

ISOTOPIC EXCHANGE BETWEEN CARBON DIOXIDE AND OZONE VIA O(<sup>1</sup>D) IN THE STRATOSPHERE

Yuk L. Yung, W.B. DeMore, and Joseph P. Pinto

**Abstract.** We propose a novel mechanism for isotopic exchange between CO<sub>2</sub> and O<sub>3</sub> via O(<sup>1</sup>D) + CO<sub>2</sub> → CO<sub>3</sub><sup>\*</sup> followed by CO<sub>3</sub><sup>\*</sup> → CO<sub>2</sub> + O(<sup>3</sup>P). A one-dimensional model calculation shows that this mechanism can account for the enrichment in <sup>18</sup>O in the stratospheric CO<sub>2</sub> observed by Gamo *et al.* [1989], using the heavy O<sub>3</sub> profile observed by Mauersberger [1981]. The implications of this mechanism for other stratospheric species and as a source of isotopically heavy CO<sub>2</sub> in the troposphere are briefly discussed.

## Introduction

Recently Gamo *et al.* [1989] reported measurements of heavy isotopes of CO<sub>2</sub> in the stratosphere over Japan. Let O = <sup>16</sup>O and Q = <sup>18</sup>O. The δQ of stratospheric CO<sub>2</sub>, as defined in Gamo *et al.*, was found to be 2‰ (two parts in a thousand) greater than that of tropospheric CO<sub>2</sub> at 19 km, and appears to continue to increase with altitude up to the highest altitude (25 km) where samples were taken. The data are reproduced in our Figure 2. The authors proposed two possible explanations, one dynamical and one chemical. The preferred mechanism was chemical and involved transferring a heavy oxygen atom from O<sub>3</sub>, which was first measured to be isotopically enriched by Mauersberger [1981], to carbon dioxide. However, the authors did not propose a detailed chemical scheme. The direct thermal exchange reaction CO<sub>2</sub> + O<sub>2</sub>Q → COQ + O<sub>3</sub> is inefficient (J.-S. Wen, private communication, 1990).

We propose, instead, that the transfer of Q from O<sub>3</sub> to CO<sub>2</sub> is initiated by O(<sup>1</sup>D) reactions. We will use a one-dimensional photochemical model incorporating reactions for exchange of Q to simulate the vertical profile of COQ observed by Gamo *et al.* The factors controlling the COQ/CO<sub>2</sub> ratio in the troposphere have been studied by Keeling [1961], Bottinga and Craig [1969], Francey and Tans [1987], and Friedli *et al.* [1987]. We will attempt to assess the importance of exchange with the stratosphere relative to these other factors.

## Chemical Kinetics

Photolysis of O<sub>3</sub> below about 305 nm produces O(<sup>1</sup>D), an excited state of the oxygen atom, by R1 (see listing and numbering of reactions in Table 1). The primary fate of O(<sup>1</sup>D) in the atmosphere is quenching by the ambient atmospheric gases (R2a). Because of the efficiency of the quenching reactions, the abundance of O(<sup>1</sup>D) in the atmosphere is very small and it has not been directly detected in the atmosphere. Nevertheless, this excited atom plays fundamental roles in the chemistry of HO<sub>x</sub> radicals [Levy, 1971; Wofsy *et al.*, 1972] and NO<sub>x</sub> species [McElroy and McConnell, 1971; Nicolet and Peetermans, 1972]. Figure 1 presents the O(<sup>1</sup>D) concentration predicted by the

<sup>1</sup>Division of Geological and Planetary Sciences, California Institute of Technology, Pasadena, California 91125.

<sup>2</sup>Jet Propulsion Laboratory, 4800 Oak Grove Drive, Pasadena, California 91109.

<sup>3</sup>AREAL, United States Environmental Protection Agency, Research Triangle Park, North Carolina 27711.

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Table 1

List of essential reactions used in the photochemical model. The units for photodissociation coefficients and two-body rate constants are s<sup>-1</sup> and cm<sup>3</sup> s<sup>-1</sup>, respectively. The coefficients for photolysis refer to mid-latitude diurnally averaged values at 30 km. All molecular kinetic data are taken from DeMore *et al.* [1990], except otherwise stated. Rate constants for isotopic species are estimated by us. Q = <sup>18</sup>O.

R1a	O <sub>3</sub> + hv → O <sub>2</sub> + O( <sup>1</sup> D)	J <sub>1a</sub> = 7.4 × 10 <sup>-5</sup>
R1b	O <sub>2</sub> Q + hv → O <sub>2</sub> + Q( <sup>1</sup> D)	J <sub>1b</sub> = 1/2 J <sub>1a</sub>
R2a	O( <sup>1</sup> D) + M → O + M	k <sub>2a</sub> = 2.0 × 10 <sup>-11</sup> e <sup>110/T</sup>
R2b	Q( <sup>1</sup> D) + M → Q + M	k <sub>2b</sub> = k <sub>2a</sub>
R3	O( <sup>1</sup> D) + CO <sub>2</sub> → CO <sub>2</sub> + O	k <sub>3</sub> = 7.4 × 10 <sup>-11</sup> e <sup>120/T</sup>
R4a	Q( <sup>1</sup> D) + CO <sub>2</sub> → COQ + O	k <sub>4a</sub> = 2/3 k <sub>3</sub>
R4b	O( <sup>1</sup> D) + COQ → CO <sub>2</sub> + Q	k <sub>4b</sub> = 1/3 k <sub>3</sub>
R5	O( <sup>1</sup> D) + CO → CO + O	k <sub>5</sub> = 4.6 × 10 <sup>-11</sup> (a)
R6a	Q( <sup>1</sup> D) + CO → CQ + O	k <sub>6a</sub> = 1/2 k <sub>5</sub>
R6b	O( <sup>1</sup> D) + CQ → CO + Q	k <sub>6b</sub> = 1/2 k <sub>5</sub>
R7	QH + CO <sub>2</sub> → COQ + OH	k <sub>7</sub> < 1 × 10 <sup>-17</sup> (b)
R8	Q + CO <sub>2</sub> → COQ + O	k <sub>8</sub> < 1 × 10 <sup>-18</sup> (c)

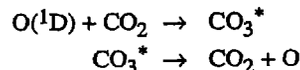
(a) Cvetanovic [1975].

(b) Greenblatt and Howard [1989].

(c) Wen and Thiemens [1990].

Caltech-JPL model [Froidevaux *et al.*, 1985], a typical diurnally-averaged one-dimensional photochemical model, along with our O<sub>3</sub> profile.

Consider the reaction between O(<sup>1</sup>D) and CO<sub>2</sub>



The existence of the CO<sub>3</sub><sup>\*</sup> complex was first proposed by Katakis and Taube [1962] in a study of the exchange between O(<sup>1</sup>D) and CO<sub>2</sub>. Jacox and Milligan [1971] favored a three-member ring structure with an O-C-O angle of 65°. The complex rapidly predissociates to CO<sub>2</sub> + O [DeMore

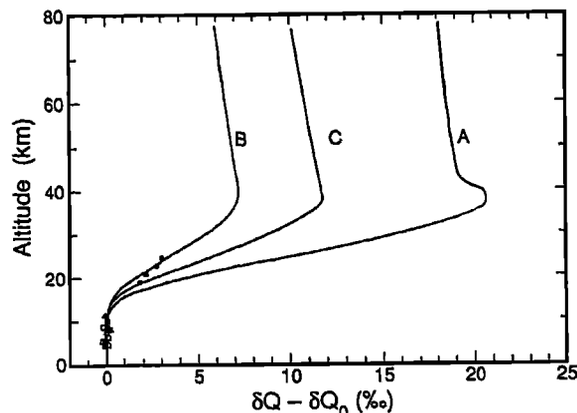


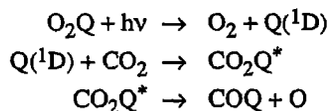
Fig. 1. Concentrations of O(<sup>1</sup>D) and O<sub>3</sub> from the Caltech-JPL one-dimensional photochemical model [Froidevaux *et al.*, 1985].

and Dede, 1970]. There has been great interest in  $\text{CO}_2^*$  in connection to the  $\text{CO}_2$  stability problem on Mars [McElroy and Hunten, 1970; Noxon, 1970], which was subsequently solved by a different scheme [Parkinson and Hunten, 1972; McElroy and Donahue, 1972].

The rapid quenching of  $\text{O}(^1\text{D})$  by  $\text{CO}_2$  [DeMore *et al.*, 1990] with rate coefficient,  $k_3 = 1.1 \times 10^{-10} \text{ cm}^3 \text{ s}^{-1}$  at  $T = 298 \text{ K}$ , undoubtedly proceeds via the  $\text{CO}_2^*$  intermediate [DeMore and Dede, 1970]. For comparison we note that the corresponding rate constants for quenching of  $\text{O}(^1\text{D})$  by Ar and Kr are  $7 \times 10^{-13}$  and  $8 \times 10^{-12} \text{ cm}^3 \text{ s}^{-1}$ , respectively [Cvetanovic, 1975]. The existence of  $\text{CO}_2^*$  would be expected to result in an exchange between the incident and emergent O atoms. Indeed, in an isotopically labelled study using  $\text{CO}_2$  photolysis at 147 nm as the source of  $\text{O}(^1\text{D})$ , Baulch and Breckenridge [1966] showed that the efficiency yield for isotopic exchange is 0.69, close to the expected value of 2/3 if the ejection of an O atom from  $\text{CO}_2^*$  were a purely statistical process. Other kinetic studies [Yamazaki and Cvetanovic, 1964; Weissberger *et al.*, 1967] have generally confirmed this result.

The startling discovery of an  $^{18}\text{O}$  enrichment in stratospheric  $\text{O}_3$  by Mauersberger [1981] has been confirmed [Carli *et al.*, 1982; de Zafra *et al.*, 1984; Mauersberger, 1983, 1987; Rinsland *et al.*, 1985; Goldman *et al.*, 1989], and studied in the laboratory [Heidenreich and Thiemens, 1986; Thiemens and Jackson, 1987; Yang and Epstein, 1987a,b; Morton *et al.*, 1989; Anderson *et al.*, 1989]. The theory of isotopic enrichment for  $\text{O}_3$  was initiated even before the atmospheric observations [Cicerone and McCrumb, 1980], but this and all subsequent theories [Kaye and Strobel, 1983; Blake *et al.*, 1984; Kaye, 1986, 1987] completely fail to account for either the atmospheric or laboratory measurements. The suggestions of Valentini *et al.* [1987] and Bates [1986] are not widely accepted. It is not the purpose of this paper to enter this great debate. We accept as empirical fact that stratospheric  $\text{O}_3$  is enhanced in  $^{18}\text{O}$  and pursue its implications for the isotopic enrichment of  $\text{CO}_2$  and other species.

Consider the following sequence of reactions initiated by the photolysis of a heavy ozone molecule,



The net result is the transfer of Q from ozone to carbon dioxide. The reverse process can also readily occur via R4b, leading to the transfer of Q from carbon dioxide back to ozone. Therefore, ozone and carbon dioxide may set up some kind of kinetic isotopic equilibrium in the stratosphere. Under the assumptions of *photochemical equilibrium* and using the somewhat idealized chemical scheme summarized in Table 1, it can be shown that  $[\text{COQ}]/[\text{CO}_2] = 2([\text{Q}(^1\text{D})]/[\text{O}(^1\text{D})]) = [\text{O}_2\text{Q}]/[\text{O}_3]$ . On the other hand, this ratio would be equal to that at the ground if the atmosphere were well-mixed (i.e., transport time  $\ll$  chemical time). In other words, the ratio  $\text{COQ}/\text{CO}_2$  in the atmosphere reflects a "tug-of-war" between photochemical equilibrium driven by UV radiation in the upper atmosphere, and reactions involving exchange with  $\text{H}_2\text{O}$  in the biosphere, the oceans, and clouds.

For our theory to be correct we must prove that other isotopic exchange reactions such as R7 with QH and R8 with Q are *not* important. Evidence for R7 was found by Kurylo and Laufer [1979], but Greenblatt and Howard [1989] reported an upper limit of  $k < 1 \times 10^{-17} \text{ cm}^3 \text{ s}^{-1}$ , making it unimportant for the stratosphere. Jaffe and Klein [1966] reported  $k_8 = 1.1 \times 10^{-12} e^{-1750/T} \text{ cm}^3 \text{ s}^{-1}$  and this would suggest that R8 is fast enough to neutralize the isotopic

enrichment produced by  $\text{Q}(^1\text{D})$ . However, we believe that the results of Jaffe and Klein are in error. First, it is extremely improbable that  $k_8 > 100 k_7$ , since OH is generally more reactive than O. Second, a qualitative upper limit for  $k_8$  may be obtained by modeling an experiment performed by Wen and Thiemens [1990]. Wen (1990, private communication) deduced  $k_8 < 1.5 \times 10^{-17} \text{ cm}^3 \text{ s}^{-1}$  at  $T = 363 \text{ K}$ . We conclude that  $k_8 < 10^{-18} \text{ cm}^3 \text{ s}^{-1}$  at stratospheric temperatures, and hence R8 is not important.

### Photochemical Modeling

The Caltech-JPL one-dimensional photochemical model is described elsewhere [Allen *et al.*, 1981; Froidevaux *et al.*, 1985; Michelangeli *et al.*, 1989]. In the present investigation we fix the concentrations of  $\text{O}(^1\text{D})$  and  $\text{O}_3$  (see Figure 1) to the values in Froidevaux *et al.* [1985]. This is justified because  $\text{CO}_2$  has negligible influence on oxygen photochemistry. The relevant photochemistry of O,  $\text{O}(^1\text{D})$ ,  $\text{O}_3$ ,  $\text{CO}_2$ , CO, and their isotopes is summarized in Table 1. The rate coefficients for the isotopic species are for the most part unknown, and we have to make reasonable guesses to arrive at the preferred values given in Table 1. Vertical transport is parameterized by eddy diffusion.

Let the standard Q/O ratio be  $f$ . Since stratospheric  $\text{O}_3$  is enriched in Q, we have  $[\text{O}_2\text{Q}]/[\text{O}_3] = 2 f E$  where  $E =$  enrichment factor. Since  $\text{O}(^1\text{D})$  is in photochemical equilibrium with  $\text{O}_3$ , we have, assuming the reactions of Table 1,  $[\text{Q}(^1\text{D})]/[\text{O}(^1\text{D})] = f E$ . The choice of  $E$  in our model represents some difficulty. The observed enrichment factors are not reproducible, either due to uncertainties in the experiments or a possible dependence on the solar cycle (Mauersberger, 1990, private communication). It must be emphasized that there is no theoretical justification for a solar cycle dependence. In case A, we set the enrichment factor  $E_A$  to be the measurements of Mauersberger [1981]. To set a lower bound for  $E$  we also use the later measurements to derive another profile  $E_B = 1 + (E_A - 1)/3$ . All other uncertainties in the model are probably smaller than that of  $E$ .

The photochemical model was run to steady state in the diurnally averaged mode. We adopt  $f = 1/500$  [Kaye, 1987]. The lower boundary conditions at the ground are  $[\text{CO}_2] = 340 \text{ ppm}$  and  $[\text{COQ}] = 1.36 \text{ ppm}$ . At the upper boundary (80 km), the fluxes of  $\text{CO}_2$  and COQ are zero. In the standard run  $E = 1.0$ . In runs A and B we use as input the profiles  $E_A$  and  $E_B$ . The results for  $\delta\text{Q} - \delta\text{Q}_0$  for  $\text{CO}_2$  are shown in Figure 2 ( $\delta\text{Q}_0 =$  tropospheric value =  $40.7\text{‰}$  relative to SMOW at  $24^\circ\text{C}$ ), along with Gamo *et al.*'s measurements. It is clear that our mechanism yields the correct trend with altitude as well as the magnitude of isotopic enrichment below 30 km. Model B appears to be closer to the observations of Gamo *et al.* [1989], although data in the higher regions were not available. The profile of  $\delta\text{Q} - \delta\text{Q}_0$  is generally what we expected. The maximum effect occurs at 40 km where  $\text{O}(^1\text{D})$  concentrations are highest. In the troposphere the effect is overwhelmed by the ocean and the biosphere. The results above 60 km may not be reliable because we have completely ignored possible isotopic effects in  $\text{CO}_2$  photolysis in the mesosphere [Allen *et al.*, 1981].

To test the sensitivity of our model to transport, we double the eddy diffusion coefficients in model C, which is otherwise the same as model A. This result, shown in Figure 2, demonstrates the greater influence of the troposphere when mixing is increased. Similar runs were made for CO. The source of CO is assumed to be isotopically normal. The exchange of Q between ozone and carbon monoxide is less efficient than the loss of CO via OH. We obtained an isotopic enrichment of CQ of the order of  $1\text{‰}$ . Unfortunately, there are no observations of CQ in the stratosphere to compare with.

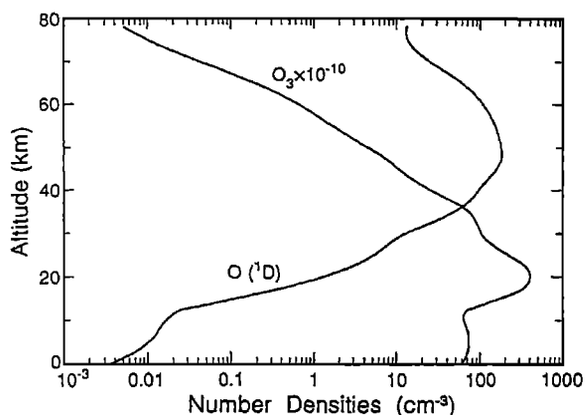


Fig. 2. Computed  $\delta Q - \delta Q_0$ . Cases A and B were computed using enrichment factors  $E_A$  and  $E_B$  for  $O_3$  described in the text. Case C is same as case A, except that the eddy diffusivity profile is doubled. The data are from Gamo *et al.* [1989].

### Discussion and Conclusion

The discovery of the  $\delta^{18}O$  anomalies in stratospheric  $O_3$  and  $CO_2$  opens a new and exciting chapter in atmospheric chemistry. For the first time we encounter chemical phenomena that are the result of subtle quantum kinetic effects which are still not fully understood. We now have further evidence on how the "heaviness" in ozone is transmitted to another species,  $CO_2$ . We believe that similar transmissions of  $^{18}O$  anomaly could occur to other stratospheric species that are connected to  $O_3$ ,  $O$ , and  $O(^1D)$ . However,  $CO_2$  is somewhat unique for enrichment by the present mechanism because of its rapid exchange with  $O(^1D)$  and its long lifetime.

It is of interest to note that not all stratospheric species can communicate isotopically. In a perceptive analysis Kaye [1990] pointed out that the enhancement of  $^{18}O$  in stratospheric  $O_3$  is *not* expected to be reflected in stratospheric  $H_2O$ , thus casting doubts on the claims of observations of enhancements of  $^{18}O$  in  $H_2O$  in the upper stratosphere [Guo *et al.*, 1989]. Recent accurate measurements of  $H_2O$  in the stratosphere by the ATMOS experiment (M. Gunson, private communication 1990) have brilliantly confirmed Kaye's insight.

As shown in the modeling section,  $\delta Q$  of  $CO_2$  is determined by transport and the abundance of  $O(^1D)$ . Thus, accurate simultaneous measurements of  $O_2Q$  and  $CO_2$ , together with precise determination of appropriate rate constants may provide a reliable method for estimating the mean concentration of  $O(^1D)$  in the stratosphere, a species that is otherwise exceedingly difficult to measure due to its low concentrations. This idea is similar to that of using methyl chloroform to estimate a mean global OH concentration in the troposphere [Singh, 1977; WMO 1986, Chapter 4]. The downward transport of  $CO_2$  may have implications for the interpretation of its abundance and latitudinal gradient in the troposphere.  $CO_2$  is observed to be heavier by a few tenths per mil at the South Pole, compared to the tropics and decreases even more rapidly towards the North Pole [Francey and Tans, 1987]. Assuming that stratospheric air entering the troposphere in tropopause folds in the Southern Hemisphere is enriched in  $^{18}O$  by about 1‰ and scaling the downward transport of  $CO_2$  to that for  $O_3$  given by Mahlman *et al.* [1980], we calculate an enrichment in  $^{18}O$  in  $CO_2$  of 0.25 to 0.5‰, in going from the tropics to the South Pole. This calculation assumes that the isotopic equilibration time with the oceans and clouds is much longer than the intrahemispheric mixing time of about 3–6 months [Friedli *et*

*al.*, 1987; Francey and Tans, 1987]. An even greater enhancement in  $CO_2$  should result in the high latitude Northern Hemisphere by this mechanism since the stratosphere-troposphere exchange in the northern hemisphere is twice as large as in the Southern Hemisphere. However,  $CO_2$  rapidly exchanges with liquid water in the terrestrial biosphere, resulting in even lower  $CO_2$  abundances at high northern latitudes compared to the tropics. The effects of stratospheric intrusions therefore may not be visible in the troposphere of the northern hemisphere except for short periods following injection. Measurements of the decay of excess stratospheric  $CO_2$  in the troposphere may give additional information regarding rates of exchange with the terrestrial biosphere compared with  $^{13}CO_2$ , which is currently used. In view of the importance of  $CO_2$  in controlling climate, the presence of another isotopomer must be welcome.

Other stratospheric species such as  $CO$  and  $N_2O$  may also exchange  $Q$  with  $O_3$ . However, in the case of  $CO$ , the calculated enrichment by this mechanism is small. The isotopic composition of stratospheric  $CO$  may be determined mainly with reactions in the methane oxidation chain.  $O(^1D)$  quenching by  $N_2O$  primarily results in a reaction. The upper limit for regenerating an oxygen atom is 4% [DeMore *et al.*, 1990]. Therefore, we expect negligible exchange of  $Q$  via this mechanism.

A number of important studies remain to be done. The chemical kinetics of isotopic species, summarized in Table 1, urgently needs to be refined by laboratory work. When properly modeled, the measurement of  $CO^{18}O$  in the stratosphere may offer the best estimate for the average value of  $O(^1D)$  concentrations in the stratosphere. Further laboratory studies are also required to clarify the mystery of  $^{18}O$  enrichment in  $O_3$  itself.

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W. B. DeMore, Jet Propulsion Laboratory, 4800 Oak Grove Drive, Pasadena, California 91109.

J. P. Pinto, AREAL, United States Environmental Protection Agency, Research Triangle Park, North Carolina 27711.

Y. L. Yung, Division of Geological and Planetary Sciences, California Institute of Technology, Pasadena, California 91125.

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