DIMETHYLSULFIDE IN THE MARINE ATMOSPHERE — FROM THE CRADLE TO THE GRAVE

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ABSTRACT

Air-sea exchange has profound implications for biogeochemical cycling and climate. Sea-to-air transfer of the biologically produced dimethylsulfide (DMS) is of particular interest because the gas is a precursor to sulfate aerosols, which affect atmospheric chemistry and earth’s radiative balance. In my dissertation, I discuss the air-sea exchange of DMS and the oxidation of DMS in the marine atmosphere. Chapter 1 provides background information on the formation of seawater DMS, theories behind air-sea exchange, oxidative destruction of DMS in the atmosphere, and the climatic impacts of sulfate aerosols. I also briefly describe our method for measuring DMS sea-to-air flux and provide an example of elevated DMS field near a front between two water masses. In Chapter 2, I compare air-sea transfer velocity measurements from the Southern Ocean to previous observations in temperate and tropical regions. I show that DMS transfer is suppressed in cold waters at high latitudes, not only due to reduced diffusivity, but also increased solubility. The transfer velocity of DMS is lower than those of the less soluble gases also because of this solubility effect. I shift my focus to biogeochemistry in the stratocumulus clouds topped Southeast Pacific in Chapter 3 and 4. In Chapter 3, based on shipboard and aircraft measurements, I estimate an average effective hydroxyl radical concentration from the budget analysis of DMS. In Chapter 4, I use a similar budget approach to analyze the diel cycles of sulfur dioxide and sulfate aerosols. I find that the dynamics of the boundary layer, which alternates from being coupled at night to decoupled during the day, significantly affects observed
chemical concentrations. I also provide examples of recent particle formation observed on the ship and aircraft, which were often associated with drizzle events and open cell structures in the clouds. Along with results from Chapter 3, I summarize the atmospheric sulfur cycling in the Southeast Pacific. While oxidized sulfur species from pollution emissions dominate the near shore region, DMS still supplies most of the sulfur mass, and at times new particle number, to the remote marine boundary layer.
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Chapter 1. Background

Over the past few decades, there have been numerous studies on the marine gas dimethylsulfide (DMS) from various disciplines, including biological regulation, atmospheric oxidation, air-sea exchange, and climatic feedback, etc. What initially spurred these interdisciplinary studies was the idea that the Earth is a self-regulating entity, where living constituents help maintain conditions favorable for life on the planet through feedback cycles in biogeochemistry and climate (Lovelock and Margulis, 1974). Such a feedback loop involving DMS was first hypothesized by Charlson et al. (1987) and coined the ‘CLAW’ hypothesis. Biologically produced in the upper ocean, DMS is ventilated to the atmosphere across the air-sea interface and constitutes ~90% of the global biogenic sulfur flux (Andreae, 1985). The oxidation of DMS in the marine atmosphere leads to sulfate aerosols that can serve as cloud condensation nuclei (CCN), which affect the formation, reflectivity, and lifetime of clouds. In the CLAW hypothesis, warmer climate leads to enhanced marine primary production, higher DMS concentration, more sulfate aerosols, and more CCN in the atmosphere. With amount of water vapor unchanged, more CCN leads to brighter and more persistent clouds, which reflect more incoming radiation back to space and induce cooling, thereby reducing biological production and completing the negative feedback loop (Falkowski et al., 1992). Verifying this hypothesis, however, has proven to be extremely challenging because it requires quantifying the biological production of DMS and the amount ventilated to the atmosphere, as well as understanding the complex chemical reactions of sulfur species in the marine air and the modifications of clouds due to sulfate aerosols on a large scale. The rest of this chapter consists of brief introductions on these various
aspects of DMS-related research. I draw particular attention to important and open questions in air-sea gas exchange and atmospheric sulfur cycling, which I attempt to address through observational and simple modeling works in Chapters 2, 3, and 4 of my dissertation.

1.1 Seawater DMS

DMS is produced by the enzymatic cleavage of dimethylsulfoniopropionate (DMSP) in the upper ocean, with typical DMSP to DMS turnover rates in the range of < 0.1~10 nM per day (Kiene and Service, 1991). Produced mostly by plankton, DMSP makes up the bulk of the reduced sulfur pool in marine systems (Simó et al, 2002). DMSP content is highly species-specific, with micro- and macro-algae among the most abundant sources. In the open ocean, phytoplankton containing the highest levels of DMSP include dinoflagellates and coccolithophores, with intracellular concentrations exceeding 100 mM (Sunda et al, 2002). In addition to a postulated osmotic role inside of phytoplankton cells (Keller et al, 1989), DMSP and its breakdown product DMS are thought to be antioxidants and protect algal cells from ultraviolet radiation (UVR) because they readily scavenge hydroxyl radicals and other reactive oxygen species (Sunda et al., 2002).

DMSP and DMS are released from phytoplankton to the water column via direct discharge and cell rupture. Direct discharge can be accelerated by stress, such as elevated UVR, high sea surface temperature (SST), and low nutrient availability (Kloster et al., 2006). Cell rupture occurs as a result of zooplankton grazing, bacterial, and viral attacks (Dacey and Wakeham, 1986; Sunda et al, 2002). Intracellular DMSP in phytoplankton is
found to be physically separated from the enzyme capable of catalyzing the DMSP cleavage reaction, suggesting healthy algal populations produce little DMS (Stefels and Boekel, 1993). Bacteria catalyze the degradation of dissolved DMSP in the water column to DMS.

The major sinks for DMS in the upper water column are bacterial degradation and photolysis, while ventilation to the atmosphere only removes a few percent of DMS (Kiene and Service, 1991; Toole et al., 2004; Bailey et al., 2008). Even though DMS is produced from algae, its concentration does not always correlate well with primary productivity or even the concentration of its precursor – DMSP. In bloom regions such as the Southern Ocean and the North Sea, mono-species phytoplankton blooms determine the DMS stocks and DMS appears to have a positive correlation with DMSP (Simó and Dachs, 2002). In contrast, in oligotrophic regions such as the Sargasso Sea, DMS appears to be uncoupled to DMSP or chlorophyll-a. Instead, SST and UVR account for most of the variability in DMS (Toole and Siegel, 2004). For such nutrient limited waters, the conversion from DMSP to DMS tends to peak during the decline of a bloom as opposed to during the active growing phase; the highest concentration of DMS is often observed in the summer despite lower levels of chlorophyll-a and DMSP than in the spring (Dacey et al, 1998).

To measure DMS in seawater, the gas is usually taken out of the water phase by either a purge-and-trap or equilibration method and then quantified in the gas phase by gas chromatography (GC) or mass spectrometry (MS). The purge and trap technique involves complete stripping of DMS from the water sample using an inert gas and trapping on an adsorbant (e.g. Bates et al. 2000). The equilibration method requires
establishing a Henry’s law equilibrium between the water and the headspace gas. Given
the complexity in the cycling of seawater DMS, it is no surprise that surface observations
show high degrees of variability both spatially and temporally (Bates et al., 1987),
presenting a challenge to numerical models. Diagnostic approaches to predict seawater
DMS concentration rely on some combination of chlorophyll-a (Anderson et al., 2001;
Aumont et al., 2002), mixed layer depth (Simó and Dachs, 2002), and solar radiation
dose (Vallina and Simó, 2007). While these algorithms capture some of the large-scale
patterns in seawater DMS, they do not reproduce local variability very well, necessitating
further in situ measurements. Updating the database assembled by Kettle et al. (1999),
Kettle and Andreae (2000) interpolated a global, monthly 1° x 1° climatology of surface
seawater DMS concentration. Significant differences between these two related
climatological datasets are found in high latitude regions, where observations are limited.
In Chapter 2, I report seawater DMS concentration measured during a cruise in March
and April of 2008 in the Southern Ocean.

1.2 Air-Sea Exchange of DMS

In order to understand the impact of DMS on climate, we first need to quantify its
sea-to-air flux. Transfer of gases across the air-sea interface has profound implications
on biogeochemical cycling and climate. The dissolution of carbon dioxide (CO₂) in the
surface ocean has increased the acidity of seawater and slowed down the build up of this
Greenhouse Gas in the atmosphere. Future prediction of climate change requires
knowing how fast the ocean can take up more CO₂. Chlorofluorocarbons, sulfur
hexafluoride, bomb ¹⁴C, and other anthropogenically produced gases dissolve in the
ocean have been used as tracers for water mass movements. DMS is ubiquitous in the surface ocean and is always emitted to the atmosphere. Previous estimates for the global sea-to-air flux of DMS range from 15 to 33 Tg yr\(^{-1}\) (Kettle and Andreae, 2000; Simó and Dachs, 2002). A more recent review by Elliot (2009) suggested an annual flux of 17~19 Tg S, with discrepancies mainly stemming from the uncertainties in the transfer velocity parameterization as a function of wind speed. In Chapter 2, I provide a comprehensive assessment on the measurements of DMS air-sea exchange in distinct oceanographic environments and over a wide range of wind speeds.

1.21 Theoretical Backgrounds in Gas Exchange

The air-water flux of a gas \( F \) is typically parameterized as the transfer velocity \( k \) multiplied by the concentration difference across the air-sea interface:

\[
F = k(C_w - \alpha C_a)
\]  

(1.1)

A positive \( F \) denotes an upward flux out of the ocean; \( k \) is in units of velocity and is by definition always positive; \( C_w \) and \( C_a \) are bulk waterside and airside concentrations, respectively; \( \alpha \) is the dimensionless Ostwald solubility of the gas and is related to the Henry’s law equilibrium (larger \( \alpha \), more soluble). Gases in the atmosphere have \( \alpha \) spanning several orders of magnitude, exemplified by the extremely insoluble sulfur hexafluoride (SF\(_6\)) and helium (He) to the very soluble amines and alcohols. Formulated from the perspective of the liquid phase, the term \( \alpha C_a \) represents the waterside concentration that would be in equilibrium with the airside.

Gas transfer velocity is limited by molecular diffusion and turbulent transport on both sides of the air-water interface. Quiescent molecular sub-layers form just above and
below the interface in which turbulent transports diminish and molecular diffusions dominate (Liss and Slater, 1974). We may conceptualize this system as an electrical circuit, with molecular diffusive sub-layers on both sides of the interface analogous to two resistors in series, concentration difference as the voltage potential, and $F$ as the current. Inversely related to resistance, total transfer velocity ($k$) is a function of transfer velocities in both the waterside and the airside ($k_w$ and $k_a$, respectively):

$$k = \left[ \frac{1}{k_w} + \frac{\alpha}{k_a} \right]^{-1}$$  

(1.2)

Transfer velocity does not depend on the concentration difference, just as the resistance of the resistors is independent of the voltage potential. If the interfacial concentration difference reverses in sign, so will the direction of the flux (i.e. into the water vs. out of the water). For gases with very low solubility, few molecules need to be transported across the interface before solubility equilibrium is achieved. Thus transport through the waterside molecular sub-layer is limiting to $k$ (i.e. $k \approx k_w$). Such gases, including carbon dioxide (CO$_2$) and SF$_6$, are said to be waterside controlled. The contribution to total resistance from the airside is only on the order of 0.2% for CO$_2$ at 20 °C. In contrast, exchange of highly soluble gases are controlled by airside resistance because reaching Henry’s equilibrium is the slow, rate-limiting step (i.e. $k \approx k_a/\alpha$), with water vapor (H$_2$O) representing complete airside control. Gases that are consumed rapidly in the surface water via chemical reactions, such as ozone (O$_3$) and sulfur dioxide (SO$_2$), can also be airside controlled. DMS is a largely waterside controlled gas, with the airside providing ~5% of the total resistance at 20 °C (McGillis et al., 2000). As solubility increases with decreasing temperature, airside resistance becomes more important. In Chapter 2, I
address the variance in DMS transfer velocity measurement due to the temperature
dependence in airside resistance.

The Schmidt number is the ratio between kinematic viscosity of a medium and molecular diffusivity of the gas in that medium. To remove variance due to the temperature dependence in waterside diffusivity or to relate one relatively insoluble gas to another, measured $k_w$ is often normalized to a reference waterside Schmidt number ($S_{cw}$) of 600 or 660 following a $S_{cw}^n$ relationship. The exponent $n$ is shown both theoretically and in the laboratory to vary from $-2/3$ for a rigid boundary to $-1/2$ for a non-rigid surface, agreeing with the measurement from Nightingale et al. (2000) in the North Sea. The reference 600 and 660 correspond to $S_{cw}$ of CO$_2$ at 20°C in freshwater and seawater, respectively.

1.22 Transfer Velocity Parameterizations

Wind stress ($\tau$) at the surface generates turbulence, which disrupts molecular sub-layers on both side of the interface and increases $k$. Most gas exchange parameterizations are functions of wind speed at 10 meters ($U_{10}$), but with significant differences in both magnitude and functionality, especially in high winds. Based on lake measurements of SF$_6$ and wind tunnel observations, Liss and Merlivat (1986) (LM86) modeled $k_w$ as three piece-wise linear functions of wind speed with increasing slope towards higher winds [cm hr$^{-1}$]:

LM86 $k_w = 0.17U_{10}(S_{cw}/600)^{-2/3}$ \hspace{1cm} $U_{10} \leq 3.6$ (1.3a)

$k_w = (2.85U_{10} - 9.65)(S_{cw}/600)^{1/2}$ \hspace{1cm} $3.6 < U_{10} \leq 13$ (1.3b)

$k_w = (5.9U_{10} - 49.3)(S_{cw}/600)^{-1/2}$ \hspace{1cm} $U_{10} > 13$ (1.3c)
Based on natural $^{14}$C disequilibrium and the bomb $^{14}$C inventory, Wanninkhof (1992) (Wa92) fitted a quadratic relationship between $k$ and ship-based $U_{10}$ [cm hr$^{-1}$]:

Wa92 $\quad k = 0.31 U_{10}^2 \left( \frac{S_{cw}}{660} \right)^{-1/2}$ (1.4)

From artificial injections of two volatile tracers ($^3$He and SF$_6$) and a non-volatile tracer (spores) in the North Sea, Nightingale et al. (2000) (N00) parameterized $k$ with both a linear and a quadratic term with respect to $U_{10}$ [cm hr$^{-1}$]:

N00 $\quad k = (0.222 U_{10}^2 + 0.333 U_{10}) \left( \frac{S_{cw}}{600} \right)^{1/2}$ (1.5)

Ho et al. (2006) (Ho06) decided on a quadratic dependence on $U_{10}$ based on $^3$He/SF$_6$ measurements in the Southern Ocean [cm hr$^{-1}$]:

Ho06 $\quad k = 0.266 U_{10}^2 \left( \frac{S_{cw}}{600} \right)^{1/2}$ (1.6)

Based on eddy covariance observations of CO$_2$ in the North Atlantic, Wanninkhof and McGillis (1999) (WM99) suggested a cubic fit between $k$ and $U_{10}$ [cm hr$^{-1}$]:

WM99 $\quad k = 0.0283 U_{10}^3 \left( \frac{S_{cw}}{660} \right)^{1/2}$ (1.7)

Using the same approach during a subsequent cruise in the Equatorial Pacific, McGillis et al. 2004 (M04) reported a weaker wind speed dependence in $k$ and a large positive intercept at $U_{10} = 0$ [cm hr$^{-1}$]:

M04 $\quad k = 8.2 + 0.014 U_{10}^3 \left( \frac{S_{cw}}{660} \right)^{-1/2}$ (1.8)

The authors attributed this discrepancy to a limited wind speed range and strong diurnal heating. From recent observations of the transfer velocity of DMS in the Northeast Atlantic, Huebert et al., 2010 (Hu10) found near linear dependences in $k$ with $U_{10}$ as well as with the friction velocity ($u_*$) in the wind speed range of 4~12 m s$^{-1}$ [cm hr$^{-1}$]:

Hu10 $\quad k = (-5.0 + 2.4 U_{10}) \left( \frac{S_{cw}}{660} \right)^{1/2}$ (1.9a)

$\quad k = (-1.5 + 58.9 u_*) \left( \frac{S_{cw}}{660} \right)^{1/2}$ (1.9b)
The linear dependence with \( u^* \) is consistent with air-sea exchange theory from Csanady (1990). The friction velocity relates to 10-m wind speed (\( U_{10} \)) through the drag coefficient (\( C_D \)): 

\[
u^* = C_D U_{10} = \frac{\tau}{\rho_a},
\]

and more directly represents wind stress, with \( \rho_a \) being the air density. The above wind speed parameterizations are plotted in Fig. 1.1.

Except for M04, the formulations agree fairly well up to a wind speed of \( \sim 8 \) m s\(^{-1} \), above which they diverge significantly (by a factor of \( \sim 2 \) in high winds). The large differences in high winds can be partly explained by the different solubility of the gases, which affect how much they can be transferred by bubbles.

Waves break and whitecaps form on the sea surface in high winds. The added surface area from injected air bubbles (Woolf, 1997) and the turbulence from breaking waves (Monahan and Spillane, 1984) can cause additional bubble-mediated gas transport (\( k_b \)). Scaling up from a single bubble model, Woolf (1997) (Wo97) parameterized \( k_b \) for CO\(_2\) [cm hr\(^{-1} \)]:

\[
Wo97 \quad k_b = 2450 f_{wh} \alpha^{-1} \left[1 + (14\alpha S_{rw}^{-1/2})^{-1/1.2}\right]^{-1.2}
\]

(1.10)

Here \( f_{wh} = 3.84 \times 10^{-6} U_{10}^{3.41} \) is the whitecap fraction (Monahan and O’Muircheartaigh, 1980). Assuming the solubility dependence in (1.10) holds for other gases, we see that \( k_b \) is larger for an insoluble gas than for a soluble gas. Total transfer velocity at high wind speed is the sum of direct transfer and \( k_b \). However, the fractional contributions to gas exchange at high winds from direct and bubble-mediated transfer are still poorly quantified in the field. I attempt to separate these two mechanisms in Chapter 2 from DMS transfer velocity observations.
Exchange of airside controlled gases is traditionally studied by looking at water vapor transfer. Based on wind tunnel measurements, Liss (1973) (L73) proposed a linear relationship between airside transfer velocity \( k_a \) and \( U_{10} \) that is not gas specific [cm s\(^{-1}\)]:

\[
L73 \quad k_a = 0.005 + 0.21U_{10}
\]  \hspace{1cm} (1.11)

Liss and Slater (1974) subsequently suggested an estimate of \( k_a \) for any gas from that of H\(_2\)O by multiplying by the square root of the ratio of the molecular weights (MW). From resistance theory, Duce et al. (1991) (D91) estimated \( k_a \) using an \( MW^{1/3} \) dependence; a linear relationship with wind speed was assumed by choosing a fixed \( C_D \) [cm s\(^{-1}\)]:

\[
D91 \quad k_a = \frac{100U_{10}}{700 + 45MW^{1/3}}
\]  \hspace{1cm} (1.12)

Derived from water vapor transfer models, the parameterization of \( k_a \) from McGillis et al. (2000) (M00) is [cm s\(^{-1}\)]:

\[
M00 \quad k_a = 0.183U_{10}(MW / MW_{H_2O})^{1/2}
\]  \hspace{1cm} (1.13)

1.23 NOAA COARE Gas Transfer Model

While wind-speed dependent parameterizations might be adequate for estimating transfer velocities of specified gases in certain environments, they tend to oversimplify complex physical mechanisms that drive gas exchange. In comparison, models that attempt to approximate the physical processes at the air-water interface are more likely to be applicable to a wider range of gases and distinct environments. Originally developed to estimate bulk heat and moisture fluxes, the NOAA COARE 3.0 air-sea bulk flux model (Fairall et al., 2003) predicts waterside and airside gas transfers separately by estimating the thickness of the respective diffusion sub-layers. Total transfer velocity is then
calculated following (1.2). The waterside transfer velocity contains terms for direct
transfer and bubble-mediated transfer in breaking waves [cm hr⁻¹]:

\[ k_w = 360000u_s(\rho_w/\rho_a)^{-1/2}[h_wS_{cw}^{1/2} + \kappa^{-1}\ln(0.5/\delta_w)]^{-1} + B(k_b) \]  \hspace{1cm} (1.14)

Here \( \rho_w \) is the density of water, \( \delta_w \) the waterside molecular sublayer thickness (~1 mm), \( \kappa \) the von Karman constant (0.4), and \( h_w = 13.3/(A\varphi) \). The second term inside the bracket
on the RHS of (1.14) is small compared to the first term. In \( h_w \), \( A \) is an empirical constant
and tuned to 1.3 by Blomquist et al. (2006) using early eddy covariance measurements of
DMS; \( \varphi \) accounts for surface buoyancy flux enhancement of the transfer and only
becomes important in low wind speed. The term \( B(k_b) \) refers to the Wo97
parameterization for bubble-mediated exchange in high winds (1.10), with \( B \)
 provisionally set to 1.0. Figure 1.2 shows \( k_w \) predicted by COARE for CO₂, DMS,
acetaldehyde, acetone, and methanol without the bubble effect \((A=1.0; B=1.3)\) at 20 °C.
Normalized to \( S_{cw}^{-0.5} \), direct waterside transfer of these gases would be essentially
identical.

The airside transfer velocity in COARE is formulated analogously to direct
transfer on the waterside, but with the molecular sublayer thickness empirically estimated
from \( C_D \) [cm s⁻¹]:

\[ k_a = 100u_s[13.3S_{ca}^{1/2} + C_D^{-1/2} - 5 + 0.5\kappa^{-1}\ln(S_{ca})]^{-1} \]  \hspace{1cm} (1.15)

Figure 1.3 shows \( k_a \) of acetaldehyde, acetone, and methanol predicted by COARE. A
heavier molecule has a lower diffusivity in air, higher airside Schmidt number \( S_{ca} \), and
thus lower \( k_a \). Compared to (1.11), (1.12), and (1.13), the COARE formulation of \( k_a \) has
a nonlinear wind speed dependence, though the magnitudes from different models for a
particular gas are fairly similar. The total transfer velocity of CO₂, DMS, acetaldehyde,
acetone, and methanol normalized to the respective temperature at which $S_{cw} = 660 \left(k_{660}\right)$ are shown in Fig. 1.4 ($A=1.0; B=1.3$). The large divergences in $k_{660}$ among these gases are principally due to their different solubility, which affects transfer velocity both through (1.2) and (1.10) in high winds. Methanol is by far the most soluble gas out of this group and has the lowest $k_{660}$. Since CO$_2$ and DMS are both largely waterside controlled, their $k_{660}$ are similar in low-to-mid winds before the onset of whitecapping. COARE 3.0 qualitatively captures the diverging curvatures of their transfer velocities in higher winds due to bubbles (Hare et al., 2004; Blomquist et al., 2006), suggesting that this physics-based model is more universal than any simple wind speed parameterization. However, quantitatively how well does the COARE model predict DMS transfer over a wide range of wind speeds? Southern Ocean Gas Exchange Experiment (SO GasEx) provided us for the first time with a large sample size of DMS transfer velocity observations up to a wind speed of 16 m s$^{-1}$, which I use to assess the performance of COARE. I also compare the DMS transfer velocity to previous observations from CO$_2$ and $^3$He/SF$_6$.

1.24 Measurements of DMS Flux and Transfer Velocity

Global DMS flux estimates rely on climatology of waterside concentration and empirical parameterizations of transfer velocity; both are coarse in spatial resolution and subject to substantial errors. We measure the sea-to-air flux of DMS directly with eddy covariance (EC) on a ship in the open ocean. Dividing flux by the concurrent air-sea concentration difference in DMS, which is largely dictated by the seawater concentration, yields the transfer velocity.
Equating turbulent flux typically measured at 10~20 m above the ocean surface with the true surface flux requires the assumption of a small vertical gradient in flux associated with near-steady state and horizontal homogeneity. With DMS emitted from the ocean surface and diluted by entrainment of free troposphere air, there is a decreasing trend in DMS concentration with height in the marine boundary layer. Blomquist (2010) showed that for typical seawater DMS fields and an 800 m deep boundary layer, measuring at 18 m only underestimates true surface flux by ~1%. The steady state assumption is justified over the ocean because the lifetime of DMS (~1 day) is much longer than the averaging time required by EC (~1 hr).

On a ship, we measure the atmospheric DMS concentration at a frequency of 20 Hz using an Atmospheric Pressure Ionization Mass Spectrometer with isotopically labeled standard (APIMS-ILS). Determination of the ambient DMS mixing ratio is based on the ratio of ambient DMS and a triply deuterated d$_3$DMS standard that is continuously added to the sample air at the inlet. The use of the labeled standard ensures that the accuracy of the measurement is largely unaffected by changes in the sensitivity of the instrument (Bandy et al. 2002). A high turbulent flow is used to preserve high frequency variability during the transit from the inlet to the detector. However, an air drier, necessary for a high ionization efficiency of DMS, adsorbs DMS slightly and limits the half-power response frequency to 1~2 Hz.

We measure winds and ship’s three-dimensional motion at the same frequency as DMS using a co-located sonic anemometer and motion sensor. Following Edson et al. (1998), we subtract ship’s motion from measured winds to obtain wind velocities along the ship ($U$), across the ship ($V$), and in the vertical ($W$). To account for mean flow
distortions caused by ship’s superstructure, the wind velocity coordinates are rotated so that both $V$ and $W$ average to zero on hourly-basis. An arbitrary ~100 s segment of DMS and motion corrected $W$ is shown in Fig. 1.5, with covariance between the two variables visible at time scales ranging from a few seconds to over a minute. The lag correlation between DMS and $W$ usually shows a maximum at ~1.5 s that corresponds to the time it takes for air to travel from the inlet through the air drier to the APIMS. Accounting for the lag time, turbulent flux is calculated as $\overline{DMS'W'}$ in 10-minute intervals and reduced to hourly averages. The primes denote instantaneous deviations from the means, with the overbar indicating time averaging. Figure 1.6 shows a typical DMS:$W$ cospectrum, with most of the flux found in the frequency range of 0.5~0.005 Hz. The quadrature spectrum, or the imaginary component of cross correlation, fluctuates around zero, as expected for two variables directly correlated. When winds are clearly out-of-sector (i.e., from a stern quadrant) or when the ship is turning, flow distortion at the inlet is increased, which results in an absence of a clear peak in the DMS:$W$ lag correlation and a non-zero mean in the quadrature spectrum; those periods are excluded from the hourly averages. An empirical inlet attenuation correction is applied to the DMS:$W$ cospectra for signal loss at high frequencies. An estimate for additional losses at high and low frequencies is obtained by fitting observation with a theoretical cospectral function for neutral conditions (Kaimal et al., 1972). Together, these corrections on flux are generally less than 10%.

In addition to sensor resolution, ambient variability (stochastic nature of eddies) contributes to the random noise of EC flux, which may be reduced through averaging (noise $\propto N^{-1/2}$, where $N$ is the number of independent measurements). At a wind speed of
~8 m s\(^{-1}\) and in a near-neutral atmosphere, Blomquist et al. (2010) showed that relative uncertainty in DMS flux is on the order of ~30% for an integration time of one hour, and further increases when the boundary layer is stable. The instrumentation details of APIMS-ILS and the measurement of DMS flux by EC are described in Blomquist et al. (2010).

There are significant deviations among transfer velocity parameterizations as functions of wind speed (e.g. Fig. 1.1). EC measurements also show amplifying scatter in high winds, which, in addition to the aforementioned sampling uncertainty, may be due to environmental variability. A number of potential controlling factors that could affect gas exchange have been proposed, including surfactants, wave properties, and boundary layer dynamics. From the Deep Ocean Gas Exchange Experiment (DOGEE) in the Northeast Atlantic, artificial surfactant patches were laid out over areas of several squared kilometers. DMS transfer velocity measured in the presence of surfactants by EC appears to be slightly suppressed compared to observations outside surfactant patches (M. Salter personal communication, 2010), though the reduction is much smaller than previously demonstrated in laboratory (e.g. Frew et al. 2004). From the same experiment, DMS transfer velocity is seen to increase with wave steepness and decrease with wave age. But because both wave steepness and age are correlated with wind speed, their effects on the DMS transfer velocity are, to first order, captured in the wind speed dependence.

The stability of the marine boundary layer, as related to the air-sea temperature difference, can also affect the precision of EC measurements. When the sea surface is warmer than the overlying air, as typical of tropical and temperate environments, the
boundary layer is statically unstable. In the surface water, besides wind shear, buoyancy-driven convection provides another mechanism for turbulence production. In contrast, when the boundary layer is statically stable (air warmer than water), shear-generated turbulence becomes intermittent or suppressed and the boundary layer tends to be shallow. With sea-to-air and entrainment fluxes unchanged, a shallower boundary layer implies a greater systematic underestimate of the true surface flux as a result of measuring at 10~20 m above the surface. In the Southern Ocean Gas Exchange Experiment (SO GasEx) near South Georgia Island, the air temperature frequently exceeded SST. In Chapter 2, I examine the effects of marine boundary layer stability on EC measurements.

1.25 Example of DMS Flux Measurement in a Region of Elevated Seawater DMS

Elevated concentration of seawater DMS was often found on the edge of an eddy or near a front between distinct water masses (T. Bates and P. Matrai, unpublished data). We encountered such a vastly enhanced DMS field during Leg 1 of the Vamos Ocean-Cloud-Atmosphere-Land Study Regional Experiment (VOCALS-REx) in October 2008. The ship sailed southward at 11 knots from Panama City, past Ecuador, to the Southeast Pacific. At ~2°S latitude, the sea surface temperature decreased by ~6°C and salinity increased by ~2.5 psu over a distance of 100 km (1.7(a)). At the interface between the warm equatorial water and the cold, nutrient-rich, upwelled water, biological productivity must have been much higher. At the southern edge of this front, seawater DMS concentration (shown every 30 minutes) increased from a typical level of ~2 nM to ~80 nM in a space of just ~20 km. The atmospheric DMS concentration (every minute) and
sea-to-air flux (every 10 minutes) also increased from ~200 pptv and ~3 μmoles m$^2$ day$^{-1}$ to ~1700 pptv and ~150 μmoles m$^2$ day$^{-1}$, respectively (1.7(b)). Because the ship was sailing into surface wind from the south, enhanced atmospheric DMS and DMS flux were detected before the seawater DMS peak and their signals appeared to be somewhat smeared out. The high correspondence between atmospheric DMS concentration and flux was aided by the fact that the front passing occurred in the nighttime, when DMS was not photochemically destroyed by OH. The temporal resolution of the semi-automated seawater DMS measurement by purge-and-trap/GC was limited to every half an hour. A much faster measurement, such as by membrane equilibration/MS, would have afforded better spatial resolution near the edges of such fronts or eddies.

1.3 Atmospheric Sulfur Cycle

In the atmosphere, the oxidation of DMS yields, among other products, sulfur dioxide (SO$_2$) and non-sea-salt sulfate aerosols (SO$_4^{2-}$). These water-soluble and acidic SO$_4^{2-}$ aerosols affect atmospheric chemistry (Charlson and Rodhe, 1982) and also influence the global radiative balance directly by scattering of incoming radiation and indirectly through their effects on cloud microphysics (Charlson et al., 1992; Charlson et al., 1987).

When ambient water vapor content reaches supersaturation, aerosols larger than a critical size act as cloud condensation nuclei (CCN) and activate to form cloud droplets (Hoppel et al., 1986). Given the same amount of liquid water, more CCN leads to a greater cloud droplet number concentration and reduced mean droplet size, which results in more reflective and possibly longer lived clouds. The impact of naturally derived
SO$_4^{2-}$ on clouds should be the most noticeable in regions lacking anthropogenic CCN (Twomey, 1991), such as over the remote ocean. Gunson et al. (2006) modeled a 1.6 °C warming / 0.8 °C cooling for a decrease / increase of DMS emission by a factor of two. Moreover, their modeled DMS flux out of the ocean responds to warming (cooling) by slightly increasing (decreasing) over time, consistent with a negative feedback. Other studies suggest less pronounced climatic effects from DMS with increasing input of anthropogenic SO$_2$ and SO$_4^{2-}$ to the atmosphere. From the modeling work of Boucher and Lohmann (1995) and Woodhouse et al. (2010), the indirect radiative forcing by SO$_4^{2-}$ aerosols is found to be much more sensitive to total SO$_4^{2-}$ and cloud droplet number concentration than to the spatial distribution of DMS flux and background SO$_4^{2-}$ level. The apparent insensitivity to DMS is in part related to the difficulty in separating the natural and anthropogenic contributions to SO$_4^{2-}$ aerosols. The climatic effect of DMS might be more apparent locally. Boucher et al. (2003) showed high spatial heterogeneity in their modeled indirect radiative forcing, varying from close to zero to -5 Wm$^{-2}$. More accurately constraining the climatic effects of DMS-derived SO$_4^{2-}$ aerosols requires being able to model their regional distributions. The complex formation and removal pathways of sulfur species in the atmosphere, however, are still not well-understood and are subject to ongoing research. Using observations from a ship and an aircraft, I closely examine the budgets of DMS-derived SO$_2$ and SO$_4^{2-}$ aerosols in the marine atmosphere of the Southeast Pacific in Chapters 3 and 4.

1.31 DMS Oxidation
About 95% of atmospheric DMS oxidation takes place in the gas phase, with the hydroxyl radical (OH) being the most important oxidant, resulting in a DMS lifetime of ~1 day (Lucas and Prinn, 2002; Chen et al., 1999). The principle oxidizing free radical towards trace gases in the atmosphere (Levy, 1971), OH is formed from the reaction between \(^1\text{D}\) O, a product of ozone (O\(_3\)) photolysis, and water vapor. The tropospheric concentration of OH peaks in midday and becomes non-existent at night due to its high reactivity and low selectivity. Globally, the highest OH levels are found in the tropics characterized by high humidity and strong actinic fluxes (Seinfeld and Pandis, 2006).

DMS is oxidized by OH through two distinctive pathways: H-atom abstraction and direct OH-addition (Hynes et al., 1996). The abstraction pathway dominates at high temperatures and leads to SO\(_2\), which can be further oxidized to SO\(_4^{2-}\). The addition pathway, in contrast, speeds up at low temperatures and leads to dimethylsulfoxide (DMSO), dimethylsulfone (DMSO\(_2\)), methanesulfonic acid (MSA), SO\(_2\), and other minor products. At 298 K, about 80% of the oxidation of DMS goes through the abstraction pathway; at 285 K the two pathways are about equal (Wallington et al., 1993). A simplified DMS oxidation scheme in the remote marine atmosphere is illustrated in Fig. 1.8, as simplified from Davis et al. (1999). Studies from Christmas Island in the remote Central Pacific have shown a pronounced diel cycle in DMS that is dictated by the reaction with OH (Bandy et al., 1996). As DMS is emitted from the ocean to the atmosphere continuously, its concentration builds up overnight in the absence of OH and reaches a maximum by sunrise. During the daytime, DMS is oxidized by OH and its concentration declines to a minimum by sunset.
At night in a polluted marine environment, DMS can also be lost through the reaction with the nitrate radical (NO₃), which is formed from the reaction between NO₂ and O₃. Because NO₃ rapidly photolyzes during the daytime, its concentration is only significant in the nighttime. The oxidation of DMS by NO₃ proceeds through the H-atom abstraction pathway and has a rate constant ~4 times lower than that of the overall OH reaction at 298 K (Sander et al., 2006). Because of its localized production and short residence time (1~4 days), NO₃ is estimated to remove more DMS in the heavily industrial Northern Hemisphere (8.7%) than in the cleaner Southern Hemisphere (6.5%), where NO₂ input is lower (Kloster et al., 2006). In the remote tropical Pacific, NO₃ accounts for only a few percent of the total DMS oxidative loss (Davis et al., 1999; Chen et al., 2000). The oxidation of DMS by NO₃ becomes more important at high latitudes where sunlight is limited in the winter, allowing for the build up of the radical.

Other oxidants of DMS that have been studied include O₃ and halogen radicals. Since the mixing ratio of O₃ generally increases with altitude, the DMS oxidation by O₃ is thought to be more important in the upper troposphere (on the order of 5%, Boucher et al., 2003). Halogen radicals produced from sea-salt aerosols, in particular bromine oxide (BrO), can catalytically oxidize DMS (Seinfeld and Pandis, 2006). The oxidation of DMS by BrO follows the addition pathway and leads to DMSO. Some authors (Sciare et al., 2000; Boucher et al. 2003) have suggested that the removal of DMS by BrO can be significant, which is supported by recent measurements of high concentrations of BrO in the tropical Atlantic (Read et al., 2008). However, techniques for measuring BrO at concentrations typically encountered in the open ocean (differential optical absorption spectroscopy and mass spectrometry) are still challenged with detection limits. Until
halogen radical concentrations can be measured accurately and routinely at ambient levels, their importance towards DMS remains mostly speculative.

What dictates the diel variability in DMS in the Southeast Pacific, in a marine region that is also influenced by pollution transport from the continent? Is OH still the main oxidant, or do pollution-derived nitrate radicals and halogen radicals play important roles in DMS oxidation? In Chapter 3, from the mass balance of DMS measured in VOCALS-REx, I estimate an effective concentration of OH in the stratocumulus-topped marine boundary layer of the Southeast Pacific. This OH-equivalent oxidant concentration is compared to estimates from complex photochemical models. By examining the diel cycles of DMS as well as the oxidants, I place constraints on the oxidation of DMS due to NO₃ and BrO radicals.

1.32 Products of DMS Oxidation

Bandy et al. (1996) first reported from measurements at Christmas Island that SO₂ increased during the day when DMS decreased due to OH oxidation. The anti-correlation suggested that DMS is indeed the main source for SO₂ in the remote marine atmosphere. The conversion efficiency from DMS to SO₂ has been found to be 70-90% (Davis et al., 1999; Chen et al., 2000; Shon et al., 2001; Faloona 2009). Among oxidation products of DMS, only SO₂ has the potential to make more new particles, while the others increase aerosol mass by condensing upon preexisting surfaces.

SO₂ is oxidized in both the gas phase and the aqueous phase. The gas phase reaction between SO₂ and OH forms sulfuric acid vapor (H₂SO₄(g)):

\[
SO_2 + OH \rightarrow HO\text{SO}_2 \rightarrow SO_3 + HO_2
\]  (1.16a)
$SO_3 + H_2O \rightarrow H_2SO_4(g)$  \hspace{1cm} (1.16b)

In most circumstances, the non-volatile $H_2SO_4(g)$ condenses upon preexisting aerosol surfaces and grows them to larger sizes, thereby increasing $SO_4^{2-}$ aerosol mass. Under specific conditions, $H_2SO_4(g)$ molecules undergo gas-to-particle conversion and form new nm-sized particles, resulting in sudden and dramatic increase in the particle number concentration (Weber et al., 1999). Environments favorable for such nucleation processes typically have high $H_2SO_4(g)$ and humidity, as well as low aerosol surface areas and temperatures. New particle formation events have been observed in clean air high in the free troposphere (Clarke 1998b), near cloud outflow regions (Hegg et al., 1990; Perry and Hobbs, 1994), in the remote marine boundary layer (Hoppel et al., 1994), and adjacent to a region of precipitation (Clarke et al., 1998a). Clouds remove aerosol surface area by scavenging and inject moisture into the surroundings. In the outflow region, elevated actinic flux enhances OH and so $H_2SO_4(g)$ production. Mixing of air is that cleaned of particles by clouds and air rich in reactive vapors creates an ideal condition for nucleation (Clarke et al, 1998a). Rainfall similarly creates favorable conditions for nucleation by removing aerosols through precipitation scavenging.

Freshly nucleated particles grow by condensation of vapor and coagulation by Brownian motion to a detectable size of ~3 nm in minutes to an hour (Weber et al., 1999) and to 10 nm with in a few hours (Hegg et al., 1992). From shipboard measurements, Covert et al. (1992) observed a ~3 hr time lag between a recent nucleation event and the increase in concentration of 20~24 nm particles. In the Southeast Pacific, we observed frequent occurrences of recent new particle formation from an aircraft as well as from a ship.

Where in the atmosphere and under what conditions did the nucleation take place? Was
the source of volatile vapors in the gas-to-particle conversion natural or anthropogenic?

In Chapter 4, I provide examples of new particle formation observed on both sampling platforms and discuss their correspondence to drizzle events and open cell structures in clouds.

The oxidation of SO₂ in the aqueous phase is much faster than in the gas phase. In an aqueous solution, SO₂ is equilibrated to SO₂.H₂O, HSO₃⁻, and SO₃²⁻ (together S(IV)) in an analogous fashion to CO₂. The equilibrium reactions are pH dependent with HSO₃⁻ being the predominant species at typical atmospheric pH. Aqueous phase oxidation of SO₂ can take place in cloud water mostly by H₂O₂, or in sea-salt aerosols by O₃, leading to SO₄²⁻ aerosols (S(IV)):

\[
\begin{align*}
S(IV) + H₂O₂ & \rightarrow S(VI) + H₂O \\
S(IV) + O₃ & \rightarrow S(VI) + O₂
\end{align*}
\]

These secondary SO₄²⁻ typically make up the bulk of the submicron (fine) aerosol mass, whereas sea-salt from wave breaking account for the majority of the supermicron (coarse) number as well as total aerosol mass. Due to the dissolution of CO₂ and other acidic species, cloud water typically has a pH under 5 (Chameides, 1984), which makes H₂O₂ the principal oxidant of SO₂ in cloud. Hegg (1985) suggested cloud processing to be the most important mechanism for the conversion from SO₂ to SO₄²⁻.

The S(IV)-H₂O₂ reaction is largely pH insensitive because the pH dependences in the equilibrium of HSO₃⁻ and in the H₂O₂ reaction cancel. The S(IV)-O₃ reaction, on the other hand, is extremely pH-sensitive. The reaction rate approximately varies inversely with the square of [H⁺] and is greatly enhanced at high pH. Sea-salt tends to be more basic than the fine mode aerosols because of the initial alkalinity of seawater (pH~8.1)
and the carbonate buffering capacity. Some authors postulated oxidation in sea-salt aerosols by O$_3$ to be a major sink of SO$_2$ and source of SO$_4^{2-}$ mass (Sievering et al., 1991; Faloona et al., 2010). However, both the uptake of SO$_2$ and the production of SO$_4^{2-}$ increase the acidity of the sea-salt, making the O$_3$ reaction self-limiting.

Aside from oxidation reactions, SO$_2$ is removed from the MBL by dry deposition to the ocean surface. The rate of dry deposition of aerosols strongly depends on particle size. Deposition velocity is very slow for the accumulation mode, which is the reason why aerosols in that size range accumulate in the atmosphere. Dry deposition is much faster for coarse particles due to gravitational settling and has been proposed to be a significant mechanism for the removal of non-sea-salt SO$_4^{2-}$ (Sievering et al., 1991; Faloona et al., 2010). However, given that little non-sea-salt SO$_4^{2-}$ mass is usually present in the coarse mode, it appears unlikely for dry deposition to be a major sink for SO$_4^{2-}$. Because of its relatively low solubility in precipitating droplets, SO$_2$ is not significantly affected by wet deposition. In contrast, the highly soluble MSA and SO$_4^{2-}$ are efficiently removed from the MBL by wet deposition. The efficiency of wet scavenging is a complex function of precipitation droplet size, rain rate, and aerosol size distribution.

In a marine atmosphere covered by stratocumulus clouds, what fractions of SO$_2$ are removed due to cloud processing and sea salt oxidation? I estimate the heterogeneous reaction rates of SO$_2$ in the Southeast Pacific in Chapter 4. I also constrain the dry and wet deposition fluxes of SO$_4^{2-}$ based on observations using a mass balance approach. In addition to being produced in situ, SO$_2$ and SO$_4^{2-}$ aerosols may be transported over a long distance in the free troposphere and mixed into the marine boundary layer via
entrainment. I use the aircraft observations to estimate the entrainment of these two sulfur species. To further examine the sources and sinks of SO$_2$ and SO$_4^{2-}$, a simple one-layer model is used to calculate their expected diurnal cycles. Along with results from Chapter 3, I summarize the atmospheric cycling of DMS, SO$_2$, and SO$_4^{2-}$ in the Southeast Pacific in Chapter 4.
Figures

Figure 1.1 Transfer velocity parameterizations of waterside controlled gases at $S_c = 660$ as functions of wind speed. While most models agree fairly well in low to moderate winds, they diverge significantly in high winds. Transfer of DMS appears to be lower than those derived from sparingly soluble tracer gases, including CO$_2$ and $^3$He/SF$_6$. 
Figure 1.2 Waterside transfer velocity ($k_w$) of CO$_2$, DMS, acetaldehyde, acetone, and methanol without the bubble effect predicted by the NOAA COARE 3.0 air-sea bulk flux model at 20 °C. $k_w$ of different gases can be related by the relationship $S_{cw}^{-0.5}$. 
Figure 1.3 Airside transfer velocity ($k_a$) predicted by the NOAA COARE 3.0 air-sea bulk flux model at 20 °C. $k_a$ decreases as the molecular weight (MW) of the gas increases.
Figure 1.4 Total transfer velocity ($k_{660}$) predicted by the NOAA COARE 3.0 air-sea bulk flux model. Soluble gases having much lower transfer velocity than insoluble gases due to both increased airside control and less partition in air bubbles.
Figure 1.5 A ~100 s time series of DMS and motion-corrected W showing covariance on different time scales, from a few seconds to over a minute. Due to the time it takes for air to travel from the inlet to the mass spectrometer, DMS usually lags behind W by ~1.5 seconds, a time accurately determined from the lag correlation analysis between these two variables.
Figure 1.6 Typical hourly cospectrum (flux) and quadrature between DMS and $W$. Most of the flux is found between 0.005~0.5 Hz. The loss of flux at very high and very low frequencies is usually only a few percent. Quadrature (the imaginary component of the cross-correlation) fluctuates around zero, as expected for a high degree of correlation between two variables.
Figure 1.7 (a) Crossing of a front from the warm and fresh equatorial water to the cool and salty water in the south during the 1st Leg of the VOCALS-REx cruise; the mean wind was perpendicular to the front (b) A large enhancement in seawater DMS (30 min), atmospheric DMS (1 min), and sea-to-air flux (10 min) on the southern side of the front crossing. As the ship sailed into the wind, elevated atmospheric DMS and flux were observed slightly before the peak in seawater concentration. A faster seawater DMS measurement would have afforded better resolution near the edges of similar fronts and eddies.
Figure 1.8. Sulfur oxidation scheme in the remote marine atmosphere simplified from Davis et al., (1999). DMS oxidation by OH has two pathways, addition and abstraction. The abstraction pathway leads to SO₂, which is further oxidized to SO₄²⁻. The addition channel leads to DMSO, DMSO₂, MSA, and also SO₂. The symbol ‘?’ denotes proposed reaction steps with no supporting kinetic or field data.
References


2010.


Chapter 2. Sea-to-air DMS Transfer Velocity

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Abstract

Dimethylsulfide (DMS), a biogenic sulfur gas produced in the surface ocean and constantly emitted to the marine atmosphere, has recently emerged as one of the key gases with which to directly quantify the air-sea transfer velocity ($k$). In the Southern Ocean Gas Exchange Experiment (SO GasEx), we measured atmospheric DMS concentration of $118\pm54$ pptv (1 sigma), DMS sea-to-air flux of $2.9\pm2.1$ μmoles m$^{-2}$ day$^{-1}$ by eddy covariance, and seawater DMS concentration of $1.6\pm0.7$ nM. Dividing flux by the concurrent air-sea concentration difference yields the transfer velocity of DMS ($k_{DMS}$). Compared to previous open ocean measurements in the equatorial East Pacific, Sargasso Sea, Northeast Atlantic, and Southeast Pacific, $k_{DMS}$ in the Southern Ocean was lower mostly because of the cold surface water. Furthermore, we found that the Schmidt number normalization for waterside diffusivity does not account for all temperature effects in $k_{DMS}$. Higher solubility of DMS at a lower temperature results in greater airside control and less transfer of the gas by bubbles formed from breaking waves. We present here a normalization of $k_{DMS}$ for the temperature dependence in solubility, in addition to diffusivity. The final normalized DMS transfer velocity shows good agreement among
tropical and high latitude cruises, and is similar to $k$ of less soluble gases such as carbon
dioxide in low-to-moderate winds but significantly lower in high winds due to the
reduced bubble-mediated transfer of DMS.

1. Introduction

Transfer of gases across the air-water interface has profound implications for the
carbon and sulfur budgets in the marine atmosphere and biosphere. The sea-to-air
transport of dimethylsulfide (DMS) is of particular interest because the gas is thought to
influence climate through the production of sulfate aerosols that could serve as cloud
condensation nuclei (Charlson et al., 1987). DMS is derived from phytoplankton in the
surface seawater ($\text{DMS}_w$) and typically has a concentration of a few nM. Because of
photochemical loss and dilution, the atmospheric DMS concentration ($\text{DMS}_a$) in the
boundary layer is orders of magnitude lower than the Henry’s Law equilibrium
concentration. As a result, the flux of DMS is always positive (upward from the ocean
surface) and is the largest natural source of reduced sulfur to the marine atmosphere
(Lovelock et al., 1972). The substantial magnitude of the sea-to-air flux and the absence
of other sources make it relatively easy to quantify the DMS transfer velocity ($k_{\text{DMS}}$):

$$k_{\text{DMS}} = \frac{\text{Flux}}{(\text{DMS}_w - \alpha \cdot \text{DMS}_a)}$$ (2.1)

Here $\alpha$ is the dimensionless Ostwald solubility of the gas (adopted from Dacey et al.
1984) for DMS. The term inside of the parenthesis is the air-sea concentration
difference that is the driving force for exchange; $\alpha \cdot \text{DMS}_a$ represents the waterside DMS
concentration that would be in equilibrium with the bulk airside concentration.

Air-sea gas transfer is controlled by molecular and turbulent diffusion on the
waterside as well as airside. Close to the interface, turbulent transport diminishes and molecular diffusion dominates. With resistance on both sides of the boundary acting in series, the total transfer velocity of a gas \((k)\) is a function of waterside transfer velocity \((k_w)\) and airside transfer velocity \((k_a)\):

\[
k = \left[ \frac{1}{k_w} + \frac{\alpha}{k_a} \right]^{-1}
\]

For exchange of sparingly soluble gases, including carbon dioxide \((\text{CO}_2)\), sulfur hexafluoride \((\text{SF}_6)\), and helium \((\text{He})\), the molecular sub-layer on the waterside provides the greatest resistance. In contrast, exchange of highly soluble gases, such as methanol, is controlled by airside resistance, with water vapor representing complete airside control. DMS is intermediate in solubility, with \(\alpha = 14.2\) at 20 °C in seawater.

Since \(k\) scales inversely to resistance, turbulence generated from wind stress \((\tau)\) that thins the molecular sub-layers speeds up transfer. Because the 10-meter wind speed \((U_{10})\) is much easier to measure than \(\tau\), most gas exchange parameterizations are based on \(U_{10}\) only (Liss and Merlivat, 1986; Wanninkhof, 1992; Nightingale et al., 2000; Ho et al., 2006). However, these models that are derived from waterside controlled gases diverge significantly in high winds. From a more detailed consideration of the physics, Csanady (1990) theorized that \(k\) has a linear dependence on the friction velocity \((u*)\), which is related to stress through the density of air \((\rho_a)\):

\[
\tau = \rho_a u_*^2.
\]

Our observations of \(k_{\text{DMS}}\) and \(u*\) in the Northeast Atlantic confirmed this linearity at moderate wind speeds (Huebert et al., 2010). However, this linear relationship might not hold in calm or heavy breaking wave conditions, when additional physical processes confound this simple picture of shear-driven direct exchange. When \(U_{10} < \sim 2 \text{ m s}^{-1}\), buoyancy-driven free convection at the ocean surface may account for more transfer than wind shear (Soloviev and Schlüssel,
Above ~7 m s\(^{-1}\), whitecaps start to form on the sea surface. The additional surface area from air bubbles (Woolf, 1997) and the associated turbulent plume (Monahan and Spillane, 1984) can increase gas exchange. Insoluble gases partition readily into bubbles and should therefore show enhanced transfer in moderate-to-high winds. In contrast, a more soluble gas like DMS exhibits only modest bubble-mediated enhancement for wind speeds up to ~10 m s\(^{-1}\) (Blomquist et al., 2006).

In the past several years, we have measured the sea-to-air flux of DMS by eddy covariance (EC) on five cruises in different oceanic environments (Fig. 2.1). Chronologically, they are 2003 Tropical Atmosphere Ocean (TAO) on the NOAA Ship *Ronald H. Brown*, 2004 Biocomplexity (hereinafter BIO) on the R/V *Steward Johnson*, 2007 Deep Ocean Gas Exchange Experiment (DOGEE) on the UK ship RRS *Discovery*, 2008 Southern Ocean Gas Exchange Experiment (SO GasEx) on *Ronald H. Brown*, and 2008 Vamos Ocean-Cloud-Atmosphere-Land Study Regional Experiment (VOCALS-REx, hereinafter VOCALS) on *Ronald H. Brown*. The location, time, DMS sea-to-air flux, seawater DMS concentration, and reference for each cruise are listed in Table 2.1.

While the environmental conditions of each cruise individually were not highly variable, together these five cruises encompass a large range in wind speed (1 ~ 21 m s\(^{-1}\)), sea surface temperature (SST, 2~30°C), and atmospheric boundary layer stability (statically unstable to stable), which is shown as histograms in Fig. 2.2. The higher winds, lower SST, and frequent occurrences of stable boundary layer set SO GasEx apart from other cruises. A compilation of \(k_{DMS}\) from these projects allows us to assess our current understanding of gas exchange in an effort to improve existing models.
2. Experimental

2.1 Background on SO GasEx

The Southern Ocean is characterized by sustained periods of high winds, low SST, and large seasonal cycles in biological productivity. Gas exchange between the ocean and atmosphere in the Southern Ocean plays an important role in the global climate through, for example, the sequestration of atmospheric CO₂ via production of polar bottom waters and through the role of DMS emissions in generation of atmospheric sulfate aerosols. SO GasEx was conceived as the third in a series of projects combining direct measurements of gas exchange with concurrent studies of related biological and physical processes, focused on the unique high wind conditions of the Southern Ocean (Ho et al., this issue a). SO GasEx included a ³He/SF₆ tracer patch study (Ho et al., this issue b), EC observations of CO₂ (Edson et al., this issue), and DMS (this paper). The measurements of atmospheric DMS concentration using an atmospheric pressure ionization mass spectrometer (APIMS) and DMS flux by EC are detailed recently by (Blomquist et al., 2010) and will not be described further here.

2.2 Measurements of Seawater DMS Concentration

Seawater DMS (DMSₗ) concentration near the surface was measured by a gas chromatograph (GC). For SO GasEx, DMSₗ was measured at ~10 minute intervals using a semi-continuous analytical system linked to an equilibration device. Seawater from the ship’s clean water supply at ~5 m depth was passed through a membrane equilibrator (Liqui-Cel®, Extra-Flow 2.5 x 8) at ~400 mL min⁻¹. Nitrogen was passed through the equilibrator at a continuously monitored flow rate of 40 mL min⁻¹ and pressure of 150
kPa. DMS was trapped on an adsorbent (Carbopack-X, Sigma-Aldrich®) at -50 ºC and measured on a Varian 3800 GC equipped with a pulsed flame photometric detector (PFPD). The system was calibrated and sensitivity monitored hourly using a permeation device (Dynacal®, Vici Metronics Inc.) delivering 120 ng DMS min⁻¹ at 30 ºC. In DOGEE, DMSₗ was sampled every 7~10 min from a towed Fish at 1~2 m below the surface and quantified using a purge-and-cryogenic trap linked to a Varian 3800 GC with PFPD. A similar purge-and-trap method was used on the TAO and VOCALS projects, but at a lower frequency of every 15~30 minutes (Bates et al., 2000). The BIO cruise was a Lagrangian water mass study following a cyclonic and an anticyclonic eddy. DMSₗ was determined from the shallowest bin (~5 m) of CTD casts (Kiene and Service, 1991) and showed little variation as a result of the Lagrangian design. Limited inter-comparisons between samples manually collected at the surface and from the ship’s clean seawater supply during DOGEE and SO GasEx suggest that there was no significant, systematic gradient in DMSₗ in the top 5 m of the water column in moderate winds. Nor was there a discernible difference in concentration caused by the different means of water sampling.

2.3 Diffusivity Normalization on kₐ₃₈

The waterside diffusivity dependence of gas transfer can be described by the Schmidt number ($S_c$) as the ratio between kinematic viscosity ($\nu$) of seawater and molecular diffusivity ($D$), both functions of temperature and (to a lesser degree) salinity (Saltzman et al., 1993). To remove variance due to diffusivity, it is common to normalize transfer velocity measured at ambient conditions to a reference $S_c$ of 660.
\[ k_{660} = k_{DMS} \left( \frac{660}{S_c} \right)^n \]  \hspace{1cm} (2.3)

The exponent \( n \) is thought to vary from -2/3 for a rigid surface to -1/2 for a rough surface; we assume the latter for the open ocean. A Schmidt number of 660 corresponds to 20 °C for CO\(_2\) and 27.2 °C for DMS in seawater. A relationship similar to (2.3) is also frequently used to account for diffusivity differences among gases, as in the derivation of \( k_{660} \) for a specific gas of interest from \( k_{660} \) of \(^3\)He/SF\(_6\) obtained from deliberate tracer studies. The analysis below suggests this may lead to bias at moderate-to-high wind speeds when the gas of interest is much more soluble than that of the tracer gas.

Figure 2.3 shows time series of atmospheric and seawater DMS concentrations, DMS flux, 10-m wind speed, transfer velocity at ambient condition and normalized to \( S_c \) of 660, as well as the DMS Schmidt number from SO GasEx. Elevated DMS flux clearly corresponded to higher DMS\(_w\) at times, such as on March 14 and 24, but not always because of variable winds. Towards the end of the experiment, the ship encountered a storm while in transit to Uruguay. Wind speed briefly exceeded 20 m s\(^{-1}\) and the surface water was much warmer (~14°C) during this period than what was typical of SO GasEx (~5°C). However, DMS flux and transfer velocity were not elevated, while DMS\(_w\) was similar to earlier periods of the project.

3. Data Refinement and Normalization

3.1 Basic Data Filtering

Uncertainty in \( k_{660} \) hinges principally on the precision and sampling variability of the flux and DMS\(_w\). There is substantial noise in EC flux observations that is independent of the measurement system. This scatter originates from sampling a
stochastic process (turbulence) over relatively short time scales where low frequency contributions to flux are poorly sampled. Even for a well-resolved scalar like water vapor, the sampling uncertainty of an hourly flux is on the order of 25% (Fairall et al., 2003). At a wind speed of ~8 m s\(^{-1}\) and in a near-neutral boundary layer, Blomquist et al. (2010) showed that relative uncertainty in hourly DMS flux is on the order of ~30%. Uncertainty further increases when the boundary layer is stable, which is discussed in the next section. For a large number of hourly observations, however, precision in the mean flux estimate improves significantly. Here, we focus on eliminating conditions that might lead to nonrandom biases in the measurement.

From the lag correlation between DMS\(_a\) and vertical wind velocity (\(w\)), we find a clear peak in correlation and a corresponding lag time of ~1.5 seconds (depending on the inlet length and flow rate) when the wind is from the bow sector. When the wind is coming from the stern quadrant, however, DMS\(_a\) and \(w\) show poor correlation due to severe airflow distortion by the ship’s superstructure. For SO GasEx, three sonic anemometers were mounted on the foremast of the *Ronald H. Brown*, with one in the middle and two others ~1 m to the port and starboard sides. The DMS sampling inlet was located near the base of the middle anemometer. We limit relative wind direction to ±60° from the bow for the middle anemometer and -120° to 0° and 0° to 120° for the port and starboard anemometers, respectively. DMS fluxes are computed at 10-minute intervals from all three anemometers and averaged when two or more anemometers are in-sector. On all other cruises with a single anemometer, a limit in relative wind direction of ±60° from the bow is applied. Airflow distortions and complex vibrations can also occur when the ship is turning rapidly. Thus 10-minute flux segments when the range in gyro
heading exceeds 30 degrees are excluded from hourly averages.

In Fig. 2.4, hourly $k_{660}$ satisfying the above criteria from all five cruises are plotted vs. $U_{10}$, including the brief high wind speed period (>20 m s$^{-1}$) from SO GasEx. During this storm encounter, the sea was characterized by fully developed long swells and the measured $k_{DMS}$ was much lower than the trend extrapolated from intermediate winds. To date, we have not identified a measurement error responsible for the low $k_{DMS}$. There may be an environmental factor involved, but pending a more detailed analysis of this event, we will not consider these points further in this section.

To more clearly examine the trend in $k_{660}$ as a function of wind speed, we plot $k_{660}$ from the five cruises averaged to $U_{10}$ bins in Fig. 2.5. While overall $k_{660}$ increases with $U_{10}$, $k_{660}$ from tropical cruises such as TAO appears to be higher than that from the high latitude SO GasEx, particularly in high winds. A similar SST trend is observed by Marandino et al. (2009) in their synthesis of $k_{DMS}$ measurements. Also included in Fig. 2.5 is the $k_{DMS}$ curve computed from the NOAA COARE gas transfer model (discussed in detail below) using parameters $A=1.3$ and $B=1.0$ at 27.2 °C.

Another source of scatter in $k_{660}$ is the variability in discretely sampled DMS$_w$. When DMS$_w$ shows high temporal or spatial variability, uncertainty in the hourly mean waterside concentration increases; the location of DMS$_w$ measurement is also less likely to correspond to the flux footprint. For a measurement height of 18 m and at typical wind speeds, the flux footprint covers a region several hundred meters upwind of the ship under neutral or unstable conditions; the footprint can be much larger under stable conditions (Horst and Wiell, 1994). We address the variability in DMS$_w$ by setting a threshold of 0.25 for the relative standard error of the mean in the hourly DMS$_w$ for all
cruises except BIO, following Huebert et al. (2010).

3.2 Effect of Atmospheric Stability

The wind speed profile as a function of height depends on boundary layer stability. At a reference height of 10 meters, wind speed is related to wind stress through $u^*$ and the 10-m drag coefficient ($C_{D10}$): $u^* = C_{D10}^{1/2} U_{10}$. Adjusted to neutral condition, the relationship becomes $u^* = C_{D10n}^{1/2} U_{10n}$, where $C_{D10n}$ and $U_{10n}$ are stability-corrected. Logic follows that some of the scatter in the relationship between $k$ and $U_{10}$ is due to the effect of stability on the logarithmic wind speed profile, and may be reduced by using $U_{10n}$. We estimate $u^*$, $U_{10n}$, $z_0$ (roughness length), and $L$ (Monin-Obukhov length scale) from bulk meteorological variables using the NOAA COARE 3.0 bulk flux model (Fairall et al., 2003).

The stability of the atmospheric surface layer can be represented by the ratio $z/L$, with $z$ being the sensor height. The ratio $z/L$ depends on the air-sea temperature difference, and is a proxy for the relative contribution to turbulence generated by buoyancy and shear. A significantly positive $z/L$ implies a statically stable boundary layer, usually caused by warm air over cold water. A negative $z/L$ indicates statically unstable, as is typical of temperate and tropical regions. Under a near neutral condition, $z/L \approx 0$. Figure 2.6 shows DMS $k_{660}$ vs. $U_{10}$ and $U_{10n}$ for SO GasEx. The degree of scatter in $k_{660}$ is reduced when $U_{10n}$ is used, with the difference most noticeable in low-to-moderate winds where buoyancy-driven convection is more important. The coefficient of determination ($r^2$) for a linear relationship between $k_{660}$ and wind speed is improved from 0.51 to 0.56 with $U_{10n}$ instead of $U_{10}$.
Besides modifying the wind speed profile, static stability in the boundary layer increases uncertainty in the flux measurement (Blomquist et al., 2010). Stable conditions lead to suppression of turbulent eddies and possibly a shallow boundary layer. Strong, warm winds over a cooler ocean surface are often found in warm sectors of frontal passages. While damping of turbulence in these conditions may not be very significant due to the high surface shear, the boundary layer can become quite shallow. With all else being constant, a shallower boundary layer implies a steeper flux gradient, which results in a greater systematic underestimate of the true surface flux, as illustrated in Fig. 2.6 by the solid circles \((z/L > 0.05)\). Hourly flux observations when \(z/L > 0.05\) are hereafter omitted. This stability threshold is intended to remove conditions unfavorable for EC measurement and does not imply a cessation of turbulence. This filter removes about 30\% of our observations from high latitude cruises (SO GasEx and DOGEE), while data from the tropical and temperate waters are largely unaffected. About 1100 hourly \(k_{DMS}\) observations from all cruises remain following screening for \(DMS_w\) and stability.

### 3.3 Flux Corrections at High and Low Frequencies

Some flux signal is lost at the lowest frequencies at 10-minute time intervals, and of the lowest frequencies that are measured, the sampling statistics are poor. Additional flux signal is lost at the highest frequencies due to an attenuation of small-scale fluctuations by the inlet system and the Nafion® air drier, which is needed to remove water from the ambient air and increase the sensitivity of the APIMS towards DMS. We partially correct hourly cospectra for high frequency inlet-attenuation based on an empirical frequency response function (a correction of a few percent, Blomquist et al.,
An estimate for additional missing flux is obtained through comparison with a theoretical cospectral function for neutral conditions (Kaimal et al., 1972). Partially corrected cospectra are normalized to DMS$_w$ and $S_e = 660$ and bin-averaged by relative wind speed ($U_R$). For each $U_R$ bin, the theoretical Kaimal function is fit to the observed data over a restricted frequency range (0.008~1 Hz). Measurement height and relative wind speed parameters in the fit are fixed to mean observed conditions. Within each bin, the difference in area between the fit and the observed cospectrum under the high and low frequency tails (0~0.005 Hz and 1~10 Hz) is an estimate of the missing flux. A cospectral correction factor ($F_C$) is computed as the ratio between the total corrected flux and the observed flux for each bin and fit to a polynomial as a function of $U_R$:

$$F_C = 1.165 - 0.020U_R + 0.001U_R^2$$

for SO GasEx. The estimated correction to the flux is less than 5% for $U_R$ of 8~16 m s$^{-1}$, and on the order 10% for $U_R$ less than 5 or greater than 20 m s$^{-1}$. The magnitude of the correction varies somewhat among cruises because of variable sampling height, inlet length, flow rates, etc.

### 3.4 Airside Resistance

Airside resistance to exchange of sparingly soluble tracer gases is small and usually neglected. However, as the solubility ($\alpha$) of a gas increases with decreasing temperature, the airside resistance becomes more important. Rearranging (2.2), McGillis et al. (2000) defined the atmospheric gradient fraction $\gamma_a$ as the fraction airside contribution to total concentration difference: $\gamma_a = \frac{1}{1 + k_a(\alpha k_w)^{-1}}$. For DMS, $\gamma_a$ is on the order of 0.05 at 20 °C, compared to only ~0.002 for the less soluble CO$_2$. Thus, the temperature dependence in airside resistance, which is not accounted for in (2.3), is more
significant for DMS than for CO$_2$. $k_{DMS}$ obtained from (2.1) represents the total transfer velocity. To account for all temperature effects when normalizing to $S_c = 660$, the airside and waterside transfer velocity need to be specified and adjusted separately. Following McGillis et al. (2000), we estimate $k_a$ of DMS as $659U_{10n}(MW_{DMS}/MW_{H_2O})^{-1/2}$ (cm hr$^{-1}$), with $MW$ representing molecular weight. To approximate $k_a$ a priori, we use the formulation from the NOAA COARE gas transfer model (cm hr$^{-1}$):

$$k_a = 360000u_s(\rho_w/\rho_a)^{-1/2}[h_wS_c^{1/2} + \kappa^{-1}\ln(0.5/\delta_w)]^{-1} + B(2450f_{wh}G)$$ (2.4)

Here $\rho_w$ is the density of water, $\delta_w$ the waterside molecular sublayer thickness, $\kappa$ the von Karman’s constant (0.4), and $h_w = 13.3/(A\phi)$. In $h_w$, $A$ is an empirical constant and tuned to 1.3 by Blomquist et al. (2006) using tropical EC measurements of DMS; $\varphi$ accounts for surface buoyancy flux enhancement of the transfer and only becomes important in wind speed less than ~2 m s$^{-1}$. The second term on the RHS of (2.4) is the parameterization of $k_b$ from Woolf (1997), where the whitecap fraction $f_{wh} = 3.84 \times 10^{-6} U_{10}^{3.41}$ (Monahan and O’Muircheartaigh, 1980), and $G = \alpha^{-1}[1 + (14\alpha S_c^{-1/2})^{-1/1.2}]^{-1.2}$. $B$ is an empirical constant, to which we assign the provisional value of 1.0 (Blomquist et al., 2006).

At typical SO GasEx temperatures (~5 °C), $\gamma_a$ for DMS is about 0.10 at a wind speed of 15 m s$^{-1}$. The airside effect calculated here is smaller than was estimated by McGillis et al. (2000) for DMS because they used the parameterization from Wanninkhof (1992) for $k_w$, which significantly overestimates $k_{DMS}$ at high wind speeds. We can now estimate the ambient waterside transfer velocity of DMS from measured $k_{DMS}$:

$$k_w = k_{DMS}/(1 - \gamma_a)$$

Applying the Schmidt number normalization to $k_w$ yields $k_{w660}$. For consistency, the airside transfer should also be adjusted to 27.2 °C ($S_c = 660$ for DMS),
such that the total normalized transfer velocity becomes:

\[ k_{660} = \left[ \frac{1}{k_{w660}} + \frac{\alpha_{660}}{k_a} \right]^{-1} \]  

(2.5)

Here \( \alpha_{660} \) represents the solubility of DMS at 27.2 °C (10.4). Compared to the \( S_c \)-only normalization specified by (2.3), separate treatment of the temperature dependence in waterside and airside transfer given by (2.5) increases \( k_{660} \) from SO GasEx by ~4%.

After screening for DMS\(_w\) variability as well as atmospheric stability, and accounting for temperature dependence in airside resistance, \( r^2 \) for a linear relationship between \( k_{660} \) from (2.5) and \( U_{10n} \) for all cruises is increased from 0.63 in Fig. 2.4 to 0.71 in Fig. 2.7. However, discrepancies in \( k_{660} \) remain among different cruises, particularly in higher winds, which might be in part related to the temperature dependence in bubble-mediated transfer.

### 4. Bubble-mediated Exchange and Solubility Normalization

When waves break, air is trapped in water and entrained to depth of a few meters in the form of bubbles, which then rise and exchange gas with the surrounding water. The amount of a trace gas partitioning into air bubbles from the bulk water likely depends on both solubility and diffusivity. At lower temperatures, \( \alpha \) increases and less gas is transferred from the bulk water to bubbles. The bubble component of \( k_{DMS} \) should be normalized to a reference temperature separately from the interfacial component, which does not depend on \( \alpha \). It is convenient to first estimate the interfacial component of gas exchange due to shear \( (k_v) \), ignoring buoyancy-driven exchange that is insignificant in moderate-to-high winds. An estimate for bubble-mediated transfer velocity \( (k_b) \) is then simply the difference between \( k_w \) and \( k_v \):
\[ k_b = k_w - k_v = k_w - k_{v660}(660/S_c)^{1/2} = (k_{w660} - k_{v660})(660/S_c)^{1/2} \]  
(2.6)

Here \( k_{v660} \) is interfacial transfer velocity at \( S_c = 660 \). We estimate \( k_{v660} \) as a linear function of the tangential component of the friction velocity \( (u_v) \) due to viscous wind shear (i.e. \( k_{v660} = C_1 + C_2u_v \), as described below). To account for temperature effects, we normalize \( k_b \) to \( S_c = 660 \) using the \( S_c \) and \( \alpha \) dependence \( (G) \) described by the Woolf (1997) model for bubble-mediated exchange. Adding the interfacial component yields the solubility and diffusivity normalized waterside transfer velocity:

\[ k_{w660}' = k_{v660} + k_b(G_{660}/G), \]  
where \( G_{660} \) is the Schmidt number-solubility dependence at 27.2 °C. We may then substitute \( k_{w660}' \) in place of \( k_{w660} \) in (2.5) to yield the final normalized total transfer velocity \( (k_{660}') \). While the procedure described above is simple, the justification and background require more detailed explanations, which will be covered in the remainder of this section.

4.1 Quantifying the Interfacial Transfer Velocity

From the two-layer model described by Liss and Slater (1974), physical processes that thin the diffusive sub-layers are conceptualized to enhance direct (interfacial) gas exchange through a reduction in resistance. Shear (viscous) stress from wind blowing tangentially to the sea surface is usually the most important of such processes, as it leads to micro-scale breaking of capillary waves and wavelets that are largely responsible for interfacial gas exchange (Frew et al., 2004). Surface renewal models from Csanady (1990) and Soloviev and Schlüssel (1994) predict that without bubbles, \( k \) should be linearly related to \( u_v \), consistent with our \( k_{DMS} \) observations from DOGEE in moderate winds (Huebert et al., 2010).
In high winds \((U_{10} > \sim 10 \text{ m s}^{-1})\) and mature seas, the ocean surface transitions from micro-scale waves to the breaking of long gravity waves. As total surface stress grows with wind speed, an increasingly larger fraction is partitioned to wave stress, which includes momentum transfer into the ocean by breaking waves and form drag. While wave breaking can lead to more gas exchange via enhanced turbulence and bubble plume formation, increasing partition of total stress to form drag, which is a result of the pressure differential developed between the front and lee sides of large waves, may suppress micro-scale breaking (Banner et al., 1989) and potentially reduce the wind speed dependence of \(k\). The net effect of these competing mechanisms on \(k\) in high winds partly depends on gas solubility. Since wave stress does not significantly contribute to thinning of the diffusive sub-layer, a linear relationship between tangential friction velocity \((u^*)\) and interfacial transfer velocity \((k_v)\) should be more general than that between \(u^*\) and \(k\) and hold true even in high winds, concurrent with the additional decrease in transfer resistance due to bubbles and whitecaps.

We determine \(u^*\) from the tangential component of wind stress. The COARE bulk flux algorithm follows Smith (1988) and partitions total roughness length as:

\[
  z_o = q u_s^2 / g + 0.11v / u_s
\]

(2.7)

Here \(q\) is a fit to drag coefficient observations and \(g\) is gravity. The first term on the RHS of (2.7) is due to waves, and the second due to smooth flow. From laboratory wind-wave studies, Banner and Peirson (1998) found that total stress in the smooth flow regime represents the upper limit for the tangential stress. Based on this result, Mueller and Veron (2009) (hereinafter MV09) estimate the 10-m tangential drag coefficient in neutral condition \((C_{D10_s})\) as a function of the smooth flow roughness length:
\[ C_{D_{10v}}^{1/2} = \kappa / \ln(10 / (0.11v / u_{*v})) \]  

Thus at a given wind speed, \( u_{*v} \), can be computed iteratively: \( u_{*v} = C_{D_{10v}}^{1/2}U_{10n} \). The result of this partition of the total wind stress to the tangential and wave component is qualitatively similar to the alternate estimate by Soloviev and Schlüssel (1996) (hereinafter SS96), as shown in Fig. 2.8. SS96 use the Keulegan number defined by Csanady (1978) as the criterion for the transition to large-scale wave breaking:

\[ Ke = u_{*w}^3 / g \nu, \] where \( u_{*w} \) is the waterside transfer velocity. Tangential stress is then estimated as a fraction of total stress: \( (1 + Ke / Ke_{cr})^{-1} \), where \( Ke_{cr} \) is the critical \( Ke \), an empirically derived threshold dependent on wave age and speculatively set to 0.45 here for illustration. Tangential stress from SS96 varies with wave age, whereas \( u_{*v} \) from MV09 does not (only the wave component does). We adopt the approach from MV09 because \( Ke \) is not required, as tangential stress is approximately independent of the specification of total drag coefficient (i.e. \( u^* \) is not required to estimate \( u_{*v} \)).

Because of the increasing importance of wave stress with wind speed, \( C_{D_{10v}} \) decreases in high winds and \( u_{*v} \) shows a slight downward curvature, with estimates from SS96 rolling off more. If \( k_v \) follows \( u_{*v} \) in high winds, it too should roll off slightly, which might not be apparent in observed total transfer of a sparingly soluble gas (e.g. \( CO_2 \)) in high winds because the additional bubble-mediated exchange term overwhelms any suppression in interfacial exchange. However, for a soluble gas like DMS with a more modest bubble enhancement, the difference between \( k_v \) and \( k \) should be smaller. Indeed, if we plot observed \( k_{DMS} \) vs. \( u_{*v} \), the trend would be best described as linear in moderate winds with a slight leveling off in high winds, qualitatively similar to the “wave attenuation” effect observed for DMS in the laboratory by Rhee et al. (2007). This roll
off suggests that the behavior of $k_{DMS}$ in high winds is related to reduced partition of total stress to the tangential component.

Let us now examine how $k_{w660}$ relates to $u^*$ in different wind speed regimes for all cruises (Fig. 2.9). In the calmest conditions ($u^* < \sim 0.1 \text{ m s}^{-1}$, corresponding to a wind speed of 2–3 m s$^{-1}$), the slope between $k_{w660}$ and $u^*$ appears to be relatively flat, likely because buoyancy-driven convection is more important than wind shear at driving gas exchange. The slope steepens with increasing wind speeds; a linear fit from 2–6 m s$^{-1}$ yields an intercept ($C_1$) of -3.8 and a slope ($C_2$) of 69.7. We interpret the extrapolation of this fit to higher winds as interfacial transfer velocity ($k_{v660}$), which becomes less than the bin-average $k_{w660}$ above $u^* \approx 0.2 \text{ m s}^{-1}$ (a wind speed of 7–8 m s$^{-1}$, or the onset of whitecap formation). Subtracting $k_{v660}$ from $k_{w660}$ in moderate-to-high winds and removing the prior $S_c$ normalization yield an estimate of $k_b$ for DMS.

### 4.2 Solubility Normalization of Bubble-mediated Exchange

Normalizing $k_b$ to a reference temperature requires a model describing the $S_c$ and $\alpha$ dependence in bubbles. Scaling up from a single bubble model, Woolf (1997) parameterizes $k_b$ of CO$_2$ to be proportional to whitecap fraction ($f_{wh}$) and $G$. A linear relationship between $k_b$ and $f_{wh}$ was proposed by Monahan and Spillane (1984) and confirmed by laboratory results of Asher et al. (1996) with optically measured bubble plume coverage as the analog for $f_{wh}$. The term $G = \alpha^{-1}[1 + (14\alpha S_c^{-1/2})^{-1/1.2}]^{-1.2}$ contains the solubility and diffusivity dependence in $k_b$. Woolf (1997) identifies two asymptotic behaviors of $k_b$ at opposing limits of gas solubility. For an insoluble gas, $G$ approximately scales as $S_c^{-1/2}$. This implies the temperature dependence in solubility has
little effect on $k_b$ for gases like SF₆ and CO₂, such that the usual normalization to $S_c = 660$ is sufficient. For a very soluble gas, $G$ approaches $\alpha^{-1}$ and the $S_c$ dependence vanishes. For DMS at 5 °C, $\alpha = 28.3$ and $S_c = 2050$; the functional form of $G$ is close to $\alpha^{-1}$, with a weak $S_c$ dependence. Multiplying $k_b$ by the factor $(G_{660}/G)$ yields the normalized bubble-mediated transfer velocity, which is then summed with the prior estimate of $k_{v,660}$ to yield the final normalized waterside transfer velocity ($k_{w,660}'$). For SO GasEx, the normalization to 27.2 °C increases $k_b$ by ~150%, or ~40% relative to $k_b$ adjusted using the $S_c$-only dependence. Together with interfacial exchange, $k_{w,660}'$ is ~6% greater than $k_{w,660}$ in high winds. While the effect of this normalization appears to be small and only secondary in importance, for a larger $k_b$ (e.g. if $k_v$ rolls off more in high winds, as in SS96), the adjustment in $k_b$ would be correspondingly greater as well.

Keeling (1993) and Asher et al. (1996) describe different dependences on $S_c$ and $\alpha$ in bubble-mediated exchange than Woolf (1997). Keeling (1993) models $k_b$ based on bubble spectra photographically recorded in laboratory experiments designed to simulate wave breaking (Monahan and Zeitlow, 1969; Cipriano and Blanchard, 1981), with a range in bubble radius from ~0.03 mm to 4 mm. The author suggests that relatively large bubbles (>0.5 mm in radius) contribute significantly to $k_b$, which should scale roughly as $\alpha^{-0.3} S_c^{-0.35}$. Asher et al. (1996) measured the exchange of multiple gases in a tipping-bucket whitecap simulation tank. Following Memery and Merlivat (1985), Asher et al. (1996) separate $k_b$ to contributions from bubbles that dissolve in or equilibrate with the surrounding water before reaching the surface and ones that never equilibrate: $k_b = a\alpha^{-1} + b\alpha^{-0.37} S_c^{-0.18}$. Theoretically, small bubbles with long lifetimes tend to be dissolved or equilibrated; $k_b$ due to these bubbles should scale as $\alpha^{-1}$ (not limited by
diffusivity). On the other hand, large bubbles that rise rapidly have insufficient time to equilibrate with the bulk water before surfacing. Gas is exchanged between the bulk water and these large bubbles during their entire lifetime, implying a dependence on both $\alpha$ and $S_c$. For evasion and through a cleaned surface, Asher et al. (1996) estimate that gas exchange due to dissolving and equilibrating bubbles is orders of magnitude smaller than that due to non-equilibrating bubbles ($a = -1 \times 10^{-4}$ and $b = 1.7 \times 10^{-2}$, in units of m s$^{-1}$), suggesting that the overall functionality of $k_b$ for evasion is approximately $\alpha^{-0.37} S_c^{-0.18}$.

If we assume a weaker solubility dependence in $k_b$ from Keeling (1993) or Asher et al. (1996), the temperature effect in $k_b$ will be smaller than what we have shown. Over the SST range of 5~27 °C, the formulation $\alpha^{-0.37} S_c^{-0.18}$ for DMS differs by a factor of ~1.8, whereas the Woolf (1997) formulation differs by a factor of ~2.5. Coincidentally, $S_c^{-0.5}$ for DMS over the same SST range also differs by a factor of ~1.8. Thus, using the formulation from Asher et al. (1996) to adjust $k_b$ leads to the essentially the same result as using the widely used $S_c$-only normalization.

The error caused by ignoring the temperature-solubility dependence in $k_b$ when normalizing transfer velocity to $S_c = 660$ is most significant for gases of intermediate solubility. Using the COARE model, we can calculate the difference between $k_b (G_{660}/G)$ and $k_b (660/S_c)^{1/2}$ for a normalization from 5 °C to the temperature at which $S_c = 660$ for a range of gases, and compute the relative error by dividing that difference by $k_{660}$. This is shown in Fig. 2.10 for DMS, CO$_2$, methyl iodide (CH$_3$I), and chloroiodomethane (CH$_2$ClI). In seawater, $S_c = 660$ at 24.0 and 27.8 °C for CH$_3$I, and CH$_2$ClI; the respective $\alpha$ values are 3.6 and 18.8 (Archer et al., 2007). It is apparent that for very insoluble gases, the relative error in transfer velocity in high winds is near zero because the
temperature dependence of $G$ is similar to $S_c^{-1/2}$. Such is the case for CO$_2$ (Fig. 2.10), as well as SF$_6$, radon, and nitrous oxide (not shown). For soluble gases, the temperature dependence of $G$ diverges more from $S_c^{-1/2}$, but the magnitude of $k_b$ also decreases, so the error in total transfer velocity is negligible. It is transfer of gases with intermediate solubility, such as organosulfur compounds, that are affected the most by the temperature dependence in solubility. For example, applying the $S_c$ normalization to CH$_3$I at SST = 5 °C underestimates $k_{660}$ by ~11% at a wind speed of 16 m s$^{-1}$.

5. Discussion

Replacing $k_{w660}$ in (2.5) with the $k_{w660}'$ yields the final normalized total transfer velocity ($k_{660}'$), which is plotted in $U_{10n}$ bins for all five cruises in Fig. 2.11. Compared to Fig. 2.5, $k_{660}'$ is greater than $k_{660}$ from (2.3) for the coldwater SO GasEx by ~7 cm hr$^{-1}$ at 15 m s$^{-1}$, whereas for the tropical TAO and BIO cruises, $k_{660}'$ is slightly lower than $k_{660}$ at most wind speeds. To more clearly demonstrate the effects of filtering for DMS$_w$ variability and atmospheric stability, normalizing for temperature dependence in airside resistance, and adjusting for both $\alpha$ and $S_c$ dependence in $k_b$, Fig. 2.12(a) shows the difference between $k_{660}$ from (2.3) and COARE model estimate at 27.2 °C in $U_{10}$ bins; (b) shows the difference between final normalized $k_{660}'$ and COARE estimate in $U_{10n}$ bins. The COARE gas transfer model estimate is used as a reference here for comparison. The divergence at high wind speeds shown in Fig. 2.12(a) illustrates the biases among cruises partly due to the residual temperature dependence not accounted for by (2.3). $k_{660}'$ from all cruises show closer agreement with each other; at 9 m s$^{-1}$, the standard deviation of the bin-averages is 3.6 cm hr$^{-1}$ for $k_{660}$ and is reduced to 2.3 cm hr$^{-1}$.
for \( k_{660}' \). Table 2.2 contains the bin-average \( k_{660} \) (from (2.5), accounting for temperature-solubility dependence in airside resistance, but not in bubbles) and \( k_{660}' \) of the five cruises, weighted by the number of points per bin for each cruise. Average \( k_{DMS} \) from SO GasEx at ambient conditions and corresponding \( u^* \) values in \( U_{10n} \) bins from 1 to 15 m s\(^{-1}\) are also presented.

We can now compare this average \( k_{660}' \) to previous observations of relatively insoluble gases and widely used wind speed-dependent parameterizations (Fig. 2.13). Based on measurements of SF\(_6\) gas exchange in a lake and wind tunnel observations, Liss and Merlivat (1986) model \( k \) as three piece-wise linear functions with increasing slope at higher winds, representing distinctive regimes of smooth surface, rough surface (capillary waves), and breaking waves, respectively. From natural \(^{14}\)C disequilibrium and the bomb \(^{14}\)C inventory, Wanninkhof (1992) fits \( k \) with a quadratic function. From artificial injections of two volatile tracers (\(^{3}\)He and SF\(_6\)) and one non-volatile tracer (spores) in the North Sea, Nightingale et al. (2000) report a parameterization of \( k \) that consists of both linear and quadratic dependences. Ho et al. (2006) fit a quadratic function to \(^{3}\)He/SF\(_6\) measurements in the Southern Ocean. Based on EC observations of CO\(_2\) during the GasEx I cruise in the North Atlantic (a CO\(_2\) sink), Wanninkhof and McGillis (1999) suggested a cubic fit between \( k \) and wind speed. However, an equivalent study during the follow-up GasEx II cruise in the Equatorial Pacific (a CO\(_2\) source) resulted in a much weaker wind speed dependence (McGillis et al. 2004). The authors attributed the GasEx II result to a limited wind speed range and strong diurnal heating.

Also shown in Fig. 2.13 are the \( k_{660} \) curves from the COARE gas transfer model for DMS (\( A=1.3; B=1.0; \text{SST}=27.2 \ \text{°C} \)) and CO\(_2\) (\( A=1.3; B=1.0; \text{SST}=20.0 \ \text{°C} \)).
Normalized for diffusivity, transfer velocities of waterside controlled gases due to buoyancy and shear are expected to be similar (Asher et al., 1996). Indeed, below 7~8 m s$^{-1}$, before the onset of whitecapping, $k_{660}$ of different waterside controlled gases are comparable. The exception is GasEx II (McGillis et al. 2004), which shows elevated $k_{660}$ in low winds. In higher winds, transfer velocity of DMS is significantly lower than that determined for CO$_2$ and from $^3$He/SF$_6$ due to the much higher solubility of DMS, and hence decreased partitioning into air bubbles. While a power law parameterization of $k$ as a function of wind speed might be adequate in application for one particular gas, it masks the complex physical mechanisms responsible for gas exchange in high winds and rough seas. To more accurately quantify the interfacial and bubble-mediated components of air-sea exchange, simultaneous high wind speed measurements of the transfer velocity of multiple gases with a large range of solubility will be needed.

In general, the COARE gas transfer model appears to predict DMS transfer velocity fairly well at the reference $S_c$. A better assessment of model performance is a comparison between the observed $k_{DMS}$ and modeled $k_{DMS}$ at ambient conditions, as normalization to a reference will introduce additional uncertainty due to assumptions in $S_c$ and $\alpha$. Such uncertainty amplifies when the SST is very different from 27.2 °C, as was the case for SO GasEx. Since temperature is a required input parameter for COARE 3.0 and the gas transfer parameterization for $k_b$ accounts for solubility and diffusivity effects, no adjustment on the model output is necessary for a direct comparison of $k_{DMS}$.

Figure 2.14 shows the hourly residual (observation minus model) in $k_{DMS}$ for all five cruises. The difference between the bin-average observation and model prediction is within ~2 cm hr$^{-1}$ across the wind speed range. In low winds, observation appears to be
less than the COARE prediction, which may be a result of measurement errors or model deficiencies. Relative uncertainty in the EC flux observations increases in low winds due to the lower signal to noise ratio and a shift of turbulent eddies towards lower frequencies that are less adequately sampled. Near surface gradients in DMSw are also more likely in calm conditions, which can lead to biases in DMSw measurements. With regard to the model, COARE uses the oceanic turbulent Richardson number and heat fluxes to determine the buoyancy contribution to gas exchange. Uncertainty in the critical turbulent Richardson number, an empirical constant that defines the threshold from free convection to forced convection, is an additional source of error. Overall, the discrepancy between observation and model at low winds should not greatly bias the predicted global mean flux because of the small magnitude of $k_{DMS}$ in calm conditions.

In moderate winds, observations and COARE prediction agree closely, implying that the empirical constant $A = 1.3$ for direct transfer is reasonable (the bin-average at 9 m s$^{-1}$ shows a small positive bias partly due to the several high $k_{DMS}$ points between 8~9 m s$^{-1}$). In high winds, the model prediction exceeds observation by ~10%, which could be due to an overestimation of bubble-mediated exchange (e.g. empirical constant $B$). Also, the model presently uses $u^*$ in the estimation of direct transfer; as $u^*$, has a flatter curvature than $u_*$, using $u^*_v$ would result in a smaller interfacial term in high winds. However, incorporating $u^*_v$ in to the COARE model requires further tuning of empirical parameters $A$ and $B$. Such fitting is best done, if a universal relationship exists, with multiple gases of varying solubility, which is beyond the scope of this paper.

6. Conclusions
Over the past several years, we have measured the transfer velocity of DMS directly in five open ocean cruises at locations ranging from the tropics to the high latitudes. To limit the influence of measurement bias, we apply a number of quality control criteria based on relative wind direction, ship maneuvers, and atmospheric stability. To account for losses of flux signal at high and low frequencies due to inlet attenuation and limited sampling time, respectively, an approximate correction using the Kaimal cospectral function for neutral conditions is used, yielding a total correction of ~5% typically and ~10% for the highest and lowest relative wind speeds.

SO GasEx is unique for the low SST, high winds, and frequent occurrence of stable atmosphere. Normalized for diffusivity only, DMS transfer velocity from SO GasEx is still lower than those from warm water cruises. The solubility of DMS increases in low SST, which results in greater airside control and reduced bubble-mediated exchange. We demonstrate here adjustments accounting for these temperature effects, which yields improved agreement among DMS transfer velocity observations from five cruises. Compared to gas exchange observations of CO₂ and SF₆, normalized transfer velocity for DMS is similar at low-to-moderate wind speeds, where shear-driven interfacial exchange dominates. In high winds, however, DMS transfer trends significantly lower than transfer of other gases due to the solubility dependence in bubble-mediated exchange. Among widely used gas transfer parameterizations (e.g. Liss and Merlivat (1986), Wanninkhof (1992), Nightingale et al. (2000), Ho et al. (2006)), the physics-based NOAA COARE model shows the closest agreement with field observations of $k_{DMS}$.

In a recent review, Elliott (2009) summarizes gas transfer parameterizations for
DMS flux and examines the global distribution of DMS emissions with a planetary level sulfur cycle model. The author lists the discrepancies between generalized gas transfer parameterizations and recent eddy correlation flux measurements, and adopts a composite DMS transfer model based on formulations from Wanninkhof (1992) and Liss and Merlivat (1986) to account for the lower transfer at higher wind speeds compared to insoluble gases. At the time of the review by Elliot (2009), model validation with direct EC observations of DMS extended only to wind speeds of ~10 m s$^{-1}$ (Blomquist et al., 2006). Furthermore, provision for the computation of friction velocity within the ocean circulation model was not implemented, limiting the gas transfer representation to a simple dependence on wind speed. More recent $k_{DMS}$ observations, especially from SO GasEx, have extended the range of wind speed to 15 m s$^{-1}$. Results from this paper suggest that the dependence on SST in $k_{DMS}$ is more complex than previously assumed. For example, in high latitude regions characterized by low temperatures and high wind speeds, estimated $k_{DMS}$ from wind speed parameterizations will be biased high if only the Schmidt number normalization is used. For these reasons, the implementation of a physics-based gas transfer scheme similar to COARE 3.0 in planetary sulfur models is clearly desirable. In schemes that may already use bulk parameterizations of heat fluxes, the additional overhead to compute friction velocity and gas transfer velocity should be minimal.

The polar seas, especially the Southern Ocean, play a key role in the global sulfur cycle and atmospheric aerosol distribution through emissions of DMS (Gabric et al., 2004). The conclusions from Elliot (2009) with respect to an overestimate of polar DMS emissions by current sulfur cycle models seems justified based on the most recent field
observations of DMS transfer velocity at moderate-to-high wind speeds. However, the influence of wave fields and sea state on surface stress, and therefore on gas exchange, is not well parameterized by models, including COARE 3.0. If the anomalously low $k_{DMS}$ observations in very high winds from SO GasEx (Fig. 2.4) are found to be related to sea state, a further reduction in the net emission of DMS from polar seas may be warranted, but currently we must consider this issue unresolved.

Acknowledgements

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## Tables

### Table 2.1 Summary of cruises

<table>
<thead>
<tr>
<th>Cruise</th>
<th>Location</th>
<th>Time</th>
<th>Flux(^1)</th>
<th>DMS(_w)(^2)</th>
<th>Publication</th>
</tr>
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<tbody>
<tr>
<td>TAO</td>
<td>Equatorial Pacific</td>
<td>Nov, 2003</td>
<td>7.1 (3.7)</td>
<td>2.6 (0.8)</td>
<td>Huebert et al., 2004</td>
</tr>
<tr>
<td>BIO</td>
<td>Sargasso Sea</td>
<td>Jul-Aug, 2004</td>
<td>6.2 (2.4)</td>
<td>2.6 (0.4)</td>
<td>Blomquist et al., 2006</td>
</tr>
<tr>
<td>DOGEE</td>
<td>Northeast Atlantic</td>
<td>Jun-Jul, 2007</td>
<td>5.2 (6.8)</td>
<td>2.2 (2.4)</td>
<td>Huebert et al., 2010</td>
</tr>
<tr>
<td>SO GasEx</td>
<td>Southern Ocean</td>
<td>Mar-Apr, 2008</td>
<td>2.9 (2.1)</td>
<td>1.6 (0.7)</td>
<td>This work</td>
</tr>
<tr>
<td>VOCALS</td>
<td>Southeast Pacific</td>
<td>Oct-Nov, 2008</td>
<td>3.4 (1.9)</td>
<td>2.8 (1.1)</td>
<td>Yang et al., 2009</td>
</tr>
</tbody>
</table>

1. Project mean DMS flux (1 sigma), μmoles m\(^{-2}\) day\(^{-1}\)
2. Project mean seawater DMS concentration (1 sigma), nM

### Table 2.2 Average DMS Transfer Velocity for SO GasEx and All Cruises

<table>
<thead>
<tr>
<th>(U_{10w}), m s(^{-1})</th>
<th>(u^*), m s(^{-1})</th>
<th>(k_{DMS}), cm hr(^{-1})</th>
<th>(k_{660}), cm hr(^{-1})</th>
<th>(k_{660}'), cm hr(^{-1})</th>
</tr>
</thead>
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<tr>
<td>1</td>
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<td>*</td>
<td>1.1 (0.1)</td>
<td>1.1 (0.1)</td>
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<tr>
<td>3</td>
<td>0.11</td>
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<td>2.8 (0.1)</td>
</tr>
<tr>
<td>5</td>
<td>0.17</td>
<td>3.8 (0.4)</td>
<td>6.4 (0.1)</td>
<td>6.4 (0.1)</td>
</tr>
<tr>
<td>7</td>
<td>0.25</td>
<td>4.9 (0.5)</td>
<td>10.7 (0.2)</td>
<td>10.9 (0.2)</td>
</tr>
<tr>
<td>9</td>
<td>0.33</td>
<td>6.6 (0.3)</td>
<td>15.8 (0.4)</td>
<td>16.3 (0.4)</td>
</tr>
<tr>
<td>11</td>
<td>0.42</td>
<td>8.7 (0.3)</td>
<td>19.2 (0.6)</td>
<td>19.8 (0.6)</td>
</tr>
<tr>
<td>13</td>
<td>0.52</td>
<td>10.1 (0.4)</td>
<td>22.8 (0.8)</td>
<td>23.4 (0.8)</td>
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<tr>
<td>15</td>
<td>0.62</td>
<td>11.2 (0.7)</td>
<td>26.2 (1.3)</td>
<td>27.6 (1.4)</td>
</tr>
</tbody>
</table>

\(k_{DMS}\): bin-average from SO GasEx (standard error of the mean) at ambient conditions
\(k_{660}\): bin-average from all cruises (standard error of the mean) from (2.5), without bubble normalization
\(k_{660}'\): bin-average from all cruises (standard error of the mean) with bubble normalization
Figures

Figure 2.1 Map showing the five cruises from which DMS flux was measured directly with eddy covariance by U. Hawaii.
Figure 2.2 Histograms of 10-m neutral wind speed, SST, and stability parameter $z/L$.

Tropical and temperate cruises were characterized by relative low wind speed, high SST, and unstable atmosphere. In SO GasEx, we encountered higher winds, lower SST, and more frequent occurrences of stable atmosphere.
Figure 2.3 SO GasEx time series of atmospheric DMS concentration and flux (top); seawater DMS concentration and $U_{10}$ (middle); transfer velocity at ambient condition, normalized to $S_c = 660$ as described in (2.3), as well as the DMS Schmidt number (bottom). Elevated DMS flux at times corresponded to higher seawater DMS concentration, such as on March 14 and 24. Near the end of the experiment, the ship encountered a storm while in transit to Uruguay. While wind speed briefly exceeded 20 m s$^{-1}$, DMS flux and transfer velocity were not elevated.
Figure 2.4 $k_{660}$ from (2.3) vs. $U_{10}$ from five cruises. The open circles in very high winds were from a storm in transit to Uruguay during SO GasEx. The $r^2$ for a linear relationship in the range of 0–16 m s$^{-1}$ is 0.63.
Figure 2.5 $k_{660}$ from (2.3) averaged to $U_{10}$ bins, along with the $k_{DMS}$ curve computed from the NOAA COARE gas transfer model (parameters $A=1.3; B=1.0$) at 27.2 °C ($S_c = 660$ for DMS). Error bars correspond to standard errors of the mean within the bins. $k_{660}$ from SO GasEx clearly trends lower than $k_{660}$ from tropical cruises, such as TAO.
Figure 2.6 $k_{660}$ from (2.3) vs. $U_{10}$ and $U_{10n}$ from SO GasEx, with solid points indicating statically stable boundary layer ($z/L > 0.05$). The $r^2$ for a linear relationship between $k_{660}$ and wind speed is increased from 0.51 to 0.56 when $U_{10n}$ is used instead of $U_{10}$. In a more stable boundary layer, greater scatter and a potentially negative bias in $k_{660}$ are apparent.
Figure 2.7 $k_{660}$ from (2.5) vs. $U_{10n}$ for all cruises. A spectral (Kaimal) correction has been applied to hours when $z/L < 0.05$, while hours when $z/L > 0.05$ are eliminated. Normalization for the temperature dependence in the airside resistance is included (Section 3.4). The $r^2$ for a linear relationship between $k_{660}$ and $U_{10n}$ for all cruises is increased from 0.63 in Fig. 2.4 to 0.71.
Figure 2.8 Partition of drag coefficient (top) and friction velocity (bottom) to tangential and wave components per Mueller and Veron (2009) (MV09) and Soloviev and Schlüssel (1996) (SS96). In high winds, departures in both parameters from total are due to increasing partition of total wind stress to wave stress. The SS96 approach results in a greater roll off in high winds for $u^*_{nv}$ than MV09; we adopt the latter because wave age is not required for the estimation of tangential stress.
Figure 2.9 $k_{w,660}$ vs. $u_{*v}$ for all cruises, with error bars corresponding to the standard errors of the mean within the bins. A linear fit in the low-to-moderate $u_{*v}$ range yields the proportionality for interfacial exchange. Bubble-mediated transfer is estimated as the difference between the extrapolation of the linear fit and total waterside transfer velocity.
Figure 2.10 Relative error resulting from the application of the $S_c$-only normalization (2.3) for several trace gases from SST = 5 °C. Gases with intermediate solubility, such as CH$_3$I and DMS, show the greatest bias, while the widely used $S_c$-only normalization appears to be adequate for less soluble gases like CO$_2$. 
Figure 2.11 Bin-average of solubility normalized $k_{660}$' vs. $U_{10n}$, along with the estimate from COARE as a reference. Error bars correspond to standard errors of the mean within the bins. Compared to Fig. 2.5, transfer velocities from warm water cruises such as TAO and BIO are decreased slightly, whereas those from colder water cruises such as SO GasEx and VOCALS are adjusted upwards due to the temperature dependence in airside control and bubble-mediated transfer. At 9 m s$^{-1}$, the standard deviation among the five cruises is 3.6 cm hr$^{-1}$ in $k_{660}$ (Fig. 2.5), which is reduced to 2.3 cm hr$^{-1}$ in $k_{660}'$. 
Figure 2.12 (a) Difference between $k_{660}$ from (2.3) and COARE gas transfer model estimate at 27.2 °C in $U_{10}$ bins for all cruises, and (b) difference between final normalized $k_{660}'$ and COARE estimate in $U_{10n}$ bins. Error bars correspond to standard errors of the mean within the bins. Filtering for DMS$_w$ variability and atmospheric stability, normalizing for temperature dependence in airside resistance, and adjusting for both solubility and $S_c$ dependence in $k_b$ results in closer agreement in transfer velocity among the five cruises. At 15 m s$^{-1}$, $k_{660}'$ is about 7 cm hr$^{-1}$ higher than $k_{660}$ for SO GasEx.
Figure 2.13 Comparison of DMS $k_{660}$' (average of all projects) with oft-used wind speed parameterizations derived from sparingly soluble gases. Transfer velocity of DMS is significantly lower in moderate to high winds than those of less soluble gases due to its much higher solubility, and thus reduced bubble mediated transfer.
Figure 2.14 Difference between observed and COARE modeled $k_{DMS}$ at ambient conditions. The difference between the bin-average observation and model prediction is within ~2 cm hr$^{-1}$ across the wind speed range.
References


Chapter 3. Oxidation of DMS by the Hydroxyl Radical


Abstract

The hydroxyl radical (OH) is an important oxidant in the troposphere due to its high reactivity and relative abundance. Measuring the concentration of OH in situ, however, is technically challenging. Here we present a simple method of estimating an OH-equivalent oxidant concentration (“effective OH”) in the marine boundary layer (MBL) from the mass balance of dimethylsulfide (DMS). We use shipboard eddy covariance measurements of the sea-to-air DMS flux from the Vamos Ocean-Cloud-Atmosphere-Land Study Regional Experiment (VOCALS-REx) in October and November of 2008. The persistent stratocumulus cloud-cover off the west coast of South America and the associated strong inversion between MBL and the free troposphere (FT) greatly simplify the dynamics in this region and make our budget estimate possible.

From the observed diurnal cycle in DMS concentration, the nighttime entrainment velocity at the inversion is estimated to be 4 mm sec\(^{-1}\). We calculate \(1.4(\pm0.2) \times 10^6\) OH molecules cm\(^{-3}\) from the DMS budget, which represents a monthly effective concentration and is well within the range of previous estimates. Furthermore, when
linearly proportioned according to the intensity of solar flux, the resultant diel OH profile, together with DMS surface and entrainment fluxes, enables us to accurately replicate the observed diurnal cycle in DMS (correlation coefficient over 0.9). The nitrate radical (NO₃) is found to have little contribution to DMS oxidation during VOCALS-REx. An upper limit estimate of 1 pptv of bromine oxide radical (BrO) would account for 30% of DMS oxidation and lower the OH concentration to 1.0 x 10⁶ OH molecules cm⁻³. Our effective OH estimate includes the oxidation of DMS by such radicals.

3.1 Introduction

While non-reactive towards the major constituents of the atmosphere, the hydroxyl radical (OH) is the principal oxidizing free radical in the troposphere for trace species (Levy, 1971). The primary source of OH is the reaction between water vapor and excited atomic oxygen (O(¹D)); the latter is a product of the photo-dissociation of ozone (O₃) by photons between 290 and 320 nm. Reduction of the hydroperoxy radical (HO₂) by nitrogen oxide (NO), and to a lesser degree by O₃, contributes to the recycling of OH (Donahue and Prinn, 1990). OH reacts readily with most trace gases and is often regenerated during the process, resulting in sustained OH concentrations on the order of 10⁶ molecules cm⁻³ globally (Prinn et al., 1995; Spivakovsky et al., 2000). The radical is consumed primarily by reacting with carbon monoxide (CO), but it also reacts with sulfur dioxide (SO₂), nitrogen dioxide (NO₂), and a wide range of hydrocarbons. Models predict higher OH concentrations for the tropics and for the southern hemisphere. The former is attributed to higher humidity and solar fluxes that promote O₃ dissociation; the
latter is presumably due to less CO south of the equator (a result of lower anthropogenic emissions; Seinfeld and Pandis, 2006).

Due to the transient nature of the radical, measuring OH in situ was not possible until the last two decades, when specialized instruments based on absorption, fluorescence, and mass spectroscopy were developed (e.g. Mount, 1992; Hard et al., 1984; Eisele and Tanner, 1991). However, direct observations of this radical cannot be made in all field experiments (such as during VOCALS-REx). Even when in situ OH measurements are available, they might not represent a temporal or regional average, which is of importance to large-scale models.

Alternatively, global and regional distributions of OH are often estimated by measuring chemicals with known lifetimes that are exclusively consumed by this radical, such as methyl chloroform (Prinn et al., 1992). Here we follow a similar approach and use a budget analysis of naturally-derived atmospheric dimethylsulfide (DMS) to constrain the equivalent OH concentration in a clean, stratocumulus-capped marine boundary layer.

Dimethylsulfide (CH$_3$SCH$_3$) in the open ocean (DMS$_w$) is derived exclusively from phytoplankton. Due to photochemical destruction and rapid dilution, the atmospheric concentration of DMS is always orders of magnitude lower than the expected Henry’s law equilibrium concentration with DMS$_w$. Consequently, DMS gas always effluxes from the surface ocean and is the largest source of reduced sulfur to the atmosphere (Lovelock et al., 1972). The substantial magnitude of the flux and the lack of other sources make DMS an ideal gas with which to study air-sea exchange. In the atmosphere, DMS is principally oxidized by OH to form SO$_2$, dimethylsulfoxide
(DMSO) and methane sulfonic acid (MSA). SO₂ may be oxidized further to form sulfate, which contributes to aerosol growth and formation of cloud condensation nuclei (CCN). The coverage and lifetime of marine clouds formed from these CCN affect the earth’s radiative balance and may in turn be linked to biological production in the ocean (Charlson et al., 1987; Kloster et al., 2006).

The oxidation of DMS by OH proceeds through two separate pathways: the hydrogen-atom abstraction at a methyl group that subsequently leads to mostly SO₂:

\[
CH_3SCH_3 + OH \rightarrow CH_3SCH_2 \cdot + H_2O \quad (3.1)
\]

and the OH addition to the sulfur atom that can lead to a wider range of products, including DMSO and MSA:

\[
CH_3SCH_3 + OH \rightarrow CH_3S(OH)CH_3 \quad (3.2)
\]

The abstraction pathway is favored under high temperatures; the Arrhenius form for this rate constant is \( k_1 = 1.1 \times 10^{-11} \exp(-240/T) \), equating to \( 4.9 \times 10^{-12} \text{ cm}^3 \text{ molec}^{-1} \text{ sec}^{-1} \) at 298 K. The addition pathway is favored under low temperatures; this oxygen (O₂) and temperature dependent rate constant is best represented by \( k_2 = 1.0 \times 10^{-39} [\text{O}_2] \exp(5820/T) / \{1+5.0 \times 10^{-30} [\text{O}_2] \exp(6280/T)\} \) (Sander et al., 2006). At 298 K, about 80% of the oxidation of DMS goes through the abstraction pathway; the two pathways are about equal at 285 K (Stickel et al., 1993, Wallington et al., 1993).

Other oxidants, including nitrate and halogen radicals, may also consume DMS. The nitrate radical (NO₃) is produced from the reaction between NO₂ and O₃. The DMS-NO₃ reaction is several times slower than the DMS-OH reaction. Unlike OH, NO₃ can only have an appreciable concentration at night because it is photolyzed rapidly during the day. Studies from Equatorial Christmas Island showed that in the remote Pacific, the
contribution to DMS loss from NO₃ is less than a few percent due to the low levels of NOₓ (NO + NO₂) away from anthropogenic activities (Davis et al., 1999; Chen et al., 2000). The same studies also suggested that the DMS-chlorine reaction, while still very poorly understood, accounts for no more than 5~10% to the oxidation of DMS. Other authors (e.g. Sciare et al., 2000) have found OH and NO₃ to be inadequate in explaining their observed DMS losses, and speculated that the naturally derived bromine oxide (BrO) might be another significant DMS oxidant.

The aforementioned Christmas Island experiments demonstrated a clear diurnal cycle of DMS in the MBL (Bandy et al., 1996; Davis et al., 1999). While the supply of DMS to the boundary layer via air-sea exchange is continuous, destruction of DMS principally takes place during the sunlit hours, when OH is available. As a result, DMS starts to increase from its daily minimum just before sunset and builds up through the night until just after sunrise, at which time photolysis causes DMS to decrease until the next cycle. This diurnal cycle in DMS can be altered in a pollution-influenced marine environment where anthropogenically produced NOₓ level is higher, as nighttime oxidation of DMS by NO₃ can be comparable in magnitude to daytime oxidation by OH (Yvon et al., 1996).

The equivalent OH we estimate here from VOCALS-REx is an effective concentration, the concentration of OH that would cause the same rate of DMS loss as if OH were the only oxidant. In all likelihood, our effective OH is slightly higher than the actual OH concentration, since it includes minor contributions from the other oxidants.

The VOCALS-REx experiment took place in October and November of 2008 in the Southeast Pacific, off the west coast of Chile and Peru
The experiment aimed to address the interactions among the ocean, atmosphere, and clouds in this tightly coupled regional climate system. The Andes mountain range to the east forces strong winds parallel to the coast. Ekman transport results in coastal upwelling of cold, nutrient-rich waters, which leads to stability in the lower troposphere. The warm, dry air over the cool water surface, together with the large-scale subsidence from the Hadley Cell, creates a widespread and persistent stratocumulus cloud deck (Bretherton et al., 2004). These thin, low-lying, and hence radiatively-cooling clouds form from CCN aerosols. While anthropogenic emissions dominate the coastal region, a significant fraction of CCN likely originates from DMS-derived sulfate in the clean marine environment offshore. From time to time, hundred-kilometer sized openings (which can be observed from satellites) form in the stratocumulus cloud-deck and advect with the mean wind. These features have been coined “pockets of open cells” (POCs) and are hypothesized to be related to removal of CCN by drizzle from the air (Stevens et al., 2003). To further our understanding in this phenomenon, we need to look at the origin and fate of the CCN-precursors, including DMS and its oxidants.

A stratocumulus marine boundary layer is characterized by vigorous mixing due to the intense longwave radiative cooling at the cloud-top (Lilly 1968). Air entrains down from the FT, which is balanced by a deepening of the MBL and wind divergence. The strong inversion in effect caps the MBL, confining surface-borne scalars like DMS. Therefore, a stratocumulus region such as the Southeast Pacific can be viewed as a natural analog to a box model, with clearly defined inputs and outputs. These simplified conditions enable us to estimate the effective OH concentration from mass balance.
3.2 Experimental and Methods

The National Oceanographic and Atmospheric Administration (NOAA) ship *Ronald H. Brown (RHB)* was deployed in the Southeast Pacific from approximately October 24th to November 30th, 2008. In addition to retrieving and maintaining the Woods Hole Oceanographic Institute’s Stratus buoy at 85°W and 20°S, a number of E-W transects were made at latitudes between 18.5°S and 21.5°S to survey eddies and upwelling features. The edges of such eddies are often associated with higher levels of biological productivity, hence more DMSw (T. Bates, P. Matrai: unpublished data). In the same general time frame, multiple aircraft, including the National Science Foundation (NSF)/National Center for Atmospheric Research (NCAR) C-130, repeatedly flew westward from its base station in Arica, Chile to the stratocumulus region offshore. The anthropogenic influence was minimal away from the coast, with average CO, O3, and SO2 concentrations of 65 ppbv, 30 ppbv, and 25 pptv, respectively, typical of a clean MBL. Most of the research flights were designed to survey the 20°S cross-section and study the formation and development of POCs.

During the experiment, two nearly identical atmospheric pressure ionization mass spectrometers (APIMS) were used to measure DMS on board of *RHB* and the C-130, the distinction being that the instrument at sea was sampling at a high rate of 20 Hz to quantify the sea-to-air flux, whereas the one aloft was measuring concentration every 10 seconds. The measurement of DMS by the APIMS with isotopically labeled standard technique has been described previously (Bandy et al., 2002; Huebert et al., 2004;
Blomquist et al., 2006). Most recently, Blomquist et al. (2009) discussed in great details about the instrument setup, flux processing, data refinement, and uncertainty estimates.

We estimate the OH-equivalent oxidant concentration from the budget analysis of DMS. The conservation equation for any chemical species, \( S \), in the marine boundary layer is represented as (with the overbar representing an average in time/space):

\[
\frac{\partial \bar{S}}{\partial t} + u \frac{\partial \bar{S}}{\partial x} + \frac{\partial \bar{S}' w'}{\partial z} = P - L
\]  

The first term on the left represents the time rate of change in the concentration of \( S \). The second term is the horizontal advection of \( S \) due to gradients along the mean wind \((u)\). The third term is the flux divergence, the difference between vertical fluxes at the bottom and top of the box model. Local (chemical) production and loss of \( S \) are represented by \( P \) and \( L \), respectively, on the right side of the conservation equation.

As described in the previous section, DMS fluxes upwards from the surface ocean to the stratocumulus-topped MBL, where it is oxidized (principally by OH), diluted by DMS-free air entrained from the FT, and transported horizontally. DMS is not produced chemically, so \( P = 0 \). Therefore the chemical loss due to oxidation can be calculated by closure if we know the DMS surface flux, entrainment flux, horizontal advection, and the time rate of change.

### 3.2.1 Surface Flux of DMS

On the ship, the inlet of our mass spectrometer was located at 18 m above the water surface on the jackstaff, where flow-distortion due to the ship’s superstructure is minimized. A precisely known amount of the triply-deuterated DMS standard was continuously injected at the inlet and combined with a high ambient air flow. The DMS
concentration was determined from the ratio between the ambient and standard DMS mass spectrometer signals. Relative wind speeds, acceleration, and rotation in three axes were recorded at the same frequency as the DMS by a Gill Sonic anemometer and a Systron-Donner Motionpak accelerometer, respectively. Ship’s motion was removed from relative winds to get true winds following Edson et al., (1998). To obtain surface flux \( F_0 \) via eddy covariance, DMS concentration was correlated with the vertical wind velocity \( w \) in the form of \( F_0 = \overline{DMS'}w' \) (primes here denote deviations from the means). Flux was initially computed in ten-minute segments that overlapped by 50%. To remove times when sampling conditions were unfavorable, any ten-minute segment with relative wind direction more than 90 degrees off the bow or gyro heading changing by over 10 degrees (ship maneuvering) was screened out. The remaining “good” segments were averaged to hourly values.

Time-series of 18-m DMS concentration and flux from the \textit{RHB} are shown in Fig. 3.1, with campaign means (±standard deviations) of 57±(26) pptv and 3.4 (±1.9) \( \mu \)moles m\(^{-2}\) day\(^{-1}\), respectively. Also shown are true wind speed and direction from the sonic anemometer at the same height. The wind direction hardly varied, almost always coming from the South/Southeast (150°). We limit our averaging to offshore observations only (73°W ~ 86°W and 18°S ~ 22°S, hereinafter ‘the VOCALS region’). Near-shore data were not included in the averaging due to anthropogenic influence as well as heterogeneities in the atmospheric DMS field caused by spikes in \( DMS_w \) on the edges of localized coastal eddies.

Even with the ship steaming at up to 12 knots, the classic sinusoidal diurnal cycle in DMS is clearly visible on a number of days (Fig. 3.2). This observation implies a
widespread horizontal homogeneity over the region with limited meso-scale variability. Continental influence, which is usually associated with high levels of NO\textsubscript{x}, and thus high NO\textsubscript{3}, was minimal.

3.2.2 Boundary Layer Structure

We determined the boundary layer structure during VOCALS-REx from shipboard radiosonde launches every six hours. Figure 3.3 shows a time series of temperature contours from the soundings, with the dark line indicating the inversion immediately above the stratocumulus cloud deck. Over the course of more than a month, the structure of the MBL was relatively constant, with an average temperature of 13°C. In addition to radiosondes, a W-band radar (24 m range gate, 50 m pulse length) from the ship measured cloud top height continuously. On average, the stratocumulus cloud top varied from ~1350 m near dawn to ~1200 m near dusk, suggesting a greater entrainment rate in the nighttime. Inversion heights determined from radiosondes are nearly identical with the cloud top estimates from the W-band radar (S. de Szoeke, personal communication).

We used DMS concentration from the C-130 aircraft (\textit{DMS}_{c130}) to infer the distribution of the scalar within the MBL and exchange with the FT. Figure 3.4 shows a typical profile of the lower troposphere during VOCALS-REx. \textit{DMS}_{c130} decreased gradually with altitude within the MBL and rapidly dropped to near-zero above the stratocumulus cloud-top in the FT. Across the thin inversion layer of only a few tens of meters, the changes in dew point and potential temperature were usually of the order of 20 and 10 K, respectively.
When the MBL is shallow and well mixed, scalars with long lifetimes are thought to be homogeneously distributed vertically. DMS in the boundary layer is emitted from the ocean surface and diluted by entrainment above; with a mixing time in the MBL (~1 hour) much shorter than its lifetime (1~2 days, estimated later), DMS concentration at the top of the MBL is usually only a few pptv lower than at the surface (Faloona et al., 2005). However, this difference was more significant in the VOCALS region, where the average inversion height was 1.3 km. Figure 3.5 shows the mean profile from the C-130 during the campaign. Altitude was normalized to the inversion height ($z_i$). DMS$_{C130}$ aloft was normalized to the concentration from the lowest leg at ~30 m (DMS$_{C130,0}$). On average, DMS$_{C130}$ immediately below the inversion was ~80% of the concentration near the surface, exhibiting a linear decrease with height. We linearly regressed DMS$_{C130}$/DMS$_{C130,0}$ against $z/z_i$ in the MBL ($r^2 = 0.8$). The fraction of DMS at height $z$ relative to the surface value can then be parameterized as $1 - \alpha (z/z_i)$, where $\alpha = 0.2$ represents the “decoupling” parameter. This formulation of the MBL structure is similar to what is used in Wood and Bretherton (2004).

3.2.3 Advective Flux of DMS

In VOCALS-REx, DMS concentration and flux measured from the ship show increasing trends away from the coast due to generally greater DMS$_w$ concentration and higher wind speeds offshore (Fig. 3.6). The ratio between DMS flux and concentration, however, was nearly constant with longitude. Bin-averaged inversion heights from radiosondes and 18-m wind speeds within the latitude range of 20±2°S are also plotted on
the same figure. The boundary layer was deeper offshore (~1.5 km at 85°W) than at the coast (~1.1 km), as is typical of this region.

We estimated the advective flux of DMS from the horizontal gradient computed from shipboard DMS concentration. Using multi-variate linear regression, DMS averaged to daily means (to remove diel variability) was regressed against Lat and Lon. The fitting coefficients were -2 pptv °Lon$^{-1}$ and -1 pptv °Lat$^{-1}$, with higher DMS towards the West/Southwest (Fig. 3.7). For an average wind speed of 6 m sec$^{-1}$ from 150°, multiplying the zonal and meridional winds (-3 and 5 m sec$^{-1}$, respectively) by the DMS gradient gave an advection of 0.04 pptv hr$^{-1}$. Integrating over the 1.3 km column led to an advective flux of only 0.05 μmoles m$^{-2}$ day$^{-1}$, less than 1% of the DMS sea-to-air flux. Advection was small because the mean wind direction was largely orthogonal to the horizontal DMS gradient; we set it to zero hereafter.

3.2.4 Entrainment Velocity

To calculate the entrainment flux of DMS, we need to know the entrainment velocity ($\omega_e$) at the inversion. While $\omega_e$ was not directly measured during VOCALS-REx, we can estimate it from the observed nighttime increase in shipboard DMS concentration. To account for the dilution effect from changing MBL depth, we integrate DMS concentration from the ship to the depth of MBL using the vertical gradient determined from Section 2.2, resulting in column concentration, $\langle$DMS$\rangle$, in units of μmoles m$^{-2}$.

The vertically integrated form of (3.3) is shown in (3.4), with the advection and production terms now dropped out. Now every term is in units of flux:
\[ \frac{\partial \langle DMS \rangle}{\partial t} = F_0 - \omega_e (DMS_{z_i} - DMS_{z_i^+}) - k_{PFO} \langle DMS \rangle \]  \hspace{1cm} (3.4)

The term on the left side of (3.4) is the time rate of change. The last term on the right represents the local loss of DMS, which has been expanded to the product of \( k_{PFO} \) (a pseudo first order reaction rate constant) and DMS column concentration. We have separated flux divergence into the first and second terms on the right hand side of (3.4), representing the DMS fluxes at the ocean surface and inversion, respectively. The entrainment flux is formulated as the entrainment velocity, \( \omega_e \), multiplied by the concentration jump across the inversion (Lilly, 1968), with subscripts \( z_i^- \) and \( z_i^+ \) denoting just below and above \( z_i \). The concentration just below the inversion can be approximated from the 18-m concentration by the fraction \((1 - \alpha)\). Taking the concentration above the inversion to be zero, (3.4) thus simplifies to:

\[ \frac{\partial \langle DMS \rangle}{\partial t} = F_0 - \omega_e (1 - \alpha)DMS - k_{PFO} \langle DMS \rangle \]  \hspace{1cm} (3.5)

The nighttime build up of DMS column concentration appears to be linear with time (Fig. 3.8). The slope over these hours approximately represents the time rate of change in \( \langle DMS \rangle \), which includes the flux divergence plus any nocturnal chemical loss. There is no OH at night, and we assume the NO\(_3\) reaction to be insignificant (this assumption is validated in the last section); the loss term thus drops out. From hour 1 to 10 UTC (or 20:00 to 05:00 local time), we calculated a time rate of change in \( \langle DMS \rangle \) of 2.6 \( \mu \)moles m\(^{-2}\) day\(^{-1}\). The mean nighttime surface flux obtained by eddy covariance was 3.2 \( \mu \)moles m\(^{-2}\) day\(^{-1}\). The nighttime entrainment flux thus needs to be 0.6 \( \mu \)moles m\(^{-2}\) day\(^{-1}\) to balance the budget, requiring \( \omega_e = 4 \) mm sec\(^{-1}\).
The entrainment velocity derived above is only a nighttime value, and might not be representative of the climatological mean since there was possibly diurnal modulation of the inversion height. Shortwave heating of cloud top stabilizes the MBL and tends to reduce entrainment during the day, while longwave cooling at night may enhance entrainment. Off the coast of California, where stratocumulus clouds are also prevalent, eddy covariance flux measurements of three conserved scalars (DMS, O₃, and total water) immediately above and below the inversion indicated a range of 1.2~7.2 mm sec⁻¹ for \( \omega_e \), with higher values at night (Faloona et al., 2005). Using a mixed-layer approach, Caldwell et al. (2005) found a 6-day average \( \omega_e \) of 4±1 mm sec⁻¹ during the East Pacific Investigation of Climate (EPIC) stratocumulus cruise in 2001. Combining satellite observations with National Centers for Environmental Prediction (NCEP) reanalysis, Wood and Bretherton (2004) determined an \( \omega_e \) of 2~6 mm sec⁻¹ in the tropical and subtropical east Pacific, with a likely value of 4 mm sec⁻¹ in the VOCALS region. While our estimated nighttime \( \omega_e \) could theoretically be an over-estimate of the diurnal average, 4 mm sec⁻¹ appears to be a reasonable value compared to previous results. We show later that our calculated effective OH concentration is not very sensitive to the choice of \( \omega_e \), because reaction with OH represents a far greater removal term for DMS than dilution due to entrainment. At the limit of zero for \( \alpha \) (i.e. no decoupling in DMS), \( \omega_e \) is lowered to 3 mm sec⁻¹, still within the range of previous estimates.

### 3.2.5 Estimating Effective OH

From the clear diurnal cycles and near-constant flux to concentration ratio in the project averages (Fig. 3.2 and Fig. 3.6), diurnally averaged DMS field in the MBL
appears to be approximately constant in space and time over the duration of the project. We thus assume DMS to be in a steady-state on that scale.

For the purpose of this derivation we first assume that the oxidation of DMS is solely due to reactions with OH. The pseudo first-order rate constant $k_{PFO}$ in (3.5) is expressed as $k_{OH}$, the total second order rate constant of the DMS-OH reaction ($k_{OH} = k_1 + k_2$), multiplied by the concentration of OH. With an average MBL temperature of 13°C during VOCALS-REx, we calculate $k_{OH}$ to be $7.9 \times 10^{-12}$ cm$^3$ molec$^{-1}$ sec$^{-1}$, with the abstraction pathway accounting for 60% of the total oxidation.

Rearranging and expanding (3.5) with no time rate of change leads to:

$$OH = \frac{F_0 - \omega_e (1 - \alpha)DMS}{\langle DMS \rangle k_{OH}} \tag{3.6}$$

Using $\omega_e = 4$ mm sec$^{-1}$ and $\alpha = 0.2$, we find an effective OH concentration of $1.4 \times 10^6$ molec cm$^{-3}$ for VOCALS-REx, somewhat higher than the estimated global average possibly due to high water content and solar radiation and low anthropogenic influence in this region. From PEM-Tropics A and B, which took place during September/October of 1996 and March/April of 1999, respectively, Olsen et al. (2001) showed that OH ranged from $1.4\sim1.8 \times 10^6$ molec cm$^{-3}$ in the latitude range of 10S~30S and in the lowest 2 km, with higher values in the austral spring. The PEM-Tropics experiments did not extend into the stratocumulus region, however, where solar flux is lowered by 10~20% relative to the normal trade wind cumulus regime. An effective OH concentration of $1.4 \times 10^6$ molec cm$^{-3}$ implies a lifetime of 1.1 days for DMS with respect to OH in the MBL, well within the range of previous estimates of 0.5~2 days (e.g., Wine et al., 1981; Hynes et al., 1986).

Allowing for a substantial uncertainty of ±2 mm sec$^{-1}$ in the entrainment velocity
(due to diel variability, for example), the relative error in the calculated OH concentration is only ~12% because dilution due to entrainment is small compared to photochemical loss. The potential of \( \omega_e \) being biased high implies that our calculated OH might be an underestimate (though this could be partially compensated by the our assumption that OH is the only DMS oxidant). At a limit of \( \alpha = 0 \), OH is lowered to \( 1.2 \times 10^6 \) molec \( \text{cm}^{-3} \).

The uncertainties in the inversion heights from radiosoundes and the W-band were about 25 m (S. de Szoeke, personal communication, 2009). Assuming an absolute uncertainty of 2 mm sec\(^{-1} \) for \( \omega_e \) and 0.1 for \( \alpha \), the propagated absolute uncertainty for the equivalent OH concentration is ~0.2 molecules \( \text{cm}^{-3} \), or 15%. One apparent benefit of estimating OH from the DMS flux and concentration is that both measurements are made by the same instrument; any calibration uncertainty in the instrument (such as flow rate or standard concentration) is thus negated, not affecting the final outcome.

### 3.3 Discussion

The above OH estimate is an average value over a month. In reality, OH is only produced during the day when photochemistry is possible; the radical is consumed completely when there is no light. The next logical step is to approximate the diel variation in OH, which has pronounced impacts on the cycling of sulfur species in the MBL, from DMS to \( \text{SO}_2 \) to sulfate.

#### 3.3.1 Diel Variability in OH

To approximate the diel profile of OH, we need to know the sources and sinks of the \( \text{HO}_x \) (OH + HO\(_2\)) system because the OH : HO\(_2\) ratio is usually a steady-state constant
(Faloona et al. 2000). To a first order, the availability solar flux, and hence the ozone photolysis frequency, $J_O(1^D)$, limits OH production since the subsequent reaction between $O(1^D)$ and water vapor is rapid. OH can also be recycled from HO$_2$ by reaction with NO and O$_3$. As for the sinks of HO$_x$, both the self-reaction of HO$_2$ and OH + HO$_2$ (when NO$_x$ is low) represent quadratic loss in HO$_x$. In a high NO$_x$ environment, the OH + NO$_2$ reaction is the major sink and a linear loss for HO$_x$.

We can approximate the dependence of steady-state OH concentration on solar flux as $J_O(1^D)^b$, with the exponent $b$ being NO$_x$-dependent. When NO$_x$ is high, OH relates linearly to $J_O(1^D)$ because HO$_2$ becomes less important (e.g. Holland et al., 1998). When NO$_x$ is low, OH is formed by the reaction with water vapor as well as from recycling of HO$_2$ by O$_3$ (and NO, if any), while the loss of HO$_x$ is largely quadratic. Overall $b$ likely takes on a value between 0.5 and 1. Berresheim et al. (2003) found that $b = 0.68$ off the coast of Crete in the Mediterranean, where the mean NO$_2$ mixing ratio was 0.4 ppbv. While there was no direct measurements of NO$_x$ during VOCALS-REx, in situ observations from 10S~30S during PEM-Tropics A and B yielded 5~10 pptv of NO$_x$ in the lowest 2 km (Olson et al., 2001). Schultz et al. (1999) showed that in the tropical South Pacific, the median NO$_x$ level was only 4 pptv in the lowest 2 km. Thus VOCALS likely falls in the low NO$_x$ regime and one may expect $b$ to be less than 1.

We first binned solar flux measured by the shipboard radiometer in the VOCALS region to the time of day, which expectedly resulted in a smooth curve with peak insolation at ~17:00 UTC. Not knowing the exact value of $b$, hourly hydroxyl concentrations (OH$_t$) were proportioned as linear and square root functions of the solar flux, with the diel average of the radical constrained by $1.4 \times 10^6$ molecules cm$^{-3}$. The
radical peaked at $\sim 5 \times 10^6$ molecules cm$^{-3}$ in the linear case and $\sim 4 \times 10^6$ molecules cm$^{-3}$ in the square root case (with a broader distribution). While this method seems rather simplistic compared to photochemical models containing many photoactive species, the general shape of our derived diel OH profile agrees fairly well with previous direct observations in the tropical/subtropical Pacific basin (e.g. Davis et al., 2001; Mauldin et al., 2001). As another comparison to our effective OH estimate from DMS, we use photolysis rates of O$_3$ and hydrogen peroxide (H$_2$O$_2$) predicted from the NCAR Tropospheric Ultraviolet and Visible (TUV) Radiation Model (http://cprm.acd.ucar.edu/Models/TUV/), reaction rate between O$_3$ and HO$_2$, and destruction rate due to CO, methane, hydrogen gas, O$_3$, H$_2$O$_2$, and HO$_2$ from Sander et al. (2006) to calculate the expected OH concentration. The TUV estimates photolysis rates based on time, latitude, longitude, and oxidant field, and accounts for clouds and aerosol scattering using climatological reflectivity from the Total Ozone Mapping Spectrometer (TOMS). The model yielded a noontime OH concentration of $\sim 6 \times 10^6$ molecules cm$^{-3}$, which is $\sim 20\%$ higher than our daytime peak in the linear scenario but likely within the range of uncertainties.

3.3.2 Reproducing the Diurnal Cycle of DMS

The oxidation of DMS in the atmosphere has been modeled previously given an estimate of OH (Shon et al., 2001; Nowak et al., 2001). Using a similar approach and building upon (3.5), the surface DMS concentration at time $t$ can be written as:

$$
\langle DMS \rangle_t = \langle DMS \rangle_{t-1} + F_{0,t-1} - \omega_t (1 - \alpha) DMS_{t-1} - k_{PFO,t-1} \langle DMS \rangle_{t-1}
$$

(3.7)
For simplicity, we used the average $\omega_e$ of 4 mm sec$^{-1}$, neglecting any potential diel variation in the entrainment velocity. The implied DMS diurnal variation is not very sensitive to the choice of $\omega_e$ for the same reason as in Section 2.5.

We first equated, $k_{PFO,t}$, the pseudo first order reaction rate constant of DMS, simply to $k_{OH} OH_t$, with $OH_t$ being the time-dependent radical concentration calculated from Section 3.1. Figure 3.9(a) shows the diel cycle of $OH$ approximated as a linear function of solar flux, DMS column concentration calculated from (3.7), as well as the observation. Figure 3.9(b) shows the same calculation except $OH$ was approximated as a square root function of solar flux. It is apparent that the calculated DMS agrees closer with the observation when the dependence on solar flux, and hence $JO(^1D)$, is linear. In the square root case, implied DMS does not decrease as rapidly as observed after sunrise because the OH distribution is too broad. The linear dependence might come as a surprise given the expectedly low level of NO$_x$; this might be due to chemistry that are not considered or well understood. In the next section we explore the importance of nitrate and bromine oxide radicals in DMS oxidation, and see whether we can further improve the fit to the observed diurnal cycle.

3.3.3 DMS Oxidation by NO$_3$ and BrO

The rate constant of the DMS-NO$_3$ reaction can be represented by $k_{NO_3} = 1.9 \times 10^{-13} \exp(500/T)$, giving $1.0 \times 10^{-12}$ cm$^3$ molec$^{-1}$ sec$^{-1}$ at 298 K (Sander et al., 2006). Studies from Allan et al. (2000) in the relatively clean MBL at Mace Head suggested that the nocturnal level of the nitrate radical hovers on the order of a pptv, with reaction with
DMS being its most important loss mechanism. Platt and Heintz (1994) estimated a globally averaged concentration of 3 pptv for NO$_3$.

Uncertain about how much NO$_3$ was present in the Southeast Pacific, we apportioned the total oxidative loss of DMS to fractions due to NO$_3$ and OH’ (the prime denotes the ‘actual’ OH concentration, assuming a linear dependence on solar flux). For example, if 10% of DMS oxidation was due to NO$_3$, the ratio between diurnal averaged $k_{NO_3}$ NO$_3$ and $k_{OH}$ OH’ should be 1:9. To crudely mimic the diel variation in NO$_3$, we assume its nighttime level to be twice the diurnal average and the daytime concentration to be zero. In reality, NO$_3$ might not achieve steady-state throughout the night (Allan et al., 2000). We arrived at the closest least-square fit between observed and calculated DMS (correlation coefficient exceeding 0.9) when 99% of DMS oxidation is due to OH, as any additional NO$_3$ would degrade the fit; this corresponds to less than 0.005 pptv of NO$_3$. In all likelihood, low levels of NO$_3$ and O$_3$ were limiting the production of NO$_3$ during VOCALS-REx.

The rate constant of the DMS-BrO reaction can be represented by $k_{BrO} = 1.4 \times 10^{-14} \exp(950/T)$, yielding $3.4 \times 10^{-13}$ cm$^3$ molec$^{-1}$ sec$^{-1}$ at 298 K (Sander et al., 2006). Read et al. (2008) measured BrO with differential optical absorption spectroscopy (DOAS) at Cape Verde in the tropical Atlantic Ocean. They found an average BrO concentration over 2 pptv during the day and zero at night, with a diurnal profile resembling a ‘top hat’ distribution.

During VOCALS-REx, with a similar DOAS technique, no significant BrO was detected, with ~1 pptv being the upper limit daytime peak concentration (R. Volkamer, personal communication, 2009). The same instrument also detected ~1 pptv of iodine
oxide (IO), a level insignificant towards DMS oxidation due to its much lower reactivity. If we crudely approximate the diurnal profile of BrO with a square root dependence on solar flux, 1 pptv of daytime maximum BrO implies a diel average concentration just under 0.4 pptv. This upper limit concentration of BrO would account for ~30% of total DMS oxidation; the OH concentration would then need to be reduced to 1.0 x 10^6 molecules cm^{-3}. The shape of implied DMS from (3.7) with this allocation of oxidants is not significantly different from what is shown in Fig. 3.9(a). Therefore from the implied DMS diurnal cycle alone, it is difficult to assess whether 1 pptv of BrO is too high. Our OH-equivalent oxidant concentration may still include other DMS oxidants, such as NO_3 and BrO.

3.4 Conclusions

We have demonstrated that a month-long average OH-equivalent oxidant concentration in a stratocumulus-capped marine boundary layer can be derived from the sea-to-air flux and concentration of DMS measured from a ship with a simple mass balance method. From aircraft profiles during VOCALS-REx, DMS concentration appeared to decrease linearly with altitude in the MBL due to decoupling. On average, DMS immediately below the inversion was ~80% of the surface concentration. DMS concentration in the MBL displayed the classic sinusoidal diurnal cycle, in which daily maxima and minima were found to be just after sunrise and just before sunset, respectively, as a result of the continuous build up via air-sea exchange and daytime consumption by photochemistry. From the nighttime build up of DMS column concentration, we estimated an entrainment velocity of 4 mm sec^{-1} from the difference
between the time rate of change and the observed surface flux. The advective flux of DMS was found to be insignificant because the horizontal gradient in DMS was largely perpendicular to the mean wind direction. The calculated OH concentration of $1.4 \pm 0.2 \times 10^6$ molecules cm$^{-3}$ from the DMS budget equation agrees well with previous estimates, and represents a regional OH-equivalent oxidant concentration. Actual OH will be less by the extent that other DMS oxidants, such as nitrate- and halogen- radicals, react with DMS. The applicability of this approach of estimating the equivalent-OH concentration hinged on the stationarity and horizontal homogeneity of the region over a long enough time scale. Areas with much greater mesoscale variability or additional source and sink terms will complicate the budget analysis.

We fitted the OH-equivalent oxidant concentration as both linear and square root functions of the observed solar flux averaged to the time of day. Using the resultant diel profiles in OH, the mean observed DMS surface flux, and an entrainment velocity of 4 mm sec$^{-1}$, we were able to accurately replicate the observed diurnal cycle of DMS in the MBL. Despite the expected low NO$_x$ concentration, a linear relationship between OH and solar flux, and hence JO(1D), yielded a better fit between observed and implied diurnal cycles (correlation coefficient over 0.9) than a square root relationship. The nitrate radical was considered in the DMS oxidation and found to be unimportant in VOCALS-REx. We cannot rule out BrO as a possible oxidant, however. An upper limit estimate of 1 pptv of BrO (daytime peak) would account for 30% of total DMS oxidation, lowering our effective OH to $1.0 \times 10^6$ molecules cm$^{-3}$.

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Figure 3.1 Time-series of DMS concentration and surface flux (top), and true wind speed and direction (bottom) from the RHB during VOCALS-REx. Elevated DMS flux usually corresponded to spikes in DMS$_w$ on the edges of localized eddies and high winds. The wind was consistently from the S/SE.
Figure 3.2 Examples of diurnal cycles in DMS concentration observed from the RHB, along with solar flux measured from a shipboard radiometer. Maxima and minima in DMS were usually observed just after sunrise and just before sunset, respectively, as a result of continuous build up of the gas via air-sea exchange and daytime photochemistry.
Figure 3.3 Time-series of temperature contours from the radiosondes from the *RHB*, with the dark line indicating the inversion. Over a ~monthly timescale, the trend in the depth of the MBL along the E-W transect was relatively constant, deeper offshore (e.g. Oct 26th and Nov 20th) than near the coast (e.g. Nov 3rd and Nov 10th). The average temperature was 13 °C in the middle of the MBL.
Figure 3.4 A typical profile of the lower troposphere determined from the C-130 (12:24~12:35 UTC on Oct 25th during Research Flight No. 5). DMS_{C130} decreased linearly with altitude in the MBL (the lower ~1300 m in this profile) and dropped to zero in the FT. A strong inversion atop the stratocumulus cloud deck (indicated by high liquid water content) was defined by sharp gradients in dew point and potential temperature.
Figure 3.5 Averaged profile from the C-130 with altitude normalized to the inversion height and DMS$_{C_{130}}$ normalized to concentration from the lowest leg. Concentration at height $z$ is related to the near-surface concentration by the parameterization $1 - \alpha (z/z_i)$, with $\alpha$ being the DMS “decoupling” parameter and on average equaling to 0.2.
Figure 3.6 VOCALS averages of atmospheric DMS concentration and surface flux from the RHB in the latitude range of 20±2°S, as well as wind speed and inversion height. The ratio between DMS flux to concentration was largely constant, suggesting minimal time rate of change in the DMS field on a monthly timescale. Both the boundary layer height and wind speed were greater offshore.
Figure 3.7 Multi-regression of daily DMS concentration from the RHB (colored circles) vs. Lat and Lon. Contours from the regression analysis show a gradient in DMS concentration that is orthogonal to the mean wind vector. The advective flux was thus minimal. The box with dashed lines defines the “VOCALS region.”
Figure 3.8 DMS surface flux, column concentration, shortwave radiation, and cloud top measured from the \textit{RHB} in the VOCALS region averaged to the time of day. The nighttime increase in DMS column concentration was approximately linear, enabling us to estimate an entrainment velocity of 4 mm sec$^{-1}$. 
Figure 3.9 Observed and implied DMS diurnal cycle: a) with OH linearly proportional to the solar flux; b) with OH proportional to the square root of the solar flux. A linear relationship between OH and solar flux yielded a better fit to the DMS diurnal cycle, despite the expected low NOx concentration. To investigate the importance of other oxidants, the OH-equivalent oxidant concentration was apportioned to OH, NO3 and BrO. Even a minuscule amount of NO3 was found to degrade the fit between the implied and observed DMS. BrO cannot be distinguished from OH based on the DMS diurnal cycle. For VOCALS-REx, the maximum DMS oxidation due to BrO was estimated to be 30%.
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Chapter 4. Cycling of SO₂ and SO₄²⁻ in the Marine Atmosphere


Abstract

Dimethylsulfide (DMS) emitted from the ocean is a natural precursor gas for sulfur dioxide (SO₂) and non-sea-salt sulfate aerosols (SO₄²⁻), which affect cloud formation in the marine atmosphere. During the VAMOS-Ocean-Cloud-Atmosphere-Land Study Regional Experiment (VOCALS-REx) in October and November of 2008, a variety of instrumentation platforms were deployed in the Southeast Pacific off the coast of Chile and Peru to study the linkage between aerosols and stratocumulus clouds. We present here observations from the NOAA Ship *Ronald H. Brown* and the NSF/NCAR C-130 aircraft along ~20°S from the coast (70°W) to a remote marine region (85°W). While SO₄²⁻ and SO₂ concentrations were distinctly elevated in the coastal marine boundary layer (MBL) due to anthropogenic influence (~800 and 80 pptv, respectively), their concentrations rapidly decreased offshore (~100 and 25 pptv). Compared to the mass entrainment fluxes of SO₄²⁻ and SO₂ from the free troposphere (0.5±0.3 and 0.3±0.2
μmoles m$^{-2}$ day$^{-1}$), the sea-to-air DMS flux (3.8±0.1 μmoles m$^{-2}$ day$^{-1}$) remained the predominant source of sulfur mass to the MBL. In-cloud oxidation was found to be the most important mechanism for SO$_2$ removal (2.1±0.4 μmoles m$^{-2}$ day$^{-1}$) and in situ SO$_4^{2-}$ production. Surface SO$_4^{2-}$ concentration in the remote region displayed pronounced and unexpected diel variability, increasing rapidly in the first few hours after sunset and then decaying for the rest of the diurnal cycle. We theorize that the increase in SO$_4^{2-}$ was due to nighttime coupling of the MBL that mixed down cloud-processed air, while sporadic precipitation scavenging and daytime decoupling were responsible for the subsequent decline in SO$_4^{2-}$.

Nucleation of new particles was surprisingly common in the remote region of VOCALS-REx, as observed on both the aircraft and the ship. Often associated with precipitation events and pockets of open cells (POCs), these new particle formation events likely took place in the cloud outflow region, where the preexisting aerosol surface area was low due to scavenging and the reactive gas concentrations were high. The low anthropogenic influence in the remote region and ample DMS concentration suggest that the source of sulfuric acid vapor for nucleation was likely natural.

4.1 Introduction

The ocean is the largest source of natural reduced sulfur gas to the atmosphere, most of which is in the form of dimethylsulfide (DMS), a volatile organic compound produced from phytoplankton. The annual global sea-to-air DMS emission is estimated to be 15~33 Tg S (Kettle and Andreae, 2000), depending on the gas exchange and wind
speed parameterization. In the atmosphere, DMS is principally oxidized by the hydroxyl radical (OH) to a number of products, including sulfur dioxide (SO$_2$). In addition to DMS oxidation, SO$_2$ is formed from fossil fuel combustion and biomass burning and removed from the MBL through deposition to the ocean surface and oxidative reactions in the gas and aqueous phase. Gas phase oxidation of SO$_2$ by OH and subsequent reactions with water vapor (H$_2$O$_{(g)}$) yield sulfuric acid vapor (H$_2$SO$_4_{(g)}$), which usually condenses upon preexisting aerosol surfaces and increases sulfate aerosol (SO$_4^{2-}$) mass. Under infrequent, specific conditions, H$_2$SO$_4_{(g)}$ may undergo gas-to-particle nucleation and form numerous new nm-sized SO$_4^{2-}$ aerosols, which then grow by condensation and coagulation.

Generally the oxidation of SO$_2$ in the aqueous phase, which can take place in cloud water or in sea-salt aerosols by hydrogen peroxide (H$_2$O$_2$) at low pH and with ozone (O$_3$) at high pH, is much faster than in the gas phase. Aqueous oxidation of SO$_2$ also leads to SO$_4^{2-}$ aerosols, which are termed non-sea-salt-SO$_4^{2-}$ to differentiate from sulfate originated in seawater. These aerosols from oxidative reactions typically make up the bulk of the submicron (fine) aerosol mass, causing the fine mode to be acidic. Sea-salt aerosols formed from wave breaking account for the majority of the supermicron (coarse) number as well as total aerosol mass, and tend to be more basic than the fine mode because of the initial alkalinity of seawater (pH~8.1) and its carbonate buffering capacity. Due to dissolution of CO$_2$ and other acids, cloud water typically has a pH under 5 (Chameides, 1984), which makes H$_2$O$_2$ the principal oxidant of SO$_2$ in cloud. Hegg (1985) suggested cloud processing to be the most important mechanism for the conversion from SO$_2$ to SO$_4^{2-}$. Because the reaction rate between SO$_2$ and O$_3$ is greatly accelerated at high pH
(Hoffmann, 1986), some authors postulated O$_3$ oxidation in sea-salt aerosols to be an important sink of SO$_2$ and source of SO$_4^{2-}$ mass (Sievering et al., 1991; Faloona et al., 2010). However, both the uptake of SO$_2$ and the production of SO$_4^{2-}$ increase the acidity of the sea-salt, making the O$_3$ reaction self-limiting.

In addition to affecting atmospheric chemistry, SO$_4^{2-}$ aerosols are climatically important because they alter the global radiative balance directly by scattering light (Charlson et al. 1992) and indirectly by controlling the optical properties, areal extent, and lifetimes of clouds via the formation of cloud condensation nuclei (CCN). At a certain level of water vapor supersaturation, aerosols larger than a critical size and with a soluble composition (e.g. SO$_4^{2-}$) act as CCN and activate to form cloud droplets. Pruppacher and Klett (1997) estimated a supersaturation level of ~0.35% for typical marine clouds with an updraft velocity of ~1 m s$^{-1}$, corresponding to a critical diameter of approximately 70~80 nm for activation. In a remote marine environment where CCN represent a large fraction of the aerosol number, a gap in this size range in the number distribution is often observed, which has been coined the ‘Hoppel minimum’ (Hoppel et al., 1986). Aerosols smaller than the critical diameter are unactivated and sometimes called Aitken particles, while aerosols larger than the critical diameter have been activated and grown into the accumulation mode (0.1~1 um).

Charlson et al. (1987) coined the ‘CLAW’ hypothesis, which postulated a negative feedback loop from enhanced phytoplankton growth and DMS efflux in a warming climate leading to a decrease in incident radiation (and hence cooling) due to greater albedo of marine clouds from more SO$_4^{2-}$ derived CCN. With liquid water content
unchanged, more CCN should increase the cloud droplet number concentration and reduce the mean droplet size, which lead to more reflective and possibly longer living clouds. The effects of naturally derived SO$_4^{2-}$ on clouds are expected to be greatest in areas deprived of CCN (Twomey, 1991), such as over the remote ocean in the relatively pristine Southern Hemisphere.

The Southeast Pacific is a region characterized by large-scale subsidence associated with subtropical anticyclonic flow, coastal upwelling of cold water driven by Ekman transport, and a large stratocumulus cloud deck capped by a strong inversion layer (Bretherton et al., 2004). The stratocumulus topped MBL is shallow near the coast (~ 1 km) with more broken clouds. Away from shore, the MBL deepens with thicker clouds and more extensive precipitation in the form of drizzle. Satellite observations show frequent hundred-kilometer sized openings in the stratocumulus cloud deck. Termed “pockets of open cells” (POCs), these features persist for a timescale of a day and advect with the mean wind (Stevens et al., 2005). The geographical gradient in cloud properties as well as the formation and evolution of POCs likely depend on the availability of aerosols that can act as CCN and suppress drizzle, and drizzle that in turn removes said aerosols. The coastal regions of Chile and Peru are characterized by pollution emissions of SO$_4^{2-}$ and SO$_2$ from fossil fuel consumption and processing of copper ores. Because the prevailing wind direction at the surface is S/SE along the Andes mountain range, anthropogenic influence decreases quickly away from shore, where a greater contribution to the SO$_4^{2-}$ burden and larger fraction of CCN likely originate from DMS.
In this paper, we will report the longitudinal and vertical distributions of SO$_2$ and SO$_4^{2-}$ measured from a ship and an aircraft during Vamos Ocean-Cloud-Atmosphere-Land Study Regional Experiment (VOCALS-REx). Two examples of new particle nucleation observed in the MBL will be discussed. Following up on the DMS budget presented in Yang et al. (2009), we will estimate the budgets of SO$_2$ and SO$_4^{2-}$ using a mass balance approach in the remote region of the Southeast Pacific. The conservation equation for any chemical species, $S$, in the MBL is represented in (4.1), with the overbar representing an average in time and/or space:

$$\frac{\partial S}{\partial t} + \frac{\partial u S}{\partial x} + \frac{\partial S w'}{\partial z} = P - L \quad (4.1)$$

The terms on the LHS are the time-rate of change in the concentration of $S$, horizontal advection of $S$ due to gradients along the mean wind, and vertical flux divergence, respectively. In situ production and loss of $S$ are represented by $P$ and $L$ on the RHS of the conservation equation.

### 4.2 Experimental

During VOCALS-REx, the National Oceanographic and Atmospheric Administration (NOAA) research vessel *Ronald H. Brown (RHB)* made multiple transects between the coastal city of Arica at 70°W and the remote marine region near the Woods Hole Oceanographic Institution (WHOI) Improved Meteorology (IMET) moored buoy at 20°S, 85°W. The ship sailed in two legs: Leg 1 from Panama to the IMET buoy and then to Arica (Oct. 20~Nov. 3), and Leg 2 from Arica to near 85°W and back to Arica (Nov. 10~Dec. 11). In addition to surveying several eddies, *RHB* stationed at 85°W and 75°W
for over 24 hours to more closely observe diel cycles.

The National Science Foundation (NSF)/National Center for Atmospheric Research (NCAR) aircraft C-130 flew 14 research flights (RF) out of Arica between Oct. 15 and Nov. 15. Out of the 14 flights, eight were dedicated to survey the 20°S line to as far as 86°W, four were designed to study the structures and evolution of POCs, and two were parallel to the coast in the N-S direction between 20°S and 30°S for examining continental influence. For the 20°S surveying flights, the aircraft typically flew 10-minute level legs near the surface, in the mid MBL, at cloud level, and above clouds. Sounding profiles from the surface to ~4 km were frequently performed.

4.2.1 RHB

The measurement of airside DMS by an atmospheric pressure ionization mass spectrometer (APIMS) during VOCALS-REx is outlined by Yang et al., (2009). DMS was sampled from the mast of the ship (18 m) along with instantaneous winds and ship motions at a high rate of 20 Hz, which allowed for the eddy covariance computation of the DMS sea-to-air flux. Instrument design, flux computation, and error analysis are detailed by Blomquist et al. (2010). Seawater DMS was taken from the ship’s non-toxic water supply at ~5.5 m below the ocean surface, extracted by a purge-and-trap method, and analyzed by gas chromatography with a sub-nM detection limit (Bates et al., 2000).

Aerosol chemical compositions and size distributions were measured by the Pacific Marine Environmental Laboratory through an isokinetic inlet on the forward deck of the ship. Submicron and supermicron aerosols were collected on a two-stage multi-jet cascade impactor (50% aerodynamic cutoff diameters of 1.1 and 10 μm) over 2–23
hours. In addition to providing gravimetric mass, collected aerosols were analyzed for soluble concentrations of sodium (Na$^+$), potassium (K$^+$), calcium (Ca$^{2+}$), magnesium (Mg$^{2+}$), ammonia (NH$_4^+$), methane sulfonic acid (MSA$^-$), chloride (Cl$^-$), bromide (Br$^-$), nitrate (NO$_3^-$), sulfate, and oxalate (Ox$^{2-}$) using ion chromatography (Bates et al., 2008). A coating of silicon grease on the first impaction stage prevented bouncing of large particles. Non-sea-salt- SO$_4^{2-}$ and Cl$^-$ deficit were calculated by subtracting sea-salt sulfate and Cl$^-$ components from aerosol concentrations. An Aerodyne quadrupole aerosol mass spectrometer (Q-AMS) was used to continuously measure submicron particulate concentrations of SO$_4^{2-}$, NO$_3^-$, NH$_4^+$, and organic matter with detection limits of 0.02, 0.02, 0.15, and 0.16 µg m$^{-3}$ respectively (Hawkins et al., 2010). The AMS measured non-refractory aerosols (those vaporizing at 600°C), thus excluded mineral dust, elemental carbon, and sea-salt. The AMS sulfate measurements agreed exceptionally well with collocated submicron filter measurements of SO$_4^{2-}$ and showed a high degree of correlation with integrated submicron aerosol volume. The number concentration of aerosols greater than ~13 nm (CN) was measured by a TSI 3010 condensation particle counter (CPC). Aerosols greater than ~5 nm (UFCN) were measured by a TSI 3025 ultrafine CPC and a water based TSI counter. Aerosol size distributions from 0.02~10 µm were obtained by a merging spectra from a differential mobility particle sizer (DMPS) and an aerodynamic particle sizer (APS) at a regulated RH of 60% (Bates et al., 2008). Aerosol scattering and ozone (O$_3$) were measured by a 3-channel TSI 3563 nephelometer and a Thermo Environmental Instruments Model 49C ozone analyzer.
Vertical structures of temperature and humidity were determined from radiosonde launched several times a day from the ship by the NOAA Earth System Research Laboratory (de Szoeke et al., 2008). During the second half of VOCALS-REx, cloud fraction and cloud top height were determined from a W-band radar. Rain rate was measured by a shipboard optical rain gauge; but considering the difficulty with quantifying drizzle, the measurement should be viewed as qualitative. The mixed layer height (MLH) from the surface was determined by a High Resolution Doppler Lidar (2 μm) from velocity variance (turbulence) profiles and aerosol backscatter gradient (Tucker et al. 2009). Liquid water path ($L_{wp}$), or the amount of liquid water integrated over the MBL column, was estimated using a microwave radiometer by University of Miami (Zuidema et al., 2009).

4.2.2 C-130

$\text{SO}_2$ was measured at 1 Hz using an APIMS by Drexel University following the method described in detail by Thornton et al., (2002). An isotopically labeled $^{34}\text{SO}_2$ was mixed with ambient air at the inlet, enabling the calculation of the ambient concentration of $\text{SO}_2$ from the ratio between ambient and labeled species. Both DMS and $\text{SO}_2$ on the C-130 were sampled from backward facing inlets, which should not be affected by cloud droplets. Thus within clouds, measurements of those gases represent interstitial concentrations.

The University of Hawaii aerosol instruments were typically located behind a low turbulence inlet (LTI) (Huebert et al., 2004). Breaking up of cloud droplets upon collision on the wall of the inlet caused a cloud shattering artifact, or an increase in the
measured number of particles. By excluding such incidents, aerosol data at cloud level were biased towards cloud free regions. A high resolution time-of-flight AMS on the C-130 measured aerosol concentration of $\text{SO}_4^{2-}$, $\text{NO}_3^-$, $\text{NH}_4^+$, and organic matter. As with the AMS on the ship, the AMS on the C-130 sampled with a near-unity efficiency for aerosols up to $\sim0.8$ $\mu$m in diameter and excluded large particles.

Aerosol size distributions from diameters of 0.01 to 10 $\mu$m were obtained by merging spectra from a radial differential mobility analyzer (RDMA, 0.01~0.20 $\mu$m), a TSI long differential mobility analyzer (LDMA, 0.10~0.50 $\mu$m), a PMS laser optical particle counter (OPC, 0.12~8.0 $\mu$m), and a TSI 3321 aerodynamic particle sizer (APS, 0.78~10.0 $\mu$m) (Howell et al., 2006). Measured at near ambient pressure and cabin temperatures, the sample air was dried to a RH of $\sim20\%$ by mixing with desiccated air to remove water from aerosols. Direct comparisons between the distributions measured on the C-130 and RHB can only be qualitative in nature because of the different sampling strategies and airspeeds. At the dried environment of C-130 measurements, MBL aerosols had most likely undergone efflorescence and been approximately halved in diameters, whereas the already dry FT aerosols remained largely unchanged. The RHB distributions, on the other hand, were measured at $\sim60\%$ RH, closer to the ambient RH of $\sim70\%$ and above the humidity threshold for efflorescence; the measured diameters were thus only $\sim10\%$ smaller than ambient diameters. Moreover, the lowest size cut was lower for the C-130 distribution (10 nm) than for the RHB distribution (20 nm); thus more small particles were detected by the instruments aflight. For the coarse mode, the isokinetic inlet on the ship had a higher sampling efficiency than the LTI on the aircraft.
Bulk cloud water was collected by Colorado State University with an Airborne Cloud Collector, which had a 50% size cut diameter of 8 μm and thus collected most cloud drops but excluded small, unactivated aerosols (K. Beam, personal communication, 2010). Cloud water was analyzed for pH, S(IV) by absorbance, peroxides by fluorescence, anions and cations by ion chromatography. Aqueous concentrations in μM can be converted to mixing ratios using cloud liquid water content.

Inversion heights ($Z_i$) determined based on potential temperature and dew point gradients from the C-130 sounding showed excellent agreement with those determined from potential temperature profiles from shipboard radiosondes. Because $Z_i$ varied both with longitude and with the time of day, to average vertical distributions, altitude ($Z$) is normalized to $Z_i$, with $Z/Z_i =1$ indicating the inversion. For FT averages along 20°S, only observations within 1 km above $Z_i$ are included.

4.3 Observations

To evaluate the relative importance of natural and pollution sulfur sources, we separate the VOCALS-REx sampling area longitudinally to the ‘coastal’ region (70°W ~ 73°W) and ‘offshore’ region (73°W ~ 86°W). We further isolate the ‘remote’ region (78°W ~ 86°W) from the offshore region. Figure 4.1 shows a map of the RHB cruise track color-coded by the DMS sea-to-air flux; the dotted and solid lines indicate the offshore and remote regions. This stratification is based on aircraft observation of CO and shipboard measurement of radon, both indicators of continental influence. The lowest MBL CO concentration was found west of 78°W at ~62 ppbv; CO increased to ~73 ppbv near the coast, corresponding to elevated $\text{SO}_4^{2-}$ concentrations. Hawkins et al.
(2010) showed that the concentration of radon (a radioactive decay product of uranium in rocks and soil) in the remote marine region was approximately half of the value by the coast. Three-day back trajectories by those authors showed that east of 75°W, airmasses had been previously in contact with land south of the VOCALS-REx sampling area, near the Chilean capital Santiago. In contrast, west of 78°W, air had originated from the remote South Pacific. Figure 4.2 show a typical 20°S survey flight by the C-130 (RF 03 on Oct 21). The flight track is color-coded by SO$_2$ concentration, with marker size corresponding to SO$_4^{2-}$. The aircraft encountered more polluted air in two flights south of ~22°S, closer to Santiago, as well as on RF 14. Greater continental influence was also observed on the RHB north of ~15°S, closer to Peru. For our ‘20°S’ averages, we limit the latitudinal range to 18°S ~ 22°S and exclude RF 14 from C-130 statistics. Because the aircraft usually took off from Arica in the early morning, reached 80~85°W at around sunrise, and returned to shore in the afternoon, spatial and temporal biases were inherent. We will thus mostly rely on ship observations for diel cycles and aircraft observations for vertical structures.

4.3.1 Mean Concentrations: Coastal vs. Marine

It was hypothesized prior to VOCALS-REx that coastal upwelling of cold, nutrient rich water stimulates growth of phytoplankton, which leads to more seawater DMS and atmospheric DMS by the shore than in the remote region. We only observed enhanced seawater DMS in isolated pockets near the coast. More often high seawater DMS was associated with the edge of an eddy or near a front between two water masses, where temperature and salinity changed markedly. On average along 20°S, seawater
DMS was not substantially different between the coastal and the remote regions. Higher atmospheric DMS concentration and sea-to-air flux were observed away from the coast principally as a result of higher wind speed. At the same air-sea concentration difference in DMS (dictated by the seawater concentration), a higher wind speed generally leads to a greater transfer velocity, and hence more DMS flux out of the ocean.

Project average concentrations of DMS, SO$_2$ and SO$_4^{2-}$ along 20°S and in the MBL and FT are shown in Fig. 4.3. Pollution emission was unquestionably the major source of sulfur mass near shore, with SO$_2$ and SO$_4^{2-}$ concentrations elevated at ~80 and ~800 pptv in the MBL, respectively. Moving offshore towards the remote region, SO$_2$ and SO$_4^{2-}$ decreased rapidly to ‘background’ levels. SO$_2$ varied from flight to flight but was relatively well mixed with a mean (standard deviation) MBL concentration of ~25 (15) pptv, about 40% of the DMS concentration. We show later that the relatively low concentration of SO$_2$ in the MBL was likely due to processing of air by stratocumulus clouds. SO$_4^{2-}$ was exceedingly variable horizontally and vertically, which in part explains the difference between the averaged MBL concentrations from the RHB 118 (85) pptv and C-130 60 (60) pptv in the remote MBL. While generally not very variable with mean remote FT concentrations of ~26 and ~40 pptv, respectively, SO$_2$ and SO$_4^{2-}$ were occasionally elevated immediately above the clouds, as was CO. CO in the FT had a range of 67–77 ppbv, with higher concentrations suggesting sporadic long distance transport of pollution-influenced air. Due to the lack of precipitation, both SO$_2$ and SO$_4^{2-}$ aerosols can have much longer lifetimes in the FT (a few weeks) than in the MBL (a few days), enabling them to be transported from continental sources to offshore regions.
O₃ averaged 26 and 52 ppbv in the offshore MBL and FT, respectively. MBL O₃ displayed a diurnal variability qualitatively similar to that of DMS. O₃ built up at night as a result of entrainment from the FT, whereas during the day, photochemical and heterogeneous reactions outpaced entrainment and led to a steady decrease in O₃. Dry deposition to ocean surface represents a small but continuous sink for O₃ at all times. As with the DMS budget (Yang et al., 2009), we can estimate the entrainment velocity ($\omega_e$) by balancing the nighttime O₃ build up with entrainment and deposition fluxes, assuming no chemical source and loss at night. Assuming a deposition velocity of 0.01 cm s⁻¹ (derived from eddy covariance measurements of O₃ during the Stratus 06 cruise in the same region, J. Hare, personal communication, 2010), we calculate a nighttime $\omega_e$ of ~6 mm s⁻¹, higher than $\omega_e = 4$ mm s⁻¹ that was derived from the DMS budget. Using a higher deposition velocity would require a greater $\omega_e$ to balance the O₃ time-rate-of-change. However, unlike the E-W orientated horizontal gradient in DMS, O₃ was generally higher in the north (~30 ppbv at 19°S) than in the south (~20 ppbv at 21°S), implying a gradient along the mean wind direction. Thus the advective flux of O₃ could be substantial and complicate the interpretation of $\omega_e$ from the O₃ budget.

4.3.2 Vertical Distributions

Lilly (1968) first theorized that longwave radiative cooling at the cloud-top causes convective instability and generates turbulence in a stratocumulus capped MBL. Liquid water content ($L_{wc}$) averaged 0.2 g m⁻³ in the ~300 m deep stratocumulus clouds during VOCALS-REx. Immediately above the cloud deck, a strong inversion layer separates the
MBL from the stable, dry, and relatively quiescent FT. With jumps of 10 and 20 K in potential temperature and dew point over a few tens of meters, the inversion layer confines surface-derived scalars to the MBL. Turbulence at the inversion irreversibly entrains air from the FT into the MBL, with the rate of entrainment varying with radiative fluxes at the stratocumulus cloud-top. At night, longwave cooling causes greater convective instability, more turbulent mixing in the MBL, and increased entrainment. In VOCALS-REx, mixed layer height measured by the shipboard lidar indicated a coupled (well-mixed) MBL at night. During the day, shortwave heating of the cloud-top partially suppresses turbulence, which reduces entrainment, causing the surface layer and the upper MBL to become decoupled (Nicholls, 1984). Evaporative cooling from falling drizzle droplets may further affect the coupling of the MBL. Lidar measurements showed that the velocity variance in the MBL was ~50% lower during the daytime, with the mixed layer height (MLH) only reaching 60~70% of $Z_i$ in the remote region, generally below the stratocumulus cloud bottom but above the lifting condensation level (LCL). While a cloud base well above the LCL is usually thought to indicate decoupling, the MLH measurement provides more direct information on the degree of mixing. Turbulence was still present, but more intermittent above the MLH. The extent of decoupling also depends on the MBL depth; as the MBL deepens further to the west (~1.6 km at 85°W) from the coast (~1 km at 70°W), the frequency and degree of decoupling increase.

Near the surface (e.g. at the ship’s level), the concentration of a tracer produced from the upper MBL or entrained from the FT and lost to the ocean surface may decrease during the day due to decoupling that shuts off the source above. Such would be the case
for \( \text{O}_3 \), and for \( \text{SO}_4^{2-} \) if most of it were produced in clouds. In contrast, surface concentration of a tracer with an oceanic source, such as DMS, may increase faster during the decoupled daytime without considering chemical loss. The mean C-130 vertical profiles of temperature, dew point, potential temperature, liquid water content, CO, \( \text{O}_3 \), \( \text{SO}_4^{2-} \), and \( \text{SO}_2 \) in the remote region are shown in Fig. 4.4. \( \text{SO}_2 \) and \( \text{SO}_4^{2-} \) concentrations appeared reduced at cloud level (~18 and 30 pptv), likely in part to due in-cloud oxidation and scavenging by clouds when measurements were taken near cloud outflow regions. Figure 4.5 shows the mean aerosol size distributions (number, surface area, and volume) at ~20% RH from the C-130 in the remote region, stratified by the normalized altitude \( (Z/Z_i) \). The low and mid MBL distributions were similar, with the Hoppel minimum at ~70 nm indicating cloud processing. The upper MBL distribution was biased towards cloud free and POC regions, as cloud shattering compromised measurements inside of clouds. The above cloud distribution was a superposition of two modes: a 30 nm mode associated with low CO (< ~64 ppbv) and an 80 nm mode associated with high CO (> ~74 ppbv). The former can be seen as a ‘background’ distribution, whereas the latter is indicative of long distance transport of pollution.

4.3.3 Precipitation

Marine clouds usually undergo many activation-evaporation cycles before aerosols and cloud droplets are deposited to the sea surface as rain. From C-130 radar and lidar measurements, Bretherton et al. (2010) estimated a mean cloud-base drizzle rate of 1~2 mm day \(^{-1} \) over the entire VOCALS project, with 80-90% evaporating before
reaching the surface, agreeing with previous estimates (Bretherton et al. 2004). An array of measurements from the RHB and C-130 all indicated that precipitation was more likely in the late evening to early morning. Cloud fraction determined from the shipboard radiosondes averaged over 90% between midnight and sunrise and decreased to 20~30% in the afternoon. LWP determined from the shipboard ceilometer peaked above 150 g m\(^{-2}\) over the same time span, more than three times the afternoon value (Fig. 4.6). The optical rain gauge on the ship detected sporadic precipitation events with some rates over ~1 mm hr\(^{-1}\), but typically at a lower value of ~0.2 mm hr\(^{-1}\). Precipitation was more intense and frequent offshore, with most of the detectable rain events taking place during Leg 1 and west of 80ºW.

4.3.4 Diurnal Cycles

SO\(_4^{2-}\) concentration measured by the AMS on the RHB displayed large diel variability, as exemplified by the period from Oct. 28 to 31 when the ship was steaming eastward at 11 knots from 85ºW to 75ºW (Fig. 4.7). SO\(_4^{2-}\) increased from 20~30 pptv to over 150~300 pptv over the first few hours after sunset, peaking at around 0400 UTC (1 am local), with O\(_3\) showing a similar trend. SO\(_4^{2-}\) then declined over the rest of the day, with the sharpest decrease in the early morning, when the frequency and intensity of drizzle peaked. Precipitation with rates of ~0.3 mm hr\(^{-1}\) were recorded on the ship at night and during the day on Oct. 28 and 29. The MBL alternated between being coupled at night and decoupled during the day, as evidenced from the difference between the \(Z_i\) and MLH. The rapid nighttime increase in SO\(_4^{2-}\) is in contrast with previous measurements in the Equatorial Pacific (Huebert et al. 1996), which shows a daytime
building up of SO$_4^{2-}$ from SO$_2$ oxidation by OH. Moreover, the observed build up in SO$_4^{2-}$ is much too fast to be caused by any chemical mechanisms alone. As O$_3$ was increasing at the same time, the most likely explanation evokes a source of SO$_4^{2-}$ in the upper MBL, such as the in-cloud oxidation of SO$_2$. This source did not add to the surface concentration in the daytime due to decoupling of the MBL. As the MBL became re-coupled after sunset, air rich in SO$_4^{2-}$ was mixed downwards, increasing the surface concentration. Precipitating droplets that evaporate while falling might have sped up the transport of SO$_4^{2-}$ from cloud level to the surface. Unfortunately, per flight plans, the C-130 did not sample the remote region at this time of the day to offer any direct evidence.

The observed diel variability in SO$_4^{2-}$ is consistent with size distribution measurements on the RHB, which are stratified according to the time of day in Fig. 4.8. In the remote region, the greatest accumulation mode was measured in the evening (2200~0600 UTC), whereas the coarse mode appeared to be the largest in the late afternoon (1400~2200 UTC), presumably due to high winds that generate more sea-salt aerosols.

4.3.5 New Particle Nucleation

During the day, H$_2$SO$_4$(g) formed from the OH oxidation of SO$_2$ usually condenses on the surfaces of preexisting aerosols and grows them to larger sizes. The total SO$_4^{2-}$ aerosol mass is increased by this process, but the aerosol number remains largely unchanged. Under specific conditions (usually high H$_2$SO$_4$(g) and H$_2$O(g) and low aerosol surface area), however, H$_2$SO$_4$(g) may undergo gas-to-particle conversion and nucleate new nm-sized particles, resulting in a sudden and dramatic increase in the number
concentration (Weber et al., 1999). These freshly formed particles grow by condensation of vapor and coagulation by Brownian motion to a detectable size of ~3 nm in minutes to an hour (Weber et al., 1999) and to 10 nm with in a few hours (Hegg et al., 1992). Nucleation events have been observed in clean air high in the free troposphere (Clarke 1998b), near cloud outflow regions (Hegg et al., 1990; Perry and Hobbs, 1994), in the remote marine boundary layer (Hoppel et al., 1994), and adjacent to a region of precipitation (Clarke et al., 1998a).

In VOCALS-REx, nucleation events were detected on numerous occasions on both the C-130 and RHB as evidenced by elevated concentrations of small particles. On the aircraft, we categorize recent nucleation events by a greater abundance of particles with diameter (at 20% RH) under 30 nm relative to over 30 nm; these events are averaged for the remote region and plotted in Fig. 4.5. The number of particles from 10~30 nm was 390 cm$^{-3}$ at times of recent nucleation, much higher than the low MBL average under normal conditions (87 cm$^{-3}$); the submicron and total aerosol surface areas (3.4 and 5.0 $\mu$m$^2$ cm$^{-3}$, respectively) were significantly reduced during nucleation compared to normal (17.1 and 22.7 $\mu$m$^2$ cm$^{-3}$). The coarse distributions were similar between recent nucleation and the upper MBL, consistent with the idea that new particle formation occurred near cloud level. C-130 profiles also showed enhanced small particles at cloud level that were volatile. Clouds scavenge aerosols, and also pump moisture and aerosol precursor gases into the surroundings. In the outflow region, elevated actinic flux enhances OH and hence H$_2$SO$_4$(g) production. Mixing of cloud outflow air devoid of particles and adjacent air rich in H$_2$O$_{(g)}$ and H$_2$SO$_4$(g) creates an ideal condition for nucleation (Clarke et al, 1998a). CO during these recent nucleation events was very low.
at ~61 ppbv, similar to the remote MBL concentration, suggesting that the source of H$_2$SO$_4$(g) was likely natural.

On the RHB, we identify times of recent nucleation by looking for high UFCN:CN ratio and greater relative abundance of particles under 35 nm diameter (at 60% RH) from size distributions. As UFCN and CN represented aerosol numbers greater than ~5 and ~13 nm respectively, the difference between the two correspond to recently nucleated aerosols in the size range of 5~13 nm. The mean distribution for recent new particle formation in the remote region is plotted in Fig. 4.8. For these nucleation events, the number of particles in 20~35 nm range was 250 cm$^{-3}$, much higher than 40 cm$^{-3}$ normally; the submicron and total surface areas were 10.4 and 25.5 μm$^2$ cm$^{-3}$, respectively, as compared to the typical 23.5 and 37.8 μm$^2$ cm$^{-3}$. While the coarse mode during nucleation was much smaller compared to normal in the C-130 distributions, this distinction was not observed in the distributions from the RHB. The ship was likely detecting particles recently nucleated at cloud level that had been subsequently mixed down to the surface.

Two examples of recent nucleation events observed on the RHB are shown as time-series in Fig. 4.9 and Fig. 4.10. The first took place on Oct. 24, when the ship was near 85ºW and 16ºS and steaming south at 11 knots. The submicron particle area decreased quickly from ~20 μm$^2$ cm$^{-3}$ to ~13 μm$^2$ cm$^{-3}$ by 0500 UTC (2 am local time), when we began to detect small particles. Over 3 hours (a distance of 60 km), the number of particles in the 5~13 nm range was consistently 200~300 cm$^{-3}$. The ship likely encountered a different airmass at 0800 UTC, coinciding with a precipitation event; the
wet scavenging of nucleation mode aerosols is very efficient due to diffusion. Further evidence of freshly formed particles, albeit less dramatic, can be seen in few hours following 0900, as the aerosol surface area continued to decline. Interestingly, our detection of recent nucleation occurred largely in the absence of light, or H$_2$SO$_4$(g) production (sunrise was not until 1000 UTC). Given the brief lifetime of H$_2$SO$_4$(g) due to its low volatility, the gas-to-particle conversion had likely taken place in the daytime of Oct. 23 rather than during 0500–0800 UTC of Oct. 24. The relatively high temperature and low RH on Oct. 24 were also not favorable for in situ nucleation. Lacking H$_2$SO$_4$(g) for condensation, the recently formed particles grew very slowly in the nighttime from coagulation alone such that many of them were still under 13 nm by 0800 UTC. With the ship closer to land at 16ºS than at 20ºS, radon was ~30% higher during this period than in the VOCALS remote region. Thus we cannot rule the possibility that the source of H$_2$SO$_4$(g) for nucleation and condensational growth was partly anthropogenic.

For our second example of nucleation, the ship was largely stationed at 85ºW and 20ºS on Oct. 27 (Fig. 4.10). Geostationary Satellite (GOES) images showed that a pocket of open cell (POC) was advecting over the area from approximately 0800 to 2200 UTC. CN steadily decreased from ~600 cm$^{-3}$ at 1000 UTC to ~100 cm$^{-3}$ by 1600 UTC, along with an increase in O$_3$. As O$_3$ started to decrease at 1700 UTC, both CN and UFCN increased rapidly in a few bursts until 1900 UTC, with a number concentration of ~250 cm$^{-3}$ in the 5–13 nm size range. The water mixing ratio also increased during these bursts of new particles, suggesting that the ship was in the downdraft region of the clouds and nucleation took place near cloud level. The submicron particle surface area was very
low compared to normal, likely in part due to precipitation scavenging. The surface area actually increased from 2 to 8 μm² cm⁻³ while CN decreased, which was a result of the dramatic loss of very small particles due to preferentially enhanced scavenging and growth of the accumulation mode by condensation and coagulation. Light drizzle events were visually observed on multiple occasions, as was the formation of new stratus clouds. Liquid water path, air temperature, RH, and dew point all showed periodicity on a timescale of ~30 minutes, which may be reflecting intermittent turbulence and cell structures. Measurements from the shipboard lidar also indicated decoupling, as at 1700 UTC the ~700 m deep MLH was only about half of $Z_i$. Similar to cloud scavenging, precipitation removes aerosols and can create favorable conditions for nucleation. As this recent nucleation event was observed in the early afternoon when the actinic flux peaked, there should be ample in situ production of H₂SO₄(g), so that the gas-to-particle conversion likely took place recently in the POC. Both SO₄²⁻ and radon were very low, while DMS peaked over 100 pptv for this day, suggesting that the source of H₂SO₄(g) was most likely natural. Interestingly, the seawater DMS concentration was ~50% higher in the POC, where the solar flux was also ~30% higher, compared to the closed-cell (cloudy) days before and after.

4.4 Discussion

We focus our budget calculations on the remote region to examine the natural cycling of sulfur in the marine atmosphere. In the following sections, we will first briefly introduce and approximate each of the rates in the budget equations of SO₂ and SO₄²⁻. At
the end, initiated with observations, we will calculate the diurnal cycles of $\text{SO}_2$ and $\text{SO}_4^{2-}$ based on those rates and adjust the least certain terms if necessary to achieve steady state mass balance. Through multi-variate fitting, Yang et al. (2009) showed that the horizontal gradient in atmospheric DMS concentration mostly lay in the E-W direction, orthogonal to the prevailing wind along the Andes (~150°); the advective flux of DMS was thus negligible. While DMS was not measured in the upwind region of the Southeast Pacific during the campaign, previous observations of seawater DMS in the South Pacific (e.g. Lee et al., 2010) were similar to those measured during VOCALS-REx, implying that the biogenic sulfur source strength in the upwind region was likely not very different. Analogous multi-variate analysis of $\text{SO}_2$ and $\text{SO}_4^{2-}$ concentrations in the remote MBL over the entire project resulted in similarly small advective fluxes on average because gradients in $\text{SO}_2$ and $\text{SO}_4^{2-}$ were also largely longitudinal. We will therefore ignore advection in our budgets below.

4.4.1 Sulfur Dioxide Budget

Observations from the Equatorial Pacific first demonstrated a quantitative anti-correlation between DMS and $\text{SO}_2$ as a result of the oxidation of DMS by the OH radical (Bandy et al., 1996). During the day, $\text{SO}_2$ is produced from DMS faster than it is photochemically destroyed in the MBL. Thus it is no surprise that MBL $\text{SO}_2$ concentration from the C-130 was greater during the day than at night in the remote region in VOCALS-REx. But due to the temporal and spatial biases, the aircraft observations might not accurately represent the actual diurnal profile of $\text{SO}_2$. It is thus informative to examine the budget of $\text{SO}_2$ by integrating (4.1) from the surface to the
inversion height. The angular brackets denote column integrals, with every term now in units of flux (e.g. μmoles m$^{-2}$ day$^{-1}$):

$$\frac{\partial \langle SO_2 \rangle}{\partial t} = F_{SO_2,0} - F_{SO_2,Z} + \langle P_{SO_2} \rangle - \langle L_{SO_2,OH} \rangle - \langle L_{SO_2,\text{cloud}} \rangle - \langle L_{SO_2,\text{aero}} \rangle$$  \hspace{1cm} (4.2)

The term on the LHS of (4.2) is the time-rate-of-change in column concentration. The flux divergence is separated into the first and second terms on the RHS, representing the SO$_2$ flux at the ocean surface (deposition) and at the inversion (entrainment). $P_{SO_2}$ is in situ production of SO$_2$ from DMS, while the last three terms represent losses due to OH, in clouds, and in aerosols, respectively.

4.4.1.1 SO$_2$ Dry Deposition

The surface flux, or the dry depositional loss, of SO$_2$ to the ocean may be related to the SO$_2$ concentration near the surface ([SO$_2$,0]) and a deposition velocity ($V_{dep}$):

$$F_{SO_2,0} = -V_{dep}[SO_2,0]$$  \hspace{1cm} (4.3)

The seawater concentration is irrelevant here for the air-sea exchange of SO$_2$ due to its high reactivity in the seawater (Fairall et al., 2007). At a typical VOCALS-REx wind speed of 6 m s$^{-1}$, models based on atmospheric resistance predict $V_{dep}$ $\approx$ 5 mm s$^{-1}$ (Wesely and Hicks, 1977; Fairall et al., 2007). Aircraft eddy covariance measurements from a recent Equatorial project yielded a somewhat lower $V_{dep}$ of 3~4 mm s$^{-1}$ at this wind speed (Faloona et al. 2010). With $V_{dep} = 4±1$ mm s$^{-1}$ and [SO$_2$,0] = 35 pptv, dry deposition flux was 0.6±0.1 μmole m$^{-2}$ day$^{-1}$, or 0.4±0.1 pptv hr$^{-1}$ for a 1.3 km deep MBL. We will show later that dry deposition was a minor term in the SO$_2$ budget, such that the uncertainty in $V_{dep}$ does not significantly alter our conclusions.
4.4.1.2 \( \text{SO}_2 \) Entrainment

The entrainment of \( \text{SO}_2 \) may be formulated as the entrainment velocity, \( \omega_e \), multiplied by the concentration jump in \( \text{SO}_2 \) across the inversion (Lilly, 1968), with \( Z_i^- \) and \( Z_i^+ \) indicating below and above \( Z_i \):

\[
F_{\text{SO}_2,zi} = \omega_e \{[\text{SO}_2]_{Z_i^-} - [\text{SO}_2]_{Z_i^+}\} \tag{4.4}
\]

From the nighttime budget analysis of DMS in the offshore region, Yang et al. (2009) estimated an \( \omega_e \) of \( \sim 4 \text{ mm s}^{-1} \), same as the diel mean estimated from climatological and mixed-layer models for this region (Wood and Bretherton, 2004; Caldwell et al. 2005). Using the Weather and Research Forecast model and solving for the change of MBL depth over time, Rahn and Garreaud (2010) estimated a similar mean \( \omega_e \) value as the residual for VOCALS-REx.

With a mean of \( \sim 25 \text{ pptv} \) from the C-130, the concentration of \( \text{SO}_2 \) in the remote MBL was higher in the day (30–50 pptv) than before sunrise (\( \sim 20 \text{ pptv} \)) due to DMS oxidation. The concentration at cloud level was reduced to \( \sim 18 \text{ pptv} \), likely due to a combination of in-cloud oxidation and incorporation of FT air by entrainment. At \( \sim 26 \text{ pptv} \), \( \text{SO}_2 \) in the FT likely originated from long distance transport of natural or anthropogenic emissions. Using a mean \( \omega_e \) of \( 4 \text{ mm s}^{-1} \) and the concentration jump between the FT and the cloud level (26 and 18 pptv), entrainment increased MBL \( \text{SO}_2 \) concentration at an approximate rate of 0.1 \( \mu \text{mole m}^{-2} \text{ day}^{-1} \), or \( \sim 0.1 \text{ pptv hr}^{-1} \).

However, as entrainment is a unidirectional process, the ‘mass flux’ of \( \text{SO}_2 \) mixed into the MBL from the FT does not depend on the concentration below the inversion.
Multiplying 4±2 mm s\(^{-1}\) by the FT concentration then yields a SO\(_2\) mass flux of ~0.3 ±0.2 μmole m\(^{-2}\) day\(^{-1}\), all of which would be converted to SO\(_4^{2-}\) aerosols, mostly in clouds as we will show later.

**4.13. SO\(_2\) Production from DMS**

In the marine boundary layer (MBL), DMS is principally oxidized through the reaction with the hydroxyl radical (OH) in two separate pathways: H-atom abstraction and OH-addition (Yin et al., 1990). The abstraction channel is favored at high temperatures and essentially all leads to SO\(_2\), while the addition channel is favored at low temperatures and leads to methane sulfonic acid (MSA), dimethylsulfoxide (DMSO), and also SO\(_2\). From DMS budget, Yang et al. 2009 estimated an effective OH concentration of 1.4±0.2 x 10\(^6\) OH molecules cm\(^{-3}\) on a diel mean, peaking at ~5 x 10\(^6\) OH molecules cm\(^{-3}\) at noontime. Representing the conversion efficiency from DMS to SO\(_2\) as \(\gamma\), the in situ production of SO\(_2\) is the DMS flux divergence multiplied by \(\gamma\), or:

\[
\langle P_{SO_2} \rangle = \gamma (k_{OH + DMS} [OH] \langle DMS \rangle)
\]  

(4.5)

Here \(k_{OH + DMS}\) is the total second order rate constant of the DMS-OH reaction (sum of the abstraction and addition channels). At an average MBL temperature of ~13°C during VOCALS-REx, \(k_{OH} \approx 7.9 \times 10^{-12}\) cm\(^3\) molec\(^{-1}\) sec\(^{-1}\), with the abstraction pathway accounting for 60% of the total oxidation (Sander et al., 2006). The coefficient \(\gamma\) typically ranges from 0.6~0.9, and is recommended to be 0.7(±0.2) by Faloona (2009).

MSA largely resides in the particulate phase; in VOCALS-REx, most of the MSA was found in the fine mode according to filter measurements. Since MSA is entirely
produced from DMS whereas $SO_4^{2-}$ can have both natural and anthropogenic origins, the MSA:SO$_4^{2-}$ ratio can be used to infer the natural contribution to the sulfur burden. In the Southern Hemisphere, a strong correlation was observed between the submicron MSA:SO$_4^{2-}$ ratio and DMS emission (Bates et al. 1992). The submicron MSA:SO$_4^{2-}$ ratio in this study varied between 0.04~0.15 and very closely followed DMS concentration (Fig. 4.11). The ratio was high when DMS peaked and was the lowest at the coast. In an environment free of anthropogenic influence, a higher MSA:SO$_4^{2-}$ ratio may also be interpreted to indicate greater importance of the addition channel of DMS oxidation (Yin et al., 1990). The mean ratio for the fine mode (0.08) and total (0.11) during VOCALS-REx were higher than in the tropical Pacific (R. Simpson, personal communication, 2010) likely in part because of the lower temperature in the Southeast Pacific. If MSA was the only other major oxidation product of DMS besides SO$_2$, and if all of the SO$_4^{2-}$ was formed from DMS, assuming equal lifetime for MSA and SO$_4^{2-}$ due to precipitation scavenging, a MSA:SO$_4^{2-}$ ratio of 0.1 implies $\gamma \approx 0.9$. Accounting for DMSO, the other major product from the addition channel but not measured during the campaign, would reduce $\gamma$. Using a range of 0.7~0.9 in $\gamma$ led to a diel average SO$_2$ gas phase production of 2.2~2.8 $\mu$moles m$^{-2}$ day$^{-1}$, or 1.8~2.3 pptv hr$^{-1}$. In midday, the SO$_2$ production rate from DMS peaked at 7.5~9.7 $\mu$moles m$^{-2}$ day$^{-1}$, or 6.1~7.8 pptv hr$^{-1}$.

4.4.1.4 SO$_2$ Gas Phase Oxidation

The rate of daytime oxidation of SO$_2$ by OH is:

$$\langle L_{SO_2,OH} \rangle = k_{OH+SO_2}[OH][SO_2]$$  \hspace{1cm} (4.6)
The second order rate constant between OH and SO\textsubscript{2}, \(k_{\text{OH-SO}_2}\), is about \(9.6 \times 10^{-13}\) cm\(^3\) molec\(^{-1}\) sec\(^{-1}\) at \(\sim 13^\circ\text{C}\) (Sander et al., 2006). At a SO\textsubscript{2} concentration of 35 pptv, the diel loss rate was 0.22 μmoles m\(^{-2}\) day\(^{-1}\), or 0.17 pptv hr\(^{-1}\). The greatest SO\textsubscript{2} loss occurred at a rate of 0.9 μmoles m\(^{-2}\) day\(^{-1}\), or 0.7 pptv hr\(^{-1}\), not during midday, but in the late afternoon.

**4.4.1.5 SO\textsubscript{2} Aqueous Phase Oxidation – In Cloud**

SO\textsubscript{2} may be removed from the MBL by reacting with H\textsubscript{2}O\textsubscript{2} and O\textsubscript{3} in cloud droplets or in sea-salt aerosols. With extensive cloud coverage in the Southeast Pacific, in-cloud oxidation of SO\textsubscript{2} is expected to be a major source of SO\textsubscript{4}\textsuperscript{2-} mass. Figure 4.12 shows SO\textsubscript{2} concentration in the MBL vs. liquid water content (\(LWC\)) for the offshore region. Compared to the concentration below cloud or in the clear sky MBL, the interstitial SO\textsubscript{2} concentration was reduced to \(\sim 18\) pptv, with perhaps slightly higher concentration during the day than at night. In an aqueous solution, SO\textsubscript{2} is equilibrated to SO\textsubscript{2}.H\textsubscript{2}O, HSO\textsubscript{3}\textsuperscript{-}, and SO\textsubscript{3}\textsuperscript{2-} (together S(IV)) and the equilibrium reactions are pH dependent:

\[
[SO\textsubscript{2} \cdot H\textsubscript{2}O] = H_{SO\textsubscript{2}}[SO\textsubscript{2}] \tag{4.7a}
\]

\[
[HSO\textsubscript{3}^-] = \frac{K_{s1}[SO\textsubscript{2} \cdot H\textsubscript{2}O]}{[H^+]} = \frac{H_{SO\textsubscript{2}}K_{s1}[SO\textsubscript{2}]}{[H^+]} \tag{4.7b}
\]

\[
[SO\textsubscript{3}^-] = \frac{K_{s2}[HSO\textsubscript{3}^-]}{[H^+]} = \frac{H_{SO\textsubscript{2}}K_{s1}K_{s2}[SO\textsubscript{2}]}{[H^+]^2} \tag{4.7c}
\]

The Henry’s coefficient \(H_{SO\textsubscript{2}} =1.23\) M atm\(^{-1}\) as well as equilibrium constants \(K_{s1}=1.3\times10^{-2}\) M and \(K_{s2}=6.6\times10^{-8}\) M at 298 K are all higher at lower temperatures with \(\sim \Delta H/R\) of 3135, 1960, and 1495 K, respectively (Hoffmann and Calvert, 1985). \(K\) at the ambient
temperature is related to the equilibrium constant at 298 K \( (K_{298}) \) by the relationship \( K = K_{298} \exp[-\Delta H/R (1/T – 1/298)] \).

In the remote region, with an average cloud water \( \text{H}_2\text{O}_2 \) concentration of 65 \( \mu \text{M} \) (~0.9 ppbv in gas phase), stoichiometrically there was enough oxidant to oxidize all of the S(IV) to SO_4^{2-}. Bulk cloud water measurements showed mean SO_4^{2-} concentration of ~45 \( \mu \text{M} \) in the remote region, which for \( L_{WC} \approx 0.2 \text{ g m}^{-3} \) converts to ~200 pptv, somewhat higher than the below cloud concentration. S(IV), on the other hand, was barely detectable in cloud water. Cloud water had an average pH of 4.2 (typically 3.5~6), consistent with the ensemble mean pH of 4.3 in stratocumulus droplets from a recent a survey of measurements over the last twenty years (Faloona, 2009). At this pH, HSO_3^- is the predominant S(IV) species.

Ignoring mass transfer limitation, the loss rate of S(IV), and hence the production rate of SO_4^{2-}, by reaction with \( \text{H}_2\text{O}_2 \) is taken from Martin and Damschen, (1981) [M s\(^{-1}\)]:

\[
- \frac{d[S(IV)]_{H_2O_2}}{dt} = k_{H_2O_2}[H^+][H_2O_2][HSO_3^-] \tag{4.8a}
\]

\[
k_{H_2O_2} = \frac{2.1 \times 10^6}{0.1 + [H^+]} \tag{4.8b}
\]

The rate constant \( k_{H2O2} \) is in units of M\(^{-2}\) s\(^{-1}\) and decreases at lower temperatures with \( -\Delta H/R = -3650 \) (Maahs, 1983). The reaction is largely pH insensitive because the pH dependences in the equilibrium of HSO_3^- and in the \( \text{H}_2\text{O}_2 \) reaction cancel. With a gas phase SO_2 concentration of 25 pptv, cloud water \( \text{H}_2\text{O}_2 \) concentration of 65 \( \mu \text{M} \), and pH of 4.2, the instantaneous loss rate of S(IV) due to \( \text{H}_2\text{O}_2 \) is \( 7.2 \times 10^{-10} \text{ M s}^{-1} \). An alternate reaction rate, which is about three times higher than the one from Martin and Damschen (1981), was presented by Hoffmann and Calvert (1985). We demonstrate later that this
alternate rate appears to be unrealistically high from the perspective of the steady state mass balance of SO$_2$. However, the large discrepancy in this reaction rate contributes to the uncertainty of our budget estimates.

The loss rate of S(IV) by reaction with O$_3$ [M s$^{-1}$] is:

$$-\frac{d[S(IV)]_{O_3}}{dt} = (k_{O_3,0}[SO_2 \cdot H_2O] + k_{O_3,1}[HSO_3^-] + k_{O_3,2}[SO_3^{2-}])[O_3]$$

(4.9)

where $k_{O3,0}$, $k_{O3,1}$, and $k_{O3,2}$ are $2.4 \times 10^4$, $3.7 \times 10^5$, and $1.5 \times 10^9$ M$^{-1}$ S$^{-1}$ at 298 K, respectively (Hoffmann, 1986). Because $k_{O3,2}$ is orders of magnitude higher than the other two reactions, the overall reaction rate approximately varies inversely with the square of [H$^+$] and is greatly enhanced at high pH when there is more SO$_3^{2-}$. At the average cloud pH and gas phase concentrations of 26 ppbv and 25 pptv for of O$_3$ and SO$_2$, the instantaneous loss rate of S(IV) due to O$_3$ is $\sim 4 \times 10^{-12}$ M s$^{-1}$, two orders of magnitude slower than the H$_2$O$_2$ reaction.

The aqueous phase oxidation rate of S(IV) in cloud [M s$^{-1}$] can be converted to an equivalent gas phase equivalent rate of SO$_2$ loss [pptv hr$^{-1}$] using $L_{WC}$:

$$L_{SO_2,cloud} = 3.6 \times 10^9 L_{WC} RT(-\frac{d[S(IV)]_{H_2O_2}}{dt} - \frac{d[S(IV)]_{O_3}}{dt})$$

(4.10)

$R$ is the gas constant (0.082 atm L K$^{-1}$ mole$^{-1}$), and $T$ is the average cloud temperature ($\sim 282$ K). With 25 pptv of SO$_2$, 65 $\mu$M ($\sim 0.9$ ppbv) of H$_2$O$_2$, and $L_{WC}$ of 0.2 g m$^{-3}$, $L_{SO_2,cloud}$ is 12 pptv hr$^{-1}$ (48% hr$^{-1}$). On average including the O$_3$ reaction affects the in-cloud oxidation rate insignificantly.

However, using the instantaneous $d[S(IV)]/dt$ (i.e. assuming an open system) to budget SO$_2$ loss over time will certainly lead to an overestimation. In a more realistic closed system, the concentration of the limiting oxidant decreases significantly as
reaction proceeds, and so will $d[S(IV)]/dt$. To calculate SO$_2$ depletion as a function of time in cloud over an hour, we start with an initial SO$_2$ concentration of 25 pptv, H$_2$O$_2$ concentration of 0.9 ppb, $L_{WC}$ of 0.2 g m$^{-3}$, and temperature of 282 K. For simplicity, since there is more than sufficient H$_2$O$_2$ to oxidize all of the SO$_2$ and this reaction is largely pH-independent, a constant pH of 4.2 and an O$_3$ concentration of zero are used. Assuming rapid equilibration between the gas phase and aqueous phase (i.e. no mass transfer limitation), at a given minute gas phase SO$_2$ is computed by subtracting $L_{SO_2,cloud}$ / 60 from SO$_2$ from the previous minute. Aqueous concentration of H$_2$O$_2$ is updated by subtracting $d[S(IV)]/dt$ x 60 from the previous time step and equilibrating with the gas phase. As seen in Fig. 4.13, 9.3 pptv (37%) of SO$_2$ is consumed after an hour in cloud, which is 0.8 times the rate calculated from the open system, while the excess H$_2$O$_2$ is only decreased slightly.

From large eddy simulations (LES), Feingold et al. (1998) showed that for a stratocumulus capped MBL, a parcel of air is in clouds for on average ~15 minutes in an hour, or 25% of the time. From Fig. 4.13, 2.8 pptv of SO$_2$ (~11%) is oxidized to SO$_4^{2-}$ after 15 minutes in cloud, or 1.9 pptv if we account for the mean cloud fraction of ~0.67. The SO$_2$ formation rate from DMS is 1.8~2.3 pptv hr$^{-1}$ (for $\gamma$ of 0.7~0.9) on a diel average. Adding of ~0.1 pptv hr$^{-1}$ due to entrainment and subtracting dry deposition of 0.4 pptv hr$^{-1}$ and OH loss of ~0.2 pptv hr$^{-1}$ leaves a daily residual of 1.3~1.8 pptv hr$^{-1}$. Thus the approximated loss rate of SO$_2$ in clouds appears comparable to the production rate. If using the much higher rate constant of the SO$_2$-H$_2$O$_2$ reaction from Hoffmann and Calvert (1985) instead, the in-cloud oxidation of SO$_2$ would far exceed SO$_2$ production from DMS, which is unrealistic and inconsistent with observations. Lacking other
evidence, the rate constant from Martin and Damschen (1981) appears to be more applicable for VOCALS-REx.

In addition to the uncertainty in the SO$_2$-H$_2$O$_2$ rate constant, the calculation above might be subject to caveats due to the simplifications with respect to mass transfer and reaction time in cloud. Mass transfer from interstitial air to a cloud droplet involves diffusion in the gas phase, interfacial transfer from the gas phase to the aqueous phase, and diffusion in the aqueous phase. If the rate of the chemical reaction is faster than any of those steps, a gradient in concentration develops away from the interface and the reaction becomes limited by the rate of mass transfer. Schwartz (1988) showed that even for 30 µm diameter cloud droplets, in-cloud reaction between SO$_2$ and H$_2$O$_2$ does not appear to be limited by mass transfer. The typical mean cloud droplet radius was 10~15 µm in the VOCALS-REx remote region from MODIS images. However, calculations assuming instantaneous equilibrium between gas and aqueous phases should still be considered upper limits. For simplicity, we applied a mean in-cloud residence time instead of a distribution of cloud residence times. We could also account for the times inside and outside of clouds by using the mean $L_{wc}$ for the entire MBL (~0.06 g m$^{-3}$), which is similar to multiplying the cloud level mean $L_{wc}$ of 0.2 g m$^{-3}$ by 25%.

Zhang et al. (1999) modeled the depletion of SO$_2$ within a single cloud pass using LES coupled with chemistry. For the case of low SO$_2$ (excess oxidants) and typical cloud pH, they showed that H$_2$O$_2$ reaction dominates over the O$_3$ reaction due to the orders of magnitude higher aqueous H$_2$O$_2$ concentration. While our calculated depletion of SO$_2$ is slower than what was simulation by Zhang et al. (1999) because they used the H$_2$O$_2$ rate constant from Hoffmann and Calvert (1985), qualitatively the results are
similar. Those authors also suggested that $\text{SO}_4^{2-}$ produced by the oxidation of $\text{H}_2\text{O}_2$ is evenly distributed across the drop-size spectrum and tends to broaden the processed CCN spectrum, which can enhance coalescence and hence drizzle. From our calculation, it appears that in-cloud oxidation was enough to account for essentially all of the $\text{SO}_2$ loss, which would render other sinks insignificant. Nevertheless, we will look at the oxidation of $\text{SO}_2$ in sea-salt aerosols for the sake of completeness.

4.4.1.6 $\text{SO}_2$ Aqueous Phase Oxidation – In Sea-salt Aerosols

As alluded to previously, the oxidation of $\text{SO}_2$ by $\text{O}_3$ only becomes important at $\text{pH} > \sim 6$. While aerosol pH was not directly measured during VOCALS-REx, we can get a sense for the acidity of the aerosol phase by looking at bulk filter measurements on the $\text{RHB}$. From ion balance of aerosols, the anion:cation molar equivalent ratio averaged 1.38 for the submicron mode away from shore, implying acidic aerosols. As 60~70% of the total aerosol surface area was submicron near the ocean surface, the fine mode was acidic due to the uptake of $\text{SO}_2$, $\text{SO}_4^{2-}$, etc. The anion:cation ratio peaked above 2.3 (most acidic) when the ship was near the coast, where $\text{SO}_4^{2-}$ and $\text{NO}_3^-$ levels from pollution were elevated. This longitudinal pattern in acidity is in agreement with shipboard AMS measurements of submicron $\text{SO}_4^{2-}$ and $\text{NH}_4^+$ (Fig. 4.14). $\text{SO}_4^{2-}$ in aerosols is typically titrated by ammonium ($\text{NH}_4^+$), which can have continental and marine sources (Zhuang and Huebert, 1996), to form ammonium bisulfate ($\text{NH}_4\text{HSO}_4$) and ammonium sulfate ($\text{(NH}_4)_2\text{SO}_4$), depending on the extend of neutralization. Beyond ~80°W, the $\text{SO}_4^{2-}:\text{NH}_4^+$ molar ratio was close to the full neutralization limit of 1:2 for ($\text{NH}_4)_2\text{SO}_4$, with additional acidic components likely from nitric and organic acids. That
the fine mode was not very acidic in the remote region might be due to the frequent wet removal by drizzle. Near the coast, the \( \text{SO}_4^{2-}:\text{NH}_4^+ \) ratio exceeded 1:1 (the ratio for \( \text{NH}_4\text{HSO}_4 \)) due to high anthropogenic emission of acidic species.

In the supermicron mode, the anion:cation ratio averaged 1.03 away from shore, less acidic than in the fine mode partly because the alkalinity of sea-salt and the carbonate buffer could initially slow down the increase in acidity from the uptake of acidic vapors. A coarse mode with pH less than 7 can also be inferred from chloride deficit calculations, which is included in Fig. 4.11. Sulfuric and nitric acid vapors react with sea-salt aerosols that are initially rich in sodium chloride, forming hydrochloric acid, which is vaporized to the atmosphere and results in increased acidity and reduced Cl\(^{-}\) in reacted sea-salt (e.g. Keene et al., 1990; McInnes et al., 1994). In the offshore region, a small Cl\(^{-}\) deficit of -0.1 \( \mu \text{g m}^{-3} \) was observed in the coarse mode. A greater deficit of -0.9 \( \mu \text{g m}^{-3} \) was seen near the coast again because of pollution, as evidenced by the much higher aerosol scattering at 550 nm. With low aerosol calcium throughout the entire project (0.1 \( \mu \text{g m}^{-3} \)), the less acidic coarse mode offshore than near the coast was due to lower uptake of acidic species rather than a titration with alkaline dust materials.

We estimate the upper limit sea-salt oxidation rate of \( \text{SO}_2 \) in an analogous fashion to in-cloud oxidation (4.10). The cloud \( L_{\text{we}} \) is replaced with the integrated supermicron aerosol mass concentration derived from the \textit{RHB} size distribution, which averaged ~8 \( \mu \text{g m}^{-3} \). At a pH of 6.0, the \( \text{SO}_2 \) reaction with \( \text{O}_3 \) is a couple orders of magnitude faster than the reaction with \( \text{H}_2\text{O}_2 \). With gas phase \( \text{O}_3 \) and \( \text{SO}_2 \) concentrations of 26 ppbv and 25 pptv, respectively, the instantaneous loss rate of S(IV) due to \( \text{O}_3 \) is ~1.2 \( \times \) 10\(^{-8} \) M s\(^{-1} \), implying a sea-salt \( \text{SO}_2 \) oxidation rate less than 0.01 pptv hr\(^{-1} \), negligible compared to dry
deposition, gas phase oxidation, and in-cloud oxidation. At a higher pH of 6.5, the oxidation rate of SO$_2$ in sea-salt is increased to 0.08 pptv hr$^{-1}$, but still less than half of the gas phase oxidation rate. By neglecting mass transfer and keeping the pH constant, the calculation above represents an overestimation because in real sea-salt aerosols, SO$_4^{2-}$ production and uptake SO$_2$ by diffusion (which could be rate limiting) would acidify the aerosols eventually even with the carbonate buffer, slowing down further O$_3$ reaction. The insignificance of O$_3$ oxidation from the perspective of MBL SO$_2$ budget is consistent with the observation that only 5~6% of the total non-sea-salt SO$_4^{2-}$ was found in the coarse mode during VOCALS-REx per filter measurements. Moreover, this percentage was relatively insensitive to longitude, whereas the pH of sea-salt aerosols likely differed substantially from near shore to the remote marine region, suggesting that mechanisms other than SO$_2$ oxidation (e.g. direct uptake of H$_2$SO$_4$) was responsible for SO$_4^{2-}$ to exist in the coarse mode.

4.4.1.7 SO$_2$ Budget Summary

To more accurately calculate the sources and sinks of SO$_2$, we look at their time-evolutions over the course of a day and compare the implied SO$_2$ diurnal cycle with observations. From (2.2), omitting sea-salt oxidation, the column integrated concentration of SO$_2$ at time $t$ can be written as:

$$\left\langle SO_2 \right\rangle_t = \left\langle SO_2 \right\rangle_{t-1} + (F_{SO_2,0})_{t-1} - (F_{SO_2,Zi})_{t-1} + (P_{SO_2})_{t-1} - (L_{SO_2,OH})_{t-1} - (L_{SO_2,cloud})_{t-1}$$  (4.11)

The last five terms on the RHS represent time-integrated rates from the previous time step. C-130 Observations of SO$_2$ in the remote region were available in the hours of 0800~2000 UTC. We initialize our calculation with the column integrated concentration
at 0900 UTC (6 am local) because that was approximately the time when the aircraft reached the west most location (80~85°W) on most 20°S survey flights. Compared to other hours, observations from 0900 to 1300 UTC should be subject to the least amount of bias because the aircraft usually stayed in the same general area. The DMS diurnal cycle is calculated as in Yang et al. (2009) for the remote region and used to estimate SO$_2$ production assuming conversion efficiencies of 0.7 and 0.9 from DMS to SO$_2$. The loss rates of SO$_2$ are largely first order (i.e. dependent on the SO$_2$ column concentration) and updated at each hour. Over one diel cycle, we consider the budget closed when SO$_2$ column concentration at the end of the calculation matches that at the beginning under the assumptions of steady state and stationarity.

In Yang et al. (2009), a constant entrainment velocity of 4 mm s$^{-1}$ was used for simplicity because entrainment was a small term in the DMS budget. From direct measurements of DMS, O$_3$, and water vapor fluxes, Faloona et al. (2005) estimated higher $\omega_e$ at night in the stratocumulus region off the coast of California due to greater radiatively driven turbulence and cloud coverage than during the day, consistent with findings from Bretherton et al. (2004) in the Southeast Pacific. We can crudely approximate the diel variability in $\omega_e$ for VOCALS-REx by linearizing $\omega_e$ with cloud fraction at each hour relative to the mean cloud fraction (0.67), which leads to 6 mm s$^{-1}$ at night and 2 mm s$^{-1}$ during the day. As seen in Fig. 4.15, entrainment, deposition, and gas phase OH oxidation are all minor losses of SO$_2$. In-cloud oxidation, the largest sink for SO$_2$, is estimated using a fixed H$_2$O$_2$ concentration of 65 µM and constant pH of 4.2 (i.e. an open system). The concentration of SO$_2$ below the inversion is adjusted from the implied time dependent MBL SO$_2$ concentration using the observed gradient from C-130
profiles. The aqueous concentration of S(IV) is equilibrated with the cloud level SO₂ mixing ratio. Instead of using $L_{WC}$ from the C-130 and approximating the time parcels spend in clouds, we use the liquid water path ($L_{WP}$) determined from the ship, which accounted for both cloudy and clear air conditions. For $\gamma = 0.9$, we find that the in-cloud oxidation loss term needs to be multiplied by a coefficient of 0.43 to balance the diel budget of SO₂. This coefficient is necessary for two reasons. First is the overestimation of in-cloud oxidation from using an open system as opposed to the more realistic closed system; the latter is not trivial to simulate over a diel cycle. Secondly, we calculate in-cloud oxidation based on implied SO₂ concentration at cloud level, which has both MBL and FT sources. For the cloud-processing term in the budget of MBL SO₂ concentration, only SO₂ originated from the MBL and lost in clouds should be included. For the lower $\gamma$, the in-cloud oxidation term needs to be further reduced. We see in Fig. 4.15 that the implied SO₂ cycle matches well with observations up to 1300 UTC (a few hours after sunrise). Comparison worsens for the subsequent hours, which is likely in part due to the large spatial variability of the C-130 observations as the aircraft was then often profiling or transiting. The major rates in the MBL SO₂ budget are summarized in Table 4.1 with both choices of $\gamma$.

### 4.4.2 Sulfate Budget

From a island-based experiment in the Equatorial Pacific, Huebert et al. (1996) measured a clear diurnal cycle in $SO_4^{2-}$ aerosols. $SO_4^{2-}$ built up slowly during the day and peaked near sunset, consistent with photochemical production from SO₂. Dilution due to entrainment of FT air was mostly responsible for the nighttime $SO_4^{2-}$ decline. In
contrast to the diurnal trend shown by Huebert et al. (1996), SO$_4^{2-}$ during VOCALS-REx was the lowest in the late afternoon, increased rapidly for the first several hours after sunset, peaked at 0400 UTC (1 am local), decreased steadily and leveled off over the rest of the diurnal cycle. As with SO$_2$, we first look at the magnitudes of the different terms in the MBL integrated SO$_4^{2-}$ budget:

$$\frac{\partial \left[ \text{SO}_4^{2-} \right]}{\partial t} = F_{\text{SO}_4^{2-},0,\text{dry}} + F_{\text{SO}_4^{2-},0,\text{wet}} - F_{\text{SO}_4^{2-},z_i} + \left< P_{\text{SO}_4^{2-}} \right>$$ \hspace{1cm} (4.12)

The term on the LHS is the time-rate of change. The first three terms on the RHS are dry deposition to the surface, wet deposition due to precipitation, and entrainment at the inversion, respectively. The last term represent in situ production of SO$_4^{2-}$ from SO$_2$.

Similar to DMS profiles shown in Yang et al. (2009), average vertical profiles of SO$_4^{2-}$ showed a slightly decreasing concentration with height in the MBL, which can be approximated as a function of $Z/Z_i$ and $\alpha = 0.2$; the latter is the ‘decoupling’ parameter derived from the profile of DMS. We thus adjust the surface SO$_4^{2-}$ concentration by $1 - 0.5 \alpha$ for the column integration. During the decoupled daytime, as SO$_4^{2-}$ above the MLH might be different from that within the MLH, this adjustment is rather crude, but represents our current best estimate in the absence of continuous measurements of SO$_4^{2-}$ above the MLH.

4.4.2.1 SO$_4^{2-}$ Dry Deposition

The surface flux of SO$_4^{2-}$ is related to the concentration near the surface ($[\text{SO}_4^{2-}]_0$):
The deposition velocity of particles, $V_{aero}$, is a strong function of particle size ($D$). Coarse particles deposit orders of magnitude faster due to gravitational settling than accumulation mode particles. At less than 0.02 $\mu$g m$^{-3}$ (~5 pptv), only ~5% of the total non-sea-salt SO$_4^{2-}$ was found in the coarse mode in the offshore region per filter measurements. Aside from the oxidation of SO$_2$ on sea-salt, which we have shown to be insignificant, other potential mechanisms for transferring SO$_4^{2-}$ to the coarse mode might be direct uptake of H$_2$SO$_4$(g) or through drizzle. If precipitating droplets evaporate before falling to the surface, as was the case 80–90% of the time, aerosols would be returned to the MBL, but likely at greater sizes than before being scavenged. While little SO$_4^{2-}$ was measured in the coarse mode, the enhanced $V_{aero}$ of supermicron aerosols suggests that it is still possible for dry deposition to be a noticeable sink for SO$_4^{2-}$. Lacking size distributed SO$_4^{2-}$ information, we rely on shipboard APS aerosol size distribution and coarse SO$_4^{2-}$ mass from filter measurements.

The deposition velocity of coarse aerosols as a function of size is estimated using the model from Slinn and Slinn (1980), with wet diameters adjusted from measured diameters at 60% RH to the ambient RH according to the sea-salt growth curve (Howell et al., 2006). Integrating the mass distribution over supermicron sizes yields a concentration that agrees well with supermicron gravimetric mass from filter measurements (~8 $\mu$g m$^{-3}$), of which ~0.5% was non-sea-salt SO$_4^{2-}$. To approximate size resolved SO$_4^{2-}$, we apply this percentage uniformly to the supermicron mass distribution (i.e. assuming coarse SO$_4^{2-}$ has the same relative distribution as sea-salt aerosols).
Previous impactor measurements in the remote Pacific showed that most of the coarse $\text{SO}_4^{2-}$ is present at 2~3 $\mu$m, or towards the smaller end of the supermicron spectrum (R. Simpson, personal communication, 2010), whereas the distribution of sea-salt tends to be broader. Allocating more $\text{SO}_4^{2-}$ to the larger end of the supermicron spectrum, where the deposition velocity is much higher, could thus overestimate deposition flux.

At the upper size cutoff of the APS (10 $\mu$m at 60% RH, or ~11 $\mu$m at ambient RH), $V_{\text{aero}}$ peaks at ~0.2 cm s$^{-1}$. Multiplying the approximated $\text{SO}_4^{2-}$ distribution by the size dependent $V_{\text{aero}}$ and summing over supermicron sizes leads to a mean surface flux of merely ~0.002 μmole m$^{-2}$ day$^{-1}$. Even with a ten-fold higher $V_{\text{aero}}$, the resultant deposition flux would still be only ~10% of the $\text{SO}_4^{2-}$ production rate from the gas phase oxidation of SO$_2$, and orders of magnitude lower than other terms in the $\text{SO}_4^{2-}$ budget, consistent with the estimate from Rodhe (1978) that the residence time of $\text{SO}_4^{2-}$ is much longer than that of SO$_2$ with respect to dry deposition. This result is unsurprising because the vast majority of the $\text{SO}_4^{2-}$ mass was found in the accumulation mode, where dry deposition is very slow.

4.4.2.2 $\text{SO}_4^{2-}$ Entrainment

The entrainment of $\text{SO}_4^{2-}$ is calculated similarly to that of SO$_2$ (4.4). From average C-130 profiles, with a $\text{SO}_4^{2-}$ concentration below the inversion ~40 pptv less than above, entrainment diluted the MBL at a rate of 0.5(±0.3) μmole m$^{-2}$ day$^{-1}$, or 0.4(±0.2) pptv hr$^{-1}$. The 20~30 pptv of $\text{SO}_4^{2-}$ observed at cloud level was biased towards clear sky, cloud outflow, and POC regions, thus not representative of $\text{SO}_4^{2-}$ concentration below the
inversion. As with SO\textsubscript{2}, considering the mass flux of SO\textsubscript{4}\textsuperscript{2-} at the inversion to be only proportional to the FT concentration (~40 pptv, which did not vary much diurnally), entrainment added 0.5(±0.3) μmole m\textsuperscript{-2} day\textsuperscript{-1} of SO\textsubscript{4}\textsuperscript{2-} to the MBL. Together with the mass flux of SO\textsubscript{2} of 0.3(±0.2) μmoles m\textsuperscript{-2} day\textsuperscript{-1}, the upper limit contribution of anthropogenic sulfur mass from the FT to the MBL is 0.8(±0.4) μmoles m\textsuperscript{-2} day\textsuperscript{-1}, ~20% of the DMS sea-to-air flux.

4.4.2.3 SO\textsubscript{4}\textsuperscript{2-} Production

The in situ production rate of SO\textsubscript{4}\textsuperscript{2-} is derived from the loss rate of SO\textsubscript{2} and includes both gas and aqueous phase oxidations.

\[
\left\langle P_{SO_4^{2-}} \right\rangle = \left\langle P_{SO_4^{2-},OH} \right\rangle + \left\langle P_{SO_4^{2-},cloud} \right\rangle \quad (4.14)
\]

To balance the SO\textsubscript{2} budget in the MBL (4.12), 2.1 μmoles m\textsuperscript{-2} of SO\textsubscript{2} needs to be oxidized in cloud over the entire day, from which we excluded the in-cloud oxidation of SO\textsubscript{2} originated from the FT (to SO\textsubscript{4}\textsuperscript{2-}). Considering the additional FT SO\textsubscript{2} contribution, 4.0 μmoles m\textsuperscript{-2} of SO\textsubscript{4}\textsuperscript{2-} was produced in-cloud over the entire day.

4.24. SO\textsubscript{4}\textsuperscript{2-} Wet Deposition

Because of its high water solubility, SO\textsubscript{4}\textsuperscript{2-} is readily taken up in clouds, particularly for aerosols above the critical size for activation. Coalescence of cloud droplets form precipitation, which removes aerosols both scavenged in-cloud and collected in the column of air below while falling. Figure 4.16 shows SO\textsubscript{4}\textsuperscript{2-} vs. rain rate reaching the surface ($P_d$) measured by the shipboard optical rain gauge, color-coded by
longitude. The heaviest precipitation events (> 1 mm hr\(^{-1}\)) occurred west of 80°W and were associated with the lowest SO\(_4^{2-}\). The average \(P_d\) of 0.01 mm hr\(^{-1}\) in the remote region (including non-precipitating periods) is consistent with the typical cloud bottom drizzle rate of ~1.5 mm day\(^{-1}\) with 15% reaching the surface estimated from radars and lidars by Bretherton et al. (2010). The wet deposition flux of SO\(_4^{2-}\) is the sum of below cloud scavenging and in-cloud scavenging (first and second terms on the RHS below):

\[
F_{SO_4^{2-},0,wet} = F_{SO_4^{2-},0,BC} + F_{SO_4^{2-},0,IC}
\]  \hspace{1cm} (4.15)

The below cloud scavenging rate of SO\(_4^{2-}\) by droplets that reach the surface depends on the average MBL concentration:

\[
F_{SO_4^{2-},0,BC} = -\Lambda_{BC}[SO_4^{2-}]H_{BC}
\]  \hspace{1cm} (4.16)

Here \(\Lambda_{BC}\) is the below cloud scavenging coefficient of SO\(_4^{2-}\), which depends on rain rate and the collection efficiency; the latter is a complex function of the droplet and aerosol size distributions. Andronache (2003) showed that small drizzle droplets are more efficient at scavenging aerosols than larger rain droplets. Scavenging by falling droplets is particularly efficient for coarse particles (by impaction) and for nuclei mode particles (by Brownian motion). In contrast, accumulation size particles lie in the ‘scavenging well’ and have the lowest efficiency. From Seinfeld and Pandis (2006), for typical drizzle diameter of 0.2 mm and 200~300 nm aerosols (mean diameter for SO\(_4^{2-}\) mass), the below cloud scavenging coefficient is about 0.01 hr\(^{-1}\) for a rain rate of 1 mm hr\(^{-1}\).

Thus we estimate \(\Lambda_{BC} [\text{hr}^{-1}]\) as 0.01 \((P_d / 1)\). For a well-mixed MBL, \(H_{BC}\) is the height from the surface to the bottom of a precipitating cloud, which we equate to \(MLH\). This approximation should be valid for the nighttime when the MBL was coupled and
shipboard lidar sensed up to the cloud bottom; it is less robust during the decoupled
daytime, but because precipitation did not reach the surface as frequently then,
uncertainty from using $H_{BC}$ should be fairly small. From (4.16), we calculate a below
cloud SO$_4^{2-}$ wet removal of 0.006 µmole m$^{-2}$ day$^{-1}$ on average and ~0.1 µmole m$^{-2}$ day$^{-1}$
for heavy rain events.

Most of the wet removal of SO$_4^{2-}$ was due to in-cloud scavenging. When cloud
forms in a rising air parcel, SO$_4^{2-}$ aerosols mostly become cloud droplets (nucleation
scavenging) and are later removed by precipitation; a minor fraction of SO$_4^{2-}$ remains
cloud interstitial aerosols. ten Brink et al. (1986) found very low levels of aerosol SO$_4^{2-}$
in cloud interstitial air compared to cloud-free air, suggesting a high efficiency for the
scavenging of SO$_4^{2-}$ by cloud droplets. The in-cloud scavenging of SO$_4^{2-}$ is formulated
similarly to below cloud scavenging, but with in-cloud scavenging coefficient ($\Lambda_{IC}$) and
concentration integrated to $Z_i$:

$$F_{SO_4^{2-},0,IC} = -\Lambda_{IC} \left( SO_4^{2-} \right)$$

(4.17)

Okita et al. (1996) found a bulk scavenging efficiency of 0.5 $P_d^{0.74}$ hr$^{-1}$ from SO$_4^{2-}$
measurements over the Sea of Japan during a winter monsoon. Following the concepts of
in-cloud scavenging from Scott (1982) and constrained by bulk precipitation
measurements from the Eastern United States, Andronache (2004) parameterized the in-
cloud scavenging efficiency of SO$_4^{2-}$ using a Marshall and Palmer (1948) precipitation
size distribution: 1.26 $P_d^{0.78}$ hr$^{-1}$. Utilizing at the moment the parameterization from
Okita et al. (1996), the in-cloud scavenging loss would be ~1.2 µmole m$^{-2}$ day$^{-1}$ and over
25 µmole m$^{-2}$ day$^{-1}$ for the heaviest precipitation events.
4.4.2.5 $SO_4^{2-}$ Budget Summary

To explain the $SO_4^{2-}$ observations from the ship, we first calculate the expected diel cycle of $SO_4^{2-}$ with sources and sinks discussed above assuming a well-mixed MBL. Omitting dry deposition, the column integrated concentration of $SO_4^{2-}$ at time $t$ can be written as:

$$\left< SO_4^{2-} \right>_t = \left< SO_4^{2-} \right>_{t-1} - (F_{SO_4^{2-}, Zi})_{t-1} + \left< P_{SO_4^{2-}} \right>_{t-1} - \left< L_{SO_4^{2-}, wet} \right>_{t-1} \quad (4.18)$$

Shipboard $SO_4^{2-}$ measurements were taken in the remote region during Oct. 24~30 and Nov. 16~17. We initiate our calculation with the observation at 0000 UTC and compute over one diel cycle. To quantify the average effect of episodic rainfall, the mean precipitation rate of 0.01 mm hr$^{-1}$ is proportioned linearly with the relative cloud fraction, as done with $o_C$. We found that an in-cloud scavenging coefficient of $1.0 P_d^{0.74}$ hr$^{-1}$, which is between the parameterizations from Okita et al. (1996) and Andronache (2004), results in a balance between the source and sink terms. As seen in Fig. 4.17, being proportional to $L_{WP}$, the in-cloud oxidation of $SO_2$ peaks in the early morning, which is largely offset by wet deposition that also peaks at the same time when cloud fraction was the highest and precipitation the most intense. Without clouds, gas phase oxidation is responsible for the slow daytime build up of $SO_4^{2-}$, which then decreases at night due to increased entrainment, similar to the observed trend from Huebert et al. (1996).

However, the processes discussed above do not capture the observed diel trend in VOCALS-REx (note that we could have started our calculation at any initial value, but
would still not obtain a rapid increase in SO$_4^{2-}$ from 0000 to 0400 UTC and decline afterwards).

The observation that surface O$_3$ (which had a FT source) always increased along with SO$_4^{2-}$ in the early evening suggests that the main source of SO$_4^{2-}$ spike was from above. High levels of SO$_4^{2-}$ were not usually observed in the remote FT, which leaves the upper MBL to be the more likely source, with SO$_4^{2-}$ produced from in-cloud oxidation of SO$_2$. During the day, due to the decoupling of the MBL, SO$_4^{2-}$ generated in cloud could be concentrated in the upper MBL and not mixed down to the surface. The re-coupling of the MBL after sunset would mix down this air rich in SO$_4^{2-}$, causing a spike in measured concentration. SO$_4^{2-}$ was then depleted by wet deposition over the rest of diurnal cycle, with the most rapid loss in the early morning. To test this hypothesis, instead of adding in-cloud SO$_2$ oxidation to the SO$_4^{2-}$ budget continuously as in (4.17), we sum the in-cloud SO$_2$ oxidation over the entire day and incorporate it to MBL SO$_4^{2-}$ budget only during the first four hours after sunset. The daily production of SO$_4^{2-}$ in cloud amounted to 4.0 μmoles m$^{-2}$, similar to the increase observed from 0000 to 0400 UTC in the column integrated concentration. Thus we add 1 μmole m$^{-2}$ of SO$_4^{2-}$ to each of these four hours and re-calculate the SO$_4^{2-}$ budget with this sudden SO$_4^{2-}$ input (Fig. 4.18). As wet depositional loss is first order, the large amount of SO$_4^{2-}$ added initially is adequately depleted over the rest of the diurnal cycle and the implied diel variability is qualitatively similar to observations. The major rates in the MBL SO$_4^{2-}$ budget are included in Table 1.
A refinement to the one layer box model utilized here for our diel calculations would be to solve the upper and lower parts of the MBL separately in the decoupled daytime; at night the two layers would join to become one as the MBL re-couples. However, not knowing the exchange between the upper and lower parts of the MBL during the day (which may be examined by eddy covariance) makes the implementation of such a dynamical model difficult for the time being.

4.5 Conclusions

In order to evaluate the effects of anthropogenic sulfur inputs to the atmosphere, we must first understand the natural background cycles. In VOCALS-REx, multiplatform observations and the relatively simple dynamics of the stratocumulus capped marine boundary layer allowed us estimate the sources and sinks of DMS, SO$_2$, and SO$_4^{2-}$ aerosols. The sea-to-air flux of DMS (3.8±0.4 μmoles m$^{-2}$ day$^{-1}$), rather than entrainment of mostly anthropogenically produced SO$_2$ and SO$_4^{2-}$ (0.8±0.4 μmoles m$^{-2}$ day$^{-1}$), was the predominant source of sulfur mass in the remote region of the Southeast Pacific. Accounting for dilution due to entrainment (0.7±0.3 μmoles m$^{-2}$ day$^{-1}$), the remaining DMS was oxidized by OH to mostly SO$_2$. The SO$_2$ production rate from DMS was calculated to be 2.2~2.8 μmoles m$^{-2}$ day$^{-1}$ using a DMS to SO$_2$ conversion of 0.7~0.9, peaking at 7.5~9.7 μmoles m$^{-2}$ day$^{-1}$ in midday. From our implied diel cycle calculations ($\gamma = 0.9$), entrainment slightly increased the MBL SO$_2$ concentration at a rate of 0.05±0.02 μmoles m$^{-2}$ day$^{-1}$, while dry deposition decreased SO$_2$ at a rate of 0.6±0.1 μmoles m$^{-2}$ day$^{-1}$. The gas phase oxidation by OH to H$_2$SO$_4$(g) (then to SO$_4^{2-}$) was about
0.2 μmoles m⁻² day⁻¹ on a diel average, peaking at 0.9 μmoles m⁻² day⁻¹ in the afternoon. The largest SO₂ loss term was the in-cloud oxidation to SO₄²⁻ at 2.1±0.5 μmoles m⁻² day⁻¹, which appears to balance the SO₂ budget and implies other potential removal mechanisms were insignificant. With a slightly acidic aerosol coarse mode, our order of magnitude calculation of SO₂ oxidation by O₃ in sea-salt using measured aerosol size distribution yielded a negligible rate. Entrainment diluted MBL SO₄²⁻ at a rate of 0.5±0.3 μmoles m⁻² day⁻¹, while dry deposition represented an insignificant sink for SO₄²⁻ because little non-sea-salt SO₄²⁻ mass was present in the supermicron mode. The SO₄²⁻ produced from SO₂ was largely removed by wet deposition to the ocean surface.

Diurnally, the implied SO₂ cycle shows a maximum in concentration at around sunset due to oxidation of DMS by OH and minimum just after sunrise as a result of in-cloud oxidation, with dry deposition being a continuous loss term. This diel variation is anticorrelated with that of DMS, consistent with previous observations in the remote Pacific. The SO₄²⁻ diel cycle observed on the ship appeared to be mostly due to boundary layer dynamics and the intricately connected in-cloud SO₂ oxidation and drizzle. SO₄²⁻ produced in cloud was confined to near cloud level during the daytime, when the upper and lower parts of the MBL were often decoupled. As increased turbulence re-coupled the MBL after nightfall, the large amount of SO₄²⁻ built up in the upper MBL was mixed downwards, increasing the concentration at the surface. After peaking in the middle of the night, SO₄²⁻ steadily declined and leveled off for the rest of the day due to wet removal. Drizzle, more frequent in the remote region than near shore, was also more
intense from late night to early morning than in the afternoon. Overall, SO$_2$ had a
time of just over half a day, while SO$_4^{2-}$ had a lifetime of ~1.5 days.

Recent particle nucleation events were observed on numerous occasions in the
MBL from the aircraft as well as on the ship during VOCALS-REx. Such events
occurred when H$_2$SO$_4(g)$ production was likely high and available aerosol surface areas
were low due to cloud or precipitation scavenging, such as in a POC. Given the ample
production rate of H$_2$SO$_4(g)$ from naturally derived SO$_2$ and low levels of pollution
indicators during times of recent nucleation, it is possible that DMS remained not only
the most important precursor gas to SO$_4^{2-}$ aerosol mass in the remote Southeast Pacific,
but also a significant source for new aerosol number.

Acknowledgement

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from http://saga.pmel.noaa.gov/data/download.php?cruise=VOCALS. Special thanks to
C.W. Fairall, R.M. Simpson, University of Hawaii at Manoa, and the crew of the R/V
Ronald H. Brown.
### Table 4.1 Diel averaged rates in MBL concentration budgets of DMS, SO$_2$, and SO$_4$$^{2-}$

<table>
<thead>
<tr>
<th></th>
<th>DMS</th>
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<tbody>
<tr>
<td></td>
<td>µmole m$^{-2}$ day$^{-1}$</td>
<td>pptv hr$^{-1}$</td>
<td>% day$^{-1}$</td>
</tr>
<tr>
<td>Sea-to-air Flux</td>
<td>3.81±0.38</td>
<td>3.09±0.31</td>
<td>118±12</td>
</tr>
<tr>
<td>Entrainment</td>
<td>-0.66±0.33</td>
<td>-0.54±0.27</td>
<td>-20±10</td>
</tr>
<tr>
<td>Loss to OH</td>
<td>-3.15±0.50</td>
<td>-2.56±0.41</td>
<td>-97±16</td>
</tr>
</tbody>
</table>

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<tr>
<th></th>
<th>SO$_2$</th>
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<tbody>
<tr>
<td></td>
<td>µmole m$^{-2}$ day$^{-1}$</td>
<td>pptv hr$^{-1}$</td>
<td>% day$^{-1}$</td>
</tr>
<tr>
<td>Prod. from DMS</td>
<td>2.84±0.45</td>
<td>2.30±0.37</td>
<td>158±25</td>
</tr>
<tr>
<td>Entrainment</td>
<td>0.05±0.02</td>
<td>0.04±0.02</td>
<td>3±1</td>
</tr>
<tr>
<td>Dry Deposition</td>
<td>-0.56±0.14</td>
<td>-0.45±0.11</td>
<td>-31±8</td>
</tr>
<tr>
<td>Loss to OH</td>
<td>-0.22±0.03</td>
<td>-0.18±0.02</td>
<td>-12±2</td>
</tr>
<tr>
<td>Loss to Clouds</td>
<td>-2.10±0.48</td>
<td>-1.70±0.39</td>
<td>-117±26</td>
</tr>
</tbody>
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<table>
<thead>
<tr>
<th></th>
<th>SO$_4$$^{2-}$</th>
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</tr>
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<tbody>
<tr>
<td></td>
<td>µmole m$^{-2}$ day$^{-1}$</td>
<td>pptv hr$^{-1}$</td>
<td>% day$^{-1}$</td>
</tr>
<tr>
<td>Prod. from SO$_2$</td>
<td>4.02±0.89</td>
<td>3.26±0.72</td>
<td>75±16</td>
</tr>
<tr>
<td>Entrainment</td>
<td>-0.51±0.26</td>
<td>-0.41±0.21</td>
<td>-9±5</td>
</tr>
<tr>
<td>Wet Deposition</td>
<td>-3.51±0.92</td>
<td>-2.85±0.75</td>
<td>-65±17</td>
</tr>
</tbody>
</table>

Positive (negative) indicate increases (decreases) in MBL concentrations.

SO$_2$ and SO$_4$$^{2-}$ rates are calculated with two separate DMS to SO$_2$ conversion efficiencies, 0.9 and 0.7. Percent daily rates are calculated with respect to observed (DMS and SO$_4$$^{2-}$) or implied (SO$_2$) daily mean concentrations.

Errors for entrainment, dry deposition, and SO$_2$ oxidation are estimated from uncertainty in $\omega$, $V_{dep}$, and [OH], with errors for other rates propagated under 95% confidence.
Figure 4.1 VOCALS-REx cruise track color-coded by the DMS sea-to-air flux measured with eddy covariance. Flux was higher offshore mostly as a result of higher wind speed. Seawater DMS concentration was variable and elevated in isolated areas mostly associated with fronts and eddies, but showed no general longitudinal trend. The dotted and solid boxes indicate the VOCALS ‘offshore’ and ‘remote’ regions, respectively.
Figure 4.2 A typical 20°S Survey flight by the C-130 (RF 03), with color indicating SO$_2$ concentration and marker size representing aerosol SO$_4^{2-}$ concentration. Both sulfur species showed higher concentrations near the coast than offshore. Also shown is the inversion height derived from C-130 profiles, which increased with distance away from shore. Level legs were typically flown near the surface, below the stratocumulus cloud deck, and above cloud, with occasional profiles to higher altitudes.
Figure 4.3 Longitudinal trends in the atmospheric concentrations of DMS, SO$_2$, and SO$_4^{2-}$ along 20(±2)$^\circ$S in the MBL and FT (project averages). SO$_2$ and SO$_4^{2-}$ were significantly elevated near shore due to pollution, and rapidly decreased away from shore, where DMS was higher. In the remote atmosphere, the sea-to-air DMS flux accounted for about three quarters of the MBL sulfur burden, with the rest due to the entrainment of SO$_2$ and SO$_4^{2-}$. 
Figure 4.4  (a) Averaged vertical profiles of temperature, dew point, CO, and O₃; (b) SO₄²⁻, SO₂ (with standard error of the mean), potential temperature, and liquid water content vs. altitude normalized to the inversion height. CO and O₃ concentrations were consistently greater above the sharp inversion (Z/Zᵢ = 1), while SO₄²⁻ was greater below the inversion. Both SO₂ and SO₄²⁻ (cloud free air and interstitial concentrations) were reduced at cloud level, most likely due to cloud processing and scavenging.
Figure 4.5 Mean VOCALS C-130 remote particle number, surface area, and volume distributions at different normalized altitudes. While the mid-MBL distribution was similar to the surface distribution, the upper MBL distribution was much reduced, likely as a result of cloud scavenging and MBL decoupling. The above cloud distribution was a superposition of two modes, with the ~30 nm mode corresponding to low CO and the ~80 nm mode corresponding to high CO. Also shown is the distribution identified by high concentrations of recently nucleated small particles and typically characterized with reduced surface area.
Figure 4.6 Diel cycles in inversion height, mixed layer height, liquid water path, and cloud fraction in the VOCALS offshore region. All of these variables were greater at night than during the day mostly due to the diel variability in the radiative properties associated with the stratocumulus cloud deck.
Unlike what photochemistry predicts, \( \text{SO}_4^{2-} \) frequently decreased during the day and peaked at night, as did \( \text{O}_3 \). MLH approached \( Z_i \) at night, suggesting a well-mixed MBL, whereas during the day MLH was only \( \sim 1/2 \) of \( Z_i \), indicating decoupling. The nighttime increase in \( \text{SO}_4^{2-} \) was likely due to MBL coupling and in-cloud production of \( \text{SO}_4^{2-} \).
Figure 4.8 Mean VOCALS RHB particle number, surface area, volume, and mass distributions stratified by the time of day in the remote region. The accumulation mode (dominated by $\text{SO}_4^{2-}$) was clearly the largest between near sunset to a few hours before sunrise (2200–0600 UTC), consistent with the observed increase in $\text{SO}_4^{2-}$ by the shipboard AMS. The coarse mode (dominated by sea-salt) appeared to be largest in the afternoon (1400–2200 UTC) likely as a result of higher wind speed. Also shown is the distribution characterized by high concentrations of recently nucleated small particles and low aerosol surface area.
Figure 4.9 Recent nucleation event observed on the night of Oct. 24, when the ship was near 85ºW and 16ºS and steaming south at 11 knots. The submicron particle area decreased drastically from ~20 μm² cm⁻³ to ~13 μm² cm⁻³ by 0500 UTC (2 am local time), when the number of particles in the 5~13 nm range increased to 200~300 cm⁻³. Assuming the airmass stayed the same, a precipitation event at 0800 UTC appeared to scavenge essentially all of these small particles. Due to the short lifetime of H₂SO₄(g), the gas-to-particle conversion had likely taken place in the daytime of Oct. 23 rather than during 0500~0800 UTC of Oct. 24. Without H₂SO₄(g) for condensational growth, the nucleation model particles grew very slowly in the nighttime such that many of them were still under 13 nm by the morning.
Figure 4.10 Recent nucleation event observed on Oct. 27, when the ship was stationed at 85°W and 20°S. The beginning and the end of the time series roughly corresponded to the leading and trailing edges of a POC, as determined from GOES images. CN steadily decreased from ~600 cm\(^{-3}\) at 0900 UTC to ~100 cm\(^{-3}\) by 1600 UTC, along with an increase in O\(_3\). As O\(_3\) started to decrease at 1700 UTC, both CN and UFCN increased rapidly in a few bursts, with a difference of ~250 cm\(^{-3}\) between the two number concentrations. The submicron particle surface area was very low (2~8 \(\mu m^2 cm^{-3}\)), likely in part due to frequent light drizzle events. Liquid water path, air temperature, and RH all showed periodicity on a timescale of ~30 minutes, which may be reflecting the intermittent turbulence and cell structures in the decoupled MBL.
Figure 4.11 Time-series of submicron MSA:SO$_4^{2-}$ ratio and scattering at 550 nm, DMS, and supermicron chloride deficit. The MSA:SO$_4^{2-}$ ratio follows closely to DMS concentration, providing another evidence that DMS was the most important sulfur source away from the coast and also implying that the lifetimes of these sulfur species were similar. Chloride deficit was the largest near the coast, where pollution emission of acidic aerosols (sulfate and nitrate) were the highest, as evidenced by the high scattering.
Figure 4.12 SO$_2$ concentration below the inversion vs. liquid water content from the C-130 for the VOCAL offshore region, stratified to daytime and nighttime. Higher SO$_2$ during the day was due to DMS oxidation. Above 0.2 g m$^{-3}$ (mean $L_{WC}$ for the stratocumulus clouds), SO$_2$ was reduced to ~18 pptv as a result of in-cloud oxidation. The interstitial SO$_2$ might be slightly higher during the day due to the initially higher MBL concentration.
Figure 4.13 Calculated oxidation of SO$_2$ to sulfate in cloud due to H$_2$O$_2$ in a closed system. Initial concentrations were set to 20 pptv and 65 μM (~0.9 ppb) for SO$_2$ and H$_2$O$_2$. The liquid water content and pH were fixed at 0.2 g m$^{-3}$ and 4.2, respectively. After ~15 minutes (typical in cloud residence time), 11% of the SO$_2$ is converted to SO$_4^{2-}$, leaving an interstitial concentration SO$_2$ of ~18 pptv.
Figure 4.14 $\text{SO}_4^{2-}$ vs. $\text{NH}_4^+$ from the shipboard AMS, color-coded by longitude. The $\text{SO}_4^{2-}:\text{NH}_4^+$ ratio lies close to the theoretical full neutralization limit of 1:2 for ammonium sulfate in the remote marine region. Closer to the coast, the ratio exceeds 1:1 (for ammonium bisulfate) due to pollution emission.
Figure 4.15 Implied SO$_2$ diel cycle, with oxidation from DMS being the principal source and in-cloud oxidation as the main sink. The implied cycle agrees well with observations (error bars correspond to standard deviation within hour bins) until 1500 UTC, with measurements in the subsequent hours likely subject to greater spatial bias.
Figure 4.16 *RHB* $\text{SO}_4^{2-}$ concentration vs. rain rate from an optical rain gauge, color-coded by longitude. While there was clearly a longitudinal trend in $\text{SO}_4^{2-}$, the lowest concentration at 85°W corresponded to high rain rates, suggesting efficient precipitation scavenging.
Figure 4.17 Implied SO$_4^{2-}$ cycle assuming a well-mixed MBL, with calculation initiated with the observation at 0000 UTC. The in-cloud production term is largely offset by wet deposition, with gas phase oxidation by OH accounting for the small increase in SO$_4^{2-}$ towards the end of the day. While the source and sink terms appear to balance, the observed diel cycle in SO$_4^{2-}$ is not captured by this calculation.
Figure 4.18 Implied SO$_4^{2-}$ cycle assuming a well-mixed MBL at night and decoupled MBL during the day. SO$_4^{2-}$ produced in-cloud is summed over the entire day and only added to the MBL budget over the first four hours after sunset as the MBL re-coupled. Initiated with the observation at 0000 UTC, the implied cycle qualitatively agrees with shipboard observations (error bars correspond to standard deviation within hour bins).
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