MEASUREMENT OF SYNTHESIS RATE OF LONG-CHAIN ACYL-COENZYME A ESTER IN BOVINE LIVER BY HIGH-PERFORMANCE LIQUID CHROMATOGRAPHY

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Summary

A high performance liquid chromatographic procedure is described for the direct determination of the picomole amount of palmitoyl-Coenzyme A and stearoyl-Coenzyme A, using a stainless steel column packed with C-18 derivatized porous silica (SPh), an isocratic elution with a mixture of 33 mM KH₂PO₄/acetoniitrile as a mobile phase and a UV detector. The long-chain acyl-Coenzyme A esters were determined in incubated microsomal fractions of a bovine liver to demonstrate the utility of this method for monitoring acyl-CoA synthesis in biological samples. The reaction rate of palmitate was higher than that of stearate. After a 60 minute incubation period, the generated amount of palmitoyl-Coenzyme A and stearoyl-Coenzyme A were approximately 70 and 20 nmol/mg microsomal protein, respectively. The advantage of this method are in that no decomposition of the CoA esters is involved, while the constituent molecular species is detected.

(Key Words: HPLC, Acyl CoA, Bovine Liver)

Introduction

The recent introduction of C-18 derivatized porous silica microparticles for reverse-phase high performance liquid chromatography (HPLC) has facilitated the assay of Coenzyme A (CoASH) in biological extracts (Ingebritsen and Farstad, 1980). The method was adequate for the measurement of free CoASH after controlled alkaline hydrolysis of the perchloride insoluble materials from biological samples, thus giving an indication of the amount of acyl-CoA derivatives in the sample (Ingebritsen et al., 1979). However, with this method, it was impossible to determine the individual molecular species of acyl portions of the derivatives and to separate CoASH from acyl-CoA derivatives without degradation of free CoASH (Ingebritsen et al., 1979). In view of these problems during determination of acyl-CoA esters in biological samples, it is important to develop a simple method which provides sensitive direct measurement, detecting the constituent molecular species. In this communication, the details of our procedure used for the determination and identification of palmityl Coenzyme A (P-CoA) and steroyl Coenzyme A (S-CoA) by HPLC is described. When combined with enzymatic synthesis, this method provides a quantitative assessment of the conversion of long chain fatty acid to the corresponding acyl-CoA ester.

Materials and Methods

Chemicals

CoASH was obtained from Kohnin Co. Ltd. (Tokyo, Japan). P-CoA and S-CoA were purchased from Pharmacia F-L Biochemicals (Uppsala, Sweden). ATP and L-cysteine were supplied by Nakarai Chemicals Ltd. (Kyoto, Japan). All other reagents were commercially available and of highest purity.

Chromatographic conditions

A constant volume high-performance liquid chromatographic pump equipped with UV detector (Model LC-4A, Shimadzu, Kyoto, Japan) was
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Figure 1. Figure 1A. The HPLC chromatogram of 230 p mol of palmitoyl Coenzyme A (P-CoA) before alkaline hydrolysis with a retention time of (tR=) 4.9 min. P-CoA (0.23 μmol) was dissolved in 10 ml of 10 mM aqueous solution of 2-mercaptoethanol and 5 ml aliquot of the solution was diluted to 10 ml with distilled water before the injection of 20 μl into a 15 cm x 0.6 cm column of 5 μm silica ultrosphere ODS. Individual components were eluted isocratically at 32°C with a flow rate 1 ml/min, using a solvent mixture comprised of 33 mM KH₂PO₄/acetonitrile (30:70, v/v). UV absorbance was monitored at 260 nm with 0.02 absorbance until full scale. Figure 1B. The eluate of the solution of figure 1A by the solvent stock comprised of 220 mM KH₂PO₄/methanol/thiodiglycol (100:12:0.05, v/v/v) at room temperature. HPLC column and flow rate were the same as Fig. 1A. UV absorbance was monitored at 254 nm. Figure 1C. Disappearance of P-CoA after alkaline hydrolysis. The counterpart 5 ml solution of figure 1A was hydrolyzed at pH 12.7 with 1 N KOH, for 60 min at 40°C. After cooling on ice water, the pH of the suspension was readjusted to 5 with 6% (v/v) perchloric acid containing 0.5 M triethanol-amine-HCl and diluted to 10 ml as described for figure 1A. 20 μl aliquot was injected and separated as described for figure 1A. Figure D. Appearance of CoA (tR=16.2 min) after alkaline hydrolysis of P-CoA. 20 μl aliquot of the solution of figure 1C was injected and separated as described for figure 1B.
Enzymatic synthesis of CoA derivatives of long chain fatty acids

A bovine liver, obtained from a 14 month old Japanese Black steer which had been fasted for 2 days, was used. Fractional collection of enzymes from the liver and the enzymatic synthesis of CoA derivatives of palmitic acid and stearic acid were performed by the method of Kornberg and Prizer (1953).

Protein determination

Protein content of the microsome fraction of bovine liver was determined by the Folin-Ciocalteu phenol reagent (Lowry et al., 1951) using bovine serum albumin as standard.

Results and Discussion

Optimization of HPLC conditions

Using a micro particulate reversed-phase column, the acetonitrile mobile phase and commercial preparation of palmitoyl-CoA (P-CoA), a possible peak of P-CoA with the retention time of (TR = 4.9 min was collected (figure 1A). Since the injected sample contained a reducing agent and different compounds that absorbed at 260 nm, peaks were always observed shortly after sample injection. When the sample was eluted by methanol mobile phase adequate for CoASH analysis, CoASH (TR=16.2 min) was slightly detected (figure 1B). The reducing agent 2-mercaptoethanol was eluted as two peaks, with the retention time of 5.3 min and 14.0 min, respectively (figure 1B). A possible peak of P-CoA was not obtained on the chromatogram during the 3 hours when the sample was eluted by the methanol mobile phase. Cockey et al. (1981), using C18 column for assay of short-chain acyl CoA, reported that the more lipoplyc compounds were more strongly absorbed in the column and required a higher methanol concentration for removal. However, in our trial with the highest methanol concentration (99%), a possible peak of the long-chain acyl-CoA was not collected within a convenient retention time, although a broadened indistinct peak was obtained after the given time. In practice, acetonitrile is more effective than methanol for the elution of long-chain acyl-CoA from C18 column. The retention time of long-chain acyl-CoA was not altered by phosphate but by acetonitrile content in the mobile phase solvent, and was shortened by increasing acetonitrile content in the solvent. It should be noted that a higher acetonitrile with an inappropriate high concentration of phosphate causes recrystallization of phospholic acid in the mobile phase during analysis, which leads to damage on HPLC column, rendering it useless. On the other hand, the phosphate buffer is indispensable in keeping mobile phase pH at approximately 5 in order to avoid

Figure 2. Separation of P-CoA and stearoyl Coenzyme A (S-CoA), P-CoA (approx. 0.75µmol) and S-CoA (approx. 0.75 µmol) were dissolved in a 10 ml of 10 mM aqueous solution of 2-mercaptoethanol. The 3 µl of the solution was injected and separated using the chromatographic condition given for figure 1A. The letters a and b refer to the P-CoA and S-CoA, respectively.
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destruction of acyl-CoA ester during analysis. The phosphate and acetonitrile content had to be balanced in the mobile phase for determining the most convenient conditions for sample analysis. S-CoA was eluted as a distinct peak (tr=8.3 min), completely separated from the proposed peak of P-CoA (tr=4.9 min) with the appropriate flow rate and phosphate buffer: acetonitrile ratio (figure 2).

Identification of acyl-CoA esters

Peak identification was accomplished not only by alkaline hydrolysis but by enzymatic synthesis of acyl CoA esters. After alkaline hydrolysis manipulation in the solution of P-CoA, a possible peak of P-CoA in figure 1A disappeared (figure 1C), whereas the peak corresponding to CoASH (tr=16.2 min) appeared (figure 1D). This suggests that the proposed peak of P-CoA in figure 1A originates from CoA derivative. Similar series of experiments were performed using samples containing S-CoA. After the alkaline hydrolysis of the

Figure 3. Chromatogram of enzymatically synthesized P-CoA (tr=4.9 min) in microsomal fraction from bovine liver. The incubation mixture contained 0.8 ml of palmitate (0.02M palmitic acid), 2.0 ml of ATP (0.02M), 1.6 ml of CoA (0.0034M, 94 percent pure), 0.6 ml of cysteine (0.2M), 0.4ml of NaF (0.5M), 0.2 ml of MgCl₂ (0.3M), 0.8 ml of phosphate buffer (0.5M, pH 7.4), 0.6 ml of lyophilized residue fraction (10.3 mg protein per ml) and 1 ml of water. Incubation was at 40°C for 60 min. Aliquots of 0.5 ml were taken at 2 min, 10 min, 60 min and heated in 60°C bath for 3 min to inactivate the enzymes, then chilled to 0°C and diluted to 2 ml with the solvent mixture used for figure 1A. After centrifugation at 1600 g for 15 min, 20μl aliquots of the supernatant were injected and separated using the chromatographic condition used for figure 1A.
sample, the proposed peak of S-CoA disappeared while CoASH peak appeared.

Definite proof of the composition of the proposed peak of P-CoA and S-CoA in the chromatogram was obtained in experiment with the enzymatic synthesis of CoA derivatives of palmitate and stearate. After the addition of palmitate into microsomal fraction and incubation at 40°C, a peak having identical retention time of the proposed peak of P-CoA appeared and continuously increased (figure 3 A,B,C). Similarly, addition of stearate instead of palmitate resulted in enlargement of the peak having identical retention time of the proposed peak of S-CoA (figure 4 A,B,C). These phenomena suggest that the proposed two peaks definitely originate from P-CoA and S-CoA, respectively.

Quantitation and measurement of long-chain acyl-CoA synthesis

The most widely used procedure for the determination of long-chain acyl-CoA in biological samples was the measurement of liberated CoASH after alkaline hydrolysis of perchloric insoluble material from the sample (Ingebritsen et al., 1979). However, in our experiment, complete hydrolysis of acyl-CoA induced negligible destruction of liberated CoASH, even when the alkaline hydrolysis was performed by adding a recommended reducing agent into the sample. Since precise measurement of acyl-CoA by alkaline hydrolysis procedure had to be performed by calculating the total amount of liberated and intact CoASH, and the unknown degradation product of CoASH, we thought that the alkaline

![Chromatogram](image)

**Figure 4.** Chromatogram of enzymatically synthesized S-CoA (tR=8.3 min) in microsomal fraction from bovine liver. The 0.8 ml of stearate (0.02M stearic acid) was added instead of palmitate to the incubation medium used for figure 3. The injected aliquots were prepared as described for figure 3 and separated using the chromatographic condition used for figure 1A.
hydrolysis procedure was not convenient for the microanalysis of long-chain acyl-CoA. The advantage of our substitute method is that no decomposition of the CoA esters is involved in the process.

Excellent linearity was obtained between the injected amount of the individual authentic acyl-CoA ester and the integrated peak area up to at least 1000 p mol (figure 5). It was necessary to dissolve the CoA esters in a 0.1 N potassium phosphate buffer (pH 5.5) in order to obtain the standard curve which can extrapolate to the origin. Apparent losses of peak area corresponding to those esters were observed when dissolved in distilled water. As seen in figure 2 with the amount of individual standard of 230 p mol, peak height of 50% full scale corresponded to 150 p mol to 300 p mol of P-CoA or S-CoA. The acyl-CoA peak of 5% full scale can be measured, so that the lower limit of detection would be approximately 30 p mol.

Quantitation of individual acyl-CoA esters in a biological sample was achieved by comparing the peak area of a known amounts of individual acyl-CoA standards with the area of peaks having retention times corresponding to those of the standards. The kinetic feature of the acyl-CoA synthesis in microsomal fraction from bovine liver was obtained by the injection of aliquots picked up at certain intervals from the incubated sample (figure 6). The reaction rate of palmitate was higher than that of stearate. After 60 min. incubation, the generated amount of P-CoA and S-CoA were approximately 70 and 20 n mol/mg microsomal protein, respectively. The specificity of acyl-CoA synthesis towards these two fatty acids is in agreement with those of Massaro and Lennartz (1965) with bacteria, Brindley and Hubscher (1966) with cat and guinea pig, Fande and Mead (1968) with rat and Tanaka et al. (1979) with purified long-chain acyl-CoA synthetase from rat liver. The synthesized amount of P-CoA, although varying slightly, is in general agreement.
with the experiment of Kornberg and Pricer (1953) which measured radioactive palmitate incorporation into P-CoA in guinea pigs.

To our knowledge, this is the first report where kinetic feature of long-chain acyl-CoA synthesis has been monitored by HPLC method. This method should provide useful information for detailed understanding of several metabolic pathways which employ long-chain acyl-CoA esters.

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Literature Cited


