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Effects of H₂O, CO₂, and N₂ Air Contaminants on Critical Airside Strain 

Rates for Extinction of Hydrogen-Air Counterflow Diffusion Flames 

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ABSTRACT 

Coaxial tubular opposed jet burners (2.7, 5.0 and 7.0 mm i.d. OJB) were used to form dish shaped counterflow diffusion flames (CFDF), centered by opposing laminar jets of H₂ (N₂ diluted in some cases) and both clean and contaminated "air" (O₂/N₂ mixtures) in an argon bath at 1 atm. Jet velocities for flame extinction (Blowoff) and restoration (Restore) limits are shown versus wide ranges of contaminant (0 to 20 mole %) and O₂ (16% to 30%) concentrations in the air jet, and also input H₂ concentration (30% to 100%). Blowoff, a sudden breaking of CFDF to a stable ring shape, occurs in highly stretched stagnation flows and is generally believed to measure kinetically limited flame reactivity. Restore, a sudden restoration of central flame, is a relatively new phenomenon which exhibits a H₂-dependent hysteresis from Blowoff. Both 2.7 and 7.0 mm OJB jet velocities at Blowoff were converted to critical radial strain rates, α; resultant αair were 3 to 5 times smaller than recently published numerical results for 100%, 50%, and 21% H₂/N₂ fuel inputs and normal air. Whereas measured αfuel doubled over the range 80 to 100% H₂, αair was constant. The αair for pure H₂ is considered a valid relative measure of kinetically controlled O₂ consumption by a H₂-air CFDF just before extinction. Furthermore, αair is a more fundamental measure of CFDF extinction than previously used fuel velocity (or strain rate), which mainly satisfies jet momentum balance for centering an OJB flame. 

For 25% O₂ "air" mixtures, mole-for-mole replacement of 25% N₂ contaminant by steam increased U(air) or "flame strength" at Blowoff by about 5%. This result is consistent with laminar burning velocity (Su) results from analogous substitution of steam (25%) for N₂ in a premixed stoichiometric H₂-O₂-N₂ (or steam) flame, shown by Koroll and Mulpuru to promote a 10% increase in experimental and calculated Su, due to enhanced third body (M) efficiency of water in: H + O₂ + M → HO₂ + M. When the OJB results were compared with Liu and MacFarlane's experimental Su of premixed stoichiometric H₂ + air + steam (where N₂ + steam > 55.7 mole %), a crossover occurred, i.e. steam enhanced OJB "flame strength" at extinction relative to Su. This finding remains consistent with Koroll and Mulpuru's extensive Su results, however, which show a crossover at 45% inert in stoichiometric H₂-O₂-inert flames; i.e. substitution of steam for N₂ (≥ 55.7% inert) decreases Su, due to dominance of thermodynamic effects at lower O₂ levels. Overall, the relative effect of steam, compared to N₂, on extinction of H₂ vs. air + steam CFDF is about one-half the incremental effect on Su of stoichiometric H₂ + O₂ + N₂ + steam mixtures over a wide range of concentrations. In contrast, CO₂ contamination significantly reduced U(air) at extinction relative to steam. 

Finally, unpublished analyses of previous OJB data on extinction of H₂/N₂ jets vs. contaminated air yielded characterizations of the U(air) intercept, at unity H₂ mole fraction, as functions of N₂, CO₂, and also NO, CO concentrations in air; results were generally consistent with recent data derived from pure H₂ jets. 

NOMENCLATURE 

X(i) = mole fraction of incoming species i in fuel or air jet. 

H₂/O₂ = incoming ratio of mole H₂/mole O₂ in air. 

M(H₂) = average mass flux, over tube cross section, of incoming H₂ in fuel mix, g H₂/cm²-s. 

C(i) = mole percentage of contaminant i in incoming air jet. 

D(i) = normalized linear percent decrease of U(air) or M(H₂) per mole percent contaminant. 

U(air) = average velocity of air jet at tube exit, cm/s. 

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BACKGROUND AND INTRODUCTION

Because of recent interest in National Aero-Space Plane technological developments, there is need to understand the influence of certain facility related (typically combustion produced) air contaminants on the ignition and extinction of H2-air diffusion flames in high speed shear flows. For example, because ground-based testing of H2-air scramjet combustors requires high enthalpy supersonic air, which typically is grossly contaminated with CO2 and/or H2O, flameholding of highly stretched H2-air diffusion flames in local recirculation regions may be perturbed relative to inlet air from the free atmosphere. Specifically, the presence of up to 30% H2O, 10% CO2, and 3% NO are of concern when H2 and hydrocarbon (HC) vitiated air, and arc jet facilities are used. For example, earlier studies of the effects of contaminants on H2 ignition and combustion, using shock tube ignition techniques, jet stirred combustor residence time studies, and laminar burning velocity measurements in conical flames have relevance to ignition and combustion limits for mixtures, but not for unpremixed combustion.

An opposed jet burner (OJB) technique was previously developed at NASA LaRC to characterize the effects of key air contaminants on strain (flame stretch) induced extinction limits of N2 diluted H2 versus air counterflow diffusion flames (CFDF). Extensive measurements of critical jet mass flow rates which led to sudden extinction of central flame formed by a N2-diluted H2 jet opposed by air, were reported for clean and contaminated airs. Resulting critical jet input velocities, U(air) and U(fuel) can be generalized for application to practical systems by estimating corresponding axial-input and radial-output strain rates (velocity gradients). Although the author's previous exclusive use of input hydrogen mass fluxes, M(H2), and molar H2/O2 input ratios was technically adequate for characterizing extinction (Blowoff) and reignition (Restore) of these flames as a function of H2 mole fraction in the fuel jet, X(H2), it tended to obscure the fundamental nature of the results, as described below.

Recent LDA axial velocity measurements using the authors' 7 mm OJB, considerations of input jet momentum balances, and reduction and crossplotting of earlier data highlighted a basic recognition that, whereas U(fuel) at extinction generally doubled over the fuel input range of 0.8 < X(H2) < 1, U(air) remained virtually constant. This led to a hypothesis that chemical kinetics control the combustion rate of O2 in this constant air velocity region. Because the molar H2/O2 input ratio greatly exceeds the stoichiometric ratio of 2 in this regime, overall H2 is in excess and the supply of hydrogen to the reaction zone (predicted and observed on the airside of the stagnation surface) is achieved by molecular and thermal diffusion of H2 and H. Thus an exponential increase in fuel jet velocity as X(H2) increases simply satisfies the fuel-air jet momentum balance to stabilize the flame centrally. As a correlative to this new interpretation, it follows that U(air) should be the preferred yardstick for measuring the effects of air contaminant concentration on CFDF extinction.

Therefore, for more dilute H2 jets, X(H2) < 0.8, diffusion of H2 in the fuel jet may become progressively important and the reaction zone may locate closer to the stagnation surface -- further velocity and composition profile measurements are needed to better define this regime so that models of CFDF can be more fully validated.

Problem Definition Regarding Effects of Steam, and Objectives of this Paper

Recently it has become clear that steam has an anomalous and extraordinary effect on flame structure and laminar burning velocity, Su, of H2-O2 mixtures. In early studies of H2 (and HC) combustion, steam was usually considered to behave as a simple diluent which lowers Su by acting as a heat sink, thereby reducing flame temperature. However beginning with Kuehl's pioneering study in 1962 of the positive effect of added water, relative to N2, on Su of H2-air mixtures, followed by recent detailed measurements of Su for heated mixtures of H2-air-steam by Liu and MacFarlane, and finally a recent and thorough experimental and theoretical study by Koroll and Mulpuru, we now know that progressive reductions in Su due to steam addition are not commensurate with changes in the heat capacity of the mixture. In fact, steam enhances Su by up to 16% when it simply replaces N2 diluent in a mixture; and furthermore we know the principal chemical kinetic reasons for this effect.

Although the Koroll and Mulpuru results offer important chemical kinetic insight on the effect of steam in premixed H2-O2 flames (discussed in greater detail below), comparable experimental and/or analytic studies of the possible effects of steam on extinction of simple or highly stretched H2-air diffusion flames have not been published, except for a recent study, by some of the present authors, of H2O effects based on changes in hydrogen mass flux.

Thus the objectives of the present paper are first: To review the relevant literature on CFDF; and summarize our recent findings that air jet velocities (strain rates) at blowoff of H2-air flames provide a more direct and fundamental measure of kinetic effects on flame extinction than fuel jet velocity, which indirectly measures strain rates. Second, new data are presented on the specific effects of H2O, CO2, N2, and O2 air additives on resultant air jet velocities (strain rates) at extinction. Finally, the extinction results are compared with published burning velocity measurements on the effects of steam in premixed flames, in an attempt to generalize the results.
Opposed Jet and Other Counterflow Diffusion Flames

Pioneering investigations by Potter et al. and Anagnostou and Potter revealed that simple, circular, coaxial tubes were suitable for determining reaction-limited opposed jet combustion rate characteristics of various fuel and oxidizer gases (expressed as Apparent Flame Strength, defined as average of cold fuel and oxidizer mass fluxes at flame Blowoff). Preliminary characterizations indicated that flame strength was essentially independent of tube separation distance (from 0.4 to 4 D) when uniform tubes (diameter D) were used and the flow profile was parabolic, but not when "flat profile" nozzle flows were tried. Flame strength was linearly proportional to D for laminar flow, and also for turbulent flow; additionally, the latter result was in conformance with Spalding's main simplifying assumption (flame within D/10 of stagnation plane for Peclet no. > 1000) and consequent prediction of linearity with D. Finally, Potter et al. concluded that pure H₂ burned at a significant distance on the air side of the stagnation point, based on Spalding's theory and their confirming observation of a 3:1 air/H₂ mass flux imbalance. Introduction of magnesia powder proved that hydrocarbon-air flames also lay on the air side of the stagnation point.

Potter's pioneering approach was supported and augmented by Spalding's theory of mixing and chemical reaction; Pandya and Weinberg's classic measurements of temperature and axial/radial velocity fields; Fendell's theory of ignition and extinction, which emphasized use of the first Damkohler number; Tsuji and Yamaoka's original temperature and flow pattern measurements; and later important theoretical developments. Further detailed observations of species profiles, coupled with temperatures, flow structures and velocities of HC-air CFDF have been reported by Tsuji and Yamaoka for propane ejected into a stream of air from a porous cylinder, and similarly for methane using the same apparatus. In 1982 Tsuji published an extensive review of CFDF. More recently, there have been several important papers on theory and application of counterflow diffusion, and also counterflow premixed, flames.

By now it is generally agreed that certain rate-limiting combustion reactions within the flame zone usually determine the extinction (Blowoff) of a counterflow diffusion flame. That is, as Blowoff is approached by gradually increasing the flows of fuel and air, it tends to occur suddenly when the combustion rate of fuel and air, initially at near-ambient temperatures, cannot keep up with the rate of convective (and some radiative) heat loss from the central flame/stagnation region. Thus for a given flame system, exceeding certain limiting flow rates does not allow the diffusing fuel and convecting air to remain in the flame zone long enough to support a flame. At this critical set of fuel and air flow velocities (which result in a critical radial velocity gradient in the stagnation region), the central portion of the flame 'blows off' catastrophically to form a stable torus or ring shaped flame (see Fig. 2). The flame is now located substantially outside of the projected tube diameter where the radial flow velocities have slowed enough to allow combustion reactions.

Early Observations Using the Authors' OJB System

When the present authors developed their OJB to begin determination of Blowoff limit conditions for H₂-diluted mixtures of H₂ opposed by air at ambient conditions, they reproduced the single highly non-symmetric result that Potter reported for pure H₂ versus air (3:1 mass flux ratio of air/H₂). The authors also observed a sudden and unique restoration of the central flame; this new critical point was named Restore.

Whenever Restore was slowly approached after Blowoff, the ring-shaped residual flame appeared to creep slightly inward at first. At some critical velocity, however (which was generally much higher than that required for autoignition of pyrophoric SiH₄/H₂ mixtures), the flame instantly "healed" the remaining circular opening, which was typically about one tube diameter in size. Hence it is argued that Restore does not appear to be an autoignition phenomenon, but instead an abrupt restoration of the central flame via radial 'inward flame propagation,' which appears to overcome a maximum outward radial velocity of gas mixture somewhere near the periphery of the jet impingement zone. Extensive characterizations of Restore indicated that, for H₂/N₂ vs. air systems: (a) Restore was readily and reproducibly observed after Blowoff; (b) the functional relationship between M(H₂) and X(H₂) was independent of OJB tube diameter; and (c) Restore exhibited a large hysteresis from Blowoff in hydrogen (but not hydrocarbon) containing flame systems.

Overall Structure of H₂-Air CFDF

The central (axial) portion of a counterflow diffusion flame system consists of two distinct parts, a nearly planar stagnation surface and a luminous flame zone (see Fig. 1). The stagnation surface is located on the fuel rich side of the luminous flame. Near the stagnation surface, axial flows of fuel and air exchange momentum and their respective axial velocities decrease to zero, and then become radial. Molecular and atomic hydrogen must diffuse to the luminous flame zone (by ordinary multicomponent and thermal diffusion); but air is transported to the flame, and resultant products pass through the flame, mainly by convection.
As the flows move radially away from the central stagnation point, their velocities first increase approximately linearly out to the tube edge and then decrease rapidly as they move away from the tube, thus giving a maximum radial velocity near the projected tube diameter. This flow is schematically illustrated in Fig. 2, where the tracing of any blowoff line, from the origin (tube center) out to the tube diameter (for either of two different tubes), shows a linear increase in radial velocity, and then an abrupt decrease in radial velocity as the flow moves away from the tube edge.

AXIAL-RADIAL FLOWFIELD FOR OJB COMBUSTION

![Schematic of laminar flowfield for a wide, flat, counterflow \( \text{H}_2\)-air diffusion flame.](image1)

Fig. 1. Schematic of laminar flowfield for a wide, flat, counterflow \( \text{H}_2\)-air diffusion flame.

Effects of Steam on Laminar Burning Velocity, \( S_u \), of Premixed \( \text{H}_2\)-\( \text{O}_2\)-\( \text{N}_2 \) or \( \text{H}_2\)O

Kuehl was the first to observe the anomalous changes in \( S_u \), using a premixed stoichiometric \( \text{H}_2\)-air flame, that occur when water vapor is substituted for part of the \( \text{N}_2 \) in air (mixtures were at 0.25 atm and initially 700 K). He used a converging rectangular nozzle technique to establish a laminar flame with essentially parallel flame fronts. Kuehl found that steam addition caused only about one-third the reduction in \( S_u \) observed with equivalent volume additions of \( \text{N}_2 \). Empirical analysis of the data clearly indicated that when steam was substituted for \( \text{N}_2 \) on a mole-for-mole basis, there was an increase in \( S_u \) which was a linear function of steam concentration; in fact, the positive slopes of interpolated data were approximately equal for six unit \( \text{O}_2 \) concentrations over the range 16 to 21 mole percent. Based partly on arguments of reduced equilibrium temperature and radical concentrations, and also on radiation heat transfer considerations, Kuehl postulated that steam accelerates burning, relative to \( \text{N}_2 \) addition, by increasing radiative heat transport from hot combustion products to water vapor in the unburnt gas.

More than twenty years later, Liu and MacFarlane measured \( S_u \) of \( \text{H}_2\)-air and \( \text{H}_2\)-air-steam mixtures as a function of temperature and composition of the unburned gases, using laser-Doppler anemometry and schlieren photography in conjunction with a constant-velocity nozzle (Mache-Hebra) burner (with 3 mm throat). A correlation equation was derived (used later in this paper) from the observed \( S_u \) data (285 measurements), covering concentration ranges of 18-65 volume percent \( \text{H}_2 \) and 0-15 volume percent steam, and an initial temperature range of 23-250 °C.

Most recently, Koroll and Mulpuru conducted a comparative study of the effects of various diluents on \( S_u \) of \( \text{H}_2\)-\( \text{O}_2 \) mixtures, and focused on the extraordinary effect of steam on \( S_u \) and flame structure. Their \( S_u \) data were obtained by the previously-used nozzle-burner (but with a more ideal 5 mm nozzle throat) cone angle method, with particle tracking by laser Doppler anemometry (velocimetry), for a wide range of stoichiometries and diluent fractions (He, Ar, \( \text{N}_2 \), and steam). The experimental technique produced an independent set of \( S_u \) measurements at 373 K for six steam concentrations and a relatively wide range of \( \text{H}_2\)-air (and \( \text{O}_2 \)) mixture ratios; the data range was extended from the previous 15% to 50% steam (inerting occurs at 55% with air, and at 79% with \( \text{O}_2 \) oxidant).
A simple correlation was obtained by Koroll and Mulpuru which accurately predicted the physical effects of inert diluents, namely flame cooling and heat transport, on burning velocity:

\[ Su(a_o/a)^{1/2} = Su(1-X/X_L), \]

where \( a \) is the thermal diffusivity, \( X \) is the mole fraction of diluent, \( X_L \) is the diluent fraction sufficient to inert the mixture, and the subscript \( o \) denotes the undiluted mixture. Whereas the correlation accurately fitted all their \( Su \) data for He, Ar, and \( N_2 \) diluted mixtures, it under-estimated their \( Su \) data for steam diluent by as much as 20%, and thus indicated that steam positively influenced the burning velocity by mechanisms other than flame cooling and heat transport.

Using a one-dimensional flame model, Koroll and Mulpuru calculated the detailed changes to flame structure caused by addition of steam, and determined that steam significantly alters the chemical kinetics and effects a redistribution of heat release in the flame. They concluded that this redistribution is caused almost entirely by the unusually high third-body efficiency of steam (6.0) in the exothermic reaction: \( H + O_2 + M \rightarrow HO_2 + M \). Because steam greatly increases the rate of this reaction (third body efficiency is higher by a factor of 15 compared to \( N_2 \) or \( O_2 \)), it also has a small additional effect on a cycle of exothermic reactions involving \( HO_2 \) with \( H \), \( OH \), and \( O \) in the preheat region of the flame. The result is shown to be a steeper temperature profile at the leading edge, a thinner flame, and an increased burning velocity. Because the postulated chemical kinetic effect appeared to account for essentially all the observed increase in \( Su \), Koroll and Mulpuru concluded that the radiant heat transport mechanism proposed by Ruehl must have a vanishingly small role.

**EXPERIMENTAL PROCEDURE**

Each OJB system (total of 3) consisted of two uniform, equal diameter (\( D \)) relatively long (50-100 \( D \)) opposed tubes. Measured mass flow rates typically ranged from 500 to 10000 \( SC\(CM \) (standard cubic centimeters per minute), and Reynolds numbers (based on \( D \)) were always less than 2000. Thus the input flows should have been laminar throughout, with a well developed parabolic velocity profile. Three sets of OJB tubes were used (2.7 mm i.d. nickel, and 5.0 and 7.0 mm i.d. pyrex) in order to include a complete range of jet velocities (strain rates) leading up to extinction of pure \( H_2 \)-air (2.7 mm). All measurements were made at atmospheric pressure in an argon-bathed plexiglass box partly open at the top (1987 set of 2.7 mm data), polycarbonate box partly open at the top (1988 set of 2.7 mm data), an alumina-silica fiberboard box with a porous sintered plate over the top (1989, 2.7 mm data), or polycarbonate box partly open at the top (7 mm). Radial movements of the tubes were mechanically restricted, but minor axial adjustments in separation distance were occasionally made (typically 2 to 3 \( D \)), which did not affect the results. The OJB systems, with tubes oriented either vertically (7 mm) or horizontally (all 3 sizes), were continuously purged with argon through a fibrous diffuser pad at the bottom of each box. Argon sometimes heated) prevented secondary combustion of excess fuel outside the central impingement zone; hence the argon bath improved visibility of the central flame, decreased local heating and buoyancy effects, and prevented attachment of a very dilute \( H_2 \) flame to the air jet.

A schematic of the flow system is shown in Fig. 3. The fuel most generally used was \( N_2 \)-diluted \( H_2 \), and the mixture was prepared using a digital mass flow metering system, using two flow meters in series for each fuel-combustible, fuel-diluent, and air flow. If the air was to be contaminated with water, the air supply flowed through a two-stage water bubbler system, equipped with stainless steel frits and immersed in a temperature-controlled bath. The air supply tube was resistively heated and insulated in order to prevent condensation of moisture within the line. The fuel tube was similarly heated to preserve continuity. The OJB tube wall temperatures were measured by thermocouples, and corresponding exit gas temperatures were usually estimated by use of prior calibrations. Final gas temperatures ranged from 85 to 100 °C (and slightly higher) for all experiments, except for the ambient temperature baseline data.

The temperature of the bubbler bath was set prior to each series of runs; the dew point of the exit flow associated with this temperature established the water content of the air mixture. An EG&G Model 911 Digital Humidity Analyzer was used under matching experimental conditions to measure the dew point and dry bulb temperature of air streams at the OJB tube exit. Dew point temperatures of the air jet were measured corresponding to the full range of bubbler bath temperatures. An analysis of the dew point data was given in Ref. 48.

For obtaining a typical pair of Blowoff and Restore data points at a fixed air contaminant level, the mass flow rate of \( H_2 \) was varied (or held at a constant value if the flow of \( N_2 \) fuel diluent was varied instead) and also the flow rate for air (plus contaminant) was simultaneously and gradually increased until Blowoff occurred; then flows were gradually decreased until Restore occurred. Critical mass flow rates of \( H_2, N_2, \) air, and contaminant were recorded, along with tube wall temperatures and (sometimes) exit gas temperatures, for both Blowoff and Restore.
CLEAN AIR, RESULTS AND DISCUSSION

Review of Early Observations using 5 mm OJB

Descriptions of flame appearance, Blowoff and Restore phenomena are given first, followed by example Blowoff/Restore data for three different OJB tube separation distances. For N2-diluted H2 opposed by air, the flame was a thin, pale blue-violet, slightly-curved dish shape (usually concave from the air side), which became fainter and flatter as X(H2) decreased. The inset in Fig. 4 contains three different prints of the same color negative (using blue, green, and red light), which shows a 5 mm OJB flame after NaCl was vaporized from a hot filament into the air jet (right tube) supply. By adjusting power to the filament and also photographic exposure, the 1 mm thick blue-violet reaction zone could be visualized inside the broader (3 mm) zone of hot gas and Na emission, which is bounded on the left (fuel side) by the curved stagnation surface. Upon reaching Blowoff, the flame shifted slightly (but suddenly) toward the fuel side as it became ring-shaped. At Restore, the flame shifted slightly (and suddenly) to the air side as it returned to its original dish-like appearance. Partly because of the low emittance of visible light at low H2 concentrations (observations required a darkened room), a CCD (charge coupled device) video camera with UV filter removed was used to image light from the near ultra violet OH bands, at wavelengths greater than 350 nm, in order to confirm naked-eye observations. Subsequent observations have shown that flame Blowoff is essentially complete in less than 2 ms; for Restore, the flame generally closes up in less than 10 ms, consistent with 'premixed-like' propagation speeds against the radially outward flow of gases.

Blowoff and Restore limits for the 5 mm OJB are shown in Fig. 4 for various axial tube separation distances (1 to 2 D). The respective Blowoff and Restore data were clearly independent of tube separation, which in other test cases (e.g. 2.7 mm OJB) was varied as much as 1 to 4 D without showing significant change.

SCHEMATIC OF OJB

Fig. 3. Schematic of Opposed Jet Burner system.

Review of Previous Characterizations of Blowoff and Restore

In the past, area-average (OJB tube) mass flux of incoming H2, M(H2), or sometimes fuel ejection velocity, U(fuel), for incoming H2 + N2 mixture, was shown as a function of H2 mole fraction in the fuel jet, X(H2). For each data set the simultaneous air input was always shown in terms of molar H2/O2 input flowrate ratio.

Previous independent data from 2.7 and 5.0 mm OJB's are shown versus X(H2) in Fig. 5, where 5 mm Blowoff M(H2)'s were linearly normalized to the 2.7 mm reference size by applying a factor of (2.7)/(5.0) to the 5 mm data in order to compare results at the same critical radial velocity gradient near the stagnation point. Whereas the Blowoff data overlapped very well after normalization, the Restore data overlapped exceptionally well without normalization. This suggested that Restore is not as strongly coupled to stagnation flow dynamics as in the case of Blowoff. Fig. 6 is the corresponding projection of the M(H2) data versus H2/O2 molar input ratio, which previously provided the only linkage between simultaneous data from the fuel and air jets.
Notice that $M(H_2)$ increased (nearly linearly) as $H_2/O_2$ ratio increased to values approaching 16, which implies that most $H_2$ in the fuel jet simply provides momentum balance for the opposing jet; from an overall system standpoint this corresponds to grossly incomplete combustion of hydrogen at high $X(H_2)$. A subsequent plot of air/fuel jet mass flux ratio, Fig. 7, confirms the 3:1 ratio Potter observed for pure $H_2$, and characterizes the full range of mass flux ratio variation with $X(H_2)$.

Recent LDA Airside Velocity Measurements and Approximation of Strain Rates

Preliminary sets of LDA data were recently obtained from three 7 mm OJB $H_2/N_2$ vs. air CFDF. From plots of the axial velocity profiles, estimations were made of the "bulk" airside strain rate, $a_{air}$, or radial velocity gradient in the vicinity of the stagnation point, based on bulk input flow. Three measures of $a_{air}$ were made directly from the axial data as follows, and values are given below for a 83% $H_2/N_2$ vs. air flame. Note that Eq. (1) for radial velocity gradient is described in applications of stagnation point boundary layer theory.

\[ a_{air} = \frac{dV}{dr} = \frac{(-dU/dx)}{2}. \quad (1) \]

Equation (2) is an expression for $a_{air}$ which stems from potential flow theory for the impingement of uniform velocity, semi-infinite jet flows at uniform temperature.

\[ a_{air} = \frac{(-dU/dx)/2}{U_{max}/(2R_{tube})} = 67.0/(2*0.35) = 95.7 \text{ s}^{-1} \quad (2) \]

\[ a_{air} = \frac{2U_{ave}/(2R_{tube})}{U_{ave}/(2R_{tube})} = 58.9/(2*0.35) = 84.1 \text{ s}^{-1} \quad (3) \]

Equation (3) is obtained from the present area-average flow rate data, i.e. $U(air)$, by assuming a parabolic axial velocity profile for the present air jet input, and a resultant axial centerline velocity equal to twice the area-average $U(air)$. Furthermore, in applying the third measure to characterize CFDF extinction, it is assumed that the relevant velocity for characterizing extinction phenomena is that which occurs in the vicinity of the stagnation streamline where the radial velocity gradient is ideally the greatest. All visual, photographic and video images to date indicate that this is a good assumption.

In summary, our recent but limited LDA data clearly demonstrate that the $H_2$-air flame resides on the air side, as predicted by earlier investigators. Before it is possible to test the one-dimensional stagnation point boundary layer assumptions generally used by CFDF modelers to determine strain rate, extensive further data will be required on both the axial and radial flowfields, for both $H_2$ and air jets.

Review of OJB Data Showing Asymptotic Limit for Clean Air Ejection Velocity

The Blowoff/Restore results in Figs. 5-7 represent the authors' original (published in 1987) set of 2.7 mm (and 5 mm) OJB $H_2$-air data, obtained over most of the usable range of $X(H_2)$. An argon-bathed OJB box, constructed of Plexiglas and partially open at the top, was used for this set. A more recent (1988) and quite independent set of data, obtained at 85 °C (nominal) using an electrically heated OJB system enclosed in a differently constructed polycarbonate box, is shown.
in Fig. 8 for $M(H_2)$ as a function of $X(H_2)$, and in Fig. 9 for $M(H_2)$ as a function of $H_2/O_2$ molar input ratio. The corresponding fuel ejection velocity is shown in Fig. 10. Note that previously derived exponential and linear least square curve fits in Figs. 8 and 9 are reasonably good; furthermore, they can be solved algebraically to show that (indirectly deduced) $U(air)$ versus $X(H_2)$ approached (approximately) a horizontal asymptotic value for pure $H_2$ (shown later). However, plots of the actual $U(air)$ data as functions of $X(H_2)$, shown in Figs. 11(a) and 11(b) for the respective 1987 and 1988 data sets, clearly indicate that a truly quasi-steady state asymptotic value was obtained for Blowoff in which $U(air)$ was effectively constant for $X(H_2) > 0.80$. Restore appears to behave similarly.

Cross plots of $U(fuel)$ versus $U(air)$, shown in Figs. 12(a) and 12(b) for the respective data sets, are even more revealing. The Blowoff curves clearly show two highly reproducible regimes: For $U(fuel)$ ranging 0 to 800 cm/s (and $0.30 \leq X(H_2) \leq 0.80$), $U(fuel)$ increases monotonically with $U(air)$, with a limiting slope of unity in the vicinity of the origin. Further increase in $U(fuel)$ causes a sharp transition to an asymptotically constant $U(air)$ value. After the transition, $U(fuel)$ more than doubled as $X(H_2)$ increased from 0.80 to 1, and $U(air)$ remained constant at 525 and 527 cm/s for the respective data sets. An alternate perspective on this highly reproducible transition of Blowoff (and Restore) data is shown in Figs. 13(a) and 13(b), where air flowrate is plotted as a function of $N_2$ diluent flowrate. The hook-shaped curve provides a unique characterization of the transition for both Blowoff and Restore. Finally, plots of air/fuel jet momentum ratios for the two sets of 2.7 mm data were not as good as those obtained with the 7 mm OJB, but the trends were quite reproducible.
In summary the tube area-average air ejection velocity, \( U(\text{air}) \), is considered more fundamental and meaningful than \( M(\text{H}_2) \), \( U(\text{fuel}) \), or \( \text{H}_2/\text{O}_2 \) input ratio for defining the extinction condition.
Empirical least-squares (LS) fits of five quasi-independent subsets of early (published in 1987) 2.7 mm extinction data for clean air were analyzed and solved algebraically to estimate critical radial strain rates for both air and fuel (30 to 100 % H₂) jets, e.g. \( \alpha_{air} = \frac{2U(air)_{ave}}{(2R_{tube})} \). Equivalent forms have conventionally been assumed and used by modelers as the basis for one dimensional stagnation point boundary layer analyses of CFDF for various hydrocarbon fuels. In the present system, where jet velocity profiles should be nearly parabolic, the assumption has been made that axial centerline velocities control extinction, and thus the centerline velocity is set equal to twice the average velocity.

The resultant LS averaged 2.7 mm \( \alpha_{fuel} \) and \( \alpha_{air} \) strain rate data, which presently are unique for the H₂-air system, are shown as open points (for \( \geq 40 \% \) H₂) in Fig. 14. Note that additional, more precise 2.7 mm results for \( \alpha_{air} \), obtained directly from the comprehensive plots of \( U(air) \) data in Figs. 11(a) and 11(b), can be included in a more precise version of Fig. 14. Finally, the very recent 7 mm strain rate results, also shown in Fig. 14 (shaded points), were derived from LS fits of extensive data. The 7mm results are considered especially reliable because the larger OJB was thoroughly characterized in three different tube orientations and was shown to be relatively ideal (i.e. larger OJB allowed adequately large flows at low strain rates, and thus led to good input jet momentum balances and absence of buoyancy effects).

Presently the \( \alpha_{fuel} \) results in Fig. 14 cannot be compared with either calculations or other data. For example existing published data, acquired for undiluted hydrocarbon flows using a cylindrical Tsuji-type burner, are not single valued at extinction in the forward stagnation region; also, extinction of H₂-rich CFDF with air has not been studied with a Tsuji burner (Personal Communication with R. Tsuji), apparently due to the very high strain rates, and very small porous cylinders, required for extinction.

Fortunately, the Fig. 14 experimental \( \alpha_{air} \) results at extinction, derived from both 2.7 and 7.0 mm OJB data, can be compared with results of a recent numerical study by Dixon-Lewis and Missaghi. As shown, the experimental \( \alpha_{air} \) results are three to five times smaller than those calculated a priori by Dixon-Lewis for input \( X(H_2) = 1.00, 0.50, \) and 0.21. Notably, the experimentally determined \( \alpha_{air} = \frac{U(air)}{R_{tube}} = 102/0.35 = 292 \text{s}^{-1} \) at \( X(H_2) = 0.21 \) is five times smaller than the \( \alpha_{air} \) calculated by Dixon-Lewis, despite the fact that fuel and air jet densities (and several other molecular properties) are very similar for 21 % H₂/N₂ versus air. Although a query regarding the factor of three difference for \( X(H_2) = 0.5 \) and 1.0 (from 2.7 mm OJB) was recently posed by one of the authors (GLP Comment in Ref. 40), reasons for the difference are unknown at present.

Finally, whereas measured \( \alpha_{fuel} \) increased by a factor of 2 over the range \( 0.8 > X(H_2) > 1 \) and \( \alpha_{air} \) was asymptotically constant in this range, the large change in \( \alpha_{fuel} \) is consistent with conservation of jet momentum, which applied approximately for the 2.7 mm OJB over a wide range of stagnation flow rates for \( 0.3 > X(H_2) > 1 \). Thus Potter's singular observation of a 3:1 mass flux imbalance for the H₂-air system \(^2\) was consistent with the present results.

**Fig. 14.** Input jet strain rates (= \( U_{ave}/R_{tube} \)) for extinction of N₂-diluted H₂ vs. air CFDF. Note numerical results by Dixon-Lewis. Exp. symbols: open, 2.7 mm; filled, 7 mm OJB.
Before the contaminated air CFDF results are presented, note that an oxygen displacement/inert gas effect occurred in all of the OJB extinction experiments when air was diluted by contaminant. The effect can be isolated and quantified by the addition of N$_2$ to air, and by using various O$_2$/N$_2$ mixtures. Nitrogen is preferred because of its predominance in air, relative chemical inertness, and also its use in premixed H$_2$-O$_2$-N$_2$-steam flames$^{4-6}$ (later comparisons). Mole fraction expressions for a simplified standard air mixture are:

\[
\begin{align*}
X(O_2) &= 0.2095 \times (1-X(\text{contaminant})), \\
X(N_2) &= (1 - 0.2095) \times (1-X(\text{contaminant})).
\end{align*}
\]

Note that 16.8 mole \% O$_2$ and 83.2\% N$_2$ are obtained when a final mixture contains 80\% dry air and 20\% added N$_2$.

Extinction of H$_2$ vs. N$_2$ + "Air"

In Fig. 15, three sets of U(air) vs. C(N$_2$) data were obtained from respective series of 2.7 mm OJB experiments at Blowoff, using pure H$_2$ opposed by N$_2$ contaminated "air" for each of three "airs" having different initial O$_2$ composition. The "airs" were laboratory service air (used exclusively to represent ambient air, assumed to contain 20.95\% O$_2$), and respective commercial cylinders (Air Products, one each) containing O$_2$/N$_2$ mixtures certified at 25.0\% and 29.9\% O$_2$. The Fig. 15 results demonstrate that O$_2$ concentration had a large effect on U(air) at flame extinction; i.e. for each "air," and also through the O$_2$ displacement effect. The linear least-squares (LS) data fits are good over the 0 to 20 \% contaminant range. Table I shows the LS (a) intercepts and (b) negative slopes normalized to 100, $D_{\text{C}\text{O}_2}$ (percent decrease in U(air) for each 1\% added N$_2$ = ($\%$ decrease)/(mole \% C added)). Respective LS slopes and intercepts are crossplotted and analyzed later (Fig. 18).

**Table I. Summary of: Least-squares intercepts for U(air)$_{C=0}$, and negative slopes, $D_{\text{C}}$, normalized to U(air)$_{C=0} = 100$.**

Thus each $D_{\text{C}}$ represents the linear sensitivity of U(air) to contaminant C in units of ($\%$ decrease)/(mole \% C added).

<table>
<thead>
<tr>
<th>Contaminant</th>
<th>Initial % O$_2$</th>
<th>20.95</th>
<th>25.0</th>
<th>29.9</th>
</tr>
</thead>
<tbody>
<tr>
<td>N$_2$, Fig. 15</td>
<td>(12/87; 4/89; 9/89)</td>
<td>D$_{N_2}$</td>
<td>1.620</td>
<td>1.451</td>
</tr>
<tr>
<td>H$_2$O, Fig. 17</td>
<td>(3/89)</td>
<td>D$_{H_2O}$</td>
<td>1.530</td>
<td>1.431</td>
</tr>
<tr>
<td>CO$_2$, Fig. 19</td>
<td>(12/87; 4/89)</td>
<td>D$_{CO_2}$</td>
<td>2.196</td>
<td>-</td>
</tr>
<tr>
<td>N$_2$, Fig. 20</td>
<td>(12/87)</td>
<td>D$_{N_2}$</td>
<td>1.727</td>
<td>-</td>
</tr>
<tr>
<td>H$_2$O, Fig. 21</td>
<td>(12/87-1/88)</td>
<td>D$_{H_2O}$</td>
<td>2.248</td>
<td>-</td>
</tr>
<tr>
<td>CO$_2$, Fig. 22</td>
<td>(12/87)</td>
<td>D$_{CO_2}$</td>
<td>2.297</td>
<td>-</td>
</tr>
<tr>
<td>N$_2$, Fig. 23</td>
<td>(summer ’86)</td>
<td>D$_{N_2}$</td>
<td>1.867</td>
<td>-</td>
</tr>
<tr>
<td>CO$_2$, Fig. 23</td>
<td>(spring ’87)</td>
<td>D$_{CO_2}$</td>
<td>2.581</td>
<td>-</td>
</tr>
<tr>
<td>clean air, Fig. 11(a)</td>
<td>(summer ’86)</td>
<td>U(air)$_{X(H_2)=1}$</td>
<td>525.0</td>
<td>-</td>
</tr>
<tr>
<td>clean air, Fig. 11(b)</td>
<td>(fall ’87)</td>
<td>U(air)$_{X(H_2)=1}$</td>
<td>526.5</td>
<td>-</td>
</tr>
</tbody>
</table>
A check on the validity of the dilution technique was made by comparing two corresponding U(air) values in Fig. 15 at equivalent final O₂ concentrations (21%). Based on LS fits of the 25% initial O₂ data, dilution to 20.95% O₂ yields U(air) = 491.3 cm/s (for C(N₂) = 16.2%); this compares closely (3% lower) with the clean air LS-intercept, U(air) = 506.3 cm/s for 20.95% O₂.

Extinction of Pure H₂ vs. N₂-Contaminated "Air" 85 and 100 deg. C, 2.7mm OJB (blowoff)

Extinction of H₂ vs. N₂ + "Air" w/Variable % O₂ 85 and 100 deg. C, 2.7mm OJB (blowoff)

Effect of O₂ Concentration on U(air) at Extinction

The N₂-contaminated air data in Fig. 15 are analyzed collectively by a linear regression of U(air) as a function of final O₂ concentration, C(O₂), as shown in Fig. 16. These data are well represented by a linear LS fit (r² = 0.972), over the range 16 - 30% O₂ in the final air mixture, by

U(air) = -285.4 + 37.59 C(O₂), cm/s

When C(O₂) is set to 20.95%, Eq. (1) gives U(air)w=N₂ = 502.0 cm/s for the clean (ambient) air intercept, and the resultant slope normalized to this intercept is 37.59/502.0 = 7.487 (% change)/(% O₂). This normalized slope is close to an early value (6.20) for the sensitivity of maximum Su of premixed H₂ + O₂ + N₂ to the concentration of O₂.⁹

Extinction of H₂ vs. Steam + "Air"

Fig. 17 shows the comparable effects of steam contamination on U(air) at Blowoff, using the same three "airs." The LS intercepts and normalized slopes, DH₂O, are given in Table I. In order to maximize the validity of subsequent data correlations for the three "airs," Blowoff data (in duplicate) were acquired sequentially for each of the three "airs" (generally within the same hour) after the bubbler bath temperature equilibrated to a given set point; in order to reduce the possibility of systematic error, successive bath set temperatures were staggered during the run series (3/89). Note that relatively small adjustments in voltage were made to resistive heating elements (coils and final OJB tubes) for both N₂, H₂O and CO₂ contaminant runs in order to compensate for the effect of fuel and "air" flowrates on (heat transfer rates) and final jet temperatures. Earlier tests indicated that OJB results were not very sensitive to these changes.

Extinction of H₂ vs. Steam + "Air" w/Variable % O₂ 100 deg. C, 2.7mm OJB (blowoff)

Comprehensive LS Empirical Fits of $U_{\text{air}}$ for $N_2$ and $H_2O$ Contaminants

In order to provide additional checks on the data, and also to obtain useful empirical fits of the results, LS intercepts and slopes from respective $N_2$ and $H_2O$ contaminant data sets (Figs. 15 and 17) were plotted in Fig. 18(a) and 18(b) as functions of initial $O_2$ concentration in the "air". Note that respective slope and intercept results from Figs. 15 and 17 also appear to be highly linear, with correlation coefficients ranging from 0.9984 to 0.9999. Based on the correlations of slopes and intercepts in Figs. 18(a) and 18(b), the following empirical equations were derived to calculate $U_{\text{air}}$ as a function of final air mole fractions $X(O_2)$, $X(H_2O)$, and $X(N_2)$ for added $N_2$:

$$U_{\text{air}, N_2} = \left(1 - 2.615 X(N_2) + 4.712 X(O_2) X(N_2)/(1 - X(N_2))\right) \times \left(-256.7 + 3624 X(O_2)/(1 - X(N_2))\right), \text{cm/s}$$ (2).

$$U_{\text{air}, H_2O} = \left(1 - 2.013 X(H_2O) + 2.317 X(O_2) X(H_2O)/(1 - X(H_2O))\right) \times \left(-357.5 + 4061 X(O_2)/(1 - X(N_2))\right), \text{cm/s}$$ (3).

When respective mole fractions for $N_2$ and $H_2O$ are set to zero in these equations, the resultant $U_{\text{air}}$ expressions for clean air agree quite closely with each other and with Eq. (1) over the range 16% to 30% $O_2$.

Extinction of $H_2$ vs. $CO_2$ + "Air"

Fig. 19 shows the effect of $CO_2$ contamination on $U_{\text{air}}$ at Blowoff for a combined data set obtained 12/87 and 4/89. Similar to that observed previously for the $N_2$ contamination set in Figs. 15, the $U_{\text{air}}$ data for $CO_2$ contamination show little scatter and belong to a common statistical population.

Extinction of Pure $H_2$ vs. $CO_2$ Contaminated Air
85 and 100 deg. C, 2.7mm OJB (blowoff)

$$U(\text{Air}), \text{Ejection Velocity, cm/sec.}$$

Fig. 19. Air jet velocity limit versus final concentration of $CO_2$ contaminant in air. Includes 12/87 and 4/89 data.
12/87 Extinction Data, H₂ vs. N₂, H₂O, and CO₂ Contaminated Air

Figures 20-22 show the respective effects of N₂, H₂O, and CO₂ air contaminants on U(air) at Blowoff for three 2.7 mm OJB data sets obtained during a one month period (12/87). Respective intercepts and normalized slopes are summarized in Table I. Note that all results in this section are designated 85 °C data. The air and fuel preheat coils, and the nickel OJB tubes, were resistively heated, but the temperature was not controlled as high as more recently (> 100 °C, 3/89 - 9/89). Occasional condensation of H₂O in the air line during the 12/87 runs had to be overcome by external heating of an aluminum block near the OJB tube end.

Extinction of Pure H₂ vs. N₂-Contaminated Air Extinction of Pure H₂ vs. Steam Contaminated Air
85 deg. C, 2.7mm OJB (blowoff) 85 deg. C, 2.7mm OJB (blowoff)

Extinction of Pure H₂ vs. CO₂ Contaminated Air
85 deg. C, 2.7mm OJB (blowoff)

Fig. 20. Air jet velocity limit versus final concentration of N₂ contaminant in air. Includes 12/87 data only.

Fig. 21. Air jet velocity limit versus final concentration of steam contaminant in air. Includes 12/87 and 1/88 data only.

Fig. 22. Air jet velocity limit versus final concentration of CO₂ contaminant in air. Includes 12/87 data only.

Extinction of N₂-Diluted H₂ vs. Contaminated Air

Our earliest studies of air contaminants focused on the extinction of 2.7 mm OJB flames formed by jets of N₂ diluted H₂ versus air containing 0 to 10 mole percent NO, N₂, CO, and CO₂. These data were originally analyzed to determine the effects of contaminant concentration on M(H₂), using exponential data fits at various X(H₂) ranging 0.6 to 1. For the present study we retrieved the original data files and plotted U(air) as a function of X(H₂). These plots (unpublished data) were very similar to Figs. 11(a) and 11(b), except that (a) asymptotic data levels for U(air) varied with air contaminant, and (b) the "apparent" U(air) = 0 intercepts on the X(H₂) axis generally became larger with increasing contaminant.
Linear LS fits of the data, for \(X(H_2) > 0.80\), were used to represent the asymptotic \(U(\text{air})\) data levels, and to calculate LS intercepts at \(X(H_2) = 1.0\). The LS intercepts, which represent \(U(\text{air})\) for pure \(H_2\), are plotted versus mole percent contaminant, \(C\), for \(N_2\) and \(CO_2\) in Fig. 23; linear LS lines were then drawn through these (previously averaged) points (5 each). The resultant intercepts, 'grand average' intercepts at \(C = 0\), and normalized slopes are tabulated in Table II.

Table II. Effects of Air Contaminants on LS-Intercept and Normalized Slope, for Extinction of \(N_2\)-Diluted \(H_2\) vs. Contaminated Air, 2.7 mm OJB.
(Data obtained on single apparatus from 4/86 to 4/87)

<table>
<thead>
<tr>
<th>Contaminant</th>
<th>(U(\text{air})_{X(H_2)=1})</th>
<th>(U(\text{air})_{C=0})</th>
<th>Normalized Slope, (\frac{dU(\text{air})}{d(C)})</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO</td>
<td>570.0</td>
<td>567.5</td>
<td>2.182</td>
</tr>
<tr>
<td>(N_2)</td>
<td>566.4</td>
<td>561.4</td>
<td>1.867</td>
</tr>
<tr>
<td>CO</td>
<td>566.4</td>
<td>559.4</td>
<td>0.941</td>
</tr>
<tr>
<td>(CO_2)</td>
<td>573.1</td>
<td>563.0</td>
<td>2.581</td>
</tr>
<tr>
<td>Clean (1987)</td>
<td>525.0</td>
<td>-----</td>
<td>-----</td>
</tr>
<tr>
<td>Clean (1988)</td>
<td>526.5</td>
<td>-----</td>
<td>-----</td>
</tr>
</tbody>
</table>

Comparison with Effects of Steam on \(H_2 + \text{air} + Su\) Combustion Limits

Fig. 24 compares the relative effects of steam on (a) \(U(\text{air})\) extinction limits for \(H_2\) vs. air + steam, to (b) laminar burning velocity \(Su\) of premixed \(H_2 + \text{air} + \text{steam}\). The comparison is based on Eq. (3) for the OJB results, and on Liu and MacFarlane's empirical fit of their \(Su\) data. To retain an "equivalent" dilution schedule for the two cases, the initial 14.7 mole % \(O_2\) (dry basis) is progressively diluted to 11.8 % \(O_2\) after steam has attained 20%. The relative effects on \(U(\text{air})\) of using 21 and 30 initial % \(O_2\) are also shown. Clearly, the relative decrease in \(Su\) due to steam addition is about 1.5 times greater than the relative decrease in \(U(\text{air})\) extinction limit up to 20% steam; and the relative decay of \(U(\text{air})\) is not very sensitive to initial % \(O_2\).

Comparison with Effects of Steam on \(H_2 + O_2 + \text{Inert Su}\) Combustion Limits

The effects of steam on \(U(\text{air})\) can also be compared with detailed \(Su\) measurements and calculations reported by Koroll and Mulpuru. For stoichiometric mixtures containing (mole basis) 50% \(H_2\), 25% \(O_2\), and either 25% \(N_2\) or 25% \(H_2O\), \(Su\) increased by 10.7% (experimental) and 10.1% (calculated) when 25% \(N_2\) was replaced by 25% \(H_2O\). These results are
compared with our OJB extinction results as follows. Use of the Eq. (3) fit of the Fig. 17 results yields a 5.0% increase in $U(\text{air})$ when $O_2$ is held constant at 25% and $H_2O$ contaminant increases from 0 to 25% (i.e. $H_2O$ displaced $N_2$). Thus the 5% increase in $U(\text{air})$ at CFDF extinction represents one-half the observed increase in burning velocity when steam is substituted for $N_2$. Based on Koroll and Mulpuru’s analysis, the increase in $S_0$ is due almost entirely to a 15 times larger third-body efficiency for $H_2O$, compared to $N_2$, for the exothermic reaction

$$H + O_2 + M \rightarrow HO_2 + M,$$

and for a subsequent cycle of exothermic low-activation energy reactions of HO$_2$ with $H$, OH and O, all of which exert a strong influence near the cool upstream boundary of the flame, where temperatures are generally below 1000 K.

EFFECTS OF STEAM ON LIMITS OF $H_2$ COMBUSTION IN AIR:

1. Extinction limit. $H_2$ vs. $O_2/N_2$ steam counterflow DF
2. Burning velocity of stoichiometric $H_2 + air + steam$

Fig. 24. Effects of Steam on Limits of $H_2$ Combustion in Air: Comparison of 2.7 mm OJB extinction limits with laminar burning velocity data from Liu and MacFarlane (1983).

Fig. 25. Reproduction of Koroll and Mulpuru's results showing effects of $N_2$ and steam diluents on burning velocity of stoichiometric $H_2-O_2$ mixtures.

Burning velocities of 2:1 $H_2-O_2$ mixtures at 373 K with the diluents $N_2(\Delta)$ and steam (O).

Fig. 26. Reproduction of Koroll and Mulpuru’s calculated flame structures for 2:1 $H_2-O_2$ mixtures at 373 K (a) containing 25% $N_2$ diluent, (b) containing 25% steam diluent.

Calculated flame structures for 2:1 $H_2-O_2$ mixtures at 373 K (a) containing 25% $N_2$ diluent, and (b) containing 25% steam diluent.
SUMMARY AND CONCLUSIONS

Coaxial tubular opposed jet burners (2.7, 5.0 and 7.0 mm i.d. OJB) were used to form dish shaped counterflow diffusion flames (CFDF), centered by opposing laminar jets of H₂ (N₂ diluted in some cases) and both clean and contaminated "air" (O₂/N₂ mixtures) in an argon bath at 1 atm. Jet velocities for flame extinction (Blowoff) and restoration (Restore) limits are shown versus wide ranges of contaminant (0 to 20 mole %) and O₂ (16% to 30%) concentrations in the air jet, and also input H₂ concentration (30% to 100%). Blowoff, a sudden breaking of CFDF to a stable ring shape, occurs in highly stretched stagnation flows and is generally believed to measure kinetically limited flame reactivity. Restore, a sudden restoration of central flame, is a relatively new phenomenon which exhibits a H₂-dependent hysteresis from Blowoff. Both 2.7 and 7.0 mm OJB jet velocities at Blowoff were converted to critical radial strain rates, \( \dot{a}_{air} \); resultant \( \dot{a}_{air} \) were 3 to 5 times smaller than recently published numerical results for 100%, 50%, and 21% H₂/N₂ fuel inputs and normal air. Whereas measured \( \dot{a}_{air} \) doubled over the range 80 to 100% H₂, \( \dot{a}_{air} \) was constant. The \( \dot{a}_{air} \) for pure H₂ is considered a valid relative measure of kinetically controlled O₂ consumption by a H₂-air CFDF just before extinction. Furthermore, \( \dot{a}_{air} \) is a more fundamental measure of CFDF extinction than previously used fuel velocity (or strain rate), which mainly satisfies jet momentum balance for centering an OJB flame.

For 25% O₂ "air" mixtures, mole-for-mole replacement of 25% N₂ contaminant by steam increased \( \dot{a}_{air} \) or "flame strength" at Blowoff by about 5%. This result is consistent with laminar burning velocity (Sₜ) results from analogous substitution of steam (25%) for N₂ in a premixed stoichiometric H₂-O₂-N₂ (or steam) flame, shown by Koroll and Mulpuru to promote a 10% increase in experimental and calculated Sₜ, due to enhanced third body (M) efficiency of water in: \( H + O₂ + M \rightarrow H₂O + M \). When the OJB results were compared with Liu and MacFarlane's experimental Sₜ of premixed stoichiometric H₂ + air + steam (where N₂ + steam > 55.7 mole %), a crossover occurred, i.e. steam enhanced OJB "flame strength" at extinction relative to Sₜ. This finding remains consistent with Koroll and Mulpuru's extensive Sₜ results, however, which show a crossover at 45% inert in stoichiometric H₂-O₂-inert flames; i.e. substitution of steam for N₂ (≥ 55.7% inert) decreases Sₜ, due to dominance of thermodynamic effects at lower O₂ levels. Overall, the relative effect of steam, compared to N₂, on extinction of H₂ vs. air + steam CFDF is about one-half the incremental effect on Sₜ of stoichiometric H₂ + O₂ + N₂ + steam mixtures over a wide range of concentrations. In contrast, CO₂ contamination significantly reduced \( \dot{a}_{air} \) at extinction relative to steam. Finally, unpublished analyses of previous OJB data on extinction of H₂/N₂ jets vs. contaminated air yielded characterizations of the \( \dot{a}_{air} \) intercept, at unity H₂ mole fraction, as functions of N₂, CO₂, and also NO, CO concentrations in air; results were generally consistent with recent data derived from pure H₂ jets.

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