CO₂ diffusion in polar ice: observations from naturally formed CO₂ spikes in the Siple Dome (Antarctica) ice core

Jinho AHN,1,2 Melissa HEADLY,1 Martin WAHLEN,1 Edward J. BROOK,2 Paul A. MAYEWSKI,3 Kendrick C. TAYLOR4

1Scripps Institution of Oceanography, University of California–San Diego, La Jolla, California 92093-0225, USA
E-mail: jinhoahn@gmail.com
2Department of Geosciences, Oregon State University, Corvallis, Oregon 97331-5506, USA
3Climate Change Institute, University of Maine, 303 Bryant Global Sciences Center, Orono, Maine 04469-5790, USA
4Desert Research Institute, University of Nevada, 2215 Raggio Parkway, Reno, Nevada 89512-1095, USA

ABSTRACT. One common assumption in interpreting ice-core CO₂ records is that diffusion in the ice does not affect the concentration profile. However, this assumption remains untested because the extremely small CO₂ diffusion coefficient in ice has not been accurately determined in the laboratory. In this study we take advantage of high levels of CO₂ associated with refrozen layers in an ice core from Siple Dome, Antarctica, to study CO₂ diffusion rates. We use noble gases (Xe/Ar and Kr/Ar), electrical conductivity and Ca²⁺ ion concentrations to show that substantial CO₂ diffusion may occur in ice on timescales of thousands of years. We estimate the permeation coefficient for CO₂ in ice is ~4 × 10⁻¹⁹ mol m⁻¹ s⁻¹ Pa⁻¹ at ~−23°C in the top 287 m (corresponding to 2.74 kyr). Smoothing of the CO₂ record by diffusion at this depth/age is one or two orders of magnitude smaller than the smoothing in the firn. However, simulations for depths of ~930–950 m (~60–70 kyr) indicate that smoothing of the CO₂ record by diffusion in deep ice is comparable to smoothing in the firn. Other types of diffusion (e.g. via liquid in ice grain boundaries or veins) may also be important but their influence has not been quantified.

INTRODUCTION
Carbon dioxide (CO₂) is the most important greenhouse gas directly impacted by human activities. Ancient air preserved in polar ice cores provides extremely important information about the functioning of the carbon cycle in the past (e.g. Etheridge and others, 1996; Fischer and others, 1999; Petit and others, 1999; Kawamura and others, 2003; Ahn and others, 2004; EPICA Community Members, 2004; Siegenthaler and others, 2005). The reconstructed records extend direct measurements of atmospheric CO₂ concentrations, which started in 1958 (Keeling, 1960), and may help us predict future climate under rapidly increasing CO₂ more accurately.

The integrity of an ice core as a reliable archive depends on the incorporation followed by the preservation of the original atmospheric signal. It is well known that atmospheric records are smoothed, due to diffusion in the firn column and gradual air trapping in the bubble close-off zone (e.g. Schwander and others, 1998; Trudinger and others, 2002; Spahni and others, 2003). However, CO₂ diffusion in ice after the air is trapped in bubbles is poorly understood, because the diffusion coefficient is too small to be precisely measured in the laboratory (Hondoh, 1996). This uncertainty also limits our understanding of rapid CO₂ changes in the atmosphere.

The permeation coefficient (solubility × diffusion coefficient) quantifies gas diffusion in solids. Recent results from a molecular dynamics simulation (Ikeda-Fukazawa and others, 2004) show that CO₂ molecules can diffuse orders of magnitude faster in ice than indicated by previous estimates that were based on an interstitial mechanism (Ikeda and others, 2000). The fast diffusion may be due to a new mechanism called the breaking-bond mechanism, where hydrogen bonds break and CO₂ hops between stable sites in ice crystals (Ikeda-Fukazawa and others, 2004). However, to date, no good observational estimate of CO₂ diffusion in polar ice cores has been reported.

In this study we take advantage of natural CO₂ spikes in an ice core associated with refrozen melt layers (hereafter melt layers) to study diffusion that occurred in the ice matrix over thousands of years. We use the gradual decrease of CO₂ concentration away from the melt layers, combined with high-resolution analyses of Ca²⁺ ion, ECM (electrical conductivity measurements), Xe/Ar and Kr/Ar of trapped air to quantify CO₂ diffusion in ice. We provide an estimate of the CO₂ permeation coefficient in the ice, and discuss the implications of the results for the preservation of CO₂ signals in ice cores.

MATERIALS AND METHODS
The Siple Dome (Antarctica) ice core was drilled between 1997 and 1999. The site is at 81.66° S, 148.82° W, with a present-day annual mean temperature of −25.4°C (Severinghaus and others, 2001) and accumulation rate of 12.4 g cm⁻² a⁻¹ w.e. The total depth of the core is 1003.8 m. While summer air temperatures were generally well below the melting point (Das and Alley, 2005), surface melting could occur during brief summer warm periods. This happened zero to two times per century during the Holocene (Das and Alley, 2008).

CO₂ measurements were made at the Scripps Institution of Oceanography (SIO) on ice containing bubbles that also included some of the visually distinctive melt layers in the Siple Dome A core (Fig. 1). In each 1 cm depth interval ice samples of 4–6 g were used, and the outer 0.5 cm of the samples was removed with a bandsaw to reduce the possibility of contamination from present atmospheric CO₂. The
gas extraction and infrared (IR) spectroscopic methods used were described by Ahn and others (2004), and are similar to previous methods (Wahlen and others, 1991; Smith and others, 1997a, b; Fischer and others, 1999). Trapped air was extracted from the ice by mechanical crushing in a double-walled crushe cooled to about –40°C using flowing cold ethanol. The liberated air was collected in small cold traps chilled by closed-cycle helium refrigerators to a temperature of ~32 K. Trapped air samples were liberated by heating the traps to –60°C and then transferred to an IR absorption cell held at constant pressure and temperature. IR absorption measurements were made several times on each gas sample with a tunable diode laser. The single-mode IR laser output was scanned across a single vibrational–rotational molecular absorption line of CO2 at Doppler resolution. To calibrate the instrument, measurements were made with three air standards of precisely known CO2 concentrations of 163, 240 and 330 ppm. The standards were introduced over three of the crushed ice samples. This calibration procedure was performed each day. The internal precision was ~2 ppm. in the concentration range 163–330 ppm.

Noble-gas ratios (δ132Xe/36Ar, δ84Kr/36Ar and δ40Ar/36Ar) were analyzed in order to quantify the amount of refrozen melt. These permit us to estimate how much CO2 originated from the meltwater. The gas extraction and measurement on a dual viscous-inlet Finnigan MAT 252 mass spectrometer at SIO have been described previously (Severinghaus and others, 2001; Severinghaus and Battle, 2006; Headly and Severinghaus, 2007). The depth resolution (4–6 cm) for the noble gases was poorer than the CO2 resolution (1 cm) because a larger ice sample was required for the noble-gas measurements. Around 40–60 g of ice and two stir bars were put in chilled glass vessels, which were attached to a vacuum line. The ambient air was evacuated from the vessel and line for 40 min, after which we closed the valve to stop pumping, and the ice was allowed to melt. The air released from the air bubbles was cryogenically (liquid He) concentrated at 10 K into a stainless-steel dip tube. After warming to room temperature, the gas sample was exposed to a Zr/Al getter at 900°C for 10 min to destroy all the reactive gases, followed by 2 min at 300°C to remove H2.

After the gettering process, the remaining noble gases were transferred into a sample tube at 10 K. Finally, ultra-high purity N2 equal to ten times the noble-gas pressure in the vacuum line was added to the sample to add bulk suitable for mass spectrometry. Samples were run against aliquots of a standard gas mixture of commercially obtained N2, Ar, Kr and Xe. δ132Xe/36Ar, δ84Kr/36Ar and δ40Ar/36Ar were normalized using clean marine air outside the laboratory. The pooled standard errors of the triplicate measurements for the Greenland Ice Sheet Project 2 (GISP2) ice core of the Holocene were 3.77‰ and 1.56‰ for δ132Xe/36Ar and δ84Kr/36Ar, respectively, for the measurement conditions. For Ca2+ ion concentration measurements, we used ~5 g of ice from the same depths used for CO2 concentration measurements. Ice was prepared at SIO and measured at the Climate Change Institute, University of Maine, using Dionex-500 ion chromatography, where calcium was measured on a CS12A cation-exchange column with 25 mM methanesulfonic acid eluent, a self-regenerating suppressor and a conductivity detector. Sample size was 500 μL.

RESULTS

Excess CO2 associated with melt layers

The annual mean temperature at Siple Dome is ~25.4°C (Severinghaus and others, 2001). However, the site has experienced occasional melting during austral summers (0–2 melt events per century during the Holocene) (Das and Alley, 2008). The meltwater formed on the snowpack surface percolates into the firm and refreezes at some depth, but rarely percolates >0.2 m, as shown in Figure 2. The snow layer where the melt refreezes is in the form of a fine-crystal size crust that was formed during the previous winter, and has a strong capillary force (Das and Alley, 2005). Melt layers preserved in the Siple Dome ice core are easily identified visually (Fig. 1) due to lower gas content and reduced bubble size, compared to normal bubbly ice.

A 30 cm long ice sample from a depth of 286.7–287.0 m was intensively studied (Fig. 1). The age of the ice is
~3.04 kyr BP (thousand years before AD 1950), and the gas age is ~2.74 kyr BP (Taylor and others, 2004; Brook and others, 2005). Annual ice layers are ~5–9 cm thick at this depth (Taylor and others, 2004). This section of ice core contains two melt layers (~1 cm thick), identified by smaller bubble sizes than in the surrounding ice (Fig. 1). The upper (left) melt layer, M1, is thicker than the lower (right) melt layer, M2 (Fig. 1). Distinct dark patches (due to less light scattering in Fig. 1) between M1 and M2 have small bubbles as seen in M1 and M2, suggesting other melt features that do not continue horizontally.

Due to the high solubility of CO2 in liquid water (Table 1), we expect CO2 concentrations in the melt layer to be higher than those in normal bubble ice (Stauffer and others, 1985). Assuming an atmospheric CO2 concentration of 278 ppm (atm/atm) (Indermühle and others, 1999) at the gas age of 2.74 kyr BP and surface pressure at Siple Dome of 937 hPa, we expect 16230 ppm CO2 (μmol CO2/mol total air) dissolved in 0°C meltwater in equilibrium, 58 times greater than in the atmosphere (Table 1). If a thin film of snowmelt attains solubility equilibrium with the atmospheric air at the surface, and then refreezes rapidly at some greater depth, the excess CO2 in the meltwater can be trapped and preserved in small bubbles in the melt layer. These bubbles can be preserved through the firm metamorphism process and incorporated in mature ice. The CO2 concentration gradient between the melt layer and the neighboring ‘normal’ bubble ice may then cause diffusion through the ice. Excess CO2 (CO2 concentration observed in ice minus atmospheric CO2 concentration) would not be preserved in the melt layer if degassing had occurred fully before the meltwater refroze. Preservation of bubbles in melt layers suggests that refreezing is not slow enough for effective degassing.

Our high-resolution (depth interval of 1 cm) study shows that the CO2 concentration in bubbles gradually decreases from 750 to 285 ppm away from M1. Another, smaller, peak is also found around M2 (Fig. 3). CO2 records from colder Antarctic ice-core sites indicate atmospheric concentrations of 278 ppm for the age of our samples (Indermühle and others, 1999). The CO2 concentrations in the Siple Dome samples are therefore greater by up to 470 ppm (excess CO2), and clearly do not represent atmospheric values.

The existence of refrozen meltwater in M1 and M2 is supported by our analyses of \(^{132}\text{Xe}/^{36}\text{Ar}\) and \(^{84}\text{Kr}/^{36}\text{Ar}\) as shown in Figure 3. \(\text{Xe}/\text{Ar}\) and \(\text{Kr}/\text{Ar}\) in glacial ice are useful indicators of extensive snowmelting and refreezing, as \(\text{Xe}\) and \(\text{Kr}\) are about four and two times as soluble in liquid water as \(\text{Ar}\), respectively (Severinghaus and others, 2003). Our data show significantly enriched \(\text{Xe}/\text{Ar}\) and \(\text{Kr}/\text{Ar}\) for ice that includes the visible melt layers, M1 and M2, relative to normal layers, indicating the existence of the refrozen meltwater in M1 and M2.

In order to quantitatively determine the excess CO2 resulting from the refrozen melt, we use the melt-sensitive isotope ratios, \(^{132}\text{Xe}/^{36}\text{Ar}\) and \(^{84}\text{Kr}/^{36}\text{Ar}\). Considering the solubility of the noble gases at 0°C, the gravitational fractionation in the firm layer and the Ar loss from the bubbles during sample handling or bubble close-off, our calculation using the \(\text{Xe}/\text{Ar}\) and \(\text{Kr}/\text{Ar}\) data gives the volume ratio of liquid water to pore air (see Appendix B for details). This is accomplished by solving three equations for three unknowns with three observed quantities: \(\text{Xe}/\text{Ar}\), \(\text{Kr}/\text{Ar}\) and \(^{40}\text{Ar}/^{36}\text{Ar}\).

### Table 1. Solubilities and diffusion coefficients of air components in fresh water for Siple Dome

<table>
<thead>
<tr>
<th>Air</th>
<th>(X_{\text{atm}})</th>
<th>(X_{\text{H,O}})</th>
<th>(D_{\text{H,O}})</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\text{N}_2)</td>
<td>100%</td>
<td>78.08%(^a)</td>
<td>1.4b</td>
</tr>
<tr>
<td>(\text{O}_2)</td>
<td>100%</td>
<td>20.95%(^a)</td>
<td>1.3b</td>
</tr>
<tr>
<td>(\text{Ar})</td>
<td>0.93%(^a)</td>
<td>1.68%(^c)</td>
<td>1.2b</td>
</tr>
<tr>
<td>(\text{CO}_2)</td>
<td>278 ppm(^b)</td>
<td>16230 ppm(^a)</td>
<td>4.21 ppm(^d)</td>
</tr>
<tr>
<td>(\text{Kr})</td>
<td>1.14 ppm(^a)</td>
<td>4.21 ppm(^d)</td>
<td>0.679 ppm(^d)</td>
</tr>
<tr>
<td>(\text{Xe})</td>
<td>0.087 ppm(^a)</td>
<td>0.88(^d)</td>
<td>0.66(^d)</td>
</tr>
</tbody>
</table>

\(X_{\text{atm}}\): composition of atmospheric air; \(X_{\text{H,O}}\): air composition dissolved in water at 0°C and 937 mbar; \(D_{\text{H,O}}\): diffusion coefficient (10\(^{-9}\) m\(^2\) s\(^{-1}\)) in water at 0°C.

For the ice sample that includes normal layer, M1 and the partial-melt layer between M1 and M2, a volume ratio of liquid water to pore air of 1.38 is obtained. Using the total air in the sample (4.7 ± 0.5 cm³ STP (standard temperature and pressure)), we calculate that the volume of refrozen melt is 6.5 ± 0.7 cm³. The total air content of the ice samples was calculated with (1) the volumes of ice from the normal and melt layers, (2) bubble-volume ratios (air-content ratios) of normal melt layers and (3) the air content of the normal layers, which was estimated from the average over the Holocene as the air content is almost constant in that period (Severinghaus and Battle, 2006). The estimated volume of refrozen melt is comparable to our visual observation. The volume of visual melt layer (defined by ice with smaller bubbles than in the normal layer) is 16.4 ± 0.7 cm³ (13.9 ± 0.7 cm³ from M1; 2.5 ± 0.7 cm³ from the partial-melt patch between M1 and M2). This visual refrozen ice volume is apparently bigger than our estimation of the volume of refrozen melt because the volume of the visual melt layer includes pre-existing ice (firm), where meltwater filled the void space and refroze. The difference between estimates based on Xe/Ar, Kr/Ar, 40Ar/36Ar and the visual observation is 40 ± 8%, close to the porosity of snow at the surface, 56% (ice density at the surface = 0.4 g cm⁻³; personal communication from J. Fitzpatrick, 2007), indicating that Xe, Kr and Ar were close to equilibrium with the ancient atmosphere and were trapped in melt layers. Therefore, we can also expect that the melt at the surface was in equilibrium with ancient atmospheric CO₂ which was trapped in M1.

For additional confirmation for our diffusion model, we estimate the total CO₂ trapped in M1 at the snow surface with the calculated volume of purely refrozen melt in M1 from Xe/Ar and Kr/Ar measurements:

\[
\text{Half of the excess CO₂ in M1 before diffusion in ice (mol cm}^{-2}\text{)} = \frac{\text{half of the melt layer thickness (cm)}}{\text{volume fraction of purely refrozen melt in M1 from Xe/Ar and Kr/Ar}} \times \text{ice density (g cm}^{-3}\text{)} \times \text{solubility of CO₂ in water at 0°C (mol g}^{-1}\text{)}
\]

\[
= (0.5 \times 1.23) \times (0.416 \pm 0.059) \times 0.917 \times 1.998 \times 10^{-8}
\]

\[
= 4.7 \pm 0.7 \times 10^{-9} \text{ mol cm}^{-2}.
\]

This agrees with the integration of the observed excess CO₂ in the upper part of M1 (including the upper half of M1) of \(~5.4 \times 10^{-9}\text{ mol cm}^{-2}\) This agreement supports the assertion that the CO₂ profile around the melt layers was formed by diffusion from the melt layer. The accuracy of the above estimate is possibly limited by different degrees of equilibration of different gases in the meltwater. For example, Xe should approach equilibrium more slowly than CO₂, due to its lower diffusivity but within the same order (Table 1). In
this estimate we assume gas diffusion in ice does not change the profile of Xe/Ar and Kr/Ar.

**CO₂ diffusion in the ice**

The gradual decrease in CO₂ away from the melt layers is consistent with CO₂ diffusion over the past 2.74 kyr, and agrees reasonably well with predictions from a molecular volume-diffusion model (Fig. 4) (Neftel and others, 1983; see Appendix A for details). For simple one-dimensional curve fitting, we utilize the CO₂ mixing ratio data from the normal and melt layers. The CO₂ concentrations from the partial-melt layer between the two melt layers have higher values than those in the fitting curve, due to additional CO₂ in the partial-melt patches. Thinning of vertical layers with increasing depth due to the flow of the ice is taken into account. In our numerical approach, we utilize an age–depth profile from CH₄ correlation with the Greenland ice core (Brook and others, 2005) and a depth–density profile (personal communication from J. Fitzpatrick, 2007) for the real depth, pressure and porosity parameters in each time-step rather than estimation based on the assumption of constant snow accumulation rate and porosity (Neftel and others, 1983). For example, the porosity profile for the numerical modeling was estimated from the density profile for the top 300 m ($\phi = 1 - \rho_{\text{bubbly ice}} / \rho_{\text{bubble-free ice}}$) rather than using confining ice pressure, because air pressure in bubbles at shallow depth is not in equilibrium with the confining hydrostatic ice pressure, as shown in the top 300 m of the Vostok (Antarctica) ice core (Lipenkov, 2000). For depths >300 m we obtained porosity from the gas content (~0.11 cm³ air (STP) g⁻¹ ice) (Severinghaus and Battle, 2006) divided by confining pressure in the atmosphere (personal communication from J. Fitzpatrick, 2007) and the density of bubble-free ice (0.917 g cm⁻³). Two approaches were used for estimating the vertical thinning factor, $\alpha$ (see Appendix A for details): (1) assuming constant strain-rate thickness and (2) utilizing the paleo-accumulation rate estimated from the isotopic temperature proxy (Brook and others, 2005). The latter estimations of $\alpha$ are 89% and 57% of the former for the last 2740 years and ~18.5–80 kyr, respectively. However, in the 2740 year modeling and curve fitting, using the two different thinning factor approaches gave no difference in the estimated permeation coefficient within the range of uncertainty of curve fitting.

We assume that the permeation coefficient (diffusion constant × solubility) is almost constant over a wide range of solubility (see Appendix A), and the same value of the permeation constant of CO₂ in ice was applied for the two melt layers. The good fit of the model to the data supports our proposition that the CO₂ originally trapped in the melt layer has diffused through the ice for thousands of years. The baseline CO₂ levels far from the melt layers are still slightly higher than in the Taylor Dome (Antarctica) and Dome C (Antarctica) ice records (Indermühle and others, 1999), by 11 ppm on average. The reason for this discrepancy is not clear, but may be due to microscale refreeze of melt (Ahn and others, 2004) or differences in laboratory standard scales for CO₂.

Alternative explanations for the excess CO₂ in the ice adjacent to the distinctive melt layers include: (1) visibly undetected micro-melt layers, (2) carbonate–acid reaction (Delmas, 1993; Anklin and others, 1995, 1997; Barnola and others, 1995; Smith and others, 1997a, b) and (3) oxidation of organic compounds abiologically (Tschumi and Stauffer, 2000) or biologically (Campen and others, 2003). If the excess CO₂ of hundreds of ppm around M1 and M2 originated from micro-melt layers, we would observe the same characteristics in the excess Xe/Ar and Kr/Ar as in the excess CO₂. However, we do not detect significantly elevated Xe/Ar and Kr/Ar above M1 and below M2, in areas where we do see significant excess CO₂ (Fig. 3). These observations strongly support our idea that excess CO₂ in the normal layer formed by diffusion from the melt layers. In between M1 and M2, the resolution prevents us from comparing the CO₂ with Xe/Ar or Kr/Ar. Different degrees of saturation of the gas species and different degassing during freezing may explain the difference between CO₂ and the Xe/Ar and Kr/Ar profiles. However, the diffusion coefficients
of these gas species in water are of the same order (Table 1), supporting our diffusion model. It is important to note that we are assuming that diffusion of Xe, Kr and Ar in ice is negligible compared to that of CO$_2$. However, recent work suggests that Ar may diffuse more rapidly in ice than Kr and Xe (Severinghaus and Battle, 2006). If Ar is more mobile than Kr and Xe, the Kr/Ar and Xe/Ar ratios will decrease outside the melt layers but increase in the melt layers. To date, the permeation coefficients of Xe, Kr and Ar are unknown. Thus, exact evaluation of this effect remains elusive. However, the good agreement of two noble-gas datasets in all layers other than melt layers would have to be fortuitous if this effect is important. For this reason we consider the scenario unlikely.

The concentration of dissolved Ca$^{2+}$ can be used to estimate the upper limit of the amount of CaCO$_3$ that potentially could have reacted with H$^+$. There is no significant correlation between the excess CO$_2$ and the dissolved Ca$^{2+}$ or ECM (a proxy for H$^+$), as shown in Figure 3. Moreover, assuming that all the Ca$^{2+}$ is produced from the reaction between CaCO$_3$ and H$^+$, the potential excess CO$_2$ is $\sim$17 ppm on average. Thus, the amount of Ca$^{2+}$ is too small to explain our observations (excess CO$_2$ of up to 470 ppm); 1 µg Ca$^{2+}$ per kilogram of ice can explain at most 7 ppm of excess CO$_2$ by the carbonate–acid reaction.

Oxidation of organic compounds has been proposed to be at least as important as acid–carbonate reactions for CO$_2$ production (e.g., 2H$_2$O$_2$ + HCHO → 3H$_2$O + CO$_2$) in some ice cores (Tschumi and Stauffer, 2000). However, H$_2$O$_2$ data for the top 100 m of the Siple Dome ice core show H$_2$O$_2$ concentrations near or below the detection limit of $\sim$0.02 µm, except at 0–2.5 m depth (McConnell, 1997). H$_2$O$_2$ is one of the major oxidants in snow. These H$_2$O$_2$ concentrations at Siple Dome are much lower than those found in other Antarctic ice cores (McConnell, 1997). The data for other important oxidants such as CH$_3$COO$^-$ and HCOO$^-$ (Tschumi and Stauffer, 2000) are not available.

However, the excess CO$_2$ due to this process may be less than in Greenland ice cores, where the dust and oxidant contents are greater than in Antarctic ice cores by an order of magnitude and several times, respectively (Tschumi and Stauffer, 2000), and the excess CO$_2$ is $\sim$30 ppm for the Holocene (Anklit and others, 1995; Barnola and others, 1995). Thus, oxidation of organic compounds cannot explain the high excess CO$_2$ of up to 470 ppm.

We conclude that the three alternative mechanisms cannot explain the excess CO$_2$ distribution around the melt layers, and that the concentration profile is most likely to have formed by the diffusion of CO$_2$ through the ice matrix from the refrozen melt (M1, M2 and partial-melt) layers. However, we cannot exclude the possibility of invisible micro-melt layers due to our unproved assumption that diffusion did not change Xe/Ar and Kr/Ar profiles. In this case, our estimation of the permeation coefficient of CO$_2$ in the following part of this section would be a maximum and provide an upper limit on the impact on CO$_2$ mobility of the ice-core records.

From the best-fit calculation, the permeation coefficient of CO$_2$ (the product of the diffusion constant and the solubility) in ice is $\sim4 \times 10^{-21}$ m$^2$ s$^{-1}$ mol Pa$^{-1}$ at $-23$°C (time-averaged temperature) as seen in Figure 4. We find higher CO$_2$ concentrations in the samples that include partial-melt layers than in the melting because the former have lower gas content than the normal bubble ice. Our result for the permeation coefficient of CO$_2$ is independent of the Xe/Ar and Kr/Ar data. We estimate that the amplitudes of the CO$_2$ variation in the measured core segment are $\sim$30% and $\sim$20% of the initial amplitude, $c_0$, for M1 and M2, respectively (Fig. 5). The different ratios are due to the different melt-layer thicknesses.

Our result for the CO$_2$ permeation coefficient is an order of magnitude greater than that estimated from the Dye 3 (Antarctica) ice core ($1.3 \times 10^{-22}$ m$^2$ s$^{-1}$ mol Pa$^{-1}$ at $-20$°C) (Neftel and others, 1983). This discrepancy is probably due to
the inaccurate assumption that the Dye 3 ice core has never changed to clathrate ice from bubbly ice (Neftel and others, 1983). The Dye 3 ice-core segment studied for diffusion was selected from the bottom of the bubble-ice-clathrate transition zone (depth = 1616 m; 31.1 kyr), where gas species go into the ice lattice and bubbles shrink and finally disappear (Neftel and others, 1983). The formation of clathrate begins at a depth of 1200 m (14.7 kyr) (Neftel and others, 1983) and possibly significantly changes the diffusion of CO₂. Salamatin and others (1998) deduced that the diffusion coefficient of air in hydrate crystals is at least two orders of magnitude less than the diffusion coefficient of air in the ice matrix. Thus, CO₂ may be bound more in clathrate crystals than in ice. In addition, the solution of gas in clathrate crystals into the ice matrix is much less dependent on hydrostatic pressure than in bubbles, following Equation (A3).

Dividing the permeation coefficient by the solubility gives the diffusion coefficient. Unfortunately, the solubility of CO₂ in ice is not well known at present, as it is too small to be measured precisely (Hondoh, 1996). We calculate the CO₂ diffusion coefficient in the ice matrix at –23°C to be >1.3 × 10⁻¹³ m² s⁻¹ for a solubility of CO₂ in ice <4.5 × 10⁻⁴ mol m⁻³ at –23°C, using the permeation coefficient divided by the modeled diffusion coefficient of 7.8 × 10⁻¹⁰ m² s⁻¹ (Ikeda-Fukazawa, 2004, table 3).

### Smoothing of the CO₂ record in polar ice

Significant diffusion for thousands of years in the ice sheet might smooth rapid atmospheric changes and this would be extremely important in interpreting ice-core records. Well-known smoothing processes include gas diffusion in the firm layer and gradual bubble close-off at the transition from firm to ice, the effect of which can be roughly approximated by a Gaussian filter corresponding to the gas-age distribution (e.g. Schwander and others, 1988; Trudinger and others, 2002; Spahni and others, 2003).

Based on our observations and modeling, the smoothing of the CO₂ concentration by diffusion in deep ice is of the order of a few centimeters in our samples and, thus, negligible compared to the smoothing by the gas-age distribution at that depth (~30 years, corresponding to diffusion in the depth interval of ~100 cm). However, at greater depths the smoothing by CO₂ diffusing through the ice matrix may become larger.

A 10 year instantaneous atmospheric CO₂ spike (corresponding to ~1.55 m at the bubble close-off depth) is modeled with the permeation coefficient obtained in the shallow Siple Dome ice. The diffusion coefficient may significantly increase with greater depth due to geothermal warming. Results from the molecular simulation suggest that the diffusion coefficient approximately doubles for each 20°C increase in temperature (Ikeda-Fukazawa and others, 2004) as shown in Table 2. At a depth of 960 m (~80 kyr) at a location near the Siple Dome core site, the ice temperature reaches ~4°C (269 K) (Engelhardt, 2004), 19°C higher than at the shallow depth where we estimated the CO₂ permeation coefficient. Combining the depth-temperature profile with the temperature dependence of the diffusion coefficient (Ikeda-Fukazawa and others, 2004), we calculated permeation coefficients for modeled depths. Solubility of CO₂ in ice was assumed constant because the solubility/temperature relation is unknown.

The results from the 80 kyr simulations for two different estimations of thinning factor suggest that diffusion in deep ice may smooth the CO₂ concentration profile on decadal timescales, and at the age of ~60–70 kyr (Siple Dome depths of ~930–950 m) may be comparable to smoothing by diffusion in firm (Fig. 5). There are no decadal CO₂ data for ice that is 80 kyr old. However, the CO₂ record from the Siple Dome ice core shows significant variation of CO₂ on millennial timescales for the past 40 kyr (Ahn and others, 2004). Ice cores from colder sites than Siple Dome would experience slower CO₂ diffusion in deep ice. The formation of clathrate ice (bubble-free ice) at depths from 500 to 1200 m (~25–65 kyr) at other Antarctic cold-drilling sites (Vostok, Dome Fuji and EPICA Dome C) is expected to result in highly reduced gas diffusion (Salamatin and others, 1998).

### DISCUSSION AND DIRECTIONS FOR FUTURE WORK

The processes of gas diffusion related to variable physical properties of ice are still not well known. Thus, our volume-diffusion model should be investigated further. As discussed in the previous section, our interpretation of the noble-gas species is limited due to our lack of knowledge of their diffusion properties. Nonetheless, our study provides an important upper limit on the CO₂ permeation coefficient in ice cores. The true value of the permeation coefficient could be lower than we estimate if micro-melt layers around the visible melt layers contribute to the excess CO₂ in our data. Processes other than volume diffusion may be important but are difficult to quantify. For example, there is evidence of the existence of melt at triple junctions of grain boundaries in polar ice (Mulvaney and others, 1988). Thus, CO₂ may dissolve and migrate in the liquid vein, while noble-gas species, with lower solubility, may mostly stay at the original sites. If this is the case, the diffusion via the liquid vein or ice grain boundaries may be governed directly by the grain-growth rate, as suggested from an ion chemistry study (Barnes and others, 2003).

Our extension of the modeling results at shallow depth (corresponding to 2.74 kyr BP) to greater depths is limited by uncertainties in basic parameters and requires further study.

### Table 2. Temperature dependence of CO₂ diffusion coefficient in ice

<table>
<thead>
<tr>
<th>Ice-core location</th>
<th>T_surface</th>
<th>D</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>°C</td>
<td>m² s⁻¹</td>
</tr>
<tr>
<td>Dye 3, Greenland</td>
<td>–19.6</td>
<td>8.52 × 10⁻¹¹</td>
</tr>
<tr>
<td>Siple Dome, Antarctica</td>
<td>–25.4</td>
<td>7.24 × 10⁻¹¹</td>
</tr>
<tr>
<td>Byrd Station, Antarctica</td>
<td>–28</td>
<td>6.71 × 10⁻¹¹</td>
</tr>
<tr>
<td>Taylor Dome, Antarctica</td>
<td>–42</td>
<td>4.34 × 10⁻¹¹</td>
</tr>
<tr>
<td>Dome C, Antarctica</td>
<td>–54</td>
<td>2.85 × 10⁻¹¹</td>
</tr>
<tr>
<td>Vostok, Antarctica</td>
<td>–55.5</td>
<td>2.70 × 10⁻¹¹</td>
</tr>
<tr>
<td>Dome Fuji, Antarctica</td>
<td>–58</td>
<td>2.45 × 10⁻¹¹</td>
</tr>
</tbody>
</table>

Note: \(D\) (diffusion coefficient) is calculated from \(D = D_o \exp(-Q/R/T)\), where \(D_o = 9.10 \times 10^{-8} \text{ m}^2 \text{ s}^{-1}\) at ~8°C, using the permeation coefficient divided by the modeled diffusion coefficient of 7.8 × 10⁻¹⁰ m² s⁻¹ (Ikeda-Fukazawa and others, 2004).
For example, the assumption of Henry’s law for the CO₂ solubility in ice may not be valid through all pressure ranges. The temperature dependence of CO₂ solubility in ice is not considered since it is unknown. The dependence of the diffusion coefficient on temperature should be constrained, based on observations. Moreover, better constraints on the porosity–depth profile, which is one of the key parameters in our model, are needed.

In addition, the permeation coefficient of CO₂ may vary from core to core and depth to depth due to the variable physical properties of the ice. Studies with ice samples from various physical conditions (e.g. temperature, pressure, crystal growth rate) with different ice cores will better constrain the permeation coefficient.

Our estimation of the CO₂ concentrations in the melt layers is based on the experimental results for ice samples that include both the melt and normal layers. We assume constant gas extraction efficiency (gas extracted ÷ gas in ice before extraction) for both the normal and melt layers. If the gas extraction efficiency varies with the size of the bubbles (small bubbles in the melt layers and large ones in the normal layers), our measurements are not precise enough to detect differences in efficiency. Other techniques that allow 100% extraction efficiency such as sublimation of ice (e.g. Gülük and others, 1998) or melting of ice (e.g. Kawamura and others, 2003) could improve the estimation.

Our results also imply the possibility of an artifact in δ¹³C of CO₂ records due to different diffusion rates of ¹²C and ¹³C. Another possible implication is change of CO₂ mixing ratio in ice cores during storage.

CONCLUSIONS

Refrozen melt layers in the Siple Dome ice core contain excess CO₂ due to the high solubility of CO₂ in the meltwater. Our analyses of samples from the Siple Dome ice core show a gradual decrease of CO₂ away from two refrozen melt layers. The excess CO₂, combined with noble-gas data (Xe/Ar, Kr/Ar) and chemical and electrical properties of the ice, suggest that an initial CO₂ spike diffused through the ice. By modeling the CO₂ molecular diffusion, we calculate the permeation coefficient (the product of solubility and the diffusion coefficient) of CO₂ in ice to be ~4 x 10⁻²² m² s⁻¹ mol Pa⁻¹ at -23°C. This rate indicates smoothing of the CO₂ record by diffusion is one to two orders of magnitude smaller than the smoothing by diffusion in the firm at the depth of 287 m (gas age ~ 2.74 kyr BP) in the Siple Dome ice, and so does not degrade the record. However, applying the permeation coefficient to greater depth (equivalent to tens of thousands of years) in the Siple Dome ice core suggests an impact on smoothing of the CO₂ records on a decadal scale. Processes other than volume diffusion may be important but are difficult to quantify. Further studies should include the mechanism of the diffusion, dependence of the diffusion coefficient on temperature and solubility of the gas in the ice. Formation of clathrate seems to significantly hinder the CO₂ diffusion and will help preserve atmospheric records.

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REFERENCES


**APPENDIX A**

**VOLUME-DIFFUSION MODEL**

Elevated CO₂ diffuses from a melt layer to bubbles within a normal layer through the ice matrix. The smoothing of the CO₂ concentration is calculated from molecular volume diffusion with constant mixing ratio, cᵢ, in a certain width of the melt layer (Neftel and others, 1983). The CO₂ flux by the diffusion is:

\[ j = -D \frac{dc_E}{dx_{eff}} \]  

where \( j \) (mol·m⁻²·s⁻¹) is the flux of CO₂ by diffusion, \( D \) (m²·s⁻¹) is the diffusion constant of CO₂ in ice, \( c_E \) (mol·m⁻³) is the concentration of CO₂ dissolved in the ice and \( x_{eff} \) (m) is the effective vertical distance accounting for thinning by ice flow. This is related to the original distance, \( x \), and the thinning factor, \( \alpha(t) \):

\[ x_{eff} = \alpha(t) x \]
as function of time and depth, \( p(z(t)) \), and \( \text{CO}_2 \) solubility in
ice, \( S \) (mol m\(^{-3}\) Pa\(^{-1}\)), according to Henry’s law:
\[
c_E = \frac{c_0 p^\phi}{RT} + c_E. \tag{A3}
\]
The total \( \text{CO}_2 \) concentration per unit volume of bubble ice, \( c_{\text{tot}} \) (mol m\(^{-3}\)), requires the porosity of the ice, \( \phi \), and is:
\[
c_{\text{tot}} = \frac{c_0 p^\phi}{RT} + c_E. \tag{A4}
\]
The diffusion equation then becomes:
\[
\frac{\partial c_{\text{tot}}}{\partial t} = \frac{1}{R \theta} \left( \frac{c_0 p^\phi}{T} \right) + \frac{1}{R \theta} \left( \frac{c_0 p^\phi}{T} \right) = - \frac{dj}{dx_{\text{eff}}}. \tag{A5}
\]
The middle part of Equation (A5) can be written as:
\[
\frac{1}{R \theta} \left( \frac{c_0 p^\phi}{T} \right) + \frac{\partial c_E}{\partial t} = \left[ \frac{1}{RT} \left( \frac{c_0 p^\phi}{T} \right) + \frac{\partial c_E}{\partial t} \right] + \left[ \frac{1}{RT} \left( \frac{c_0 p^\phi}{T} \right) \right] - \left[ \frac{1}{RT^2} \left( \frac{c_0 p^\phi}{T} \right) \right] + \left[ \frac{1}{RT^2} \left( \frac{c_0 p^\phi}{T} \right) \right]. \tag{A6}
\]
where we assume \( S \) is constant. The second and third terms
in square brackets in Equation (A6) are smaller than that in
the first square bracket with order of \( >O(10^2) \) and \( >O(10^4) \), respectively. Therefore, we simplify Equation (A6):
\[
\frac{1}{R \theta} \left( \frac{c_0 p^\phi}{T} \right) + \frac{\partial c_E}{\partial t} \approx \frac{1}{R \theta} \left( \frac{c_0 p^\phi}{T} \right) \tag{A7}
\]
As we found \( \frac{1}{R \theta} (\frac{c_0 p^\phi}{T}) \) is greater than the other terms in
Equation (A7) with order of \( O(10) \) for most times and
locations in ice for the 2740 year simulation, Equation (A6) can be written as:
\[
\frac{1}{R \theta} \left( \frac{c_0 p^\phi}{T} \right) + \frac{\partial c_E}{\partial t} \approx \frac{1}{R \theta} \left( \frac{c_0 p^\phi}{T} \right). \tag{A8}
\]
Also, the right-hand side of Equation (A5) becomes:
\[
- \frac{dj}{dx_{\text{eff}}} = - \frac{d}{dx_{\text{eff}}} \left( -D \left( \frac{\partial c_E}{\partial t} \right) \right) = D \left( \frac{d^2 c_E}{dx_{\text{eff}}^2} \right) = \frac{Dc_E}{\phi \alpha(t)^2} \left( \frac{d^2 c_E}{dx^2} \right). \tag{A9}
\]
Thus, Equation (A5) becomes:
\[
\frac{\partial c_E}{\partial t} = \frac{DSRT}{\phi \alpha(t)^2} \frac{d^2 c_E}{dx^2}. \tag{A10}
\]
For each time-step, we calculate the \( \text{CO}_2 \) mixing ratio in
bubble air for each \( n \)th position (\( C_n^{t+1} \)) with a constant
distance interval (\( \Delta x = 1 \) mm for the 2740 year simulation;
0.155 m for the 80 kyr simulation). Equation (A10) can be discretized as:
\[
C_n^{t+1} = \frac{DSRT \alpha(t)}{\phi \alpha(t)^2} (\Delta x)^2 (C_n^{t+1} - 2C_n^{t} + C_{n-1}^{t}) + C_n^{t}. \tag{A11}
\]
In the finite-difference equation, realistic \( \alpha(t) \), \( \phi \) and \( p \) are
estimated for each time-step. The thinning factor, \( \alpha(t) \), is
determined by two different methods: (1) assuming a
constant strain rate with depth, which is
\[
\alpha(t) = \frac{H - z(t)}{H}, \tag{A12}
\]
where \( H \) is the thickness of the ice sheet and \( z(t) \) is depth
below the surface, which is estimated from the independent
depth–gas age profile (Brook and others, 2005), and
(2) utilizing the paleo-accumulation rate estimated from
the isotopic temperature proxy (Brook and others, 2005):
\[
\alpha(t) = \frac{\text{annual thickness at depth from a depth–age profile}}{\text{(ice density at depth)}} \times \text{(snow accumulation rate (w.e.))}. \tag{A13}
\]
A realistic \( \phi \) is estimated from the gas content in ice of
\(~0.11 \text{ cm}^3 \text{g}^{-1} \text{STP} \) (Severinghaus and Battle, 2006) and
accumulated ice load at depths >300 m (personal
communication from J. Fitzpatrick, 2007), assuming that
the pressure of air in bubbles equilibrates with the confining
pressure. At shallow depth (<300 m), this assumption is not
valid and \( \phi \) is estimated using ice-density data (personal
communication from J. Fitzpatrick, 2007) at time \( t \):
\[
\phi(t) = 1 - \frac{p(t)}{\rho_{\text{ice}}} \left( \frac{\rho_{\text{ice}}}{917 \text{ kg m}^{-3} \text{ for bubble-free ice}} \right). \tag{A14}
\]
Hydrostatic pressure, \( p \), is estimated from depth–density
profiles (personal communication from J. Fitzpatrick, 2007).
For the 80 kyr modeling, we use a diffusion coefficient, \( D \),
which varies with temperature (Ikeda-Fukazawa and others,
2004, table 3).

The constant parameters used for the Siple Dome ice
studied are:
\[
H = \text{thickness of the ice sheet} = 1003.8 \text{ m}
\]
\[
R = \text{gas constant} = 8.314 \text{ J mol}^{-1} \text{ K}^{-1}
\]
\[
T = \text{absolute temperature} = 250 \text{ K} \text{ (–23°C)}
\]
\[
S = \text{solubility of } \text{CO}_2 \text{ in ice} = 6.45 \times 10^{-11} \text{ mol m}^{-3} \text{ Pa}^{-1}
\]
\[
\Delta t = \text{time interval} = 3.16 \times 10^7 \text{ s (1 year) for 2740 year simulation}
3.16 \times 10^9 \text{ s (10 years) for 80 kyr simulation}
\]
\[
\Delta x = \text{distance interval} = 0.001 \text{ m for 2740 year simulation}
0.0155 \text{ m for 80 kyr simulation}
\]

Thickness of \( \text{CO}_2 \) spike (or melt layer) at surface
\[
= 0.018 \text{ m for M1}
0.012 \text{ m for M2}
\]

Thickness of 10 year \( \text{CO}_2 \) spike for 80 kyr simulation
\[
= 1.55 \text{ m}.
\]

APPENDIX B

VOLUME RATIO OF REFROZEN LIQUID WATER
TO BUBBLE AIR

We have modeled the effects of the formation of melt layers
on \( \delta \text{Kr/Ar, } \delta \text{Xe/Ar} \) and \( \delta^{36}\text{Ar/}^{38}\text{Ar} \) in air bubbles in ice cores.
The detailed model will be reported elsewhere. Briefly, we
assume that \( ^{39}\text{Ar, } ^{40}\text{Ar, Kr and Xe in air measured in an ice-
core sample are affected only by gravitational settling, melt} \)
and gas loss as follows (units are moles of gas):

\[
36\text{Ar}_{\text{sample}} = 36\text{Ar}_{\text{air bubble}} + 36\text{Ar}_{\text{dissolved}}/C_0 36\text{Ar}_{\text{lost}}
\]

\[
40\text{Ar}_{\text{sample}} = 40\text{Ar}_{\text{air bubble}} + 40\text{Ar}_{\text{dissolved}} - 0.993(36\text{Ar}_{\text{lost}})
\]

\[
K_{\text{r sample}} = K_{\text{r air bubble}} + K_{\text{r dissolved}}
\]

\[
X_{\text{e sample}} = X_{\text{e air bubble}} + X_{\text{e dissolved}}
\]

We also assume that all the dissolved gas is retained during the refreezing process. We use this model to invert the $\delta$Kr/Ar, $\delta$Xe/Ar and $\delta^{40}$Ar/$^{36}$Ar measurements for the firn diffusive column depth, $z$, gas loss during ice-core storage, $F$, and the volume ratio of refrozen liquid water to bubble air at ambient pressure, $\gamma$. We use the equations above, as well as the ideal gas law ($pV = nRT$) and the barometric equation ($p = p_0 \exp(mgz/RT)$), to derive the following system of equations, which can be solved by iteration for $\gamma$, $F$ and $z$ (all variables defined below):

\[
\frac{\delta^{40}\text{Ar} / ^{36}\text{Ar}}{1000} + 1 \text{ measured} = \frac{(1 - 0.993F) e^{m_{40}gz/RT} / RT + S_{40} \gamma}{(1 - F) e^{m_{36}gz/RT} / RT + S_{36} \gamma}
\]

\[
\frac{\delta\text{Kr} / \text{Ar}}{1000} + 1 \text{ measured} = \frac{e^{m_{40}gz/RT} / RT + S_{40} \gamma}{(1 - F) e^{m_{36}gz/RT} / RT + S_{36} \gamma}
\]

$S = \text{solubility (mol L}^{-1} \text{Pa}^{-1})$

$\gamma = \text{volume ratio of liquid water to pore air}$

$z = \text{depth of diffusive column (m)}$

$R = \text{gas constant (L Pa mol}^{-1} \text{K}^{-1})$

$T = \text{temperature (250 K)}$

$F = \text{fraction of Ar lost by gas loss when Kr and Xe are not being lost}$

$m = \text{mass (kg mol}^{-1})$

0.084 for Kr

0.040 for $^{40}$Ar

0.036 for $^{36}$Ar

0.132 for Xe

$g = \text{gravitational acceleration (9.82 m s}^{-2})$. 

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