Protein Components of a Cytochrome P-450 Linalool 8-Methyl Hydroxylase*

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The cytochrome P-450 heme-thiolate monooxygenases that hydroxylate monoterpene hydrocarbon groups are effective models for the cytochrome P-450 family. We have purified and characterized the three proteins from a P-450-dependent linalool 8-methyl hydroxylase in *Pseudomonas putida* (incognita) strain PpG777. The proteins resemble the camphor 5-exohydroxylase components in chemical and physical properties; however, they show neither immunological cross-reactivity nor catalytic activity in heterogenous recombination. These two systems provide an excellent model to probe more deeply the heme-thiolate reaction center, molecular domains of substrate specificity, redox-pair interactions, and the regulation of the reaction cycle.

Microbial catabolism of mono- and sesquiterpenoid hydrocarbons has focused primarily on reaction pathways (1-3), e.g. acyclic (4-6), alicyclic (7-9), and analogous aromatic structures (6). The initial hydroxylation of a methyl or methylene group provides substrates poised for catabolic pathways yielding carbon and energy for growth (15).

Due to the facility with which cytochrome P-450 monooxygenases catalyze the stereo- and regiospecific incorporation of an oxygen atom into "inert" hydrocarbon groups, the activation of terpenoid substrates is frequently P-450-mediated (9-11). These heme-thiolate proteins are broadly distributed in both pro- and eucaryotes (12-14) and share remarkable similarities in electronic and resonance spectral properties (12-16). Despite these similarities, P-450s are often extremely specific in substrate selection and electron transfer protein requirements. The selectivity of the substrate binding sites and requirements for a specific redox partner displayed by these monooxygenases are areas requiring further clarification.

The camphor 5-exo hydroxylase (EC 1.14.15.1) isolated

from Pseudomonas putida PpG786 (ATCC 29607) has provided a relevant model for the P-450 family through extensive physical and chemical characterization (15–20). It has also led the studies of P-450 structure and reaction mechanism, yielded primary structures via a combination of protein and DNA techniques (21–25), and a tertiary structure by x-ray analysis at 1.6-Å resolution (26, 27). Intermediate reaction states have been characterized by a variety of chemical and spectral techniques (15–20). The operon array and regulation have been documented (23–25).

This paper reports the purification and characterization of a second soluble P-450 monooxygenase. The linalool 8-hydroxylase, first detected in collaboration with P. K. Bhattacharyya, has been purified from the strain of P. putida (incognita) isolated by Madyastha et al. (6). The monooxygenase catalyzes, in two reaction cycles, the formation of 8-oxolinalool via the 8-alcohol. The component proteins of the linalool 8-hydroxylase are similar in general properties to those of the camphor-5-exo-hydroxylase, but little activity is obtained in heterogenous reconstitutions. Differences in the active site environments of the two P-450s have been indicated in preliminary publications (24, 54-56).

MATERIALS AND METHODS¹

RESULTS AND DISCUSSION

P. putida (incognita) strain PpG777 (formerly known under work number PH651), when grown on a phosphate ammonium salt (28) basal medium with (±)-linalool as the sole carbon and energy source, produces a P-450-dependent monoxygenase that catalyzes the first step in the catabolism of linalool. The whole cell content of LIN² P-450 reaches 2.5 mg of P-450/g of cells, wet weight, about 60% of the level of P-450_{cam} produced by P. putida, strain PpG786. Both strains release substantial amounts of the soluble protein components upon freeze-thaw autolysis, allowing ready purification on a large scale.

Protein Purification—The purification protocols detailed in the methods were derived from the procedures established for the CAM system (28) and are equally suited for the LIN and CAM components. Principal changes in the procedure for the reductase include use of the second DEAE ion exchange column before the Bio-Gel P-100 filtration, eluting the third DEAE column with a step gradient, and the addition of two affinity columns (Affi-Gel Blue and Matrix Gel Blue A) as the final steps in the purification. The redoxin purification

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¹ The "Materials and Methods" 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.

² The abbreviations used are: LIN, linalool; CAM, camphor; SDS, sodium dodecyl sulfate; PAGE, polyacrylamide gel electrophoresis.

TABLE I
Pseudomonas (LIN/CAM) reductase purifications

Purifications were normalized to 1 kg cell paste.

Step	Volume	Protein	Absorbance ratio	Activity ^b	Yield
	ml	g		units	%
Extract	2670/1690	161/90		42/620	100
DEAE-I	900/430	90/6.2		35/410	83/66
$(NH_4)_2SO_2$, 36–65%	100/90	75/5.5		34/380	81/61
DEAE-II	180/	89/		32/	76/
Bio-Gel P-100	220/270	15/2.5		29/340	71/55
DEAE-III	235/490	$2.8/0.5^{\circ}$	/10	17/210	41/34
Affi-Gel Blue	180/	0.3/	18.0/	15/00	37/
Matrix Gel Blue A	55/360	0.02°/0.21°	8.0/7.4	14/115	34/19

^a Determined in 50 mm Tris·Cl, pH 7.4, 5% glycerol (v/v); A₂₇₅/A₄₅₆, oxidized state.

Table II Pseudomonas (LIN/CAM) redoxin purifications Purifications were normalized to 3 kg cell paste.

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Step	Volume	Protein	Absorbance ratio	Redoxin	Yield
	ml	g		g	%
Extract	4040/3850	310/240		$0.85^{b}/1.8^{b}$	100
DEAE-I	830/1400	5.6/10.5	0.03/0.08	$0.60^{b}/1.4^{b}$	71/78
$(NH_4)_2SO_4$, 0-85%	53/70	3.7/5.5	0.05/0.46	$0.49^{b}/1.35^{b}$	58/75
Bio-Gel P-30 (P-100)	295/450	1.1/1.5	0.12/0.59	$0.42^{b}/0.90^{b}$	49/50
DEAE-II	204/330	0.34/0.65	0.90/0.71	0.34°/0.65°	39 [′] /36

 $^{^{\}circ}A_{324}/A_{280}$, oxidized state, in 50 mm Tris·Cl, pH 7.4, 10 mm 2-mercaptoethanol.

Table III $Pseudomonas\ (LIN/CAM)\ cytochrome\ P\text{-}450\ purifications}$ Purifications were normalized to 3 kg cell paste.

Step	Volume	Protein	Absorbance ratio	Cytochrome P450	Yield
	ml	g		g ^b	%
Extract	3730/3850	260/240	0.04/0.05	3.4/6.0	100
DEAE-I	985/1100	76/17	0.07/0.30	2.8/4.5	82/75
(NH ₄) ₂ SO ₄ , 36–60%	180/90	47/12	0.12/0.77	2.6/4.2	76/70
Bio-Gel P-100	130/480	5.3/5.5	0.66/1.05	1.7/3.3	50/55
DEAE-II	330/420	1.3°/2.3°	1.42/1.45	1.1/2.1	32/33

^a In 50 mM K-P_i, pH 7.0, 100 mM KCl, 200 μ M linalool or camphor; A_{391}/A_{280} , substrate-saturating oxidation state.

was modified by replacing Bio-Gel P-100 with Bio-Gel P-30 for the filtration step. The procedure for purification of the P-450 component remains essentially unchanged. All three protocols include minor changes in buffer composition.

The data in Tables I-III show the results of typical purifications of LIN reductase, redoxin, and P-450. For comparison the data for the purification of the corresponding CAM components are included in the tables. The data have been normalized to standard weights of cell paste (1 kg for the reductase procedure; 3 kg for the redoxin and P-450 components). Due to the high spectral background, purifications of the reductase and redoxin were monitored by NADH oxidase activity (see "Materials and Methods"). The P-450 purification was monitored by specific content (nmol of P-450/mg of protein) determined from the Fe^{II}CO-Fe^{II} difference spectrum (28). Lin P-450 and redoxin levels are ~50% of their CAM system counterparts, while LIN reductase is present at ~7% of the level for CAM reductase. Whether this difference arises from levels of expression or enzyme stability is unknown. Purification of LIN reductase shows an increased yield (2fold) relative to the CAM system. Most of this difference arises with the first DEAE column and the Matrix Gel Blue A column (Table I). Yields of CAM and LIN redoxin and P-450 are essentially equivalent (Tables II and III). The purification achieved with each step varies for the two systems due to differences in the content of the enzymes and the background in cell extracts from the two bacterial strains.

Criteria of Purity—Purity of the proteins in the final pools was estimated from SDS-PAGE and correlated with characteristic spectral parameters of the holoenzymes. LIN reductase with $A_{275}/A_{456} = 8.5$ was estimated to be >85% homogenous in total protein; LIN redoxin with $A_{324}/A_{280} = 0.90$ was >90% pure; and LIN P-450, in the presence of saturating substrate, had $A_{392}/A_{280} = 1.43$ and was >90% homogenous.

Storage and Stability—Following ultrafiltration or dialysis, the concentrated LIN redoxin reductase (0.16 mm), LIN redoxin (1 mm), and LIN P-450 (1 mm) solutions were packaged in 3-ml aliquots and stored in liquid nitrogen. The enzyme stocks responded differently to storage and freezethaw cycles. LIN redoxin reductase and LIN P-450 demonstrates.

^b Unit = 1 nmol of NADH oxidized/s.

 $^{^{}c}E_{454} = 10 \text{ mM}^{-1} \text{ cm}^{-1} \text{ for } M_{\rm r} = 43,500.$

^b Estimated from activity measurements.

^c Estimated with $E_{412} = 11.3 \text{ mM}^{-1} \text{ cm}^{-1}$ (LIN), $11.1 \text{ mM}^{-1} \text{ cm}^{-1}$ (CAM), $M_r = 11,600$.

 $^{^{}b}E_{448-490} = 93 \text{ mM}^{-1} \text{ cm}^{-1} \text{ for } m_{co}^{rs} - m_{co}^{rs}; M_{r} = 45,000 (28).$

 $^{^{}c}E_{280} = 63 \text{ mM}^{-1} \text{ cm}^{-1}.$

Table IV
Pseudomonas flavin (FAD) reductase properties

Property	LIN Reductase	CAM Reductase
Molecular weight, M_r	***	
Analytical data	$43,700^a$	$42,700^{a,b}$
SDS-PAGE	44,000	$47,000^{c}$
Gel filtration	45,000	$45,000^{\circ}$
Isoelectric point, pI	•	
Oxidized	4.66	4.38°
UV-visible, nm/mM		
Oxidized	275, 381, 456, 480,	275, 378, 454, 480,
	72, 9,2, 10.0, 8.3	72, 9.7, 10.0, 8.5
Turnover number, d s	-1	, , ,
Homologous	43	25
Heterologous	ND^e	2
Immunological reaction	on ^f	
Homologous	+++	+++
Heterologous	_	_

^a Determined from amino acid composition and FAD group.

TABLE V
Pseudomonas iron-sulfur (Fe₂S₂*) redoxin properties

Pseudomonas i	ron-sugar (Fe ₂ S ₂ °) rea	oxin properties
Property	LIN redoxin	CAM redoxin
Molecular weight, M_r		
Analytical data	12,800°	$11,594^{a,b}$
SDS-PAGE	10,700	11,700°
Gel filtration	11,000	11,000
Isoelectric point, pI	·	
Oxidized	<4.5 forms apopro-	<4.5 forms apopro-
	tein	tein
UV-visible, nm/mm ⁻¹		
Oxidized	277, 280, 324, 412,	277, 280, 328, 410,
	455, 17.8, 17.5,	455, 22.7, 21.9,
	15.7, 11.3, 9.9	15.6,
		11.1, 10.4
EPR , ^{d}g values		,
Reduced	2.021, 1.938	2.022, 1.938
Turnover number, e s-1	-,	,
Homologous	2.0	2.0
Heterologous	\mathbf{ND}^f	ND

^a Determined from amino acid composition and Fe₂S₂*Cys₄ group.

strated no measurable loss of enzymatic activity from prolonged storage at $-196\,^{\circ}\mathrm{C}$ nor from repeated freeze-thaw cycles. LIN redoxin, however, loses the prosthetic group during standing at 0 $^{\circ}\mathrm{C}$ in the absence of an active reducing agent, as evidenced by the decrease in absorbance in the iron sulfide chromophore; reagent grade dithiothreitol (5 mm) was found to stabilize the redoxin by retarding apoprotein formation.

Physical and Chemical Characteristics—Pertinent physical and chemical characteristics for the three components of the LIN monooxygenase are reported in Tables IV-VII; properties of the CAM system enzymes are included for comparison. In addition to similarities of the three LIN proteins and their

TABLE VI
Pseudomonas cytochrome P-450 (heme-thiolate) properties

	TOILE 1 -400 (Helle-th	
Property	LIN P-450	CAM P-450
Molecular weight, M_r		
Analytical data	44,800°	$46,820^{a,b}$
SDS-PAGE	47,000	$50,500^{c}$
Gel filtration	45,000	$44,000^{c}$
Isoelectric point, pI		
Oxidized, m ^{os}	4.84	4.59°
UV-visible, nm/mm ⁻¹		
cm ⁻¹		
Oxidized		
$m^{ m o}$	280, 360, 417, 538,	280, 360, 417, 535,
	569, 63.9, 30.6,	569, 68.3, 36.7,
	102, 10.5, 11.2	115, 11.6, 11.9
$m^{ m os}$	280, 392, 510, 540,	280, 391, 510, 540,
	646, 62.5, 92.0,	646, 63.3, 102,
	11.7, 9.9, 4.9	13.0, 11.2, 5.4
Reduced		
m^{r}	411, 540, 75.6,	480, 540, 76.7,
	13.6	15.1
$m_{ m co}^{ m rs}$	411, 542, 73.7,	409, 542, 86.5,
	12.9	16.0
$m_{ m co}^{ m rs}$	447, 550, 120, 17.2	446, 550, 120, 14.0
$m_{ m co}^{ m rs}-m^{ m rs}$	446-490, 84.0	446-490, 92.8
EPR, g values		
Oxidized, m^0	1.91, 2.27, 2.44,	1.91, 2.26, 2.45,
	8.13, 3.56, 1.68,	7.85, 3.96, 1.77,
	1.98, 2.25, 2.41	1.97, 2.24, 2.41
K_D (substrate), c μ M		
Oxidized	3.6	2.9^c
Turnover number, d s ⁻¹		
Homologous	32	34
Heterologous	1.2	ND^e
Immunological reac-		
$tion^f$		
Homologous	+++	+++
Heterologous		

^a Determined from amino acid composition and protoheme IX group.

CAM counterparts in chromatographic behavior, there is also marked similarity in size, chemical composition, and spectroscopic properties. SDS-PAGE and gel filtration data indicate molecular weights of reductase (44,000) redoxin (11,000), and P-450 (46,000) very close to the values for the CAM hydroxylase components. The isoelectric points for the LIN reductase and P-450 were, respectively, 4.66 and 4.84, showing both to be less acidic than the corresponding CAM proteins. These are in agreement with the amino acid compositions of the proteins. The LIN reductase has 8 more basic residues than CAM reductase (His + Lys + Arg) and probably less acidic residues (Asx + Glx, -12; amide content unknown). Similarly, the LIN P-450 has 4 more basic residues and less acidic residues (Asx + Glx, -5) than P-450_{cam} (Table VII). As with CAM redoxin, the LIN iron sulfide protein loses the prosthetic group to acid decomposition (pH < 4.5) before reaching its isolectric point.

Despite the physical and chemical similarity of the two systems, the P-450s lack cross-reactivity in immunodiffusion assays with antisera raised against the purified proteins (Tables IV and VI).

Enzymatic Activity—Catalytic properties for each of the LIN monooxygenase components and the corresponding

^b Tsai et al. (52).

^c Gunsalus and Wagner (28).

^d Substrate-stimulated NADH oxidation using excess amounts of redoxin and P-450 from the native (homologous) or related (heterologous) *Pseudomonas* system.

[&]quot; Not detected.

^fRabbit antibody raised against reductase from native (homologous) or related (heterologous) *Pseudomonas* system.

^b Amino acid sequence, Tanaka et al. (53).

^c Gunsalus and Wagner (28).

 $[^]d\,50$ mM Tris, pH 7.4, 10 mM 2-mercaptoethanol, 25 °C; X-band measurements at 7 K, dithionite-reduced.

^e Substrate-stimulated NADH oxidation using excess amounts of reductase/P-450 from the native (homologous) or related (heterologous) *Pseudomonas* system.

^{&#}x27;ND, Not detected.

^b Amino acid sequence, Haniu et al. (21, 22, 24).

^c Gunsalus and Wagner (28), CAM P-450 only.

^d Substrate-stimulated NADH oxidation using excess amounts of redoxin/reductase from the native (homologous) or related (heterologous) *Pseudomonas* system.

e Not detected.

[/]Rabbit antibody raised against P-450 from native or related (heterologous) *Pseudomonas* system.

Table VII

Amino acid composition of LIN/CAM monooxygenase proteins

	Re	Reductase		Redoxin		Cytochrome P-450	
	LIN	CAM ^a	LIN	CAM ^b	LIN	CAM ^{c,d}	
Asx	28	40(15)°	10	13(3)e	40	35(13)°	
Thr	16	20	4	5	14	20	
Ser	28	18	9	7	13	24	
Glx	40	$40(24)^{e}$	15	$10(5)^{e}$	46	56(23)°	
Pro	13	18	7	4	26	30	
Cys	6	6	4	6	5	8	
Gly	46	33	12	8	32	25	
Ala	47	48	9	9	30	31	
Val	40	34	7	14	32	24	
Met	4	6	3	3	6	10	
Ile	20	24	6	6	17	26	
Leu	39	42	15	6	45	42	
Tyr	11	6	1	3	7	9	
Phe	10	10	4	1	19	18	
His	10	6	2	2	10	13	
Lys	7	13	1	3	12	12	
Arg	34	24	7	5	33	26	
Trp	3	3	1	ĺ	6	5	
- *P	402	391	$1\overline{17}$	106	393	414	

- ^a Tsai et al. (52).
- ^b Tanaka et al. (53).
- ^c Haniu et al. (21, 22).
- ^d Corrected via cloned camC DNA.
- ^e Values in parentheses are, respectively, Asn or Gln.

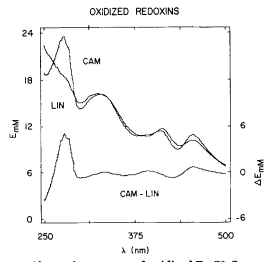


FIG. 1. Absorption spectra of oxidized Fe₂S*₂Cys₄ proteins. Conditions as specified in Table V and under "Materials and Methods."

CAM proteins are summarized in Tables IV-VI. Each turnover number (nmol of NADH oxidized/s/nmol of enzyme) was determined in reconstitutions with the other system components in saturating excess. The P-450 and redoxin components from each system are equivalent, while the LIN reductase turnover number is 1.7 times that of its CAM counterpart (Table IV).

While the similarity in chemical, physical, and catalytic properties between the components of the LIN and CAM monooxygenases suggests similar structures, little activity is observed in assays using heterologous reconstitutions (Tables IV-VI). CAM reductase couples with LIN P-450 and redoxin to hydroxylate linalool at <10% of the native system turnover; the activity observed for LIN P-450 supported by the CAM redox proteins attains <5% of the native turnover. These observations indicate that the LIN redox proteins cannot support reactions by the CAM components, suggesting as a

first hypothesis that the redox potentials of the LIN reductase and redoxin may be greater than those of the corresponding CAM proteins.

Absorption Spectra—The extinction coefficients at various wavelength maxima are reported for oxidized LIN reductase and LIN redoxin in Tables IV and V. The reductase spectrum is typical of a flavoprotein in showing maxima at 275, 381, 456, and 480 nm; the extinction coefficients were essentially identical to those of CAM reductase. The iron sulfide center of LIN redoxin, like the CAM redoxin, shows maxima at 324, 412, and 445 nm with similar extinction coefficients, whereas in the UV at 270 nm the aromatic absorption is markedly less (Fig. 1, Table VII). The extinction coefficients for LIN P-450 in oxidized and reduced states with various ligands are reported in Table VI. The heme content of the P-450 holoenzyme was determined by the pyridine hemochrome measurement of iron protoporphyrin IX (49). In the oxidized substrate-free state, LIN P-450 shows a Soret maximum at 417 nm, characteristic of low spin ferric heme. Upon addition of substrate, the spin equilibrium is shifted in favor of the high spin form, with a Soret maximum at 392 nm. The ferrous-CO complex of LIN P-450 has a maximum at 447 nm with extinction coefficient equal to that for CAM P-450.

LIN Monooxygenase Reaction Products—Substrate oxidation was monitored by gas chromatography of sample extracts taken at intervals from a reconstitution of LIN monooxygenase with NADH and linalool. Two products were detected, with product 1 (P1) appearing as early as 1 min into the reaction and product 2 detected after ~6 min. In order to identify these products, a large-scale reaction mixture was prepared as described under "Materials and Methods." The two products, purified by column chromatography, were then characterized by ¹H NMR, IR, and mass spectra. Spectral data and assignments are reported under "Materials and Methods."

The first product was identified as 8-hydroxylinalool on the basis of the mass spectral molecular ion, m/z = 170 (linalool + 0); IR data supporting -OH as the only oxygen function; ¹H NMR consistent with -CH₂OH at C8 or C9 and nuclear Overhauser enhancement refinement to assign the CH₂OH as cis to C6-H, i.e. 8-hydroxylation. The second product was identified as 8-oxolinalool on the basis of the mass spectral molecular ion, m/z = 168 (²H oxidation of P1), IR support for both carbonyl oxygen and aldehydic proton, and NMR analysis supporting the aldehyde assignment cis to C6-H. The identifications of P1 and P2 were further supported by comparison with synthetic compounds. Thus, LIN monooxygenase catalyzes two sequential oxidations of linalool, producing first 8-hydroxylinalool and further oxidizing this product to 8-oxolinalool. The oxidation of the alcohol to the aldehyde in the LIN system parallels the conversion of 5-exo-hydroxycamphor to 5-ketocamphor by the CAM monooxygenase (Fig. 2). In both systems, each oxidation requires the three protein components, NADH, and O₂. Both reactions were fully inhibited by CO; the preparations were free of detectable alcohol dehydrogenase activity. The occurrence of P-450-catalyzed alcohol oxidation is interesting, since both strains have substantial alcohol dehydrogenase activities for these substrates, with the corresponding enzymes having been purified to homogeneity.3 It is probable that a second hydroxylation by the P-450 gives rise to transient gem diol adducts that spontaneously dehydrate to the more stable carbonyl compounds. Purification of a P-450-dependent linalool 8-hydroxylase from Pseudomonas incognita, strain PpG777, contrasts with the reports by Bhattacharyya et al. (6, 50, 51) of a series of

³ A. J. H. Ullah and I. C. Gunsalus, unpublished results.

CAMPHOR-5-exo-

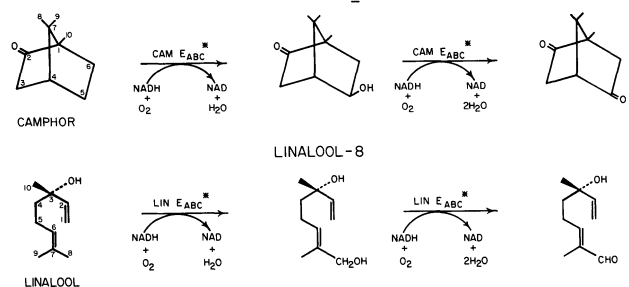


Fig. 2. Reactions catalyzed by the P-450_{cam} and P-450_{lin} terpene monocygenases. ${}^*E_{ABC} = FAD$ reductase, Fe₂S₂Cys₄ redoxin, heme thiolate (P-450) monooxygenase.

products indicating also a 10-hydroxylation of linalool from the freshly isolated P. incognita culture. The 10-methyl hydroxylation was not observed in the Psuedomonas Hardy isolate. The elucidation of the very interesting genetic control of these strains is continuing (55).

The initial characterization of the LIN and CAM monooxygenases indicates that despite the chemical and physical similarities, the component interactions clearly define two unique systems. Spectral characteristics also delineate active site differences in the P-450s substrate binding domains and in the water structure associated with the sixth ligand position (54-57). Current studies focus on the primary and tertiary structure and functional aspects of these two P-450 systems and on the control of the protein expression.

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Supplemental Material to "Protein Components of a P450 Linalool-8-methyl Hydroxylase"

A.J.H. Ullah, R.I. Murray, P.K. Bhattacharyya, G.C. Wagner, and I.C. Gunsalus

Materials and Methods

BACTERIAL CULTURE

BAUTERIAL CULTURE

FREUDOMORAS incognita strain PpG777 (work #PH651), a P. <u>putida</u>
variant, selected from subcultures of the Madyastha isolate (6) by P.
Harder, was maintained on minimal phosphate-ammonium-inorganic salts (PAS)
agar plates and fed by vapor from 0.1 ml of neat (±)-linalool spread in the
petri dish top. Stock cultures were transferred monthly and were stored at
4°C.

Harder, was maintained on minimal phosphate-ammonium-inorganic saits (PAS) agar plates and fed by vapor from 0.1 ml of neat (1-1 inalool spread in the petri dish top. Stock cultures were transferred monthly and were stored at 4°C.

The basal PAS medium for liquid culture was prepared from a phosphate-ammonium solution (25 mM KH₂PO₄, 50 mM K₂HPO₄, 150 mM NH₂Cl) supplemented with inorganic saits: 100x saits I containing (g/l): MgSO₂, 9.75; NMSO₄H₂O, 5.0: FeSO₄7H₂O, 5.0; CaCl₂·2H₂O, 0.3: and ascorbic acid, 1.0: 1000x saits II, containing (g/l): (NH₂)₂RO₂O₂, 0.185; CuSO₄·5H₂O, 3.92; CC(NO₃)₂·10H₂O, 5.0; ascorbic acid, 1.0. The medium was also supplemented with Bacto yeast extract (DifcO) and FNN (Signa) to final concentrations of 0.1t (v/v) and 2 uM, respectively. The L-broth contained (g/l): Bactotryptone (DifcO), 10.0; Bacto yeast extract (DifcO), 5.0; and glucose, 1.0: and was supplemented with Cb (100 ug/ml) immediately prior to innoculation.

Tube and flask cultures were incubated at 30°C in a New Brunswick shaker for the specified times. Liquid cultures were started by loop innoculation from the stock plate into 5 ml L-broth. The cultures were incubated 8 hours and transferred at 4t innoculation to 24 tubes containing 5 medium. Growth was monitored turbidometrically at 660 mm with a Cary 219 spectrometer, using a 3 nm spectral band width. When the flask cultures reached Ago = 0.4, growth on linalool was initiated by addition of 200 ul/flask (2)-linalool neat (concentration of 2.25 mM). After 5 n the cultures were actively growing on linalool was initiated by addition of 200 ul/flask (2)-linalool neat (concentration of 2.25 mM). After 5 n the cultures were actively growing on linalool was initiated by addition of 200 ul/flask (2)-linalool neat (concentration of 2.25 mM). After 5 n the cultures were actively growing on linalool was initiated by addition of 200 ul/flask (2)-linalool neat (concentration of 2.25 mM). After 5 n the cultures were actively growing on linalool

PROTEIN PURIFICATION PROCEDURES

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Buffers. Two stock buffers were used: 50 mM Tris-cl, pH 7.4 at 25°C, termed buffer T: and 20 mM potassius phosphate, pH 7.0 (buffer P). The ionic strengths of the buffers were increased for elution of ion exchange columns by addition of KC1; mM KC1 concentrations in buffer T are indicated as T-20, T-50,... T-600. A similar nomenclature is used for the phosphate buffers. For the purification of P450; all buffers contained 175 mM linalool. Buffers used in the purification of LIN-redoxin were supplemented with a 2-mercaptoethanol at 10 mM, while those used for the purification of LIN-reductase were made 5 or 10% in glycerol as noted in the text.

Assays. Analytical measurements followed the procedures of Gunsalus and Magner (28) with the following modifications. Total protein was estimated with Coomassie Brilliant Blue C-250 according to Sedmark and Grossberg (29) with an ovalbumin standard. After partial purification, protein concentration were estimated from the 280 m absorption with concentration were estimated from the 280 m absorption with concentration were retimated from the 280 m absorption with concentration were settimated from the 280 m absorption with concentration were estimated from the 280 m absorption with concentration were settimated from the 280 m absorption with concentrations were estimated from the 280 m absorption with concentrations were estimated from the 280 m absorption with concentrations were estimated from the 280 m absorption with concentrations were estimated from the 250 m absorption with concentration. The concentration of the 250 mm absorption with concentration were estimated for the 250 mm the 250 mm absorption with concentration. The concentration of the 250 mm absorption with concentration were estimated for the 250 mm the 250 mm absorption with concentration. The concentration of 150 mm absorption with concentration were estimated for the concentration of 150 mm absorption with concentration were estimated for the concentration of 150 mm abso

Purification Protocols

The modifications of the Gunsalus and Wagner procedure (28) reported here, provide a simplified method with improved yields and are equally well suited for the separation and purification of the CAN and LIN Components. The control of the provide as implified method with the propagation of the cell-free extract and the first ion exchange throwatography. The single exception is that for purification of the reductase, all buffers contain 10% glycerol (y/v), and for purification of the StO and redown buffers contain 10% united and 10% and 2-see Stop 1: Cell Free Extract. Storage of cells at -20°c for a minimum of one week before thaving at room temperature is considered to aid in the rupture of the cells and release of protein. A 3 kg portion of cell paste buffer T containing phenylechyleulfoldy flouride (1 mg), as the cell mass thaved the temperature of the suspension dropped rapidly to 3°C.

Manipulation with a spatul cawe a homegnous solution. The mixture was then a state of the suspension dropped rapidly to 3°C.

Manipulation with a spatul cawe a homegnous solution in the mixture was then a state of the suspension dropped rapidly to 3°C.

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Manipulation with a spatul cawe a homegnous solution in the mixture was then a state of the suspension of the supplement of the palled of the supplement of the palled of the palled. The poll was then subjected to DEAR ion exchange chromatography.

Manipulation of the squeens phase of the pellet. The clarified solution of Tris (free base). The pool was then subjected to DEAR ion exchange chromatography uses two columns in sequence to allow rapid elution of the subject of the supplement of the squeens placed to the pellet. The clarified solution of Tris (free base). The pool was then subjected to DEAR ion exchange chromatog

Step 2: Affi-Gel Blue Affinity Chromatography. The dialyzed enzyme fraction, from the previous step, was applied to an Affi-Gel Blue (Bio-Rad) column (3.3 x 22.0 m), equilibrated with buffer T. The column was developed with the equilibration buffer at a flow rate of 60 ml h⁻¹. The remaining LIN P450 was eluted very early, while LIN redoxin reductase activity was slightly retarded by the affinity column and was eluted in the later fractions. The active LIN redoxin reductase fractions were combined and concentrated to 15.0 ml by the ultrafiltration methods described in Step 6.

and concentrated to 15.0 ml by the ultrafiltration methods described in Step 6. Step 8: Matrex Gel Blue A Affinity Chromatography. The reductase from the Affi-Gel Blue column (Step 7), was applied to a Matrex Gel Blue A (Amicon) affinity column (1.9 x 38 cm), equilibrated with buffer T. The column was developed with the equilibration buffer at a flow rate of 30 ml h $^{-1}$. Elution of LIN redoxin reductase was retarded on this affinity column. The active fractions were combined and concentrated to 7 ml by ultrafiltration methods as described in Step 6, and stored in liquid nitrogen.

nitrogen.

LIN Redoxin:

Step 1:

O°C, Solid: (N₁₄)₂S₂6 (enzyme grade, Schwarz_Mann) was added to the redoxin pool from step 2 to a level of 560 g liter 1 (88 of saturation). After stirring for an additional 30 min at O°C, the precipitate was collected by centrifugation. The precipitate was suspended in buffer T-20 to a final volume of -50 ml and subjected to rapid dialysis under an argon atmosphere to achieve complete dissolution of the proteina.

Step 2: Gel Filtration Chromatography. The dialyzed solution was clarified by centrifugation and applied to a Bio-Gel P-30 gel filtration column (5 x 80 cm), equilibrated with buffer T-20. x 50 ml portion of buffer T-20 containing 20% sucrose was applied before the sample to reduce gravitational effects, and the column was eluted with the equilibration buffer at a rate of 80 ml h⁻¹.

Step 2: DE-52 Ion Exchange Chromatography II. The LIN redoxin pool from Step 4 was applied at a flow rate of 300 ml h⁻¹ to a DE-52 cellulose column (6.2 x 19 cm) equilibrated with buffer T. The column was developed with a linear gradient (3.6 liters) of buffers T-150 to T-550 at a flow rate of 130 ml h⁻¹. Fractions were combined and concentrated by ammonium sulfate precipitation of the protein as described in Step 3. The precipitate recovered by centrifugation was suspended in buffer T and dialyzed under an argon atmosphere to achieve complete dissolution and eliminate ammonlum sulfate. The dialyzed redoxin pool was made 5 ml in dithiothreitol prior to storage in liquid nitrogen.

LIN PASO

Step 1: Ammonium Sulfate Fractionation. The PASO pool from column a (Step 1) was made 1 mM, in linalcol and fractionated with ammonium sulfate by adding 200 g liter 1 (36 saturation) of finely meshed (NM,),SO, (enzyme grade, Schwarz-Mann) over a 2 h period at 0°C. The precipitate recovered by centrifugation was discarded and a second fraction was obtained from the supernatant by further addition of (NM,),SO, (140 g liter 1, 60 to an argon atmosphere against the same buffer.

Sigp 1: Gal Filtration Chromatography. The dialyzed PASO fraction was applied to a Bio-Gel P-100 gel filtration column (8.8 x 100 cm) and pulled to a Bio-Gel P-100 gel filtration column (8.8 x 100 cm) of buffer P-20 containing 201 (w/v) sucrose. The column was then developed in the ascending mode with the equilibration buffer at 200 ml h of buffer P-20 in Exchange Chromatography II. The PASO pool from Step 2: DE-52 Ion Exchange Chromatography II. The PASO pool from Step 2: DE-52 Ion Exchange Chromatography II. The PASO pool with a containing 2.01 (w/v) sucrose. The column was then developed in the ascending mode with the equilibration buffer at 200 ml h on DE-52 cellulose column (7.0 to was papplied at a flow rate of 300 ml h to a DE-52 cellulose column (7.0 to was papplied at a flow rate of 300 ml h to a DE-52 cellulose column (7.0 to was papplied at a flow rate of 300 ml h to a DE-52 cellulose column (7.0 to was papplied at a flow rate of 300 ml h to a DE-52 cellulose column (7.0 to was papplied at a flow rate of 300 ml h to DE-52 cellulose column (7.0 to was papplied at a flow rate of 300 ml h to DE-52 cellulose column (7.0 to was papplied to 30 ml h to DE-52 cellulose column (7.0 to was papplied to 30 ml h to DE-52 cellulose column (7.0 to was papplied to 50 ml h to DE-52 cellulose column (7.0 to was papplied to 50 ml h to DE-52 cellulose column (7.0 to ml h to DE-52 cellulose column (7.0 to was papplied to 50 ml h to DE-52 cellulose column (7.0 to Ml h to DE-52 cellulose column (7.0 to Ml h to DE-52 cellulose column (7.0 to D

ANALYTICAL METHODS

SDS polyacrylamide gel electrophoresis was according to Laemmli (30) with st total acrylamide monomer, and isoelectric focusing used: polyacrylamide gels (5 x 100 mm) containing 2.2% (v/v) ampholine (pH 3.5-5.0) according to Catterall (31), with ovalbumin as standard. Immunodiffusion assays used the methods of Davis et al. (32). Antisera were raised to purified P450 and reductase (CAM and LIN) in female New Zealand rabbits (32) and were purified according to standard procedures (33).

Amino Acid Analyses - The amino acid compositions of the LIN proteins were determined on a Beckman 119CL analyser at the University of Illinois, Urbana facility. Supplementary data was provided by Dr. G.E. Tarr (14,35) of the University of Michigan Protein Sequencing Pacility. And an operation of The University of Michigan Protein Sequencing Pacility. And by Dr. K. T. Yasunobu (21,22) of the University of Hawaii, Monoa campus.

All amino acid analyses were removed after precipitation of the apoprotein at 0°C by addition of 20t (w/v) trichloroacetic acid and standing for 0.5 h. Heme was removed from P450 in by the acid butanone method used for P450 (and (36)). The P450 apoprotein was then precipitated with 20t trichloroacetic acid. The apoproteins were washed twice with diethyl ether/acetone (1:1), dried at ambient temperature, and hyrolyzed for 24 h in yacug, with 4 N methane sulfonic acid at 115°C. Cysteine content was ectimated as cysteic acid in samples oxidized with performic acid (37) prior to hydrolysis.

Amino acid mole fractions were averaged from duplicate measurements. The nearest integer values were calculated from the composition estimated by averaging least squares refinement (38) and an integer composition (prior products) in the composition for the composition of the products in Chicle, extracts were based on G methods used for metabolites in the camphor system (39). Compounds were separated on columns (1/8 inch x 6 feet, 80/100 Supelcoport as support) with 33 ov-17 (140°C) or 104 carbowax 200 (180°C) as stationary phase. Structures of products from the LIN-monoxygenase were determined using samples purified from a scaled-up reconstitution assay. A reaction volume of 160 ml contained 50 mM potassium phosphate (Ph.7.0), 2.2uM each of LIN-Patos and LIN-reductions, 1010 products and product (150 ether) and product (150 ether) and product (150 ether) and product (150 ether) and prod

and P2 samples, which started from 8-oxolinalyl acetate, will be presented elsewhere.

Instrumentation: Protein optical spectra were recorded at 25°C on a Cary 21°S spectrophotometer operating with a 1 ms spectral band width. EPR spectra were measured on a Brucker 9 GHz spectrometer (ER 200D) equipped with an Oxford cryostat and helium transfer system. Magnetic fields were calibrated from the six Mn* hyperfine lines in SFO powder (42-44).

Protein EPR spectra of frozen solutions (1 mk) were recorded with 5-10 G modulation amplitude and 100 kHz modulation frequency at non-asturating microwave power levels (45-47). Computational analyses, including redoxin optical spectra, hand-digitized in 1 mm increments, and amino acid composition data, were processed on a Horizon Northstar computer interfaced (8-bir) to an X-*P piotter. Proton MRS spectra were recorded on a Varian 220 kHz PT-spectrometer (HR220) and Nuclear Overhauser Effect (NCE) experiments (48) were carried out on a Nicolet 360 kHz FT-spectrometer (NT360). Positive chemical shifts (ppm) were downfield from tetramethylsilane (TMS). MRS mamples were prepared in CCC1₂ as 54 (w/*) solutions that were carefully degassed prior to measurement. Infrared spectra were obtained for neat liquids on a Nicolet 7199 PT-spectrometer. Mass spectra were deteranded on a Varian MARCH-5. Gas Chromatography was performed on a Hewlett-Packard series 5700 Research Gas Chromatograph, equipped with a flame ionization detector.