

**Hydrogen peroxide
and chemical
instability**

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The annual cycle of hydrogen peroxide: is it an indicator of chemical instability?

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Received: 11 March 2004 – Accepted: 25 March 2004 – Published: 6 April 2004

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Abstract

A box model has been used to study the annual cycle in hydrogen peroxide concentrations with the objective of determining whether the observed difference in summer and winter values reflects instability in the underlying photochemistry. The model is run in both steady-state and time-dependent modes. The steady-state calculations show that, for some range of NO_x background levels, two stable solutions to the continuity equations exist for a period of days in spring and fall. The corresponding time-dependent model indicates that, for sufficiently high background NO_x concentrations, the spring and fall changes in H_2O_2 concentration may be interpreted as a forced transition between the two underlying stable regimes. The spring transition is more rapid than that in fall, an asymmetry that becomes more marked as background NO_x increases. This asymmetry is related to the different time scales involved in chemical production and loss of H_2O_2 . Observations of the spring increase in H_2O_2 concentration may therefore provide a better measure of the change in the underlying chemical regime than does the fall decrease.

1. Introduction

Several publications (e.g. White and Dietz, 1984; Stewart, 1993; Kleinman, 1994) have shown that the continuity equations that describe tropospheric photochemistry can exhibit bifurcations and consequent multiple steady-state solutions. These phenomena occur in the transition region between what may be characterized as low- NO_x and high- NO_x chemical regimes. It is not known whether these mathematical properties exhibited by steady-state models correspond to observable phenomena in the time-dependent world. This paper addresses the question of observable consequences.

Steady-state and time-dependent calculations over an annual cycle are carried out showing that bifurcations in the steady-state solutions have an analogue in the time-dependent case. Above a certain background NO_x level rapid changes in oxidant con-

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centrations occur in time-dependent calculations. These correspond to bifurcations in the steady-state case. The rapidity of the transition increases with increasing background NO_x .

It is obvious that such transitions do not occur on the global scale, a predominately low- NO_x environment. Their importance lies in the fact that, should they occur on more polluted regional scales, they can involve large changes in oxidant levels with a corresponding change in pollutants over a short time period of several days. Hydrogen peroxide, the focus of the discussion in this paper is an especially sensitive indicator of the low- NO_x , high- NO_x transition.

1.1. The observed annual cycle of H_2O_2

This paper will discuss the annual cycle of H_2O_2 in the context of the low- NO_x and high- NO_x chemical regimes described by Kleinman (1991). A basic difference in these regimes is the concentration and fate of oxidants. In the high- NO_x regime OH reacts predominantly with NO_2 to form nitric acid. Nitric acid is mostly removed by deposition before it can react to return its components to the atmosphere and its formation is thus a permanent loss for odd hydrogen ($\text{HO}_x = \text{OH} + \text{HO}_2$). Since H_2O_2 is formed through reaction of two HO_2 radicals it is particularly sensitive to the changes in concentration of HO_x components that occur in the transition from low- NO_x to high- NO_x conditions. We will explore the possibility that observed mixing ratio changes in the tropospheric boundary layer might, under some circumstances, be interpreted as resulting from instability in the underlying steady-state chemistry. Due to the sensitivity of hydrogen peroxide to such instability it is reasonable to concentrate on its annual cycle.

Most measurements of H_2O_2 have been made over relatively limited time spans (see the review by Lee et al., 2000). Measurements over an annual cycle have been made by Ayers et al. (1996) at Cape Grim, Tasmania (41°S) and Sakugawa and Kaplan (1989) in and near Los Angeles, CA. The Ayers et al. (1996) measurements represent unpolluted marine air and show monthly means varying from 0.16 to 1.4 ppb. The measurements of Gnauk et al. (1997) were carried out over a 3-year period, but are

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only reported for selected days or over short intervals. Peroxide observations were made over a four-month period during the Tropospheric Ozone Production about the Spring Equinox (TOPSE) experiment (Snow et al., 2003). Although this time period is of special interest to the present study, the NO_x mixing ratios were too low to contribute to low to high- NO_x transitions (Snow et al., 2003; Tie et al., 2003).

The body of observations covers different locations and times of year and is useful for deducing general characteristics of the annual cycle of H_2O_2 . The property of the H_2O_2 annual cycle of most interest in this study is the change from a winter minimum to a summer maximum under various levels of background pollution as represented by the NO_x mixing ratio.

1.2. Nonlinearities

The nonlinearity of the continuity equations describing various aspects of atmospheric photochemistry gives rise to a number of interesting consequences. This paper will focus on multiple solutions in tropospheric models, but these have also been discussed with regard to the stratosphere (Prather et al., 1979; Fox et al., 1982; Konovalov et al., 1999) and mesosphere (Yang and Brasseur, 1994).

A relatively simple consequence of nonlinearity that has been discussed in the context of tropospheric models for some time is 'turnaround' behavior. This refers to a change in sign of the rate of change of a species concentration, such as ozone, with respect to a monotonically changing control parameter, such as a nitric oxide source. Stewart et al. (1977) described the change in O_3 response with increasing NO_x emission. They showed that $[\text{O}_3]$ first increased, then decreased as NO emission increased, with a turnaround occurring when the NO_x background reached roughly 500 ppt. Hameed et al. (1979) discussed the importance of the NO_x background on OH behavior and on the effect of increased CO source levels in the troposphere. The hydroxyl radical, like ozone, exhibits a turnaround as we go from a low to a high- NO_x background. It was shown that an increasing CO source could either increase OH and decrease CH_4 or have the opposite effect as NO_x background changes. Poppe et

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al. (1993) discussed the nonlinear OH response to changes in several model parameters.

White and Dietz (1984) were the first to describe a more startling aspect of nonlinearity in a tropospheric model, the existence of multiple steady states. These authors showed that the NO_x removal rate is a non-monotonic function of NO_x concentration. This implies that multiple concentration values can exist for each NO_x source value within a specific range of NO_x emission rates. Stewart (1993, 1995) described methods for computing multiple solutions and exhibited them for a variety of assumptions regarding both the reactions included in a model and the source values for key species. Multiple solutions occur when a control parameter, typically NO source, is varied past a critical value at which a bifurcation from one to three solutions occurs. Further variation in the same direction leads to another bifurcation in which the number of solutions is reduced to one. These bifurcation points bound a bistable region in the state space of the system, i.e. a region in which two stable steady states exist. It has not been clear how, or if, the bistability exhibited in simple models relates to observable phenomena. Kleinman (1991) suggested that the difference in summer and winter H_2O_2 concentrations observed in the northeastern United States might reflect the difference in underlying low- NO_x (summer) and high- NO_x (winter) photochemical states. Jacob et al. (1995) suggest that measurement of H_2O_2 , and several other species, show evidence of a September transition from low- NO_x (NO_x -limited) to high- NO_x (hydrocarbon-limited) conditions. Their data, along with other observations noted in the paper, suggest a roughly factor of 5 decrease in H_2O_2 during September.

1.3. Chemical instability

The terms “chemical instability” and “unstable solution” are used throughout this paper. Both steady-state and time-dependent solutions of continuity equations are presented to illustrate the phenomena described by these terms and possible atmospheric consequences of such phenomena. For corresponding parameter values, the steady-state solutions are expected to serve as “attractors” for the time-dependent model. For exam-

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ple, the solutions of a diurnally forced model should remain close to the corresponding solutions of a diurnally averaged one. In this sense, steady state models can provide an approximation to averaged conditions in the atmosphere. An exception occurs if the steady-state solution is unstable.

5 An unstable solution refers to a steady-state model in which the Jacobian matrix, evaluated at a solution, has at least one eigenvalue with a positive real part. The Newton-Raphson method used in solving the steady-state model (Stewart, 1993) uses the Jacobian, but not its eigenvalues. In a steady-state model there is thus no computational difference between stable and unstable solutions. If we now solve the same
10 equations, but use a time-dependent method to study the behavior of the solutions near a steady state, the Jacobian eigenvalues appear as arguments of exponential functions that describe the model's temporal evolution. If the real parts of all eigenvalues are negative, the solution simply relaxes to the steady state, but any positive real part will drive the time-dependent solution away from the steady state. This is
15 the meaning of instability and the consequence is that unstable steady-state solutions are inaccessible to a time-dependent model and cannot provide an approximation to conditions in the atmosphere.

Our interest in instability is in determining how a time-dependent model behaves when underlying steady-state attractors, differing only by a small value of a parameter,
20 are nonetheless widely separated in the state space of the system. This will occur in bistable regions referred to above. An example in the following is provided by steady states calculated at noon on successive days of the year. The small change in solar zenith angle that this entails will, under some conditions, result in large changes in species concentrations. When these steady-state changes have a time-dependent
25 analog we expect that, to the extent that the time-dependent model provides some approximation to the atmosphere, there may be observable consequences.

2. Background

2.1. Tropospheric chemical regimes

In both stratosphere and troposphere the source of ozone is molecular oxygen. In the stratosphere a single UV photon having wavelength $\lambda \leq 242$ nm suffices to dissociate molecular oxygen and form O_3 via reaction of the product O atoms with O_2 . Since these photons do not penetrate to the troposphere, ozone production utilizes two less energetic photons and a more involved sequence of chemical reactions. The first photon, $\lambda \leq 340$ nm, photolyzes a seed molecule of O_3 yielding an excited-state oxygen atom, $O(^1D)$. This may then react with water vapor to form OH. OH oxidation of a hydrocarbon (HC), represented by CO and CH_4 in the following, releases H atoms or methyl radicals, respectively. These capture O_2 to form HO_2 or CH_3O_2 and, with sufficient NO present, an O atom is subsequently transferred to NO forming NO_2 . The second photon, $\lambda \leq 423$ nm, now photolyzes NO_2 releasing O to form O_3 as in the stratosphere. Since this process only consumes HC and H_2O and not NO_x or HO_x there is no need for further O_3 photolysis to continue O_3 production. Ozone in the troposphere is thus its own precursor and its production is autocatalytic (Stewart, 1995). These ideas are elaborated in the remainder of this section.

A simplified photochemical scheme will be used to illustrate the behavior of the model used in this study. Such a simplification is often used to demonstrate basic properties of tropospheric chemistry and variants have appeared often in the literature (Tinsley and Field, 2001; Field et al., 2001; Kalachev and Field, 2001; Prather, 1994). The actual model chemistry is described later, but the simplified scheme will be a useful reference for interpreting model results and understanding conditions in which bifurcations in the steady-state model and low- NO_x , high- NO_x transitions in the time-dependent model occur. Many authors have discussed the reactions described below (e.g. Levy, 1971; Crutzen and Zimmermann, 1991; Kleinman, 1991; Lelieveld et al., 2002).

The photochemical system may be regarded as consisting of three sets described as follows: 1) Initiation, 2) Chain propagation, and 3) Termination. The initiation phase

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begins with the photolysis of an ozone molecule by solar UV radiation in the 300–340 nm range. This produces excited-state O(¹D) atoms, some of which react with water vapor to form two OH radicals. These initial reactions may result in ozone loss or production, depending on subsequent reactions.

5 2.1.1. Initiation



The fate of oxidants after the initiation phase begins with oxidation of CO. Carbon monoxide serves here as a surrogate for CH₄ and other hydrocarbon species. Since NO_x and HO_x are not consumed in the following they are referred to collectively as chain propagation reactions.

10 2.1.2. Chain propagation



For simplicity only the following chain termination reactions are considered.

2.1.3. Termination



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In a low- NO_x regime the net chemistry may result in either ozone loss or production. If it results in production, large numbers of ozone molecules may be produced for each one initially photolyzed. Depending on the amount of CO or hydrocarbon (HC) available reactions (R3)–(R5) may occur several times before a termination reaction occurs that removes radicals or NO_x from the system. Suppose that reactions (R3)–(R5) occur m times before a termination reaction occurs. Further, within this sequence, suppose that reaction R4a occurs m_1 times and (R4b) m_2 times with $m_1+m_2=m$. Then the net effect of (R1)–(R5) is:

2.1.4. Net



The above system behaves in the following way:

- Net production of O_3 : $m_1 > m_2 + 1$
- O_3 neutral: $m_1 = m_2 + 1$
- Net destruction of O_3 : $m_1 < m_2 + 1$.

The transition from ozone destruction to ozone production in the low- NO_x regime occurs when the reaction rates (R4a) and (R4b) are approximately equal, i.e. $k_{4b}[\text{O}_3] = k_{4a}[\text{NO}]$.

Once there is sufficient NO in the system to provide net chemical production, we see from the net reaction that this production is autocatalytic with $m_1 - m_2$ ozone molecules produced for each one destroyed. For CH_4 or NMHC consumption, the potential ozone production is greater, but details will not be considered here.

Dominance of (R4b) over (R4a), if it occurs, does not imply that $\text{HO}_2 + \text{O}_3$ is itself a major ozone loss mechanism. The competition between O_3 and NO for HO_2 simply provides a switch that, if O_3 is favored, allows ozone photolysis to act as the principal ozone loss mechanism rather than as the first step in the autocatalytic production

of ozone. This process was described as inhibition (of ozone production) in Stewart (1995).

The effects of these reactions are summarized in Fig. 1. The circles represent species playing the role of sources (HC) or sinks (H_2O_2 and HNO_3) for the more reactive species shown in the boxes. In this figure, the light-blue boxes and circles and dark-blue arrows indicate the chemistry of the low- NO_x , ozone loss regime. OH production through reactions (R1) and (R2) is followed by conversion to HO_2 via hydrocarbon oxidation and then loss of HO_x and O_3 after reaction (R7) forms peroxide. The path (R3) followed by (R7) shows the loss of ozone and odd hydrogen. Net production of oxidants does not occur in this system.

Adding NO_x to the system provides the mechanism for autocatalytic production of ozone. This is shown by adding the brown boxes and green arrows in Fig. 1. The dotted green arrows in the diagram show the recycling of NO_x and HO_x . The arrows connected to small filled circles in the HO_2 box indicate the alternate pathways that may be followed by HO_2 . The low- NO_x switch from ozone loss to production occurs when the (R4a) path dominates (R4b) as described above.

Finally, when enough NO_x is present, OH will react preferentially with NO_2 to form nitric acid. This establishes the high- NO_x regime shown by adding the red circle (HNO_3) and arrows on the left side of Fig. 1. Once this occurs the low- NO_x paths on the right of the figure are substantially shut down and production of oxidants, including H_2O_2 , becomes negligible. The small filled circles in the OH box indicate the switch between low and high- NO_x chemistry.

2.2. Model description

2.2.1. Methods and parameters

Many model details are the same as described in Stewart (1993, 1995). A box model (Stewart et al., 1983) is used to compute species concentrations. A Newton-Raphson method (Press et al., 1992), modified to enhance its robustness (Stewart, 1993), is

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used to compute steady-state solutions and the LSODE integrator (Hindmarsh, 1983) is used for time-dependent runs. Methane is held fixed at a mixing ratio of 1.7 ppm. Temperature is assumed to vary through the year from 273K to 297K similar to the model of Kleinman (1991). Water vapor varies with an annual cycle consistent with the temperature change and the assumption of a fixed 60% relative humidity. Fixed ozone burdens of 300 D.U. and boundary layer height of 1 km. have been used in the results reported below. Test calculations have been made with annual variations of these last parameters, but without significant change in the results.

There are 26 variable species (Table 1) undergoing 65 reactions (Table 2). In addition to their chemical interactions, several species shown in Table 1 are assumed to have physical sources and/or sinks. The physical source of O₃ is transport from the stratosphere. The mean value used for the O₃ flux is in the range discussed by Holton and Lelieveld (1994). The CO source is based on a global anthropogenic source value (Olivier et al., 1999). Sources for ethene and ethane are assumed that give values of these species consistent with observations (Bottenheim and Shepherd, 1995) over some range of model runs. Except for H₂CO, deposition velocities are from the model study of Hauglustaine et al. (1994). The formaldehyde deposition rate is assumed the same as that for CH₃CHO.

Some experimentation has been carried out with all of these model parameters. The main qualitative features discussed below, namely the existence of spring and fall transitions between high and low-NO_x regimes in both steady-state and time-dependent models, have not been sensitive to the values used.

2.2.2. Chemistry

Table 2 presents the chemical reactions used in this study. Photolysis rates are computed from the Madronich (1987) model for clear sky conditions at 40° N. Reaction rates are mostly taken from Demore et al. (1997) and Atkinson et al. (1992). A few tests were made with updated values given in the compilation of Sander et al. (2003) with no significant change in results. Some of the reactions in Table 2 have products

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that react with O_2 . In such cases, the products shown are those subsequent to the O_2 reaction. Reaction 1 showing ozone yielding OH directly is a composite reaction having an effective rate coefficient that depends on water vapor concentration as well as total number density. The effective rate is similar to that discussed in Stewart (1995), but is now treated as a photolysis rate rather than a bimolecular rate involving H_2O . Hydrolysis of N_2O_5 (R41 in Table 2) follows the discussion of Dimitroulopoulou and Marsh (1997) using an effective rate that accounts for both heterogeneous and homogeneous reactions. Reactions (R50) and (R51) following the oxidation of C_2H_4 by OH (R49) are based on the oxidation sequence given in Barker (1995). The rate for (R50) is a guess taken to be of a magnitude similar to the rates for HO_2+NO and $CH_3O_2 + NO$. The thermolysis rate for (R51) is also a guess taken to be the same as the rate for HNO_4 thermolysis. The reactions (R52) and (R53) are based on Barker (1995) as well as the discussion of Paulson (1995).

3. Results

3.1. The computed annual cycle of H_2O_2

The primary control parameter in this study is the day of the year. Steady-state concentrations for the variable species given in Table 1 are calculated at noon on successive days throughout the year. This implies a corresponding small variation in solar zenith angle between runs and it is also reasonable to consider zenith angle as the primary control parameter. Noon photolysis rates are multiplied by the daylight fraction for each day to make the results more representative of diurnal averages obtained from a time dependent model than are noon values. The secondary control parameter is the nitric oxide source, S_{NO} . A range of source fluxes from 4×10^9 to $4 \times 10^{12} \text{ cm}^{-2} \text{ s}^{-1}$ has been used in the calculations. These are converted to source values by dividing by the boundary layer height (1 km). A time-dependent model has been run for a two-year period with annual and diurnal forcing for several of the S_{NO} values used in the

steady-state model. The annual cycles of H_2O_2 from the steady-state model and from the second year of the time-dependent model are compared in this section.

In panels (c) and (d) of the following Figs. 2–5, the displayed rates of OH and HO_2 reaction are keyed to the discussion in Sect. 2.1 and the mechanism shown in Fig. 1.

In some cases, at the higher NO_x levels, the formation of nitrous or pernitric acid is actually among the top three rates. Rapid photolysis or thermolysis however will make these “do nothing” reactions and their exclusion is justified. In the models displayed in the following figures, reaction of OH with NMHC is not competitive with the rates shown.

Figure 2 shows steady-state calculations (a), the corresponding time-dependent results (b), the time-dependent variation of noon values of OH loss due to reaction with CO and CH_4 and with NO_2 throughout the year (c), and the HO_2 loss due to reactions with NO and O_3 along with the H_2O_2 production rate (d). The NO source value in these calculations is the lowest considered and results in a NO_x background that ranges from about 40 to 80 ppt. A regional characterization based on this range would be remote tropical forest (National Research Council, 1992). This, and similar NRC characterizations noted later, refers only to NO_x levels and not to other aspects of the chemistry. Smaller NO_x sources would result in an annual low- NO_x cycle characteristic of the remote marine troposphere. Both steady-state and time-dependent H_2O_2 mixing ratios vary from a winter low of about 25 ppb to a summer high of about 2 ppb. The annual cycle of H_2O_2 shown in Fig. 2b is qualitatively similar to the observations of Ayers et al. (1996), but with a higher summer maximum as would be expected for higher $[\text{NO}_x]$ mixing ratio. Figure 2c indicates that low NO_x conditions prevail throughout the year in this model since nitric acid formation is never competitive with hydrocarbon oxidation. A similar comparison of HO_2 loss rates (d) shows that the chemistry results in net ozone destruction throughout most of the year. In this model ozone photolysis removes odd oxygen and peroxide formation removes odd hydrogen.

The effect of increasing S_{NO} from that used in this model is to depress the winter mixing ratio and increase the summer high. The chemistry remains low- NO_x through-

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out the year until, at $S_{\text{NO}} \approx 3.7 \times 10^5 \text{ cm}^{-3} \text{ s}^{-1}$, a narrow region of bistability first appears that spans a single day, 8 February in late winter and 25 November in fall. Further increases in S_{NO} increase the bistable ranges and move their locations towards spring and early fall.

Figure 3 shows the same series of calculations as Fig. 2 with an NO source of $2 \times 10^6 \text{ cm}^{-3} \text{ s}^{-1}$. This S_{NO} value was chosen because it is in a range characteristic of the northeastern US and is similar to values used by Kleinman (1991) and Jacob et al. (1995). The range of background NO_x mixing ratios in this calculation is 0.5 to 5 ppb, which may be classified as a rural environment (National Research Council, 1992). In this figure and those following the H_2O_2 units in the steady-state calculation, part (a), will be shown as Log_{10} concentration rather than mixing ratio, as in Fig. 2a, to show the full range of summer to winter variation. Mixing ratios will be retained for the time-dependent profile, Fig. 3b, since this will better exhibit the transitions of interest. The H_2O_2 detection limit depends on measurement method, but is of the order of several to several tens of ppt (Lee et al., 2000). Mixing ratios along the abscissa in Fig. 3b and subsequent figures may be regarded as unobservable.

The steady-state profile, Fig. 3a, now exhibits a mushroom shape (Grey and Scott, 1990). In this model the mushroom results from the symmetry of the spring/fall calculation and consists of two conjoined hysteresis curves. The bistable regions extend from days 78 to 91 in the spring (19 March to 1 April and days 275 to 288 in the fall (2–15 October). The stable high- NO_x (winter) and low- NO_x (summer) regions are separated by sequences of unstable states shown in red. The instability in the steady-state mode corresponds relatively well to the times at which the dominant pathway for OH reaction, Fig. 3c, switches from nitric acid formation to hydrocarbon oxidation in the spring (day 80, 21 March) and back to nitric acid formation in the fall (day 290, 17 October). The spring increase in time-dependent mixing ratio, Fig. 3b, is most rapid between days 78 and 86 (19–27 March) in reasonable agreement with the switch from high to low- NO_x conditions indicated by the steady-state model (Fig. 3a) and the comparison of OH reaction rates (Fig. 3c). During this time the peroxide mixing ratio increases from

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about 1 ppt to 300 ppt. Thereafter a slower increase to the summer maximum of about 2.5 ppb takes place. In the fall, the most rapid decrease in peroxide mixing ratio occurs between days 280 and 310 (2 October to 6 November) and is roughly the opposite of the spring increase, but over a longer time span. HO₂ loss rates shown in Fig. 3d indicate that the summer low-NO_x regime is ozone producing throughout. It also indicates, in contrast with the model in Fig. 2, that significant H₂O₂ production occurs only in summer.

As S_{NO} is increased from the value used in Fig. 3, the bistable regions in the steady-state calculation lengthen and the neck of the mushroom is pinched inward.

The changes resulting from further increase in S_{NO} are seen in Fig. 4a where the spring and fall bistable regions extend from days 95 to 158 (5 April to 7 June) and 205 to 268 (24 July to 25 September). The NO source used in the model shown in Fig. 4 is S_{NO}=3.73×10⁶. This value is chosen for display since it is the highest S_{NO} considered for which the time-dependent model shows high to low-NO_x transitions.

It is a consequence of the model design that the steady-state profile is symmetric about mid-summer. However, the steady-state calculation suggests that its time-dependent analogue will not be. As days progress through the seasons, a transition from winter to summer conditions will not be forced until 7 June and the corresponding return to winter not forced until 25 September. We thus expect a shift of the time-dependent profile towards the fall side of summer, the shift being greater the more pronounced the pinch in the neck of the mushroom. This is apparent in Fig. 4b along with a much more dramatic asymmetry in the rates of the spring increase and fall decrease of [H₂O₂]. The high to low-NO_x transition in the time-dependent H₂O₂ profile begins on day 195 (14 July) and is completed in 3 days. The mixing ratio increase in this period is from 0.5 ppt to 1.7 ppb. The much less abrupt fall transition occurs between days 250 and 280 (7 September to 7 October). The mixing ratio decreases from 0.4 ppb to 0.002 ppt during this 30 day interval. The switch between nitric acid formation and hydrocarbon (CO and CH₄) oxidation by OH occurs on days 195 and 248 (Fig. 4c) corresponding to the transitions described for the H₂O₂ mixing ratio. The

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chemistry, as expected, is ozone producing throughout the short low-NO_x portion of summer (Fig. 4d). The NO_x range in this model is about 70 ppb (winter/spring) to 2 ppb (summer), characteristic of rural to urban-suburban regions (National Research Council, 1992). The correspondence between steady-state and time-dependent models in the time of the spring transition is not as close as that shown in Fig. 3. Although, according to the steady-state model, a transition should be forced by 7 June it does not occur in the time-dependent case until 14 July.

Figure 5 shows the model calculations for $S_{\text{NO}}=4.0 \times 10^6 \text{ cm}^{-3} \text{ s}^{-1}$ and a corresponding summer to winter [NO_x] of 75 to 130 ppb. This is the next higher source value from that considered in Fig. 4. The mushroom configuration seen in Figs. 3a and 4a has now become a continuous branch of high NO_x states (Fig. 5a) that persists throughout the year and an isolated branch consisting of low-NO_x summer values and unstable states. An isolated branch of solutions as shown in Fig. 5a is called an “isola” (Grey and Scott, 1990).

For this and higher NO source and [NO_x] background values, the summer low-NO_x states are inaccessible from the time-dependent model. This model remains in a high-NO_x state throughout the year as indicated by the low H₂O₂ mixing ratios (Fig. 5b) and the dominance of nitric acid formation over hydrocarbon oxidation (Fig. 5c). The units in Fig. 5b are parts per quadrillion (ppqd). The HO₂ ozone Production/Loss switch remains in the production position (Figs. 5d and 1), but this is irrelevant since oxidant production has been effectively cut off by the switch from low to high-NO_x chemistry (Fig. 1, OH box). Figure 5d shows the negligible peroxide production rate. As S_{NO} increases further, the isola shown in Fig. 5a shrinks and finally disappears at $S_{\text{NO}} \approx 1.5 \times 10^7 \text{ cm}^{-3} \text{ s}^{-1}$.

The NO_x mixing ratios in this model are characteristic of the urban troposphere (National Research Council, 1992), but the overall model chemistry is not since no significant NMHC are included. In a more comprehensive chemical model, NMHC would result in greater HO_x production and defer an annually dominant high-NO_x regime to higher background NO_x levels. For example, the model study of Honoré et al. (2000)

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contains a more realistic urban chemistry. These authors employ a chemistry-transport box model and, for stagnant conditions, find two distinct diurnal equilibria for NO_x emissions \sim a few times $10^7 \text{ cm}^{-3} \text{ s}^{-1}$ (this estimate is based on their reported maximum NO_x emissions, NO_x emission weight and their maximum mixed layer height of 1.5 km).

5 Their bistable regime occurs at NO_x source values that would result in only high- NO_x solutions in the current model. This effect is also seen in many other models of urban photochemistry in which contour plots of O_3 (or O_3 production) as a function of NO_x and VOC (volatile organic compounds) mixing ratios show the low- NO_x (or NO_x -sensitive or NO_x -limited) to high- NO_x (or VOC-sensitive or VOC-limited) transition moving to higher NO_x values as VOC increases. Examples may be found in Sillman (1999) and Tonnesen (1999).

The progression of steady-state solutions shown in Figs. 2a–5a are slices through a three-dimensional (Day, S_{NO} , $[\text{H}_2\text{O}_2]$) surface for selected (Day, $[\text{H}_2\text{O}_2]$) planes perpendicular to the NO source axis. The full set of solutions is shown by the bifurcation surface in Fig. 6. In this figure the Day axis ranges from days 0 through 400 increasing away from the vertex. The NO source axis, labeled $\text{Log}(S/\text{cm}^3 \text{ s})$, ranges from 8 to 4 decreasing away from the vertex and the $\text{Log}(\text{H}_2\text{O}_2/\text{cm}^3)$ concentrations range through twelve orders of magnitude. The top part of this figure, furthest from the day axis, represents the purely low- NO_x solutions corresponding to situations like that shown in Fig. 2a. Since the annual range of low- NO_x peroxide values is only about 2 ppb, this portion of Fig. 6 has a flat appearance. Slices taken with a plane parallel to the plane defined by the (Day, $\text{Log}(\text{H}_2\text{O}_2)$) axes generate the profiles in Figs. 2a through 5a. The rightmost such plane generates the purely low- NO_x profile of Fig. 2a. As this plane is moved towards the front of the figure, the pure low- NO_x solutions give way to mushrooms with increasingly pinched necks, then to a combination of high- NO_x solution branches and low- NO_x isolas and finally to a single branch of high- NO_x solutions. These last are along the lower part of the figure nearest the day axis.

As stated in the preceding text, to test the robustness of the results some runs have been made with changes in selected parameter values and model reactions. Interest-

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ing variants on the above model are provided by cases in which, as S_{NO} is increased from low values, a high- NO_x isola, lying under the branch of low- NO_x states, first develops in winter. This isola grows and merges with the low- NO_x states to form a single mushroom branch, similar to Figs. 3a and 4a. The neck of the mushroom then pinches off to leave a continuous branch of high- NO_x states and a low- NO_x summer isola as in Fig. 5a. An example of such a system is a model consisting only of reactions (R1)–(R7) supplemented by HNO_3 deposition and H_2O_2 deposition, photolysis and reaction with OH. The results regarding the correspondence between steady-state bifurcations and time-dependent transitions in the model remain unaltered.

3.2. Annual variation of selected species

Table 3 shows the approximate winter to summer change in noon mixing ratio for selected species in the time-dependent models shown in Figs. 2b–5b. The corresponding labels in the table are F2–F5. These models are characterized by increasing assumed NO source values and hence by increasing NO_x mixing ratios.

Model F2 is typical of a low- NO_x , ozone destruction environment. Ozone in this case is higher in winter than summer since the enhanced summer photolysis acts to destroy it. This type of annual ozone cycle is seen in the remote marine troposphere at Cape Grim (Ayers et al., 1996). In this environment the seasonal variation of O_3 is negatively correlated with H_2O_2 and positively correlated with CO.

4. Discussion and summary

A series of steady-state and time-dependent models has been run with the objective of determining whether seasonal changes in H_2O_2 mixing ratios are indicative of an underlying instability in the photochemical system.

The annual progression of steady-state values shown in Figs. 2a–5a, begins with an annual cycle that is entirely low- NO_x (Fig. 2a). The time-dependent analogue, Fig. 2b,

shows a similar variation. This model, with a winter to summer NO_x range of 80 to 40 ppt, is characteristic of the relatively clean remote troposphere (National Research Council, 1992). The annual H_2O_2 mixing ratio is qualitatively similar to the monthly-averaged Cape Grim observations (Ayers et al., 1996), but with somewhat higher summer values.

As the assumed NO source is increased the annual progression of steady-states evolves from a purely low- NO_x cycle to one having winter high- NO_x and summer low- NO_x solutions separated by spring and fall transition regions. A region of bistability that contains a run of unstable solutions characterizes the transitions. The branch of solutions shown in Figs. 3a and 4a is known as a mushroom (Grey and Scott, 1990). As S_{NO} increases, the neck of the mushroom narrows and finally pinches off to leave a continuous branch of stable high- NO_x states that persists throughout the year and an isolated branch of summer low- NO_x states consisting of stable and unstable solutions (Fig. 5a). As S_{NO} is increased from the value used in the model shown in Fig. 5, the isola shrinks and ultimately disappears leaving only an annual progression of high- NO_x states. These are shown by the left-most profiles, parallel to the day axis, in Fig. 6.

The time-dependent analogues of the steady states, Figs. 2b–5b, do not precisely reflect the transitions in the latter, but there are many qualitative similarities. The most interesting feature of the time-dependent solutions is the asymmetry in spring and fall transitions. This is noticeable in Fig. 3b and obvious in Fig. 4b. The difference in rapidity of spring and fall transitions is a consequence of the different mechanisms involved in H_2O_2 production (spring) and loss (fall) mechanisms. In spring, peroxide production is limited by the rapid radical reactions in the chain propagation phase, reactions (R3)–(R5) in Sect. 2.1. In the fall its loss is limited by photolysis and reaction with the OH radical. These loss reactions are included in the model, but are not part of the simplified scheme given by reactions (R1) to (R7).

With reference to Fig. 1 and the simplified reaction set (R1)–(R7), as the NO_x background increases a point is reached at which the O_3 P/L “switch” in the HO_2 box is thrown from ozone loss to production. The autocatalysis indicated by reaction (R8)

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then results in an explosive growth of O_3 , OH and HO_2 . The rapid increase of H_2O_2 follows suit. In the model shown in Fig. 3b the H_2O_2 production rate resulting from reaction R7 increases from about .1 to .7 ppb/day in the 8-day interval of most rapid H_2O_2 growth from 19–27 March. In the model shown in Fig. 4b the corresponding production rate increases from .007 to 22 ppb/day over the three-day interval from 14–17 July. The corresponding fall losses occur between 2 October and 6 November (Fig. 3b) and 7 September to 7 October (Fig. 4b). Loss rates in the former case decrease from .56 to .004 ppb/day and from 1.4 to 3.5×10^{-6} ppb/day in the latter.

5. Conclusions

This paper has explored the relationship between the chemical instability seen in steady-state tropospheric models and corresponding time-dependent behavior. The objective has been to see whether the observed changes in H_2O_2 concentrations from summer to winter may, under some circumstances, be interpreted as a manifestation of chemical instability. The rapid growth of $[H_2O_2]$ predicted for spring high to low- NO_x transitions would be a more definitive indicator of instability than would the slower $[H_2O_2]$ decay in the fall low to high- NO_x transition. For NO source values greater than $2 \times 10^6 \text{ cm}^{-3} \text{ s}^{-1}$, the spring transition in the model reported here results in $[H_2O_2]$ increases from <10 ppt (i.e. unobservable) to a few hundred ppt in periods of 3 to 7 days. Shorter transition times correspond to higher S_{NO} , and thus higher background NO_x values. Ideally, observations would be carried out about the spring equinox in a region in which winter $[NO_x]$ is sufficient to govern a high- NO_x regime. A persistent interval, days to weeks, of below detection limit peroxide concentrations followed by a change over an interval of a week or less to a period of detectable and rapidly increasing concentrations would suggest that bifurcations in the chemistry might be invoked as an explanation. On the other hand, a relatively smooth transition lasting a month would indicate that such a nonlinear effect does not occur.

The model used in this study, in common with many models, gives results that are

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concise and readily interpretable. Nature is not so accommodating. Clouds and fogs form and dissipate, rain falls and winds blow. All such events influence the concentration of a moderately long-lived and soluble species such as H_2O_2 . The result is a degree of variability that is sure to complicate any attempt to relate observations to the consequences of chemical influences. The H_2O_2 observations during TOPSE (Snow et al., 2003) illustrate potential problems. The greatest difficulty in carrying out a program of H_2O_2 observations would probably be separating the effects of transport from local photochemistry. Should such a difficulty prove insurmountable it might still be possible to look for evidence of chemical instability in the transitions of a shorter-lived species, such as OH, that would quickly relax to the local chemistry following transport events.

Chemical instability is a mathematical characteristic of photochemical models. Due to the magnitude of the changes induced by such instability the question of the existence of atmospheric consequences should be pursued.

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Table 1. Variable species included in the present model.

No.	Species	Flux ($\text{cm}^{-2} \text{s}^{-1}$)	Dep. velocity (cm/s)
1	O ₃	7×10^{10}	0.4
2	OH		
3	HO ₂		
4	H ₂ O ₂		0.50
5	NO	variable	0.016
6	NO ₂		0.10
7	HNO ₃		4.0
8	CO	1×10^{11}	0.03
9	CH ₃ OOH		0.1
10	CH ₃ O ₂		
11	CH ₂ OOH		
12	H ₂ CO		0.25
13	CH ₃ OH		
14	NO ₃		0.1
15	N ₂ O ₅		4.0
16	HNO ₂		
17	HNO ₄		4.0
18	C ₂ H ₄	2×10^9	
19	C ₂ H ₄ OHO ₂		
20	C ₂ H ₄ OOH		
21	CH ₂ O ₂		
22	C ₂ H ₆	2×10^9	
23	C ₂ H ₅ O ₂		
24	CH ₃ CHO		0.25
25	CH ₃ CO ₃		
26	PAN		

Table 2. Reactions used in the current model. The temperature used for rate evaluation is 278 K. This is the temperature at noon on 1 January in the model. Rate numbers in parentheses correspond to the numbers in Sect. 2.1.

No.	Name	Rate(cm ³ s ⁻¹)
R1 (R1+R2)	O ₃ + hv → 2OH	2.03D-07
R2 (R5)	NO ₂ + hv → NO + O ₃	6.42D-03
R3	H ₂ O ₂ + hv → OH + OH	3.45D-06
R4	HNO ₃ + hv → OH + NO ₂	1.97D-07
R5	PAN + hv → CH ₃ CO ₃ + NO ₂	2.10D-07
R6	OH + HO ₂ → H ₂ O + O ₂	1.18D-10
R7	H ₂ O ₂ + OH → HO ₂ + H ₂ O	1.63D-12
R8	OH + O ₃ → HO ₂ + O ₂	5.45D-14
R9 (R4b)	HO ₂ + O ₃ → OH + 2O ₂	1.82D-15
R10 (R4a)	NO + HO ₂ → OH + NO ₂	8.60D-12
R11	NO + O ₃ → NO ₂ + O ₂	1.30D-14
R12 (R3)	CO + OH → CO ₂ + HO ₂	2.44D-13
R13	OH + OH + M → H ₂ O ₂ + M	6.15D-12
R14 (R6)	OH + NO ₂ + M → HNO ₃ + M	1.05D-11
R15 (R7)	HO ₂ + HO ₂ + M → H ₂ O ₂ + O ₂ + M	5.33D-12
R16	H ₂ CO + hv → 2HO ₂ + CO	1.66D-05
R17	H ₂ CO + hv → CO + H ₂	2.80D-05
R18	CH ₃ OOH + hv → CH ₃ O ₂ + OH	3.10D-06
R19	CH ₄ + OH → CH ₃ O ₂ + H ₂ O	4.15D-15
R20	CH ₃ O ₂ + HO ₂ → CH ₃ OOH + O ₂	6.74D-12
R21	CH ₃ OOH + OH → CH ₃ O ₂ + H ₂ O	5.46D-12
R22	CH ₃ OOH + OH → CH ₂ OOH + H ₂ O	2.30D-12
R23	CH ₂ OOH + M → H ₂ CO + OH	1.0DD-12
R24	CH ₃ O ₂ + CH ₃ O ₂ → 2H ₂ CO + 2HO ₂	1.95D-13
R25	CH ₃ O ₂ + CH ₃ O ₂ → H ₂ CO + CH ₃ OH + O ₂	2.92D-13
R26	CH ₃ OH + OH → H ₂ O + CH ₃ O ₂	8.15D-13
R27	NO + CH ₃ O ₂ → NO ₂ + H ₂ CO + HO ₂	8.21D-12
R28	H ₂ CO + OH → CO + HO ₂	1.00D-11
R29	NO ₃ + hv → NO ₂ + O ₃	1.56D-01
R30	NO ₃ + hv → NO + O ₂	2.04D-02
R31	N ₂ O ₅ + hv → NO ₂ + NO ₃	1.23D-05
R32	HNO ₃ + OH → NO ₃ + H ₂ O	1.96D-13
R33	NO ₂ + O ₃ → NO ₃ + O ₂	1.80D-17
R34	NO ₃ + NO ₂ → NO + NO ₂ + O ₂	5.39D-16
R35	NO ₃ + NO → 2NO ₂	2.76D-11
R36	NO ₃ + NO ₃ → 2NO ₂ + O ₂	1.56D-16

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Table 2. Continued.

No.	Name	Rate(cm ³ s ⁻¹)
R37	OH + NO ₃ → HO ₂ + NO ₂	2.30D-11
R38	HO ₂ + NO ₃ → HNO ₃ + O ₂	2.05D-12
R39	HO ₂ + NO ₃ → OH + NO ₂ + O ₂	3.55D-12
R40	NO ₃ + H ₂ CO → HNO ₃ + HO ₂ + CO	4.32D-16
R41	N ₂ O ₅ + H ₂ O → HNO ₃ + HNO ₃	9.85D-22
R42	NO ₃ + NO ₂ + M → N ₂ O ₅ + M	1.35D-12
R43	N ₂ O ₅ → NO ₃ + NO ₂	3.97D-03
R44	HNO ₂ + hν → OH + NO	2.50D-03
R45	HNO ₄ + hν → HO ₂ + NO ₂	2.03D-06
R46	NO ₂ + HO ₂ + M → HNO ₄ + M	1.62D-12
R47	OH + NO + M → HNO ₂ + M	7.90D-12
R48	HNO ₄ → HO ₂ + NO ₂	8.21D-03
R49	C ₂ H ₄ + OH + M → C ₂ H ₄ OHO ₂	8.16D-12
R50	C ₂ H ₄ OHO ₂ + NO → C ₂ H ₄ OOH + NO ₂	2.00D-12
R51	C ₂ H ₄ OOH + M → 2H ₂ CO + HO ₂	1.96D-02
R52	C ₂ H ₄ + O ₃ → H ₂ CO + CH ₂ O ₂	5.85D-19
R53	C ₂ H ₄ + O ₃ → H ₂ CO + HO ₂ + OH	5.85D-19
R54	CH ₂ O ₂ + HO ₂ → H ₂ CO + OH	1.00D-11
R55	C ₂ H ₆ + OH → C ₂ H ₅ O ₂ + H ₂ O	1.86D-13
R56	C ₂ H ₅ O ₂ + NO → NO ₂ + HO ₂ + CH ₃ CHO	9.66D-12
R57	CH ₃ CHO + hν → CH ₃ O ₂ + HO ₂ + CO	2.21D-06
R58	CH ₃ CHO + hν → CH ₃ CO ₃ + HO ₂	9.71D-07
R59	CH ₃ CHO + OH → CH ₃ CO ₃ + H ₂ O	1.48D-11
R60	CH ₃ CHO + NO ₃ → CH ₃ CO ₃ + HNO ₃	1.77D-15
R61	CH ₃ CO ₃ + NO ₂ → PAN	9.70D-12
R62	PAN → CH ₃ CO ₃ + NO ₂	2.05D-05
R63	CH ₃ CO ₃ + HO ₂ → M	1.51D-11
R64	CH ₃ CO ₃ + CH ₃ CO ₃ → 2CH ₃ O ₂	1.75D-11
R65	CH ₃ CO ₃ + NO → NO ₂ + CH ₃ O ₂	1.93D-11

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Table 3. Average winter (W) and Summer (S) values in selected species mixing ratios (all in ppb). Models F2 through F5 correspond to the calculations shown in Figs. 2–5.

Model	F2		F3		F4		F5	
Season	W	S	W	S	W	S	W	S
O ₃	75	55	70	85	20	100	40	40
NO _x	.08	.04	5	0.5	70	2	130	75
CO	95	70	90	55	130	55	130	125

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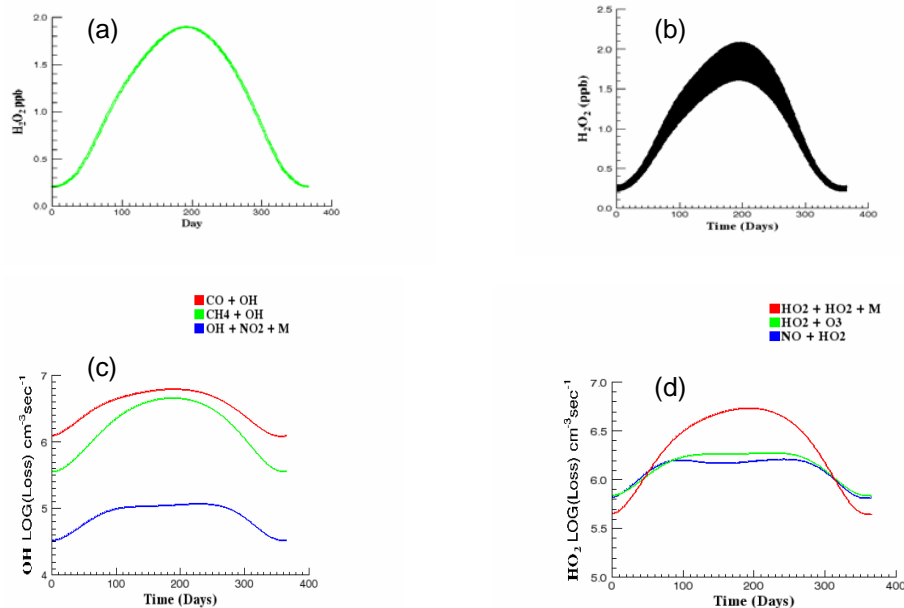


Fig. 2. Steady-state (a) and time-dependent (b), (c), (d) calculations for the lowest S_{NO} considered. The NO source here is $4 \times 10^4 \text{ cm}^{-3} \text{ s}^{-1}$ and results in a summer to winter range of background NO_x of 40 to 80 ppt. Low NO_x conditions prevail throughout the year as shown by the dominance of CO (red) and CH_4 (green) oxidation over HNO_3 formation (blue) in panel (c). Panel (d) indicates that the net chemistry is ozone destroying throughout the year. The color scheme in parts (c) and (d) is based on the integrated loss by a reaction over an annual cycle, not on the reaction name. In some of the following figures the reactions will be represented by different colors.

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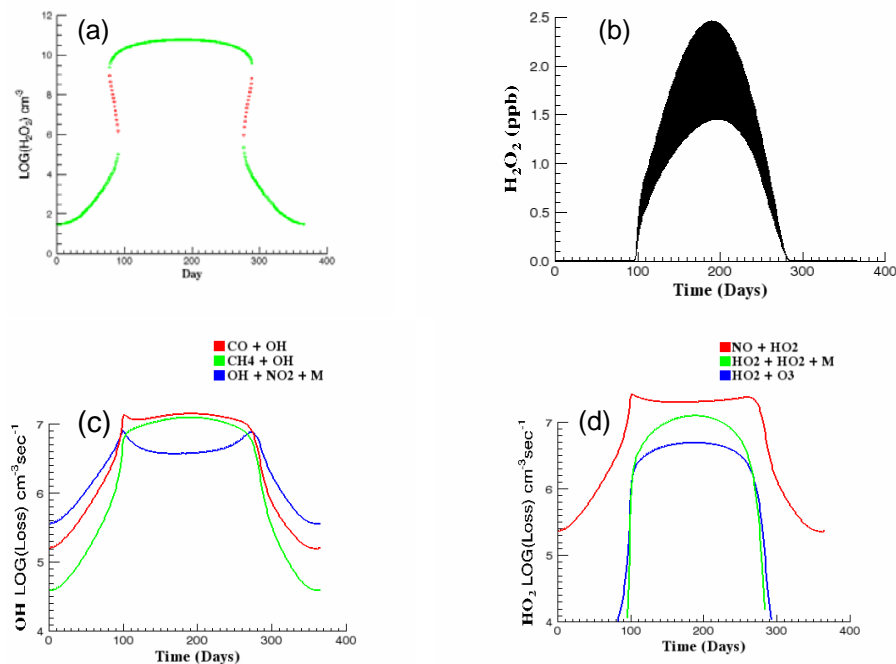


Fig. 3. Steady-state (a) and time-dependent (b) and (c) calculations for S_{NO} of $2 \times 10^6 \text{ cm}^{-3} \text{ s}^{-1}$. The summer to winter NO_x range in this model is 0.5 to 5 ppb. Part (c) shows that HNO_3 formation (blue) dominates CO (red) and CH_4 (green) oxidation in winter, while panel (d) shows the HO_2 switch (Fig. 1) favors ozone production (red) over loss (blue).

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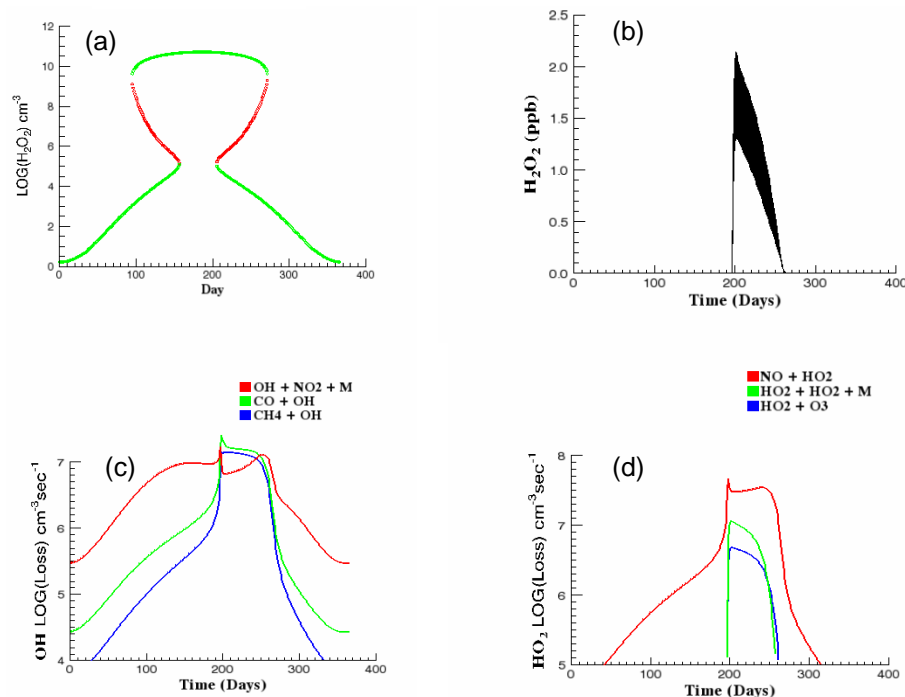


Fig. 4. Similar to Figs. 2 and 3, but with an assumed S_{NO} of $3.73 \times 10^6 \text{ cm}^{-3} \text{ s}^{-1}$. The summer to winter range of NO_x is 2 to 70 ppb. Part (c) shows that HNO_3 formation (now in red) dominates CO (green) and CH_4 (blue) oxidation except for a brief period of about 52 days in summer. In panel (d) the dominance of HO_2 reaction with NO (red) rather than O_3 (blue) facilitates ozone production in the low NO_x regime.

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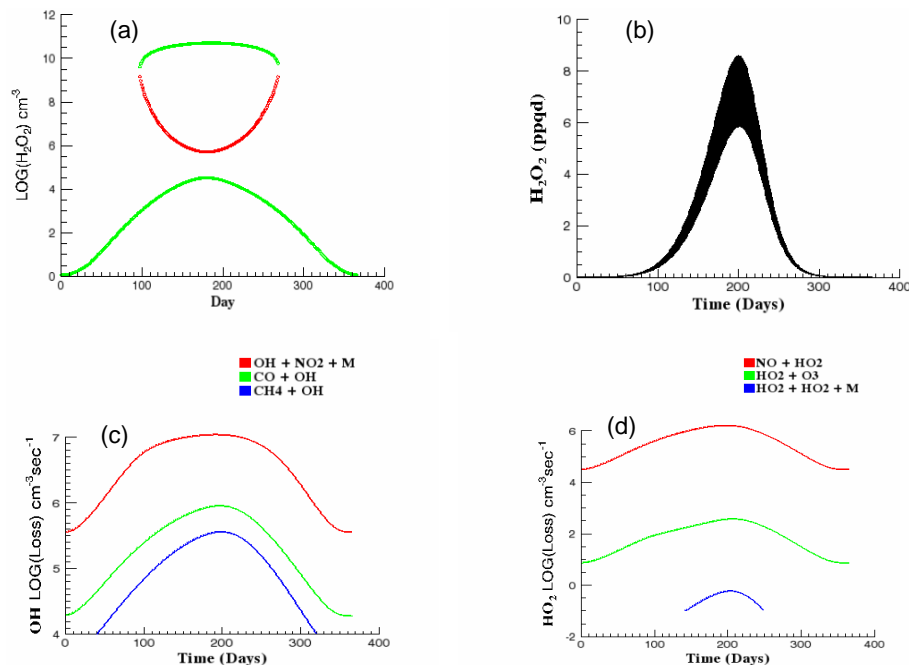


Fig. 5. Similar to Figs. 2–5 but for an assumed $S_{\text{NO}} = 4.0 \times 10^6 \text{ cm}^{-3} \text{ s}^{-1}$. The summer to winter NO_x range is 75 to 130 ppb. The OH reaction rates in part (c) indicate high- NO_x dominance throughout the year. In this high- NO_x regime H_2O_2 production is negligible (d).

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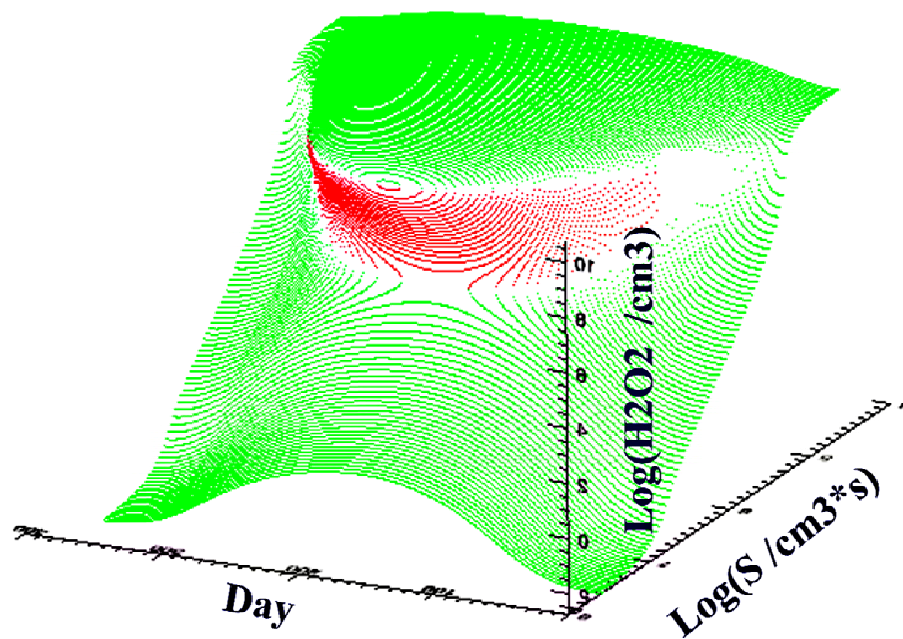


Fig. 6. H₂O₂ bifurcation surface. Selected slices through this surface parallel to the plane defined by the Day, Log(H₂O₂) axes generate the profiles shown in Figs. 2–5.

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