# Premixed Carbon Monoxide–Nitrous Oxide–Hydrogen Flames: Measured and Calculated Burning Velocities with and without Fe(CO)<sub>5</sub><sup>‡</sup>

# GREGORY T. LINTERIS,\* MARC D. RUMMINGER,† AND VALERI I. BABUSHOK

Building and Fire Research Laboratory, National Institute of Standards and Technology, Gaithersburg, MD 20899

The burning velocity of premixed carbon monoxide–nitrous oxide flames (background water levels of 5 to 15 ppm) has been determined experimentally for a range of fuel–oxidizer equivalence ratio  $\phi$  from 0.6 to 3.0, with added nitrogen up to a mole fraction of  $X_{N_2}=0.25$ , and with hydrogen added up to  $X_{H_2}=0.005$ . Numerical modeling of the flames based on a recently developed kinetic mechanism predicts the burning velocity reasonably well, and indicates that the direct reaction of CO with  $N_2O$  is the most important reaction for CO and  $N_2O$  consumption for values of  $X_{H_2} \le 0.0014$ . The calculations show that a background  $N_2O$  is the most important reaction for CO and  $N_2O$  consumption for values of  $N_2O$  compared to the bone-dry case. Addition of iron pentacarbonyl, Fe(CO)<sub>5</sub>, a powerful flame inhibitor in hydrocarbon–air flames, increases the burning velocity of the CO– $N_2O$  flames significantly. The promotion is believed to be due to the iron-catalyzed gas-phase reaction of  $N_2O$  with CO, via  $N_2O+M=N_2+MO$  and CO + MO = CO<sub>2</sub> + M, where M is Fe, FeO, or FeOH. Published by Elsevier Science Inc.

### INTRODUCTION

The present investigation was conducted for the dual purpose of studying the effectiveness of the flame inhibitor iron pentacarbonyl in nonbranching chain flames of CO and N<sub>2</sub>O, and to investigate the role of the direct reaction of CO with N<sub>2</sub>O at flame temperatures with controlled amounts of hydrogen. The most effective chemical flame inhibitors are believed to act through catalytic cycles that recombine radicals in the flame. These inhibitors, however, have not been tested in systems without chain branching. In addition, we are also responding to the need described by Allen et al. [1] for burning velocity measurements in the CO-N<sub>2</sub>O system to assess the role of the direct reaction under dry conditions at flame temperatures.

The high-temperature gas-phase reactions of carbon monoxide and nitrous oxide are an

The CO-N<sub>2</sub>O reaction has been studied in shock tubes [5–7] and flow reactors [1]. Loirat et al. [8] measured the critical ignition pressure of CO-N<sub>2</sub>O mixtures in a cylindrical reactor. In flame studies, Dindi et al. [9] measured the stable species mole fractions in low-pressure premixed flames using gas chromatography, and Vandooren et al. [10] recently used mass spectrometry to measure the structure of CO-N<sub>2</sub>O-H<sub>2</sub> flames. Cor and Branch [11] measured the stable species profiles in low-pressure counterflow CO-N<sub>2</sub>O-N<sub>2</sub> diffusion flames. These studies have provided data for determining elementary rates and for testing comprehensive

important and well-studied system. These reactions occur in the gas-phase region during combustion of nitramine-based solid rocket propellants [2-4], and they are also important for understanding the combustion emission characteristics of stationary and mobile power plants. The direct reaction of CO with N<sub>2</sub>O is of fundamental interest because it is one of the simplest examples of an exchange reaction between saturated molecules. For inhibition studies, the reactant mixture provides a nonchain mechanism involving oxygen atoms, so that the significance of catalytic O-atom recombination cycles of iron species from the inhibitor can be tested, as well as those of H and OH when trace hydrogen is added as a reactant.

<sup>\*</sup>Corresponding author: Gregory T. Linteris, National Institute of Standards and Technology, Bldg. 224, Room B356, Stop 8651, Gaithersburg, MD 20899-8651, USA. E-mail: linteris@nist.gov

<sup>&</sup>lt;sup>†</sup> National Research Council/NIST Postdoctoral Fellow, 1996–1999; current address: Ceryx, Inc., Santa Paula, CA 93060.

<sup>\*</sup> Official contribution of the National Institute of Standards and Technology, not subject to copyright in the United States.

mechanisms. In many of the studies, the possible interference of H-atom reactions from impurities has been described, but not always quantified.

The burning velocity of premixed CO-N<sub>2</sub>O flames has been measured previously in three investigations. Van Wonterghem and Van Tiggelen [12] measured the flame speed of lean, stoichiometric, and rich flames, some with nitrogen dilution, having estimated hydrogen impurities of less than 2000 ppm<sup>1</sup> in the CO (but not noted for the N2O). Kalff and Alkemade [13] provided data on stoichiometric and rich flames with up to 10% added water vapor, and a minimum hydrogen content estimated to be less than 500 ppm. Simpson and Linnett [14] investigated quite rich systems ( $\phi = 2.0$ ), diluted by nitrogen, with unquantified, but low, levels of hydrogen impurities. The burning velocity of premixed CO-N2O flames has been calculated [1, 15], but the absence of data for flames with low hydrogen content was noted in both studies.

Since the levels of hydrogen are somewhat high or unquantified in previous experiments, additional experiments are required to understand the importance of the direct reaction at flame temperatures. We report burning velocity measurements for stoichiometric CO-N<sub>2</sub>O flames with added H<sub>2</sub> mole fractions from 0 ppm to 6800 ppm. For the driest conditions (5 to 15 ppm H<sub>2</sub>O), we also report flame speeds for equivalence ratios from 0.6 to 3.0, and for stoichiometric flames with nitrogen dilution up to 25% of the total volumetric flow. For all conditions, the flame structure is numerically calculated using a detailed chemical kinetic mechanism, providing an estimate for the rate of the direct reaction at the flame temperature, and allowing assessment of the relative importance of the different reaction routes for consumption of N<sub>2</sub>O and CO.

The CO-N<sub>2</sub>O flames were also used to study the inhibition mechanism of Fe(CO)<sub>5</sub>. There is an urgent need to find replacements for the effective and widely used fire suppressant CF<sub>3</sub>Br and related compounds [16]; however, a replacement with all of the desirable properties

of CF<sub>3</sub>Br is proving difficult to find and research has intensified [17]. Certain metallic compounds have been found to be substantially more effective flame inhibitors than halogencontaining compounds [18-20]. Iron pentacarbonyl is among the most effective flame inhibitors ever identified [18], up to two orders of magnitude more effective than CF<sub>3</sub>Br, and recent progress has been made in understanding its mechanism [21]. Although Fe(CO)<sub>5</sub> is flammable and highly toxic and could never be used as a fire suppressant itself, other iron-containing compounds have been identified which are far less toxic and flammable, and which also demonstrate strong flame inhibition [22]. If means can be identified to safely introduce metal species into a fire, they may be beneficial components in a fire suppressant blend, particularly for unoccupied spaces. Further, the fundamental understanding gained from study of these nearideal inhibitors, which catalytically recombine radicals, provides a theoretical basis for development of new inhibitors.

A detailed chemical kinetic mechanism for iron-species inhibition of flames has been introduced [23], and modeling with the mechanism supports the premise that the inhibition is primarily a gas-phase phenomenon. Numerical calculations using the mechanism predict many of the properties of the flames examined; nonetheless, some of the features of the flames are not well-described, and much work remains to be done to test and validate the mechanism. In particular, inhibition in lean flames (where Oatom reactions are much more important) is not accurately modeled by the mechanism.

In previous research, oxides of nitrogen have been used as the oxidizer in studies of the effectiveness of flame inhibitors [24]. Since systems using nitrogen oxides instead of  $O_2$  undergo non-chain-branching radical reaction sequences, the relative effectiveness of agents believed to act through catalytic radical recombination cycles may be very different, affording new opportunities to understand the inhibition by  $Fe(CO)_5$ .

In the absence of hydrogen, the oxidation mechanism of  $CO-N_2O$  flames is believed to proceed through either the direct reaction or through the thermal decomposition of  $N_2O$  followed by O-atom reaction with  $N_2O$  or CO.

 $<sup>^{1}</sup>$ All references to ppm in this paper are on a volume basis and refer to  $\mu$ l/L.

G. T. LINTERIS ET AL.

Recent research with Fe(CO)<sub>5</sub> inhibition of CO-O<sub>2</sub>-N<sub>2</sub>-H<sub>2</sub> flames has shown that in some systems a new O-atom radical recombination cycle may dominate radical destruction [25]. Also, Kaufman [26] has shown that Fe(CO)<sub>5</sub> addition to a flow tube causes very strong reduction in O-atom concentration. The extent to which Fe(CO)<sub>5</sub> affects dry CO-N<sub>2</sub>O flames will depend upon the predominance of the O-atom route for N<sub>2</sub>O and CO consumption and the effectiveness of the iron-catalyzed Oatom radical recombination cycle. Because gasphase iron is also believed to catalyze the decomposition of N<sub>2</sub>O (as discussed below), the relative efficiency of this reaction will also affect the influence of iron in this flame. In summary, in the flame inhibition experiments and modeling, we seek to determine the effectiveness of Fe(CO)<sub>5</sub> in a system that is non-chain branching, to test the postulate that for these systems, radical recombination—even by a very powerful catalytic agent—is not as effective. The results may have implications for the practical problem of the suppression of propellant fires.

### **EXPERIMENTAL**

A Mache-Hebra nozzle burner (inner diameter  $1.02 \text{ cm} \pm 0.005 \text{ cm}$ ) [27] with a schlieren imaging system [28] provides the average burning velocity of these Bunsen-type flames using the total area method [29]. The experimental system has been described in detail previously [30]. Calibrated mass flow controllers meter the gas flows, and a two-stage saturator adds Fe(CO)<sub>5</sub> as a vapor. The burner produces straight-sided schlieren images that are captured by a video frame-grabber board in a computer. The fuel gas is carbon monoxide (Matheson<sup>2</sup> Ultra High Purity (UHP), 99.97% CO, for which a batch analysis by the manufacturer shows the concentration of H<sub>2</sub>O and CH<sub>4</sub> to be less than 0.3 ppm and 1.0 ppm, respectively, and an in-house Fourier transform infrared radiation [FTIR] analysis showed  $\rm H_2O$  and HCs to be present at less than 1 ppm each). For some tests, the fuel stream contains added hydrogen (Matheson UHP, 99.999%  $\rm H_2$ , with sum of  $\rm N_2$ ,  $\rm O_2$ ,  $\rm CO_2$ ,  $\rm CO$ , Ar,  $\rm CH_4$ , and  $\rm H_2O < 10$  ppm). The oxidizer stream consists of nitrous oxide (Matheson UHP, 99.99%  $\rm N_2O$ , with sum of  $\rm N_2$ ,  $\rm O_2$ ,  $\rm CO$ ,  $\rm CO_2$ , and  $\rm CH_4$  less than 100 ppm, which according to an in-house FTIR analysis had 25 ppm of  $\rm H_2O$ , and less than 4 ppm of the sum of hydrocarbons up to  $\rm C_4$ ). Added nitrogen is boil-off from liquid  $\rm N_2$ .

The burning velocity in Bunsen-type flames is known to vary at the tip and base of the flame and is influenced by curvature and stretch; however, these effects are most important over a small portion of the flame. In order to minimize the influence of curvature and stretch on interpretation of the action of the chemical inhibitor, we present the burning velocity of inhibited flames as a normalized parameter: the burning velocity of the inhibited flame divided by the burning velocity of the uninhibited flame at the same flow conditions.

## **Uncertainty Analysis**

The uncertainty analysis consists of calculation of individual uncertainty components and root mean square summation of components [31], and has been described in detail in a previous publication [32]. All uncertainties are reported as expanded uncertainties:  $X \pm U$  (level of confidence approximately 95%), or relative uncertainties:  $U/X \cdot 100\%$ . The primary sources of uncertainty in the average burning velocity measurement are the (1) accuracy of the flow controllers, (2) determination of the flame area, (3) effect of flame base location on flame area, and (4) location of the schlieren image relative to the cold gas boundary. In general, uncertainty increases with burning velocity. A major source of uncertainty arises from selection of the flame location. Traditionally, the burning velocity of the flame is the flow velocity of the unburned reactants; however, since the schlieren image corresponds to a surface with a temperature above that of the unburned gas [29], use of the schlieren image as the flame area can lead to an underestimation of the burning velocity [33], as compared to the area at the cold-gas boundary.

<sup>&</sup>lt;sup>2</sup>Certain commercial equipment, instruments, or materials are identified in this paper to adequately specify the procedure. Such identification does not imply recommendation or endorsement by the National Institute of Standards and Technology, nor does it imply that the materials or equipment are necessarily the best available for the intended use.

Previous researchers [29, 33, 34] have attempted to adjust the flame area to the 298 K isotherm (T<sub>298</sub>), although this generally requires knowledge of the 2-D temperature field in the flame. We do not make such corrections in this paper, and note that it has not yet been done using experimental (rather than calculated) temperature profiles in a flame. Further, some researchers argue for the use of the visible flame area rather than the schlieren area (which would tend to reduce the burning velocities in the present work) [35]. We retain the traditional approach of using the schlieren area, but note that differences on the order of 10% can result from the choice of image to use in the analyses.

Inaccuracies in the flow controllers result in uncertainties of 1.4% for the equivalence ratio, 1.1% for the nitrogen mole fraction, and 1.2% for the hydrogen mole fraction. For stoichiometric mixtures of  $N_2O$  and CO, trace hydrogen (as  $H_2O$ ) is present in the reactant gases at about 13 ppm. Neglecting the uncertainty (unspecified) in the vapor pressure correlation of Gilbert and Sulzmann [36], the uncertainties in the bath temperature, ambient pressure, and carrier gas flow rate yield an Fe(CO)<sub>5</sub> mole fraction uncertainty of 6.5%.

#### NUMERICAL

One-dimensional, freely propagating premixed flames are simulated using the Sandia flame code Premix [37], the Chemkin subroutines [38], and the transport property subroutines [39]. For all of the calculations the absolute tolerance is  $10^{-14}$ , the relative tolerance is  $10^{-9}$ , GRAD is 0.15, and CURV is 0.35. The computational domain is 0 to 50 cm (except when noted). The initial temperature is 298 K and the pressure is one atmosphere. The moist CO-N<sub>2</sub>O flames are modeled using the mechanism and thermodynamic data set of Allen et al. [1], from which the species NCN, NCO, CNO, CH, CH<sub>2</sub>, CH<sub>3</sub>, C<sub>2</sub>H, HCCO, HNCO, HOCN, HCNO, and H<sub>2</sub>CN have been removed (they have been found to be unimportant for the present flames). The resulting reduced mechanism has 20 species and 92 reactions. For all calculations (except where noted) we use the rate for  $CO + OH \leftrightarrow CO_2 + H$  from Yu et al.

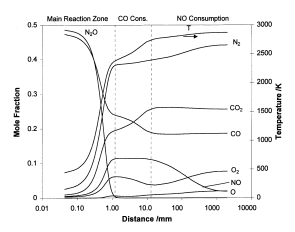


Fig. 1. Calculated major species mole fraction and temperature profiles in a stoichiometric premixed dry CO-N<sub>2</sub>O flame (note log distance scale).

[40], and for CO +  $N_2O \rightarrow CO_2 + N_2$  from Milks and Matula [6] since they provide reasonable agreement with our data. The chemical mechanism for Fe(CO)<sub>5</sub> inhibition of flames (12 species and 55 reactions) and necessary thermodynamic data are compiled from a variety of sources as described in Rumminger et al. [23].

## RESULTS AND DISCUSSION

### **General Features**

Prior to discussing the inhibited flames, we first present findings for the uninhibited flames. Calculations show that the pure CO-N<sub>2</sub>O flame has a very large domain due to the slow reaction of CO and NO downstream of the main heat release region. A plot of the major species profiles and temperature of a stoichiometric CO-N<sub>2</sub>O flame is shown in Fig. 1. There are three distinct regions in the calculated results. In the first, which extends a few millimeters from the start of the temperature rise, there is very fast reaction of CO with N2O, and thermal decomposition of N2O followed by reaction of N<sub>2</sub>O with O atoms. In the second region, [NO] is constant, but CO is consumed through its slow reaction with O and O<sub>2</sub>. In the third region, NO is consumed. Although no practical flame can support such weak reaction over a domain of 2 meters, it is of interest to determine how the features of the calculated results over the

Reactant Conditions			Temperature (K)			Peak Radical Mole Fraction within Flame (ppm)			
φ	Mole % H <sub>2</sub>	Mole % N <sub>2</sub>	$v_{o, num}$ (cm/s)	A.F.T.	At Point of 99.5% N <sub>2</sub> O Consumption	At peak of CO+N <sub>2</sub> O Reaction	0	OH (ppm)	Н
CO-N	2O flame								
1	0	0	24.5	2870	2323	1770	2833	0	0
1	0.012	0	25.9	2872	2377	1773	3524	79	2
1	0.68	0	45.0	2866	2589	1896	4524	1811	38
0.6	0	0	20.1	2789	2303	1693	2690	0	0
1	0	0	24.5	2860	2324	1770	2833	0	0
1.3	0	0	25.7	2867	2319	1789	2710	0	0
3.2	0	0	22.1	2378	2139	1658	1109	0	0
1	0	0	24.5	2860	2323	1770	2833	0	0
1	0	25	16.6	2559	2155	1737	1590	0	0
CH <sub>4</sub> -a	ir flame								
1	_	_	40.0	2230	_	_	3150	7660	6740
CO-ai	r-H <sub>2</sub> flame								
1	1.0	_	35.8	2376	_	_	14,000	5200	2900

TABLE 1

Calculated CO–N<sub>2</sub>O Flame Properties for Various Reactant Streams<sup>a</sup>

entire domain influence the comparisons with experimental results from a Bunsen-type flame, which remains quasi-one-dimensional only for a few millimeters.

Table 1 presents some calculated burning velocities  $v_{o,num}$ , temperatures and peak radical mole fractions for dry, undiluted CO-N2O flames over a range of  $\phi$  and for stoichiometric flames, over a range of added N2 and H2. The adiabatic flame temperature (A.F.T.) is given, as well as the temperature at the location of 99.5% consumption of N<sub>2</sub>O, which roughly corresponds to the end of the region of rapid heat release. The temperature at the location of the peak rate of the CO + N<sub>2</sub>O reaction is also listed. For pure CO-N<sub>2</sub>O flames with  $\phi = 1.0$ and hydrogen addition up to 6800 ppm, or dry flames with  $0.6 \le \phi \le 1.3$  (the first six lines of Table 1), the A.F.T. is 2830 K  $\pm$  40 K, whereas for the richest flames ( $\phi = 3.2$ ) and stoichiometric flames with nitrogen dilution, the A.F.T. is significantly lower, 2378 K and 2559 K respectively. As Table 1 shows, however, the calculated peak temperature at the point of 99.5% N<sub>2</sub>O consumption is 240 K to 550 K lower than the adiabatic flame temperature. For flames rich in CO or with added N<sub>2</sub>, the adiabatic flame

temperature is much lower because of dilution, but the temperature at the point of N<sub>2</sub>O consumption is only slightly lower. For flames with added hydrogen, the temperature in the main reaction zone is closer to the final temperature far downstream, since the faster overall reaction rate allows more of the CO to be consumed before the N<sub>2</sub>O is gone. Similarly, at the peak of the CO +  $N_2$ O reaction, the temperature is about 1800 K; for flames with 6800 ppm of hydrogen it is about 100 K higher, for rich or lean flames it is about 100 K lower, and with 25% dilution nitrogen is about 40 K lower. Hence, the temperature at the peak of the  $CO + N_2O$  reaction is less sensitive to changes in  $\phi$ ,  $X_{N_2}$  or  $X_{H_2}$  than is the adiabatic flame temperature (which is not nearly achieved in the experimental flames). In the present experiment which has a flame thickness of about 1 mm, the calculated temperatures in the main reaction zone, where most of the fuel and oxidizer consumption occur, are more relevant than the adiabatic flame temperatures (which would only be reached nearly 3 m downstream).

The significant but gradual temperature rise far downstream of the main reaction zone is caused by the slow reaction of the remaining

<sup>&</sup>lt;sup>a</sup> Values for stoichiometric CH<sub>4</sub>-air and CO-air-H<sub>2</sub> flames are provided for comparison.

CO with O or  $O_2$ , and from NO consumption. Nonetheless, this additional temperature rise does not have much consequence for the main reaction zone; the thickness is too great to provide a high rate of heat feedback, and the flow field in the experiment does not support such a large, one-dimensional downstream region. Numerical experiments for  $0.6 \le \phi \le 3.2$ show that turning off the CO + O and  $CO + O_2$ reactions, and the even slower NO consumption reactions, provides a burning velocity within 2% of those using the entire domain with all reactions included. Calculations were also performed over the entire range of  $\phi$  and  $X_H$ ; for each condition, domains of 2 mm, 3 mm, or 3000 mm all yield calculated burning velocities within 1% of each other, providing further evidence that the slow reactions in the very large domain do not affect the measured or calculated burning velocities of smaller, practical flames.

Unlike CO or CH<sub>4</sub> flames, these CO-N<sub>2</sub>O flames do not demonstrate the characteristic radical super-equilibrium. The O, H, and OH mole fractions, X<sub>O</sub>, X<sub>H</sub>, and X<sub>OH</sub>, rise monotonically throughout the computational domain up to the point where the N<sub>2</sub>O is 99.5% gone. The peak in-flame X<sub>O</sub> is about the same as in stoichiometric CH<sub>4</sub>-air flames, and about a factor of 4 lower than in CO-air flames with 1% H<sub>2</sub>. Even with about 0.7% of added H<sub>2</sub>, the peak in-flame X<sub>H</sub> in the CO-N<sub>2</sub>O flames is still more than two orders of magnitude lower than that for the CH<sub>4</sub> or CO flames, clearly demonstrating the straight-chain character of these flames, and suggesting their utility for testing the effectiveness of chemical inhibitors in nonbranching systems.

## **Burning Velocity of Uninhibited Flames**

Figure 2 presents the burning velocity of the premixed CO– $N_2$ O flames from the experiments (symbols) as a function of the hydrogen concentration in the reactants. The "dry" burning velocity is measured to be 23.1 cm/s  $\pm$  0.6 cm/s, and this value increases to about twice that value with 0.5% of hydrogen. The marginal effect of added  $H_2$  is smaller at higher values of the hydrogen mole fraction. For the reader's

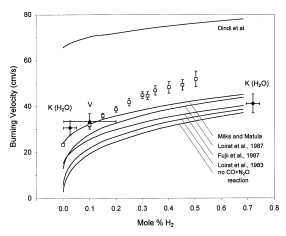


Fig. 2. Burning velocity of stoichiometric CO– $N_2O$  flames as a function of hydrogen mole percent. Points (squares) are experimental data, and the solid lines are the modeling results using the rate of the CO +  $N_2O$  from the reference indicated in the figure. The data of Kalff and Alkemade [13] with  $H_2O$  (not  $H_2$ ) are indicated by "K( $H_2O$ )" and diamonds, and the datum of Van Wonterghem and Van Tiggelen [12] by "V" and a triangle.

convenience, Table 2 contains the experimental data and uncertainties.

The figure also shows the calculated burning velocity (lines) for various values of the specific reaction rate constant for the reaction CO +  $N_2O \rightarrow CO_2 + N_2$ . The bottom curve does not

TABLE 2

Measured Burning Velocity of CO-N<sub>2</sub>O Flames.<sup>a</sup>

φ	<i>v<sub>o, exp</sub></i> (cm/s)	Mole % $H_2$ ( $\phi = 1.0$ )	<i>v<sub>o, exp</sub></i> (cm/s)	Mole % $N_2$ ( $\phi = 1.0$ )	<i>v<sub>o, exp</sub></i> (cm/s)
0.60	16.3 ± 0.4	0.00	$23.4 \pm 0.6$	0.0	$23.7 \pm 0.6$
0.75	$19.9 \pm 0.5$	0.10	$31.9 \pm 0.8$	2.5	$22.2 \pm 0.6$
0.80	$20.8\pm0.5$	0.15	$35.9\pm1.1$	5.3	$22.8 \pm 0.6$
0.85	$21.6\pm0.5$	0.20	$38.7 \pm 1.2$	8.1	$22.2 \pm 0.6$
0.90	$22.2\pm0.5$	0.25	$41.8\pm1.6$	11.1	$21.7 \pm 0.6$
0.95	$22.7\pm0.6$	0.30	$44.8\pm1.5$	14.3	$20.9 \pm 0.6$
1.0	$23.4\pm0.6$	0.32	$44.6 \pm 1.7$	17.7	$20.2 \pm 0.6$
1.1	$24.4\pm0.6$	0.35	$46.9 \pm 2.0$	21.3	$19.3 \pm 0.5$
1.2	$25.5\pm0.6$	0.40	$48.2\pm2.5$	25.1	$17.9 \pm 0.6$
1.3	$26.3\pm0.7$	0.45	$49.3 \pm 2.3$		
1.5	$27.6\pm0.7$	0.50	$51.8 \pm 3.4$		
1.8	$28.4\pm0.7$				
2.1	$28.8 \pm 0.7$				
2.5	$28.4 \pm 0.7$				
3.0	$27.3\pm0.7$				

 $<sup>^</sup>a$  Data are presented for dry flames at varying equivalence ratio  $\phi$  and for stoichiometric flames with varying %  $H_2$  and %  $N_2.$ 

include the direct reaction in the mechanism, and indicates a burning velocity of 2.9 cm/s with no added hydrogen, clearly illustrating that this reaction is required to reproduce our data. The remaining curves in the figure show the calculated result using the rate expressions of Dindi et al. [9], Milks and Matula [6], Loirat et al. [41], Fujii et al. [7], and Loirat et al. [8]. The value recommended by Dindi et al. overpredicts the burning velocity, with too weak of a dependence on X<sub>H.</sub>, while the other expressions underpredict it for all values of X<sub>H</sub>. The expression of Milks and Matula [6] provides a burning velocity which is very close to the present data for low values of  $X_{H_2}$ , but which deviates for  $X_{H_2} \ge$ 0.2%. Also shown in the figure are the data of Van Wonterghem and Van Tiggelen [12] (< 2000 ppm of H<sub>2</sub>), and the data of Kalff and Alkemade [13] (<500 ppm H<sub>2</sub>O). The present data are in good agreement with the low-hydrogen data of both previous investigations, and provide the additional advantage of a very low background level of hydrogen with a controlled level of added hydrogen. Note that although the data of Kalff and Alkemade are presented on the same figure, the experiments contained H<sub>2</sub>O rather than H<sub>2</sub>. Based on calculations with either added H<sub>2</sub> or H<sub>2</sub>O, we estimate that H<sub>2</sub> addition, as compared to H<sub>2</sub>O addition, lowers the burning velocity about 10% at 250 ppm and raises it about 10% at 7200 ppm. Thus we find that the data of Kalff and Alkemade are in excellent agreement with the present data at low hydrogen content, and approximately within the uncertainty of both experiments if our data are extrapolated to their conditions at 7200 ppm  $H_2O$ .

Because the CO + OH reaction becomes increasingly important as the hydrogen content increases, the rate of the CO + OH reaction has a large influence on the ability of the model to predict the experimental data at higher X<sub>H<sub>2</sub></sub>. Figure 3 presents the calculated burning velocity of the CO–N<sub>2</sub>O flames with the direct reaction rate of Milks and Matula [6], but using the CO + OH rate expression of Baulch et al. 1973 [42], Yu et al. [40], Baulch et al. 1992 [43], and Wooldridge et al. [44]. The recommendation of Baulch et al. 1992 [43] significantly overpredicts the burning velocity for our data, while that of Baulch et al. 1973 is slightly lower. The rates of

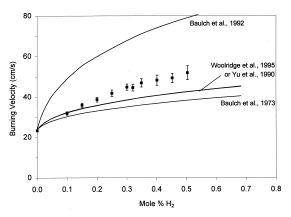


Fig. 3. Calculated burning velocity (lines) of  $CO-N_2O$  flames with values of the  $CO + OH \rightarrow CO_2 + H$  rate from Baulch et al. [42], Yu et al., [40], Wooldridge et al. [44], and Baulch et al. [43], together with data from the present study (symbols).

Yu et al. and Wooldridge et al. provide results that are nearly identical for the present conditions, and that are very close to the measured burning velocities. Although this rate is important for predicting the dependence of the burning velocity on the added hydrogen, it is not important for predicting the burning velocity of the driest flames here (less than 15 ppm  $H_2O$ ). For the remainder of the calculations in the present paper, we retain the rate of Yu et al. for CO + OH.

The suggested rate expressions for the direct reaction include those with a high activation energy (e.g. [5, 8, 41]) and those with a low value (e.g. [6, 7, 45]). An Arrhenius plot for some of the literature values of the direct reaction is shown in Fig. 4. Since the flame temperature in the main reaction zone of the flames varies by about 260 K, the different activation energies can affect the shape of the burning velocity curves in Fig. 2 (comparing for example the curves for Loirat et al. [41] and Fujii [7], which are fairly close at  $X_{H_2} = 0$ ). In order to obtain some temperature variation while avoiding the complications from uncertainty in the CO + OH rate, we examine dry  $(5-15 \text{ ppm H}_2)$  flames with inert additive or over a range of  $\phi$ .

Figure 5 presents the burning velocity of the dry flames with added nitrogen up to about 25% together with calculations using the rate expressions of Loirat et al. [41] and Milks and Matula

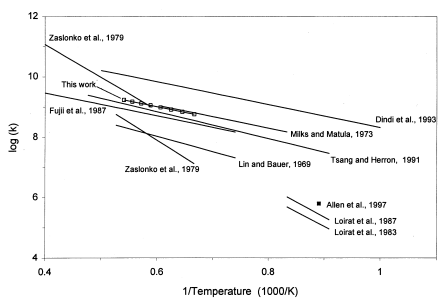


Fig. 4. Arrhenius plots of the rate of the direct exchange reaction  $CO + N_2O \rightarrow CO_2 + N_2$  from various investigators (k in cm, K, mole, s).

[6]. Also shown in the figure are the predictions when the preexponential factors of the direct reaction rates have been adjusted to provide agreement at  $X_{N_2} = 0$  (dotted lines). The higher activation energy expression appears to yield too low of a burning velocity with added nitrogen, whereas the lower activation energy expres-

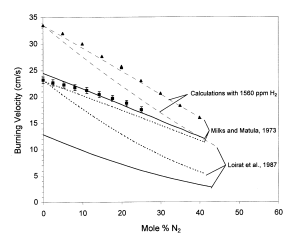


Fig. 5. Measured burning velocity (squares) and calculated results (lines) of CO–N<sub>2</sub>O flames with added N<sub>2</sub>. The solid lines correspond to the as given CO + N<sub>2</sub>O rate of Milks and Matula [6] or Loirat et al., 1987 [41]; the dotted lines have the preexponential adjusted for agreement at  $X_{\rm N_2}=0$ . Triangles are data from Ref. 12 and dashed line has 1560 ppm  $\rm H_2$  and adjusted preexponential factor.

sion provides a closer temperature dependence. Van Wonterghem and Van Tiggelen [12] also presented data on the flame speed as a function of  $X_{N_a}$ . Although the background hydrogen levels in their experiment are somewhat high (and not measured), their data can still be used to test the present mechanism. (Even with 1000 ppm of H<sub>2</sub>, half the CO consumption occurs via the direct reaction, and the burning velocity is very sensitive to the rate of the direct reaction.) We estimate the background level of H<sub>2</sub> in their reactant mixtures to be that value of X<sub>H<sub>2</sub></sub> (1560 ppm) which provides a calculated burning velocity which matches their experimental burning velocity for  $\phi = 1.0$  and  $X_{N_2} = 0$ . (In the calculation, we use the  $CO + N_2^TO$  rate of Milks and Matula with a 10% lower preexponential, which gives the best fit to our data.) We also show the calculated result using the rate of Loirat et al., again matched for agreement at  $X_{N_0} = 0$ . The triangles and the dashed lines in Fig. 5 show the data of Van Wonterghem and Van Tiggelen and our calculations. The lower activation energy again provides a closer agreement.

The dependence of the burning velocity on the fuel-oxidizer equivalence ratio  $\phi$  is shown in Fig. 6. The experimental results (symbols) for

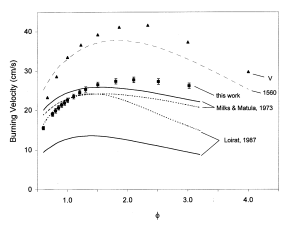


Fig. 6. Measured burning velocity (squares) and calculated results (lines) of CO–N<sub>2</sub>O flames as a function of fuel-oxidizer equivalence ratio  $\phi$ . The solid lines correspond to the CO + N<sub>2</sub>O rate of Milks and Matula [6] or Loirat et al. [41] as given; the dotted lines have the preexponential adjusted for agreement at  $\phi = 1.0$ . Triangles are data from Ref. [12] and dashed line has 1560 ppm H<sub>2</sub> and adjusted preexponential factor.

dry flames are presented along with the calculated results (solid lines), again using the direct reaction rate expressions of Loirat et al. [41] and Milks and Matula [6]. As shown previously in Fig. 2, the expression of Loirat et al. [41] (along with most of the others) provides a burning velocity significantly lower than the present data for dry flames. When the preexponential factor in the direct exchange reaction rate is adjusted to give agreement at  $\phi = 1.0$ (dotted lines in the figures), the high activation energy rate of Loirat et al. underpredicts the burning velocity for richer flames (which have lower temperature) more than does the lowactivation energy rate of Milks and Matula (note from Table 1 that at  $\phi = 3.2$ , the peak flame temperature within the primary reaction zone drops by about 200 K compared to its peak at  $\phi = 1.0$ ). Nonetheless, the rate expression of Milks and Matula overpredicts the burning velocity by about 18% at  $\phi = 0.6$ , and underpredicts it by about 25% at  $\phi = 3.0$ . The data of Van Wonterghem and Van Tiggelen [12] for flames with  $X_{H_2} \le 2000$  ppm are also shown in Fig. 6. The dashed line shows the calculated burning velocity for 1560 ppm (as described above). As with our data, the calculations slightly overpredict the flame speed for lean flames and underpredict it for rich flames.

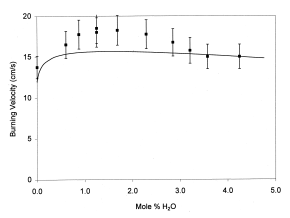


Fig. 7. Experimental burning velocity of CO– $N_2O$  flames as a function of the mole percent  $H_2O$  in the reactants, from [14] for  $\phi=2.0$  and  $X_{N_2}=0.25$ , together with numerically calculated prediction.

For the given mechanism, over the range of  $\phi$  in Fig. 6, the burning velocity is only significantly sensitive to the rate of the exchange reaction (as described below). We have attempted to modify the rate of the next most sensitive reactions (including the rates of the reactions: N<sub>2</sub>O + M, N<sub>2</sub>O + O, and CO + O) to improve agreement over  $\phi$ ; however, adjustment of these rates within their uncertainty does not improve agreement in Fig. 6. Also, addition of H<sub>2</sub> impurity at 100 ppm, while raising the burning velocity 10% to 15% at all values of  $\phi$ , does not affect the qualitative variation with  $\phi$ .

Finally, Fig. 7 presents the experimental burning velocity data of Simpson and Linnett [14] for rich ( $\phi$  = 2.0), slightly preheated ( $T_{in}$  = 323 K) CO–N<sub>2</sub>O flames with 25% dilution nitrogen and added water vapor up to about 5% (error bars estimated here based on Ref. 29). The calculated results for the same initial conditions are shown by the solid line. The agreement is reasonable considering possible differences in the straight tube/flame angle method of determining the burning velocity relative to the nozzle burner/total area method used here.

The numerical modeling of our experimental flame requires the use of the direct reaction for prediction of burning velocity, and the flame results of earlier researchers are also reasonably modeled using the rates suggested in the present work. The kinetic mechanism of Allen et al. [1] provides reasonable agreement with the flame data, while use of the CO + OH rate

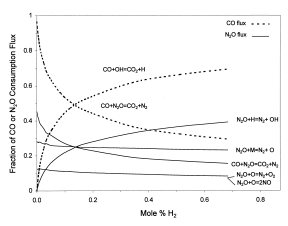


Fig. 8. Calculated flux of important CO (dotted lines) and  $N_2O$  reactions (solid lines) in a stoichiometric  $CO-N_2O$  flame as a function of mole percent of hydrogen.

of Yu et al. [40] or Wooldridge et al. [44] and the CO + N<sub>2</sub>O rate of Milks and Matula [6] provide improved agreement. Lowering the preexponential factor of the CO + N<sub>2</sub>O reaction rate constant of Milks and Matula by about 10% provides the best agreement with our data for dry conditions. Using this rate, the temperature dependence is reasonably modeled, the burning velocity of lean flames ( $\phi = 0.5$ ) is overpredicted by about 18%, and that of rich flames  $(\phi = 3.0)$  is underpredicted by about 25%. The flame results imply a rate of 10<sup>9.2</sup> cm<sup>3</sup>/mol s at 1800 K for the direct reaction, with an activation energy near 71 kJ/mol. Because the direct reaction dominates the reaction system even more so under rich conditions (as described below), it is perhaps more reasonable to infer the direct reaction rate from the burning velocity at  $\phi =$ 3.0. Doing this implies a reaction rate of  $10^{9.3}$ cm<sup>3</sup>/mol s at 1800 K, or about 30% higher than the Milks and Matula rate.

The important reactions in the CO–N<sub>2</sub>O flames with and without added hydrogen can now be examined using the kinetic mechanism described above. These calculations use the recommendation of Milks and Matula [6] for the direct reaction of CO with N<sub>2</sub>O. In Fig. 8, the fractional flux of CO (dotted lines) is shown for each important reaction as a function of the mole percent hydrogen. The fractional consumption flux is the fraction of the total consumption of a species occurring due to a given reaction. The total flux for a species is deter-

mined by integrating each consumption reaction over the domain of interest (the cold boundary to the point of 99.5% N<sub>2</sub>O consumption), and summing the contribution from each reaction. The fractional flux is the flux for a specific reaction divided by the total flux. As indicated, the direct exchange reaction accounts for about 96% of CO consumption for the dry condition, decreasing rapidly to about 50% at 1000 ppm, with a more gradual decrease for greater hydrogen concentrations. Correspondingly, the fraction of CO consumption due to reaction with OH increases. For the reasonably small domain of these calculations (about 2 mm), reactions of CO with O or  $O_2$  (not shown) are at most a few percent of the total consumption flux for CO. The reaction of N<sub>2</sub>O is more varied, and the fractional consumption flux of N<sub>2</sub>O is shown by the solid lines in Fig. 8. Thermal decomposition accounts for about 30%, and is weakly dependent on the hydrogen concentration. Likewise, reactions of N<sub>2</sub>O with O atom to form 2NO or O<sub>2</sub> and N<sub>2</sub> account for about 13% each, whereas reactions with NO (not shown) account for a few percent; all of these reactions vary little with X<sub>H<sub>2</sub></sub>. The contribution of the reaction of N<sub>2</sub>O with hydrogen radical (produced from the  $CO + OH \rightarrow CO_2 + H$  reaction) increases rapidly as  $X_{H_2}$  approaches 1000 ppm and more slowly above that value, with the reaction of N<sub>2</sub>O with CO again decreasing correspondingly. A main feature of these flames is that while hydrogen has a large effect on the reaction mechanism, the direct reaction still accounts for half of the CO consumption and a quarter of the  $N_2O$  consumption at  $X_{H_2} = 1000$  ppm.

In Fig. 9, a similar plot is shown for CO and  $N_2O$  consumption as a function of  $\phi$ . In these dry flames (5 to 15 ppm  $H_2O$ ) the direct reaction of CO and  $N_2O$  is always responsible for about 96% of the CO consumption; however, for  $N_2O$  consumption, its contribution varies from about 33% for lean flames to 80% for the rich flames. Similarly, the contributions of other reactions for  $N_2O$  consumption decrease when CO is present in abundance.

## Effect of Fe(CO)<sub>5</sub> on CO-N<sub>2</sub>O Flames

In order to study the action of chemical inhibitors in systems without radical chain branching,

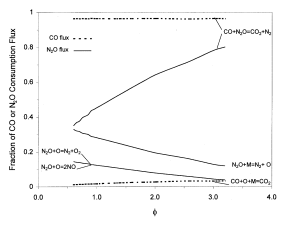


Fig. 9. Calculated flux of important CO (dotted lines) and  $N_2O$  reactions (solid lines) in a dry, stoichiometric CO– $N_2O$  flame as a function of  $\phi$ .

previous researchers have employed nitrogen oxides as the oxidizers. In studies of H<sub>2</sub>-N<sub>2</sub>-N<sub>2</sub>O flames inhibited by CF<sub>3</sub>Br and HBr, Dixon-Lewis et al. [46] found the nitrous oxide flames to be inhibited much more weakly than hydrogen-air flames. Similarly, Rosser et al. [24] found that neither CH<sub>4</sub>-NO<sub>2</sub> nor C<sub>2</sub>H<sub>4</sub>-NO<sub>2</sub> flames were significantly inhibited by HBr. They further postulated that any flames in which NO<sub>2</sub> is present in significant quantities will not be inhibited by HBr. In studies of moist CO-O<sub>2</sub> and CO-N<sub>2</sub>O flames with chlorine, Palmer and Seery [47] predicted that after initial inhibition, further addition of Cl2 would accelerate the N<sub>2</sub>O decomposition, and that CO would then react with ClO; however, they provided no burning velocity data for such flames, and stated that the flames are still ultimately inhibited by chlorine. To further understand the effect of Fe(CO)<sub>5</sub> on radical recombination, particularly in nonbranching systems, it is of interest to study its effect in dry and moist CO-N<sub>2</sub>O flames. Since Fe(CO)<sub>5</sub> is about 200 times more effective as a flame inhibitor than is Cl, it is of interest to determine if the region of inhibition that Palmer and Seery suggested would occur for low Cl2 mole fraction, as well as the ultimate inhibition that they described, may be modified for Fe(CO)<sub>5</sub>.

Figure 10 shows the experimental data (symbols) for the burning velocity of  $N_2O$  flames with added  $Fe(CO)_5$ , normalized by the burning velocity with no inhibitor. Data are provided for

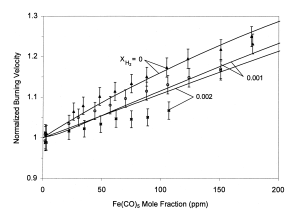


Fig. 10. Normalized burning velocity of stoichiometric CO– $N_2O$  flames with  $X_{H_2}=0.0,\,0.001$ , and 0.002 for increasing quantities of Fe(CO)<sub>5</sub>. The symbols are the experimental data, the lines are the calculated results.

added  $H_2$  mole fractions  $X_{H_2}$  of 0, 0.001, and 0.002 (note that the background water is 15 ppm). Unlike flames of  $CH_4$ -air, and CO-air which are inhibited by  $Fe(CO)_5$ , the overall reaction rates of these  $CO-N_2O$  flames are increased. As the figure shows, for hydrogenfree flames with 171 ppm of  $Fe(CO)_5$ , the burning velocity is increased by 25%; as the amount of hydrogen is increased, the rate of burning velocity increase is smaller. In order to understand the reasons for the promotion of the reactions, and also the reduction in this promotion, the flames were modeled as described below.

A mechanism for Fe(CO)<sub>5</sub>-inhibition of hydrocarbon and carbon monoxide flames [23, 25] indicates that for those systems, the inhibition occurs due to gas-phase catalytic cycles in which H- or O-atoms are recombined by iron oxide and hydroxide species. Using this mechanism, calculations for the present flames do not show promotion. Recent research in atmospheric chemistry, however, suggests that interactions between Fe-species, N<sub>2</sub>O, and CO may be important. West and Broida [48] observed that nitrous oxide destruction is catalyzed by Fe, producing chemiluminescent FeO. Rates for the reaction Fe +  $N_2O \rightarrow FeO + N_2$  have been measured by Campbell and Metzger [49] and Plane and Rollason [50]. This reaction alone does not provide promotion; however, with subsequent reaction of FeO and CO, it does. Kappes and Staley [51] proposed a catalytic

TABLE 3

Reactions in the Iron Catalytic Cycle for the CO-N<sub>2</sub>O

System, and Their Estimated Reaction Rates\*

Reaction	A	b	Ea/R
$\overline{Fe + N_2O = FeO + N_2}$	1.40 E+14	0	5940
$FeO + N_2O = FeO_2 + N_2$	3.00 E+13	0	5033
$FeOH + N_2O = FeOOH + N_2$	1.30 E+14	0	4530
$FeO + CO = Fe + CO_2$	1.80 E + 12	0	3522
$FeO_2 + CO = FeO + CO_2$	1.18 E+13	0	4530
$FeOOH + CO = FeOH + CO_2$	6.00 E+13	0	4026

<sup>\*</sup>  $k_f = A T^b \exp(-E_a/RT)$ , and units are cm, K, mole, s.

oxidation cycle involving ions: Fe $^+$  + N<sub>2</sub>O  $\rightarrow$  FeO $^+$  + N<sub>2</sub> and FeO $^+$  + CO  $\rightarrow$  Fe $^+$  + CO<sub>2</sub>, but reaction of neutral FeO with CO is also possible, and would provide a gas-phase catalytic cycle, with the net reaction: CO + N<sub>2</sub>O  $\rightarrow$  CO<sub>2</sub> + N<sub>2</sub>

Previous experimental work suggests such catalytic effects. In a shock tube study, Matsuda [52] reports that addition of Fe(CO)<sub>5</sub> to CO- $O_2$ -Ar mixtures promotes the oxidation of CO. Also, in a fluidized-bed study of the interactions of compounds of iron with those of nitrogen, Hayhurst and Lawrence [53] argue that iron acts as a catalyst to convert N2O to N2 and CO to CO<sub>2</sub> (but this appears to be a heterogeneous effect). In addition to the cycle above with FeO, estimates of the heats of reaction indicate that cycles with FeO<sub>2</sub> and FeOOH are possible. The rates for the reactions in the iron-catalyzed CO-N<sub>2</sub>O system are listed in Table 3. The rate of the first reaction in the table is from Plane and Rollason [50], while the others are our estimates. These reactions are added to the iron-species inhibition mechanism described in Ref. 23. The thermodynamic data for FeOOH (and the related rate expressions) have been updated based on the recent calculations of Kellogg and Irikura [54]. Calculations of the flame structure using this mechanism have been performed for the N<sub>2</sub>O-H<sub>2</sub>-Fe(CO)<sub>5</sub> flames and are described below.

Figure 10 also shows the calculated burning velocity (normalized by the calculated burning velocity in the absence of iron pentacarbonyl) as a function of the initial  $Fe(CO)_5$  mole fraction  $X_{Fe(CO)_5}$  for added hydrogen mole fractions of 0, 0.1%, and 0.2%. The calculated results show the correct qualitative behavior for  $Fe(CO)_5$ 

addition to the dry  $N_2O$  flames. For the two cases with added hydrogen, a slight difference in the promotion effect is predicted for  $X_{\rm H_2} = 0.001$  or 0.002, but the variation is greater in the experiments. The calculated burning velocity in the moist flames with Fe(CO)<sub>5</sub> is very sensitive to the rates of the catalytic steps involving FeOOH. While changes in these rates do affect the separation of the curves for  $X_{\rm H_2} = 0.001$  and 0.002, they cannot provide agreement between experiment and calculation, and further research is required.

Careful investigation of Fig. 10 reveals that for all values of X<sub>H</sub>, the slope of the experimental curve decreases as  $X_{Fe(CO)_5}$  increases, but that the calculated curve is linear. The curvature is mild, and ordinarily might not be of interest given the other discrepancies between the measured and predicted flame speeds with added Fe(CO)<sub>5</sub>. This reduction in the marginal influence of the Fe(CO)<sub>5</sub> (reaction promotion) as  $X_{Fe(CO)_s}$  increases, however, is much like the loss of effectiveness of Fe(CO)<sub>5</sub> observed in flames of CH4 or CO with air, in which the marginal inhibition is reduced at higher Fe(CO)<sub>5</sub> mole fractions. In previous work, the loss of inhibition effectiveness was argued to be due to the formation of condensed-phase particulates, which serve as a sink for the active gas-phase iron-containing inhibiting species [21]. Through the use of classical laser scattering and extinction measurements with phasesensitive detection, it was later shown that a large increase in the particle scattering signal was well correlated with the point where the inhibitor lost its marginal effect [55, 56]. It is possible that the reduction in the promotion effect in the present flames is also due to loss of the active iron intermediates through condensation. In order to test this hypothesis, we performed laser scattering measurements on premixed CO-N<sub>2</sub>O flames with 0% and 0.2% added hydrogen and for 0 and 180 ppm of added Fe(CO)<sub>5</sub> using the apparatus described in detail in Refs. 55 and 56. The experiments show that with addition of iron pentacarbonyl, there was no significant particle scattering signal in the main reaction region for either level of H<sub>2</sub> in the reactants, unlike the hydrocarbon flames in which a significant scattering signal was detected that clearly increased as the mole fraction of Fe(CO)<sub>5</sub> increased. Hence, we cannot conclude that the reduction in the promotion effect near  $X_{Fe(CO)_5} = 60$  ppm for  $X_{H_2} = 0.002$  is due to formation of particulates in the flame.

Although agreement between the measured and calculated normalized burning velocity in Fig. 10 is not perfect, it is still of interest to investigate the numerical results to determine the reasons for the promotion of the reaction (and the lower promotion when hydrogen is present). Reaction flux and sensitivity analyses are used to provide insight. For each important species in the mechanism, Table 4 shows the fractional flux through the reactions contributing more than 1% to their consumption or production; Table 5 shows the first-order sensitivity coefficient of the burning velocity with respect to the reaction rate constant  $[d(\ln v)]$  $d(\ln k) / d(\ln v)/d(\ln k)|_{\max}$ . In both of these tables, calculated results are provided for  $X_{H_2}$  = 0 and 0.002, and for added Fe(CO)<sub>5</sub> of 0 ppm and 213 ppm.

## **Dry Flames**

The properties of the dry flames without Fe(CO)<sub>5</sub> are as described above. As Table 4 shows, 94% of the CO is consumed by the direct reaction, while N<sub>2</sub>O consumption is roughly equally portioned between thermal decomposition, the direct reaction, and reaction with O atoms. In Table 5, the burning velocity of the iron-free dry flames is most sensitive to the rate of the direct reaction, followed by the decomposition of N<sub>2</sub>O, and to a lesser extent its reaction with O. Upon addition of iron species, 32% of the CO is consumed through reaction with FeO or FeO<sub>2</sub>, which increases the burning velocity. Likewise, 18% of the N<sub>2</sub>O is consumed through the related reactions of Fe or FeO with N<sub>2</sub>O, reducing the N<sub>2</sub>O consumption through both the direct reaction and reaction with Oatoms. While reaction of N<sub>2</sub>O through the catalytic route with iron species or through the direct reaction with CO proceeds at roughly the same rate, reaction of N<sub>2</sub>O with O-atoms is less exothermic than the direct reaction (because of the slow consumption of NO as shown in Fig. 1), so reducing the importance of the O-atom reactions with N<sub>2</sub>O increases the burning velocity. Nonetheless, the effect of iron as a catalytic agent to reduce radical mole fractions is secondary in these flames.

As described in Ref. 25, Fe(CO)<sub>5</sub> in moist CO-O2-N2 flames acts as a catalyst to recombine O atoms (as opposed to methane-air flames, where H-atom recombination appears to be the important inhibition reaction). These dry N<sub>2</sub>O flames appear to be similar to the CO-H<sub>2</sub>-O<sub>2</sub>-N<sub>2</sub> flames in that iron species do serve to recombine O atoms. For example, with 213 ppm of Fe(CO)<sub>5</sub>, about 23% of the O-atom consumption occurs through reactions with iron intermediates. In this flame, however, a reduction in O-atom mole fraction leads to a slightly higher overall reaction rate. Very little of the CO consumption occurs via reaction with O atoms, but a reduced mole fraction of O reduces its reaction with N<sub>2</sub>O. Nonetheless, the effect of O-atom recombination is minor, and the main influence of adding iron pentacarbonyl to the dry flames is to promote the N2O reaction through the iron-catalyzed reaction sequence described above.

## **Moist Flames**

The moist N<sub>2</sub>O flames without Fe(CO)<sub>5</sub> are described above and in Tables 4 and 5. Carbon monoxide is consumed by reaction with OH (60%) and N<sub>2</sub>O (38%), and N<sub>2</sub>O is consumed roughly equally by reaction with CO, H atom, O atom, and by thermal decomposition. With addition of iron species, the catalytic route described above again contributes to CO and N<sub>2</sub>O consumption, but it is only about two-thirds as important as in the dry case (primarily because of the dominant role of the CO + OH and  $N_2O + H$  reactions). The iron species do enter into catalytic recombination reactions for H and O atoms, but these cycles are not significant. For H and OH, less than 1% of the flux of each involves iron-species reactions. For O atom, although iron-species reactions account for about 8% of its consumption, the sensitivities (not shown in Table 5) indicate that consumption of O atoms by reaction with iron species increases the burning velocity as described above. (For reference, Fe reactions in CO flames with 1% hydrogen cause 30% and 70% of the H- and O-atom destruction at about 200

			Fractional Flux (%)			
		$X_{H_2}$ :	0.	000	0.	002
Species	Reaction	$X_{Fe(CO)_5}$ (ppm):	0	213	0	213
СО	Destruction					
	CO + OH ←	$\rightarrow$ CO <sub>2</sub> + H	_	_	60	48
	$CO + N_2O$	$\leftrightarrow$ CO <sub>2</sub> + N <sub>2</sub>	94	68	38	30
		$M) \leftrightarrow CO_2 (+M)$	3	0	1	0
	,	$\leftrightarrow$ CO <sub>2</sub> + NO	2	_	_	_
	_	$\leftrightarrow$ Fe + CO <sub>2</sub>	_	9	_	3
		$\leftrightarrow$ FeO + CO <sub>2</sub>	_	23	_	6
	_	$CO \leftrightarrow FeOH + CO_2$	_	_	_	12
N <sub>2</sub> O	Destruction	30 11 10011 1 002				12
1120		$\leftrightarrow$ CO <sub>2</sub> + N <sub>2</sub>	40	30	19	16
		$\Rightarrow N_2 + O(+M)$	31	30	26	26
			31	30	28	20
	$N_2O + H \Leftrightarrow$	<del>-</del>	13	10	11	
	$N_2O + O \Leftrightarrow$					10
	$N_2O + O \Leftrightarrow$		13	10	11	10
	NH + NO «	=	0	0	3	3
	_	$\leftrightarrow$ NO <sub>2</sub> + N <sub>2</sub>	2	1	1	1
	_	$\leftrightarrow$ FeO <sub>2</sub> + N <sub>2</sub>	_	12	_	5
	$Fe + N_2O \leftarrow$	-	_	6	_	4
	_	$O \Leftrightarrow FeOOH + N_2$	_	_	_	6
O	Creation					
	$N_2O(M+) \leftarrow$	$\rightarrow$ N <sub>2</sub> + O(+M)	99	99	95	88
	$H + O_2 \leftrightarrow O_2$	O + OH	_	_	4	11
	Destruction					
	$N_2O + O \Leftrightarrow$	$O_2 + N_2$	44	36	42	36
	$N_2O + O \Leftrightarrow$	· 2NO	44	36	42	36
	$H + O_2 \leftrightarrow O_2$	O + OH	_	_	6	2
	$NH + O \leftrightarrow NO + H$		_	_	4	4
	$CO + O(+M) \Leftrightarrow CO_2(+M)$		4	2	2	2
	$NO2 + O \Leftrightarrow O_2 + NO$		5	3	1	1
	$NO + O(+M) \Leftrightarrow NO_2(+M)$		2	_	_	_
	$O + H_2 \leftrightarrow I$		_	_	1	1
	$Fe + O_2 \Leftrightarrow FeO + O$		_	12	_	9
	$FeO_2 + O \Leftrightarrow FeO + O_2$		_	11	_	7
Н	Creation	- 2				
	CO + OH ←	$\rightarrow$ CO <sub>2</sub> + H	_	_	87	89
	$H + O_2 \leftrightarrow O_2$	-	_	_	4	2
			_	_	3	3
	$NH + O \leftrightarrow NO + H$				3	3
	$ NH + NO \leftrightarrow N_2O + H  O + H_2 \leftrightarrow H + OH $		_	_	1	1
	Destruction Destruction	1 + 011			1	1
	$N_2O + H \leftrightarrow$	N + OH			83	72
			<del></del>	_	10	73
	NH + NO «		_	_		10
	$H + O_2 \leftrightarrow O_2$		_	_	3	12
OH	NO2 + H ←	NO + OH	_	_	3	3
OH	Creation	N + OH			07	
	$N_2O + H \leftrightarrow$	-	_	_	87	77
	$H + O_2 \leftrightarrow O + OH$		_	_	3	13
	NH + NO «		_	_	3	3
	$NO_2 + H \Leftrightarrow$		_	_	3	3
	$H_2O + O \Leftrightarrow$		_	_	1	2
	$O + H_2 \leftrightarrow I$	H + OH	_	_	1	1

-	ΓABL	E	4
(	Contin	u	ed

			Fractional Flux (%)				
		$X_{H_2}$ :	0.000		0.002		
Species	Reaction	$X_{Fe(CO)_5}$ (ppm):	0	213	0	213	
ОН	Destruction						
	CO + OH «	$\rightarrow$ CO <sub>2</sub> + H	_	_	93	95	
	$H + O_2 \leftrightarrow$	O + OH	_	_	5	2	
	H <sub>2</sub> O + O ←	→ 2OH	_	_	1	_	
Fe	Creation						
	FeO + CO	$\leftrightarrow$ Fe + CO <sub>2</sub>	_	53	_	35	
	Fe + $O_2 \leftrightarrow$	FeO + O	_	45	_	54	
	$Fe + O_2(+M) \leftrightarrow FeO_2(+M)$		_	_	_	6	
	$FeO + H \leftrightarrow Fe + OH$		_	_	_	3	
	Destruction						
	$Fe + N_2O \leftarrow$	$\rightarrow$ FeO + N <sub>2</sub>	_	85	_	88	
	Fe + $O_2(+1)$	$M) \leftrightarrow FeO_2(+M)$	_	14	_	11	
FeO	Creation						
	$Fe + N_2O \Leftrightarrow FeO + N_2$		_	33	_	42	
	$FeO_2 + O \Leftrightarrow FeO + O_2$		_	16	_	20	
	$FeO_2 + CO \Leftrightarrow FeO + CO_2$		_	51	_	35	
	Destruction						
	$FeO + CO \Leftrightarrow Fe + CO_2$		_	21	_	17	
	$FeO + N_2O \Leftrightarrow FeO_2 + N_2$			62		53	
	$Fe + O_2 \Leftrightarrow FeO + O$		_	18	_	26	
	$FeO + H \leftrightarrow Fe + OH$		_	_	_	2	
	$FeO + H_2O \Leftrightarrow Fe(OH)_2$		_	_	_	1	

<sup>&</sup>lt;sup>a</sup> Results are given for  $X_{H_2} = 0.0$  and 0.002, and for  $X_{Fe(CO)_5} = 0$  and 213 ppm.

ppm of Fe(CO)<sub>5</sub> [25], and about a 30% reduction in the flame speed.)

In the present moist CO–N<sub>2</sub>O flames, oxygen atom is not a significant species for CO consumption or in chain reactions. Hydrogen and hydroxyl radicals are important for consumption of N<sub>2</sub>O and CO, but the system is straightchain rather than chain-branching, and neither radical reaches very high concentrations (their estimated peak mole fractions in CH<sub>4</sub>, O<sub>2</sub>, and the present flames are listed in Table 1). The sensitivity of the burning velocity to the rate of the branching reaction O + H<sub>2</sub> is also low, ranking as the twelfth most sensitive reaction, as compared to methane-air or moist CO-air flames, where the burning velocity is very sensitive to the rate of the branching reaction H + O<sub>2</sub> (which is also not important in the present flames).

Hence, while there is some inhibition, the effect of the iron species is mostly to promote the overall reaction through the iron-catalyzed reactions of CO and N<sub>2</sub>O described above. With

 $\rm H_2$  addition the promotion is less pronounced, not because of significant hydrogen radical recombination by the iron species, but because the moist system is dominated by the fast OH + CO reaction, so the iron-catalyzed reactions account for less of the CO and  $\rm N_2O$  consumption.

## **CONCLUSIONS**

The first measurements of the burning velocity of  $CO-N_2O$  flames with low (13 ppm) levels of hydrogen-containing impurities have been obtained, for  $0.6 \le \phi \le 3.0$  and with added nitrogen up to  $X_{N_2} = 0.25$ ; data have also been collected for flames with added hydrogen up to  $X_{H_2} = 0.005$ . The measured burning velocity of pure stoichiometric flames is 23.1 cm/s  $\pm$  0.6 cm/s, and the measurements with added hydrogen are in good agreement with those of other researchers which were obtained at higher hydrogen mole fractions. The present data and those of earlier investigations were numerically

TABLE 5 Sensitivity of Burning Velocity to the Specific Reaction Rate Constant for Stoichiometric CO–N<sub>2</sub>O Flames with  $X_{\rm H_2}=0$  and 0.002, and for  $X_{\rm Fe(CO)_5}=0$  ppm and 213 ppm.<sup>a</sup>

		d (ln v)/d (ln k) d (ln v)/d (ln k) $ _{max}$					
$X_{H_2}$ :		)	0.002				
$X_{Fe(CO)_5}$ (ppm):	0	213	0	213			
Reaction							
Dry Reactions							
$CO + N_2O \leftrightarrow CO_2 + N_2$	1.00	1.00	1.00	1.00			
$N_2O(+M) \leftrightarrow N_2 + O(+M)$	-0.13	-0.13	0.19	-0.10			
$N_2O + O \Leftrightarrow 2NO$	-0.07	-0.07	-0.34	-0.23			
$N_2O + O \leftrightarrow O_2 + N_2$	0.00	0.04	0.04	0.06			
Moist Reactions							
$CO + OH \leftrightarrow CO_2 + H$			0.83	0.63			
$N_2O + H \leftrightarrow N_2 + OH$			0.32	0.31			
$H_2O + O \Leftrightarrow 2OH$			-0.10	-0.06			
$O + H_2 \leftrightarrow H + OH$			0.09	0.09			
Iron Reactions							
$FeO_2 + CO \Leftrightarrow FeO + CO_2$		0.30		0.28			
$FeO + CO \Leftrightarrow Fe + CO_2$		0.09		0.04			
$FeO + N_2O \leftrightarrow FeO_2 + N_2$		0.14		-0.05			
$Fe + N_2O \leftrightarrow FeO + N_2$		0.02		0.04			
$FeOH + N_2O \Leftrightarrow FeOOH + N_2$				0.44			
$FeOOH + CO \leftrightarrow FeOH + CO_2$				0.25			
$FeO + H_2O \leftrightarrow FE(OH)_2$				0.21			
$FeOH + H \leftrightarrow FeO + H_2$				0.20			
$Fe(OH)_2 + H \leftrightarrow FeOH + H_2O$				0.13			
$FeOH + O \Leftrightarrow FeO + OH$				-0.08			
$FeOOH + OH \leftrightarrow FeO_2 + H_2O$				-0.04			

<sup>&</sup>lt;sup>a</sup> Sensitivities are normalized by the value for the maximum sensitivity, which is the direct CO +  $N_2$ O reaction.

modeled using a mechanism based on Allen et al. [1] with a CO + OH rate from Yu et al. [40], and the  $CO + N_2O$  direct reaction rate of Milks and Matula [6]. Modeling of the flames requires the use of the direct reaction, and the present results imply a rate of  $10^{9.2}$  cm<sup>3</sup> / mol s at 1800 K, which corresponds to a 10% decrease in the preexponential factor of the Milks and Matula rate. Experiments with nitrogen dilution and over a range of  $\phi$  suggest an activation energy near 71 kJ/mole. For dry flames with  $0.6 \le \phi \le$ 3.0, the mechanism overpredicts the burning velocity by about 18% for lean flames and underpredicts it by about 25% for rich flames. For the moist flames, the CO + OH rate also has a strong effect on the predicted burning velocity, and the rates of Yu et al. [40] or Wooldridge et al. [44] provide good agreement with our data.

Iron pentacarbonyl, which is the most effec-

tive flame inhibitor identified for hydrocarbonair flames, is not effective in N<sub>2</sub>O flames; in fact, 213 ppm of Fe(CO)<sub>5</sub> actually increases the burning velocity of the dry flames by about 25%. The promotion is believed to be due to the iron-catalyzed gas-phase reaction of N<sub>2</sub>O with CO, via  $N_2O + M = N_2 + MO$  and CO +  $MO = CO_2 + M$ , where M is Fe, FeO, or FeOH. The rate expression of Plane and Rollason [50] for the former reaction with M = Fe, together with estimates of the rates of other reactions provide reasonable agreement with the present data. For moist CO-N<sub>2</sub>O flames, the promotion provided by the iron pentacarbonyl is less pronounced, not because of radical recombination by the inhibitor, but because the iron-catalyzed reaction of CO and N2O is of lesser importance relative to CO and N2O reaction with OH and H, respectively.

The present results show that the extraordi-

nary effectiveness of iron pentacarbonyl may be limited to systems in which the oxidizer is O<sub>2</sub>. The findings emphasize that unlike thermal diluents, the effect a chemical "inhibitor" will have on the overall reaction rate is highly dependent upon the chemical system involved. For example, for some propellant flames that release CO and N<sub>2</sub>O in the gas phase, the most effective inhibitor found for hydrocarbon–air flames [Fe(CO)<sub>5</sub>] would likely accelerate the burning, as may halogen-based inhibitors.

We are grateful to Dr. Richard Yetter for helpful discussions of  $N_2O$  kinetics and for providing his  $N_2O$  mechanism, to Dr. Pamela Chu for the FTIR analysis of  $N_2O$  and CO, and to Ms. Nikki Privé for assistance with the data acquisition and uncertainty analysis programs. The helpful conversations and encouragement of Dr. Wing Tsang of NIST contributed much to this research.

#### REFERENCES

- Allen, M. T., Yetter, R. A., and Dryer, F. L., Combust. Flame 109:449 (1997).
- Cor, J. J., and Branch, M. C., J. Propul. Power 11:704 (1995).
- Litzinger, T. A., Fetherolf, B. L., Lee, Y. J., and Tang, C. J., J. Propul. Power 11:698 (1995).
- Yetter, R. A., Dryer, F. L., Allen, M. T., and Gatto, J. L., J. Propul. Power 11:683 (1995).
- Zaslonko, I. S., Losev, A. S., Mozzhukhin, E. V., and Mukoseev, Y. K., Kinet. Catalysis 20:1144 (1979).
- Milks, D., and Matula, R., Fourteenth Symposium (International) on Combustion, The Combustion Institute, Pittsburgh, 1973, pp. 84.
- Fujii, N., Kakuda, T., Takeishi, N., and Miyama, H., J. Phys. Chem. 91:2144 (1987).
- 8. Loirat, H., Caralp, F., and Destriau, M., *J. Phys. Chem.* 87:2455 (1983).
- 9. Dindi, H., Tsai, H. M., and Branch, M. C., *Combust. Flame* 87:13 (1991).
- 10. Vandooren, J., Van Tiggelen, P. J., and Pauwels, J. F., *Combust. Flame* 109:647 (1997).
- 11. Cor, J. J., and Branch, M. C., *Combust. Sci. Technol.* 127:71 (1997).
- 12. Van Wonterghem, J., and Van Tiggelen, A., *Bull. Soc. Chim. Belg.* 64:780 (1955).
- Kalff, P. J., and Alkemade, C. Th.J., Combust. Flame 19:257 (1972).
- Simpson, C. J. S. M., and Linnett, J. W., Sixth Symposium (International) on Combustion, Reinhold, New York, 1957, pp. 257–265.
- Linteris, G. T., and Williams, F. A., Twenty-Fourth Symposium (International) on Combustion, The Combustion Institute, Pittsburgh, 1992, pp. 803–811.

- 16. Andersen, S. O., Fire J. 81:56 (1987).
- Gann, R. G., Ed., Fire Suppression System Performance of Alternative Agents in Aircraft Engines and Dry Bay Laboratory Simulations, National Institute of Standards and Technology, NIST SP 890, 1995.
- Lask, G., and Wagner, H. G., Eighth Symposium (International) on Combustion, Williams and Wilkins, Baltimore, 1962, pp. 432–438.
- Vanpee, M., and Shirodkar, P., Seventeenth Symposium (International) on Combustion, The Combustion Institute, Pittsburgh, 1979, pp. 787–795.
- Miller, D. R., Evers, R. L., and Skinner, G. B., *Combust. Flame* 7:137 (1963).
- 21. Reinelt, D., and Linteris, G. T., *Twenty-Sixth Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1996, pp. 1421–1428.
- Linteris, G. T., Rumminger, M. D., Babushok, V. I., and Tsang, W., Twenty-Eighth Symposium (International) on Combustion, The Combustion Institute, Pittsburgh, 2000 (submitted).
- Rumminger, M. D., Reinelt, D., Babushok, V., and Linteris, G. T., Combust. Flame 116:207 (1999).
- Rosser, W. A, Inami, S. H., and Wise, H. (1959). Study of the Mechanisms of Fire Extinguishment of Liquid Rocket Propellants. WADC Technical Report 59-206.
- Rumminger M. D., and Linteris, G. T., Inhibition of premixed carbon monoxide–hydrogen–oxygen–nitrogen flames by iron pentacarbonyl, *Combust. Flame*, 120:451 (2000).
- Kaufman, F., Proc. R. Soc. London, A A247:123 (1958).
- 27. Mache, H., and Hebra, A., Sitzungsber. Österreich. Akad. Wiss. IIa:150:157 (1941).
- Van Wonterghem, J., and Van Tiggelen, A., *Bull. Soc. Chim. Belg.* 63:235 (1954).
- Andrews, G. E., and Bradley, D., Combust. Flame 18:133 (1972).
- Linteris, G. T., and Truett, L., Combust. Flame 105:15 (1996).
- Taylor, B. N., and Kuyatt, C. E. (1994). Guidelines for Evaluating and Expressing the Uncertainty of NIST Measurement Results. National Institute of Standards and Technology, NIST Technical Note 1297.
- Rumminger, M. D., and Linteris, G. T. (1999). Inhibition of Premixed Carbon Monoxide–Hydrogen–Oxygen–Nitrogen Flames by Iron Pentacarbonyl, National Institute of Standards and Technology, NIST IR 6360.
- Weinberg, F. J., Optics of Flames, Butterworth, London, 1963.
- Dunn-Rankin, D., and Weinberg, F., Combust. Flame 113:303 (1998).
- 35. Fristrom, R. M., Phys. Fluids 8:273 (1965).
- Gilbert, A. G., and Sulzmann, K. G. P., J. Electrochem. Soc. 121:832 (1974).
- Kee, R. J., Grcar, J. F., Smooke, M. D., and Miller, J. A. (1991). A Fortran Computer Program for Modeling Steady Laminar One-dimensional Premixed Flames, Sandia National Laboratories Report, SAND85-8240.
- 38. Kee, R. J., Rupley, F. M., and Miller, J. A. (1989). CHEMKIN-II: A Fortran Chemical Kinetics Package for

- the Analysis of Gas Phase Chemical Kinetics. Sandia National Laboratory, SAND89-8009B.
- Kee, R. J., Dixon-Lewis, G., Warnatz, J., Coltrin, R. E., and Miller, J. A. (1986). A Fortran Computer Package for the Evaluation of Gas-Phase, Multicomponent Transport Properties. Sandia National Laboratory, SAND86-8246.
- Yu, C.-L., Wang, C., and Frenklach, M., Eastern States Section of the Combustion Institute, The Combustion Institute, 1990, pp. 20-1–20-4.
- Loirat, H., Caralp, F., Destriau, M., and Lesclaux, R., J. Phys. Chem. 91:6538 (1987).
- Baulch, D. L., Drysdale, D. D., Horne, D. G., and Lloyd, A. C., Evaluated Kinetic Data for High Temperature Reactions, Vols. 1 and 2, Butterworths, London, 1973.
- Baulch, D. L., Cobos, C. J., Cox, R. A., Esser, C., Frank, P., Just, T., Kerr, J. A., Pilling, M. J., Troe, J., Walker, R. W., and Warnatz, J., J. Phys. Chem. Ref. Data 21:411 (1992).
- Wooldridge, M. S., Hanson, R. K., and Bowman, C. T., *Twenty-Fifth Symposium (International) on Combus- tion*, The Combustion Institute, Pittsburgh, 1994, pp. 741–748.
- Lin, M. C., and Bauer, S. H., J. Phys. Chem. 50:3377 (1969).
- Day, M. J., Stamp, D. V., Thompson, K., and Dixon-Lewis, G., Thirteenth Symposium (International) on

- Combustion, The Combustion Institute, Pittsburgh, 1971, pp. 705–712.
- 47. Palmer, H. B., and Seery, D. J., *Combust. Flame* 4:213 (1960).
- 48. West, J. B., and Broida, H. P., *J. Chem. Phys.* 62:2566 (1975).
- Campbell, M. L., and Metzger, J. R., Chem. Phys. Lett. 253:158 (1996).
- Plane, J. M. C., and Rollason, R. J., J. Chem. Soc., Faraday Trans. 92:4371 (1996).
- Kappes, M. M., and Staley, R. H., J. Am. Chem. Soc. 103:1286 (1981).
- 52. Matsuda, S., J. Phys. Chem. 57:807 (1972).
- Hayhurst, A. N., and Lawrence, A. D., *Combust. Flame* 110:351 (1997).
- Kellogg, C. B., and Irikura, K. K., J. Phys. Chem. A 103:1150 (1999).
- Rumminger, M. D., and Linteris, G. T., Halon Options Technical Working Conference, Albuquerque, NM, 1999, pp. 511–521.
- Rumminger, M. D., and Linteris, G. T., An experimental study of the role of particles in flame inhibition by iron pentacarbonyl, *Combust. Flame*. (submitted).

Received 17 August 1999; revised 11 January 2000; accepted 23 January 2000