General Disclaimer

One or more of the Following Statements may affect this Document

- This document has been reproduced from the best copy furnished by the organizational source. It is being released in the interest of making available as much information as possible.

- This document may contain data, which exceeds the sheet parameters. It was furnished in this condition by the organizational source and is the best copy available.

- This document may contain tone-on-tone or color graphs, charts and/or pictures, which have been reproduced in black and white.

- This document is paginated as submitted by the original source.

- Portions of this document are not fully legible due to the historical nature of some of the material. However, it is the best reproduction available from the original submission.

Produced by the NASA Center for Aerospace Information (CASI)
CATALYTIC COMBUSTION FOR THE AUTOMOTIVE GAS TURBINE ENGINE

by David N. Anderson, Robert R. Tacina, and Thaddeus S. Mroz
National Aeronautics and Space Administration
Lewis Research Center
Cleveland, Ohio 44135

TECHNICAL PAPER to be presented at the
Fourth International Symposium on Automotive Propulsion Systems
sponsored by the NATO Committee on the Challenges of Modern Society
Washington, D. C., April 17-22, 1977
ABSTRACT

A study of fuel injectors to provide a premixed prevaporized fuel-air mixture and an evaluation of commercial catalysts were performed as part of a program leading to the demonstration of a low-emissions combustor for an automotive gas turbine engine. At an inlet temperature of 800 K, a pressure of $5 \times 10^5$ Pa and a velocity of 20 m/s a multiple-jet injector produced less than ±10 percent variation in Jet-A fuel-air ratio and 100 percent vaporization with less than 0.5 percent pressure drop. Fifteen catalytic reactors were tested with propane fuel at an inlet temperature of 800 K, a pressure of $3 \times 10^5$ Pa and inlet velocities of 10 to 25 m/s. Seven of the reactors had less than 2 percent pressure drop while meeting emissions goals of 13.6 gCO/kg fuel and 1.64 gHC/kg fuel at the velocities and exit temperatures required for operation in an automotive gas turbine engine. NOx emissions at all conditions were less than 0.5 ppm. All tests were performed with steady-state conditions.
CATALYTIC COMBUSTION FOR THE AUTOMOTIVE
GAS TURBINE ENGINE

by David N. Anderson, Robert R. Ticina,
and Thaddeus S. Mroz

National Aeronautics and Space Administration
Lewis Research Center
Cleveland, Ohio 44135

SUMMARY

A study of fuel-air premixing-prevaporizing systems and commercial
catalysts was performed as part of a program leading to the demonstration
of a low-emissions combustor for an automotive gas turbine engine. The
goals included a fuel preparation system which would supply a fuel-air
mixture which was uniform to within ±10 percent of the mean fuel-air
ratio, with 90 percent fuel vaporization and with no autoignition. The
catalytic reactor was required to produce emissions which were low
enough to meet the most stringent proposed U.S. automotive standards.
The overall pressure drop for both systems was to be less than 3 per-
cent, with 1 percent allowed in the fuel-air preparation system and
the remainder in the catalytic reactor.

The engine conditions included a combustor inlet temperature range
of 970 K at full speed to 1210 K at idle, a combustor exit temperature
of 1310 K at all speeds, and a pressure range of 1.5x10^5 Pa at idle to
4.5x10^5 Pa at full speed. The maximum available inlet temperature in
the test rig was 800 K; however, it was possible to duplicate the other
operating parameters.

For the fuel-air preparation studies, a pressure of 5x10^5 Pa was
considered the most likely to cause autoignition. Four fuel injector
configurations were tested at this pressure, at a temperature of 800 K,
and a reference velocity of 20 m/s with Jet-A fuel. Jet-A is a typical
gas turbine fuel. A multiple-jet injector produced the required fuel-
air distribution and vaporization with only 0.5 percent pressure drop.
No autoignition occurred in a 35-cm length at these conditions; however,
at the higher inlet temperatures of the regenerative gas turbine engine,
autoignition will probably be the most serious problem to be solved.

For the catalyst evaluation, a pressure of 3x10^5 Pa was chosen to
represent the regenerative gas turbine engine pressure range, and propane
fuel was used to avoid autoignition and vaporization problems. Fifteen
catalytic reactors were tested at an inlet temperature of 800 K and with
reference (inlet) velocities of 10 to 25 m/s. Seven reactors met the
steady-stage emissions goals of 13.6 g CO/kg fuel and 1.64 g HC/kg fuel
with less than 2 percent pressure drop at the velocities and exit tem-
peratures required for engine operation. Inlet temperatures of 500 to
825 K were required before the catalysts were sufficiently active to

STAR category 45
combust the propane fuel.

INTRODUCTION

This paper describes studies which are intended to lead to the demonstration of a catalytic combustor for an automotive gas turbine engine.

The Lewis Research Center has established an ERDA-supported program with the goal of demonstrating an improved automotive gas turbine engine which operates with the combustor conditions shown in table 1 (ref. 1). With a turbine inlet temperature of 1310 K for all engine speeds, extensive use of ceramic components should not be required. The engine conditions of table 1 were defined by a computer analysis which predicted a combined EPA city-highway driving cycle fuel economy of 26.2 miles/gallon in a compact car (ref. 1). Also included in table 1 are reference velocities computed from the combustor inlet conditions for several combustor diameters. U. S. automotive emissions standards are defined for operation over a city driving cycle; for this cycle, the engine of reference 1 would have a fuel economy of about 22 miles/gallon. The most stringent standards proposed by the 1970 Clean-Air Act are 0.4 g NOx/mile, 3.4 g CO/mile, and 0.41 g HC/mile. The Lewis program includes the evaluation of the catalytic combustor with the goal of achieving emissions which are half those of the most stringent proposed standards.

Catalytic combustion has the principal advantage over gas-phase combustion that lower combustion temperatures and therefore, lower NOx emissions are possible. The importance of combustion temperature to NOx emissions is illustrated by figure 1 which is based on the prediction of a well-stirred reactor computer program (ref. 2). The assumptions made in preparing figure 1 included a vehicle fuel consumption of 0.0775 kg/km (about 22 miles/gallon fuel economy), a combustor operating with an inlet temperature of 970 K at a pressure of 5×10^5 Pa and no nitrogen bound to the fuel. Thermal NOx emissions are shown as a function of the maximum combustion temperature achieved before dilution to an exit temperature of 1310 K. It is clearly advantageous to burn at the lowest possible temperature to limit NOx emissions. The lowest temperature at which combustion can take place is determined by the lean flammability limit, which for hydrocarbon fuels occurs between flame temperatures of 1600 K (ref. 3) and 1800 K (ref. 4). Thus, for good stability, peak flame temperature should probably be maintained near 1500 K. Figure 1 shows that at this temperature, steady-state NOx emissions should be only half the 0.4 g/mi proposed as a standard; lean premixed combustors have in fact been demonstrated with emissions low enough to meet this standard (ref. 5).

A way to react fuel and air at even lower flame temperatures is to use catalysts to augment the reactions. Catalytic combustion at reaction temperatures of 1300 to 1400 K has been shown to give good com-
bustion efficiency and produce thermal NOx emissions which were several orders of magnitude below those of conventional combustors (refs. 6 to 9).

A schematic view of one possible catalytic combustor configuration is shown in figure 2. Fuel is injected upstream of the reactor to vaporize and mix with the inlet air. Autoignition must be avoided in this premixing section because the flame would tend to seat on the fuel injector and burn with locally high temperatures; high NOx emissions and possible component damage could result.

The catalyst bed, or reactor, may consist of several sections, each made of a different kind of catalyst; it is desirable to use a catalyst which is active at low temperatures for the inlet while subsequent sections need to be selected for good oxidation efficiency. All sections must be durable at the reactor operating conditions. Downstream of the catalytic reactor a thermal reaction zone may be provided to permit catalytically-initiated reactions to continue. Such reactions have been shown to contribute as much as 20 percent of the total reactor temperature rise for some conditions (ref. 9). Finally, dilution of the reacted products to achieve the turbine inlet conditions may be necessary. Some method of starting the combustor will also be required, although none is shown in figure 2.

Successful fuel-air premixing systems have been demonstrated in earlier studies (ref. 10) and the potential of catalysis applied to low-emissions combustion has been shown (ref. 6 to 9 and 11). However, the development of the technology required to demonstrate a complete catalytic combustor has yet to be achieved, and an evaluation of both fuel-air preparation and catalyst performance in terms of particular engine requirements has not been made. The Lewis catalytic combustion program includes both fuel-air preparation studies and catalyst testing. The emissions goals are to be achieved while operating at the conditions shown in table 1 and with a combustor pressure loss of no more than 3 percent. Arbitrarily, a goal of 90 percent fuel vaporization and a fuel-air ratio uniform to within ±10 percent with no more than 1 percent pressure drop across the fuel preparation section was chosen.

Steady-state tests were performed in a 12-cm diameter combustion test rig at inlet temperatures of 450 to 800 K, pressures of 3 and 5x10^5 Pa, catalyst inlet velocities of 10, 15, 20, and 25 m/s and fuel-air ratios from 0.01 to 0.026. Jet-A, a typical gas-turbine fuel, was used in the fuel-air preparation studies, while vaporized propane fuel was used in the evaluation of catalysts to avoid the vaporization and auto-ignition problems associated with typical gas-turbine fuels.

FUEL PREPARATION STUDIES

Test Configuration

Fuel preparation systems were tested in the rig shown schematically in figure 3. The inlet airflow rate was measured with an ASME standard
square-edged orifice. The air was heated to temperatures as high as 900 K in a non-vititating preheater. The test section inlet duct was 10 cm in diameter with a 7.6 cm diameter insert to increase the velocity of the air. This section was followed by a straight-walled diffuser to increase the diameter to 12 cm. The Jet-A fuel flow was measured by two turbine flowmeters in series; the fuel was injected into the highest velocity region of the inlet duct (see fig. 3) to provide good atomization.

The fuel-air mixture was sampled 35.6 cm downstream of the fuel injector. Two probes were traversed along perpendicular diameters to obtain the spatial fuel distribution by iso-kinetic sampling at several positions. The sample from each probe was allowed to react in a catalyst bed maintained at 1050 K in an oven. The reaction products were then analyzed to determine the concentrations of CO, CO₂ and HC. These concentrations were used to compute the local fuel-air ratio at the sampling position. The degree of vaporization was also measured with these same probes using the spillover technique described in references 10 and 12.

Eighteen meters of electrically heated (410 to 450 K) 0.5 cm diameter stainless steel tubing connected the sampling probes with the exhaust gas analyzers. CO and CO₂ concentrations were measured with Beckman Model 315B nondispersive infrared analyzers and unburned hydrocarbon concentration was measured with a Beckman Model 402 flame ionization detector. Water vapor was removed with a Hankinson Series E refrigeration-type dryer before the sample was analyzed for CO or CO₂, and corrections were made to obtain the actual, wet-basis, concentration. The concentrations of unburned hydrocarbons and CO were negligible.

The temperature and static pressure were measured upstream of the fuel injector and at the sampling station as shown in figure 3. A single Chromel-Alumel thermocouple was used in both locations.

Downstream of the sampling station, hydrogen was injected into the fuel-air stream to produce a flammable mixture. The mixture was burned to avoid discharging fuel into the atmosphere. A water quench cooled the exhaust before it discharged through a back-pressure valve.

Test Injectors

Four fuel injectors were evaluated: a multiple-jet injector, a splash-groove injector, a simplex pressure atomizer, and a Sonicore nozzle. The multiple-jet and splash-groove injectors are air-blast atomizers; that is, they rely on the relative momentum between fuel and air for atomization. The simplex pressure atomizer uses fuel pressure for atomization. The Sonicore nozzle is a commercially-available injector which depends on a high-velocity external air stream for atomization.
The multiple-jet cross-stream injector used 28 orifices of 0.37 mm diameter injecting into zones of approximately equal duct cross-sectional areas (see fig. 4(a)). The splash-groove injector (fig. 4(b)) was developed by Ingebo (ref. 13). Fuel is injected through 21 passages to impinge upon conical surfaces of the nozzle. Fine droplets are formed which are rapidly atomized by the air flow. The simplex nozzle was a Monarch 70-degree angle spray nozzle with a flow rating of 0.013 m³/hour at 6.8×10⁵ Pa differential pressure. The Sonicore nozzle (fig. 4(c)), Model 125MA, used an external air supply at 6.5×10⁵ Pa pressure to provide a high-velocity airstream. The airstream impinges on the resonator cap, and fuel injected into the airstream forms a cone-shaped finely-atomized spray. The fuel-air ratio distribution provided by the splash-groove injector, the simplex nozzle, and the Sonicore nozzle were determined both with and without the concentric air swirler shown in figure 4(d). The 30-degree vane angle induced a swirl in the air flow to improve mixing between the fuel and the air.

Test Conditions

Tests were performed at inlet temperatures varying from 450 K to 800 K, a pressure of 5×10⁵ Pa, fuel-air ratios of 0.01 and 0.025 and a reference velocity of 20 m/s. The reference velocity is computed from the inlet air temperature, pressure, and mass flow rate using the 12 cm diameter; thus, this velocity corresponds with that at the inlet to the catalytic reactor. The pressure and reference velocity are representative of those in table 7; however, the inlet temperature was not duplicated due to preheater limitations. The test pressure was chosen to be slightly higher than that in Table 7 in order to simulate the most severe condition for autoignition.

Results

The spatial fuel distribution, degree of vaporization and pressure drop were determined for each injector.

The fuel distribution along one diameter of the duct for the splash-groove injector is given in figure 5. The injector alone produced a very non-uniform profile; however, when the 30-degree-vane air swirler was mounted concentric with the injector, the fuel distribution became nearly uniform. Local values of the fuel-air ratio, calculated from the gas-analysis carbon balance, agreed within 10 percent of the average value determined from the metered fuel and air flows. Results obtained over the traverse perpendicular to that shown were essentially the same, and a nearly uniform profile was also obtained with the simplex and Sonicore nozzles when tested with the 30-degree-vane swirler. The multiple-jet injector gave a uniform profile without the necessity for air swirl. Tests conducted at inlet air temperatures of 500 and 600 K at both 0.01 and 0.025 fuel-air ratios suggested that these variables had no effect on the
uniformity of the mixture profile.

The effect of inlet air temperature on the degree of vaporization at the center of the duct is given in figure 6 for all four nozzles tested. None of the vaporization data were obtained with swirler-assisted nozzles. At any temperature, the best performance results were obtained with the multiple-jet and splash-grove injectors, although for the inlet temperatures of interest for this application, vaporization should approach 100 percent for all injector configurations.

The multiple-jet cross-stream injector, which used no air swirler, achieved the required fuel-air mixture uniformity and vaporization with only 0.5 percent pressure drop. The pressure loss for each of the other injectors was also about 0.5 percent when no air swirler was used, but was about 1 percent when the 30° air swirler was added.

**CATALYST EVALUATION**

**Test Configuration**

Catalysts were evaluated in the same 12 cm diameter combustion test facility used to test fuel-air preparation systems. Commercial grade propane fuel (91 percent pure by volume) was introduced as a vapor into the airstream through a single-orifice injector (ref. 9) 150 cm upstream of the test section. The resulting fuel-air ratio profile at the test section inlet was found to be uniform within ±10 percent of the mean.

Inlet and exit pressures were measured at wall static taps located 9 cm upstream and 12 cm downstream of the test section, respectively. For the test conditions of these experiments, total pressures are only about .2 percent higher than the static so no total pressure measurements were made. The pressure drops across the test section was measured with a differential pressure transducer between the inlet and exit static location.

A description of the catalyst evaluation test section is given in figure 7. The inlet temperature to the test section was taken as the average reading of the thermocouples in the 8-thermocouple array labelled number 1 in figure 7. Individual catalyst elements were located between pairs of the thermocouple arrays labelled 4-8 as shown in figure 7. The housing containing the catalyst elements was uncooled to limit heat loss; a downstream section containing a gas sampling probe and an additional 12-thermocouple array was water cooled, however.

The gas-sampling probe was also water cooled and had five 1.5 mm-diameter sampling orifices located in the center of equal cross-sectional areas as shown in figure 7. The use of small orifices along with copious
water cooling was found in previous testing to be necessary to insure that the CO concentration in the probe and sampling line is frozen at the sampling-station value.

In addition to the gas analysis instruments described in the Fuel-Air Preparation Section, these experiments also used a Thermo-Electron Model 10A Chemiluminescent analyzer to measure total NOx emissions.

Test Catalysts

Twelve catalysts from four manufacturers were evaluated either singly or in combination to form a total of fifteen reactors. A description of the reactors is given in table 2 and specifications of the individual catalyst elements appear in table 3. Typical catalyst elements are pictured in figure 8. The two Johnson catalysts were 7.6 cm long and used a corrugated metal substrate wound into a cylinder as shown in figure 8; the remaining elements were all 2.5 cm long and had ceramic substrates. All catalysts were either platinum, palladium, or a mixture of the two.

Tests Performed

Two series of tests were made on the reactors: (1) a determination of the catalyst activation temperature (i.e., the inlet temperature at which the catalyst became sufficiently active to react most of the fuel), and (2) measurement of reactor emissions and pressure drop.

The catalyst activation temperature was found by increasing the inlet-air temperature in increments of 25 K, starting at 500 K, with a pressure of 2x10^5 Pa and a catalyst inlet velocity of 10 m/s. This pressure and velocity correspond approximately to those existing when the engine is being cranked during start-up. At each temperature, propane was introduced at an equivalence ratio of 0.3 while the reactor thermocouples were monitored. For inlet temperatures below the catalyst activation temperature, a small temperature rise across the bed (of as much as 15-20 K) was generally observed; however, when the inlet temperature was increased to the catalyst activation temperature, the bed and exit temperatures began to increase steadily for a period of about 2 or 3 minutes until a temperature rise of several hundred degrees across the bed was achieved.

To evaluate catalyst performance, exhaust emissions and pressure drop were measured with an inlet propane-air mixture temperature of 800 K, a pressure of 5x10^5 Pa and a range of inlet velocities of 10 to 25 m/s. It was not possible to simulate the combustor inlet temperature shown in table 1; however, the pressure and velocities are fairly representative of those at all engine speed settings.
Results

The catalyst activation temperatures are given in Table 2. The GLP and G24D reactors were not available for this part of the testing. Testing was discontinued at 625 K with the OCL6 and OCDS reactors when they produced only 15-20 K temperature rise. Activation tests performed on reactors which had only one kind of element showed that activation occurred at relatively low temperatures (500-550 K) with Pt catalysts (EL1 and JM1 reactors) or when the catalyst was a mixture of equal amounts of Pt and Pd (G24 reactor). Higher temperatures were required for activation when the catalyst was either Pd (reactors E14, G17, and JM2) or a mixture with more Pd than Pt (reactors OCL6 and OCDS). Reactors E1, E2, E3, E4, and G1 had low-activation temperature elements at the front of the reactor and high-activation-temperature elements following. Table 2 shows that these combinations had the low activation temperatures characteristic of their leading elements. Because propane was used for these tests, the activation temperatures which were determined may be higher than would result if Diesel, gasoline, or Jet-A had been used.

When the reactor performance and emissions were determined at 800 K inlet temperature, NOx emissions were found to be less than 0.5 ppm for all test conditions. Actual values were difficult to measure accurately because the lowest range of the analyzer was 2.5 ppm full scale, while it was necessary to use a 100 ppm calibration gas. NOx emissions should be well below levels required to meet the 0.4 g NOx/mile standard, at least for propane fuel which has no fuel-bound nitrogen. Only the conditions for which the HC and CO emissions met the standards remained to be found.

Each reactor can be characterized by a minimum operating exit temperature. This temperature is the adiabatic reaction temperature (established by the fuel-air ratio and inlet temperature) at which the reactor must be operated to achieve the CO and HC emissions goals. Reaction temperatures higher than this minimum result in lower emissions of CO and HC. Although the automotive exhaust emissions standards are established for operation over a specific cycle which includes start-up and transient operation, conditions not duplicated in these tests, reference emission indexes can be computed from the standards to provide a guide in determining the minimum operating temperature. Assuming a vehicle fuel consumption of 0.0775 kg/km (about 22 miles/gallon fuel economy), 3.4 g CO/mile translates to 27.3 g CO/kg fuel and 0.41 g HC/mile becomes 3.29 g HC/kg fuel. The emissions goals for the program were half of these reference values, or 13.6 g CO/kg fuel and 1.64 g HC/kg fuel. A combustion efficiency of about 99.5 percent is required to achieve these goals.

Both pressure drop and minimum exit temperature are functions of the inlet velocity; therefore, the tradeoff between pressure drop and minimum exit temperature can be illustrated by plotting the re-
ults for each reactor as shown in figure 9. The pressure drop in
figure 9 is that which was measured at a reaction temperature of
1400 K. The lines representing each inlet velocity were drawn
arbitrarily through the results for the bulk of the best performing
reactors tested. For inlet velocities of 10 and 15 m/s, the per-
formance variation among all except the Ell and G24D reactors sug-
gested only the effects of the tradeoff between pressure drops and
minimum operating temperature. At higher velocities greater dif-
ferences appeared. At 25 m/s the minimum exit temperature for
reactors Ell, E4, G4, OC16 and OCDS was in excess of 1550 K and
was not determined.

One of the functions of a plot like that of figure 9 is to
help in determining if the reactor can be operated at conditions
which are consistent with both the program goals and the engine
requirements. For example, a combustor with no dilution air flow
and a 16 cm diameter reactor has catalyst inlet velocities of
15-17 m/s (table 1). The reactor exit temperature is required to
be 1310 K with no dilution flow. Figure 9 shows that none of the
reactors tested have minimum exit temperatures as low as 1310 K
at 15 m/s and with less than 2 percent pressure drop. However,
if 25 percent of the combustion air is used to dilute the products
from the reactor, the reactor exit temperature increases and the
inlet velocity decreases to the values shown in table 4. From
figure 9 it can be determined that seven of the reactors tested
would meet the program goals for both pressure drop and emissions
while operating at the reactor conditions of table 4.

The results for reactor Ell (which used a Pt catalyst) il-
istrates that a reactor which activates at a low temperature
(table 2) will not necessarily have good performance at higher
temperatures (see fig. 9). In contrast, reactor El4 (which used
a Pd catalyst) required a relatively high temperature to become
active but performed well at higher temperatures. The best
attributes of both reactors were obtained when elements from each
reactor were combined to form reactor El.

DISCUSSION

This study has successfully demonstrated both fuel-air prepa-
ration systems and catalytic reactors capable of operating within
the pressure drop and emissions restraints of an improved automotive
gas turbine engine. However, the results obtained cannot be applied
directly to engine operation because of a number of considerations.

The inlet temperature at which the tests were conducted was
limited by preheat capabilities to 800 K while the engine combustor
inlet temperature varies from 970 to 1210 K. This higher temperature
can be expected to affect the performance of the combustor in at
least two ways. The first effect is the increased tendency for autoignition of the fuel upstream of the catalyst. Although the fuel-air ratios of interest are well below the lean flammability limit and thus should not be susceptible to autoignition, locally richer mixtures exist near the fuel injector before complete mixing has taken place. Thus, mixing will have to be achieved in less than the ignition delay time. The autoignition delay time decreases exponentially with increasing inlet temperature and linearly with pressure (ref. 14). For the combustor conditions of table 1, the maximum inlet temperature occurs at the minimum pressure; as a result, the predicted range of ignition delay times is only 10.8 ms at 100 percent speed to 12.2 ms at idle. To keep the residence time with a flammable mixture below these values at the velocities shown in table 4, it is necessary that the mixing distance be less than 13.9 cm.

The second effect of operating at higher inlet temperatures should be a reduction in the reactor pressure loss. The pressure drop through the reactor is primarily due to frictional losses which are proportional to the product of mass flowrate and average reactor velocity. As the inlet temperature increases, the mass flowrate decreases because the reactor temperature rise is less for the same exit temperature. This effect of inlet air temperature may be offset by the losses incurred by the mixing of dilution air with products downstream of the reactor, however.

None of the catalytic reactors used in this study were subjected to durability tests. Each reactor was tested for 2 to 3 hours, and only the G12P reactor showed loss of activity after that time. Final selection of catalysts will have to be made on the basis of both performance and durability, however.

An evaluation of catalyst durability involving catalytic combustion at 1500 K exit temperature for a period of 1000 hours has been performed by Engelhard Industries under contract to Lewis (ref. 18). Parametric tests were performed with propane fuel at the start of durability testing and with both propane and Diesel fuel at the completion of 1000 hours of operation. The tests showed that after a period of high-temperature operation combustion of Diesel fuel may remain very efficient at conditions for which propane combustion efficiency has decreased drastically. The combustor exit temperatures shown in table 1 are lower than those tested by Engelhard, so loss of activity may not be as severe as that suggested by the Engelhard results. Tests of selected catalysts at the required reactor conditions for the desired lifetime will be needed before final catalyst selection can be made.

The Engelhard results suggest that Diesel fuel may be easier to combust than propane fuel. An additional advantage of Diesel fuel over propane is that its catalyst activation temperature is lower (ref. 16). On the other hand, NOx emissions with Diesel fuel
may be higher as a result of fuel-bound nitrogen (ref. 17). Clearly, catalyst evaluation will have to be performed with typical gas turbine fuels like Jet-A or Diesel.

No transient tests were made in these studies. The ability of catalytic reactors to respond to changes in inlet conditions will be crucial to the success of the concept. It is likely that catalytic substrate materials which have high thermal conductivities and low thermal capacities such as metal or silicone carbide will be required to insure both rapid warm-up and quick response. Tests will also have to be made to determine if engine transients will induce conditions favorable to autoignition in the fuel-air mixing section.

The emissions produced during start-up and transient operation may be considerably higher than those produced at the steady-state conditions described in this paper. Because the emissions standards are based on operation over a cycle involving both start-up and transient conditions, a full evaluation of catalytic combustor feasibility cannot be made until tests can be performed at these conditions.

The practice operation of a catalytic combustor in an engine will require a method of heating the catalyst to its activation temperature for cold-engine-start conditions. Possible approaches include the use of a preburner or upstream torch with a spark igniter, an electrical heater to raise the catalyst substrate temperature directly, a downstream burner with a spark igniter, and possibly even the use of hydrogen fuel for starting. Whatever system is chosen would operate only until the combustor inlet temperature reached the catalyst activation temperature.

While this study has shown that catalytic combustion may be feasible for steady-state operation, final evaluation of the concept will depend on engine demonstration. For a full engine demonstration to take place considerably more technology development will be required, and engine demonstration with start-up and transient operation may well reveal additional problem areas requiring attention.

REFERENCES


TABLE 1. - AUTOMOTIVE REGENERATIVE GAS TURBINE ENGINE:

HYPOTHETICAL COMBUSTOR INLET CONDITIONS FOR AN IMPROVED ENGINE (Ref. 1)

<table>
<thead>
<tr>
<th>Speed, % of full</th>
<th>Airflow, kg/s</th>
<th>Fuel flow, $10^{-3}$kg/s</th>
<th>Pressure, $10^5$Pa</th>
<th>Combustor inlet temperature, K</th>
<th>Combustor exit temperature, K</th>
<th>Reference velocity (m/s) for possible combustor diameter</th>
</tr>
</thead>
<tbody>
<tr>
<td>50 (idle)</td>
<td>0.127</td>
<td>0.455</td>
<td>1.43</td>
<td>1210</td>
<td>1310</td>
<td>19.8  15.1  12.0</td>
</tr>
<tr>
<td>60</td>
<td>0.204</td>
<td>1.135</td>
<td>1.91</td>
<td>1140</td>
<td>1310</td>
<td>22.4  17.2  13.6</td>
</tr>
<tr>
<td>70</td>
<td>0.254</td>
<td>1.705</td>
<td>2.28</td>
<td>1100</td>
<td>1310</td>
<td>22.6  17.3  13.6</td>
</tr>
<tr>
<td>80</td>
<td>0.336</td>
<td>2.77</td>
<td>2.92</td>
<td>1045</td>
<td>1310</td>
<td>22.1  16.9  13.4</td>
</tr>
<tr>
<td>90</td>
<td>0.409</td>
<td>3.84</td>
<td>5.56</td>
<td>1005</td>
<td>1310</td>
<td>21.2  16.3  12.8</td>
</tr>
<tr>
<td>100</td>
<td>0.517</td>
<td>5.41</td>
<td>4.50</td>
<td>970</td>
<td>1310</td>
<td>20.5  15.7  12.4</td>
</tr>
</tbody>
</table>
### Table 2. - CATALYTIC REACTOR DESCRIPTION

Note: The JMPt and JMPd elements are 7.6 cm long. All other elements are 2.5 cm long. A complete description of each element is given in Table 3.

<table>
<thead>
<tr>
<th>Reactor designation</th>
<th>Catalyst elements</th>
<th>Catalyst activation temperature, K</th>
</tr>
</thead>
<tbody>
<tr>
<td>E11</td>
<td>El128</td>
<td>E1128</td>
</tr>
<tr>
<td>E14</td>
<td>El1412</td>
<td>E1412</td>
</tr>
<tr>
<td>E1</td>
<td>El128</td>
<td>E1412</td>
</tr>
<tr>
<td>E2</td>
<td>El128</td>
<td>JMPd</td>
</tr>
<tr>
<td>E3</td>
<td>El128 G512:3.2</td>
<td>G524:290</td>
</tr>
<tr>
<td>E4</td>
<td>El128 OC3.6:N</td>
<td>OC1.8:1.6</td>
</tr>
<tr>
<td>G12P</td>
<td>G512:P</td>
<td>G512:P</td>
</tr>
<tr>
<td>G17</td>
<td>G517:290</td>
<td>G517:290</td>
</tr>
<tr>
<td>G24</td>
<td>G524:290</td>
<td>G524:290</td>
</tr>
<tr>
<td>G24D</td>
<td>G524D:200</td>
<td>G524D:200</td>
</tr>
<tr>
<td>G1</td>
<td>G512:5.2</td>
<td>G524:200</td>
</tr>
<tr>
<td>JM1</td>
<td>------</td>
<td>JMPt</td>
</tr>
<tr>
<td>JM2</td>
<td>------</td>
<td>JMPd</td>
</tr>
<tr>
<td>OCL6</td>
<td>OCL1.8:1.6</td>
<td>OCL1.8:1.6</td>
</tr>
<tr>
<td>ODDS</td>
<td>OC3.6:S</td>
<td>OC3.6:S</td>
</tr>
</tbody>
</table>

N.A.: Not available for these tests.
### TABLE 3. - DESCRIPTION OF CATALYST ELEMENTS

All elements are 12 cm diameter

<table>
<thead>
<tr>
<th>Element designation</th>
<th>Catalyst manufacturer</th>
<th>Manufacturers designation</th>
<th>Catalyst</th>
<th>Catalyst loading, kg/m²</th>
<th>Substrate manufacturer</th>
<th>Substrate composition</th>
<th>Cell density, cm⁻³</th>
<th>Cell shape</th>
<th>Element length, cm</th>
</tr>
</thead>
<tbody>
<tr>
<td>E1128</td>
<td>Engelhard</td>
<td>EVD 1128</td>
<td>Pt</td>
<td>*</td>
<td>Corning</td>
<td>Cordierite</td>
<td>35</td>
<td>Sine Wave</td>
<td>2.5</td>
</tr>
<tr>
<td>E1412</td>
<td>Engelhard</td>
<td>EVD 1412</td>
<td>Pt</td>
<td>*</td>
<td>Corning</td>
<td>Cordierite</td>
<td>35</td>
<td>Sine Wave</td>
<td>2.5</td>
</tr>
<tr>
<td>GS12:1:2</td>
<td>W. R. Grace</td>
<td>Davex 512A</td>
<td>Pt/Pt/1/1</td>
<td>2.0</td>
<td>General Refactories</td>
<td>Silicon Carbide</td>
<td>30</td>
<td>Sine Wave</td>
<td>2.5</td>
</tr>
<tr>
<td>GS12:1:2</td>
<td>W. R. Grace</td>
<td>Davex 512A</td>
<td>Pt/Pt/1/1</td>
<td>2.0</td>
<td>General Refactories</td>
<td>Silicon Carbide</td>
<td>30</td>
<td>Sine Wave</td>
<td>2.5</td>
</tr>
<tr>
<td>GS17:200</td>
<td>W. R. Grace</td>
<td>Davex 517</td>
<td>Pt</td>
<td>2.5</td>
<td>W. R. Grace</td>
<td>Cordierite</td>
<td>45</td>
<td>Square</td>
<td>2.5</td>
</tr>
<tr>
<td>GS24:200</td>
<td>W. R. Grace</td>
<td>Davex 554A</td>
<td>Pt/Pt/1/1</td>
<td>3.5</td>
<td>W. R. Grace</td>
<td>Cordierite</td>
<td>45</td>
<td>Square</td>
<td>2.5</td>
</tr>
<tr>
<td>GS24:200</td>
<td>W. R. Grace</td>
<td>Davex 554A</td>
<td>Pt/Pt/1/1</td>
<td>3.5</td>
<td>W. R. Grace</td>
<td>Cordierite</td>
<td>45</td>
<td>Square</td>
<td>2.5</td>
</tr>
<tr>
<td>GS24:200</td>
<td>W. R. Grace</td>
<td>Davex 554A</td>
<td>Pt/Pt/1/1</td>
<td>3.5</td>
<td>W. R. Grace</td>
<td>Cordierite</td>
<td>45</td>
<td>Square</td>
<td>2.5</td>
</tr>
<tr>
<td>JM2t</td>
<td>Johnson Matthey</td>
<td>--------</td>
<td>Pt</td>
<td>5.3</td>
<td>Johnson Matthey</td>
<td>Metal</td>
<td>62</td>
<td>Sine Wave</td>
<td>7.6</td>
</tr>
<tr>
<td>JM2n</td>
<td>Johnson Matthey</td>
<td>--------</td>
<td>Pt</td>
<td>5.3</td>
<td>Johnson Matthey</td>
<td>Metal</td>
<td>62</td>
<td>Sine Wave</td>
<td>7.6</td>
</tr>
<tr>
<td>OCL 8:1:6</td>
<td>Oxy-Catalyst</td>
<td>Oxy-Catalyst</td>
<td>Pt/Pt/1/2</td>
<td>1.0</td>
<td>General Refactories</td>
<td>Silicon Carbide</td>
<td>34</td>
<td>Circular</td>
<td>2.5</td>
</tr>
<tr>
<td>OCL 8:1:6</td>
<td>Oxy-Catalyst</td>
<td>Oxy-Catalyst</td>
<td>Pt/Pt/1/2</td>
<td>1.0</td>
<td>General Refactories</td>
<td>Silicon Carbide</td>
<td>34</td>
<td>Circular</td>
<td>2.5</td>
</tr>
<tr>
<td>OCL 8:1:6</td>
<td>Oxy-Catalyst</td>
<td>Oxy-Catalyst</td>
<td>Pt/Pt/1/2</td>
<td>1.0</td>
<td>General Refactories</td>
<td>Silicon Carbide</td>
<td>34</td>
<td>Circular</td>
<td>2.5</td>
</tr>
<tr>
<td>OCL 8:1:6</td>
<td>Oxy-Catalyst</td>
<td>Oxy-Catalyst</td>
<td>Pt/Pt/1/2</td>
<td>1.0</td>
<td>General Refactories</td>
<td>Silicon Carbide</td>
<td>34</td>
<td>Circular</td>
<td>2.5</td>
</tr>
</tbody>
</table>

*Proprietary
TABLE 4. - CATALYTIC REACTOR INLET CONDITIONS FOR IMPROVED ENGINE

DOWNSTREAM DILUTION, 25% OF TOTAL COMBUSTOR AIRFLOW
REACTOR DIAMETER, 16 cm
COMBUSTOR EXIT TEMPERATURE, 1310 K

<table>
<thead>
<tr>
<th>Speed % full</th>
<th>Total air flow, kg/s</th>
<th>Total fuel flow, 10^-3 kg/s</th>
<th>Reactor air flow, kg/s</th>
<th>Reactor f/a</th>
<th>Reactor inlet temperature, K</th>
<th>Reactor exit temperature, K</th>
<th>Pressure, 10^5 Pa</th>
<th>Reactor inlet velocity, m/s</th>
</tr>
</thead>
<tbody>
<tr>
<td>50 (idle)</td>
<td>0.127</td>
<td>0.455</td>
<td>0.0952</td>
<td>0.00478</td>
<td>1210</td>
<td>1350</td>
<td>1.43</td>
<td>11.4</td>
</tr>
<tr>
<td>60</td>
<td>0.204</td>
<td>1.135</td>
<td>0.153</td>
<td>0.00742</td>
<td>1140</td>
<td>1375</td>
<td>1.91</td>
<td>12.9</td>
</tr>
<tr>
<td>70</td>
<td>0.254</td>
<td>1.705</td>
<td>0.190</td>
<td>0.00885</td>
<td>1100</td>
<td>1385</td>
<td>2.28</td>
<td>12.9</td>
</tr>
<tr>
<td>80</td>
<td>0.336</td>
<td>2.77</td>
<td>0.252</td>
<td>0.0110</td>
<td>1045</td>
<td>1400</td>
<td>2.92</td>
<td>12.7</td>
</tr>
<tr>
<td>90</td>
<td>0.409</td>
<td>3.84</td>
<td>0.307</td>
<td>0.0125</td>
<td>1005</td>
<td>1405</td>
<td>3.56</td>
<td>12.2</td>
</tr>
<tr>
<td>100</td>
<td>0.517</td>
<td>5.41</td>
<td>0.388</td>
<td>0.0140</td>
<td>970</td>
<td>1425</td>
<td>4.50</td>
<td>11.8</td>
</tr>
</tbody>
</table>
**Figure 1.** Predicted thermal NOx emissions from a well-stirred reactor (ref. 2). Combustor inlet temperature, 970 K; combustor exit temperature, 1310 K; combustor pressure, 5x10^5 Pa; vehicle fuel consumption, 0.0775 kg/km.

**Figure 2.** Schematic representation of a catalytic reactor.
Figure 3. - Rig schematic. (Dimensions in cm.)
WITH 30-DEGREE VANE AIR SWIRLER (FIG. 4(d))

WITHOUT AIR SWIRLER

Figure 5. - Spatial fuel distribution. Inlet temperature, 800 K; pressure, 5x10^5 Pa; reference velocity, 20 m/s; mean (metered) fuel-air ratio, 0.61; splash-groove fuel injector.

Figure 6. - Effect of inlet air temperature on degree of vaporization. Pressure, 5x10^5 Pa; reference velocity, 20 m/s; fuel-air ratio, 0.010.
Figure 7. - Catalyst evaluation test section. (Dimensions in cm.)
Figure 8. - Typical catalyst elements.
Figure 9. - Tradeoff between pressure drop at 1400 K and minimum required exit temperature. Inlet mixture temperature, 800 K; pressure, 3x10^5 Pa; propane fuel.