Dissimilatory sulfate reduction and methane production in Gotland Deep sediments (Baltic Sea) during a transition period from oxic to anoxic bottom water (1993–1996)

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ABSTRACT: During a transition period from oxic to anoxic conditions in the bottom water, rates of sulfate reduction and methane production, methane fluxes, as well as concentration profiles of sulfate, sulfide and methane were measured in sediments at a central site of the Gotland Deep (Stn AL 93, 241 m depth), which is regarded as representative for the deepest part of this basin. During this period from 1993 to 1996 oxic conditions in the bottom water prevailed from spring 1994 until summer 1995 with oxygen concentrations decreasing progressively with time. In the sediments methane production occurred primarily in layers below 1 m depth and flux rates of methane to the sediment surface were characterized by a steep concentration gradient from approx. 5 mM at 4 m depth to values close to 30 μM at the surface, determined by diffusion processes and anaerobic oxidation of methane. Both processes were independent of changes at the sediment surface. Differences in the flux rates of methane between the deeper part with a mean value of 259 nmol m⁻² d⁻¹ and the upper layers with a mean of 47.7 nmol m⁻² d⁻¹ indicate that a considerable proportion of the methane is oxidized within the anoxic horizon of the sediment (71 to 86% in the layer from 40 to 70 cm). Low rates of methane production found within the top 20 cm of the sediment during periods of oxic bottom water increased after depletion of oxygen and resulted in a clear maximum of the methane concentration in the top 2 cm. Sulfate concentrations declined exponentially from values of 11.5 mM in June 1994 and 8.5 mM in October 1995 at the sediment surface to values of 2.5 mM at 20 cm depth and of less than 0.5 mM at 50 to 60 cm depth. High sulfate reduction rates (150 to 250 nmol cm⁻³ d⁻¹) in the upper part of the sediment (8 to 13 cm) coincided with maxima of sulfide concentrations. During the time period of this investigation an increase of maximum sulfate concentrations in the sediment from 1 to 10 mM was measured together with decreasing oxygen concentrations in the deep water. At the same time sulfate reduction established a small but distinct maximum at the top layer of the sediment (0 to 2 cm). The relative importance of sulfate reduction and methanogenesis in the carbon budget of the Gotland Deep sediments is calculated on the basis of the actual measurements.

KEY WORDS: Sulfate reduction · Sulfide · Methane production · Methane fluxes · Gotland Deep · Baltic Sea

INTRODUCTION

The Gotland Deep is one of the major deep basins of the Baltic Sea. It is located within the Eastern Gotland Basin between the island of Gotland (Sweden) and Latvia, and has a maximum water depth of 241 m. A permanent thermo-/halocline at 60 to 80 m water depth largely inhibits water exchange between surface water (7 to 8% salinity) and deep water (11 to 13%). Long anoxic stagnation periods of the deep water were only interrupted by irregular, lateral intrusions of water masses from the North Sea with higher salinity and moderate oxygen content. The investigation period of this study (1993 to 1996) was preceded by an unusually long anoxic period from 1979 to 1993 and a massive
salt water intrusion into the Baltic Sea starting in January 1993. During this anoxic time period stable gradients of sulfide and methane from the sediment to the chemocline at 120 to 130 m water depth were established in the water column. Maximum concentrations near the sediment surface were more than 150 μM sulfide and 800 nM methane (Brettar & Rheinheimer 1991, Schmaljohann unpubl. results).

In the natural environment methanogenesis and sulfate reduction are competing processes in the anaerobic decomposition of organic matter. Competition is governed by the redox potential, the availability of electron acceptors and the affinity for common substrates such as acetate and hydrogen. Though methanogenesis is a dominant process in fresh water sediments, in the marine environment methanogenic bacteria are quite effectively outcompeted by sulfate reducers which are nearly unlimited by the availability of their common electron acceptor and have a greater affinity to hydrogen and acetate (Ward & Winfrey 1985).

There is considerable information on the relevance of sulfate reduction and methanogenesis in marine and freshwater sediments (see reviews by Ward & Winfrey 1985, Skyring 1987, Widdel 1988). Comparatively less is known about brackish water habitats, which exhibit intermediate salinities and sulfate concentrations compared to the 2 former types of ecosystems. Investigations have been largely restricted to shallow coastal areas and inlets (Iversen & Blackburn 1981, Whiticar 1982, Heyer et al. 1990, Jørgensen et al. 1990, Schmaljohann 1996). First studies concerning rates of methanogenesis and sulfate reduction in sediments of deep brackish water bodies in the Baltic Sea were published by Lein et al. (1981) and Lein (1983).

This first salt water intrusion after 14 yr of permanent anoxic conditions in the bottom water of the Gotland Basin provided the opportunity to investigate the influence of oxic water on anaerobic processes (i.e. sulfide and methane production) in a generally anoxic sediment and the re-establishment of anoxic, sulfidic conditions. We quantified these processes in a sediment of the Gotland Deep, calculated their proportions in the anaerobic decomposition of organic matter, and determined changes on the sediment biogeochemistry during the transition period from oxic to anoxic conditions in the bottom water.

**MATERIALS AND METHODS**

**Sampling**. Samples were taken and measurements done during 7 expeditions to the Gotland Deep with the RV 'Alkor' (April 1993, June and August 1994, May and October 1995, May and October 1996) at Stn AL 93 (57°18.51' N, 20°06.88' E) located in the deepest part of the Gotland Deep (241 m water depth). Sediment cores of up to 1 m length were obtained using a small gravity corer (Rumohr-Lot) fitted with liners (7.4 cm inner diameter). Two undisturbed cores were used for electrode measurements and for concentration and rate measurements, respectively, in order to obtain sufficient sediment for the measurements and to minimize disturbance due to subsampling. A third core was taken to determine grain size, water content and organic matter. Data on concentrations and rates during June 1994, May 1995, October 1995 and October 1996 were determined in identical cores.

**Chemical analysis**. Immediately after sampling pH and redox potential were measured simultaneously in the sediment cores using Pt electrodes with a small diameter (Mettler-Toledo, Steinbach) and a pH/mV meter (WTW, Weilheim). Feeding of electrodes into the sediment was performed at steps of 0.5 or 1.0 cm. Readings of redox and pH values were constant after 3 and 1 min, respectively.

Grain size distribution was obtained by dry sieving using a standard series of sieves (2000, 1000, 500, 250, 125, 63 μm). Water content was measured by weight loss after drying at 75°C for 24 h (Holme & McIntyre 1984) and organic matter was determined by weight loss of the dried sample after 6 h in a 550°C furnace (loss on ignition).

Pore water from sediments was recovered by centrifuging sediment slices for 10 min at 3500 x g and 4°C. In order to minimize loss of hydrogen sulfide the sediment was transferred rapidly to centrifuge tubes which were completely filled and sealed by gas-tight viton stoppers.

For determination of hydrogen sulfide an aliquot of pore water was fixed in 5% zinc acetate (w/v) immediately after centrifugation and determined according to Cline (1969).

Sulfate concentrations were measured using single column ion chromatography (modified after Gjerde et al. 1980) under isocratic conditions (eluent: 3 mM phthalic acid, pH 4.9; flow rate: 2 cm³ min⁻¹, HPLC pumping system 420, Kontron, Milano). Anions were separated at 40°C on a silica-based column (AnionS, 250 × 4.6 mm, Wescan, Deerfield). Pore water samples had to be diluted 1:20 in suprapure water (Milli-Q-UF-water system, Millipore) before injection (injection volume: 100 μl) to clearly resolve peaks of chloride and sulfate. Conductivity was used for detection and integration of the chromatograms was performed by PC-integration software (Kontron, Milano). Calibration curves showed linear correlation between r² = 0.96 and r² = 0.98.

Methane concentrations in sediment samples were measured using a headspace technique salting out of the gas. Subcores (5 cm³) of each depth were taken with cut-off syringes. The syringe contents were injec-
was used to estimate the rate of anaerobic methane oxidation and the percentage of the upward diffusing methane consumed by this process. Flux rates were calculated from the vertical profile of pore water methane concentrations using Fick's first law:

\[ J_D = -\phi D_{sed} \frac{dC}{dz} \]

where \( J_D \) is the diffusive flux (\( \mu \text{mol CH}_4 \text{ m}^{-2} \text{ d}^{-1} \)), \( \phi \) is the porosity of the sediment, \( D_{sed} \) is the sediment diffusion coefficient, calculated from the molar diffusion coefficient (\( D \)) by the empirical equation \( D_{sed} = D \phi^2 \) (Lerman 1979) using the \( 5^\circ C \) value for \( D \) (0.87 \( \times 10^{-5} \text{ cm}^2 \text{ s}^{-1} \); Iversen & Witherspoon 1970), and \( dC/dz \) is the measured methane concentration gradient in the considered sediment layer (\( \mu \text{mol dm}^{-3} \text{ cm}^{-1} \)).

Sulfate reduction rates (SSR) were calculated using the theoretical model of Berner (1964), which describes the concentration changes of sulfate with time in a given depth below the sediment surface as a function of diffusion processes, sediment accumulation, and DSR. In order to derive the depth-dependent rate of sulfate reduction a diffusion coefficient for sulfate of \( D_S = 0.46 \times 10^{-5} \text{ cm}^2 \text{ s}^{-1} \) (\( 4^\circ C \), \( \phi = 0.90 \); Iversen & Jørgensen 1985), a sedimentation rate in the Gotland Deep of 1 mm yr\(^{-1} \) (Niemestø & Voipio 1974), and the corresponding exponential function of the sulfate gradient in the sediment were used.

**RESULTS**

**General observations**

While oxygen transported into the deep water of the Gotland Basin by the first significant inflow in 1993 was consumed within a rather short time, further intrusions of oxic water in December 1993 and March 1994 resulted in a complete removal of sulfide from the water and a completely oxic water column during 1994 and the first half of 1995. Oxygen concentrations in the deep water reached values as high as 4.3 mg dm\(^{-3} \) (Nehring et al. 1995, K. Gocke, Kiel, pers. comm.). During 1995 oxygen in the deep water was depleted again, and in October 1995 low concentrations of sulfide were present in the water layer up to 2 m above the bottom (Fig. 1). A minor inflow in May 1996 again supplied low amounts of oxygen to the deep water which were consumed rapidly, however, allowing methane and sulfide to penetrate into the water column up to 215 m depth in October 1996.

An optical documentation of the sediment surface of the Gotland Deep by underwater video during the period of oxic bottom water (in June 1994 and May 1995) revealed white or yellowish bacterial mats on its surface. These bacterial mats contained filamentous
sulfur-oxidizing bacteria resembling *Beggiatoa* species. During 1994 and early 1995 numerous individuals of *H. sarsi* (Polychaeta) were also present on the sediment surface. The bacterial mats disappeared and Harmothoe died off when conditions in the bottom water close to the sediment surface became anoxic (October 1995). No macrofauna and no tubes indicating bioturbation were observed within the sediments. The lack of bioturbation resulted in a fine layering of the sediment in the deep part of the Gotland basin over very long time periods, as could be seen in sediment cores down to 4 m depth (data not shown).

**Characterization of the sediment at Stn AL 93**

Sediments consisted of fine silty material without sandy components. The material of the top 15 to 20 cm was of brownish fluffy consistency followed by mushy black fine silt down to 35 cm depth and a dark greyish zone between 35 to 60 cm that merged to a black-ribboned sediment further down. The water content of more than 90% (w/v) in the first 20 cm decreased to 75% from 20 to 30 cm and did not change significantly in deeper layers (Fig. 2). The content of organic matter, determined as loss on ignition, was between 23 and 27% within the first 20 cm at all sampling dates, decreasing to 12 to 14% in deeper layers (Fig. 2).
Redox potential and pH

A relatively constant negative redox potential in the vertical profile was observed in June 1994, with low Eh values of -200 mV at the sediment surface decreasing to -295 mV at 60 cm depth (Fig. 3). During the following cruises the redox potential in the uppermost centimeter increased to 0 and +30 mV in May and October 1995, respectively, before it decreased again to -120 mV in May 1996. At these sampling dates the redox potential decreased to values between -200 and -250 mV with depth. A high oxidation state of the sediment at 10 to 15 cm depth was still recorded in October 1996, when reducing conditions were already re-established at the sediment surface. The vertical profile of the pH showed a minimum in the sediment layer from 6 to 10 cm, a few centimeters above maxima of SRR and sulfide concentrations, and was most pronounced in May 1995 but slowly disappeared later (Fig. 3).

Concentrations of sulfate and sulfide and rates of dissimilatory sulfate reduction

Sulfate concentrations decreased exponentially with sediment depth. The steepest decline was always observed between 10 and 15 cm. At depths of 20 and 60 cm concentrations were quite constant, approximately 2.5 and 0.5 mM respectively, at all sampling dates. During the study period sulfate concentrations in the top 2 cm decreased from 11.5 to 8.5 mM indicating a net sulfate consumption by highly active sulfate reduction in this layer. Apparently, sulfate that had penetrated into the sediment down to 2 to 4 cm during the period of high oxygen concentrations in the bottom water and negligible sulfate reduction in the top sediment layer was consumed much more slowly causing a sulfate maximum at 2 to 4 cm depth in October 1995 and October 1996.

The sulfide concentration always had maxima at 10 to 20 cm (Fig. 4). In October 1995 and October 1996 an additional maximum developed at 0 to 2 cm, which at both sampling dates coincided with lower sulfate concentrations beneath the sediment surface (Fig. 4). The deeper as well as the near-surface sediment horizons with maxima of sulfate concentrations were also characterized by maxima of sulfate reduction (see Fig. 5). A double peak of sulfate concentrations was characteristic for this sediment and was observed repeatedly between 10 and 20 cm (e.g. June 1994 and October 1995). Maxima of sulfide concentrations increased from about 1 mM (June 1994 and May 1995) to approximately 3 mM (October 1995), to more than 7 mM (May 1996, data not shown) and 10 mM (October 1996).

SRR at Stn AL 93 revealed similar distribution patterns at different sampling dates (Fig. 5) with maximal rates of 150 to 250 nmol cm\(^{-3}\) d\(^{-1}\) at a depth of 8 to 13 cm while SRR in adjacent layers did not exceed 50 nmol cm\(^{-3}\) d\(^{-1}\). In May and October 1995 the development of a second maximum between 0 and 2.5 cm depth was observed, although at that time this sediment layer did not show a pronounced negative redox potential. Integrated SRR for the upper 20 cm based on \(^{35}\)S-sulfate measurements were 3.8 (June 1994), 7.6 (May 1995) and 7.2 (October 1995) mmol m\(^{-2}\) d\(^{-1}\), whereas the calculation using Berner's diffusion model (Berner 1964) led to corresponding values of 0.9, 1.8 and 1.7 mmol m\(^{-2}\) d\(^{-1}\) respectively.
Methane production and methane concentrations

Methane concentrations steadily increased from the surface to a depth of at least 80 cm, where values between 0.7 and 1.8 mM were found (Fig. 4). Saturation concentrations were not reached at that depth and there was no indication of in situ formation of gas bubbles. In one instance it was possible to obtain a sediment sample from the end of a 4 m long ‘Kastenlot’ corer, which had a methane content of 5.1 mM. The increase of methane concentrations was nearly linear from the sediment surface to a depth of about 50 cm and became exponential below that depth. The transition between these different curve sections of the methane profile corresponded to sulfate concentrations in the sediment of about 1 mM.

During periods with oxic bottom water (spring 1994 to summer 1995) a continuous gradient with decreasing methane concentrations towards the surface and October 1995 small maxima of methane concentrations were also found at 0 to 2 cm depth (Fig. 6). Integrated methane production rates in the upper 20 cm increased from 3.3 μmol m⁻² d⁻¹ in June 1994 to values of 4.2 μmol m⁻² d⁻¹ in August 1994, 8.6 μmol m⁻² d⁻¹ in May 1995, and 27.9 μmol m⁻² d⁻¹ in May 1996.

DISCUSSION

The sediments of the Central Gotland Basin are a location of intensive anaerobic degradation of organic matter and are subjected to repeated changes from oxic to anoxic conditions at the sediment surface due to irregular inflow events. We were the first to have the opportunity to measure the significant changes that occurred in the sediment during a transition period from oxic to anoxic bottom water, the first such event existed with concentrations as low as 8 to 10 μM in the pore water of the top layer (0 to 2 cm; Fig. 6). During anoxic periods (October 1995 and later), with active methane formation but in the absence of aerobic methane oxidation at the sediment surface, methane concentrations were higher in the 0 to 2 cm layer (30 to 34 μM) than at 2 to 4 cm depth.

The fact that methane concentrations increased without any evidence of saturation to sediment depths of at least 80 to 90 cm is an indication that the main methane production occurred below that depth. As it was not possible with the available sampling gear to obtain sediment samples of depths of more than 1 m, an estimation of the rate of methanogenesis was carried out by calculation of the methane flux rate in layers between 62.5 and 92.5 cm. The methane flux rates varied between 146 and 396 μmol m⁻¹ d⁻¹ with a mean value of 259 μmol m⁻² d⁻¹ compared to 42 to 55 μmol m⁻² d⁻¹ (mean value: 47.7 μmol m⁻² d⁻¹) in the near-surface, linear region of the methane profile (Table 1).

Fig. 5. Depth distribution of sulfate reduction rates (mean of duplicates) at Stn AL 93 in June 1994 (left), May 1995 (middle), and October 1995 (right). The thickness of the bars is adapted to the sampled sediment horizon (space between bars; not determined). Calculated SRR (sulfate reduction rates) using Berner’s diffusion model are depicted as dotted lines.
Table 1. Methane flux rates and calculated anaerobic methane oxidation in the sediment of Stn AL 93 in the Gotland Deep

<table>
<thead>
<tr>
<th>Date</th>
<th>16 June 1994</th>
<th>27 April 1995</th>
<th>5 October 1995</th>
<th>12 October 1996</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sediment horizon (cm)</td>
<td>22.5-1.0</td>
<td>22.5-1.0</td>
<td>22.5-1.0</td>
<td>22.5-1.0</td>
</tr>
<tr>
<td>Porosity (average)</td>
<td>0.973</td>
<td>0.954</td>
<td>0.970</td>
<td>0.983</td>
</tr>
<tr>
<td>Max. methane conc. (µM)</td>
<td>182.0</td>
<td>147.2</td>
<td>197.7</td>
<td>153.4</td>
</tr>
<tr>
<td>Min. methane conc. (µM)</td>
<td>10.0</td>
<td>8.0</td>
<td>34.4</td>
<td>30.0</td>
</tr>
<tr>
<td>Methane flux rate (µmol m⁻² d⁻¹)</td>
<td>55.4</td>
<td>42.3</td>
<td>52.1</td>
<td>41.0</td>
</tr>
<tr>
<td>Sediment horizon (cm)</td>
<td>92.5-62.5</td>
<td>92.5-72.5</td>
<td>62.5-72.5</td>
<td>87.5-62.5</td>
</tr>
<tr>
<td>Porosity (average)</td>
<td>0.899</td>
<td>0.913</td>
<td>0.913</td>
<td>0.920</td>
</tr>
<tr>
<td>Max. methane conc. (µM)</td>
<td>1863</td>
<td>1008</td>
<td>1868</td>
<td>2012</td>
</tr>
<tr>
<td>Min. methane conc. (µM)</td>
<td>634.6</td>
<td>496.3</td>
<td>1175.0</td>
<td>857.4</td>
</tr>
<tr>
<td>Methane flux rate (µmol m⁻² d⁻¹)</td>
<td>223.6</td>
<td>145.8</td>
<td>396.4</td>
<td>270.1</td>
</tr>
<tr>
<td>Anaerobic methane oxidation rate (µmol m⁻² d⁻¹)</td>
<td>168.2</td>
<td>103.5</td>
<td>344.3</td>
<td>229.1</td>
</tr>
<tr>
<td>Percentage of anaerobic methane oxidation (%)</td>
<td>75.2</td>
<td>71.0</td>
<td>86.9</td>
<td>84.8</td>
</tr>
</tbody>
</table>

after more than 14 yr. Changes in the sediments were demonstrated on the basis of (1) elevated sulfide and methane concentrations as indicators of terminal oxidation of organic carbon by DSR and methanogenesis, and (2) direct measurements of SRR and methane production rates. It became obvious that the hydrographic conditions of the bottom water had considerable impact on the biogeochemistry of the surface layers of the investigated sediments.

Stn AL 93 is representative of the central part of the Gotland Deep, where sedimentation rates were reported to be of the order of 1 mm yr⁻¹ (Niemesto & Voipio 1974). Echosounding profiles through the Gotland Deep showed that at Stn AL 93 the thickness of the post-Ancylus deposits was considerably higher than at shallower stations, which indicates a higher sedimentation rate during the past 7000 yr (Schmaljohann et al. 1998). This corresponds to higher amounts of organic matter (loss on ignition), sulfide and methane in the sediment as well as higher SRR and a much more pronounced decrease of sulfate concentrations with sediment depth (data not shown).

During the whole period of these investigations the sediment at Stn AL 93 and in its close vicinity appeared quite homogeneous and gave no indication of small-scale variations in structure and function. The comparison of different cores during each sampling period revealed similar sequence and thickness of the sediment layers and parallel measurements of the redox potential and pH in several cores during June 1994 and October 1995 confirmed shape and absolute values of the vertical profiles. A detailed analysis of the horizontal distribution of sulfide and methane in the deep part of the Gotland Basin around Stn AL 93 revealed a simultaneous increase of both concentrations during 1996 (Schmaljohann et al. 1998). On the basis of these observations and because of the sequential appearance of the changes it is quite conclusive that our data reflect time dependent changes rather than spatial heterogeneity.

Re-establishment of anoxic conditions

The comparatively high redox potential and steep redox gradient in May and October 1995 in the upper 15 cm of the sediment suggest a transition to suboxic conditions in the subsurface sediment due to the prolonged exposure to oxygen-containing bottom water. Apparently, oxidation processes in the sediment followed the establishment of oxic or anoxic conditions at the sediment surface after a considerable delay. This may be concluded from the low redox potential in the upper 15 cm in June 1994 when oxic bottom water was already in close contact with the sediment for more than a year and vice versa from the relatively
high redox potential when oxygen in the bottom water was consumed already, as in October 1995 and 1996. The transition from oxic to anoxic conditions in the bottom water was accompanied by (1) a drop of the redox potential in the top layer of the sediment; (2) an increase of sulfide concentrations in the pore water and (3) a temporary increase of integrated SRR. Furthermore, at the sediment surface a zone of active methane production was newly established and led to increased methane concentrations in the top layer of the sediment.

Sulfide

Sulfide accumulated from maximum concentrations of 1 mM in 1994 to 10 mM in October 1996. Parallel measurements at several stations in the deep central part of the Gotland Basin demonstrated an increase of sulfide concentrations over the time, which was most evident during 1996 (Schmaljohann et al. 1998). The mechanism for the significant increases also in the deeper layers of the sediment down to 80 cm remains unclear. Because of the lack of bioturbation, one would assume that diffusion processes dominate in the investigated system. However, considering theoretical aspects of diffusion in marine sediments (Duursma & Hoede 1967), the distance of sulfide diffusion from the main source of sulfide (peak of SRR at 8 to 13 cm sediment depth) is expected to lie in a range between 3 and 5 cm yr⁻¹. Therefore the increase of sulfide concentration in deeper sediment layers may be explained by increased sulfide production and decreased consumption, rather than by transport processes. Indeed, between June 1994 and May 1995 there was an average increase of SRR from 5 to 24 nmol cm⁻³ d⁻¹ in the sediment layer from 20 to 80 cm, which would be sufficient to explain an increase of the sulfide concentration of 7 mM during a year, if there is no consumption or loss. In view of the low sulfate concentrations, it is unlikely, however, that reduction of sulfate alone was responsible for such an increase. High concentrations of elemental sulfur at this station with maximal concentrations up to 7 mM and values of 2 mM down to 60 cm depth during June 1994 and May 1995 (Schneider & Imhoff 1996) would provide an alternative source for sulfide, though at present we have no indication for reduction of elemental sulfur in these horizons. It remains an open question how SRR in the deep layers can be influenced by processes at the sediment surface.

An important factor regulating the concentrations of free sulfide in the pore water of Baltic Sea sediments is the iron concentration, which is the limiting factor for the precipitation of sulfide as iron sulfide in Gotland Deep sediments (Boesen & Postma 1988). Therefore, the produced sulfide is supplied to the pool of free sulfide and causes immediate changes of the concentrations of free sulfide concentrations.

Rates of dissimilatory sulfate reduction

SRR at Stn AL 93 lie in the range of SRR in shallower parts of the Baltic and the North Sea (Limfjord: Jørgensen 1977; Limfjord, Aarhus Bay, Kysingfjord: Thode-Andersen & Jørgensen 1989; southwestern Baltic coast: Piker & Reichardt 1997) and exceed SRR in sediments of the Kattegat and the Belt Sea (Jørgensen & Revsbech 1989, Jørgensen & Bak 1991) by an order of magnitude. Repeatedly observed maxima of SRR in a sediment layer between 7 and 13 cm coincide with maxima of sulfide concentrations. However, neither the redox potential nor the sulfate concentration, the measured regulating factors of DSR, corresponded to these distribution patterns. Apparently, the redox potential as rate controlling factor involved in the regulation of DSR had largely been replaced by other environmental variables. This became most evident when a second SRR maximum could be detected in the upper sediment layer at relatively high redox potential. Concentrations of low-molecular-weight fatty acids (as preferred carbon sources for SRB) in the pore water also showed no striking coherence with SRR maxima. However, concentrations of dissolved free amino acids showed a negative correlation to these distribution patterns indicating a possible role as precursors of low-molecular-weight fatty acids (Piker 1995).

Integrated SRR (0 to 20 cm sediment depth) were in the same order of magnitude as SRR at a comparable station in the Gotland Basin investigated earlier (4.85 mmol m⁻² d⁻¹; recalculated from µg S kg⁻¹ dry weight, sum of integrated rates in sediment depths of 0 to 7 and 25 to 40 cm; Lein et al. 1982). Lein (1983) inferred from low SRR at a sediment depth of 0 to 7 cm that the main activity of anaerobic microorganisms must be in deeper regions of stable anoxic conditions. He found maximal SRR at a depth of 25 to 40 cm but did not analyze the sediment layer at 7 to 13 cm where maxima of SRR were found in this study. This view is in contrast to our findings of elevated SRR in the upper 2.5 cm sediment depth in May and October 1995, where rather suboxic conditions were present (0 to -100 mV). Apparently, sulfite reduction in the top sediment layer was inhibited by oxygen only in the first period with high concentrations of oxygen in the bottom water. However, with decreasing concentrations of oxygen in the bottom water, sulfate reduction activity in the subsurface layer increased as soon as oxygen content in the water phase was low. This led to the
establishment of a second maximum of SRR in the uppermost sediment layer in May 1995, a time when oxygen was still present in the bottom water. It even appears that the oxic period in the top sediment layer stimulated sulfate reduction rates after oxygen concentrations were reduced.

In several studies with shelf and open ocean sediments the model of Berner (1964) described the decomposition of organic matter by DSR quite well (Westrich & Berner 1984). An assumption of this model is that sulfate-reducing bacteria oxidize organic matter according to first-order kinetics. Furthermore, the application of the model requires that the sediment system is in steady state, which means that diffusion coefficient, rate of sedimentation and depth-dependent rates of sulfate reduction are independent of time. It could not describe the situation we found in Gotland Deep sediments (broken line in Fig. 5). Applied to these sediments, the model underestimates the SRR approx. 4- to 5-fold compared to the measured values. The observed deviation from the model indicates a dynamic rather than a steady-state situation in the investigated sediment, which could be seen in the development of concentration profiles of sulfate, sulfide and methane.

Methane

By comparing the methane flux rates in the near-surface region and in the zone of steep increase in the sulfate-free layers (Table 1) a remarkable difference can be noticed between Stn AL 93 and offshore marine sediments with oxic conditions in the overlying water column (e.g. Kattegat and Skagerrak: Iversen & Jørgensen 1985; or the Santa Barbara Basin: Kostur & Warford 1979). In the Gotland Deep sediment there is a widely linear increase in methane from the surface to a depth of about 50 cm, gradually changing into an exponential increase below that depth; the other offshore marine sediments show constant low or only threshold concentrations with no discernible gradient from the surface to the depth of sulfate depletion and a steep increase below. This difference can be quantified by calculating the methane flow rates in the near-surface horizons and in the zone of steep increase in the sulfate-free layers (Table 1). At Stn AL 93 the difference in the flux rates of the 2 layers, representing the proportion of anaerobically oxidized methane, is lower (71 to 86% ; Table 1) than in other investigated marine sediments, resulting in a comparatively higher methane flux to the surface. Data reported for anaerobic consumption of methane in other marine sediments range between 90 and 100% of the upward diffusing methane (Reeburgh 1980, Devol 1983, Iversen & Jørgensen 1985, Reeburgh & Heggie 1977).

Methane gradients extending to the sediment surface were found in some permanently anoxic marine basins, such as the Black Sea (Reeburgh et al. 1991), and in very productive coastal sediments, where methane production occurs close to the sediment surface, as in Saanich Inlet (Kuivila et al. 1990), Cape Lookout Bight (Hoehler et al. 1994) or in Kiel Harbour (Schmaljohann 1996).

Methane production rates

The experimentally determined methane production rates of the upper 20 cm of the sediment are within the range found in offshore marine sediments of the Black Sea (Heyer 1990), Bering Sea (Griffiths et al. 1982) or open Baltic Sea (Lein et al. 1981). They also fit very well to the data obtained by Lein et al. (1981) in the Gotland Deep (Stn 2622) with the radiotracer method (reduction of $^{14}$CO$_2$). These authors determined a rate of methanogenesis of 5.4 μmol m$^{-2}$ d$^{-1}$ within the top 15 cm (calculated after the given data), which is in the range rates in our study (3.0 to 8.5 μmol m$^{-2}$ d$^{-1}$). Coastal shallow brackish water sediments in the southern part of the Baltic Sea showed rates which range between 553 and 2016 μmol m$^{-2}$ d$^{-1}$ for open areas and $1.1 \times 10^5$ μmol m$^{-2}$ d$^{-1}$ for creeks and basins (Heyer et al. 1990). In Kiel Harbour, another site in the western part of the Baltic Sea with salinities of 15 to 22%, methane production rates in the sediments seasonally varied between 302 and 1963 μmol m$^{-2}$ d$^{-1}$. The fact that only negligible net production was detected by the applied headspace method at depths below 25 cm could be due to the fact that anaerobic methane oxidation consumed the produced methane. As discussed above on the basis of the methane concentration profiles this process is expected to occur in the depth range in question. However, Lein et al. (1981), applying the tracer method for measuring methanogenesis in Gotland Basin sediments of different depths, also found no or only minimum activity below the surface maximum and increasing activity at depths of 65 cm or more. In comparison with methane flux rates in the surface layers (Table 1), the integrated in situ methane production (0 to 20 cm) contributed 6 to 20% to the surface methane flux and may considerably influence the diffusion from the sediment to the water column during anoxic periods. The inverted methane gradients in the surface layers (Fig. 6) are indirect evidence of methane formation in the top sediment layer during anoxic periods. Though methane production at the sediment surface was superimposed by high vertical diffusion from deeper layers, production of methane and lack of aerobic methane oxidation during the anoxic period resulted in a 3-fold increase of concentrations in the 0 to 2 cm layer and a relative increase compared to the 2 to 4 cm layer.
Importance of sulfate reduction and methane production in the carbon budget of the Gotland Deep

The carbon consumption during sulfate reduction and methanogenesis can be calculated according to Ingvorsen & Brock (1982) based upon the consumption of 2 mol organic matter (CH₂O), corresponding to 24 g C, per mol H₂S and CH₄ produced:

**Oxidation half reaction:**
Oxidation of organic matter:
2(CH₂O) + 2H₂O → 2CO₂ + 8e⁻ + 8H⁺

**Reduction half reaction:**
Sulfate reduction: SO₄²⁻ + 8e⁻ + 9H⁺ → HS⁻ + 4H₂O
Methane formation: CO₂ + 8e⁻ + 8H⁺ → CH₄ + 2H₂O

Based on an average SRR of about 6 mmol m⁻² d⁻¹ (in the upper 20 cm of Stn AL 93 in June 1994, May and October 1995) the total annual oxidized organic carbon by DSR amounts to 52.5 g C m⁻² yr⁻¹. This is 44% of the total average primary production in the Central Baltic (120 g C m⁻² yr⁻¹; Gocke 1996). As Stn AL 93 is representative of only the deepest part of the Gotland Basin, this calculation is valid only for this part and would overestimate the average annual oxidized organic carbon by DSR, if applied to the whole basin.

On average methane production rates in the deeper sediment layers (>80 cm depth) were 259 μmol m⁻² d⁻¹ (Table 1), corresponding to 2.7 g C m⁻² yr⁻¹ and integrated rates of methane production in the sulfate-containing, surface layers (0 to 20 cm depth) were 5.8 μmol m⁻² d⁻¹ (corresponding to 0.05 g C m⁻² yr⁻¹). These calculations show that methanogenesis is not important in the main sulfate reduction layer, where the percentage of carbon consumption is only 0.1% of that consumed by DSR. Including the deeper layer this proportion rises to 4.6%.

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**LITERATURE CITED**


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