

Title	STUDIES ON THE ENZYMATIC REACTION CONCERNING LIBERATION OF ETHANOLAMINE IN ASCITES HEPATOMA OF RAT
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# STUDIES ON THE ENZYMATIC REACTION CONCERNING LIBERATION OF ETHANOLAMINE IN ASCITES HEPATOMA OF RAT

- 1. Substrate of the Enzymatic Reaction Concerning
  Liberation of Ethanolamine in Ascites Hepatoma
  of Rat
- Properties of Acetone Insensitive Phosphatase from Ascites Hepatoma of Rat
- 3. Effect of Acetone on Alkaline Phosphatase Activity

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As reported in the previous paper (1), the apparent contents of ethanolamine of hepatoma cells and regenerating liver of rat were about eight to ten times greater than that of normal liver, and this difference was attributed to the effect of acetome, which was used in the determination process of ethanolamine.

Namely, acetome inhibited the enzymatic reaction system concerning the liberation of ethanolamine in normal liver homogenate but not in others.

In the present work, isolation and identification of the substrate of this reaction in hepatoma cells were attempted. It was found that phosphorylethanolamine (PE)\*\* is the direct precursor of ethanolamine in hepatoma cells, AH 130, and that the ensymmtic hydrolysis of PE in the tumor cells was not

<sup>\*</sup> A part of this work was presented at the 34th Congress of the Japanese Biochemical Society held at Osaka in November, 1961.

<sup>\*\*</sup> Follwing abbrevations are used throughout this paper: FE:

phosphorylethanolamine, El: ethanolamine, GPE: glyceryl
phosphorylethanolamine, CDP-E: cytidine-5'-diphosphate ethanol
amine, Pd-E: phosphatidyl ethanolamine, P-Ser: phosphorylserine,

TCA: trichloroscetic acid.

inhibited by acetome.

It has been shown by several investigators (2-4) that PE was present in a large amount in some neoplastic tissues, and the present data agree with their results in that the PE contents of AH 130 and regenerating liver of rat are much larger than that of normal liver.

#### MATERIALS AND METHODS

Male Sprague-Dawley strain rats weighing approximately

200 g. were used. Partial hepatectomy was operated by the method
of Higgins and Anderson (5), by which about two thirds of the
liver was removed. Animals were fed ad libitum before and
after the operation. AH 130 cells were harvested eight to ten
days after transplantation and washed with isotonic seline
before use. The homogenetes were made with Teflon-pestled
Potter-Elvehjem homogenizer in 0.154 H KCl solution. The
soluble fraction was obtained from above homogenetes by centrifugation at 105,000 x g for 60 minutes. FE was synthesized by
the method of Outhouse (6), and glycerylphosphorylethanolamine
(GPE) was prepared from crude phosphatidylethanolamine (Pd-E)
by mild alkaline hydrolysis according to the method of Dawson
(7) and purified by cellulose powder column chromatography

(water saturated phenol: glacial acetic acid: water = 100 : 10 : 12 (v/v)). Cytidine-5'-diphosphate ethanolamine (CDP-E) was purchased from Sigma Chemical Co., St Louis, U.S.A..

Quantitative determination of ethanolamine was made by dinitrophenylation, ether extraction and Amberlite CG-50 column chromatography as previously described (1). Inorganic phosphate was determined by the Fiske-SubbaRew's method (8).

column Chromatography --- (1) Isolation of a precursor of ethenological from AH 130 cells: A 2.5 x 34 cm. column of Amberlite OG-120 (H<sup>+</sup> form, 200-300 mesh) was used. Aliquots of a concentrated dislysate of soluble fraction (15-25 ml.) were applied to the column, and eluted with deionised water. Effluent was collected in 10 ml. fractions. One ml. of samples was taken from each fraction for the analysis of minhydrin positive substances by the method of Yemm and Cocking (2).

(2) Estimation of phosphorylethanolamine in tissues:

Aliquots of Equeous homogenates of a tissue were treated with
an equal volume of 10 per cent trichloroscetic soid (TCA) and
the precipitate was washed twice with 5 per cent TCA. The
supernatent solution and washings were combined and concentrated
to 10 ml. in yaquo in a boiling water bath after removal of TCA

with diethyl ether. One ml. of the sample was applied to an 1 x 30 cm. column of Amberlite CG-120 (H<sup>+</sup> form) or Dowex 50 (H<sup>+</sup> form) and the column was eluted with deionised water. Every one ml. of effluent was collected and PE was quantitatively determined by the minhydrin method (9). Minhydrin colour yield of PE was determined by a synthetic sample in each experiment.

#### RESULTS

Ethanolamine libration in the homogenates of AH 150--- In the previous paper (1), it was found that ethanolamine was produced linearly for at least 5 hours when aqueous homogenates of both normal and regenerating liver were incubated at 37°C, and that the liberation of free ethanolamine was almost completely suppressed in normal liver homogenates, when accetone was added to the homogenates at the concentration of 50 per cent (v/v), while in homogenates of regenerating liver the suppression by accetone was only about 50 per cent. As shown in fig. 1, ethanolamine was also liberated linearly for 5 hours when the homogenates of AH 150 cells were incubated at 37°C (Fig. 1 (A), solid line) and the reaction was not suppressed by 50 per cent of accetone as the case of regenerating liver (1) (Fig. 1 (A),

dotted line). The same phenomena were also observed in the case of the supermatant obtained by centrifugation of the homogenates of AH 130 at 105,000 x g for 60 minutes (soluble fraction) (Fig. 1 (B)). The recovery of the activity in slouble fraction was approximately one third of the original homogenates. Separation of substrate and enzyme --- Separation of a precursor of ethanolamine from the enzyme system concerning liberation of ethanolamine was attempted with soluble fraction of AH 130 cells, as follows: Soluble fraction was dialysed throughly against deionized water with use of a cellophane tube in a cold room, and the dislynate was concentrated to a small volume in vacuo in a boiling water bath. As shown in Table I, ethanolamine was liberated only when dislysed soluble fraction was incubated together with the concentrated dialysate at 37°C. This fact suggests that a substrate of the ethanolamine liberating reaction was separated in the dialysate from the enzyme system. The possibility that a compound separated in the dialysate was not a substrate but a cofactor for the reaction, was ruled out by the experiment described below. Isolation and identification of substrate of the reaction---The concentrated dialysate of soluble fraction was subjected to a column (2.5 x 34 cm.) of Amberlite CG-120 (H form).

and elution was carried out with deionised water. Fig. 2 shows an example of the chromatogram. Authentic phosphorylserine and taurine were eluted at a column volume but authentic sample of PE was eluted just in the Same position as the 2 md peak in the chromatogram. The elunte corresponding to,2 nd peak was concentrated and the concentrate was incubated with dislysed soluble fraction of AH 130 cells for 2 hours at 37°C. The liberated ethanolamine and inorganic phosphate were estimated by DMP-method (1) and the method of Piske-Subballow (8). As shown in Table II, only when both concentrated solution of the 2nd peak and dialysed soluble fraction were incubated, liberation of ethanolamine and inorganic phosphate was observed. In this case, the ratio of liberated ethanolamine to liberated phosphate was found to be nearly 1 (Table II). Under the same condition, liberation of ethanolamine was not observed with use of substances in the first peak of the chromatogram in Fig. 2.

The substance of the 2nd peak was compared with an authentic sample of PE by means of paper-chromatography and paper-electropheresis, as shown in Fig. 5. All maps showed that the pursued substance gave one minhydrin- and molybdate reagent (1)- positive spot and the position of the spot coincided with the reference one; So other minhydrin-positive spot was detected. Furthermore, the substance in the 2nd peak was analysed for contents of

amine group and phosphorous (Table III). It was found that the molar ratio of amine group to phosphorous was nearly 1.

In oder to examine whether there is snother compound, beside PE, which liberates ethanolamine in AH 150 cells. following experiment was undertaken: The concentrated dislysate and its chromatographically purified fraction (2nd peak in Fig. 2) were incubated respectively with dislysed soluble fraction, until liberation of ethanolamine stopped. Assuming that amounts of precursors of ethanolamine were equal to amounts of ethanolamine liberated under the condition described above. amounts of the precursors in the dislysate and its purified fraction were compared. It was found that almost all amount of the precursor of ethanolamine found in the dielysate was recovered in the 2nd peak, PE fraction of chromatography on Amberlite 06-120. Therefore, it is thought that there is no other compound, beside PB, which participates in liberation of ethanolamine in soluble fraction of AH 130 cells. Namely, the direct precursor of ethanolamine in soluble fraction of AH 130 cells is considered to be phosphorylethanolamine. Effect of acetone on the reaction --- With use of dislysed soluble fraction of AH 130 cells and normal liver as ensyme sources, the rates of enzymatic hydrolysis of authentic PE

were compared in the absence and presence of acetone (Table IV). It can be seen in Table IV that the enzymatic activity of AH 130 is much greater than that of normal liver and that the activity is not suppressed by 50 per cent acetone for both AH 130 and normal liver.

When GFE, CDP-E and crude cephaline were added to the enzyme solution (dialysed soluble fraction of AH 130 cells), the rates of ethanolamine liberation from these compounds were small, compared with the case using PE as substrate, and the liberation of ethanolamine from these compounds were inhibited by 50 per cent acetone in a greater extent (Table IV).

Phosphorylethanolamine contents in AH 130 cells, regenerating and normal liver essentiated in AH 130 cells, regenerating and normal liver of rat were estimated as described in Materials and Methods. As shown in Table V, AH 130 cells contained PE about eight times and regenerating liver (48 hours after partial hepatectomy) about three times greater than normal liver.

#### DISCUSSION

In the previous paper (1), it was shown that an enzymatic reaction system concerning the liberation of ethanolamine is present in homogenates of normal and regenerating liver of rat.

The present work revealed that this system also exists in both homogenates and soluble fraction of an ascites hepatoma of rat (AH 130).

By means of dialysis, the precursor of ethanolamine was isolated from soluble fraction of AH 130 cells and proved to be phosphorylethanolamine. No other compounds which participate to the liberation of ethanolamine, beside FE, were found in the soluble fraction. Furthermore, the presence of a phosphatase hydrolysing FE was also proved in soluble fraction of both AH 130 cells and normal liver. Hydrolysis of FE by this enzyme was scarcely inhibited by 50 per cent of acetone in the case of normal liver as well as AH 130 cells (Table IV). Therefore, the fact that the ethanolamine liberation in normal liver homogenates was completely suppressed in 50 per cent acetone in contrast to homogenates of AH 130 cells (1), would be explained as mentioned below. For the ethanolamine liberation, two pathways are considered in mammalian tissues as follows;

- (1) hydrolysis of PE by phosphatase (10)
- (2) hydrolysis of GPE by phosphodiesterase (11), which is produced from Pd-E by phosphatidase A and lysophosphatidase (12). Reaction (1) was proved to be acetome insenstitive (Table IV) but reaction (2) may be acetome sensitive because the hydrolysis of GPE by dialysed soluble fraction of AH 130 was inhibited by acetome (Table IV). Hence, ethanolamine liberated

in homogenates of normal liver and AH 130 would be due to both reaction (1) and (2) in the absence of acetome. When acedome is added to the homogenates, ethanolamine would be liverated only by reaction (1), since reaction (2) is suppressed by acetome. Thus, the difference in the ethanolamine liberation between normal liver and regenerating liver or AH 130 cells, observed when these homogenates were incubated with acetome, would be attributed to the quantitative difference of acetomesensitive (reaction (2)) and acetome-insensitive (reaction (1)) enzyme systems. In fact, the amount of PE was much larger in AH 130 than in normal liver, and phosphatase is also present in a large amount in hepatoms, as reported by Greenstein (13).

As for the formation of PE in tissues, two pathways are known as follows:

- (3) phosphorylation of ethanolamine by phosphokinase (14,15)
- (4) hydrelysis of Pd-E by two-steps reactions by PE-

glyceride transferase and FE-cytidyl transferase (16,17).

The reactions (4) are considered to play a synthetic role for Pd-E rather than a degradative role. The fact that PE content in hepatoma is much larger than normal liver, would be explained by two ways; stimulation of reaction (3) in hepatoma, or

suppression of the formation of Pd-E from PE by reactions (4), and the physiological significance of this elevated PE content of hepatoma is now under investigation.

The properties of the enzyme hydolysing PE will be reported in a forthcoming paper.

#### SUMMARY

- 1. The ethanolamine liberating ensymmtic system was also present in AH 130 cells.
- 2. The main precursor of free ethanolamine in an ascites hepatoms of rat (AH 130) was identified as phosphorylethanolamine.
- 3. Phosphatase which hydrolyses phosphorylethanolamine was not inhibited at all even in the presence of 50 per cent of acetone in the case of normal liver as well as AH 130 cells.
- 4. Phosphorylethanolamine contents of ascites hepatoma and regenerating liver of rat were larger than that of normal rat liver.
- 5. The difference between normal liver and regenerating liver or tymour cells of rat in the reaction of formation of ethanolamine (1) was discussed from the results decribed above.

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Table I

Liberation of Ethanolamine from a Compound Isolated in the Dialysate of Soluble Fraction of AH 130 Cells.

Soluble fraction of AH 130 cells was dialysed against deionized water and the dialysate was concentrated to a small volume as described in the text. Dialysed soluble fraction was incubated with and without the concentrated dialysate at 37°C for 2 hours in 0.05 M veronal buffer (pH 8.5) and 1 mM MgCl<sub>2</sub>. The reaction was stopped by heating in a boiling water bath for 5 minutes and ethanolamine was estimated by DNP-method (1).

	Ethanolamine (proles/flask)		
- 1	1	2	3
Dialysed Soluble Fraction (without incubation)	0.00	0.00	en de
Dialysate (without incubation)	0.878	0,190	0,190
Dialysed soluble Fraction (incubated)	0,00	0.00	0,009
Dialysed Soluble Fraction Plus Dialysate (incubated)	2,40	1,21	0.833

Table II

Enzymatic Digestion of the Compound Isolated in the 2nd

Peak of the Chromatogram of Fig. 2.

		El**	Pi**	Pi/El
(1)	Concentrated Fraction of the 2nd Peak in the Chromatogram of Fig. 2 (without incubation)	0.00	0.00	
(2)	Dialysed Soluble Fraction of AH 130 (incubated*)	0.00	0.00	***
(1)	+ (2) (incubated)	3,09	3.25	1.05

- \* Incubation was carried out at 37°C for 2 hours in 0.05M veronal-carbonate buffer (pH 9.5) and in 1 mM MgCl<sub>2</sub>.
- \*\* Ethanolamine was determined by DNP-method  $(\underline{1})$ , and inorganic phosphate by Fiske-SubbaRow's method  $(\underline{8})$ .

Table III

## Contents of Amino Group and Phosphorous in the Compound Isolated in the 2nd Peak of the Chromatogram

Exp. No.	Amino Group* (µmoles/ml.)	Phosphorous (µmoles/ml.)	P/Amino Group
1	2,40	2.60	1,08
2	3.17	3.31	1.04

<sup>\*</sup> Contents of amino group were estimated as PE by the ninhydrin method .

Table IV

#### Effect of Acetone on Enzymatic Hydrolysis of Various Compounds

Dislysed soluble fractions of normal rat liver and AH 130 cells were used as enzyme sources. Incubation was carried out at  $37^{\circ}$ C for 1.5 hours in 0.05 M veronal-carbonate buffer (pH 9.5) and 1 mM MgCl<sub>2</sub> in the absence and presence of acetone (50 per cent (v/v)).

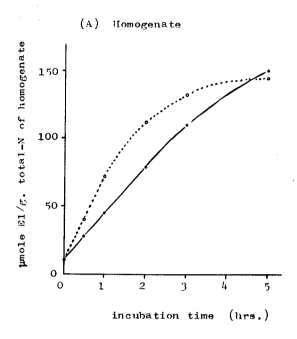
Ensyme Source	Substrate	Specific Activity (El umoles/g-N/hr.)		Acetone Effect*	
	(Alima Ouis)	+Acetone	-Acetone	(%)	
Normal Rat Liver	PE (1.Comm)	32.6	23.5	72.0	
	PS (1.5aM)	23.4	18.1	74.2	
AH 150 Cells	PS (1.51M)	1405.	1558.	109,	
	PB (2.4mM)	2840.	3335.	117.	
	GFE (0.69m	M) 20.2	2.2	10.9	
	CDP-E (0.75mM	278.	41.6	15.0	
	Cephalin (4.3mg/	35.6	18.9	53.2	

<sup>\*</sup> Acetone effect = Specific Activity (+Acetone) x 100
Specific Activity (-Acetone)

Table V Phosphorylethanolamine Contents in Various Tissues.

		number   n
Formal ret liver	0.428 ± 0.023	16.1 ± 0.43
Regenerating rat liver (48 hrs after partial hepatector	y) 1.17 ± 0.19	44.8 ± 9.0
AH 130 cells	2.56 ± 0.68	131 ± 22

<sup>\*</sup> Mean value and standard error of three experiments.



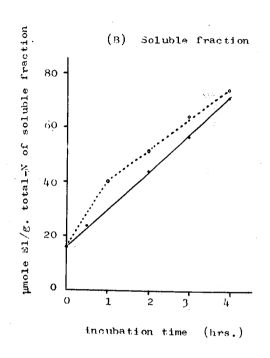


Fig. 1 Time curves of the reaction liberating free ethanolamine in homogenate(A) and soluble fraction(B) of rat ascites hepatoma cells. Incubation was carried out at  $57^{\circ}$ C, in the absence (solid line) and presence (dotted line) of 50 per cent acetone( $\sqrt{v}$ ).

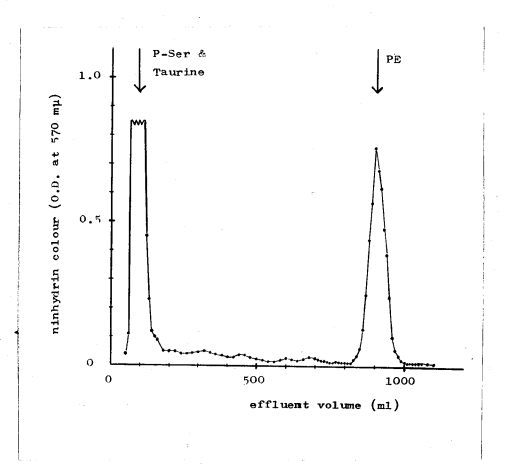


Fig. 2 Column chromatography of the concentrated dislysate of soluble fraction of AH 130 cells.

Fifteen ml. of the concentrated dialysate of soluble fraction ( prepared from 45 ml. of packed cells of AH 130 ) was applied to a 2.5 x 34 cm. column of Amberlite CG-120 (H<sup>+</sup>form) and elution was carried out with deionized water. Arrows indicate the elution positions of authentic samples.

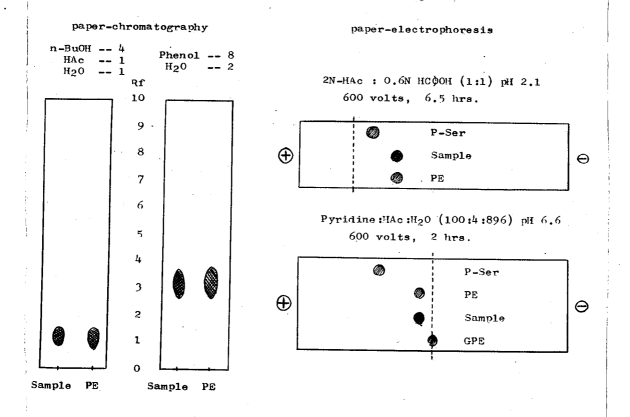


Fig. 3 Patterns of paper-chromatography and paperelectrophoresis of a compound isolated in the 2nd peak of the chromatogram of Fig. 2.

#### PROPERTIES OF ACETONE INSENSITIVE PHOSPHATASE PROM ASCITES HEPATOMA OF RAT

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### PROPERTIES OF ACETONE INSENSITIVE PHOSPHATASE FROM ASCITES HEPATOMA OF RATY

#### MASAYORI INOUYE

Ensymatic reaction in which ethanolamine is produced from phosphoethanolamine occurs in homogenates of ascites hepatoma of rat and of normal rat liver (1.3). The activities in the two preparations differ in that the reaction is inhibited by the presence of organic solvents in 50 per cent concentration in normal liver homogenates but not in hepatoma homogenates (1.2).

In the present work the enzyme which hydrolyses phosphoethanolamine has been partially purified from rat ascites
hepatoma. Some properties of this enzyme and the effect of
organic solvents on its activity have been studied. It was
found that this enzyme is the same enzyme known as non-specific
alkaline phosphatase. Whereas it was active toward phosphoethanolamine even in the presence of organic solvents, it
was markedly supressed by organic solvents in its activity
toward f-glycerophosphate, the usual substrate of alkaline
phosphatase. Furthermore, the level of the enzymatic activity
and the effect of acetome on the activity of preparations
from hepatoma cells, regenerating rat liver, and normal rat
liver were compared.

<sup>\*</sup> A part of this work was presented at the 9 th Kinki Local Meeting of the Japanese Biochemical Society held in Mara in June, 1962.

#### MATERIALS AND METHODS

Rats----Male Sprague-Dawley strain rats weighing approximately 150 g. were used.

Hepatoma ---- Ascites hepatoma cells, AH 130, were harvested by laparotomy 8 to 10 days after transplantation and washed with cold isotonic saline.

Regenerating Liver ---- Rats were partially hepatectomized as described in the previous paper (1). Livers, 48 hours after the operation, were used.

Homogenization ---- Normal liver and regenerating liver were perfused with cold isotonic saline and homogenized in 3 volumes of saline with a Teflon-peatled Potter homogenizer. Hepatoma cells were also homogenized as described above. Ten ml. of homogenates were dialyzed overnight against about 2 liters of deionized water at 3°C, befor enzyme assay.

Ensyme Assay ----- Activity was estimated from the liberated inorganic phosphate by the method of Fiske-SubbaRow (4).

The usual assay system consisted of 0.3 ml. of the enzyme solution, 0.1 ml. of 0.1 M substrate, 0.1 ml. of 10<sup>-2</sup> M MgCl<sub>2</sub> and 0.5 ml. of 0.1 M veronal-carbonate buffer, pH 9.45, in a total volume of 1.0 ml.. The reaction was started by adding the enzyme solution, and the mixture was incubated at 37°C. The reaction was stopped by adding 1 ml. of 10 per cent trichloro-acetic acid, and the mixture was centrifuged. One ml. of the supernatant was taken for determination of inorganic phosphate. The amount of inorganic phosphate liberated was corrected for

the amount liberated from the enzyme preparation in the absence of substrate. When inhibitor was used, 0.1 Ml. of inhibitor solution was added instead of MgCl. In the experiments with organic solvents, 1 ml. of organic solvent or of organic solvent plus water was added to 1 ml. of the reaction mixture described above; for instance, 0.5 ml. of selvent and 0.5 ml. of water were added to 1 ml. of the reation mixture to make the final concentration of 25 per cent solvent. Since the color yield in phosphate estimation changed according to organic solvents involved in the reaction mixture, the amount of inorganic phosphate liberated was corrected for each case. Phosphoethanolamine and  $\underline{\beta}$ -glycerophosphate were used as substrates. One unit was defined at the amount of enzyme which liberates 1 pmcle of inorganic phosphate per hour under the conditions described above. The specific activity was expressed as units per mg. of total nitrogen of ug. of phosphate of DNA. \*\*\* Nitrogen contents were determined by the micro Kjeldahl method. DMA contents were estimated according to the method of Schneider (5).

<u>Materials</u>————Phosphoethanolamine was synthesized by the method of <u>Schmidt</u> (6) (mp. 233-4°C, uncor.), and proved to be paper-chromatographically pure.

Pertial Purification of the Ensyme from Ascites Hepstoms Cells

+---The ensyme was extracted from the homogenate of tumour

cells with use of n-butanol as described for alkaline phos
phatase by Morton (7) as follows: Half volume of n-butanol

<sup>\*\*\*</sup> The following abbreviations are used: DMA; deoxyribonucleic scid, EDTA; ethylenediamine tetrascetic scid, PCMB; p-chloromercuribensoate.

was added to the homogenate of tumour cells, and the mixture was incubated at 37°C for 1.5 hours with occasional stirring.

After incubation, the mixture was centrifuged at 4,000 r. p.

m. for 15 minutes, and the residue was extracted twice with was water. The supermatants were collected, and the n-butanol layer was removed with a pipet (Fr. I in Table I). Cold acetone was added to the supermatant, and the precipitate formed between 25 to 50 per cent of acetone was collected and dissolved in a small amount of water (Fr. II in Table I). Before the ensymatic activity was assayed, the fractions were always dialyzed two times against about 50 volumes of deionized water at 3°C.

About 13 fold purification was achieved. The procedure for purification is summarised in Table I.

(Table I)

#### results

pH-Activity Curves----Fig. 1 shows pH-activity curves with use of the homogenate of tumour cells. Activities at each pH are expressed as per cent of the activity at pH 9.40 measured with β-glycerophosphate as substrate. As shown in Fig. 1, the pH optima are 9.4 for both phosphoethanolasine and β-glycerophosphate under the conditions used. Similar results were obtained for the engyme partially purified from tumour cells. At acidic pH, 5.8 and 4.5, the activities, measured with phosphoethanolasine, were only 0.45 and 0.76 per cent, respectively, of the activity at pH 9.40. These facts suggest that the ensyme

hydrolysing phosphoethanolsmine in the homogenate of tumour cells (3) is what is known as non-specific alkaline phosphatase.

(#18. 1)

Effect of Various Compounds on the Reaction ---- With use of the partially purified ensyme from tumour cells (Fr. I in Table. I), the effect of magnesium ion and various other compounds on the reaction were investigated. As shown in Fig. 2, magnesium ion acts as activator of alkaline phosphatase from tumour cells, and its presence at 10<sup>-5</sup> μ concentration results in about 45 per cent activation for both substrates. Ratios of the activity measured with β-glycerophosphate to that with phosphoethanolamine differ considerably according to the concentration of magnesium ion (Fig. 2-A). Similar phenomena were also observed in experiments with other inhibitors, as shown in Table II.

(Fig. 2)

(Table II)

It can be seen in Table II that EDTA and potassium cyanide markedly inhibit both activities of the ensyme. Zinc ion at

10<sup>-3</sup> M inhibits about 90 per cent of the activity for both substrates. Cadmium ion is also inhibitory, whereas calcium ion is without effect, under the conditions used. PCMB does not inhibit the activities at 10<sup>-3</sup> M. Ethanolamine and glycine stimulate the enzymatic activities. These results, mentioned above, are similar to those obtained for several alkaline phosphatases from different species (8-12).

Effect of Acatone on the Reaction ---- The time curves of the reaction in the presence or absence of 50 per cent acetone are shown in Fig. 3, with use of the enzyme of Fr. I in Table I. It can be seen in Fig. 3-A that the activity measured with phosphoethanolamine is scarecely affected by 50 per cent acetone up to 40 minutes incubation, while the activity measured with β-glycerophosphate is markedly suppressed by 50 per cent acetone and is completely inhibited after 20 minutes produced incubation. In order to determine whether there are two kinds of alkaline phosphatases, acetone-sensitive and acetone-insensitive, the following experiment was undertaken: The ensyme solution was

#### (Fig. 3)

incubated at 37°C in the presence of 50 per cent acetone and  $10^{-3}$  M of Mg<sup>++</sup> without added substrate, and at intervals of 5 minutes 1 ml. of the solution was taken and added to 0.1 ml. of 0.1 M substrate. After 9 minutes incubation at 37°C, inorganic phosphate liberated was measured as mentioned in Methods.

The results are shown in Fig. 4. In the case of phosphoeithanolamine the activity gradually decrease just as in the case of  $\beta$ -glycerophosphate. By 20 minutes preincubation in the presence of 50 per cent acetone without added substrate, the activities were almost completely suppressed for both substrates. Furthermore, no inorganic phosphate was liberated from phosphoethanolamine, when it was added to the reaction mixture with use of  $\beta$ -glycerophosphate in 50 per cent acetone after 20 or 30 minutes incubation (broken line in Fig. 3-B). These facts suggest that phosphoethanolamine has a protective effect on the enzyme protein, not possessed by  $\beta$ -glycerophosphate.

The enzyme solution was incubated for 30 minutes at 37°C in 50 per cent acetone in the absence of substrate and then centrifuged. The precipitate was dissolved in a small amount of water, and this solution was assayed after overnight dislysis at 3°C against 50 volumes of deionized water. The recovery of the activities for phosphoethanolamine and g-glycerophosphate were 95.3 and 81.6 per cent of the original activities, respectively. Therefore, inactivation by acetone appears to be reversible.

Effect of Various Organic Solvents ---- Fig. 5 shows the effects of various organic solvents on the reaction with use of phosphoethanolamine as substrate. It can be seen that dioxan, acetone and dimethylformamide increase the ensyme activity. Notably, about 80 per cent increase in activity was obtained with 30 per cent dioxen. In contrast to ribonuclease (13), which is activated by dimethylformemide and 2-chloroethanol, the ensyme of the present study was inhibited markedly by dimethylformswide at 50 per cent concentration and by 2chloroethanol at much lower concentration. n-Propanol significantly increases the activity of the ensyme toward phosphoethenclamine and ethanol has a similar effect to a lasser extent in contrast to their inhibitory effect on ribonuclease (13). Methanol shows increase of inhibition with increase of concentration. The effect of chain length of alcohols on inhibition or activation was previously observed with ribonuclease (13) and the enzyme of the present study (2).

#### (Fig. 5)

Fig. 6 represents the effect of various organic solvents on the reaction with use of g-glycerophosphate as substrate. In contrast to the results obtained with phosphoethanolamine, all solvents used show increase of inhibition with increase of their concentration.

Effect of Acetone on the Reaction in the Case of Regenerating
Liver and Normal Liver of Rat----Pig. 7 shows the effect of
acetone on the activity of alkaline phosphatase from regenerating liver and normal liver of rat. The ensyme was partially
purified by the same method described for alkaline phosphatase
from tumour cells (see in "Methods"). About 15 and 4 folds
purification was achieved for regenerating and normal liver,
respectively.

(Pig. 7)

The similar results as for alkaline phosphatase from tumour cells (Fig. 3) were obtained for those from both regenerating and normal liver. Namely, 50 per cent scattone scarcely affected the hydrolysis of phosphoethanolamine but inhibited markedly the hydrolysis of  $\beta$ -glycerophosphate.

Alkaline Phosphatase Activity of Tumour Cells. Regenerating
Liver and Normal Liver of Rat----As shown in Table III,

(Table III)

the enzyme activity referred to DNA-P and total mitrogen was about 5 and 18 times, respectively, greater in tumour cells than in normal rat liver. The activity of regenerating liver was also larger than that of normal liver as already reported (14-16).

#### DISCUSSION

In previous papers (1-3), it was revealed that an ensymptically catalysed reaction for the liberation of ethanolamine occurs in homogenates of rat ascites hepatoms cells and normal rat liver, and that this reaction occurs in the homogenate of tumour cells even in the presence of 50 per cent acetome but not in the homogenate of normal liver. Furthermore, the substrate of the reaction was shown to be phosphoethanolamine (2). In the present paper, several properties of the reaction were investigated with use of the enzyme partially purified from tumour cells. From the fact that pH optimum of the reaction was 9.4, which was the same as that measured with g-physero-phosphate as substrate, and the results of experiments with various activators and inhibitors, it is concluded that the enzyme is the same as the enzyme known as non-specific alkaline phosphatase.

As shown in Figs. 3 and 7, 50 per cent acetone scarcely inhibited the hydrolysis of phosphoethanolamine by enzymes prepared from regenerating liver and normal liver as well as by the enzyme prepared from tumour cells. Therefore, the difference between tumour cells and normal liver, as mentioned

above, is not due to a difference in the acetone insensitive property of the ensyme, but may be due to differences in the contents of alkaline phosphatase and its substrate, phosphoethanolamine, in the two tissues. In fact, the content of phosphoethanolamine in ascites hepatoma, AH 130, was about 8 times as much as that of normal ret liver (3), and alkaline phosphatase activity of tumour cells was also 18 times as much as that of normal liver on the base of total nitrogen (Table III).

Was found in the present paper that alkaline phosphatase was quite active in several organic solvents, of even activated by these solvents, when phosphoethanolamine was used as substrate. But 2-chloroethanol and dimethylformamide, especially the former, inhibited the activity, in contrast to the marked activation of ribonuclease by these solvents as reported by E1341 (13).

Therefore it is thought that the folding of the ensyme protein, caused by dimethylformamide and 2-chloroethanol as reported on ribonuclease by Tang et al. (17) and Webb et al. (18) is not suitable for the ensymatic action of alkaline phosphatase. Since organic solvents may affect physicochemical properties of the ensyme protein, the mechanism of the activation by organic solvents will be unknown until the changes of the ensyme protein caused by solvents are elucidated.

It is also interesting that alkaline phosphatese activity was markedly suppressed by all solvents examined, when geglycerophosphate was used as substrate, in contrast with phosphoethanolamine. The causes of this difference are under investigation.

#### SUMMARY

- The properties of the enzymatic hydrolysis of phosphoethanolamine were examined with use of the enzyme partially purified from AH 130 ascites hepatoma of rat.
- 2. From its pH optimum and experiments with various activators and inhibitors, it was concluded that the enzyme is the same as non-specific alkaline phosphatase.
- 3. The effect of various organic solvents on the activity was investigated. It was found that the hydrolysis of phosphoethanolamine was activated by several organic solvents, but that the hydrolysis of β-glycerophosphate was markedly suppressed by all solvents examined.
- 4. Partially purified alkaline phosphatases from regenerating liver and normal liver of rat behaved similarly to enzyme the from hepatoma cells in the presence of acetone.
  - 5. The difference between normal liver, and regenerating liver and tumour cells of rat in the reaction of formation of ethanolamine (1-3) was discussed from the results described above.

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Table I

# Summary of the Purification Process of Alkaline Phosphatase from Tumour Cells

The enzyme assay was carried out at  $37^{\circ}$ C with use of  $\underline{\beta}$ -glycerophosphate and phosphoeths-nolamine as substrates. Each fraction was dialyzed before use. In each experiment, a time curve was constructed and the activity was estimated from an initial velocity.

		Substrate							
Praction	Operation	E-grac	erophos	phate(GP)	Phosphoe	thanola	nine(PE)	GP/PB	
		Total	Activity		Total	Activit;			
		Unite	Yield(%)	- Specific ) Activity (Units/mg.		Yield(% (U	- Specific ) Activity aits/mg. N)		
Homogenate		8.99x104	100	18.8	8.18x10	100	4.	1.10	
	tration with n-butanel	7.03×10	78.3	162.	6.39x10	78.2	147.	1.10	
	ectionstics with acetone	6.00210	66.7	238.	6.00×10	73.3	<b>238.</b>	1.00	

Table II

Effect of Various Compounds on the Reaction with Use of Phosphoethanolamine and  $\underline{\beta}$ -Glycerophosphate as Substrates

The enzyme assay was carried out at pH 9.45 and 37°C under the conditions shown in the table. The enzyme partially purified from tumour cells (Fr. II in Table I) was dislysed overnight against 50 volumes of deionised water at 3°C befor use. Incubation was carried out for 15 minutes with use of 3.1 units of the enzyme for g-glycerophosphate.

		Inhi	bition(%)	GP/PB
Compounds	Concentration(N)	GP <sup>1)</sup>	PE <sup>2</sup> )	
None				1.00
edta	10-2	73.4	79.5	1.30
	10-3	29.4	45.6	1.29
KCW	10*2	100.	100.	
	10-3	72.2	39.5	0.46
ZnCl <sub>2</sub>	10**3	90.8	93.0	1.31
	10**4	78.2	66.0	0,64
CaCl <sub>2</sub>	10-3	5.0	24.0	1,25
	10**4	-0.2	16.0	1,22
Ca(CH <sub>3</sub> COO) <sub>2</sub>	10-3	0	<b>o</b> 10 1	1.00
NaF	10-1	27.6	44.0	1.29
	5x10 <sup>-2</sup>	13.2	21.0	1.10
POMB	10-3	0	0 14	1.00
Ethanolamine	10-2	-22.0	-7.0	1.14
Glycine	10-2	-10.5	-10.0	1.06

<sup>1)</sup> β-glycerophosphate

<sup>2)</sup> phosphoethanolamine

Table III

Alkaline Phosphatase Activity of Ascites Hepatoma, Regenerating Liver and Normal Liver of Rat

The ensyme assay was carried as described in the text. In each experiment, a time curve was constructed and the values indicated was estimated from its initial velocity.

Tissues		β <b>-Gly</b> (	cerophosphate(GP)	Phosphoethanolamine (Pi	E)
	No. of Experiments	Units/ug.	Units/mg. Units/g. Total-W. Tiesue	Units/ug. Units/mg. Units DNA-P Total-N. Tis	GP/PE
AH 130 Asci- tes Hepatoms		0.622 ±0.096 <sup>1</sup> )	18.8 ±4.5 <sup>1</sup> ) 275 ±45 <sup>1</sup> )	0.565 ±0.096 <sup>1</sup> ) 17.2 ±3.6 <sup>1</sup> ) 25	59.8 <sup>1)1.08</sup> ±0.045 <sup>1)</sup>
Regenerating Liver	<b>3</b>	0.550 ±0.048	4.63 112 ±0.48 ±7.4		3.0 1.14 6.7 <u>+</u> 0.014
Normal Liver	4	0.132 ±0.047	1.04 25.6 ±0.25 ±6.9		0.8 1.27 6.8 <u>+</u> 0.158

<sup>1)</sup> Standard deviation

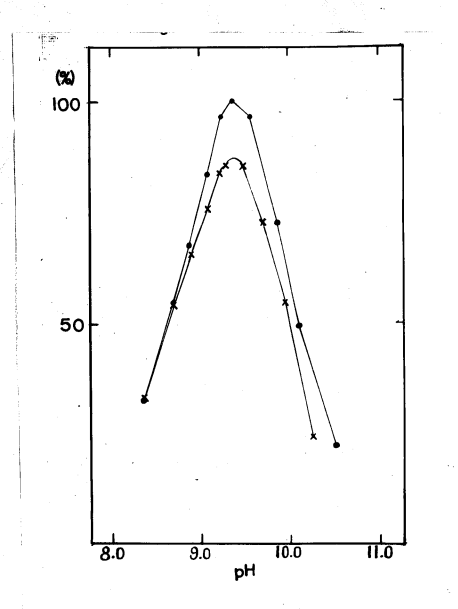
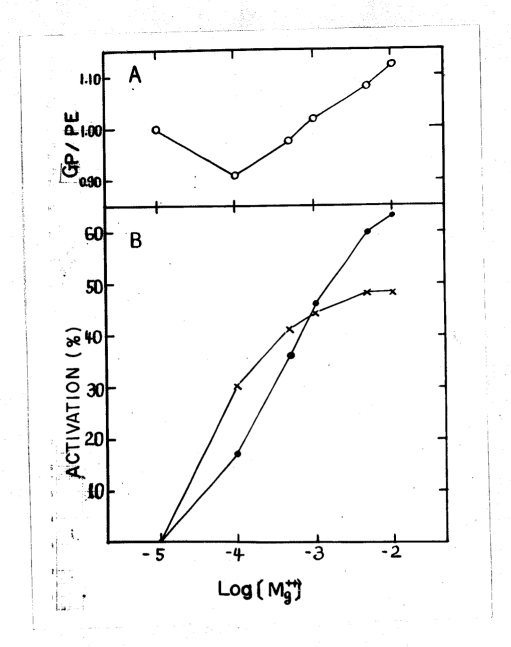
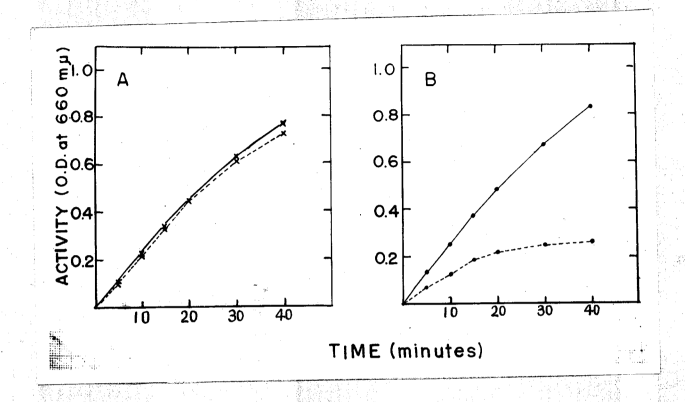


Fig. 1. pH-activity curves in the case of homogenate of tumour cells. The enzyme assay was carried out as described in the text. Activities indicated are expressed as per cent of the activity at pH 9.45, measured with  $\underline{\beta}$ -glycerophosphate as substrate.

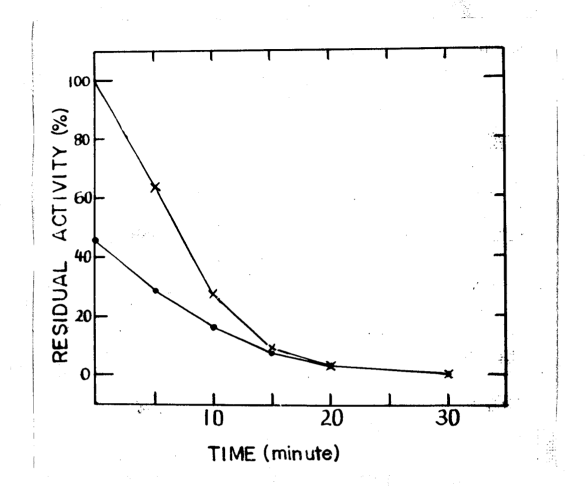
----- : Activities measured with A-glycerophosphate as substrate

. Activities measured with phosphoethanolamine as substrate





Pig. 3. Time curves in the absence and presence of 50 per cent acetone with use of phosphoeths—
nolamine or β-glycerophosphate as substrate. The enzyme assay was carried out as described
in the text. Activities were measured, A; with phosphoethanolamine, and B; with β-glycerophosphate, as substrates. Solid lines express the activities in the absence of acetone,
and broken lines express the activities in the presence of 50 per cent of sectore.



of phosphoethanolamine and β-glycerophosphate as substrates. The ensyme solution was incubated at 37°C in 50 per cent acetone without added substrate, and at each time indicated an aliquot was taken for the assay of residual activity as described in the text. Activities indicated are expressed as per cent of the activity at 0 time measured with phosphoethanolamine as substrate. Activity with β-glycerophosphate at 0-time preincubation corresponds to the initial velocity shown as the broken line in Fig. 3, B.

Residual activities with use of phosphoethanolamine

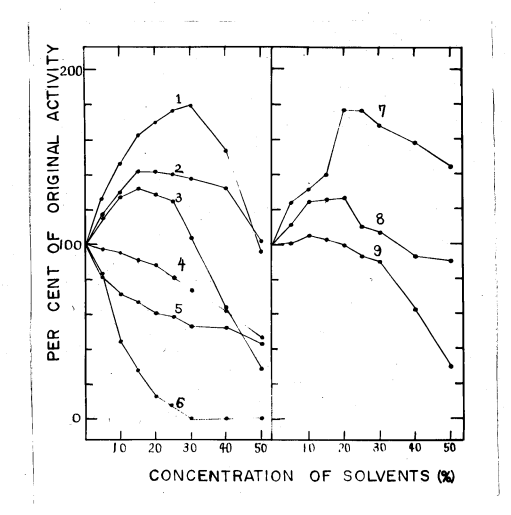


Fig. 5. Effect of various organic solvents on the reaction with use of phosphoethanolamine as substrate. The enzyme assay was carried out as described in the text. 1; activities in dioxan, 2; acetone, 3; dimethylformamide, 4; ethylene glycol, 5; glycerol, 6; 2-chloreethanol, 7; n-propanol, 8; ethanol, 9; methanol.

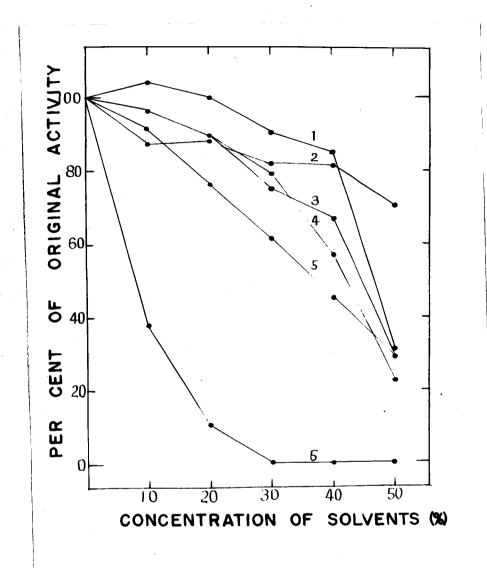
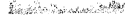


Fig. 6. Effect of various organic solvents on the reaction with use of β-glycerophosphate as substrate. The ensyme assay was carried out as described in the text. 1; Activities in dioxan, 2; n-propanol, 3; dimethylformamide, 4; acetone, 5; ethylene glycol, 6; 2-chloroethanol.



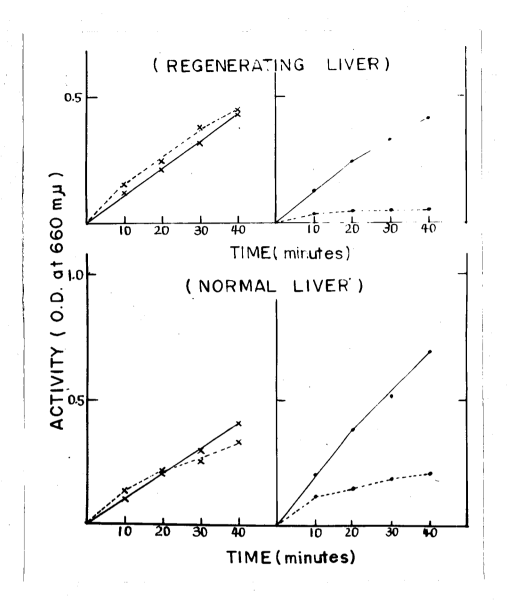


Fig. 7. Time curves in the absence and presence of 50 per cent acetone in the cases of regenerating liver and normal liver of rat. The enzyme assay was carried out as described in the text. Solid lines express the activities in the presence of 50 per cent acetone.

. Activities with use of β-glycerophosphate as substrate

# EFFECT OF ACETONE ON ALKALINE PHOSPHATASE ACTIVITY

MASAYORI INOUYE

#### EFFECT OF ACETONE ON ALKALINE PHOSPHATASE ACTIVITY

#### MASAYORI INOUYE

Alkaline phosphatase activity with phosphoethanolamine as substrate is increased by several organic solvents such as acetone, diozan, dimethylformamide and n-propanel, while the activity with figlycerophosphate is suppressed by these solvents (1).

In the present work, the effect of acetone on the activity with some other substrates was studied with partially purified ensyme from rat ascites hepatoma (AH 130). It was found that the activity with p-nitrophenyl phosphate was suppressed more strongly than with p-glycerophosphate not only by acetone but also by other solvents which caused activation with phosphoethanolamine (1). With phosphoethanolamine and p-nitrophenyl phosphate as substrates the kinetics of the reaction was analysed in the absence and presence of acetone and the difference in the effect of acetone on the activities with the two substrates is discussed.

#### MATERIALS AND METHODS

Ensyme Assay —— The activities with phosphoethanolamine

(PE\*), 8-glycerophosphate (GP\*) and phospho-Di-serine (PS\*) were estimated from the inorganic phosphate liberated during the reactions as described in the previous paper (1), except that the whole reaction mixture was used for the estimation of inorganic phosphate, since no precipitate was formed by adding 10 per cent trichloroacetic acid to purified enzyme in the present work. Activity with p-nitrophenyl phosphate (PNPP) was assayed from the p-nitrophenol liberated. This was estimated spectrophotometrically by measuring the change in optical density at 430 mm in 0.25 N NaOH. The usual assay system consisted of 0.1 ml. of substrate solution, 0.1 ml. of 10-2 M MgCl., 0.5 ml. of 0.1 M veronal-carbonate buffer, pH 9.50, and 0.3 ml. of enzyme solution in a total volume of 1.0 ml.. In experiments with organic solvents. 1.0 ml. of organic solvent and/or water was added to the reaction mixture described above, in a total volume of 2.0 ml.. After incubation at 37°C, the reaction was stopped by adding 0.5 ml. of 10 per cent of trichloroacetic acid for the activities with PE, GP, and PS or 2.0 ml. of 0.5 N NaOH for the activity with PNPP. In experiments with low concentrations of PNPP, 3.0 ml. of reaction mixture, containing the same constituents as above in the same ratio

(Fig. 1)

<sup>\*</sup> The following abbrevations are used in the present paper:

PE, phosphoethanolamine; GP, g-glycerophosphate; SP, phospho
Di-serine; PNPP, p-nitrophenyl phosphate.

and the reaction was stopped by adding 1.0 ml. of 1.0 M NaOH.

The color yield was corrected in both inorganic phosphate and pnitrophenol estimations in experiments with organic solvents, since
the color changed with the amount and type of solvent, added to the
reaction mixture.

In each experiment, a time curve was constructed and Fig. 1 shows an example obtained using PE and PNPP as substrates in the absence and presence of acetone. As shown in Fig. 1, linear results were obtained at least during the first 15 minutes of the incubation period even in the presence of acetone.

One unit of activity is definited at the amount of ensyme which liberates I umole of inorganic phosphate or p-altrophenol per hour, when 0.01 M of PE or 0.002 M of PNPP are used as substrate in 0.001 M MgCl<sub>2</sub> at 37°C and pH 9.50.

Ensyme ——Alkaline phosphatase was partially purified from rat ascites hepatoma (AH 130) as described in the previous paper (j). In the present work, partially purified preparations, (Fr. II in Table I of the previous paper (j)) were purified further with ribonuclease, bacterial protease digestion and gal filtration through Sephadex G-75 and G-200 (Pharmacia, Uppsala), as follows: A preparation extracted with butanol and fractionated with acetone (Fr. II in Table I) obtained from 365 g. of tumour cells, was incubated with 2 mg. of ribonuclease of bovine pancreas (Sigma Chemical Co.) at room temperature for 20 hours and the digest was fractionated with acetone. A fraction precipitating between 25 and 55 per cent acetone was collected by contrifugation and dissolved in water (Fr. III in Table I). Next,

the solution was digested with 4 mg. of Nagase (bacterial protease, Nagase Industrial Co.) at neutral pH at 57°C for 2 hours. After 2 hours incubation, 4 mg. more Nagase was added and the incubation was continued for 2 hours more. To the resulting solution, acctone was added to a concentration of 60 per cent and the precipitate formed was collected and dissolved in a small amount of water. This solution was subjected to gel filtration through a Sephadex G-75 column (3.5 x 23 cm.) in 0.1 M sodium acetate. The fractions eluted in the column volume were collected and concentrated by adding acetone (0 to 60 per cent) (Fr. IV in Table I). The procedure for purifica-

## (Table I)

tion is summarized in Table I. As can be seen, about 140 fold purification was achieved. Further purification using ion exchangers such as DEAE- and CM- cellulose and Duclite A2 was not successful. However, by gel filtration of Fr. IV through Sephadex G-200, a further purified preparation was obtained, as shown in Fig. 2. Fraction Nos. 33 - 38 were collevted. The recovery of protein (optical density at 280 m $\mu$ ) and alkaline phosphatase activity were 19.9 and 65.5 per cent, respectively. Therefore the ensyme had been purified about 3.5 fold over Fr. IV in Table I.

(Fig. 2)

Substrate specificity and pH-activity curves with PE and PNPP

as substrates are shown in Table II and Fig. 3, respectively. In the present work, Fr. IV in Table I was used for all experiments.

(Table II)

(Fig. 3)

#### RESULTS

Various Substrates ——Pig. 4 shows the effect of acctone on the ensyme activities with PE, PNPP, GP and SP as substrates. It can be seen that acctone activates the ensyme with PE at concentrations of 10 to 30 per cent, but not with PNPP, as already reported in the previous paper (1). In the case of SP, a slight activation is observed at concentrations of 10 to 20 per cent acctone, while the activity with PNPP is markedly suppressed and the inhibitory effect increase with

## (Fla 4)

increase in the acctone concentration. Table III shows the effects of some organic solvents at concentrations of 25 per cent on the ensyme activity with PE and PNPP. It can be seen that all the solvents tested inhibited the activity with PNPP in contrast to the acti-

vity with PE.

#### (Table III)

Identity of the Ensyme Which Hydrolyses PE and PNPP—
From the results mentioned above, the question arises as to whether the same ensyme participates in the hydrolysis of both PE and PNPP.
Although there was a little difference between the two pH-activity curves (Fig. 3), the ratio of the activity with PE to the activity with PNPP was constant through the purification procedure, as shown in Table I and Fig. 2. This suggests that both activities are due to the same ensyme. Fig. 5 shows the heat inactivation curve. The ensyme solution was incubated at each of the temperatures indicated in Fig. 5, at pH 9.50 for 3 minutes, and after cooling in an ice-water bath the residual activities were measured with PE, PNPP, GP and PS as sub-

## (Fig. 5)

strates. Inactivation of the activities toward all substrates began from 50°C and almost all the activities with all substrates were lest at 60°C.

The kinetics of the reactions was analyzed to examine this problem further. As shown in Fig. 6, PE acts as a competitive inhibitor of the activity with PNPP. Ki, the dissociation constant of the ensyme-inhibitor complex, was calculated from Fig. 6-B, since the slopes in the presence of PE are equal to  $\frac{KB}{Vmax}$ .  $\left(1 + \frac{I}{KI}\right)$ ,

where Km, Vmax., and I are the Michaelis constant for PNPP (1.13 x 10<sup>-4</sup> moles/liter, obtained from the line in the absence of PE in Pig. 6-B), the maximum velocity and the concentration of inhibitor (PE), respectively. The Ki's were found to be 1.90 x 10<sup>-3</sup> and 2.08 x 10<sup>-3</sup> moles/liter in the presence of 0.0050 M and 0.0025 M PE, respectively. These values are in quite good agreement with the Km for PE, 1.93 x 10<sup>-3</sup> moles/liter, which was obtained from a Lineweaver-Burk plot (3) for PE (Pig. 6-A).

#### (Fig. 6)

From these results, it is evident that the same enzyme or the same active site reacts with both PE and PNPP.

Effect of Substrate Concentration on the Activation and Inhibition Caused by Acetone——Fig. 7 shows the effect of substrate concentration on the activation and inhibition caused by acetone. With PE as substrate, the activation caused by acetone increases with decrease in the concentration of PE. With 0.0010 M PE, more than 70 per cent activation was observed at a concentration of 40 per cent acetone. With PNPP as substrate, the inhibition caused by acetone

## (Fig. 7)

increases with decrease in the concentration of PNPP. It should be noted that the suppression of the activity with PNPP is not due to the denaturation of the ensyme protein caused by acctone, since time curves in the presence of acctone were linear for at least the first 15 minutes of the incubation period at 37°C, as shown in Fig. 1.

Therefore it is thought that acctone acts as an activator for PE and as an inhibitor for PNPP.

Fig. 8 shows Lineweaver-Burk's plots for PE and PNPP in the absence and presence of acetone. As expected from Fig. 7, the slopes decrease with increase in the acetone concentration in the case of PE, and both lines cut the vertical axis very near the point

## (Fig. 8)

given in the absence of acetone (Fig. 8-A). The apparent Michaelis constants were 1.42 x  $10^{-3}$ , 1.11 x  $10^{-3}$ , 0.81 x  $10^{-3}$  and 0.45 x  $10^{-3}$  moles/liter at 0, 5, 10 and 25 per cent acetone, respectively. With PNPP as substrate, on increase in the acetone concentration, the apparent Km values increased as follows: 0.97 x  $10^{-4}$ , 1.37 x  $10^{-4}$ , 1.61 x  $10^{-4}$  and 1.66 x  $10^{-4}$  moles/liter in 0,5, 10 and 25 per cent acetone, respectively. Furthermore, intercepts on the vertical axis increased with increase in acetone concentration, which means that the apparent  $k_2$  values, the rate constants for dissociation of the ensyme-substrate complex to the ensyme and the product, decreased.

Effect of Acetone on the Activities with PE and PNPP in the Presence of PNPP and PE, Respectively—Table IV shows the effect of acetone on the hydrolysis of PE and PNPP by the ensyme in the presence of PNPP and PE, respectively. When both subst-

rates are present in the reaction mixture, they will be hydrolysed at the same time by alkaline phosphatase, the one substrate competing tively inhibiting the hydrolysis of the other substrate, and if the change of the ensyme protein caused by a certain concentration of acetone in the presence of PE is different from the change caused by the same concentration of acetone in the presence of PNPP, the increase in the activity with PE and the inhibition of the activity with PNPP caused by acetone will decrease in the presence of both substrates. However, as can be seen in Table IV, the activity with PE was increased by acetone even in the presence of PNPP, while the activity with PNPP was inhibited by acetone even in the presence of PE. Therefore, it is thought that the effects of acetone on the ensyme activities with PE and PNPP are independent.

(Table IV)

(Fig. 9)

#### DISCUSSION

In the previous paper (1), it was shown that alkaline phosphatase activity with PE as substrate was activated by several organic solvents, while ensyme activity with GP as substrate was suppressed by these solvents. The present paper shows that beside GP, the activity with PNPP is inhibited more strongly than the activity with GP by several organic solvents. This paper shows that the name ensyme or the same active site participates in the hydrolysis of both PE and PNPP, and that the suppression of the activity with PNPP by acetone is not due to denaturation of the ensyme protein, since time curves for PNPP in the presence of acetone were linear, at least for the first 15 minutes of the incubation period at 37°C (Fig. 1). Therefore acetone acts as an activator with PE and as an inhibitor with PNPP.

Kinetical analysis (Fig. 8), showed that acctone decreased the apparent Km, Michaelia constant, for PE and increased it for PNPP. This fact suggests that in the presence of acctone the affinity of PE for the ensyme increased and that of PNPP for the ensyme decreased. It was also found that the rate constant for dissociation of the ensyme-substrate complex to the ensyme and product was scarcely changed by acctone in the case of PE, while it was decreased by acctone when PNPP was the substrate.

These differences between the activities with PE and PNPP in the effect of acetone appear to be due to the same change in the ensyme protein. This change is caused by acetone in the presence of either

PE or PNPP, since similar phenomena caused by acetone were observed even in the presence of both PE and PNPP (Table IV). Although PE had more of a protective effect than GP on the inactivation caused by acetone as shown in the previous paper (1), it is unknown whether this protective effect of PE has any relation with the phenomena described above. It would also be possible to explain the phenomena as due to changes in the substrates caused by acetone, rather than as due to a change in the ensyme protein caused by acetone. In fact, the activities with compounds having: amino groups such as PE and PS, were increased by acetone, in contrast to those with other compounds (Fig. 4).

Effects of organic solvents on ensyme activities have been reported in various papers (4 - 10), and <u>Brahms et al</u>. (10) have shown a relationship between the ensyme activity of myosin A and its conformational changes caused by organic solvents. They mentioned that the activation by solvents would be due to a partial unfolding of myosin in the region of the active site. If the same is true for alkaline phosphatase, a change in the ensyme protein caused by acctone would be inhibitory with PNPP but activatory with PE. These problems will be elucidated by examining conformational changes in alkaline phosphatase caused by organic solvents, using more purified preparations.

### SUMMARY

1. The effects of acetone on the activities of alkaline phospha-

tase with various substrates were investigated. It was found that the activity with g-nitrophenyl phosphate (PNPP) was markedly suppressed by several organic solvents, which increased the activity with phosphoethanolamine (PE).

- It was shown that the same ensyme or the same active site participates in the hydrolysis of both PE and PNPP.
- 3. The kinetics of the reactions with PE and PNPP were analyzed in the absence and presence of acetone. It was found that the apparent Km, Michaelis constant, was decreased by acetone with PE but was increased with PNPP.
- From these results, the effect of acetone on alkaline phosphatase activity are discussed.

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Table I
Summary of the Purification of Alkaline Phosphatase from Ascites Repatoma of Rat

The ensyme assay was carried out as described in the text. Each fraction was dialysed against deionized water before use.

Fraction		4 : 								
	Volume	PE (0.01 M)			GP (0.01 M)		PNPP(0.002 <u>M</u> )		GP/PE	PNPP/PE
	<b>≇</b> .	Total Act. <u>Units</u>	Yield Z	Specific Act. Units/Total Bills	Total Act. Units	Yield Z	Total Act. <u>Vaita</u>	Yield Z		
iomogenate of 365 g. tumour	1,300	9.60x10 <sup>4</sup>	100.	17.1	10,6x10 <sup>4</sup>	100.	8,72x10 <sup>4</sup>	100.	1.10	0.91
Pr.I. extraction with g-butanol	440	7.00x10 <sup>4</sup>	73.0	97.5	8,40x10 <sup>4</sup>	79.2	6.55×10 <sup>4</sup>	75.0	1,20	0.94
Pr.II, fractio- nation with acctone	300	6.40x10 <sup>4</sup>	66.6	134.0	7.05x10 <sup>4</sup>	66.5	6.21x10 <sup>4</sup>	71.2	1.10	0.97
Pr.III, ribonu- clease diges- tion	95	5.29x10 <sup>4</sup>	55.0	. <b>290.</b> 0	5.61x10 <sup>4</sup>	54,8	5.01x10 <sup>4</sup>	57.5	1.10	0.95
Pr.IV, Nagase digestion and gel-diltration	24	5,00x10 <sup>4</sup>	52.1	2400.	5.45x10 <sup>4</sup>	51.5	4.75×10 <sup>4</sup>	54.3	1,09	0.95

Table II
Substrate Specificity of Alkaline Phosphatase from Ascites
Hepatoma of Rat

Activities are expressed as a percentage of the activity measured with PE as substrate at pH 9.50. Activities with ATP\*, AMP\*\* and G1P\*\*\* as substrates were assayed according to the method described for glycogen phosphorylase activity (2).

	Substrate	Activity at pH 9.50
	0,01 <u>M</u> PE	100.
	0.01 <u>M</u> GP	110.
	0.01 M G15	13.
· · · · · · · · · · · · · · · · · · ·	0.01 M PS	107.
	0.002 M PNPP	95.0
	0.005 M ATP	6.0
	0.005 <u>M</u> AMP	87.0

\* ATP: adenosine triphosphate

\*\* AMP: adenosine monophosphate

GIP: glucose-1-phosphate

Table III

# Effect of Various Organic Solvents on Alkaline Phosphatase Activities with Phosphoethanolamine (PE) and p-Nitrophenri Phosphate (PNPP)

Activities were assayed in 25 per cent (v/v) concentrations of solvents and 0.005 M PE or 0.001 M PNPP under the following conditions; 0.0005 M MgCl<sub>2</sub>, at 37°C and pH 9.50. Activities are expressed as a percentage of the activity with PE as substrate.

Solvent (25%)	Substrate					
	78	PNPP				
None	100.	100.				
Acetone	116.	35.0				
Dioxan	134.	32.8				
Dimethylformamide	105.	24.7				
n-Propanol	112.	22,6				

Table IV

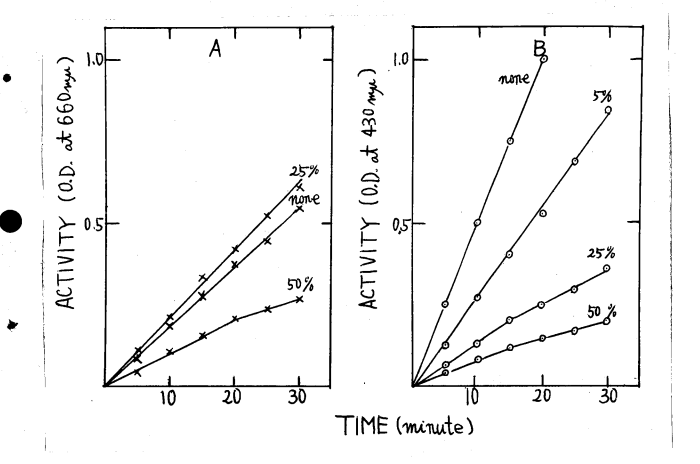
## Effect of Acetone on the Activities with PE and PNPP in the presence of both PNPP and PE

Activities with PNPP in the presence of PE were measured from the liberated g-aitrophenol as described in the text. Activities with PE in the presence of PNPP were estimated as follows; inorganic phosphate (iP) liberated from PE) = (Total iP liberated from both PE and PNPP) = (iP liberated from PNPP, which was estimated from the g-aitrophenol liberated from PNPP). Ensyme assay was carried out at pH 9.50 in 0.0005 M MgCl<sub>2</sub> in the absence and presence of acetone. Activities are expressed as a percentage of the activity in the absence of acetone.

Acetone Yolume		Substrate										
			PE			PNPP						
	:	0,0025		0.0050 M			0,0005 <u>M</u>			G,0010 M		
	None	PNPP 0,0005 M	PNPP 0,0010 <u>M</u>	None	PNPP 0.0005 <u>M</u>	PNPP 0.0010 <u>M</u>	None	PE 0.001 M	PE 0.005 M	None	PE 0.001 M	PE 0.005 M
•	100. (100.) <sup>2</sup>	100. (26.2) <sup>8</sup>	100 <b>.</b> (13.3) <sup>a</sup>	100. (100.)ª	100. (43.8) <sup>2</sup>	100. (25.0)a	100. (100.) <sup>b</sup>	190. (93.0)b	100. (66.0) <sup>b</sup>	100. (100.) <sup>b</sup>	100. (93.5) <sup>b</sup>	100. (72.0) <sup>3</sup>
<b>5</b>	110.	105.	170,	110,	104.	125.	78.0	73.2	74.5	79.0	75.5	71.0
10	127.	159.	185.	125.	131.	144.	56.5	55.8	49.5	57.5	55.8	51.0
25	152.	215.	315.	137.	156.	221.	37.7	34.1	25.7	39.2	39.1	25.1

a) per cent of the activity in the absence of PNPP

b) per cent of the activity in the absence of PE



Pig. 1 Time curves of alkaline phosphatase activity in the absence and presence of acetone. Phosphatase activity was assayed as described in the text. A; PE (0.01 M) and B; FEFF (0.002 M) were used as substrates in the absence and presence of acetons.

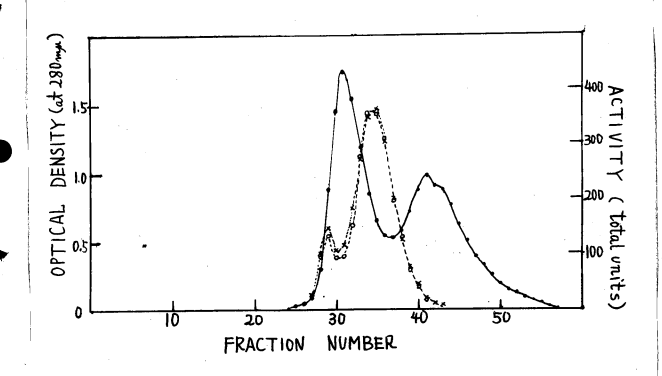
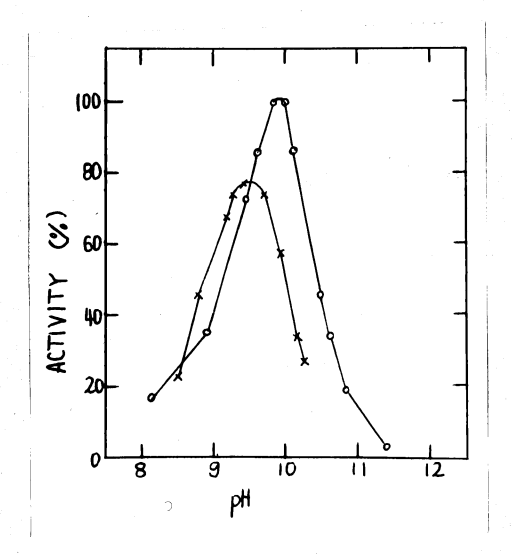


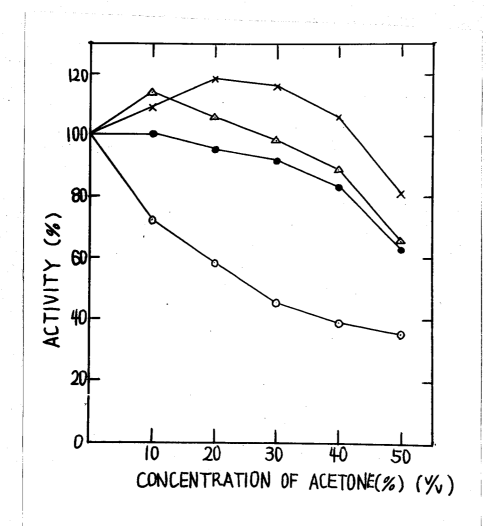
Fig. 2 Gel filtration of Fr. IV in Table on Sephadex G-200. Two ml. of Fr. IV were applied to a column of Sephadex G-200 (3.5 x 25 cm.) in 0.1 M sodium acetate. The column was eluted with 0.1 M sodium acetate and 2 ml. fractions of effluent were collected.

. sabsorbancy at 280 mu

\* sectivity with PE (0.01 M)

: activity with PMPP (0.002 M)





Pig. 4 Effect of acetome on alkaline phosphatase activities with various substrates. The enzyme assay was carried out as described in the text. Activities are expressed as a percentage of the activity in the absence of acetome.

\*\*\times : PE (0.01 M), 0-0 : PNPP (0.01 M), \times : PS (0.01 M) and \times : GP (0.01 M) were used as substrates.

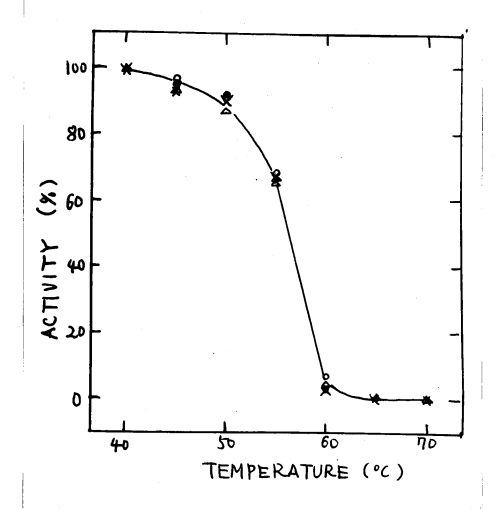


Fig. 5 Heat inactivation of alkaline phosphatase.

The ensyme solution was incubated at the temperatures indicated at pH 9.50 for 3 minutes and after cooling in an ice-water bath the residual activities were assayed with XX PE (0.01 M), 0-0; PMPP (0.002 M), 6-0; GP (0.01 M) 6-6; and PS (0.01 M) as substrates.

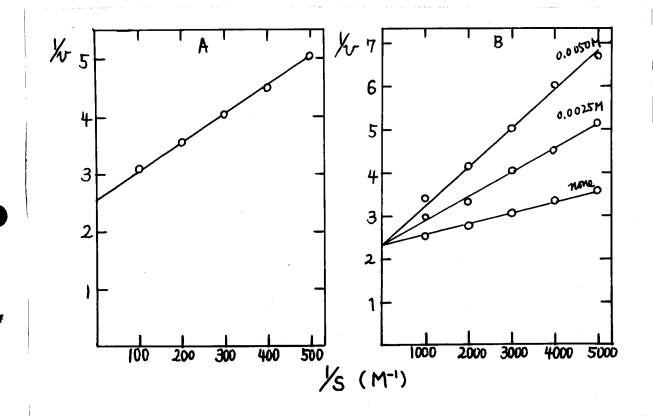


Fig. 6 Lineweaver-Burk plots with PE and PEPP in the absence and presence of PE.

A; Lineweaver-Burk plots with PE and B; with PHPP in the absence and presence of 2.5 mM and 5.0 mM of PE at pH 9.50 in 1 mM MgCl<sub>2</sub>.

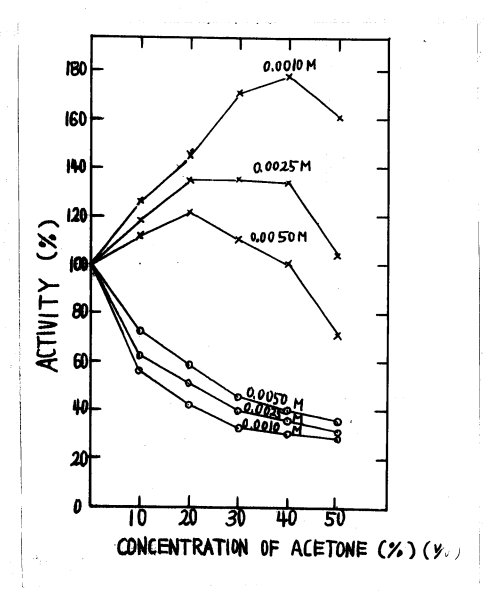


Fig. 7 Effect of substrate concentration on activation and inhibition by acctone. The ensyme assay was carried out as described in the text. Activities are expressed as a percentage of the activity in the absence of acctone.  $\times$  ; PE and  $\circ$ — $\circ$ ; PNPP were used as substrates at the various concentrations indicated in the figure .

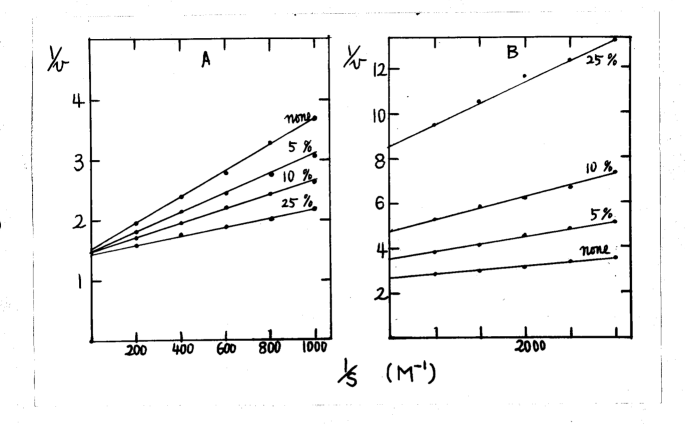


Fig. 8 Lineweaver-Burk plots with PE and PNPP in the absence and presence of acetone.

A; Lineweaver-Burk plots with PE and B; with PNPP in the absence and presence of scetone at pH 9.50 in 5 mM MgCl<sub>2</sub>. The concentrations of acetone are indicated in the figure as volume per cent.

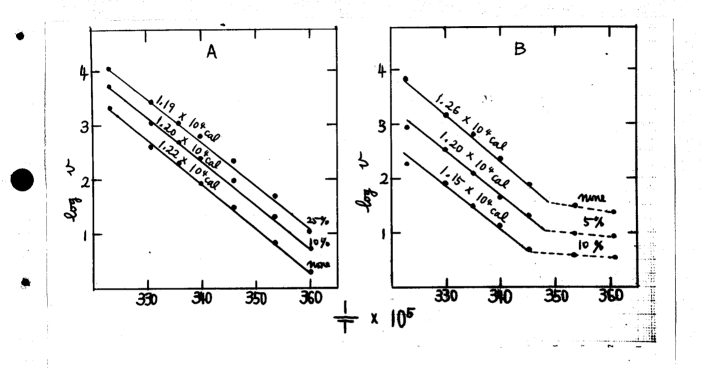


Fig. 9 Arrenius plots for alkaline phosphatese in the absence and presence of scetone.

Ensyme assay was carried out at pH 9.50 in 0.5  $\underline{m}\underline{M}$  MgCl<sub>2</sub>, A; with PE (0.01  $\underline{M}$ ) and B; with FMFP (0.002  $\underline{M}$ ) in the absence and presence of acetone. The concentrations of acetone are indicated in the figure as volume per cent.