Differential protein expression during growth on linear versus branched alkanes in the obligate marine hydrocarbon-degrading bacterium *Alcanivorax borkumensis* SK2T

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Summary

*Alcanivorax borkumensis* SK2T is an important obligate hydrocarbonoclastic bacterium (OHCB) that can dominate microbial communities following marine oil spills. It possesses the ability to degrade branched alkanes which provides it a competitive advantage over many other marine alkane degraders that can only degrade linear alkanes. We used LC–MS/MS shotgun proteomics to identify proteins involved in aerobic alkane degradation during growth on linear (n-C14) or branched (pristane) alkanes. During growth on n-C14, *A. borkumensis* expressed a complete pathway for the terminal oxidation of n-alkanes to their corresponding acyl-CoA derivatives including AlkB and AlmA, two CYP153 cytochrome P450s, an alcohol dehydrogenase and an aldehyde dehydrogenase. In contrast, during growth on pristane, an alternative alkane degradation pathway was expressed including a different cytochrome P450, an alcohol oxidase and an alcohol dehydrogenase. *A. borkumensis* also expressed a different set of enzymes for β-oxidation of the resultant fatty acids depending on the growth substrate utilized. This study significantly enhances our understanding of the fundamental physiology of *A. borkumensis* SK2T by identifying the key enzymes expressed and involved in terminal oxidation of both linear and branched alkanes. It has also highlights the differential expression of sets of β-oxidation proteins to overcome steric hinderance from branched substrates.

Introduction

*A. borkumensis* SK2T is a model strain of a group of organisms known as obligate hydrocarbonoclastic bacteria (OHCB) which grow on a highly restricted spectrum of substrates, predominantly alkanes and their derivatives, with carbon chain length from n-C9 to at least n-C32 (Schneiker et al., 2006; Yakimov et al., 1998; 2007; Naether et al., 2013). *Alcanivorax* spp. are found in low abundances in unpolluted marine environments but can multiply and grow rapidly in oil-polluted waters, where they can constitute 80%–90% of the microbial community (Harayama et al., 1999; Kasai et al., 2002; Syutsubo et al., 2001). Previous reports have also shown *Alcanivorax* spp. become abundant in field and mesocosm experiments involving the addition of nitrogen and phosphorus fertilizers to stimulate microbial degradation of oil (Cappello et al., 2007; McKew et al., 2007).

*Alcanivorax* spp. has a cosmopolitan distribution with isolations or detection of its 16S rRNA gene sequences from many different oil-impacted marine environments (Yakimov et al., 2007). *Alcanivorax borkumensis* SK2T (DSM 11573) was the first OHCB to have its genome sequenced revealing a plethora of genes accounting for its wide hydrocarbon substrate range and efficient oil-degradation capabilities (Reva et al., 2008; Schneiker et al., 2006). Analysis revealed proteins required to oxidize n-alkanes up to the corresponding acyl-CoA derivative are coded by the *alkSBGJH* operon which has >80% amino acid similarity to the corresponding well-characterized alkane degradation components in *Pseudomonas putida* Gpo1 (Kok et al., 1989; van Beilen et al., 2001; van Beilen et al., 2003; van Beilen et al., 2004). However, genomes only give insight into the genetic potential of an organism. Several transcriptomic studies hint even more at this potential. A microarray-based study led to the identification of up-regulated genes predicted to be involved in the terminal oxidation of n-hexadecane (Sabirova et al., 2011). These included genes coding for two alkanes monoxygenases (*alkB1* and *alkB2*), three flavin-binding monoxygenases (*ABO_0282, ABO_1097, ABO_2107*),

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an alcohol dehydrogenase (ABO_2483) and an aldehyde reductase (AKR1A1/ABO_2414). A subsequent microarray analysis identified genes that distinctly respond to solvent stress by the addition of 1-octanol, a known toxic intermediate of \( n \)-alkane degradation (Naether et al., 2013). In accordance with Sabirova et al., 2011, the expression level of \( \text{alkB1} \) (ABO_2707) was increased during growth on the \( n \)-hexadecane. In contrast, RNA-sequencing surprisingly showed that only 1% of genes were up-regulated while \( A. \text{borkumensis} \) was growing on \( n \)-dodecane and none of these were involved in alkane metabolism (Barbato et al., 2016). The mRNA abundances presented in such studies may not directly correlate to the amount of protein within a cell, due to many factors. Large changes in gene expression may not necessarily result in similar changes in translated proteins as there can be a requirement for gene co-regulation, controls on post-translational modification or protein degradation, or protein translation may be highly dependent on translation initiation features and recruitment of ribosomes (Vogel and Marcotte, 2012). The proteins are the functional molecules in cells and it is the proteome, rather than the genome or transcriptome, that is, therefore most related to the phenotype of an organism, yet there is little knowledge or quantification of protein biosynthesis in \( A. \text{canivorax} \) during alkane degradation. A previous study of \( A. \text{borkumensis} \) using two-dimensional gel electrophoresis identified up-regulation of two alkane hydroxylases \( \text{AlkB1} \) and \( \text{AlkB2} \) (Sabirova et al., 2006), which have been shown to oxidize medium-chain alkanes in the range of \( n \)-C\(_5\)-n-C\(_{12}\) and \( n \)-C\(_6\)-n-C\(_{16}\) (van Beilen et al., 2004), respectively, three P450 cytochromes, and a flavin-binding monoxygenase with a strong similarity to the \( \text{AlmA} \) enzyme, which hydroxylates long-chain alkanes (\( n \)-C\(_{22}\)-n-C\(_{36}\)) in \( A. \text{dieselolei} \) and \( A. \text{hongdengensis} \) (Liu et al., 2011; Wang and Shao, 2012b, Wang and Shao, 2014). A downside of this proteomic technique is that it is only semiquantitative and often a relatively small number of proteins are detected (e.g. only 97 proteins were excised, sequenced and identified in Sabirova et al., 2006).

The ubiquity of \( A. \text{canivorax} \) and its success as an alkane degrader may result from the ability to also efficiently utilize branched alkanes as the sources of carbon, which provides a competitive advantage over other \( \text{OHCB} \) which are unable to utilize these more difficult to degrade substrates (Pirnik et al., 1974; Hara et al., 2003). In the previous reports, branched alkanes, such as pristane and phytane, which are the abundant branched alkanes present in many crude oils, induce the expression of \( \text{AlkB1} \) and the long-chain alkane monoxygenase \( \text{AlmA} \) in \( A. \text{hongdengensis} \) A-11-3 and \( A. \text{dieselolei} \) B-5 (Liu et al., 2011; Wang and Shao, 2012b). Sevilla et al., 2017 demonstrated through transcriptional fusions of the three cytochrome P450 genes from \( A. \text{borkumensis} \) to a green fluorescent protein reporter gene and fluorescence assays that expression of P450-3 was higher on pristane compared to \( n \)-alkanes or pyruvate. However our knowledge of the proteins involved branched alkane degradation remains very limited. This study is first to compare the differences in protein expression during linear and branched alkane degradation in \( A. \text{borkumensis} \) SK2\(^T\). It provides insights into the metabolic pathways in this environmentally relevant hydrocarbon degrader.

We quantified changes in the proteome using liquid chromatography–tandem mass spectrometry (LC–MS/MS) identifying proteins significantly up-regulated, while growing on either linear alkanes (tetradecane, \( n \)-C\(_{14}\)) or pristane (the branched alkane 2,6,10,14-Tetramethylpentadecane) looking at the key differences in the expression patterns on each substrate.

### Results

**Overview of LC–MS/MS shotgun proteomic analysis**

Nine LC–MS/MS runs were performed consisting of three independent biological replicates of three treatments (linear alkane (\( n \)-C\(_{14}\)), branched chain alkane (pristane), non-hydrocarbon control (pyruvate)) resulting in 112,095 spectral counts that were assigned to 1309 proteins, representing 48% of the total protein-coding genes on the \( A. \text{borkumensis} \) SK2\(^T\) genome (Supporting Information Table S1). Over half (52%) of the spectral counts were assigned to the 100 most abundantly detected proteins, and 90% were assigned to the 500 most abundant proteins. The remaining 809 proteins (representing 7% of the total normalized spectral counts) were detected with only a very low number of spectral counts.

A total of 381 proteins were significantly differentially expressed during growth on \( n \)-C\(_{14}\) compared to pyruvate, with 80% of these being up-regulated on \( n \)-C\(_{14}\) (Fig. 1A). In contrast, only 23 proteins were differentially expressed during growth on pristane compared to pyruvate, 39% of which were up-regulated on pristane (Fig. 1B). Between the linear (\( n \)-C\(_{14}\)) and branched (pristane) chain alkanes, 288 proteins were significantly differentially expressed, with 91% of these being up-regulated on \( n \)-C\(_{14}\) (Fig. 1C). Overall the total proteomes differed markedly between the three growth substrates with highly similar proteomes between replicates (Fig. 1D).

**Terminal oxidation of linear \( n \)-alkanes**

Several proteins involved in terminal oxidation of linear \( n \)-alkanes were significantly differentially expressed during growth on \( n \)-C\(_{14}\) compared to pyruvate (Fig. 2). The
expression of two P450 CYP153 cytochromes (ABO_0201/2288, \( P = 0.028 \)), which share the same amino acid (100% similarity) and their corresponding genes share 99.6% nucleotide identity, was three-fold and two-fold greater while growing on \( n\)-C14 and pristane, respectively (Fig. 2A). This suggests the proteins may have a broad substrate specificity being able to utilize both linear and branched alkanes or is constitutively expressed due to expression also on pyruvate. The cytochrome P450s are part of a putative operon made up of genes coding for a ferredoxin (ABO_0200, not detected), which shuttles electrons to the cytochrome P450 (ABO_0201/2288), an alcohol dehydrogenase (ABO_0202) and a ferredoxin reductase (ABO_0203) (Schneiker et al., 2006). The alcohol dehydrogenase, AlkJ2 (ABO_0202, \( P = 0.008 \)), which converts alcohols generated by the oxidation of an alkane to its corresponding aldehyde, was highly up-regulated during growth on \( n\)-C14 and pristane but absent on pyruvate (Fig. 2A). The expression of a ferredoxin reductase (ABO_0203, \( P = 0.049 \)), which oxidizes NAD(P)H to NAD(P)\(^+\) capturing electrons to transfer to the ferredoxin, was seven-fold and five-fold greater during growth on \( n\)-C14 and pristane, respectively, compared to pyruvate. Multiple alkane monoxygenases were also detected. The alkane 2-monooxygenase, AlkB2 (ABO_0122) was exclusively expressed only during growth on \( n\)-C14 (Fig. 2B). The expression of another putative monoxygenase (ABO_2107, \( P = 0.011 \)) was seven-fold greater on \( n\)-C14 than pristane with no expression on pyruvate (Fig. 2B). Domain analysis revealed the protein is a member of the FMO (flavin-binding monoxygenase)-like family (PF00743), which specializes in the oxidation of xenobiotics. The rubA (ABO_0163) and rubB (ABO_0162) genes code rubredoxin and rubredoxin reductase, respectively, and are arranged in a putative operon. RubA (ABO_0163), which transfers electrons to the alkane monooxygenase from the rubredoxin reductase, was not detected in this data set and the expression of RubB (ABO_0162), which transfers electrons from NAD(P)H to the rubredoxin, had 5-fold greater expression on \( n\)-C14 compared to pristane (Fig. 2B). Two aldehyde dehydrogenases that catalyse the oxidation of the aldehyde produced from the alcohol dehydrogenase (ABO_0962/ABO_2414) were significantly (\( P < 0.0001 \)) differentially expressed on \( n\)-C14 (Fig. 2B). The combination of these alkane monoxygenases or CYP153 cytochrome P450 with an alcohol dehydrogenase and an aldehyde dehydrogenase makes up two alternative
pathways for the terminal oxidation of \( n \)-alkanes to their corresponding acyl-CoA derivatives (Fig. 2A and B).

**Terminal oxidation of the branched alkanes**

The expression of an AlmA-type monooxygenase (ABO_0282) was only present in \( n \)-C14 and pristane (Fig. 3). It has a 90\% identity to the long-chain flavin-binding monooxygenase AlmA from *Bacillus halodurans* (AGW21778.1) and a 63\% identity to AlmA from *A. dieselolei* B-5 (B5T_02052). In addition to the cytochrome P450 (ABO_0201/ABO_2288) as previously mentioned, another cytochrome P450 (ABO_2414) was expressed only when growing on pristane (Fig. 3). A GMC oxidoreductase family protein (ABO_1174, \( P = 0.010 \)) was exclusively expressed during growth on pristane and predicted as an alcohol oxidase in I-TASSER and TM-align, which identified an alcohol oxidase from *Pichia pastoris* (5i68A) in PDB as structurally close to ABO_1174 (TM score-0.822). The FunFam assignment

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in CATH for matching regions to the ABO_1174 identified the long-chain-alcohol oxidase FAO1 (Evalue-2.5e−60). The protein is FAD binding (GO:0050660) suggesting it is an alcohol oxidase, which oxidize primary alcohols into a primary aldehydes and hydrogen peroxide, as they require flavin-based cofactors, while alcohol dehydrogenases require NAD-based cofactors. An alcohol dehydrogenase (ABO_0061, P = 0.041) was also exclusively expressed during growth on pristane. This suggests A. borkumensis can use either and alcohol oxidase or dehydrogenase to convert pristanol to pristanal. No aldehyde dehydrogenase was significantly differentially expressed during growth on pristane, but ABO_0087 was expressed on all substrates tested. The combination of these enzymes makes up a complete pathway for the terminal oxidation of branched alkanes (Fig. 3B).

**β-Oxidation of acyl-CoA derivatives**

The fatty acids generated by oxidation of n-alkanes are further metabolized by β-oxidation. The β-oxidation proteins detected have putatively been assigned functions for their potential role in straight-chain alkane degradation (Fig. 4 A, C, E and G, left side) or branched-chain alkane degradation (Fig. 4B, D, F and H, right side) based on their pattern of expression. Before a fatty acid can enter any metabolic pathway, it must first be activated to form acyl-CoA. This reaction is catalysed by acyl-CoA synthetases, generating AMP. No acyl-CoA synthetase was differentially expressed on n-C14, compared to pristane. ABO_0985 was initially annotated as an acid thiol ligase but a BLASTP search found it is identical (100% identity) to long-chain acyl CoA synthetases in multiple *Alcanivorax* sp. (97CO-5; 97CO-6; NBRC 101098). This protein was expressed equally across the three growth
substrates with high spectral counts (>75, Fig. 4A) and could potentially be constitutively expressed, suggesting it can potentially use branched and linear fatty acids as substrates. Two acyl-CoA synthetases potentially catalysing this reaction, fadD/ABO_0367 ($P = 0.041$) and alkK/ABO_2748 ($P = 0.046$) were differentially expressed on pristane compared to $n$-C$_{14}$ but also similarly expressed on pyruvate (Fig. 4B). Acyl-CoA dehydrogenase creates a double bond between the second and third carbons down from the CoA group on acyl-CoA producing trans-delta 2-enoyl CoA and in the process, uses FAD as an electron acceptor reducing it to FADH$_2$. Four acyl-CoA dehydrogenases were significantly differentially expressed during growth on pristane, including ABO_0571 ($P = 0.024$), ABO_1264 ($P = 0.023$), acdA/ABO_2223 ($P = 0.046$) and ABO_2453 ($P = 0.017$) (Fig. 4D), while three other acyl-CoA dehydrogenases were significantly differentially expressed during growth on $n$-C$_{14}$ (Fig. 4C). These include fadE/ABO_2098 ($P < 0.0001$), ABO_2102 ($P = 0.004$) and ABO_2739 ($P = 0.001$). Enoyl CoA hydratase removes the double bond in trans-delta 2-enoyl CoA adding a hydroxyl group to the third carbon down from the CoA group and a hydrogen on the second carbon down from the CoA group producing L-3-hydroxyacyl CoA differentially expressed on $n$-C$_{14}$ (E) and pristane (F). G and H. The FadAB2 (G) and FadAB (H) operon consisting of $\beta$-oxidation multifunctional enzyme complexes with 3-hydroxyacyl-CoA dehydrogenase (removes the hydrogen in the hydroxyl group added to L-3-hydroxyacyl CoA producing 3-ketoacyl CoA) and ketoacyl-CoA thiolase (attaches a CoA group on the third carbon down from the CoA group producing acetyl-CoA and acyl-CoA which is two carbons shorter) activities were differentially expressed on $n$-C$_{14}$ and pristane, respectively.
to the third carbon down from the CoA group and hydroxyl on the second carbon down from the CoA group producing L-3-hydroxyacyl CoA. Four enoyl CoA hydratases were significantly differentially expressed during growth on \( n-C_{14} \) compared to pristane (Fig. 4E), including/ABO_0526 \( (P = 0.005) \), ABO_1238 \( (P = 0.006) \), fadC/ABO_1645 \( (P < 0.001) \) and eno/ABO_2556 \( (P = 0.020) \). No specific enoyl CoA hydratases were differentially expressed on pristane. However, fadB \( (ABO_1566, P < 0.0001) \), which was up-regulated on pristane, from the \( fadAB \) operon codes the \( \alpha \)-subunit of a \( \beta \)-oxidation multifunctional enzyme complex possessing both 3-hydroxyacyl-CoA dehydrogenase and enoyl-CoA hydratase activities (Fig. 4F). 3-hydroxyacyl CoA dehydrogenase removes the hydrogen in the hydroxyl group added to L-3-hydroxyacyl CoA producing 3-ketoacyl CoA, using NAD as an electron acceptor reducing it to NADH. In the final step of \( \beta \)-oxidation ketoacyl-CoA thiolase attaches a CoA group on the third carbon down from the CoA group resulting in the formation of two molecules, an acetyl-CoA and an acyl-CoA that is two carbons shorter. Only one ketoacyl-CoA thiolase was detected \( (ABO_2452, P = 0.446) \), which was not expressed on pristane but was not significantly differentially expressed on either \( n-C_{14} \) and pristane due to detection of very low spectral counts. The \( fadAB \) operon consisting of \( fadB \) \( (ABO_1566, P < 0.0001) \) previously mentioned above and \( fadA \) \( (ABO_1567, P = 0.024) \), which codes for the \( \beta \)-subunit of the complex with 3-ketoacyl-CoA thiolase activity, was significantly differentially expressed on pristane (Fig. 4H). The \( fadAB2 \) operon consisting of \( fadB2 \) \( (ABO_1652, P = 0.011) \), which was differentially expressed on both \( n-C_{14} \) and pristane, and \( fadA \) \( (ABO_1653, P = 0.022) \), which was significantly differentially expressed on \( n-C_{14} \) (Fig. 4G).

Discussion

Terminal oxidation of linear \( n \)-alkanes

Bacteria degrading linear alkanes typically contain a non-heme alkane monoxygenase system including a membrane bound alkane monooxygenase, a rubredoxin and a rubredoxin reductase and/or a heme-thiolate cytochrome system including a soluble cytochrome P450, a ferredoxin and ferredoxin reductase (Teimoori et al., 2011; Nie et al., 2014). It is rather common to find bacteria that contain more than one alkane oxidation system as each have different substrate ranges or different induction patterns. \( A. borkumensis \) has two alkane monooxygenases on its genome, which are homologous to well characterized the alkane hydroxylase AlkB from \( Pseudomonas putida \) GPO1, which oxidizes alkanes from \( n-C_5 \) to \( n-C_{16} \) (Schneiker et al., 2006; van Beilen and Funhoff, 2007).

\( AlkB_1 \) \( (ABO_2707) \) and \( AlkB_2 \) \( (ABO_0122) \) oxidizes alkanes in the range of \( n-C_9 \) to \( n-C_{12} \) and \( n-C_9 \) to \( n-C_{16} \), respectively (Hara et al., 2004; van Beilen et al., 2004). \( AlkB_2 \) was exclusively expressed during growth on \( n-C_{14} \). Electrons are transferred to the AlkB active site via a rubredoxin. There are two rubredoxin genes, \( alkG \) \( (ABO_2708) \) and \( rubA \) \( (ABO_0163) \) in the \( A. borkumensis \) genome (Schneiker et al., 2006). Both rubredoxins were however not detected in this data set. Rubredoxin reductases oxidizes \( \text{NAD(P)}H \) to transfer electrons to rubredoxin reducing it allowing electron transfer to the monooxygenase. There is one rubredoxin reductase gene in the \( A. borkumensis \) genome, \( RubB \) \( (ABO_0162) \), which has been shown to reduce \( AlkB \) when both proteins were cloned and functionally overexpressed in \( E. coli \) (Teimoori et al., 2011). This activity compensates for a lack of \( AlkT \), also a rubredoxin reductase, missing in the \( A. borkumensis \) genome (Schneiker et al., 2006). \( RubB \) expression was greatest during growth on \( n-C_{14} \), although expression was observed while growing on all three substrates at levels that were not significantly different. \( alkB \)/\( alkB2 \) expression is strictly \( n \)-alkane dependent, as confirmed here by the expression of \( AlkB_2 \) only while growing on \( n-C_{14} \), but \( RubA/RubB \) are constitutively expressed in other bacteria (Geissdörfer et al., 1999; Tani et al., 2001; Marin et al., 2003). The unbalanced expression between monooxygenase (\( AlkB \)) to rubredoxin/rubredoxin reductase (\( RubA/RubB \)) suggest that the molecules of membrane-bound monooxygenases share a limiting number of rubredoxin and rubredoxin reductase molecules, which are soluble proteins that can probably associate and dissociate from the membrane-bound monooxygenase, as seen in \( P. aeruginosa \) (Marin et al., 2003).

\( A. borkumensis \) contains alternative enzyme systems to \( AlkB \) for terminal oxidation of alkanes (Figs. 2 and 3), including three CYP153 cytochrome P450s; \( ABO_0201 \) \( (P450-1)/P450-c) \), \( ABO_2288 \) \( (P450-2/P450-b) \) and \( ABO_2384 \) \( (P450-3) \) (Schneiker et al., 2006). The expression of \( ABO_0201 \) and \( ABO_2288 \) was higher on \( n-C_{14} \) than pyruvate, suggesting it can terminally oxidize medium-chain alkanes. \( ABO_2384 \) was not expressed on \( n-C_{14} \) suggesting it may act on different substrates. A similar trend has been observed in other bacteria with multiple CYP153s. For example, \( Sphingomonas \) sp. HNX200 possesses five CYP153 genes, three of which show activity towards \( n-C_7 \) to \( n-C_{10} \), whereas no affinity for these substrates was observed for the other two genes, suggesting that these genes are either pseudogenes or act on different substrates (Van Bogaert et al., 2011). P450-1 and P450-2 are phylogenetically grouped in one branch with CYP153A1 from \( Acinetobacter \) sp. EB104, which was experimentally confirmed to catalyse terminal alkane oxidation (Maier et al., 2001; Schneiker et al., 2006; van Beilen et al., 2006).
The coexpression of AlkB and CYP153 may cooperatively ensure that A. borkumensis can utilize a broader range of linear substrates.

The expression of an AlmA homologue (ABO_0282) was at least 4-fold greater during growth on n-C14 differentially expressed compared to pyruvate (Fig. 2). Although AlmA is a long-chain monooxygenase, it has been shown in other bacteria to have some activity on shorter alkanes but has much lower activity compared to growth on long-chain alkanes, for example, >n-C20. For example, almA gene expression of M. koreensis, M. alkaliphilus and M. lipolyticus responded to alkanes ranging in length from n-C9 up to n-C36 (Wang and Shao, 2012a). AlmA in A. dieselolei was shown to have activity against alkanes ranging in length from n-C10 up to n-C36. This suggests that n-C14 is within the substrate range of ABO_0282 and therefore the protein is expressed. However, if A. borkumensis was grown on longer alkanes the expression of ABO_0282 would potentially be much higher.

**Terminal oxidation of branched alkanes**

The reason branched alkanes are harder to degrade may be either that the alkyl branches hinder the uptake of the hydrocarbons into the cell or that the branches are not susceptible to the enzymes of the β-oxidation pathway (Schaeffer et al., 1979). One of the conditions necessary for aerobic degradation of branched hydrocarbons in nature is that the bacteria must possess genes and enzymes for initial oxygenation (Britton, 1984; Beguin et al., 2003). The degradation pathways of pristane have been established based on the analyses of the metabolic intermediates formed from pristane using Brevibacterium erythrogenes (Pimik et al., 1974), Nocardia globulera (Alvarez et al., 2001), and several strains such as Corynebacterium, Mycobacterium and Nocardia (McKenna and Kallio, 1971; Pimik et al., 1974; Beguin et al., 2003). Pristane is degraded by at least three catabolic pathways: monoterminus, diterminus and subterminal oxidation (Nhi-Cong et al., 2009). The metabolites produced from monoterminus oxidation of pristane are pristanol, pristyl aldehyde and pristane acid (Nakajima et al., 1985).

The initial hydroxylation reaction converting pristane to pristanol could be catalysed by the AlmA homologue (ABO_0282), P450-1/2 (ABO_0201/2288) and/or P450-3 (ABO_2384) (Fig. 3). AlmA expression was significantly higher on pristane compared to pyruvate, but the expression was highest on n-C14. The expression of P450-1/2 was at least 2-fold higher on both n-C14 and pristane compared to pyruvate. P450-3 was exclusively expressed on pristane. AlmA has previously been implicated in pristane degradation in other Alcanivorax strains. In A. dieselolei B-5 pristane activated the expression of AlmA (Liu et al., 2011). Enzymatic assays showed it converted pristane to its corresponding primary alcohol (Wang and Shao, 2014). In A. hongdengensis A-11-3 pristane selectively activated the expression almA (Wang and Shao, 2012b). In other OHCB expression of AlmA was enhanced in four Marinobacter strains (M. koreensis L53-1-2, M. alkaliphilus L53-10-4, M. alkaliphilus L52-11-18 and M. lipolyticus 342-2) when grown in the presence of pristane (Wang and Shao, 2012a). P450-1 from A. borkumensis is expressed from the promoter of the upstream gene (ABO_0199), and this promoter was more active when n-C9n-C18 or pristane were assimilated than when pyruvate was available (Sevilla et al., 2017). This is consistent with our results as P450-1 is differentially expressed in n-C14 and pristane compared to pyruvate suggesting it can use either linear or branched alkanes as substrates. P450-3 is phylogenetically distant from the other two cytochromes in A. borkumensis suggesting it may have a different substrate specificity, that is, branched alkanes (Wang and Shao 2012b; Sevilla et al., 2017). Further evidence of this has been shown by Schneiker et al. (2006), who confirmed that the branched alkane phytane strongly induce the expression of P450-3 in A. borkumensis SK27. In A. hongdengensis A-11-3 pristane selectively activates the expression of P450-3 (Wang and Shao, 2012b), and the expression of P450-3 in A. borkumensis was much higher when cells assimilated pristane in contrast to when n-alkanes or pyruvate were available (Sevilla et al., 2017). This was confirmed in our study as P450-3 was exclusively expressed on pristane.

The conversion of pristanol to pristyl aldehyde could be catalysed by either the alcohol oxidase (ABO_1174; exclusively expressed on pristane) or alcohol dehydrogenase (ABO_0061; two-fold up-regulated on pristane), which utilize two different biological modes of alcohol dehydrogenation (Fig. 3). Alcohol dehydrogenases are nicotinamide dependent, while alcohol oxidas are flavin dependent and generate hydrogen peroxide (Geissler and Hemmerich, 1981; Kemp et al., 1988). Following terminal oxidation, the alcohols generated are normally oxidized to the corresponding aldehyde by means of alcohol dehydrogenases in bacteria, although alcohol oxidas have been reported in alkane degrading yeasts and moulds, for example, Candida spp. where they functionally substitute alcohol dehydrogenases (Blasig et al., 1988; Kemp et al., 1988; Hommel and Ratledge, 1990).

For the conversion of pristyl aldehyde to pristanic acid no aldehyde dehydrogenases were significantly differentially expressed during growth on pristane. An aldehyde dehydrogenase (ABO_0087) was expressed on all substrates tested meaning it may be constitutively expressed. Also, catalysis of sequential oxidation reactions is usual in cytochrome P450 reactions particularly...
with xenobiotics (Guengerich et al., 2011) (Fig. 3). P450 enzymes catalyse a notably diverse range of oxidative transformations on a very wide range of substrates (Meunier et al., 2004). P450 enzymes can oxidize aldehyde to carboxylic acids through a well understood mechanism (Newcomb et al., 2003) and examples include the oxidation of aliphatic α,β-unsaturated aldehydes and anthraaldehyde to the corresponding acids by the latter enzyme (Guengerich et al., 2011). This means the cytochrome P450 could potentially functionally substitute an aldehyde dehydrogenase generating a fatty acid, which is further degraded through β-oxidation.

**β-oxidation**

β-Oxidation of acyl-CoA derivatives

A. borkumensis expressed a different set of enzymes for β-oxidation depending on the growth substrate (branched vs. linear alkane) utilized (Fig. 4). Fatty acids generated from the oxidation of alkanes are transported through the cytoplasmic membrane coupled with the activation to acyl-CoA thioesters catalysed by the acyl-CoA synthetase with concomitant hydrolysis of ATP (Fiedler et al., 2002). Acyl-CoA synthetases are widely distributed in both prokaryotic and eukaryotic organisms and exhibit a broad substrate specificity (Ruth et al., 2008). ABO_0367 encodes a fadD homologue (Fig. 4B). The product of the fadD gene is a long-chain fatty acyl-CoA ligase that converts exogenous long-chain fatty acids (LCFAs) into acyl-CoA thioesters when they are transported in bacteria (Black et al., 1992). The mechanism behind FadD substrate specificity is not fully understood. Structures of a homologous long chain acyl-CoA synthetase from Thermus thermophilus and a medium chain acyl-CoA synthetase from Homo sapiens suggest that it is the length of the FadD fatty acid binding pocket that determines specificity (Hisanaga et al., 2004; Kochan et al., 2009). A consensus sequence (‘DGWLTGDIGWXPGXLKIIDDRKK’)

which is common to all fatty acyl-CoA synthetases and represents a FACS signature motif (Black et al., 1997). Mutational studies revealed subtle changes in this region result in marked changes with respect to substrate recognition (e.g. D22 and K26) defining the fatty acid specificity of the enzyme (Black et al., 1997). For example, fatty acyl-CoA synthetases from Pseudomonas oleovorans and Saccharomyces cerevisiae, which lack the lysyl residue at the position equivalent to K24 have specificity to medium-chain fatty acids (van Beilen et al., 1992; Knoll et al., 1994). ABO_0367 (three-fold higher expression on pristane compared to n-C14) has a valine residue in the K24 position and a glycine in the D22 in the FACS signature motif, which may contribute to its preference to branched fatty acids over straight-chain fatty acids. ABO_2748, which had 3-fold higher expression on pristane compared to n-C14, is an AlkK homologue (Fig. 4B). The gene product of alkK in Pseudomonas putida encodes a medium-chain acyl CoA synthetase, which catalyses the final reaction in alkane oxidation, converting medium-chain length fatty acid to their CoA derivatives (van Beilen et al., 1992). The enzyme was able to complement a fadD mutation in E. coli K-12 verifying its predicted role (van Beilen et al., 1992). Assays using branched hydroxalkanoates as substrates confirmed AlkK is also capable of converting branched substrates to their corresponding acyl-CoA thioesters (Satoh et al., 2005). As AlkK is capable of utilizing branched substrates this may account for the higher spectral counts during growth on pristane.

The A. borkumensis genome contains two fadAB operons, consisting of β-oxidation multifunctional enzyme complexes with 3-hydroxyacyl-CoA dehydrogenase and ketoacyl-CoA thiolase activities that were detected in our data set (Fig. 4G and H). They exhibited different expression patterns depending on the substrate utilized (fadAB/ABO_1566/1567 was expressed on the branched pristane, and FadAB2/ABO_1652/1653 was expressed on the linear alkane). It has been widely reported that different FadAB isofoms have different substrate specificities and multiple FadAB operons exist within a microbes genome. For example, Escherichia coli possess an anaerobic paralog of the canonical FadAB, YtcYX (Campbell et al., 2003). There are multiple sets of FadAB complexes described in the model strain Pseudomonas putida KT2440, which accounts for the huge metabolic versatility of this strain (Nelson et al., 2002; Ouyang et al., 2007). More restricted substrate specificity was reported for Mycobacterium tuberculosis FadB, which needs the action of a trans-acting enoyl-CoA isomerase in order to completely degrade cis-unsaturated fatty acids (Srivastava et al., 2015). In the case of Streptomyces coelicolor FadAB complexes, low efficiency or narrow substrate specificity range of the complexes accounted for lower growth rates exhibited by two of the three complexes during growth on oleic acid as the sole carbon source. (Menendez-Bravo et al., 2017).

**Conclusions**

In conclusion, A. borkumensis SK2T, an important OHCB that can dominate microbial communities following marine oil spills, possesses the ability to degrade branched alkanes, which provides it a competitive advantage over many other marine alkane degraders that can only degrade linear alkanes. This study has significantly enhanced our understanding of the fundamental physiology of A. borkumensis SK2T by the identification of the expressed enzymes for degrading linear and branched alkanes. It has also highlighted the differential expression...
of sets of β-oxidation proteins to overcome steric hindrance from branched substrates.

Experimental procedures

**LC–MS/MS shotgun analysis of A. borkumensis SK2**

**Culture conditions.** A. borkumensis SK2 (DSM 11573) was grown from a frozen stock in sterile 160 ml Nunc Cell Culture-Treated Flasks containing 100 ml of ONR7a media (Dyksterhouse et al., 1995). Cultures were supplemented with tetradecane (n-C14), pristane or pyruvate as the sole carbon source. The starting inoculum was 20 µl of exponentially growing cells (from a 3 day old preculture of cells growing on either n-C14, pristane or pyruvate that were in the exponential growth phase, i.e., the first measurable day of growth by OD). Cultures were incubated in an orbital shaker (16°C, 60 rpm). Growth curves were highly similar on all three substrates, and cells were harvested for protein extraction after 4 days, which represented cells in early exponential growth phase (determined by growth curves measured by OD at 600 nm, that showed a two day lag phase, followed by exponential growth measurable by an increase in OD from day 3, with cells entering stationary phase after 14 days).

**LC–MS/MS proteomics.** Cells were harvested from 50 ml of each culture by centrifugation (4600 × g, 15 min) and washed in 2 ml of phosphate buffered saline. Total protein was immediately extracted by resuspending the cell pellet in 75 µl of extraction buffer (62.5 mM TRIS–HCl pH 6.8, 10% glycerol v/v, 12 mM dithiothreitol (DTT), 2% sodium dodecyl sulfate (SDS) v/v and one Pierce Protease Inhibitor Tablet per 50 ml), heating in a water bath (95°C, 12 min) and then centrifuging (10,500 × g, 5 min) to remove cell debris. Proteins extracts were visualized by SDS-PAGE and trypsin digestion, and LC–MS/MS with a ThermoFisher hybrid high-resolution LTQ Orbitrap instrument was performed as previously described (McKew et al., 2013).

**MS/MS analysis.** MS/MS analysis was performed in MaxQuant (Cox and Mann, 2008). The LTQ Orbitrap raw data files were first converted to MSM files with the MaxQuant ‘Quant’ module. The open-source search engine Andromeda, which is integrated into MaxQuant, was used to identify peptides in sequence databases by their fragmentation spectra (Cox et al., 2011). Peptides and proteins were filtered at 0.3% false discovery rate (FDR) to obtain the final data sets. Proteins were quantified by counting the number of MS/MS spectra matched to the corresponding proteins. Uniprot protein sequences from the A.borkumensis SK2 genome (Schneiker et al., 2006) were used to perform protein identification. Proteins were validated using the default settings in MaxQuant and Andromeda with a minimum of at least one peptide, but that any such protein had to unambiguously identified by peptides that were unique to that protein. Spectral counts were normalized to total spectral counts that account for differences between runs (total spectral counts varied between 11,015 and 15,856 per run).

**Statistical and bioinformatic analysis.** Differential expression analysis was performed by analysis of variance (ANOVA) and Tukey’s HSD test with Benjami-Hochberg post-hoc corrections (Benjami and Hochberg, 1995) within the XLSTAT-Premium Version 2016.1 (Addinsoft) ‘OMICs’ package. All proteins significantly (P < 0.05) up-regulated during growth on hydrocarbons were subjected to a BLAST (Basic Local Alignment Search Tool) (Altschul et al., 1990) against the NCBI nr database. Protein family and domain analysis was carried out in Pfam v30.0 (Finn et al., 2016). Proteins were assigned to functional families by hierarchical classification of protein domains based on their folding patterns in CATH v4.1 (Class, Architecture, Topology, Homology) (Sillitoe et al., 2015). Full length secondary and tertiary structure predictions, functional annotations on ligand-binding sites, enzyme commission numbers and gene ontology terms were generated using the I-TASSER SERVER (Zhang, 2008).

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### Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

**Table S1** Mean spectral counts (*n* = 3) of detected proteins during growth on a linear alkane (*n*-C<sub>14</sub>), a branched alkane (Pristane) and a non-hydrocarbon control (Pyruvate) in *Alcanivorax borkumensis* SK2<sup>T</sup>, with differential expression analysis (Anova with Benjamini Hochberg FDR correction (P-value) and Tukey HSD Post Hoc Test (a,b,c in parentheses indicate significant differences by Tukey HSD test))