Involvement of Coenzyme M during Aerobic Biodegradation of Vinyl Chloride and Ethene by *Pseudomonas putida* Strain AJ and *Ochrobactrum* sp. Strain TD†

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The involvement of coenzyme M in aerobic biodegradation of vinyl chloride and ethene in *Pseudomonas putida* strain AJ and *Ochrobactrum* sp. strain TD was demonstrated using PCR, hybridization, and enzyme assays. The results of this study extend the range of eubacteria known to use epoxyalkane:coenzyme M transferase.

Elucidating the pathway for vinyl chloride (VC) biodegradation is likely to improve methods for predicting the fate of this compound in natural and engineered environments. Under aerobic conditions, metabolism of VC and ethene, the nonchlorinated counterpart of VC, begins with epoxide formation (4, 8, 11, 20). The involvement of epoxyalkane:coenzyme M transferase (EaCoMT) in the processing of VC epoxide and ethylene oxide (EtO) has recently been demonstrated with several strains of *Mycobacterium* and one strain of *Nocardioides* (5, 6, 15). The EaCoMT gene is located on large linear plasmids in *Mycobacterium* strains and *Nocardioides* sp. strain JS614 when they are grown on VC (5, 15). The objective of this study was to determine if coenzyme M (CoM) is involved in the pathway of two other microbes that also grow on VC and ethene, *Pseudomonas putida* strain AJ and *Ochrobactrum* sp. strain TD (8).

Chemicals, media, and growth conditions are described elsewhere (7). Ethene, EtO, and VC consumption was quantified by gas chromatography (20). Growth rates were determined based on plots of protein concentration (measured by UV [4, 7, 12]) versus time (4). The EaCoMT assay was performed (6) with minor modifications (7). Cells were harvested at the early to mid-exponential phase (optical density at 600 nm, 0.2 to 0.35), washed, and resuspended in buffer (2, 6, 7).

Agarose plugs were prepared and pulsed-field gel electrophoresis (PFGE), DNA elution, and PCR were performed as described previously (5–8). Southern hybridization and radiolabeling of probe DNA were performed (3, 17) using a probe that was from the 891-bp EaCoMT gene fragment from strain AJ, using an Ambion DECAprime II labeling kit. Partial CoM gene sequences (834 bp) for various *Mycobacterium* strains were obtained from GenBank (accession no. AJ243034 to AJ243043).

The rates of EtO consumption by extracts of EtO-grown strain AJ increased from 57 ± 26 nmol min⁻¹ mg protein⁻¹ with no CoM added to 670 ± 70, 1,300 ± 80, and 2,600 ± 280 with 1, 10, and 100 mM CoM added, respectively (Fig. 1). For EtO-grown strain TD, the rates of EtO consumption were 86 ± 72, 1,800 ± 230, 3,500 ± 280, and 7,200 ± 710 nmol min⁻¹ mg protein⁻¹ with 0, 1, 10, and 100 mM CoM added, respectively.

EaCoMT activities were also determined following growth of strains AJ and TD on VC and ethene (Table 1). The activities with 10 mM CoM were 1 to 2 orders of magnitude higher than the activities without CoM. There were statistically significant differences among the EaCoMT activities with CoM for strains AJ and TD grown on VC, ethene, and EtO (α = 0.05, as determined by analysis of variance). Tukey’s range test indicated that the EaCoMT activity of strain AJ when it was grown on EtO (1,300 nmol min⁻¹ mg protein⁻¹) was statistically lower than the other activities (range, 3,400 to 4,600 nmol min⁻¹ mg protein⁻¹).

Table 1 also shows the EaCoMT activities with 10 mM CoM for Luria-Bertani broth (LBB)-grown strains AJ and TD following growth on VC, ethene, and EtO. Growth in LBB caused the EaCoMT activities to decrease by 1 order of magnitude, although the activities of LBB-grown cells were greater than the activities of controls with no cells. The EaCoMT activities of LBB-grown cells with CoM were higher than the activities of LBB-grown cells without CoM, although the value for only one of the LBB-grown treatments with CoM was statistically greater (α = 0.05, as determined by the t test) than the value for the corresponding treatment with no CoM (i.e., strain TD grown in LBB following growth on ethene).

CoM is a methyl carrier in the final step of methanogenesis (18, 21), which is inhibited by bromoethanesulfonate due to its reactivity with methyl-CoM reductase (10). Since bromoethanesulfonate inhibits reductase activity, it should not inhibit aerobic metabolism of VC, ethene, and EtO, which involves only transferase activity. Adding bromoethanesulfonate did not alter the EaCoMT activities with extracts of ethene-grown strain AJ containing 10 mM CoM (Fig. 2).

The EaCoMT assay was performed with glutathione in place.
of CoM. There was no difference in the EaCoMT activities between extracts with and without glutathione, indicating that glutathione is not involved in catabolism of EtO in strains AJ and TD (data not shown). Previous studies have shown that glutathione is a cofactor in epoxide degradation (19).

The maximum specific substrate utilization rates for strain AJ were 18, 22, and 62 nmol min\(^{-1}\) mg protein\(^{-1}\) when VC, ethene, and EtO were used as the substrates, respectively. For strain TD, the maximum specific substrate utilization rates were 20, 15, and 84 nmol min\(^{-1}\) mg protein\(^{-1}\) with VC, ethene, and EtO, respectively. Since the EaCoMT activities of strains AJ were 18, 22, and 62 nmol min\(^{-1}\) mg protein\(^{-1}\) and EtO, respectively. Since the EaCoMT activities of strains AJ and TD (Table 1) were at least 10 times higher than the maximum substrate utilization rates measured for whole cells, it was evident that EaCoMT is not a limiting factor during growth. The maximum substrate utilization rate determined in this study for strain AJ growing on VC was higher than the maximum rate reported previously (1.41 μmol VC mg biomass\(^{-1}\) day\(^{-1}\) or 1.8 nmol mg protein\(^{-1}\) min\(^{-1}\)) (8). Differences in the incubation temperature (22°C in the previous study and 30°C in this study) and in the assay procedures (steady-state reactor biomass in the previous study and exponentially growing cells in this study) likely contributed to the different rates. Variations in kinetic parameters are often a consequence of differences in the physiological state of the cells used (9).

DNA was eluted from plugs prepared from strains AJ and TD following growth on VC, ethene, and EtO. The DNA was subjected to PCR using primers based on conserved regions of the EaCoMT gene. Cells grown on VC, ethene, and EtO yielded a single PCR product of the expected size (891 bp), as did ethene-grown *Mycobacterium* sp. strain JS60. No product was obtained with a control containing no DNA (data not shown).

Growth of strains AJ and TD in LBB cures the large linear plasmid(s) required for use of VC and ethene as growth substrates (7), as well as for use of EtO by strains AJ (8) and TD (data not shown). Following growth of both strains in LBB, the cultures were also subjected to PCR, and a band of the expected size was recovered (data not shown), confirming the presence of the EaCoMT gene even after the large linear plasmids were cured. Sequencing of the 891-bp fragment confirmed the presence of the EaCoMT gene. However, there were differences in the sequence of the EaCoMT gene following growth in LBB compared to growth on VC (differences in four amino acids for strain AJ and in five amino acids for strain TD, based on the 891-bp fragment).

Southern hybridization was performed to determine if the EaCoMT gene was located on the plasmid. Agarose plugs of strains AJ and TD grown on VC and in LBB were subjected to contour-clamped homogeneous electric field PFGE (Fig. 3A) and then exposed to the EaCoMT probe (Fig. 3B). A hybridization signal for VC-grown strains AJ and TD corresponded to an approximately 260-kb plasmid (Fig. 3B, lanes 7 and 9) and then exposed to the EaCoMT probe (Fig. 3B). A hybridization signal for VC-grown strains AJ and TD corresponded to an approximately 260-kb plasmid (Fig. 3B, lanes 7 and 9). The strongest hybridization signal for VC-grown strain TD did not correspond to a plasmid from PFGE (lane 9). In addition, a hybridization signal was observed for both strains (lanes 7 and 9) in the compression zone (i.e., large, intact, high-molecular-weight DNA). The compression zone for strains AJ and TD following growth in LBB also exhibited hybridization

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**TABLE 1. EaCoMT activity in cell extracts of strains AJ and TD**

<table>
<thead>
<tr>
<th>Strain</th>
<th>Substrate</th>
<th>EaCoMT activity(^a) (nmol min(^{-1}) mg protein(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>10 mM CoM added</td>
</tr>
<tr>
<td>AJ</td>
<td>VC</td>
<td>4,400 ± 580</td>
</tr>
<tr>
<td></td>
<td>Ethene</td>
<td>3,400 ± 690</td>
</tr>
<tr>
<td></td>
<td>EtO</td>
<td>1,300 ± 80</td>
</tr>
<tr>
<td></td>
<td>LBB, growing on VC</td>
<td>250 ± 46</td>
</tr>
<tr>
<td></td>
<td>LBB, growing on ethene</td>
<td>190 ± 37</td>
</tr>
<tr>
<td></td>
<td>LBB, growing on EtO</td>
<td>120 ± 79</td>
</tr>
<tr>
<td>TD</td>
<td>VC</td>
<td>3,700 ± 700</td>
</tr>
<tr>
<td></td>
<td>Ethene</td>
<td>4,600 ± 700</td>
</tr>
<tr>
<td></td>
<td>EtO</td>
<td>3,500 ± 280</td>
</tr>
<tr>
<td></td>
<td>LBB, growing on VC</td>
<td>270 ± 89</td>
</tr>
<tr>
<td></td>
<td>LBB, growing on ethene</td>
<td>300 ± 40</td>
</tr>
<tr>
<td></td>
<td>LBB, growing on EtO</td>
<td>160 ± 16</td>
</tr>
<tr>
<td>No cells</td>
<td>None</td>
<td>25 ± 5</td>
</tr>
</tbody>
</table>

\(^a\) The values are means ± standard deviations of triplicate measurements.
Retention of the EaCoMT gene even after curing of the plasmids is consistent with the higher levels of EaCoMT activity in cells grown in LBB and exposed to 10 mM CoM than in cells not exposed to CoM, even after the abiotic activity with no thiol. Alternatively, plasmid DNA may have integrated into chromosomal DNA via insertion elements. Strain TD also showed the strongest hybridization signal that did not correspond to a plasmid (lane 9). Similar results were observed with VC-grown Mycobacterium sp. strain JS616 (5).

The sequences of the EaCoMT gene even after curing of the plasmids is consistent with the higher levels of EaCoMT activity in cells grown in LBB and exposed to 10 mM CoM than in cells not exposed to CoM, even after the abiotic activity with no thiol. Alternatively, plasmid DNA may have integrated into chromosomal DNA via insertion elements. Strain TD also showed the strongest hybridization signal that did not correspond to a plasmid (lane 9). Similar results were observed with VC-grown Mycobacterium sp. strain JS616 (5).

The sequences of the EaCoMT gene of VC-grown strains AJ and TD were compared to the sequences of various strains of Mycobacterium, Rhodococcus, Xanthobacter, and Nocardioides. Strains AJ and TD clustered with Mycobacterium strains JS60 and JS621 (see Fig. S1 in the supplemental material). The high degree of homology among EaCoMT sequences is interesting when how geographically diverse the microbes are is considered (4, 8), although there is considerable variability among EaCoMT rates (5, 6).

This study demonstrated that EaCoMT is used by P. putida strain AJ and Ochrobactrum sp. strain TD during metabolism of VC, ethene, and EtO, thereby extending the range of bacteria known to use CoM during catabolism of alkenes (1, 4, 5, 13–15).

Nucleotide sequence accession numbers. Sequences of the EaCoMT genes in strain AJ grown on VC (accession no. AY858984) and in LBB (accession no. DQ370435) and in strain TD grown on VC (accession no. AY858985) and in LBB (accession no. DQ370436) have been deposited in the GenBank database.

Jim C. Spain (Department of Civil & Environmental Engineering, Georgia Institute of Technology) generously provided Mycobacterium sp. strain JS60. William C. Bridges, Jr. (Department of Applied Economics and Statistics, Clemson University) provided assistance with the statistical analyses.

REFERENCES