NONEQUILIBRIUM CO CHEMISTRY IN THE SOLAR ATMOSPHERE

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ABSTRACT

Investigating the reliability of the assumption of instantaneous chemical equilibrium (ICE) for calculating CO number density in the solar atmosphere is of crucial importance for the resolution of the long-standing controversy over the existence of "cool clouds" in the chromosphere and for determining whether the cool gas owes its existence to CO radiative cooling or to a hydrodynamical process. Here we report the first results of such an investigation in which we have carried out time-dependent gas-phase chemistry calculations in radiation hydrodynamical simulations of solar chromospheric dynamics. We show that while the ICE approximation turns out to be suitable for modeling the observed infrared CO lines at the solar disk center, it may substantially overestimate the heights of formation of strong CO lines synthesized close to the edge of the solar disk, especially concerning vigorous dynamic cases resulting from relatively strong photospheric disturbances. This happens because during the cool phases of the hydrodynamical simulations, the CO number density in the outer atmospheric regions is smaller than what is stipulated by the ICE approximation, resulting in decreased CO opacity in the solar chromosphere. As a result, the cool CO-bearing gas that produces the observed molecular lines must be located at atmospheric heights not greater than ~700 km. We conclude that taking into account the nonequilibrium CO chemistry improves the agreement with the available on-disk and off-limb observations but that the hydrodynamical simulation model has to be even cooler than anticipated by the ICE approximation, and this has to be the case at the "new" (i.e., deeper) formation regions of the rovibrational CO lines.

Subject headings: astrochemistry — molecular processes — radiative transfer — Sun: chromosphere

1. INTRODUCTION

Thirty years ago, Noyes & Hall (1972) inferred very low brightness temperatures ($T_b \approx 3700$ K) from their discovery of strong rovibrational CO lines at 4.7 $\mu$m observed close to the edge of the solar disk. Several years later, after such surprising observational results had been confirmed by Ayres & Testerman (1981), it was suggested that the low chromosphere might not be hot at all but could instead be permeated by CO-cooled "clouds" at altitudes between 500 and 1000 km above continuum optical depth unity (Ayres 1981). This led to controversy, because other (UV and submillimeter) diagnostics had suggested the existence of a uniformly hot chromosphere with a minimum temperature of about 4400 K near 500 km and a temperature rise above this temperature-minimum region. The controversy over the existence of cool gas in the low chromosphere continues today (see Kalkofen 2001; Ayres 2002; Avrett 2003), after the publication of an abundance of literature on the subject with theoretical investigations concluding that the CO lines have LTE source functions (Ayres & Wiedemann 1989; Uitenbroek 2000a) and with new observations showing off-limb CO emission protruding hundreds of kilometers into the chromosphere (Solanki, Livingston, & Ayres 1994; Clark et al. 1995), but also with the discovery that far-UV chromospheric lines observed on the solar disk always remain in emission at all positions and times (Carlsson, Judge, & Wilhem 1997).

Over the last few years, it has become increasingly evident that the next crucial step toward a better understanding of the enigmatic thermal structure of the solar chromosphere is to carefully investigate the reliability of the assumption of instantaneous chemical equilibrium (ICE), which is currently used for calculating the molecular number densities in stellar atmospheres (see, in particular, Uitenbroek 2000a, 2000b; see also Ayres & Rabin 1996; Avrett et al. 1996; Asensio Ramos & Trujillo Bueno 2003; Avrett 2003). Actually, both observations and simulations indicate that the solar chromosphere is a highly inhomogeneous and dynamic region of low-density plasma whose thermal, dynamic, and magnetic properties we need to decipher for unlocking new discoveries. If the ICE approximation turns out to be adequate for modeling the strongest CO lines close to the edge of the solar disk, then the available CO observations would really be indicating the existence of cool gas in the solar chromosphere. Otherwise, a natural resolution of the current chromospheric temperature discrepancy could perhaps emerge if the CO concentration turns out to be sufficiently lower than would be expected on the basis of chemical equilibrium. This Letter addresses this challenging issue by showing the first results of a detailed investigation on the nonequilibrium CO chemistry in the solar atmosphere.

2. FORMULATION OF THE PROBLEM

Our strategy consists in performing chemical evolution calculations in the radiation hydrodynamical simulations of solar chromospheric dynamics described by Carlsson & Stein (1997, 2002) that do not include CO cooling in the energy equation. Therefore, at each time step of the hydrodynamical simulation, we have fixed the ensuing thermodynamic conditions and calculated the corresponding CO number density by following the chemical evolution starting from the molecular concentrations of the previous time step.

The chemical evolution problem consists in calculating the molecular number densities at each time step of the hydrodynamical simulation by taking into account all relevant chemical reactions with their ensuing rates. In order to solve this
problem, one has to consider the following evolution equation for each chemical species "i" included in the model

\[
\frac{dn_i}{dt} = \sum_{A} \sum_{B} \sum_{C} k_{ABC}n_An_Bn_C + \sum_{A} \sum_{B} k_{AB}n_An_B \\
+ \sum_{A} k_A - \sum_{B} \sum_{C} k_{ABC}n_An_Bn_C \\
- \sum_{A} k_{A}n_A - k_n, \tag{1}
\]

where three-body reactions (first and fourth terms), two-body reactions (second and fifth terms), and one-body reactions (third and sixth terms) have been taken into account. When all the reaction rates for three-body (\(k_{ABC}\)), two-body (\(k_{AB}\)), and one-body reactions (\(k_A\)) are given, this set of equations for all the species included in the model form a set of first-order ordinary differential equations that must be solved numerically. Because of the stiffness of the system of equations (produced by the enormous differences in the abundances and their temporal variability), an implicit scheme must be used. Two key ingredients must be taken into account: the number of chemical species and the reaction rates for all possible reactions. After a careful investigation, we found that at least the following set of 13 species, which includes the most abundant diatomic molecules, is required to achieve a suitable description of the relevant chemical processes: H, C, O, N, He, CH, CO, H2, OH, NH, N2, NO, and CN. We have verified that the inclusion of ionic species does not significantly affect the CO concentration for the typical astrophysical conditions encountered at heights \(h \leq 1000\) km, although ideally they should be taken into account for very strong shocks capable of producing sizable changes in the degree of ionization. Concerning the reaction rates, we have used a reaction database created for the study of combustion mechanisms\(^5\) that seems to be appropriate for the physical conditions in the solar atmospheric plasma. Other databases such as UMIST (Bennett 1988) have been used for the study of the interstellar chemistry, but they suffer from the lack of certain important reactions that can efficiently take place in the Sun’s atmosphere (e.g., the catalytic three-body formation of molecular hydrogen: \(\text{H} + \text{H} + \text{He} \rightarrow \text{H}_2 + \text{He}\)). We have also investigated the possible influence of \(\text{CO}\) photodissociations, which are one-body reactions. They are produced mainly by discrete photon absorptions at wavelengths between the Lyman cutoff (912 Å) and the dissociation threshold (1120 Å). By using the photodissociation rates of Mamon, Glassgold, & Huggins (1988), we have verified that the contribution of photo-dissociation processes to the total CO concentration is negligible for the radiation field in the solar atmosphere.

3. RESULTS AND DISCUSSION

We used two time series of snapshots from the above-mentioned radiation hydrodynamical simulations, each one lasting about 3600 s and showing the upward propagation of acoustic wave trains growing in amplitude with height until they eventually produce shocks. The first one corresponds to a relatively strong photospheric disturbance showing well-developed cool phases and pronounced hot zones at chromospheric heights (see Carlsson & Stein 1997). The strongest dynamic cycle of this simulation produces a peak-to-peak line-core brightness temperature of 1000 K, concerning the strong 3–2 R14 CO line synthesized at disk center with the ICE approximation. However, the brightness temperature variations in most of the cycles of this simulation are \(~400\) K, which is similar to the observed values found by Uitenbroek, Noyes, & Rabin (1994) under excellent seeing conditions but larger than those inferred from the temperature histograms in Ayres & Rabin (1996). We refer to this as the strongly dynamic case.\(^6\) The second simulation corresponds to a much less intense photospheric disturbance (see Carlsson & Stein 2002), which for strong CO lines synthesized at disk center produces a peak-to-peak line-core brightness temperature fluctuation that is always smaller than 400 K. We refer to this as the weakly dynamic case.

Starting from the molecular concentration given by the ICE approximation, we have followed the chemical evolution in order to obtain the temporal variation of the CO number density \((N_{\text{CO}})\) at each height in the simulated solar atmosphere. As expected, the CO concentration is anticorrelated with the local temperature variations, yielding relatively low \(N_{\text{CO}}\) values in the hot phases and relatively high \(N_{\text{CO}}\) values in the cool phases. However, the amplitude of the local \(N_{\text{CO}}\) fluctuation is smaller than that given by the ICE approximation. In relatively low-density regions characteristic of the outer atmospheric layers (e.g., around \(h = 1000\) km), the ICE approximation does a fairly good job during the hot phases, but it overestimates the CO number density during the cool phases. In contrast, in relatively high-density regions characteristic of photospheric layers (e.g., around \(h = 400\) km), the ICE approximation underestimates the \(N_{\text{CO}}\) values during the hot phases. Thus, the CO abundance does not react instantaneously to the changes in the temperature because of the finite reaction rates. If one carries out a linear analysis by introducing a small temperature perturbation in a medium of given density and temperature and calculates the relaxation time needed to recover the original CO concentration, one finds that, for a given density, the relaxation time is larger the cooler the medium where the temperature perturbation is introduced. Similarly, for a given temperature of the unperturbed medium, the relaxation time increases rapidly with decreasing density. Short relaxation times are typical of high-temperature and high-density media (e.g., \(t_{\text{relax}} \approx 0.006\) s for \(n_{\text{H}} = 10^{16}\) \(\text{cm}^{-3}\) and \(T = 6000\) K), while long relaxation times are characteristic of low-temperature and low-density situations (e.g., \(t_{\text{relax}} \approx 400\) s for \(n_{\text{H}} = 10^{14}\) \(\text{cm}^{-3}\) and \(T = 4000\) K). One would find a broad range of relaxation times at a fixed height (e.g., \(h = 700\) km) in the dynamic atmosphere that results from the above-mentioned hydrodynamical simulations, simply because the existing rarefraction, compressions, and temperature fluctuations are continually changing the atmospheric conditions. Obviously, the situation is highly nonlinear and the relaxation time concept, although useful, loses its meaning in the real Sun. Therefore, any firm conclusion must be achieved via detailed numerical simulations.

Figure 1 refers to the strongly dynamic case. The left panel shows the temporal variation in the brightness temperatures of the line-core emergent intensities at \(\mu = 1\) and 0.1 in the strong 3–2 R14 CO line. The right panel gives the ensuing fluctuations of the atmospheric height where the line-core optical depth is unity, which we use as an indicator of the “representative height” where the CO line-core radiation originates. As seen in the figure, the ICE approximation does a sufficiently good job concerning the synthesis of the emergent CO spectrum at the solar disk center (\(\mu = 1\)), but it largely underestimates the line-core emergent intensities at \(\mu = 0.1\) during the cool phases.

\(^5\) See http://homepages.vub.ac.be/~akonnov.

\(^6\) This is similar to that used by Uitenbroek (2000a) and Ayres (2002) in their ICE modeling, although they considered only a segment of 190 s corresponding to the most dynamic cycle of the full simulation.
of the hydrodynamical simulations, producing brightness temperatures that are typically 500 K lower than those computed with the nonequilibrium CO concentration. Clearly, this is because during the cool phases, the ICE approximation overestimates the “heights of line formation” by about 300 km, concerning the synthesis of strong CO lines at $\mu = 0.1$ in the strongly dynamic case. This happens because during such cool episodes, the CO number density in the outer atmospheric regions is smaller than what is stipulated by the ICE approximation, resulting in decreased CO opacity in the solar chromosphere. Interestingly, in the weakly dynamic case that has smaller kinetic temperature fluctuations (but still larger than the fluctuations of observed brightness temperatures), the ICE approximation does a much better job, even at $\mu = 0.1$.

The reader may find surprising our conclusion that the ICE approximation is suitable for modeling the CO spectrum at the solar disk center, given that Uitenbroek (2000b) found that the spatially averaged line cores of weak CO lines synthesized in the three-dimensional hydrodynamical model of Stein & Nordlund (1989) are overly dark compared to the observed ATMOS spectrum described by Farmer & Norton (1989). As we will show in a forthcoming publication (A. Asensio Ramos & J. Trujillo Bueno 2003, in preparation), this is because such a three-dimensional hydrodynamical model of the solar photosphere is too cold in the CO line-forming region. In fact, our ICE synthesis of the 7–6 R68 CO line in the improved three-dimensional hydrodynamical model of Asplund et al. (2000) shows a notable agreement with the observed ATMOS spectrum, which constitutes an additional indication of the realism of the most recent three-dimensional hydrodynamical simulations of solar surface convection (see Shchukina & Trujillo Bueno 2001 concerning the iron spectrum in such a three-dimensional hydrodynamical model).

Figure 2 contrasts the time-averaged CO concentration obtained assuming ICE at each time step of the strongly dynamic simulation case with that resulting from the chemical evolution. Note that the ICE approximation leads to a significant overabundance of CO in the outer layers of the atmosphere (i.e., above 700 km). Thus, the CO number density can be relatively low in such outer layers, in spite of the fact that the temporally averaged temperature of the Carlsson & Stein (1997) simulations decreases outward and does not show any chromospheric temperature rise. As expected from the previously reported results, the ICE approximation does a sufficiently good job concerning the modeling of the temporally averaged CO spectrum at the solar disk center ($\mu = 1$). In contrast, the emergent CO spectrum computed close to the edge of the solar disk (i.e., at $\mu = 0.1$) shows significantly stronger CO lines when the ICE approximation is used, especially concerning strong CO lines such as the 3–2 R14 one, for which the line-core brightness temperature is about 100 K lower than that obtained using the nonequilibrium CO number densities.
Finally, we turn our attention to the modeling issue of the off-limb CO emission, which we have carried out by solving the radiative transfer equation in spherical coordinates at each time step of the hydrodynamical simulation.\(^7\) The off-limb distances where (time-averaged) CO emission has been observed depend on the line (see Ayres 2002 for a summary of the available off-limb observational results): the off-limb emission extension of strong lines such as the 3–2 R14 line lies between 0.55 and 0.77 above the 4.7 \(\mu\)m continuum limb, while weaker lines such as the 7–6 R68 line extend to \(~0.4\). Figure 3 refers to the strongly dynamic case. It shows that the atmospheric region where we can have a significant off-limb emission is extremely large when the ICE approximation is used, i.e., much larger than when the spectral synthesis is carried out using the nonequilibrium CO number densities. The dashed and solid lines show the corresponding height variation of the temporally averaged off-limb emission at the core of the strong 3–2 R14 CO line. They show that the nonequilibrium CO chemistry improves the agreement with the available off-limb observations. First, as seen in Figure 3, the ICE approximation predicts that appreciable emission in the 3–2 R14 line should continue to relatively large off-limb distances, while partial eclipse measurements by Clark et al. (1995) show a rapid disappearance of CO emission at heights above 700 km, which is more in line with our chemical evolution calculations. Second, the representative off-limb emission extension where the normalized intensity falls to 50% of the on-disk value is larger when the nonequilibrium CO concentration is used, which helps to improve the agreement with the observations of translimb emission extensions, although they still seem to be below the most recently observed values (see Ayres 2002).

4. CONCLUDING COMMENT

Our results indicate that the CO line radiation observed close to the edge of the solar disk comes from atmospheric heights not greater than \(~700\) km, and that the gas in these regions of the low chromosphere must be much cooler than indicated by the cool phases of the Carlsson & Stein (1997) hydrodynamical simulations. Lower temperatures will probably increase the relaxation times needed to reach the molecular equilibrium concentrations. In a forthcoming publication, we will show what happens when the hydrodynamical simulations themselves are carried out taking the CO cooling into account, in a way consistent with the nonequilibrium evolution of the molecular number densities.

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