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Original Citation:
The complete amino acid sequence of the low molecular weight cytosolic acid phosphatase / G. Camici; G. Manao; G. Cappugi; A. Modesti; M. Stefani; G. Ramponi In: THE JOURNAL OF BIOLOGICAL CHEMISTRY ISSN 0021-9258 STAMPA 264:(1989), pp. 2560-2567.
Availability: This version is available at: 2158/323440 since:
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The Complete Amino Acid Sequence of the Low Molecular Weight Cytosolic Acid Phosphatase*

(Received for publication, February 1, 1988)

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This paper presents the complete amino acid sequence of the low molecular weight acid phosphatase from bovine liver. This isoenzyme of the acid phosphatase family is located in the cytosol, is not inhibited by L-(+)-tartrate and fluoride ions, but is inhibited by sulfhydryl reagents. The enzyme consists of 157 amino acid residues, has an acetylated NH2 terminus, and has arginine as the COOH-terminal residue. All 8 halfcystine residues are in the free thiol form. The molecular weight calculated from the sequence is 17,953. The sequence was determined by characterizing the peptides purified by reverse-phase high performance liquid chromatography from tryptic, thermolytic, peptic, Staphylococcus aureus protease, and chymotryptic digests of the carboxymethylated protein. No sequence homologies were found with the two known acylphosphatase isoenzymes or the metalloproteins porcine uteroferrin and purple acid phosphatase from bovine spleen (both of which have acid phosphatase activity). Two half-cystines at or near the active site were identified through the reaction of the enzyme with [14C] iodoacetate in the presence or in the absence of a competitive inhibitor (i.e. inorganic phosphate).

Ac-AEQVTKSVLFVCLGNICRSPIAEAVFR
KLVTDQNISDNWVIDSGAVSDWNVGRSP
NPRAVSCLRNHGINTAHKARQVTKEDFV
TFDYILCMDESNLRDLNRKSNQVKNCRA
KIELLGSYDPQKQLIIEDPYYGNDADFET
VYQQCVRCCRAFLEKVR-OH

Acid phosphatases (orthophosphoric-monoester phosphohydrolases (acid optimum), EC 3.1.3.2) are ubiquitous in nature and often occur in multiple forms differing in M_{τ} , substrate specificity, and sensitivity to inhibitors (1-4). In addition, most of these enzymes are glycoproteins and some are also metalloproteins (such as the iron-containing acid phosphatases, porcine uteroferrin and purple acid phosphatase from spleen (5) and bone (6, 7), and the manganese-containing acid phosphatases isolated from some plants (8, 9)). The presence in mammalian tissues of low M_{τ} isoenzymes was clearly demonstrated by Heinrikson (2) who purified the enzyme from bovine liver. Subsequently, De Araujo et al. (4) localized the low M_{τ} acid phosphatase in the cytosol.

Lawrence and Van Etten (10) have recently reinvestigated

the low M_r acid phosphatase from bovine liver. They reported that they had purified the enzyme to homogeneity and criticized Heinrikson's (2) data, particularly with respect to the amino acid composition and specific activity, which they found to be approximately twice as high.

The substrate specificity of the low M_r acid phosphatases is more restricted than that of the high M_r acid phosphatases in that the former efficiently hydrolyzes only p-nitrophenyl phosphate and riboflavin phosphate (2, 11). In 1980, we reported that the enzyme isolated from bovine liver is catalytically very active with acylphosphates such as carbamoyl phosphate and benzoyl phosphate (12). Taga and Van Etten (11) also found that the low M_r isoenzyme from human liver has a high activity on acetyl phosphate and suggested a similarity between the low M_r acid phosphatases and another class of enzymes called acylphosphatases (EC 3.6.1.7). The latter enzymes were extensively studied in our laboratory (13-16): although they have in common a subcellular localization in the cytosol and similar molecular weights, we found that the specificity of the two known isoenzymes of acylphosphatase is limited to acylphosphates and that they do not hydrolyze orthophosphoric-monoesters. In addition, the amino acid sequence was determined for several acylphosphatases from skeletal muscle of vertebrate species (17-23) and for the isoenzyme from human erythrocytes (24).

Recently, Chernoff and Lee (25) demonstrated that the major phosphotyrosyl-protein phosphatase from bovine heart is associated with a low M_r acid phosphatase. They reported that this enzyme appears to be similar to the low M_r acid phosphatases from other tissues, including the liver enzyme. In fact, all these enzymes have similar M_r , pH optima, and K_m values for p-nitrophenyl phosphate, and all are insensitive to inhibition by L-(+)-tartrate and fluoride ions. Furthermore, Boivin and Galand (26) purified two isoenzymes from human red cell cytosol that efficiently dephosphorylate the membrane protein band 3, previously phosphorylated on a specific tyrosine residue by a tyrosine phosphokinase present in the red cell membrane.

EXPERIMENTAL PROCEDURES AND RESULTS¹

DISCUSSION

The complete amino acid sequence of the low molecular weight acid phosphatase (cytosol) from bovine liver is presented in Fig. 1, together with the peptides used to delineate the primary structure. The protein, consisting of 157 amino acid residues, is acetylated at the NH₂ terminus, and has Arg

^{*} This work was supported by grants from the Italian Ministero della Pubblica Istruzione and from the Consiglio Nazionale delle Ricerche, Gruppo Nazionale di Coordinamento "Struttura e Funzione delle Macromolecole Biologiche." The costs of publication of this article were defrayed in part by the payment of page charges. This article must therefore be hereby marked "advertisement" in accordance with 18 U.S.C. Section 1734 solely to indicate this fact.

¹ Portions of this paper (including "Experimental Procedures." "Results," Figs. 2–11, and Tables I–VI) are presented in miniprint at the end of this paper. Miniprint is easily read with the aid of a standard magnifying glass. Full size photocopies are included in the microfilm edition of the Journal that is available from Waverly Press.

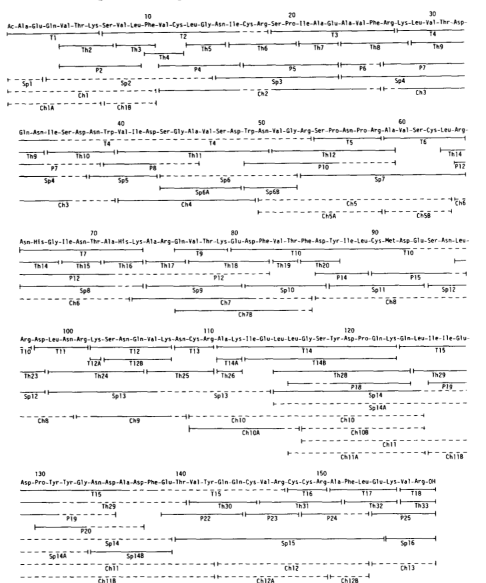


Fig. 1. The complete amino acid sequence of the low molecular weight acid phosphatase from bovine liver. The solid lines indicate the amino acid sequences determined for peptides obtained from trypsin (T), thermolysin (Th), pepsin (P), S. aureus protease (Sp), and chymotrypsin (Ch)cleavages. The notations A and B refer the NH2- and COOH-terminal subfragments, respectively, of a peptide which was also found unbroken. Dashed lines indicate sequence information which was inferred from the amino acid composition of the peptide and from data on sequence analysis of other peptides. Ac, acetyl.

as the COOH-terminal residue. All 8 half-cystines in the acid phosphatase are present as free sulfhydryls. The calculated minimum molecular weight is 17,953.

The protein was first reduced and carboxymethylated to stabilize the cysteine residues. The sequence was determined by analyzing the peptides obtained from five different enzymatic digestions (trypsin, thermolysin, pepsin, S. aureus protease, and chymotrypsin). Peptides were purified by HPLC² on Aquapore RP 300 with a trifluoroacetic acid/acetonitrile-based solvent system. Peaks containing more than one peptide were rechromatographed on the same column with a different solvent system and/or different elution programs.

The sequences were analyzed by the manual Edman degradation. The structure of T1, that is, the NH₂-terminal blocked peptide, was obtained by the combination of FAB mass spectrometry, enzymatic digestions, and Edman degradation, as described in the Miniprint.

The COOH-terminal Arg was determined by treatment of

the Cm-protein with carboxypeptidase B. All cleavage points in the protein were overlapped by peptides obtained from one or more of the other digests.

No homology emerged when the sequence of the low M_r acid phosphatase from bovine liver was compared with that of acylphosphatase from bovine skeletal muscle (22), which indicates that these enzymes are expressed by different genes. Nor was there any homology between acid phosphatase and the isoenzyme of acylphosphatase isolated from human erythrocytes (24). The latter enzyme differs from that of human skeletal muscle in about 44% of the amino acid positions, but they clearly have originated from a common ancestral gene (24). Both of these isoenzymes show a strict specificity for acylphosphates and do not hydrolyze orthophosphoric-monoesters. Thus, although the low M_r acid phosphatase from bovine liver hydrolyzes similar acylphosphate substrates as other acylphosphatases, these share no structural similarities. Hunt et al. (5) have studied the sequences of two metalloglycoproteins, uteroferrin from porcine uterus and purple acid phosphatase from beef spleen, both of which exhibit acid phosphatase activity. Although their sequence data were incomplete, they demonstrated that the sequence homology between these two proteins was >90%. Comparison of these

² The abbreviations used are: HPLC, high performance liquid chromatography; Cm, carboxymethyl; PITC, phenylisothiocyanate; TFA, trifluoroacetic acid; PAGE, polyacrylamide gel electrophoresis; SDS, sodium dodecyl sulfate; FAB, fast atom bombardment; PTH, phenylthiohydantoin; Ac, acetyl.

partial sequences (accounting for about 90% of the entire molecules) with the sequence of the low M_r acid phosphatase from bovine liver shows virtually no sequence homologies. Furthermore, the two proteins (uteroferrin consists of a single polypeptide of 35 kDa, whereas purple acid phosphatase consists of two polypeptide chains of 20 kDa and 15 kDa) have a low cysteine content: uteroferrin contains 2 cysteine residues per molecule, whereas the 20-kDa purple acid phosphatase chain contains only 1, and the 15-kDa purple acid phosphatase chain contains 2. In contrast, the low M_r acid phosphatase contains 8 cysteines per molecule ($M_r = 17.953$). The sequences around cysteine residues in uteroferrin and purple acid phosphatase are different from those around the 8 cysteines of the low Mr acid phosphatase. Uteroferrin and the two chains of purple acid phosphatase have free α -NH₂ groups at the NH_2 termini, whereas the low M_r acid phosphatase has an α-N-acetylated NH₂-terminal residue. Acetylation at the NH₂ terminus has been postulated to be characteristic of proteins synthesized on free polysomes in the cytosol (27). Thus, our results agree with the data of De Araujo et al. (4) on the cytosolic localization of the low M_r acid phosphatase. We found that iodoacetate causes the inactivation of the enzyme and that the competitive inhibitor P_i protects the low M_r acid phosphatase against inactivation (Fig. 10). In agreement with the data of Lawrence and Van Etten (10), our results indicate that at least 1 half-cystine residue is present at or near the active site because 85% inactivation of the enzyme occurred with the carboxymethylation of 0.9 residue of half-cystine per molecule of enzyme (Table I); furthermore, the competitive inhibitor Pi reduced the rate of inactivation by preventing the iodoacetate reaction with active site sulfhydryl group(s). Because of the difference in inactivation of the enzyme by iodoacetate in the presence and absence of P_i (Fig. 10), differential modification by [14C]iodoacetate (28) was used to distinguish essential active site half-cystines from others that might be modified at the same time at other sites of the molecule. We found that Cys-12 and Cys-17, both labeled by [14C]iodoacetate, are protected to the same extent by Pi, so that these two half-cystine residues are at or near the active site of the enzyme. Nevertheless, Cys-12 reacts with iodoacetate 3.5 times faster than Cys-17 (see the specific radioactivity of Cys-12- and Cys-17-containing peptides in the Miniprint). Thus, Cys-12 contributes most to enzyme's inactivation. In this paper, we present the first complete amino acid sequence for an acid phosphatase.

Acknowledgments—We thank the Centro di Spettrometria di Massa of the Medical School of the University of Florence for FAB mass spectrometric analysis.

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Supplementary Material to:

"The Complete Amino Acid Sequence of the Low Molecular Weight Cytosolic Acid Phosphatase" by Guido Camici, Giampaolo Manao Alessandra Modesti, Massimo Stefani and Giampietro Ramponi

EXPERIMENTAL PROCEDURES

Materials. Low M acid phosphatase was prepared essentially as described by Lawrence and Van Etten (10). With their technique however, we were not able to obtain a pure protein; a small quantity of a contaminating protein (about 14 KDa) always remained. Therefore we purified the enzyme to homogeneity by means of MPLC on an Aquapore RP 300 (Brownlee Labs. Inc.) column with a TFA/acetonitrie gradient system and checked its purity by SDS-PAGC or by PAGE in urea-containing gels, as described by Manao et al. (14).

gradient system and checked its purity by SDS-PAGE or by PAGE in urea-containing gels, as described by Manao et al. (14).
Diphenylcarbamylchloride-treated trypsin, pepsinogen, carboxypeptidase B, disopropylfluorophosphate-treated carboxypeptidase B and G-chymotrypsin were obtained from Signa. S.surcus VB protease was obtained from Miles. Thermolysin was purchased from Merck (Darmstadt). Reagents and solvents (sequenal grade) for sequence determination by the Edman degradation technique were obtained from Fluka A.O. Todo [2- C] acetic acid was from Amersham Int., with a specific radioactivity of 56 mCi/mmol.All other reagents used were of the highest purity commercially available.

Determination of free sulfhydryl groups. Total sulfhydryl group content of acid phosphatase was determined both by spectrophotometric titration with 5.5°-dithobis(2-nitrobenzoate). Ellman's reagent (29), and by carboxymethylation of the enzyme with iodoacettae in the presence of 6 M guanidinium chloride but in the absence of reducing agents. The Spectrophotometric titration was carried out by dissolving the protein in 1 ml of 0.1 M Tris-HCl buffer, pH 8.O. containing 0.01 M EDTA and 6 M guanidinium chloride; the mixture was incubated at room temperature for 20 min pijor to addition of Ellman's reagent. E412 = 13,600 M fc cm was used for calculation. Protein concentration was determined by amino acid analysis. The carboxymethylation of the enzyme in the absence of reducing agents was carried out as follows: the enzyme (29 mol) was dissolved in 40 µl of 0.1 M tris-HCl buffer, pH 8.50, containing 8 M guanidinium chloride and the reaction vessel was flushed with nitrogen. After 20 min incubation at room temperature, 40 µl of 64.5 mN indoacetic acid solution (adjusted to ph 8.70 with tris base) was added. The reaction was dissolved in 40 µl of 0.1 M tris-HCl buffer, pH 8.50, containing 8 M guanidinium chloride and the reaction vessel was flushed with nitrogen. After 20 min incubation at room temperature, 40 µl of 64.5 mN indoacetic acid so

carpoxynethylated protein was purified from these and reagents by qelf liferation on a column (2 x 40 cm) of Sephadex S 25 superfine equilibrated with 0.2 K ummonja solution.

***Sinolacid **analysis**. Amino acid manalyses were carried out on 0.5-2 mool of peptides or protein by means of a Carlo no.5-2 mool of peptides or protein by means of a Carlo Erba 3429 amino acid analyzer equipped with an SP 4100 computing integrator (20). Cysteine was determined as Cmcysteine and tryptophan was massayed by the method of Penke et al. (31). Alternatively, amino acid analysis of peptides (70-500 pmol) was performed by analyzing the phenylthiocarbamyl-derivatives of amino acid by HBLC on a Waters 2:oc. Tag write acid analysis column (3.2 x 150 nm, 4 µs), using an sectonitrile gradient.

Enzymic hydrolysis. Before the various enzymatic diagestions, the carboxymethylated moid phosphatase solution in a strew-cap sealed vial was immersed for 5 nm in a boiling water bath and then chilled in ice.

**Tryptic, peptic, 5, aureup protease, and thermolytic digestions were carried out as previously described (241. Chymotryptic digestions were carried out as previously described (241. Chymotryptic digestion of Gm-ard phosphatase (40 nmol) was carried out with a 2.5 % (w/w) chymotryptic in no.2 % amsonium bicarbonate buffer, pH 8.50, for 4 h at 37 °C. With regard to the digestion with carboxypeptidases, Co-acid phosphatase (20 nmol) was dissolved in 0.2 M N-ethyl-morpholline-acctate buffer, pH 8.50, and nixed with carboxypeptidase B at a carboxypeptidase/substrate ratio of 0.02 units per nnol. For the digestion with carboxypeptidase X at a carboxypeptidase/substrate molar ratio of 1.76. The mixtures were incubated at 30 °C aliquous Proteins and chynotryptic peptides were purified by reverse phase HPCC on an Aquapore RP 300 column (4.6 x 250 mm, 10 ym; guard column; 4.6 x 30 om, 10 ym; surfice was purified by reverse phase HPCC on an Aquapore RP 300 column (4.6 x 250 mm, 10 ym; surfice of the condition of peptides of column; 4.6 x 30

Nu_terminal analysis, and sequence determination. Edman degradation: protein (6 nmol) or peptides (1-10 nmol) were submitted to Edman degradation carried our by the samular technique devised by Tarr (32), modified as follows: the conversion from anilinothiaunlinones to PTH-derivatives was performed by incubating the dried thiszolinone extract at 80°C for 10 min with 0.2 ml of 1 N aqueous HCL, containing ethanethiol 11 % vil; then exit of the PTH-derivatives were extracted twice with 0.5 ml of ethylacetate. The organic and aqueous phases were separately dried and analyzed PTH-derivatives present in the organic phase were analyzed by HPLC on a Beckman Ultrosfere 00% collant (4.6 x 250 mm, 10 μ m) according to Bhown et al. (31). The analyses of FTH-His and PTH-Arg faqueous phase) were carried out by HPLC on a small column of Aquapore NP 300 (4.6 x 250 mm, 10 μ m). The elution can be successful to the secretic mode (90 % 0.04 M sodium accetate buffer, pH 4.40, and 10 % nethanol, at a flow rate of) elization. Nug-terminal qualysis and sequence determination. Edman

lecular Weight Lampaolo Manno, stro Ramponi Manno, stro Ramponi

Nomenclature of peptides. Peptides are indicated by prefixes corresponding to the type of cleavage by which they are produced: T. tryptic; Th, thermolytic; P., peptic; Sp. 5. aureus proteose; T. tryptic; Th, thermolytic; P., peptic; Sp. 5. aureus proteose, and Ch, chymatryptic peptides. Chemical modification with indoacetate. Before using, the enzyme was precipitated with 75 % ammonium sulfate, centricitized at 10.000xg for 10 sin, and the precipitate was dissolved in the buffer used for enzyme modification. The specific activity was 110 units/mg of protein (none unit of activity is defined as the amount of enzyme which catalizes the hydrolysms of 1 µmol of p-nitrophenyl phosphate per min at 37 °Cl. Enzyme modification experiments with indoacetate acid were performed in 0.13 M cacedylate buffer, pN 6.30, in a final volume of 2.37 ml and 25°C. Four incubation mixtures were prepared. One was 8.4 mM in indoacetate, a second was 8.4 mM willier in indoacetate and 8.4 mM in indoacetate and 8.4 mM in indoacetate and second was 8.4 mM willier in indoacetate and Pl, and a third was 8.4 mM willier in indoacetate and Pl, and a third was 8.4 mM willier in indoacetate and Pl, and a third was 8.4 mM willier in indoacetate and Pl, and a third was 8.4 mM willier in indoacetate and Pl, and a third was 8.4 mM willier in the control contained only enzyme and buffer, shout 20 mol of enzyme was used for each mixture. At various time intervals, 5 µl aliquots were taken and the activity was measured at pH 5.50 and 37°C, with 4 mM p-nitophenylphosphate as substrate. Isolation of Castboxymethylated peptides. The enzyme 40 mmol was inactivated as described above but with Caindoacetate whose specific radinactivity was bu, 59 Aci/amol. After 140 min, the residual activity was about 15 % and the enzyme was purified from reagents by Sophadex G 25 chromatography (2 x 30 cn column). The protein fraction were pooled, freeze-drued, and tested for amino acid composition and radioac

thread with 4% thermolysin (w/w) at 37°C for 210 min. The thermolytin subfragments were separated by MPLC on Aquapper RP 300. All fractions were tested for 145-radiuactivity. RESULTS

The low Mr acid phosphatase used for the study of the prisory structure was a pure protein as enecked by PAGE and SOG-PAGE. The amino acid composition and the NH— and COOH-terminal analyses of the Ca-maywe are presented in Table 1. The amino acid composition is similar to that reported by lawrence and Van Etten for the same engage (10) and is in good agreement with the amino acid composition calculated from the sequence which shows 8 half cystines. The litration with 5,5'-dithiobis[2-nitrobenoic acid revealed 7.6 equivalents of free sulfnydryl groups per nolecule of enzyme. The carboxymethylation of the enzyme in the presence of 6 M guandinium chloride but in the absence of reducing agents agreed closely to the results obtained with Elinan's reagent (Table I). Both theme findings indicate that all eight half cystine residues occur in the form of Gree thick side chains. Furthermore, Table I shows that the NH_-terminus is not available for the PITC reagent. Arginine was found as the COOH-terminal residue by carboxyperpidase B. Lime-course analysis. The results, expressed as mal ratios, were as follows: 10 min, 0,24 Arg; 30 min, 0,86 Arg; 2 h, 0,89 Arg; 17 h, 1,00 Arg. These results agree with the asino and carboxy termin found in the reconstructed primary structure of the enzyme (Fig. 1). Putificalizing and characterization of, peptides obtained by entities bydrolysis. Fig. 2 shows the preparative WELC finger print of tryptic peptides. The chromatogram was developed by a gradient of 0.01 M TFA in acetonitrile as indicated in the figure legend, using an Aquapper RP 300 column. All peaks were rechromatographed on the same column but with a solvent system consisting of: a) 0,02 M annonium bicarbonate, pH 7.0; b) acetonitrile. The pradient program for acetonitrile concentration of sont of the tryptic peptides in the proparative minerati

large initial peaks. Tables IV-VI report the amino acid compositions of these last three peptide series, whose sequence information is presented in Fig 1. Two peptides (Sp2 and Sp15) partially precipitated during the digestion and were separated prior to fractionation of the mixture on the Aquapore RP 300 HPLC column. The precipitate was dissolved in 6 M guanidinium chloride; thus Sp2 and Sp15 were purified by MPLC on Aquapore RP 300 using the TFA-acetonitrile solvent system and the same gradient elution used for other Sp peptides. Two anomalous cleavages were observed for P20 (Asp 120 Pro 131) bond) and P19 (Asp 136 Pro 131) due to the low pR value and the relatively long incubation time at 37 C during peptid digestion. In fact Hermodomo (34) reported that aspartyl-prolyl bonds are (in some cases) susceptible to cleavage in acidic solutions. As for the partial cleavage at the Asp-Pho bond was probably derived from non-specific digestion, as indicated by its low yield. Kasper (35) has reported that the pepsin susceptible bonds in proteins are those formed by the carboxyl group of all L-amino acids except proline. On the basis of the sequence information from the various peptide series and the amino acid composition of all purified peptides we were able to reconstruct the complete amino acid sequence (Fig. 1).

Structure of Il. This peptide represents the NH, blocked (Edmann-agative) fragment obtained by tryptic hydrolysis of the Ca-enzyme. Its structure was obtained by the combination of the following techniques: i) FAB mass spectrometry of the entire peptide; the FAB mass spectrum in Fig. 9 shows a protonated molecular ion at m/r 717, a value which corresponds to a M. of 716; ii) digestion of II with 5_aureus protease which generated the NH, acylated T15p1 and the T15p2 fragments. These peptides were separated by a small column (3 x 30 mm) of AGSOW-X4 (H-form) fragment obtained by thich corresponds to a M. of 716; ii) digestion of If NH 18_aureus protease which generated the NH, acylated T15p2 was equinced by

In conclusion the structure of Tl was: Ac-Ala-Glu-Gln-Val-Thr-Lys.
Glutamine found at position 3 agrees with the data on the M_p of Tl, calculated from the PAB mass spectrum.
Active site <u>nodification</u>. Fig. 10 shows the inactivation experiments of the low M acid phosphatase by lodeacetate. The kinetic is pseudo-first order both in the presence and in the absence of Pl, a competitive inhibitor of the enzyme. Inorganic phosphate protects the enzyme against inactivation and these results agree with those reported by Lavence and Van Etten (10) on the same enzyme, from which we conclude that at least one half cystine residue occurs at the active site of the enzyme. When 85 % inactivation had occurred (after about 140 min.) further modification was prevented by quickly separating the enzyme from reagents and salts with gel filtration on Sephadex 0.25. Then amino acid analysis was carried out on the modified protein. The results reported in Table 1 indicate that 85 % of enzyme activity loss was related to the carboxymethylation of 0.9 residues of half cystine per molecule of enzyme, while no loss of other amino acid residue was observed. This seems to indicate that one half cystine is involved in the active site of the enzyme. We therefore proceeded too localing the half cystine residue(s) that could be labeled with C-iodoacetate at about 85 % inactivation of the enzyme. The labelled enzyme was then

completely carboxymethylated by unlabelled indoacetate and the Co-protein was purified by MFLC as described in the CE-protein was purified by MFLC as described in the CE-protein was gigested with trypsin. Only 72 incorporated C-radioactivity. This peptide contains both Cys_{1,2} and Cys_{1,2}. Then digested with thermolysin and the subfrageents were separated by MFLC. Figures 11A.8 show the chromatogram and the radioactivity, respectively. Only three peaks were capacitied by MFLC. Figures 11A.8 show the chromatogram and the radioactivity, respectively. Only three peaks were 12 C-labelled; these were hydrolyzed and their amino acid composition and sequence revealed that T2Th1 (specific radioactivity - 0.36 nCl/nnol) corresponds to Val-Cys_{1,2}. T2Th2 (specific radioactivity - 0.38 nCl/nnol) corresponds to Pau-Cys_{1,2} and Cymal) corresponds to Dau-Cly-Asm-(lee-Cys_{1,2}-Arg. The mean specific radioactivity of the Cys_{1,2} and Syscific radioactivity of the Cys_{1,2} and Syscific radioactivity of the Cys_{1,2} and demonstrate that the modification of Cys_{1,2} was responsible for the loss of most of the enzymatic activity, although Cys_{1,2} also reacted but to a lower extent. Cys_{1,2} and Cys_{2,3} are close enough to each other so that it appears possible for the loss of most of the enzymatic activity, and Cys_{2,3} are close enough to each other so that it appears of the enzyme. To clarify this point we labelled the active site of the enzyme was processed in the way described above for the completely against insctivation by inducetate (Fig. 20). The thornolytic poptides from T. were checked for radioactivity and the results, reported in Fig. 11C. clearly demonstrate an efficient protective action by Pi which competes at it indoacetate at the active site of the enzyme. This mentioned was comparable for all three peaks. Lince the calinactivity abound to peptides TThi, TTEL, and TTB3 was obtained in the absence of Pi.

Table I. Amino Acid Composition and Terminal Residues of Sovine Liver Low Molecular Weight Cm-Acid Phosphatase .

Amino acid [®]	Reduced and carboxymethylated	Carboxymethylated without reduction	Sequence values	Modified ^e	Modified Pi
Cm-Cysteine	7.9	7.6	(8)	0.9	0.2
Aspartic acid	21.0	23.2	(24)	23.3	22.7
Threonine	5.6	6.2	(6)	5.9	5.7
Serine	9.0	9.8	(10)	9.7	10.3
Glutamic acid	15.8	16.7	(16)	16.8	16.9
Proline	4.9	4.8	(5)	n.d.	n.d.
Glycine	5.5	6.2	(6)	6.4	6.5
Alanine	9.2	9.5	(10)	9.1	9.4
Valine	13.1	14.2	(15)	15.2	14.5
Methionine	0.8	1.0	(1)	1.1	1.1
Isoleucine	7,8	8.5	(9)	8.5	8.2
Leucine	10.7	11.3	(11)	11.0	11.0
Tyrosine	5.1	4.9	(5)	5.1	5.0
Phenylalanine	6.4	6.1	(6)	6.2	5.8
Lysine	9.2	9.1	(9)	9.0	8.8
Histidine	2.0	2.0	(2)	2.1	2.2
Arginine .	12.7	12.2	(12)	11.7	11.6
Tryptophan	1.8	n.d.	(2)	n.d.	n.d.

NH₂-Terminus: Edman^c : none CCOH-Terminus: Carboxypeptidase B^d : Arg

The results are expressed as residues per molecole of enzyme. Values for Ser and Thr were determined by hydrolyses at 110°C for 22 and 70 h in duplicate and extrapolation to zero time. Determined according to (29). The NN_-terminal analysis was carried out on 5 meol of Cm-acid phosphatase. The carboxypeptidase B digestion was performed on 30 meol of Cm-acid phosphatase. 85 % inactivated by indomentate in 140 min. 1.1 the presence of 42 mM Pl, 12 % inactivated by indomectate in 140 min. n.d., not determined.

Table II. Amino Acid Composition of Tryptic Peptides from Savine Liver Low Molecular Weight Cm-Acid Phosphatese*.

																	_		Yield %	Sequence position.
Tì			1.1		1.6			0.8							1.0				33	1-6
TZ	1.0	1.2		0.8	0.3		1.0		2.2		1.0	1.0	0.2	1.0			1.0		35	7-18
73				٥.€	1.0	1.0		1.8	1.0		1.1			1.0			1.0		70	19-27
74		6.5	1.1	3.1	1.3		2.4	1.1	4.0		1.8	1.2			1.1		1.2	1.8	56	28-53
75		1.1		0.7		1.6											1.0		57	54-58
76	1.2			0.0				0.8	1.0			0.9					1.0		43	59~64
†7		1.7	1.3				1.0	0.9			1,1				1.0	1.0			56	65-73
T10	1.1	3.8	1.1	1.0	2.0				1.1	0.9	0.9	2.0	1.0	2.0			1.0		39	80-97
T11		1.9										1.0					1.1		55	98-101
T14		0.9		0.8	2.0	1.0	1.0	1.0			0.8	2.0	0.9		2.0				12	111-123
F149		0.9		0.4	2.0	1.0	1.1				0.9	2.0	1,0		1.1				44	113-123
T15	1.1	4.2	1.3		5.3	1.1	1.3	1.0	2.0		1.6	1.1	3.1	1.0			1.0		11	124-147
717					1.1			0.9				1.0		1.1	1.2				63	151-155

The digestion was carried out on 50 med of Co-sold phosphates. The values are expressed as solar ratios. Values for Ser and The were corrected for 15 % and 5 % destruction, respectively. Values of contaminating among acts at a level of less than 15 were not reported. The hydrophills pergisses 75, 172, 1723, 1723, 1713, 1714, 1716 and 1716 early 172 were correctly active with PITC and than experised as phesylthicarhiespyl-derivatives (see Fig. 3), and successively sequenced without determining their cannot cooperation. Taild was calculated as notes that now for protein city of protein city.

Low Molecular Weight Acid Phosphatase Sequence

Table III. Amino acid composition of thermolytic peptides from bovine liver low molecular weight Cm-mcid phomphatame.

																-		rieta X	Sequence poestion
Th3									1.3				1.0					35	8-10
154	1.2								1.0				1.0					72	10-12
ThS		c.a					1.1				1.0							73	13-15
The	1.2			1.0		1.0				0.9						1.0		73	16-20
The								0.9	0.9				0.9	1.1		1.0		41	24-28
The		2.1	1.0		1.0				1.0		1.0							96	29-34
Thio		2.0		1.1						0.9							0.9	50	35-39
Thii		2.7		1 - 9			1.c	1.0	:.8	0.8							1.0	36	40-50
Th12		t.5		0.9		2.0	1.0	0.9	1.2							2.3		60	51-59
Th14		0.9					1.0				0.9				1.0	1.0		41	63-67
This		1.1	1.0		1.1				1.0				0.9	1.1				72	77-82
1450		1.0										0.9	1.0					7.	85-87
Th23		0.9									1.0					1.1		53	96-98
4754		2.0		1.0	1.0						1.0			1.1		1.0		14	99-105
Th25	1.2	1.0							9.9					1.2		1,1		34	106-110
Th28		4.1		1.0	2.1	1.0	ι.ο				2.0	0.9		1.0				53	115-124
Th29		4.1	1.0		2.1	1.2	1.1	1.0		1.6	1.0	1.9	1.0					35	125-140
Th 30	1.1				2.1				1.5			0,9						84	141-145
7h31	2.1							:.0	1.5							2.1		12	146-151
Th 32					1.0						1.0		1.0	1.0				41	152-155

The digestion was performed on 90 maol of Co-acid phosphetawa. The hydrophilic puptides Th2, Th10, Th17, Th13, Th20 and Th31 (see Fig. 4) were decivated with P2TC and then apparated as phosphicacramopyl-acravatives (see Fig. 5), and excessively exquenced without determining their manus acid composition (thrus cetain excessively excessive excessiv

Table IV. Amino Acid Composition of Pentic Pentides from Movine Liver Low Molecular Weight Cm-Acid Phosphetase.

																	-		Y:eld %	Sequence physician
P2			0.9						1.0			1.3			1.2				67	4-9
P4	0.9	1.0							:.0		1.5	1.0							16	11-16
PS	1.0		a.3	1.0	;.0	ē. 3		c.5	¢.2		¢.8						Ç.9		36	17-23
P4								1.0	1.0					1.0					13	24~26
P7		3.9	0.9	1.1	1.0				1,2		0.9	1.0			1.5		1.1		15	27-38
P0		1.0		1.0			1.1	1.c	9.9		1.0							3.9	41	39-45
F10		2.1		.,0		1.9	1.3	1.1	1.0								1.9	:.5	19	49-59
P12		3.2	1.7		2.1		1.0	1.9	1.1		1.0			0.9	2.1	1.8	2.0		1.4	64-82
P14		1.0									1.0	1.1	1.0						5.2	36-69
P15	0.9	2.0		1.2	1.1					¢.7		1.2							17	90-96
Ple		1.3		0.9	2.1	1.1	1.0					1.0	1.9		1.0				37	116-124
P19		3.0			1.0	1.1	1.0				1.5		1.9						12	126-135
P20		2.9				0.9	1.3	0.9					1.7						24	130-137
P22			1.0		3.0				1,1				0.8						60	139-144
P23	1.9								1.0								1.0		47	145+148
724	1.1							1.0				1.0		1.0			1.0		39	149-153
P25					1.1				0.9						1.0		1.1		70	134-157

The digestion was carried out on 40 mAol of Cm-acid phosphatase. Other details are the same as in Table II.

Table V. Asino acid composition of \$1 sureue V8 protease peptides from bovine liver low molecular weight Co-acid phosphatase.

																				Sequence
	Ca-Cys													Phe	Cys				Yveld %	pos) tion
Sp1					1.0			1.0											95	1-2
5p2 h	1.2	c. 1	4.5	1.2	1.3		1.4	7	2.6			2.0		1.0	0.1				14	3-14
Sp3	1.0	C . B		0.8	1.0	0.7		1.2			2.2						0.7		21	15-23
Sp4		3.0	0.9	1.0	1.0			0.9	2.0		0.9	1.0		1.0	1.0		1.0		33	24-37
SpS		2.1							1.6		0.9							1.0	33	38-42
5p6		1.7		2.0			2.0	1.0	2.1									0.9	24	43-52
5 p 6 A		1.3		1.9	0.2		1.0	1.0	0.9	0.2									26	43-40
Sp&B		1.0					1.0		1.0									0,9	14	49-52
Sp7	0.8	1.3		2.0		2.1		0.9	1.1			1.0					3.3		11	53-64
598		2.3	1.2				1.0	0.9			1.0				1.2	2.0			5	65-73
Spe			1.0		2.0			1.0	1.2						1.0		1.2		21	74-80
5 p10		2.0	1.0						1.0					1 . 9					69	81-06
Spli	1.1	1.0			1.0					1.0	1.0	1.0	0.9						35	67-93
Sp12		2.0		1.0								1.0					1.5		54	94-98
3p13	1.2	3.3		0.7	5.5			1.0	1 - 1		1.0	1.1			3.0		1.7		46	99-114
Sp14		5.0		1.0	4.0	1.9	2.0	1.2			2.3	3.8	2.7	0.9	0.9				20	115-139
5p14A		2.1									1.7	3.0	2.9		1.0				13	115-133
Sp148		2.9						1.9						1.2					1.4	134-139
5p15 ^b	3.4		1.2		3.0			1.0	1.9			0.9	0.8	0.7			2,1		22	140-154
5p16									1.0						1.0		1.1		75	155-157

The digestion was performed on 40 sool of Cu-sold phosphetase. Betwee poptides partially precipitated during digestion. Other details are the case as in Table II.

- 1	abie VI.	Amino	ecid	composition o	of chymotryptic	peptides	from	bovine	liver	low molecular	weight	Conserve
		••										

	C#-Cys	Asp																	Yould %	Sequence
'ni				1.0				1.0				1.0		1.1					23	i-10
hia			1.0		2.1			1.0	1.0						3.1				27	1-6
h1 8		0.2		1.1					1.0			1.0		1.0					27	7-10
h2	1.8	1.0		1.0	1.1	0.9	1.1	1.6	2.0		2.0	1.1		1.1			1.0		48	11-26
k 3		3.7	0.9	1.0	1.0				1.0		1.0	1.0			1.e		1.0	1.0	34	27-39
14		1.9		2.0			1.2	1.1	1.9		0.8							0.9	58	40-49
h5	1.0	2.0		2.0		2.0	1.0	0.9	1.8			1.0					2.0		19	50-63
hSA		2.0		1.0		1.9	1.0		1.0								8.0		23	50-56
655	1.1			1.1				1.0	1.0			1.1							15	59-63
.6		2.0	1.0				1.1	1.0			1.0					2.1	1.0		14	64-72
h 7		1.1	1.8		1.9			1.0	1.9					2.0	2.2		1.1		6	73-85
h78		1.1	1.9		2.0				1.8					1.9	1.1				42	76-85
	1.0	4.7		1.0	1.0					1.0	0.9	2.9	0.9				1.0		14	86-100
		2.0		1.0	1.0				1.1						2.0		1.0		•	101-106
10	0.7	1.2		1.2	3.0	0.9	1.1	1.0			1.1	3.1	0.9		2.2		1.0			109-125
104	1.0				1.0			0.9			1.0	2.0			1.0		1.0		13	109-116
108		1.0		1.1	2.0	1.0	1.0					1.0	1.0		1.0				20	117-125
111		4.5	0.9	1.2	3.9	1.9	2.2	1.0	1.2		1.6	1.0	3.7	1.5	1.0				7	117-142
11A		1.0		1.1	1.9	1.0	1.1					2 . D	1.0		1.0				12	116-125
51 1 8		3.9	0.9	0.3	1.9	0.9	1.1	1.0	1.0		1.7		2.7	1.1					37	126-142
112	3.3				2.0			1.0	0.9			1.0		1.0			2.0		10	143-153
124	3.2				2.0				1.1								2.0			143-150
,128								0.9				1.0		1.0					13	151-153
.13					1.0				1.1						1.0		0.9		56	154-157

The digestion was carried out on 40 mool of CW-ecid phosphatase. Other details are the same as in Table II.

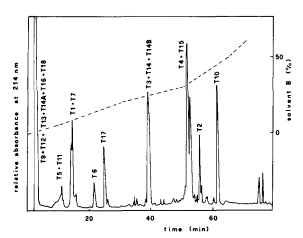


Fig. 2. The HPLC separation of tryptic peptides from 50 nmol of the Cm-acid phosphatase. Column: Aquapore RP 300, 10 µm, 4.6 x 250 mm; guard column: Aquapore RP 300, 10 µm, 4.6 x 250 mm; guard column: Aquapore RP 300, 10 µm, 4.6 x 30 mm, 50/vent A: 10 mM TFA in water: Solvent B: 10 mM TFA in acctonitrile. Flow rate, 1.5 ml/min; {—1, absorbance; {—1, elution gradient. The overlapped peptides were rechromatographed at different conditions. The peptides contained in the big peak near the origin was first derivatized with FITC and then the phenylthiocarbamoyl-derivatives separated by HPLC (see Fig. 3).

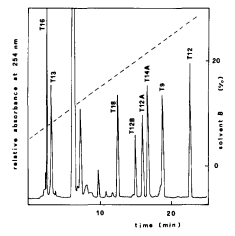


Fig. 3. The HPLC separation of the phenylthiocarbamoylderivatives of the hydrophylic tryptic peptides.

Details are the same as in Fig. 2, except: solvent
A: 0.00 M ammonium bicarbomate, pH 7.0;solvent B: sectonitrile.

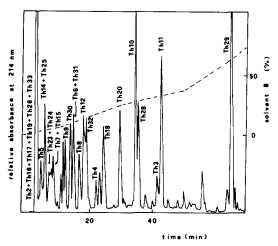


Fig. 4. The HPLC separation of thermolytic peptides from 50 mmol of the Cm-acid phosphatase. Details are the same as in Fig. 2.

The overlapped peptides were rechromatographed at different conditions. The peptides in the big peak near the origin were first derivatized with PITC and then separated by HPLC (Fig.5).

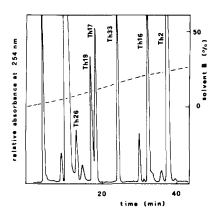


Fig. 5. The HPLC separation of the phenylthiocarbamoylderivatives of the hydrophylic thermolytic peptides. Details are the same as in Fig. 3.

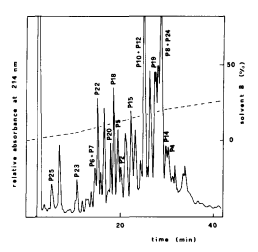


Fig. 6. The HPLC separation of peptic peptides from 40 mmol of the Cm-acid phosphataus. Petails are the same as in Fig. 2. The overlapped peptides were rechromatographed at different conditions.

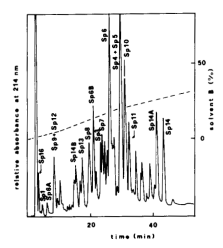


Fig. 7. The HPLC separation of S. aureus protesse peptides from 60 meol of the Ca-acid phosphatase. Details are the same as in Fig 2. The overlapped peptides were rechromatographed at different conditions.

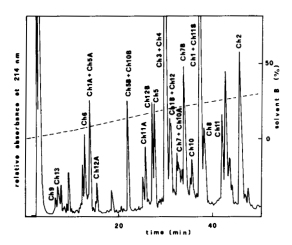


Fig. 8. The HPLC separation of chymotryptic peptides from 40 mmol of the Ca-acid phosphatams. Details are the same as in Fig. 2. The overlapped peptides were rechromatographed at different conditions.

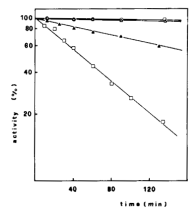


Fig. 10. Inactivation of the low M acid phosphatase by iodoacetate in the presence and in the absence of Pi. Incubations were performed in 0.13 M carndylate buffer, pH 6.3, containing 1 am EDTA at 20°C.

O--O, control: O--O, 8.4 eM iodoacetate;

A---A, 8.4 M indoacetate and 8.4 eM Pi;

A---A, 8.4 M indoacetate and 42 eM Pi.

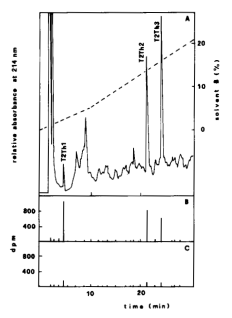


Fig. 11.HPLC separation of the thermolytic subfragments of the peptide 72. A, details are the same as in Fig. ?: B. "C-radioactivity measured on the chromatografic fractions obtained from the enzyme modified at the getive site by "C-indoacetate without Pi; C. "C-radioactivity measured on the chromatographic fractions obtained from the enzyme incubated with ²C-indoacetate but in the presence of 40 mM Pi.

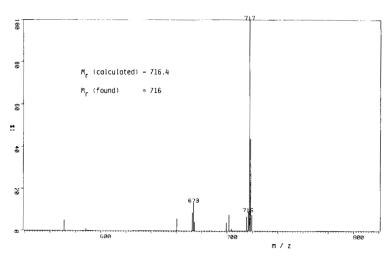


Fig. 9. Positive FAB mass spectrum of Tl.