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Anaerobic Oxidation of Ethane, Propane, and Butane by Marine Microbes: A Mini Review

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The deep ocean and its sediments are a continuous source of non-methane short-chain alkanes (SCAs) including ethane, propane, and butane. Their high global warming potential, and contribution to local carbon and sulfur budgets has drawn significant scientific attention. Importantly, microbes can use gaseous alkanes and oxidize them to CO₂, thus acting as effective biofilters. A relative decrease of these gases with a concomitant ¹³C enrichment of propane and n-butane in interstitial waters vs. the source suggests microbial anaerobic oxidation. The reported uncoupling of sulfate-reduction (SR) from anaerobic methane oxidation supports their microbial consumption. To date, strain BuS5 isolated from the sediments of Guaymas Basin, Gulf of California, is the only pure culture that can anaerobically degrade propane and n-butane. This organism belongs to a metabolically diverse cluster within the Deltaproteobacteria called Desulfosarcina/Desulfococcus. Other phylotypes involved in gaseous alkane degradation were identified based on stable-isotope labeling and fluorescence in-situ hybridization. A novel syntrophic association of the archaeal genus, Candidatus Syntrophoarchaeum, and a thermophilic SR bacterium, HotSeep-1 was recently discovered from the Guaymas basin, Gulf of California that can anaerobically oxidize n-butane. Strikingly, metagenomic data and the draft genomes of ca. Syntrophoarchaeum suggest that this organism uses a novel mechanism for n-butane oxidation, distinct from the well-established fumarate addition mechanism. These recent findings indicate that a lot remains to be understood about our understanding of anaerobic SCA degradation. This mini-review summarizes our current understanding of microbial anaerobic SCA degradation, and provides an outlook for future research.

Keywords: Gulf of Mexico, short-chain alkanes, sulfate reduction, anaerobic oxidation, Desulfosarcina/Desulfococcus

INTRODUCTION

Microbes drive fundamental processes in marine sediments, including the oxidation of organic matter, production of methane and other hydrocarbons, and the removal of sulfate from oceans (Jørgensen, 1982; D’Hondt et al., 2004; Hinrichs et al., 2006; Kallmeyer et al., 2012). The marine biosphere represents a major reservoir for microbial life on Earth. Kallmeyer et al. (2012) estimated the global subseaﬂoor sedimentary microbial abundance to be 2.9 × 10²⁹ cells, corresponding to ~0.6% of Earth’s total living biomass. Unlike photosynthetic processes, metabolic strategies in dark oceans are based on chemotrophy, where reduced organic and inorganic compounds...
including methane are the dominant electron donors (Orcutt et al., 2011; Colwell and D’Hondt, 2013). Non-methane short-chain alkanes (SCAs), [ethane (C₂), propane (C₃), butane (C₄); only microbial n-butane consumption is noted thus far] represent additional substrates for primary productivity and play an important role in marine ecosystems. Considerable amounts of SCAs are produced continuously from the ocean via biotic and abiotic processes (0.54 Tg year⁻¹ ethane, 0.35 Tg year⁻¹ propane, and 0.11 Tg year⁻¹ butane) (Plass-Dülmer et al., 1995). Their subsequent escape into the atmosphere significantly contributes to the formation of ozone and organic aerosols (Etiope and Ciccioli, 2009; Pozzer et al., 2010). However, aerobic and anaerobic hydrocarbon-degrading microorganisms can dramatically lower the amount of SCAs reaching the atmosphere (Head et al., 2006; Atlas and Hazen, 2011; Callaghan, 2013). Aerobic microorganisms that can oxidize SCAs are well-characterized (Dworkin and Foster, 1958; Kinnaman et al., 2007; Yakimov et al., 2007; Redmond et al., 2010). Recent studies in cold marine seeps and marine hydrothermal vents have shed light on the microbial aerobic oxidation of SCAs (Kniemeyer et al., 2007; Adams et al., 2013; Bose et al., 2013; Kleindienst et al., 2014; Dowell et al., 2016; Laso-Pérez et al., 2016). Nevertheless, research on the anaerobic oxidation of ethane is still in its infancy. Microbiologically mediated anaerobic ethane oxidation linked to SR has been reported for the Gulf of Mexico (GoM) and Middle Valley (MV) sediments (Adams et al., 2013; Bose et al., 2013). However, identification of individual isolates/consortia and the mechanisms involved are still unknown and await discovery. This mini-review summarizes our current understanding of anaerobic microbial SCA degradation, and provides an outlook for future research.

SHORT-CHAIN ALKANES AS CARBON AND ENERGY SOURCE IN THE MARINE ECOSYSTEMS

SCAs are chemically the least reactive compounds due to their non-polar C-H σ-bonds (Carey, 2007). Despite this, microbes can oxidize them aerobically or anaerobiocally (Leahy and Colwell, 1990; Maeng et al., 1996; Heider et al., 1999; Callaghan et al., 2006; Rojo, 2009; Callaghan, 2013). Aerobes activate alkanes by cleaving C-H bonds via monoxygenase or dioxygenase enzymes (Callaghan et al., 2006). Their terminal oxidation results in an alkanoic acid by dehydrogenases to aldehydes, then to fatty acids followed by β-oxidation (Rabus et al., 2001; Callaghan et al., 2006). The key role that oxygen plays in aerobic alkane transformations led to the belief that alkanes are biologically inert under anoxic conditions. However, research conducted over the years has shown that activation of hydrocarbons can also occur under such conditions. In marine ecosystems, anaerobic SCA degradation is linked to only SR unlike terrestrial ecosystems where nitrate and chloride act as electron acceptors for hydrocarbon (>C₄) degradation (Wilkes et al., 2003; Melboob et al., 2009; Zedelius et al., 2011; Adams et al., 2013; Bose et al., 2013; Kimes et al., 2013; Chanton et al., 2015).

Emission of oil and gas from hydrocarbon seeps are widespread along continental margins. This gas is primarily composed of methane, a potent greenhouse gas; and marine hydrocarbon seeps are estimated to contribute 20 Tg year⁻¹ methane to the atmosphere, representing about 5% of the total atmospheric flux (Fung et al., 1991; Judd, 2004). Due to the high concentration of methane in the atmosphere, bacterial oxidation of methane under aerobic and anaerobic conditions has received considerable attention. Over the past four decades, studies focusing on anaerobic oxidation of methane (AOM) have revealed the diversity and distribution of methane oxidizers, and the underlying biochemical processes (Reeburgh, 2007; Knittel and Boetius, 2009; Callaghan, 2013; Haroon et al., 2013). In addition to methane, these hydrocarbon seeps also release an estimated 0.45 Tg year⁻¹ ethane and 0.09 Tg year⁻¹ propane into the atmosphere (Etiope and Ciccioli, 2009). The amount reaching the atmosphere would be substantially larger if not for microbial oxidation in the sediments and water column (Reeburgh, 2007). Although, microbial aerobic and anaerobic consumption of SCAs from marine and terrestrial environments is widely known (Redmond et al., 2010; Mbadinga et al., 2011; Callaghan, 2013; Musat, 2015), to the best of our knowledge, no quantitative approaches have been used to clearly define the partitioning of SCAs between their atmospheric emission and oxidation. The lack of such data makes it difficult to estimate the influence of SCAs to global carbon budgets and their potential effect on climate. Future research on SCA degradation would help fill this large knowledge gap.

In marine ecosystems, anaerobic oxidation of C₂-C₄ alkanes can significantly contribute to community bioenergetics (Lorenson et al., 2002; Formolo et al., 2004; Sassen et al., 2004; Alain et al., 2006; Bose et al., 2013), while competing with AOM for sulfate, an electron acceptor shared by these processes (Joye et al., 2004; Orcutt et al., 2010; Bowles et al., 2011; Adams et al., 2013; Bose et al., 2013). Indeed, at the GoM cold seeps, SR rates are higher than can be accounted for by AOM alone, indicating that SR is potentially linked to the oxidation of non-methane SCAs or higher petroleum hydrocarbons (Joye et al., 2004; Musat et al., 2009; Orcutt et al., 2010; Bowles et al., 2011). The microbial oxidation of SCAs is confirmed by ¹³C-enriched propane and n-butane in the sediment interstitial water relative to gas in the hydrates, and the carbonate alkalinity around them (Sassen et al., 2004). Overall, studies on the microbial degradation of SCAs in marine settings are motivated by questions of how such processes affect global carbon and sulfur cycling, and who the participating organisms are.

Anaerobic oxidation of C₂-C₄ alkanes with SR has been demonstrated in anoxic, marine settings (Kniemeyer et al., 2007; Savage et al., 2010; Adams et al., 2013; Bose et al., 2013; Kleindienst et al., 2014; Laso-Pérez et al., 2016). These studies have identified novel and metabolically diverse microbes thriving on C₂-C₄ alkanes. Sulfate-reducing bacteria (SRB) from the GoM and Guaymas Basin (GB), Gulf of California sediments oxidized C₃-C₄ alkanes to CO₂ (Kniemeyer et al., 2007). These authors tested different temperature regimes (12, 28, and 60°C) using various substrates (methane, ethane, propane,
$n$-butane, iso-$n$-butane, alcohols, or carboxylic acids). A pure culture (BuSS) isolated from 28°C enrichments, anaerobically oxidized C$_3$-C$_4$ alkanes (Kniemeyer et al., 2007). The strain was determined to be a *Deltaproteobacterium* within the *Desulfosarcina/Desulfococcus* (DSS) cluster. Further, Kleindienst et al. (2014) showed the presence of distinct DSS clades in two seep sediments from the Mediterranean Amon mud volcano and Guaymas Basin degrading $n$-butane and dodecane. Jaekel et al. (2013) enriched microbial populations from the GoM and Hydrate Ridge marine cold seeps that degraded C$_3$-C$_4$ alkanes. Similar to the previous observation by Kniemeyer et al. (2007), the enrichment cultures degraded propane and $n$-butane simultaneously, but not methane, ethane, iso-$n$-butane, or pentane. They also identified DSS cluster members as the responsible phylotypes. Using *ex-situ* sediment slurries, Bose et al. (2013) demonstrated the anaerobic oxidation of C$_1$-C$_4$ alkanes coupled with SR. Interestingly, these authors observed ethane consumption comparable to methane, propane, and $n$-butane. This is in contrast to the study by Kniemeyer et al. (2007), who reported extremely slow rates of ethane-driven SR. A notable difference between these studies is the use of sediment slurries by Bose et al. (2013) compared to enrichment techniques used by Kniemeyer et al. (2007). Community analyses suggested the enhancement of *Deltaproteobacteria* in SCA amended reactors. Deltaproteobacterial sequences from ethane incubations were closely related to the isolate BuSS, and the enrichment culture Butane 12-GMe (both isolated from marine sediments by Kniemeyer et al., 2007 and Bose et al., 2013).

Anaerobic SCA degradation was also demonstrated in metalliferous, organic-poor Middle Valley hydrothermal vent sediments at 25, 55, and 75°C. Sediment slurries showed degradation of C$_1$-C$_4$ alkanes under SR conditions (Adams et al., 2013). Comparison of bacterial communities, suggested the presence of *Deltaproteobacteria* mediating the anaerobic oxidation of C$_1$-C$_4$ alkanes. This implied that, anaerobic alkane degraders exist in both cold marine seeps and high temperature hydrothermal vent systems. Importantly, anaerobic oxidation of SCAs is not restricted to only the *Deltaproteobacteria*. For example, an enrichment from GB sediments with propane at 60°C was dominated by Gram positive, SRB closely related to *Desulfitomaculum*, a commonly found cluster of bacteria in the subsurface biosphere within the *Peptococcaceae* (Ollivier et al., 2007; Wang et al., 2008; Aüllo et al., 2013).

The diversity of SCA degraders in marine environments was recently demonstrated by the discovery of syntrophic $n$-butane degraders from a thermophilic enrichment culture from the GB vent area (Laso-Pérez et al., 2016). Syntrophic association of the archaeal genus, *Candidatus Syntrophoarchaeum* and a thermophilic SRB, HotSeep-1 completely oxidized $n$-butane to CO$_2$. A subsequent study on GB hydrothermal mound sediments used bacterial and archaeal 16S rRNA gene clone libraries and V6 tag pyrosequencing to show the co-occurrence of archaeal groups (such as, anaerobic methane-oxidizing archaea ANME-1, ANME-1Guaymas, and ANME-2) with bacterial groups (such as, SEEP-SRB2 and HotSeep-1) (Dowell et al., 2016). This corroborates that an archaeal-bacterial syntrophic community mediates alkane degradation in a GB hydrothermal mound.

The HotSeep-1 group was also detected in thermophilic SR enrichments with $n$-butane from GB at 60°C (Kniemeyer et al., 2007), and in SR enrichments inoculated with MV hydrothermal sediments amended with C$_2$-C$_4$ alkanes at 55°C (Adams et al., 2013). It is likely that the organisms in the HotSeep-1 group do not oxidize these alkanes directly but function as versatile syntrophs that serve as electron/hydrogen sinks within different consortia (Zengler et al., 1999; Dowell et al., 2016). Together, these findings reflect the diversity of anaerobic microorganisms thriving on non-methane SCAs in marine environments. These studies also suggest that the processes mediated by these organisms contribute to ocean chemistry and community bioenergetics via both sulfate and SCA removal.

Despite informative studies over the past decade on anaerobic SCA degradation, very little is known about ethane-oxidizing phylotypes, and how they interact with other organisms in deep-sea ecosystems. Kniemeyer et al. (2007) reported ethane dependent SR in an enrichment from the GoM at 12°C. However, the reported rate was orders of magnitude slower than the oxidation rates of C$_3$-C$_4$ alkanes. Notably, Bose et al. (2013) observed ethane consumption approximately two orders of magnitude higher than those reported by Kniemeyer et al. (2007) in *ex-situ* slurry incubations of GoM sediments incubated at 7°C. It is possible that SRB closely related to the C$_3$-C$_4$ degrading DSS cluster might be associated with ethane degradation in these incubations, though this remains to be investigated. These authors also demonstrated carbon flux dynamics of ethane oxidation using $\delta^{13}$C of DIC and alkanes from their enrichment experiments.

Batch incubations with sediments from MV hydrothermal vent systems showed modest rates of ethane dependent SR at 25, 55, and 75°C (Adams et al., 2013). In accordance with the observed stoichiometries, SR coupled to the anaerobic oxidation of C$_2$-C$_4$ proceeded at a faster rate than AOM at mesophilic and thermophilic temperatures (25 and 55°C, respectively). These faster rates might indirectly limit AOM. These findings suggest that microbial anaerobic SCA degradation affects local carbon and sulfur cycles. In contrast to C$_3$-C$_4$ alkane degradation, anaerobic ethane oxidation has not yet been reported by sediment-free cultures or isolates consortium. This makes ethane oxidation the least understood among the SCA degradative processes. This is in part due to the slow rate of microbial anaerobic ethane oxidation observed in enrichment studies (Kniemeyer et al., 2007). Anaerobic ethane oxidation needs to be investigated in further detail due to its abundance in marine seeps (Plass-Dülmer et al., 1995; Sassen et al., 1998; Etiope and Ciccioli, 2009), its contribution to tropospheric chemistry (Singh et al., 1994; Katzenstein et al., 2003), and the demonstrated ability of microbial populations to degrade ethane linked to SR (Adams et al., 2013; Bose et al., 2013). Studies should focus on the precise nature and extent of this process; the responsible microbes; and the associated biochemistry.
BIOCHEMISTRY OF ANAEROBIC SHORT-CHAIN ALKANE OXIDATION

Fumarate addition is noted as the biochemical mechanism for aromatic hydrocarbon and n-alkane activation by anaerobes (Biegert et al., 1996; Beller and Spormann, 1997; Kropp et al., 2000; Rabus et al., 2001; Wilkes et al., 2002; Callaghan et al., 2006, 2008, 2012; Grundmann et al., 2008). In this pathway, n-alkanes are activated by fumarate addition to the double bond at the sub-terminal or terminal carbon producing 2-(1-methylalkyl)succinates (or 2-alkylsuccinates). Degradation of 2-(1-methylalkyl)succinates involves carbon skeleton rearrangement, and decarboxylation yielding branched fatty acids followed by β-oxidation (Widdel and Grundmann, 2010; Agrawal and Gieg, 2013; Callaghan, 2013; Musat, 2015) (Figure 1). C₃-C₄ alkane activation by strain BuS5 and, in the marine enrichment Propane60-GuB is suggested to occur via the same pathway (Kniemeyer et al., 2007). Based on the metabolites detected in both cultures, activation of n-butane presumably occurs at the secondary carbon yielding (1-methylpropyl) succinate. Interestingly, it was suggested that propane activation occurs at both secondary and primary carbon atoms producing isopropyl- and n-propylsuccinate respectively (Kniemeyer et al., 2007) (Figure 1). Although initially considered a side reaction, the second pathway was substantiated by incubations of strain BuS5 with position-specific deuterium-labeled propane (Jaekel et al., 2014). Results showed that the activation of propane at the secondary carbon is more significant, accounting for an estimated 70% of the activation events, with 30% of activation occurring at the primary carbon. Based on these findings, activation of ethane would likely yield ethylsuccinate. Although, metabolites analyses in ethane-degrading laboratory batch reactors are lacking, ethylsuccinate is reported from hydrocarbon-rich settings such as, in crude oil processing facilities and production wells (Duncan et al., 2009), oilfields (Gieg et al., 2010), and coal beds (Wawrick et al., 2012).

At the biochemical level, this process involves the abstraction of an H atom from the alkane substrate by the glycyl radical enzyme (GRE), 1-methylalkyl succinate synthase (MAS) (Grundmann et al., 2008) also known as alkylsuccinate synthase (ASS) (Callaghan et al., 2008). These enzymes utilize free radicals to catalyze fumarate addition to form a succinate radical intermediate (Callaghan et al., 2008, 2010; Bharadwaj et al., 2013; Von Netzer et al., 2013; Gittel et al., 2015).

The genes encoding an alkane activating GRE have been identified in the SRB Desulfatibacillum alkenivorans AK-01 (Callaghan et al., 2008) and Desulfoalaba alkanexedens ALDCT (Callaghan et al., 2010) and in nitrate reducing strains HxN1 (Grundmann et al., 2008) and OcN1 (Zedelius et al., 2011), all affiliated to the Deltaproteobacteria. Recent genome analysis of strain BuS5 identified a single putative masD gene, suggesting that one MasD is involved in the activation of both propane and n-butane (Musat, 2015). Phylogenetic reconstruction of translated full-length and partial masD/assA/bssA/nmsA homologs from selected isolates, as well as pristine and seepage-impacted metagenomes showed that there is a wide diversity of organisms that can degrade short-, long-chain hydrocarbons, and cyclic aromatic hydrocarbons (Figure 2). Gittel et al. (2015) designed novel PCR primers to the masD/assA gene to determine the diversity and distribution of anaerobic alkane degraders in pristine and seepage-impacted Danish coastal sediments. Seepage-impacted sediments were dominated by a single masD/assA gene cluster, which indicates an occurrence of a substrate-adapted community. In contrast, pristine sediments harbored a diverse range of masD/assA phylogenotypes including those present in seepage-impacted sediments. This comprehensive cultivation-independent survey of the diversity and distribution

![FIGURE 1](image-url)  
FIGURE 1 | Anaerobic activation of propane at the sub-terminal (A) and terminal (B) carbon atom (marked with stars) via fumarate addition yielding isopropylsuccinate and n-propylsuccinate, respectively. A similar activation mechanism exclusively at the sub-terminal carbon atom is proposed for the anaerobic oxidation of n-butane.
FIGURE 2 | Maximum likelihood tree of translated full-length and partial masD/assA/nmsA homologs from selected isolates as well as pristine and seepage-impacted metagenomes obtained from GenBank (accession numbers are shown in parentheses). Tree was inferred using the Le_Gascuel_2008 model (Le and Gascuel, 2008) and involved 85 amino acid sequences and a total of 210 positions. All positions with less than 95% site coverage were eliminated. Full-length glycerol dehydratase (dhaB1) from Clostridium butyricum was used as an outgroup. Node circles denote bootstrap value percentages from 100 replicate trees. Scale bar represents 20% estimated sequence divergence. Evolutionary tree was constructed in MEGA7 (Kumar et al., 2016).
of anaerobic alkane degraders highlighted the relevance of masD\text potential influences on sulfur cycling and microbial diversity. Front. Microbiol. 4:110. doi: 10.3389/fmicb.2013.00110


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AUTHOR CONTRIBUTIONS

RS performed all the necessary literature searches and data compilation. MG performed the phylogenetic analysis. RS and AB wrote the manuscript.

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