

# ACYL-CoA OXIDASE [ACOD]

from *Arthrobacter* sp.  
(Acyl-CoA: oxygen 2-oxidoreductase, EC 1.3.3.6)



## Preparation and Specification

Appearance : Yellowish amorphous powder, lyophilized  
Specific activity : More than 20 U/mg solid

## Properties

Substrate specificity	: See Table 1	
Molecular weight	: 210 kDa (Sephadex G-150)	
Isoelectric point	: pH 4.7	
Michaelis constant	: Palmitoyl-CoA $2.0 \times 10^{-5}\text{M}$	
Optimum pH	: Serum Acyl-CoA 7.5	Figure 1
	: Palmitoyl-CoA 8.5	Figure 1
pH stability	: 6.0-7.5 (37°C, 60 min)	Figure 2
Thermal stability	: Stable at 40°C and below (pH 7.0, 10 min)	Figure 3
Storage stability	: At least one year at -20°C	Figure 4
Effects of various chemicals	: See Table 2	
Stabilizer	: FAD	
Activator	: Triton X-100	

## Applications for Diagnostic Test

This enzyme is useful for enzymatic determination of **fatty acid** when coupled with Acyl-CoA synthetase (T-16).

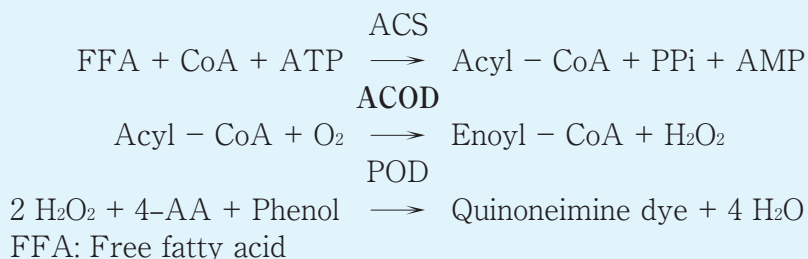


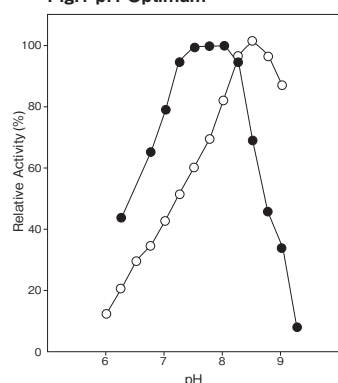
Table 1. Substrate specificity of ACOD

Substrate	Relative activity (%)	Km value (10 <sup>-5</sup> M)
Hexanoyl-CoA (6:0)	11	
Octanoyl-CoA (8:0)	60	
Decanoyl-CoA (10:0)	73	
Dodecanoyl-CoA (12:0)	87	
Tetradecanoyl-CoA (14:0)	99	
Hexadecanoyl-CoA (16:0)	53	2.0
Octadecanoyl-CoA (18:0)	16	3.8
Icosanoyl-CoA (20:0)	7	
9-Tetradecenoyl-CoA (14:1)	100	
9-Hexadecenoyl-CoA (16:1)	65	
9,12-Hexadecadienoyl-CoA (16:2)	55	
<i>cis</i> -9-Octadecenoyl-CoA (18:1)	45	4.0
<i>trans</i> -9-Octadecenoyl-CoA (18:1)	31	
<i>d</i> -12-Hydroxy- <i>trans</i> -9-octadecenoyl-CoA (18:1)	7	
<i>cis</i> -9, <i>cis</i> -12-Octadecenoyl-CoA (18:2)	31	3.8
<i>cis</i> -6, <i>cis</i> -9, <i>cis</i> -12-Octadecenoyl-CoA (18:3)	95	1.67
15-Tetracosenoyl-CoA (24:1)	7	

Table 2. Effect of various chemicals on ACOD activity

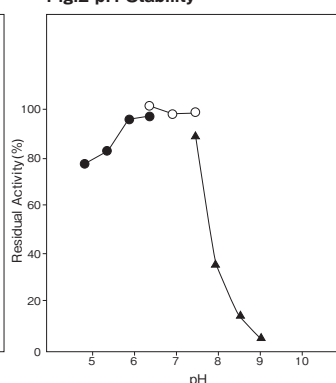
Additive	Concentration	Relative activity (%)
None	-	100
NaCl	1mM	102
KCl	1mM	104
LiCl	1mM	101
NH <sub>4</sub> Cl	1mM	101
MgCl <sub>2</sub>	1mM	138
BaCl <sub>2</sub>	1mM	147
CaCl <sub>2</sub>	1mM	137
MnCl <sub>2</sub>	1mM	155
ZnCl <sub>2</sub>	1mM	99
CoCl <sub>2</sub>	1mM	121
FeCl <sub>3</sub>	1mM	101
EDTA	1mM	74
Triton X-100	0.1%	140
Adekamol SO-120	0.1%	147
Sodium laurylbenzene sulfonate	0.1%	54
Sodium laurylsulfate	0.1%	43
Deoxycholate	0.1%	106

Fig.1 pH Optimum



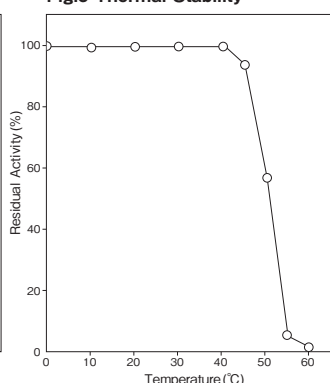
pH 6 – 7 Phosphate buffer  
pH 7 – 9 Tris-HCl buffer  
○ : Palmitoyl-CoA  
● : Serum Acyl-CoA

Fig.2 pH Stability



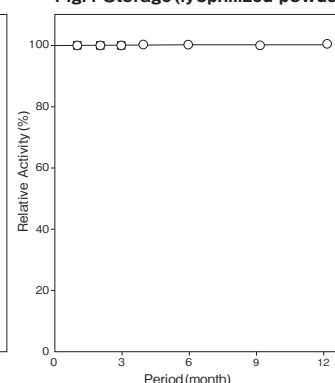
● : 3,3-Dimethylglutarate-NaOH buffer  
○ : Phosphate buffer  
▲ : Tris-HCl buffer

Fig.3 Thermal Stability



pH 7.0, 10 min.  
Phosphate buffer

Fig.4 Storage (lyophilized powder)

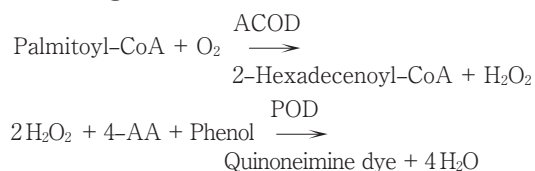


○ : -20°C  
□ : 5°C

## Assay

### Principle

The assay is based on the increase in absorbance at 500 nm as the formation of quinoneimine dye proceeds in the following reactions:



### Unit definition

One unit is defined as the amount of enzyme which generates 1 μmole of H<sub>2</sub>O<sub>2</sub> per minute at 37°C under the conditions specified in the assay procedure.

### Reagents

- Reaction mixture
 

0.2 M Tris-HCl buffer pH 8.0	0.20 ml
15 mM 4-AA solution	0.10 ml
0.2% (W/V) Phenol solution	0.10 ml
50 U/ml POD solution <sup>1)</sup>	0.10 ml

- |   |         |
|---|---------|
| 1% (W/V) Triton X-100 solution            | 0.10 ml |
| 5 mM Palmitoyl-CoA solution <sup>2)</sup> | 0.10 ml |
| Distilled water                           | 0.30 ml |

#### 1): 50 U/ml POD solution

Dissolve 500 U (PPU) of POD with 10 ml of distilled water.

#### 2): 5 mM Palmitoyl-CoA solution

Dissolve 50.3 mg (purity calculation) of palmitoyl-CoA with 10 ml of 10 mM KH<sub>2</sub>PO<sub>4</sub>-NaOH buffer pH 7.0.

#### 2. Enzyme dilution buffer

10 mM KH<sub>2</sub>PO<sub>4</sub>-NaOH buffer (pH 7.0) containing 3 mM ATP and 10 μM FAD

#### 3. Reagents

4-AA: NACALAI TESQUE, INC. Special grade #01907-52  
POD: Sigma Chemical Co. Type II # P-8250  
Triton X-100: The Dow Chemical Company  
Palmitoyl-CoA: Asahi Kasei Pharma Corporation  
ATP (2Na · 3H<sub>2</sub>O): Kyowa Hakko Co., Ltd.  
FAD (2Na): Kyowa Hakko Co., Ltd.  
ATP: Adenosine triphosphate  
FAD: Flavine adenine dinucleotide

## ■ Enzyme solution

Accurately weigh about 20 mg of the sample and add enzyme dilution buffer to make a total of 20ml. Dilute it with enzyme dilution buffer to adjust the concentration to within 0.2-0.5 U/ml.

## ■ Procedure

1. Pipette accurately 1.0 ml of reaction mixture into a reaction cuvette (1 ml volume black cuvette) and preincubate at 37°C.
2. After 5 min, add 20  $\mu$ l of enzyme solution and mix to start the reaction at 37°C.  
※ In the case of a test blank, add 20  $\mu$ l of enzyme dilution buffer in place of enzyme solution.
3. After starting the reaction, measure the rate of increase in absorbance at 500 nm. The rate must be measured within the linear portion of the absorbance curve.

Absorbance sample : As/min  
blank : Ab/min

$$\Delta A/\text{min} = (A_s/\text{min} - A_b/\text{min}) \leq 0.060 \text{ Abs/min}$$

## ■ Calculation

$$\text{Activity (U/mg of powder)} = \frac{\Delta A/\text{min}}{12.0 \times 1/2} \times \frac{1.02}{0.02} \times \frac{1}{X}$$

12.0 : millimolar extinction coefficient of quinoneimine dye at 500 nm ( $\text{cm}^2 / \mu\text{mole}$ )

1/2 : a multiplier derived from the fact that 2 mol of  $\text{H}_2\text{O}_2$  produce 1 mol of quinoneimine dye  
1.02 : final volume (ml)  
0.02 : volume of enzyme solution (ml)  
X : concentration of the sample in enzyme solution (mg/ml)

## Storage

Storage at  $-20^\circ\text{C}$  in the presence of a desiccant is recommended. Enzyme activity will be retained for at least one year under this condition (Figure 4).

## References

1. Shimizu, S., Yasui, K., Tani, Y. and Yamada, H. (1979) Biochem. Biophys. Res. Commun., **91** (1), 108-113.
2. Kikuchi, T., Ogawa, M., Ando, M. and Nakagiri, Y. (1979) Proceedings of Japanese Conference on Biochemistry of Lipids, **21**, 144-147.
3. Hosaka, K., Kikuchi, T. and Mitsuhida, N. (1979) Proceedings of the Symposium on Chemical Physiology, **19**, 180.
4. Shimizu, S., Inoue, K., Tani, Y. and Yamada, H. (1980) Anal. Biochem., **107**, 193-198.
5. Hosaka, K., Kikuchi, T., Mitsuhida, N. and Kawaguchi, A. (1981) J. Biochem., **89**, 1799-1803.
6. Kawaguchi, A., Tsubotani, S., Seyama, Y., Yamakawa, T., Osumi, T., et al. (1980) J. Biochem., **88**, 1481-1486.

## ACOD 活性測定法 (Japanese)

### I. 試薬液

1. 反応試薬混合液
 

0.2M トリス -HCl 緩衝液 pH8.0	0.20 ml
15mM 4-AA 溶液	0.10 ml
0.2% (W/V) フェノール溶液	0.10 ml
50U/ml POD 溶液 <sup>1)</sup>	0.10 ml
1% (W/V) トリトン X-100 溶液	0.10 ml
5mM パルミトイル-CoA 溶液 <sup>2)</sup>	0.10 ml
精製水	0.30 ml

1): 50U/ml POD 溶液

POD500 単位 (PPU) を精製水 10ml で溶解する。

2): 5mM パルミトイル-CoA 溶液

パルミトイル-CoA 50.3mg (純度換算) を 10mM  $\text{KH}_2\text{PO}_4$ -NaOH 緩衝液 pH7.0 10ml で溶解する。

### 2. 酵素溶解希釈用液

3mM ATP と 10  $\mu$ M FAD を含む 10mM  $\text{KH}_2\text{PO}_4$ - $\text{K}_2\text{HPO}_4$  緩衝液 pH7.0

### 3. 試薬

4-AA : ナカライテスク製 特級 #01907-52

POD : シグマ製 Type II #P-8250

トリトン X-100 : Dow Chemical 製

パルミトイル-CoA : 旭化成ファーマ製

ATP (アデノシン三リン酸  $\cdot 2\text{Na} \cdot 3\text{H}_2\text{O}$ ):

協和発酵製

FAD (フラビンアデニンジヌクレオチド  $\cdot 2\text{Na}$ ):

協和発酵製

### II. 酵素試料液

検品約 20mg を精密に量り、酵素溶解希釈用液に溶解して全容 20ml とする。

その液を酵素溶解希釈用液で 0.2~0.5U/ml 濃度となるように適宜希釈する。

### III. 測定操作法

1. 反応セル (1ml 用ブラックセル) に反応試薬混合液 1.0ml を正確に分注して 37°C で予備加温する。
2. 5分経過後、酵素試料液 20  $\mu$ l を加えて混和し、37°C で反応を開始する。  
※ 盲検は酵素試料液の代わりに酵素溶解希釈用液 20  $\mu$ l を加える。
3. 反応開始後、500nm における吸光度を測定して直線的に反応している 1分間当たりの吸光度変化を求めらる。  
求められた吸光度変化を試料液は As/min、盲検液は Ab/min とする。

$$\Delta A/\text{min} = (A_s/\text{min} - A_b/\text{min}) \leq 0.060 \text{ Abs/min}$$

### IV. 計算

$$\text{活性 (U/mg)} = \frac{\Delta A/\text{min}}{12.0 \times 1/2} \times \frac{1.02}{0.02} \times \frac{1}{X}$$

12.0 : キノンイミン色素の 500nm におけるミリモル分子吸光係数 ( $\text{cm}^2 / \mu\text{mole}$ )

1/2 :  $\text{H}_2\text{O}_2$  2 モルからキノンイミン色素 1 モルが生成することによる係数

1.02 : 反応総液量 (ml)

0.02 : 反応に供した酵素試料液量 (ml)

X : 酵素試料液中の検品濃度 (mg/ml)