DISTRIBUTION OF GASEOUS  ${}^{12}$   $\infty_2$ ,  ${}^{13}$   $\infty_2$ , AND  ${}^{14}$   $\infty_2$  IN THE SUB-SOIL UNSATURATED ZONE OF THE WESTERN US GREAT PLAINS

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ABSTRACT. Data on the depth distribution of the major atmospheric gases and the abundance of gaseous  ${}^{12}\Omega_2$ ,  ${}^{13}\Omega_2$ , and 14 CO<sub>2</sub> in the subsoil unsaturated zone have been obtained from several sites in the western Great Plains of the United States. Sample profiles range from land surface to depths of 50m. Although each site must be considered on an individual basis, several general statements can be made regarding the profiles. 1) Diffusion of these gaseous molecules through the unsaturated zone is an important transport mechanism. 2) As predicted by diffusion theory, depth profiles of the various isotopic species of  $\Omega_2$  differ substantially from one another, depending on individual sources and sinks such as root respiration and oxidation of organic carbon at depth. 3) In general, post-bomb (> 100% modern)  $^{14}$ C activities are not observed in the deep unsaturated zone, in contrast to diffusion model predictions. 4) In spite of generally decreasing  $^{14}$ C activities with depth, absolute partial pressures of  $14 \, \mathrm{co}_2$  in the subsoil unsaturated zone are 1-2 orders of magnitude higher than the partial pressure of  $^{14}$   $^{12}$ in the atmosphere.

## INTRODUCTION

JL Kunkler (1969) performed the first <sup>14</sup>C analyses of  $CO_2$  gas from the deep unsaturated zone in the Bandelier Tuff at the Los Alamos National Laboratory. His data showed <sup>14</sup>C activities > 100% modern carbon (pmc) at depths of 24m and 86m, indicating that post-bomb  $CO_2$  had penetrated the tuff to at least that depth. Kunkler's research remained unique until recent studies on the unsaturated zone by Reardon, Allison, and Fritz (1979) and Reardon, Mozeto, and Fritz (1980). Data are presented here on the abundance and distribution of unsaturated-zone  ${}^{12}CO_2$ ,  ${}^{13}CO_2$ , and  ${}^{14}CO_2$ from several sampling sites in North Dakota and Texas.

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Three of the North Dakota sites were sampled repetitively over a five-year period; two of these sites are discussed in a preliminary attempt to evaluate the role of gaseous diffusion in unsaturated zone processes at these locales. All of the unsaturated zone  $\Omega_2$  data collected to date by the authors are presented. Although site-specific process models have not been formulated for each site, some important geochemical generalizations can be made.

## PHYSICO-CHEMICAL PRINCIPLES

For chemical modeling and equilibrium calculations involving gases, the gas partial pressure is the most useful variable. Alternatively, transport modeling and calculation of diffusive fluxes from Fick's laws require the abundance of diffusing species to be expressed in units of mass/volume (generally mol-cm<sup>-3</sup>). These statements hold for any isotopic species of the same gas, as well as for different gases, eg,  $1^4 \text{CO}_2$  will diffuse in response to its own concentration gradient regardless of the distribution of  $1^2 \text{CO}_2$  and  $1^3 \text{CO}_2$  in the gas mixtures<sup>\*</sup>. Outlined below are the methods for calculating partial pressures and concentrations from the original analytical data. For purposes of this paper, the behavior of all gases can be assumed to be ideal.

MAJOR GASES. The analytical data for the unsaturated zone gases are available as volume percent for N<sub>2</sub>, O<sub>2</sub>, Ar, and  $\Omega_2^{**}$ , as  $\delta^{13}C(^{O}/oo)$  for  $^{13}\Omega_2$ , and as A<sup>14</sup>C in percent modern carbon (pmc) for  $^{14}\Omega_2$ . Dissolved  $\Omega_2$  is initially reported as partial pressure. For N<sub>2</sub>, O<sub>2</sub>, Ar, and  $\Omega_2$  the volume percent equals mol percent, and the mol fraction of a given gas, X<sub>i</sub>, is simply X<sub>i</sub> = (vol%i/100).

<sup>\*</sup> The fact that different isotopes of a single chemical species must respond only to their own concentration gradients is perhaps best visualized through the randomwalk model for diffusive processes (Feynman, Leighton, and Sands, 1963, Chap 43). The independence of isotopic diffusion is the basis for tracer diffusion experiments and is implicit in the general equations for calculating the self-diffusion coefficients of one isotopic species in another (Bird, Stewart, and Lightfoot, 1960; Jost, 1960; Li and Gregory, 1974).

<sup>\*\*</sup> For simplicity, we use  $CO_2$  to refer to  ${}^{12}O_2 + {}^{13}O_2 + {}^{14}O_2$ (there is no isotopic separation in the gas chromatograph columns).

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The partial pressure of i is given by

$$P_{i} = X_{i} P_{total}^{\dagger}.$$
 (1)

Then, from the ideal gas law  $n_i/V = P_i/RT$ ,

$$C_{i}(mol-cm^{-3}) \equiv \frac{n_{i}(mol)}{V(cm^{3})} = \frac{X_{i}P_{total}(atm)}{R(cm^{3}-atm-K^{-1}-mol^{-1})T(K)} .$$
(2)

 $^{13}\text{C}$ . The molar ratio, R, of  $^{13}\ensuremath{\varpi}_2$  to  $\ensuremath{\varpi}_2$  in the PDB standard ( $\delta^{13}\ensuremath{\mathsf{C}}\xspace$  = 0) is 0.0111 (Landgren, 1954). Given this value, concentrations of  $^{13}\ensuremath{\varpi}_2$  can be calculated from  $\delta^{13}\ensuremath{\mathsf{C}}\xspace$  and total  $\ensuremath{\varpi}_2$  concentrations as follows:

$$\delta^{13}C \equiv \left\{ \frac{R_{sample}}{R_{reference}} - 1 \right\} 1000 = \left\{ \frac{R_{sample}}{0.0111} - 1 \right\} 1000.$$
(3)

Rearranging,

$$R_{sample} = \frac{C(^{13}\omega_2)}{C(\omega_2)} = 0.0111 \{ 1 + (\delta^{13}C/1000) \}, (4)$$

and

$$c(^{13}\omega_2) = [c(\omega_2)](0.0111) \{ 1 + (\delta^{13}c/1000) \}$$
. (5)

The relative gradients of  $^{13}\, \odot_2$  and  $\odot_2$  will therefore differ only by the factor (1+ $\delta^{13}\,C/1000$ ). Total variation of this factor through the range of  $\delta^{13}\,C$  values observed in this study amounts to < 2%, which is of about the same magnitude as the uncertainty in  $\odot_2$  collection and analysis. Therefore, transport modeling of  $^{13}\,\odot_2$  will produce results essentially identical to those for  $\odot_2$ , with calculated fluxes multiplied by 0.0111. The utility of the  $^{13}C$  signature as a means of identifying sources of carbon in the system remains undiminished, although the  $^{13}\,C$  variations are not emphasized in this report.

<sup>&</sup>lt;sup>†</sup> For the North Dakota sites, P<sub>total</sub> = 0.91 atm; at the Texas sites P<sub>total</sub> = 0.88 atm; and at Los Alamos, P<sub>total</sub> = 0.77 atm

14 C. For <sup>14</sup>C, the ratio  $R_{14}_{CO_2} = (\text{moles } ^{14} \varpi_2)/(\text{moles } \varpi_2)$ must be obtained from analytical values of  $A^{14}C(\text{pmc})^*$ . The quantity  $R_{14}_{CO_2}$  can be obtained from the decay equation  $-dN/dt = \lambda N$  where N is the number of <sup>14</sup>C atoms per gram of carbon, and the decay constant  $\lambda$  has the value  $3.84 \times 10^{-12}$ sec<sup>-1</sup>.

For a 100 pmc sample, the decay rate is 0.226 dps/g-C. In this case

$$N = \frac{-dN/dt}{\lambda} = \frac{0.226}{3.84 \times 10^{-12}} = 5.89 \times 10^{10} \text{ atoms } {}^{14}\text{C/g-C}, \quad (6)$$

or, introducing Avogadro's number and the atomic weight of carbon,

$$R_{14} co_2 = \frac{5.89 \times 10^{10}/6.023 \times 10^{23}}{1 / 12.011} = 1.17 \times 10^{-12} \frac{mol^{14}c}{mol c} \text{ at } 100 \text{ pmc.}$$
(7)

Then for a 1 pmc sample,  $R_{14}_{CO_2}$  would be  $1.17 \times 10^{-14} \text{ mol}^{14} \text{ C per}$  mol C, and for an arbitrary  $^{14}\text{C}$  activity

$$R_{14}_{CO_2} = (1.17 \times 10^{-14}) (A^{14} C(pmc)) mol^{14} C/mol C.$$
 (8)

The mole fraction of  ${}^{14} \infty_2$  in the original gas sample is then

$$X_{14_{CO_2}} = (R_{14_{CO_2}})(X_{CO_2}) = (1.17 \times 10^{-16}) (A^{14} C(pmc))(vol\% \ O_2).$$
(9)

that the  $\ensuremath{\mathfrak{O}_2}$  content of the gas phase is constant.

<sup>\*</sup> Reardon, Mozeto, and Fritz (1980) discussed the distribution of  $^{14}\, \varpi_2$  in the unsaturated zone as a function of  $R_{14}$ . Their conclusions will be correct to the extent  $\varpi_2$ 

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Partial pressures will be given by

$$P_{14} \omega_2 = X_{14} \omega_2 P_{total}$$
 (10)

Concentrations of  ${}^{14}$   $\infty_2$  will be given by combining equations (2) and (9), with R =  $82.05 \text{ atm-cm}^3\text{-mol}^{-1}\text{-K}^{-1}$ :

$$c_{14} c_{02} = \frac{\sum_{RT}^{X_{14}} c_{02}^{P_{total}}}{RT} =$$
 (11)

$$= \frac{(1.43 \times 10^{-18})(A^{14} C(pmc))(Vo1\% \Omega_2)(P_{total}(atm))}{T(K)} \text{ in } \frac{mo1^{14} \Omega_2}{cm^3}.$$

The values of  $A^{14}C(pmc)$  to be used in equation (11) must be uncorrected because no age estimation is intended, and the chemical parameters of interest require only raw data on the isotopic abundances.

## DIFFUSION MODELING

As mentioned above, each isotopic species of  $CO_2$  within the soil gas diffuses through the unsaturated zone according to its own concentration gradient as described by Fick's Second Law. However, for movement in the unsaturated zone, Fick's Second Law must be modified to account for the effects of the porous medium structure on the rate of diffusion and for reactions between the gas and the liquid and solid phases of the medium. With these modifications, the Law becomes

$$\tau \Theta_{\mathbf{D}} \mathbf{D}_{\mathbf{A}\mathbf{B}} \frac{\partial^2 \mathbf{C}_{\mathbf{A}}}{\partial \mathbf{x}^2} = \Theta_{\mathbf{D}} \frac{\partial \mathbf{C}_{\mathbf{A}}}{\partial \mathbf{t}} + \Theta_{\mathbf{w}} (\Theta_{\mathbf{T}} - \Theta_{\mathbf{D}}) \frac{\partial \mathbf{C}_{\mathbf{A}}^{\dagger}}{\partial \mathbf{t}} + \frac{\partial \overline{\mathbf{C}}_{\mathbf{A}}}{\partial \mathbf{t}} + \alpha_{\mathbf{T}} , \quad (12)$$

where

 τ = a tortuosity factor accounting for the added resistance to diffusion imposed by the structure of the porous medium (dimensionless);

- θ<sub>D</sub> = drained or gas-filled porosity (dimensionless);
- D<sub>AB</sub> = molecular diffusion constant for diffusion of gas A into gas B (cm<sup>2</sup>/sec);
- $C_A$  = concentration of gas A (mol/cm<sup>3</sup>);
- x = dimension increasing with depth; = 0 at land surface (cm);
- t = time (sec);
- $\rho_w$  = density of soil water (g/cm<sup>3</sup>);
- $\Theta_{T}$  = total porosity (dimensionless);
- CA = concentration of substance A transferred to the solid phase (mol/[cm<sup>3</sup> of medium], where [cm<sup>3</sup> of medium] refers to the space occupied by solids + liquids + gases);
- and  $\alpha_T$  = a production term for substance A [mol/(cm<sup>3</sup> of medium)/sec].

Equation (12) states that the rate of one-dimensional diffusion of gas A toward a given point minus the rate of diffusion from the point is equal to the combined rate of change in concentration in 1) the gas phase, 2) the liquid phase, as dissolved gas and any dissolved chemical species containing the gas molecule, 3) in the solid phase, plus 4) the amount of the gas produced or consumed at that point per unit time.

THE DIFFUSION COEFFICIENT. The diffusivity of  $\Omega_2$  in free air has been measured to be 0.144 cm<sup>2</sup>/sec at STP (Bird, Stewart, and Lightfoot, 1960), which is corrected within the program to ambient conditions by the equation

$$D_{AB} = D_{AB}^{\circ} (P_{O}/P) (T/T_{O})^{1.823} *, \qquad (13)$$

<sup>\*</sup> Equation (13) is based on Bird, Stewart, and Lightfoot (1960), p 505, eq 16.3-1.

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where  $D^{\circ}_{AB}$  = diffusion constant at STP (cm<sup>2</sup>/sec);

P<sub>o</sub> = standard atmospheric pressure (one atmosphere);

P = mean atmospheric pressure at site (atmospheres);

- T = mean annual air temperature at site (K);
- and  $T_0$  = temperature at standard conditions (273.2 K).

EVALUATION OF  $\partial C_A^{\dagger}/\partial t$  AND  $\partial \overline{C}_A^{\dagger}/\partial t$ . To evaluate the partitioning of  $\omega_2$  between the gas and liquid phase requires a functional relationship between  $P_{\omega_2}$  and the total dis-

solved  $\Omega_2$ . A simple function that can be used in the transport program becomes available if the aqueous chemistry in the unsaturated zone can be ascribed to few or one reaction. For this model, it is assumed that calcite is present throughout the unsaturated zone, and that carbonate chemistry in the water is governed by  $\Omega_2$  - calcite equilibria via reaction (14):

$$CaCO_{3(s)} + O_{2(g)} + H_{2}O_{(k)} = Ca^{2+}_{(aq)} + 2HO_{3(aq)}.$$
 (14)

The equilibrium constant for this reaction is

$$K_{eq} = \frac{a_{Ca}^{2+} a_{HOO_3}^2}{P_{OO_2}}, \qquad (15)$$

and if this is the only reaction considered,

$$m_{Ca}^{2+} = \frac{1}{2} m_{HOO_3}$$
 (16)

If we assume ideality  $(a_i = m_i)$ ,

$$K_{eq} = \frac{m^3_{H} \omega_3^{-}}{2 P_{\omega_2}}$$
, (17)

or,

$$m_{HCO_{3}(aq)} = (2 K_{eq} P_{CO_{2}})^{1/3} = 1.260 K_{eq}^{1/3} P_{CO_{2}}^{1/3}, (18)$$

where

 $m_i$  = molality and  $P_{OO_2}$  = partial pressure of  $OO_2$  in atm \*.

Use of the partitioning coefficient concept to model transport in the gas phase relies on determining the amount of the subject gas going into solution due to a change in concentration in the gas phase. In reaction (14) 1/2 of the aqueous  $CO_2$  comes from solid-phase carbonate dissolution, and the other half by solution from the gas phase. Thus, in equation (12),

$$\frac{\partial \overline{C}_{A}}{\partial t} = -\frac{1}{2} \rho_{w} (\Theta_{T} - \Theta_{D}) \frac{\partial C_{A}}{\partial t} , \qquad (19)$$

where the minus sign indicates that  $CaCO_{3(s)}$  decreases as  $CO_{2(g)}$  increases. The relation  $\partial C_A^{\dagger}/\partial t = (\partial C_A^{\dagger}/\partial C_A)(\partial C_A^{\dagger}/\partial t)$ , in conjunction with equation (19), allows equation (12) to be expressed only in terms of  $C_A$ , as derived below. From equation (18),

$$m_{HCO_3}^{-} = (2 K_{eq} P_{CO_2})^{1/3}$$
 (20)

However, the concentration in mol/cm<sup>3</sup> is essentially equal to  $m_{\rm HOO_3}$ -/1000. Thus,

$$c_{\rm A}^{\dagger} = 10^{-3} (2 \ {\rm K_{eq}} \ {\rm P}_{\rm OO_2})^{1/3}$$
 (21)

To express all terms as a function of  $C_A$ , rather than  $P_{O_2}$ , the gas law is employed as in equation (2):

<sup>\*</sup> This model is a simplified version of the calculation of carbonate equilibria at a given  $P_{OO}$  (see, eg, Garrels and Christ, 1965, p 81-83).

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$$C_{A} = \frac{n}{V} = \frac{P \omega_2}{RT}$$
,

or

$$P_{0} = RT C_A$$

With P in atm,  $C_A$  in mol/cm<sup>3</sup>, and T in kelvins, R = 82.05 atm-cm<sup>3</sup>-mol<sup>-1</sup>-K<sup>-1</sup>. Therefore

$$C_{\rm A}^{\dagger} = 10^{-3} (2K_{\rm eq})^{1/3} (82.05TC_{\rm A})^{1/3},$$
 (22)

•

$$C_{A}^{\dagger} = 5.475 \times 10^{-3} (K_{eq}T)^{1/3} C_{A}^{1/3}$$
, (23)

and thus

$$\frac{\partial c_{A}^{\dagger}}{\partial c_{A}} = 1.825 \times 10^{-3} (K_{eq}T)^{1/3} c_{A}^{-2/3} .$$
 (24)

Combining equations (24) and (19), expressing the fact that only 1/2 of the change in  $C_A^{\dagger}$  is due to  $\infty_2$  from the gas phase, equation (12) becomes

$$\tau \Theta_{\rm D} D_{\rm AB} \frac{\partial^2 C_{\rm A}}{\partial x^2} =$$

$$= \{\Theta_{\rm D} + 9.125 \times 10^{-4} (\Theta_{\rm T} - \Theta_{\rm D}) (K_{\rm eq} T)^{1/3} C_{\rm A}^{-2/3} \} \frac{\partial C_{\rm A}}{\partial t} + \alpha_{\rm T} . \qquad (25)$$

The same equation can be expressed in terms of  $P_{CO_2}$  by substituting  $P_{CO_2}/RT$  for  $C_A$  (from the gas law) in equation (25) and multiplying by RT to give

$$\tau \Theta_{\rm D} D_{\rm AB} \frac{\partial^2 P_{\rm CO}_2}{\partial x^2} = (26)$$

$$= \{\Theta_{\rm D} + 9.125 \times 10^{-4} (\Theta_{\rm T} - \Theta_{\rm D}) (K_{\rm eq} T)^{1/3} [P_{\rm CO}_2/RT]^{-2/3} \} \frac{\partial^P \Theta_2}{\partial t} + RT \alpha_{\rm T},$$
or
$$\tau \Theta_{\rm D} D_{\rm AB} \frac{\partial^2 P_{\rm CO}_2}{\partial x^2} = (27)$$

$$= \{\Theta_{\rm D} + 0.01723T(\Theta_{\rm T} - \Theta_{\rm D}) \kappa_{\rm eq}^{1/3} P_{\rm O2}^{-2/3}\} \frac{\partial^{\rm P} \Theta_{\rm 2}}{\partial t} + 82.05T\alpha_{\rm T}.$$

Either equation (25) or equation (27) may be used for modeling purposes, depending on convenience or preference.

THE PRODUCTION TERM. The high concentration of  $\Omega_2$  in the unsaturated zone relative to that in the atmosphere results from root respiration, microbial activity, and oxidation of organic carbon in the soil zone. All of these activities usually occur mainly near the surface, and in our model were <u>assumed</u> to occur uniformly with depth from land surface to a depth of lm, but not to occur below that depth. Attempts to include additional  $\Omega_2$  production at greater depths (see Site 4, fig 6) have not yet been made.

$$\frac{1}{\alpha} = \frac{2\Theta_{D}\tau D_{AB}(C_{depth} - C_{atm})}{a^2} , \qquad (28)$$

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where  $\overline{\alpha}$  = the average production rate of  $\infty_2$  in the root zone (mol/cm<sup>3</sup> of medium/sec);

$$C_{depth} = \omega_2$$
 concentration at depth (mol/cm<sup>3</sup>);

 $C_{atm} = \omega_2$  concentration in the atmosphere (mol/cm<sup>3</sup>);

a = depth of root zone (cm);

and other symbols are defined above.

To model seasonal variations in  $P_{CO_2}$  with depth, it is

assumed that the  $\[mathcal{O}\]_2$  production rate in the root zone is proportional to that given by a sine curve during the growing season, and that the rate is zero during the rest rest of the year. The amplitude of the production curve is calculated to provide the same total production for the year as that determined from the average production rate. Based on integration of the area under the sine curve, this amplitude is

$$\Omega = \frac{\pi}{2} \left( T_{tot} / T_{P} \right) \overline{\alpha} , \qquad (29)$$

where

 $\Omega$  = the maximum rate of  $\infty_2$  production, (mol/cm<sup>3</sup> of medium/sec);

 $T_{tot}$  = total time (one year);  $T_P$  = time of  $OO_2$  production (in fraction of a year);

and other symbols are as defined above. Thus, the seasonal production rate may be expressed in equation form

$$\alpha_{\mathrm{T}} = \frac{\pi \, \mathrm{T}_{\mathrm{tot}}}{2 \, \mathrm{T}_{\mathrm{P}}} \, \frac{\pi \, (\mathrm{T} - \mathrm{T}_{\mathrm{o}})}{\mathrm{asin}[\frac{\pi \, (\mathrm{T} - \mathrm{T}_{\mathrm{o}})}{\mathrm{T}_{\mathrm{P}}}], \, \mathrm{T}_{\mathrm{o}} \leq \mathrm{T} \leq (\mathrm{T}_{\mathrm{o}} + \mathrm{T}_{\mathrm{P}}); \quad (30)$$

and

$$\alpha_{T} = 0, \quad T < T_{o} \quad \text{or} \quad T > (T_{o} + T_{P}); \quad (31)$$

where

 $\alpha_T$  = production rate at time T (mol/cm<sup>3</sup> of medium/sec);

T = time (years);

T<sub>o</sub> = time at which seasonal production starts (years).

## DATA PRESENTATION

Table 1 presents the various derived parameters  $P_{OO_2}$ ,  $X_{OO_2}$ , and  $P_{14}_{OO_2}$  for all samples collected. The more standard  $\delta^{13}$ C and  $A^{14}$ C data are provided for comparison.  $P_{13}_{OO_2}$ was not tabulated because for all practical purposes  $P_{13}_{OO_2}$   $\cong 10^{-2}P_{OO_2}$ . Table 2 includes additional compositional data for the principal atmospheric gases. Seasonal variations of  $P_{OO_2}$  in individual probes are shown in figure 1, seasonal variation of  $P_{14}_{OO_2}$  in figure 4, and depth profiles of  $P_{OO_2}$ and  $P_{14}_{OO_2}$  in figure 6.

#### MODEL APPLICATION

Only two sites provide sufficient data to attempt seasonal modeling (North Dakota Sites 4 and 6); only Site 6 contains calcite throughout the unsaturated zone. The modeling efforts presented here are thus limited to North Dakota Site 6. The location and hydrogeologic characteristics of this site are described in Haas et al (1983). Models are presented first for the time and depth distribution of  $\Omega_2$ , then for the time and depth distribution of  $^{14}\Omega_2$ .

 $CO_2$  concentrations in the unsaturated zone were modeled for this study using a numerical solution to the finitedifference form of equation (27), as described by Weeks, Earp, and Thompson (1982). Briefly, for modeling purposes, the entire unsaturated zone is divided into equal nodal increments, and a finite-difference equation is written for each node. Variations in media properties are accounted for

$\begin{array}{ccc} v_{o1} & r & \alpha_2^{a/} \\ \omega_2 & & a_{1m} \times 10^2 \end{array}$			0.033 0.030						1.63		2.00		1.28 1.16	6/17		1.42	1.32		1.54	1.56 1.42	1.1.1		60.1		1.21		1.14 1.04					
δ <sup>13</sup> ω <sub>2</sub> <sup>b/</sup> °/₀₀		- 0	1.0- 1.0-	-23.5	-18.3	-24.6	-24.0		. ,	-24.2	-22.6	-22.6	1	c. 22-	ı		-21.8	-20.1	-20.8	-21.7	6.02-			-18.5	1	-18.2	-17.8	-17.9				
A <sup>14</sup> ω <sub>2</sub> pmc		00 361	129.05	118.08	103.62	79.07 122.67	124.83		, ,	114.13	113.26	112.79	112.52	112.33			90.90	86.24	86.07	85.47	80 <b>.</b> 84		1.1	51.16	53.07	51.28 52.45	52.30	52.96				
$x_{14} \frac{c}{\omega_2}$ x 10 <sup>14</sup>		0,0 0	0.050	,	,		I		, ,	1.44	2.65	1.95	1.68	2.30	,		1.40	1.76	1.75	1.56	08°T		1.1	.62	.75	0.71	0.70	.74				
<sup>P</sup> 14 <sub>002</sub> 4/ atm x 1014		370 0	0.046	,	,	. ,	,		, ,	1.31	2.41	1.77	1.53	2.14			1.28	1.60	1.41	1.42	1 • 0 4		, ,	0.56	0.68	0.66	0.64	0.67				
c <sub>14</sub> m <sub>2</sub> e/ x 10 <sup>20</sup>			1.96	,	ı		,			56.7	104.2	76.8	66.3	5.26			55.2	0.69	1.60	61.3	/.0/		, ,	24.5	29.5	28.3	27.4	29.2				
Date of collection	Site 6 (continued)	Water table 2	50uth 4 well (15.9m) 6/28/81 10/04/81	North 4" well (	8/17/80	12/04/80 3/05/81	6/25/81 10/06/81	10 100 101	Site 4, ND	Surface 8/	3/05/81	6/26/81 10/05/81	Totolot	Probe 1 (2.7m) 6/29/77(6b)	5/23/79	0/ 10/ 00/ 03/	3/05/81	6/26/81(6a)	T9/cn/nT	Probe 2 (5.8m)	8/23/78(8b)	5/23/79	8/16/80(8a) 12/05/80	3/05/81	6/26/81(6a)	10/05/81	Probe 3 (9.1m)	6/29/77(6b 8/23/78/8h)	5/23/79	8/16/80(8a)	3/05/81	6/26/81(6a)
vol 2 002	G	14.5m b/	(#6. CT	/q/=, 217	(mt.) (T						ı	, ,		0.73	0.44	0.56	0**0	0.71	60°T	:	4.90	5.63	5.34 6.69	4.63	4.66	5.53		17.8	15.7	18.8	17.0	18.3
$P_{\infty_2}^{a/a}$ atm x $10^2$			2.3		2.4	2.1	2.2	7 . 7			,	1.1	ı	0.66	0.40	0.51	0.36	0.65	66.0		1.20	5.12	4.86	4.21	4.24	5.03		16.2	14.3	17.1	15.5	16.7
δ <sup>13</sup> ω2 <sup>b/</sup> °/oo			-6.8 -7.3		-5.3	-6.7	- 4 - 9 - 9	t. 0			-19.7	-23.9	0.77-	ı	-27.9	-20.3	-21.5	-20.8	-21.3		1 1	-27.9		-23.3	-23.9	-22.8		ı	-29.4	-24.0	-24.0	
Α <sup>14</sup> ω <sub>2</sub> ) pmc			17.29 18.84		21.24	20.33	19.99	1 ** 07			115.17	123.64	//- 911		103.00	95.13 83 77	81.52	87.38	91.70		1.)	10.00	10.92	8.82	9.29	10.71		2.4	- 1.75	1.43	2.35	00.0
x <sub>14</sub> α <sub>2</sub> α/ × 1014			/q		, Ч		I	,			ı	,	1		0.53	1.02	0.38	0.73	1.17		, ,	0.66	0.68	0.69	0.51	0.69		.50	. 6	.31	-54	
${}^{P}_{14}\omega_{2}^{d/}$ atm x 1014			0.47 <sup>h/</sup>		/q_09-0	0.45	0.51	0.53			1	,	'		0.48	0.93	00	0.66	1.06		'	0.60	0.62	0.63	97.0	0.63		0.45	0 20	0.29	0.44	
<sup>C</sup> 14 002 <sup>E</sup> / x 10 <sup>20</sup>			/q								1	1	ı		20.8	40.3	0.12	28.5	76.0		ľ	25.9	26.8	27.0	0.01 9.91	27.2		9.61	7 61	12.4	19.1	1

TABLE 1. Carbon dioxide and carbon isotopes at various depths from sites in southwestern North Dakota, west Texas, and New Mexico

												55					
c <sub>14</sub> m <sub>2</sub> e/ x 10 <sup>20</sup>			12.8	5.3 9.51	7.9		\d					. 65.8		- 911	. 601	- 117 -	
$P_{14} \frac{d}{\omega_2} / \frac{1}{2}$			0.34	0.12 0.32	0.18		1	- 2.62	1		' '	-		, c	2.52	2.70	11311
x <sub>14</sub> 2/ 		1	0.32	0.13	0.20		<b>∕</b> ¶,		ı		' '	-	1 *	- 46	- 2.77	2.96	
А <sup>14</sup> 002 ртс		ι,	2.1 20.35	9.04 3.65	3.12 -			37.98	1		I I	-		- 02	12.2	-	11151
6 <sup>13</sup> ω2 <sup>b/</sup> °/00		I	-17.28 -16.6	-15.4 -17.4	-16.0			-11.7	-11.2		-21.64	- -21.63	1.1	-12 01	-11.68		1 1 1 1 1
P <sub>.</sub> α2 <sup>a/</sup> atm x 10 <sup>2</sup>	area and a second second as	17.2	12.0 1.45	1.16	4.09 1.21		6.9	6.3 5.9	4.4		4.95 5.92	5.28 7.22	10.3	10.2	22.9 17.7	22.0 17.5	1.66 0.95 0.08 0.12
vol 2 002					5.49 1.33	-0 <b>-</b>	- /	ı			5.42 6.51	5.81 7.93	11.3 11.6	11.2	25.2 19.4	24.2 19.2	1.89 1.08 0.09 0.14
Date of collection	Site 1 (continued)	Probe 5 (13.7m) 9/10/76	6/28/77 8/13/80	12/06/80 3/06/81	6/28/81 10/08/81	Water table ≅ 14	4" well (16.5m) 8/18/80	3/07/81	10/02/81	Site 2, ND Probe 1 (2.7m)	9/13/76 6/28/77	Probe 2 (5.8m) 9/13/76 6/28/77	Probe 3 (8.8m) 9/13/76 8/10/80	Site 3, ND Probe 1 (1.2m) 7/03/75	Probe 2 (3.4m) 9/13/76 7/03/77	<u>Probe 3</u> (6.1m) 9/13/76 7/03/77	Lamb         Stee         1           10/73778         #5 (5.8m)         #5 (5.8m)           #5 (5.8m)         #4 (11.0m)         #4 (11.0m)           #3 (17.1m)         #1 (20.5m)         #1 (20.5m)
c <sub>14</sub> 002 <sup>e</sup> / × 10 <sup>20</sup>				16.1 17.8	18.6 22.0	20.5		/q_	, ,			1.1	,	- 14.3 14.3	,	16.9	- 17.9 14.6 12.3 12.3 12.3 10.4 8.2
$P_{14} \frac{d}{\omega_2}^{d/}$ atm x 10 <sup>14</sup>			, .	0.37	0.43	- 0.48		/q1210	0.62	0.57		1.1	1.1	- .46 0.33 0.45	pumped water) -	0.39	- - 0.35 0.35 0.35 0.19
14 002 c/ × 10 <sup>14</sup>			1.1	14.	.56	- 53		<b>`</b> व_				1.1		0.50 0.36 0.49	is sample -	0.43	- 0.45 0.31 0.38 0.38 0.28
A <sup>14</sup> CD 2 Pmc			, ,	2.09	2.32	2.40		6 73	6.51	5.29 5.89 5.01		121.5 117.95	- (small isotope	- 3.21 2.99 5.46	- (89	6.03	2.1 1.85 1.81 1.81 1.85 1.57
δ <sup>13</sup> ω2 <sup>b/</sup> °/00					-24.0	-23.1 -23.9		1 21-	-16.7	-16.6 -16.6 -16.6		-23.4 -19.4	-34.8	- 1.8.1  - 17.9	, , ,	-17.5	- 17.68 - 17.0 - 18.9 - 17.5 - 17.5
$P_{002}^{a/3}$ atm x 10 <sup>2</sup>			15.1	15.3	15.8	16.7 17.0		0	8.1	8.3 8.3 10.0			0.073	10.3 12.2 9.46 7.01	0.51 6.38 10.6	5.55	14.9 16.8 13.5 13.5 11.1 11.1
Vol 2 002							E.S.	1					0.056	11.3 13.4 7.71	0.56 7.01		16.4 18.5 17.2 14.8 15.9 12.3
Date of collection		<pre>Site 4 (continued) Probe 4 (12.8m)</pre>	6/29/77(6b) 8/23/78(8h)	5/23/79 8/16/80(8=)	3/05/80	6/26/81(6a) 10/05/81	Water table = 17	(19.4m)	0/10/00 12/05/80	3/05/81 6/26/81 10/05/81	Site # 1, ND	Surface 6/28/81 10/08/81	Probe 1 (3.0m) 9/10/76 6/28/77	8/13/80 3/06/81 6/28/81 10/08/81	Probe 2 (5.2m) 9/10/76 6/28/81 6/28/81	10/08/81 Probe 3 (7.6m) <sup>1</sup> /	Probe 4 (10.9m) 9/10/76 6/28/77 8/13/80 12/06/80 3/06/81 6/28/81 6/28/81 10/08/81
	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$ \begin{array}{cccccc} v_{01} & v_{02} & v_{10} &$	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$														

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# Hydrology

. Footnotes to Table l	0.88, and 0.77atm at North Dakota, Texas, and New Mexico emites, respectively.	D. FUB standard c. Galculated from eq (9)	d. Calculated from eq (10); P <sub>total</sub> as in footnote a	e. Calculated from eq (11); " ";	<sup>T</sup> North Dakota = 283K, Trexas = 289K, TNew Mexico = 283K	f. The inlet for the air sample was $2m$ above land surface.	g. Volume X $\infty_2$ (and thus, $P_{\varpi_2}$ ) were not measured in the soil zone. The surface	samples represent $\infty_2$ that diffused out of the soll zone.	h. These analyses represent dissolved gases in shallow groundwater. The values	. reported for ${}^P\varpi_2$ represent the partial pressure of dissolved $\varpi_2$ , ${}^P\varpi_{2(aa)}$ .	It can be shown that $P_{14} = (R_{14} + C_{02}) + (R_{14} + C_{03})$ . Because there is no	gas phase present, $X_{14}^{4}$ , $X_{14}^{4}$ , are not relevant for these samples.	1. No samples were obtained; periodic attempts produced a vacuum whenever	this probe was pumped.	j. For additional data on ${ m CO}_2$ contents of the Texas sites, see Petraltis (1981).	k. These data (Kunkler, 1969) are from the unsaturated zone in the Bandeller	Tuff near Los Alamos, New Mexico.	1. These data are from the unsaturated zone in a forested area underlain by	calcareous sand at Trout Creek, Ontario, Canada. The numbers cited are	estimated average values. For further details, see Reardon et al (1979;	1980).
2 e/ 20			40							2 20 7		5			<b>~</b>		4				
<sup>c</sup> 14∞2 <sup>€/</sup> × 10 <sup>20</sup>		143.	84.4 99.0	106.					;	i g :	31.3	23.		5	18.3	ŝ	4. 45		22.		
P <sub>14</sub> 00_4/ atm x 10 <sup>14</sup>		3.39	2.00	2.48					2	44.0	0.74	0.57			64.0		76.0		0.51		
x <sub>14</sub>		3.85	2.27 2.66	2.82						0.50	0.84	0.64		2	9C.U		1.20		0.51		
A <sup>14</sup> ω <sub>2</sub> X <sub>1</sub> pmc >		1.811	111.5	108.6				, ,		97.18 62.91	63.71 63.71	56.18			101		CTT		~ 110		
هه1 <sup>3</sup> ∞2 <sup>1</sup> / °/۰۰۰		-20.6	-22.9 -18.7	-21.2						-10.03	-23.1	-25.9			C-81-		6. /T-		_k/		
P ω <sub>2</sub> 8/ atm x 10 <sup>2</sup>		2.65	1.53	1.94			0.50	0.05		0.59	- <b>66</b> .0	0.86			95.0		69.0		4.0		
v₀1 <b>x</b> ∞2	( pa	2.91	1.74	2.22			1.01	0.06		0.68	1.13	86.0			14.0	0	68.0	<sub>آ آ</sub> (ر	4.0		
Date of collection	Lamb Site (continued)	4/11/79 #5 (5.8m)	#4 (11.0m) #3 (17.1m)	#2 (21.4m)	Glenn Site	5/79	#5 (13.4E) #3 (25.3E)	#2 (36.3m) #1 (44.5m)	10/79	#5 (/.00) #5 (13.40)	#4 (19.8m) #3 (25.3m) #2 /2/ 2->	(⊞C.OC) 27 (⊞C.04) 1≇	Kunkler (1969) <u>k</u> /	8/24/67	IA-22-22 83.0-88.8m	8/5/67	TA->2-23 23.8-25.3≖	Reardon et al (1980) $\frac{1}{2}$	3-7m		

(Table 1 cont'd)

02		1.11			96.	ę.		17	10.				10.1	4.51		6 2	11.6			.43	.15																			
Ar 	ne X	0.1			1.0	1.1		-	1.1				1.0	1.0		1.1	1.1			1.2	1.2																			
N2	2m) Volume X	75.5		(=7	72.6	1.61		73.0	0.67			7m)	82.9	7.61	(=0	86.7	79.5		8m)	87.0	87.7																			
Date of collection	Site 3, ND Probe 1 (1.2m)	9/13/76		Probe 2 (3.4m)	9/13/76	1/03/11		Probe 3 (0.18)	7/03/77		Site 2, ND	Probe 1 (2.7m)	9//EI/6	0/ 28/ 11	B-obo 2 (5 8=)	9/13/76	6/28/77		Probe 3 (8.8m)	7/04/76	9/13/76																			
02 		1.8 20.8	10.6	5.0	15.0	1		16.7	1.4	12.5	16.0		ò	٥ <u>،</u>		2 C S	3.7	6.9	9.7		č	1)	7.4	0.02	0.41	14.2	20			leres .	100.>	100 2	1001	200.						
Ar	Volume X	1.3	1.0	6.	1.0	1.0			1.9	1.0	1.0			+ · ·		9	1.0	1.1	1.1						6.	1-1	1.0			, atmospheres	600.	200								
N2		96.8 78.1	77.1	80.7	73.5	/9		81 7	89.7	74.9	77			1.20	1.00	0.07	79.5	1.11	78				4.18	0.01	76.8	79.3	78			) Pressure	1.22	6	00							
Date of collection	Site 1, ND Probe 1 (3.0m)	7/10/76 6/28/77	8/13/80	3/06/81	6/28/81	10/08/81		CLODE 2 (2.4m)	6/28/77	6/28/81	10/08/81		Probe 4 (10.9m)	0//nT/6	11/07/0	12/06/80	3/06/81	6/28/81	10/08/81		Probe 5 (13.7m)	9//nT/6	0/727/0	00/CT/0	3/06/81	6/28/81	10/08/81			4" well (16.5m)	8/18/80	6/28/81	10/03/81	TO/CO/OT						
02		19.3 19.5	21.3	21.3	20.0	19-8	50		18.4	15.0	14.0	20.1	16.8	• •	16.7	2		.15	·06	1.9	60 <sup>-</sup>	• •	۰, م	90. V	5		.11	•00	r,	90.	U				eres	100	.002	.003	< .001	.002
Ar 	Volume X	1.0	1.0	1.0	1.0	1.0	1.1		1.0	1.1	1.0	1.0	o, (	ŗ.		1.1		1.1	1.1	1.1	1.1	0.T			A.1		1.1	1.1	1.1	1.0	1.0			1.1	re. atmospheres	010			.011	
N2		79.0	76.8	77.2	78.5	78.4	78		79.2	79.0	79.4	73.5	75.6	8-9/	6 F					81.4	80.1	80.8	81./	80.7	00	~	82.2	80.9	81.8	80.3	81.2		0.00	00			.94	.93	.98	96.
Date of collection	Site 4, ND Probe 1 (2.7m)	6/29/77	8/16/80	12/05/80	3/05/81	6/26/81	10/05/81	1-0 3/ 6 -10	6/29/77	8/23/78(8b)	5/23/79	8/14/80	12/05/80	3/05/81	19/72/9	10/00/01	Prohe 3 (9.1m)	6/29/17	8/23/78(8b)	5/23/79	-8/16/80(8a)	12/05/80	3/05/81	19/72/01	10/00/01	Probe 4 (12.8m	6/29/77 8	8/23/78(8b)	5/23/79	8/16/80(8a)	12/05/80	10/00/0	10/07/0	19/cn/n1	(m9.4m) (19.4m)	00/11/0	12/05/80	5/05/81	6/26/81	10/02/81
02 		15.2	16.6	20.2	20.3	20.0	19.5	19		18.1	18.1	17.4	19.9	19.7	19.0	0.61	17		18.8	17.6	18.2	19.6	19.6	18./	10.11	2	nospheres	< 001	100.			100. >	700.	700.	020	070.				
Ar 	Volume X	1.1	1.1	1.0	6.	¢.	6.	1.0		1.0	1.0	1.0	1.0	1.0	e	0.0	1.0		1.0	1.0	1.0	1.0	6.	1.0	0.1		ssure, att	.010	.011			110.	010.	210	710	110.				
N2	Δ	82.1	81.3	76.8	77.3	78.1	78.3	78		2.67	78.9	80.2	77.4	77.6	78.6	4.8/	6/		79.2	80.4	1.61	78.2	78.2	79.2	8.8/		5.9m) Pre	.92	.97		(=+. []	11-11	4. 1	66. ·	01-1	10.1				
Date of collection	Site 6, ND Beeks 1 (3 0m)		6/73/76(0D) 5/22/79					10/06/81	Broke 2 (5 8m)			5/22/79		_			T0/00/01	Prohe 3 (8.5m)	7/06/77			8a)	_		6/25/81		South 4" well (1	6/28/81 .92 .010 < .001	10/04/81		North 4" well (17.4m)	8/T//80	12/04/80	3/02/81	18/57/0	TO/00/01				

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and in west Texas

southwestern North Dakota

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contents of

TABLE 2. Nitrogen, argon, and oxygen

Hydrology

02	20.6	19.5	19.5	22.1	21.5	20.8	19.7	20.0	19.4
Ar	1.0	1.1	1.0	1.0	1.0	1.0	1.0	1.0	1.1
N2	77.8	78.4	78.6	76.8	17.7	78.1	78.9	1.91	78.7
Date of collection	$\frac{\text{Glenn site, TX}}{\frac{4/11/79}{\text{Probe}}5}$	(25.3m) Probe 2	(36.3m) Prohe 1	<u>10/79</u>	$\frac{Probe \ 6}{(7.0m)}$	(13.4m)	(19.8m)	$\frac{(25.3\pi)}{(25.3\pi)}$	( <u>36.3m</u> )
02	1.11	20.7	20.8	17.8	17.5	16.5	15.6	15.5	
Ar	1.2	1.0	1.1	1.1	1.0	1.0	1.1	1.1	
N2	86.6	77.6	77.3	6.08	78.8	80.5	80.5	81.1	
Date of collection	Lamb site, TX 10/23/78 Probe 4 (11.0m)	Probe 3 (17.1m) Probe 2	(21.4m)	$\frac{(26.5\pi)}{4/79}$	Probe 5 (5.8m)	(11.0m)	(17.1m)	(21.4m)	

(Table 2 cont'd)

by dividing the unsaturated zone into layers, each containing several nodes and within which the properties of the medium, including its tortuosity, drained porosity, and total porosity, are assumed constant. Each screened interval is considered to comprise a layer.

GAS AND MEDIA PROPERTIES. Application of the model requires the input of various parameters to describe the diffusive properties of the gas and medium; the interaction of  $CO_2$ among the gas, liquid, and solid phases within the medium; the production term; boundary conditions at the top and bottom of the unsaturated zone column; and initial conditions at the start of the transient simulation. The values of the parameters used to model North Dakota Site 6 are described below.

Based on observed groundwater temperatures, the mean annual temperature at Gascoyne, North Dakota, is approximated as 10°C. The total pressure is 0.91 atm. The diffusion coefficient of  $CO_2$  in air is calculated internally in the computer program (equation (13)) with these parameters. The equilibrium constant for equation (15) has the value log  $K_{eq}(10°C) = -5.65^*$ . At 283 K, equation (27) thus reduces to

$$\tau \Theta_{\mathbf{D}} \mathbf{D}_{\mathbf{A}\mathbf{B}} \frac{\partial^2 \mathbf{P}_{\mathbf{CO}_2}}{\partial \mathbf{x}^2} =$$

$$= \{\Theta_{\rm D} + 0.0638(\Theta_{\rm T} - \Theta_{\rm D}) P_{\rm CO_2}^{-2/3}\} \frac{\partial^{\rm P} \Theta_2}{\partial t} + 2.322 \times 10^4 \alpha_{\rm T} . \quad (32)$$

The dependence of the rate of mass transfer of  $CO_2$  between the gas and liquid phases upon the gas-phase partial pressure of  $OO_2$  makes equation (32) nonlinear. However, because the value of  $P_{CO_2}^{-2/3}$  varies relatively slowly with  $P_{CO_2}$ , no particular problems arise in the numerical solution of the equation. For this study,  $P_{CO_2}^{-2/3}$  for a given time step was evaluated from the concentration at the previous time step. Small enough time steps (3 days) were used to

\* Calculated from the data of Plummer and Busenberg (1982)

$$Gaseous \stackrel{12}{\sim}CO_2, \stackrel{13}{\sim}CO_2, \text{ and } \stackrel{14}{\sim}CO_2 \text{ in Sub-Soil} \qquad 333$$

ensure that the largest error was  $\leq 1\%$ , as determined by trial and error.

The materials comprising the unsaturated zone at North Dakota Site 6 include sandy clay from a depth of 0 to 10m, and fine-grained sand below that depth. Based on these textural descriptions, the total porosity was assumed to be 0.35 for the entire profile, as this is a common value for unconsolidated sediments. Drained porosity was assumed to be 0.15 above 10m, and 0.20 below that depth. Tortuosities for the two layers were computed from the relationship (Lai, Tiedje, and Erickson, 1976)

$$\tau = \Theta_{\rm D}^{4/3}$$
, (33)

resulting in a value of .08 for the upper layer and of .12 for the lower layer.

THE  $O_2$  MODEL. Boundary and initial conditions: The concentration of  $O_2$  at land surface (the upper boundary) is assumed to be specified as a function of time. For the seasonal modeling,  $P_{CQ_2}$  was assumed constant at .0003 atm,

and the water table, which is at a depth of ca 15m at Site 6, was assumed to be a no-diffusion boundary. Assumed initial conditions are that the partial pressure of  $\infty_2$  is 0.014 atm throughout the profile.

Assuming that the average  $P_{O2}$  at depth is 0.014 atm as measured at 6m at Site 6, that  $P_{O3}$  in the atmosphere is

.0003 atm, and that the root zone depth is lm, the average rate of production in the root zone at Site 6 is 2.4 x  $10^{-13}$  mol/cm<sup>3</sup> of medium/sec, a value that is calculated internally in the computer program from user-supplied data. The variation of P<sub>CQ2</sub> with depth is based on a seasonal production

rate (equations (29) to (31)) that assumes a five-month growing season that begins on May 1 each year.

RESULTS. The most important influence on  $\infty_2$  chemistry in the unsaturated zone at Site 6 appears to be chemical reactions in the soil zone with downward propagation of  $\infty_2$  by vapor phase diffusion. The well defined annual cycles of  $P_{\Omega_2}$  at the 3m probe, the attenuation of the cycles with

depth, and the shifts of the maxima and minima to later dates with depth are all consistent with downward diffusion

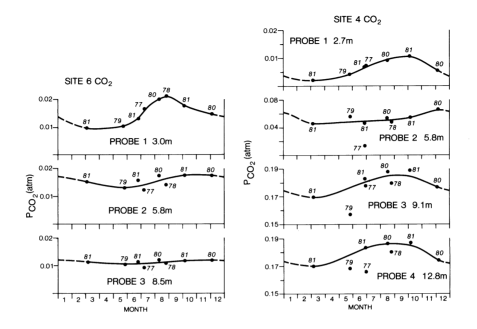


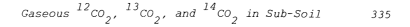
Fig 1. Seasonal variation of PCO at North Dakota Sites 6 and 4. Data are plotted <sup>2</sup>by collection date, independent of year, with the year of collection noted at each data point. Data from table 1.

from a seasonally varying source of  $\Omega_2$  in the soil zone (fig 1). Some details of the modeling results are described below.

Once specific data for Site 6 were incorporated into the mathematical model, the seasonal distribution of  $P_{CQ_h}$  with

depth was computed, based on the assumption that the  $\Omega_2$  production history exactly repeats itself every year, an assumption that appears well-justified based on the seasonal cycles in the shallow probes at both Site 6 and Site 4 (fig 1). Five years were simulated to ensure steady cyclic conditions for the final year of simulation. Results, in terms of  $P_{\Omega_2}$  vs depth during each month of the growing sea-

son, are shown in figure 2. These curves are compared (fig 2) with the detailed depth-time  $CO_2$  data from Trout Creek, Ontario (Reardon, Allison, and Fritz, 1979). Qualitative



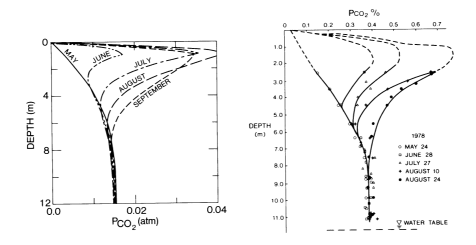


Fig 2. Seasonal profiles of  $P_{CO}$  vs depth in the unsaturated

zone. A. Model calculated for North Dakota Site 6 B. Data from Trout Creek, Ontario; from Reardon, Allison, and Fritz (1979).

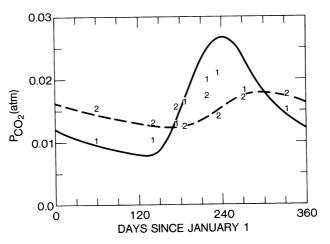


Fig 3. Comparison of calculated seasonal variation of  $P_{OO}$  at North Dakota Site 6 with observed data. Solid 2 line: probe 1 model; dashed line: probe 2 model. Numbers are measured  $P_{OO}$  's for probes 1 and 2, respectively. Probe 1 is 2 screened at 3.0m, probe 2 at 5.8m.

agreement is quite good, indicating that diffusion theory adequately explains the Trout Creek data, even without incorporating site-specific parameters for their site.

The time variation of  $P_{O_2}$  at the depths of probes 1 and

2, Site 6, from this simulation were also calculated and are compared with the measured values for Site 6 in figure 3. The comparison shows slightly less predicted seasonal variation for probe 1 than was observed; for probe 2, the predicted seasonal variation is somewhat greater than observed. Trial-and-error simulations (not shown) made by varying the tortuosity indicate that the amplitude and timing of the measured  $P_{CO_2}$  seasonal pattern for probe 1 can be very well

simulated using a tortuosity of 0.07 ( $\tau = 0.08$  is used in fig 3). Alternatively, an excellent match between measured and simulated results can be obtained for probe 2 using a tortuosity of 0.09. These tortuosities are both well within the plausible range for Site 6. However, an implausibly large value for tortuosity is required to force a simultaneous good fit to the data for both probes.

The inability to obtain a satisfactory simultaneous fit to the measured values for probes 1 and 2 at Site 6 suggests that the actual diffusion process may not be truly onedimensional. A plausible explanation is that  $CO_2$  moves by a somewhat less tortuous path from land surface to probe 2 than from land surface to probe 1, possibly due to a very local clay lens in the top three meters. However, the goodness of fit is adequate to strongly suggest that mainly vertical diffusion and calcite equilibrium are indeed the dominant mass-transport and mass-transfer mechanisms affecting the  $CO_2$ distribution in the unsaturated zone at Site 6.

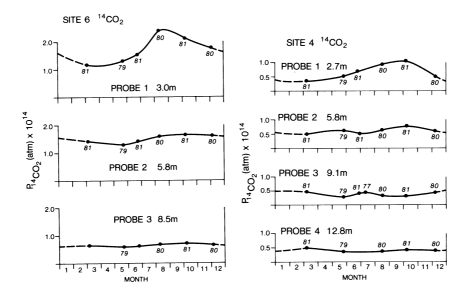
THE  $^{14}$  CO $_2$  MODEL. Boundary and initial conditions: The welldefined seasonal variation of  $P_{14}$  in the shallow probes  $CO_2$ 

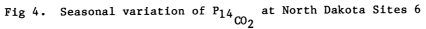
at Sites 6 and 4 (fig 4) suggests that diffusion is also important in the transport of  $^{14}\Omega_2$ . However, unlike  $\Omega_2$ , atmospheric P<sub>14</sub> has shown dramatic changes during the last  $CO_2$ 

30 years as a result of atmospheric nuclear testing in the 1950's and 1960's. Because each species diffuses according to its own gradient, this post-bomb  $^{14}$ C should have migrated at least tens of meters into the unsaturated zone by now. To test this idea, an attempt was made to simulate  $P_{14}$  at  $CO_2$ 

Site 6, with the same media parameters used to simulate the seasonal effects described above.

$$Gaseous \begin{array}{c} {}^{12}CO_2, \\ {}^{13}CO_2, \\ {}^{and} \end{array} \begin{array}{c} {}^{14}CO_2 \\ {}^{in} Sub-Soil \\ {}^{337} \end{array}$$





and 4. Data are plotted by collection date, independent of year, with the year of collection noted at each data point. Data from table 1.

To model  ${}^{14}$   ${}^{\circ}{}_{2}$ , information is needed regarding the production of  ${}^{14}$   ${}^{\circ}{}_{2}$  in the root zone.  ${}^{14}$   ${}^{\circ}{}_{2}$  production was calculated from equation (34):

$$\overline{\alpha}_{14}_{CO_2}(t) = [\overline{\alpha}_{14}_{CO_2}(1981)] \times \frac{A^{14}C(t)}{A^{14}C(1981)}$$
(34)

where t = time;

$$\overline{\alpha}_{14}$$
 (t) = average production rate of  ${}^{14}$   $\Omega_2$  in the root  
zone at time t, mol/cm<sup>3</sup> of medium/sec;

and  $A^{14}C(t)$  = activity (pmc) of  ${}^{14}C$  in the atmosphere at time t (fig 5A).

The atmospheric  $^{14}$ C activity through 1975 was estimated from data presented in figure 4 of Broecker, Peng, and Engh (1980),

and was then extrapolated to our measured values in 1981.  $\overline{\alpha}_{14}$  (1981) was computed to produce an average  $P_{14}$  of 1.6  $\varpi_2$ x 10<sup>-14</sup> atm, the average Site 6 value measured at 3m. A 30year period, starting in 1951, was simulated. Seasonal effects were ignored and 30-day time steps were used in the simulation. The proportionality postulated between atmospheric <sup>14</sup>C activity and soil-zone <sup>14</sup>C activity represents a major assumption that cannot be rigorously documented at present; the assumption is supported by the near-atmospheric <sup>14</sup>C activities in most of the surface  $\varpi_2$  collections.

The mass transfer of  $^{14}$ CO  $_2$  between the gas and liquid phases was also handled differently in this simulation. The  $^{14}$ C mass transfer was assumed to occur at a rate proportional to that of CO  $_2^*$ . The soil gas was assumed to have an average partial pressure throughout the column of .014 atm. Using this value, the partitioning term in equation (32) reduces to  $\{\Theta_D + 1.10(\Theta_T - \Theta_D)\}$ , thus linearizing equation (32) and making large time steps possible.

The lower boundary condition was also modified for this simulation. Diffusion in the liquid phase could be significant during the long time span covered; hence, another layer was added to represent the saturated material from 15m to 25m. As a mathematical artifice, these materials were assumed to have total and drained porosities of 0.35, and the tortuosity was assumed to be .0001. This approach allows gas diffusion through the fully-saturated medium to be approximated without reprogramming.

This, in turn, leads to aqueous dilution or enrichment factors that overshadow the changes in  $P_{14}$ .

Hence, the dependence on  $P_{OO_2}$ , rather than  $P_{14}_{CO_2}$ .

<sup>\*</sup> This is not an obvious assumption. It is derived from a chemical model developed by the authors, the presentation of which is beyond the space limitations of this paper. The fundamental assumption is that dissolution and precipitation of calcite is governed by changes in  $P_{CO_2}$ .

$$Gaseous \stackrel{12}{\sim}CO_2, \stackrel{13}{\sim}CO_2, \text{ and } \stackrel{14}{\sim}CO_2 \text{ in Sub-Soil} 339$$

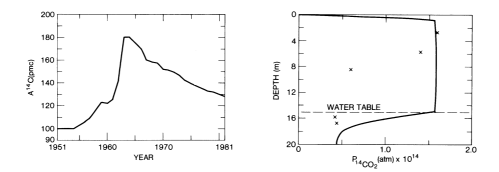


Fig 5. A. Atmospheric  ${}^{14}C$  activity, adapted from Broecker, Peng, and Engh (1980); see text for discussion. B. Calculated depth profile for P<sub>14</sub> at North Dakota  $CO_2$ 

Site 6; see text for discussion.

Initial conditions to be assumed in the simulation of  $^{14}\mathrm{CO}_2$  are problemmatical, as the  $^{14}\mathrm{CO}_2$  profiles themselves suggest that they cannot be adequately explained by diffusion theory alone. For this simulation, it was assumed that  $^{\mathrm{P}}_{14}\mathrm{_{CO}}_2$  in 1951 was equal to 0.48 x 10<sup>-14</sup> atm, which is the

same as that measured in the ground water in 1981.

RESULTS. The long-term  $^{14}$ C simulation is shown in figure 5B. The calculated profile is almost vertical, with a slight bulge about mid-depth in the unsaturated zone arising from the effect of currently declining production rates following the 1964 peak in  $^{14}$ C activity in the atmosphere. Note that the curve departs radically from the average measured partial pressures, symbolized by X's in the figure. The measured values show a sharp dropoff that cannot be explained by the gaseous diffusion-calcite equilibrium model. Moreover, according to theory, the dissolved  $^{14}$ CO<sub>2</sub> should have established a steep gradient to a depth of ca 3m into the water table during the 30-year period. The near equality of  $^{14}$ CO<sub>2</sub> partial pressures in the lower gas probes and groundwater samples at different depths (fig 6) shows that this has not been the case.

In spite of the clearly defined seasonal profile in the shallow probes (fig 4) application of diffusion theory to measured  $^{14}$   $\infty_2$  profiles in the deep unsaturated zone indicates that some as yet unexplained mechanisms severely retard the migration of  ${}^{14}\omega_2$  with depth in the deep unsaturated zone. The general decreases in  $\text{CO}_2$  and  $^{14}\text{CO}_2$  cannot be readily explained with a steady state diffusion model because of the relatively large fluxes that can be achieved with gaseous diffusion. The depth distribution of  $\omega_2$  suggests transient processes on a fairly short time scale - not exceeding a few tens of years. The  $^{14}$   $\Omega_2$  is more rapidly attenuated with depth and shows a greater overall decrease than does  $\omega_2$ , suggesting a selective sink for  $^{14}CO_2$  at Site 6. A time lag might also exist between changes in atmospheric <sup>14</sup>C activity and the <sup>14</sup>C activity in the root zone. However, if this were the only faulty assumption in the model, the decrease in P14 CO2 from 3m to 9m should not exceed ~30%, rather than

the observed decrease of a factor of 2.

A particularly interesting observation is that soil gasshallow groundwater  $^{14}$   $^{16}$ 

the selective attenuation of  ${}^{14}$ CO<sub>2</sub> with depth in the unsaturated zone at Site 6. No obvious processes to account for this phenomenon are apparent. However, whatever these mechanisms are, they can profoundly affect the  ${}^{14}$ C activity of recharged ground water, and suggest that conclusions concerning the residence time of ground water in the unsaturated zone based solely on  ${}^{14}$ C activities are extremely suspect.

An estimate of the relative magnitudes of diffusive and advective fluxes of  $\ensuremath{\mathbb{C}}_2$  is needed. The following example, based on North Dakota Sites 4 and 6, provides a numerical comparison that might approximate Great Plains environments. Groundwater alkalinities in the area average  $\sim 10 \text{ meq}/1$ ; recharge in Bowman County is estimated as  $\sim 1~{\rm cm/yr}$  (Croft, 1978). A recharge rate of lml H<sub>2</sub>O/cm<sup>2</sup>-yr (ignoring media effects) yields a  $\Omega_2$  flux to the water table of  $10 \mu mo1/cm^2$ yr. In the gas phase a gradient of 1% CO2 over 10m, including the moderating effects of porosity and reaction, provides a calculated diffusive flux of  $\sim 15 \mu mol/cm^2$ -yr. It thus appears that the diffusive fluxes due to the large, rapidly changing gradients associated with near-surface seasonal OO2 fluctuations should overshadow recharge effects, while if smaller gradients occur in the deeper unsaturated zone the fluxes might approach each other in magnitude. Rough estimates suggest that sub-soil diffusion should pre-

$$Gaseous \begin{array}{c} {}^{12}CO_2, \\ {}^{13}CO_2, \\ {}^{20}CO_2, \\ {}^{20}CO_2, \\ {}^{20}CO_2 \\ {}^{20}CO_2$$

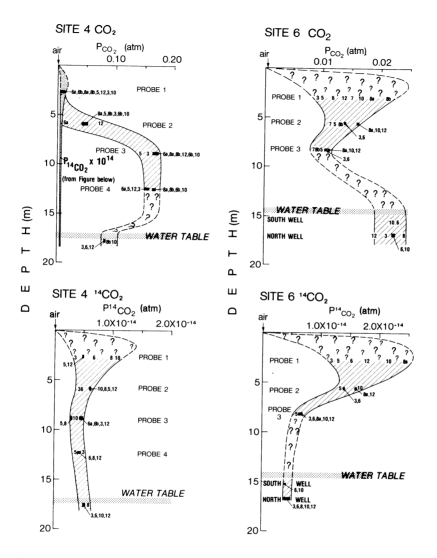


Fig 6.  $P_{CO_2}$  and  $P_{14}$  vs depth at North Dakota Sites 4 and 6. Numbers refer to month of collection date, independent of year, and correspond to entries in table 1.

dominate for  $\[mathbb{M}_2\]$  and  $\[mathbb{14}]{14}\[mathbb{M}_2\]$  at Sites 6, and  $\[mathbb{M}_2\]$  at Site 4; the relative magnitude of  $\[mathbb{14}]{14}\[mathbb{M}_2\]$  diffusion below ~ 5m at Site 4 is much less.

The data tables show that a wealth of information is available in the accumulated  $^{13}$ C data for the unsaturated zone gases at all sites, both in terms of inter-site comparisons and in the differences between gases and the water table wells. No attempt has been made in this paper to incorporate these  $^{13}$ C data into chemical models of unsaturated zone process. Also, no quantitative transport modeling for the other sites has yet been attempted. In the following discussion, important general aspects of the data at other sites are briefly pointed out.

#### DESCRIPTION OF OTHER SITES

NORTH DAKOTA SITE 4. (See Haas et al, 1983, for detailed description). This site provides an excellent example of the independent behavior of  $^{12}\text{CO}_2$ ,  $^{13}\text{CO}_2$ , and  $^{14}\text{CO}_2$ . The major source of  $^{12}\text{C}$  and  $^{13}\text{C}$  in the system occurs at a depth of 7.5 to 9.5m, where oxidation in the upper portion of the lignite reduces oxygen pressure to near zero and produces CO<sub>2</sub> contents approaching 20 volume%. This CO<sub>2</sub> then diffuses upward, and masks the seasonal cycle in P<sub>CO<sub>2</sub></sub> in all but the

shallowest probe. The observed gradient in PCQ, suggests

that the  $\Omega_2$  produced in this zone also diffuses downward towards the water table, which appears to be acting as a sink for  $\Omega_2$  at this site.

In spite of  $P_{OO_2}$  -depth variation greater than an order

of magnitude in the unsaturated zone, and three orders of magnitude difference relative to the atmosphere, the partialpressure depth profiles of  ${}^{14}$   $\Omega_2$  are the most uniform observed in this study. The  ${}^{14}$   $\Omega_2$  depth profile is a better approximation to the elementary diffusion model (compare fig 6B with fig 1A) than either  $\Omega_2$  or  ${}^{14}\Omega_2$  from any other North Dakota sampling site. The surprising observation is again made that partial pressure equilibrium between the gas phase and the water table is approached more closely for  ${}^{14}\Omega_2$  than for  $\Omega_2$ . Note that here the partial pressure of  $\Omega_2$  is lower in the water - opposite the situation present at North Dakota Site 6. The existence of the  $\Omega_2$  gradient from the base of the lignite to the water table-if steady state-implies a large flux of  ${}^{12}\Omega_2$  and  ${}^{13}\Omega_2$  to the ground water.

$$Gaseous \stackrel{12}{\sim} CO_2, \stackrel{13}{\sim} CO_2, \text{ and } \stackrel{14}{\sim} CO_2 \text{ in Sub-Soil} \qquad 343$$

NORTH DAKOTA SITES 2 AND 3. Both of these sites are located in lignite-rich spoils, and both show some of the general characteristics of North Dakota Site 4 - namely high and variable  $O_2$  contents due to lignite oxidation, and relatively uniform  $^{14}O_2$  profiles. Although an investigation of these sites was originally one of the motivations for this work, because of their hydrologic variability due to mine pumpage, we decided early to concentrate on natural processes in the undisturbed sites.

NORTH DAKOTA SITE 1. At Site 1 a stringer of lignite is present just above the water table and a perched water table (the gas probes pumped water) was present at depths of 3m to 8m until at least 1977. Prior to this, the deep samples at Site 1 resembled samples from similar depths at Site 4 in both  $\Omega_2$  and  ${}^{14}\Omega_2$  content. The perched water lens disappeared sometime between 1977 and 1980. From the latter date to the present the data at all depths show extreme variability; it was initially believed that the grouting had cracked or that some other sampling problem existed. However, more recent analysis of the data shows that the partial pressure of  ${}^{14}\Omega_2$  is generally (not always) nearly constant, suggesting a lack of atmospheric influence. No plausible model for processes at this site is available at present.

THE TEXAS SITES. The sampling nests at both Texas sites were constructed as part of an artificial recharge study in the Ogallala aquifer conducted by the US Geological Survey (Weeks, 1978). At the Glenn and Lamb sites, located near Lubbock, Texas, the Ogallala formation consists of interbedded sands, silts, and clays, with local development of massive caliche. The water tables at the sites are at depths of 51m (Glenn) and 77m (Lamb). For additional geologic and hydrologic information, see Weeks (1978) and Petraitis (1981); much additional CO<sub>2</sub> and  $\delta^{13}$ C data for these sites are presented in Petraitis (1981).

THE GLENN SITE. Two sets of samples were collected from this site (table 1). CO<sub>2</sub> partial pressures are 0.01 atm or less; highest CO<sub>2</sub> pressures occur at intermediate depths. In the single set of  $^{14}\text{CO}_2$  samples collected, the relative variation of  $\text{P}_{14}_{\text{CO}_2}$  is less than that of  $\text{P}_{\text{CO}_2}$ , but the greatest

value is found at an intermediate depth. The partial pressure of  $^{14}$   $\mathrm{CO}_2$  at this site is quite similar in magnitude to the values observed at North Dakota Site 4. No  $^{14}$   $\mathrm{CO}_2$  data are available at the water table.

THE LAMB SITE. Two sets of  $\Omega_2$  data are available, but in this case, they are quite dissimilar; only the shallow probes show fairly consistent values of  $P_{\Omega_2}$ . If the variation is

due to sampling error, it is most likely that the initial samples were collected without sufficient initial pumping.

This site is quite interesting in that it is the only site for which our data definitively show post-bomb  $^{14}$   $\Omega_2$  activities in deep samples, in spite of a perched water table at 28m. Combined with the relatively high  $\Omega_2$  partial pressures, the resulting  $^{14}$   $\Omega_2$  partial pressures at the Lamb site are equalled only by the lignite spoils at North Dakota Site 3. The  $^{14}$   $\Omega_2$  partial pressures are nearly constant in the deeper probes.

#### GENERAL OBSERVATIONS

A number of general observations can be made regarding the behavior of  $^{14}$   $^{14}$   $^{14}$   $^{14}$  in the unsaturated zone that are independent of sampling location or date.

1) The geochemistry of  $^{14}$ CO<sub>2</sub> in the unsaturated zone must be evaluated in terms of both the partial pressure of  $^{14}$ CO<sub>2</sub> and the  $^{14}$ C activity of the CO<sub>2</sub> gas samples. A value

of  $A^{14}C$  >100pmc still remains the only unequivocal signature of post-bomb carbon. However, the general lack of correlation between the  $A^{14}C$  and  $P_{14}CO_2$  depth profiles emphasizes the

need for evaluation of both parameters in unsaturated zone  $^{14}$ C studies.

2) Diffusion is a major mechanism of gas transport in the sub-soil unsaturated zone. The various isotopes of  $CO_2$ each diffuse in response to their own sources and sinks, as best illustrated by the data from North Dakota Site 4. The diffusion models must consider individually the absolute concentration of each isotopic species in units of mass/volume or the corresponding gas partial pressures. Diffusive fluxes of  $CO_2$  and its isotopes appear to predominate over advective fluxes in the shallow (< 5m) unsaturated zone. No generalizations regarding greater depths can be made.

3)  $^{14} \[mu]_2$  and  $^{12} \[mu]_2$  are biologically generated in the shallow soil zone. Penetration of the  $^{14} \[mu]_2$  into the unsaturated zone and observed activities >100pmc in surface samples imply relatively rapid diffusion of this species in the subsurface. Considering the relatively large fluxes that can be generated by diffusive transport of gases, the variable

$$Gaseous \stackrel{12}{\sim} CO_2, \stackrel{13}{\sim} CO_2, \text{ and } \stackrel{14}{\sim} CO_2 \text{ in Sub-Soil} \qquad 345$$

shapes of the  $P_{CO_2}$  - and  $P_{14}$  depth profiles and the partial  $CO_2$ 

pressure disequilibrium between the gas and aqueous phases suggest that steady-state processes are very unlikely at any of the sites studied.

4) The measured concentration, in mass/volume of gas, of  $^{14}\mathrm{OO}_2$  in the unsaturated zone is without exception 10 to 100 times greater than the concentration of  $^{14}\mathrm{OO}_2$  in the atmosphere.

## ACKNOWLEDGMENTS

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Finally, the existence of much of the data is due in large part to the continual good humor and competence--sometimes under extreme conditions--of Dave Stannard, whose help in the field is much appreciated.

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