# NGP RESEARCH ON FIRE SUPPRESSION CHEMISTRY

Gregory T. Linteris Fire Research Division, National Institute of Standards and Technology Gaithersburg, MD 20899 USA Tel: 301-975-2283; Fax: 301-975-4052; <u>linteris@nist.gov</u>

## ABSTRACT

Several NGP projects studied the mechanisms of chemical suppressants, including those containing halogens, phosphorus, and metals. From the work, several general principles of chemical flame inhibition are outlined. The present paper describes the conditions for which a chemically active agent can be effective, and when it is most effective, and when it may not be effective. These general principles are demonstrated with numerical and experimental data and analyses for flame inhibition by various metals, halogens, phosphorus, and inert compounds, in premixed, counterflow diffusion, and cup-burner flames.

## **INTRODUCTION**

The NGP research examining the effects of agent chemistry and flame inhibitor efficiency is outlined in references [1-13], and the present paper draws its material from those references. From that research, one can assemble some general principles of flame inhibition, and those basic principles are outlined here.

Two words commonly used to describe the influence of chemical additives on flames are inhibition and suppression. The term "suppression" refers to the extinguishment of a flame or fire, while the term "inhibition" describes a weakening of the flame, which may or may not lead to extinguishment. Whether a flame is only weakened or is completely extinguished through addition of an agent depends upon the amount of the agent applied, and how close the flame is to extinguishment to start with. That is, there is a balance between the time available for chemical reaction and the flow time, which are described by the extinction Dahmköhler number. Chemical additives typically work by increasing the chemical reaction time (i.e., lowering the overall reaction rate) so that the flame cannot be supported in the existing flow field [14]. While addition of any amount of chemical additive will weaken a flame, only a high enough concentration will extinguish it. As will be described below, the effectiveness of chemical additives is generally non-linear with agent concentration. Hence, the agent concentration of most interest is that which completely extinguishes the flame.

## **RESULTS AND DISCUSSION**

Flame inhibitors generally work through two mechanisms: physical and chemical. Physical mechanisms involve dilution of the reactants, a change in the average specific heat of the reactant mixture per mass of fuel burned (which typically is the more important effect). Chemical effects change the overall reaction rate due to either trapping of the reactive chainbranching radicals (for example with fluorinated hydrocarbons), or catalytic cycles (for example with HBr) which effectively recombine the radicals into less reactive species. Agents that enter

into catalytic cycles are more interesting, since far lower concentrations of these species are required to suppress fires. One might reasonably ask the question: "When is a catalytic inhibitor effective?" Answering this question is the goal of the present paper.

Catalytic flame inhibitors are effective when they get to the right place in the flame, at the right time, in the right form, and can then lower the volume fraction of chain-carrying radicals (O, H, and OH). Since we are dealing here with steady flames, the time part is not relevant (but could be in other situations). Each of these requirements is discussed below.

## AGENT LOCATION FOR EFFECTIVE INHIBITION

#### **Overview**

The most effective location for a flame inhibitor is where the radicals are. This is where the catalytic agents recombine the radicals, and consequently is where the active species need to be delivered. From both experimental measurements (see Figure 1, Figure 2, and Figure 3) and calculations (see Figure 4, bottom frame), it is known that catalytic agents reduce radical concentrations. Also, catalytic agents are most effective when their active species overlap in space with the peak radical volume fractions (see Figure 5).



Figure 1. Normalized OH LIF profile areas versus inhibitor agent delivery concentrations propaneair counterflow diffusion flame. Data legend: (□)

N<sub>2</sub>, (O)FM-200, ( $\Delta$ )FE-36, ( $\nabla$ )PN, ( $\diamond$ )CF<sub>3</sub>Br, ( $\blacksquare$ )DMMP, and ( $\bullet$ )Fe(CO)<sub>5</sub>. Insert: plot of the PN, CF<sub>3</sub>Br, DMMP, and Fe(CO)<sub>5</sub>, data for agent concentration up to 0.75 % volume. From ref. [10].



Figure 2. Dependence of normalized maximum OH LIF intensity on inhibitor concentration in a low-pressure CH<sub>4</sub>-air counterflow diffusion flame. Symbols:  $(\Box)$ Fe(CO)<sub>5</sub>,  $(\Delta)$ CF<sub>3</sub>Br, (O)N<sub>2</sub>. From ref. [11].



Figure 3. Effect of 572 ppm of DMMP on calculated OH, H, and O concentration profiles in Flame 1 (CH<sub>4</sub>/O<sub>2</sub>/N<sub>2</sub>). Dark lines are undoped profiles and thin lines are with DMMP addition. From ref. [8].



Figure 4. Burning velocity, temperature, and super-equilibrium ratio of flame radicals computed for atmospheric pressure stoichiometric methane/air mixtures inhibited by sodium hydroxide. From ref. [12].



Figure 5. Variation of flame speed (left axis) for four types of perturbation: reduction of the H+O<sub>2</sub> $\leftrightarrow$ OH+O or CO+OH $\leftrightarrow$ CO<sub>2</sub>+H reaction rate, and inhibition by a perfect agent or CF<sub>3</sub>Br. The "CF<sub>3</sub>Br baseline" refers to the flame speed when the halogen chemistry is damped by 10<sup>-4</sup>. The band width  $\Delta$  is 300 K, so the band extends 150 K below and above the temperature shown on the x-axis. The calculated volume fraction of OH, H, and O (right axis) is shown for an uninhibited stoichiometric methane-air flame. From ref. [13].

## Superequilbrium

The effectiveness of a given agent at the location of peak radical volume fraction is dependent upon the degree of radical super-equilibrium. Catalytic agents speed up the movement of a chemical system that is out of equilibrium to equilibrium but do not modify the equilbrium conditions themselves. As Figure 4 (bottom frame) shows, the uninhibited premixed methaneair flame has superequilibrium ratios of 2.5 to 18, and the degree of inhibition (marginal change in slope of the burning velocity in top frame) is decreased as the super-equilibrium ratio decreases. Also, as shown in Figure 6 (for premixed CH<sub>4</sub>-air flames at various fuel-air ratios and oxygen volume fraction, which change the super-equilibrium ratio), as the degree of radical super-equilibrium increases, the inhibition effect for a given amount of agent is also larger. For counterflow diffusion flames, the amount of strain (i.e., gas velocities) affects the degree of radical super-equilibrium is shown in Figure 7.

A consequence of the need for radical super-equilibrium is that the effectiveness of catalytic agents varies with the temperature of the flame. This is illustrated indirectly in Figure 6 (since the super-equilibrium is highest in the lowest temperature flames), and directly in Figure 8, which shows the measured effectiveness (from extinction strain rate or [OH] reduction) in counterflow diffusion flames as a function of flame temperature. Nonetheless, to reach the location of peak radical volume fraction in a system with radical super-equilibrium, effective agent transport is required.





Figure 6. Variation of inhibition paramter  $\Phi_0$  [15] with degree of H-atom super equilibrium in premixed methane-air flames with added Fe(CO)<sub>5</sub>. Data from ref. [16].

Figure 7. Degree of H-atom super-equilibrium and peak temperature as a function of strain rate for a  $CH_4$ -air counterflow diffusion flame.



Figure 8. Temperature dependence of phorsphorus agent effectiveness. Open circles are effectiveness data defined in terms of reduction of global extinction strain; filled squares are effectiveness data in terms of reduction in OH concentration) From ref. [8].

#### AGENT TRANSPORT

In order for an chemical agent to reach the region of peak radical volume fraction, it must be transported there. The efficiency of this process depends on the phase of the agent (solid, liquid or gas), and the location of agent addition relative to the flame and peak radical volume fractions.

The agent transport occurs primarily through convection or diffusion, and can be helped or hindered by thermophoresis (or thermal diffusion for gases). Figure 9 shows a counterflow diffusion flame with the flame (indicated by the blue line) on either the fuel or oxidizer side of the stagnation plane (indicated by the horizontal dotted line). This is achieved in practice by diluting the fuel and air stream (and is described by the varying stoichiometric mixture fraction). If agent is added to the side of the stagnation plane where the flame is, the agent convects through the flame; otherwise, it can only reach the flame by gaseous diffusion. If the agent is in the form of



Figure 9. Counterflow diffusion flame with flame on fuel or oxidizer side of stagnation plane.



Figure 10. Flame suppression effectiveness of 25,000 ppm argon as an oxidant-side or fuel-side additive vs. stoichiometric mixture fraction. Effectiveness is defined as  $((a_{q_0} - a_q)/a_{q_0})$ . Flames are methane/nitrogen vs. oxygen/nitrogen;  $a_{q_0} = 350 \pm 10 \text{ s}^{-1}$ . From ref. [7]

particles, they cannot diffuse, are entrained by the flow, and are susceptible to thermophoresis and momentum effects if they are large. The dependence of agent efficiency on the stoichimetric mixture fraction (i.e., flame location) is illustrated for a thermally acting agent (Ar) in Figure 10.

## ACTIVE MOITIES FOR FLAME INHIBITION

#### Formation

To be effective, the agent must also be present as the correct chemical compounds (which usually are not the form in which the agent is added). This means that the agent must break down in the flame, form species which are active in the catalytic cycle, and then not be lost to other non-active forms. For iron, the for inhibition has been shown to occur in the gas phase, and is believed to procede as indicated in Figure 11. First, the inhibitor molecule itself (in this case  $Fe(CO)_5$ ) must break down. Often, the rate of decomposition of the initial inhibitor molecule is not limiting, over a wide range of activation energy (see Figure 12). Second, in addition to being transported to the location of peak radicals, the breakdown products must be able to react with other species to form those active in the inhibition cycle. For iron, in the usually more important H-atom cycle, the species FeO, FeOH, and Fe(OH)<sub>2</sub> must be present in the gas phase in the region of the flame with highest radical volume fraction. Thus, Fe must be able to react with O<sub>2</sub> to form FeO<sub>2</sub>, which must then have access to O to form FeO. In some cases [17] this can be a limitation. Third, for inhibitors added as particles, they must have sufficient residence time to evaporate in the high-temperature region of the flame to deliver the active species [18].



Figure 11. Schematic of iron inhibition cycle.



Figure 12. The normalized burning velocity of stoichiometric  $CH_4/O_2/N_2$  flames at 400 ppm of ferrocene as a function of the activation energy of the one-step ferrocene decomposition reaction. From ref. [19].

#### **Properties of Active Species**

For the most effective chemical flame inhibition, the reaction rates within the catalytic cycle must be very fast (approaching gas-kinetic rates, with near zero activation energy, and near thermally neutral). This has also been dicussed in the context of the bond energies of the inhibiting species [20]. It also helps if there are multiple catalytic cycles for a given element [21]. Finally, the cycle must be complete, in that the inhibiting species are regenerated [22].

## **Loss Mechanisms**

To remain effective, the active species must not be subject to loss mechanisms. These can include condensation to particles [23, 24], loss to inactive gas-phase species [19], or shifting of the equilibrium constant for the active species [25-27]. The loss of effectiveness at higher concentrations and concurrent formation of particles for Fe(CO)<sub>5</sub> added to premixed flames is shown in Figure 13, and in counterflow diffusion flames in Figure 14. The loss of effectiveness of NaOH in water droplets (possible due to condensation of NaOH) is shown in Figure 15. The loss of iron compounds to Fe<sub>x</sub>F<sub>y</sub> compounds in flames with added Fe(CO)<sub>5</sub> and CF<sub>3</sub>H is shown in Figure 16. Finally, the shifting in the equilibrium products for an Mn-inhibited flame is shown in Figure 17, in which the volume fraction of Mn(OH)<sub>2</sub>, a necessary intermediate species in the catalytic cycle, is shown to decrease in volume fraction at higher temperature, which leads to lower effectiveness, (but greater effectiveness at lower temperature as compared to iron).



Figure 13. Normalized burning velocity (from Ref. [16]) and maximum  $Q_{\nu\nu}$  for  $\phi=1.0$  CH<sub>4</sub> flame with  $X_{O_{2}ox} = 0.21$  and 0.24 (from ref.: [23]).

Figure 14. Correlation between inhibition effect and maximum scattering signel  $Q_{\nu\nu}$ . Filled points are experimental normalized  $a_{ext}$ , solid line is calculated  $a_{ext}$  ([17]). Open symbols connected by dotted lines are maximum measured  $Q_{\nu\nu}$ . Particle data collected at 75 % of  $a_{ext}$  (from ref.: [24]).



Figure 15.  $Y_0$  and mole fraction of NaOH in air as a function of  $y_{NaOH}$ , for the extinction strain of 125 s<sup>-1</sup>. From ref. [28].



Figure 16 – Equilibrium mole fraction of active inhibiting species (Fe, FeO, FeOH, Fe(OH)<sub>2</sub>) and iron-fluorine species with 1 % to 4 % CF<sub>3</sub>H (containing 0.35 % ferrocene) added to a stoichiometric methane-air reaction mixture. From ref. [19].



Figure 17. Fraction of Mn-species at equilibrium as a function of temperature in methane-air flames. From ref. [27].

#### THE PERFECT FLAME INHIBITOR

A model was developed of a perfect flame inhibitor [22]. In it, the inhibitor is added in its effective form, is a gas, is non-condensing, and reacts only with the chain-branching flame radicals (i.e., it is non-reacting with all other species). The perfect inhibitor model is written in terms of the following two reactions: X + In = InX, and X + InX = X2 + In, where X is the major chain carrier and In is the inhibiting species. Both scavenging and regeneration reactions are termination processes. Thus the prefect catalytic recombination cycle is the most effective one due to termination of chain carriers at each reaction step of the catalytic cycle, and due to gas-kinetic rates for these reactions. Similarly, a perfect heterogeneous model was developed for flame inhibition by particles [23]. In it, all collisions of radicals with the particles effectively recombine the radicals into stable species. Figure 18 shows the predicted burning velocity reduction caused by a perfect gas-phase inhibitor, or a perfect heterogeneous inhibitor, together with experimental data for Fe(CO)<sub>5</sub>. As shown, the perfect inhibitor model will always have a higher efficiency than the perfect heterogeneous model. Interestingly, the efficiency of Fe(CO)<sub>5</sub> is relatively close to that of the perfect gas-phase inhibitor. As indicated in Figure 5, the perfect inhibitor model shows that for premixed flames, the region where innhibition is most effective is where the radicals are located. For CF<sub>3</sub>Br, the region of maximum effectiveness is also close to the region of peak radical volume fraction, but it's effectiveness there is limited by the equilibrium relationship for the reaction  $HBr + H = H_2 + Br$ , which shifts to the right at higher temperature, so that HBr is nolonger available for the key inhibition step.



Gas-Phase Precursor Volume Fraction (µL/L)

Figure 18. Calculated normalized burning velocity for several diameters d<sub>m</sub> of ideal heterogeneous inhibitor. Also shown are Fe(CO)5 data, and calculated normalized burning velocity using the perfect gas-phase inhibitor mechanism. From ref. [23]

#### CONCLUSIONS

Several properties which make chemical flame inhibitors effective or ineffective fire suppressants have been outlined. These have to do with both the properties of the flame, as well as of the inhibitor. Chemical suppressants have the potential to be up to two orders of magnitude more effective than  $CF_3Br$ . The difficult challenge is to devised methods to effectively deliver the active moieties to the radical production region of flames, and then keep them from condensing to particles, which are not as effective flame inhibitors.

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