THEORETICAL AND EXPERIMENTAL INVESTIGATION OF SUPERSONIC COMBUSTION

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ABSTRACT

A theoretical and experimental investigation of mixing and supersonic combustion is presented. A review of the problem under investigation is given in Part I. Then an analysis of inviscid flow fields with a finite rate chemistry for a hydrogen-air reaction is presented in Part II. In Part III, an analysis of turbulent mixing for flows with large density gradients with no chemical reaction is presented. The analysis is compared with the experimental results. In Part IV, the results of experiments in supersonic combustion are presented.

LIST OF SYMBOLS

a = radius of the jet

 $a_{\epsilon} = \text{effective area}$

A = area

 A_0 = arbitrary flow area at x = 0

 $c_p = \text{specific heat}$

D = turbulent diffusion coefficient

 G_k = quantity defined by Eq. (II-2.7)

H = total enthalpy

k = constant given by Prandtl as 0.025 for Eq. (III-4.2); mass concentration

 $K_c = \text{mass concentration at the center line}$

 $K_{c,j}$ = equilibrium constant based on molar concentration

 k_i = specific rate constant

L = number of elements

 M_i = chemical formula of the species i

 $M_i = Mach number of jet$

n = coordinate normal to streamline

N = number of species

p = pressure

r = radial coordinate of jet

 $(r_{1/2})_i$ = width of the mixing zone in the incompressible plane where the velocity equals $(u_{\text{max}} + u_c)^{1/2}$

 R_0 = universal gas constant

s = coordinate along streamline

 $S_t = Schmidt number$

T = temperature

 $U = u/u_i$

v =velocity components

V = velocity

 w_i = rate of formation of species i

W = mean molecular weight of the mixture

 W_i = molecular weight of ith species

x =coordinate along the axis of the jet

 $Y_1 = \text{mass fraction of } O_2$

 Y_2 = mass fraction of H_2

 $Y_3 = \text{mass fraction of } H_2O$

 Y_4 = mass fraction of N_2

 $Y_5 = \text{mass fraction of } O$

 Y_6 = mass fraction of H

 $Y_7 = \text{mass fraction of HO}$

 $Y_i = \text{mass fraction of } i\text{th element}$

 Y_i = element mass fraction of *i*th element

 ϵ = compressible eddy viscosity

 $\theta = \text{flow deflection}$

 μ = kinematic viscosity; Mach angle

 μ_{ij} = number of atoms of element j in chemical species i

 ν_{ij} = stoichiometric coefficients for the jth reaction

 ξ = coordinate in transformed plane

 $\rho = density$

 τ_D = characteristic time associated with diffusion

 τ_c = characteristic time associated with chemical reaction

 au_i = the time at which the computed temperature starts to rise signifi-

 $\tilde{\psi} = \frac{\text{cantly } (\mu \text{sec})}{\text{stream function}}$

Subscripts

c = center

e = external flow

inc = incompressible

j =for the jet

°K = degree Kelvin

s = stagnation

PART I. INTRODUCTION

In recent years the problem of combustion in a stream which moves at supersonic velocity has attracted the attention of several groups of investigators. One of the practical motivations for research on this problem is the hypersonic ramjet employing supersonic combustion. In the studies of the detailed processes involved in the release of chemical energy in a supersonic stream two modes of combustion have been considered: detonations which are controlled by a wave mechanism and supersonic diffusion flames which are controlled by mixing of the fuel and oxidizer.

Consider briefly these two modes; in detonation the oxidizer and fuel are completely premixed upstream of the combustor in a region of the flow wherein the static pressure and temperature are sufficiently low so that essentially no chemical reaction takes place. When the stream of premixed gases arrives at the combustor, it encounters waves which produce an increase in pressure and temperature and which initiate chemical reaction. Initially the waves are generated by the geometry of the combustor; however, the wave pattern is directly influenced by the heat release so that under steady operating conditions the geometry, upstream flow conditions and heat release are intimately related. Indeed, after steady combustion is established the geometry of the combustion could possibly be altered so that a self-sustaining wave pattern is achieved.

This detonation mode of combustion in supersonic flows has been studied by Gross and Chnitz, I-1 J. A. Nicholls, I-2 and more recently by Rhodes and coworkers. I-3, I is related from a fundamental point of view to detonation waves which have been studied for roughly 80 years (cf. Ref. I-5 for a recent, excellent review of the research pertaining thereto). While this mode of combustion is of considerable interest for the investigation of the fundamentals of chemically reacting flows in general and of chemical kinetics in particular, it seems from the available experimental results that it will be difficult to apply in an operating engine system primarily because of the aforementioned interdependence of combustor geometry and of aerodynamic and heat release effects. Indeed operating limits associated with combustion instability may be expected in an engine system employing this mode.

The second mode of combustion, the supersonic diffusion flame, involves a fuel jet in a supersonic oxidizing stream, as shown schematically in Fig. 1. The processes of mixing and chemical reaction are in general intimately connected, the reactants at first mixing and then combining so as to release chemical energy. It will be recognized that this mode is analogous to the fuel jet in low-speed flow, apparently first treated in 1928 by Burke and Schumann (Refs. I-6, I-7). As in all viscous flows involving chemical reaction it is instructive to consider two characteristic times, one associated with diffusion τ_D and one with the chemical reaction, τ_c . The diffusion time can be associated with the time required by a fluid element with the external stream velocity u_c to travel the distance in which the velocity at the axis is close to that in the external stream. The chemical time can be estimated from the kinetic rates under the assumption of infinitely fast mixing. In the treatment of low-speed fuel jets it is implicitly assumed that

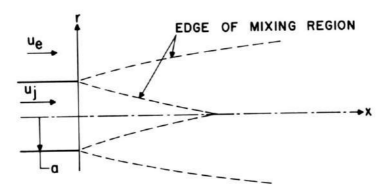


Fig. 1. Schematic representation of flame, showing coordinate system.

 $\tau_D\gg\tau_c$ so that effectively equilibrium chemistry prevails, i.e., so that diffusion is rate controlling. In a practical supersonic diffusion flame τ_D and τ_c can be of the same order for some flow conditions; therefore, in the analysis of the combustion process the interplay of chemical and fluid dynamic effects must be considered.

There is a variety of fluid mechanical and chemical problems which must be considered in achieving an understanding of the basic features of this mode of combustion. There has been under way for several years research on these problems at the Aerodynamics Laboratory of the Polytechnic Institute of Brooklyn (PIBAL) and the General Applied Science Laboratories (GASL). It is the scope and purpose of this paper to review some of the fundamental aspects of this research effort including the progress to date, and to indicate problem areas requiring additional research.

For purposes of orientation it may be instructive to consider some of the overall aspects connected with the hypersonic ramjet employing supersonic combustion. In the discussion of Ref. I-8 at the First ICAS, the first author suggested for the first time the use of deflagration or diffusion flame in a supersonic combustor. This idea was developed further in Ref. I-9, where the first author presented a detailed discussion of a hypersonic ramjet employing hydrogen as a fuel with this mode of combustion. It was shown therein that such a ramjet can produce a large thrust per unit frontal area and can operate at a high specific impulse even at hypersonic speeds. Figure 2 is taken from Ref. I-9 and provides the results significant for present purposes. Other studies of various aspects of hypersonic ramjets employing supersonic combustion are discussed in Ref. I-10, I-11.

It is fundamental to the operation of such a ramjet to decelerate the flow in an inlet from the flight Mach number to a supersonic value such that at the entrance to the mixing region the static temperatures are in the range of 1000 to 2000°K. The hydrogen fuel is injected parallel to, and in the downstream direction with respect to, the air flow. Mixing and heat release occur in the combustor; by controlling the mixing and by changing the cross-sectional area

available to the flow, the pressure distribution in the mixing region can be controlled so as to avoid shock formation. A nozzle connected to the combustor expands the combustion products to obtain gross thrust from the engine. In view of the specific impulse which can be realized with hydrogen as a fuel, research at PIBAL and GASL has been confined to the hydrogen-air system; only this system will be discussed explicitly herein although many aspects of the discussion would apply to other fuels.

The investigation of supersonic diffusion flames is complicated by the simultaneous occurrence of fluid mechanical and chemical effects. After mixing of fuel and oxidizer, reaction occurs with large heat release and, therefore, produces large changes of density and pressure. In a supersonic combustion process the pressure variation propagates in the flow through a wave mechanism in a diffused medium. A complete analysis of such processes is complex and, therefore, many simplifications are required to increase the understanding of the phenomena. For a large range of static pressure and temperatures important for practical applications, the reaction process is much faster than the diffusion process. In this case the effect of transport properties during the time in which the chemical process takes place can be neglected in the first approximation, and the concentration of all elements along a given streamline can be assumed to be constant, unaffected by the diffusion process. Then the chemical behavior along the streamlines can be considered an inviscid phenomenon. Pressure and density can change along the streamline; therefore, chemical reaction with variable pressure must be analyzed. The results of this type of analysis will indicate for what

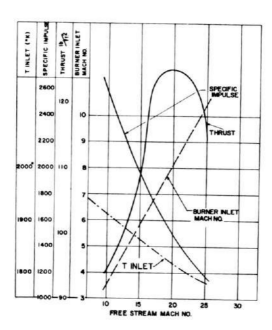


Fig. 2. Ramjet. Inlet static temperatures, thrust, and specific impulse, for various free stream Mach numbers (hydrogen fuel). Altitude 150,000 ft.

range of problems this assumption can be accepted. If this assumption is sufficiently accurate, the mixing process can also be analyzed in a simplified form by assuming that the chemical reaction is infinitely fast with respect to the mixing process. Then the flow everywhere is in chemical equilibrium; however, the mixing may occur with pressure gradients in all directions. The combustion process that is of practical interest is one which avoids formation of shock waves in the flow; therefore the analysis of heterogeneous mixing, i.e., of fuel and oxidizer, with constant pressure is of great importance. However, the mixing rate and heat release must be controlled and related to the shape of the combustor in order to achieve constant pressure.

Following this line of investigation this paper is organized as follows: Part II is devoted to the analysis of inviscid flows with finite rate chemistry, to a discussion of the mechanisms in the hydrogen-air reaction, and to the description of the relation between combustor shape and mixing process required to avoid shock formation. Part III is concerned with turbulent mixing in flows with large density gradients due to large differences in temperature and in concentration of light gases, but with either no chemical reaction or with equilibrium chemical behavior. Finally, the results of experiments, which employ the hypersonic facility of PIBAL to investigate the combustion of hydrogen in a supersonic stream, are discussed in Part IV.

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PART II. INVISCID FLOWS WITH CHEMICAL REACTION

As part of the investigation of supersonic diffusion flames with the hydrogenair system, there is considered herein the mechanisms and kinetics involved when mixing and diffusive effects are absent. These considerations correspond to the assumption, either of premixed fuel and oxidizer, or to the other limit wherein chemical reaction is so fast with respect to the diffusion process, so that the diffusive effects taking place during the combustion process can be neglected.

The problem of inviscid gas flows with nonequilibrium chemical behavior has been studied intensively in the past several years. Primary attention has been devoted to nonequilibrium air flows around bodies in hypersonic flight and through nozzles of hypersonic test facilities. Recently, nozzle flows involving combustion products of interest in high-performance propulsion devices have been considered because of the practically important differences in gross thrust associated with the limiting cases of frozen and equilibrium flow. Olson¹¹⁻¹ provides a recent review of these latter flows.

In all of the analyses of nonequilibrium behavior for propulsion the flow fields and the thermodynamic behavior of the gases have been idealized. The flow is treated either as one-dimensional with the area distribution of the nozzle specified or is considered along streamlines with the pressure distribution specified. As will be discussed in detail below, the latter idealization appears to have several advantages. With respect to the thermodynamic behavior of the combustion products, there will be assumed thermodynamic equilibrium, i.e., full rotational and vibrational equilibration of all molecules at a local translational temperature which is the same for all species. In addition, the forward and reverse rate constants are assumed to be related by the appropriate equilibrium constant. It is generally considered that these assumptions are satisfactory in view of the inaccuracy with which the mechanisms and rates associated with practical fuel-oxidizer systems are known.

REVIEW OF RECENT RESEARCH

Consider now the available results of theoretical calculations applicable to the hydrogen-air system. The fundamental paper is that of Duff¹¹⁻² who carried out calculations of the reaction profile behind a detonation wave satisfying the Chapman-Jouget condition in a hydrogen-oxygen mixture. Duff attributes the individual reaction steps and the relevant rate data to N. Davidson. This research was continued at Los Alamos by Schott et al.;^{11-3, 4} experimental research confirmed in general the overall accuracy of the chemistry. In Ref. II-5, Libby, Pergament, and Bloom presented the results of reaction histories for hydrogenair mixtures at constant pressure using the chemistry of Duff and nitrogen treated as an inert diluent.* Recently, Westenberg and Favin¹¹⁻¹ carried out one-dimensional nozzle calculations for the hydrogen-air system and for a second mixture corresponding to rocket exhaust gases. A range of pressures in the settling chamber, a single stagnation temperature of 3000°K and two equivalence

^{*} Some of the results of these and similar calculations will be discussed in detail below.

ratios were assumed. There was obtained the expected effect of pressure, namely, that the flow tends to remain near equilibrium when the pressure is high and to depart from equilibrium, indeed to freeze effectively at some point downstream of the throat, when the pressure is low. Also of interest was a result, found independently by Westenberg and Favin and by the present authors as discussed below, concerning the behavior of the intermediates, atomic hydrogen and the hydroxyl radical, under essentially frozen conditions. It is found that in an expanding flow at sufficiently low pressures so that the three-body reactions are frozen, the intermediates do not necessarily have concentrations between their frozen and equilibrium values.

Finally, Montchiloff et al.¹¹⁻¹ carried out calculations for constant area combustion of hydrogen and air. A range of initial conditions of pressure, temperature, velocity and equivalence ratio were covered although to reduce computing time, the reaction profiles were not carried to equilibrium. In this work several nitrogen-oxygen reactions leading to the formation of nitric oxide (NO) were included in the reaction steps but were found to be unimportant below 2200°K, justifying in part the assumption employed herein of nitrogen as a diluent.

As discussed in more detail below, the reaction steps and the rate constants for the above references are essentially the same as those given by Ref. II-2. The rate constants differ somewhat but probably only within the accuracy to which such data are known.†

ANALYSIS ALONG STREAMLINES—GENERAL CONSIDERATIONS

To indicate in more detail the characteristics of the hydrogen-air reaction in a flowing system and the methods of analysis for inviscid flows, the pertinent gas dynamical equations, the reaction steps, the rate constants, and some numerical results will be discussed. Consider the inviscid, steady flow along streamlines of a chemical reacting gas involving N species and L elements. The conservation of momentum, energy, species and elements can be stated as follows:

Momentum:

$$\rho V \frac{dV}{ds} = -\frac{dp}{ds} \tag{II-2.1}$$

Energy:

$$\frac{dH}{ds} = 0, \quad H \equiv (V^2/2) + \sum_{i}^{N} Y_i h_i$$
 (II-2.2)

[†] After completion of this manuscript the attention of the authors was called to the recent paper —Fowler, R. G., "A Theoretical Study of the Hydrogen-Air Reaction for Application to the Field of Supersonic Combustion," Proceedings of the 1962 Heat Transfer and Fluid Mechanics Institute, Stanford University Press. There is presented constant-density calculations employing chemical mechanisms and rates similar to those described herein.

Species:

$$\rho V \frac{dY_i}{ds} = \dot{w_i}, \quad i = 1, 2, \dots N-L$$
 (II-2.3)

Elements:

$$\frac{d\widetilde{Y}_{j}}{ds} = 0, \quad j = N - L + 1, \dots N$$

$$\widetilde{Y}_{j} \equiv \sum_{i=1}^{N} \mu_{ij} W_{j} Y_{i} / W_{i}$$
(II-2.4)

These equations are supplemented by algebraic equations: An equation of state

$$p = \rho R_0 T \sum_{i=1}^{N} Y_i / W_i = \rho R_0 T / W \qquad (II-2.5)$$

the creation terms as functions of composition and state; and the species enthalpy-temperature relations $h_i = h_i(T).\ddagger$

The form of the creation terms is

$$\dot{w}_{i} = W_{i} \sum_{j=1}^{K} (\nu_{ij}'' - \nu_{ij}') k_{j} \rho^{n_{j}} \prod_{i=1}^{N} (Y_{i}/W_{i})^{v_{ij}'} G_{j}$$
 (II-2.6)

where

$$G_{j} = 1 - \rho^{m_{j}} K_{c,j}^{-1} \prod_{i=1}^{N} (Y_{i}/W_{i})^{v_{i}j'' - v_{i}j}$$

$$m_{j} = \sum_{i=1}^{N} (\nu_{ij}'' - \nu_{ij}')$$
(II-2.7)

and where k_j is the specific is rate constant, $K_{c,j}$ the equilibrium constant based on molar concentrations, and ν_{ij} and ν_{ij} are the stoichiometric coefficients for the jth reaction

$$\sum_{i=1}^{N} \nu_{ij}' M_i \xrightarrow{k_j} \sum_{i=1}^{N} \nu_{ij}'' M_i; \qquad j = 1, \dots K$$
 (II-2.8)

Basically, these equations have been employed to date in all inviscid calculations with nonequilibrium chemical behavior. In nozzle flows, the system of equations is usually completed by the assumption of one-dimensional flow so that $\rho VA \equiv m$ with the area distribution prescribed. This approach does not permit generalization to more complex pow field calculations, for example, performed in connection with the method of characteristics. In such analyses the physical properties such as pressure and density are determined along any given

[‡] In the calculations reported here linear relations between h_i and T have been assumed. This corresponds to taking an average $c_{p,i}$ for the temperature range of interest and leads through Eq. (II-2.2) to a convenient, explicit relation for temperature $T = T(Y_i, V^2)$.

streamline, and not streamtube area. For these reasons herein the pressure distribution is specified along the streamline under consideration as suggested by Bloom and Steiger¹¹⁻⁸ in connection with nonequilibrium dissociation and recombination in external, hypersonic flows. This also completes specification of the problem, i.e., given initial conditions at a generic piont (s=0) the distribution of flow composition and state variables with respect to s can be found by integration.

It is perhaps worth while at this point to make some remarks about several numerical problems which arise in the treatment of the above equations. The number of differential equations which must be integrated with respect to the space variable s can be reduced by employing the element conservation equations [Eqs. (II-2.4)]. These can be integrated immediately and provide Lalgebraic equations among the N species, so that only N-L equations involving creation terms need be considered. However, in many calculation the mass fractions of the intermediates can be many orders of magnitudes less than the mass fractions of the major constituents. As a result the solution of the algebraic equations can involve a loss of accuracy due to the practical problem of small differences in relatively large numbers. Similar difficulties can be encountered when the flow is near-equilibrium; the creation term in Eqs. (II-2.3) represents in general the sum of differences between the forward and backward rates associated with a series of reaction steps. At equilibrium in a flowing system with pressure and temperature changes, the net result of the sum is nonzero but can involve small differences and thus a loss in accuracy. Another numerical problem is related to starting the integration in some cases; in nozzle calculations the combustion products in the settling chamber or combustor are usually assumed to be in equilibrium; but at equilibrium the creation terms must be considered indeterminant in that a large reaction rate multiplies a quantity which is the equilibrium condition, i.e., a quantity which is essentially zero. The product yields a value of w_i maintaining the species concentration at its equilibrium value at each point in the flow. Thus integration from an equilibrium state involves a singular perturbation such as has been discussed by Bloom and Ting,11-9 Hall and Russo,11-10 and Cincenti.11-11 However, it has been found possible in actual calculations as per the present report and in Refs. II-6 and II-12 to start the integration from a nominal equilibrium state with an initial step-size which is smaller than that usually required by integration accuracy. After a few integration steps, the effect of the numerically uncontrolled start is found to disappear and the solution of the differential equations becomes a well-behaved approximation to equilibrium flow, which is, of course, described by a solution to a set of algebraic equations. Finally, it is remarked that computing time with standard accuracy requirements for the selection of step-size can become excessive even for high speed computing equipment when the flow is near-equilibrium. This is due to a tendency for the solution to oscillate about the equilibrium solution.

[§] A detailed discussion of numerical difficulties associated with calculations of nonequilibrium nozzle flows is given by Emmanuel and Vincent. 11-12 The current state of development for air calculations is indicated in Ref. II-13.

THE REACTION STEPS AND REACTION RATES

The reaction steps and rate constants used here are derived from the work of Duff and co-workers. 11-2-11-4 The same chemical reactions suggested by Duff have been used independently by the several investigators cited above; thus it currently appears to represent the best available set of reactions for the hydrogen-oxygen system. Here and in Ref. II-6 nitrogen is treated as an inert diluent; this assumption as shown in Ref. II-7 is considered valid for temperatures below roughly 2500°K. It is worth mentioning at this point that the temperature range of interest in supersonic combustion appears to be below roughly 3000°K; this is within the range covered by the shock tube experiments of Schott et al. 11-3. 11-4 and only somewhat above the temperature obtained in flame experiments, e.g., Nicholls 11-14 and in laminar flame experiments, e.g., Fine. 11-14 These experiments do not verify and establish the individual reaction steps and reaction rates put have verified partially the overall consistency of the ensemble required for practical analysis.

The reaction rates considered here are as follows:

The temperatures are in ${}^{\circ}K$; the forward rate constants, i.e., for the reaction proceeding to the right are given in $(\text{mole/cc})^{-1} \sec^{-1}$ for second order reactions (1-4) and in $(\text{mole/cc})^{-2} \sec^{-1}$ for third order reactions (5-8); the symbol M is used to denote any molecule or atom serving as a third body. As will be discussed in detail below, the rate constants for reaction steps 5 and 6 play an important role at low pressures in determining the chemical history of the flow; therefore, several values thereof have been selected. Those denoted RR No. 1 and RR No. 2 represent the range of probable values estimated by Schott, 11-3 while RR No. 3 are values considered possible on the basis of more recent combustion results. The rate constants for the reverse reactions are obtained from the equilibrium constants approximated as functions of the temperature in the form $Be^{A/T}$.

To indicate the range of values of the reaction rate constants employed in related theoretical studies there are compared in Table II-1 the rate constants for reaction 5 employed therein; this reaction is selected as being an important but typical one in the array of reaction steps. It will be noted that in Refs. II-3, II-6, and II-7 a different relation between the temperature and $k_{\rm 5}$ has been assumed. However, in the temperature range of 1000 to 3000°K the numerical

TABLE II-1. REACTION RATES FOR HYDROGEN RECOMBINATION

 $(2H + M \rightarrow H_2 + M)$

Present Report	$k_5 \text{ (moles/cc)}^{-2} \text{ sec}^{-1}$
RR No. 1	1015
RR No. 2	5(1015)
RR No. 3	(1016)
Ref. II-3	1015-5(1015)
Ref. II-6	$2(10^{18})T^{-1}$
Ref. II-7	$5(10^{18})T^{-1/2}$

values given by the different expressions are within an order of magnitude; this is indicative of the accuracy with which these physical-chemical data are known.

The reaction steps described in Eqs. (II-3.1) can be separated into two groups: the second-order or "shuffling" reactions (1-4) which proceeding to the right deplete the population of molecular oxygen and hydrogen and lead to the formation of water and of the intermediates, atomic oxygen and hydrogen and the hydroxyl radical; and the third-order reactions (5-8) which proceeding to the right deplete the intermediates and lead to the formation of water and molecular species. This complex array of forward and reverse reactions makes difficult, if not impossible, a priori estimates of chemical behavior in general and of chemical times in particular.

NUMERICAL RESULTS FOR CONSTANT PRESSURE

It is instructive for an understanding of the kinetics involved in the supersonic combustion of hydrogen to consider the reaction history of the following flow model: a mixture of hydrogen and air at time zero at an initial temperature of 1000°K or higher. The pressure is assumed constant with time.

In the calculations presented here, some of these results are taken from Ref. II-5; the additional calculations have been performed by H. Pergament and will be reported "in extenso" elsewhere, the initial state of the gas is assumed to correspond to a partial equilibrium between the atomic and molecular species of oxygen and between the atomic and molecular species of hydrogen. The results are not significantly altered if the initial concentrations of the atomic species are changed somewhat; relatively large concentrations of atomic hydrogen would of course change the early phase of the reaction histories.

Consider the temperature histories for some typical cases based on this model as shown in Figs. 3a–c and as computed by a standard integration program on an IBM 7090 computer. Presented are the results for three pressures, for several equivalence ratios^{1/2} and for the three rates of reactions 5 and 6 as discussed in Part II, p. 1099 above. These temperature histories are characterized by two time periods; there is an initial time interval, usually termed the induction time and denoted τ_i , during which the temperature remains sensibly constant,

^{//} The equivalence ratio, denoted as η , is defined by the chemical equation for the reactants; namely, $O_2 + 2\eta H_2 + 3.76N_2 \rightarrow$ products.



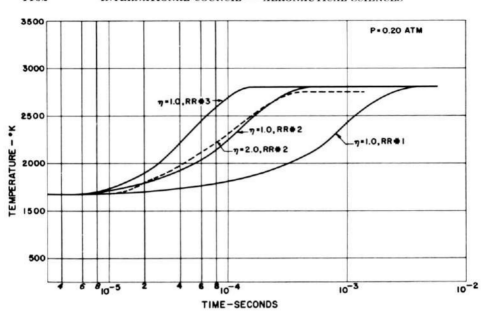


Fig. 3a. Temperature-time histories of hydrogen-air mixtures at constant pressure.

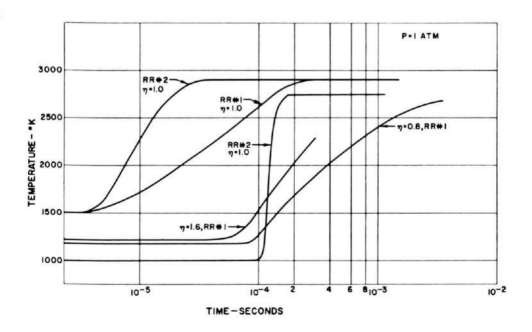


Fig. 3b. Temperature-time histories of hydrogen-air mixtures at constant pressure.

and a later phase, during which the temperature increases from its initial value T_i to the final temperature T_f ; the latter corresponds in this model to the adiabatic flame temperature (Ref. II-17). The time associated with the second phase is termed here the energy release time and is denoted τ_r ; thus $\tau_c = \tau_i + \tau_r$. Under conditions of high initial temperature and low pressure the second time interval dominates while under conditions of relatively low initial temperature (1000°K) and high pressure (4 atmospheres) the two time intervals are of roughly equal duration. This last result appears not to have been observed previously.

The changes in composition during these periods are shown in respresentative cases in Figs. 4a–f. Of greatest thermodynamic significance because of its high species enthalpy per unit mass is the behavior of the atomic hydrogen; during the induction phase the concentration of H and the other intermediates O and OH increases rapidly and reaches a maximum corresponding to mass fractions of the order of several percent or less. At the same time the molecular species, O_2 and O_2 are depleted and water is formed. However, the chemical energy released by the formation of water is effectively absorbed by the atomic hydrogen so that no signifficant change in temperature occurs. During the second time period the intermediates decay to their appropriate equilibrium values releasing chemical energy and leading to the temperature rise seen in Figs. O_2 and O_3 and O_4 are depleted and water is effectively absorbed by the atomic hydrogen so that no signifficant change in temperature occurs. During the second time period the intermediates decay to their appropriate equilibrium values releasing chemical energy and leading to the temperature rise seen in Figs. O_3 and O_4 are depleted and water is effectively absorbed by the atomic hydrogen so that no significant change in temperature occurs. During the second time period the intermediates decay to their appropriate equilibrium values releasing chemical energy and leading to the temperature rise seen in Figs. O_4 and O_4 are depleted and water is effectively absorbed by the atomic hydrogen second time period the intermediates decay to their appropriate equilibrium values releasing the second period the decay of the intermediates leads to only secondary adjustments in the concentrations of the major constituents.

This behavior is explicable in terms of the reaction steps given by Eqs. (II-3.1). During the induction phase the two-body or "shuffling" reactions are relatively fast and lead to an overproduction of the intermediates; during

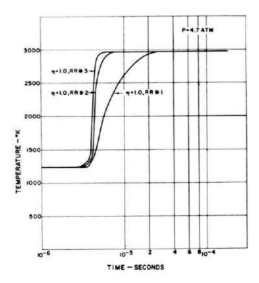


Fig. 3c. Temperature-time histories of hydrogen-air mixtures at constant pressure.

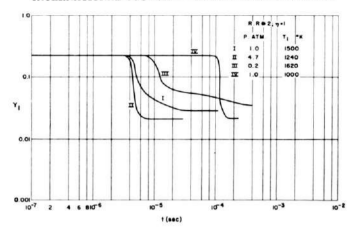


Fig. 4a. Composition-time histories of hydrogen-air mixtures at constant pressure, O2.

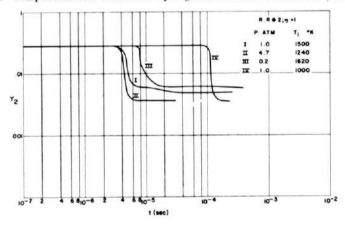


Fig. 4b. Composition-time histories of hydrogen-air mixtures at constant pressure, H2.

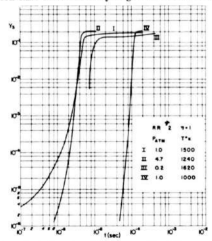


Fig. 4c. Composition-time histories of hydrogen-air mixtures at constant pressure, H2O.

the second period, the third-order reactions lead to recombination and depletion of the intermediates with the accompanying energy release. This characteristic of the hydrogen-oxygen reaction was apparently published first by Duff. ¹¹⁻² The results presented here confirm this behavior over a variety of initial conditions.

It is of interest to note that at low pressures the overall reaction time is sensitive to the recombination reaction rate, e.g., for p=0.2 atmospheres the overall reaction time is 15 times less for RR No. 3 than for RR No. 1. On the contrary, for p=4.7 atmospheres these times differ by only a factor of three. It must also be noted that for static pressure of the order of one atmosphere or higher, the energy release time is short; therefore the difference between the results for different reaction rates is not important for practical applications. However, this is not the case at low pressures.

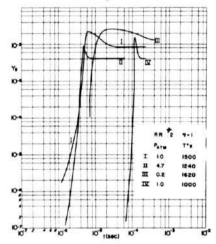


Fig. 4d. Composition-time histories of hydrogen-air mixtures at constant pressure, O.

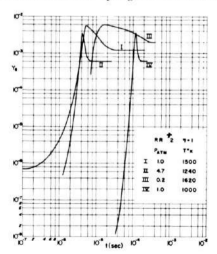


Fig. 4e. Composition-time histories of hydrogen-air mixtures at constant pressure, H.

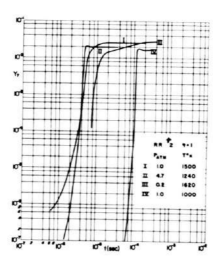


Fig. 4f. Composition-time histories of hydrogen-air mixtures at constant pressure, HO.

THE INDUCTION PERIOD

In the past considerable attention has been devoted to the time associated with the induction period; for example, Schott and Kinsey¹¹⁻⁴ and Nicholls¹¹⁻¹⁴ present the results of experiments, employing respectively techniques associated with the shock tube and standing detonation waves, to show the dependence of τ_i on pressure, temperature and composition; in both references theoretical considerations based on the reaction steps were employed to facilitate correlation of the data.

A convenient formula which can be used for estimating τ_i within a factor of two or three for mixtures with equivalence ratios on the order of unity can be obtained by simplifying the functional form suggested by Nicholls and by employing coefficients selected from the present numerical results. With τ_i defined as the time at which the computed temperature starts to rise significantly, there is obtained

$$p_{\tau_i} \cong 45(10^{-4})e^{10^4/T_i}$$
 (II-5.1)

where τ_i is in μ sec, p in atmospheres and T_i in °K. In Fig. 5 the numerical results are compared with the predictions of Eq. (II-5.1); it will be seen that within the cited factor of two or three the correlation is satisfactory.

From Figs. 3a-c and from figure 5 it will be noted that the induction time is not greatly altered by changes in the equivalence ratio nor in the reaction rates for recombination. By assuming the third-order reactions to have a negligible role and the temperature to be constant during the induction period, it is possible to make estimates of the composition at the end of the induction period. Such estimates were made by Schott¹¹⁻³ and were shown to be in good agreement with more complete calculations and with shock-tube experiments.

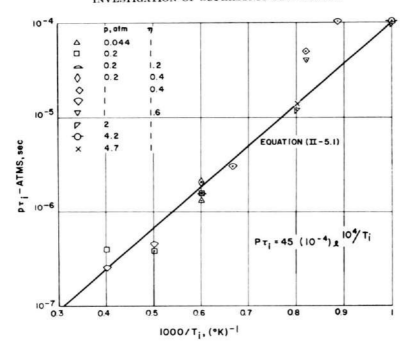


Fig. 5. Correlation of compiled induction times.

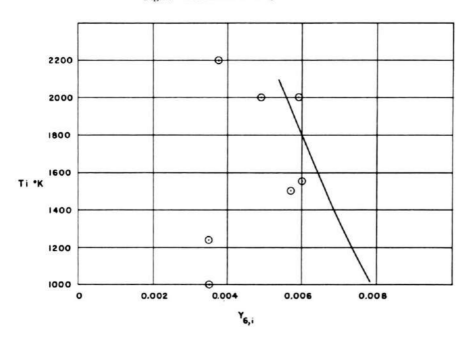


Fig. 6. Dependence of mass fraction of atomic hydrogen at the end of the induction period on initial temperature.

The calculation assumes that at the end of the induction phase there exists a partial equilibrium state corresponding to equilibrium of any three of the two-body reactions. ¹⁻⁴In addition, since these dominant reactions involve no change in moles of gas, and since the temperature is approximately constant, it can be assumed that the molecular weight of the mixture remains constant during the induction period. The resulting three equilibrium conditions, the statement concerning molecular weight, and element conservation determine the mass fractions of all species. The parameters involved in such a calculation are η and T_i ; the pressure does not enter.

In Fig. 6 the mass fractions of atomic hydrogen at the end of the induction period as computed by this scheme are shown for a range of initial temperatures. The calculations were carried out on a Bendix G-15 computer and involved solution by a combination of iteration and Newton-Raphson methods of a set of algebraic, nonlinear equations. Mr. H. Fox carried out these calculations. Also shown are the maximum values of Y_6 obtained from the computer program. The satisfactory agreement will be noted; similar agreement is found for the remaining constituents.

APPLICATION OF PARTIAL EQUILIBRIUM FOR ESTIMATES OF THE ENERGY RELEASE PERIOD

As a result of the estimates of the induction time and of the scheme described above for approximating the composition at the end of the induction period, it is possible to consider a further approximation applicable to the energy release period. The basis for the approximation resides in the coexistence during this period of second- and third-order reactions, i.e., those corresponding to reactions 1-4 and to 5-8, respectively; if it is assumed that the former are sufficiently fast relative to the latter so as to maintain during the entire second period a partial equilibrium state; an approximate description of the reaction history can be obtained by a single quadrature. It is noted that the concept of partial equilibrium can be applied only when the pressure is sufficiently high so that the assumed equilibration of the two body reactions can occur. Analyses of this type have been carried out by Westenberg and Favin 11-6 and Zinman and Romano. II-18 Care must be exercised in connection with the treatment of partial equilibrium in flowing systems in that there are introduced indeterminaties which correspond to an infinite reaction rate multiplying a zero corresponding to the equilibrium condition for the fast reaction step. The second author discussed this difficulty in general in Ref. II-19 and showed how it may be overcome.

Consider the analysis of the second period according to a partial equilibrium concept. Take for the three independent, two-body reactions those denoted 1, 2, and 3 in Eq. II-3.1 and take reactions 5 and 6 to be the most important three-body reactions.* Now the assumption that the former are relatively fast implies from Eq. (II-2.7)

$$G_1 \equiv 1 - (W_6 W_1 / W_7 W_5) K_{c,1}^{-1} (Y_7 Y_5 / Y_6 Y_1) \simeq 0$$
 (II-6.1)

[#] The analysis is not complicated in principle by the inclusion of more three-body reactions, although numerically the complexity is increased thereby.

$$G_2 \equiv 1 - (W_5 W_2 / W_7 W_6) K_{c,2}^{-1} (Y_7 Y_6 / Y_5 Y_2) \simeq 0$$
 (II-6.2)

$$G_3 \equiv 1 - (W_7 W_2 / W_3 W_6) K_{c,3}^{-1} (Y_3 Y_6 / Y_7 Y_2) \simeq 0$$
 (II-6.3)

The element mass fractions for the oxygen-hydrogen system with nitrogen treated as a diluent are from Eqs. (II-2.4)

$$\tilde{Y}_1 = Y_1 + (W_1/2W_3)Y_3 + Y_5 + (W_1/2W_7)Y_7 = \text{Const.}$$
 (II-6.4)

$$\tilde{Y}_2 = Y_2 + (W_2/W_3)Y_3 + Y_6 + (W_2/2W_7)Y_7 = \text{Const.}$$
 (II-6.5)

$$Y_4 = \text{Const.}$$
 (II-6.6)

Thus Eqs. (II-6.1)–(II-6.6) represent six equations in seven mass fractions and in the temperature and must be supplemented by a single reaction equation and by an energy equation. Since the atomic hydrogen is thermodynamically the most important intermediate, consider the species conservation thereof; prior to making the assumption of partial equilibrium, i.e., that $G_k \simeq 0$, k = 1, 2, 3, Eqs. (II-2.3), (II-2.6), and (II-2.7) for i = 6, give

$$\frac{dY_6}{dt} = \frac{\dot{w}_6}{\rho W_6} (W_6) = \frac{W_6}{\rho} \left[-k_1 (Y_6/W_6) (Y_1/W_1) \rho^2 G_1 + k_2 (Y_5/W_5) (Y_2/W_2) \rho^2 G_2 + k_3 (Y_7/W_7) (Y_2/W_2) \rho^2 G_3 \right]$$
(II-6.7)

$$-2k_{5}(Y_{6}/W_{6})^{2}(\rho^{3}/W)G_{5}-k_{6}(Y_{6}/W_{6})(Y_{7}/W_{7})(\rho^{3}/W)G_{6}]\equiv F^{-1}$$

Now with partial equilibrium assumed, the products $k_j \rho^{nj} \prod_{i=1} (Y_i/W_i)^{\nu_i j'} G_j$ must be interpreted as an infinite rate k_j multiplied by zero, namely G_j , such that partial equilibrium prevails. Accordingly, let σ_j denote these terms and consider σ_j , j=1, 2, 3 as unknowns but σ_b and σ_b as determinant values. If $G_k \simeq 0$, k=1, 2, 3, then Eqs. (II-6.1)–(II-6.3) upon total differentiation imply

$$\left(\rho \frac{dT}{dt}\right) \frac{\partial G_k}{\partial T} = -\sum_{i=1}^7 \frac{\partial G_k}{\partial Y_i} \dot{w}_i, \qquad k = 1, 2, 3$$

$$= \rho \frac{dT}{dt} \left(\frac{d}{dT} \ln K_{\epsilon,k}\right)$$
(II-6.8)

At this point it is necessary to consider the energy equation; for constant pressure flow the energy equation is simply h = constant so that

$$\rho \frac{dT}{dt} = -\left(\sum_{i=1}^{7} h_i \dot{w}_i\right) / c_p \tag{II-6.9}$$

In Eqs. (II-6.8) and (II-6.9), w_i can be expressed in terms of the σ_i 's according to

$$\dot{w}_i = W_i \sum_j (\nu_{ij}^{"} - \nu_{ij}^{"}) \sigma_j, \qquad j = 1, 2, 3, 5, 6$$
 (II-6.10)

so that Eqs. (II-6.8) and (II-6.9) yield

$$\sum_{i=1}^{7} \sum_{j} \left(\frac{\partial G_{k}}{\partial Y_{i}} - \frac{h_{i}}{c_{p}} \frac{d}{dT} \ln K_{e,k} \right) (\nu_{ij}'' - \nu_{ij}') \sigma_{j} W_{i} = 0,$$

$$j = 1, 2, 3, 5, 6$$

$$k = 1, 2, 3$$
(II-6.11)

Equations (II-6.11) permit σ_j , j = 1, 2, 3, to be expressed in terms of σ_5 and σ_6 so that the right-hand side of Eq. (II-6.7) involves only determinant values. Moreover, Eq. (II-6.7) can be integrated to yield

$$t = \tau_i + \int_{Y_{6,i}}^{Y_6} F dY_6'$$
 (II-6.12)

where the constant of integration is selected so that when $t = \tau_i$, Y_6 is equal to that given by the analysis of the induction period, namely $Y_{6,i}$.

The calculations involved in applying this analysis can be carried out on a small-scale computer; e.g., the results reported here were obtained on a Bendix G-15 computer. The procedure is as follows: The value of Y₆ varies from the maximum computed according to partial equilibrium applied to the induction period to that prevailing at equilibrium at the final temperature T_f . This latter value can be estimated by an equilibrium calculation or from available tables of the combustion products of hydrogen and air. This range of values is divided into a number of intervals appropriate to the integration of Eq. (II-6.12). With a generic value of Y_6 selected Eqs. (II-6.1)-(II-6.6) and the energy equation h = constant are employed to determine, e.g., by a combined iteration and Newton-Rapheson method, the temperature T and the mass fractions Y_i , i = 1, 2, 3, 5, and 7 corresponding thereto. Then Eqs. (II-6.11) are solved by matrix inversion to obtain σ_j , j=1,2,3; the integrand F in Eq. (II-6.12) can then be evaluated and the integration for the time t carried out. Note that when Y_6 equals the equilibrium value $F \to \infty$ so that in reality this final value does not have to be known a priori.

In Fig. 7 there is presented a comparison of the approximate and exact reaction history for one case; shown are the distributions of Y_3 , Y_6 and T and the initial conditions. It will be seen that in general the agreement is satisfactory. As a result of these considerations it can be concluded that a reasonable understanding of the interaction of the complex mechanisms involved in the hydrogenoxygen reaction is available; of course, the characterization of the reaction history into two time periods and the approximations employed in the simplified treatment of each have been deduced from more accurate calculations. It is noted that the extension of the analysis of this secton to nozzle flows according to either the streamline or one-dimensional point of view is straightforward.

COMPARISON WITH EXPERIMENT FOR NOZZLE EXPANSION

As mentioned previously there are available limited experimental results which can be used to verify the overall reaction mechanism and reaction rates employed in this and similar studies of the hydrogen-oxygen system. In addition to the cited experimental research of Schott et al. 11-3, 4 there are available the tests of nozzle flows of the combustion products of hydrogen and air carried out by Lezberg and Lancashire. 11-16 Of particular interest herein are the measurements of the temperature along the axis of the nozzle. In order to carry out a comparison between experiment and theory the pressure distribution along the axis of the nozzle had to be estimated; the approximate transonic theory for the flow in the throat of a nozzle and the measured pressure distribution along the nozzle wall were employed to obtain the estimated pressure distribution shown in Fig. 8. The composition in the settling chamber of the nozzle was taken to be close to equilibrium at the specified stagnation pressure and temperature and with the specified equivalence ratio. The equations of Sec. II, p. 1097, with the reaction mechanism of Eqs. (II-3.1) with RR No. 2 selected were integrated from upstream of the throat. The composition, velocity and temperature distributions along the axis were computed. Mr. H. Perament performed these calculations. Shown on Fig. 8 are the temperature distributions corresponding to equilibrium and frozen chemical behavior taken from Ref. II-16 and corresponding to finite rate chemistry computed here. Also shown are the two experimentally determined temperatures from Ref. II-16. In view of the approximations attendant upon the estimation of the pressure distribution and in view of the possible inaccuracies in measurement, the agreement with respect

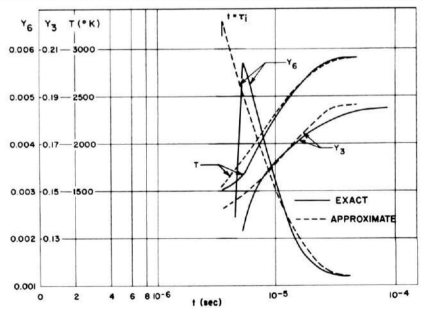


Fig. 7. Comparison of approximate and exact composition and temperature histories.

to computed and measured temperatures is considered satisfactory. Clearly additional experimental data would be valuable in validating the reaction mechanism and rates.

An interesting result concerning the concentrations of the intermediates in a nonequilibrium flow of combustion products is shown in Figs. 9a-c. Shown there are the distributions of the mass fractions of the species along the axis of the nozzle from the aforementioned calculation. It will be noted that recombination of H and O takes place in the initial part of the expansion; however, downstream of the point where the temperature deviates from the equilibrium distribution and where the mass fraction of the major constituents is frozen, the mass fractions of intermediates continues to change, of particular interest is the increase in H and the decrease in OH. This is somewhat unexpected; in the extensive studies of the expansion of dissociated air through nozzles and around bodies in hypersonic flight it was always found that the concentrations of all species were between those for complete equilibrium and complete frozen flow. The behavior here for the hydrogen-air system appears to be due to the relatively fast two-body reactions (1-4) which can maintain a partial equilibrium of the intermediates while the major constituents are essentially frozen. This indicates that perhaps the partial equilibrium analysis discussed above may provide reasonable estimates of the behavior of combustion products during nozzle expansion. As noted above, this behavior of H and OH was found independently by Westenberg and Favin. 11-6

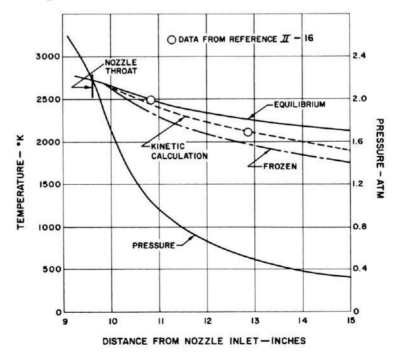


Fig. 8. Pressure and temperature distribution along the axis of nozzle.

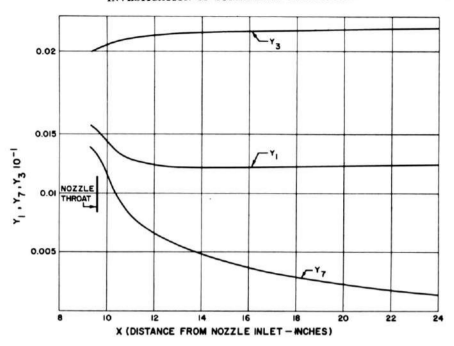


Fig. 9a. Distribution of mass fraction along the axis in nozzle flow.

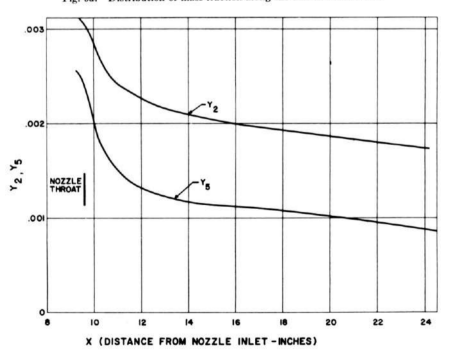


Fig. 9b. Distribution of mass fraction along the axis in nozzle flow.

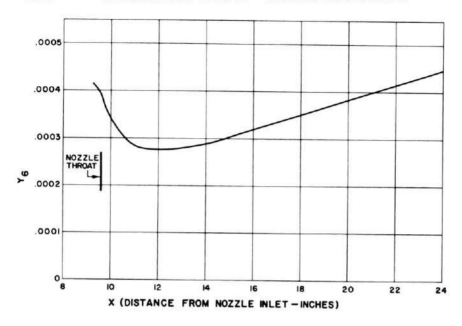


Fig. 9c. Distribution of mass fraction along the axis in nozzle flow.

PRELIMINARY CONCLUSIONS OF THE ANALYSIS FOR CONSTANT PRESSURE CHEMICAL REACTION

The results of the analysis described above for constant pressure show that the induction time decreases rapidly when static temperature and pressure increase. Such times of the order of 10^{-5} sec can be easily obtained for high Mach number flight conditions even at high altitudes. As a consequence, in a practical combustor, the ignition zone will be limited to a few inches after mixing, and the energy release zone will be of the same order for pressures between 0.5 and 1 atm. On this basis the assumption of inviscid flow for the combustion region, where large variations of pressure may occur, represents a good approximation.

For such conditions the analysis of the diffusion and combustion process along a given streamline can be divided into two parts: a mixing region where transport properties are important, and a combustion region where large variation of state occurs, but where viscous effects can be neglected. In general, both processes occur with pressure variations.

APPLICATION OF INVISCID ANALYSIS TO COMBUSTOR DESIGN

It is of interest to consider the application of the streamline analysis discussed above to actual combustor design. The objective herein is to indicate how shock waves, which could be associated with heat release, can be avoided by proper design of the combustor, i.e., by expanding the flow area in a manner compatible with the progress of combustion. For this purpose a constant pressure flow will be discussed, although similar considerations apply to the more general case of pressure varying along the streamline.

Consider a premixed hydrogen-air mixture corresponding to an equivalence ratio of unity, with a constant pressure of 4.7 atm, a uniform velocity of 11,000 ft/sec, and an initial temperature of 1,240°K. Then the numerical integration for the reaction history as discussed above can be employed to compute inter alia the spatial distribution of mass density which can be used to compute the distribution of the area ratio $(A \sim \rho^{-1})$ required to maintain constant pressure flow. The area ratio A/A_0 where A_0 is an arbitrary flow area at x=0 is shown in Fig. 10; it will be seen that there is a relatively small region in which the density decreases rapidly and in which the flow area must increase if constant pressure is to be maintained. The total increase in cross-sectional area is 2.15. Note that if a different analysis, one based on constant area were employed, a significant pressure rise and velocity decrease would occur in the region of rapid heat release; this is shown, e.g., in Ref. II-7.

Now consider Fig. 11 which indicates how this result can be employed qualitatively in the design of a practical two-dimensional combustor. Assume that the total inlet flow area is divided into streamtubes and that the mixing between hydrogen and air is controlled; in particular, assume that the hydrogen can be distributed by mixing nonuniformly across the inlet to the combustor as would be the case if a "swept" injector were provided. Now select the length AB to correspond to the length in which no significant change in area ratio is required; at B turn the flow by an angle α so that at C the original streamtube has increased in area by a factor of 2.15, where the length AC corresponds to the length obtained from Fig. 10, beyond which the area is again essentially constant.

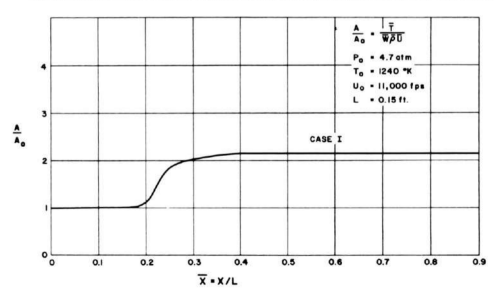


Fig. 10. Distribution of flow area for constant-pressure combustion.

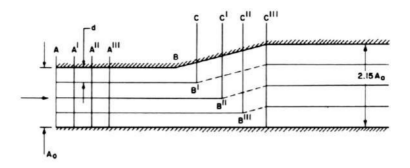


Fig. 11. Schematic representation of a constant-pressure combustion.

Note that $\alpha \simeq 1.15 (d/\Delta_z)$ where d is the original streamtube height and Δ_z is the length over which the change in cross-sectional area is effectively required. Now for this second streamtube assume that the origin for the chemical history is shifted by the amount BC so that its required increase in cross-sectional area occurs downstream of point C and is complete in the length $\Delta_z = B'C' = BC$. If this process is carried out for each streamtube, there will be obtained the duct configuration which is shown in Fig. 11 and which involves a final cross-sectional area 2.15 times greater than the original one. It will be recognized that this treatment corresponds to a rough streamtube analysis for a combustor. This analysis can be improved and refined by a two-dimensional treatment of the flow field with finite-rate chemistry. An analysis thereof is discussed in the following subsection.

METHOD OF CHARACTERISTICS FOR TWO-DIMENSIONAL FLOWS

Consider now the analysis by the method of characteristics of two-dimensional supersonic flows with finite-rate chemistry.** It will be shown that the streamline analysis forms an integral part of such a method. Although for practical reasons, two-dimensional flow is considered explicitly herein, the extension to the axisymmetric case is straightforward. For completeness some of the equations of Part II, p. 1097, are repeated here.

The equations of motion in coordinates along a streamline s and in the normal direction n are

$$\rho V \frac{\partial V}{\partial s} + \frac{\partial p}{\partial s} = 0 \tag{II-10.1}$$

$$\rho V^2 \frac{\partial \theta}{\partial s} + \frac{\partial p}{\partial n} = 0 \tag{II-10.2}$$

^{**} The method of characteristics for nonequilibrium gas flows corresponding to either dissociation or vibrational relaxation has been treated in Refs. II-20 to II-23. The analysis here is more general.

The equations of species conservation are

$$\rho V \frac{\partial Y_i}{\partial s} = \dot{w}_i \tag{II-10.3}$$

and the overall continuity equation for two-dimensional flows is

$$\frac{\partial}{\partial s} (\rho V) + \rho V \frac{\partial \theta}{\partial n} = 0 \tag{II-10.4}$$

These equations are supplemented by the requirements of energy conservation [Eq. (II-2.2)]; by the equation of state [Eq. (II-2.5)]; by the description of the creation terms [Eqs. (II-2.6) and (II-2.7)]; and finally by the enthalpy-temperature relations for the species.

Now following the usual procedure in the method of characteristics (cf., e.g., Refs. II-24 and II-25). Eqs. (II-2.2) and II-2.5) are differentiated with respect to s; the resulting equations and Eqs. (II-10.1) and (II-10.4) can be used to obtain

$$\frac{1}{p}\frac{\partial p}{\partial s}\left[\left(1-\frac{R_0}{c_pW}\right)-\frac{R_0T}{V^2W}\right]+\frac{\partial \theta}{\partial n}=\frac{W}{\rho V}\sum_{i=1}^N\frac{\dot{w}_i}{W_i}\left(1-\frac{h_iW_i}{c_pTW}\right)\equiv F \quad (\text{II-10.5})$$

which involves only the partial derivatives $\partial p/\partial s$ and $\partial \theta/\partial n$ and finite quantities. As usual there are considered the total differentials

$$\frac{1}{p}\frac{dp}{ds} = \frac{1}{p}\frac{\partial p}{\partial s} + \frac{1}{p}\frac{\partial p}{\partial n}\left(\frac{dn}{ds}\right)$$
 (II-10.6)

$$\frac{d\theta}{ds} = \frac{\partial \theta}{\partial s} + \frac{\partial \theta}{\partial n} \left(\frac{dn}{ds} \right)$$
 (II-10.7)

The Eqs. (II-10.2) and (II-10.5)–(II-10.7) are formally considered to yield the four partial derivatives in terms of finite quantities and of $d \ln p/ds$, $d\theta/ds$ and dn/ds. Select dn/ds so that the determinant of these equations vanishes; there are obtained two values of dn/ds; namely,

$$\frac{dn}{ds} = \pm \left[\frac{V^2}{T} \frac{W}{R_0} \left(1 - \frac{R_0}{W c_p} \right) - 1 \right]^{-1/2}$$
 (II-10.8)

The condition for dn/ds to be real is evident. Moreover, the degeneration of this relation to the special case usually treated in gas dynamics, i.e., constant composition and constant specific heats, is also evident. Finally, it is interesting to note that the speed of sound, which arises quite naturally from the analysis, is associated with frozen composition and frozen vibrational energy: namely,

$$a_{\ell}^{2} = (R_{0}T/W) [1 - (R/c_{p}W)]^{-1}$$
 (II-10.9)

Additional restrictive assumptions must be made a priori in order that there enter the analysis other limiting speeds of sound, i.e., those associated with equilibrium behavior in some sense.

Denote the values of dn/ds corresponding to the plus and minus signs by $(dn/ds)_{\rm I}$ and $(dn/ds)_{\rm II}$, respectively; introduce $\mu_{\rm I}$ and $\mu_{\rm II}$ where

$$\tan \mu_{\rm I} = (dn/ds)_{\rm I}$$

$$\tan \mu_{\rm II} = (dn/ds)_{\rm II}$$
(II-10.10)

The vanishing of the determinant leading to the special values of dn/ds implies along the line $(dn/ds)_{\rm I}$ that

$$\frac{1}{p}\frac{dp}{d\xi} + \left[\frac{V^2W}{RT}\left(\frac{dn}{ds}\right)_{\rm I}\right]\frac{d\theta}{d\xi} = \left(\frac{V^2W}{RT}\right)\left[\left(\frac{dn}{ds}\right)_{\rm I}\right]^2 \\
\left\{1 + \left[\left(\frac{dn}{ds}\right)_{\rm I}\right]^2\right\}^{1/2} F = \left(\frac{V^2W}{RT}\right)F\frac{\sin^2\mu_{\rm I}}{\cos^3\mu_{\rm I}} \tag{II-10.11}$$

and along the line $(dn/ds)_{II}$ that

$$\frac{1}{p}\frac{dp}{d\eta} + \left[\frac{V^2W}{RT}\left(\frac{dn}{ds}\right)_{II}\right]\frac{d\theta}{d\eta} = \left(\frac{V^2W}{RT}\right)\left[\left(\frac{dn}{ds}\right)_{II}\right]^2 \\
\left\{1 + \left[\left(\frac{dn}{ds}\right)_{II}\right]^2\right\}^{-1/2} = \frac{V^2W}{RT}F\frac{\sin^2\mu_{II}}{\cos^3\mu_{II}} \tag{II-10.12}$$

where $d\xi$ and $d\eta$ are differential elements along each characteristic line. In the nonreactive case $F \equiv 0$ so that Eq. (II-10.11) and (II-10.12) reduce to those usually prevailing in gas dynamics. The right-hand sides of Eqs. (II-10.11) and (II-10.12) are equivalent to the terms which arise due to rotationality. In addition to the two characteristic lines related to $\mu_{\rm I}$ and $\mu_{\rm II}$ the streamlines are also characteristics along which the reation equations must be applied.

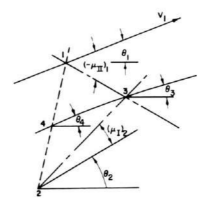


Fig. 12. Unit problem for method of characteristics.

The application of these equations to a unit problem within the flow field as shown in Fig. 12 follows the usual numerical procedure. Assume that at two adjacent points 1 and 2 all the flow and state variables are known. Then from Eqs. (II-10.11) and (II-10.12) in the usual way two finite difference equations for p_3 and θ_3 can be established provided the finite quantities in these equations are evaluated from conditions at points 2 and 1 respectively. Thus the location of, and the values of p and θ at point 3 can be found to a first approximation. Now construct point 4 on the streamline, inclined at θ_3 passing through point 3, and at θ_4 passing through 4, where linear interpolation between θ_2 and θ_1 is assumed; further, by linear interpolation estimate V, p and Y_i, $i = 1, 2 \dots N - 1$ at point 4. Thus, if the pressure is assumed to vary linearly from points 4 to point 3, these estimates plus the energy equation permits the conservation Eqs. (II-10.1) and (II-10.3), and the supplemental equations relative to the state, kinetics and species enthalpy-temperature to be integrated along s to yield the composition at point 3 in the same fashion as in the usual streamline calculations described above. Iteration can also be used if required.

It is pointed out that the procedure outlined here assumes that requirements for accuracy in integration for the species conservation equations along the streamline are more severe than the requirements for accuracy in the flow variables p, θ , V, etc. If this is not the case, then finite difference approximations for integration along the streamline from point 4 can be readily employed.

In addition to the unit problem for the flow field, special unit problems for points involving shock wave and for boundary points must be established. For the flow across shock fronts use of the von Newman-Doring-Zeldovitch jump conditions would be consistent with the inviscid flow assumption. No special difficulty with respect to these special unit problems would appear likely in principle, but as in the method of characteristics for classical gas dynamics, difficulties associated with secondary shocks, numerical accuracy, programming logic, etc., are certain to arise and to require solution. An analysis employing the method of characteristics with finite-rate chemistry along the lines described here will be valuable in combustor and exhaust-nozzle design and analysis.

It is interesting to note that the effects of viscosity, diffusion and heat conduction normal to the streamlines can be incorporated in the above analysis in an approximate manner, if these effects are slowly varying with respect to combustion processes. For example, in the species conservation equations the effect of diffusion normal to the streamlines is to add an additional term $\partial/\partial n(\rho D\partial Y_i/\partial n)$; an average value thereof for the increment along the streamline from point 4 to point 3 can be evaluated in finite-difference form from a knowledge of the state at points 1, 2, and 4. Similarly, the energy equation, which in this case must be treated in differential form, will have an additional term arising from energy transport due to conduction and to diffusion and a dissipation term due to viscosity; again a finite-difference representation can be employed. Finally, the viscosity contribution to the streamwise momentum equation can be taken into account. For laminar flow the transport coefficients are clearly defined, while, as is well known, for turbulent flow effective exchange coefficients must be introduced.

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PART III. HETEROGENEOUS TURBULENT MIXING

It appears that under flight conditions for hypersonic vehicles of practical sizes, the mixing process between the fuel jet and the air stream will be predominantly turbulent. Accordingly, the present discussion of the mixing problems connected with supersonic diffusion flames will be exclusively devoted to turbulent flows. It will be realized that at the origin of mixing there will always exist on the fuel injector, boundary layers, one in air and one in the hydrogen fuel; in considering only turbulent processes, it is assumed that either one of these boundary layers is turbulent at the origin of mixing or that transition from laminar to turbulent flow occurs in a negligibly short distance downstream from the origin of mixing.

GENERAL REMARKS

The mixing processes in supersonic diffusion flames play an important role in determining the characteristics of a combustor, since the fuel and oxidizer must be brought together by mixing prior to chemical reaction. As mentioned in the Introduction the relative rates of the mixing and of the chemical processes taking place simultaneously in the fuel jet determine the characteristics of the flow; if the chemical reactions are relatively fast, i.e., for the hydrogen-air system of interest herein, if the initial static temperatures and the static pressures are high, then diffusion will be rate controlling. In a limiting sense under these conditions the behavior corresponds to local chemical equilibrium and heat release occurs close to the exit of the fuel injector. If, on the contrary, the chemical reaction is relatively slow, the mixing of the fuel and air close to the exit will correspond to a heterogeneous, nonreactive jet; far downstream the fuel and oxidizer will be completely mixed and if conditions are suitable will react in an essentially inviscid manner. This second case in a limiting sense corresponds to "frozen" mixing.

The two limiting cases of equilibrium and frozen chemical behavior are the easiest to treat analytically so that most analyses of viscous reacting flows have considered only such cases. However, for combustor design it is necessary to determine quantitatively the conditions under which the two limiting cases prevail and the characteristics of flows intermediate to the limiting cases, i.e., of flows with finite-rate chemistry. In the analysis of mixing with chemical reactions the interaction between mixing, combustion and boundary conditions introduces severe complications. The combustion process locally affects temperature, density, and pressure of the flow. If the flow is supersonic pressure variations propagate in the flow through a wave mechanism, and local changes of static pressure can influence reaction rates and mixing rates elsewhere in the flow. These coupling effects can be important in practical applications. In the case of supersonic diffusion flames the mixing process is the controlling mechanism and a strong increase in pressure must be avoided by careful selection of the boundary conditions. Therefore it appears logical to analyze in the first step

mixing problems where the pressure can be assumed to be practically constant through the entire flow field.

Even in the case of a constant pressure process complications are introduced by several effects; e.g., by the details of the flow at the origin of mixing; in the boundary layer on the air side of the fuel injector there will exist regions of low velocity and high static temperature because of aerodynamic heating. Now it is known from analyses such as those presented in the previous section that the reaction history for hydrogen-air mixtures is sensitive to initial temperature as well as to static pressure. Thus the chemical times and lengths associated with the static temperatures and velocities near the exit of the fuel jet may actually be several orders of magnitude smaller than those based on the static temperatures and velocities in the streams external to the viscous region. If this is the case, reaction may initiate close to the fuel jet and spread rapidly so as to cause extensive amounts of heat release in flow distances shorter than those expected on the basis of the usual overall considerations of diffusion and chemical times for idealized conditions.

In addition to the effect of initial boundary layers the conditions under which limiting chemical behavior can be expected, are complicated by the wide range of flow velocities which can occur in supersonic diffusion flames of practical interest. For example, consider the case wherein the fuel jet has a small velocity compared to that in the external stream. In this case chemical reaction may first appear near the axis of symmetry, where the lowest velocities occur, and may spread rapidly outward in contrast to the previous discussion.

The problem of treating analytically the simultaneous effects of turbulent mixing and finite-rate chemistry is formidable if some realism is desired. The preliminary research reported here considers first the turbulent mixing of heterogeneous gas streams. Both theoretical and experimental research will be described. In addition, some remarks concerning the chemical kinetics of turbulent reacting flows will be made to indicate an essential difficulty in proceeding to the analysis of practically interesting cases wherein the effects of finite-rate chemistry are important; clearly these cases are the intermediate ones cited above.

The problem of describing quantitatively turbulent-shear flows has occupied the attention of fluid mechanicians for many years. It is generally agreed that even for the simplest case of incompressible flow the available knowledge is incomplete. With respect to turbulent-shear flows involving significant gradients of density the present state of knowledge is poor. In view of this situation research directed toward establishing some of the engineering fundamentals of turbulent mixing of heterogeneous gases has been undertaken at PIBAL and at GASL. The flow configuration has been idealized in both theory and experiment to correspond to an axisymmetric fuel injection in an essentially infinite oxidizing stream. In a practical combustor the presence of walls, of other asymmetries and of nonuniformities in the oxidizing stream, e.g., due to gradients in stagnation pressure, will complicate the flow so that only a general but essential understanding of the phenomena can be expected from the study of idealized configurations such as are considered herein.

THEORETICAL ANALYSIS—GENERAL EQUATIONS

Consider first an engineering analysis of a turbulent jet of a fuel such as hydrogen in an air stream; assume that the mean turbulent quantities are described by the equations for laminar flow if the transport coefficients replace their molecular counterparts. Then (cf. Refs. III-1 and III-2) for constant pressure flows the following conservation equations are obtained:

Conservation of Momentum

$$\rho u \frac{\partial u}{\partial x} + \rho v \frac{\partial u}{\partial r} = r^{-1} \frac{\partial}{\partial r} \left(\epsilon \rho r \frac{\partial u}{\partial r} \right)$$
 (III-2.1)

Conservation of Energy

$$\rho u \frac{\partial H}{\partial x} + \rho v \frac{\partial H}{\partial r} = r^{-1} \frac{\partial}{\partial r} \left[\left(\rho \epsilon p_{t}^{-1} r \frac{\partial H}{\partial r} \right) + (p_{t} - 1) \rho \epsilon p_{t}^{-1} r u \frac{\partial u}{\partial r} + (L_{\epsilon_{t}} - 1) \epsilon \rho r p_{t}^{-1} \sum_{i=1}^{N} h_{i} \frac{\partial Y_{i}}{\partial r} \right]$$
(III-2.2)

Global Conservation of Mass

$$\frac{\partial}{\partial x} (\rho u) + r^{-1} \frac{\partial}{\partial r} (\rho v r) = 0$$
 (III-2.3)

Species Conservation

$$\rho u \frac{\partial Y_i}{\partial x} + \rho v \frac{\partial Y_i}{\partial r} = r^{-1} \frac{\partial}{\partial r} \left(\rho \epsilon S_t^{-1} r \frac{\partial Y_i}{\partial r} \right) + \dot{w}_i, \quad i = 1, 2, -N - L \text{ (III-2.4)}$$

$$\rho u \frac{\partial \widetilde{Y}_{j}}{\partial x} + \rho v \frac{\partial \widetilde{Y}_{j}}{\partial r} = r^{-1} \frac{\partial}{\partial r} \left(\rho \epsilon S_{t}^{-1} r \frac{\partial \widetilde{Y}_{j}}{\partial r} \right), \quad j = 1, 2, -L \quad (III-2.5)$$

Eqs. (III-2.1) through (III-2.5) are derived from the general equations of turbulent motion by introducing the concept of eddy viscosities that permits to determine the contribution of the fluctuating quantities of the flow as a function of the local gradients of the average properties. On this basis the transport properties due to the turbulent motion can be expressed in the same form as the transport properties due to the molecular motion. However, the coefficients of the terms that represent the molecular motion are functions only of local physical properties of the gas, while the coefficients related to the turbulent motion usually are related to gradients of physical properties of the gas. When the contribution due to molecular motion is small and can be neglected, then the eddy viscosities are the same for all the transport properties; this corresponds to the turbulent Prandtl and Schmidt number equal to unity. In the case that the contribution of the molecular transport is important and must be retained, then it is represented in the equations in an average form, by introducing correcting factors to the different transport properties. This is done by

assuming that the turbulent Prandtl and Schmidt numbers are different from unity but constant at any given cross section of the flow. On the equations presented, a single turbulent diffusion coefficient D corresponding to a single turbulent Schmidt number $S_t = \epsilon/D$ has been used.

Fundamental to the application of these equations to the problem of a turbulent jet is a requirement to describe, at least approximately, the compressible eddy viscosity ϵ . In the past theoretical analyses of turbulent compressible mixing (see Refs. III-3 and III-4) have carried over the Prandtl hypothesis generally accepted for incompressible flow—namely, that the eddy viscosity is essentially uniform across the mixing region and varies at most with the streamwise coordinate.

A different analysis for compressible mixing problems has been presented by the second author in Ref. III-1. In the analytical work presented here the approach presented in Ref. III-1 will be followed and the same formal dependence of the eddy viscosity on spatial coordinates introduced in Ref. III-1 will be used. However, a different expression that relates the eddy viscosity to the physical properties of flow will be introduced. At the same time, more general solutions will be obtained and simplifications will be presented. The results of such analyses will be compared with some experimental results.

ANALYSIS OF MIXING

Consider a jet of gas exhausting into an infinite stream, in general of foreign composition. Assume that the gases are nonreactive. Now following Refs. III-1 and III-5, introduce a stream function ψ so that

$$\rho ur = \rho_i u_j \widetilde{\psi} \frac{\partial \widetilde{\psi}}{\partial r}$$
 (III-3.1)

$$\rho vr = -\rho_{j}u_{j}\widetilde{\psi}\frac{\partial\widetilde{\psi}}{\partial x}$$
 (III-3.2)

and employ the von Mises transformation, i.e., $(x, r) \rightarrow (x, \psi)$, then Eq. (III-2.1) becomes without approximation

$$\frac{\partial u}{\partial x} = (u_j \widetilde{\psi})^{-1} \frac{\partial}{\partial \psi} \left[\left(\frac{\epsilon \rho^2 r^2 u}{\rho_j^2 u_j \psi^2} \right) \widetilde{\psi} \frac{\partial u}{\partial \psi} \right]$$
 (III-3.3)

where from Eq. (III-3.1)

$$r^{2} = \int_{0}^{\widetilde{\psi}} \left(\frac{2\rho_{j}u_{j}}{\rho u} \right) \psi d\psi \qquad (III-3.4)$$

and at the axis or at infinity $\partial(\rho u)/\partial\psi=0$. Therefore near the axis

$$r^2 = \frac{\rho_j u_j}{\rho u} \widetilde{\psi}^2 \tag{III-3.5}$$

and

$$\frac{\epsilon \rho^2 r^2 u}{\rho_i^2 u_i \psi^2} = \epsilon \rho \simeq (\epsilon \rho)_c \tag{III-3.6}$$

Assume $(\epsilon \rho)_c = f(x)$, then near the axis $\epsilon \rho^2 r^2 u/\rho_j^2 u_j \psi^2 = f(x)$ and Eq. (III-3.3) becomes

$$\frac{\partial u}{\partial x} = \frac{(u_j)}{\psi} \frac{(\epsilon \rho)_c}{\rho_j} \frac{\partial}{\partial \psi} \left(\widetilde{\psi} \frac{\partial u}{\partial \psi} \right)$$
 (III-3.7)

if $U = u/u_j$ and

$$\xi = \int_0^x \frac{(\epsilon \rho)_c}{\rho_i u_i a} dx$$
 (III-3.8)

then

$$\frac{\partial U}{\partial \xi} = \frac{a}{\psi} \frac{\partial}{\partial \psi} \left(\widetilde{\psi} \frac{\partial U}{\partial \psi} \right) \tag{III-3.9}$$

The expression [Eq. (III-3.6)] is accurate near and far away from the axis, and is only an approximate expression moving away from the axis. The error introduced is due to the fact that terms of the order $\{\partial^2(\rho u)/\partial\psi^2\}\psi^2$ are neglected with respect to one in Eq. (III-3.5). Indeed

$$\rho u = (\rho u)_{\epsilon} \left[1 + \frac{1}{4} \frac{1}{(\rho u)_{\epsilon}} \left(\frac{\partial^{2} (\rho u)}{\partial \psi^{2}} \right)_{\epsilon} \widetilde{\psi}^{2} + \dots \right]$$
 (III-3.10)

Then, if the second term is neglected in Eq. (III-3.5), Eq. (III-3.6) gives

$$\epsilon \rho^2 u r^2 \sim (\epsilon \rho)_{\epsilon} \widetilde{\psi}^2 = f(x) \widetilde{\psi}^2$$
 (III-3.11)

This expression leads to Eq. (III-3.9). The solution of Eq. (III-3.9) has been indicated in Refs. III-1 and III-5, and is independent of the energy equation.

EVALUATION OF THE EDDY VISCOSITY

In Ref. III-1 the suggestion has been made that the expression

$$\rho_{e}\bar{\epsilon} = k(u_{\text{max}} - u_{\text{min}})\rho_{e}(r_{1/2})_{i}$$

be used for definition of the eddy viscosity ϵ , where $(r_{1/2})_i$ is the width of the mixing zone in the incompressible plane wherein the velocity changes from u_{max} to $(u_{\text{max}} + u_c)1/2$. The value suggested for the constant k was 0.025 as given by Prandtl, then

$$\rho_{e}\bar{\epsilon} = 0.025u_{e} \mid 1 - u_{e} \mid (r_{1/2})_{inc} \rho_{e}$$
 (III-4.1)

This expression indicates that the turbulence would decay when the two jets have the same velocities even if the two streams have different temperature or composition. This is not consistent with the fact that in these cases a dissipative mechanism still exists due to heat conduction or concentration changes that would sustain turbulence. It would also indicate that a sharp decrease of mixing rates would occur for $u_j/u_e \sim 1$ with rapid changes of mixing rates in both sides of the conditions $u_j/u_e \gtrsim 1$. A modification of the expression for the eddy viscosity has been suggested by the first author in Ref. III-6. The expression used here has the form

$$\rho \epsilon = (\rho \epsilon)_c = k r_{1/2} | \rho_{\epsilon} u_{\epsilon} - \rho_{c} u_{c} | \qquad (III-4.2)$$

where k is the constant given by Prandtl and is assumed to be equal to 0.025. This expression still follows the basic concept of mixing length of Prandtl and the basic concept of Ref. III-1 with respect to the dependency of eddy viscosity on special coordinates. In this expression the quantity $r_{1/2}$ is the width of the mixing zone in the physical plane, defined as the value where the quantity ρu has an average value between the value at the axis and the value in the outside flow. Therefore

$$(\rho u)_{r_{1/2}} = (\rho_e u_e + \rho_e u_e)^{1/2}$$
 (III-4.3)

It has been found that this expression is in better agreement with the experimental results presented here.

THE FREE JET IN A QUIESCENT ATMOSPHERE

Consider next the problem of free jet in a quiescent atmosphere. This problem is of interest because it has been investigated experimentally for a large variety of cases. Assume that the boundary layer at the origin of mixing $(x = \xi = 0)$, is negligible. Then $U(\xi, \psi)$ is to satisfy the conditions.

$$U (0, \tilde{\psi} = 1 \quad 0 < \tilde{\psi} < a$$

$$= u_{\epsilon}/u_{j} \quad \tilde{\psi} > a \qquad (III-5.1)$$

$$\lim_{\widetilde{\psi} = a} U (\xi, \tilde{\psi}) = u_{\epsilon}/u_{j}$$

The solution of the problem specified by Eqs. (III-3.11) and (III-5.1) is a standard one in the theory of heat conduction (cf. Refs. III-7 and III-8). The solution is given by the equation

$$\widetilde{U} = \frac{[U - (u_{\epsilon}/u_{j})]}{[1 - (u_{\epsilon}/u_{j})]} = (2\xi/a)^{-1} \exp\left[-(\widetilde{\psi}/a)^{2}/4(\xi/a)\right]$$

$$\int_{0}^{1} \exp\left[-r'^{2}/4(\xi/a)\right] I_{0} \left[r'(\widetilde{\psi}/a)(2\xi/a)^{-1}\right] r' dr'$$
(III-5.2)

and has been tabulated by Masters (III-7) in terms of the function P^* where $U = P^*(Z/\gamma, R/\gamma)U_c$ with $Z/\gamma = (2\xi/a)^{-1/2}$, $R/\gamma = (\psi/a)(Z/\gamma)$. Of particular

interest in the present discussion is the velocity decay along the axis in terms of ξ ; there is obtained from the solution that

$$\tilde{U}_{e} = 1 - \exp(-a/4\xi)$$
 (III-5.3)

For the case $u_c = 0$,

$$(\epsilon \rho)_c - f(x) = k \rho_e u_e r_{1/2} \tag{III-5.4}$$

$$(\rho u)_{r_{1/2}} = \frac{1}{2} \rho_e u_e$$
 (III-5.5)

Then the quantity $r_{1/2}$ is defined by

$$(r_{1/2})^2 = \left[\int_0^{\widetilde{\psi}_{1/2}} \frac{2\rho_j u_j}{\rho u} \psi' d\psi' \right]$$
 (III-5.6)

Asymptotically $(\xi \to \infty)$ this expression becomes equal to

$$r_{1/2} \longrightarrow \left(\frac{\rho_j}{\rho_\epsilon}\right)^{1/2} \xi$$
 (III-5.7)

The quantity u_c asymptotically becomes

$$u_{\epsilon} \longrightarrow \frac{u_{j}a}{\xi}$$
 (III-5.8)

and

$$[(\rho \epsilon)_c]_{as} = k \rho_c u_j a \left(\frac{\rho_j}{\rho_c}\right)^{1/2} = \text{Const.}$$
 (III-5.9)

Mr. G. Kleinstein has found from the analysis of numerical and experimental data that the quantity $(\epsilon\rho)_c$ remains constant in a large zone of the mixing and that the asymptotic expression can be used with good approximation, also in the region of the flow near the exit of the jet. On the basis of this observation, the evaluation of Eq. (III-3.9) becomes very simple. If the quantity k is assumed to be the constant given by Prandtl, by introducing Eq. (III-5.9) in Eq. (III-3.10) it results

$$\frac{x}{a} = 40 \left(\frac{\rho_j}{\rho_e}\right)^{1/2} \frac{\xi}{a} + \text{Const.}$$
 (III-5.10)

Denote the constant in this equation as x_i/a , then Eq. (III-5.3) becomes

$$u_{\epsilon} = 1 - \exp\left\{-10\left[\frac{x}{a_{\epsilon}} + \frac{x_{i}}{a_{\epsilon}}\right]^{-1}\right\}$$
 (III-5.11)

where a, is an effective area of the jet,

$$a_{\epsilon} = a \left(\frac{\rho_{j}}{\rho_{\epsilon}}\right)^{1/2} \tag{III-5.12}$$

This result is in accord with the observation of Thring and Newby,¹¹¹⁻⁹ that a homogeneous compressible jet has the same behavior as an incompressible one, provided the actual jet radius a is replaced by an effective radius a_e .

The accuracy of this simple result can be assessed as follows: Consider experiments with $u_e = 0$ yielding the decay of center line velocity $u_c = u_c(x/a)$; compute from Eq. (III-5.3) for each u_c the value of ξ/a and compare the computed values with the measured values plotted in terms of $0.025(\rho_e/\rho_j)^{1/2}(x/a)$. This has been done in Fig. 13; there have been considered data for homogeneous, isothermal (i.e., incompressible) jets, ^{111-9-11, 111-13} isothermal jets of carbon dioxide, ¹¹¹⁻¹³ heated homogeneous jets, ¹¹¹⁻¹¹ and high-speed homogeneous jets. ¹¹¹⁻¹⁰ A density range of $1/2 \lesssim \rho_e/\rho_j \lesssim 2$ is represented by the data. With the single exception of the high Mach number data of Warren ($M_j = 2.60$) the correlation of the results is as good as can be expected. It is noted that a similar discrepancy with respect to these data was found in Ref. III-1 and remains unexplained, but that for $M_j = 1.51$ the agreement is satisfactory herein.

The results of Fig. 13 imply that the constant in Eq. (III-5.10) is approximately -0.175 independently of the ratio ρ_e/ρ_j ; thus the transformation from the ξ , ψ plane to the x, ψ plane is for this case of $u_e = 0$ simply

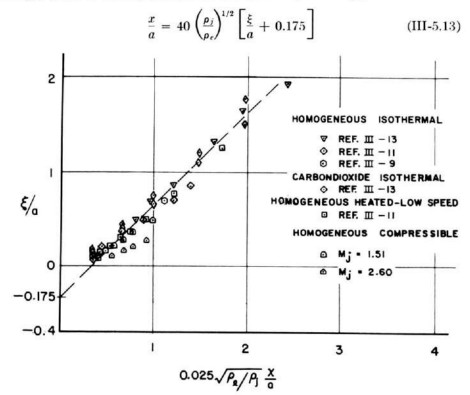


Fig. 13. Relation between ξ/a and x/a from available experimental results and comparison with the relation given by the expression $d\xi/dx = 0.025 \sqrt{\rho_e/\rho_j}$.

Note that the final transformation from x, ψ to x, r involves application of Eq. (III-3.4) and requires specification of the density distribution at least in terms of ξ and ψ .

In Figs. 14, 15, 16, and 17, some comparisons between measured and calculated quantities for the case of heterogeneous jets, investigated experimentally in Ref. III-13, are shown.

Figures 14, 15, and 16 give the decay of velocity and concentration along the axis for nitrogen, carbon dioxide, and helium jets. In the case of helium jet the experimental velocity distribution shown does not agree with the analytical value, however, it is stated in Ref. III-13 that "velocity data could not be obtained for the jet of helium because of the low impact pressures that it produced."

In Fig. 17, a radial profile distribution for the helium experiment is shown. The agreement with the analysis is good.

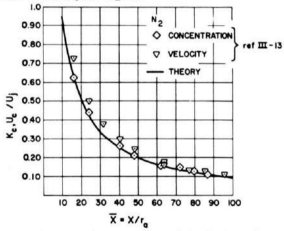


Fig. 14. Comparison of theoretical and experimental distributions of center-line velocity and concentration for nitrogen jet.

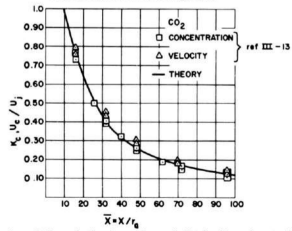


Fig. 15. Comparison of theoretical and experimental distributions of center-line velocity and concentration for carbon dioxide jet.

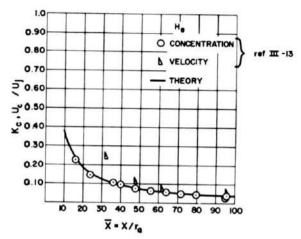


Fig.16. Comparison of theoretical and experimental distributions of center-line veolocity and concentration for helium jet.

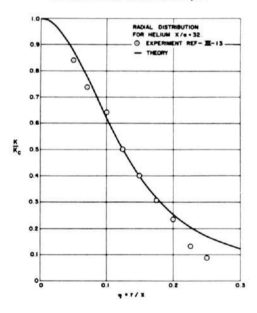


Fig. 17. Comparison of theoretical and experimental radial distributions of concentration for helium jet.

ANALYSIS OF MIXING FOR P, AND S, DIFFERENT FROM UNITY

In order to determine concentration and temperature distribution in the flow, Eqs. (III-2.4), (III-2.5) and (III-2.2) must be solved. These equations involve the turbulent Prandtl and Schmidt numbers.

If these numbers are unity, then for jet flows, generalized Crocco relations exist among the velocity stagnation enthalpy and element mass fractions so

that the velocity distribution determines the entire flow. If the flow is nonreacting Eqs. (III-2.4) and (III-2.5) are identical, therefore, the same relations are valid also for species mass fractions.

In the case of jets discharging in quiescent atmosphere u, is zero and if P_t and S_t are unity the ratio u/u_j and K_{e_j} (concentration of the constituent of the jet) are exactly the same.

For the case when P_t and S_t are different from unity, solutions of Eqs. (III-2.4)–(III-2.5) can still be carried over directly to obtain concentration and temperature provided P_t and S_t are constant or at most functions of the streamwise coordinate.

Consider a solution to Eq. (III-2.4) applied to one foreign species, i.e., for $Y_i \equiv K$ and for $w_i \equiv 0$, i.e., for no chemical reaction. Application of the von Mises transformation and the previous considerations concerning ϵ leads with no further approximation to

$$\frac{\partial K}{\partial x} = \frac{(\epsilon \rho)_{\epsilon}}{\rho_{j}} \widetilde{\psi}^{-1} \frac{\partial}{\partial \psi} \left(S_{t}^{-1} \widetilde{\psi} \frac{\partial K}{\partial \psi} \right)$$
 (III-6.1)

Now the available data on turbulent Schmidt numbers, S_t , for a variety of mixtures indicates that an sssumption of a constant value therefore, is reasonable (cf. Refs. III-12 and III-14). Accordingly, for these jet flows assume S_t to be constant and introduce a new independent variable ϵ_1 such that*

$$\xi_1 = S_t^{-1} \int_{\epsilon}^{x} \frac{(\epsilon \rho)_e}{\rho_i u_i} \frac{dx'}{a} = S_t^{-1} \xi$$
 (III-6.2)

Then Eq. (III-6.1) becomes

$$\frac{\partial K}{\partial \xi_1} = \frac{a}{\psi} \frac{\partial}{\partial \psi} \left(\widetilde{\psi} \frac{\partial K}{\partial \psi} \right) \tag{III-6.3}$$

Eq. (III-6.3) has the same form as Eq. (III-3.11). For a jet discharging in quiescent atmosphere

Then the solution for K is the same as that for U given by Eqs. (III-5.2) and (III-5.3) provided ξ is replaced by $\xi_1 = S_t^{-1}\xi$; clearly if $S_t \simeq 1$, $K \equiv U$, i.e., a Crocco relation exists between velocity and concentration of foreign gas. The concentration along the axis is from the solution given by

$$K_c = 1 - \exp[-S_t/4(\xi/a)]$$
 (III-6.5)

while from Ref. III-8 the concentration profiles are given as $K/K_c = P^*(Z'/\gamma, R'/\gamma)$, where $Z'/\gamma = (2\xi_1/a)^{-1/2}$, $R/\gamma = (\psi/a)Z'/\gamma$.

^{*} Note that with no difficulty S, could be taken to be a function of x.

For isothermal, isobaric conditions $\rho W^{-1} = \text{constant}$ or more precisely

$$\rho/\rho_j = [K + (W_j/W_e)(1 - K)]^{-1}$$
 (III-6.6)

Thus from Eqs. (III-6.5) and (III-6.6) evaluated at the center line, Eq. (III-3.5) can be used to carry out the transformation back to the x- ψ plane. Eq. (III-6.2) for $u_{\epsilon} = 0$ and the approximate relation

$$(\rho\epsilon)_c = 0.025 u_j a \rho_\epsilon \left(\frac{\rho_j}{\rho_\epsilon}\right)^{1/2}$$

gives

$$\xi_1 = S_t^{-1} [0.025 \left(\frac{\rho_e}{\rho_i}\right)^{1/2} x - \text{Const.}]$$
 (III-6.7)

Therefore, from experimental data of $K_c = K_c(x/a)$ it is possible to determine experimentally the value of S_t . The results of these considerations, applied to the jets of helium and carbon dioxide of Figs. 15 and 16, indicate that in this case $S_t \sim 1$. Note that a change in density ratio ρ_e/ρ_j of over an order of magnitude is represented here.

A similar analysis can be performed when the temperature of the jet is different from the temperature of the surrounding flow. In this case

$$\rho/\rho_i = T_i/T$$

so that the temperature field must be computed. For a constant turbulent Prandtl number, not unity in general, the previous analysis for K can be carried through to yield the temperature field; the dependent variable becomes

$$\theta \equiv (T - T_{\epsilon})(T_{i} - T_{\epsilon})^{-1} \tag{III-6.9}$$

while the independent variable becomes

$$\xi_2 \equiv \sigma_t^{-1} \int_0^x (\rho_{\epsilon}/\rho_j) (\bar{\epsilon}/u_j a) dx' = \sigma_t^{-1} \xi$$
 (III-6.10)

The solution with $\theta \equiv 1$ at $x = \xi_2 = 0$ is again given by Eqs. (III-5.2) and (III-5.3), provided ξ is replaced by ξ_2 and U by θ .

EXPERIMENTS CONCERNING HETEROGENEOUS MIXING OF TWO MOVING STREAMS

In order to obtain fundamental data on the mixing processes involved when there is an external flow, a series of experiments concerning the axisymmetric mixing of hydrogen and air have been carried out by Mr. L. Alpinieri at PIBAL in a mixing rig utilizing the supersonic facility at PIBAL; the static temperatures were roughly ambient so that no chemical reaction occurred. Two sets of experiments were performed; in one a subsonic hydrogen jet exhausted into an axisymmetric stream of air with a Mach number of 3.0. The experimental setup is

shown in Fig. 18. In the second a subsonic jet of hydrogen exhausted into a subsonic air stream; the experimental rig is shown in Fig. 19. In both sets of tests the distributions of static and total pressures and of hydrogen concentration throughout the mixing region were measured. The pressures were measured by standard techniques while the hydrogen concentration was measured by a thermal conductivity cell which was calibrated against known hydrogenair mixtures. In the subsonic-supersonic tests the stagnation pressure of the air stream was 1.3 atmospheres. In both sets of tests the stagnation temperatures of all streams was approximately 290°K. The experimental results and comparisons with theory will be given later in extenso; for the present only the results indicating some important points will be presented, i.e., those pertaining to the importance of initial boundary layers, to isovelocity mixing, and pressure gradients.

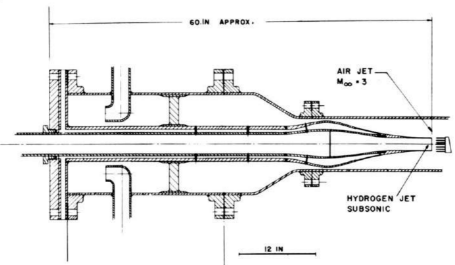


Fig. 18. Experimental apparatus for the investigation of subsonic-supersonic mixing.

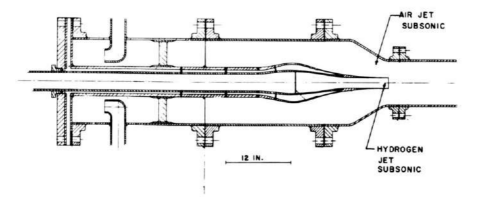


Fig. 19. Experimental apparatus for the investigation of subsonic-subsonic mixing.

For this purpose consider first the distributions of concentration obtained in the subsonic-supersonic tests; shown in Fig. 20 is the distribution of mass fraction of hydrogen denoted K along the center line for four different values of the velocity ratio, $u_j/u_e \equiv U_j$. Shown in Fig. 21 are the radial distributions of K for the same velocity ratios and for two downstream stations x/a

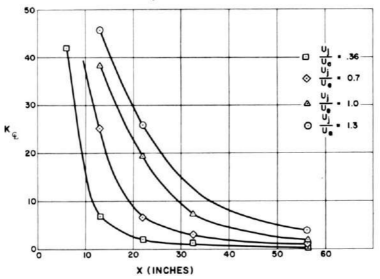


Fig. 20. Distribution of center-line concentration for various velocity ratios; subsonicsupersonic mixing.

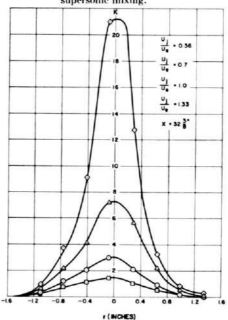


Fig. 21. Radial concentration profiles for various velocity ratios; subsonic-supersonic mixing x/a = 32.4.

= 13 and 32.4. It will be noted first from these figures that the mixing of hydrogen and air is rapid, i.e., K_c drops to 0.1 with a distance corresponding to $x/a \leq 12$ for $u_j = 0.36$ u_e . These results are qualitatively in agreement with the analysis discussed above. For a given u_e , as u_j increases from zero the theory predicts, at a given station x/a, an increase in K_c . For the form of the eddy viscosity proposed here the increase of K_c is gradual in the region of $u_j = u_e$, however no changes in sign of the variation of K_c and u_j is predicted. This is not true for the form of the eddy viscosity proposed in Ref. III-1, which would indicate an increase in K_c until $u_j = u_e$, and then a decrease of K_c for values of $u_j > u_e$.

A direct comparison between these experiments and analysis requires the development of analyses that takes into account the presence of the boundary layer on the air side at the origin of the mixing. These analyses can be derived approximately by extending the present analysis if the thickness of the boundary layer is not too large. The presence of the air boundary layer for the experiments involving mixing of air and hydrogen is very important, because the momentum flux in the air boundary layer is very large with respect to the momentum flux of the hydrogen stream, due to the low density of the hydrogen. In these experiments for the data presented here, the boundary layer is very large, the thickness is of the order of one-half inch, therefore the momentum flux deficit in the boundary layer is larger than the momentum of the hydrogen. It is difficult to reach even qualitative conclusions from these experiments, and the results shown in Figs. 20 and 21 should not be interpreted in terms of step distributions of velocity at the origin of mixing. Indeed, it is of interest to note the following: The great density difference between hydrogen and air implies that the boundary layer in the air stream at the origin of mixing will generally play an important role in the mixing processes as well, as mentioned previously, in the chemical kinetics under practical conditions of interest in supersonic diffusion flames.

One of the main reasons for setting up the rig for subsonic-subsonic mixing was to reduce significantly the initial boundary layers. Indeed, the results of subsonic-subsonic mixing between hydrogen and air indicate that the qualitative agreement between theory and experiment is not due to the effects of the initial boundary layer. In this case the boundary layer in the air stream at the origin of mixing is thin; consequently, the effect of the initial boundary layers is negligible at a small distance downstream of the origin of the mixing region.

A comparison between experimental and analytical results for axial distributions of concentration of hydrogen for three ratios of jet to free-stream velocities is presented in Fig. 22. In Fig. 23 a comparison of radial distributions for a velocity ratio $u_j/u_e = 1$ is shown for values of x/R = 5 and 10.5.

The trend given by the analysis is in agreement with experiments. The concentration at any given station x = constant increases gradually with u_j changes from 0.7 to 1.3 when the mixing extends to the axis. The concentration initially changes very rapidly with distance and then when small values of concentration are reached changes slowly with distance. Both results indicate that the mixing process is very rapid also for values of $u_j/u_e \sim 1$ and the shape of $\partial K/\partial x$ given by the analysis appear to be in agreement with experiments.

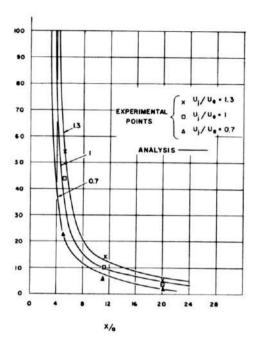


Fig. 22. Comparison between experimental and analytical results for axial distributions of concentration of hydrogen.

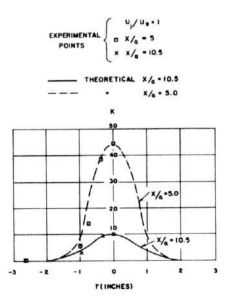


Fig. 23. Comparison between experimental and analytical results for radial distributions of concentration of hydrogen.

EFFECT OF TURBULENCE ON REACTION RATES

In order to establish properties of mixing, reacting flows, it is necessary to combine mixing analyses of the type described herein with descriptions of the chemical kinetics in turbulent flows. Formally, expressions for the creation terms w_i in the species conservation equations are required. It will be recalled that the equations employed here to investigate mixing are in terms of time-averaged quantities so that the desired descriptions are of the time-averaged mass rate of production of each species per unit volume per unit time. While the effect of turbulence in increasing the mixing of fuel and oxidizer is understood from a phenomenological point of view, the effect of turbulence on kinetics is not clear.

The effect of turbulence on the rate of flame propagation has been extensively studied but apparently the fundamentals of the effect are still subject to debate. In Ref. III-15, Hawthorne, Weddell, and Hottel considered the effect of turbulence to be described by an "unmixedness factor" which indicates the degree of inhomogeneity of fuel and oxidizer; they considered, however, that the actual kinetic rates of chemical behavior were infinite so that fuel, oxidizer and products were in equilibrium locally with respect to space and time. Scientific investigation of the chemical kinetics of turbulent flows according to statistical theory has been initiated by turbulence specialists (cf. Corrsin^{111-16, 111-17}) but the idealizations necessary for progress are so drastic as to provide little hope for application to chemical systems of practical interest in the near future.

It is the purpose of this discussion to indicate qualitatively the influence of turbulence on chemical kinetics. For this purpose it is useful to recall the basis for the rate constants employed in the analysis of Part II and shown explicitly in connection with Eqs. (II-3.1). These rate constants have been obtained by theoretical and/or experimental means under the cited assumption that thermodynamic equilibrium roughly prevails and thus that nonequilibrium prevails only with respect to chemical composition. Therefore, it is assumed that equipartition of energy among particles and internal degrees of freedom exists at least approximately. Now in high-speed turbulent flows these assumptions may be open to question; therefore, substituting classical rate expressions into the creation terms interpreted as instantaneous rates and the time averaging thereof in order to obtain time-averaged values for w, may not be valid. However, it appears that the following statements can be made: If at a fixed point in the flow, the absolute velocity squared fluctuations which may arise from either velocity or temperature fluctuations are significant with respect to $2kT/m_i$, then these fluctuations influence significantly the collision processes. This appears to be the case for mixing processes of very high velocity streams having very large ratio of static to total enthalpy. In this case the enthalpy connected with the fluctuating process is of the same order of the static enthalpy. Therefore the mechanism of collisions and the number of a given type of collision can be quite different from the values given by statistical mechanics on the basis of the thermodynamic equilibrium. The effect of the turbulence on the mechanism of collision can be discussed qualitatively by considering a reaction step following an Arrhenius law. The turbulence effects the value of T on the exponent of the expression and the functions of T which multiply the exponential function. Consider, for example, a reaction step following an Arrhenius law [reactions 1–4 to the right or to the left and reactions 5–8 to the left in Eq. (II-3.1)]. The reaction rate in this case increases greatly with the relative velocity between colliding particles as evidenced by the exponential dependence of the rate constant on temperature. Therefore, it can be expected that a large fluctuation in the absolute velocity squared will increase the reaction rate for reaction steps following an Arrhenius law. On the other hand, for reactions such as reactions 5–8 in Eqs. II-3.1 proceeding to the right where the rate constant does not have an exponential term it can be expected that turbulence would affect the reaction rates in the sense of increasing the probability of useful collisions. These statements imply that all of the reactions of interest in the hydrogen-oxygen reaction except the recombination reactions (5–8 to the right) will be accelerated by turbulence.

It would be of fundamental interest to conduct experiments, e.g., in nozzles expanding the products of hydrogen-oxygen combustion with various degrees of turbulence in the stream, to determine the effect of turbulence on reaction rates of practical importance in supersonic combustion.

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PART IV. EXPERIMENTAL RESULTS OF COMBUSTION

In order to investigate experimentally the process of supersonic combustion controlled by mixing of hydrogen and air, a series of tests is in progress at the Aerodynamics Laboratory of the Polytechnic Institute of Brooklyn. Some of the results of such investigation are presented here.

EXPERIMENTAL APPARATUS

The tests presented here were performed in an axially symmetric jet. The hydrogen was injected axially through subsonic jet at the axis of the air jet (see Fig. 24). The air Mach number was chosen equal to 1.54 in order to change sufficiently some of the parameters which are important for chemical reaction and which affect significantly the reaction time. The maximum air stagnation temperature available in the facility was roughly 1660° K; therefore, the maximum static temperature of the air could be changed for M = 1.54 between room temperature and 1110° K. With this Mach number it was possible to change the static pressure in the mixing region from values of the order of 1/20 atm to several atms. However, for safety reasons the maximum pressure, which permits optical observation through the glass windows, was limited to 1.3 atm. The temperature of the hydrogen was also variable between 1110° K and 324° K.

The hydrogen and air jets discharge into a large chamber where the external pressure can be carefully controlled. Therefore, the mixing and combustion process take place in an open jet region, which presents several advantages.

- (a) It permits the static pressure along the external part of the jet to be kept constant, independently of the combustion process.
 - (b) It simplifies the optical observation of the flow.

The measurements made were pressure, velocities and temperature of the two streams. The combustion process was observed by direct photographs and

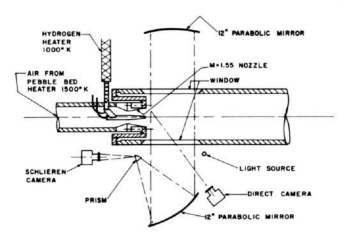


Fig. 24. Schematic representation of combustion apparatus; supersonic-subsonic mixing.

schlieren photographs. Temperature survey was also made. Direct photographs which are important in understanding the processes in the flow were taken by a long time exposure (about 1–7 sec) of the flame.

The purpose of the tests was to obtain some experimental information on the chemical and mixing process. The mixing process in the tests is complex because of the presence of the boundary layer outside of the hydrogen injector; therefore, initial tests were made in order to determine the effects of the boundary layer on the combustion tests. The presence of the air boundary layer is important for two reasons: (a) it increases the mixing process, and (b) the boundary layer air has regions where the static temperature is higher and the velocity is lower than the static temperature and velocity of the air jet. It must be noted that in all tests the walls of the hydrogen jet are preheated close to the stagnation temperature of the air.

The first part of the test program has been performed at a static pressure of 0.9 atm. For this condition the chemical behavior is described by the data of Fig. 25, which gives the induction and energy release times as functions of local temperature. In the range of temperatures, and for the pressure chosen for the tests, these times are practically independent of the chemical composition of the mixture (value of η); therefore, it is possible to obtain indications on the process of mixing and reaction also, if the air and hydrogen concentrations of the mixture are not constant and are not exactly known along any given streamline of the flow. Figure 25 also shows that the energy release time is very short, and the induction time represents the larger portion of the total combustion time. Rapid changes of reaction time take place in the range between 800° and 900°K. For example, a variation of temperature between 940°K and 830°K changes the reaction time between 2.0×10^{-4} to 10×10^{-4} , which corresponds to variations of distance in the flow of several inches. This large variation of travel length permits an interpretation of the data obtained from photographic observations.

Several series of tests were performed for the cited air Mach number of 1.54 and for hydrogen Mach number variable but less than one. In each series of

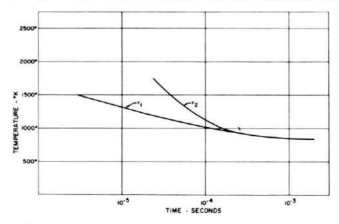


Fig. 25. Reaction times as a function of initial temperature for stoichiometric mixtures.

tests the stagnation temperature of the air and of the hydrogen were kept constant, and the velocity of the hydrogen u_i was changed.

Several series of tests were performed corresponding to different combinations of stagnation temperature of the air and of the hydrogen.

The mixing analysis does not permit accurate determination of the mixing process; however, it does permit obtaining some relation between velocity stagnation temperature and static temperature as function of concentration. If ρ_e and ρ_j are the partial mass densities of air and hydrogen at a given position where the mass density is ρ , then the average velocity and stagnation temperature can be expressed as

$$\rho u = \rho_e u_e + \rho_j u_j \tag{IV-1}$$

where u_e and u_j are the initial velocities of air and hydrogen stream and the effects of viscous losses are neglected. This approximation is satisfactory in view of the fact that the exchange of momentum between the air and hydrogen stream represents only a small percentage of the momentum of the air stream, and therefore the viscous losses related to this exchange, that depends on the momentum exchanged can be neglected, and

$$H_0 = c_p T_0 = \rho_e c_{p_e} T_{s_e} + \rho_j c_{p_j} T_{s_j}$$
 (IV-2)

As it has been discussed in Part III, these expressions are consistent with the concept of the mixing length introduced by Prandtl when molecular transport phenomena can be neglected. Therefore, if $K = \rho_i/\rho$,

$$T_0 = \frac{H_e + K(H_j - H_e)}{c_{p_e} + (c_{p_j} - c_{p_e})K}$$
 (IV-3)

$$u = u_e + (u_i - u_e)K (IV-4)$$

and

$$T = \frac{c_{p_e} T_{s_e} + K(c_{p_i} T_{s_i} - c_{p_e} T_{s_e}) - \frac{1}{2} u^2}{(c_{p_i} - c_{p_e}) K + c_{p_e}}$$
(IV-5)

From these relations, and from the relation between reaction time and static temperature T, the effect of the variation of H_j and H_e and of the concentration on the flame front can be investigated.

In the first series of tests the stagnation temperature for the air was chosen to be 1170° K corresponding to an air velocity of 2,870 ft/sec and to a static temperature of 834°K. The stagnation temperature of the hydrogen was chosen equal to 985°K. The Mach number of the hydrogen stream is low, and the static temperature changes only slightly when u_j changes, and is close to the stagnation temperature; therefore, in this series of tests the static temperature of the hydrogen is larger than the static temperature of the air. For these conditions the reaction time was expected to be of the order of 10^{-4} . During this time the air in the boundary layer has time to travel several inches and, therefore, because

of the small boundary-layer thickness, it could be expected that the boundary layer would be mixed with the main flow before combustion starts. Some of the results of such series of tests are presented in the next figures.

Figure 26a presents a direct photograph of the process of combustion for low velocity ratio, $u_j/u_e = 0.3$. A drawing of the flame front for the same condition is given in Fig. 26b. For this condition the velocity of the hydrogen jet is equal to 860 ft/sec. The flame starts at the axis at a distance of 3.5 in. downstream of the exit of the hydrogen injector; it propagates rapidly across the central region of the mixing zone and then slows down sharply when the flame reaches the outside region of the jet. For a short period the flame width increases very slowly. At a distance of the order of 7 in. from the exit of the ejector the slope of the flame front changes, and the velocity of propagation in a direction normal to the jet increase again. Downstream of this point the flame width grows approximately at a constant rate, and the flame front is inclined at an angle of approximately 7° -8° with respect to the axis of the jet.

The shape of the flame is consistent with what could be expected from the results of an analysis. In Fig. 27 the flow travel required for combustion is given as a function of local average concentration for the conditions of this test. This flow travel has been calculated by means of Eqs. (IV-4) and (IV-5) on the basis of the data of Fig. 25, and does not take into account the effect of the boundary layer.

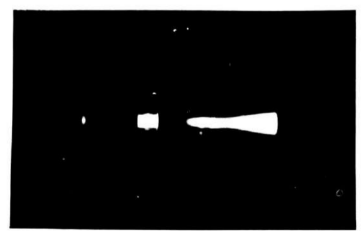


Fig. 26a. Direct photograph of air-hydrogen combustion; $T_{s_i} = 1170^{\circ}\text{K}$; $T_{s_j} = 985^{\circ}\text{K}$;

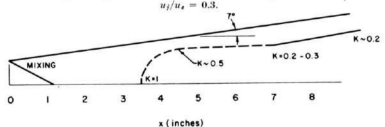


Fig. 26b. Schematic representation of mixing versus length.

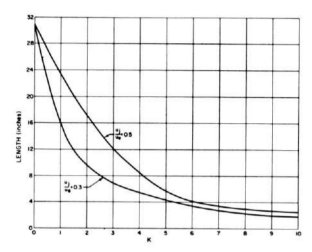


Fig. 27. Length versus mass concentration for combustion; T_s = 1170°K; T_s = 970°K.

At the center of the flame the hydrogen concentration is very high; therefore, the static temperature is high and the velocity is low. The flow travel for combustion given by Fig. 27 here is of the order of 1.9 in. Therefore, about 1.6 in. of flow travel are required before some air reaches the center of the jet. When the concentration decreases, the flow travel required for combustion increases; however, the mixing length decreases. Hydrogen concentrations of the order of 0.50 or lower are reached almost immediately downstream of the injector at the outer edge of the mixing region; therefore, for those conditions the total flow travel between the end of the ejector and the flame front corresponds roughly to the flow travel required for combustion. The flow travel required for combustion for K=0.50 corresponds to 4.4 in. This explains the shape of the flame in the central region of the mixing. When the concentration decreases the flow travel for combustion becomes very large; therefore, the velocity of propagation of the flame across the jet decreases very rapidly.

The region of the flame, where the width of the flame remains practically constant, can be attributed to the presence of the boundary layer in the air stream, generated along the external surface of the injector. The air boundary layer near the wall of the injector has larger static temperature and lower velocity than the flow outside of the boundary layer. As a consequence, in the region where the mixture contains boundary-layer air, combustion times and flow velocities are smaller; therefore, flow travels for combustion are much smaller. Then the flame front propagates fast into regions of low concentrations. When the air velocity increases and static temperature decreases, the flow travel for combustion for a given value of concentration increases and, therefore, the flame front moves into regions of higher concentrations than for the case of a mixture having boundary-layer air. As a consequence, the transversal dimension of the flame increases very slowly because the hydrogen concentration must increase along the flame front. When the flame reaches a region where the air has free stream conditions, the flame front propagates along a line of roughly

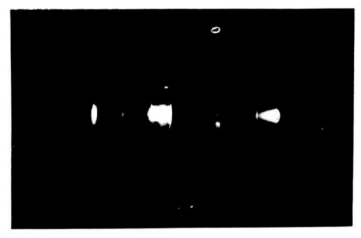


Fig. 28. Direct photograph of air-hydrogen combustion; $T_{s_e}=1190^{\circ}\mathrm{K};\ T_{s_j}=970^{\circ}\mathrm{K};\ u_i/u_e=0.5.$

constant concentration. On this basis the flame front permits information to be obtained on the mixing process. Mixing analysis indicates that the mixing for these conditions propagates at angles of the order of 7°.

When the velocity of the jet increases, then the flame changes shape, as shown in Fig. 28. The flame moves downstream of the injector and then propagates in similar fashion as before. Again the shape of the flame, the distance from the discharge section of the injector, is consistent with the mixing process and the calculated reaction rates.

In the photograph shown the velocity of the hydrogen u_j is equal to 1,430 ft/sec and $u_j/u_e = 0.5$. The stagnation temperature of the air is 1190°K, and of the hydrogen is 970°K. The relation between flow travel for combustion and concentration for these conditions is given also in Fig. 27. The flame starts at the axis of the mixing zone at a distance of roughly 7.5 in. There the travel for combustion corresponds approximately to 4.5 in.; therefore, the mixing travel is equal to 3 in. If u_j continues to increase, the flame moves gradually downstream of the exit of the injector. For high u_j/u_e the mixing process becomes slower and the combustion starts at the outer edge of the mixing region.

In Fig. 29 the distance x in inches measured from the discharge section of the injector is plotted as a function of u_i/u_e . For the case of $u_i = u_e$ the velocity of the hydrogen is 2,820 ft/sec, and the static temperature of the hydrogen is 890°K. For these conditions the temperature of the mixing is also of the order of 872°K, and the combustion will occur after 4.5×10^{-4} sec. Then the distance would be of the order of 13 in. This value does not agree exactly with the experimental results shown in Fig. 29 which gives a distance of the order of 10 in. The difference is due to the presence of the boundary layer that slightly increases locally the static temperature and decreases the velocity of the mixture.

If the velocity u_j increases to a value of $2u_e$, then the static temperature of the hydrogen becomes equal to 834°K and the reaction time becomes of the order of 6.7×10^{-4} ; therefore the length must become of the order of 24 in.

The experimental data agree approximately with the values of the distances given by the analysis.

In order to understand the interplay between mixing and combustion phenomena of hydrogen and air, it can be useful to discuss two other series of tests having different stagnation conditions. In the second series the stagnation temperature of the hydrogen is the same as the static temperature of the air, and this temperature is somewhat higher than in the first series; therefore, the reaction rates are faster. In the third series of tests the stagnation temperature of the hydrogen is much lower than the static temperature of the air.

$$T_{s_e} = 1320^{\circ} \text{K}$$
 $T_{s_i} = 954^{\circ} \text{K} - 1000^{\circ} \text{K}$ (series 2)
 $T_{s_e} = 1320^{\circ} \text{K}$ $T_{s_i} = 362^{\circ} \text{K}$ (series 3)

For the second series, because the static temperature of the hydrogen and of the air are about the same. The reaction rate is constant and of the order of 1.7×10^{-4} independent of concentration. Photographs for values of $u_j = 0.28$, 0.92, $1.21u_s$ are shown in Figs. 30, 31, and 32, respectively.

For low values of u_j the combustion starts first at the center, Fig. 30. The distance is roughly 2.6 in. from the exit of the jet. In this test the hydrogen temperature is equal to 1000° K, and the reaction time at the axis is 1.0×10^{-4} . The velocity is 860 ft/sec; therefore, the reaction length is of the order of 1 in.; and the mixing length is of the order of 1.6 in., that is, equal to the value determined in the preceding test for about the same value of u_j/u_e . The flame shape is

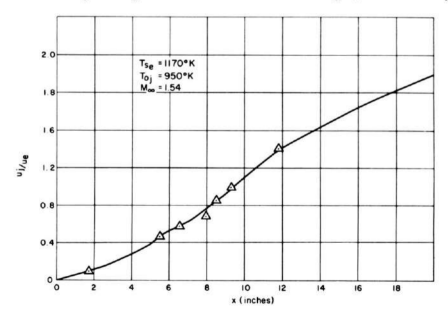


Fig. 29. Initiation of combustion mixing region for various values of u_i/u_e : $T_{e_e}=1170\,^{\circ}\mathrm{K}$; $T_{e_i}=950\,^{\circ}\mathrm{K}$.

as shown schematically in Fig. 33. The flame propagates rapidly across the jet. The flame front depends only on the velocity distribution because the temperature of the mixture is approximately constant everywhere. The velocity outside is 3,100 ft/sec, therefore the flame reaches the outside of the mixing at a distance of the order of about 6 in., which corresponds to a reaction time of 1.7×10^{-4} . This agrees with the measured value. The shape of the flame downstream of this point gives the measure of the velocity of turbulent diffusion of the hydrogen normally to the stream.

If the velocity u_j increases, then the time required for initiation of the combustion at the axis increases; however, the outside conditions do not change. This is shown from photographs in Figs. 31 and 32. If $u_j = 0.92u_e$, the static temperature, and therefore, the reaction time are about the same as before; however,

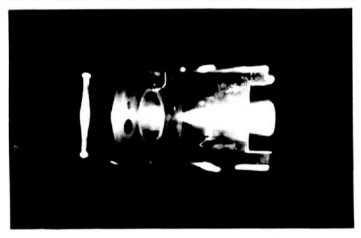


Fig. 30. Direct photograph of air-hydrogen combustion; $T_{s_e} = 1300^{\circ}\text{K}$; $T_{s_j} = 970^{\circ}\text{K}$; $u_j/u_e = 0.28$.

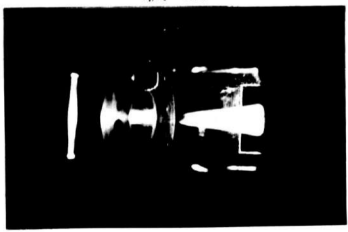


Fig. 31. Direct photograph of air-hydrogen combustion; $T_{s_e} = 1300^{\circ}\text{K}$; $T_{s_f} = 970^{\circ}\text{K}$; $u_i/u_s = 0.92$.

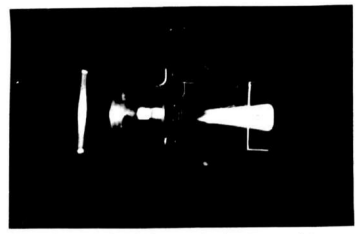


Fig. 32. Direct photograph of air-hydrogen combustion; $T_{s_e}=1300\,^{\circ}\mathrm{K};\ T_{s_j}=960\,^{\circ}\mathrm{K};$ $u_j/u_e=1.21.$

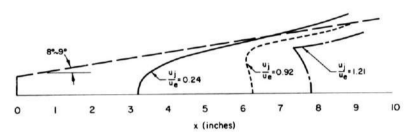


Fig. 33. Schematic representation of combustion for various values of u_j/u_e .

the reaction length at the center is larger because of the increase in velocity. The combustion starts at a distance of the order of 6 in. The flame outside has the same shape as before.

For the condition $u_j = 1.21u_e$ the hydrogen temperature is slightly lower. The flame has the same general configuration as for $u_j = 0.92u_e$; however, it starts slightly downstream of the preceding case. The difference is of the order of 1 in. This would indicate a variation or mixing length of the order of 13 percent; this is in qualitative agreement with the mixing tests. All the photographs indicate that the velocity of propagation of the flame normal to the stream increases moving downstream with the flow. This can be attributed to a gradual axial pressure rise connected with the heat release due to combustion. These tests indicate that the reaction time calculated is in agreement with experiments. It also confirms the results of the mixing experiments which indicate that the mixing is fast when $u_j < u_e$, and decreases only gradually when u_j increases without any significant changes at $u_j = u_e$.

In all the tests presented before, the process of heat release is relatively slow, and therefore the local pressure rise due to combustion is small and gradual. The schlieren pictures for such flows indicate that the compression generated by combustion waves does not form an envelope, and shocks are not found in the flow. However, the absence of shock is directly dependent on the mechanism of heat release and on the dimensions and type of boundary of the air jet. Shocks can be produced either by changing the geometry of the air jet or by accelerating and increasing the mechanism of heat release. Photographs in Figs. 34 and 35 are proof of such possibilities. Both photographs are a composition obtained by superimposing a schlieren photograph to a flame photograph taken from the same experiment.

The windows for the two photographs are not of the same dimensions and the schlieren axis is perpendicular to the axis of the flow, while the flame photograph is inclined with respect to the axis of the flow as shown in Fig. 24. Therefore a difference of angle exists for the two data. However, the superposition is qualitatively correct and gives a correct impression of the relation between heat release and shock formation.

Two different tests have been performed in order to indicate the effect of the two parameters, mechanism of heat release and boundary conditions.

The photograph of Fig. 34 corresponds to stagnation conditions $T_{s_e} = 1440^{\circ}$ K, $T_{s_j} = 935^{\circ}$ K, and $u_j = 0.28u_e$. For these conditions the reaction rate is about



Fig. 34. Direct and schlieren photographs of air-hydrogen combustion; $T_{\epsilon_e} = 1440$ °K; $T_{\epsilon_i} = 935$ °K; $u_i/u_e = 0.28$.

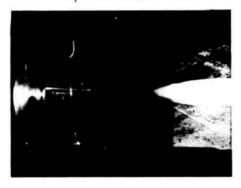


Fig. 35. Direct and schlieren photographs of air-hydrogen combustion; $T_{s_a}=1280\,^{\circ}\mathrm{K}$; $T_{s_a}=380\,^{\circ}\mathrm{K}$; $u_i/u_*=0.27$; $p_a/p_b=1.4$.

twice as fast as for the case of Fig. 30 and therefore the heat release is faster. At the same time, the mixing rate increases when combustion takes place because of the pressure rise due to heat release; therefore, as soon as the combustion rate increases the mixing rate also increases, and consequently the heat release becomes much faster. The faster rate of heat release produces stronger pressure rise and, therefore, produces shocks. For the case shown in Fig. 35 the stagnation temperature of the air is equal to 1170°K and the hydrogen temperature to 362°K, therefore, the combustion rate for a given concentration is lower. However, in this test the mixing process has been strongly augmented by creating a positive pressure gradient along the axis of the mixing stream.

The hydrogen jet at the exit of the injector is subsonic, while the air jet is supersonic, therefore, for a short length the central part of the mixing region is subsonic. If a pressure rise is produced in the flow, in some region not too far from the exit of the injector, this disturbance can travel upstream through the subsonic central part and from the center propagates into the supersonic region. Then the velocity of the central part of the mixing tends to decrease because of the pressure rise, while the mixing tends to produce the opposite effect. As a consequence the mixing is strongly increased, specifically if the velocity at the center is low. In the present test, the static pressure rise has been produced by increasing the static pressure in the discharge chamber to a value of 1.4, the static pressure of the air stream at the exit of the nozzle. Then a shock is produced at the end of the nozzle that travels toward the axis and reaches the axis somewhat downstream of the region where combustion starts. The shock produces a pressure rise in the central part of the mixing where the stream is still subsonic. The pressure rise decreases the velocity and the mixing sharply increases the heat release. While shocks would not be formed in this region in the absence of combustion, the rapid combustion process produces sufficient changes in density and pressure to create shocks as shown by the photograph.

Consider now test series 3. For this series the hydrogen is cold, therefore, combustion can occur only when the concentration is low. The flow travel for combustion as function of concentration increases very rapidly for values of concentrations between 0 and 0.1, because this change corresponds to a temperature change between 942°K and 600°K for $u_i = 0.1u_e$ and between 942°K and 560°K for $u_i = 0.78u_e$. Therefore combustion can take place only in regions where the concentration is somewhat less than 0.10. The velocity of the jet for K = 0.1 changes from 2,667 ft/sec to 2,983 ft/sec when u_i changes from 0.1 to $0.78u_a$. This change of u is small and therefore it can be assumed that the jet velocity at K < 0.1 is about the same independently of the value of u_j . Figs. 36, 37, and 38 show the flame shape for $u_i/u_e = 0.78$, 0.525, and 0.1, respectively. For high velocity of the hydrogen jet $(u_i/u_e = 0.78 \text{ and } 0.525)$ the concentration below 0.1 is reached first in the outside region of the mixing zone, and the flame initiates there. For the temperatures and velocities of the experiments the flow travel for combustion for K < 0.1, determined on the basis of the calculated reaction rates, is equal to 6.5 in. The photographs of Figs. 36 and 37 give for this distance a value of 5.5 in. The smaller value of flow travel for combustion given by the experiments is justified by the presence of the boundary layer.

For very low values of u_j , $u_j/u_e = 0.1$, the combustion starts at the center. This indicates that a very short flow travel is required to completely mix hydrogen and air when the velocity of the hydrogen is small, so that very low hydrogen concentration is obtained at the axis. Again this rapid mixing can be attributed to the presence of the boundary layer in the air stream. For very low u_j the mass of the hydrogen jet is of the same order as the mass of the air boundary-layer, then the mixing is fast and the travel is short, because of the lower velocity of the boundary-layer air.

The last experimental investigation on combustion presented here is related to the study of effects of pressure variation on reaction rates. For this series of tests the values of u_j/u_e and the stagnation temperatures of hydrogen and air have been kept constant while the static pressure has been changed. The mixing

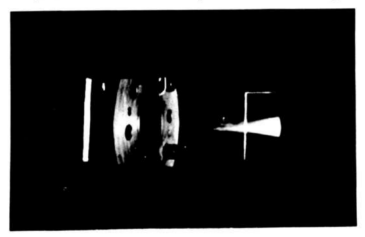


Fig. 36. Direct photograph of air-hydrogen combustion; $T_{s_e} = 1320^{\circ}\text{K}$; $T_{s_j} = 330^{\circ}\text{K}$; $u_j/u_e = 0.78$.

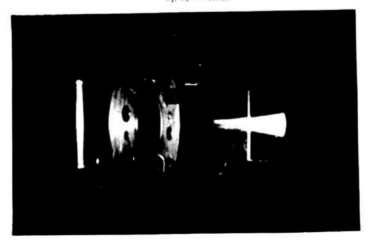


Fig. 37. Direct photograph of air-hydrogen combustion; $T_{s_e} = 1320^{\circ}\text{K}$; $T_{s_j} = 340^{\circ}\text{K}$; $u_j/u_e = 0.525$.

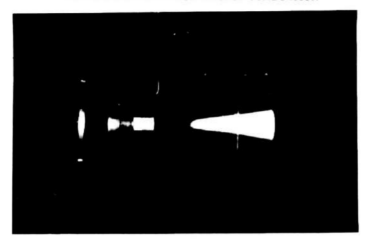


Fig. 38. Direct photograph of air-hydrogen combustion; $T_{s_e}=1320\,^{\circ}\mathrm{K};\ T_{s_j}=380\,^{\circ}\mathrm{K};$ $u_j/u_e=0.1.$

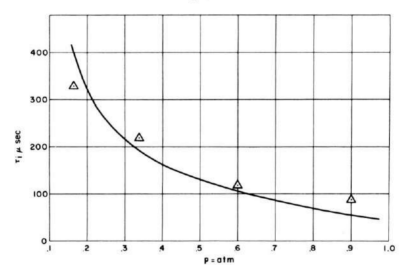


Fig. 39. Comparison of theoretical and experimental induction times.

process is only slightly affected by the pressure changes because the flow is turbulent, therefore, the variation of flame position can be attributed to the effect of the variation of the static pressure on the reaction rate. In Part II an approximate equation has been derived for the ignition delay as function of temperature [Eq. (II-5.1)]. This equation relates the induction time to the values of the static pressure and temperature. The relation is a function of the values of the chemical reaction rates. The experiments permit measuring the distance between the exit of the jet and the initiation of the flame; therefore it permits checking the approximation of such an equation and of the reaction rates used. In order to have an accurate indication of the mixing length, the tests were

performed with air at 1440° K and hydrogen cold, 380° K, so that the combustion will start only for very low values of concentration (below K=0.05). For such conditions the velocity of the mixture is equal to the velocity of the air, and the travel required for mixing is negligible, because of the rapid mixing process and of the low values of concentration required for combustion.

An analysis of the photographs indicates that the flame front has a discontinuity. This discontinuity is due to the presence of the boundary layer in the air stream. As it has been explained before, the small amount of combustion that occurs before the discontinuity is due to the low velocity part of the boundary layer. The distance used for the determination of the reaction time is the distance between the end section of the hydrogen injector and the discontinuity on the flame front which corresponds to a mixture having a velocity equal to the free stream velocity of the air. Figure 39 shows a comparison between the analytical results and the combustion times determined from the distances given by the experiments. The comparison indicates that the analysis is in good agreement with the experimental results.

PART V. CONCLUSIONS

The theoretical and experimental investigation reported herein indicates that for the range of conditions investigated the analysis of chemical reactions between hydrogen and air is sufficiently accurate to permit fluid dynamic analysis of the flow field and interpretation of experimental data connected with supersonic diffusion flames. The fluid dynamic problem is complex and requires a better quantitative understanding of mixing problems.

The experimental results indicated that turbulent mixing of hydrogen air stream is fast and at present can be predicted, at least qualitatively, by analyses; however, variations to the incompressible type of expression of eddy viscosity must be introduced. Such modifications seem called on for physical ground. The presence of air boundary layer strongly affects the mixing and must be taken into account.

On the basis of these results it appears that additional work along the lines indicated in the report is required to improve the analysis of the fluid dynamic aspects of supersonic diffusion flames; the chemical aspects of the problem are well understood for the range of parameters investigated in the experiments reported here and in related work elsewhere. The question of the effect of turbulence on the chemical kinetics is still open.

