A General Circulation Model Study of Atmospheric Carbon Monoxide

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The carbon monoxide cycle is studied by incorporating the known and hypothetical sources and sinks in a tracer model that uses the winds generated by a general circulation model. Photochemical production and loss terms, which depend on OH radical concentrations, are calculated in an interactive fashion. The computed global distribution and seasonal variations of CO are compared with observations to obtain constraints on the distribution and magnitude of the sources and sinks of CO, and on the tropospheric abundance of OH. The simplest model that accounts for available observations requires a low latitude plant source of about $1.3 \times 10^{13}$ g yr$^{-1}$, in addition to sources from incomplete combustion of fossil fuels and oxidation of methane. The globally averaged OH concentration calculated in the model is $7 \times 10^{5}$ cm$^{-3}$. Models that calculate globally averaged OH concentrations much lower than our nominal value are not consistent with the observed variability of CO. Such models are also inconsistent with measurements of CO isotopic abundances, which imply the existence of plant sources.

INTRODUCTION

Carbon monoxide is the third most abundant carbon species in the atmosphere (after carbon dioxide and methane), but its roles in the atmospheric and biogeochemical cycles of carbon are not quantitatively understood. An understanding of the CO cycle is important in its own right, for the information it implies with regard to the global dispersal of atmospheric pollutants, and for an assessment of perturbations of tropospheric chemistry by man's activities. Early work on carbon monoxide included its discovery in the infrared solar absorption spectrum by Mottle [1949] and an investigation of its potential sources and sinks by Bates and Witherspoon [1952]. The subject remained dormant for nearly 2 decades until the importance of the hydroxyl radical in tropospheric chemistry was recognized [Levy, 1971; McConnell et al., 1971]. The hydroxyl radical is derived from

$$O_3 + h\nu \rightarrow O_2 + O(1D) \quad (R1)$$

The reaction

$$O(1D) + H_2O \rightarrow 2OH \quad (R7)$$

provides the only important sink for CO in the gas phase and is responsible for determining the chemical lifetime of CO in the atmosphere. In addition to destroying CO, the hydroxyl radical also destroys CH$_4$

$$CH_4 + OH \rightarrow H_2O + CH_3 \quad (R16)$$

Reaction (R16) initiates a sequence of reactions that ultimately leads to the production of CO. As was first pointed out by McConnell et al. [1971], (R16) represents a major source of CO, especially in the clean troposphere. It was argued [see Wofsy, 1976] that this source, together with that derived from incomplete combustion of fossil fuels [Bates and Witherspoon, 1952], could account for the budget of atmospheric CO. However, later experiments [see, for example, Biermann et al., 1978] showed that reaction (R18), which is primarily responsible for removing CO, is pressure-dependent in the presence of oxygen and should proceed nearly twice as fast as previously thought. A renewed search for sources of CO suggested that oxidation of nonmethane hydrocarbons (NMHC), especially isoprenes and terpenes emitted by trees [Zimmerman et al., 1978] and biomass burning [Crutzen et al., 1979] could be important. However, it is difficult to quantify the magnitudes of these sources (see Logan et al. [1981] for a detailed catalog and assessment of the uncertainties).

The intense interest in CO in the last decade resulted in the generation of an extensive data base concerning its spatial distribution [see, for example, Seiler, 1974] and seasonal variation [Seiler et al., 1976; Dianov-Klokov et al., 1978]. Previous theoretical studies of the CO cycle include photochemical models that neglected horizontal transport [Wofsy, 1976; Crutzen and Fishman, 1977; Logan et al., 1981] and Hameed and Stewart's [1979] model, which considered north-south mixing by diffusion. However, the chemical lifetime of CO is believed to be at least several weeks, and, hence, both chemistry and transport should play a role in determining the CO concentration as a function of height, latitude, and season. A study of the CO cycle with a three-dimensional global model, which explicitly accounts for atmospheric motions, should therefore be useful for helping to identify and quantify the sources and sinks of CO in a self consistent manner. Greater weight should also be given to tropospheric OH calculations that are consistent with the major features of the global CO cycle. Because OH determines the CO lifetime, and the strength of the methane oxidation source, and reaction with CO is the major loss for OH, we must calculate OH concentrations in an interactive manner. The nature of this coupling between CO and OH is such that additional CO sources require higher OH concentrations and vice versa. We will use three independent pieces of information in the CO data base to constrain our predicted OH concentrations; these are (1) the latitudinal variation of CO, (2) its vertical variation, especially in the tropics, and (3) its seasonal variation. Since we can estimate the magnitude of anthropogenic emissions to better than a factor of 2, there is essentially one remaining free parameter in our model, the magnitude of plant sources in the tropics. A large number of other parameters have also been tested, but they were shown to be unimportant. Since we have used measurements of CO variability as a measure of its chemical lifetime, we regard our derived OH concentrations as empirical values that satisfy...
these observations. Their validity should therefore transcend the assumptions of our simplified chemical model.

In the following section we briefly describe the three-dimensional transport model used in our studies. In the subsequent section we describe several experiments used to investigate the CO cycle. In the final two sections we discuss the results and implications of the experiments.

**DESCRIPTION OF THE MODEL**

The numerical experiments were performed with the general circulation model described by Hansen et al. [1983]. The general circulation model is global in extent, with a horizontal resolution of 8° × 10° (latitude x longitude). Seven layers, evenly spaced in σ coordinates, were employed between the surface and 10 mbar (31 km), with at least one layer entirely within the stratosphere at all locations. The model was run for 5 years, and the necessary dynamical variables for the fifth year stored off line for use in a 'tracer model.'

Compared with climatological data, the three-dimensional wind fields, temperatures, and humidity, have a rough correspondence with reality. An indication of the representativeness of the model's wind fields is presented in Figure 1a for the surface winds in February and in Figure 1b for the jet stream winds at 315 mbar level. The surface wind pattern shows the expected latitudinal variation, with easterlies at low latitudes and cyclonic and anticyclonic circulations at high latitudes. The southeast and northeast trade winds converge near the equator, the pattern being more apparent over the ocean, where frictional effects are less than that over land. Anticyclonic patterns dominate over the oceans near 30° latitude, typical of the subtropical high pressure system. At about 50°N latitude, the Aleutian low in the mid-Pacific and the Icelandic low in the North Atlantic produce cyclonic wind patterns, while prevailing westerlies occur over the United States and Asia. In the southern hemisphere, strong west winds are evident near 60°S. All these features predicted by our model also exist in observations. At the jet stream level, strong winds are evident from 30° to 60°N, associated with the subtropical and the polar jets. Essential features include the enhanced wind velocities near Japan and the east coast of the United States and a strong poleward component over the far North Atlantic. At low latitudes the jet stream level winds are weak and predominantly easterly, consistent with observations.

For purposes of this work, however, the most relevant dynamical quantities are the meridional transport of water vapor, heat, and angular momentum. Comparison of these quantities predicted by our GCM and observations [e.g., Oort and Rasmusson, 1971] indicate agreement generally better than 75%. This provides strong evidence that the GCM can simulate the transport of trace species to a comparable accuracy, which should be adequate for the present investigation. Another relevant question is the accuracy of the coarse resolution model used in this work compared with a finer resolution model. A comparative study [Hansen et al., 1983] performed with a GCM similar but not identical to this one indicates that the
transport properties of the coarse resolution (8° × 10°) model are similar to those of a model with twice the horizontal resolution (4° × 5°).

At hourly time steps the instantaneous mass distribution and the averaged horizontal mass fluxes are fed into the tracer program. These mass fluxes are averaged over 6 hour intervals [Mahlman and Moxim, 1978]. The vertical mass fluxes are derived from the horizontal mass fluxes and the continuity equation. We also included monthly averaged vertical mixing by dry and moist convection. Advection of a tracer is accomplished by a novel finite differencing technique. For each grid box we predict four prognostic quantities: the mean concentration of the tracer $n$ and its spatial gradient $Vn$. The scheme determines the four quantities for the next time step by using an upstream method. This algorithm is comparable in accuracy to a fourth-order scheme [Mahlman and Sinclair, 1977], but it has much less numerical noise. The mathematical details are presented by Russell and Lerner [1981].

The chemical sources and sinks of CO in the tracer continuity equation are provided by a simplified chemical model, which is described in greater detail below. The concentration of the OH radical at each grid point is calculated by using the current abundance of CO. We then use this value of OH to calculate the methane oxidation source (R16) as well as the chemical sink (R18) for CO. The values of OH are updated monthly, which is generally adequate, as indicated by comparisons with test runs in which the updating is done every 10 days. The nonphotochemical sources and sinks are assumed to be time independent and are modeled as constant release rates or destruction coefficients over the relevant surfaces.

Each experiment starts with a uniform distribution of CO, with a volume mixing ratio $1 \times 10^{-7}$. The tracer model is then run for 4 years, repeatedly by using one year's wind data generated by the general circulation model. That the tracer model has reached steady state in the fourth year is indicated by the lack of change in the global CO distributions between the third and fourth years. An a posteriori check on the mass balance in the tracer model reveals that it conserves mass to better than 1% per year, which is adequate for the present investigations.

Photochemical production and loss rates in the troposphere are constructed, using the set of reactions shown in Table 1a. The zonally averaged CO distribution is used to generate photochemical equilibrium solutions for the free radicals OH, HO$_2$, H$_2$O$_2$, O(1D), NO, and NO$_2$. Distributions of water vapor mixing ratios and temperatures are taken from the general circulation model. Ozone concentrations and their seasonal variations are taken from the observations of Hering and Borden [1964]. The concentrations of CH$_4$ and H$_2$ are set equal to 1.4 and 0.5 ppm, respectively, throughout the troposphere. The distribution of NO$_x$ (NO + NO$_2$) is taken from Crutzen and Fishman [1977], with surface values of 0.1 ppb at 40°N and northward and 0.05 ppb southward, along with a scale height of 2 km. Because of a lack of data, no seasonal variations are

![Jet Stream Winds](image-url)

**Fig. 1b.** Jet stream level winds predicted by general circulation model.
prescribed for the NOx concentrations. Recently, Kley et al. [1981] reported somewhat lower NOx levels in the lower troposphere and higher values in the upper troposphere than we have used. If we had assumed that these values were representative of the background troposphere, our calculated OH levels in the lower troposphere would have been marginally affected while upper tropospheric values would have been higher. However, since most of the CO mass loss occurs in the lowest troposphere and the lifetime of CO is much longer in the upper troposphere, the impact of this change on our calculated budgets would be very small. The diurnally averaged photodissociation rates are calculated in the usual way, with corrections for Rayleigh scattering [Yung et al., 1980]. The effect of cloud cover is approximated by dividing all dissociation rates in the lower troposphere below the cloud heights (5 km) by a factor of 2, since the average cloud cover is about 50%.

In arriving at the simplified chemistry shown in Table 1a we have used several approximations regarding the interactions between the hydroxyl radical, nitrogen oxides, and methane and formaldehyde. Since we fix total NOx (= NO + NO2), the only HOx - NOx couplings are via the ratio,

\[
\frac{NO}{NO_2} = \frac{J_7}{J_4} \frac{k_{16}[O_3]}{[H_2CO]} + \frac{k_8}{k_9} \tag{1}
\]

and the formation of nitric acid,

\[
NO_2 + OH + M \rightarrow HNO_3 + M \tag{R12}
\]

Since the primary fate of HNO3 formed in (R12) is loss by rainout, it is a good approximation to assume that (R12) is a sink for HOx. The coupling between HOx and radicals derived from the oxidation of methane is more complicated. We note that to an excellent approximation the destruction of each CH4 molecule leads to the production of one H2CO molecule. Hence, the net result of methane chemistry in HOx can be parameterized by the following expressions,

\[
P(H) = J_4[H_2CO] \approx \gamma_2[OH] \tag{2}
\]

\[
P(HO_2) = k_{20}[H_2CO][OH] + J_4[H_2CO] \approx \gamma_1[OH]^2 + \gamma_2[OH] \tag{3}
\]

\[
L(OH) = k_{20}[H_2CO][OH] \approx \gamma_1[OH]^2 \tag{4}
\]

where

\[
\gamma_1 = k_{20}k_{16}[CH_4]/(J_4 + J_3 + k_{20}[OH])
\]

\[
\gamma_2 = J_4\gamma_1/k_{20}
\]

and \(P(x)\) and \(L(x)\) denote the production and loss rate (cm\(^{-3}\) s\(^{-1}\)) for species \(x\), respectively.

With these approximations we can calculate the photochemical equilibrium concentrations of the hydroxyl radicals (H, OH, HO2, and H2O2) by balancing the production and loss rates for these radicals, as given in Table 1a and expressions (2)-(4). This involves solving four simultaneous equations.

\[\text{Input to Chemical Model} \quad \text{Output from Chemical Model}\]

<table>
<thead>
<tr>
<th>Input to Chemical Model</th>
<th>Output from Chemical Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>(T = 281^\circ K)</td>
<td>([O^1D] = 1.7 \times 10^{-3})</td>
</tr>
<tr>
<td>(M = 2.3 \times 10^{-19})</td>
<td>([\text{H}] = 1.5 \times 10^4)</td>
</tr>
<tr>
<td>([H_2O] = 2.4 \times 10^{-17})</td>
<td>([\text{OH}] = 5.9 \times 10^3)</td>
</tr>
<tr>
<td>([O_2] = 8.4 \times 10^{-11})</td>
<td>([H_2O_2] = 1.8 \times 10^6)</td>
</tr>
<tr>
<td>([\text{NO}] = 1.6 \times 10^9)</td>
<td>([\text{H}_2\text{CO}] = 3.5 \times 10^{10})</td>
</tr>
<tr>
<td>([\text{CO}] = 4.4 \times 10^{12})</td>
<td>([\text{NO}] = 3.6 \times 10^8)</td>
</tr>
<tr>
<td>([\text{NO}_2] = 1.2 \times 10^9)</td>
<td></td>
</tr>
</tbody>
</table>

The kinetic rate coefficients are taken from Table 1a. All number densities are given in units of cm\(^{-3}\).
However, using the expressions
\[
[H] = \frac{k_{18}[CO] + k_{19}[H_2] + \gamma_2}{k_{11}[O_2]} \cdot M \quad (5)
\]
and
\[
[H_2O_2] = \frac{k_{14}}{J_3 + k_{15}[OH] + J_{22}} [HO_2]^2 \quad (6)
\]
further simplifies the mathematics to seeking the solution for only two nonlinear algebraic equations in two unknowns (OH and HO$_2$). The solution is then used to improve the estimate of NO/NO$_2$ by (1). This procedure converges in a few iterations to better than 10$^{-4}$. Table 1b gives a representative example of the concentrations of important radical species calculated by this simplified chemical model. The approximate scheme (1)-(6) calculates HO$_x$ concentrations that are within 10% of those computed by using the more extensive set of reactions in our one-dimensional photochemical model [Yung et al., 1980].

The stratospheric photochemical parameters are taken from two-dimensional model calculations by using the reaction scheme of Yung et al. [1980], in which the latitudinal distribution of the long-lived species (Cl$_x$, NO$_x$, CH$_x$, etc.) are prescribed by projecting the one-dimensional model results along preferred mixing surfaces [McElroy et al., 1976]. The distribution of stratospheric water vapor is taken from Wofsy [1980].

**EXPERIMENTS**

For this investigation, we performed a total of eight experiments. They were designed to consider a number of hypothetical sources and sinks in order to derive a budget consistent with atmospheric observations. The input parameters for these models are discussed below, and a brief summary appears in Table 2.

In experiment T1 we included an anthropogenic source of 6.3 x 10$^{14}$ g/yr, [Seiler, 1974]. The latitudinal distribution of this source was prescribed by assuming that emissions are proportional to energy consumption in a given latitude belt. The data for energy consumption were taken from Darmstadter et al. [1971]. No seasonal variation in the anthropogenic source was included, as this effect is thought to be small [Hall et al., 1975]. We also included a source from methane oxidation and a sink due to the oxidation of CO by reaction with OH. Both processes were distributed throughout the troposphere and stratosphere.

Tracer experiment T2 included the same source and sink terms as T1 except that the source from methane oxidation and loss by reaction with OH radicals were both arbitrarily reduced after each calculation to ensure a mean global abundance close to observations. This was done to determine the sensitivity of the model to the OH value.

Experiment T3 included, in addition to the sources and sinks used in T1, a plant source of magnitude 4.3 x 10$^{14}$ g/yr, equal to the minimum estimate of Zimmerman et al. [1978], while in experiment T4 we used their upper limit of 1.3 x 10$^{15}$ g/yr. The latitudinal distribution of these sources was also taken from Zimmerman et al. for both experiments. Although the equatorial ground source has been parameterized by using values estimated by Zimmerman et al. [1978] for NMHC oxidation, there are other possibilities. For example, Crutzen et al. [1979] suggested that the burning of vegetation either by forest fires or slash and burn agricultural practices, which are concentrated mainly in the tropics, might be a large CO source.

In experiment T5, we used the anthropogenic source and included the methane oxidation source and OH oxidation sink only in the stratosphere, to investigate the importance of transport to the stratosphere as a sink [Newell, 1977] and to isolate the effects of tropospheric photochemistry. Based on previous experience with this combination, we anticipated that an additional loss term would be necessary; thus, in experiment T6 we parameterized a soil sink by using a deposition velocity of 4 x 10$^{-2}$ cm/s, as estimated by Liebl and Seiler [1976].

In experiment T7, we incorporated an ocean source, which might be associated with marine biological activity. While estimates of a potential oceanic source vary widely, we have used

**TABLE 2. Summary of Sources and Sinks Used in the Construction of the Tracer Models**

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Sources</th>
<th>Sinks</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1</td>
<td>anthropogenic (a)</td>
<td>CO + OH</td>
</tr>
<tr>
<td>T2</td>
<td>anthropogenic (a)</td>
<td>1/2 (CO + OH)</td>
</tr>
<tr>
<td>T3</td>
<td>anthropogenic (a)</td>
<td>CH$_4$ oxidation (b)</td>
</tr>
<tr>
<td>T4</td>
<td>anthropogenic (a)</td>
<td>plant source MIN (c)</td>
</tr>
<tr>
<td>T5</td>
<td>anthropogenic (a)</td>
<td>CH$_4$ oxidation (b)</td>
</tr>
<tr>
<td>T6</td>
<td>anthropogenic (a)</td>
<td>CO + OH stratosphere (e)</td>
</tr>
<tr>
<td>T7</td>
<td>anthropogenic (a)</td>
<td>CH$_4$ oxidation stratosphere (e)</td>
</tr>
<tr>
<td>T8</td>
<td>anthropogenic (a)</td>
<td>ocean (g)</td>
</tr>
<tr>
<td></td>
<td>anthropogenic (b)</td>
<td>CO + OH stratosphere (e)</td>
</tr>
<tr>
<td></td>
<td>CH$_4$ oxidation stratosphere (e)</td>
<td>CO + OH</td>
</tr>
<tr>
<td></td>
<td>CH$_4$ oxidation (b)</td>
<td>plant source max (d)</td>
</tr>
<tr>
<td></td>
<td>CO + OH stratosphere (e)</td>
<td></td>
</tr>
</tbody>
</table>

a. 6.3 x 10$^{14}$ g/yr$^{-1}$ [Seiler, 1974]; b. based on OH calculated in the chemical model (see text); c. 4.3 x 10$^{14}$ g/yr$^{-1}$ distributed according to Zimmerman et al. [1978], however, we do not distinguish between NMHC and vegetation burning sources; d. 1.3 x 10$^{15}$ g/yr$^{-1}$, otherwise same as (c); e. these reactions take place in the stratosphere only, and there is no tropospheric chemistry; f. deposition velocity over soil = 4 x 10$^{-3}$ cm s$^{-1}$; g. 4 x 10$^{13}$ g/yr$^{-1}$ [Seiler and Schmidt, 1975].
the (conservative) value of $0.4 \times 10^{14}$ g/yr, estimated by Seiler and Schmidt [1975], which was distributed uniformly over the oceans.

Finally, in experiment T8 we included the same sources as in T4, except that the anthropogenic source was excluded in order to investigate the impact of an anthropogenic releases on the global distribution.

The classification of sources into three groups—anthropogenic, CH oxidation, and plant—is somewhat arbitrary and is done solely for convenience. For instance, there would be a large anthropogenic component in the plant source, if a major component of that was due to slash and burn agricultural practices.

RESULTS AND DISCUSSION

Figures 2a and 2b depict the vertically and zonally averaged CO concentrations, as a function of latitude, calculated in the eight experiments, along with the observed values [after Seiler, 1974]. It is clear from a comparison of the model results with the observations that experiments T2 and T4 could produce a meridional distribution similar to that observed. Both models could also satisfy the mean global abundance equally as well. Experiment T5 (which did not consider tropospheric photochemistry at all) produced values roughly twice as large as observed. Unless transport to the stratosphere is substantially underestimated in the model, this result indicates that the stratosphere is not an important sink in the global CO cycle. When the soil sink was added, as in T6, concentrations were reduced to values lower than observed. Obviously, it would be possible to 'tune' the deposition velocity by varying it within the uncertainty range of Liebl and Seiler [1975] to force better agreement, and, therefore, definitive conclusions about the strength of a soil sink cannot be drawn from this experiment alone. However, as will be discussed later, measurements of the isotopic content of CO [Stevens et al., 1972] can be used to place constraints on the overall importance of a soil sink for the atmospheric CO budget. In general though, models (T2, T5, and T6) which calculated OH values lower than those in T4 produced interhemispheric ratios that differed from the observed by 20% or more. While the meridional distributions produced in experiments T4 and T5 are qualitatively similar, they have different causes. In T4, the CO abundance in the

| TABLE 3. Summary of Results Obtained From the Eight Tracer Experiments |
|------------------------|---|---|---|---|---|---|---|---|---|
| Experiments           | T1 | T2 | T3 | T4 | T5 | T6 | T7 | T8 | Observations |
| CO(ppb)               | 46 | 95 | 62 | 107| 241| 59 | 48 | 65 | 110 |
| L(OH + CO, $10^{14}$g/yr) | 15 | 8.9| 18.6| 25.9| 5.7 | 1.8 | 15.3 | 20.6 |
| P(CH$_4$ oxidation)   | 8.7| 2.6| 7.9 | 6.3 | 0.3 | 0.3 | 8.6 | 7.4 |
| P(Ind)                | 6.3| 6.3| 6.3 | 6.3 | 6.3 | 6.3 | 6.3 | 6.3 |
| P(NMHC)               | 6.3| 6.3| 6.3 | 6.3 | 6.3 | 6.3 | 6.3 | 6.3 |
| P(Soil)               | —  | —  | —  | —  | —  | —  | —  | —  |
| L(Ocean)              | —  | —  | —  | —  | —  | —  | —  | —  |
| OH(cm$^{-3} \times 10^{-5}$) | 10.0| 3.0| 9.0 | 7.1 | —  | —  | 9.9 | 8.4 |
| CO(NH)/CO(SH)         | 2.6| 3.0| 2.5 | 2.3 | 1.8 | 3.4 | 2.5 | 1.3 |
| OH(NH)/OH(SH)         | 0.7| 0.7| 0.7 | 0.7 | —  | —  | 0.7 | 0.8 |

CO concentrations are given in parts per billion by volume. Units for production and destruction mechanisms are $10^{14}$ g CO/yr. The abbreviation 'Ind' refers to industrial and NMHC to nonmethane hydrocarbon.
The latitudinal asymmetry in this model reflects the greater portion of land areas in the northern hemispheric tropics relative to the southern hemisphere. The important assumptions and results of experiments T1-T8 are summarized in Table 3.

The computed seasonal variations of CO at 55°N is shown in Figures 3a and 3b along with observations at Zvenigorod, USSR [Dianov-Klokov et al., 1978]. Figures 3c and 3d show the model-derived seasonal variations at 20°N. The data are based on the observations of Seiler et al. [1976] at Mauna Loa. Both data sets were shown to be relatively free of local pollution influences, and should be representative of conditions in the clean troposphere. In general, all the models that had produced...
OH levels in the troposphere much lower than those calculated in experiment T4 could not satisfactorily reproduce the observed amplitude and phase of the seasonal variation at 50°N.

The observations of CO at 55°N and 20°N show little or no phase difference. This behavior is in striking contrast to the seasonal variations of CO2, which show significant differences between latitude belts [Junge and Czeplak, 1968]. The seasonal variations of CO2 are driven mainly at high latitudes; at high northern latitudes the minimum occurs in late summer, which is 6 months different from the southern hemispheric summer minimum. At low northern latitudes, with the intrusion of air from the southern hemisphere, the phase is delayed. However, the observed seasonal variations of CO appear to be driven uniformly with latitude, indicating a shorter lifetime and a dominance of local sources and sinks. This is consistent with seasonal variations in the OH concentration as the primary mechanism responsible for the seasonal variations of CO. The concentration of OH depends strongly on the duration and intensity of UV sunlight. Model-derived OH values showed little or no variation in the times when maximum or minimum concentrations occur at different latitudes. The seasonal variations of CO observed by Seiler et al. [1976] and Dianov-Klokov et al. [1978] likewise showed no difference in phase, a difference which would have been expected if the seasonal forcing were to arise from some feature such as a seasonal variation in the anthropogenic input. There is, however, about a 30% discrepancy in the amplitude observed by Seiler and the results of model T4.

If the seasonal CO data are accurate, this suggests that local chemistry may be more important and the atmospheric OH may be even higher than it is in the model. We have examined the sensitivity of the results in a model in which the OH concentrations were increased by one third over the values in T4. The results showed that both the amplitudes and the phases of the seasonal variations were in much better agreement with observations. Alternatively, there may be a seasonal variation in the strength of the low latitude source. As there is little seasonality in primary productivity in tropics, which is dominated by rain forests, we would expect no seasonal variation in the strength of the tropical source, if the scaling of hydrocarbon emissions to productivity used by Zimmerman et al. [1978] is adopted. There may be a seasonal variation in the CO source derived from vegetation burning. In the northern hemispheric tropics, the dry season is in winter [Richards, 1979]. Enhanced forest fires in this season would contribute a source that could improve the comparison between the model and observations.

T5 and T6, which do not include tropospheric chemistry, show little seasonal variability, while the seasonal variations produced in T2 are somewhat smaller than observed. The comparisons in Figures 2–3 imply that both tropospheric chemistry and the anthropogenic source are necessary components of the CO budget.

We also investigated the latitude-height distribution of CO. Figure 4 shows an estimate of the observed distribution [Seiler, 1974]. The low level maximum at upper mid-latitudes, presumably due to the anthropogenic source, is apparent, as is a lower latitude upper tropospheric bulge in the CO isopleths. At 10°S, values increase by more than 75% between the surface and 12 km. All the model experiments fail to produce this feature except for T3 and T4, the experiments that include the plant CO course. Figure 4b shows the results from year 4 for T2. The omission of a low latitude source is primarily responsible for the absence of the equatorial bulge in this model. Figure 4c presents the results for T4. The upper tropospheric bulge is apparent as well as the maximum at high latitudes. The distribution above the tropopause is unrealistic due to the failure of a seven-layer model to adequately resolve the tropopause and thus to model accurately troposphere-stratosphere exchange.

It is of interest to understand how the upper troposphere bulge is produced in T4. As was indicated in the above discussion, T4 concentrations are largely produced locally, and this is true for the vertical distribution as well. The biologically produced CO at low latitudes is transported vertically, mainly by the model's small-scale convection, in agreement with the mechanism proposed by Falconer and Pratt [1980]. Indication that a local ground source is responsible for the observed bulge comes from the high altitude measurements by Gauntner et al. [1979] of CO and Aitken nuclei over the tropics. Their results
show that high CO concentrations are associated with high Aitken nuclei counts, indicating a common source for both of these quantities. The upper troposphere data of Seiler [1974] were taken over the Atlantic and are similar to those observed by Gauntner et al. [1979] over Africa. In addition, the data of Gauntner et al. show higher values at equatorial latitudes over Africa than over the Pacific. The results of T4 are consistent with these observations. The longer distance from the major northern hemispheric source regions result in lower values over the equatorial Pacific. In T2, the longer lifetime of CO allows for its more complete redistribution around the globe. The Gauntner et al. data were obtained on only one flight, so more data are needed to check if these are persistent features of the CO distribution, and to examine the seasonal variability of the interhemispheric transport of CO.

Based on the complete set of experiments, we conclude that T4 is the most successful combination of sources and sinks tried for simulating the observed CO distribution. It is of interest to understand why the results of experiments T2 and T4 could both be reasonably successful in satisfying constraints imposed by (1) the observed global mean abundance and (2) the observed latitudinal distribution, since their chemical lifetimes differ by almost a factor of 3. The continuity equation may be written as

$$\frac{\delta M}{\delta t} = H + V + P - L + S$$

(7)

where $M$ is the mass of tracer, $H$ the horizontal convergence associated with transport by eddies and the mean circulation, $P$ and $L$ the chemical production and loss terms, $V$ the vertical transport into the troposphere from the stratosphere, and $S$ the ground source. The results are shown in Figures 5a and 5b for T4 and T2, to emphasize the roles of the various terms in different experiments. In experiment T4, much of the CO is governed by a local cycle of sources and sinks, with concentrations peaking where the sources are a maximum and in the intervening 30° latitude belt, where the horizontal convergence from the neighboring zones is significant. In T2, the distribution at lower latitudes is more strongly influenced by the horizontal convergence. In this case, the long lifetime of CO allows for its long range advection, but in so doing it also reduces much of its variability.

The results of these experiments imply that the globally averaged OH concentration is approximately $7 \times 10^5$ cm$^{-3}$ and cannot be greatly less than this value. Singh [1977] has used observations of the interhemispheric difference of methyl chloroform (CH$_3$CCl$_3$) to deduce globally averaged OH concentrations of at most $4 \times 10^5$ cm$^{-3}$. It is of interest to note that the globally averaged OH concentration calculated in experiment T2 is approximately the same as derived by Singh in his analysis. However, two recent studies [Jeong and Kaufman, 1979; Kurylo et al., 1979], using independent techniques, have derived a revised rate coefficient for the reaction CH$_3$CCl$_3$ + OH. This value is about a factor of 2 less than that used by Singh in his analysis. These revisions imply that his derived OH concentrations should be increased by a factor of 2 and are consistent with values obtained in experiment T4. Isopleths of the mean concentrations of OH in units of 10$^5$ molecules cm$^{-3}$ for the month of January and July computed in experiment T4 are shown in Figures 6a and 6b. These results for OH are in good agreement with those derived recently by Volz et al. [1981] on the basis of $^{14}$CO studies.

Experiment T8 is designed to reconstruct the state of the atmosphere prior to the industrial era. The input parameters for T8 are the same as those for T4 except for the industrial source, which is absent in T8. A comparison between T4 and T8 suggests that there may be a 65% change in the northern hemispheric CO concentration due to industrial pollution and a much smaller change in the southern hemisphere. Indeed, a comparison between Shaw's [1958] data taken in Ohio during 1952-1953 and Dianov-Klokov's data in the 1970's in the USSR indicates that there might have been a substantial increase in the CO concentration during this period. The mean value of OH in T8 is about 20% higher than that in T4, which suggests that there has been an increase in the tropospheric lifetimes of species such as CH$_4$, CH$_3$Cl, CH$_3$Br, and CHCl$_3$. In our current investigation we have fixed the source strength of
CH$_4$, the shorter lifetime would have implied an increase of CH$_4$ concentration in the atmosphere. Indeed, the recent observations by Rasmussen et al. [1981] provide strong evidence for an increase of atmospheric CH$_4$.

A primary objective of this paper is to derive OH concentrations from the behavior of CO in the model atmosphere. This study is, therefore, more like a diagnostic study of CO and OH than a prognostic one. It should be pointed out that the GCM simulation of the CO distribution and its variations provides the best check for the OH concentrations computed by our chemical model. Owing to the intrinsic difficulty of defining the key inputs such as 0$_3$, H$_2$O, NO$_x$, cloud cover, and heterogeneous processes, all of which exhibit great spatial and temporal variability, no chemical model of the troposphere can claim to compute OH concentrations from first principles to better than a factor of 2. Note that the test of the quality of the OH calculations is based on the behavior of CO in the model and is largely independent of the uncertainties in our chemical model.

Global measurements of the $^{18}$O/$^{16}$O ratio by CO by Stevens et al. [1972] typically show values much lower than would be derived from the incomplete combustion of fossil fuels. Fractionation by destruction mechanisms cannot be invoked to explain the data. Loss by reaction with OH radicals (R20) would favor the lighter isotope [Weinstock and Chang, 1974]. Stevens et al. [1972] have measured the fractionation in CO consumption by soil bacteria and found that it favors the lighter isotopes. All of their northern hemispheric measurements indicate a predominance of light isotopic species during the summer when soil scavenging would be at its maximum, which is contrary to what would be expected if the soil were acting as a major sink. It would also be difficult to explain the data by assuming equilibrium of CO with respect to uptake and release by the soil. By analogy with CO$_2$, release from the soil and from plants may involve no fractionation or again favor the lighter isotope [Keeling, 1973].

The models that do not include vegetation sources have difficulty in explaining the isotopic pattern found in the northern hemisphere, as they predict northern hemispheric CO to be mainly anthropogenic throughout the year. Experiment T6, which assumes the only sink to be soil scavenging, is also inconsistent with the Stevens et al. [1972] results as the CO in the northern hemisphere would be even heavier than that released from combustion. As was mentioned previously, the seasonal pattern of the $^{18}$O/$^{16}$O ratio would be exactly opposite to what is observed. These results are consistent with the view that the soil plays a relatively minor role in the CO cycle.

Since carbon 13 values from fossil fuel CO overlap those from natural sources, these measurements may not be particularly useful as a diagnostic of source types. Measurements of carbon 14, on the other hand, might be more useful. The use of carbon 14 and oxygen 18 data would then minimize the uncertainties due to mixing of air masses. Figure 7 shows the annually averaged concentrations contributed by each of the sources we have included in experiment T4. They were derived by using each source separately with the OH field derived by T4 to determine the loss rate. The figure shows that virtually none of the anthropogenic CO emitted in the northern hemisphere reaches the southern hemisphere. The results also show that a sizeable fraction of the CO at middle to high northern latitudes is not anthropogenic. Our results are in qualitative agreement with the major conclusions reached by Weinstock and Niki [1972] and Stevens et al. [1972]. More definitive statements cannot be made until more clean air measurements are available. The two sets of measurements reported above were subject to local anthropogenic influences.
In this paper, we have performed and studied an extensive set of experiments in search for an understanding of four major features of atmospheric CO: (1) the meridional distribution, (2) the seasonal variation, (3) the equatorial bulge, and (4) the isotopic composition. The results demonstrate the power and versatility of the general circulation model approach, as compared with more traditional analyses [Wofsy, 1976; Crutzen and Fishman, 1977; Hameed and Stewart, 1979]. The most successful experiment, in reproducing all of the above observed features, T4, requires a large surface source of order $1.3 \times 10^{15}$ g/yr$^{-1}$, in addition to anthropogenic and CH$_4$ oxidation sources. This additional source must exist at low latitudes in the Hadley cell regime, in order to produce the observed equatorial bulge.

Our model is consistent with a globally averaged OH concentration of about $7 \times 10^5$ cm$^{-3}$. At this OH level, we are able to simulate the main features of the seasonal variation, observed by Dianov-Klokov et al. [1978] in Zvenigorod (55°N). The seasonal data obtained by Seiler et al. [1976] at Mauna Loa (20°N) could be simulated by an even higher concentration of OH at lower latitudes than that in our T4 experiment or, alternatively, a possible seasonality in the plant source. However, based on our current understanding of the CH$_3$CCl$_3$ distribution [Singh, 1977; Chang and Penner, 1978; Jeong and Kaufman, 1979; Kurylo et al., 1979], the global mean OH concentration cannot significantly exceed that calculated in T4. We also conclude, from our model studies, that the globally averaged OH concentration cannot be significantly lower than that calculated in experiment T4. The observed variability of CO is not obtained in models T2, T5, and T6, implying that a shorter chemical lifetime and hence additional sources are required. The results of the latter models are also inconsistent with observations of the isotopic content of CO. Of course the accuracy of quantitative inferences based on the lifetime of CO requires. The results of the latter models are also inconsistent with observations of the isotopic content of CO. Of course the accuracy of quantitative inferences based on the lifetime of CO are dependent on the general circulation model's ability to simulate atmospheric transport. To provide an independent assessment of tropospheric OH, we are planning to study CH$_3$CCl$_3$ including tests with improved vertical resolution for the general circulation model.

No doubt a different and perhaps more elaborate combination of sources and sinks and OH concentrations could also reproduce the observed features of CO. However, we have attempted to construct a model with the fewest number of assumptions possible, and within the current small set of assumptions we are confident of the uniqueness of our results. We expect to improve and refine our models as more CO measurements become available.

Measurements of the isotopes of CO formed by various mechanisms such as vegetation burning and the oxidation of hydrocarbons such as isoprenes and terpenes could be valuable in estimating the relative roles of these processes. Measurements of the seasonal variation of total CO concentration, and the isotopic composition at remote locations could be used with data on the source characteristics to further quantify the magnitudes of the sources. Locations already in use for monitoring CO$_2$, N$_2$O, and chlorofluorocarbons could be used for this purpose.

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Conclusions

In this paper, we have performed and studied an extensive set of experiments in search for an understanding of four major features of atmospheric CO: (1) the meridional distribution, (2) the seasonal variation, (3) the equatorial bulge, and (4) the isotopic composition. The results demonstrate the power and versatility of the general circulation model approach, as compared with more traditional analyses [Wofsy, 1976; Crutzen and Fishman, 1977; Hameed and Stewart, 1979]. The most successful experiment, in reproducing all of the above observed features, T4, requires a large surface source of order $1.3 \times 10^{15}$ g/yr$^{-1}$, in addition to anthropogenic and CH$_4$ oxidation sources. This additional source must exist at low latitudes in the Hadley cell regime, in order to produce the observed equatorial bulge.

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