

Photolysis of pyruvic acid in ice: Possible relevance to CO and CO₂ ice core record anomalies

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[1] The abnormal spikes detected in some CO and CO₂ polar ice core records indicate persistent chemical activity in glacial ice. Since CO and CO₂ spikes are correlated, and their amplitudes scale with reported CO/CO₂ yields for the photolysis of dissolved natural organic matter, a common photochemical source is implicated. Given that sufficient actinic radiation is constantly generated throughout ice by cosmic muons (Colussi and Hoffmann, 2003), it remains to be shown that the photolyses of typical organic contaminants proceed by similar mechanisms in water and ice. Here we report that the photodecarboxylation of pyruvic acid (PA, an ubiquitous ice contaminant) indeed leads to the same products nearly as efficiently in both media. CO₂ is promptly released from frozen PA/H₂O films upon illumination and continues to evolve after photolysis. By analogy with our studies in water (Guzmán et al., 2006b), we infer that ³PA* reacts with PA in ice producing CH₃C(O)C(O)O· and (CH₃ C(OH)C(O)OH) radicals. The barrierless decarboxylation, CH₃C(O)C(O)O· → CH₃C(O)· + CO₂, accounts for prompt CO₂ emissions down to ~140 K. Bimolecular radical reactions subsequently ensue in fluid molecular environments, both in water and ice, leading to metastable intermediates that decarboxylate immediately in water, but protractedly in ice. The overall quantum yield of CO₂ production in the λ ~313 nm photolysis of PA in ice at 250 K is ~60% of that in water at 293 K. The in situ photolysis of natural organic matter is, therefore, a plausible explanation of CO and CO₂ ice core record anomalies.

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1. Introduction

[2] Gas bubbles trapped in deep ice could be intact remnants of Earth's paleoatmospheres [Brook, 2005; EPICA, 2004; Fischer et al., 1999; North Greenland Ice Core Project Members, 2004; Stauffer, 2000; Wolff, 2005]. If so, the correlation of CO₂ concentrations in ancient ice bubbles with the coeval ambient temperatures reported by ¹⁸O or ²H abundance in the enclosing ice [Brook, 2005; Landais et al., 2006] would be a significant (although perhaps not definitive) [Boucot and Gray, 2001; Mudelsee, 2001; Pearson and Palmer, 2000] precedent to the potential climatic effects of contemporary anthropogenic CO₂ emissions.

[3] Air extracted from polar ice registers atmospheric CO₂ concentrations that have risen exponentially (from preindustrial ~280 ppmv levels) during the last two centuries. Notice that even higher CO₂ concentrations attended the transition from the last glaciation period to the Holocene [Tschumi and Stauffer, 2000]. Ominously, however, some records appear to be tainted by uncharacterized in situ processes. For example, Siple Dome CO₂ ice core records contain anomalous sections

that are not replicated in other Antarctic sites [Ahn et al., 2004]. Furthermore, Greenland records dating from the last two millennia display CO₂ concentrations that exceed (by up to ~20 ppmv) those in coeval Antarctic sites [Anklin et al., 1995; Haan and Raynaud, 1998]. These discrepancies (1) are localized in thin sections representing less than two annual periods, i.e., much shorter than the decadal intervals elapsing between snow deposition and gas encapsulation, (2) are larger than the precision of measurements (±3 ppmv), and (3) cannot be ascribed to interhemispheric gradients [Stauffer et al., 2002]. The international scientific community justifiably awaits a plausible explanation of this outstanding paleoclimatic issue.

[4] The irregular records reflect chemical activity under conditions that apparently exclude such possibility, because the production of carbon oxides requires their precursors to react while frozen at approximately -35°C, at hundreds of meters below the surface, in the complete absence of sunlight. Several mechanisms have been proposed to account for this phenomenon. It has been speculated that CO₂ could originate from the acidification of carbonate [Delmas, 1993; Stauffer et al., 1998] or the oxidation of organic matter by coextensive oxidizers, such as H₂O₂ [Tschumi and Stauffer, 2000].

[5] The likelihood that CO₂ spikes originate from the acidification of carbonates can be tested by comparing CO₂ versus carbonate profiles in ice cores, as proposed by Anklin

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et al. [1995] Such test has now been performed at Dome Fuji, East Antarctica, where CO₂ was found to be uncorrelated with either Ca²⁺, a proxy for carbonate, or acidity [Kawamura *et al.*, 2003]. Kawamura *et al.* estimate, from their measurements, that only 2–4% of the 10–20 ppmv excess CO₂ detected in 17–22 kyr old sections could have originated from the in situ acidification of carbonate [Kawamura *et al.*, 2003].

[6] The putative thermal oxidation of easily oxidizable aldehydes by H₂O₂ [Haan and Raynaud, 1998; Tschumi and Stauffer, 2000] is chemically unlikely for kinetic and mechanistic reasons. The uncatalyzed oxidation of formaldehyde by H₂O₂ (1) with an activation energy $E_a = 36 \text{ kJ mol}^{-1}$ is a slow reaction [$\exp(E_a / RT) = 1.4 \times 10^8$ at 240 K] [Do and Chen, 1994] and (2) in aqueous solution yields carboxylic acids (e.g., formic acid from formaldehyde) that are quite resilient to further oxidation, instead of carbon oxides [McElroy and Waygood, 1991; Satterfield and Case, 1954]. The detection of unreacted H₂O₂ [Hutterli *et al.*, 2003; Sigg and Neftel, 1991] and organic matter in close proximity within ice cores after thousands of years further undermines this proposal [Tschumi and Stauffer, 2000]. Finally, the large activation energy ($E_a \sim 110 \text{ kJ mol}^{-1}$) [Price and Sowers, 2004] associated with the metabolic activity of psychrophilic microbes drastically caps production rates of catabolic gases (such as CO₂ and CH₄) in anomalous ice core sections, which are much shallower and colder ($T < -35^\circ\text{C}$ down to $\sim 1000 \text{ m}$ depths) than Lake Vostok dome (nearly 4000 m deep) [Price *et al.*, 2002].

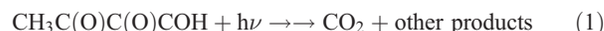
[7] Since CO cannot be produced by acidifying carbonate, the fact that some CO records are also exceedingly irregular [Haan and Raynaud, 1998] is a significant clue. The nearly constant $\sim 50 \text{ ppbv}$ CO levels detected in the Vostok (Antarctica) ice core are not replicated by Eurocore (Greenland) readings, which display greater variability and reach up to 180 ppbv in layers antedating $\sim 1500 \text{ AD}$ [Haan *et al.*, 1996; Haan and Raynaud, 1998; Tschumi and Stauffer, 2000]. Even more revealingly, Eurocore ice core CO and CO₂ profiles exhibit similar trends [Haan and Raynaud, 1998]. The latter observation strongly suggests a common source of contamination that, as a further constraint, should generate CO and CO₂ in $\sim 1:100$ ratios.

[8] At variance with thermal bimolecular thermal reactions, photochemical transformations are temperature-independent unimolecular events. The possibility of photochemistry in deep ice has been unwittingly disregarded in the past, however, due to the lack of conceivable sources of in situ actinic radiation. The detection of ultraviolet radiation generated by penetrating muons of cosmic origin in deep Antarctic ice [Andres *et al.*, 2001; Burden and Hieftje, 1998; Cowen *et al.*, 2001] and the finding that Greenland ice cores are significantly contaminated with organic matter from medieval boreal forest fires [Kawamura *et al.*, 2001; Savarino and Legrand, 1998] lend verisimilitude to a photochemical mechanism [Colussi and Hoffmann, 2003].

[9] Humic and humic-like substances [Gao and Zepp, 1998; Johannessen and Miller, 2001; Kieber *et al.*, 1989; Mopper *et al.*, 1991; Moran and Zepp, 1997; Opsahl and Zepp, 2001; Valentine and Zepp, 1993; Zuo and Jones, 1996], widespread in natural waters, atmospheric aerosol, and polar ice [Calace *et al.*, 2001; Grannas *et al.*, 2006; Grannas *et al.*, 2004; Grannas *et al.*, 2002; Kawamura *et al.*,

et al., 2005; Kawamura *et al.*, 2003; Kawamura *et al.*, 2001] are known to be photodegraded by sunlight into CO and CO₂ in $\sim 1:50$ ratios, which are tantalizingly commensurate with their relative excesses in anomalous ice core records (vide supra). The preceding analysis suggests that the photodecomposition of humic-like chromophores sporadically injected as contaminants into glacial ice is potentially able to simultaneously explain CO and CO₂ anomalous spikes.

[10] Organic photochemistry in ice is, however, a largely unexplored subject. More specifically, information is lacking about whether a typical photodecarboxylation reaction would proceed by the same mechanism and produce CO₂ in similar quantum yields in ice and water. Here we report a comparative study of the photodecarboxylation of the environmentally ubiquitous pyruvic acid (2-oxopropanoic, PA) [Kawamura *et al.*, 2001] in fluid and frozen aqueous solutions, reaction (1) [Budac and Wan, 1992; Guzmán *et al.*, 2006a; Guzmán *et al.*, 2006b; Wan and Budac, 1995]:



which provides factual support to our hypothesis.

2. Experimental Section

[11] One hundred millimolar PA (Aldrich 98.0%, bidistilled at reduced pressure) solutions in Milli-Q water were acidified to pH 1.0 with perchloric acid (Mallinckrodt, 70% analytical reagent) prior to photolysis to ensure that all (>98%) of the photogenerated CO₂ would be released into the gas-phase after thawing the photolyzed samples. It should be emphasized that the photolysis of PA ($\text{p}K_a = 2.39$) actually involves the undissociated acid rather than its anion [Leermakers and Vesley, 1963]. We confirmed that quantum yields of PA [Closs and Miller, 1978] and benzoylformic acid (BF) [Gorner and Khun, 1999] photodecarboxylations are independent of photolyte concentration. Fluid or frozen solution samples (4 mL) were photolyzed in a reactor consisting of a cylindrical chamber fitted with an axial silica finger housing a UV pencil lamp (Hg Pen-Ray model CPQ 8064, emitting at $313 \pm 20 \text{ nm}$) and connected to the infrared cell (CaF₂ windows, 10-cm optical path) of a Bio-Rad Digilab FTS-45 FTIR spectrometer for the continuous monitoring of CO₂ via online absorption infrared spectrophotometry at 2349 cm^{-1} . The gases contained in the (reactor + IR cell) assembly were recirculated (by means of a Schwartzel model 135 FZ micropump) every $\sim 4 \text{ s}$. The reactor was immersed in a thermostat/cryostat (Neslab ULT-80) for temperature control. Photolyses involved irradiation of 1-mm-thick ice layers produced by freezing aqueous solutions (4 mL, previously sparged with 1 atm N₂ or O₂ at 293 K for 30 min) onto the reactor chamber walls. Ice layers were maintained at target temperatures for at least 2 hours prior to photolysis. Sample absorption and lamp emission spectra were recorded with a HP 8452A diode array spectrophotometer. Quantum yield (ϕ_{CO_2}) measurements were based on the concerted photodecarboxylation of 10 mM BF (Aldrich 97%, used as received) reaction (2):



as chemical actinometer [Gorner and Khun, 1999]. The concertedness of reaction (2) makes its course independent of the medium. Aqueous solutions of the radical scavenger 2,2,6,6-tetramethylpiperidin-1-oxyl (TEMPO, Aldrich 99%, purified by vacuum sublimation) were used in some experiments [Guzmán *et al.*, 2006b]. Control dark experiments confirmed that PA and BF are thermally stable under present reaction conditions.

[12] Organic product analysis were performed via high performance liquid chromatography (HPLC) at ambient temperature, using UV and electrospray ionization-mass spectrometric detection (Agilent 1100 series Model 1100 Series HPLC-MSD System), immediately after sample melting, as described in detail elsewhere [Guzmán *et al.*, 2006b].

3. Results and Discussion

3.1. CO₂ Photogeneration

[13] Figure 1 shows the CO₂ released during and after irradiation of frozen aqueous PA layers maintained at 248 K. Additional CO₂ is liberated upon melting the ice layers. The CO₂ values plotted in Figure 1 correspond to moles of CO₂ divided by the fluid volume of PA layers (4×10^{-3} L). We observed that CO₂ is immediately released and continues to evolve at nearly constant rates, upon irradiation at temperatures down to ~ 130 K. The latter is an upper limit imposed by instrumental sensitivity and corresponds to the temperature at which the sublimation pressure of solid CO₂ falls below detection limits [Guzmán *et al.*, 2006a]. Postirradiation CO₂ releases display single exponential growth kinetics: $\text{CO}_2(t) = \text{CO}_2(\infty) \times [1 - \exp(-k_D t)]$, in isothermal experiments (Figure 2). Neither CO nor CH₄ were detected as gas-phase products in these experiments. Rate constants k_D (Figure 3) display Arrhenius temperature dependence:

$$\log(k_D/\text{s}^{-1}) = 1.08 - 1191/T \quad (3)$$

between 227 and 269 K which corresponds to an apparent activation energy: $E_{a,D} = 22.8$ kJ mol⁻¹, and A-factor: $A_D = 12.1$ s⁻¹.

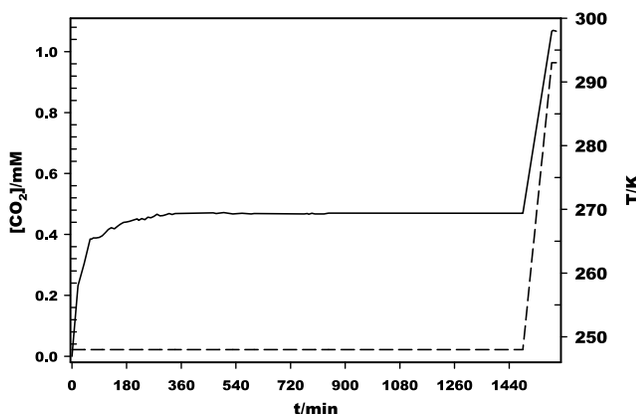


Figure 1. Solid trace (left axis), CO₂ released in the $\lambda \sim 313$ nm photolysis of frozen, deaerated aqueous pyruvic acid (4 mL, 100 mM, pH 1.0) solutions. CO₂ is released during the initial 60-min irradiation period and also in the post-illumination stage. Additional CO₂ is liberated upon thawing. Dashed trace (right axis), sample temperature.

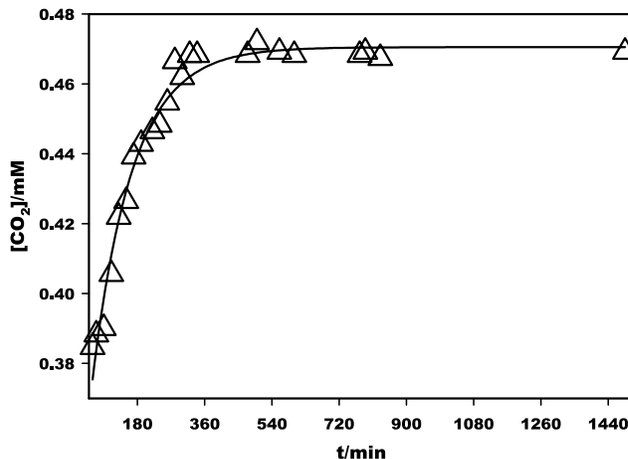


Figure 2. (Open triangle) A zoom of postirradiation CO₂ data from the experiment of Figure 1. Solid trace: $(\text{CO}_2) = A + B(1 - \exp(-k_D \times t))$; $A = 0.304$ mM; $B = 0.167$ mM; $k_D = 9.307 \times 10^{-3}$ min⁻¹.

[14] In a typical experiment, ~ 1.54 μmoles of CO₂ are released at the end of photolysis at 248 K. Additional CO₂ (up to a total of 1.87 μmoles of CO₂) is slowly released from irradiated frozen samples maintained at the same temperature. Further CO₂ evolves after thawing. Thus, only $\sim 1\%$ of the 400 μmoles of PA contained in the frozen layers is photolyzed in these experiments. About 50% of the photo-generated CO₂ ultimately escapes from frozen layers via a porous network open to the atmosphere, while the rest remains indefinitely trapped in the solid. The CO₂ amounts released after each stage are plotted as function of temperature in Figure 4. The apparent enthalpy change associated with overall CO₂ production, i.e., with the total CO₂ amounts recovered after melting, $\Delta H = (6.4 \pm 1.6)$ kJ mol⁻¹, is suggestively similar to the enthalpy of ice melting: $\Delta H_M = 6.3$ kJ mol⁻¹.

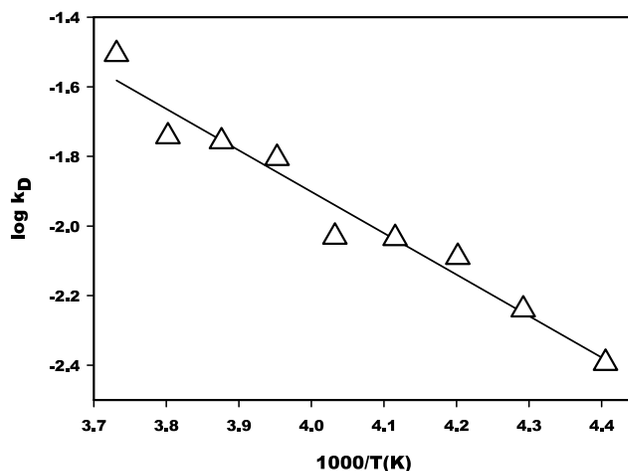


Figure 3. (Open triangle) Experimental first-order rate constants k_D for the release of CO₂ from previously photolyzed frozen pyruvic acid solutions. Solid trace: $\log(k_D/\text{s}^{-1}) = 1.08 - 1191/T$.

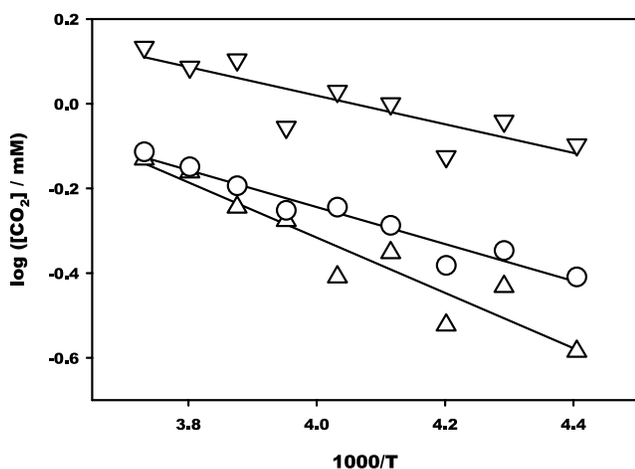


Figure 4. CO_2 released in the photolysis of frozen, deaerated aqueous pyruvic acid (4 mL, 100 mM, pH 1.0) solutions as function of temperature. (Open triangle) Immediately after photolysis; (Open circle) before sample thawing; (Open inverted triangle) after sample thawing.

[15] Post-illumination CO_2 emissions from irradiated frozen PA solutions are not due to the slow desorption of preformed CO_2 but arise instead from an ongoing thermal reaction. This conclusion follows from the fact that post-illumination CO_2 emissions are absent in the photolysis of frozen BF solutions (Figure 5). The implication is that PA photolysis in ice proceeds, as in fluid solutions [Guzmán *et al.*, 2006b], via reactive intermediates and/or unstable species. In other words, unless further CO_2 were produced in the dark, the CO_2 photogenerated in ice cavities connected to the gas-phase is released fully and promptly (i.e., within a few seconds). The stepwise warming of pre-irradiated frozen PA samples further reveals that the CO_2 plateaus preceding each temperature jump in Figure 5 do not imply that the source of dark CO_2 has been exhausted,

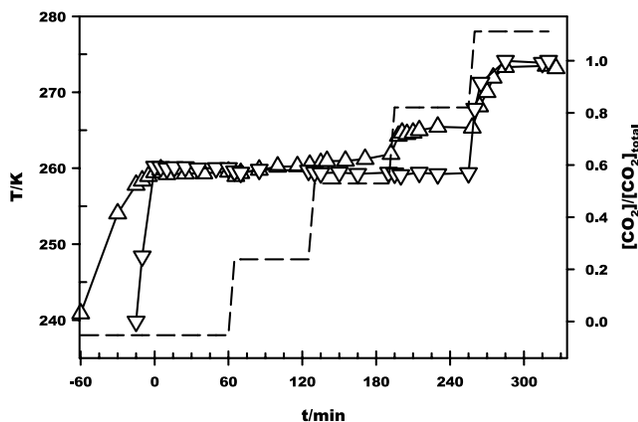


Figure 5. Symbols and solid traces (right axis), CO_2 fractions released immediately after (open triangle) 60 min photolysis of frozen, deaerated aqueous pyruvic acid (4 mL, 100 mM, pH 1.0) and (open inverted triangle) 15 min photolysis of frozen, deaerated aqueous benzoylformic acid (4 mL, 10 mM, pH 1.0). Dashed trace (left axis), sample temperature.

because additional CO_2 is chemically generated upon rising the temperature.

[16] Another mechanistic test is provided by the differential effect of radical scavengers on PA photolysis in ice and water. The photolysis of PA in water is known to be partially inhibited by radical scavengers [Guzmán *et al.*, 2006b], such as the persistent nitroxide 2,2,6,6-tetramethylpiperidin-1-oxyl (TEMPO) [Beckwith *et al.*, 1992; Bowry and Ingold, 1992; Fischer, 2001]. The fact that photochemical CO_2 production rates in aqueous PA solutions drop by a factor of two for $[\text{TEMPO}] \geq 1.7$ mM [Guzmán *et al.*, 2006b] proves that half the CO_2 is produced in a fast pathway that cannot be intercepted by TEMPO, while the remainder arises from a slower process involving scavengeable radical intermediates. In contrast, the addition of TEMPO to pre-frozen PA solutions has no effect whatsoever on CO_2 production rates during and after the photolysis of frozen PA solutions frozen under 1 atm N_2 (Figure 6). The efficiency of TEMPO as a radical scavenger is seemingly forfeited in frozen media by mobility restrictions. However, $\text{O}_2(\text{g})$ still behaves as a weak inhibitor both in fluid [Gorner and Khun, 1999] and frozen solutions; CO_2 production rates in the photolysis of frozen PA solutions performed under 1 atm O_2 are $\sim 25\%$ slower than under 1 atm N_2 at 253 K.

[17] Quantum yields for the overall production of CO_2 in reaction (1), ϕ_{CO_2} , in fluid and frozen aqueous PA solutions were determined from (1) measured rates of CO_2 evolution above aqueous solutions under continuous illumination and (2) the CO_2 amounts recovered after thawing frozen solutions that had been irradiated at constant temperature for specified periods of time, respectively. The effective absorption coefficients, ϵ_{eff} 's, of PA and BF in these experiments were evaluated from the convolution of their experimental absorption spectra in aqueous solution with the emission spectrum of the lamp. Since the extent of the

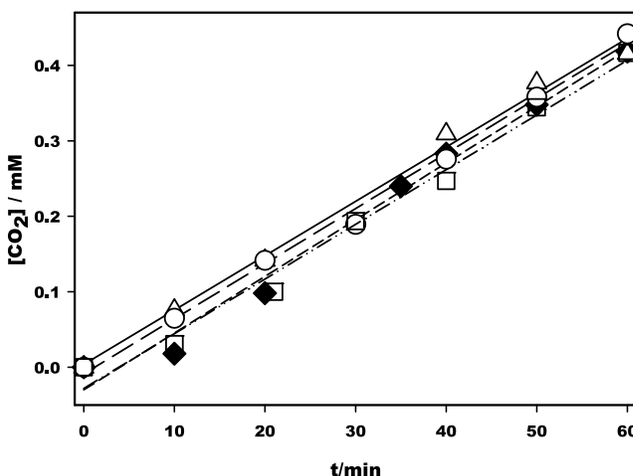
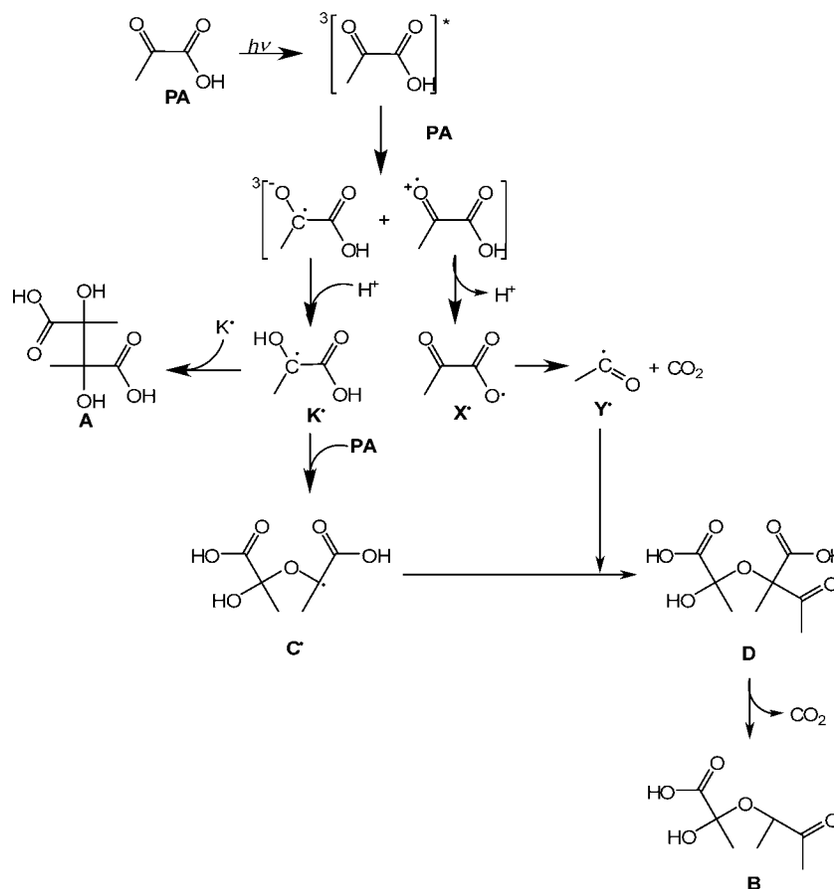


Figure 6. CO_2 released during irradiation of deaerated aqueous pyruvic acid (4 mL, 100 mM, pH 1.0) doped with TEMPO frozen at 253 K. (Open triangle) without TEMPO; (solid diamond) (TEMPO) = 0.253 mM; (open circle) (TEMPO) = 0.994 mM; (open square) (TEMPO) = 2.403 mM. Although TEMPO partially inhibits PA photolysis in water in this concentration range [Guzmán *et al.*, 2006b], it has no effect on PA photolysis in ice.



Scheme 1. Mechanism of pyruvic acid photolysis in ice.

partial hydration of keto-PA (*k*-PA), the actual chromophore, into transparent (above $\lambda > 300$ nm) 2,2-dihydroxypropanoic acid (*h*-PA) is an inverse function of temperature, the ϵ_{eff} of PA determined in water at 293 K was scaled down [$\epsilon_{\text{eff}} (<270 \text{ K}) = 0.69 \times \epsilon_{\text{eff}}(293 \text{ K})$] in frozen solutions according to the experimental ratios: $[h\text{-PA}]/[k\text{-PA}] = 2.54$ (293 K) and 4.10 ($T < 270 \text{ K}$) measured by $^1\text{H-NMR}$ and reported elsewhere [Buschmann *et al.*, 1982; Buschmann *et al.*, 1980; Guzmán *et al.*, 2006c]. BF is barely hydrated under present conditions [Fleury *et al.*, 1977]. The assumption can be made that the quantum yield of CO_2 formation in the concerted photodecarboxylation of BF ($\phi_{\text{CO}_2, \text{BF}} = 0.39$ at pH 3.0, 298 K) [Gorner and Khun, 1999; Kuhn and Gorner, 1988] does not change with physical state or temperature (c.f. with the intramolecular photoisomerization of 2-nitrobenzaldehyde) [Dubowski *et al.*, 2001]. The only reason for using PA and BF solutions of dissimilar concentrations is that they lead to comparable photolysis rates under present experimental conditions. Photolysis experiments in fluid and frozen BF or PA solutions led to: $\phi_{\text{CO}_2} = 0.78 \pm 0.10$ in water at 293 K (in excellent agreement with Leermakers and Vesley [1963], but a factor of two larger than our previously reported value) [Guzmán *et al.*, 2006b] and to the expression:

$$\log \phi_{\text{CO}_2} = (0.81 \pm 0.33) - (338 \pm 81)/T \quad (4)$$

in frozen PA solutions down to 227 K. Equation (4) leads to an extrapolated value of $\phi_{\text{CO}_2} = 0.45$ at 293 K, i.e., only

$\sim 40\%$ smaller than in water ($\phi_{\text{CO}_2} = 0.78$). These results confirm that the photodecarboxylation of PA, a typical ice core organic contaminant, proceeds with similar efficiency in ice and water.

3.2. Mechanism of Reaction

[18] The similar ϕ_{CO_2} values measured above and below the freezing point, and the fact that the organic products detected after thawing photolyzed frozen aqueous PA solutions are identical to those previously reported for the photolysis of fluid solutions, i.e., 2,3-dimethyl tartaric (A) and 2-(3-oxobutan-2-yloxy)-2-hydroxypropanoic acid (B) (Scheme 1) [Guzmán *et al.*, 2006b], indicate that PA photolysis follows the same course in fluid and frozen solutions. The initial chemical transformation results from electron transfer between the $^3\text{PA}^*$ excited state and an adjacent PA molecule [Guzmán *et al.*, 2006a], followed by fast proton transfers that lead to the ketyl $\text{K}\cdot$ and carbonyloxy $\text{X}\cdot$ radicals. While $\text{K}\cdot$ is stable and can engage in bimolecular reactions, the decarboxylation of the acetyl-carbonyloxy radical $\text{X}\cdot$ into acetyl radical $\text{Y}\cdot$:



occurs within a few vibrational periods even at low temperatures [Abel *et al.*, 2003; Bockman *et al.*, 1997]. The prompt release of CO_2 in the photolysis of frozen PA solutions down to at least $\sim 130 \text{ K}$ [Guzmán *et al.*, 2006a]

and the existence of a CO₂ fraction that cannot be scavenged by TEMPO in aqueous solution are both accounted for by reaction 5. Paramagnetic species disappear irreversibly from irradiated frozen aqueous PA solutions above ~190 K [Guzmán *et al.*, 2006a], i.e., all radicals ultimately recombine within nominally frozen matrices. While the direct recombination (K· + K· → A) leads to a 2,3-dimethyl tartaric acid, the addition of K· to PA yields yet another radical: K· + PA → C·. The recombination (C· + Y· → D) produces the C₈ β-oxocarboxylic acid D, whose slow decomposition gives rise to delayed CO₂ emissions.

[19] Scheme 1 comprises bimolecular events that entail large amplitude molecular motions, such as librations and center-of-mass translations, which can only take place in sufficiently fluid reaction media. The fact that ~2 mM TEMPO is not able to inhibit CO₂ emissions in 100 mM PA photolysis in ice, although it is effective in water, suggests that the allowed translational motions have a rather short range and only involve close molecular neighbors. The question of whether the requisite bimolecular reaction can be still invoked at the vastly smaller overall concentrations of organic matter present in natural ice cores deserves closer scrutiny (*vide infra*).

[20] The Arrhenius parameters: $E_{a,D} = 22.8 \text{ kJ mol}^{-1}$, $A_D = 12.1 \text{ s}^{-1}$, associated with k_D , the pseudo first-order rate constant for the decomposition of D in ice, are much smaller than those typical of the thermal decarboxylation of β-oxocarboxylic acids in aqueous solution ($E_a \sim 95 \text{ kJ mol}^{-1}$, A-factor $\sim 10^{12} - 10^{13} \text{ s}^{-1}$) [Guthrie, 2002]. $A_D = 12.1 \text{ s}^{-1}$ represents an impossibly large negative activation entropy for an elementary reaction: $\Delta S \sim -230 \text{ J K}^{-1} \text{ mol}^{-1}$ [Andraos, 2000; Benson, 1976]. Notice that $E_{a,D} = 22.8 \text{ kJ mol}^{-1}$ is nearly identical to the strength of the hydrogen bond in ice ($\sim 21 \text{ kJ mol}^{-1}$) [Cotton and Wilkinson, 1980; Galwey *et al.*, 2001]. However, as pointed out above, these parameters must be associated with a chemical reaction (the thermal decarboxylation of D) rather than with physical desorption of preformed CO₂. The production and release (from prephotolyzed frozen PA solutions) of limited amounts of CO₂ after discrete temperature jumps (Figure 4) are typical of the kinetic behavior displayed by intrinsically fast chemical reactions hindered by a cryogenic matrix [Andraos, 2000; Spath and Raff, 1992]. Since the decarboxylation of β-ketoacids is a highly concerted process involving four (or more) internal coordinates [Guthrie, 2002; Huang *et al.*, 1997], it is quite conceivable that a rigid ice matrix could block its decomposition. That the decomposition of D is actually controlled by the ice matrix is confirmed by the fact that the half-life, $\tau_D = 0.69 / k_D > 10 \text{ min}$, in water at 293 K extrapolated from equation (3), should have led to measurably slow CO₂ emissions from thawed irradiated samples, at variance with observations.

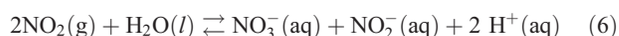
3.3. Reaction Environments in Natural Versus Laboratory Ices

[21] An anonymous reviewer has argued that conclusions derived from present experiments are inapplicable to natural ices because (1) glacial ice is produced by the sintering of snow rather than by freezing aqueous solutions, (2) organic matter is likely to settle onto glacial ice carried by the atmospheric aerosol, and (3) the typical micromolar concentrations of individual organic acids in thawed natural

ices [Savarino and Legrand, 1998] are orders of magnitude smaller than that used in this study.

[22] Natural and laboratory ices may differ in many ways, but what really matters in the present context are the molecular (nanoscopic) domains sensed by the solutes themselves rather than the macroscopic/mesoscopic ice textures and morphologies perceived by observers or revealed by microscopy at submicrometer resolution [Baker *et al.*, 2005; Killawee *et al.*, 1998]. Furthermore, the purported disparity in overall concentrations is rather incidental to ice photochemistry because it effectively refers to the disparity of solute concentrations measured in the fluids obtained by thawing natural versus laboratory ices, rather than to the local concentrations in the molecular domains where solutes reside and where photochemical processes actually take place in ice. In fact, in the presence of ice, the activity of water (and, hence, of the solute) in the unfrozen portion becomes an exclusive function of temperature, independent of the initial concentration of the aqueous solution [Koop, 2002]. In other words, when dilute solutions are cooled to (externally imposed) subfreezing temperatures, they freeze out more water than concentrated solutions (the lever rule [Reisman, 1970]), and vice versa, to comply with this thermodynamic constraint [Koop, 2002]. Intensive variables, such as solute concentration in the unfrozen portion, depend on temperature and pressure. Only the volume fraction of the unfrozen portion is proportional to total amount of solute in the sample. We recently confirmed that the activity of water, as probed by magic angle spinning ¹H-NMR measurements of the “gem”-diol PA/keto PA ratio in frozen aqueous PA solutions, becomes independent of the initial concentrations of aqueous PA solutions cooled below ~250 K [Guzmán *et al.*, 2006c]. The contention that solute concentration in the unfrozen portion somehow depends on the initial concentration of the aqueous solution is, therefore, untenable. Practical considerations (such as duration of experiments × detection limits) dictate the use of larger solute concentrations in laboratory experiments.

[23] Size restrictions allow only small species to be incorporated as substitutional impurities into the ice phase. Most organic and inorganic solutes [Killawee *et al.*, 1998] (particularly the macromolecular humic-like substances relevant to this discussion) will be segregated from the ice phase into highly concentrated fluid or glassy media upon freezing [Boxe *et al.*, 2005; Boxe *et al.*, 2006; Boxe *et al.*, 2003; Heger *et al.*, 2005]. Recently, it was found that by slowly freezing 0.1 μM aqueous methylene blue (MW = 374 Da, a fair surrogate of dissolved natural organic matter) solutions, the solute is concentrated by at least 6 orders of magnitude in the unfrozen portion remaining at 243 K [Heger *et al.*, 2005]. In a related study, we found that the second order hydrolysis of the NO₂ produced in situ during the photolysis of nitrate in ice, reaction 6:



is still significant in frozen micromolar nitrate solutions [Boxe *et al.*, 2006]. Therefore most molecular ice impurities are generally expected to be located in low-dimensional fluid phases at concentrations that are vastly larger than

those measured in thawed ice core samples. The acidities experienced in ice at the molecular level are also remotely related to those of the thawed samples and are determined by specific ion combinations [Heger *et al.*, 2006; Robinson *et al.*, 2006].

[24] Still, as previously pointed out, most solutes probably land on surface snow (and become embedded in glacial ice after several decades) effectively encapsulated in prefrozen atmospheric aerosol microdroplets. If these microbeads were to remain intact, solute molecules would not be incorporated as isolated impurities in the ice matrix by this mechanism either, because typical 1- μm radii aqueous droplets consist of $\sim 2 \times 10^{11}$ H_2O molecules or $\sim 2 \times 10^5$ molecules of a solute present at ppm levels (PA > 18 ppm in urban aerosol at 50% relative humidity [Guzmán *et al.*, 2006b; Kawamura *et al.*, 2005; Kawamura and Yasui, 2005]). Furthermore many organic solutes are expected to crowd during freezing, driven by intermolecular interactions such as hydrophobic forces in the case of nonpolar species or hydrogen bonding in the case of PA dimers. At the modest droplet cooling rates prevailing in the atmosphere ($\sim 1 \text{ K min}^{-1}$ [Haag *et al.*, 2003] versus the 10^5 K min^{-1} rates required to hyperquench water into amorphous glassy water [Bednarek *et al.*, 1996]) departures from equilibrium solute distributions are expected to be minor [Haag *et al.*, 2003; Stuart and Jacobson, 2006] and relax during the subsequent firnification process. Notice that the extended hydraulic networks filled by concentrated solutions detected in glacial ice [Nye, 1989] could only be created from a material consisting of randomly distributed concentrated spots (such as that produced by the deposition of frozen aerosol beads on ground snow) via strain-induced melting and recrystallization. The existence of quasi-fluid aqueous domains of nanoscopic dimensions containing significant numbers of solute molecules in close proximity, as strictly required by the mechanism of Scheme 1, seems ensured, therefore, under natural and laboratory conditions.

3.4. Photochemical Origin of CO and CO₂ Ice Core Records Irregularities

[25] It has been generally assumed that given the prevalent low temperatures [Price *et al.*, 2002], and the fact that sunlight is completely blocked within a few meters below the surface by the highly reflective snow/firn cover) [Belzile *et al.*, 2000; Gerland *et al.*, 1999; Glendinning and Morris, 1999; Phillips and Simpson, 2005] species immobilized in deep ice core layers, i.e., below the firn line, will indefinitely retain their chemical identity. Thus the fact that CO and CO₂ levels in Greenland ice core records dating from the last two millennia exceed those in contemporaneous Antarctic records by up to 130 ppbv and 20 ppmv (i.e., well beyond their experimental uncertainties), respectively [Haan and Raynaud, 1998; Tschumi and Stauffer, 2000], is an unexpected finding that seems to undermine the archival value of polar records and demands a plausible explanation. The anomalies are not restricted to interhemispheric differences because CO₂ concentrations in Siple Dome, Antarctica, from certain periods of the Holocene are also larger than in other Antarctic sites by up to 20 ppmv [Ahn *et al.*, 2004]. The fact that outliers appear as sharp spikes superimposed on otherwise normal time series rules out analytical errors and militates against diffusional processes within the ice matrix.

At stake is the issue of whether these anomalies are exceptional and, if so, why.

[26] Several hypothesis have been advanced to account for the abnormal records [Monnin *et al.*, 2001]. “The sudden occurrence of high CO concentrations at about 155 m depths suggests the presence of organic species that are able to produce carbon monoxide” [Haan and Raynaud, 1998]. “The most probable explanation for CO₂ peak values in thin core sections representing only a few annual layers, or even less than an annual layer, specially in Greenland ice, is production of CO₂ by chemical reactions between impurities in the cold ice” [Tschumi and Stauffer, 2000]. Possible CO₂ production mechanism include [Ahn *et al.*, 2004] (1) acidification of carbonates [Anklin *et al.*, 1995; Barnola *et al.*, 1995; Delmas, 1993], (2) chemical [Tschumi and Stauffer, 2000], or (3) biological oxidation of organic compounds [Campen *et al.*, 2003; Mader *et al.*, 2006; Price and Sowers, 2004].

[27] Both polar ice caps are known to be contaminated by organic material or continental or marine origin [Desideri *et al.*, 1998; Gayley and Ram, 1985; Kawamura *et al.*, 1995], but Greenland ice cores are conspicuously spoiled by the scavenging of organic gases and deposition of the organic aerosol released by major, epochal boreal forest fires [Domine and Shepson, 2002; Kawamura *et al.*, 1995; Kawamura *et al.*, 2001; Savarino and Legrand, 1998]. Full speciation of the organic matter present in polar ice remains a daunting analytical challenge [Grannas *et al.*, 2006], but it is generally agreed that organic contaminants largely consist of humic-like substances, such as those globally present in natural waters, the simpler products of their solar photodegradation [Gao and Zepp, 1998; Miller and Zepp, 1995; Moran and Zepp, 1997; Opsahl and Zepp, 2001; Valentine and Zepp, 1993], as well as the products of the photooxidation of atmospheric organic substances, such as dicarboxylic, α -oxocarboxylic and ω -oxocarboxylic acids, and α -dicarbonyl species [Kawamura *et al.*, 2005; Kawamura *et al.*, 1996; Kawamura *et al.*, 2001; Wang *et al.*, 2006]. Pyruvic acid, 2-oxopropanoic acid, is a ubiquitous representative of these species.

[28] The photochemical degradation of dissolved organic matter in the atmosphere, surface waters, and snow [Grannas *et al.*, 2004; Osburn *et al.*, 2001] yields CO [Mopper *et al.*, 1991; Schade *et al.*, 1999; Valentine and Zepp, 1993] and CO₂ [Johannessen and Miller, 2001]. Quantum yields of CO₂ and CO photoproduction, which depend somewhat on the origin of dissolved organic matter, decrease exponentially at longer wavelengths [Johannessen and Miller, 2001; Valentine and Zepp, 1993]. Relative yields vary in the range $20 \leq \phi_{\text{CO}_2} / \phi_{\text{CO}} \leq 50$. The possibility that photochemical processes similar to those occurring in surface waters could occur in deep ice, and therefore contaminate paleorecords, has been implicitly discounted because of the high reflectivity of the snow cover to ultraviolet light, the lack of a conceivable in situ source of actinic radiation and the absence of documented examples of photoinduced decarboxylations/decarbonylations in frozen aqueous media. Still, “even the highly improbable must be considered over the long timescales involved in ice core chemistry. Organic acids and other C-containing species present in sufficient quantities have the potential

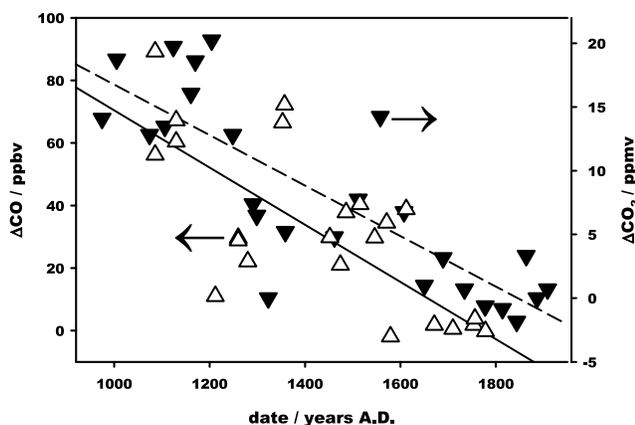


Figure 7. (Solid inverted triangle) Differences between CO_2 (ppmv) readings from Greenland (Eurocore, GRIP) and Antarctic (South Pole, D47, D57) ice core records (right axis) versus date (Anklin *et al.*, 1995). (Open triangle) Differences between CO (ppbv) readings from Greenland (Eurocore, GRIP) relative to the constant value (89 ppbv) for the period 1640–1870 AD, (left axis) versus date [Haan and Raynaud, 1998].

to influence CO and CO_2 , if a reasonable mechanism were proposed” [Wolff, 1996].

[29] We have recently proposed and quantified the likelihood that the dissolved organic matter trapped in ice cores could be photodegraded by the ultraviolet Čerenkov radiation generated by cosmic muons [Colussi and Hoffmann, 2003]. By assuming that the quantum yields of decarboxylation of dissolved organic matter were similar in ice and water, the timing and magnitude of the CO anomalies in Greenland versus coeval Antarctic ice core bubbles dating from the Middle Ages could be plausibly correlated with the occurrence of boreal fires [Savarino and Legrand, 1998]. This analysis revealed that significant photochemistry could be induced in deep ice by Čerenkov radiation over geological periods. Despite the small local photon production rates, the exceptional transparency of pristine glacial ice between 200 and 500 nm [Askebjør *et al.*, 1997a; Askebjør *et al.*, 1997b] allows chromophoric point impurities, such as those created by sporadic contamination from dissolved organic matter, to be photochemically processed not only by locally generated photons but also from those created elsewhere and collected onto the absorbing layer after diffusing through the ice matrix. A strongly absorbing sink effectively acts as an antenna whose collection efficiency is proportional to the ratio of the thickness of the pristine layer to the effective thickness of the layer containing the organic chromophore. This ratio depends, in general, on dust levels in the ice, as well as on chromophore concentration and absorptivity [Colussi and Hoffmann, 2003].

[30] A comparison between CO [Haan and Raynaud, 1998] and CO_2 excesses [Anklin *et al.*, 1995] in Eurocore sites versus Antarctic profiles as function of gas age (Figure 7) shows that they not only follow similar trends, but the ratio of the slopes: $S_{\text{CO}_2} / S_{\text{CO}} \sim 200$, falls in the range of the ratio of quantum yields, $\phi_{\text{CO}_2} / \phi_{\text{CO}}$, for the production of dissolved inorganic carbon (that should appear as CO_2 in ice) and CO in the solar photolysis of dissolved organic

matter in natural waters [Gao and Zepp, 1998; Johannessen and Miller, 2001; Kieber *et al.*, 1989; Mopper *et al.*, 1991; Moran and Zepp, 1997; Opsahl and Zepp, 2001; Valentine and Zepp, 1993; Zuo and Jones, 1996]. A more stringent comparison should, of course, involve ϕ 's for the photolysis of dissolved organic matter in ice by (Čerenkov) radiation in the range $200 < \lambda/\text{nm} < 400$, rather than in water under solar ($\lambda > 300 \text{ nm}$) radiation. Notice that increasing deviations in older ice deposits support continuous photochemistry under an in situ incessant radiation source, such as that provided by Čerenkov radiation, but rules out photochemistry in the upper ice layers for the limited periods during which they are exposed to sunlight. Present results and analysis, together with the data of Figure 7, represent strong evidence that the in situ photochemistry of organic ice contaminants is a physically and chemically plausible mechanism that could account, in principle, for the observed ice core records anomalies.

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