Controlled Combustion—An Approach for Reducing Irreversibilities in Energy Conversion

Investigators
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Introduction
In conventional energy conversion devices (e.g., gas turbines and piston engines) useful work is produced in a two-step process: conversion of chemical energy to sensible energy by combustion, followed by work extraction by expansion. The first of the two processes typically occurs on a very short time scale and is essentially complete before the second process begins. The reaction between fuel and oxygen in these devices, yielding product species (H₂O, CO₂) and heat, is unrestrained and highly irreversible, resulting in a second law efficiency that is significantly less than unity. If this irreversibility can be reduced (i.e., modifying the process such that it is more reversible) then the efficiency of energy conversion is improved, resulting in fuel savings and decreased emission of CO₂. One approach to accomplish this is by closely matching the rates of heat release and expansion, effectively extracting work while the reaction proceeds. The simultaneous reaction and work extraction is much like the fuel cell process. Matching heat release and expansion rates requires control of the combustion process.

Controlled combustion may be achieved by carefully controlling the temperature and composition of the reactants, thereby tailoring the rate at which fuel and oxygen react. This type of combustion, where the reaction rate is dominated by elementary reaction steps as opposed to heat or mass transport, may be achieved by a combination of waste heat recovery and exhaust gas recirculation. In addition to the improved thermal efficiency, this process decreases the peak flame temperature which can decrease NOₓ emissions by orders of magnitude. The reduction in peak temperature may also help reduce heat losses in combustion devices, further improving efficiency. While various forms of controlled combustion have been investigated, commercial application has been limited to industrial furnaces.

Controlled combustion, which occurs at conditions of high preheat and low O₂ concentration, is the result of chemical processes which are fundamentally different from conventional combustion processes. Detailed kinetic models, which are currently optimized for conventional combustion, are needed to model implementation of controlled combustion in low-irreversibility engines. Development of an engine that utilizes controlled combustion will rely heavily on detailed kinetic models developed and validated specifically for these conditions. Since the engine combustion process would occur at elevated pressure, the influence of pressure on the reaction kinetics should also be investigated. The influence of product species, especially CO₂, on reaction rates as well as the formation of intermediate species also are of interest.
Figure 1: The regimes of combustion with variations in O$_2$ concentration and preheat temperature.

Background

Figure 1 shows the regimes of combustion processes as determined by preheat temperature and the concentration of O$_2$ in the oxidant mixture, with controlled combustion occurring at low O$_2$ concentration and high air preheat. Controlled combustion is being experimentally investigated in a high pressure flow reactor facility where these parameters, along with fuel composition and concentration, can be varied independently at pressures up to 50 bar and temperatures up to 1400 K. This facility, shown in Figure 2, has been developed specifically for the investigation of combustion chemistry at high pressure. Fuels being investigated include methane, ethane, and methane-ethane mixtures (to simulate natural gas).

Axial temperature profiles in the reactor are measured using a thermocouple, and the concentrations of stable gaseous species in the reactor are sampled by extraction through a water-cooled quartz probe. Concentrations of key reactant, intermediate, and product species in the sample gas are measured online. The measured concentration profiles can be compared with profiles predicted by detailed reacting flow models.

The starting reaction mechanism used to plan the experiments and analyze the data, GRI-Mech 3.0, was developed by the Gas Research Institute for conventional combustion of natural gas$^6$. This model, which includes 325 elementary reactions among 53 species, is used in combination with codes from the CHEMKIN-III$^7$ suite. For a particular set of reactor conditions and system constraints, the reaction rates and the resulting evolution of species concentration over time can be predicted.
Gas Analyzers

Figure 2: Schematic of the flow reactor in the high pressure facility.

Results
Comparisons of measured profiles of stable species concentrations and temperature for controlled combustion of $\text{C}_2\text{H}_6$ at ambient pressure with predicted profiles are shown in Figures 3 and 4 for a pressure of 1 bar. Figure 5 shows this comparison for a pressure of 2 bar. Measured and predicted temperature, $\text{O}_2$ and $\text{CO}_2$ profiles are in good agreement in all of the experiments; however, discrepancies between the measured and predicted profiles of the fuel, $\text{C}_2\text{H}_6$, and $\text{C}_2\text{H}_4$ and $\text{CO}$ are observed. In addition to $\text{C}_2\text{H}_4$, trace amounts of $\text{CH}_4$ are observed; both of these species are products of the initial stages of the fuel oxidation reaction. One significant feature of the fuel oxidation process is that the oxidation rate decreases as pressure increases, a result not observed under normal combustion conditions for small hydrocarbons.

To determine the reactions that are important in initial fuel oxidation and CO formation, sensitivity analyses were carried out using Senkin. Two features of the CO profile were used in these analyses – the time-to-peak CO concentration and the half-width of the CO profile. Representative results from these sensitivity analyses are shown in Figure 6. For the conditions of the experiments in Figures 3 – 5, reactions involving the peroxy radical, $\text{HO}_2$, are among the most important in the initial fuel oxidation. Because of the relative stability of this radical, its role in the reaction mechanism tends to be inhibitive. At a pressure of 2 bar, predicted $\text{HO}_2$ levels are higher than at a pressure of 1 bar, which is a partial explanation for the decreased reaction rate with increasing pressure. The rate parameters for many of the reactions involving $\text{HO}_2$ in the starting oxidation mechanism have relatively large uncertainties, and this is an area for future improvement of the mechanism.
Figure 3: Comparison of measured and predicted time profiles of temperature and stable species. Pressure = 1 bar; initial temperature = 1170K; 1450 ppm C$_2$H$_6$; 3.0% O$_2$.

Figure 4: Comparison of measured and predicted time profiles of temperature and stable species. Pressure = 1 bar; initial temperature = 1140K; 1450 ppm C$_2$H$_6$; 0.5% O$_2$.

Figure 5: Comparison of measured and predicted time profiles of temperature and stable species. Pressure = 2 bar; initial temperature = 1250K; 1050 ppm C$_2$H$_6$; 0.5% O$_2$. 
Figure 6: Results from sensitivity analysis for CO for the conditions of Figure 3 (left) and Figure 4 (right).

Figure 7 shows measured and predicted temperature profiles for ethane and methane at the same initial temperature and O$_2$ concentration. The oxidation rate for methane under these conditions is significantly slower than for ethane. The agreement between measured and calculated temperature profiles is good.

Progress
Controlled combustion of ethane and methane has been demonstrated in the flow reactor. The data obtained indicate that the combustion reaction can be inhibited sufficiently such that the combustion process can be spread over tens of milliseconds.
Initial model predictions show reasonable agreement with the measured temperature, CO$_2$ and O$_2$ profiles. However, predicted CO and stable hydrocarbon species profiles are significantly delayed compared to the experimental profiles. Sensitivity analyses have identified reactions of the hydroperoxyl radical, HO$_2$, as important elementary reactions that impact the initial stages of the reactions and, given that the rate parameters of these reactions are uncertain, some refinement of the chemical model will be required. A factorial design of the experimental is being completed in order to establish the minimum number of experiments required to establish the dependence of the controlled combustion process on the major independent variables.

**Future Plans**

Over the next few months, experiments at other pressures, O$_2$ concentrations and fuel types are in progress. Experimental data will be used in conjunction with the detailed chemical reaction mechanism to optimize the mechanism for use in studies of the application of controlled combustion to low-irreversibility engine cycles. These cycles are the subject of another project under GCEP support.  

**References**


[http://www.me.berkeley.edu/gri_mech/](http://www.me.berkeley.edu/gri_mech/)


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