

Modelling the contribution of sea salt and dimethyl sulfide derived aerosol to marine CCN

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Abstract. The concentration of cloud condensation nuclei (CCN) in the marine boundary layer (MBL) was estimated from dimethyl sulfide (DMS) flux, sea salt (SS) emission, and aerosols entrained from the free troposphere (FT). Only under clean air conditions, did the nucleation of DMS derived sulfur (DMS CCN) contribute significantly to the MBL CCN. The accommodation coefficient for sulfuric acid mass transfer was found to be a very important parameter in the modeling the contribution of DMS to MBL CCN. The relationship between seawater DMS and MBL CCN was found to be non-linear mainly due to the transfer processes of sulfuric acid onto aerosols. In addition, sea salt derived CCN (SS CCN) and entrained aerosol from the FT (FT CCN) affected the MBL CCN directly, by supplying CCN, and indirectly, by behaving as an efficient sink for sulfuric acid. The SS CCN explained more than 50% of the total predicted MBL CCN when wind speeds were moderate and high. Sea salt and FT aerosol may often be more efficient sources of MBL CCN than DMS.

1 Introduction

Aerosols in marine air influence planetary albedo both indirectly, by forming cloud condensation nuclei CCN and increasing cloud-top reflectivity, and directly, by backscattering incoming solar radiation. Because the global radiation budget is sensitive to the amount of cloud and their reflectivity towards solar radiation, the CCN, on which cloud droplets can form, play an important role in the Earth's climate. Charlson et al. (1987) suggested the existence of a feedback mechanism between climate change and the flux of oceanic DMS. This proposed climate feedback mechanism is commonly known as the CLAW hypothesis after the

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four founding authors. Dimethyl sulfide produced by marine phytoplankton is ventilated into the MBL, and oxidized to form sulfur compounds and ultimately sulfate aerosols. These aerosols form CCN in the remote marine atmosphere. These CCN affect the cloud albedo and global radiation budget. The change in cloud albedo results in global temperature perturbation which affects the productivity of marine eco-systems and hence the concentration of oceanic DMS.

In this hypothesis, authors assumed that the oxidation of DMS as the major source of CCN over the oceans. Charlson et al. (1987) set up their argument on the strong belief that the non-sea salt (NSS) sulfate aerosols, which are final products of atmospheric DMS oxidation, are the major source of CCN in the remote marine air. They argued that sea salt cannot act as CCN because the concentration of sea salt particles at cloud level is typically not more than 1 cm⁻³ (Hobbs, 1971). This assumption in the CLAW hypothesis that sea salt cannot act ac CCN in the marine air has been challenged since its initial statement (Blanchard and Cipriano, 1987; O'Dowd and Smith, 1993). Moreover, the nucleation ability of DMS derived sulfuric acid in the MBL is still under debate (Katoshevski et al., 1999; Pirjola et al., 2000).

The role of sea salt as a major CCN source has been supported by numerous laboratory and field studies in the last decade (Blanchard and Cipriano, 1987; Cipriano et al., 1987; O'Dowd and Smith, 1993; Murphy et al., 1998; O'Dowd et al., 1999a, O'Dowd et al. 1999b). Blanchard and Cipriano (1987) argued that the background concentration of sea salt particle is $15 - 20 \text{ cm}^{-3}$, which can serve as CCN. They suggested this value based on field measurements and argued that the biological regulation of the climate is less obvious than CLAW hypothesis. More recently, using a volatility technique, O'Dowd and Smith (1993) and O'Dowd et al. (1999a) succeeded in distinguishing non-sea salt accumulation mode aerosol from sea salt CCN and quantified the accumulation mode sea salt aerosol concentrations as a func-



Fig. 1. The structure of the marine boundary layer CCN formation from DMS and sea salt flux for this work (the original figure was adapted from Raes, 1995).

tion of wind speeds up to 17 m s^{-1} . They concluded that sea salt can act as major CCN source. They also argued that sea salt aerosols are well mixed throughout the MBL because these aerosols have similar radii to sulfate aerosols. As a result it is likely that sea salt aerosols formed via film and jet drops may act as important sources of CCN in the MBL and they could dominate the concentration of total CCN under moderate and high wind speeds. Murphy et al. (1998) also supported these arguments. They measured chemical composition of accumulation mode marine aerosols and found that almost all particles larger than 0.13 μ m contained sea salt under unpolluted conditions.

Katoshevski et al. (1999) and Pirjola et al. (2000) have challenged the dependency of marine CCN concentration on DMS flux. Katoshevski et al. (1999) argued that aerosol concentration in MBL is very sensitive to the entrainment fo free troposheric aerosols and sea salt emission. They proposed that a prediction of the relative contribution to MBL CCN by sea salt and DMS derived sulfate is important to elucidate the relationship between marine biota and climate. Pirjola et al. (2000) also elucidated that DMS flux in MBL alone cannot explain new particle formation, instead they suggested that additional condensable species other than DMS-derived sulfuric acid is required to help thermodynamically stable sulfate clusters grow into detectable particle size.

As highlighted, the MBL CCN act as one of the key parameters in the Earth radiation balance. It is likelihood that SS is an efficient CCN source in the MBL, means that its role within the CLAW should be investigated. It is also important to investigate an effect which could arise from the interaction between DMS derived sulfur compounds and aerosols in the MBL. The main goals of this study are to model the MBL CCN concentration from various sources, such as DMS emission from seawater, SS flux, and entrainment of aerosols from the free troposphere.

2 MBL CCN model - description and considerations

In this research, MBL CCN were assumed to originate from marine DMS and SS. In addition, the entrainment of FT aerosols into the MBL was also considered as a source of CCN in the MBL. Some direct sources to the MBL, such as volcanic emissions, shipping emissions and transport from continents, are not explicitly treated. This simplified approach may underestimate the CCN concentration in the MBL because anthropogenic contributions to the MBL aerosol were ignored at the present stage. Nevertheless this simplification is a reasonable method in assessing the contribution of natural sources to the MBL aerosol system.

The structure of the MBL aerosol model for this work is outlined in Fig. 1. The accumulation mode aerosols which can act as CCN are categorised into two types according to their origins, SS CCN and NSS CCN. The direct emission of SS CCN into the marine atmosphere was modelled following O'Dowd et al. (1997). The model for NSS CCN formation from DMS was adopted and modified from Pandis et al. (1994) and Russell et al. (1994) (hereafter these two models will be cited as the P&R model). In addition to these two processes, the mass transfer of sulfuric acids onto existing aerosols were modelled by adopting the H_2SO_4 mass accommodation coefficient from Jefferson et al. (1997). The entrainment of free tropospheric gases and aerosols into the MBL are also considered as sources of MBL CCN.

2.1 Sea salt and CCN

A relationship between the film drop concentration and wind speed by O'Dowd et al. (1977) was used to estimate the SS CCN concentration in this work. These concentrations were described using a logarithmic relationship between wind speed and film drop SS:

$$\log N_{\rm SS-CCN} = 0.095U_{10} + 0.283\,,\tag{1}$$

where $N_{\text{SS-CCN}}$ is the SS CCN number concentration (cm⁻³) and U_{10} the wind speed at 10 m above the sea surface (m s⁻¹).

In this research, the contribution of SS to the MBL aerosol over a long time period needed to be considered because this work has assumed a steady state MBL CCN concentration. The parameter suggested by O'Dowd et al. (1997) as shown in relationship (1) was based on the average wind speed for one hour (O'Dowd, personal communication, 2000). In order to estimate a steady state SS CCN concentration from climatological wind data, such as global monthly mean wind speed distribution, the fluctuation of wind speed should be taken into account. This fluctuation of wind speed was approximated in this work using the Weibull distribution of wind speed probability.

The climatological SS CCN concentration (\overline{N}_{SS-CCN}) was estimated using the relationship:

$$\overline{N}_{\rm SS-CCN} = \int N_{(U10)} f_{(U10)} dU$$
⁽²⁾

Here, $N_{(U10)}$ is directly estimated from mean wind speed using Eq. (1) and $f_{(U10)}$ represents the wind speed as the Weibull probability density function:

$$f(U_{10}) = \alpha \beta U_{10}^{\ \beta - 1} \exp(-\alpha U_{10}^{\ \beta}).$$
(3)

This Weibull distribution is frequently used to describe the wind speed distribution over the ocean (Wanninkhof, 1992; Erickson and Taylor, 1989). In this study, the shape parameter (β) was assumed to be 2.0 (Palutikof, personal communication, 2000) and α was calculated using Eq. (4):

$$\alpha = \left(\frac{\Gamma(1+\beta^{-1})}{U_{10}}\right)^{\beta},\tag{4}$$

where U_{10} stands for the mean wind speed and Γ the Gamma function.

Figure 2 shows the SS CCN concentration derived from Eq. (1) after considering the Weibull probability density



Fig. 2. Directly estimated SS CCN concentration from wind speed from O'Dowd et al. (1997)'s relation (dotted line) and climatological SS CCN estimated considering the Weibull wind distribution (solid line). Note that the wind speed means are averaged for a period longer than 1 month and extremely high mean speeds are very rare.

function (Eq. 2). Though the probability of extremely high wind speed is very low, the SS CCN concentration at this speed is very high because of the logarithmic relationship. This makes the re-estimated SS CCN, shown as a solid line in Fig. 2, steeply increase at high mean wind speeds. This high value, up to 500 cm^{-3} , may be misleading because the relationship shown in Eq. (1) has only been tested for wind speeds lower than 17 m s^{-1} . For the default case simulation, a mean wind speed of 8 m s^{-1} was used. The climatological SS CCN concentration, which considered the fluctuation of wind speeds over the ocean, was calculated as 18 cm^{-3} . This was higher than the SS CCN concentration (11 cm^{-3}) derived from Eq. (1) considering a constant average wind speed.

2.2 MBL-FT exchange

The P&R model only dealt with the MBL as a closed system. An exchange between the MBL and the FT is believed to be possible even during inversions, through turbulent diffusion and microscopic phenomena (Kritz, 1983). The entrainment of aerosols from the FT into the MBL may provide a source of aerosols in the MBL (Raes, 1995; Clarke et al., 1998b; Raes et al., 2000). Clarke (1992) and Clarke et al. (1998b, 1999) reported, based on the experiments on the upper- tropospheric aerosol, new particles and sulfuric acid concentrations in the FT. Weber and McMurry (1996) also reported aerosol size distribution in the FT based on the observations at Mauna Loa Observatory. Their results suggest that the aerosol concentration in the FT shows a consistent shape. Clarke et al. (1998b) could find no evidence for particle production in the MBL in the first aerosol characterization experiment (ACE-1) experiment, but they reported enhanced 'new' particle formation in the FT. They suggested that the gases enter into the FT through cloud pumping, and then nucleation occurs. These particles subside into the MBL via entrainment process and contribute a quasi steady state particle and mass concentration in the MBL. Recently, Johnson et al. (2000) performed a Lagrangian experiment tracing a smart, constant balloon over the North Atlantic Ocean as part of the ACE-2 (The second aerosol characterization experiment) project. They concluded that sea salt aerosols, scavenging by cloud droplets, and the entrainment of FT air could affect the condensation nuclei concentration in the MBL. Furthermore, according to the report of Clarke et al. (1998a), the classic binary nucleation theory predicts no nucleation under the observed MBL conditions.

The exchange of gases and aerosol particles between the MBL and the FT was considered following Raes (1995):

$$\frac{dC_{\rm MBL}}{dt} = S_{\rm MBL} - \frac{Ve}{H}(C_{\rm MBL} - C_{\rm FT}),\tag{5}$$

where C_{MBL} and C_{FT} represent the concentration of a certain species C in the MBL and the FT, respectively, and S_{MBL} means the sources and sinks in the MBL. The entrainment velocity (Ve) is assumed to be 0.36 cm s^{-1} following Katoshevski et al. (1999) and Huebert et al. (1996) although this value is dependent on the spatially varying meteorological conditions. Recently, the entrainment rate between the MBL and the FT was reported as $0.7 \,\mathrm{cm \, s^{-1}}$ for the eastern North Atlantic (Sollazzo et al., 2000). The MBL height (H) was assumed to be 1000 m as a default. These entrainment of aerosols from the FT into the MBL was added to the P&R model as Eq. (5). Katoshevski et al. (1999) adopted the FT aerosol concentration from Weber and McMurry, and this data was also used in our research. The approach in our study assumed that there are fluxes of DMS out of MBL and other DMS derived sulfur compounds have same concentration for both MBL and FT.

2.3 DMS flux and CCN

The DMS flux was estimated using Turner et al. (1996)'s method, which could then be used to estimate DMS concentration in the MBL. We modelled the gaseous reactions of DMS, SO_2 and H_2SO_4 from the P&R model. The two main features that have been modified are the FT entrainment (Eq. 5) and the mass transfer processes of sulfuric acids onto NSS and SS aerosols.

Concentrations for DMS and SO_2 in the MBL were adapted from P&R including parameters which affect production and removal rates of these gases (see reactions (9) and (10) in Katoshevski et al., 1999). Concentration of gaseous sulfuric acid in the MBL was modified from P&R as:

$$\frac{d(\mathrm{H}_{2}\mathrm{SO}_{4})_{g}}{dt} = k_{\mathrm{SO}_{2}(\mathrm{OH})_{g}(\mathrm{SO}_{2})_{g}} - L_{\mathrm{nucl}} -K_{mt}^{1}N_{1}(\mathrm{H}_{2}\mathrm{SO}_{4})_{g} - K_{mt}^{2}N_{2}(\mathrm{H}_{2}\mathrm{SO}_{4})_{g} -K_{mt}^{3}N_{3}(\mathrm{H}_{2}\mathrm{SO}_{4})_{g} - \left(K_{\mathrm{dep}}^{\mathrm{H}_{2}\mathrm{SO}_{4}}(\mathrm{H}_{2}\mathrm{SO}_{4})_{g}\right) / H -R_{\mathrm{cloud}}^{\mathrm{H}_{2}\mathrm{SO}_{4}} + (\mathrm{Ve} / H) \Big[(\mathrm{H}_{2}\mathrm{SO}_{4})_{\mathrm{FT}} - (\mathrm{H}_{2}\mathrm{SO}_{4})_{g} \Big]$$
(6)

where $(H_2SO_4)_g$ is H_2SO_4 concentration, L_{nucl} the mass loss by nucleation and ignored at this stage, and N_i represents aerosol concentration for *i*th species. Here, i = 1: Aitken mode aerosol, i = 2: NSS CCN, and i = 3: SS CCN. The sulfuric acid deposition velocity ($K_{dep}^{H_2SO_4}$) was assumed as 1 cm s⁻¹ (equivalent to 0.86 d⁻¹ with 1000 m height). The term $R_{cloud}^{H_2SO_4}$ is H_2SO_4 mass transfer to cloud droplets and calculated using the relation, { $f_{cloud} \times (H_2SO_4)_g$ }.

The sectional mass transfer coefficient (K_{mt}^i) for *i*th species to the aerosol phase by condensation was calculated from Pandis et al. (1994)'s work as:

$$K_{mt}^{i} = \frac{2\pi D}{x_{i+1} - x_{i}} \int_{x_{i}}^{x_{i+1}} 10^{x} F(x) A(x) dx,$$
(7)

where $x_i = \log_{10}(D_i)$, $x_{i+1} = \log_{10}(D_{i+1})$. Here D_i and D_{i+1} mean diameter of particle in micrometer, D is diffusivity of sulfuric acids in air (0.1 cm² s⁻¹), and F is a coefficient correcting for free molecular effects and can be calculated as:

$$F(kn) = \frac{1+kn}{1+1.71\,kn+1.33\,kn^2}.$$
(8)

In this relation, kn represents the Knudsen number. In Eq. (7), A is a coefficient correcting for interfacial mass transport and calculated using the equation:

$$A = \left[1 + 1.33 \, kn F(kn) \left(\frac{1}{a_e} - 1\right)\right]^{-1}.$$
(9)

The term a_e is an accommodation coefficient of sulfuric acid onto aerosols.

Figure 3 shows the mass transfer coefficient calculated using Eq. (7). The equation was integrated from 0.03 to 0.1 μ m for Aitken mode aerosol (N_1), and 0.1 to 0.6 μ m for accumulation aerosol (N_2). It is clear from this figure that selection of accommodation coefficient plays an important role in modelling the amount of sulfuric acid transferred onto aerosols. Pandis et al. (1994) and Russell et al. (1994) used 0.02 as the accommodation coefficients from Van Dingenen and Raes (1991), and Fitzgerald and Hoppel (1998) used 0.1 as a default value for their MBL aerosol model. Recently, Jefferson et al. (1997) reported higher values compared with previous studies. They suggested an accommodation coefficient of H₂SO₄ onto (NH₄)₂SO₄ and NaCl as 0.73 and 0.79,



Fig. 3. Calculated mass transfer constant of sulfuric acids onto Aitken (dotted line) and accumulation mode (solid line) marine aerosols with various accommodation coefficients.

respectively. In this study, 0.70 was selected as a default value following Jefferson et al. (1997).

The aerosol size distribution spectrum was adopted from the P&R's original study. They used bimodal distribution of marine aerosol and a series of step functions. Aerosols in the accumulation mode (0.1 μ m 0.6 μ m) were defined as CCN for this study. The Aitken and accumulation modemarine aerosol concentration from DMS oxidation and heterogeneous nucleation was modeled using the reactions in P&R (see reactions (4) and (5) in Pandis et al., 1994).

3 MBL CCN model - results and discussion

3.1 Default case simulation

A modified P&R model was used to simulate the concentration of MBL CCN derived from DMS flux, SS flux and FT entrainment. The initial marine air was assumed free of gaseous DMS, SO₂, and H₂SO₄, and aerosols. Every reaction equation was integrated for 35 days to let these systems have enough time to reach their steady state. A test of this model showed that 35 days are enough time to stabilize. Default seawater DMS concentration was selected as 2 nmol 1^{-1} . The DMS flux from this assumption was calculated to be 4.7 μ mol m⁻² d⁻¹ for this base case simulation. During the integration, a wind speed of 8 m s⁻¹ was assumed as a mean value. Marine boundary air entered into the cloud once per a day, and a rain event occurred once every 10 days following the P&R. Values from the 11th to 30th day were used to calculate the average concentration of each species.

The DMS concentration reached a stable diurnal cycle from the 4th day of integration. This reached a maximum value of 110 ppt at 0700 and a minimum of 61 ppt at 1600. The average concentration for 20 days was 85 ppt. A (DMS)g concentration measurement by Andreae et al. (1995) reported a similar result over the tropical regions. The modelling result here also showed same (DMS)g diurnal variations reported by Yvon et al. (1996). The work by Yvon et al. (1996) reported higher DMS concentration than the result from this work because their sampling was done for the high seawater DMS concentrations ranging from 4 to 8 nmol 1^{-1} . The DMS diurnal cycle was mainly dominated by the concentration of OH radical. The OH concentration was assumed to show a maximum at noon and zero at night. During the night, DMS accumulated in the atmosphere because reactions with other species were ignored in this run for remote clean marine air. As the OH concentration increases until mid-day, their reactions with DMS become more active and the concentration of DMS decreases dramatically until 1600. If the entrainment term was ignored, the mean DMS concentration was 115 ppt. This means that the net flux of DMS from the MBL to the FT reached up to 26% of the total oxidized amount. This process was not considered in the P&R's approach although it now appears that this DMS loss by entrainment may suppress the formations of NSS CCN in the MBL.

The SO₂ diurnal cycle showed an inverse phase relationship with DMS. This is because, in this simulation, DMS oxidation processes were assumed to be the only source of SO_2 . The mean concentration of SO_2 was 45 ppt and the diurnal variation ranged between 27 and 62 ppt. A measurement by Yvon and Saltzman (1996) also showed that the SO₂ diurnal variation showed a negative correlation with the variation of (DMS)g concentration. In this default case simulation, the mean sulfuric acid concentration was 0.05 ppt. This gas phase H₂SO₄ concentration had 1-day and 10-day cycles. After the Sun rises, OH concentration increases and its reaction with SO₂ starts and H₂SO₄ begins to form. The sulfuric acid concentration reached its maximum around noon. Along with this production route, gaseous H₂SO₄ was transferred to the surface of aerosols, or deposited to the sea surface. A measurement by Weber et al. (1998) also reported the typical sulfuric acid concentration in MBL as approximately 0.1 ppt. Though other measurements by Weber et. al. (1997) are available from continental sites, the sulfuric acid concentration showed similar diurnal variation to our result. Besides this 1-day cycle, the concentration of sulfuric acid showed a steep increase every 10 days. Rain events were assumed to occur every 10 days in this calculation. These rain events cleaned the MBL of gases and particles. As a result, the transfer of sulfuric acid onto aerosol particles was less effective which explains the sudden increase of sulfuric acid following a rain event.

The sulfuric acid concentration was sensitive to the accommodation coefficient because it determined the effectiveness of the transfer of sulfuric acid onto existing aerosols.



Fig. 4. Predicted variations in (a) Aitken and (b) accumulation mode aerosols. The entrainment terms were included in this case. The averaged concentration for N_1 and N_2 were 198 and 126 cm⁻³, respectively.

When this base case was re-simulated using an accommodation coefficient of 0.02 (a value used by the P&R), the concentration of sulfuric acid rose to 0.34 ppt. This result was a factor of seven higher than the base case and affected the predicted concentration of MBL CCN. This will be discussed in the following section.

The estimated concentration variations for Aitken and accumulation mode MBL aerosol are shown in Fig. 4. There was a noticeable 10-day cycle for both modes because of the way the model simulates scavenging. As shown in Fig. 4a, Aitken mode aerosol occurred when MBL air was clean especially, after a rain event. As time proceeds, these Aitken mode aerosols grew to accumulation aerosols and their concentration decreased gradually until the next rain event. The 20 day averaged concentration of these small aerosols was 198 cm^{-3} .

The variation of accumulation mode aerosol is shown in Fig. 4b. It reached a maximum value of 146 cm^{-3} before a rain event. During a rain event, this concentration dropped dramatically to 78 cm^{-3} but increased gradually when the event finished. The average concentration for these CCN was 126 cm^{-3} .

These two size mode aerosols had an inverse phase diurnal cycle. In the case of Aitken mode aerosols (Fig. 4a), a maximum concentration was predicted at 0700 and a minimum at 1700. By contrast, accumulation mode aerosols showed a maximum value at 1700 and minimum at 0700 (Fig. 4b), and this is because of the growth of Aitken mode aerosols into CCN.

With the mean wind speed as 8 m s^{-1} , the SS CCN alone was estimated to be 18 cm^{-3} . This accounted for 14% of the predicted total MBL CCN concentration. These wind derived SS CCN affected the MBL CCN simulation directly, by supplying CCN, and indirectly, as a sink for sulfuric acid.

3.2 Sensitivity tests

3.2.1 Sensitivity to the accommodation coefficient

As highlighted in Sect. 2.3, one of the main features to be modified from the P&R model was the accommodation coefficient ($a_e = 0.70$) for sulfuric acid transfer onto aerosols. The results of sensitivity test for this selection are shown in Fig. 5. With an accommodation coefficient of 0.02, the mean CCN concentration was predicted to be 238 cm^{-3} . This value was higher than the base case estimation by a factor of two. There was no change in the mean wind speed from the base case simulation, so the SS CCN concentration remained the same for these two simulations. Figure 3 showed that the mass transfer constant for accumulation mode aerosol was very sensitive to the choice of the accommodation coefficient. As a result, the lower accommodation coefficients made the transfer of sulfuric acid onto aerosol less efficient, and the nucleation process became more active. As shown in Fig. 5b, the concentration of H_2SO_4 was seven fold higher than the base case, and the mean concentration of sulfuric acid was calculated as 0.34 ppt when the lower coefficient of 0.02 was used. This increase in sulfuric acid concentration accelerated the nucleation rate again, and as a result, the



Fig. 5. (a) MBL CCN and (b) H_2SO_4 variation with $\alpha = 0.02$ and $\alpha = 0.70$. The entrainment terms in the reactions were included in this case. The averaged concentrations for MBL CCN and (H_2SO_4) g were 238 cm⁻³ and 0.34 ppt, when an accommodation coefficient of 0.02 was used instead of 0.70.



Fig. 6. CCN concentration with entrainment (solid line), without entrainment (dotted line), and with entrainment of clean air from FT (dashed line). The mean values for these simulation were 126, 63 and 39 cm^{-3} , respectively.

MBL CCN concentration was much larger when using the P&R's original concepts.

When the SS CCN contribution and FT entrainment were ignored, the DMS CCN concentration was predicted as 64 cm^{-3} with $a_e = 0.70$, and 412 cm^{-3} with $a_e = 0.02$, respectively. The only source of CCN in this simulation was assumed to be the nucleation of sulfuric acid, so the error caused by selecting a wrong accommodation coefficient became larger when other physical processes that supply CCN into the MBL were ignored.

3.2.2 Sensitivity to the entrainment processes

The role of entrainment of FT aerosols into the MBL was tested performing simulations with two assumptions; firstly there was no exchange between the MBL and the FT, secondly there was an exchange but the FT air was free of aerosols. The results of CCN variation for these experiments are shown in Fig. 6. For the three cases, including the default case, the diurnal variation patterns were similar, but the concentrations were lower for the two tests. When no entrainment was assumed, the mean CCN concentration was $63 \,\mathrm{cm}^{-3}$. This value was half of the result from the default case simulation. This means that the aerosol entrainment from the FT may explain at least half of the CCN formation in MBL. A change in the entrainment velocity (Ve) did not affect the predicted MBL CCN concentration - 50% increase or decrease in this velocity did not affect the calculated CCN seriously. The dashed line in Fig. 6 shows the

Change in

CCN/

Tested

	D_2	$0.6\mu{ m m}$	1.0	116	
	Aitken nuclei deposition velocity	$0.04{ m cms^{-1}}$	0.02	127	
			0.08	125	
	CCN deposition velocity	$0.06{ m cms^{-1}}$	0.03	131	
			0.12	118	
	Nucleation empirical coefficient	10	5	126	
			15	139	
case when FT arristed. In this case only 39 cm^{-3} . A from SS CCN (12 plained only 21 c 3.2.3 Sensitivit	was free of aerosols but the entrain e, MBL CCN concentration was pre- bout half of this concentration wer 8 cm^{-3}), so the nucleation process cm^{-3} CCN concentration in this ca- y to each parameters	iment ex- edicted as e e derived c alone ex- a se. M c p o	For the base case sn mpirical enhancemer leation, but also terna variation in this coe ABL CCN concentra oncentration of sulfu- rocesses and the nucl f MBL CCN.	nulation, a va it factor to re ry nucleation fficient could tion seriously ric acid was v eation alone c	flu fle ef l n y. vei cai
The results of se	nsitivity tests for each parameter	that were	The choice of a re	liable CCN	ac

e of 10 was used for an ect not only binary nuffects. It was found that not affect the modelled This was because the ry small due to transfer nnot be the main source

The choice of a reliable CCN activation diameter was found to be very important. When $0.05 \,\mu m$ for this parameter, modelled potential CCN concentration was increased by 51%, but this meant that the concentration of Aitken mode aerosols was decreased by this factor. The activation diameter of CCN is, in practice, dependent on the supersaturation, so the lower activation diameter limit means more CCN

cussed in detail in Sect. 3.3.

used in the default case simulation are listed in Table 1. The

two most sensitive factors that affected the concentration of

MBL CCN were seawater DMS concentration and the mean

wind speed, because these provided the direct sources. The

relationships between CCN and these two factors are dis-

Table 1.	Change in the	predicted 2	20-days av	erage M	BL CCN	concentration	for	variations	in each	parameter	used	for the	default	case
simulation	n													

Default

value	value	$\rm cm^{-3}$	CCN/%
$2 \mathrm{nmol}\mathrm{l}^{-1}$	1	105	-17
	4	152	+21
$8{ m ms^{-1}}$	4	76	-40
	12	194	+54
1000 m	500	126	0
	1500	120	-5
$0.36{ m cms^{-1}}$	0.18	122	-3
	0.54	123	-2
0.8	0.7	119	-6
	0.9	141	+12
1 / 10 days	1/5	113	-10
$1.0 day^{-1}$	0.5	137	+9
	1.5	120	-5
5×10^{6} molecule cm ⁻³	2.5×10^{6}	104	-17
	7.5×10^{6}	142	+13
0.9	0.7	118	-6
	1.0	130	+3
$0.5 {\rm cm s^{-1}}$	0.25	130	+3
	1.00	121	-4
$1.0 \mathrm{cm}\mathrm{s}^{-1}$	0.5	126	0
	1.5	126	0
0.70	0.35	120	-5
	0.90	129	+2
$0.023\mu\mathrm{m}$	0.001	77	-39
$0.1\mu\mathrm{m}$	0.05	190	+51
0.6 μm	1.0	116	-8
$0.04 \mathrm{cm}\mathrm{s}^{-1}$	0.02	127	+1
	0.08	125	-1
$0.06{ m cms^{-1}}$	0.03	131	+4
	0.12	118	-6
10	5	126	0
	15	139	+10
	value 2 nmol 1 ⁻¹ 8 m s ⁻¹ 1000 m 0.36 cm s ⁻¹ 0.8 1 / 10 days 1.0 day ⁻¹ 5×10 ⁶ molecule cm ⁻³ 0.9 0.5 cm s ⁻¹ 1.0 cm s ⁻¹ 0.70 0.023 μ m 0.1 μ m 0.6 μ m 0.04 cm s ⁻¹ 10	value value $2 \text{ nmol } 1^{-1}$ 1 4 4 8 m s^{-1} 4 8 m s^{-1} 4 12 1000 m 500 0.36 cm s^{-1} 0.18 0.36 cm s^{-1} 0.18 0.36 cm s^{-1} 0.18 0.36 cm s^{-1} 0.54 0.8 0.7 0.9 1/5 1.0 day^{-1} 0.5 $5 \times 10^6 \text{ molecule cm}^{-3}$ 2.5×10^6 7.5×10^6 0.7 0.9 0.7 1.0 0.5 0.9 0.7 1.00 1.5 0.7 1.0 0.5 m s^{-1} 0.25 1.00 1.0 1.0 m 0.05 $0.6 \mu \text{m}$ 1.0 $0.023 \mu \text{m}$ 0.001 $0.1 \mu \text{m}$ 0.02 0.08 0.06 \text{ cm s}^{-1} 0.03 0.021 0.12 10	valuevaluecm ⁻³ $2 \text{ nmol } 1^{-1}$ 1105 8 m s^{-1} 4152 8 m s^{-1} 476121941000 m5001260.36 cm s^{-1}0.181220.36 cm s^{-1}0.181220.36 cm s^{-1}0.181220.36 cm s^{-1}0.181220.36 cm s^{-1}0.541230.80.71190.91411/10 days1/51131.0 day^{-1}0.5137.5120 5×10^6 molecule cm ⁻³ 2.5×10^6 1.01.00.5 cm s^{-1}0.251.0 m s^{-1}0.51.0 m s^{-1}0.50.700.351.200.023 μ m0.0010.770.1 μ m0.081250.06 cm s^{-1}0.030.121181051261.51371.01310.121411.0151.39

Parameter



Fig. 7. Predicted distributions of (a) gaseous H_2SO_4 (ppt), and (b) MBL CCN (cm⁻³) concentration. This simulation used the default parameters except for seawater DMS concentration and mean wind speeds.

available in the MBL with a lower supersaturation. This effect was not included in the modelling work reported here.

3.3 MBL CCN concentration for various seawater DMS concentrations and wind speeds

The estimation of MBL CCN for various wind speeds and seawater DMS concentrations was performed to show more complete picture of these relationships. These results for base case simulations are shown in Figs. 7 and 8. Figure 7a and b show the default case concentrations for predicted sulfuric acid and MBL CCN, respectively. In the case of the sulfuric acid, the concentration distribution was non- linear with respect to DMS concentration and wind speed. At low seawater DMS concentration, the concentration of sulfuric acid was very low because the DMS flux was also very weak. When seawater was highly concentrated with DMS, the flux affected the sulfuric acid amount more strongly, via wind speeds and transfer processes. The maximum sulfuric acid concentration occurred when the DMS concentration was high (around 4 nmol l^{-1}) and mean wind speed was between 10 and 12 m s^{-1} . Predicted MBL CCN is shown in Fig. 7b. This pattern differed from that of sulfuric acid because of the SS contribution. The MBL CCN concentration was more sensitive to mean wind speed than the seawater DMS concentration. For example, at a wind speed of $8 \,\mathrm{m \, s^{-1}}$, CCN concentration was not very variable as DMS concentration changed, but at DMS concentrations of 2 nmol 1^{-1} , the CCN concentration increased sharply as wind speed increased. In case of constant seawater DMS concentration, wind speed affected the CCN formation system by affecting both the DMS flux and the SS CCN production. As a result, at extreme high wind speeds, CCN concentration was not affected by DMS concentration. The DMS concentration variation affected CCN concentration only when the mean wind speed was weak.

This NSS CCN distribution was plotted (Fig. 8). to examine the CCN derived from NSS sources alone. The SS CCN contribution to the MBL CCN was omitted in this simulation so this figure represented the DMS CCN and the FT



Fig. 8. Predicted CCN concentration (cm^{-3}) when these are assumed to be composed of NSS CCN only. The flux of SS CCN was omitted in this simulation.

CCN distribution only. This pattern was similar to the sulfuric acid distribution because of their nucleation processes. The concentration of the NSS CCN increased as wind became stronger, but at high wind speeds, it decreased again. This arose from the lack of sulfuric acid under these conditions, where the transfer processes of sulfuric acid onto aerosol were very active. The maximum NSS CCN concentration did not exceed 150 cm⁻³, and occurred at high DMS concentrations and moderate wind speed.

This NSS CCN accounted for less than 10% of the total predicted CCN at high wind speed regardless of DMS concentration. Nevertheless the NSS CCN played a major component in marine CCN system when wind speed was less than 6 m s^{-1} . For these wind speeds, the NSS CCN accounted for more than 70% of the predicted total MBL CCN. The SS CCN accounted for more than 80% of the total predicted CCN when wind speed exceeded 12 m s^{-1} .

3.4 Comparison with other results

The MBL CCN model results were compared with the observed CCN concentration at Cape Grim (40° S, 144° E). Monthly CCN data at Cape Grim (Ayers et al., 1997; Ayers and Gras, 1991; Gras, 1989, 1990) were collected and the MBL CCN for the location were estimated for each month. For this comparison, monthly DMS concentration was extracted from Kettle et al. (1999)'s global data and monthly mean wind speeds were adopted from da Silva et al, (1994). Other parameters are the same as the base case simulation.

The observed and modelled CCN concentrations are shown in Fig. 9. The climatological wind data (da Silva et al., 1994) showed that the annual fluctuation for the loca-



Fig. 9. Comparison of modelled results with Cape Grim CCN data. The dashed line is from the simulation. The solid and dotted line represent the CCN concentration for Cape Grim at 1.2 and 0.23% supersaturation, respectively.

tion was smaller than DMS concentration variation (Kettle et al., 1999). The weakest monthly wind speed was 8 m s^{-1} in February, and the maximum was 9 m s^{-1} in August. The predicted SS CCN concentration varied from 16 to 31 cm^{-3} . In contrast, the concentration of seawater DMS showed a minimum (0.5 nmol 1^{-1}) in October and a maximum (3.1 nmol 1^{-1}) in March. Because the monthly mean wind speed was less variable than the seawater DMS concentration, the estimated MBL CCN were mainly affected by the variation of seawater DMS, shown as a dashed line in Fig. 9. The predicted MBL CCN showed a maximum in austral summer and minimum in winter, respectively. The maximum MBL CCN concentration was simulated as 142 cm^{-3} in March, and the minimum was 96 cm⁻³ in October. This seasonal change was predicted to be similar to the observed CCN, but the predicted CCN for the austral winter season was about 10% higher than the observed value. This is mainly because of the OH radical concentration. In this simulation, OH maximum value was assumed to be same all the year round, and this simplified assumption may over predict the DMS oxidation in the winter and underestimate for the summer. The activation diameter of CCN is dependent on the supersaturation, and the lower supersaturation means larger activation diameter limit and this leads less CCN available in the MBL as shown in Cape Grim data. This effect was not included in the work reported here. Nevertheless, the modelled MBL CCN concentration lay between 1.20% and 0.23% supersaturation CCN data (solid and dotted lines) for the other seasons.

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The model was run with DMS flux as a variable. This DMS flux was also dependent on DMS concentration and wind speeds so this aim was to compare the results with other empirical and model relations between DMS flux and CCN concentration. Figure 10 shows the simulated results for various wind speeds, such as 2, 4, 8, 12, and 16 m s^{-1} . For moderate and weak wind speeds, predicted MBL CCN lay between 150 and 200 cm⁻³ as the DMS flux changed. For the high mean wind speeds, for example at 16 m s^{-1} , MBL CCN concentration was 590 cm^{-3} and showed almost no slope with DMS flux increases because SS CCN was the major component above this extremely high wind speed. These estimations are compared with the empirical relationship of Lawrence (1993) and model results of Pandis et al. (1994), in Fig. 10.

The present MBL CCN model results are in accord with these two relationships for various wind speeds as shown in Fig. 10. Even for a high mean wind speed, such as 14 m s^{-1} , the modelled MBL CCN agreed well with the relationship of Lawrence (1993). For this wind speed, DMS flux less than 5 μ mol m⁻² d⁻¹ is very rare unless the seawater DMS concentration is very small. When compared with the steady state model results of Pandis et al. (1994), the simulated CCN concentration was larger for the weak DMS flux regime, and smaller for the strong DMS flux. These results suggests that the role of DMS flux in formation of MBL CCN is weaker than offered by Pandis et al. (1994)'s because of the SS contributions, sulfuric acid transfer onto aerosols, and FT entrainment processes.

3.5 Discussions

The P&R model, which we adapted and modified for this study, is composed of only two aerosol size bins, and this simplified approach has been discussed in detail between Reas and Van Dingenen (1995) and Pandis et al. (1995). Because of the simplified aerosol size distribution, some dynamical information might have not been included in this study. This approach might not be able to fully reflect the nucleation of new particles and their growth from 0.001 μ m to the first bin (0.023 - 0.1 μ m), though we tried to tune the nucleation rate by adopting an empirical enhancement factor 10^{10} – see reaction (4) in Pandis et al. (1994) – from Katoshevski et al. (1999). As already been discussed by Katoshevski et al. (1999) and Pirjola et al. (2000), theoretical binary nucleation rate have failed to reproduce the MBL aerosol. Pirjola et al. (2000) also elucidated the even ternary nucleation can hardly contribute to the general MBL aerosol concentration exept for the case when there is a significant removal of MBL aerosol such as due to precipitation. It is likely that whether a classical nucleation or tuned rate is used, this does not effect the MBL CCN vatiation seriously when compared with the effects due to mean wind speed. For example, a sensitivity test of this empirical factor showed 10% increase in CCN concentration with the value 10^5 , re-



Fig. 10. Modelled MBL CCN concentration as a function of DMS flux changes and other empirical suggestions. The solid line is the result of Pandis et al. (1994), and the shaded area represents the empirical relationship suggested by Lawrence (1993).

DMS flux / micromol m^-2 d^-1

spectively. By contrast, changes in wind speed from 8 m s^{-1} to 4 m^{-1} and from 8 m s^{-1} to 12 m^{-1} showed CCN concentrations to decrease by 40% and increase by 54%, respectively. It is obvious that the contribution of DMS to MBL CCN is suppressed by wind speed because of direct contribution of sea salt to CCN concentration. Latham and Smith (1991) have already suggested a negative feedback process between global warming and increased wind generated sea salt particles. Even though physico-chemical processes of the recently formed ultra-fine particles and growth of Aitken mode particles into CCN were simplified in our approach due to the limited aerosol size distribution, this important role of sea salt CCN should b seriously projected into the hypothesis between climate change and feedback due to marine biological activity and sea salt concentration change.

The result here not only highlights a direct sea salt contribution to CCN concentration, but also elucidates an additional role for sea salt particles in suppressing the action of DMS in global climate change scenatios – the mass transfer process of sulfuric acid onto wind generated SS CCN. As discussed in Sect. 3.3, NSS CCN concentration showed nonlinear dependency on wind speed and seawater DMS concentration. The role of DMS in CCN formation seems seriously decreased when wind speeds are moderate and high. This means that sea salt CCN act as an efficient sink of DMS derived sulfuric acid. The DMS derived SO₂ also efficiently react in sea salt cloud droplets (O'Dowd et al. , 1997), then reduce the production rate of NSS CCN. Though DMS flux as a function of wind speed was not dealt with in O'Dowd et al. (1999b) as in our approach, they also postulated a reduced role of DMS in cloud droplet formation. Even more, condensation of sulfuric acid onto super micron jet drop sea salt particles, which was not considered in this study, is nearly three times greater than that onto accumulation mode distribution according to a calculation by O'Dowd et al. (1997). This means that even our study might have overestimated the role of DMS in CCN formation in MBL due to the lack of sulfuric acid condensation onto super-micron sea salt particles. This fact also supports the importance of selection of accommodation coefficient when modeling the CCN concentration in marine environment as suggested by Pandis et al. (1995).

4 Conclusions

For completeness, the model and the concept used in this study should further include other sources of marine CCN, such as anthropogenic contributions from shipping (Capaldo et al., 1999; Huebert, 1999), volcanic emission (Graf et al., 1997), and transport of aerosols from continents. In case of considering other CCN sources to the marine atmosphere, a more complicated chemical-transport scheme should be added. In addition, nucleation processes in the MBL, and the relation between SS CCN and the wind speed should be further investigated. Each of these factors could yield an uncertainty with a factor of at least two.

Though many uncertainties remain in this study, some important findings and hypotheses result from this work. The sea salt flux from the sea surface is an important factor in the MBL CCN formation. The SS CCN can *directly* affect the system by providing more than 70% of total CCN observed in the MBL, especially for winter seasons over middle and high latitude regions. The modelling study in this work also suggests the possibility of an *indirect* SS aerosol effect. This indirect effect provides a sink for gaseous sulfuric acid through mass transfer suppressing the efficiency of nucleation of sulfuric acid in the MBL.

Sulfuric acid nucleation in the MBL is highly dependent on the mass transfer process onto existing aerosols. As a result, selection of an appropriate accommodation coefficient is a key issue in modelling the relationship between the DMS flux and the MBL CCN concentration. A selection of lower accommodation coefficients tends to yield the result that the amount of MBL CCN is highly dependent on the DMS flux. By contrast, this paper, which used 0.7 as the accommodation coefficient, showed that the relationship between MBL CCN concentration and DMS flux (or seawater DMS concentration) is not linear. This non-linearity arose from the direct CCN sources, such as SS and entrained accumulation mode aerosol, and the active transfer of sulfuric acid onto aerosols. In general, the FT aerosols and SS CCN similarly contribute to the MBL CCN concentration. The two direct source of CCN, SS and FT aerosol may be more important than the aerosol formed from the oxidation of MBL (DMS)g.

The key parameter which affects MBL CCN concentration and, in turn, the indirect radiative effect, may be wind speed, on which the SS emission rate and the DMS flux are highly dependent. A change in the global wind speed could influence the MBL CCN system and ultimately global radiation budget. This process could be more efficient than the CLAW hypothesis which focused on the relationship between temperature and DMS flux changes.

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