Phenylacetic Acid Catabolism and Its Transcriptional Regulation in Corynbacterium glutamicum

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The industrially important organism Corynbacterium glutamicum has been characterized in recent years for its robust ability to assimilate aromatic compounds. In this study, C. glutamicum strain AS 1.542 was investigated for its ability to catabolize phenylacetic acid (PAA). The paa genes were identified; they are organized as a continuous paa gene cluster. The type strain of C. glutamicum, ATCC 13032, is not able to catabolize PAA, but the recombinant strain ATCC 13032/pEC-K18 mob2::paa gained the ability to grow on PAA. The paaR gene, encoding a TetR family transcription regulator, was studied in detail. Disruption of paaR in strain AS 1.542 resulted in transcriptional increases of all paa genes. Transcription start sites and putative promoter regions were determined. An imperfect palindromic motif (5’-ACTNACGNCCGNNCGG-3’; 22 bp) identified in the upstream regions of paa genes. Electrophoretic mobility shift assays (EMSA) demonstrated specific binding of PaaR to this motif, and phenylacetyl coenzyme A (PA-CoA) blocked binding. It was concluded that PaaR is the negative regulator of PAA degradation and that PA-CoA is the PaaR effector. In addition, GlxR binding sites were demonstrated specific binding of PaaR to this motif, and phenylacetyl coenzyme A (PA-CoA) blocked binding. It was concluded that PaaR is the negative regulator of PAA degradation and that PA-CoA is the PaaR effector. In addition, GlxR binding sites were found, and binding to GlxR was confirmed. Therefore, PAA catabolism in C. glutamicum is regulated by the pathway-specific repressor PaaR, and also likely by the global transcription regulator GlxR. By comparative genomic analysis, we reconstructed orthologous PaaR regulons in 57 species, including species of Actinobacteria, Proteobacteria, and Flavobacteria, that carry PAA utilization genes and operate by conserved binding motifs, suggesting that PaaR-like regulation might commonly exist in these bacteria.

Phenylacetic acid (PAA) is an important intermediate during the microbial degradation of aromatic compounds such as styrene, ethylbenzene, 2-phenylethylamine, tropic acid, phenylacetaldehyde, phenylacetyl esters, and amides (26), and PAA degradation pathways have been detected in a wide range of bacterial species (7, 10, 27, 30, 31). The initial reaction of PAA degradation is the activation of PAA to phenylacetyl coenzyme A (PA-CoA). Subsequently, PA-CoA is degraded via the function of a PA-CoA catabolon core (26) (see Fig. 1). Recently, it was discovered that PA-CoA is epoxidized and isomerized to form a seven-membered C-O-heterocyclic enol ether (oxepin-CoA), followed by hydrolytic ring cleavage and β-oxidation-like steps yielding tricarboxylic acid cycle intermediates (45, 46).

Genes encoding the PAA degradation pathways are diversely organized in bacteria (26). In Pseudomonas putida strain U, 15 paa genes are organized in five contiguous operons and are grouped in five functional units, including PAA transport, PAA activation, ring hydroxylation and cleavage, β-oxidation-like catalysis, and regulation (26, 28). In Escherichia coli, 14 paa genes are organized in three transcription units that function in PAA activation, ring hydroxylation and cleavage, β-oxidation, and regulation (10, 26). The regulation of the paa genes has been investigated in several bacterial species. In E. coli and Pseudomonas species, a GntR family transcriptional regulator named PaaX represses the transcription of the paa genes (7, 9). In Burkholderia cepacia K56-2 and Thermus thermophilus HB8, a transcriptional regulator of the TetR family, named PaaR, has been reported as a repressor of the PAA degradation pathway (13, 38). Instead of PAA, PA-CoA acts as the effector for all the regulatory proteins identified so far (9, 11, 38, 48).

C. glutamicum is a well-studied, high-GC-content, Gram-positive soil bacterium of industrial importance. The ability of C. glutamicum to metabolize a variety of aromatic compounds has been studied for its type strain, ATCC 13032 (5, 15, 41–43, 49). However, no degradation of PAA by C. glutamicum was reported. In this study, we investigated the abilities of various strains of C. glutamicum to degrade PAA. The results showed that strain ATCC 13032 is not able to utilize PAA but that a serine-auxotrophic strain, AS 1.542 (formerly Corynbacterium crenatum), isolated from winery soil in southeast China (6), is able to grow on PAA. The paa gene cluster in strain AS 1.542 was functionally characterized, and its regulation by a TetR-type PaaR protein was studied.

MATERIALS AND METHODS

Bacterial strains, growth conditions, primers, and plasmids. The bacterial strains, primers, and plasmids used in this study are listed in Table S1.
in the supplemental material. C. glutamicum strains were grown at 30°C in minimal medium (20) with 5 mM glucose or PAA as the sole carbon source. For strain AS 1.542, minimal medium was supplemented with 1 mM serine. E. coli was grown at 37°C in Luria-Bertani (LB) broth or on LB agar plates (39). When needed, antibiotics were added at the following concentrations: 50 μg/ml kanamycin or 200 μg/ml ampicillin for E. coli and 25 μg/ml kanamycin for C. glutamicum. Bacterial growth was monitored by measuring the optical density at 600 nm (OD_{600}) or by using a Nephelostar Galaxy instrument (BMG Laboratories) (37).

**DNA extraction and manipulation.** C. glutamicum chromosomal DNA was isolated with the GenElute kit (Sigma-Aldrich). E. coli plasmid isolation and transformation, DNA manipulation, ligation, and agarose gel electrophoresis were carried out using standard protocols (39). Transformation of C. glutamicum with plasmids was performed according to the method of Taux et al. (44).

**Identification and annotation of the paa gene cluster.** The genome sequence of strain AS 1.542 was determined with the GS FLX system (45 Life Sciences), and the paa gene cluster was identified after the initial sequence assembly. Annotation and functional assignment were carried out with GenDB software (29) and according to similarity to the paa genes of E. coli K-12 (45) using the NCBI BLAST interface (17).

**Disruption of paaR.** paaR was disrupted by the gene splicing by overlap extension (GeneSOEing) method (14), removing a DNA fragment overlapping the putative helix-turn-helix (HTTH) domain (amino acid residues 8 to 102) of PaaR. The primers used for gene SOE are listed in Table S1 in the supplemental material. The disrupted paaR gene (paaR_{ΔHTTH}) was ligated into EcoRI/BamHI-treated pK18mob, generating pK18mob:paaR_{ΔHTTH}. This plasmid was used to perform an allelic exchange of paaR in the chromosome of strain AS 1.542, according to the work of Schafer et al. (40). An AS 1.542/paaR_{ΔHTTH} mutant was obtained, and the disruption of paaR was confirmed by PCR amplification.

**Introduction of the paa gene cluster into C. glutamicum strain ATCC 13032.** One-step isothermal DNA assembly (12) was used for the construction of plasmid pEC-K18mob2::paa. Primers (see Table S1 in the supplemental material) were designed to allow for a 40- or 41-bp overlap between the ends of the pEC-K18mob2 plasmid (22) and the amplified paa gene cluster. PCR products from the plasmid and the paa gene cluster were first purified and then treated with a DNA-blunting enzyme (CloneJET PCR cloning kit; Fermentas). After one-step isothermal DNA assembly, the assembled pEC-K18mob2::paa plasmid was isolated and introduced into C. glutamicum ATCC 13032.

**RT-qPCR measurements of transcription levels.** For the preparation of total RNA, bacterial cultures were grown in minimal medium with either 5 mM glucose or PAA as the sole carbon source, and cells from a 10-ml culture at exponential phase (OD_{600} = 1.0) were harvested by centrifugation at 14,000 rpm for 1 min. The supernatant was discarded, and the pellet was immediately frozen in liquid nitrogen. Subsequent total-RNA isolation and purification were performed using the RNeasy Mini kit (Qiagen) along with the DNeasy kit (Roche) and the RNeasy-free DNease set (Qiagen) (16). Reverse transcription-quantitative PCR (RT-qPCR) was carried out using the SensiMix SYBR one-step kit (Quantece) and the Opticon Monitor system (Bio-Rad). The specificity of RT-qPCR products was verified by gel electrophoresis and melting curve analysis. For all conditions, measurements were taken for a total amount of 300 ng RNA with two biological duplicates and two technical replicates for each sample. The difference in the cycle threshold between each strain and the wild-type strain AS 1.542 grown in 5 mM glucose was expressed as a ratio calculated by 2^{-ΔC_{P}}, where ΔCP is the difference between the measured crossing points (CP).

**Identification of transcription start sites (TSS) and promoter regions by RNAseq.** RNAseq. RNA was pooled from equal amounts of RNAs from strain AS 1.542 grown on glucose, PAA, and LB broth. A 5’-end-enriched RNAseq library was constructed according to the following procedures. (i) Depletion of stable tRNA and enrichment of mRNA molecules were performed using the Ribo-Zero RNA removal kit for Gram-positive bacteria (Epipicentre Biotechnologies). (ii) The enriched mRNA was fragmented by magnesium potassium acetate (MgKOAc) hydrolysis. Four volumes of RNA solution was mixed with 1 volume of MgKOAc solution (100 mM KOAC and 30 mM MgOAc in 200 mM Tris-HCl [pH 8.1]), and the mixture was incubated for 2.5 min at 94°C. The reaction was stopped by adding an equal volume of 1× TE (10 mM Tris, 1 mM EDTA [pH 8]) and chilling on ice for 5 min. (iii) The fragmented RNA was precipitated by the addition of 3 volumes 0.3 M NaAc in ethanol with 2 μl glycygen and overnight incubation at −20°C. (iv) The precipitated RNA fragments were dissolved in water, and the 5’-end RNA fragments were enriched by using Terminator 5’-phosphate-dependent exonuclease (Epipicentre Biotechnologies). (v) After RNA precipitation (see step iii above), the triphosphates were removed using RNA 5’-end phosphatase (Epipicentre Biotechnologies). (vi) After RNA precipitation (see step iii above), the 5’-enriched, monophosphorylated RNA fragments were used to construct a cDNA library by using the Small RNA Sample Prep kit (Illumina).

The fragmentation of RNA molecules (fragment sizes, 200 to 500 bp) and the RNA concentration were measured using the RNA 6000 Pico assay on an Agilent 2100 bioanalyzer (Agilent).

The cDNA library was sequenced on the GA IIx platform (Illumina). The resulting reads were aligned to the paa cluster genomic sequence by using the mapping software SARUMAN (4). By combining published information about promoter regions in C. glutamicum (34) with 5’-end-enriched RNAseq data, probable TSS and promoter regions were deduced for the paa gene cluster in C. glutamicum AS 1.542.

**Bioinformatic analysis of PaaR binding sites and regulons.** For the prediction of PaaR binding sites, we searched for a common motif in the upstream sequences of all genes in the paa cluster using the de novo motif extraction program MEME (1). Based on the distribution of known regulator binding sites in C. glutamicum (2), upstream regions were defined as noncoding sequence ranging from position +20 relative to the translation start codon up to the next coding sequence (CDS), with a maximum size of 620 bp and a minimum size of 40 bp. CDSs smaller than 150 bp were ignored in determining the maximum length of the upstream sequences (23). Most TetR family transcription regulators reportedly bind with 15-bp DNA sequences (35), so parameters for MEME were set to detect 15- to 22-bp palindromic motifs. Genomewide detection of putative binding sites in genomic upstream regions was carried out using PoSSumSearch (3). A positional weight matrix (PWM)-based model of the respective binding motif was generated from known binding sites for both PaaR (this study) and GtxR (18, 23, 24).

Comparative genomic identification of PaaR binding sites and reconstruction of PaaR regulons in a reference set of 57 bacterial species were performed according to the approach established previously (36), implemented in the RegPredict Web server tool (regpredict.illum.org) (33). The PaaR regulons were reconstructed separately in each taxonomic group containing a PaaR ortholog. In each group of genomes, a PaaR binding motif was identified, and a PWM was constructed using the motif discovery program in the RegPredict server. Then the taxonomically specific PWMs were used to scan all reference genomes in the respective groups and to predict novel genes in the PaaR regulon. Scores of candidate sites were defined as the sum of positional nucleotide weights. The details of the reconstructed PaaR regulons are captured and displayed in the RegPrecise database of bacterial regulons inferred by the comparative genomics approach (regprecise.illum.org) (32).

**PaaR purification and EMSA.** E. coli strain ER2566 carrying the expression plasmid pTXB1::paaR was grown in LB broth at 37°C with 200 μg/ml ampicillin. Isopropyl-β-D-thiogalactopyranoside (IPTG) was added at a final concentration of 0.4 mM when the cell density reached an OD_{600} of 0.7, and the temperature was adjusted to 18°C, followed by overnight incubation. PaaR was purified using the IMPACT system (New England BioLabs). For electrophoretic mobility shift assays (EMSA) with
PaaR, Cy3-labeled DNA probes of genomic upstream regions were obtained by PCR and column-based purification. The 20-μl reaction mixtures contained 20 mM Na2HPO4, 75 mM KCl, 0.1 mM EDTA, 5% glyc erin, 25 μg/ml salmon sperm blocking DNA, 0.1 pmol DNA probe, and purified PaaR at specified concentrations. In case of competition EMSA with PaaR, an unlabeled 40-bp fragment in a gradient of 0, 0.1, 0.5, 1, 5, and 10 pmol was also added to the reaction mixture. Reaction mixtures were incubated at 30°C for 20 min before being loaded onto a cooled 1.5% (wt/vol) agarose gel. The gel buffer and running buffer contained 20 mM Na2HPO4. Hexahistidyl-tagged GlxR was purified, and EMSA were performed, as described previously (23,25). Gel electrophoresis was performed for 40 min at a voltage of 90 V, and the resulting band shifts were visualized using a Typhoon 8600 imager (Amersham Bioscience).

RESULTS

Bioinformatic and functional identification of a paa gene cluster from C. glutamicum AS 1.542. Genome sequencing of C. glutamicum strain AS 1.542 revealed a gene cluster of 15 genes (EBI-EMBL accession no. HE649965) that were deduced to be responsible for PAA transport, metabolism, and regulation (Fig. 1). Except for the putative regulator (paaR) and transporter (paaT) genes, these genes show significant amino acid sequence similarities (32 to 66% [Fig. 1]) to the PAA degradation genes of E. coli (45). The paaT gene, encoding a putative transporter, is located at the beginning of the paa gene cluster in strain AS 1.542 but is absent in E. coli K-12. In contrast to the GntR-type regulatory gene paaX in E. coli, the paa cluster in strain AS 1.542 contains a putative TetR-type regulatory gene, paaR. The genetic organization of the paa gene cluster in C. glutamicum is apparently different in general from that in E. coli (Fig. 1), although a coherent paaABCDE cassette was observed in E. coli and in C. glutamicum, as well as in other bacteria (27).

paa gene clusters were also identified in the genomes of C. glutamicum strain R and Corynebacterium efficiens YS-314, with high sequence similarity and identical genetic organization to the paa gene cluster of strain AS 1.542 (Fig. 1). But no such paa gene cluster was found in C. glutamicum strain ATCC 13032 (19). The paa gene cluster of AS 1.542 was cloned into pEC-K18 mob2, and the plasmid constructed, pEC-K18 mob2::paa, was successfully mobilized into strain ATCC 13032. The transformant, ATCC 13032/pEC-K18 mob2::paa, obtained the ability to grow on PAA as the sole carbon source (see Fig. S1 in the supplemental material), indicating that the paa gene cluster of AS 1.542 is sufficient for PAA degradation.

Transcriptional organization of the paa gene cluster in strain AS 1.542. A 5′-end-enriched RNAseq library was constructed, and 6 probable TSS and promoter regions were identified by mapping sequence reads to the paa gene cluster of strain AS 1.542 (Table 1; see also Fig. S2 in the supplemental material). The identified TSS and promoter regions were manually checked and were adjusted according to the nucleotide distribution within the C. glutamicum promoter sequences (34). These results suggest that the paa gene cluster of strain AS 1.542 is organized into six transcription units: paaTK, paaABCDEGFJHI, paaR, paaI, paaY, and paaZ.
**TABLE 1 Transcriptional organization of the paa gene cluster in**
**C. glutamicum strain AS 1.542**

<table>
<thead>
<tr>
<th>Operon</th>
<th>Strand</th>
<th>TSS positiona</th>
<th>TSS base</th>
<th>Sequence</th>
<th>Match coverageb</th>
</tr>
</thead>
<tbody>
<tr>
<td>paaTK</td>
<td></td>
<td>−213</td>
<td>A</td>
<td>TAGGGT to TGGACA</td>
<td>336, 913</td>
</tr>
<tr>
<td>paaABCDEGJFH</td>
<td>+</td>
<td>−44</td>
<td>G</td>
<td>TATCCT to TTTGACC</td>
<td>27, 209</td>
</tr>
<tr>
<td>paaR</td>
<td>−</td>
<td>−76</td>
<td>A</td>
<td>TACAGT to TTCCCGG</td>
<td>1, 7</td>
</tr>
<tr>
<td>paaI</td>
<td>+</td>
<td>−60</td>
<td>A</td>
<td>TAGACT to GTGAAA</td>
<td>22, 59</td>
</tr>
<tr>
<td>paaY</td>
<td>−</td>
<td>−54</td>
<td>C</td>
<td>TAATAT to TACACT</td>
<td>0, 35</td>
</tr>
<tr>
<td>paaZ</td>
<td>−</td>
<td>−57</td>
<td>A</td>
<td>TATATC to TTCCAG</td>
<td>21, 170</td>
</tr>
</tbody>
</table>

*a* Relative to the translation start codon.

*b* Perfect match coverage and best match coverage are defined according to Blom et al. (4).

**paaR** encodes a transcription regulator repressing the transcription of **paa** genes in strain AS 1.542. The **paaR** gene was annotated as a putative TetR family transcription regulator. The growth of strain AS 1.542 and its mutant AS 1.542/paaRth was observed on 5 mM PAA as a carbon source. The mutant AS 1.542/paaRth exhibited significantly faster adaptation to PAA as a carbon source than did AS 1.542. Furthermore, the induction of the **paa** genes and the influence of **paaR** disruption on the transcriptional control of the **paa** genes were determined by RT-qPCR (Fig. 2). The results showed that the mRNA levels of all **paa** genes of strain AS 1.542 increased when this strain was grown on PAA rather than glucose as a carbon source. The levels of the six transcription units, **paaABCDEGJFH**, **paaTK**, **paaR**, **paaI**, **paaY**, and **paaZ**, increased 178- to 896-, 36- to 64-, 13-, 7-, 59-, and 49-fold, respectively. When the mutant AS 1.542/paaRth was grown on glucose, the mRNA levels of all **paa** genes were comparable to those in AS 1.542/paaRth grown on PAA. These results clearly demonstrate that **PaaR** negatively regulates the transcription of **paa** genes in AS 1.542.

**PaaR** binds to the upstream regions of the **paa** gene cluster, and phenylacetyl-CoA (PA-CoA) is the effector. EMSA of the upstream regions of **paaK** (271 bp), **paaA** (258 bp), **paaYZ** (203 bp), and **paaRI** (247 bp) were performed with Cy3-labeled DNA probes. The addition of 5 pmol of **PaaR** (a 50-fold excess over the DNA probe) resulted in clear shifts with the DNA probes of **paaK**, **paaA**, and **paaYZ** but not with the DNA probe of **paaRI** (Fig. 3A). A distinct shift of the **paaRI** DNA probe was observed only at a higher concentration (60 pmol) of **PaaR**, and a blurred smear was observed for the negative-control **hemH** probe (114 bp) under these conditions (Fig. 3B).

The effects of **PAA**, acetyl-CoA, and PA-CoA at a concentration of 0, 10, 25, 50, 100, 250, or 500 μM on the binding between **PaaR** and the **paaA** DNA probe were determined. The results showed that **PAA** and acetyl-CoA did not affect the binding between **PaaR** and the DNA probe of **paaA** (see Fig. S3 in the supplemental material). It was observed that 250 μM PA-CoA abolished the binding of **PaaR** to the probes for **paaK**, **paaA**, and **paaYZ** (Fig. 3A). For the upstream region of **paaRI**, which required 60 pmol **PaaR** to introduce an observable shift, the effector identified above, **PA-CoA**, did not result in a clear abortion of **PaaR** binding at 250 μM or even at 3 mM PA-CoA. Instead, smeared bands were observed under these conditions (Fig. 3C).

**Determination of the PaaR binding sites.** The upstream regions of the **paa** genes were searched with the MEME software for a putative **PaaR** binding motif, and a 22-bp palindromic motif (ACTnACCnnGnnCGnnCGGTnAGT) was identified at each of the upstream loci of **paaK**, **paaA**, and **paaYZ**. A putative **PaaR** binding site at the upstream region of **paaRI** (Fig. 4) was observed, and the sequence of this site was less similar to the 22-bp palindromic motif identified. Forty-base-pair fragments, each carrying the 22-bp motif and an additional 9-bp flanks at both sides (thus totaling 40 bp), were synthesized and used for competition EMSA. The results showed that 0.5 pmol (i.e., a 5-fold excess) of the respective 40-bp fragment competed effectively for **PaaR** binding with the Cy3-labeled DNA probes of the upstream regions of **paaK**, **paaA**, and **paaYZ** (Fig. 3D). But 0.5 pmol (i.e., a 5-fold excess) or even 6 pmol (i.e., a 60-fold excess) of the 40-bp **paaRI** competition DNA fragment resulted in only partial competition for **PaaR** binding with the upstream DNA probe of **paaRI** (Fig. 3C).

**GlxR binds to the upstream regions of the paa gene cluster.** It was found that the **paa** gene transcription levels in AS 1.542/paaRth were lower when this mutant was grown on glucose than when it was grown on PAA (Fig. 2). This indicated that a second transcriptional regulator was involved in **paa** gene regulation in AS 1.542. It was reported that **GlxR** acts as a global regulator for aromatic-compound degradation in *C. glutamicum* ATCC 13032, and the consensus sequence TGTGAnnTAnnTCACA was identified (23). By application of the PoSSuMsearch tool, possible **GlxR** binding sites were searched for in the upstream regions of the **paa** genes from *C. glutamicum* strain AS 1.542.
well as from *C. efficiens*. GlxR binding sites were found in the upstream regions of the *paaK*, *paaA*, and *paaYZ* genes of all three *Corynebacterium* strains. In *C. glutamicum* strain AS 1.542 and strain R, less-conserved GlxR binding sites were also found in the upstream region of *paaRI*. They were located around the putative promoter regions of the *paa* genes and upstream of the PaaR binding sites (Fig. 4). EMSA were performed to confirm the putative GlxR binding sites in strain AS 1.542. The results showed that the upstream region of *paaRI*. They were located around the putative promoter regions of the *paa* genes and upstream of the PaaR binding sites (Fig. 4). EMSA were performed to confirm the putative GlxR binding sites in strain AS 1.542. The results showed that the

![Electrophoretic mobility shift assays of PaaR binding to the upstream regions of the *paaK*, *paaA*, *paaYZ*, and *paaRI* genes. The upstream region of *hemH* from *C. glutamicum* AS 1.542 was tested in parallel as a negative reference (A and B). Cy3-labeled DNA probes were added at 0.1 pmol, and PaaR protein was added at 5 pmol (A and D), at 60 pmol (C), or as indicated. The effector PA-CoA was added at 250 μM (A) or as indicated (C). For competition EMSA experiments with the *paaRI* DNA probe (C) or with the *paaK*, *paaA*, or *paaYZ* DNA probe (D), unlabeled 40-bp DNA fragments carrying the corresponding putative binding sites were added. As a negative control, an unlabeled fragment (40 bp) of the *argC* gene was added to an EMSA mixture with the Cy3-labeled *paaA* probe (D).](http://aem.asm.org/)

![Regulatory regions of the *paa* gene cluster in strain AS 1.542. Upstream DNA sequences of the *paaK* (I), *paaA* (II), *paaYZ* (III), and *paaRI* (IV) genes are shown in 5’→3’ orientation relative to the translation start codons of the respective genes. The determined transcription start site is in boldface, and bent arrows indicate the direction of transcription; the derived –10 and –35 regions are underlined. PaaR binding sites are indicated by boxes with solid borders, and GlxR binding sites are indicated by dotted boxes with dotted borders.](http://aem.asm.org/)
40-bp fragments of the upstream regions of paaK, paaA, and paaYZ containing the previously identified GlxR binding motif (23) specifically bound with GlxR in the presence of cyclic AMP (cAMP) (Fig. 5).

Comparative-genomics reconstruction of orthologous PaaR regulons in bacteria. In addition to the PaaR protein from C. glutamicum, two PaaR proteins from T. thermophilus and B. cenocepacia had been identified previously (13, 38). By using these functionally characterized PaaR proteins, PaaR orthologs were identified by BLAST searches against the nonredundant set of sequenced bacterial genomes and were used to analyze the phylogenetic relationships between these proteins (Fig. 6). As a result, a single copy of the paaR gene was detected in the reference set of 57 bacterial genomes from the following taxonomic groups: Actinobacteria, Alpha- and Betaproteobacteria, Deinococcus-Thermus, and Flavobacteria. In most of these genomes, the paaR gene is located in the vicinity of the paa genes; the only exception is 3 Deinococcus spp. For inference of PaaR regulons in these genomes, we applied the comparative-genomics approach implemented in the RegPredict Web server (33, 36).

Phylogenetic analysis of the PaaR-like proteins detected (Fig. 6) provided a basis for transcription factor binding site (TFBS) identification among different phyla. First, we collected training sets of prospective target genes for each of 10 defined taxonomic groups possessing PaaR orthologs. These training sets were defined by the analysis of chromosomal colocalizations of paaR genes and information on the known PaaR binding sites from strain AS 1.542 from this study, as well as from T. thermophilus and B. cenocepacia (13, 38). Using the motif recognition program applied to each training set of upstream gene regions, we identified conserved PaaR binding motifs, constructed a positional weight matrix for each motif identified, and searched for additional PaaR-binding sites in the genomes analyzed. Except for three Deinococcus species, most of the PaaR binding sites detected were located adjacent to their target genes (see the “TFBS” sheet of Table S2 in the supplemental material; see also Discussion). The resulting PaaR binding motifs are well conserved between the Actinobacteria and Proteobacteria phyla, whereas the PaaR binding motifs in the Deinococcus-Thermus group and Flavobacteria have several substituted nucleotides in their consensus sequences (Fig. 6). The reconstructed PaaR regulons are summarized in Table S2 in the supplemental material, and the detailed information on the predicted PaaR-binding sites and downstream regulated genes are available in an updated version of the RegPrecise database (32).

**DISCUSSION**

C. glutamicum strain AS 1.542 possesses a complete paa gene cluster, conferring a PAA degradation pathway and its regulation. This paa gene cluster is subjected to regulation by the TetR-type regulator PaaR. In contrast to those in P. putida (11), E. coli (9, 10), B. cenocepacia (13), and T. thermophilus (38), the paa gene cluster in C. glutamicum consists of 6 transcription units. The differences in the transcription levels of the paa genes suggest that PaaR strongly regulates the transcription of paaABCDGJFH, paaI, paaZ, and paaTK, while it weakly regulates the transcription of paaA and paaY. EMSA results from this study indicated that a 12-times-higher concentration of PaaR was needed to retard the probe of the paaR upstream region than to retard the other probes. Therefore, it is proposed that PaaR exerts a weak self-repression and a strong repression of the other genes in the cluster in C. glutamicum.

The PAA catabolic pathway is widespread in the bacterial domain (45). Bacteria have evolved two different systems to regulate this PAA catabolism, namely, the TetR-type regulation represented by the PaaR proteins of C. glutamicum (this study), T. thermophilus (38), and B. cenocepacia (13) and the GntR-type regulation represented by PaaX of E. coli (9), P. putida, and other Pseudomonas species (7, 11). The binding consensus sequence of GntR-type regulator PaaX was identified as TGATTCC(N$_2$)$_8$GAATCA in E. coli (21), and a similar inverted repeat sequence was found in Pseudomonas strain Y2 (7). The binding sequences of the TetR-type regulators from the PaaR proteins of T. thermophilus HB8 (38) and B. cenocepacia K56-2 (13) show little similarity to the binding consensus sequences of GntR-type regulators from E. coli (21) and Pseudomonas species (7). In this study, the identified binding sequence of the TetR-type PaaR protein from C. glutamicum strain AS 1.542 shows some similarity to the binding sequences of PaaR from B. cenocepacia K56-2 (13). Moreover, analysis of the respective paa gene clusters from C. glutamicum strain R and C. efficiens reveals a binding consensus sequence of the TetR-type PaaR protein in Corynebacterium species, which is characterized by an imperfect palindromic sequence (ACTnACGGnCGnnCCGnTnAGT; 22 bp).

Based on these findings and by integration of a comparative-genomics approach, we were able to define PaaR regulons in 57 distinct bacterial species from the Actinobacteria, Alphaproteobacteria, and Betaproteobacteria, Flavobacteria, and Deinococcus-Thermus phyla. PaaR regulation was restricted to paa catabolic and transport genes in a chromosomal cluster, and the sizes of predicted PaaR regulons range from one target operator (e.g., in Flavobacteria) to nine target operators (as in Deinococcus). Phylogenetic analysis of PaaR proteins and the observed conservation of the PaaR binding motifs within each taxonomic group (Fig. 6) indicate that the PaaR regulators and motifs coevolved via presumably vertical evolution in Bacteria. Interestingly, in Deinococcus species, the deduced PaaR binding motifs are not associated with PAA utilization genes, and there are no PaaR binding motifs adjacent to paa genes. Therefore, how the paa gene cluster is regulated in the Deinococcus species is still not clear.

A recent study has identified in vivo GlxR binding sites at the corresponding paaK, paaA, and paaYZ promoter regions in C. glutamicum.
FIG 6  Phylogenetic tree of a selection of PaaR orthologs marked with taxonomic groups and identified PaaR-binding DNA motifs (generated by [http://weblogo.berkeley.edu](http://weblogo.berkeley.edu)) for each group. The tree was constructed using the neighbor-joining method with 1,000 bootstraps. For the identities of the PaaR orthologs shown here, please see Table S2 in the supplemental material.
glutamicum strain R (47); of these, the paa gene cluster is almost identical to that of strain AS 1.542. This is in agreement with our EMSA results for GlxR in strain AS 1.542. As proposed in this study and previous reports (23, 24), GlxR regulates PAA degradation, although the mechanism of this regulation needs more investigation. It is known that GlxR is a cAMP-binding global regulator (18, 23, 24). The cAMP receptor protein (CRP), a homolog to GlxR in C. glutamicum, senses the intracellular cAMP levels and superimposed the PaaX-mediated repression in E. coli (9). Thus, it would be interesting to determine if phenylacetic acid catabolism is related to the intracellular cAMP level. The involvement of the global regulator GlxR and a pathway-specific regulator, PaaR, of the PAA catabolic pathway in strain AS 1.542 exemplifies the proposal of two levels of transcriptional regulation for aromatic degradation (8). It would be of interest to investigate exactly how GlxR and PaaR coordinate in the regulation of aromatic-com- pound degradation in C. glutamicum and other bacteria.

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