Activity and abundance of methane-oxidizing bacteria in secondary forest and manioc plantations of Amazonian Dark Earth and their adjacent soils

Amanda B. Lima¹, Aleksander W. Muniz² and Marc G. Dumont¹ *

1 Department of Biogeochemistry, Max Planck Institute for Terrestrial Microbiology, Marburg, Germany
2 Department of Soil Microbiology and Biogeochemistry, Brazilian Agricultural Research Corporation, Manaus, Brazil

INTRODUCTION

Most soils in the Amazon region have low fertility. Typically, Amazonian soils are acidic, have low P contents, low cation exchange capacity and high levels of Al at levels that can be toxic to crops (Cochrane and Sanchez, 1982). In contrast, Amazonian Dark Earth (ADE) soils, also known as terra preta de índio, are fertile soil patches found dispersed throughout the Amazon that were formed by the ancient Amazonian indigenous populations. It is believed that these soils were unintentionally or intentionally formed by long-term habitation with casual addition of domestic refuse and by long-lasting agricultural activity based on the clearing of vegetation and the incomplete combustion of organic material (Smith, 1980; Denevan, 1998; Glaser, 1999). Unlike their adjacent soils, ADE have high contents of P, Ca, Mg, Zn, Mn, and stable organic matter (Costa and Kern, 1999; Woods and McCann, 1999).

Differences in bacterial community structure and composition have been observed under different land use systems in Amazonian soils (Jesus et al., 2009; Navarrete et al., 2010; Taketani et al., 2013), which will in turn influence ecosystem processes such as the decomposition of organic matter and nutrient mineralization or its immobilization (Neher, 1999). In addition, the bacterial communities in ADE soils were shown by 16S rRNA tag sequence analysis to be distinct from their adjacent soils, particularly when compared at taxonomic levels lower than phylum (Taketani et al., 2013). One of the possible influences on the microbial communities of ADE soils is the presence of large amounts of biochar, which have prompted research into the effect of biochar application on microbial community structure and composition (Anderson et al., 2011; Khodadad et al., 2011). Replicating the high carbon and biochar contents of ADE in other soils has been suggested as a mechanism of CO₂ sequestration (Sombrero et al., 2003; Lehmann, 2007); however, the presence of these relatively large amounts of carbon in ADE then raises concern whether changes in climate and land use may result in increased emissions of CO₂ and CH₄. One possible mechanism of increased CH₄ emissions would be the decomposition of labile components of biochar to form substrates for methanogens (Knoblauch et al.,...
Although this activity is found in a wide variety of upland soils, ADE soils are a potential sink for atmospheric CH4 and secondly to Alten (2008). Numerous studies have shown that the conversion of pristine land to agriculture lowers the oxidation capacity of the soil (Keller et al., 1990; Mosier et al., 1991; Hüttsch et al., 1994; Jensen and Olsen, 1998; Priemé and Christensen, 1999; Kief et al., 2005; Levine et al., 2011). Factors associated with agriculture have been shown to inhibit atmospheric CH4 oxidation, including soil compaction, acidification and fertilization (Dunfield, 2007). Conversely, the abandonment of agriculture can also lead to at least partial recovery of methanotroph populations and atmospheric CH4 uptake (Levine et al., 2011). ADE soils are commonly found on well-drained areas and oxic, have an important role in the global CH4 cycle by acting as a sink for atmospheric CH4 (King, 1992), which globally is estimated at more than 30 Tg y\(^{-1}\) (Denman et al., 2007).

Upland soils, defined as those that are typically well-drained and oxic, have an important role in the global CH4 cycle by acting as a sink for atmospheric CH4 (King, 1992), which globally is estimated at more than 30 Tg y\(^{-1}\) (Denman et al., 2007). Although this activity is found in a wide variety of upland soils, pristine forest soils have been identified as the most efficient biological sinks of atmospheric CH4 (Dunfield, 2007; Dalal and Allen, 2008). Numerous studies have shown that the conversion of pristine land to agriculture lowers the oxidation capacity of the soil (Keller et al., 1990; Mosier et al., 1991; Hüttsch et al., 1994; Jensen and Olsen, 1998; Priemé and Christensen, 1999; Kief et al., 2005; Levine et al., 2011). Factors associated with agriculture have been shown to inhibit atmospheric CH4 oxidation, including soil compaction, acidification and fertilization (Dunfield, 2007). Conversely, the abandonment of agriculture can also lead to at least partial recovery of methanotroph populations and atmospheric CH4 uptake (Levine et al., 2011). ADE soils are commonly found on well-drained areas of the Amazon region (terra firme), and may also be sinks for atmospheric CH4.

Methanotroph diversity and activity has been assessed in different upland soils exhibiting atmospheric CH4 oxidation (Dunfield, 2007; Kolb, 2009). The diversity of atmospheric CH4 oxidizers is typically assessed by the detection of the pmoA gene, which encodes the β-subunit of methane monoxygenase (pMMO) enzyme (McDonald et al., 2008; Semrau et al., 2010). For the most part, as yet uncultivated microorganisms mediate atmospheric CH4 oxidation and are characterized by their pmoA gene sequences alone. In addition, phospholipid fatty acids have been used to identify atmospheric CH4 oxidizers (Bodelier et al., 2009). The USCγ pmoA clade is widely distributed in upland soils (Kief et al., 2003) and based on gene analyses are believed to belong to Alphaproteobacteria most closely related to Methylocapsa (Ricke et al., 2005). The USCγ pmoA clade is another associated with upland soils exhibiting atmospheric CH4 uptake, and appear to favor neutral or somewhat alkaline soils (Kief et al., 2003). Another clade termed JR3, initially identified in grassland soil (Horz et al., 2005) was found to dominate in desert soils with atmospheric CH4 oxidation capacity (Angel and Conrad, 2009). Methylocystis-related species have been shown to use CH4 at relatively low concentrations (Kief and Dunfield, 2005; Kief et al., 2006; Baani and Liesack, 2008), but whether they are important consumers of atmospheric CH4 in upland soils is not clear.

To our knowledge, no studies have previously examined CH4 oxidation or the diversity of methanotrophs in ADE soils. The primary objective of this study was to determine the extent to which ADE soils are a potential sink for atmospheric CH4 and secondly to determine how the methanotroph community structure and their CH4 uptake potential compares between forested and agricultural sites.

**MATERIALS AND METHODS**

**STUDY AREA, SOIL SAMPLING, AND SOIL ANALYSIS**

Soil samples were collected from two different areas, Caldeirão and Barro Branco. The Caldeirão experimental research station from Embrapa Amazônia Ocidental is located in Iranduba County in the Brazilian Central Amazon (03°26′00″ S, 60°23′00″ W). The other sampling area near the Barro Branco community is located in the Manacapuru County in the Brazilian Central Amazon (03°18′12″ S, 60°31′45″ W). ADE soils and their adjacent soils were collected from both areas. In both cases, the distance between the ADE soil zone and the adjacent soil zone was ∼2 km.

The soils were classified based on the World Reference Base for Soil Resources (FAO, 1998). ADE soils were classified as Hortic Anthrosol (i.e., reference horizon that results from prolonged cultivation with casual additions of domestic organic refuse and cultural materials). The adjacent soil from Caldeirão was classified as Haplic Acrisol (i.e., clay-rich soils with low fertility and toxic amounts of Al). The adjacent soil from Barro Branco was classified as Oxisol (i.e., red or yellowish soils with <10% weatherable minerals and low cation exchange capacity). At both areas, ADE soil and adjacent soil were sampled from secondary forest sites and agricultural sites cultivated with manioc (Manihot esculenta). The forested ADE and adjacent soil sites at Caldeirão were under ∼40-year-old secondary forest stands. At Barro Branco, the secondary forests were about 20 years-old. The agricultural sites in ADE and adjacent soils at Caldeirão had been used for manioc cultivation for at least 40 years, whereas the sites at Barro Branco had been deforested 5-years previously for conversion to plantations.

Soil samples were collected in February 2013. Three environmental replicates were collected from each sampling site. The sample plot (location) at each site was determined by choosing a random point, and from this reference point three sampling points (sublocations) 5 m apart were chosen for the collection of intact soil cores of 5 cm in diameter and 15 cm in length. Soil samples were collected in triplicate from each sublocation, which were subsequently homogenized to produce a composite soil sample for each sublocation. A total of 24 samples corresponding to the four sites (forested ADE, cultivated ADE, forested adjacent, and cultivated adjacent) from each of the two areas (Caldeirão and Barro Branco) were prepared. The samples for DNA extraction were transported from the field to the laboratory in an insulated box with dry ice. Approximately 1 kg of soil samples were collected from each of the 24 sublocations and sent to the department of Soil and Plant Nutrition of Embrapa Western Amazon. The frozen and unsieved soil samples were used for DNA extraction, whereas the 1 kg samples of fresh soil were sieved (2 mm mesh diameter) and used for the determination of soil chemical properties and CH4 oxidation potentials. Soil pH (H2O, 1:1), soil extractable Al, Ca, Fe, K, Mg, Mn, P, Zn, soil organic carbon (SOC), total C, total N, and cation exchange capacity were determined according to the methods described by Embrapa (1997).

**CH4 OXIDATION**

Potential CH4 oxidation rates were measured using soil from each sampling point (sublocation). Ten grams of fresh sieved soil was...
placed into a 120 ml serum vial in duplicate (Bull et al., 2000; Horz et al., 2002; Shrestha et al., 2012). The bottles were sealed with butyl rubber stoppers, and final mixing ratios of 10, 100, 1000, and 10 000 ppmv of CH₄ was injected into the gas headspace of the vials. The incubation of soil microcosms was performed at 25°C in the dark with shaking at 150 rpm for up to 19 days. CH₄ concentrations were measured on a daily basis by gas chromatography with a flame ionization detector using 0.5 ml gas samples from the bottle headspaces, as described previously (Shrestha et al., 2012). CH₄ oxidation rates were calculated by linear regression of CH₄ consumption versus time for the incubations with 10 ppm CH₄.

**DNA EXTRACTION FROM SOIL SAMPLES**

Soil DNA extractions were carried out in triplicate from 0.3 g wet weight subsamples of each soil sample. Extractions were performed using the Nucleospin soil DNA extraction kit (Macherey-Nagel, Düren, Germany) according to the manufacturer’s instructions. DNA was quantified using a Qubit dsDNA HS Assay (Molecular Probes, Invitrogen, USA). The triplicate DNA extracts of each sampling sublocation were pooled.

**REAL-TIME QUANTITATIVE PCR ASSAYS**

Real-time quantitative PCR (qPCR) with three technical replicates for each sublocation DNA sample was performed to determine the copy numbers of the pmoA genes. The qPCR assay using the primer set A189f-mb661r was used to target the conventional pmoA genes of *Methylocystaceae* and *Methylococcaceae* methanotrophs (Costello and Lidstrom, 1999; Kolb et al., 2003). The assay using primers A189f-Forested675r was used to target USC* pmoA* genes (Kolb et al., 2003). The qPCRs were performed with the SYBR Green JumpStart Taq ReadyMix System (Sigma, Taufkirchen, Germany) on an iCycler instrument (Bio-Rad, Munich, Germany). The data were analyzed using Bio-Rad CFX Manager (version 3.0) software. PCR mixtures and thermal cycling conditions were performed as described previously by Kolb et al. (2003). Briefly, the A189f-Forested675r assay was performed in 25 μl reaction mixtures containing 12.5 μl of SYBR Green Jump-Start Taq Ready Mix (Sigma), 1 μM of each primer, 50 ng of BSA (Roche, Mannheim, Germany), and 4 mM MgCl₂ (Sigma). The assay for the abundance of conventional pmoA genes (A189f-mb661r) was performed in 25-μl reaction mixtures containing 12.5 μl of SYBR Green Jump-Start Taq Ready Mix (Sigma), 0.667 μM of each primer and 4 mM MgCl₂. Standards for qPCR were generated by serial dilution of stocks of a known number of plasmids containing a single cloned copy of a *Methylococcus pmoA* gene or a USC* pmoA* gene, according to the assay. All samples from an experiment were run on a single plate.

**HIGH-THROUGHPUT SEQUENCING AND ANALYSIS**

PCR was performed using the primers A189f and A682r that amplify a broad range of pmoA, amoA, and related sequences (Holmes et al., 1995; Lük and Frenzel, 2011). The PCR components and conditions were identical to that described previously (Angel and Conrad, 2009). Briefly, the 50 μl reaction contained 5 μl of 10x AccuPrime™ PCR Buffer II (Invitrogen, Karlsruhe, Germany), additional 1.5 mM MgCl₂ (to a final concentration of 3 mM), 0.5 mM of each primer (Sigma), 50 ng of BSA (Roche) and 1 μl of Taq DNA polymerase (Invitrogen). All ADE samples could be amplified directly with the barcoded primer set; however, it was not possible to obtain amplicons of the expected size for the adjacent soil samples using these primers. Therefore, a 2-step PCR procedure in which conventional primers (i.e., without barcodes) was used in the first step followed by a successive low-cycle-number amplification using the barcoded primers, as described by Berry et al. (2011). This approach successfully produced PCR ampli- cons of the expected size. To allow comparisons, the same 2-step PCR approach was used for all samples. Five replicate PCR reactions were performed for each sample. After amplification, PCR reactions were pooled and loaded on 1% agarose gel stained in GelRed™ (Biotium Inc., Hayward, CA, USA). The DNA fragment of the correct size was excised from the agarose gel and eluted in 30 μl H₂O using the QIAquick gel extraction kit (Qiagen, Hilden, Germany). The purified PCR products from all samples were mixed in a 1:1 ratio and sequenced at the Max Plank-Genome-Centre Cologne (Cologne, Germany) using a Roche 454 Genome Sequencer FLX system.

A detailed description of the procedures used for sequence analysis was described previously (Dumont et al., 2014). In this study, only sequences with read lengths longer than 300 bp were used for further analysis. The sorting of sequences according to barcodes, trimming and quality filtering were processed using mothur version 1.29.2 (Schloss et al., 2009). Chimeric sequences were identified and removed using uchime (Edgar et al., 2011) implemented in mothur. Classification of pm oA sequences was performed using standalone TBLASTN version 2.2.26+ against a curated database of pm oA sequences and the lowest common ancestor (LCA) algorithm in MEGAN version 4.70.4 (Huson et al., 2011), as described previously (Dumont et al., 2014). A total of 110,437 sequences were obtained. 42,213 reads (a range from 9022 to 2977 reads per library) remained after basic quality filtering. The amplification of non-target sequences is common with these primers (Bourne et al., 2001) and these contaminants were identified by an absence of similarity to the reference database and removed from further analysis. The contaminants corresponded to an average of 57% from ADE samples and 87% from adjacent soil samples. A total of 13,595 reads remained after removing these contaminant sequences, corresponding to an average of 2802 reads from ADE and 597 from adjacent soil samples.

Representative sequences from each pm oA clade identified during the sequence analysis were selected for further analysis. These reads were translated into amino acid sequences and added to a reference pm oA/amoA phylogenetic tree using parsimony in ARB (Ludwig et al., 2004).

Sequences are available through the Metagenomics Rapid Annotation (MG-RAST) server with accession numbers 4577576.3 (TPISFBB2), 4577577.3 (TPISFBB3), 4577578.3 (TPISFBB4), 4577570.3 (TPIMBB2), 4577571.3 (TPIMBB3), 4577572.3 (TPIMBB5), 4577565.3 (ADJSFBB2), 4577566.3 (ADJSFBB3), 4577560.3 (ADJMBB2), 4577561.3 (ADJMBB3), 4577562.3 (ADJMBB4), 4577579.3 (TPISFC3), 4577580.3

1http://metagenomics.anl.gov/
ADE soils from Barro Branco had similar properties to those at and higher Al and Fe. The adjacent soils (i.e., Haplic Acrisol and Oxisol) had lower pH and Zn compared to their adjacent soils. These characteristics indicate the potential for high agricultural productivity. In contrast, the CH4 oxidation rates were more than one-order of magnitude lower in both plantations in adjacent soil and the ADE plantation with relatively high pH, Ca, CEC, K, Mg, Mn, P, SOC, and Zn compared to their adjacent soils. These characteristics indicate the potential for high agricultural productivity. In contrast, the adjacent soils (i.e., Haplic Acrisol and Oxisol) had lower pH and higher Al and Fe.

RESULTS

SOIL CHEMICAL PROPERTIES

The soil chemical properties are presented in Table 1. As previously reported, the measured soil chemical properties at the Caldeirão Experimental Station showed a clear distinction between ADE and adjacent soil samples (Taketani et al., 2013; Brossi et al., 2014). ADE soils from Barro Branco had similar properties to those at Caldeirão, with relatively high pH, Ca, CEC, K, Mg, Mn, P, SOC, and Zn compared to their adjacent soils. These characteristics indicate the potential for high agricultural productivity. In contrast, the adjacent soils (i.e., Haplic Acrisol and Oxisol) had lower pH and higher Al and Fe.

SOIL CH4 OXIDATION POTENTIALS

CH4 oxidation was immediate at concentrations of 10 and 100 ppmv, but a lag phase of 6–10 days was observed for concentrations of 1000 and 10,000 ppmv (results not shown). Relatively high rates of high-affinity CH4 oxidation (10 ppmv CH4) were observed in all soils from the forested sites and the ADE soil used for manioc cultivation at the Barro Branco area (Table 2). In contrast, the CH4 oxidation rates were more than one-order of magnitude lower in both plantations in adjacent soil and the ADE plantation soil at Caldeirão. The precise history of these soils is not available, but members of the local communities indicated that manioc has been cultivated in ADE soil at the Caldeirão site for living memory (>40 years), whereas the Barro Branco ADE soil was only recently (5 years) converted from forest to agriculture by slash-and-burn.

ABUNDANCE OF METHANOTROPHS

Quantitative real-time PCR assays were used to determine the copy numbers of pmoA genes in ADE and adjacent soils from both secondary forest and the manioc cultivation sites (Figure 1). The pmoA qPCR assay with primers A189f-A682r targets methanotrophs belonging to the Methylococcales and Methyloctysinales families and generally has poor specificity for the genes from other families of methanotrophs. The abundance of genes detected with this assay (Figure 1A) was not significantly affected by soil type or land use. Based on the diversity of pmoA genes detected in the soils (Figure 2), these results correspond to Methylocystis pmoA genes. Another qPCR assay was used to specifically enumerate USCα pmoA, which are a common uncultivated group associated with atmospheric CH4 oxidation. In ADE soils, the abundances of USCα pmoA (Figure 1B) were more than two-orders of magnitude higher than Methylocystis pmoA genes (Figure 1A). USCα were below the detection limit (1 × 10^6 copies g^-1 dry weight soil) in the plantations of adjacent soils. Taking the data from Barro Branco and Caldeirão sites together, the abundance of USCα pmoA was significantly higher in ADE than adjacent soil (ANOVA, p < 0.0001), but the difference in abundance based on land use (forested versus cultivated) was not significant (ANOVA, p = 0.77).

COMPOSITION OF METHANOTROPH COMMUNITIES

The methanotroph communities in the soils were analyzed by pmoA gene pyrosequencing. PCR using the A189f-A682r primer combination retrieves diverse pmoA-related genes, including the proteobacterial pmoA genes and those from uncultivated methanotrophs believed to be responsible for atmospheric CH4 uptake in upland soils (McDonald et al., 2008). A known problem with these primers is a tendency to co-amplify non-specific sequences, which can make clone libraries useless (Bourne et al., 2001). Non-specific amplification with these primers was also observed in our pyrosequencing data, with an average of 87% of reads from adjacent soils corresponding to non-target reads. The advantage of relatively high number of reads obtainable by pyrosequencing compared with clone libraries meant that sufficient numbers of genuine pmoA sequences were still available to allow for comparisons in pmoA diversity between the samples.

Almost all sequences passing the quality-filtering steps were assigned to seven clades, which were defined and described previously (Lüke and Frenzel, 2011). Representative sequences from each of these clades were added to a database of pmoA and amoa sequences and are shown in a simplified phylogenetic tree (Figure 2). The most abundant clades identified were USCα, tropical upland soil cluster (TUSC) and Methylocystis. The other less abundant clades were RA21, M84-P105, AOB-rel, and the AOB-like group. AOB-rel is also referred to in the literature as Cluster 1 (Köhl et al., 2005).

The relative abundance of the clades from each of the sites is shown in Figure 3. A test of proportions indicated that, with the exception of AOB-like sequences, the relative abundances of these clades were significantly different (p < 0.05) between the ADE and the adjacent soils (Table S1).

COMPARISON OF RELATIVE pmoA GENE ABUNDANCES OBTAINED BY qPCR AND PYROSEQUENCING

Data from the pmoA qPCR assays and amplicon pyrosequencing approaches provided independent numbers to compare the relative abundance of pmoA clades in the soils. Based on the diversity of pmoA detected by pyrosequencing, Methylocystis was the only group present that was a target for the A189f-mb661r pmoA qPCR assay. Therefore, the abundance of pmoA detected with this qPCR assay was taken as the abundance of Methylocystis pmoA genes. Calculating the relative abundance of Methylocystis and USCα from the qPCR assays (Figure 4A) and the pyrosequencing dataset (Figure 4B) showed relatively good agreement.
Table 1 | Soil chemical properties of Amazonian Dark Earth (ADE) and their adjacent (ADJ) soils under secondary forest and manioc cultivation.

| Soil properties | Amazonian Dark Earth | Adjacent soil | Statistics |
|-----------------|----------------------|---------------|------------|
|                 | Secondary forest     | Manioc plantation |             |            |
|                 | Barro Branco Caldeirão | Barro Branco Caldeirão |             |            |
|                 | Secondary forest     | Manioc plantation |             |            |
|                 | Barro Branco Caldeirão | Barro Branco Caldeirão |             |            |
| **A**           | 0.03 ± 0.03³        | 0.01 ± 0      | 0.09 ± 0.08 | 0.16 ± 0.13 | 2.01 ± 0.26 | 1.80 ± 0.08 | 2.51 ± 0.35 | 1.37 ± 0.08 | ** ***c |
| **Ca**          | 5.19 ± 1.35         | 2.79 ± 0.50   | 3.45 ± 0.41 | 3.36 ± 0.11 | 0.17 ± 0.06 | 0.09 ± 0.04 | 0.23 ± 0.17 | 0.32 ± 0.06 | ***          |
| **CEC[^]**      | 6.64 ± 1.24         | 3.49 ± 0.51   | 4.50 ± 0.47 | 5.64 ± 0.29 | 2.32 ± 0.23 | 2.06 ± 0.11 | 2.90 ± 0.17 | 1.89 ± 0.03 | ***          |
| **Fe**          | 5.00 ± 1.00         | 26.33 ± 1.53  | 13.00 ± 3.46| 51.33 ± 3.51| 112.33 ± 19.30| 313.00 ± 53.26| 75.33 ± 31.21| 259.67 ± 26.84| **            |
| **K**           | 21.00 ± 1.00        | 44.67 ± 5.03  | 20.33 ± 3.79| 20.33 ± 6.50| 15.33 ± 1.15| 19.67 ± 5.69| 16.33 ± 0.58| 1767 ± 3.51| *            |
| **Mg**          | 136 ± 0.09          | 133 ± 0.26    | 0.90 ± 0.16 | 1.08 ± 0.31 | 0.09 ± 0.01 | 0.08 ± 0.03 | 0.09 ± 0.03 | 0.15 ± 0.01 | ***          |
| **Mn**          | 48.65 ± 9.40        | 52.1 ± 12.04  | 31.63 ± 3.24| 22.02 ± 2.85| 3.9 ± 0.04  | 1.88 ± 0.75 | 4.93 ± 0.33 | 1.84 ± 0.29 | **            |
| **P**           | 26.00 ± 9.54        | 56.33 ± 12.06 | 51.33 ± 13.86| 73.33 ± 4.50| 6.00 ± 0.64 | 5.67 ± 1.53 | 5.33 ± 2.08 | 133 ± 0.58 | ** ***       |
| **pH<sub>water</sub>** | 5.63 ± 0.13          | 5.85 ± 0.27   | 5.53 ± 0.13 | 5.30 ± 0.29 | 4.33 ± 0.06 | 3.84 ± 0.09 | 4.38 ± 0.08 | 4.33 ± 0.13 | **            |
| **SOC**         | 29.90 ± 1.39        | 32.31 ± 3.36  | 24.7 ± 2.50 | 13.15 ± 1.34| 19.23 ± 0.97| 11.72 ± 1.88| 21.2 ± 1.11| 10.84 ± 0.25| **            |
| **Total C**     | 3.23 ± 0.45         | 2.99 ± 0.32   | 3.07 ± 0.33 | 2.14 ± 0.05 | 2.71 ± 0.43 | 2.12 ± 0.36 | 2.33 ± 0.55 | 1.81 ± 0.26 | *            |
| **Total N**     | 0.24 ± 0.08         | 0.25 ± 0.04   | 0.24 ± 0.02 | 0.16 ± 0.01 | 0.19 ± 0.02 | 0.17 ± 0.02 | 0.16 ± 0.02 | 0.14 ± 0.02 | ns            |
| **Zn**          | 10.74 ± 1.96        | 6.76 ± 0.49   | 4.40 ± 0.69 | 2.32 ± 0.28 | 5.63 ± 1.15 | 0.32 ± 0.03 | 5.63 ± 1.15 | 0.51 ± 0.16 | **            |

[^]: Cation exchange capacity.

[^a]: Al, Ca, CEC, and Mg are expressed in centimoles per cubic decimeter; Fe, K, Mn, P, and Zn are expressed in milligram per cubic decimeter; soil organic C (SOC) is expressed in gram per kilogram; Total C and Total N in percentage.

[^b]: Values are means (n = 3) followed by the standard deviation.

[^c]: ANOVA, n = 12, *p < 0.05, **p < 0.01, ***p < 0.001, ns indicates p ≥ 0.05.
Table 2 | CH₄ oxidation rates in Amazonian Dark Earth and their adjacent soils under secondary forest and manioc cultivation.

| Soil type and site | Land use             | CH₄ oxidation rate [pmol of CH₄ (g dw)⁻¹ h⁻¹] |
|--------------------|----------------------|---------------------------------------------|
| Amazonian Dark Earth |                     |                                             |
| Barro Branco       | Secondary forest     | 33.5 ± 1.6                                  |
| Caldeirão           | Secondary forest     | 48.1 ± 4.5                                  |
| Barro Branco       | Manioc plantation    | 50.0 ± 1.4                                  |
| Caldeirão           | Manioc plantation    | 6.0 ± 1.1                                   |
| Adjacent soil       |                     |                                             |
| Barro Branco       | Secondary forest     | 31.0 ± 1.9                                  |
| Caldeirão           | Secondary forest     | 21.0 ± 1.7                                  |
| Barro Branco       | Manioc plantation    | 8.0 ± 1.9                                   |
| Caldeirão           | Manioc plantation    | 6.1 ± 2.2                                   |

*Errors are standard deviation (n = 3).

The major difference between these data was that USCα in the cultivated adjacent soils was below the detection limit of the qPCR assay (Figure 1B) and therefore its relative abundance was calculated as 0 (Figure 4A); however, USCα sequences were detected of ~20% of Methylocystis in the pyrosequencing dataset from these samples (Figure 4B).

DISCUSSION

Many processes, such as CH₄ oxidation, are crucial for soil ecosystem functioning and have an impact on global biogeochemistry. Forest soils in particular have been identified as an efficient sink for atmospheric CH₄ and are highly sensitive to land use change (Dunfield, 2007). Here, we have characterized methanotrophs in ADE and their adjacent soils (Haplic Acrisol and Oxisol) under two different land uses (i.e., secondary forest and manioc cultivation). These approaches showed two major outcomes with respect to ADE soils: (1) high CH₄ oxidation rates were observed in three of four ADE soils examined, and (2) high relative and absolute abundances of methanotrophs belonging to the USCα pmxA cluster associated with atmospheric CH₄ oxidation in upland soils were observed in all ADE soil samples, independent of land use.

CH₄ OXIDATION POTENTIALS

The CH₄ oxidation rates were relatively high in forested sites. This is in agreement with other studies of tropical forests soils (Verchot et al., 2000; Veldkamp et al., 2008; Zhang et al., 2008; Dörr et al., 2010). Surprisingly, the ADE soil at the Barro Branco site under manioc cultivation showed a CH₄ oxidation rate similar to that of the forested sites. Many studies have shown that conversion of forest to agriculture diminishes CH₄ uptake. For example, after 2 years of agriculture a Norwegian soil showed a fivefold decrease in CH₄ oxidation rate (Jensen and Olsen, 1998). At the time of sampling, the ADE soil at Barro Branco had been used for manioc cultivation for ~5 years, suggesting that it too should have shown a decreased CH₄ oxidation potential. The ADE soil at the manioc plantation at the Caldeirão area, which has a longer history of cultivation, showed a decreased CH₄ oxidation potential. The cultivated ADE site at Barro Branco had been burned to clear the land, which may have also influenced in CH₄ oxidation capacity as in some cases fire has been shown stimulate atmospheric CH₄ oxidation (Jaatinen et al., 2004).

ABUNDANCE AND COMMUNITY COMPOSITION OF METHANOTROPHS

Differences in the methanotroph communities were found between ADE and adjacent soils under secondary forest and manioc cultivation, indicating that the methanotrophic community is altered depending on soil type and land use. USCα were the predominant methanotrophs in all ADE soils and the forested adjacent soils. This group is as yet uncultivated, but is believed to be responsible for atmospheric CH₄ consumption in many forest soils (Dunfield, 2007; Kolb, 2009; Nazaries et al., 2013). The abundance of USCα pmxA genes was ~1 × 10⁷ per gram dry weight in the ADE soils, which was one-order of magnitude higher than in the forested sites of the adjacent soils. In comparison, the same assay used to quantify USCα in a German forest soil detected ~1 × 10⁶ gene copies per gram dry weight of soil (Kolb...
et al., 2005), suggesting that their abundance in ADE was relatively high. It was surprising that USCα abundances were equally high in the cultivated and forested ADE soils (Figure 1B). This pattern was different for the adjacent soils where they were below the qPCR detection limit in the cultivated soils, indicating abundances at least two-orders of magnitude lower than the forested sites. In comparison, the manioc plantation in ADE soil at Caldeirão has a long history of agriculture use, yet the USCα abundance was only threefold lower than in the corresponding forested soil. To the best of our knowledge, this is the first study to detect a high absolute and relative abundance of USCα in agricultural soils. Priemé et al. (1997) showed that CH4 oxidation rates took more than 100 years to reach pre-cultivation levels and that the highest rates were in the oldest (200 years) woodlands. The apparent resilience of USCα populations in ADE soil compared with other upland soils, possibly from a protective property of ADE, suggests that recovery of CH4 oxidation capacity after agricultural abandonment might be faster in ADE than other types of upland soil.

Also of note in this study was that the CH4 uptakes rates were relatively low in cultivated ADE soil at Caldeirão, but USCα abundance in this soil was relatively high. One possible explanation for this lack of correlation is that USCα methanotrophs can incorporate acetate and possibly other organic carbon substrates (Pratscher et al., 2011), suggesting that CH4 oxidation is a facultative trait in these organisms and CH4 is oxidized only under certain conditions. Evidence that USCα are not obligate methanotrophs include reported failures to sufficiently label their nucleic acids with 13CH4 for stable isotope probing (Bengtson...
methylotrophs; however, the unconventional M84-P105 \( pnxA \) sequences, which have been shown to belong to members of the \textit{Methylococcus} (Tavormina et al., 2011), were detected in cultivated adjacent soils suggesting a low abundance of these methylotrophs in some soils (Figure 3). Although the relative abundance of \textit{Methylocystis} was high in the adjacent soils from manioc plantation sites (Figure 3), no difference in their absolute abundance between ADE and adjacent soil, or between forested and cultivated sites was observed at this sampling time (Figure 1). \textit{Methylocystis} have been shown to be important consumers of CH\(_4\) in hydromorphic soils under dry conditions when CH\(_4\) concentrations are relatively low (Knief et al., 2005). These \textit{Methylocystis} possess an unconventional pMMO gene, termed pMMO2 (Ricke et al., 2004), which is expressed under low CH\(_4\) (Baani and Liesack, 2008). We only detected two \( pmoA \) gene sequences in our pyrosequencing dataset (data not shown), suggesting that conditions in these Amazonian soils at the time of this analysis were not favorable for pMMO2 possessing oligotrophic \textit{Methylocystis} species.

Other \( pmoA \)-related gene sequences were detected, such as TUSC, AOB-rel and AOB-like groups. The AOB-like sequences correspond to the \textit{amoA} genes of \textit{Nitrosospira} and \textit{Nitrosonomas} (Figure 2). In ADE soils, these \textit{amoA} sequences were only detected in plantation soil, which is likely a consequence of enrichment by ammonium fertilizer applied to the soil for manioc cultivation. The TUSC and AOB-rel groups have not been linked to cultivated organisms and the function of the enzyme encoded by these genes is not known (Lüke and Frenzel, 2011). TUSC or “tropical upland soil cluster” is also termed “Cluster 2” elsewhere (Knief et al., 2005). As the name implies, they were found to be abundant in some tropical upland soils (Knief et al., 2005), but have also been detected in temperate forest soil (Knief et al., 2003). It is noteworthy that the relative abundance of TUSC tended to mirror USC\(\alpha\) in these Amazonian soils. One possibility to explain this correlation is that TUSC sequences are a divergent \( pmoA \) gene found in USC\(\alpha\) methylotrophs, such as the case with M84-P105 \( pnxA \) in \textit{Methylomonas} and \( pmoA \) in \textit{Methylocystis}; however, other studies have not observed a correlation between USC\(\alpha\) and TUSC relative abundances (Kolb, 2009; Dörr et al., 2010).

**CONCLUSION**

This study has shown that ADE soils are a potential sink for atmospheric CH\(_4\). The relatively high rate of “high-affinity” CH\(_4\) uptake by the ADE soil with a 5-year history of agriculture contradicts many studies showing the process to be sensitive to land use change. All the ADE soils examined had a high abundance of USC\(\alpha\) methanotrophs (~10\(^7\) \( pmoA \) genes g\(^{-1}\) soil), which was particularly surprising for the ADE soil at the Caldeirão site that had a long history of manioc cultivation. In comparison, the abundance of USC\(\alpha\) methanotrophs was up to 1000-fold lower in adjacent than ADE soil, and both the adjacent soils used for agriculture displayed relatively low CH\(_4\) uptake rates. This raises the question if USC\(\alpha\) methanotrophs are indeed more resistant to disturbance in ADE than in other upland soils and whether this apparent resilience of ADE extends to the protection of other groups of vulnerable microorganisms and their associated functions.
**ACKNOWLEDGMENTS**

We thank R. B. Correa and I. G. Braga for assistance with the fieldwork and Prof. R. Conrad for helpful discussions. We also would like to thank W. G. Teixeira and G. C. Martins for soil descriptions and members from the community of Barro Branco. This research was supported by the Max Planck Society, CNPq (Conselho Nacional de Desenvolvimento Científico e Tecnológico), FAPEAM (Fundação de Amparo à Pesquisa do Estado do Amazonas), and EMBRAPA Western Amazon. Amanda B. Lima received a postdoctoral scholarship from CNPq within the program Science Without Borders (CsF).

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://www.frontiersin.org/journal/10.3389/fmicb.2014.00550/abstract

**REFERENCES**

Anderson, C. R., Condon, L. M., Clough, T. I., Fiers, M., Stewart, A., Hill, R. A., et al. (2011). Biochar induced soil microbial community change: implications for biogeochemical cycling of carbon, nitrogen and phosphorus. *Pedobiologia* 54, 309–220. doi: 10.1016/j.pedobi.2011.07.005

Angel, R., and Conrad, R. (2009). In situ measurement of methane fluxes and analysis of transcribed particulate methane monoxygenase in desert soils. *Environ. Microbiol.* 11, 2598–2610. doi: 10.1111/j.1462-2920.2009.01984.x

Baani, M., and Liesack, W. (2008). Two isozymes of particulate methane monoxygenase with different methane oxidation kinetics are found in *Methylocystis* sp. Strain SC2. *Proc. Natl. Acad. Sci. U.S.A.* 105, 10203–10208. doi: 10.1073/pnas.0702643105

Bengtson, P., Basílio, N., Dumont, M. G., Hills, M., Murrell, J. C., Roy, R., et al. (2009). Links between methanotroph community composition and CH4 oxidation in a pine forest soil. *FEMS Microbiol. Ecol.* 70, 356–366. doi: 10.1111/j.1574-6949.2009.00751.x

Berry, D., Mahfoudh, K. B., Wagner, M., and Loy, A. (2011). Barcoded primers used in multiplex amplicon pyrosequencing bias amplification. *Appl. Environ. Microbiol.* 77, 7846–7849. doi: 10.1128/AEM.05220-11

Bodelier, P. L. E., Gillissen, M.-J. B., Hordijk, K., Damsté, J. S. S., Rijpstra, W. I. C., Geenens, W. H., et al. (2010). A reanalysis of phospholipid fatty acids as ecological biomarkers for methanotrophic bacteria. *ISME J.* 3, 606–617. doi: 10.1038/ismej.2009.6

Bourne, D. G., McDonald, I. R., and Murrell, J. C. (2001). Comparison of pmoA PCR primer sets as tools for investigating methanotroph diversity in three Danish soils. *Appl. Environ. Microbiol.* 67, 3802–3809. doi: 10.1128/AEM.67.9.3802-3809.2001

Brossi, M. J. L., Mendes, L. W., Germano, M. G., Lima, A. B., and Taisi, S. M. (2014). Assessment of bacterial *ipn* gene in Amazonian Dark Earth and their adjacent soils. *PLoS ONE* 9:e99597. doi: 10.1371/journal.pone.0099597

Bull, I. D., Parekh, N. R., Hall, G. H., Ineson, P., and Evershed, R. P. (2000). Detection and classification of atmospheric methane oxidizing bacteria in soil. *Nature* 405, 175–178. doi: 10.1038/35012061

Cochrane, T. T., and Sanchez, P. A. (1982). *Land Resources, Soils and their Management in the Amazon Region: A State of Knowledge Report.* Cali: Centro Internacional de Agricultura Tropical-CIAT.

Costa, M. L., and Kern, D. C. (1999). Geochemical signatures of tropical soils with archaeological black earth in the Amazon. *J. Geochim. Explor.* 66, 369–385. doi: 10.1016/S0375-6742(99)00038-2

Costello, A. M., and Lidstrom, M. E. (1999). Molecular characterization of functional and phylogenetic genes from natural populations of methanotrophs in lake sediments. *Appl. Environ. Microbiol.* 65, 5066–5074.

Dalal, R. C., and Allen, D. E. (2008). Greenhouse gas fluxes from natural ecosystems. *Aust. J. Bot.* 56, 369–407. doi: 10.1071/BT07127

Denman, K. L., and Frenzel, P. (2014). Classification of pmoA amplicon pyrosequencing using BLAST and the lowest common ancestor method in MEGAN. *Front. Microbiol.* 5:34. doi: 10.3389/fmicb.2014.00034

Dunfield, P. F. (2007). “The soil methane sink,” in *Greenhouse Gas Sinks*, eds D. Reay, C. N. Hewitt, K. Smith, and J. Grace (Wallingford: CABI Publishing).

Edgar, R. C., Haas, B. J., Clemente, J. C., Quince, C., and Knight, R. (2011). UCHIME improves sensitivity and speed of chimera detection. *Bioinformatics* 27, 2194–2202. doi: 10.1093/bioinformatics/btr381

Embrapa. (1997). *Manual de Métodos de Análise de Solo de Rio de Janeiro: Embrapa-CNPS.*

FAO. (1998). *World Reference Base for Soil Resources.* World Soil Resources Report 84. FAO. Rome: UN Food and Agriculture Organization.

Glasner, B. (1996). Eigenschaften und Staabilität des Humuskörpers der Indianer-schwarzerden Amazoniens. *Bayreserter Bodenkundliche Ber.* 68, 1–196.

Glaser, B. (1999). Eigenschaften und Staabilität des Humuskörpers der Indianer-schwarzerden Amazoniens. *Bayreserter Bodenkundliche Ber.* 68, 1–196.

Haas, B. J., Kissner, K. M., Beck, J. A., and Beever, D. A. (2010). Community composition of methanotrophic bacteria in a soil from a California agricultural field. *Appl. Environ. Microbiol.* 76, 7530–7537. doi: 10.1128/AEM.00282-09

Keller, M., Mitre, M. E., and Stallard, R. F. (1990). Consumption of atmospheric methane in soils of central Panama: effects of agricultural development. *Global Biogeochem. Cycles* 4, 21–27. doi: 10.1029/90GC00574

King, G. M. (1992). Ecological aspects of methane consumption, a key determinant of global methane dynamics. *Adv. Microb. Ecol.* 12, 451–468. doi: 10.1016/0066-6864(92)80021-9

Kuehn, C., and Dunfield, P. F. (2005). Response and adaptation of methanotrophic bacteria to low methane concentrations. *Environ. Microbiol.* 7, 1307–1317. doi: 10.1111/j.1462-2920.2005.00814.x

Kuehn, C., Kolb, S., Bodelier, P. E. L., Lipski, A., and Dunfield, P. F. (2006). The active methanotrophic community in hydromorphic soils change in response to changing methane concentration. *Environ. Microbiol.* 8, 312–333. doi: 10.1111/j.1462-2920.2005.00898.x

Kuehn, C., Lipski, A., and Dunfield, P. F. (2003). Diversity and activity of methanotrophic bacteria in different upland soils. *Appl. Environ. Microbiol.* 69, 6703–6714. doi: 10.1128/AEM.69.11.6703-6714.2003

Kuehn, C., VanIterson, S., Harvey, N. W., Conrad, R., Dunfield, P. F., and Chidhasnong, A. (2005). Diversity of methanotrophic bacteria in tropical upland soils.
soils under different land use. *Appl. Environ. Microbiol.* 71, 3826–3831. doi: 10.1128/AEM.71.7.3826-3831.2005

Klob, S. (2009). The quest for atmospheric methane oxidizers in forest soils. *Environ. Microbiol. Rep.* 1, 336–346. doi: 10.1111/j.1758-2229.2009.00047.x

Kolb, S., Knief, C., Dunfield, P. F., and Conrad, R. (2005). Abundance and activity of uncultured methanotrophic bacteria involved in the consumption of atmospheric methane in two forest soils. *Environ. Microbiol.* 7, 1150–1161. doi: 10.1111/j.1462-2920.2005.00791.x

Kolb, S., Knief, C., Stuhner, S., and Conrad, R. (2003). Quantitative detection of methanotrophs in soil by novel pmoA-targeted real-time PCR assays. *Appl. Environ. Microbiol.* 69, 2423–2429. doi: 10.1128/AEM.69.5.2423-2429.2003

Lehmann, J. (2007). Bio-energy in the black. *Front. Ecol. Environ.* 5:381–387. doi: 10.1890/1540-9295(2007)5[381:BTB]2.0.CO;2

Levine, U. Y., Teal, T. K., Robertson, G. P., and Schmidt, T. M. (2011). Agricultural impact on microbial diversity and associated fluxes of carbon dioxide and methane. *ISME J.* 5, 1683–1691. doi: 10.1038/ismej.2011.40

Ludwig, W., Strunk, O., Westram, R., Richter, L., Meier, H., Yadskurukam, et al. (2004). ARB a software environment for sequence data. *Nucleic Acids Res.* 32, 1363–1371. doi: 10.1093/nar/gkh293

Luke, C., and Frenzel, P. (2011). Potential of pmoA amplicon pyrosequencing for methanotroph diversity studies. *Appl. Environ. Microbiol.* 77, 6305–6309. doi: 10.1128/AEM.05355-11

McDonald, I. R., Bodrossy, L., Chen, Y., and Murrell, C. (2008). Molecular ecology techniques for the study of aerobic methanotrophs. *Appl. Environ. Microbiol.* 74, 1305–1315. doi: 10.1128/AEM.02233-07

Mosier, A., Schimel, D., Valentine, D., Bronson, K., and Parton, W. (1991). Methane and nitrous oxide fluxes in native, fertilized and cultivated grasslands. *Nature* 350, 330–332. doi: 10.1038/350330a0

Navarrete, A. A., Cannavan, F. S., Taketani, R. G., and Tsai, S. M. (2010). A molecular survey of the diversity of microbial communities in different Amazon agricultural model systems. *Diversity* 2, 787–809. doi: 10.3390/d20405078

Nazaries, L., Pan, Y., Bodrossy, L., Bags, E. M., Millard, P., Murrell, J. C., et al. (2013). Evidence of microbial regulation of biogeochemical cycles from a study on methane flux and land use change. *Appl. Environ. Microbiol.* 79, 4031–4040. doi: 10.1128/AEM.00895-13

Neher, D. A. (1999). Soil community composition and ecosystem processes compares to native methane-producing systems. *Agrofor. Syst.* 45, 159–185. doi: 10.1023/A:1006299100678

Pratscher, J., Dumont, M. G., and Conrad, R. (2011). Assimilation of acetate by the methanotroph Methylocapsa acidiphila. *Appl. Environ. Microbiol.* 77, 1032-1037. doi: 10.1128/AEM.00791-10

Ricke, P., Eikel, C., Kube, M., Reinhardt, R., and Liesack, W. (2004). Comparative analysis of the conventional and novel pmoA (particulate methane monoxygenase) operons from Methylosinus strain SC2. *Appl. Environ. Microbiol.* 70, 3055–3063. doi: 10.1128/AEM.70.5.3055-3063.2004

Ricke, P., Kube, M., Nakagawa, S., Eikel, C., Reinhardt, R., and Liesack, W. (2005). First genome data from uncultured upland soil cluster alpha methanotrophs provide further evidence for a close phylogenetic relationship to Methylocapsa acidiphila B2 and for high-affinity methanotrophy involving particulate methane monoxygenase. *Appl. Environ. Microbiol.* 71, 7472–7482. doi: 10.1128/AEM.71.11.7472-7482.2005

Schloss, P. D., Westcott, S. L., Ryabin, T., Hall, J. R., Hartmann, M., Hollister, E. B., et al. (2009). Introducing mothur: open-source, platform-independent, community-supported software for describing and comparing microbial communities. *Appl. Environ. Microbiol.* 75, 7537–7541. doi: 10.1128/AEM.01541-09

Semrau, J. D., DiSpito, A. A., and Yoon, S. (2010). Methanotrophs and copper. *FEMS Microbiol. Rev.* 34, 496–531. doi: 10.1111/j.1574-6976.2010.00212.x

Shrestha, P. M., Kammann, C., Lenhart, K., Dam, B., and Liesack, W. (2012). Linking activity, composition and seasonal dynamics of atmospheric methane oxidizers in a meadow soil. *ISME J.* 6, 1115–1126. doi: 10.1038/ismej.2011.179

Smith, N. J. H. (1980). Anthrosols and human carrying capacity in Amazonia. *Ann. Assoc. Am. Geogr.* 70, 553–566. doi: 10.1111/j.1467-8306.1980.tb01332.x

Sombroek, W., De Lourdes Ruivo, M., Fearnside, P., Glaser, B., and Lehmann, J. (2003). “Amazonian dark earths as carbon stores and sinks,” in Amazonian Dark Earths: Origin, Properties, Management. eds I. Lehmann, D. C. Kern, B. Glaser, and W. I. Woods (Dordrecht: Kluwer Academic Publishers), 141–158.

Taketani, R. G., Lima, A. B., Jesus, E. C., Teixeira, W. G., Tiedje, E. C., and Tsai, S. M. (2013). Bacterial community composition of anthropogenic biochar and Amazonian anthrosols assessed by 16S rRNA gene 454 pyrosequencing. *Antonie van Leeuwenhoek* 104, 235–242. doi: 10.1007/s10482-013-9942-0

Tamas, I., Smirnova, A. V., He, Z., and Dunfield, P. F. (2014). The (d)evolution of methanotrophy in the Beijerinckiacae—a comparative genomics analysis. *ISME J.* 8, 369–382. doi: 10.1038/ismej.2013.145

Tavormina, P. L., Orphan, V. J., Kalyuzhnaya, M. G., Jetten, M. S. M., and Klotz, M. G. (2011). A novel family of functional operons encoding methane/ammonia monoxygenase-related proteins in gammaproteobacterial methanotrophs. *Environ. Microbiol. Rep.* 3, 91–100. doi: 10.1111/j.1758-2229.2010.00192.x

Veldkamp, E., Purbopuspito, J., Corre, M. D., Brumme, R., and Murdiyarso, D. (2008). Land use change effects on trace gas fluxes in the forest margins of Central Sulawesi, Indonesia. *J. Geophys. Res.* 113, G02003. doi: 10.1029/2007JG005522

Verchot, L. V., Davidson, E. A., Cattinio, J. H., and Ackerman, I. L. (2000). Land-use change and biogeochemical controls of methane fluxes in soils of Eastern Amazonia. *Ecosystems* 3, 41–56. doi: 10.1007/s1002100000009

Woods, W. I., and McCann, J. M. (1999). The anthropogenic origin and persistence of Amazonian Dark Earths. *Yesh. Conf. Lat. Am. Geogr.* 25, 7–14.

Zhang, W., Mo, J., Zhou, L., Gunderen, P., Fang, Y., Lu, X., et al. (2008). Methane uptake responses to nitrogen deposition in three tropical forests in southern China. *J. Geophys. Res. Atmos.* 113, D11116. doi: 10.1029/2007JD009193

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.