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## **Biodegradation**

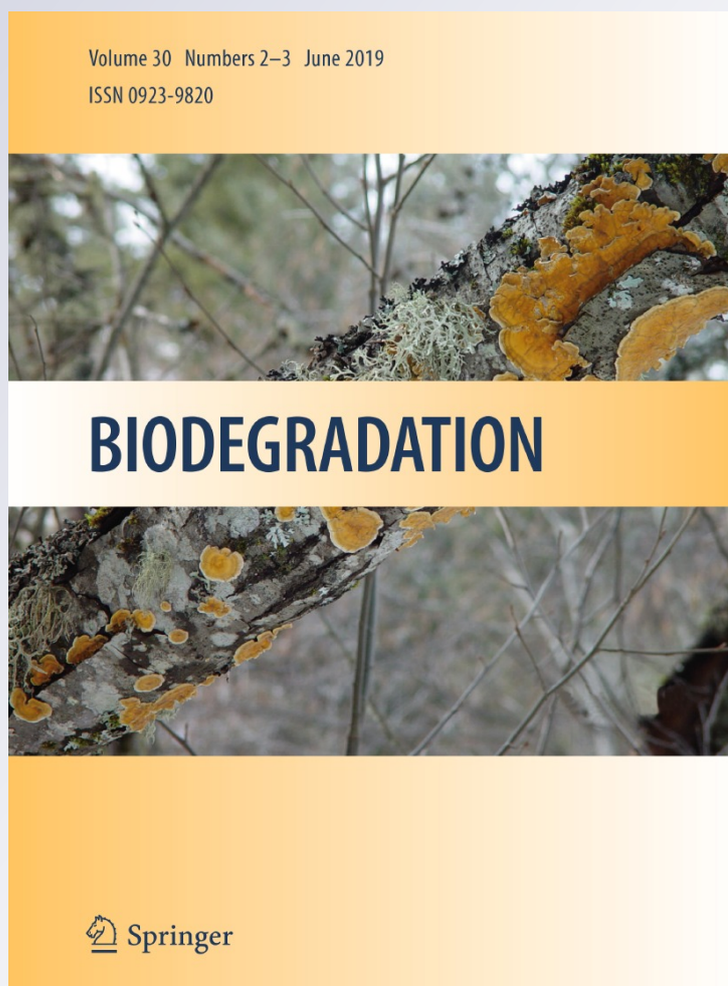
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# Identification of naphthalene carboxylase subunits of the sulfate-reducing culture N47

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**Abstract** Expanding industrialization and the associated usage and production of mineral oil products has caused a worldwide spread of polycyclic aromatic hydrocarbons. These pollutants accumulate and persist under anoxic conditions but little is known about the biochemical reactions catalyzing their anaerobic degradation. Recently, carboxylation of naphthalene was demonstrated for the sulfate-reducing culture N47. Proteogenomic studies on N47 allowed the identification of a gene cluster with products suggested to be involved in the initial reaction of naphthalene degradation. Here, we performed comparative proteomic studies with N47 proteins extracted from naphthalene versus 2-methylnaphthalene-grown cells on blue native PAGE. The

analysis led to the identification of subunits of the naphthalene carboxylase of N47. Moreover, we show that the identified subunits are encoded in an operon structure within the previously mentioned naphthalene carboxylase gene cluster. These findings were supported by a pull-down experiment revealing in vitro interaction partners of a heterologously produced GST-tagged naphthalene carboxylase subunit. Based on these lines of evidence, naphthalene carboxylase is proposed to be a complex of about 750 kDa. Naphthalene carboxylase can be seen as a prototype of a new enzyme family of UbiD like de-/carboxylases catalyzing the anaerobic activation of non-substituted polycyclic aromatic hydrocarbons.

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## Introduction

Polycyclic aromatic hydrocarbons (PAHs) are ubiquitous and persistent environmental pollutants originating from mineral oil products and spills, or from incomplete combustion. The high chemical stability and low aqueous solubility of PAHs make them very recalcitrant, especially in anoxic habitats (Meckenstock et al. 2016). Naphthalene represents the simplest polycyclic aromatic hydrocarbon and due to its

relatively high solubility in water (240  $\mu\text{M}$  at 25 °C) compared to larger molecular weight PAHs, it is often used as a model compound for studying PAHs degradation. In the last two decades, microbial degradation of naphthalene was reported under sulfate-reducing (Galushko et al. 1999; Meckenstock et al. 2000), iron-reducing (Kleemann and Meckenstock 2011; Lovley et al. 1994; Marozava et al. 2018), and methanogenic conditions (Christensen et al. 2004) (for review see Meckenstock and Mouttaki 2011).

Metabolite studies with sulfate-reducing cultures revealed 2-naphthoic acid as a potential intermediate of anaerobic naphthalene degradation and the incorporation of [ $^{13}\text{C}$ ] bicarbonate into the carboxyl group indicated a direct carboxylation of naphthalene (Meckenstock et al. 2000; Zhang and Young 1997). More reduced metabolites such as tetrahydro-2-naphthoic acid suggested a subsequent ring reduction to initiate ring cleavage (Meckenstock et al. 2000). Later, proteogenomic studies on the sulfate-reducing cultures N47 and NaphS2 allowed the identification of gene clusters specifically induced by growth with naphthalene as electron and carbon source (Bergmann et al. 2011b; DiDonato et al. 2010). Some of the gene products of one cluster exhibited strong similarity to a gene encoding ubiquinone decarboxylase UbiD of *Escherichia coli* and subunits of phenylphosphate carboxylase, one of the key enzymes in anaerobic phenol degradation (Bergmann et al. 2011b; DiDonato et al. 2010; Schühle and Fuchs 2004). Furthermore, the putative naphthalene carboxylase genes showed homology to genes putatively coding for the anaerobic benzene carboxylase detected in the iron-reducing enrichment culture BF (Abu Laban et al. 2010) (Fig. 1). Recently, Mouttaki et al. (2012) provided the first biochemical evidence confirming the naphthalene carboxylation reaction in the sulfate-reducing culture N47.

In order to identify the subunits of the naphthalene carboxylase protein complex, a recombinant production of the putative subunits was attempted in various *E. coli* strains with differing protein tags but yielded, for the most part, insoluble protein. During attempts to purify the naphthalene carboxylase complex under anaerobic conditions directly from N47 cells, all enzyme activity was lost (Kölschbach, unpublished results).

As such, it was unclear which of the gene products found to be differentially abundant in the naphthalene

grown N47 culture, were involved in the naphthalene carboxylase protein complex (Bergmann et al. 2011b).

The aim of the present study was, thus, to identify which genes belong to the putative naphthalene carboxylase operon and which polypeptides constitute the naphthalene carboxylase protein complex of N47.

## Materials and methods

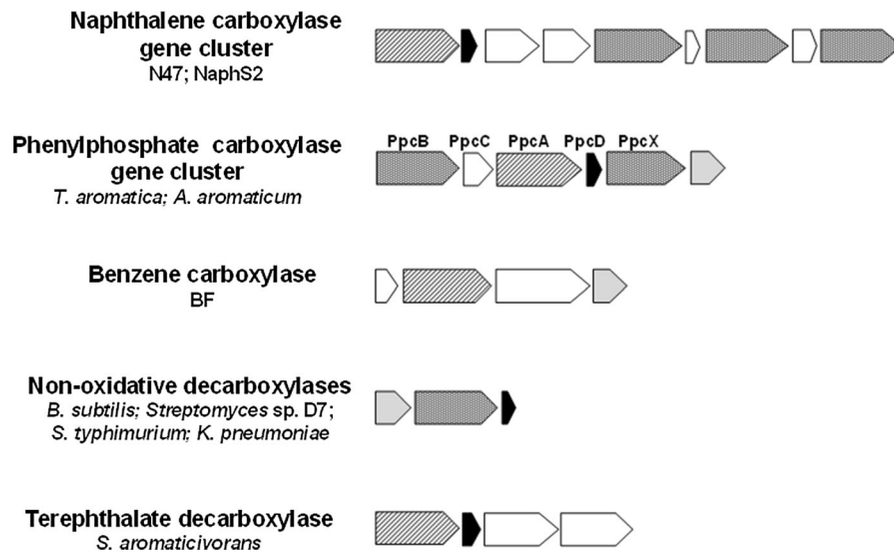
### Growth of culture N47 and preparation of cell-free extracts

The sulfate-reducing culture N47 was cultivated at 30 °C in serum bottles containing bicarbonate-buffered freshwater medium using naphthalene as carbon and electron source and sulfate as electron acceptor (Meckenstock et al. 2000). Naphthalene or 2-methylnaphthalene was added as a 1.5% (w/v) solution in 2,2,4,4,6,8,8-heptamethylnonane (20 mL/L culture volume). For the preparation of cell-free extracts, cells were harvested by centrifugation (3200 $\times g$ , 4 °C, 30 min) under anoxic conditions in an anaerobic chamber (Labstar, MBRAUN, Garching, Germany) under 100%  $\text{N}_2$  atmosphere. The anaerobic chamber, where all anaerobic experiments took place, is equipped with an internal  $\text{O}_2$  sensor to ensure that the oxygen concentration does not reach levels above 3 ppm. The cells were harvested in the mid-exponential growth phase, which had been determined in previous experiments (Meckenstock et al. 2000; Musat et al. 2009), after 6 weeks of cultivation. The cells were disrupted by 3–4 cycles of sonication (15–20 s/cycle; 30 kHz; 50% pulse; 40% amplitude; UP50H, Hielscher Ultrasonics GmbH, Teltow, Germany) in a pre-chilled metallic block followed by a 20 min centrifugation step (13,000 $\times g$ , 4 °C). The supernatant was amended with 1  $\times$  complete protease inhibitor cocktail (Roche, Basel, Switzerland) to prevent protein degradation.

### Operon mapping (transcriptional analysis)

#### RNA extraction

N47 cells were harvested by centrifugation of 150 mL culture. The cell pellets were resuspended in a buffer containing 50 mM Na-Acetate, pH 5.3 and 10 mM Na-EDTA. RNA was extracted by bead beating in the



**Fig. 1** Organization of gene clusters encoding UbiD-like proteins involved in anaerobic degradation of aromatic compounds. Open reading frames are represented by arrows. *UbiD*-like genes are shown in a pattern fill, whereas striped arrows

represent *PpcA*-like genes. *PpcD*-like genes are indicated by dark grey and *UbiX*-like genes in light grey. Further details are mentioned in the text

presence of sodium phosphate buffer and sodium dodecyl sulfate as reported by Schmitt et al. (1990). After centrifugation (5 min, 13,000×g, 4 °C), the supernatant was extracted with equal volumes of phenol-chloroform-isoamylalcohol [24:24:1 (v/v/v)] and chloroform-isoamylalcohol [24:1 (v/v)] prior to precipitation (30 min, 13,000×g, 4 °C) with 2 μL glycogen (Roche, Basel, Switzerland) and 2 volumes of polyethylene glycol (Griffiths et al. 2000). Subsequently, the nucleic acid pellet was washed with 70% ethanol and resuspended in RNase-free water (Promega, Fitchburg, WI). The co-extracted DNA was digested with RQ1 RNase-free DNase (Promega, Fitchburg, WI) according to the manufacturer's instructions. The RNA was visualized by standard agarose gel electrophoresis and analyzed using NanoDrop (NanoDrop Technologies, Inc., Wilmington, DE) to estimate the quantity. RNA was stored at – 80 °C until further use.

#### RT-PCR

The RNA was reverse transcribed to cDNA by GoScript™ reverse transcriptase using random hexamer primers (0.5 μg/reaction; Thermo Fisher Scientific, Waltham, MA) or gene-specific reverse primers (Table 1; 50199r, 48240r, 46360r, 44481r; 5 pmol

each/reaction) as described by the manufacturer's protocol (Promega, Fitchburg, WI). These primers were designed manually and synthesized by Eurofins Genomics (Ebersberg, Germany). As negative controls, reactions were performed without the reverse transcriptase to detect eventual residual genomic DNA contamination.

#### PCR amplification

Intergenic regions were amplified via PCR using the primer sets listed in Table 1 with the following program: 5 min at 94 °C initial denaturation, 35 cycles of amplification (30 s at 94 °C, 15 s at 55 °C, 1 kb/60 s at 72 °C) and 5 min at 72 °C of final extension. The PCR reaction used 1 × Taq PCR buffer, 1.5 mM MgCl<sub>2</sub>, 0.2 mM dNTPs, 0.5 μM primer (Eurofins Genomics, Ebersberg, Germany) and 1.25 U Taq polymerase (Thermo Fisher Scientific, Waltham, MA).

#### Cloning, production and purification of GST-NcA

Genomic DNA from the sulfate-reducing culture N47 was extracted with the QIAamp DNA Mini kit (Qiagen, Hilden, Germany) as described by the manufacturer. Gene *N47\_K27540* encoding one of

**Table 1** Primer sets designed and utilized for transcriptional analysis (operon mapping)

Primer set	Primer name	Sequence (5'–3')	Amplicon size (bp)
– A1	53306f	TATTAACGTAAAGGCAGGC	1239
	52068r	TTTCCAAGAGCTCAATAAACT	
– A	52641f	ATTATCAGAAGGGCTGTTTAA	574
	52068r	TTTCCAAGAGCTCAATAAACT	
A	50736f	AAAGATGTTTCAGGAAAAGGT	538
	50199r	GCACTTCATCATGATGTTG	
B	49346f	GAAGTACAGGAGGAGAAAAGC	1107
	48240r	CTTCATTGACTTCCACAAGT	
C	46832f	AGATCGACTGGCACAAGAAGT	473
	46360r	TGCGACATGGCACAGATAG	
D	45060f	TAGAGATCGTGTGGTGGCTTT	580
	44481r	TTCGCCTGAATGATTGAGG	
E	43268f	AAGATAGGCTTGGATGCCAC	913
	42356r	TCCAGGATAACACCCCTGATAT	
E1	43268f	AAGATAGGCTTGGATGCCAC	496
	42773r	GGTCGAGGTAACAATGCCTATT	
A1	pGEX6P1_SalI_52110f	GTAGTCGACTGATGGCGTTTAAAGATTTGAG	1923
	50199r	GCACTTCATCATGATGTTG	
B1	pGEX6P1_BamHI_50243f	GTAGGATCCATGGAAGAACGTGAAATACT	1837
	48416r	AAATCAATGACGCTCTTGCCTATCTT	
C1	48303f	ATGGACACTAGAGAATTGGT	1944
	46360r	TGCGACATGGCACAGATAG	
D1	pGEX6P1_SalI_46472f	GTAGTCGACTGATGAGTTCATTAAGAGAA	2003
	44481r	TTCGCCTGAATGATTGAGG	

the naphthalene carboxylase subunits (NcA) (Bergmann et al. 2011b), was amplified from N47 genomic DNA using the primer set ncA\_for (GTAGTCGACTGATGGCGTTTAAAGATTTGAG) and ncA\_rev (GTAGCGGCCGCGAATCCATATTTATTCCAA TTG). The PCR product was purified using the PCRextract kit (5Prime, Hamburg, Germany) according to the manufacturer's protocol. Standard molecular cloning procedures were applied. Enzymes were purchased from Thermo Fisher Scientific. The amplicon was inserted into the expression vector pGEX-6P-1 (GE Healthcare Europe, Freiburg, Germany) containing an IPTG-inducible *lac*-promotor and a GST-tag. Chemically competent *E. coli* Rosetta<sup>TM</sup> 2(DE3)pLysS (Merck, Darmstadt, Germany) were transformed with the plasmid pGEX-6P-1<sub>ncA</sub> as described previously (Inoue et al. 1990). For production of the GST-tagged fusion protein GST-NcA, 1 L of LB-medium (Carl Roth®, Karlsruhe, Germany)

containing ampicillin (100 µg/mL) was inoculated with the recombinant *E. coli* Rosetta<sup>TM</sup> 2(DE3)pLysS including the plasmid pGEX-6P-1<sub>ncA</sub> and incubated at 37 °C. Gene expression was induced by addition of 0.5 mM IPTG when the culture reached an optical density of 0.5 at 578 nm, and the cells were shifted to room temperature for an additional 3 h. The cells were harvested at 3200×g for 15 min at 4 °C and the pellet was stored at – 20 °C until further use. GST-NcA was purified using batch purification with Glutathione Sepharose 4B beads as recommended by the manufacturer with the following modifications (GE Healthcare Europe, Freiburg, Germany). The stored pellet was resuspended in binding buffer (50 mM Tris/HCl, pH 7.5; 1 M NaCl; 1 mM EDTA) and the cells were disrupted by sonication as described above. After sonication, 1% IGEPAL® CA-630 (Sigma-Aldrich, St. Louis, MO) was added. For purification, 1 mL of Glutathione Sepharose 4B slurry was used. The beads

were incubated with *E. coli* Rosetta<sup>TM</sup> 2(DE3)pLysS cell-free extract containing GST-NcA on an end-over-end rotator at 4 °C for 2 h and subsequently 3 times washed with washing buffer (50 mM Tris/HCl, pH 7.5; 140 mM NaCl). The GST-NcA coupled to the Sepharose beads was used for subsequent pull-down experiments. The purified protein was stored in 20 mM Tris/HCl, pH 7.5 with 50% glycerol at – 20 °C.

#### *In vitro* protein interaction analysis (pull-down experiment)

Purified GST-NcA coupled to the Glutathione Sepharose beads was washed with 1x PBS buffer (140 mM NaCl, 2.7 mM KCl, 10 mM Na<sub>2</sub>HPO<sub>4</sub>, 1.8 mM KH<sub>2</sub>PO<sub>4</sub>, pH 7.3) and 120 µL of the slurry was incubated for 4 h at 4 °C with 500 µL of the respective cell-free extract. After three washing steps (1 × PBS, 5 min, 4 °C), 30 µL beads were combined with 5 × SDS-PAGE loading dye and analyzed via SDS-PAGE. Protein bands were excised from the SDS-PAGE and submitted for proteomic analysis.

#### Blue native PAGE

Blue native PAGEs (BNPs) were performed in an anoxic chamber with N<sub>2</sub>-atmosphere using NativePAGE<sup>TM</sup> Novex<sup>®</sup> 4–16% Bis-Tris gels in the XCell SureLock<sup>®</sup> Mini Cell (Life Technologies, Carlsbad, CA, USA) according to the manufacturer's protocol. Activity of the naphthalene carboxylase was confirmed by an activity assay before samples were loaded on the gel. The activity assay contained a saturated naphthalene solution, 25 mM NaHCO<sub>3</sub> and 5 mM ATP in 100 mM MOPS/KOH, 15 mM MgCl<sub>2</sub>, pH 7.3 buffer. The assay was started by the addition of cell-free extract and stopped after 0, 15, 45, 90 min with 10% formic acid.

#### Proteomic analysis

##### *In-gel tryptic digest*

Coomassie-stained excised gel-pieces were digested as described previously (Merl et al. 2012). The eluted peptides were dried in a speed vac and stored at – 20 °C until further use.

#### Mass spectrometric measurements

Dried digested samples were thawed and dissolved in 45 µL of 2% acetonitrile/0.5% trifluoroacetic acid. After brief centrifugation (13,000×g, 15 min), LC-MS/MS analysis was performed on an Ultimate3000 nano HPLC system (Dionex, Sunnyvale, CA) online coupled to a LTQ OrbitrapXL mass spectrometer (Thermo Fisher Scientific, Waltham, MA) by a nano spray ion source as described previously (Hauck et al. 2010; Merl et al. 2012). The samples were analyzed with and without the addition of dimethylsulfoxide (DMSO) to the buffers. Samples were loaded onto the C18 trap column at a flow rate of 30 µL/min in 7% acetonitrile/0.1% formic acid (3% buffer B (73% ACN/3% DMSO/0.1% formic acid (FA) in HPLC-grade water) and 97% buffer A (2% ACN/3% DMSO/0.1% FA)) (Hahne et al. 2013). After 5 min, the peptides were eluted from the trap column and separated on the analytical column by a 135 min gradient from 7 to 32% acetonitrile in 0.1% formic acid (3 to 35% of buffer B) at 300 nL/min flow rate followed by a short gradient from 32 to 93% acetonitrile in 0.1% formic acid (35 to 95% buffer B) in 5 min. Between each sample, the gradient was set back to 7% acetonitrile in 0.1% formic acid (3% buffer B) and left to equilibrate for 20 min.

From the MS pre-scan, the 10 most abundant peptide ions were selected for fragmentation in the linear ion trap if they exceeded an intensity of at least 200 counts and if they were at least doubly charged. During fragment analysis a high-resolution (60,000 full-width half maximum) MS spectrum was acquired in the Orbitrap with a mass range from 300 to 1500 Da.

#### Label-free quantitative analysis

The acquired raw data of each dataset was loaded in the Progenesis LC-MS software (version 2.5, Nonlinear) for label free quantification and analyzed as described previously (Hauck et al. 2010; Merl et al. 2012). Briefly, profile data of the MS scans were transformed to peak lists with respective peak m/z values, intensities, abundances (areas under the peaks) and m/z width. MS/MS spectra were treated similarly. After reference selection, the retention times of the other samples were automatically aligned to a maximal overlay of all features. Features with only one

charge or more than seven charges were excluded from further analyses. After normalization and assignment of the samples to the respective groups, all MS/MS spectra were exported as Mascot generic file (mgf) and used for peptide identification with Mascot (version 2.4) in the N47 protein database (version 4, 1302846 residues, 5001 sequences). Search parameters used were: 10 ppm peptide mass tolerance and 0.6 Da fragment mass tolerance, one missed cleavage allowed, carbamidomethylation was set as fixed modification, methionine oxidation and asparagine or glutamine deamidation were allowed as variable modifications. Searches were performed with a Mascot ion score cut-off of 30 and an appropriate significance threshold  $p$ , in order to reach a maximum false discovery rate of 1%. Search results were re-imported into the Progenesis LC–MS software. The abundances of all peptides allocated to each protein were summed up and exported. The resulting normalized protein abundances were used for calculation of fold-changes of proteins and for calculation of significance using an unpaired both-sided Student's  $t$  test in Excel.

#### Protein determination

Protein concentrations were determined by the method of Bradford (1976) using Quick Start™ Bradford 1 × Dye Reagent (Bio-Rad Laboratories, Hercules, CA) and bovine serum albumin as standard.

#### SDS-PAGE

SDS polyacrylamide gel electrophoresis was performed as described previously (gel concentration 15%) (Laemmli 1970). PageRuler prestained protein ladder was used as a marker (Thermo Fisher Scientific, Waltham, MA).

## Results

#### Determining the structure of the naphthalene carboxylase operon

Messenger RNA was extracted from N47 cells grown with naphthalene in order to identify genes of the putative naphthalene carboxylase gene cluster belonging to a transcriptional unit. The RNA was reverse

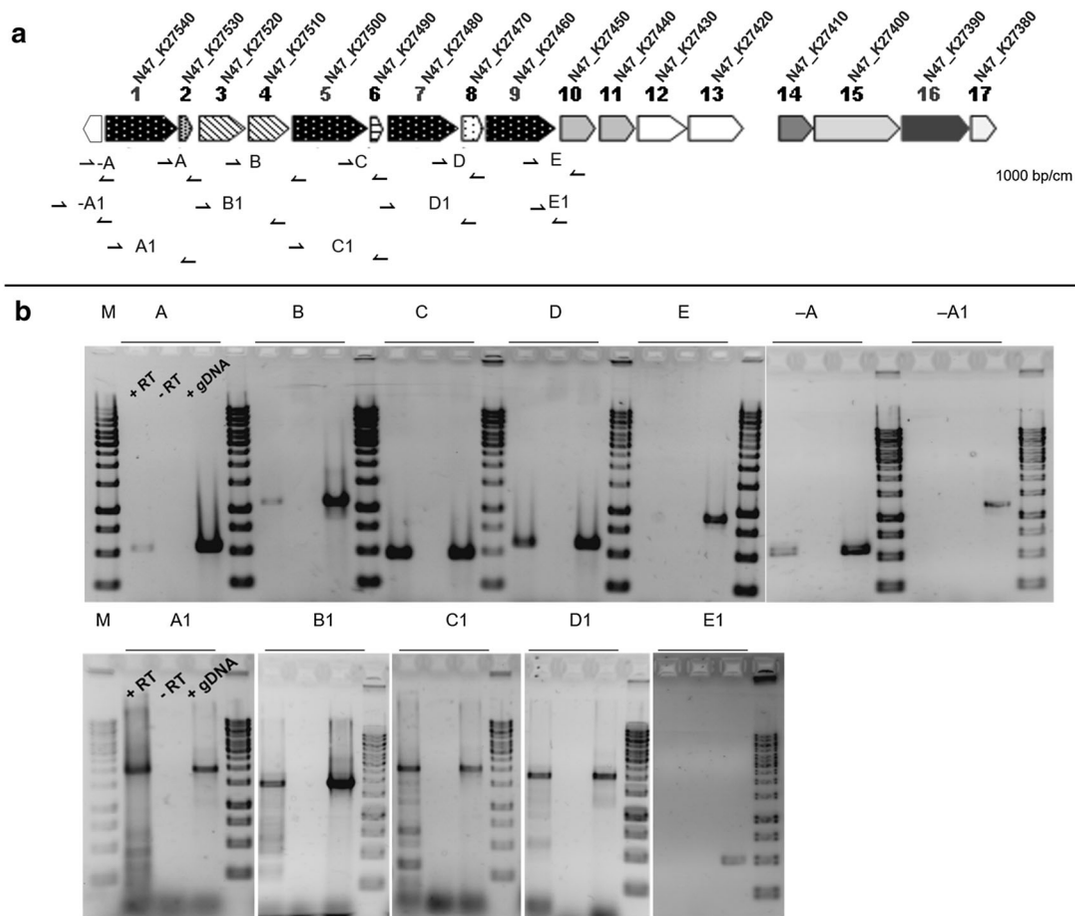
transcribed into cDNA and PCR was performed using primers covering genes and/or intergenic regions to identify the potential operon (Table 1). Two genes were assigned to belong to the same operon (same mRNA) if a forward and a reverse PCR primer hybridizing to one of two adjacent genes, respectively, produced a PCR amplicon. Gene annotations were taken from Bergmann et al. (2011b). Amplicons were obtained between genes *N47\_K27540* and *N47\_K27520*, *N47\_K27520* and *N47\_K27500*, *N47\_K27500* and *N47\_K27480*, and between *N47\_K27480* and *N47\_K27460*. The PCR-products indicated that the naphthalene carboxylase gene cluster of strain N47 is transcribed as an operon of around 8.9 kb containing 9 genes from genes *N47\_K27540* to *N47\_K27460* (Fig. 2).

#### Determining the structure of the naphthalene carboxylase protein complex

Blue native PAGEs (BNPs) and LC–MS/MS analysis of the bands were performed to determine the size and polypeptide composition of the naphthalene carboxylase complex. Bands were excised from the gel and digested in-gel with trypsin. Then, the samples were dried and LC–MS/MS analysis was performed to determine the peptide composition of the different bands. The BNPs were performed under anoxic conditions to ensure that molecular oxygen did not impair the native conformation of the protein complexes.

Cell-free extracts of N47 cells grown with naphthalene exhibited a prominent band with a size of 750 kDa (Fig. 3), which was not present in cells grown with 2-methylnaphthalene. Furthermore, two prominent bands of around 560 kDa and 400 kDa were more intense from cells grown with naphthalene (Fig. 3, Suppl. Table 1).

The three regions of the gels showing the differentially induced bands with sizes of about 750, 560, and 400 kDa were excised and analyzed by label-free quantitative proteomics (Suppl. Table 2). The 750 kDa band contained a total of 288 proteins, 29 of which were differentially abundant with a ratio of  $\geq 2$  in cells grown with naphthalene compared to 2-methylnaphthalene (Suppl. Table 2A). The ratio was defined as the protein ratio calculated by normalized abundances from naphthalene to 2-methylnaphthalene samples. Furthermore, for identification a

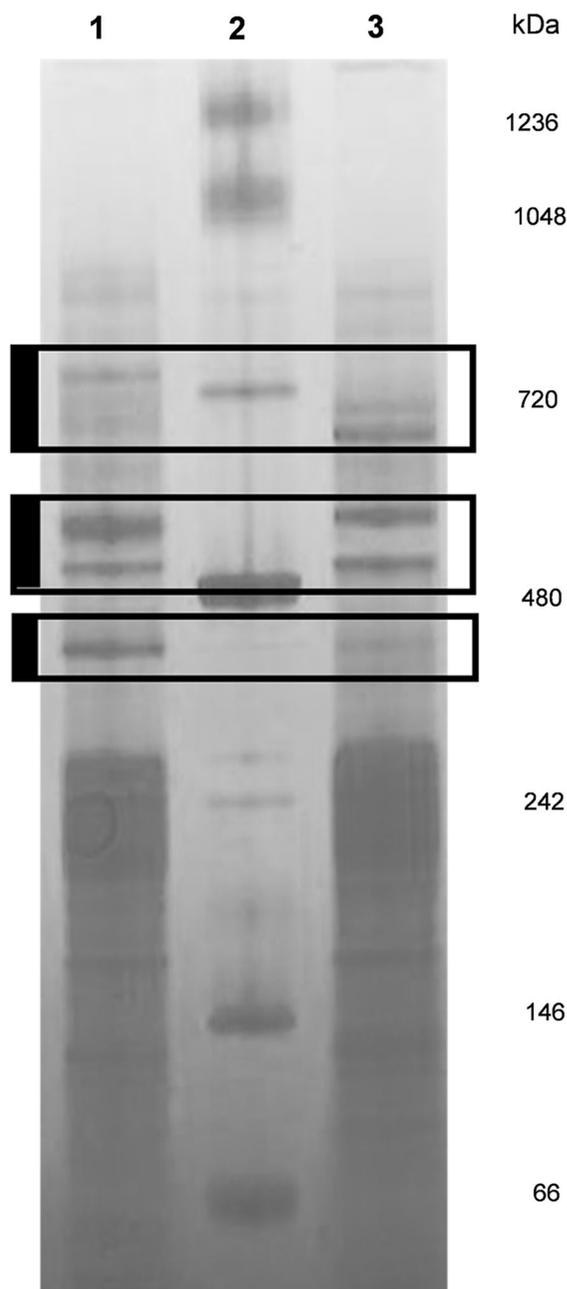


**Fig. 2 a** Organization of the gene cluster encoding enzymes potentially involved in the initial carboxylation reaction of naphthalene degradation in the sulfate-reducing *Deltaproteobacteria* N47 (GI:308273914 to GI:308273898). Open reading frames which products show similarity to UbiD-like carboxylases are filled in black, non-carboxylase-like subunits are shown in grey. Genes which are co-transcribed are indicated by a pattern fill. Arrows represent primers used for transcriptional analysis by gene-specific PCR. Annotated function of the gene products: 1, putative phenylphosphate carboxylase, alpha subunit; 2, putative phenylphosphate carboxylase, gamma subunit; 3, MRP, Fer4\_NifH superfamily; 4, ParA/MinD ATPase like, MRP, Fer4\_NifH superfamily; 5, UbiD family decarboxylase; 6, conserved hypothetical protein; 7, UbiD family decarboxylase; 8, conserved hypothetical protein; 9, UbiD family

p-value of  $\leq 0.01$  and a confidence score of  $> 30$  were applied as cut-off. In all three bands excised from the gel, the most abundant polypeptides were UbiD carboxylase-like gene products of the putative carboxylase gene cluster (Suppl. Table 2A–C). Almost

decarboxylase; 10 & 11, HAD hydrolase; 12, membrane protein involved in aromatic hydrocarbon degradation; 13, IS4 transposase; 14, succinate dehydrogenase and fumarate reductase iron-sulfur protein; 15, putative succinate dehydrogenase flavoprotein subunit; 16, UbiD family decarboxylase; 17, pyridoxamine 5'-phosphate oxidase family protein. **b** Transcriptional analysis by RT- and subsequent gene-specific PCR. Results of gene-specific PCRs using three different templates per primer set. A 1.5% agarose gel stained with GelRed was used. M, DNA ladder; numbers, name of used primer set; lane 1, gene-specific PCR on cDNA template (addition of reverse transcriptase during cDNA synthesis); lane 2, control for potential genomic DNA contamination in cDNA template (no addition of reverse transcriptase during RT-PCR); lane 3, gene-specific PCR on genomic DNA (positive control). Random hexamer primers were used in the RT-PCR

all of the gene products encoded in the naphthalene carboxylase operon were detected in the 750 kDa protein complex. Only the gene product N47\_K27460 was not found. This gene product showed high similarity to PpcX, a phenol-induced UbiD like



**Fig. 3** Differential protein induction analysis of N47 on blue native PAGE. Boxes indicate slices of the gel which were cut out for proteomic analysis. Lane 1, cell-free extract of naphthalene-grown N47 cells; lane 2, molecular mass standard (1236–66 kDa); lane 3, cell-free extract of 2-methylnaphthalene-grown N47 cells

protein from *T. aromatica* and *A. aromaticum*, which is encoded in the phenol gene cluster but is not part of the phenylphosphate carboxylase complex (Schühle

and Fuchs 2004). Three UbiD-like proteins N47\_K27540 (52.4 kDa), N47\_K27500 (57 kDa) and N47\_K27480 (54.5 kDa) were part of the naphthalene carboxylase complex. Furthermore, N47\_K27530 (9.8 kDa) which contained no conserved domains and the ParA/MinD ATPase-like proteins N47\_K27510 (29.4 kDa) and N47\_K27520 (33.6 kDa) were more abundant in naphthalene-grown N47 cells and therefore are proposed to represent subunits of the complex. Moreover, N47\_K27490 (9.2 kDa) as well as N47\_K27470 (16.6 kDa) were identified and both annotated as hypothetical proteins.

In the 560 kDa band, 23 proteins among 133 detected were differentially abundant with naphthalene as substrate (Suppl. Table 2B). The most abundant and differentially induced proteins were the carboxylase-like subunits N47\_K27540, N47\_K27500, N47\_K27480 and N47\_K27530 together with the heterodisulfide reductase subunits (N47\_J04330, N47\_J04340, N47J04350 and N47\_G39330, N47\_G39340). Moreover, N47\_E46900 a protein annotated as K(+)-insensitive pyrophosphate-energized proton pump was detected and differentially produced.

In the 400 kDa band, 10 polypeptides out of 125 detected were differentially abundant in naphthalene-grown cells (Suppl. Table 2C). The most abundant and at the same time differentially produced proteins were represented by the carboxylase-like proteins N47\_K27500 and N47\_K27480. The ATPase-like protein N47\_K27520 as well as proteins annotated as heterodisulfide reductase subunits (N47\_J04330, N47\_J04340, N47J04350) were differentially abundant. The K(+)-insensitive pyrophosphate-energized proton pump N47\_E46900 was again identified.

#### In vitro protein interaction assays

In vitro interaction pull down assays were performed in order to identify subunits of naphthalene carboxylase interacting with the phenyl phosphate carboxylase (ppc)-like polypeptide N47\_K27540 (Fig. 4, Suppl. Fig. 1), encoded by the first gene of the naphthalene carboxylase operon (Fig. 2a).

The most abundant proteins interacting with GST-NcA from naphthalene compared to 2-methylnaphthalene-grown cells were the carboxylase-like proteins N47\_K27500 and N47\_K27480 with enrichment factors of 23.6 and 31.0, respectively (Table 2A, B),

and the two ATPase-like proteins N47\_K27520 and N47\_K27510 with enrichment factors of 50.4 and 56.2 (Table 2D). These proteins were also identified in the native conformation of the naphthalene carboxylase complex analyzed with blue native gels. Furthermore, the carboxylase-like protein N47\_K27460 showed interaction with GST-NcA. The corresponding gene is also part of the naphthalene carboxylase operon but the polypeptide could never be detected in the blue native gels. Moreover, N47\_K27400 a succinate dehydrogenase-like flavoprotein was identified in one of the sliced bands with an enrichment factor of 44.6 (Suppl. Table 1).

The genes of several other proteins which are not part of the naphthalene carboxylase gene cluster also interacted in the pull-down assay (Table 2). For example, N47\_A07150, a FAD-binding domain protein, was enriched with a factor of 103.2 (Table 2C). Moreover, the ATPase-like protein N47\_K27600, which was also detected in the 750 kDa band of the blue native gels, interacted with an enrichment factor

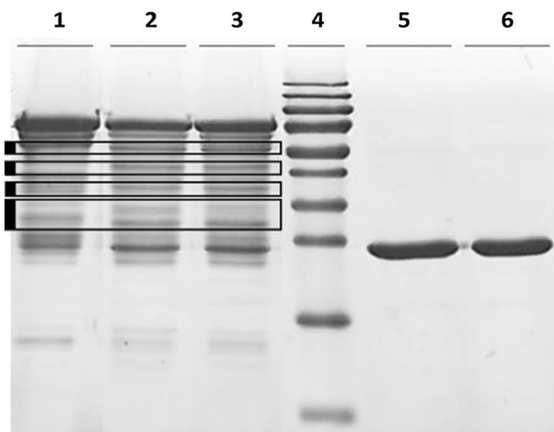
of up to 84.9 (Table 2C, D). Furthermore, a sulfate adenylyltransferase (N47\_J04360) showed an enrichment factor of 30.6 (Table 2D). This protein was not identified in the blue native gels but it is encoded in close vicinity to the putative heterodisulfide reductase subunits N47\_J04330, N47\_J04340 and N47\_J04350, which showed up in naphthalene grown cells in the blue native gels.

## Discussion

The carboxylation of naphthalene is an unprecedented biochemical reaction. In proteogenomic and transcriptomic analyses a conserved gene cluster encoding carboxylase-like proteins was identified in the sulfate-reducing culture N47 (Bergmann et al. 2011b; DiDonato et al. 2010). Due to differential abundance with naphthalene, the gene products were proposed to be involved in the carboxylation of naphthalene. In the present study, we analyzed the structure of the respective operon and the composition of the naphthalene carboxylase protein complex.

### Organization of the naphthalene carboxylase operon

The transcriptional analysis of the putative naphthalene carboxylase gene cluster revealed that the naphthalene carboxylase is transcribed as one operon ranging over 9 genes from N47\_K27540 to N47\_K27460. It contains the genes N47\_K27540, N47\_K27500, N47\_K27480 and N47\_K27460 encoding putative UbiD carboxylase like subunits. In *E. coli*, UbiD catalyzes the decarboxylation of an isoprenylated 4-hydroxybenzoate derivative in ubiquinone biosynthesis (Cox et al. 1969; Meganathan 2001). The identified subunits in N47 showed up to 46% sequence identity to the alpha subunit of the phenylphosphate carboxylase PpcA of *T. aromatica* and up to 48% to a putative anaerobic benzene carboxylase AbcA of the iron-reducing enrichment culture BF (Abu Laban et al. 2010; Schühle and Fuchs 2004). Two ParA/MinD ATPase-like proteins, N47\_K27510 and N47\_K27520, are encoded in the putative naphthalene carboxylase operon. Proteins with a ParA/MinD domain are involved in plasmid segregation and cell division (Gerdes et al. 2000) but are also found in iron-sulfur cluster biosynthesis



**Fig. 4** SDS-PAGE of a protein interaction assay with GST-NcA in cell-free extracts of N47 grown with naphthalene and 2-methylnaphthalene. Lane 1, purified GST-NcA coupled to glutathione Sepharose beads (input sample, 30  $\mu$ L); lane 2, purified GST-NcA coupled to glutathione Sepharose after 4 h of incubation with cell-free extract of naphthalene-grown N47 cells (30  $\mu$ L); lane 3, purified GST-NcA coupled to glutathione Sepharose beads (lane 1) after 4 h of incubation with cell-free extract of 2-methylnaphthalene-grown N47 cells (30  $\mu$ L); lane 4, molecular mass standard (170, 130, 100, 70, 55, 40, 35, 25, 15 and 10 kDa); lane 5, GST coupled to glutathione Sepharose beads after 4 h of incubation in cell-free extracts of naphthalene-grown N47 cells; lane 6, GST coupled to glutathione Sepharose beads after 4 h of incubation in cell-free extracts of 2-methylnaphthalene-grown N47 cells. Vertical boxes indicate parts of the gel which were cut and send for proteomic analysis

**Table 2** List of GST-NcA interacting polypeptides with an enrichment  $\geq 15$  and a p-value  $\leq 0.01$  detected in a pull-down assay and identified by LC-MS/MS

UniProtKB entry name	Peptides used for quantitation	Confidence score	Anova (p)	Description	Annotated function	ratio Naph/2MN
A.						
E1YIW4_9DELTA	18	1162	0.000633	<b>N47_K27480</b>	<b>UbiD family decarboxylase</b>	<b>31.0</b>
E1YIW6_9DELTA	13	819	0.000597	<b>N47_K27500</b>	<b>UbiD family decarboxylase</b>	<b>23.6</b>
E1YIV6_9DELTA	14	855	0.001726	<b>N47_K27400</b>	<b>Putative succinate dehydrogenase flavoprotein</b>	<b>44.6</b>
E1YAB2_9DELTA	4	217	0.000485	N47_H23280	Fumarate hydratase class I, anaerobic	28.0
E1YAB2_9DELTA	4	217	0.000485	N47_F14190	Uroporphyrin-III C-methyltransferase	155.1
E1YIW2_9DELTA	2	89	0.026798	<b>N47_K27460</b>	<b>UbiD family decarboxylase</b>	<b>5171.5<sup>a</sup></b>
B.						
E1YKJ5_9DELTA	15	745	0.005969	N47_E41400	Acetyl-CoA acetyltransferase/Thiolase	17.1
E1YIW4_9DELTA	10	608	0.001111	<b>N47_K27480</b>	<b>UbiD family decarboxylase</b>	<b>16.7</b>
C.						
E1YIX6_9DELTA	6	305	0.006848	N47_A07150	FAD binding domain protein	103.2 <sup>a</sup>
E1Y816_9DELTA	5	213	0.002488	N47_K27600	MinD ATPase like, Fer4_NifH superfamily	46.0
E1YFW1_9DELTA	2	65	0.009352	N47_F14520	CRISPR-associated protein	21.3
D.						
E1YIW7_9DELTA	21	1147	0.00746	<b>N47_K27510</b>	<b>MinD ATPase like, Fer4_NifH superfamily</b>	<b>56.2</b>
E1YIW8_9DELTA	9	866	0.00195	<b>N47_K27520</b>	<b>Fer4_NifH superfamily</b>	<b>50.4</b>
E1YFW1_9DELTA	9	392	0.00015	N47_J04360	Sulfate adenyltransferase	30.6
E1YIX6_9DELTA	7	323	0.00336	N47_K27600	MinD ATPase like, Fer4_NifH superfamily	84.9
E1YFR3_9DELTA	4	153	0.00014	N47_J03880	Putative electron transfer flavoprotein, FixB	25.8

The proteins shown in bold letters, correspond to the ones which are encoded in the naphthalene carboxylase gene cluster. A. SDS gel region with a size of 60–50 kDa. B. SDS gel region with a size of 45–40 kDa. C. SDS gel region with a size of 38–35 kDa. D. SDS gel slice with protein size of 35–28 kDa

<sup>a</sup>Coefficient of variation (CV %)  $\geq 50$ ; in 2-methylnaphthalene samples

(ApbC/Nbp35 homologs). More recent studies showed that the ParA/MinD family is further involved in the positioning of large cytoplasmic protein complexes such as carboxysomes or chemotaxis clusters (Lutkenhaus 2012). A similar function of the proteins N47\_K27510 and N47\_K27520 is possible for the quite large carboxylase complex. Additionally, the genes *N47\_K27530*, *N47\_K27490*, and *N47\_K27470* annotated as hypothetical proteins are encoded within the naphthalene carboxylase operon. Due to the small

size and the lack of conserved domains these genes might encode linker proteins of the naphthalene carboxylase complex.

Two subsequent genes were not part of the operon, *N47\_K27450* and *N47\_K27440*. They are similar to genes coding for HAD-like polypeptides (cd01427; conserved domain database, National Center for Biotechnology Information) (Bergmann et al. 2011a; Selesi et al. 2010). The HAD-like superfamily is composed of hydrolases comprising phosphatases,

among others. For example, the HAD-like protein PpcC is part of the phenylphosphate carboxylase in *Thauera aromatica*, a key enzyme in anaerobic phenol degradation (Schühle and Fuchs 2004). PpcC was proposed to bind and dephosphorylate the substrate phenylphosphate. As there was no indication for such a phosphorylated intermediate in the carboxylation of naphthalene, a phosphatase-like subunit might not be required in the functional naphthalene carboxylase complex.

Furthermore, the naphthalene carboxylase gene cluster was surrounded by viral genes in N47 (N47\_K27300, N47\_K27370, N47\_K27640 putative transposases). Considering the high sequence similarity of the gene clusters and the presence of genetic elements which are involved in gene mobility, horizontal gene transfer of the genes for naphthalene degradation is likely.

#### Naphthalene carboxylase complex and its subunits

In comparative native proteomic studies with blue native gels, three prominent naphthalene-induced bands with a size of around 750, 560, and 400 kDa were identified. Almost all polypeptides detected in the 750 kDa band, which were differentially abundant with naphthalene versus 2-methylnaphthalene as growth substrate, were encoded in the naphthalene carboxylase operon. The prominent bands at 560 and 400 kDa contained less polypeptides indicating dissociated sub complexes. This is supported by the fact that the 750 kDa protein complex was only detectable from freshly harvested cells grown with naphthalene as carbon and electron source. The 750 kDa band was absent in 2-methylnaphthalene-grown cells where naphthalene carboxylase activity could hardly be detected. Thus, our indirect evidence indicates that the 750 kDa band most likely represents the native naphthalene carboxylase complex. At the present state however, we were not able to purify an active naphthalene carboxylase enzyme complex. Considering the proteins contained in the 750 kDa band, the native naphthalene complex putatively consists of three UbiD-like carboxylase subunits N47\_K27540 (52.4 kDa), N47\_K27500 (57 kDa) and N47\_K27480 (54.4 kDa), the ParA/MinD ATPase-like proteins N47\_K27510 (29.4 kDa) and N47\_K27520 (33.6 kDa). Furthermore, the subunits N47\_K27530 (9.8 kDa), N47\_K27490 (9.2 kDa), and

N47\_K27470 (16.6 kDa) are part of the naphthalene carboxylase complex as putative linker proteins.

Therefore, all subunits encoded in the naphthalene carboxylase operon, except for the carboxylase-like subunit N47\_K27460, seem to be part of the naphthalene carboxylase complex. Our blue native gel experiments indicated that the naphthalene carboxylase complex has a size of around 750 kDa in native conformation. If all the subunits encoded in the naphthalene carboxylase operon would be involved in the complex only once, it would have a size of around 314 kDa. Without the UbiD-like subunit N47\_K27460, which might not be part of the naphthalene carboxylase, the complex is around 263 kDa.

Similar to the naphthalene carboxylase complex, the phenylphosphate carboxylase from *T. aromatica* represents a new type of UbiD-like carboxylase (Schühle and Fuchs 2004). The purified enzyme is composed of four subunits- of two UbiD-like proteins ( $\alpha\beta$ , 54 and 53 kDa), a hypothetical protein ( $\gamma$ , 10 kDa) and a phosphatase-like protein ( $\delta$ , 18 kDa). The authors suggested a trimeric structure ( $\alpha\beta\gamma\delta$ )<sub>3</sub> for this carboxylase complex.

Assuming a similar trimeric structure analogous to the phenylphosphate carboxylase, the naphthalene carboxylase complex would have a molecular mass around 943 kDa or 787 kDa with or without N47\_K27460, respectively. The latter matches the observed molecular mass of 750 kDa.

#### Interaction of naphthalene carboxylase with other proteins

The carboxylase-like protein N47\_K27460 was encoded within the naphthalene carboxylase operon and an interaction with the other carboxylase-like proteins was shown in a pull-down assay, but it was not detected within the naphthalene-induced bands on BNPs. The interaction between the carboxylase complex and N47\_K27460 could have either been destroyed during sample preparation or N47\_K27460 is not part of the naphthalene carboxylase complex. The UbiD-like protein N47\_K27460 was also not detected on 2D PAGEs (Bergmann et al. 2011b). In a phylogenetic tree of UbiD/PpcA-like subunits involved in anaerobic degradation of aromatic compounds, the protein formed a cluster with PpcX from *T. aromatica* and *A. aromaticum* strain EbN1 (Suppl. Fig/ 2, Suppl. Table 3). PpcX is a

phenol-induced UbiD-like protein and its gene is encoded within the phenol gene cluster but it is not part of the phenylphosphate carboxylase complex (Schühle and Fuchs 2004).

Moreover, proteins which are annotated as heterodisulfide reductases were identified within the naphthalene-induced bands in BNPs. Heterodisulfide reductase-like proteins are also involved in anaerobic degradation of monoaromatic compounds. The BamDE subunits of the class II benzoyl-CoA reductase of *Geobacter metallireducens* share sequence similarity to heterodisulfide reductases (Fuchs et al. 2011; Kung et al. 2009). Here, benzoyl-CoA is reduced in an electron bifurcation process via ferredoxin<sub>red</sub> coupled to the exergonic reduction of NAD(P)<sup>+</sup> using another molecule of the same ferredoxin<sub>red</sub>. However, involvement of a reduction step to carboxylate naphthalene is not implicit and the appearance of heterodisulfide reductase-like proteins cannot be explained at the present state of knowledge.

#### UbiD-like (de)carboxylases involved in anaerobic degradation of aromatic compounds

In *E. coli*, the enzymes UbiD and UbiX facilitate the decarboxylation of 3-octaprenyl-4-hydroxybenzoate to 2-octaprenylphenol in ubiquinone biosynthesis (Gulmezian et al. 2007; Zhang and Javor 2003). Recently, UbiX and UbiD were identified as flavin mononucleotide (FMN) containing proteins working in a two-component system. UbiX synthesizes a prenylated FMN by adding a non-aromatic fourth ring to the flavin mononucleotide. UbiX enzyme activity is independent from metal cofactors and utilizes dimethylallyl-monophosphate as prenylating substrate (Payne KAP 2015; White et al. 2015). The prenylated FMN<sub>pr</sub> is transferred to UbiD, which performs the decarboxylation reaction.

In the N47 genome, a UbiX-like gene was not part of the naphthalene carboxylase cluster but two completely identical copies *N47\_B20630* and *N47\_H21380* were encoded elsewhere in the genome. In proteomic analyses, differentiation between these two proteins was not possible but one or both were differentially induced in cells grown with naphthalene.

At the present state of knowledge, the presence of a cofactor in the naphthalene carboxylase remains elusive. However, the amino acid sequences suggested potential binding sites for FMN in the carboxylase-

like subunits N47\_K27540, N47\_K27500, N47\_K27480, N47\_K27460 (GO0010181; UniProt-GO annotation database, European Bioinformatics Institute) and nucleotide-binding domains in the ATPase-like proteins N47\_K27510 and N47\_K27520 (cd02037; conserved domain database, National Center for Biotechnology Information).

Various UbiD-like (de)carboxylases involved in the degradation of aromatic compounds have been identified in recent years (Fig. 1). A gene cluster putatively encoding enzymes involved in the degradation of terephthalate was identified. One gene product showed similarity to an UbiD-like protein and was suggested to be a terephthalate decarboxylase (Lykidis et al. 2011; Wu et al. 2013). Therefore, analogous to the syntrophic, terephthalate-degrading *Pelotomaculum* spp. the authors proposed that terephthalate is first activated by a ligase and subsequently decarboxylated by a complex of an UbiD-like protein and a phenylphosphate  $\gamma$ -subunit-like protein to produce benzoyl-CoA. The UbiD-like subunit showed more than 40% sequence identity to N47\_K27540. This reaction could recently be shown with phthalate as substrate in cell extracts of the strains *Thauera*, *Azoarcus* and '*Aromatoleum*' (Ebenau-Jehle et al. 2017; Junghare et al. 2016).

## Conclusion

Non-substituted PAHs like naphthalene need an initial activation in order to be further degraded. This first important reaction is carried out by a carboxylase in the case of naphthalene (Mouttaki et al. 2012). The naphthalene carboxylase complex consists of eight proteins, which are encoded in one operon. The protein N47\_K27460 is also encoded in the operon but is not part of the complex. Comparison to the phenylphosphate decarboxylase suggests a similar trimeric structure for the naphthalene carboxylase complex with a size of 750 kDa. Three of the subunits building the protein complex were identified as UbiD-like proteins, putting the naphthalene carboxylase in the growing family of de-/carboxylases using a prenylated flavin cofactor.

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