INTRODUCTION

Higher global temperatures are capable of causing massive releases of methane gas ($\text{CH}_4$) to the atmosphere, from both marine and freshwater ecosystems, with the potential effect of being a further aggravation and progression of the climate change. One of the key processes regulating the flows of $\text{CH}_4$ into the atmosphere is the anaerobic oxidation of methane (abbreviated to AOM) – which is considered responsible for a reduction of 7–25% of the global $\text{CH}_4$ production per year [Knittel and Boetius 2009; Hou et al. 2017]. AOM proceeds in line with the reduction of various electron acceptors, including sulphates, nitrites, nitrates and metals. AOM, coupled with the reduction of $\text{NO}_3^-$/$\text{NO}_2^-$ ions (known as anaerobic oxidation of methane combined with denitrification, or DAMO), exerts a simultaneous impact on the key global cycles involving carbon and nitrogen, while reducing the emissions of $\text{CH}_4$ and $\text{N}_2\text{O}$ in line with the reaction [Hou et al. 2017; Weiwei et al. 2018]:

$$5\text{CH}_4 + 8\text{NO}_3^- + 8\text{H}^+ \rightarrow 5\text{CO}_2 + 4\text{N}_2 + 14\text{H}_2\text{O}$$

$$3\text{CH}_4 + 8\text{NO}_2^- + 8\text{H}^+ \rightarrow 3\text{CO}_2 + 4\text{N}_2 + 10\text{H}_2\text{O}$$

The evidence for the presence of DAMO in freshwater ecosystems first appeared in 2004 [Islas-Lima et al. 2004], and first laboratory-scale research was carried out in 2006 [Raghoebarsing et al. 2006]. Until recently, the DAMO process in freshwater sediments was not given much consideration due to the limited availability of $\text{NO}_3^-$ (a consequence of either excessively slow nitrification or excessively rapid denitrification).
However, the research carried out worldwide has demonstrated the process occurring in such freshwater ecosystems as boreal lakes [Norði and Thamdrup 2014], Lake Constance [Deutzmann et al. 2014] or wetlands [Hu et al. 2014]. Hu et al. [2014] estimated that about 2–6% of current global CH₄ production is the result of the oxidation reactions taking place in wetlands [Bhattacharjee et al. 2016, Chen et al. 2014, Hu et al. 2014, Liping et al. 2018, Ru et al. 2017].

Since the available literature has very little data on the above-mentioned process as it occurs in the sediments of dam reservoirs, this paper presents the research on DAMO in the sediments of Rzeszów, Maziarnia and Nielisz Reservoirs – all located in SE Poland. The said research saw the sediments incubated at the authentic in-situ temperature, along with a 13CCH₄ isotope marker and NO₃⁻ as an electron acceptor, with a view to determining the rate of any DAMO reaction taking place.

MATERIALS AND METHODS

Study area

The researched Rzeszów and Maziarnia Reservoirs are located in SE Poland’s Podkarpackie Voivodship (Province-Region), while Nielisz Reservoir is in Lubelskie Voivodship. Completed in 1974, Rzeszów Reservoir involved the damming of the Wisłok some 64 km along its course. The other tributary feeding this body of water is Strug. While the main purpose of the reservoir construction was the proper operation of a water intake for the city of Rzeszów, the body of water in question also has an important sporting and recreational role to play. A proportionally large reduction in the overall volume (of 0.7M m³) has affected the Reservoir over the last 40 years, with many areas (in the upper part especially) silted up and gradually transforming into land. The catchment area in this case is of 2050 km², while the Reservoir’s immediate surroundings include wasteland, arable fields and a few scattered buildings. Away from several industrial centers, there is a predominantly agricultural area [Koszelnik and Tomaszek 2002; Gruca-Rokosz 2015].

Maziarnia Reservoir is also a dam reservoir, in this case along the River Łęg, the catchment of which in its upper part features podsolic soils, while the valley bottom is covered by sandy muds and formations reflecting the river accumulation. Another part of the catchment area is covered with soils arising out of loose sands, or else lightly loamy or loamy sands. Given the lack of industrial centers in the vicinity, the catchment of the Łęg is only polluted by domestic sewage and the surface runoff from agricultural areas. While this Reservoir was originally to meet the needs of local waterworks, irrigation and farms with ponds, it today has a retention function, above all [Gruca-Rokosz 2015].

The Nielisz dam reservoir is situated along the middle course of the River Wieprz. Surrounded by a strip of coniferous forest, it consists of a main reservoir and a primary reservoir. There is a 362 kW hydropower plant on the dam. The main purposes of this body of water are thus protection against flooding, as well as energy production, R&R and amateur angling.

The reservoir sediments were sampled at a station located near the dam (of Rzeszów Reservoir) or else near the tributary (Maziarnia and Nielisz Reservoirs), in the light of recent studies indicating that the CH₄ emissions there might be considerable, at 29.13±67.87 mmol·m⁻²·d⁻¹ (Rzeszów), 372.43±328.75 mmol·m⁻²·d⁻¹ (Maziarnia) and 209.55±155.13 mmol·m⁻²·d⁻¹ (Nielisz), on average [Gruca-Rokosz 2015].

The locations of the Rzeszów, Maziarnia and Nielisz Reservoirs and the sampling stations are shown in Figure 1.

Sediment sampling and preparation

The reservoir sediments collected in autumn (September) 2018 were found to be at a temperature of 10°C. Five cores were in fact taken, using a specially-designed sediment corer (No. P.428513), and immediately transported to the laboratory, along with the overlying water. The sediment cores were then pushed progressively from the bottom of Plexiglas tubes by a piston, in such a way that it was possible to study separately the layers present at depths of 0–5, 5–10 and 10–15 cm.

The pore-water was centrifuged from three layers of one core in a laboratory centrifuge (MPW-352), so that determinations could be made for organic carbon (TOC), as well as nitrite, nitrate and sulphate ions (NO₃⁻; NO₂⁻ and SO₄²⁻). The pore-water from a second core was squeezed directly into a gastight vial using a modified pore-water squeezer (Reeburgh 1967), in order for
contact with the atmosphere to be avoided, so that a determination for trivalent iron (Fe$^{3+}$) could be carried out.

The three sediment layers from the third core were dried to constant weight at room temperature, and subsequently at 60 °C, before the sediments were milled prior to determining parameters like pH, organic matter (OM), total organic carbon (TOC), total nitrogen (TN) and humic substances (HS).

The sediments from the fourth and fifth cores were used to research the process of the anaerobic oxidation of methane (AOM).

**Incubation experiment**

In the experiment, 7 g of wet sediment from each layer was placed into a gastight incubation vessel with 14 mL of distilled water deoxygenated by nitrogen purging (99.999% Air Products) using a laboratory scrubber [Gupta 2011]. All operations were carried out in a glove box (830-ABB Compact Glove Box Plas-Labs, Inc. USA). Prior to the laboratory work, the air in the glove box was replaced with helium (99.9999% Air Products), 7–8 times. The prepared samples were then subjected to pre-incubation by placing them in the dark in a thermostat for 10 days at in-situ temperature (10°C – autumn), to allow for the removal of residual oxygen by microorganisms, via biochemical processes. It was only after this that the $^{13}$CH$_4$ isotope marker (99%, Sigma-Aldrich 490229–1L-EU) – and in the incubations with the addition of electron acceptor NO$_3^-$ – 1 ml of Ca(NO$_3$)$_2$·H$_2$O at concentration of NO$_3^-$ $2.1$ mmol·l$^{-1}$ – were added to appropriate samples, so that the final concentration in the samples was $100$ µmol·l$^{-1}$. The incubations were carried out at a temperature corresponding to the in-situ sediment temperature in autumn (10 °C). After a certain time (incubation times of 0, 3, 10, 20, 30, 40 and 50 days), 1 cm$^3$ of the headspace was withdrawn using a gastight syringe (1001SL Hamilton) and analyzed for the CO$_2$ concentrations (ppm) using a Shimadzu GC-2010 Plus gas chromatograph equipped with a Barrier Discharge Ionization Detector (BID) and a Shin Carbon ST column (2 m, 1.00 mm ID, Mesh 100/120). The samples were injected manually into a GC apparatus with the detector working at 250°C. The temperatures were 150°C for the injection port and 60°C for the column (the latter being maintained for 13 minutes). The carrier gas was helium (99,99999% Air Products) at a 50 ml·min$^{-1}$ flow rate. The carbon isotopic composition of CO$_2$ was determined using gas chromatograph combustion isotope mass spectrometry (GC-CIII-IRMS Deltaplus Finnigan), by withdrawing 0.1 cm$^3$ of the headspace using a syringe (1700, Hamilton). The results of stable isotope analyses are presented in the form of $\delta^{13}$C expressed in [%] relative to the PDB (Pee Dee Belemnit) standard (Eq. 1):

$$\delta^{13}C = \left( \frac{^{13}C/^{12}C_{(\text{sample})}}{^{13}C/^{12}C_{(\text{std})}} - 1 \right) \times 1000.$$

(1)

The results obtained for the CO$_2$ concentrations (ppm) and $\delta^{13}$C-CO$_2$ (%) were used to calculate the rate at which the anaerobic oxidation of methane had taken place. The CO$_2$ concentration at time “zero” was subtracted from each CO$_2$ concentration measured after 3, 10, 20, 30, 40 and 50 days of incubation.

Fig. 1. Location of Rzeszów, Maziarnia and Nielisz Reservoirs, and the sampling stations
Sediment and pore-water analysis

The reaction was determined potentiometrically (MultiLine P4, WTW, Germany) in slurry with 1N KCl [Ostrowska et al. 1991]. The OM content was determined as the loss after drying of sediments within 4 h at 550°C [Holmer et al. 2005]. The contents of TOC and TN were determined at 1020°C using a CN elemental analyser (Flash EA 1112, ThermoQuest). The quality analysis made use of the standard samples of known amounts of carbon and nitrogen (acetanilide). Prior to the determination of the TOC content, the dried and ground sediment samples were placed in a desiccator with concentrated HCl vapor [Zimmermann et al. 1997] for 72 h, to ensure the removal of carbonates. Prior to the analysis, the sediment sample was again dried to constant weight at 60°C. The modified method of Griffith and Schnitzer (1975) was used in determining the content of humic substances (HS). The procedure for the determination of the HS is described in detail in Bartoszek [2019].

Ions – i.e. nitrite (NO$_{2}^{-}$), nitrate (NO$_{3}^{-}$) and sulphate (SO$_{4}^{2-}$) – were analysed in pore-water after first being centrifuged and filtered through 0.22 μm syringe filters using DIONEX ICS-5000 ion chromatography. Total organic carbon was determined using a Shimadzu TOC-V CPN Total Organic Carbon Analyzer, while the concentration of Fe$^{2+}$ was determined spectrophotometrically using ferrozine, after Viollier et al. [2000]. Absorbance was measured using a Hach UV-VIS spectrometer at 562 nm.

Calculations

The AOM and DAMO rates (nmol·g$^{-1}$·h$^{-1}$) and the amount of CH$_4$ oxidised (nmol·g$^{-2}$) were calculated using a formula developed by reference to the available literature [Gupta 2011; Shi et al. 2017] (2) and (3). δ$^{13}$C-CO$_2$, was converted to $^{13}$C atom percent ($^{13}$C AT%) in line with Eq. (4):

\[
AOM / DAMO (\text{nmol} \cdot g^{-1} \cdot h^{-1}) = \frac{\delta^{13}CAT\%}{100} \cdot \left[\frac{[CO_2]_{ppm}}{1000} \right] \cdot \left(\frac{m(g) \cdot t(h)}{V_g (dm^3)}\right)
\]

(2)

\[
CH_4 \text{oxidised (nmol} \cdot g^{-1}) \equiv AOM \cdot t(h)
\]

(3)

\[
\delta^{13}CAT = \left(\delta^{13}CPDB + 1000 \delta^{13}CPDB + 1000 \cdot \frac{1000}{R_{so}} \right) \cdot 100 = \left(\delta^{13}c_{CO} - \delta^{13}c_{CO} \right) \cdot 100 = \left(\delta^{13}c_{CO} \right) \cdot 100
\]

(4)

where $^{13}$CAT - $^{13}$C atom percent (%);

$CO_2$ – carbon dioxide concentration at time $t$ (ppm);

$V_g$ – volume of headspace in vial (0.096156 dm$^3$);

$V_m$ – molar volume of ideal gas at 25°C (24.45 dm$^3$·mol$^{-1}$);

$m$ – dry sediment weight (g);

$t$ – incubation time (h);

$R_{so}$ – absolute ratio constants in the standard (0.0111803).

RESULTS

Characteristics of sediments and pore water

Sediment characteristics

The results for selected chemical parameters characterising the sediments of Rzeszów; Mazimiarna and Nielisz Reservoirs are summarised in Table 1.

The reactions of the Rzeszów and Nielisz Reservoir sediments were slightly alkaline, with

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Unit</th>
<th>Rzeszów</th>
<th>Mazimiarna</th>
<th>Nielisz</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH</td>
<td>–</td>
<td>7.20</td>
<td>7.20</td>
<td>6.55</td>
</tr>
<tr>
<td>OM</td>
<td>%</td>
<td>10.84</td>
<td>9.09</td>
<td>6.46</td>
</tr>
<tr>
<td>TOC</td>
<td>%</td>
<td>3.62</td>
<td>2.89</td>
<td>1.82</td>
</tr>
<tr>
<td>TN</td>
<td></td>
<td>0.37</td>
<td>0.27</td>
<td>0.17</td>
</tr>
<tr>
<td>C:N</td>
<td>-</td>
<td>9.86</td>
<td>10.69</td>
<td>10.81</td>
</tr>
<tr>
<td>HS</td>
<td>mg·g$^{-1}$·d.w.</td>
<td>31.7</td>
<td>27.5</td>
<td>17.6</td>
</tr>
</tbody>
</table>

Table 1. Selected chemical parameters characterising the reservoir sediments
the pH values varying across the narrow ranges 7.20–7.39 (Rzeszów) and 7.21–7.26 (Nielisz). The sediments of Maziarnia Reservoir had much lower pH values, in the 5.57–6.55 range.

The contents of organic matter (OM) in the reservoir sediments ranged from 3.27 to 13.42%. The highest values were noted for the Nielisz sediments, as followed by those from the Rzeszów site. The average content of organic matter was greater than 8%, but the highest value did not exceed 11%. It was typical to note lower OM contents at greater depths in sediments, and in Rzeszów and Maziarnia Reservoirs, this depletion assumed certain significance, with the 10–15 cm layer depleted by about 5% in comparison with the surface layer. In contrast, the sediment profile at Nielisz Reservoir was characterised by higher OM contents in deeper layers.

The content of organic carbon (TOC) in the uppermost sediment layer varied across a 0.89–5.51% range. The tendencies for TOC analogous to those for OM were observed in all cases. The highest TOC content was obtained in the sediments sampled from Nielisz Reservoir. At all sites, organic carbon accounted for about 30% of values for OM.

As with OM and TOC, it was sediments from Nielisz Reservoir that had highest values for total nitrogen (TN) content. The lowest values, in turn, characterised the sediments sampled from Maziarnia Reservoir. All observed trends for the changes in TN were analogous to those noted for OM and TOC.

In Rzeszów Reservoir, the C:N ratios down the sediment profile were in the 9.86–15.26 range, with slight depth-related change. Similar values were obtained for the sediments from the other reservoirs, with the C:N ratios of 10.92–11.07 (Maziarnia) and 9.83–10.40 (Nielisz). Lower C:N ratios recorded for the 0–5 cm layer point to a greater share of autochthonous matter. In turn, organic matter in the 10–15 cm sediment layer is mainly of allochthonous origin, with sources taken to include macrophytes and plants rich in cellulose [Gruca-Rokosz 2015].

The content of humic substances in the sediments ranged from 9.0 to 47.8 mg g\(^{-1}\) d.w. In all cases, the tendencies were analogous to those observed for the parameters referred to above (OM, TOC and TN), with the highest values again noted for the sediments from Nielisz Reservoir.

Table 2.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Unit</th>
<th>Rzeszów</th>
<th>Maziarnia</th>
<th>Nielisz</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Sediment layer [cm]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0–5</td>
<td>5–10</td>
<td>10–15</td>
<td>0–5</td>
</tr>
<tr>
<td>NO(_2^–)</td>
<td>mg dm(^{-3})</td>
<td>0.02</td>
<td>0.01</td>
<td>0.02</td>
</tr>
<tr>
<td>NO(_3^–)</td>
<td>mg dm(^{-3})</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
</tr>
<tr>
<td>SO(_4^{2–})</td>
<td>mg dm(^{-3})</td>
<td>0.23</td>
<td>0.04</td>
<td>n.d.</td>
</tr>
<tr>
<td>Fe(^{3+})</td>
<td>µmol dm(^{-3})</td>
<td>380.57</td>
<td>162.60</td>
<td>95.40</td>
</tr>
</tbody>
</table>

Pore-water characteristics

The values for pore-water concentrations of the NO\(_2^–\), NO\(_3^–\), SO\(_4^{2–}\) and Fe\(^{3+}\) ions are as shown in Table 2.

Comparable NO\(_2^–\) and NO\(_3^–\) concentrations were obtained for all dam reservoirs, with values in the ranges 0.01–0.02 and 0.02–0.03 mg dm\(^{-3}\), respectively. In the case of sulphate ions, the ranges differed from one reservoir to another. The lowest concentrations characterised Rzeszów Reservoir (0.04–0.23 mg dm\(^{-3}\)), while the values on average three or more than ten times as high noted for Nielisz and Maziarnia Reservoirs, respectively. The Fe\(^{3+}\) concentrations varied from 95.40 µmol dm\(^{-3}\) (Rzeszów) to 580.66 µmol dm\(^{-3}\) (Nielisz).

Rates of anaerobic oxidation of methane

All sediments sampled for the purposes of the research proved capable of oxidising methane under anaerobic conditions. The CO\(_2\) enrichment with the \(^{13}\)C isotope was noted in the headspace, indicating oxidation of the \(^{13}\)CH\(_4\) isotope marker added. A value for the \(^{13}\)C atom percent (\(^{13}\)C AT\%\) was arrived at by calculation, using \(\delta^{13}\)C-CO\(_2\) (‰) readings obtained for the headspace in the course of incubation, and the results are shown graphically in Fig. 2.

During the experiment, the \(^{13}\)C AP values ranged from 1.31 to 37.22%, and from 1.38 to 46.68%, in incubations carried out in the presence of \(^{13}\)CH\(_4\) or \(^{13}\)CH\(_4\)+NO\(_3^–\), respectively. A steady increase in the concentrations of CO\(_2\) in...
Fig. 2. Change of $^{13}$C AP and CO$_2$ over time for incubations conducted with the addition of $^{13}$CH$_4$ (panel a, c, e) or $^{13}$CH$_4$+NO$_3^-$ (panel b, d, f)
the headspace was noted up to day 40 (with $^{13}$CH$_4$ and $^{13}$CH$_4$+NO$_3^-$). However, incubation with added NO$_3^-$ gave rise to maximum CO$_2$ concentrations of 7300 ppm, as opposed to 6500 ppm where incubation was with the $^{13}$CH$_4$ marker alone.

The rates of anaerobic oxidation of methane with or without added NO$_3^-$ are shown graphically in Figure 3, along with the amounts of CH$_4$ oxidised. The anaerobic oxidation of methane proved most effective in the sediments (of all layers) sampled from Nielisz Reservoir, in both incubations with $^{13}$CH$_4$ and $^{13}$CH$_4$+NO$_3^-$. The rates were lower in the sediments from Rzeszów ($^{13}$CH$_4$) and Maziarnia ($^{13}$CH$_4$+NO$_3^-$). The rates for the process taking place under both types of incubation and with sediments from all reservoirs usually peaked on day 20. However, at that time (with $^{13}$CH$_4$+NO$_3^-$), the figure obtained for Nielisz Reservoir was three times as high on average as the ones noted for Rzeszów and Maziarnia.

Where the amounts of methane oxidised were concerned, the peak values for sediments from all reservoirs were reached on day 40 of incubation (with either $^{13}$CH$_4$ or $^{13}$CH$_4$+NO$_3^-$). Thus, the amount of CH$_4$ oxidised had actually decreased by the last day of the research (day 50). For the incubation with added $^{13}$CH$_4$+NO$_3^-$, the amounts of CH$_4$ oxidised on day 40 were almost 250 nmol·g$^{-1}$ (Rzeszów), 300 nmol·g$^{-1}$ (Maziarnia) and 600 nmol·g$^{-1}$ (Nielisz). In the case of incubation with the $^{13}$CH$_4$ isotope marker alone, the amounts of CH$_4$ oxidised (after 40 days) did not exceed 90 nmol·g$^{-1}$ in the Rzeszów sediments, while the values for Maziarnia Reservoir remained below 300 nmol·g$^{-1}$, compared with those for Nielisz reaching 400 nmol·g$^{-1}$.

**DISCUSSION**

The research reported here confirms the presence of the DAMO (denitrifying anaerobic methane oxidation) process in the freshwater sediments from certain dam reservoirs in Poland. $^{13}$CO$_2$ was present in the headspace as all sediment samples were incubated, denoting an origin in the oxidation of the $^{13}$CH$_4$ with which the sediments were dosed. The AOM rate in the incubations carried out with the isotope marker alone was in most cases significantly lower than where NO$_3^-$ was also added. Slight departures were observed in just two cases of the 10–15 cm layer of sediment from Rzeszów Reservoir and the 0–5 cm layer from Maziarnia. Such results point to a significant impact of nitrate on the dynamics of methane’s anaerobic oxidation.

Out of the sediments sampled, the most and least rich in organic matter (including TOC and TN) were those from Nielisz and Maziarnia Reservoirs, respectively. The rates of the anaerobic oxidation of methane were highest in Nielisz sediments, and lowest in those from Maziarnia Reservoir. Additionally, as statistically significant correlations between the rate of DAMO and OM, TOC and TN were obtained ($R^2 = 0.65$), it may be the availability of C and N that determine whether or not DAMO takes place in the sediments of dam reservoirs.

The DAMO rates in Maziarnia Reservoir sediments were the subject of an upward trend with layer depth, though the reverse relationship applied to the Rzeszów and Nielisz sites. While anaerobic oxidation may be occurring, CH$_4$ is also being produced, in the amounts that become significant in deeper sediment layers in particular. Methanogenesis entails the decomposition of “old” carbon in deeper layers of sediment, with some of the carbon then available for anaerobic oxidation in the presence of alternative electron acceptors. In addition, the CH$_4$ produced in deeper sediment layers begins to diffuse upwards, to areas near the sediment-water interface where electron acceptors are increasingly available [Smemo and Yavitt 2007; Broman E. 2013].

The reported DAMO rate usually peaked on day 20. The subsequent decrease may reflect the gradual depletion of NO$_3^-$ over time [Fan et al. 2019]. Furthermore, by the 50th day of the research there had actually been a decrease in the amount of methane oxidized compared with the previous situation, denoting the increased relative importance of the methane production process probably occurring simultaneously throughout the trial, but proving less tangible earlier on.

Other researchers [Shi et al. 2017] have likewise determined the amounts of methane oxidised in sediments under anaerobic conditions. On day 3 of incubation, these ranged from 97.08 to 143.69 nmol·g$^{-1}$. Our values closest to this range were obtained in the 0–5 cm layer of sediment from Nielisz Reservoir (on day 3). On average, the values only one-fifth as high were obtained for other layers and other reservoirs analysed.

The literature offers very little information on the DAMO rates in freshwater ecosystems, and no data whatever for dam reservoirs. Most
existing studies relate to wetlands and peat bogs, with incubation times for sediment samples in the range from a mere 36 hours to even several weeks or months. All of our studied reservoirs yielded lower DAMO values than are present in the literature, given that 36-hour incubations of Jinchuan peatland sediments (from Jilin Province, NE China) reported on by Shi et al. [2017] were associated with average DAMO rates of 1.96–2.29 nmol·g$^{-1}$·h$^{-1}$; while Gupta et al. [2013] obtained the values of 5.94 nmol·g$^{-1}$·h$^{-1}$ after 21 days of incubation. Lower values 0.38 nmol·g$^{-1}$·h$^{-1}$ were recorded by Zhu et al. [2012], in relation to a 3-month incubation of sediments from the Brunssummerheide peatland.

The DAMO process has also been examined in the sediments from various sites in China (the Xiazhuhu and Xixi wetlands and a paddyfield). In these cases, rates were in the 0.01–0.23 nmol·g$^{-1}$·h$^{-1}$ range [Hu et al. 2014].

Fig. 3. Changes through time in AOM rate and amount of CH$_4$ oxidised, in the course of incubations of $^{13}$CH$_4$ and $^{13}$CH$_4$+NO$_3^-$ in 0–5, 5–10 and 10–15 cm layers of sediment collected from Rzeszów, Maziarnia and Nielisz Reservoirs.
CONCLUSIONS

• The research provides a direct evidence of the denitrifying anaerobic methane oxidation (DAMO) process taking place in the freshwater sediments of Polish dam reservoirs. On average, irrespective of the sediment being sampled, incubation with added NO₃⁻ was associated with doubled rates of CH₄ oxidation under anaerobic conditions.

• The most effective DAMO process was that characterising Nielisz Reservoir, above all sediments of its (0–5 cm) surface layer. In most cases, the process proved less intense in deeper layers, though no significant differences in the rates of CH₄ oxidation attributable to added NO₃⁻ were observed in either the 10–15 cm layer of sediment from Rzeszów Reservoir, or the 0–5 cm layer from Maziarnia. In most cases, the DAMO rates peaked on day 20 of incubation.

• The availability of C and N may determine whether DAMO does or does not take place in dam-reservoir sediments. The highest contents of these in our experiment characterised the sediments from Nielisz Reservoir, and this could be seen to translate into highest DAMO rates. The converse situation also applied, in that sediments from Maziarnia Reservoir, with their lowest TOC and TN contents, also reported the lowest DAMO rates.

• A comparison of results with those reported for other aquatic ecosystems revealed the DAMO rates (on days 10 or 20) respectively only one-fifth or one-tenth as high in Nielisz Reservoir sediments, and in those from Rzeszów and Maziarnia Reservoirs, as in certain Chinese peatland sediments. However, the rates we noted were noticeably higher than those reported from certain other wetlands in China (about twice as high in the case of Rzeszów and Maziarnia, and almost 3 times as high for Nielisz).

• As DAMO is a relatively complex matter, a better understanding of the process occurring in dam reservoirs will require further research, inter alia to recognise the DAMO-associated microorganisms and to assess the impact of abiotic factors to a greater extent.

Acknowledgements

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