Bacterial Acyl-CoA Mutase Specifically Catalyzes Coenzyme B<sub>12</sub>-dependent Isomerization of 2-Hydroxyisobutyryl-CoA and (S)-3-Hydroxybutyryl-CoA

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Background: Carbon skeleton rearrangements of acyl-CoA esters are catalyzed by coenzyme B<sub>12</sub>-dependent mutases. Results: A bacterial mutase specifically catalyzes the isomerization of 2-hydroxyisobutyryl- and (S)-3-hydroxybutyryl-CoA. Conclusion: Substrate affinity and enzyme activity depend strongly on the active site amino acid Ile<sup>346</sup>. Significance: This is the first characterization of an enzyme isomerizing hydroxylated short chain carboxylic acids.

Coenzyme B<sub>12</sub>-dependent acyl-CoA mutases are radical enzymes catalyzing reversible carbon skeleton rearrangements in carboxylic acids. Here, we describe 2-hydroxyisobutyryl-CoA mutase (HCM) found in the bacterium Aquincola tertiaricarbonis as a novel member of the mutase family. HCM specifically catalyzes the interconversion of 2-hydroxyisobutyryl- and (S)-3-hydroxybutyryl-CoA. Like isobutyryl-CoA mutase, HCM consists of a large substrate- and a small B<sub>12</sub>-binding subunit, HcmA and HcmB, respectively. However, it is thus far the only acyl-CoA mutase showing substrate specificity for hydroxylated carboxylic acids. Complete loss of 2-hydroxyisobutyric acid degrada
tion capacity in hcmA and hcmB knock-out mutants established the central role of HCM in A. tertiaricarbonis for degrading substrates bearing a tert-butyl moiety, such as the fuel oxygenate methyl tert-butyl ether (MTBE) and its metabolites. Sequence analysis revealed several HCM-like enzymes in other bacterial strains not related to MTBE degradation, indicating that HCM may also be involved in other pathways. In all strains, hcmA and hcmB are associated with genes encoding for a putative acyl-CoA synthetase and a MeaB-like chaperone. Activity and substrate specificity of wild-type enzyme and active site mutants HcmA 190V, 190F, and 190Y clearly demonstrated that HCM belongs to a new subfamily of B<sub>12</sub>-dependent acyl-CoA mutases.

The tertiary carbon-bearing 2-hydroxyisobutyric acid (2-HIBA)<sup>2</sup> is rarely found in Nature and is not an intermediate of the main metabolic pathways. However, it is one of the urinary organic acids found in humans with lactic acidosis (1), and several metabolic sequences can be proposed leading to this unusual short chain carboxylic acid (Fig. 1). A major natural source of 2-HIBA might be the plant cyanoglycoside linamarin (2), as the nitrile corresponding to 2-HIBA is an intermediate of linamarin biosynthesis and catabolism. In addition, 2-HIBA could be produced during degradation of isobutane and isobutene via oxidation of the corresponding aliphatic and diol metabolites. Besides, anthropogenic sources of 2-HIBA exist, because it is a pharmaceutical intermediate and by-product of industrial processes, e.g., the production of poly(methyl methacrylate) (PMMA) (3). It has also been identified as a metabolite in the bacterial degradation of the gasoline additive methyl tert-butyl ether (MTBE) (4), which is used at large scale since the 1990s as a fuel oxygenate for reducing carbon monoxide emissions and now threatens drinking water resources due to its persistence in contaminated aquifers (5). Although biodegradation of 2-HIBA and its methyl ester has already been observed in one early study in 1984 when investigating the bacterial degradation of wastewater compounds of a PMMA plant (6), convincing enzymatic steps for its conversion to common metabolites have not been proposed for >20 years.

Recently, we found a coenzyme B<sub>12</sub>-dependent rearrangement reaction in the bacterial degradation pathway of MTBE via 2-HIBA, likely catalyzed by a novel acyl-CoA mutase (7). In this enzymatic step, the CoA ester of 2-HIBA is converted to 3-hydroxybutyryl-CoA (Fig. 1). Thus, in a single reaction the branched-chain α-hydroxy carboxylic acid is rearranged into an easily metabolizable linear isomer, now possessing an oxidizable secondary hydroxyl group at the β position. Consequently, the proposed 2-hydroxyisobutyryl-CoA mutase (HCM) would play a central role in MTBE metabolism enabling its complete mineralization. Similar rearrangements in carboxylic acids have already been observed with several acyl-CoA mutases (8, 9, 10), namely methylmalonoyl-CoA mutase (MCM), isobutyryl-CoA mutase (ICM) and ethylmalonyl-CoA mutase (ECM). Recently, a variant of ICM (ICMF) has been characterized as a fusion of the ICM sequence with the G protein chap-
erone Meal (11), a paralog of the MCM-associated MeaB protein. However, HCM would be distinct from all other known acyl-CoA mutases because it would catalyze the conversion of hydroxylated carboxylic acids.

This study aimed at elucidating the biochemistry of the acyl-CoA mutase activity from the MTBE-degrading bacterium *Aquincola tertiaricarbonis* L108. It was found that HCM consists of a large acyl-CoA-binding and a small B12-binding subunit, HcmA and HcmB, respectively. Destroying the corresponding genes *hcmA* and *hcmB* by insertional mutation resulted in complete loss of 2-HIBA-degrading capability. The wild-type mutase genes were cloned in *Escherichia coli* strains, and enzyme activity of heterologously expressed subunits was characterized. Catalysis by purified recombinant HCM was specific for 2-hydroxyisobutyryl- and (S)-3-hydroxybutyryl-CoA. Sequence comparison with known acyl-CoA mutases identified a single active site amino acid residue present in HcmA likely being important for determining substrate specificity. To understand the function of this residue, we analyzed activities of three mutant HcmA subunits having relevant amino acid substitutions.

### EXPERIMENTAL PROCEDURES

#### Materials

Tert-butyl alcohol (TBA) (99%), 2-HIBA (>98%), tert-amyl alcohol (>99%), and anhydrides of butyric (98%) and isobutyric (98%) acids were purchased from Merck Schuchardt. 2-Methylpropan-1,2-diol was from Taros Chemicals at the highest purity available. Coenzyme B12 (≥97%), CoA (≥93%), methylmalonyl-CoA (≥96%, racemic mixture), and succinyl-CoA (≥94%) were purchased from Sigma. Enantiopure (R)- and (S)-3-hydroxybutyryl-CoA were provided by Evonik Industries (53–54% acyl-CoA enantiomer, >99.9 and 98.9% enantiomeric excess, respectively; 40% NaCl; 6–7% N-hydroxysuccinimide).

#### Syntheses

**Synthesis of Acyl-CoA Esters**—Acyl-CoA esters not commercially available were synthesized according to two methods. Isobutyryl-CoA and butyryl-CoA were prepared from their anhydrides (12). 2-Hydroxyisobutyryl-CoA was synthesized from its free acid via thiophenyl ester by the method of Padmakan et al. (13).
2-Hydroxyisobutyryl-CoA Mutase

Identification of Synthesis Products—Identity of synthesized acyl-CoA esters was verified by electrospray ionization-tandem mass spectrometry (ESI-MS/MS) using a 4000 QTRAP LC/MS/MS system consisting of a liquid chromatograph and a high performance triple quadrupole/linear ion trap mass spectrometer (Applied Biosystems). Acyl-CoA esters were detected directly by ESI-MS/MS. Isocratic elution was achieved using a mobile phase containing 17.5 mM ammonium acetate, 0.5 volume % acetic acid, and 30 volume % acetonitrile (14). Characteristic mass spectra for the respective acyl-CoA compounds were obtained (supplemental Fig. S1).

Bacterial Strains and Growth Conditions

Strain A. tertiaricarbonis L108 isolated from an MTBE-contaminated aquifer at Leuna, Germany (7, 15) was cultivated in liquid mineral salt medium (MSM) containing MTBE at concentrations of 0.3 g liter⁻¹ as described previously (16). For growth and resting-cell studies, also tert-amyl alcohol, TBA, 2-methylpropan-1,2-diol, and 2-HIBA were supplied as sole source of carbon and energy at 0.5 g liter⁻¹. E. coli TOP10 and ArcticExpress (DE3) were grown in Luria-Bertani broth. Growth was monitored by measuring optical density (OD) of cultures at 700 or 550 nm as indicated.

Sequencing of Genomic DNA from A. tertiaricarbonis

For sequencing a larger fragment of the hcm gene region of wild-type strain L108, genomic DNA was extracted using MasterPure DNA Purification kit (Epicenter) and sequenced by Illumina HiSeq 2000 technology (GATC Biotech). The obtained DNA sequences were analyzed for open reading frames using Rast (Rapid Annotation using Subsystem Technology). A 4.5 kb sequence including both hcm genes was obtained.

Cloning and Heterologous Expression of hcmA and hcmB from A. tertiaricarbonis in E. coli

The gene encoding for the large subunit of HCM (hcmA) was amplified from strain L108 genomic DNA by applying the forward primer 5’-AATG ACC TGG CTT GAG CCG CAG A-3’ and reverse primer 5’-TCCC GAA GAC CGG GTC TCG CGG-3’. PCR was accomplished with Pfu DNA polymerase (Promega) for 30 cycles, including denaturation at 94 °C for 1 min, annealing at 57 °C for 1 min, and extension at 72 °C for 2 min. The PCR product was cloned into expression vector pASG-IBA43 (IBA Goettingen) and transformed into E. coli TOP10 according to the protocol of IBA Goettingen. Induction was performed at OD₅₅₀ of 0.5 with 200 μg liter⁻¹ anhydrotetracycline for 3 h at 30 °C. Cells were centrifuged and suspended in Tris buffer (100 mM Tris, 150 mM NaCl, pH 8.0) for further analysis. The cloning vector pPR-IBA1::hcmB for the small subunit of HCM was purchased from DNA2.0 and transformed into 100 μl of electrocompetent E. coli ArcticExpress (DE3) (Novagen) at 300 mV for 5 ms in chilled 0.1-cm cuvettes in a MicroPulser (Bio-Rad). Induction was performed with 0.5 mM isopropyl 1-thio-β-D-galactopyranoside for 20 h at 12 °C, after growth at 30 °C in Luria-Bertani medium containing 20 mg liter⁻¹ gentamycin and 50 mg liter⁻¹ kanamycin to an OD₅₅₀ of 0.4 and further incubation at 12 °C to an OD₅₅₀ of 0.5. Then, cells were centrifuged and suspended in Tris buffer.

Purification of Recombinant Proteins

For the purification of recombinant mutase subunits, crude extracts of induced E. coli cells were prepared by disruption using a mixer mill (MM 400, Retsch GmbH, Germany) with glass beads (212–300 μm, Sigma) at 30 s⁻¹ for 30 min. The recombinant HcmA and HcmB subunits were purified with the help of their His and Strep tags, respectively. All purification steps were performed at 12 °C.

HcmA Subunit—Crude extracts of E. coli TOP10 pASG-IBA43::hcmA were loaded on a nickel-nitrilotriacetic acid Superflow 10-ml column (IBA Goettingen). After washing with 20 column volumes of imidazole buffer (50 mM sodium phosphate, 300 mM NaCl, 20 mM imidazole, pH 8.0), HcmA was eluted with the same buffer containing 250 mM imidazole. Fractions containing HcmA were concentrated via viva spin columns (30 kDa; GE Healthcare) and diluted with conservation buffer (50 mM potassium phosphate, 10% glycerol, pH 7.4).

HcmB Subunit—Crude extracts of E. coli ArcticExpress (DE3) pPR-IBA1::hcmB were loaded on a Strep-Tactin Superflow high capacity 10-ml column (IBA Goettingen). After washing with 20 column volumes of Tris buffer, HcmB was eluted with elution buffer (Tris buffer containing 2.5 mM desthiobiotin). Fractions containing HcmB were concentrated using viva spin columns (10 kDa) and diluted with conservation buffer.

Expression and Purification of Site-specific HcmA Mutants

The hcmA mutant genes cloned into pASG-IBA43 were purchased from GeneCust Europe. Site-directed mutagenesis resulted in HcmA I90V, I90F, and I90V mutants (with the point mutations a268t plus t269a, a268t, and a268g, respectively). The vectors were transformed into E. coli TOP10. Expression and purification were performed as described for the wild-type recombinant HcmA.

Quantitative Enzymatic Measurements of Recombinant Enzyme

HCM Activity—Enzyme activity was routinely measured in 1–2 ml of 50 mM potassium phosphate buffer, pH 6.6, containing 10% glycerol, 833 μM coenzyme B₁₂, and 10 mM MgCl₂. The reaction was started by adding acyl-CoA substrates. As variation of HcmA and HcmB ratios (up to 5:1 and 1:5, respectively) did not result in increased activities, the recombinant subunits were added at equimolar ratios (3 μM) throughout the study. After a 5-min preincubation in the presence of coenzyme B₁₂, the reaction was started by adding acyl-CoA substrates. Throughout the experiments, oxygen concentrations were minimized by permanently flushing the incubation vials with nitrogen. For determination of pH optimum, pH values of the phosphate buffer and a phosphate/acetate buffer (phosphate buffer plus 50 mM sodium acetate buffer) were adjusted to values between 5.0 and 7.8. The temperature optimum was determined by incubating at temperatures between 20 and 55 °C.

HPLC Analysis of Acyl-CoA Esters—Concentrations of acyl-CoA ester substrates and products were quantified by ion-pair
chromatography using an HPLC system (Shimadzu) with a Nucleosil 100–5 C18 column (250 mm × 3 mm, 5 μm; Macherey-Nagel) and a mobile phase of 14.5 vol% acetonitrile, 10 mM tetrabutylammonium hydrogen sulfate and 100 mM sodium phosphate at pH 4.5 for the separation of 2-hydroxyisobutyryl-, 3-hydroxybutyryl-, methylmalonyl-, and succinyl-CoA, with retention times of about 35, 24, 30, and 40 min, respectively. For separation of isobutyryl- and butyryl-CoA, acetonitrile content was adjusted to 0.6 ml of the eluent was 21.6 vol%, resulting in retention times of 35 and 40 min, respectively (17). Eluent flow was adjusted to 0.6 ml min⁻¹, and column oven temperature was 30 °C. Absorbance at 260 nm was used for quantifying acyl-CoA esters. In addition, spectra from 190 to 280 nm were recorded for distinguishing peaks of CoA esters from unspecific signals (18). Detection limit at 260 nm was 0.5 μM CoA ester. For stopping enzymatic degradation, we generated knock-out mutants of strain A. terriericarbonis L108 by electroporation of 1 μl of EZ-Tn5<ΔKAN-2>Tp transposome (Epicerenter Biotechnologies) to 70 μl of electrocompetent L108 cells in chilled cuvettes at 1.8 kV for 1 ms (MicroPulser; Bio-Rad). Transformed cells were rescued in 5 ml of MSM amended with 10 mM fructose for 6 h at 30 °C (MicroPulser; Bio-Rad). Transformed cells were rescued in 5 ml of MSM amended with 10 mM fructose for 6 h at 30 °C and 150 rpm. Dilutions were plated on MSM fructose agar containing 50 μg ml⁻¹ kanamycin and incubated for 2 days at 30 °C. As the transposome integrates randomly into the DNA, all colonies obtained had to be analyzed for loss of their capability to grow on MSM agar containing 0.5 g liter⁻¹ TBA. In addition, copies of the colonies were maintained on MSM fructose agar for further analysis. Colonies with restricted or even lost TBA degradation potential were transferred on MSM 2-HIBA agar plates. The mutants that failed to grow were further analyzed.

The exact integration site of the kanamycin cassette into the genomic DNA was determined by direct DNA sequencing using flanking KAN-2 primers (Epicerenter Biotechnologies) of 1 mg liter⁻¹ high concentrated genomic DNA (MasterPure DNA Purification kit). Conditions were 4 min initial denaturation at 95 °C followed by 60 cycles of 30 s at 95 °C and 4 min 60 °C, finally cooled down to 8 °C. The products were cleaned with Centri-Sep columns (Applied Biosystems) and sequenced using an ABI PRISM 3100 Genetic Analyzer with the BigDye Terminator v1.1 Cycle sequencing kit (Applied Biosystems). The resulting sequences were analyzed with BLAST (19) and aligned via Sequencher 5.0 software (Gene Codes Corporation).

**RESULTS**

**Insertional Inactivation of hcmA and hcmB, the Putative Genes Encoding for HCM Large and Small Subunits, and Characterization of Mutant Strains**—Sequencing of genomic DNA of wild-type A. terriericarbonis L108 revealed that a 4.5-kb fragment comprised both hcmA and hcmB separated by two genes encoding for a putative acyl-CoA synthetase and a MeaB-like G protein chaperone, respectively (Fig. 2). A highly similar sequence (>97% identity) having the same gene organization has already been found in the genome of the MTBE-degrading strain Methylibium petroleiphilum PM1 (20), supporting the hypothesis that HCM is involved in degradation of the MTBE metabolite 2-HIBA (7). However, the organization of hcm-like genes interrupted by genes likely encoding for a acyl-CoA synthetase and MeaB-like chaperone is also present in other bacteria, such as *Rhodobacter sphaeroides* ATCC 17029 and *Nocardioides* sp. JS614 (Fig. 2), thus far not related to fuel oxygenate ether degradation. As has already been suggested on the basis of transcriptome analysis of MTBE-grown cells of strain PM1 (21), the acyl-CoA synthetase could be involved in CoA activation of 2-HIBA (Fig. 1). The G protein chaperone may play a role in HCM assembly and stabilization, as has already been shown for MeaB and MeaI associated with MCM and ICM/IcmF, respectively (11, 22). Therefore, we propose for the MeaB/MeaI paralog associated with HCM the name MeaH (Fig. 2).

For precisely studying the role of HCM in bacterial 2-HIBA degradation, we tried to create hcmA and hcmB knock-out mutants of strain L108 by a site-specific mutagenesis approach via homologous recombination. However, even after several trials corresponding knock-out mutants were not obtained. On the other hand, unspecific transposon-mediated mutagenesis...
and screening for loss of capability to grow on 2-HIBA resulted in two hcm mutants, L108(ΔhcmA)K5 and L108(ΔhcmB)K7, bearing stable insertions of a functionally active kanamycin resistance gene in the hcmA and hcmB genes, respectively (Fig. 2). In contrast to the wild-type L108 strain, both mutant strains were not able to grow on TBA and 2-methylpropan-1,2-diol, precursors of 2-HIBA in the tert-butyl alkyl ether degradation pathway (Fig. 1), establishing the postulated central role of HCM in the degradation of organic compounds possessing the tert-butyl moiety (7). Accordingly, TBA and 2-methylpropan-1,2-diol were stoichiometrically converted to 2-HIBA by both knock-out mutants in resting-cell experiments (Fig. 3), indicating that only the 2-HIBA-metabolizing enzymatic activity was affected by the hcm knockouts. Contrarily, the mutants still grew well on tert-amyl alcohol, suggesting an alternative degradation pathway for this C5 homolog of TBA (23).

Expression of hcm Genes in E. coli and Purification of HCM Subunits—For biochemical characterization, wild-type hcmA and hcmB as well as hcmA mutant genes were heterologously expressed in E. coli. N-terminal His-tagged and C-terminal Strep-tagged proteins, respectively, were purified from crude extracts by one-step affinity chromatography. Denaturing gel electrophoresis established the presence of the mutase subunits in crude extracts of induced E. coli cells and successful purification of the tagged proteins (supplemental Fig. S2). Yields of purified HcmA (HcmA Ile90 and HcmA mutants) and HcmB were about 100 and 80 mg liter−1 of E. coli culture, respectively.

Characterization of Reconstituted Hcm Enzyme Activity—As has already been shown for the small and large ICM subunits (9), neither recombinant HcmA nor HcmB alone was able to catalyze an acyl-CoA mutase reaction. However, a combination of both purified subunits resulted in an enzymatic rearrangement of 2-hydroxyisobutyryl- into 3-hydroxybutyryl-CoA (Fig. 4). Under the experimental conditions applied, a linear increase in 3-hydroxybutyryl-CoA for about 10 min could be achieved, thus, enabling determination of turnover rates by tracing changes in substrate and product concentrations with HPLC (supplemental Fig. S3). Within a pH range of 5–8, significant enzymatic conversion of 2-hydroxyisobutyryl-CoA was observed, having a maximal activity at pH 6.6 (supplemental Fig. S4). The temperature optimum of the mutase activity was in the mesophilic range at about 30 °C. At 20 °C, however, still 50% of the maximal activity was achieved, whereas a complete loss of catalysis was shown at 45 °C (supplemental Fig. S5), which corresponds well to the temperature spectrum described for growth of wild-type strain L108 on TBA (24). Interestingly, as demonstrated by analyzing the 3-hydroxybutyryl-CoA esters produced, after derivatization to propyl esters and enantioselective separation on a chiral GC column (supplemental Fig. S6), 2-hydroxyisobutyryl-CoA was predominantly converted to (S)-3-hydroxybutyryl-CoA, and only minor amounts of about 20% of the (R)-enantiomer were obtained, indicating stereospecific catalysis.

Kinetic Parameters of HCM Catalysis—Possible enzymatic rearrangement of substrates other than 2-hydroxyisobutyryl-CoA was tested at optimal pH and temperature conditions (Table 1 and Fig. 5). As expected, purified wild-type HCM subunits were able to convert (S)-3-hydroxybutyryl-CoA at high
TABLE 1
Kinetic parameters of reconstituted HCM subunits HcmA (wild-type Ile90 and mutant I90V) and wild-type HcmB incubated with various acyl-CoA substrates at pH 6.6 and 30 °C (Fig. 5).

| Subunit and substrate | $V_{max}^{a}$ | $K_m$ | $k_{cat}$ | $k_{cat}/K_m$ | $V_{max}/K_m$ |
|-----------------------|---------------|-------|-----------|---------------|---------------|
|                       | mmol min⁻¹ mg⁻¹ | μM    | min⁻¹     | liters min⁻¹ mg⁻¹ |
| **HcmA Ile90**        |               |       |           |               |
| (S)-3-Hydroxybutyryl-CoA | 140 ± 6.6 (100)$^b$ | 128 ± 22 | 12 ± 0.5 | 90 ± 16 | 1,080 ± 190 |
| 2-Hydroxyisobutyryl-CoA | 36 ± 2.8 (26)  | 104 ± 25 | 3.0 ± 0.2 | 29 ± 7.3 | 350 ± 87  |
| (R)-3-Hydroxybutyryl-CoA | 2.4 ± 0.15 (2) | 1,660 ± 367 | 0.20 ± 0.01 | 0.12 ± 0.03 | 0.4 ± 0.3 |
| Butyryl-CoA            | 2.4 ± 0.26 (2) | 3,540 ± 880 | 0.20 ± 0.02 | 0.06 ± 0.02 | 0.7 ± 0.2 |
| Isobutyryl-CoA         | 0.67 ± 0.03 (0.5) | 550 ± 140 | 0.06 ± 0.003 | 0.10 ± 0.03 | 1.2 ± 0.3 |
| **HcmA I90V**         |               |       |           |               |
| (S)-3-Hydroxybutyryl-CoA | 24 ± 1.1 (17)  | 1,760 ± 240 | 2.0 ± 0.1 | 1.1 ± 0.16 | 14 ± 2.0  |
| 2-Hydroxyisobutyryl-CoA | 15 ± 1.1 (11)  | 1,840 ± 370 | 1.2 ± 0.1 | 0.7 ± 0.14 | 8.1 ± 1.7 |

$^a$ Activities obtained with the wild-type HcmAB for methylmalonyl- and succinyl-CoA and with the HcmA I90V mutant reconstituted with wild-type HcmB for all acyl-CoA esters tested, except for (S)-3-hydroxybutyryl- and 2-hydroxyisobutyryl-CoA, were below detection limit (<0.01 nmol min⁻¹ mg⁻¹).

$^b$ For calculating relative activities, $V_{max}$ obtained with wild-type HcmAB for (S)-3-hydroxyisobutyryl-CoA was set to 100.

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Retention of Configuration during HCM Catalysis—As already found with MCM and ECM, only converting (R)-methylmalonyl- and (R)-ethylmalonyl-CoA (10, 27), respectively, HCM catalysis is stereospecific, favoring the rearrangement of (S)-3-hydroxybutyryl-CoA versus the (R)-enantiomer. This specificity of catalysis may also shed light on the orientation of the hydroxycarboxylic acid at the catalytic site of HCM. The rearrangement by MCM and ECM proceeds strictly with retention of configuration (Fig. 8). In the case of ICM, although its natural substrates isobutyryl- and butyryl-CoA do not possess chirality, catalysis is also predominantly stereospecific with retention of configuration, as has been revealed by testing labeled substrates in vivo (28) and with partially purified enzyme (29). Now, due to the structural similarities between the 2-hydroxyisobutyryl and isobutyryl substrate residues, in HCM an ICM-like orientation of the two methyl groups of the branched-chain carboxylic acid could be expected (30) which would result in catalysis predominantly toward (R)-3-hydroxybutyryl-CoA (Fig. 8). However, considering the observed stereoselectivity of HCM catalysis favoring the (S)-enantiomer of 3-hydroxybutyryl-CoA, it can be concluded that in HCM the hydroxyacyl residue of its substrates is mainly oriented in the same way as the carboxyacyl substrate moiety in MCM and ECM. In HCM, the polar but uncharged hydroxyl group would specifically interact with the amido function of Gln208 (numbering as in HcmA of strain L108), whereas in MCM/ECM the negatively charged carboxyl group is close to the guanidinium group of the HCM acyl substrates. However, the ICM-like orientation of the methyl groups of the 2-hydroxyisobutyryl residue is not completely excluded, but takes place in about 2% of substrate binding events, as can be deduced from the transformation rates of (R)-3-hydroxybutyryl-CoA. This deviation from complete stereospecificity may also explain the low but significant conversion of the ICM substrates isobutyryl- and butyryl-CoA (Table 1), as it can be assumed that these substrates and (R)-3-hydroxybutyryl-CoA have a similar orientation in the substrate binding site of HCM (Fig. 8). Both HCM and ICM/IcmF share the uncharged Gln residue (Gln198 and Gln733 in Fig. 8), suggesting that this amino acid is not specifically interacting with the substrate in HCM due to the replacement of the aromatic Phe80/Phe589 postulated to be specifically interacting with the substrate in ICM/IcmF (9, 11) with the smaller aliphatic Ile90 (Fig. 8). Accordingly, the conservative substitution I90V still possess activity toward (S)-3-hydroxybutyryl- and 2-hydroxyisobutyryl-CoA and (S)-3-hydroxybutyryl-CoA (A) and (R)-3-hydroxybutyryl-, butyryl-, and isobutyryl-CoA by wild-type HcmAB (B). C, conversion of 2-hydroxyisobutyryl- and (S)-3-hydroxybutyryl-CoA by HcmA I90V mutant reconstituted with the wild-type HcmB. All enzyme activities were measured at pH 6.6 and 30 °C. By using nonlinear regression analysis (Graph Pad Prism 5.0 software), $K_m$ and $V_{max}$ values were revealed (Table 1), applying either the Michaelis-Menten equation (A and B) or an allosteric sigmoidal model (C).

FIGURE 5. Kinetic plots of acyl-CoA rearrangement activities catalyzed by reconstituted HcmA and HcmB. A and B, conversion of 2-hydroxyisobutyryl- and (S)-3-hydroxybutyryl-CoA by wild-type HcmB. All enzyme activities were measured at pH 6.6 and 30 °C. By using nonlinear regression analysis (Graph Pad Prism 5.0 software), $K_m$ and $V_{max}$ values were revealed (Table 1), applying either the Michaelis-Menten equation (A and B) or an allosteric sigmoidal model (C).
tyrlyl-CoA, whereas the nonconservative mutations I90F and I90Y completely lost rearrangement activity.

In contrast to the observed deviation in the predominant substrate orientation at the catalytic sites, HCM is phylogenetically close to bacterial ICM, together forming a cluster within archaeal MCM sequences (10). In addition, HCM and ICM share the same subunit organization consisting of a small B12- and acyl-CoA binding domains encoded by two distinct genes, whereas in bacterial MCM and ECM the B12- and acyl-CoA binding domains are both located on the same subunit (8, 10). The high sequence and structural similarity between HCM and ICM may allow studying substrate binding and catalysis of specific mutant enzymes. Comparable single amino acid mutations in ICM and MCM resulted mainly in loss of enzymatic activity (31), possibly due to the larger sequence and structural deviations between these mutase subfamilies. However, the activities of the amino acid mutants analyzed in this study clearly show that even in HCM single active site residue substitutions are not sufficient to change substrate specificity, e.g. achieving ICM activity with an HcmA I90F mutant.

Occurrence and Evolution of HCM—As already outlined, the HCM enzyme is, besides ICM, the second representative within the B12-dependent acyl-CoA mutase subfamilies thus far identified to be organized as small and large subunits, binding the coenzyme B12 and the acyl-CoA ester substrate, respectively. In addition, HCM activity is different, as this enzyme specifically catalyzes the rearrangement of hydroxylated carboxylic acids which has not been observed with other mutases. However, the origin of HCM is enigmatic. BLAST analysis revealed a quite heterogeneous group of bacteria, including phylogenetically distant proteobacteria and a Gram-positive strain, which all possess the four genes encoding for the HCM subunits plus a putative acyl-CoA synthetase and the chaperone MeaH organized in an operon-like structure (Figs. 2 and 7). The high sequence similarity and the identical gene grouping suggest an interchange of the complete hcm operon by horizontal gene transfer. Unlike strains A. tertiaricarbonis L108 and M. petroleiphilum PM1 (15, 32), the other bacterial strains with an hcm operon have not been associated with the degradation of fuel oxygenates MTBE and TBA. An alternative source for compounds bearing the tert-butyl group is the industrial production of PMMA (3, 6), as the corresponding wastewater contains 2-HIBA and its methyl ester. Accordingly, it has already been demonstrated that strains R. sphaeroides ATCC 17029, Xanthobacter autotrophicus Py2 and Nocardioides sp. JS614 are
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able to grow on 2-HIBA (33). However, worldwide production of PMMA at large scale started not earlier than in the late 1930s (34), whereas the hcm operon bearing strain Starkeya novella DSM 506 has already been isolated from uncontaminated agricultural soil in 1934 (35), indicating that other drivers than this kind of anthropogenic contamination for the evolution of HCM may exist. The other known B12-dependent acyl-CoA mutases are widely distributed among bacteria playing central roles in primary and secondary carbon metabolism, e.g. in branched-chain amino acid catabolism (8) and in the recently discovered ethylmalonyl-CoA pathway for acetate assimilation (10) as well as in the synthesis of macroclide and polyether antibiotics (9). In contrast, HCM would be the first mutase exclusively employed for the dissipilatory degradation of a single substrate, i.e. 2-HIBA. In this connection, several not yet identified mutases have been postulated to be involved in bacterial degradation pathways for the mineralization of alkanes, ethylene, and the quaternary carbon-bearing pivalic acid (36, 37). Hence, it could be speculated that all of these mutases employed in dissimilatory pathways may have the same origin being adapted to their specific substrates by a moderate variation of the amino acid residues in the substrate binding site. The previously described close phylogenetic relationship of bacterial HCM and ICM sequences with archaeal mutases (10) is also interesting, as it suggests an origin of HCM outside the domain Bacteria.

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2-Hydroxyisobutyryl-CoA Mutase

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