The annual cycle of the production and fate of DMS and DMSP in a marine coastal system

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The production of DMSP by phytoplankton and the fate through the marine system of both DMSP and DMS were followed for a period of 21 months in the natural environment by studying a North Sea tidal inlet. The major production and emission of DMS appeared to be limited to a period of only 2 months, which was closely linked to the presence of phytoplankton blooms. The production of DMS in the water column was not well correlated with chlorophyll a concentrations or plankton species composition. It was however, found to be closely related to the start of a Phaeocystis sp. bloom early in spring. Contrary to our expectations, the senescence phase of this Phaeocystis sp. bloom did not result in high DMS accumulation. It is postulated that this phenomenon could be the result of a bacterial consumption which increased with time in combination with a decreasing DMSP-lyase enzyme activity of Phaeocystis sp. A simple model, into which the overall DMSP consumption was introduced, could very well imitate the measured concentrations of DMS in the water column of the natural system studied.

INTRODUCTION

The production of dimethylsulphide (DMS) and its release into the air constitutes one of the most important biogenic sources of sulphur to the atmosphere. It is formed from the precursor DMSP (β-dimethylsulphonio propionate), which has a presumed osmoregulating or cryoprotecting function in marine algae (Dacey et al. 1987, Grüne & Kirst 1992, Baumann et al. 1994). In the atmosphere, DMS can be oxidized to non sea-salt sulphates and methane sulphonious acid which can be responsible for formation of cloud condensing nuclei, or CCN, which influence cloud formation and the backscattering of solar radiation and, thus, may have an influence on climate (Charlson et al. 1987, Caldeira 1989, Legrand et al. 1991, Malin et al. 1992).

There appears to be no direct correlation between chlorophyll a (chl a) and DMS concentration in oceanic surface waters (Dacey & Wakeham 1986, Turner et al. 1989, Bürgermeister et al. 1990, Leck et al. 1990). This makes it difficult to predict global DMS quantities by means of biomass estimations from remote sensing data. DMSP is associated with some algal species but there is a wide variation in the potential for algae to produce it. Diatoms are generally considered to be unimportant producers of DMSP, while other algal classes, including Dinophyceae and Prymnesiophyceae, are considered to produce relatively large amounts of DMSP and DMS (Keller et al. 1989, Gibson et al. 1990, Matrai & Keller 1993, 1994, Stefels & Van Boekel 1993). Factors which control the release of DMS into the water are not clearly understood, but possible mechanisms are: metabolic excretion, algal senescence and zooplankton grazing activity (Dacey & Wakeham 1986, Nguyen et al. 1988, Belviso et al. 1990, 1993, Stefels & van Boekel 1993, Levasseur et al. 1994). Loss factors for DMSP and DMS from the water column are bacterial consumption of DMS and DMSP, ventilation into the atmosphere and

Kwint & Kramer (1995, chapter 2) showed, in several experiments carried out in marine pelagic mesocosms, that a maximum release of DMS occurred directly after a phytoplankton bloom, during the senescence phase. In order to extrapolate these results to larger systems, like the North Sea, we monitored over a period of 21 months (November 1991 to July 1993) thereby taking in 2 phytoplankton spring blooms in the Marsdiep channel, a tidal inlet from the North Sea to the Dutch Wadden Sea. Our main goal was to see whether predictions are possible on a larger (natural) scale for the release of DMS through this extrapolation, taking into account the seasonal cycle of DMS and DMSP. Due to the length of the monitoring period, the results form a unique data set. In addition, mesocosm experiments were carried out with water from the same location (Kwint & Kramer 1995, chapter 2).

**MATERIALS AND METHODS**

Samples were collected in the Marsdiep tidal inlet from November 1991 to November 1992 (Series I) and from January to June 1993 (Series II) a few hundred metres off the coast of Den Helder, The Netherlands. Sampling was intended to take place at 10:00 h and at high tide in order to rule out any tidal or diurnal effects. This automatically implied a sampling interval of about 2 weeks. After it became clear, that changes in DMS concentration could occur very rapidly, the frequency was changed. Sampling was attempted at high tide as often as was logistically possible from the start of the phytoplankton spring bloom in 1993 (March to June).

Water samples were collected in a 10 l bucket that was rinsed thoroughly with seawater prior to sampling. Subsamples for chl<sub>a</sub>, phytoplankton, DMS and DMSP were immediately taken from this large sample. Chl<sub>a</sub> subsamples were stored in one litre polyethylene bottles in the dark, until analysis, which took place within two hours. One litre samples were filtered over glass fibre filters (Whatmann GF/C), extracted with 90% acetone and analyzed on a spectrophotometer according to standard procedures (Parsons et al. 1984).

Samples for the phytoplankton species identification were preserved and stored with Lugol's until further examination. Phytoplankton were identified and counted in a 5 ml Kolkwicz chamber under an inverted microscope. For each sample at least 20 fields were counted (magnification 400×).

Subsamples for the determination DMS and DMSP were taken in 1 l, dark, glass stoppered bottles and stored on ice until further treatment. The samples were treated immediately after returning to the laboratory (usually within 2 hours). A subsample of 10 to 25 ml was gently filtered over a Whatmann GF-C filter. It was considered important
that no pressure or suction was applied in order to minimize interference resulting from damaged cells. DMS was purged from the filtered water sample using high grade helium. The helium, containing the volatile compounds and water vapour, was dried by means of a Nafion permeation drier (Perma Pure Products, Farmingdale, USA) with nitrogen as the drying gas. After drying, the volatile compounds were collected on a cryogenic trap consisting of a straight glass tube containing Tenax-ta (Chrompack, Middelburg, The Netherlands) according to Kwint & Kramer (1995, chapter 2). After collection, the Tenax tube was closed with Swagelock caps and stored in liquid nitrogen until analysis (adapted from Lindqvist 1989). Storage tests with calibration gas showed that samples can be stored thus for at least 8 weeks without detectable change (chapter).

To determine the concentration of dissolved DMSP (DMSP\textsubscript{d}) a 10 ml sub-sample from the purged filtrate was placed in a 20 ml vial, which, after addition of 1 ml 10 M NaOH (Dacey & Blough 1987), was immediately closed with a Teflon coated septum. The vials were stored at 4°C until analysis. After hydrolysis by the NaOH, the sample was moved from the vial to the purge vessel in a closed system by means of two hypodermic needles and the helium flow (see Figure 1). The complete sample could be analyzed without any loss of DMS to the gas phase by using this procedure.

A 10 ml unfiltered seawater sample was hydrolysed in a similar way in order to assess the total amount of DMSP + DMS. The DMSP\textsubscript{p} concentration was calculated by subtracting the DMS and DMSP\textsubscript{d} concentration. Thus in principle DMSP\textsubscript{p} consists of all intra cellular DMSP, DMSP adsorbed to detritus and that incorporated into zooplankton and zooplankton faecal pellets.

DMS was analyzed according to Lindqvist (1989) on a Varian 3500 gas-chromatograph equipped with 10 m Poraplot-U capillary (0.53/0.7 mm inside/outside diameter) column (Chrompack) and a photo-ionization detector (10.2 eV, HNU Systems Inc., Newton, MA, USA). Hydrogen was used as the carrier gas. The detection limit was 1.5 pmol DMS. The coefficients of variation (CV’s) for the DMS analyses for independent analyses were no larger than 5% for samples with a concentration over 100 nM, to 17% for samples with a concentration below this value (Kwint & Kramer 1995, chapter 2).

Fluxes of DMS to the atmosphere were calculated from the water concentrations and from wind speeds (daily averages) obtained from a meteorological station near the sampling site, using the Liss/Slater model (Liss & Slater 1974).
RESULTS AND DISCUSSION

Figure 2 shows the results for the chl \( a \) and DMS measurements for both series I and II. During the winter period, DMS and chl \( a \) in these coastal waters showed relatively low background values as compared to the spring/summer values (chl \( a \) about 2.5 µg l\(^{-1}\), DMS below 0.5 nM) for both years. Each spring (March), the chl \( a \) as well as the DMS concentration started to rise and a maximum was reached within 1 month. During series I (1992), maximum chl \( a \) (20 µg l\(^{-1}\)) and DMS concentrations (17 nM) coincided. During series II we found similar values for chl \( a \) (18 µg l\(^{-1}\)) and DMS (18 nM) around the same date. The sampling frequency was increased in this period and it was found that a peak in the chl \( a \) concentration (30 µg l\(^{-1}\)) preceded the DMS peak by about 5 days. The maximum chl \( a \) and DMS concentrations in both years (DMS measured in duplicate) appear to be in surprisingly good agreement. Also, the very steep increase in the concentration of DMS was found for both years. Furthermore, these results agree well with the seasonal variations measured by Leck et al. (1990) for spring/summer data in the Baltic and with those measured by Liss et al. (1994) during a series of cruises in the southern North Sea. Our observations, especially those related to the dynamics of the DMS concentrations, are also in accordance with measurements in enclosure experiments with North Sea plankton (Kwint et al. 1992, chapter 3, Kwint & Kramer 1995, chapter 5, Kwint et al. 1997, chapter 4). The rapid changes in the DMS concentration mean that the major part of the production of DMS which may escape into the atmosphere is limited to a period of only about 2 months, i.e. in the spring and early summer. The limitation to such a short period of DMS fluxes to the atmosphere has major implications for climate modelling. Obviously the sampling frequency of the first series was not sufficient to cope with the dynamics of the phytoplankton and DMS related processes. For this reason we will focus our attention on the observations made in 1993 (series II) in this discussion.

Figure 3 compares the results of the DMS measurements with the calculated flux to the atmosphere during series II. The flux to the atmosphere appears to follow the same trend as the water concentration. Differences in the ratio between the calculated flux and DMS water concentration can be attributed to differences in wind speed. The wind
speeds observed during the measurement period varied but did not change to such an extent that major differences were found. As daily averaged wind speeds were used in these calculations, it would be possible that relatively large deviations would affect the actual daily fluxes. However, the maximum wind speeds encountered in the period of observation never exceeded 6 m\(^{-1}\)s. As a result only minor variations in fluxes are to be expected over shorter time scales, since the data are still in the rather horizontal part of the Liss-Slater (1974) model. Therefore we used an equilibrium model rather than a dynamic model. The latter would have been more appropriate under more variable wind speed conditions.

A comparison between the results obtained by a micro-meteorological measurement (gradient method) and by the Liss-Slater model calculation agreed well, at least for the wind speeds observed during these tests (A. Baart, pers. comm.). It is demonstrated in Figure 3 that the emission of DMS exceeds the background values during a period of only 6 weeks. In the period 5 April to 10 May 1994 we observed that the calculated flux of DMS ranged from about 10 to 15 µmol m\(^{-2}\) d\(^{-1}\). Assuming that the observations are representative for this period of the year, this implies that phytoplankton activity in these coastal waters during the spring period resulted in a total DMS emission of 0.4 to 0.6 mmol m\(^{-2}\) into the atmosphere. The total emission per year was calculated to be about 1.17 mmol m\(^{-2}\) yr\(^{-1}\). According to Andreae & Raemdonk (1983) and Bates et al. (1987b) coastal waters in the temperate zone have an emission rate of 1.1 to 1.8 mmol m\(^{-2}\) yr\(^{-1}\). These findings underline the importance of the DMS production during the short spring bloom period.

In order to enhance the resolution of the DMS and chl \(\alpha\) concentration development in Series II as depicted in Figure 2, these results are shown in more detail in Figure 4. Figure 4a clearly shows that the DMS concentration increases from approximately 2 to 14 nM in only 1 week. The DMS concentration remained at the same level for another week, with a maximum of approximately 18 nM and then decreased to half this value within 2 weeks. The chl \(\alpha\) development showed two successive peaks with an interval of about 2 weeks. The highest DMS peak followed the first chl \(\alpha\) peak only.
The phytoplankton composition during this period is presented in Figure 4b. After a first diatom bloom (consisting of 72 % *Skeletonema costatum*, 22 % *Nitzchia closterium* and 6 % other diatoms) there was a succession towards *Phaeocystis* sp. (mainly present in colonial form) causing the second peak in chl a. It is commonly found in these coastal waters that the initial diatom bloom is succeeded by a *Phaeocystis* development (Gieskes & Kraay 1975, Reid et al. 1990). During our observations the *Phaeocystis* bloom reached a maximum of $42 \times 10^6$ cells l$^{-1}$ in about 3-4 weeks (Figure 4b), which appears to be a normal value for these waters (Cadee & Hegeman 1991). Along with the development of the *Phaeocystis* cell numbers, a parallel development in DMSP$_p$ could be observed. Diatoms are generally considered not to contain a large amount of DMSP but *Phaeocystis* sp. is known to produce considerable amounts of DMSP (Keller et al. 1989, Liss et al. 1994).

In Figure 4c the concentrations of DMSP$_p$ and DMSP$_d$ are given. Starting in April, DMSP$_d$ increased from detection limits (0.06 nM) to 65 nM. After an initial drop, the DMSP$_d$ concentration came back to this level, after which it decreased to about 20 nM. DMSP$_p$ rose from 7 to no less than 1500 nM in 4 weeks after which a rapid decrease followed to about 500 nM in a few days. This maximum concentration of DMSP$_p$ is to be considered high for open sea values, but not uncommon in coastal waters with massive plankton (*Phaeocystis*) blooms (Liss et al. 1994, Stefels et al. 1995). The DMSP$_p$ peak coincided with the formation of a peaking of DMSP$_d$, with a ratio of DMSP$_p$ to DMSP$_d$ of about 20. The DMSP$_d$ production also coincided with the development of the DMS maximum (Figure 4a). It is known that the early stages of a *Phaeocystis* bloom enhance the transformation of DMSP$_d$ to DMS through the action of a DMSP-lyase enzyme (Stefels & Van Boekel 1993). It was also demonstrated in that paper that DMS was produced only after the bloom had peaked (senescence) in (axenic) *Phaeocystis* cultures, which is not consistent with our findings as we did not find a major release of DMS after the *Phaeocystis* bloom.

In order to get some idea about the DMSP$_p$ per unit of biomass, the ratio of DMSP$_p$ chl a, together with the chl a concentration is shown in Figure 5a. It is clear that the ratio of DMSP$_p$ chl a changes with the succession of the plankton species, and that this ratio increases steeply with the early development of the *Phaeocystis* bloom. Once the effects of the contribution of the diatoms to the chl a signal had ceased, a rather constant ratio of DMSP$_p$:chl a of about 20 nmol µg$^{-1}$ was established. This appears even to be true for the situation where the *Phaeocystis* bloom had passed its maximum. It is well known from the literature that *Phaeocystis* contain a relatively high amount of DMSP (per
biovolume) e.g. in comparison to diatoms (Keller et al. 1989, Liss et al. 1993b).
This is represented in Figure 5b. Here the amount of DMSP$_p$ is calculated based on cell
densities (Figure 4b) and the amounts DMSP$_p$ cell$^{-1}$ reported in the literature. For
diatoms (Keller et al. 1989) and Phaeocystis (Kwint et al. 1996, chapter 4) 17 fmol cell$^{-1}$
and 34 fmol cell$^{-1}$ respectively were used (note the difference in scales for the y axes).
It is highly unlikely, that the DMSP$_p$ in the diatoms was responsible for the release of
DMSP$_d$ observed in Figure 4c. Otherwise the diatoms should be producing large amounts
of DMSP that was immediately excreted without use. The Phaeocystis bloom is a
resultant of fast growing and decaying cells and we argue here, that the increase in
DMSP$_d$ can be attributed to the activity of Phaeocystis alone.
Since the formation of DMS is linked to the existence of DMSP, either by direct
transformation or via a route involving the formation of DMSP$_d$, one would expect that
a DMSP$_p$ peak would eventually yield a high DMS concentration, potentially much
higher than the peak observed after the diatom bloom. Surprisingly, the peak in
DMSP$_p$ during the Phaeocystis bloom was not followed by a major DMS development.
DMSP$_p$ may be transformed directly to DMS by Phaeocystis produced DMSP-lyase. Stefels & Van Boekel (1993) showed that this
process is particularly important in the log-phase of a culture, and not during
senescence. This process may have resulted in the relatively small but significant
increase in DMS concentration in the initial Phaeocystis growth phase. Although some
increase could be observed in the formation of DMS after the Phaeocystis bloom (Figure
4a), the amount does not reflect the large amount total DMSP that was potentially
available once the Phaeocystis bloom had collapsed in the last week of April. The drop
of 1000 nM in DMSP$_p$ concentration (and the drop of 30 nM DMSP$_d$) in the last week
of April was not at all counterbalanced by the 3 nM DMS increase (Figure 4a) and/or
the calculated flux (Figure 3). A comparison of the DMSP$_p$ loss per m$^2$ and the calculated
sea-air exchange of DMS, assuming an average water depth of 10 m, indicates that the

Figure 5. (a) Development of the DMSP$_p$ concentration in Phaeocystis sp. (◇) and diatom cells (●) (note different scales for Phaeocystis sp.
and diatoms)
(b) Comparison of the ratio DMSP$_p$:chl a (nmol µg$^{-1}$) (◇) and
the development in chl a (●) during series II (note different scales).
DMS flux consists of only an insignificant fraction in this period: DMSP\textsubscript{p} \text{loss} = 10 \text{ mmol m}^{-2} \text{ day}^{-1}  \quad \text{DMS flux} = 17 \text{ µmol m}^{-2} \text{ day}^{-1}.

A link is obviously missing in the step leading from DMSP\textsubscript{p} to either DMSP\textsubscript{d} or DMS. It is not likely that the sampling frequency applied could have caused a major DMS or DMSP\textsubscript{d} peak to be missed considering the width of the DMS peak before the Phaeocystis bloom. The missing sink may have been the result of the (bacterial) transformation of DMSP\textsubscript{p} directly into sulphur compounds that were not analyzed, export of DMSP\textsubscript{p} out of the system, e.g. by sedimentation of the particulate material (phytoplankton, faecal pellets), or by a fast turnover of DMS and DMSP\textsubscript{d}.

Physico-chemical processes (photochemical oxidation) are not considered to be very important (Brimblecombe & Shooter 1986). Of course the export of DMSP\textsubscript{p} cannot be excluded. However, the Marsdiep tidal inlet, the adjacent coastal waters and the Wadden Sea are turbulent systems, and it is not likely, therefore, that this mechanism can solely account for the export effect observed.

Therefore, either the direct or the indirect bacterial transformations seem to be the most dominant routes of DMS and DMSP conversion. Aerobic routes for DMSP transformation include the direct formation of acrylate and DMS (De Souza & Yoch, 1995), the formation of methylmercaptpropionate (MMPA) and the subsequent break down to acrylate and methanethiol, or the DMSP to MPA (mercaptpropionate) route via MMPA (Visscher & Taylor 1994, Kwint et al. 1996, chapter 4). Furthermore, a rapid utilization of DMS by bacteria could prevent the accumulation of this compound. DMS consuming bacteria (producing sulphate and CO\textsubscript{2} or DMSO) have been isolated (Hansen et al. 1993).

As a result we must assume that the rate of the transformation processes of DMSP\textsubscript{d} and DMS to other, not measured compounds is high. Otherwise we would have detected a large increase in DMS or DMSP\textsubscript{d} concentration in the samples upon release by or senescence of the Phaeocystis. Due to these fast transformation rates it is difficult to estimate which process and/or route is the most important. Kiene & Service (1991) and Bates et al. (1994) estimate that only a maximum of 30\% of the DMSP\textsubscript{d} will actually be converted to DMS. Of this DMS pool a major part can be consumed by bacteria (Kiene 1992, Kwint et al. 1996, chapter 4). The question remains, as to why this process of rapid bacterial conversion does not take place during the period when DMS and DMSP accumulates after the diatom bloom. It is postulated here that this may be the result of a change in the bacterial activity and populations.

Bacterial consumption is largely dependent upon the available substrate. For DMS consumers this means that, although they will always be present in low densities, the bacterial population will start to increase when sufficient DMS becomes present in the water column (Kwint et al. 1996, chapter 4). In the initial phase of DMS production, as a result of the diatom senescence, the bacterial population is not yet well developed.
Therefore, a relatively large proportion of the particulate and dissolved DMSP will lead to the formation of DMS (supported by a high DMSP-lyase activity from the initial Phaeocystis bloom). The presence of algal exudates enables a bacterial population to develop rapidly. A well developed bacterial population will be more effective in the facultative transformation of DMS and DMSPd. This may result in a situation which was reflected by our field observations: a DMS peak following a diatom bloom (or following the increase of Phaeocystis numbers) and the absence of such high DMS concentrations after the Phaeocystis bloom. The existence of such a process was confirmed in a mesocosm experiment where the actual bacterial DMS and DMSP consumption rate was measured, and where numbers of DMS-utilizing bacteria were measured (Kwint et al. 1996, chapter 4). In order to elucidate the possible role of bacteria in the conversion of DMS(P) a simple model was constructed. Based on a description of the fraction DMSPp, lost from the water column, a curve is fitted assuming a logistic growth curve. This curve is used to simulate bacterial activity during the period of the plankton bloom in the model situation.

The proportion of DMSPp that is converted (by bacteria) to other sulphur compounds than DMS can be defined as \((\text{DMSP}_p - \text{DMS})/\text{DMSP}_p\). This represents a net result, DMS may be well produced but turned over rapidly itself. In Figure 6 the field data calculated for the Marsdiep channel in April and May are presented. The data points show that not only the overall DMSPp-loss increases, but also the proportion of DMSPp lost from the watercolumn until almost 100% of the DMSPp is consumed at the end of the bloom without DMS accumulation. Assuming a bacterial process, the curve fitted through the field data points in Figure 6 represents a logistic growth curve not uncommon in microbiology studies, with the equation: \(N(t) = \frac{K}{K(a) + e^{-at}}\) (Schlegel & Schmidt 1985), with \(N\) representing bacterial numbers, \(K\) representing the carrying capacity of the system, \(a\) the growth constant and \(t\) the time. The simulated growth curve should be considered as a mathematical representation of total loss factors (of which bacterial activity is the major component).

To illustrate our hypothesis that the proportional increase of DMSP consumption is related to bacterial growth more clearly, we apply the logistic increase of DMSP consumption on a curve fitted through the measured DMSP data points in Figure 7 (solid line, compare with figure 4c). When the fraction (percentage) of DMSPp that is

![Figure 6. Proportion of DMSPp that is lost from the watercolumn without accumulation of DMS (●) in the Marsdiep channel in series II. A logistic growth curve (for bacteria) is fitted to the collected data (—).](image)
transformed, as identified by the logistic growth curve (dashed line), is subtracted from this DMSP$_p$-curve, a new curve appears that represents that part of the DMSP$_p$ that accumulates as DMS (broken line). When the actually measured concentrations of DMS from the field are added, a good agreement can be observed.

Although our model represents a rather simple and speculative approach, we think that it clearly illustrates the potential effects of varying bacterial activities, resulting in an outcome that reflects our findings in the natural system. It suggests a mechanism where a high density of Phaeocystis sp. does not automatically lead to a large production of DMS. Due to the nature of these complex interactions, small shifts in e.g. delayed bacterial activity may easily result in a situation where a massive DMS production follows a Phaeocystis bloom, as has been observed before (Liss et al. 1994).

CONCLUSIONS
The idea is emerging that only during special circumstances does the DMS concentration rise above normal background concentrations. Our results indicate that a considerable part (30% to 50%) of the annual emission of DMS over the sea/air interface takes place in the time span of about 6 weeks, closely related to the phytoplankton bloom. The DMS concentration is not controlled by the DMS(P) producers alone but by the balance between production and consumption of DMSP and DMS. Small shifts in bacterial activity or populations can lead to either an increase or decrease in the amount of DMS emitted to the atmosphere.

Although our quantitative results were collected in coastal waters, with a dominant role of Phaeocystis sp., we feel that the mechanisms involved have a universal application. That the major flux of DMS to the atmosphere may be limited to only a short period could have consequences for climate modelling. Furthermore, the fact that bacterial consumption of DMS and DMSP may play the major role in determining the amount of DMS available for ventilation to the atmosphere makes it difficult to predict whether a climate feedback mechanism (Charlson et al. 1987) exists via DMS formation in the oceans.