A Computational Study of Extinguishment and Enhancement of Propane Cup-Burner Flames by Halon and Alternative Agents

FUMIAKI TAKAHASHI*
Department of Mechanical and Aerospace Engineering
Case Western Reserve University
10900 Euclid Avenue
Cleveland, OH 44106, USA

VISWANATH KATTA
Innovative Scientific Solutions, Inc.
7610 McEwen Road
Dayton, Ohio, 45459, USA

GREGORY LINTERIS and VALERI BABUSHOK
National Institute of Standard and Technology
100 Bureau Drive
Gaithersburg, MD 20899, USA

ABSTRACT
Computations of cup-burner flames in normal gravity have been performed using propane as the fuel to reveal the combustion inhibition and enhancement by the CF₃Br (halon 1301) and potential alternative fire-extinguishing agents (C₂HF₅, C₂HF₃Cl₂, and C₃H₂F₃Br). The time-dependent, two-dimensional numerical code used includes a detailed kinetic model (up to 241 species and 3918 reactions), diffusive transport, and a gray-gas radiation model. The peak reactivity spot (i.e., reaction kernel) at the flame base stabilizes a trailing flame, which is inclined inwardly by a buoyancy-induced entrainment flow. As the volume fraction of agent in the coflow increases gradually, the premixed-like reaction kernel weakens, thus inducing the flame base detachment from the burner rim and blowoff-type extinguishment eventually. The two-zone flame structure (with two heat-release-rate peaks) is formed in the trailing diffusion flame. The H₂O formed in the inner zone is converted further, primarily in the outer zone, to HF and CF₂O through exothermic reactions most significantly with the C₂HF₅ addition. The total heat release of the entire flame decreases (inhibiting) for CF₃Br but increases (enhancing) for the halon alternative agents, particularly C₂HF₅ and C₂HF₃Cl₂. Addition of C₃H₂F₃Br results in unusual (non-chain branching) reactions.

KEYWORDS: aircraft cargo-bay fire suppression; halon 1301 replacement; numerical simulation; diffusion flame stabilization; reaction kernel

1. INTRODUCTION
In accordance with the Montreal Protocol to protect the stratospheric ozone layer, the use of the effective fire suppressant CF₃Br (bromotrifluoromethane, Halon 1301) has been discontinued except for certain critical applications such as the suppression of cargo-bay fires in aircraft. Halon alternative agents must pass a mandated Federal Aviation Administration (FAA) test [1,2], in which a simulated explosion of an aerosol can, caused by a fire, must be suppressed by the agent. Unlike CF₃Br, some replacement agents, including C₂HF₅ (pentafluoroethane, HFC-125) and C₃H₂F₃Br (2-bromo-3,3,3-trifluoropropene, 2-BTP), when added at any concentration less than that required for inerting, created a higher over-pressure in the test chamber and thus failed the test.

Recent work [3-6] employing thermodynamic equilibrium and perfectly stirred-reactor calculations (for premixed systems) revealed that higher overpressures in the FAA aerosol can tests might be due to higher heat release from reaction of the inhibitor itself. Nonetheless, the agents should still reduce the overall reaction rate and inhibit the reaction. For diffusion flames, however, the flame structure, combustion inhibition, and enhancement processes are not yet fully understood. In previous papers [7,8], the authors reported the results of comprehensive numerical simulations for zero- and normal Earth-gravity cup-burner flames using the FAA aerosol can test [ACT] fuel mixture with CF₃Br, C₂HF₅, C₂HF₃Cl₂ (2,2-dichloro-1,1,1-trifluoroethane, HCFC-123), and C₃H₂F₃Br added to the coflowing air. In the halon-replacement-
agent molecules as compared to CF$_3$Br, additional numbers of carbon and fluorine atoms, as well as a double bond (C$_3$H$_2$F$_3$Br), represent potential energy contributions at a fixed concentration if they burn to COF$_2$ and HF. The ACT fuel mixture is somewhat unusual, however, in that it contains a large portion of water (which is an important reactant with the halogenated species). The objectives of this study are to investigate the effects of fire-extinguishing agents (with different numbers of carbon atoms and types of halogen) on the diffusion flame and to determine if the enhanced heat release found for the previous simulations with the ACT fuel occur with a more typical hydrocarbon fuel (propane).

2. COMPUTATIONAL METHOD

A time-dependent, axisymmetric numerical code (UNICORN) [9,10] is used for the simulation of coflow diffusion flames stabilized on the cup burner. The code solves the axial and radial ($z$ and $r$) full Navier-Stokes momentum equations, continuity equation, and enthalpy- and species-conservation equations on a staggered-grid system. A clustered mesh system is employed to trace the gradients in flow variables near the flame surface. The thermo-physical properties such as enthalpy, viscosity, thermal conductivity, and binary molecular diffusion of all of the species are calculated from the polynomial curve fits developed for the temperature range 300 K to 5000 K. Mixture viscosity and thermal conductivity are then estimated using the Wilke and Kee expressions [11], respectively. Molecular diffusion is assumed to be of the binary-diffusion type, and the diffusion velocity of a species is calculated using Fick's law and the effective-diffusion coefficient of that species in the mixture. A simple radiation model [12] based on the optically thin media and gray-gas assumptions was incorporated into the energy equation. Radiation from CH$_4$, CO, CO$_2$, H$_2$O, HF, COF$_2$ and soot was considered in the present study. The Plank-mean absorption coefficients are obtained from the literature for the first four species [12] and HF [13]; or calculated for COF$_2$ [13] and soot [14]. The finite-difference forms of the momentum equations are obtained using an implicit QUICKEST scheme [9], and those of the species and energy equations are obtained using a hybrid scheme of upwind and central differencing.

A comprehensive reaction mechanism was assembled for the simulation of propane or ACT fuel flames with CF$_3$Br, C$_2$HF$_5$, C$_2$HF$_3$Cl$_2$, or C$_3$H$_2$F$_3$Br added to air from four mechanisms: the four-carbon hydrocarbon mechanism of Wang and co-workers [15,16] (111 species and 1566 one-way elementary reactions), detailed reactions of ethanol (5 species and 72 reactions) of Dryer and co-workers [17,18], the bromine and chlorine parts of the mechanism of Babushok et al. [19-22] (10 additional species and 148 reactions), and a subset (51 species and 1200 reactions) of the National Institute of Standards and Technology (NIST) HFC starting mechanism [23, 24]. The final chemical kinetics model (187 species, 3198 reactions for CF$_3$Br, C$_2$HF$_5$, and C$_3$H$_2$F$_3$Br; or 241 species and 3918 reactions for C$_2$HF$_3$Cl$_2$) and a soot model [14] are integrated into the UNICORN code. Transport data for 139 species are available in the literature; for the remaining 38 species, data are constructed by matching these species with the nearest species (based on molecular weight) with known transport data.

The ACT fuel is a propane-ethanol-water mixture [1,2] with the volume fractions of the components: $X_{C_3H_8} = 0.159$, $X_{C_2H_5OH} = 0.454$, and $X_{H_2O} = 0.387$. Table 1 shows the minimum extinguishing concentrations (MECs) of fire-extinguishing agents for n-heptane and propane fuels in the literature [25-34] using the cup-burner method [25,26]. The calculated MECs obtained for the ACT fuel and propane in our previous and present studies are also listed and discussed in the results section. There are neither experimental data for the ACT fuel nor calculated results for n-heptane available.

The boundary conditions are treated in the same way as reported in earlier papers [7,8]. The computational domain is bounded by the axis of symmetry, a chimney wall, and the inflow and outflow boundaries. Calculations are made on a physical domain of 200 mm by 47.5 mm using a 251×101 non-uniform grid system that yielded 0.2 mm by 0.2 mm minimum grid spacing in both the $z$ and $r$ directions in the flame zone. The burner outer diameter is 28 mm and the chimney inner diameter is 95 mm. The burner wall (4-mm long and 1-mm thick tube) temperature is set at 600 K and the wall surface is under the no-slip velocity condition.
Table 1. Measured and calculated minimum extinguishing concentrations.

<table>
<thead>
<tr>
<th>Agent</th>
<th>Chemical</th>
<th>Formula</th>
<th>Measured n-Heptane MEC (%)</th>
<th>Measured Propane MEC (%)</th>
<th>Calculated ACT Fuel MEC (%)</th>
<th>Calculated Propane MEC (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Halon 1301</td>
<td>Bromotrifluoromethane</td>
<td>CF₃Br</td>
<td>3.0 to 3.2 [26-27]</td>
<td>3.8 to 4.3 [27,29,30]</td>
<td>2.26</td>
<td>2.64</td>
</tr>
<tr>
<td>HFC-125</td>
<td>Pentfluorothane</td>
<td>C₂HF₅</td>
<td>8.7 to 9.3 [24-28]</td>
<td>10.2 to 10.4 [27,29]</td>
<td>8.40</td>
<td>7.65</td>
</tr>
<tr>
<td>HCFC-123</td>
<td>2,2-dichloro-1,1,1-Trifluoroethane</td>
<td>C₂HF₃Cl₂</td>
<td>7.1 to 7.4 [31]</td>
<td>N/A (4.90b) (4.60b)</td>
<td>N/A (1.87b) (2.50b)</td>
<td></td>
</tr>
<tr>
<td>2-BTP</td>
<td>2-bromo-3,3,3-Trifluoropropene</td>
<td>C₃H₂F₃Br</td>
<td>2.6 [32] 4.7 [33]ᵃ</td>
<td>N/A (4.90b) (4.60b)</td>
<td>N/A (1.87b) (2.50b)</td>
<td></td>
</tr>
</tbody>
</table>

ᵃThe fuel temperature: 50 °C
ᵇA concentration above which the calculation was unable to obtain the solution

The mean fuel velocities for the ACT fuel and propane are 0.853 cm/s and 0.307 cm/s, respectively, and the mean velocity of the oxidizer (“air” [21 % O₂ in nitrogen] with added agent) is 10.7 cm/s at 294 K.

Validation of the code with the kinetic model was performed through the simulation of opposing-jet diffusion flames. The predicted extinction strain rates for propane-air flames (no agent) were within 7.5% of the measured values (with an error margin of 9 %) by Zegers et al. [35]. The predicted extinction agent concentrations for CF₃Br and C₂HF₅ are within 4 % of the measured concentrations in weakly stretched flames and within 25 % in highly stretched flames. Although cup-burner data using the ACT fuel are unavailable for a direct comparison, computation with the assembled reaction mechanism should provide insights into the detailed flame structure.

3. RESULTS AND DISCUSSION

3.1 Internal flame structure

The flame base supports a trailing flame and controls the flame attachment, detachment, and oscillation processes [36,37]. Small variations in the agent volume fraction in the coflowing oxidizing stream (Xₐ) results in profound changes near the extinguishment limit. Figure 1 shows the calculated structure of near-limit propane flames in air with added agent: (a) CF₃Br, Xₐ=0.0258; (b) C₂HF₅, Xₐ=0.0765; (c) C₂HF₃Cl₂, Xₐ=0.0455; and (d) C₃H₂F₃Br, Xₐ=0.0246. The variables include the velocity vectors (v), isotherms (T), and heat-release rate (q̇). The base of the agent-added flames are detached and drift inward a few mm away from the burner rim by the nearly horizontal entrainment flow. In contrast to zero-gravity (0gₙ) flames [7], which are formed vertically, the normal gravity (1gₙ) flame inclines inwardly due to the streamline shrinkage in the accelerating buoyancy-induced flow. The contours of the heat-release rate show a peak reactivity spot (i.e., the reaction kernel [36]) at a height from the burner rim, zₖ = 0.8 mm to 1.8 mm. The chain propagating radical species (H, O, and OH) as well as heat diffuse back against the oxygen-rich flow at the flame base (edge), thus promoting vigorous reactions to form the reaction kernel.

Unlike the flame with CF₃Br (Fig. 1a), the other near-limit flames, particularly with C₂HF₅ (Fig. 1b) and C₃H₂F₃Br (Fig. 1d), show distinct “two-zone” flame structure as evident from nearly parallel zones of the high heat-release rate contours.

Figure 2 shows the radial variations of the calculated temperature and heat-release rate in propane flames in air with agent: (a) across a trailing flame (zₖ + 5 mm); (b) across the reaction kernel: zₖ = 1.8 mm (CF₃Br), 1.2 mm (C₂HF₅), 0.8 mm (C₂HF₃Cl₂), and 1.4 mm (C₃H₂F₃Br). The trailing flames (Fig. 2a) are characterized by the two-zone flame structure (inner and outer) as evident from two heat-release rate peaks most prominently for C₂HF₅ and least significantly for CF₃Br. Although the temperature peak is closer to the inner reaction zone, formed by the hydrocarbon-O₂ combustion, the larger heat-release rate peak for C₂HF₅ is in the outer zone by highly exothermic reactions. The temperature and heat-release-rate profiles in the propane flame with C₂HF₅ (Fig. 2a) are similar to those obtained previously [8] for the ACT fuel with C₂HF₅.
The outer heat-release-rate peak in the trailing flame in 1g (Fig. 2a) is more evident, compared to the 0g case [7], due to increased convective fluxes of reactants (i.e., the blowing effect [36]) by the buoyancy-induced incoming flow.

At the reaction kernel in the flame base region (Fig. 2b), the peak heat-release-rate for each agent slightly on the airside of each temperature peak is several times larger than that in the trailing flame. The peak temperature for C\textsubscript{3}H\textsubscript{2}F\textsubscript{3}Br is much higher than other agents, suggesting that additional number of carbon in the agent molecule, compared to CF\textsubscript{3}Br, represent potential energy contributions at a fixed concentration.

Figure 3 shows the radial variations of the species volume fractions (X\textsubscript{i}) crossing the trailing flame with C\textsubscript{2}HF\textsubscript{5} (added at X\textsubscript{a}=0.0765) at z= 6.2 mm. Oxygen penetrates through the outer zone and a pool of chain
carrier radicals (H, O, and OH) is formed in the middle of the two zones at relatively high concentrations ($X_a \approx 10^{-3}$), thus contributing to both reaction zones. The initial hydrocarbon fuel ($C_3H_8$) diffuses from the fuel side, decomposes to fragments ($CH_4$, $C_2H_4$, and $C_2H_2$) and reacts with the chain carrier radicals in the inner zone. In the outer zone, the agent ($C_2HF_5$) from the air side decomposes to many fluorinated species ($C_2F_6$, $CF_2$, $CHF_3$, etc.), which react with the radicals. The $H_2O$ (formed by hydrocarbon-O2 reaction) diffuses to the outer zone, where it is converted to HF through highly exothermic reactions. The $H_2O$ nearly vanishes in the outer zone in the propane flame, whereas that in the ACT fuel flame with $C_2HF_5$.

Fig. 2. Calculated radial variations of the temperature and heat-release rate in propane flames in air with agent: (a) across a trailing flame (at $z_k + 5 \text{ mm}$); (b) across the reaction kernel (at $z_k$). $CF_3Br$, $X_a = 0.0258$, $z_k = 1.8 \text{ mm}$; $C_2HF_5$, $X_a = 0.0765$, $z_k = 1.2 \text{ mm}$; $C_2HF_2Cl_2$, $X_a = 0.0455$, $z_k = 0.8 \text{ mm}$; and $C_3H_2F_3Br$, $X_a = 0.0246$, $z_k = 1.4 \text{ mm}$.

Fig. 3. Calculated structure of a propane flame in air with added $C_2HF_5$ at $X_a = 0.076$ and $z = 5.8 \text{ mm}$.
reported previously [8], remains at a $X_a \approx 10^{-3}$ level even outside the outer zone due to its high content ($X_{H2O} = 0.387$) in the fuel. The CF$_2$O peak ($X_{CF2O} = 0.031$) in the outer zone in the propane flame is lower than that ($X_{CF2O} = 0.048$) in the ACT fuel flame. The final products (CO$_2$, HF, and CF$_2$O) are distributed radially in a wide range. Low levels of C$_2$HF$_5$ on the fuel side and H$_2$ on the air side in Fig. 3 are due to leakage in the opposite directions through the quenched zone below the flame base. These species’ contributions to overall reactions in the opposite zones must be insignificant.

Figure 4 shows the radial variations of the calculated production (+) or consumption (-) rates (Fig. 4a) and heat-release rates (Fig. 4b) of species $i$ crossing the trailing flame at $z=5.8$ mm in a propane flame in air with C$_2$HF$_5$ at $X_a=0.076$. In the inner zone, H$_2$, CO, and the chain carrier radicals (H, O, and OH) are formed and consumed, O$_2$, and CF$_2$O are consumed, and H$_2$O, HF and CO$_2$ are formed. In the outer zone, C$_2$HF$_5$ and O$_2$ are consumed and HF, CF$_2$O, and CO are formed. The major contributors to the overall heat-release rate (Fig. 4b) are the formation of H$_2$O, CO, CO$_2$, HF in the inner zone, with HF, CF$_2$O and CO in the outer zone. Although the production rates and heat-release rates of HF and CF$_2$O in the propane flame are significantly less than those in the ACT fuel flame [7], the resultant heat-release rate profiles are very similar for the two flames. The highly exothermic reactions with the heats of reactions ($\Delta H$) include:

- $\ce{H2 + OH -> H + H2O} (\Delta H = -61 \text{ kJ/mol})$ (R5)
- $\ce{CO + OH -> CO2 + H} (\Delta H = -102 \text{ kJ/mol})$ (R61)
- $\ce{H2O + F -> OH + HF} (\Delta H = -74 \text{ kJ/mol})$ (R1685)
- $\ce{H2 + F -> H + HF} (\Delta H = -135 \text{ kJ/mol})$ (R1679)
- $\ce{CF2 + OH -> CF2O + H} (\Delta H = -268 \text{ kJ/mol})$ (R1849)
- $\ce{CF3 + OH -> CF2O + HF} (\Delta H = -493 \text{ kJ/mol})$ (R1669)
- $\ce{CF2 + O -> CFO + F} (\Delta H = -150 \text{ kJ/mol})$ (R1849)
- $\ce{CF3 + O -> CF2O + F} (\Delta H = -342 \text{ kJ/mol})$ (R1663)
- $\ce{CF3 + H -> CF2 + HF} (\Delta H = -215 \text{ kJ/mol})$ (R1719)

The reactions to form CF$_2$O are particularly exothermic because of its exceptionally low (negative) heat of formation (-640 kJ/mol).
3.2 Flame characteristics

Figure 5 shows the effects of the agent volume fraction in the coflowing oxidizer on the calculated axial ($z_k$) and radial ($r_k$) positions of the reaction kernel from the burner exit on the axis in propane flames. In the present unsteady calculations, as $X_a$ was increased incrementally, the flame-stabilizing reaction kernel in the flame base detached from the burner rim and moved downstream (i.e., the inward and upward direction) gradually and then more steeply as the extinguishment limit approached. For each $X_a$, a stable stationary flame was obtained.

For CF$_3$Br (and, to a lesser extent, C$_2$HF$_3$), the flame base oscillated, until finally, blowoff-type extinguishment occurred, whereas for C$_2$HF$_3$Cl$_2$ and C$_3$H$_2$F$_3$Br, the calculation abruptly diverged at $X_a=0.046$ and $X_a=0.025$, respectively. The radial location of the reaction kernel decreased (inward) with $X_a$, thereby more premixing occurred over the standoff distance. For propane, the MECs of CF$_3$Br and C$_2$HF$_3$ (see Table 1) are: $X_a \approx 0.04$ and $\approx 0.1$, respectively (measured); and $X_a=0.0264$ and 0.0765, respectively (calculated). By considering technical difficulties, including the stiffness in the computation, complex combustion and inhibition chemistries, and transient blowoff phenomena with occasional flame-base oscillations, the calculated MECs are in fair agreement ($\approx 30\%$) with the measurements.

The incoming flow velocity around the flame base is important in diffusion flame stability as it represents the reciprocal of the residence time through the reaction kernel. Figure 6 shows the effects of the agent volume fraction in the oxidizer on the calculated total ($|v_k|$), axial ($U_k$), and radial velocity ($V_k$) at the reaction kernel. For all agents, as $X_a$ was increased, the absolute values of $|v_k|$ increased and decreased moderately, and then it increased steeply (for CF$_3$Br and C$_3$H$_2$F$_3$Br) as the flame lifted off. Figure 7 shows the temperature, heat-release rate, and a ratio of the heat-release rate and the total velocity at the reaction kernel. For all agents, as $X_a$ was increased, the reaction kernel weakened (lower heat-release rate), but the flame stabilized at higher temperature and velocity ($|v_k|$). Nevertheless, the quantity $q/|v_k|$ (which relates to a ratio of the residence time and the reaction time, i.e., local Damköhler number, at the reaction kernel [36]) decreased mildly to a minimum level ($\approx 3$), which was comparable to the chemically passive agents [38, 39]. This result suggests that the reaction kernel shifted gradually downstream (inward) to seek a location where a subtle balance between the residence time and the reaction time can be achieved. As the flame lifted higher, it became more difficult to obtain the balance, thus leading to blowoff eventually.
3.3 Combustion enhancement

Figure 8 shows the maximum temperature in the trailing diffusion flame and the total heat-release rate ($q_{\text{total}}$) integrated over the entire flame and over the flame base region ($q_{<z=k+3 \text{ mm}}$). Thus, both the heat-release rate per unit volume along the flame and the flame size affect $q_{\text{total}}$. Unlike chemically passive agents [38,39], which work thermally to reduce the flame temperature by dilution, the maximum flame temperatures in the present work are nearly constant ($\approx 1800$ K) for C$_2$HF$_5$ or mildly increased for CF$_3$Br, C$_3$H$_2$F$_3$Br and C$_2$HF$_3$Cl$_2$ as $X_a$ increased until extinguishment. There is a striking difference in $q_{\text{total}}$ over the entire flame between CF$_3$Br and the other agents: $q_{\text{total}}$ decreased (i.e., inhibition) with added CF$_3$Br, whereas it increased (i.e., combustion enhancement) with C$_2$HF$_5$ or C$_3$HF$_3$Cl$_2$. It is neutral for C$_3$H$_2$F$_3$Br. In contrast, for all agents, $q_{<z=k+3 \text{ mm}}$ was nearly constant as $X_a$ increased. Thus, the combustion...
enhancement occurred only in the trailing flame. In fact, the heat release in the trailing flame ($\dot{q}_\text{total} - \dot{q}_{<z_k+3\text{ mm}}$) tripled with added C$_2$HF$_5$ (at $X_a \approx 0.08$). This enhancement is $\approx 1.5x$ larger than the zero-gravity flames studied previously [7], because of much higher incoming flow velocity in normal gravity, resulting in higher reactants (agent and oxygen) influx into the flame zone. Although the volumetric heat-release rate in the trailing flame was an order-of-magnitude smaller than the peak $\dot{q}_k$, integration over the entire trailing flame zone made the total value much larger. This result suggests the significant implication that even if the reaction kernel, with premixed-like flame structure, is weakened by halogenated agent addition toward the flame stability limit, the trailing diffusion flame can burn more reactants (including the agent itself) because of the additional heat release to form HF and CF$_2$O in the aforementioned “two-zone” flame structure.

4. CONCLUSIONS

By using propane as the fuel, in addition to the ACT fuel studies previously, the physical and chemical effects of Halon 1301 (CF$_3$Br) and halon-replacement fire-extinguishing agents (C$_2$HF$_5$, C$_2$HF$_3$Cl$_2$, and C$_3$H$_2$F$_3$Br) are studied numerically to gain better understanding of the flame structure, combustion inhibition/enhancement, and blowoff extinguishment of cup-burner flames. Addition of agent to the coflowing air weakens the flame attachment point (reaction kernel) at the flame base, thereby inducing the detachment, lifting, and blowout extinguishment. With added agent, the calculated maximum flame temperature remains nearly constant ($\approx 1800$ K) for C$_2$HF$_3$ or mildly increases for CF$_3$Br, C$_3$H$_2$F$_3$Br, and C$_2$HF$_3$Cl$_2$, while the reaction kernel temperature increased for all agents. Moreover, the total heat release of the entire flame increases with agent addition for C$_2$HF$_3$ and C$_2$HF$_3$Cl$_2$ (by up to a factor of 2.5). In the trailing flame, H$_2$ and H$_2$O (from hydrocarbon combustion) are converted to HF and CF$_2$O by exothermic reactions, enhancing an inner flame zone, while reactions of the inhibitor, also forming of HF and CF$_2$O, created a large outer heat-release zone. In contrast, CF$_3$Br reduced the total heat release.

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