METHANOTROPHS RESPONSIBLE FOR METHANE OXIDATION IN NATURAL PEATS FROM POLESIE LUBELSKIE REGION

Anna Szafranek-Nakonieczna, Zofia Stepniewska, Agnieszka Wołoszyn, Jakub Ciepielski

Department of Biochemistry and Environmental Chemistry, The John Paul II Catholic University, Al. Krasinska 102, 20-718 Lublin, Poland
e-mail: anna.szafranek@kul.lublin.pl

Abstract. The potential of methanotrophic activity (MTA) has been investigated under laboratory conditions in three types of peatland profiles: high (H), transition (T) and low (L) originating from Polesie Lubelskie Region. Selected peat samples differed in respect of pH, TOC, von Post index and moisture. The experiment was conducted at natural moisture (198-719 %w/w) with different ranges of both, temperature (5, 10 and 20°C) and CH4 enrichment (1 and 5%v/v). The highest MTA (19.69-155.79 mg CH4 kg D.W.⁻¹ d⁻¹) was observed at 20°C. Regardless of temperature, MTA was lower (1.38-51.16 mg CH4 kg D.W.⁻¹ d⁻¹) when peat samples were incubated in atmosphere enriched in 1% than in 5% CH4 v/v (4.75-191.26 mg CH4 kg D.W.⁻¹ d⁻¹). Strong influence of temperature and sampling sites on MTA was also noted. Total DNA was isolated from the most active (20°C, 5% CH4 v/v) peat samples from each site and the PCR (polimerase chain reaction) amplifying of genes pmoA (primers A189f/mb661r) and sequence 16S rRNA (primers Type If /Type Ir and Type IIf/Type IIr) specific for methanotrophic bacteria were carried out. Positive results of PCR with primers of pmoA gene after sequencing confirmed that methanotrophs from L point belong to family Methylococcaceae, while 16S rRNA gene sequences from microorganisms inhabiting H peat demonstrated the highest similarity to genus Methylocystis and Methylosinus.

Keywords: methanotrophic activity, methanotrophs, peat soil

INTRODUCTION

Methanotrophs, microorganisms involved in aerobic methane oxidation, focused a great interest of scientists and thus extended our knowledge about global methane cycling. Those bacteria grow on methane as a sole source of carbon and

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energy. The important role of methanotrophs in the environment results from the fact that up to 90% of methane produced by methanogens (methane producing Archaea) may be oxidised by methanotrophs before its release into the atmosphere. Biological methane oxidation is found in many places such as wetlands, soils, freshwaters, sediments, rice plantations and some extreme environments (Chen and Murrell 2010). A large group of environments favourable for methanotrophs are peatlands, especially their aerobic parts.

Methanotrophs can be broadly divided into Type I (Gamma-Proteobacteria) and Type II (Alpha-Proteobacteria). The first can be further subdivided into two classes: Type Ia and Type Ib, also referred to as Type X (Dumont et al. 2011). The types of methanotrophs differ in many characteristics including (a) carbon assimilation pathways (Type Ia uses ribulose monophosphate pathway, Type Ib ribulose monophosphate or serine pathway, while Type II uses only serine pathway) and (b) the arrangement of intracellular membranes (Heyer et al. 2002, Murrell 2010). Type Ia is represented by genera Methylococcus, Methylocaldum, Methylosporomonas, Methylophilum, Methylobacter, Methylosphaera, Methylosarcina. Type Ib contains the genera of Methylocaldum and Methylococcus, while genera of Methylocystis, Methylosinus, Methylocella, and Methycapsa are representatives of Type II (Dumont et al. 2011).

Even though the Type I group is more phylogenetically diverse, the members of Type II are more often isolated from natural systems, or detected using cultivation independent molecular techniques (Heyer et al. 2002). Quantitative studies suggest that methanotrophic microorganisms from Alpha-Proteobacteria are numerically predominant also in peatlands (Dunfield and Dedysh 2010).

Methane is converted by methanotrophs to methanol through catalysis by the methane monooxygenase enzyme (MMO), subsequently oxidised by methanol dehydrogenase, formaldehyde dehydrogenase and formate dehydrogenase to produce CO₂ and H₂O. Generally, methanotrophs can express a membrane-bound MMO termed particulate methane monooxygenase (pMMO), however when copper ion is unavailable, some strains are able to express an iron-containing soluble methane monooxygenase (sMMO) to catalyse the hydroxylation of methane to methanol (Hua et al. 2007). The pMMO is therefore the universal enzyme of all methanotrophs, and recognised genes encoding that enzyme are used as phylogenetic markers. One of them is the pmoA gene which encodes the active-site-containing subunit of the particulate methane monooxygenase enzyme (Heyer et al. 2002). The next most obvious marker for detecting methanotrophs in various environments is the 16S rRNA gene, due to the large database of sequences available.

In relation to preferred methane concentration, methanotrophs are divided into two groups: methanotrophs that can survive under atmospheric concentrations of methane (high affinity methanotrophs) and methanotrophs surviving only in methane presence at >40 ppmv (low affinity methanotrophs) (Chen and Murrell 2010).
The aim of the present work was examination the peat soil profiles (0-80 cm) originating from three different types of peatland characteristic for Poland in respect of their ability of methane oxidation. The second objective was an attempt to identify methanotrophs responsible for these activities. The sequences of pmoA and 16S rRNA genes specific primers were used for amplification.

MATERIALS AND METHODS

Soil characteristics

The three plots used in this study were selected in Polesie Lubelskie Region (Eastern Poland). The first site, defined as a low moor peatland (L) with numerous pools, fed by shallow groundwater was situated west of Garbatówka village (51°21’ N, 23°7’ E). Its vegetation mainly consisted of Carex spp., Juncus spp., Schoenoplectus spp. and various mosses.

The next studied point (51°25’ N, 23°3’ E) was placed close to Orłowskie Peatland, representing a transition moor peat (T), fed by ground water and rainfall, and adjoining with Lake Łukie. Plant cover was represented by Poa pratensis, Festuca rubra, Deschampsia caespitose, Rumex acetosa and locally shrubs with Salix lapponum and Salix myrtilloides.

The last site (51°27’ N, 23°6’ E), a high moor peat (H), was located on the south of Lake Moszne, and characterised by lacustrine genesis, with water supply only by precipitation. Therefore, in hot and drought seasons the groundwater table considerably decreases. The vegetation was composed of Vaccinium uliginosum, Ledum palustre, Eriophorum vaginatum, a great number of peat mosses (e.g. Sphagnum acutifolium, Sphagnum magellanicum), shrubs and trees: Pinus sylvestris, Betula pubescens, Quercus robur, Populus tremula, Betula pendula (Wojciechowski and Szczurowska, 2002; Radwan, 2003). Additional soil characteristics are presented in Table 1.

Soil moisture was determined gravimetrically by drying peat samples at 105°C (24h). Soil pH was determined potentiometrically, in soil–water mixture (1/1, w/w) using a pH electrode and pIONeer 65 meter (Radiometer Analytical S.A., France). Total organic carbon was determined in dry peat samples by combustion and the analysis of evolved CO2 by means of TOC-VCSH with module SSM-5000A (Shimadzu, Japan).

Methane and oxygen concentrations were measured using a gas chromatograph (GC 3800, Varian, USA) equipped with flame ionisation (FID) and thermal conductivity (TCD) detectors, according to methods described by Szafranek-Nakonieczna and Bencicelli (2010).
Table 1. Characteristic of investigated peat materials, ±SD

<table>
<thead>
<tr>
<th>Site</th>
<th>Peat layer (cm)</th>
<th>pH (H2O)</th>
<th>Von Post ranking (%)</th>
<th>Moisture (% w/w)</th>
<th>Bulk density (g cm⁻³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>L</td>
<td>0-20</td>
<td>6.93±0.02c</td>
<td>H₂a</td>
<td>592.9±0.1c</td>
<td>0.708±0.02a</td>
</tr>
<tr>
<td></td>
<td>20-40</td>
<td>6.80±0.02b</td>
<td>H₂a</td>
<td>530.1±0.1a</td>
<td>0.743±0.03b</td>
</tr>
<tr>
<td></td>
<td>40-60</td>
<td>6.83±0.02b</td>
<td>H₂b</td>
<td>582.4±0.2b</td>
<td>0.859±0.05c</td>
</tr>
<tr>
<td></td>
<td>60-80</td>
<td>6.75±0.02a</td>
<td>H₂b</td>
<td>667.1±0.3d</td>
<td>0.879±0.01c</td>
</tr>
<tr>
<td>T</td>
<td>0-20</td>
<td>6.17±0.02a</td>
<td>H₂b</td>
<td>284.1±0.1a</td>
<td>1.05±0.01a</td>
</tr>
<tr>
<td></td>
<td>20-40</td>
<td>6.44±0.02b</td>
<td>H₂b</td>
<td>400.0±0.1b</td>
<td>1.055±0.01a</td>
</tr>
<tr>
<td></td>
<td>40-60</td>
<td>6.52±0.02c</td>
<td>H₂c</td>
<td>563.2±0.1c</td>
<td>1.189±0.03b</td>
</tr>
<tr>
<td></td>
<td>60-80</td>
<td>6.53±0.02c</td>
<td>H₂d</td>
<td>573.1±0.1d</td>
<td>1.227±0.06b</td>
</tr>
<tr>
<td>H</td>
<td>0-20</td>
<td>2.87±0.01a</td>
<td>H₃a</td>
<td>198.1±0.2a</td>
<td>0.578±0.02a</td>
</tr>
<tr>
<td></td>
<td>20-40</td>
<td>2.98±0.01b</td>
<td>H₃b</td>
<td>315.2±0.1b</td>
<td>0.717±0.04b</td>
</tr>
<tr>
<td></td>
<td>40-60</td>
<td>3.36±0.01c</td>
<td>H₃c</td>
<td>667.1±0.2c</td>
<td>0.736±0.01b</td>
</tr>
<tr>
<td></td>
<td>60-80</td>
<td>4.43±0.01d</td>
<td>H₃d</td>
<td>719.1±0.1d</td>
<td>0.824±0.01c</td>
</tr>
</tbody>
</table>

- numbers characterising the properties of peat from the one location, followed by the same letter, do not differ significantly (P>0.05).

**Incubation experiments**

The effects of elevated levels of CH₄ (1 and 5%CH₄ v/v), temperature (5, 10, 20°C) and position in peat profile (from top down to 80 cm) on the potential of peat for methane oxidation were tested via laboratory incubation experiments. The peat samples were collected from separated layers (0–20, 20–40, 40–60 and 60–80 cm) of plots L, T and H. Next, they were placed (10 g of fresh peat) in dark incubation bottles (60 ml), tightly sealed with rubber plug and aluminium cap. For each combination three replicates were prepared. Next, the headspace of each unit was enriched with methane (1 or 5% CH₄ v/v). Incubation was performed until substrate depletion. At the end of the incubation, MTA was calculated and expressed as mg CH₄ kg D.W.⁻¹ d⁻¹.

**DNA extraction and electrophoretic check of the amount of isolated DNA**

Total DNA was extracted from the samples (0.5 g of peat soil) selected from material the most active in methane oxidation. The extraction procedure was performed stepwise according to the method described by Sambrook and Russel (2001).
Electrophoresis was performed in agarose gel 1% (w/v) in 1-fold concentrated TAE buffer and 1-fold TAE buffer, at a voltage of 5 V/cm of distance between the electrodes. As a marker, DNA Ladders MassRuler were used. The gel was stained in 0.01% aqueous solution of ethidium bromide and observed under UV lamp (Vilber Larmart, TFX-20M, 6x15W-312 nm tube, France) (Sambrook and Russel 2001).

**PCR reaction**

In order to confirm the presence of the *pmoA* gene encoding the active site of subunit molecular methane monooxygenases (pMMO) a universal enzyme of methanotrophs and 16S rRNA gene sequences characteristic for methanotrophs of Type I and Type II, pairs of primers were used (as is shown in Table 2).

**Table 2. Primers used in the study**

<table>
<thead>
<tr>
<th>Target gene</th>
<th>Primer</th>
<th>Sequence (5’-3’)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mb661r</td>
<td>CCG GMG CAA CGT CYT TAC C</td>
<td>Costello and Lidstrom 1999, Liebner <em>et al.</em> 2009</td>
</tr>
<tr>
<td>16S rRNA</td>
<td>Type II</td>
<td>GGG MGA ATA ATG ACG GTA CCW GGA</td>
<td>McDonald <em>et al.</em> 2008</td>
</tr>
<tr>
<td>methanotrophs</td>
<td>Type IIr</td>
<td>GTC AAR AGC TGG TAA GGT TC</td>
<td>McDonald <em>et al.</em> 2008</td>
</tr>
<tr>
<td>Type I</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Type II</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16S rRNA</td>
<td>Type I</td>
<td>ATG CTT AAC ACA TGC AAG TCG AAC G</td>
<td>McDonald <em>et al.</em> 2008, Chen <em>et al.</em> 2007</td>
</tr>
<tr>
<td>methanotrophs</td>
<td>Type I</td>
<td>CCA CTG GTG TTC CTT CMG AT</td>
<td>McDonald <em>et al.</em> 2008, Chen <em>et al.</em> 2007</td>
</tr>
</tbody>
</table>

Polymerase chain reaction (PCR) amplifications were performed in 50 µl (total volume) using PCR master Mix (Fermentas) with 3.5 µl of template DNA in MJmini thermal cycler (Bio-Rad). Amplification was carried out as follows: initial denaturation (96°C for 4 minutes), followed by 30 cycles of denaturation (94°C for 2 min), primer annealing (56°C for Type I, 55°C for Type II and 53°C for *pmoA* each for 1 min), elongation (72°C for 1 min). Final elongations at 72°C were performed for 3 minutes (Sambrook and Russel 2001).
Sequencing

The sequencing of PCR products was performed in the Laboratory of DNA Sequencing and Oligonucleotide Synthesis of the Institute of Biochemistry and Biophysics, Polish Academy of Science, in Warsaw. The sequencing of PCR products were carried out with pairs of primers used for amplification.

Analysis of similarities

In order to determine the similarity of sequences of obtained fragments *pmoA* and 16S rRNA genes with other known methanotrophic microorganisms, interactive tool from the NCBI, BLASTN was used (Blast Local Alignment Search Tool).

RESULTS

Methane consumption ranged from 1.38 to 191.3 mg CH₄ kg D.W.⁻¹ d⁻¹ in the aerobic treatments. The highest oxidation was observed in the peat from site H, in the layer of 40-60 cm, incubated at 20°C. Horizontal distribution of MTA in all investigated peat profiles was observed. It varied depending on substrate availability and temperature of incubation (Tab 3).

Samples incubated with 1% CH₄ (v/v) showed the highest activity in the 20-40 cm layer of L location, independently of temperature, whereas MTA decreased in lower layers. The highest MTA characterised the layer of 20-40 cm at 10 and 20°C in T location, while at 5°C the maximal value, above 10 mg CH₄ kg D.W.⁻¹ d⁻¹, was observed in the layer of 60-80 cm. A different situation was found in site H. At 5°C MTA increased with depth while at 10 and 20°C it rose only to 40-60 cm and next it declined slightly, but significantly (Tab. 3).

We assumed that the increase of methane availability modified MTA. In the samples from L site MTA decreased with depth when temperature was higher. The highest MTA was noted in layers of 0-20 cm and 40-60 cm for temperatures of 5 and 10°C, respectively. However, when the temperature increased to 20°C the deepest (60-80 cm) part of the peat profile demonstrated the highest MTA. In the material from site T the highest activity was noted in the surface layer (0-20 cm), irrespective of the temperature, while in H it was found in the layer of 40-60 cm.

Peat samples from particular locations differ in their total organic carbon (TOC) content. The highest value was measured in H, while the lowest in L (Fig. 1A). The sampling site determined also the peat ability for methane oxidation. The highest potential, exceeding 50 mg CH₄ kg D.W.⁻¹ d⁻¹, characterized the L location (data from depth of 0-80 cm). In T and H sites MTA was at comparable level of 30-35 mg CH₄ kg D.W.⁻¹ d⁻¹ (Fig. 1B).
METHANOTROPHS RESPONSIBLE FOR METHANE OXIDATION

Table 3. Methanotrophic activity in investigated peat profiles under selected environmental conditions

<table>
<thead>
<tr>
<th>Site</th>
<th>Depth (cm)</th>
<th>1% CH₄ (v/v)</th>
<th>5% CH₄ (v/v)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>5°C</td>
<td>10°C</td>
</tr>
<tr>
<td>L</td>
<td>0-20</td>
<td>7.80a</td>
<td>15.98b</td>
</tr>
<tr>
<td></td>
<td>20-40</td>
<td>32.46d</td>
<td>45.51c</td>
</tr>
<tr>
<td></td>
<td>40-60</td>
<td>12.65b</td>
<td>13.99b</td>
</tr>
<tr>
<td></td>
<td>60-80</td>
<td>17.95c</td>
<td>5.51a</td>
</tr>
<tr>
<td>T</td>
<td>0-20</td>
<td>5.12b</td>
<td>6.48b</td>
</tr>
<tr>
<td></td>
<td>20-40</td>
<td>8.43c</td>
<td>16.28d</td>
</tr>
<tr>
<td></td>
<td>40-60</td>
<td>4.08a</td>
<td>7.63c</td>
</tr>
<tr>
<td></td>
<td>60-80</td>
<td>10.15d</td>
<td>3.34a</td>
</tr>
<tr>
<td>H</td>
<td>0-20</td>
<td>1.38a</td>
<td>2.96a</td>
</tr>
<tr>
<td></td>
<td>20-40</td>
<td>3.44b</td>
<td>7.41c</td>
</tr>
<tr>
<td></td>
<td>40-60</td>
<td>5.40c</td>
<td>23.45d</td>
</tr>
<tr>
<td></td>
<td>60-80</td>
<td>5.48c</td>
<td>4.36b</td>
</tr>
</tbody>
</table>

- numbers characterising the properties of peat from the one location, followed by the same letter, do not differ significantly (P>0.05).

Statistical analysis of all the data confirmed that the raise of temperature stimulated considerably MTA. The most evident increase was observed when the temperature rose from 10 to 20°C; under these conditions MTA was 2.8 times more efficient (Fig. 1C). Statistical analysis of all collected data demonstrated that MTA was significantly modified by substrate availability. Increase of CH₄ concentration from 1 to 5% (v/v) induced the increase of MTA from 20 to almost 60 mg CH₄ kg D.W.⁻¹ d⁻¹ (Fig. 1D).

DNA was isolated from the samples characterised by the highest MTA. In general, methane oxidation was the most efficient in the samples incubated at 20°C and in CH₄ concentration of 5% (v/v). The most active layers were: 60-80 cm in site L, 0-20 cm in T, and 40-60 cm in H.
Identification of genes specific for methanotrophs

In the case of pmoA gene positive results for used pairs of primers A189f/mb661r in material from L were obtained (500 bp), while in the case of 16S rRNA gene and primers Type IIf/Type IIr we got a PCR product (525 bp) in the material from H. No results were found after using primers Type IIf/Type IIr. In samples from transition moor peat (T) none of the primers used gave products.

DISCUSSION

Methanotrophic activity

The MTA of investigated peatlands from Polesie Lubelskie Region was in a wide range from 1.38 to 191.3 mg CH$_4$ kg D.W.$^{-1}$ d$^{-1}$. According to literature data, higher values of MTA were determined in peat bogs (59.2-980.8 mg CH$_4$ kg D.W.$^{-1}$ d$^{-1}$) of north-east England (Chen et al. 2008) and in soil from trans-arctic climate (17.3-311 mg CH$_4$ kg D.W.$^{-1}$ d$^{-1}$) of Siberia (Knoblauch et al. 2008).
Comparable MTA data to those obtained in our study were found in acidic forest soils (38.4-169 mg CH₄ kg D.W.⁻¹ d⁻¹) from the area of England (Radajewski et al. 2002) and the state of Iowa (111.4 mg CH₄ kg D.W.⁻¹ d⁻¹) in the U.S. (Chan and Parkin 2001). Most of the wetlands described in literature were characterised by lower values, as in boreal peatlands of Finland (Kettunen et al. 1999), Canada (Moore and Dalva 1997), North Carolina (Meganigal and Schlesinger 2002) and some bog and forest soils of Alaska (Whalen and Reeburgh 1996) where maximal MTA level did not exceed 13 mg CH₄ kg D.W.⁻¹ d⁻¹.

Obtained sequences of pmoA gene fragment demonstrated the highest similarity (99%) to uncultured representatives of the family Methylococcaceae which was found in freshwater sediment in a region of Russia and in the range of this family especially to representatives of the genus Methylobacter sp. (98%) which was noted in the sediments of lake Washington (USA) (Auman et al. 2000). 99-95% similarity was also reported to an uncultured methanotrophic bacterium found in roots of rice, or soils in which rice is practiced (Lüke et al. 2010, Qiu et al. 2008, Shrestha et al. 2008, Qiu et al. BLASTN nucleotide databases) and methanotrophic microorganisms (98%) found in methane seep along the North American continental margin (Tavormina et al. 2008).

In the case of obtained fragments of 16S rRNA gene from microorganisms inhabiting high moor peatland, 98% of similarity to two species of Methylosinus and Methylocystis was estimated. Similar Methylosinus sp. were revealed in such environments as a dystrophic peatland lake (Neuglobsow, Germany), oxic surface sediment of lake Kinneret (Israel) or soils in Germany (Heyer et al. 2002, Knief and Dunfield 2005). Likewise, Methylocystis sp. species in lake sediments, Syracuse (USA), wetland soils (Norway, Russia, Germany) were detected (Wartiainen et al. 2006, Heyer et al. 2002, Costello and Lidstrom 1999, Galchenko et al. 1980). Particular data are presented in Table 4.

Sequencing analysis indicated that in low moor peatland most common are representatives of γ-Proteobacteria, exactly Type I, while in high moor peatland α-Proteobacteria, that is Type II. This might be caused by pH regime. Earlier works indicated that acidic peat is colonised mainly by methanotrophic representatives of the α-Proteobacteria: Methylocystis spp., Methylocella spp. and Methylocapsas pp. (Dunfield et al., 2003, Berestovskaya et al. 2005, Dedysz 2009). However, species from Methylobacter genera are also found in peatlands. It was found that Type I of methanotrophs were more abundant than Type II and members of the genus Methylobacter were predominant (Vecherskaya et al. 1993, Trotsenko and Khmenina 2005).

The results should be considered as preliminary and further identification experiments of peatland methanotrophs from Polesie Lubelskie Region are needed.
Table 4. Similarity of received sequences to other methanotrophic bacteria

<table>
<thead>
<tr>
<th>Fragments of compared sequences</th>
<th>Similar microorganisms</th>
<th>Similarity (%)</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>pmoA gene from low moor peat (A189f/emb661r)</td>
<td>Uncultured <em>Methylococcaceae</em> EU722420.1</td>
<td>99</td>
<td>BLASTN nucleotide databases</td>
</tr>
<tr>
<td></td>
<td>Uncultured methanotrophic bacterium FN600075</td>
<td>99</td>
<td>Lüke <em>et al.</em> 2010</td>
</tr>
<tr>
<td></td>
<td><em>Methylobacter</em> sp. AY007286.1</td>
<td>98</td>
<td>Auman <em>et al.</em> 2000</td>
</tr>
<tr>
<td>16S rRNA from high moor peat (Type II f/ Type IIr)</td>
<td>Uncultured methanotrophic bacterium EU444867</td>
<td>98</td>
<td>Tavormina <em>et al.</em> 2008</td>
</tr>
<tr>
<td></td>
<td>Uncultured methanotrophic bacterium FM986052.1</td>
<td>96</td>
<td>Qiu <em>et al.</em> 2008</td>
</tr>
<tr>
<td></td>
<td>Uncultured methanotrophic bacterium AM849714.1</td>
<td>96</td>
<td>Shrestha <em>et al.</em> 2008</td>
</tr>
<tr>
<td></td>
<td>Uncultured methanotrophic bacterium FM986052</td>
<td>95</td>
<td>Qiu <em>et al.</em>, BLASTN nucleotide databases</td>
</tr>
<tr>
<td></td>
<td><em>Methylosinus</em> sp. strain L6 AJ868422.1</td>
<td>98</td>
<td>Knief and Dunfield 2005</td>
</tr>
<tr>
<td></td>
<td><em>Methylosinus</em> sp. strain F10V2a AJ458504.1</td>
<td>98</td>
<td>Heyer <em>et al.</em> 2002</td>
</tr>
<tr>
<td></td>
<td><em>Methylosinus</em> sp. strain KS8a AJ458493.1</td>
<td>98</td>
<td>Heyer <em>et al.</em> 2002</td>
</tr>
<tr>
<td></td>
<td><em>Methylocystis parvus</em> strain 57, AJ458508.1</td>
<td>98</td>
<td>Costello and Lidstrom 1999</td>
</tr>
</tbody>
</table>
CONCLUSIONS

1. In all of the tested peat samples methanotrophic activity (MTA) was found and it ranged from 1.38 to 191.26 mg CH₄ kg D.W.⁻¹ d⁻¹.

2. Methanotrophic activity, among tested temperatures, was the highest at 20°C and the maximum values reached were 46-86 (1% CH₄ v/v) and 145-191 (5% CH₄ v/v) mg CH₄ kg D.W.⁻¹ d⁻¹. The most active was the peat from low moor peatland.

3. MTA of peat was 3-times higher when the initial concentration of methane was 5% in comparison to the combination with 1% CH₄ (v/v).

4. Fragments of 16S rRNA gene from microorganisms inhabiting site H are similar to methanotrophs from the genera Methylocystis and Methylosinus (Type II of methanotrophs).

5. Fragments of pmoA gene from microorganisms inhabiting peat from L location showed similarity to the family Methylococcaceae and the genus Methylobacter (Type I of methanotrophs).

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METANOTROFY ODPOWIEDZIALNE ZA UTLENIANIE METANU
W NATURALNYCH TORFOWISKACH POLESA LUBELSKIEGO

Anna Szafranek-Nakonieczna, Zofia Stępniewska, Agnieszka Wołoszyn,
Jakub Ciepielski

Katedra Biochemii i Chemii Środowiska, Katolicki Uniwersytet Lubelski Jana Pawła II
Al. Kraśnicka 102, 20-718 Lublin
e-mail: anna.szafranek@kul.lublin.pl

Streszczenie. Potencjalna aktywność metanotroficzna (MTA) została wyznaczona w warunkach
laboratoryjnych, w torfach pochodzących z torfowisk reprezentujących typy: wysokie (H), prze-
jściowe (T) i niskie (L), zlokalizowanych na obszarze Polesia Lubelskiego. Badane torfowiska różniły się między sobą
pod względem: pH, zawartości TOC, indeksem von Posta oraz wilgotności. Inkubacje przeprowadzono
w następujących warunkach: wilgotność w stanie naturalnym (198-719 %w/w), temperatury: 5, 10 i 20°C,
oraz atmosfera wzbogacona w 1 oraz 5% CH4(v/v). Niezależnie od temperatury, niższe wartości MTA
(1,38-51,16 mg CH4 kg D.W.-1 d-1) wyznaczono dla torfu inkubowanego w atmosferze wzbogaconej o 1%
CH4(v/v). Na MTA istotny wpływ wykazywała również temperatura oraz lokalizacja punktu poboru prób.
Z najaktywniejszych metanotroficznie torfów (20°C, 5% CH4 v/v) izolowano całkowite DNA , na którym
przeprowadzono reakcję PCR powielające fragment genu pmoA (startery A189f/mb661r) oraz sekwencję
16S rRNA (startery Typ If /Typ Ir oraz Typ IIIf /Typ IIr), specyficzne dla bakterii metanotroficznych.
Pozytywny wynik reakcji PCR ze starterami genu pmoA otrzymano dla materiału pochodzącego ze
stanowiska L, sekwenkowanie wskazało na obecność w tym materiale metanotrofów w największym
stopniu podobnych do gatunków z rodziny Methylococcaceae, natomiast na podstawie sekwencji genu 16S
rRNA pochodzącego z mikroorganizmów zasiadających stanowisko H stwierdzono ich duże podobieństwo
do przedstawicieli rodzaju Methylocystis i Methylosinus.

Słowa kluczowe: aktywność metanotroficzna, metanotrofy, torf