6		6.1	
. Threonine	0.05		0.04	0.8		. 1.6		1.5	
Serine	0.04	0.9	0.05	1.0	0.07	1.2	0.07	1.5	
Asparagine	0.07	1.6	0.24	4.6	0.19	3.3	Trace		
Glutamic Acid			0.73	14.1	0.92	16:1	0.65	14.1	
Glutamine	0.00			2.7				.0.0	
Proline	0.18	4.1	0.44	8.5	0.33			6.3	
Glycine	0.09	2.0	0.07	1.4	0.07	1.2		1.5	
Alanine	0.15		0.23	4.4		7.7		2.8	
Citruline	0.00		0.00	0.0	0.00	0.0		0.0	
Valine ·	. 0.09	2.0	0.10	: 1.9	0.14	2.5		1.5	
Cysteine	0.00	0.0	0.00	0.0	0.00			0.0	
Methionine	0.00	0.0	. 0.03	0.6	0.07	1:2	0.00	0.0	
Cystathionine	Trace	2	0.00	0.0	0.00	0.0	0.00	0.0	
Isoleucine-	. 0.07	1.6	0.12	2.3	0.14	2.5	0.08	1.7	
Leucine -	0.74	16.7	0.51	9.8	0.41	.7.2	. '0.67	14.5	
Tyrosine	0.18	4.1	0.29	. 5.6	0.37	6.5	0.40	8.7	
Phenylalanine	0.39	8.8	0.26	5.0	0.14	2.5	0.60	13.0	
Y-Aminobutyric	acid 0.00	0.0	0.00	0.0	0.00	. 0.0	0.00	.0.0	
Tryptophan.	. 0.00	0.0	0.00	0.0	0.00	0.0	0.00	0.0	
Ornithine .	0.07	1.6	0.05	1.0	0.09	1.6	0.06	1.3	
Lysine	0.31	7.0	. 0.49	9.5	. 0.88	15.4	10.23	5.0	
Histidine .	0.07	1.6	0.10	1.9	0.07	1.2	0.07	1.5	
Arginine	0.23	5.2	0.17	3.3	0.24	4.2	0.15	3.3	
Taurine	0.00	0.0	0.00	0.0	. 0.00	0.0	0.00	0.0	
6-alanine	0.00	0.0	0.00	0.0	0.00	0.0	0.00	0.0	
11 1									
	4 4 4			2	e 71		1. 64		

Appendix G. Free amino acid composition of citrate-HCl extracts from 30-week old Cheddar cheeses.

Amino Acid	Calf R	ennet	· Crude	GP	Protess	e A	From	ase	
	μ mole/g	. 1	µmole/g	Z,	µmole/g	2	umole/g	7.	
Cysteic-acid -	0.9	2.3	1.4	4.3	1.4_	4.2	0.8	2.3	
Glycerophospho-					See .				
ethanolamine	1.5	3.9	1.1	3.4	_1.5 "	4.5	1.3	3.7	
Aspartic acid	1.6	4.1	1.5	4.6	1.4	4.2	0.9	2.6	
Threonine	0.6	1.5	0.9	2.8	. 0.6	1.8	0.8	2.3	
Serine	1.2	3.1	0.5	1.5	1.3	3.9	1.7	4.8	
Asparagine	2.5	6.4	1.9	5.8	1.9	5.8	2.4 4	6.8	
Glutanic Acid	4.4	11.3	4.2	12.8		13.9	. 4.3	12.3	
Glutamine	1.4	3.6	2.2	6.7 -	1.8	5.5		5.1	
Proline -	0.7	1.8	0.4	1.2	0.7	2.1	0.8	2.3	
Glycine	0.9	2.3	0.8	2.4	0.7	2.1		1.7	
Alanine	1.0	2.6	1.5	4.6	1.4	4.2		2.8	
Citruline	0.2	0.5	Trace		0.1	0.3		0.3	
Valine	2.6	6.7	1.3	4.0	1.2	3.6		8.0	
Cysteine	Trace		. Trace		Trace		Trace		
Methionine	1.0	2.6	1.2	3.7	1.2	3.6		3.9	
Cystathionine	Trace		0.0		Trace		0.0		
Isoleucine	0.5	1.3	0.4	1.2	0.5	1.5	0.6	1.7	
Leucine	7.7	19.8	6.1	18 (7		16.7	4.2 .	12.0	
Tyrosine	1.4	3.6	1.4	4.3	1.3	3.9	1.5	: 4.3	
Phenylalanine	4.3	.11.1	2.3	7.0	2.2	.6.7	3.8	10.8	
Y-Aminobutyric ac	id 0.1	0.2	0.1	0.3	0.1	0.3	0.0	0.0	
Tryptophan	0.1	0.2	0.1	0.3	0.1	0.3		0.3	
. Ornithine	0.3	0.8	0.1	0.3	0.2	0.6		1.1	
Lysine	1.2	3.1	. 1.2	3.6	1.4	4-2		2.6	
Histidine	0.2	0.5		0.6	0.2	0.6		0.6	
Arginine	2.6	6.7	1.9	5.8		5.2	1.2	3.4	
Taurine			0.0		0.0		0.0	0.0	
β-alanine	,		- ==		=		1.5	4.3	
	00.0								

Appendix H. Total amino acid composition of citrateHC1 extracts from one-week old . Cheddar cheeses!

Anino Acid	0-16 P	ennet	Crude	con				
WHITHO WELD	Carr	ennec	trude	SUP	Protes	se A	From	ase
THE REST	Tu moleile		umale/e	7 1	umole/g	7	mole/g	
	, /		Munici 8		Winder, S		M	
Cysteic acid	Presen	t	Present		Present		Present	
Glycerophospho-				1.50		61.	7	11 1
ethanolamine								-
Aspartic scid	9.77	7.38	.: 9.55	7.3	8.99	7.4	12:44	7.1
Threonine	4.16	3.1	4.08	3.1	3.56	3.0	5.58	3.2
Serine	8.96	6.7	7.76	5.9	7.40	6.1	9.91	5.7
Asparagine							A	1
Glutamic Acid	29.17	21.8	28.73	22.0	26.98	22.3	138.68	22.1
Glutamine	-		*					****
Proline	13.77	10.3	13.38	10.2	12.18	10.1	20.49	11.7
Glycine	4.10	3.1	4:02	3.1	. 3.74	3.1		2.7
Alanine	4.23	3.2	3.90	3.0	3.61	3.0	5.05	2.9
Citruline		es		-				
Valine	9.37	. 7.0	9.44.	7.2	8.78	7.3	13.72	7.8
Cysteine	0:13	0.0	0.00	0.0	0.00	0.0	0.00	0.0
. Methionine	2.58	1.9	2.30	1.8	1.70	1.4	3.36	1.9
Cystathionine			1				-	1
Isoleucine	6.88	5.1	7.37	5.6	6.71	.5.6	10.26	5.9
Leucine	12.56	9.4 -	13.07	10.0	11.82	9.8	17.17	9.8
Tyrosine	3.15	2.4	2.95	2.3	2.71	2.2	3.86	2.2
Phenylalanine	5.17	3.9	5.20	4.0	4.54	3.8	7.42	4:2
Y-Aminobutyric ac	id							
Tryptophan	() man 2		2					-
Orni thine	-	-		-				
Lysine	11.44	. 8.6.	10.76	. 8.2	10.49	8.7	12.36	7.1
Histidine	3.71	2.8	3.76	2.9	3.36	2.8	4.92	2.8
Arginine	4.53	3.4	4.57	3.5	4.16	3.4	5.28	3.0
Taurine			· 6				,	1
B-alanine	0.00		0.00		0.00		0.00	
The second second second							V 6 0	
Total :	133.68	7 Sec.	130.84	27.7	127.70	4.1	175.26	9
								60

Note (a) Total amino scid (scid hydrolysate).

old Cheddar cheeses.1

Amino Acid '	Calf R	ennet	Crude S	SGP	Protes	se A	From	ase '
					22-10-1			
	umole/g	Z	umole/g	. %	umole/g	. %	Umole/g	Z
Aller and the	1		M		1			
Cysteic acid	0.5	0.1-	0.3	0.1	0.3	0.1	0.3	0.1
Glycerophospho-	,		- 17					
ethandlamine	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
Aspartic acid	22.3	7.9	16.1	7.3	16.0	7.6	23.2	- 7.4
Threonine .	9.8	3.5	6.8	3.1	6.6	3.1	12.0	3.8
Serine	17.9	6.4	13.2	6.0	13.0	6.2	20.4	6.5
Asparagine			*				. *	
Glutamic Acid	56.8	20.2	44.3	20.1	443.6	20.8	61.2	19.4
Glutamine	*				*		*	
Proline	30.4	10.8	23.0	10.4	21.8	10.4	38.3	12.1
Glycine	8.9	3.2	6.5	3.0	6.5	3.1	'8.6	2.7
Alanine	9.1	3.2	: 6.2	2.8	6.2	3.0	8.5	2.7
Citrulline							-	
Valine	13.0	4.6	22.1	10.0	14.7	.7.0	23.1	7.3
Cysteine	- 0.3.	0.0	0.2	0.0	.0.2	0.0	0.4	-0.1
Methionine	4.8	1.7	3.6 .	1.6	3.4	1.6	6.1	1.9
Cystathionine	0.1	0.0	0.0	. 0.0	0.0	0.0	0.1	
Isoleucine	16.0	5.7	11.3	5.1	11.6	5.5	17.5	5.6
Leucine .	26.4	9.4	20.5	9,3	20.1	9.6	28.4	9.0
Tyrosine .	9.2	.3.3	6.5	3.0	6.7	3.2		3.1
Phenylalanine "	: 11.3	4.0	8.2	3.7	7.7	3.7	13.1	4.2
Y-Aminobutyric ac	id		· · ·					
Tryptophan	-							
Ornithine	0.2	0.0	0.1	0.0	0.1	0.0	0.3	. 0.0
Lysine	24.2	8.6	16.6	7.5	17.4	8.3	24.6	7.8
Histidine .	.,7.9.	2.8	5.8	2.6	5.6	2.7	. 8.4	2.7
Arginine	9.5:	3.4	7.1	3.2	5.6	. 2.7	9.1	2.9
Taurine	2.3	0.8	1.7	0.8	2.3	1.1	1.8	0.6
β-alanine	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
11.00								
	280.9		220.1		209.4		315.3	

Note: 1 Total amino acids (Hydrolysate). *Converted to corresponding acids during HCl hydro

Appendix J. Amino acid composition of citrate-HCl extracts from "Forfar"

A	Free		Total	1 1
Amino acid	umole/g	ż	·umole/g	7:
	Maore's.		hangre! 8	. "
Cysteic acid	7.7	3.8	0.9	0.1
Glycerophospho-				
ethanolamine	0.0	0.0.	0.0	0.0
Aspartic acid	3.3	1.6	17.3	2.8
Threonine	7.3	3.6	22.8.	3.7
Serine	4.9	2.4	33.4	5.4
Asparagine	11.6	5.7	*	-
Glutamic Acid	36.7	17.9	144.4	23.3
Glutamine	7.1	3.5	*	
Proline	5.2	2.5	68-1	11.0
Glycine	6.3	3.1	19.5	3.2
Alanine	7.7	3.8	20.0	3.2
·Citrulline ···	3.2	1.6	. 1.5	0.2
Valine	13.4	6.5	46.4	7.5
Cysteine	0.2	0.0	2.1	0.3
Methionine	- 6.1	3.0	10.7	1.7
Cystathionine	. 0.0	0.0	0.3	0.0
Isoleucine	5.0	2.4 .	36.1	5.8
Leucine	. 33.5	16.3	52.9	8.5
Tyrosine	7.6	3.7	. 15.1	2.4
Phenylalanine	11.0	5.4	24.9	4.0
Y-Aminobutyric acid	.0.4	0.2	. 0.1	0.0
Tryptophan 3	0.7	0.3	0.0	0.0
Ornithine	4.4	-2.1	5.6	0.9
Lysine	16.2	7.9	.53.9	8.7
Histidine	2.6	1.3	18.4	3.0
Arginine	2.9	1.4	13.4 .	2.2
Taurine	0.0	0.0	11.0	1.8
\$-alanine	0.0	0.0	0.0	0.0
Total	205.0		618.8	

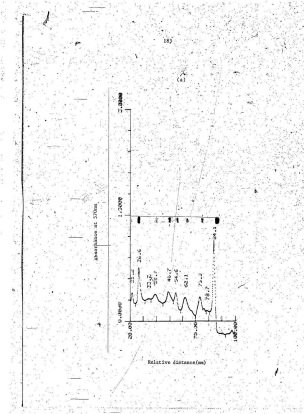
converted to corresponding acid during HCl hydrolysis.

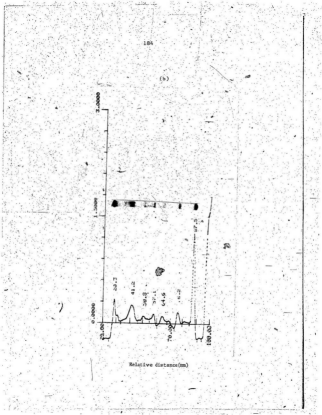
Appendix K. Amino Acid Composition of Whole Casein

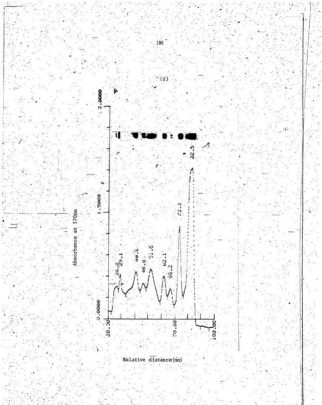
Amino Acid	Percentage Compositi
Aspartic Acid	8.8
Methionine sulphone	0.0
. Threonine	4.6
Serine 3	6.9
Glutanic Acid	18.3
Proline	9.8
Glycine	3.0
, Alanine	4.3
Valine	6.1
Gysteine	0.5
Methionine	4.1
Cystathionine	0.0
Isoleucine	4.5
Leucine	9.2
Tyrosine	4.3
Phenylalanine	3.8
Ornithine	0.0
Lysine	6.8
Histidine	2.3 —

Appendix L. Protein band patterns of citrate-HCl extracts.

Citrate-HCl extracts of 30-week -01d cheeses were prepared following the method of Yakaleria and Price (1959) and logd of extracts were webjected to polyserylamide gel electrophoresis at pil 8.3 (Davis, 1964) and the gels stained with coomassis blue and destained. The gels were then scanned in DUB spectrophotometer. (a) extract from cheese made with processe A, (b) crude SGP, (c) call rennet, (d) Fromase.







Appendix M. Starch gel electrophoresis of Berridge substrate clotted by calf chymosin and protease A.

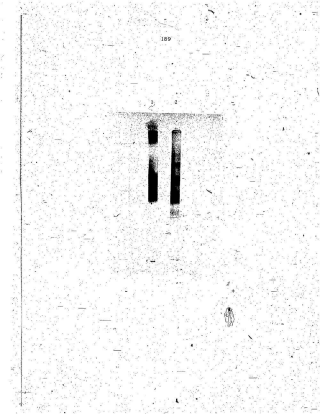
A 0.5 ml sample of Berridge substrate, equilibrated at 00°C was treated with calf chymosin or processe A so that the substrate would clot in about 10 sinutes. The clotted substrate was then dissolved in urea and analysed by starch gel electrophoresis according to Green (1972).

A, calf chymosin-treated, and B, processe A-treated Berridge substrate, C, protesse A-control.

Appendix N. Folyacrylamide gel electrophoresis of zymogen A and its active form.

Polyacrylamide gel electrophoresis in the absence of sodium dodecyl sulphate was performed as described in Materials and Methods.

1. zymogen A purified by affinity chromatography; 2, protesse A prepared by activation of zymogen A.



Judge	Protease	Control.	Fromase	Crude
Number	. A.	rennet		SGP
4		7 10 14 6		
2		7	7	8
3 4	, -	. 5		
4	4	6		
5	14 11 17	6	. 6	. 5
6	1	7	8	7
7.	1 1 1	6	6: "	8
8	6	8	5	8
9	8	4	1	9
10	1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1	6	. 8	7.
11	8	3	8	
12	6 . –	8	5	. 9
14	100	6	3	
15	Τ.	6	4	6
16	8	2 4	2.	5
17	. 8	7.	5	. 8
18		8	7	6
- 19	7	3	4	6
20	1	8	9	
21 22		8		3
23	7	7	5	
24	8	7	5	7
25	4.	5	5	4
26	. 8	9	7	7
27.	-1	6	4	. 3
28		. 6	8	. 4
29*		7	. 6	3
30	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	9		2 ., "
Mean + S.D.	6.87+1,11	6.37+1.71	5.50+1.73	5.77+1 .87
Liked by:	26	24	13 .	16
L. V. V.				
Neither li	ked .		(

nor disliked

Disliked by

9 point hedonic scale.

Appendix P. Sensory preference scores* of 30-week-old Chedder cheese

UDGE	and the state of the	Control	crude
MBER	Protease A	rennet Fromase	SGP
		sensory score	1. 2.
- 0	4. 4. 4.		10
1		5 6	6.
2	1.7	6 7	, 9
3	8	. 8	8
4 =	8	9 8	-6
5	3	7. 2	7.
6	A	8 6	3
7.			8
-8	2	.7 6.	29
19	. 6	-4	- 8-
10	6	- 6. 8 · 5	7
11	. 5	3 4	9
12	5	2 5	7.
13	1 8 4 1	8 6	7
14	5	2 6 5	- 8
15	6	6 4	7
16	. 8	5	. 75
17	1	4 8	9,
8	3	3 2	7.
19	8,	6 7.	5
20	5	6 7	4.
1	6	7	8
2	4	. 7 6	3
3	7	-8 7	8
1	6	5 5	7.
5	. 8	. 4 7	7
5	7	7 7 7 7	6
27 - :	6	7 5	
18	7	7 7	7
9	. 5	3 5	5
0	7.	7	
		Marie Carrier and Artist	
1 0	D. 5.83+1.64	. 5.67+1.97 5.90+1.5	6.83+1

Liked by 18 Judges 18 Judges 20 Judges 26 Judges Netther liked/
Netther liked/
Disliked by 6 Judges 3 Judges 6 Judges 3 Judges
Disliked by 6 Judges 3 Judges 4 Judges 3 Judges

*Rated on a 9 point hedonic scale.

Appendix . O . Ouestionnaire for hedonic scale

S			
Name	Date		
14 19 14 14	1	The Contract of	
You are provided with sample	s of Cheddar cheeses	. Taste the samples and	check how much
you like or dislike each one.	1 0 02		
		The same of the same	
like extremelylike	extremely	like extremely	like extremely
	The second secon		like very much.
		like very much	The same of the sa
		like moderately	like moderately
	/	like slightly	_like slightly
		neither like or dislike	** ***
	manuscriptor and the second	dislike slightly	_dislike slightly
	ike moderately	dislike moderately	dislike moderately .
	The state of the s	dislike very much	_dislike very much
dislike extremelydisl	ike extremely	dislike extremely	_dislike extremely
	I Maria Del Maria	per de la company de la co	
Comments:			
		A 14.	
100		and the second second	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1







