Evolutionary Relationships Among γ-Carboxymuconolactone Decarboxylases

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 γ -Carboxymuconolactone decarboxylase (EC 4.1.1.44) from Azotobacter vinelandii resembled the isofunctional enzymes from Acinetobacter calcoaceticus and Pseudomonas putida. All three decarboxylases appeared to be hexamers formed by association of identical subunits of about 13,300 daltons. The A. vinelandii and P. putida decarboxylases cross-reacted immunologically with each other, and the NH₂-terminal amino acid sequences of the enzymes differed in no more than 7 of the first 36 residues. In contrast, the A. calcoaceticus decarboxylase did not cross-react with the decarboxylase from A. vinelandii or P. putida; the NH₂-terminal amino acid sequences of these enzymes diverged about 50% from the NH₂-terminal amino acid sequence of the A. calcoaceticus decarboxylase.

Immunological comparisons conducted with protocatechuate oxygenase (EC 1.13.11.3) (2) y-carboxymuconolactone decarboxylase and (EC 4.1.1.44) (1), enzymes of the protocatechuate branch of the β -ketoadipate pathway, indicated that the Azotobacter species enzymes are evolutionarily homologous to isofunctional enzymes formed by fluorescent Pseudomonas species. As described here and elsewhere (1, 10, 13), isofunctional enzymes formed by members of other bacterial genera appear to be immunologically distant, and these organisms govern expression of the enzymes with induction patterns unlike those shared by Azotobacter and Pseudomonas species (12). The conclusions drawn from immunological evidence are supported by chemical and physical data presented here. The results show that the γ -carboxymuconolactone decarboxylases of Azotobacter and Pseudomonas species are closely related to each other and more distantly related to the evolutionarily homologous y-carboxymuconolactone decarboxylase of Acinetobacter species.

MATERIALS AND METHODS

Bacterial strains and growth conditions. Azotobacter vinelandii OP was obtained from Paul E. Bishop, North Carolina State University, Raleigh, N.C. Large-scale growth of *A. vinelandii* was accomplished at 30°C in a New Brunswick Fermacell CF-130 fermentor containing 100 liters of modified Burk medium (14). *Pseudomonas putida* PRS 2260 (4) and

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Acinetobacter calcoaceticus ADP 152 (17) were described recently. For immunodiffusion studies, the appropriate strains of A. vinelandii, P. putida, and A. calcoaceticus were grown in 500-ml Erlenmeyer flasks containing 150 ml of mineral medium (5, 14) supplemented with 10 mm ρ -hydroxybenzoate.

Purification of γ -carboxymuconolactone decarboxylase from A. vinelandii. All purification procedures were performed between 0 to 4°C. Buffer A was 20 mM Tris-hydrochloride (pH 7.5) containing 25 µM dithiothreitol; buffer B was 10 mM Na₂HPO₄-KH₂PO₄ (pH 7.0); and buffer C was 20 mM Trishydrochloride containing 0.1 M NaCl. Cell pastes with a wet weight of 200 g were suspended in three volumes of buffer A and disrupted by passage through an American Instrument continuous-flow French pressure cell at 12,000 lb/in². Unbroken cells and debris were removed by centrifugation at $40,000 \times g$ for 20 min. The resultant crude extract (Table 1 step 1) was brought to 30% saturation by the addition of ammonium sulfate followed by centrifugation at $40,000 \times g$ for 20 min. Ammonium sulfate treatment was repeated on the supernatant fraction until 75% saturation was reached. After centrifugation, the protein pellet was dissolved in buffer A (Table 1, step 2) and dialyzed against three changes of this buffer over 48 h. The dialysis was applied onto a DEAE-cellulose column (5 by 30 cm) previously equilibrated with buffer A. The column was washed with three volumes of the same buffer, after which a continuous linear gradient, constructed from 0 to 0.3 M NaCl in buffer A in a total volume of 6 liters, was applied. Fractions of 15 ml were collected at a flow rate of 60 ml/h, and those containing y-carboxymuconolactone decarboxylase activity, which eluted between 0.16 to 0.19 M NaCl, were pooled (Table 1, step 3). The DEAE-celluose eluate was treated with ammonium sulfate, and the protein pellet that precipitated between 45 to 75% saturation was dissolved in buffer B (Table 1, step 4) and dialyzed thoroughly against the same buffer. The dialysate was

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TABLE 1. Purification of γ -carboxymuconolactone decarboxylase from A. vinelandii

	Step	Volume (ml)	Total activity (U)	Total protein (mg)	Sp act (U/mg)	Yield (%)	Purification (fold)
1.	Crude extract	610	34,000	17,670	1.92	100	1.0
2.	First 30 to 75% saturated ammonium						
	sulfate fraction	310	30,130	8,290	3.63	89	1.9
3.	First DEAE-cellulose eluate	500	32,000	1,575	20.3	94	10.6
4.	Second 45 to 75% saturated ammo-						
	nium sulfate fraction	33	31,350	726	43.2	92	22.5
5.	Second DEAE-cellulose eluate	120	26,400	356	74.2	78	38.6
6.	Third 40 to 65% saturated ammonium						
	sulfate fraction	7	23,920	248	96.5	70	50.3
7.	Sephadex G-200 eluate	32	19,840	30.4	653	58	340
8.	Sephadex G-100 eluate	35	12,600	10.0	1,260	37	656
9.	Quaternary aminoethyl-Sephadex		,		-		
_	eluate	12	6,040	4.8	1,258	18	655

applied onto a second DEAE-cellulose column (1.6 by 20 cm) previously equilibrated with buffer B. The column was washed with 2 volumes each of 0.01, 0.05, and 0.1 M sodium potassium phosphate buffer, and then a continuous linear gradient, constructed from 0.1 M sodium potassium phosphate buffer to the same buffer containing 0.3 M NaCl in a total volume of 300 ml, was applied. Fractions of 3 ml were collected at a flow rate of 20 ml/h, and those containing the decarboxylase activity were pooled (Table 1, step 5) and fractionated with ammonium sulfate. The protein pellet that precipitated between 40 to 65% saturation was dissolved in buffer A (Table 1, step 6) and applied onto a Sephadex G-200 column (2.5 by 100 cm) previously equilibrated with buffer A. Fractions of 3 ml were collected at a flow rate of 14 ml/h, and those containing a specific activity of the decarboxylase greater than 600 U/mg were combined (Table 1, step 7) and concentrated with ammonium sulfate (0 to 70%saturation). The resultant pellet was dissolved in buffer A and loaded onto a Sephadex G-100 column (2.5 by 40 cm) previously equilibrated with buffer A. The eluate containing the decarboxylase activity (Table 1, step 8) was concentrated with ammonium sulfate (0 to 70% saturation), dialyzed thoroughly against buffer C, and applied onto a quaternary aminoethyl-Sephadex column (0.9 by 8 cm) previously equilibrated with buffer C. The column was washed with 10 volumes of the same buffer, and then a continuous linear gradient, constructed from 0.1 to 0.4 M NaCl in 20 mM Tris-hydrochloride (pH 7.2) in a total volume of 50 ml, was applied. Fractions of 2 ml were collected at a flow rate of 10 ml/h, and those containing the decarboxylase activity were pooled (Table 1, step 9). The quaternary aminoethyl-Sephadex eluate was stored at 4°C in the presence of ammonium sulfate at 70% saturation.

Polyacrylamide gel electrophoresis. Polyacrylamide gel electrophoresis was performed with 10% disc gels (9). The subunit size of *A. vinelandii* decarboxylase was estimated by sodium dodecyl sufate-gel electrophoresis (15) with previously described standard proteins (17).

Amino acid analysis. The amino acid composition of *A. vinelandii* decarboxylase was determined by a previously described procedure (4) and analyzed with a computer program (8) for the minimum molecular weight, which is the subunit size, of the enzyme. NH₂-terminal amino acid sequence determination. Previously published procedures (16) were used for determination of the NH₂-terminal amino acid sequence of *A. vinelandii* decarboxylase.

Serological studies. Antisera against A. vinelandii, P. putida, and A. calcoaceticus decarboxylases were prepared (2), and the method of Stanier et al. (13) was used to examine serological cross-reaction on Ouchterlony double-diffusion plates (7).

Chemicals. Chemicals were described previously (2, 17). Quaternary aminoethyl-Sephadex was obtained from Pharmacia Fine Chemicals.

RESULTS

Purity of A. vinelandii decarboxylase. The most purified preparation of A. vinelandii decarboxylase (Table 1, step 9) possessed a specific activity of 1,260 U/mg. When subjected to electrophoresis on 10% polyacrylamide gels, the decarboxylase preparation migrated as a major band and a slight minor band (Fig. 1). The minor band may have been an electrophoretically different form of the decarboxylase because, as described below, the preparation was immunologically homogeneous, and the NH₂-terminal amino acid sequence of the enzyme was determined without detectable contamination.

Molecular weight and subunit size determinations. The molecular weight of the A. vinelandii decarboxylase, close to the molecular weights of the A. calcoaceticus and P. putida decarboxylases, was estimated to be 85,500 by gel filtration (Fig. 1). The size of the A. vinelandii decarboxylase subunit was determined as 13,300 by sodium dodecyl sulfate-gel electrophoresis and as 13,460 by computer-aided analysis of the amino acid composition of the enzyme (8).

Serological properties. Antisera prepared against the purified A. vinelandii decarboxylase formed a single precipitin band when diffused against a crude extract of A. vinelandii cells in which the enzyme had been induced (Fig. 2). This band formed a spur with a precipitin band formed by a P. putida extract containing the



FIG. 1. Insert depicts a stained 10% polyacrylamide gel on which 50 μ g of A. vinelandii decarboxylase (Table 1, step 9) had migrated electrophoretically. The graph shows data indicating the molecular weights of decarboxylases determined by filtration on a standardized Bio-Gel agarose A 1.5m column (2.6 by 100 cm).

decarboxylase (Fig. 2). No precipitin band was formed with extracts of uninduced *A. vinelandii* and *P. putida* cells (not shown) or with *A. calcoaceticus* extract containing the enzyme (Fig. 2). Similar precipitin patterns were formed when purified decarboxylase was substituted for crude extract in the outer wells (Fig. 2). Thus, the *A. vinelandii* decarboxylase appears to be an immunologically homogeneous preparation that cross-reacts strongly with the *P. putida* decarboxylase and not at all with the *A. calcoaceticus* decarboxylase.

The conclusion that the A. vinelandii and P. putida decarboxylases resemble each other more closely than they resemble the A. calcoaceticus decarboxylase is fortified by the immunodiffusion patterns shown in Fig. 3. Antisera prepared against the P. putida decarboxylase form a precipitin band with the P. putida and A. vinelandii decarboxylases, but not with the A. calcoaceticus decarboxylase (Fig. 3). Antisera prepared against the A. calcoaceticus decarboxylase formed a precipitin band with this enzyme, but not with the corresponding decarboxylase from A. vinelandii or P. putida (Fig. 3).

Amino acid composition. The amino acid composition of the A. vinelandii decarboxylase is shown in Table 2. Quantitative comparison of amino acid compositions may be achieved by measurement of SAQ, the sum of the square of the difference in mole fraction of each amino acid that can be readily quantitated in a protein



FIG. 2. Double-diffusion plates showing the immunological cross-reactions of enzymes with A. vinelandii decarboxylase. The center wells received 0.2 ml (containing approximately 3 mg of protein) of antiserum against the A. vinelandii decarboxylase. The outer wells of the top plate received crude extracts of cells in which the decarboxylase had been induced by growth with p-hydroxybenzoate. The outer wells of the bottom plate received the indicated amounts of purified decarboxylases.



FIG. 3. Double-diffusion plates on which purified decarboxylases from different bacterial genera were diffused against antiserum prepared against P. putida decarboxylase (center well, top) and against antiserum prepared against A. calcoaceticus decarboxylase (center well, bottom).

hydrolysate. In over 5,000 pairwise comparisons, Marchalonis and Weltman (3) found that an $S\Delta Q$ of less than 50 invariably reflected a structural similarity that was revealed by comparison

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TABLE 2. Amino acid composition of A. vinelandii _Y-carboxymuconolactone decarboxylase

Amino acid	Amino acid residues per 13,460 daltons ^a
Asx	11.26
Thr	5.95
Ser	6.10
Glx	15.14
Pro	4.95
Gly	9.86
Ala	13.17
Cvs ^b	1.14
Val	9.04
Met	3.05
Ile	6.21
Leu	11.70
Tvr	2.63
Phe	3.66
His	4.12
Lus	4.76
Arg	7.91
Trp ^c	1.18

^a From computer-aided analysis of amino acid composition (8).

^b Estimated as cysteic acid after performic acid oxidation (4).

^c Determined after hydrolysis of a protein sample with 3 N mercaptoethanesulfonic acid (4).

of amino acid sequences. With the A. vinelandii decarboxylase as reference we found that $S\Delta Qs$ for the amino acid compositions of the A. calcoaceticus and P. putida decarboxylases were 9 and 24, respectively.

 NH_2 -terminal amino acid sequence. Quantitative measurements for the first 49 cycles in the NH_2 -terminal amino acid sequence determination of the *A. vinelandii* decarboxylase are shown in Table 3.

DISCUSSION

Relationships among γ -carboxymuconolactone decarboxylases from A. calcoaceticus, A. vinelandii, and P. putida. y-Carboxymuconolactone decarboxylases from the three bacterial genera shared some properties: the enzymes appeared to be hexamers formed by association of identical subunits of about 13,300 daltons (11), and the amino acid compositions of the proteins were similar (17). On the other hand, the specific activity of the A. calcoaceticus decarboxylase (140 U/mg) was substantially lower than the specific activities of the A. vinelandii and P. putida decarboxylases (1,260 and 1,310 U/mg, respectively), and the latter two decarboxylases cross-reacted immunologically with each other, but not with the A. calcoaceticus decarboxylase. Thus, it appears that all three decarboxylases were derived from a com-

TABLE 3. Automated sequence analysis of A. vinelandii γ -carboxymuconolactone decarboxylase: identification of amino acid residues

Cycle	PTH ^e	Me ₃ Si	BH-ORG [®]	BH-AQU ^e	Residue
1	M (193) ^d	M (181)			Met
2		D (125)	D (73)		Asp
3		E (166)	E (68)		Glu
4			K (37)		Lys
5		E (125)	E (68)		Glu
6			R (4)	R (17)	Arg
7	Y (18)	Y (101)	Y (46)		Tyr
8		D (65)	D (47)		Asp
9	A (131)	A (115)	A (53)		Ala
10	G (129)	G (106)	G (66)		Gly
11	M (66)	M (82)			Met
12		Q (20)	E (49)		Gln
		E (36)	11 (00)		
13	V (208)	V (153)	V (33)	-	Val
14			R (3)	R (8)	Arg
15			R (3)	R (14)	Arg
16	A (114)	A (104)	A (47)		
17	V (97)	V (75)	V (33)		VAL
18	G (99)	G (00)	L (35)		Leu
19	G (38)	G (33)	G (26)		Gly
20	A (70)	D (20)	D(31)		Asp
21	A (72)	A (65)	A (43)	H (7)	Ala Ula
22	V (990)	V (145)	V (96)	п(/)	Vel
23	V (230)	V (140)	V (20)		A ano
24		D (40)	B (2)	D (7)	Ara
20			1((2)	R (7)	(Cvs or
20					Ser)"
27	LI' (181)	LI (119)	L (26)		Leu
28			K (10)		Lys
29		D (11)	D (22)		Asn ^e
30	LI (163)	LI (117)	L (25)		Leu
31	-		T (14)		Thr
32	P (39)	B (04)	P (11)		PTO Db.
33	F (32)	F (34)	F (14)		Pne
34		D (7)	D (20)		Asn
35		E (40) E (49)	E (23) E (95)		Chu
30	F (20)	E (42) E (91)	E (20) F (11)		Dhe
37	г (32)	r (31) O (5)	F (11) F (22)		Gln
30		E (26)	E (44)		Jui
30		E (30)	E (24)		Ghi
40	M (9)	M (14)	13 (64)		Met
4U 41	LT (97)	LI (59)	I (11)		Пе
42	(01)	LII (00)	T (12)		Thr
43			$\mathbf{R}(\mathbf{i})$	R (3)	Arg
44			(-)	H (3)	His
45		A (45)	A (19)	,	(Ala)
46		,			(Cys or
					Ser)*
47		G (16)	G (12)		Gly
48		D (10)	D (11)		Asp
49	LI (138)	LI (67)	I (8)		lle

^e PTH. Phenylthiohydantoin.

^b BH-ORG, Back hydrolysis-extracted organic phase.

BH-AQU, Back hydrolysis-remaining aqueous phase.

^d Results indicate the single-letter amino acid designation and (within parentheses) the number of nanomoles recovered.

[•] The only two amino acid residues that could not be identified by the procedures used in this sequence determination are cysteine and serine.

¹PTH-leucine and PTH-isoleucine were coeluted by the gas chromatographic technique.

⁴ PTH-asparagine was distinguished from PTH-asparatic acid by high-pressure liquid chromatography.

mon ancestor; the structural genes for the A. vinelandii and P. putida decarboxylases seem to have diverged recently relative to their divergence from the A. calcoaceticus decarboxylase. These conclusions are strengthened by comparison of the NH₂-terminal amino acid sequences of the decarboxylases. The data allow a threeway comparison of 34 of the first 36 residues. The A. vinelandii and P. putida sequences were identical in 83% of the compared positions (Fig. 4), whereas comparison of the A. vinelandii sequence with the A. calcoaceticus sequence reveals an identity of 47% (Fig. 5).

Early events in the evolution of γ -carboxymuconolactone decarboxylases and muconolactone isomerases. γ -Carboxymuconolactone decarboxylases and muconolactone isomerases, enzymes that mediate analogous biochemical reactions, appear to share a common ancestral gene. Alignment of the NH₂-terminal

<u>Asotobacter</u>	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18
Decarboxylase	HET	ASP	GLU	LYS	GLU	ARG	TYR	ASP	Ala	GLY	Met	GLN	VAL	ARG	ARG	ALA	VAL	LEU
<u>Pseudomonas</u>	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18
Decarboxylase	HET	ASP	GLU	LYS	GLJI	ARG	TYR	ASP	Ala	GLY	Met	GLM	VAL	ARG	ARG	ALA	VAL	LEU
																		_
<u>Asotobacter</u> Decarboxylase	19 GLY	20 ASP	21 ALA	22 HIS	23 VAL	24 ASP	25 ARG	26 CTS (7)	27 LEU	28 LYS	29 ASM	30 LIEU	31 THR	32 PRO	33 Phe	34 ASH	35 GLU	36 GLU
<u>Pseudomonas</u>	19	20	21	22	23	24	25	26	27	28	29	30	31	32	33	34	35	36
Decarboxylase	GLY	ASP	ALA	HIS	VAL	ASP	ARG	CYS	LBU	GLY	L¥S	LEU	ASM	ASP	PHE	ASH	GLY	GLU

FIG. 4. Comparison of the NH₂-terminal amino acid sequences of A. vinelandii and P. putida γ -carboxy-muconolactone decarboxylases. Identical residues are enclosed in boxes.

Asotobacter	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18
Decarboxylase	HET	ASP	GLU	LYS	GLU	ARG	TTR	ASP	ALA	GLY	Met	GLM	VAL	ARG	ARG	ALA	VAL	LEU
Acinetobacter	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18
Decarboxylase	HET	ASH	ASP	GLU	GLH	ARG	TTR	LYS	GLM	GLY	Leu	GLU	VAL	ARG	THR	GLU	VAL	LEU
Asotobacter Decarboxylase Acinetobecter Decarboxylase	19 GLI 19 GLI	20 ASP 20 GLU	21 ALA 21 LYS	22 NIS 22 NIS	23 VAL 23 VAL	24 ASP 24 ASM	25 ABG 25 ABG	26 CYS (1) 26 SER	27 LIBU 27 LIBU	28 LTS 28 GLU	29 ASH 29 ASH	30 LIEU 30 LIEU	31 THR 31 ASP	32 PRO 32 ASP	33 PHE 33 PHE	34 ASU 34 ABC	35 GLU 35 GLM	36 GLU 36 ASP

FIG. 5. Comparison of the NH_2 -terminal amino acid sequences of A. vinelandii and A. calcoaceticus γ -carboxymuconolactone decarboxylases. Identical residues are enclosed in boxes.

Acinetobacter	1	2	3	4	-4-	5	6	7	8	9	10	11	12
Decarboxylase	Met	ASN	ASP	GLU		GLN	ARG	TYR	Lys	GLN	GLY	Leu	GLU
<u>Azotobacter</u>	1	-۵-	2	3	4	5	6	7	8	9	10	11	12
Decarboxylase	Met		ASP	GLU	L¥S	GLU	ARG	TYR	ASP	Ala	GLY	Met	Gln
<u>Pseudomonas</u>	1	-4-	2	3	4	5	6	7	8	9	10	11	12
Decarboxylase	Met		ASP	GLU	LYS	GLN	ARG	TYR	ASP	ALA	GLY	Met	GLN
Acinetobacter	24	25	26	27	28	29	-4-	30	31	32	33	34	35
Isomerase	L¥S	SER	VAL	GLU	Lys	ALA		TYR	SER	GLN	GLU	LEU	Glin
Pseudomonas	24	25	26	27	28	29	-4-	30	31	32	33	34	35
Isomerase	Lys	ALA	ASP	GLU	LYS	GLU		LEU	Ala	GLN	ARG	LEU	GLN

FIG. 6. A portion of the NH₂-terminal amino acid sequence of γ -carboxymuconolactone decarboxylase appears to be conserved within the primary structure of muconolactone isomerase. Numbers indicate the positions of residues in the primary sequences of the proteins. Boxes enclose residues at positions where identical residues are found in the decarboxylase-isomerase comparison.

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amino acids of the enzymes reveals sequence similarity suggesting a low overall homology (17). In addition, the NH₂-terminal amino acid sequence of the decarboxylase appears to be conserved within the primary structure of the muconolactone isomerase: the tetrapeptide extending from residues 2 through 5 in the A. *vinelandii* decarboxylase is represented at residues 26 through 29 in the amino acid sequence of the P. putida muconolactone isomerase (Fig. 6). This is consistent with the proposal that, as genes for the β -ketoadipate pathway became established, oligonucleotide substitution mutations placed sequences coding for peptides in novel structural contexts (6, 18).

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