The Early Phase of Spark Ignition

by

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ABSTRACT

In this dissertation, some practical ignition techniques are presented that show how some problems of lean-burn combustion can be overcome. Then, to shed light on the effects of the ignition techniques described, the focus shifts to the more specific problem of the early phase of spark ignition. Thermal models of ignition are reviewed. These models treat the energy provided by the electrical discharge as a point source, delivered infinitely fast and creating a spherically symmetric ignition kernel. The thesis challenges the basis of these thermal models by reviewing the work of many investigators who have clearly shown that the temporal characteristics of the discharge have a profound effect upon ignition. Photographic evidence of the early phase of ignition, as well as other evidence from the literature, is also presented. The evidence clearly demonstrates that the morphology of spark kernels in the early phase of development is toroidal, not spherical as suggested by thermal models. A new perspective for ignition, a fluid dynamic point of view, is described. The common ignition devices are then classified according to fluid dynamics. A model describing the behaviour of spark kernels is presented, which extends a previously established mixing model for plasma jets, to the realm of conventional axial discharges. Comparison of the model behaviour to some limited data is made. The model is modified by including the effect of heat addition from combustion, and ignition criteria are discussed.

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Dedication

For my father, Philip Donald Pitt, on the occasion of his seventy-fifth birthday, June 14, 1995.
Introduction

1.1 Environmental Impact on Combustion Engine Design

Combustion has been used by humans as a source of energy for a long time. It was Prometheus who, it is claimed, stole fire from the gods and gave it to Earth's more common inhabitants. Unfortunately, for his efforts Prometheus wound up chained to a rock where he was lashed by the sea and burned by a relentless sun. Perhaps Prometheus' ultimate fate foretold of dire consequences should humans misuse fire. We have. Are we and future generations to wind up chained to a decaying economic and social system and lashed by shifting climatic change, only to be burned in a global greenhouse? Thank you, Prometheus.

There is growing awareness of the environmental effects of energy use by society. Consequently, there is an increasing demand to design systems that use energy more efficiently and, at the same time, render the by-products of energy use more environmentally benign. To help society realize these new system design goals, details of the basic physical processes involved in energy use need to be understood. It is one such detail that is examined in this thesis.
In 1988, various estimates placed the global demand for energy near 318 exajoules (one exajoule = \(10^{18}\) joules). Of this amount, 88% was satisfied by the combustion of coal, oil or gas. Essentially an oxidation reaction, combustion is of practical interest only because of its exothermicity (Oppenheim 1985). The internal combustion (IC) engine is a familiar application of combustion. Used primarily for transportation, the IC engine consumes one out of three barrels of oil and contributes 15% of the carbon dioxide emissions in North America. In a typical year, IC engines in North America produce more than \(10^{16}\) piston combustion events, each of which lasts approximately 10 msec and releases about 1 kJ of energy. Even modest improvements in our understanding of the single piston combustion event may lead to improvements of efficiency that are measurable when multiplied so many times. Understanding the energy use details of the IC engine is thus very important in the context of energy systems and their environmental impact.

This thesis will examine the initiation process of combustion, more commonly known as ignition, in spark-ignited IC engines. At the time the work reported in this thesis was begun (late 1983), a senior executive of the General Motors Research Laboratory (Agnew 1984) stated, "If there is a need today for scientific contributions to practical spark ignition, it is in the area of the flame initiation period...." Engine designers recognized that a complete understanding of the spark-ignition process was necessary if significant improvements in the efficiency and performance of spark-ignited (SI) engines were to be realized.
Ignition, at first glance, appears to be an "old" subject, well addressed in the literature and not worthy of further investigation by a late-twentieth century apprentice scientist hoping to gain acceptance in the established scientific community. Surprisingly, however, the subject is still vigorously pursued, spurred recently by engine research. It must be remembered, nevertheless, that ignition of combustible mixtures is a fundamental problem in combustion science and its relevance goes beyond ignition in engines to include safety issues involving the storage of flammable or explosive substances.

1.2 Improvements in Engine Performance: Avenues for the Engine Designer

Present-day IC engine ignition systems have evolved over many years through a trial-and-error form of engineering. Because normal operating conditions for IC engines remained constant for nearly half a century, the spark-Ignition systems that had been developed seemed to perform adequately. However, once operating conditions necessary for high energy efficiency and lower emissions were explored, ignition problems arose. Examples of these new IC operating conditions include: states of very lean operation; high levels of exhaust gas recirculating; stratified charge mixtures; highly turbulent mixtures; and use of fuels other than gasoline.

The motivation for some of these new operating conditions comes from environmental pressure to produce more efficient engines that meet more demanding emission standards. For example, lean mixtures produce
much lower CO and NO emissions than do the normal stoichiometric mixtures prevalent for so many years. Also, the ratio of specific heats increases for lean mixtures, causing an increase in the thermodynamic efficiency of the combustion process. With increasing leanness, however, the flame propagation speed begins to degrade to the point where some of the fuel-air mixture fails to be consumed before the exhaust valve opens. This causes loss of output power and increased hydrocarbon emissions. Highly turbulent combustion chambers have been developed to augment the flame speed and to help offset the problems of lean-burn combustion. Cleaner burning alternative fuels, such as methane (natural gas), also introduce similar combustion problems when used with existing engine technology.

For one to understand better the problems faced by the IC engine designer, it is useful to review the thermodynamic cycle of the IC engine. Figure 1.1 shows the classic ideal "Otto" cycle. Superimposed is a real engine cycle, which shows the departure from the ideal case. A real engine cycle suffers from several loss mechanisms: pumping losses, timing losses, heat transfer losses and friction losses.
Figure 1.1 Ideal and real engine cycles.

Figure 1.2 indicates the approximate distribution of the chemical energy available to an engine under dynamic (equivalent "road") conditions. The dynamic efficiency (work output) of 16% improves to about 30% for steady-state operation of the same engine. An engine providing motive power to a road vehicle also has to overcome aerodynamic drag, transmission coupling losses, and dynamic friction. Further reduction of drag does not appear to be feasible, as most designs are nearing basic limits and vehicle weight reduction is limited by safety considerations. Therefore, design attention is now focusing on improvements in the real efficiency of engines. What can an engine designer do to improve engine efficiency?
A major design parameter of an engine is the compression ratio, $C_R$. The theoretical thermal efficiency of an air Otto cycle is

$$\eta = 1 - C_R^{\gamma}$$

where $\gamma$ is the ratio of specific heats.

Equation 1.1 is plotted in Figure 1.3, where apparently high compression ratios are definitely more efficient. Studies of real engines (Durbin 1980) indicate that fuel consumption improves by as much as 20% for a change in compression ratio from 7.5 to 9.6. Thus, increasing the compression ratio is the principal tool available to the engine designer for improving the efficiency of engines. For example, diesel engines, which generally operate with compression ratios of 15-22:1, are superior fuel consumers compared to spark-ignited gasoline fueled engines, which typically operate at compression ratios of <10:1.
1.3 The Fuel-Engine Relationship

The compression ratio of an engine is limited by the ability of the fuel used to resist detonation or "knock." Knock refers to the situation where the fuel-air mixture experiences a sudden energy release. This is caused by the bulk fuel-air mixture being heated when the piston compresses the fuel-air mixture, and by the initial release of energy shortly after ignition, to a point where the end-gas mixture detonates. In Figure 1.4, the combustion chamber pressure shows normal combustion and knock. The oscillatory nature of the pressure signal results from the shock wave...
produced by the detonation reflecting from the chamber surfaces. This is the source of the audible "ping" associated with knock.

Knock is a detrimental condition because the sudden energy release causes a high over-pressure inside the combustion chamber, which places a severe mechanical load on the chamber, piston, and associated bearings. For a given fuel, the onset of knock occurs above a certain compression ratio. The Research Octane Number (RON) is a measure of the knock resistance of a fuel. For higher RON, the fuel is more knock resistant and the compression ratio limit can increase.

Reduction of the knock susceptibility of a fuel at the high end-gas temperatures is accomplished at the refinery in further cracking stages. However, this additional treatment has costs in efficiency of fuel production, because more cracking stages result in fewer litres of gasoline from the original barrel of crude oil.
Knocking can also be impeded by the use of additives in the fuel. For many years, lead has been used to prevent knock. Lead, however, is environmentally—and now legislatively—unacceptable. Other additives, such as water and more air, are also well-known ways to reduce the knock susceptibility of fuel-air mixtures.

The relationship between the fuel characteristics—ignition delay, combustion time, and knock susceptibility—and the engine design parameters is crucial to the efficiency of an engine. The limitation on compression ratio due to knock is a major factor influencing the overall fuel efficiency of the world's entire automotive fleet.

The fuel-engine relationship demands that changes in the characteristics of fuel type be reflected in the engine design parameters and vice versa. This relationship has major implications, though it is often overlooked when viewed from an industry-wide perspective. The engine designers and manufacturers are looked upon as an industry separate from that of the fuel makers (oil companies). Instead, all the players should be seen as complementary components of a single transportation industry. Just how important the fuel-engine relationship is to our transportation system is illustrated in the next section.

1.4 The Need for a Holistic Systems Approach

In the previous sections, we have seen that increasing the compression ratio is the prime method for improving engine efficiency and that high
compression engines require special high-octane fuels or special methods to prevent the occurrence of knock. Up to 1970, the average compression ratio for the gasoline-fueled fleet worldwide was approximately 9.5:1. About that time, legislation was introduced in California aimed at reducing air pollution caused by automobile emissions. The legislation established stringent emission standards that would require engines to reduce significantly the emissions of unburnt hydrocarbons and oxides of nitrogen (NOx).

The engine manufacturers realized that their engines could not meet these new standards, but the engine designers knew how it could be done: redesign engines to operate under very lean conditions. This task, however, would require a major effort and a very large capital investment to cover, among other things, development costs and the cost of retooling plants. The deadline to achieve these new emission standards was imminent and, in practical terms, the engine manufacturers could not redesign and manufacture proven engines to meet this deadline.

An alternative approach, and one that did not require much time or money, was the use of the catalytic converter. These, by now familiar, objects appeared to keep the pre-1970 engine design intact by cleaning up the exhaust products. This "quick fix" approach initially seemed attractive to the engine manufacturers, except there was a serious problem. It didn't work. The reason was that a constituent of the fuel rendered the converter inoperative as soon as it was used. The guilty constituent was lead, which poisoned the active element of the converter,
platinum. The engine manufacturers literally told the fuel makers to "get the lead out" and unleaded fuels emerged into the market place along with catalytic converters. Lead, as noted in the previous section, was primarily an anti-knock agent and its removal forced the fuel makers to produce more stable (less knock-susceptible) fuels by means of further refining (cracking) processes. However, these additional refining steps not only make the final product more expensive, they also result in further waste so that ultimately there are fewer litres of unleaded gasoline produced from the original barrel of crude oil.

The unleaded fuels produced in the early 1970's caused the engine manufacturers to reduce the average industry compression ratio dramatically to 7.5:1 to reduce the tendency of the new fuels to knock. In other words, to meet the California emission standards, the catalytic converter solution required more costly unleaded fuels as well as a reduction in the engine compression ratio. The compression ratio reduction from an industry average of 9.5:1 to 7.5:1 represents a theoretical thermal efficiency reduction of 20%.

The combination of less efficient engines and the need for more highly refined fuels required an estimated increase of 15% in crude oil consumption. Unfortunately, the original aim of the California legislation, although well intended, resulted in increased oil consumption. Knowing what we do today about hydrocarbon fuel combustion and CO₂ emission, we might conclude that the introduction of catalytic converters and their associated unleaded fuels was a blunder.
Clearly it was not the fault of the original California legislation, but rather of management's decision to employ the catalytic converter as the means to achieve the new emission standards.

The decision to use the converters appears to have ignored the fuel-engine relationship, and demonstrates how major changes taken independently by engine manufacturers, forced by legislation, can have serious consequences for energy use. Instead, the fuel producers and engine manufacturers must be viewed as interdependent parts of a transportation system. It is refreshing to note that movement towards this new approach is taking place. In the August 31, 1991, issue of the Economist, a special report on Energy and the Environment observed that "America's main car manufacturers and oil companies have been talking for the past year about the way oil and engines react. Such co-operation is a novelty." This "holistic systems" approach is rooted in the fuel-engine relationship discussed in the previous section.

History is perhaps ready to repeat itself, though. Again California legislation is requiring 2-10% of the new fleet sold in that state in the year 1999 to be made up of zero emission (i.e., electric) vehicles. A laudable goal, but will nuclear power plants mushroom to meet the increased electrical demand?
1.5 Outline of this Thesis

The thrust of this thesis, as mentioned in Section 1.2, is to study the ignition process, because of the environmentally motivated movement towards the use of leaner burning engines and alternative fuels such as compressed natural gas. These changes have led, and will inevitably continue to lead to ignition problems.

In the thesis I will address the ignition problem in a two-fold way. First, in Chapter 2, I will look at how some of the characteristics of lean combustion manifest themselves. Some practical ignition techniques will be presented to show that some of the lean-burn combustion problems can be overcome. Then, to shed light on the effects of the ignition techniques described in Chapter 2, I will shift the focus of the thesis to the more specific problem of the early phase of spark ignition. To this end, Chapter 3 reviews a thermal model of ignition from a simple ignition source. This model has features found in many ignition theories described in the literature, some of the more important ones of which are briefly reviewed. Most thermal theories of ignition treat the energy provided by the electrical discharge as a point source, delivered infinitely fast and creating a spherically symmetric ignition kernel. Chapter 4 challenges this position by reviewing the work of many investigators who have clearly shown that the temporal characteristics of the discharge have a profound effect upon ignition. Photographic evidence of the early phase of ignition, as well as other evidence from the literature, is also presented, clearly demonstrating that the morphology of spark kernels in
the early phase of development differs from that held in the dominant view of the past 50 years.

Chapter 5 presents a new perspective for ignition, a fluid dynamic point of view. I will show how the common ignition devices of Chapter 2 can be classified according to fluid dynamics. I will then present a model, which extends a previously established mixing model for plasma jets, to the realm of conventional axial discharges, and compare the model behaviour to some of the limited data available.

In Chapter 6, the effect of heat addition from combustion on the evolution of spark kernels will be included in the model of Chapter 5, and ignition criteria discussed. Chapter 7 presents the conclusions of this thesis.
2 Practical Ignition Techniques for Lean-Burn or Alternative Fueled Engines

2.1 Combustion in a Box

We begin our study of spark ignition by examining a very simple closed combustion system, illustrated in Figure 2.1. The system consists of an enclosed vessel that can be pressurized up to four atmospheres with any desired mixture ratio of fuel and air. Ignition devices to be studied are attached to the vessel and various parameters—such as stored electrical energy, discharge current, and voltage—are measured. The ignition device is "fired" and a piezoelectric pressure transducer records the interior pressure of the vessel during the combustion event.

![Figure 2.1 Simple closed combustion system.](image-url)
A sample pressure trace from a combustion event, shown in Figure 2.2, uses a conventional spark ignition source, standard automotive industry electronics, and propane as the fuel. Propane has a flame speed similar to that of gasoline and is chosen for that purpose. The figure also helps to define two combustion parameters of interest: ignition delay time and combustion duration.

![Pressure trace from simple combustion system. The percentages refer to the peak pressure.](image)

An experiment is now performed where the only parameter varied is the normalized air:fuel ratio. The normalized air:fuel ratio, $\lambda$, is obtained by taking the ratio of the air and fuel volumes (or the masses) present in the mixture and dividing by the chemically correct air:fuel ratio. For example,
the chemically correct or stoichiometric ratio for an air-methane mixture is 9.55:1. Stoichiometric mixtures would thus have a normalized air:fuel ratio of $\lambda = 1.0$. Lean mixtures would have $\lambda > 1.0$ and rich mixtures would have $\lambda < 1.0$. In the experiment, the normalized air:fuel ratio is changed from stoichiometric ($\lambda = 1.0$) to increasingly leaner conditions by increments of 0.1. Results of such an experiment are shown in Figure 2.3. Most notable in the results is the increase in both the ignition delay time and combustion duration as the air:fuel ratio becomes leaner.

![Figure 2.3 Pressure-time histories as the mixture becomes increasingly leaner. Numbers refer to normalized air:fuel ratio $\lambda$.](image)

A second experiment is now performed with this simple combustion system, first by using a stoichiometric mixture of propane and air, and second by substituting the propane-air mixture with a stoichiometric methane-air mixture. The purpose of this experiment is to illustrate the effect on the combustion system of changing fuels. Methane's flame speed is significantly slower than propane's. The results (Figure 2.4) show that
switching between propane (and, by inference, gasoline) and methane has an effect similar to that of going to leaner propane-air (or gasoline-air) mixtures.

From these simple experiments, it is evident that lean-burn conditions or alternative fuels like methane present dramatically altered operating conditions. For combustion systems such as IC engines that require rapid combustion processes, output power may be severely degraded under these new operating conditions.

### 2.2 Ignition Techniques for Lean-Burn Conditions

A conventional ignition system for IC engine applications consists of a small (1 - 2 mm) gap separating two electrodes which, when triggered, discharge a few millijoules of stored electrical energy over a period of a few
milliseconds. This type of discharge, as shown in the previous section, results in slow combustion speeds when mixtures are lean. In this section, we look at some techniques that reduce ignition delay and combustion duration in lean mixtures. First, we examine results of experiments employing these ignition techniques in quiescent combustion bombs. We then follow the transition of the observed effects in bombs as these techniques are used in more realistic IC engine combustion chamber environments.

2.2.1 Plasma Jet Ignition

Ignition by plasma jet has received much attention during the past decade, since the results of the first experiments using this technique were reported by Topham et al. (1975) and Wienberg et al. (1978). The basic design of a plasma jet system consists of coaxial electrodes, one of which is recessed into a blind cavity. Stored electrical energy is discharged through this gap in 20 - 50 μsec. Reviews of the published work on plasma jet ignition can be found in Dale and Oppenheim (1981) and Clements (1984).

Orrin et al. (1981) have argued that plasma jet ignition derives its impressive effects from the production of hydrogen radicals that augment the combustion process. Their work was extended by the following series of experiments. A plasma jet igniter was modified to accept different gases bled into the cavity. The details of the experiment are contained in Ridley et al. (1984) (see Appendix I) The results are discussed here.
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Lean methane-air mixtures in a quiescent bomb were ignited by the modified bleed plasma jet. Pulsed shadowgraph photographs were used to estimate the flame kernel's mean area as a function of time. These results are shown in Figure 3 of Ridley et al. (1984) (see Appendix I) where the radius of the growing kernel is plotted vs time for three types of ignition devices: a conventional igniter, a plasma jet, and the modified bleed plasma jet. It can be seen that adding different gases to the cavity of the plasma jet has a measurable effect on the growth rate of the kernel.

Measurements of the rate of pressure rise (dP/dt) in the bomb after ignition with the bleed plasma jet are shown in Figure 4 of Ridley et al. (1984) (see Appendix I), where the maximum dP/dt observed for a given bleed gas is plotted vs the heat of combustion of the bleed gas. The measurements are for a very lean ambient mixture (\( \lambda = 2.0 \)). The rate of pressure rise appears to be strongly correlated to the heat of combustion for the bleed gas and not to the number of H atoms per mole of the associated bleed gas.

The plasma jet ignition technique seems to be promising for lean-burn conditions. However, the large energy requirement (1 J per firing) and, more particularly, the serious erosion of the electrodes due to the high current levels in the discharge suggest that practical implementation of this technique may be difficult.

2.2.2 Puff Jet Ignition

Once the plasma jet's plume was understood theoretically (Topham et al. 1983), a mechanical analogue of the plasma jet was developed (Pitt et al.
This technique, known as "puff jet ignition," attempts to produce the positive ignition qualities of the plasma jet, but without the high power requirement or the concomitant electrode erosion.

A puff jet ignition experiment, in a combustion bomb configuration, is shown in Figure 2.5. In the puff jet technique, a very small amount of gaseous fuel (e.g., methane) is puffed into the combustion chamber through a fast-acting valve. The puff rapidly mixes with the lean, ambient mixture as the puff advances into the chamber. Conventional electrodes are placed downstream from the valve orifice (distance 0.5-1.0 cm) and the conventional discharge circuit is triggered to fire at a time $\Delta t$ after the valve opens. A more detailed description of this technique is available in Appendix I.

![Figure 2.5 Schematic of puff jet system.](image)

Figure 3 of Pitt et al. (1984) (see Appendix I) shows a schlieren image of the turbulent element produced by the puff jet compared with that produced by
a plasma jet in air. The similarity of the two turbulent elements is striking. Results of an experiment when lean ($\lambda = 1.3$) mixtures of methane-air are ignited by three ignition sources are shown in Figure 2.6. In this figure, the pressure-time histories of the plasma jet and puff jet are very similar. Therefore, both ignition delay and combustion time of lean mixtures can be reduced using an ignition device that resembles a plasma jet but does not require 1 J of stored energy, or suffer a limited life due to electrode erosion.

![Figure 2.6 The relationship of bomb pressure $p$ and time $t$ for the puff jet, plasma jet and conventional spark at a starting bomb pressure of 5000 kPa and a normalized air:fuel ratio of 1.3.](image-url)
2.2.3 Fast-Discharge Ignition

A series of experiments into the physics of the spark discharge and its ignition properties was carried out by investigators at the University of Stuttgart. Significant progress was made towards understanding the energy exchange between a discharge and the local mixture.

Maley et al. (1978) showed striking results of ignition produced by conventional electrode configurations employing fast-discharge (<100 ns) circuits with stored electrical energies of approximately 30 mJ. The authors observed rapid growth rates of ignition kernels in lean mixtures.

The reduced ignition delay and combustion times in lean mixtures are similar to those times produced by the previous ignition devices.

2.2.4 Conclusions from Combustion Bomb Experiments

The results reviewed above demonstrate that ignition delay and combustion times of very lean mixtures can be significantly reduced with practical techniques using apparently different approaches. These results, however, are for quiescent, homogeneous mixtures. What happens in conditions of higher pressures and varying degrees of ambient turbulence is most relevant to ignition device application.

In the next section, actual engine experiments using the ignition devices discussed here are reviewed. We investigate how the results obtained from
combustion bomb experiments translate to combustion chamber conditions more typical of engines.

2.3 Ignition Experiments in Engines

The primary difference between engine combustion chambers and the simple quiescent combustion bomb is the level of ambient mixture motion. Residual turbulence from the intake of a fresh air-fuel mixture into the combustion chamber is subsequently dominated by turbulence produced by the rapid upward motion of the piston. Modern combustion chamber design focuses on enhancing the intake (swirl) turbulence and the turbulence produced by the piston motion. The motivation for enhancing the level of ambient turbulence is to speed up the combustion process, especially for lean air-fuel mixtures.

Instead of a comprehensive review of work investigating the effects of a multitude of engine parameters on the combustion process, we will now review some selected work to illustrate the transition from combustion bomb experiments to experiments involving real engines. Observations and conclusions drawn from quiescent bomb experiments must be mitigated by the important influences of turbulent flows of various scales inside real engine combustion chambers.
2.3.1 Plasma Jet Ignition in an Internal Combustion Engine

In Section 2.2.1, we discussed the dramatic effects observed when a quiescent mixture, especially a lean mixture, is ignited by a plasma jet. In a series of experiments motivated by the interest in methane as an alternative fuel, the effect of a plasma jet igniter on a simple, single cylinder engine was examined (Pitt and Clements 1983). The engine was run at a constant speed, with a wide open throttle. The air:fuel ratio was varied by altering the air flow but keeping the methane flow constant at all times. Further details can be found in the Appendix I. A comparison between brake horse power (BHP) produced using a plasma jet and a standard auto industry capacitive discharge ignition system (CDI) is shown in Figure 1 of Pitt and Clements (1983) (see Appendix I).

The most interesting result obtained was from an analysis of the combustion chamber pressure-time histories. An example, shown in Figure 2.7, illustrates the change in the product $pV'$ where $p$ is the cylinder pressure, $V$ is the cylinder volume, and $\gamma$ is the ratio the mixture's specific heats. This product is constant for isentropic processes (compression or expansion), but changes dramatically during the combustion process. The log $pV'$ curves for the plasma jet and the capacitive discharge ignition system are almost identical, except that the latter was fired more than 2 ms before the plasma jet in the example shown. It therefore appears that the primary effect of the plasma jet is to reduce the ignition delay. However,
dramatic reduction of the combustion time is not observed, unlike in combustion bomb experiments.

![Graph](image)

Figure 2.7 Log $pV^\gamma$ vs crank angle for $\lambda = 1.0$ and optimal time range for both ignition sources. Engine speed is 2000 rpm.

2.3.2 Puff Jet Ignition in Internal Combustion Engines

The application of the puff jet ignition to IC engines was first reported by Pitt *et al.* (1984). An example of combustion chamber pressure from a test engine using puff jet, plasma jet, and conventional ignition systems is shown in Figure 2.8. These results indicate that puff jet ignition characteristics are comparable with those created by plasma jet ignition, as seen in the previous combustion bomb experiments. Continuous running engine tests (Fisher *et al.* 1986) further demonstrate the similarity of puff jet and plasma jet ignition.
Figure 2.8 Variation of average pressure waveforms as a function of normalized air:fuel ratio. Ignition timing fixed at 50° BTDC. Vertical scale: 440 kPa/div; horizontal scale: 36° CA/div. TDC is located at the centre line of the grid. Ignition sources denoted by: PJ, plasma jet; PF, puff jet; SG, extended electrode spark gap. (a) $\lambda = 1.0$; (b) $\lambda = 1.3$; (c) $\lambda = 1.5$. Note in (a), the puff jet and spark gap sources produced averaged pressure waveforms that are identical.
2.3.3 Fast-Discharge Ignition

Maly et al. (1983) reported experiments on test engines using fast-discharge techniques. Reduction of ignition delay for lean mixtures and extension of the lean misfire limit were observed. These effects are identical to those observed under similar conditions using the plasma jet and puff jet techniques.

2.4 Conclusion

Practical means to ignite lean mixtures do exist. The techniques discussed all produce similar results. Nevertheless, we must conclude from the preceding results that moving from the quiescent combustion bomb to the more turbulent combustion chamber environment of an engine reduces the impact of these ignition techniques. The predominant effect of the techniques over conventional ignition systems is the reduction of ignition delay. Combustion duration is not affected. The bulk turbulence of the engine combustion chamber determines the overall flame propagation speed, and hence the combustion duration. The ignition device can, however, affect the early (1-2 ms) evolution of the combustion event in a real engine. It is for this reason that we shift the focus of this thesis to that early phase of ignition.
3 The Thermal Approach to Spark Ignition

The focus of the thesis will now move from a discussion of ignition device examples and the practical problems associated with lean burn or alternative fueled IC engines to the more specific problem of the early phase of spark ignition. There are two reasons for doing so: the need to examine the divergent electrical discharge ignition theories in the literature; and the need to understand, from basic principles, the ignition devices discussed in Chapter 2.

We begin by examining the simplest electrical discharge ignition device—the axial electrode gap. The basic ignition problem can then be stated: How does the spark energy deposited in the gap become distributed from the narrow breakdown channel into the spark kernel structure observed, and what processes govern the evolution of the structure in a combustible mixture? To begin to answer these questions, we review a physical model that is basic to most models of spark ignition.

3.1 Basic Thermal Model

A recent edition of a well-known monograph on combustion (Glassman 1987) includes a theoretical description of spark ignition based on the thermal approach of Zeldovich et al. (1940). The thermal model assumes that the energy of a capacitive spark discharge is delivered to the combustible mixture as a point heat source, and that thermal diffusion
will lead to an exponential temperature distribution throughout a spherically symmetric spark kernel. The model, though simple, represents current common understanding of the spark ignition process and it is therefore important to review. We will show in Chapters 4 and 5, however, that this viewpoint misrepresents the basic physical process involved.

For ignition to happen, the spark kernel should achieve or exceed a certain radius in a time faster than the cooling (due to conduction) can reduce its temperature below the laminar flame temperature. It is instructive to review the basic assumptions and equations of this theory.

First, it is assumed that the energy \( E = \frac{1}{2} C(V_1^2 - V_2^2) \), where \( V_1 \) is the initial voltage and \( V_2 \) is the voltage after the discharge, is deposited in the spark gap as a point source. This energy then spreads according to the spherically symmetric heat equation:

\[
\left( \frac{\partial}{\partial t} - \Omega \frac{\partial^2}{\partial r^2} \right) T = 0 \tag{3.1}
\]

where \( \Omega = k / c_p \rho \) and the boundary conditions are \( T = T_0 \) at \( r = \infty \) and \( \frac{\partial T}{\partial r} = 0 \) at \( r = 0, \infty \).

The solution of (3.1) with these boundary conditions is:

\[
T - T_0 = \frac{Q}{c_p \rho (4\pi \Omega t)^{\frac{3}{2}}} \exp \left( \frac{-r^3}{4\Omega t} \right) \tag{3.2}
\]
The criterion for ignition is that the time required for the on-axis temperature \(T(r=0)\) to drop by an amount \(\phi\) must be greater than the reaction time in the combustion zone of a laminar flame. The temperature difference \(\phi\) is taken to be the difference between the laminar flame temperature \(T_f\) and an estimated ignition temperature \(T_i\).

The cooling time \(t_c\) can be estimated from

\[
t_c \approx \frac{\phi}{\left(\frac{dT_r}{dt}\right)_{r=r_i}}
\]  
(3.3)

Before evaluating (3.3), the energy \(Q\) in (3.2) must be defined. This energy is taken to be the energy required to heat a spherical volume of radius \(r_f\) from \(T_0\) to \(T_f\). Thus:

\[
Q = \frac{4}{3} \pi r_f^3 c_p \rho (T_f - T_0)
\]  
(3.4)

Evaluation of (3.3) using (3.4) yields:

\[
t_c = 0.14 \left[\frac{\phi}{T_f - T_0}\right] (r_f^3 / \Omega)
\]  
(3.5)

An estimate for the reaction time \(t_r\) can be obtained from knowledge of the laminar flame speed \(S_L\) and thickness \(\Delta\). The reaction time is related to those quantities through:
The laminar flame speed, $S_L$, can be determined through the calculation first presented by Mallard and Le Chatelier (1881). It is based on the assumption that the heat conducted from zone I (see Figure 3.1) would equal that amount necessary to raise the temperature of the unburnt gases to the ignition temperature $T_I$.

The enthalpy balance across the flame is:

$$\Delta = S_L t_r$$  \hspace{1cm} (6)

$$\frac{dm}{dt} c_p (T_f - T_0) = k \frac{dT}{dx} A$$  \hspace{1cm} (3.7)
where \( A \) is the cross-sectional area perpendicular to the flame front. The mass rate \( \frac{dm}{dt} \) is the rate at which the combustion wave consumes the unburnt gas. The combustion wave is propagating at the laminar flame speed \( S_L \) and this is related to the mass rate by:

\[
\frac{dm}{dt} = \rho S_L A
\]  

(3.8)

The temperature derivative can be approximated by assuming the temperature profile to be linear:

\[
\frac{dT}{dx} = \frac{(T_f - T_i)}{\Delta}
\]  

(3.9)

Substituting (3.8) and (3.9) into (3.7) yields:

\[
S_L = \frac{\Omega}{\Delta} \frac{(T_f - T_i)}{(T_f - T_o)}
\]  

(3.10)

Using (3.10) in (3.6), we obtain an expression for the reaction time \( t_r \) in terms of the temperatures difference \( \phi \):

\[
t_r = \frac{\Omega}{S_L} \frac{\phi}{(T_f - T_o)}
\]  

(3.11)

where \( \phi / (T_f - T_o) \ll 1 \) has been used.

This reaction time is a factor of two faster than that calculated by Zeldovich et al. (1940), but it illustrates the basic assumptions involved.
Now, with an expression for $t_c$ and $t_r$—equations (3.5) and (3.11)—the ignition criterion, $t_c \geq t_r$, can be calculated explicitly as:

$$r_j^3 \geq \Omega^2 / (0.14S^2_L)$$  \hspace{1cm} (3.12)$$

The ratio $\Omega / S_L$ has the dimension of length and can be interpreted as being related to the flame thickness because equation (3.10) can be put into the form:

$$\frac{\Omega}{S_L} = \left( T_f - T_0 \right) \Delta / \phi$$  \hspace{1cm} (3.13)$$

Therefore:

$$r_j \geq 2.67 \left( T_f - T_0 \right) \Delta / \phi$$  \hspace{1cm} (3.14)$$

The physical implication of (3.14) is that the equivalent heat radius $r_f$ must be significantly greater than the laminar flame thickness $\Delta$.

Furthermore, the concept of minimum ignition energy can be introduced. The value $r_{\text{min}}$, as defined by the right-hand side of the inequality (3.12), is:

$$r_{\text{min}} = 7.1 \frac{\Omega}{S_L}$$  \hspace{1cm} (3.15)$$

Therefore

$$Q_{\text{min}} = \frac{4}{3} \pi r_{\text{min}}^3 c_p \rho (T_f - T_0)$$
The main result of the thermal approach to ignition is the estimate of \( Q_{\text{min}} \) given by (3.16) and its apparent dependence on physical parameters, (such as the thermal diffusivity), and chemical parameters, (such as flame speed). Much effort has gone into measuring \( Q_{\text{min}} \) for various fuel-oxidizer mixtures, but it was recognized early on that the experimental values of \( Q_{\text{min}} \) were strongly affected by spark gap length. There existed a spark gap length \( d_q \), below which ignition of the mixture was not possible, and this length was termed the "quenching distance". The interpretation was that excessive cooling by the electrodes caused quenching of the ignition kernel at spark gap lengths less than \( d_q \). It is interesting to associate the quenching distance \( d_q \) with \( r_{\text{min}} \) given by (3.15), which would imply that \( Q_{\text{min}} \) is proportional to \( d_q^3 \).

3.2 Theory of Lewis and von Elbe

Another thermal theory, that of Lewis and von Elbe (1961), was widely quoted for many years. It is based on an "excess enthalpy" assumption which presumes that in a thin shell ahead of a spherically symmetric flame front, the enthalpy is greater than in the surrounding burned or unburned gases. Again, as in the theory of Zeldovich et al. (1940), a critical diameter for the ignition kernel must be attained such that the excess enthalpy is supplied by the heat content of the burned gases in
the kernel, allowing the combustion wave to propagate. The ignition source must provide the minimum ignition energy to achieve this critical kernel size. Lewis and von Elbe (1961) calculate it to be:

\[ Q_{\text{min}} = n d_0^2 \frac{k}{S_L} (T_b - T_0) \]  

(3.17)

where \( T_b \) is the burned gas temperature. This relationship was tested by Lewis and von Elbe and by Calcote et al. (1952) who found \( Q_{\text{min}} \) to be proportional to \( d_0^2 \). They measured minimum ignition energies for quiescent, room-temperature hydrocarbon mixtures at atmospheric pressure. Typical U-shaped ignition curves were obtained, indicating minimum ignition energies in the range of 0.1-1 mJ, and these occurred at equivalence ratios of 1.2-1.5. The excess enthalpy assumption necessary for flame propagation in their model has been criticized by Linnett (1952) and others, as experimental efforts to confirm the existence of the excess enthalpy "bump" preceding a flame front have failed to detect such an effect.

3.3 Ignition in Flowing Gases: Theory of Swett

Swett (1957) examined the situation of ignition in flowing gases and calculated \( Q_{\text{min}} \) based on the assumption that, unlike in the previous models, the ignition kernel would suffer heat loss primarily through eddy diffusion. Swett derived a complex expression for \( Q_{\text{min}} \) which was substantiated by limited experimental data. His results indicated an increase in \( Q_{\text{min}} \) with increase in mixture flow velocity or turbulence
Intensity, but showed that the scale of the turbulence had little effect on $Q_{\text{min}}$. DeSote (1971) confirmed these results.

### 3.4 Theory of Ballal and Lefebvre

Probably the most extensive investigation into spark ignition in recent years has been performed by Ballal and Lefebvre. Although their interest was investigating conditions more typical of gas turbine systems (i.e., turbulent flow at sub-atmospheric pressure), their approach and results have had much influence.

Their theory, based on observation of flame propagation in flowing mixtures, revealed that the heat release mechanism in the flame zone was very dependent on the level of turbulence (Ballal and Lefebvre 1975). The interpretation of this observation was that, under conditions of low turbulence, burning rates are enhanced by the effects of increased flame surface area caused by the wrinkling of the flame due to turbulence. Ballal and Lefebvre (1975) claim that under conditions of high turbulence, the intersection of the turbulent eddies, which have fine scale structure, creates a very large total flame surface. These model calculations are relatively simple and yet the physical basis has had a lasting impact on many other investigators.

The minimum ignition energy is obtained by assuming, as the criterion for ignition, that the principal dimensions of the kernel should everywhere equal or exceed the quenching distance. A sphere of
diameter $d_q$ heated to the adiabatic flame temperature is the minimum volume for a spark kernel, and this can be used to compute an estimate of $Q_{min}$:

$$Q_{min} = \frac{\pi}{6} c_p \Delta T \, d_q^3 \quad (3.18)$$

Using the expressions derived for $d_q$ for the two cases discussed, we obtain:

$$Q_{min} = \frac{\pi}{6} c_p \Delta T \, A \, \Omega \left( S_L - 0.16 \, u' \right)^{-1} \quad \text{low turbulence} \quad (3.19)$$

$$Q_{min} = \frac{\pi}{6} c_p \Delta T \, B \, \Omega \left( S_T - 0.63 \, u' \right)^{-1} \quad \text{high turbulence} \quad (3.20)$$

Experiments testing these relationships were conducted by Ballal and Lefevbre (1977) and excellent agreement was obtained. However, all their data were at sub-atmospheric pressures. Unlike Lewis and von Elbe (1961), Ballal and Lefevbre (perhaps gleefully) pointed out that their results predicted $Q_{min}$ proportional to $d_q^3$ as in the approach by Zeldovich et al. (1940). Clearly then, a basic difference between theories exists, but experiments supporting both have been reported.

### 3.5 Thermal Diffusion Model of Ko, Arapaci and Anderson

Recent work by Ko et al. (1991 b) continues the investigation of spark ignition from the point of view of a thermal diffusion process. The model
assumes that a spherically symmetric flame kernel evolves by heat
diffusion alone. Solutions to the convection-free energy conservation
equations are presented, based on the asymptotic solution method of
Champion et al. (1986) and Deshaies and Joulin (1983). The solutions
are noteworthy because of the existence of an unstable equilibrium flame
radius which is a function of the physiochemical properties of the
mixture. An ignition criterion is developed that requires successfully
propagating flames to exceed this critical radius. Supporting
experimental work (Ko et al. 1991 a) showing measurements of flame
kernel radii in quiescent mixtures demonstrates the model behaviour
well.

3.6 Discussion

The thermal model discussed in this chapter has features common to
most thermal theories. For example, the dominant process in the early
development of a flame is a balance between the heat released by the
exothermic reactions and the heat lost by conduction to the unburnt
mixture. The spark energy is used only to raise a small volume to the
flame temperature; the heat balance is left to do the rest. Temporal
characteristics of the discharge largely ignored in this approach and, if
included in some approaches, they appear through initial conditions.

The concept of minimum ignition energy is a central theme in most
thermal models and it is based largely on the laminar flame speed and
the previously mentioned balance between heat release and conduction.
The laminar flame speed is a number associated with a fully developed flame, and its use in the very early phase of spark kernel development may be inappropriate. Furthermore, data published in the literature are conflicting about such quantities as $Q_{\text{min}}$ and the optimal discharge times that fail to differentiate between thermal models.

In the next chapter, the effect of the spark discharge's temporal characteristic on kernel growth is examined. We also review imaging data of real spark kernels in order to clearly establish their structure.
4 Spark Kernel Dynamics and Morphology

The relevant literature shows us that spark kernels have been the subject of scrutiny since 1945. Our purpose is not to review all this work; instead, we will direct our attention to two main aspects of spark kernels. First, what are the most important characteristics of electrical discharges that produce the spark kernels? Second, what is the observed structure of spark kernels produced by electrical discharges?

Chapter 3 addressed the above two questions in the following way: the main characteristic of the electrical discharge, from the point of view of ignition, was its energy. Secondly, the type of spark kernel structure produced was spherical. These two features lie at the basis of most thermal ignition models and have permeated much of the thinking about ignition for the past 40 or 50 years. In this chapter we challenge this point of view.

We begin by reviewing work which clearly shows how the temporal characteristics of the discharge have a profound effect on ignition. Next we present imaging data of spark kernels, both new and from the literature, and demonstrate that the morphology of spark kernels, in the early phase of development, is distinctly non-spherical by nature.
4.1 Characteristics of the Spark Discharge

In the thermal ignition models of the previous chapter, the only attribute of the spark discharge taken into account is the energy deposited in the gap. This energy was assumed to be deposited as a point source, infinitely fast.

There have been several attempts to determine which characteristics of the spark discharge are important for ignition. The results of these experimental investigations have been rather confusing, and contrary results have been published at different times. For example, results indicating an optimal discharge time show a wide variation of such times.

As far as the IC engine designer is concerned, standard engineering practice has dictated that the spark be a low energy (10 mJ), long duration (several ms) discharge. The trial-and-error engineering approach reasoned that a long discharge guaranteed ignition at some point and the low energy seemed to be what they could "get away with." This design approach failed when operating conditions moved away from the normal stoichiometric operating point, as in the move towards lean-burning engines.

Extensive investigations on the influences of spark discharge characteristics on ignition have been carried out during the past two
decades. Ballal and Lefebvre (1975) examined the effect of breakdown voltage, spark duration, energy deposition, gap width, electrode geometry, and electrode material in low pressure (sub-atmospheric) flowing gases.

Their results indicated that, for quiescent stoichiometric mixtures, minimum ignition energy was obtained for a discharge of 60 μs. Other results indicated that higher minimum ignition energies are obtained with electrodes materials having higher boiling temperatures. Ballal and Lefebvre (1975) also obtained results that would be expected on the basis of arc physics, namely that Pachen's Law (see for example Kuffel and Zaengl, 1984) was valid. It says that breakdown voltage is proportional to the product of pressure and gap length, and that the spark energy, $E_s$, is:

$$E_s = d_g \exp \left( \frac{P d_g}{\exp P} \right)$$  \hspace{1cm} (4.1)

where $d_g$ is gap length, and $P$ is the pressure. Thus, $E_s$ is most sensitive to gap length.

Good progress towards understanding the physics of spark ignition has been contributed by Maly and his colleagues from the University of Stuttgart. They examined the complete discharge process with time-resolved interferometry and spectroscopic techniques. Their results would warm the hearts of most plasma physicists. The distinct nature of breakdown, arc and glow phases of the discharge and the importance of
each phase to ignition of combustible mixtures were first recognized by these investigators.

The voltage and current characteristics of a discharge are shown schematically in Figure 4.1. The breakdown phase results in a rapid ionization of the spark channel (diameter typically 10-40 μm), with peak currents of a few hundred amps for a period of 10-100 nsec. The fully ionized channel has peak temperature of ~ 6 × 10^4 °K, and the resulting high peak pressure produces a shock wave.

![Figure 4.1 Discharge current and voltage characteristics](image)

The arc phase is characterized by a lower sustaining voltage, 50-100 V, the current is typically a few amps, and the duration is 10-100 μs. The arc continues to expand mainly due to conduction and diffusion. Heat loss, especially to the electrodes, limits the kernel to about 6 × 10^3 °K. In the glow phase, the current drops to ~ 200 mA, with a duration of 100
-1000 μsec; there is a high cathode fall of 300-500 V; and the temperature drops to $3 \times 10^3$ °K, resulting in a very low level of ionization. The bulk of the discharge time of typical igniting systems is in the glow phase. Erosion of the electrodes takes place primarily during the arc phase, more modestly during the glow phase, and negligibly during the breakdown phase.

Maly and Vogel (1978) examined the spark kernel dynamics of each of the three phases and presented striking results that indicated the breakdown phase was important to the development of the spark kernel. The radial extent of the kernel develops very rapidly as the energy in the breakdown phase is enhanced. Experiments in a combustible mixture showed that the breakdown type of discharge (10-100 ns) was most effective, and that these types of discharges created a much larger kernel than arc or glow discharges of equivalent energies.

Maly (1981) developed a thermal model for these ultra fast discharges. The shell-like structure suggested by the measured temperature profiles, indicated in Figure 4.2, is the basis for Maly's model. He assumes a radially symmetric geometry, constant pressure, and conduction as the only transport mechanism. The last assumption is taken into account by having laminar and turbulent heat fluxes lumped into a single transport coefficient $k_D$. Maly's ignition criterion is expressed as:

$$ (x_q - \Delta H_f) P_{\phi} \frac{dr}{dt} [2 R_k c_T (T_f - T_0)]^{-1} \geq 1 \quad (4.2) $$
where $x_q$ is the molar heat of combustion, $\Delta H_p$ is the average molar enthalpy required to raise the gas to the flame temperature, and $r_1$ and $d r_1/dt$ are the flame front position and velocity, respectively. This ignition criterion is most notable due to the appearance of the velocity of the leading edge of the kernel, $d r_1/dt$. It presents first example of the dynamics of the spark kernel manifesting itself in the ignition process. Maly also derived expressions for the quenching distance and minimum ignition energy, but this calculation based on the dubious excess enthalpy assumption of Lewis and von Elbe.

![Figure 4.2 Measured temperature profile of a spark kernel, after Maly and Vogel (1978).](image)

Ziegler et al. (1984) continued this investigation into the ignition qualities of the arc and glow discharge phases. Their results indicate that energy dissipated in the anode and cathode fall is completely lost to the electrodes, whereas the energy in the positive column is what is
effective for ignition by predominantly arc or glow phase discharges. This observation was originally pointed out by Swett (1957), and the results are particularly important because most previous investigations that determined the $Q_{\text{min}}$ experimentally did so by measuring the spark gap voltage and current, and then computing $Q_{\text{min}}$ by $\int \text{V} \text{d}t$. Thus, the results of Ballal and Lefebvre and others overestimate $Q_{\text{min}}$, and it is interesting that any correlation with $d_q$ was possible.

4.3 Time-Dependent Ignition Models

Other workers have attacked the problem of spark ignition with the idea of somehow incorporating the time dependence of the spark energy input.

Aldeman (1981) calculated the flow field behind the shock front produced by a discharge whose energy input is time dependent. The spherically symmetric model is notable because it is basically non-thermal, neglecting conduction as a loss mechanism. Unfortunately, his results would only compare well with data obtained by Lichfield (1961) if he chose a spark duration different from the experimental one.

The numerical calculations of Dixon-Lewis and Shepard (1975) solved the time-dependent conservation equations, incorporating multi-step reaction kinetics in both spherical and cylindrical coordinates, to examine the growth of H$_2$-air flames. Blast wave effects were ignored, however. The authors' results indicated that, for a constant total
ignition energy, flame initiation was enhance by increasing the proportion of energy supplied as H atoms rather than as thermal energy.

Similarly, Oran and Boris (1982) solved the same equations assuming spherical symmetry, but they used a complex 48-step reaction mechanism to describe the kinetics. Their results are somewhat disappointing, given the complexity of their model, as the ignition energy could not be computed with any accuracy.

Rafeal and Sher (1985) solved, numerically, the cylindrically symmetric conservation equations for methane-air mixtures at constant pressure. The radial profile of the time-dependent power input from the discharge was computed from the plasma conductivity given by Plooster (1979) and radiation losses where taken into account. The reaction kinetics were modelled by an 18-step mechanism. The breakdown energy of the discharge was used as an initial condition. The results of Rafeal and Sher's (1985) calculations show clear manifestations of combustion in the radial temperature profile as early as 10 µs after the initiation of the discharge.

Akindele et al. (1982) solved the spherical symmetric conservation equations with turbulent transport coefficients. Their calculations indicate the trends observed in their experiments, but no quantitative results were obtained for $Q_{\text{min}}$. 
4.4 Experimental Observations of Spark Kernels

In all the thermal theories reviewed here, the basic geometrical structure for the spark kernel was spherical. The numerical models assumed either spherical or axial symmetry. Surprisingly, experimental observations of the kernels produced by spark discharges, dating back to Olsen et al. (1952), have consistently shown a definite toroidal structure. This suggests that the mechanism involved in determining the underlying dynamics of spark kernels is responsible for this type of structure, and that it may be fundamental to the geometry of an axial arc discharge.

The schlieren photographs taken by Kono et al. (1976) of kernels are most striking evidence of the toroidal structure that evolves after the discharge. Chomiak (1979) was the first to note this toroidal structure and he developed a phenomenological model of the spark kernel. He suggested that hydrodynamic effects are dominant in determining the evolution of spark kernels. His model suggested a circulation pattern that is very similar to that for the ignition kernel model proposed for plasma jets by Topham et al. (1981). Plasma jets as ignition sources have been reviewed by Clements (1984), and the plasma jet ignition model was presented by Topham et al. (1984).

We have obtained schlieren and shadowgraph images of axial spark kernels in air for a wide range of discharge conditions, using a high speed video camera and a laser diode light source. Figures 4.3 and 4.4 show selected images from sequences taken of two similar discharges, viewed
along and normal to the discharge axis, respectively. A distinctly toroidal structure with sharp boundaries is clearly visible in both sets of images. The major and minor axes of the torus expand in time and, at about 500 - 700 μs after the beginning of the discharge, the structure becomes unstable. Figure 4.5 shows examples of single shadowgraph images of spark kernels produced by an electrical discharge of higher energy and longer discharge time—1.0 J stored energy, discharged over 150 μs. The kernel structure is again clearly toroidal, but developed over much longer time scales than the spark kernels of Figure 4.4. The structure appears to remain well defined and laminar out to 11 ms, the longest time interval imaged. Our results support the observations of Kono et al. (1976), which show a well-defined toroidal region of hot gases with a sharp boundary produced by axial discharges.

More evidence of a toroidal structure with a sharp boundary is provided by the temperature profiles of axial discharge spark kernel obtained by Maly and Vogel (1977), Akindele et al. (1982), Borghesse et al. (1988), and Haley and Smy (1989). These results suggest that the toroidal structure is an inherent characteristic of the aftermath of an axial spark discharge.

Experimental efforts have shown that the temporal characteristics of the spark discharge are very important for ignition. In particular, the initial spark gap breakdown energy is critical in spark kernel development. Other experiments clearly show that the kernel structure is markedly
different from the traditional theoretical view of a spherical region initiated by a point heat source.

In the next chapter, we develop a new model of spark kernel development that follows from the observations of the toroidal kernel structure. The energy delivered to the spark gap produces hydrodynamic flows that result in the well-defined kernel structures observed. Links to plasma jet kernels are made, giving spark kernel development from electrical discharge sources a common physical thread.

Figure 4.3 Schlieren images of a spark discharge, viewed along the discharge axis. The initial frame is taken 91 μs after the discharge begins. The subsequent frames are 167μs apart and the 62 mJ discharge is 50μs long.
Figure 4.4 Schlieren images of spark discharge, viewed normal to the discharge axis for the same discharge and frame times as Figure 4.3.

Figure 4.5 Shadowgraph images of spark kernels; energy 1.0 J, discharge time 150μs, and frame times (a) 3 ms, (b) 8 ms, (c) 11 ms after the beginning of the discharge.
5 A Unified Approach to Spark Kernel Development: Mixing Models

In this chapter we examine spark ignition by concentrating on the underlying fluid dynamics in the critical early phase of a spark kernel's development. From this fluid dynamic point of view, we can then show how the basic ignition process is common to the major practical ignition sources. The material in this and the following chapters first appeared in Pitt et al. (1991), which can be found in Appendix I. Some differences in notation and scaling are used.

The previous chapter showed clearly that the conventional view of a simple point heat source, diffusion model of ignition fails to predict the observed spark kernel structure. If we go further and include the major practical ignition devices discussed in Chapter 2, we have, from our literature, a picture of ignition that looks like that shown in Table 5.1.

<table>
<thead>
<tr>
<th>Igniter type</th>
<th>Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plasma jet</td>
<td>Exotic radicals (Orrin et al. 1981); turbulent mixing elements (Topham et al. 1984)</td>
</tr>
<tr>
<td>Puff jet</td>
<td>Turbulent mixing elements (Pitt et al. 1983)</td>
</tr>
<tr>
<td>Conventional axial discharge</td>
<td>Thermal diffusion (Ballal and Lefebvre 1975, Ko et al. (1991a)</td>
</tr>
<tr>
<td>Surface discharge</td>
<td>No models, some experimental data</td>
</tr>
<tr>
<td>Fast discharge</td>
<td>Modified thermal model (Maly and Vogel 1978)</td>
</tr>
</tbody>
</table>
Of the five practical devices in Table 5.1, only the plasma jet and puff jet are related by a common physical mechanism—hydrodynamic mixing. The remaining sources appear to be unrelated to the first two.

This chapter establishes the basis for a new point of view of spark ignition. The spark kernel is to be viewed as a distinct entity—a fluid element with distinct dynamic properties. From this point of view, a classification scheme is proposed that will group the practical ignition sources of Table 5.1 into two classes, based on either the spherical or toroidal symmetry of the spark kernel produced. Toroidal mixing elements are discussed in some detail, and then, in Section 5.4, a toroidal element mixing model is developed. In the final sections of the chapter, we compare observed spark kernel behaviour with the model presented and speculate on a possible mechanism to generate the necessary flow.

5.1 Underlying Dynamics of Spark Kernels

The observations described in the preceding chapter revealed an important characteristic of spark kernels from axial discharges: the existence of toroidal fluid structure with sharp boundaries. Much of the experimental evidence, then, does not support the conventional model of a point heat source creating, solely by diffusive processes, a spherically symmetric region with an exponentially decreasing radial temperature profile.

Figure 5.1 compares the temperature profiles and morphology predicted by the basic thermal model and experimental observations. The internal flow
necessary for the observed toroidal structure of kernels suggests that mechanisms such as entrainment and vortex dynamics play an essential role in the early evolution of spark kernels. This means that spark kernels produced by axial discharges belong to a class of self-propagating free-mixing elements governed by hydrodynamic flows. They therefore have certain features in common with the kernels produced by plasma jet and surface discharge devices. This common thread of fluid dynamics provides a basis for classifying various igniter types as is done in the next section.
Figure 5.1 Basic thermal model features and experimental observations.
5.2 Proposed Classification of Ignition Sources

Ignition sources used in most electrical discharge ignition systems are of three basic types, characterized by the geometry of the electrodes forming the spark gap of the ignition source. The three source types—axial discharge, surface discharge, and plasma jet—are shown schematically in Figure 5.2.

Figure 5.2 Basic igniter types showing imparted momentum (arrows) and corresponding flow patterns: a) axial discharge, b) surface discharge, c) plasma jet.

We propose a classification scheme for these ignition devices to link the basic physical processes that govern the development of the spark kernels produced by axial discharges, surface discharges, plasma jet discharges, and puff jets. It is not only the energy which is important for the
establishment of ignition, but also the manner by which it is supplied. This latter aspect leads to the classification scheme that differentiates between ignition sources on the basis of the energy, impulse, and initial vorticity distribution imparted to the spark kernel.

Experimental observations of kernels produced by plasma jet sources suggest that the kernels resemble fluid structures similar to atmospheric thermals (Topham et al. 1982). Measurements of the momentum imparted to the hot gases expelled by such sources have been reported (Smy et al. 1985). Ignition kernels produced by surface discharge sources have not received much attention. However, schlieren imagery (Bradley and Critchley 1974) indicates that the morphology of surface discharge kernels is similar to that of plasma jet kernels. Also, the momentum imparted to surface discharge kernels has been measured (Smy et al. 1985) and was found to be less than that for a plasma jet source using the same discharge current.

A model describing the evolution of spherical turbulent mixing elements (spark kernels) produced by plasma jets is well established (Topham et al. 1984). This model illustrates the role that momentum, imparted to the spark kernel, plays. Figure 5.2 indicates the axially directed momentum of spherical spark kernels and the basic circulation pattern. The initial hot kernel expelled by the plasma jet begins to entrain a cooler, ambient mixture as the kernel propagates away from the plasma jet exit. For a given discharge energy, plasma jet kernels with higher imparted momentum mix and dilute more rapidly.
The turbulent spherical mixing element model has been extended to include a simple Arrhenius type of heat addition term, to simulate combustion in an element propagating into a combustible mixture (Topham et al. 1986). In this case, the introduction of a cold mixture into the interior of the element acts to control the rate of heat release by chemical reactions. The model solutions divide into two distinct types: those for which the mean density finally approaches the ambient level, termed the "extinction branch"; and those for which the density rapidly attains a low value which is maintained at a constant level. This self-sustaining, high temperature state is called the "combusting branch." Ignition criteria based on this model are discussed in the next chapter.

The puff jet ignition source of Chapter 2 used the spherical turbulent mixing element model to predict where and when a small turbulent puff of methane would mix and dilute to near stoichiometric conditions suitable for easy spark ignition (Pitt et al. 1984). This ignition source is clearly related to plasma jet ignition kernels through a common physical process.

Surface discharge spark kernels, as noted previously, appear to be similar to plasma jet kernels, but have lower imparted momentum. Although no data are available to compare these kernels to the spherical turbulent mixing element model, we propose that these kernels should belong to this class of mixing elements.
In the case of the axial discharges, such as conventional ignition sources or fast discharges, the observed toroidal nature of the spark kernels produced suggests that the momentum imparted to the toroidal kernel is radially directed, as shown in Figure 5.2, where the circulation pattern is also indicated.

We propose that there are two main classes of fluid flow associated with the momentum imparted to a spark kernel. First, there is the axially directed momentum imparted by plasma jets, surface discharge, and puff jets, which create spherically symmetric mixing elements. Second, there is the radially directed momentum imparted by axial discharges of either conventional or fast discharge types, which create toroidal mixing elements. We suggest differentiating the two classes of ignition sources by the morphology of their resulting spark kernels. Spark kernels produced would then be of either the spherical or toroidal class.

Regardless of the class, all spark kernels evolve according to similar hydrodynamic mixing processes involving entrainment and vortex dynamics. Not only is the energy imparted to the element important, so is the momentum.

The ignition sources of Table 5.1 can now be rearranged according to the two main classes of spark kernels produced (Table 5.2).

<table>
<thead>
<tr>
<th>Table 5.2 Classification of ignition sources</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Spherical spark kernel</strong></td>
</tr>
<tr>
<td>plasma jet</td>
</tr>
<tr>
<td>puff jet</td>
</tr>
<tr>
<td>surface discharge</td>
</tr>
<tr>
<td><strong>Toroidal spark kernel</strong></td>
</tr>
<tr>
<td>conventional axial</td>
</tr>
<tr>
<td>fast discharge</td>
</tr>
<tr>
<td>discharge</td>
</tr>
</tbody>
</table>
5.3 Toroidal Mixing Elements

Having linked the basic physical processes underlying the development of spark kernels from plasma jets to those produced from axial discharges, we still must face the problem of developing a suitable model for the toroidal mixing elements. The fluid mechanics of the toroidal flow is considerably more complicated than that assumed for the self-similar entraining spherical elements to model plasma jet kernels.

Our model assumes that the toroidal element contains the double circulation pattern first suggested by Chomiak (1979) and shown in Figure 5.2. As the torus expands radially, the internal vorticity field stretches, which in turn influences the radial expansion velocity of the element itself. In addition, the small physical size of the elements results in low Reynolds number flows for which molecular diffusion can be important. In general, the effects of molecular diffusion, combined with the rapid adjustment forced on the internal flows by the stretching, precludes the simple similarity assumption by Maxworthy (1972) and later by Chomiak (1979) to understand the hydrodynamics of spark kernels.
5.4 Toroidal Mixing Model: Non-combusting Solutions

We now develop a model to examine the behaviour of toroidal spark kernels. The first step is to model the non-combusting case. This would be applicable to experiments involving spark kernels in air. In the next chapter, we extend the model to combustible mixtures.

Our model is an extension of the spherical mixing model applied to plasma jets by Topham et al. (1984), which itself was an adaptation of an atmospheric thermal model. The foundation of the model is that the entrainment assumption of Morton et al. (1956) is applicable. This requires the following assumptions:

- there is no loss of momentum or heat from the element to a wake;
- the mixing element has a toroidal shape with uniformly distributed properties;
- the mixing takes place at constant pressure;
- the mixing element and ambient environment are perfect gases;
- ambient fluid is entrained into the element at a rate proportional to its velocity and surface area. The constant of proportionality, $\alpha$, is fixed for a given mixing element, but can vary from one element to another.
Figure 5.3 shows the toroidal coordinates, \( r \) and \( b \), respectively the major and minor axes of the mixing element. In this orientation, the spark electrodes lie along the \( b \)-axis.

![Diagram of toroidal coordinates](image)

**Figure 5.3 Toroidal coordinate system**

The basic conservation equations are then:

**Mass:**

\[
\frac{d}{dt}(2\pi^2r^2b^3\rho) = \alpha 4\pi^2\rho_0rb \frac{dr}{dt}
\]  

\( (5.1) \)

**Momentum:**

\[
2\pi^2rb^3(\rho + k\rho_b)v = I_0
\]  

\( (5.2) \)

**Energy:**

\[
2\pi^2rb^3\rho(h - h_0) = H_0
\]  

\( (5.3) \)

**Radial velocity:**
where the initial conditions are:

\[ \beta = \frac{\rho}{\rho_0} = 0 \]
\[ b = b_0 \]
\[ r = r_0 \]  

It is convenient to find scaling quantities that will put the conservation equations into non-dimensional form. The initial conditions (5.5) provide such quantities. From the energy equation, (5.3), we get, at \( t=0 \):

\[ r_0 b_0^2 = \frac{H_0}{2\pi^2 \rho_0 b_0} \]  

We now choose the following scale lengths for the coordinates:

\[ r^* = \frac{r}{r_0} \]
\[ b^* = \frac{b}{b_0} \]  

where \( r_0 \) and \( b_0 \) must satisfy (5.6).

The momentum equation, (5.2), gives, at \( t=0 \):

\[ r_0 b_0^2 \left( \frac{dr}{dt} \right)_{t=0} = \frac{l_0}{2\pi^2 \rho_0} \]
which leads us to choose as the velocity scale:

\[ v_0 = \frac{I_0}{r_0^2 h_0^2 2\pi \rho_0} = \left(\frac{h_0}{H_0}\right) I_0 \]  

(5.9)

For scaling the time, we choose:

\[ t_m = \frac{r_0 H_0}{h_0 I_0} \]  

(5.10)

<table>
<thead>
<tr>
<th>Table 5.3 Scaling Quantities</th>
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<tbody>
<tr>
<td></td>
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<tr>
<td>Scale length</td>
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<tr>
<td></td>
</tr>
<tr>
<td>Scaled coordinate</td>
</tr>
<tr>
<td>Initial conditions</td>
</tr>
</tbody>
</table>

With the scaling quantities from Table 5.3, the normalized form of the conservation equations are:

\[ \frac{d}{dt} (b^* b^*^{2} \beta) = 2\alpha \frac{r_0}{b_0} r b^* \frac{dr^*}{dt} \]  

(5.11)

\[ r^* b^*^{2} (\beta + k) \frac{dr^*}{dt} = 1 \]  

(5.12)

\[ r^* b^*^{2} (1 - \beta) = 1 \]  

(5.13)
The solution of these scaled equations is straightforward and the details can be found in Appendix II. Solutions of equations (5.11) to (5.13), which also satisfy the initial conditions (5.5), are:

\[
\begin{align*}
    b^* &= \Psi r^* + (1 - \Psi) r^* \frac{1}{r^*} \\
    \chi &= \Psi^2 (r^{*3} - 1) + 2\Psi (1 - \Psi) \left( r^* \frac{2}{3} - 1 \right) \\
    t^* &= \frac{1}{4} (1 + k) \Psi^2 (r^{*4} - 1) + \frac{4}{5} \Psi (1 + k) (1 - \Psi) \left( r^* \frac{2}{3} - 1 \right) + \\
    &+ \left( k - (1 + k) \Psi (2 - \Psi) \right) (r^* - 1)
\end{align*}
\]

where

\[
\begin{align*}
    \Psi &= \frac{2}{3} \alpha \frac{r_0}{b_0} \\
    \chi &= \frac{\beta}{1 - \beta}
\end{align*}
\]

Figures 5.4a - 5.4c show the behaviour of the non-combusting toroidal element solutions, where the numbers shown adjacent to each plot denote a different value of the parameter \( \Psi \). The quartic asymptotic behaviour of the \( r^* \) vs \( t^* \) solution is very similar to that in the spherical element case. This is not surprising, as the basic conservation equations are functionally very similar. In the next section, we compare these solutions to observations of toroidal spark kernels.
Figure 5.4a. Non-combusting model; equation 5.14. Numbers adjacent each plot denote the value of $\Psi$.

Figure 5.4b. Non-combusting model; equation 5.15. Numbers adjacent each plot denote the value of $\Psi$. 
Figure 5.4c. Non-combusting model; equation 5.16. Numbers adjacent each plot denote the value of $\Psi$.

5.5 Comparison of the Toroidal Mixing Model with Observed Kernel Behaviour

The behaviour of spherical spark kernels produced by plasma jets has been interpreted by a turbulent mixing model (Topham et al. 1984). The fit to the model solutions was good. The mixing model was further confirmed by Soloman (1988).

In contrast, fitting data to the toroidal element model solutions is made more difficult because of our lack of knowledge of both the entrainment rate constant, $\alpha$, and the ratio of the initial major and minor radii, $r_0/b_0$. In the spherical case, measurement of the element radius as a function of propagation distance yields a value for the entrainment constant. For the
toroidal case, simultaneously fitting data to equations (5.14) and (5.16) yields suitable values for $a$ and $r_0/b_0$.

We now look at some examples of kernel behaviour and compare these with the model just described. In general, published data on kernel morphology lack accurate measurement of the energy deposited into the electrode gap. Usually recorded is the stored energy available to the whole discharge circuit. Portions of the energy dissipated in other circuit elements must be accounted for. Furthermore, the momentum imparted to the gas is not normally measured, both because the measurement is difficult and the diffusion-based theories have no need for it. However, Haley and Smy (1989) provide estimates of the momentum of radially expanding kernels as a function of energy, discharge time, and gap length.

The first candidate for comparison, which we will call Example 1, is the spark kernel data from Figure 4.4 in Chapter 4. The second candidate, to be called Example 2, is from data published by Maly and Vogel (1978). The parameters that were used to scale the data are shown in Table 5.4. The momentum values are estimates from Haley and Smy (1989).

Examples 1 and 2 are compared to the non-combusting toroidal mixing theory in Figures 5.5a, 5.5b, and 5.6, respectively, where again the numbers adjacent to each plot denote values of the parameter $\Psi$. For Example 2, data for $b^*$ vs $r^*$ were not available.
Figure 5.5a. Example 1 data (*) and the non-combusting model; equation 5.14. Numbers adjacent each plot denote the value of $\Psi$.

Figure 5.5b. Example 1 data (*) and the non-combusting model; equation 5.16. Numbers adjacent each plot denote the value of $\Psi$. 
Figure 6.6. Example 2 data (*) and the non-combusting model; equation 5.14. Numbers adjacent each plot denote the value of $\Psi$.

<table>
<thead>
<tr>
<th>Example</th>
<th>$H_0$ (mJ)</th>
<th>$I_0$ (kgm/sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5</td>
<td>$2.4 \times 10^{-7}$</td>
</tr>
<tr>
<td>2</td>
<td>30</td>
<td>$6 \times 10^{-7}$</td>
</tr>
</tbody>
</table>

The results are both encouraging and puzzling. First, the coherent toroidal structures of Figure 4.4 resemble laminar vortex rings, at least before the onset of the observed instability. This suggests that a modification of the laminar vortex ring theory of Maxworthy (1972) would be more appropriate rather than the turbulent mixing model just presented. However, the simple entrainment assumption appears to be applicable. A possible explanation is that for these coherent structures, the entrainment is dominated by a single, large-scale circulation pattern.
of the type demonstrated by the flow visualizations of Haley and Smy (1989). If the kinetic energy remains in this scale, its distribution being unchanged either by molecular diffusion or by transport to smaller scales over the time of interest, the self-similar scaling implied by the entrainment assumption may still hold.

Second, the data of Maley and Vogel (1978), used as Example 2, are compared to the non-combusting mixing model even though the data were obtained in a combustible mixture. Reasons for using the non-combusting model in this case are discussed in the next chapter. Even though the data for the non-combusting case are sparse and far from conclusive, the results are encouraging, given the uncertainties involved in determining the values of the energy and especially the momentum imparted to the elements.

5.6 Mechanism for Generating the Toroidal Flow Field

It is interesting to consider a mechanism that could result in the types of circulation patterns suggested by the toroidal structure of spark kernels. Chomiak (1979) considered magneto-hydrodynamic forces to be important in establishing counter flowing axial jets that collide in the mid-gap region, resulting in a net radial flow with counter rotating vortices. However, estimates of the mhd forces indicates that they are dominated by temperature and pressure gradients.
Kono et al. (1988) and Haley and Smy (1989) suggest that the transition of the shock wave from an initial cylindrical configuration to its final spherical form imparts the observed momentum to the kernel fluid. Vorticity is generated by the cross products of density and pressure gradients. Perhaps a spherically symmetric pressure field in the wake of the shock, together with the predominantly cylindrically symmetric temperature field of the initially hot gas kernel, is enough to generate the observed structures. Borghese et al. (1988) use similar hydrodynamic arguments, but suggest it is the high degree of curvature of the temperature and velocity fields near the electrode tips that is responsible for generating the required vorticity.

In the next chapter, we extend the toroidal model to combustible mixtures. We explore some of the model behaviour, discuss ignition criteria in light of the mixing element point of view, and examine mixing elements in turbulent ambient conditions.
6 A Unified Approach to Spark Kernel Development: Combustible Mixtures

In the previous we chapter presented a mixing model describing the evolution of a toroidal spark kernel in a non-combusting environment. We now extend the model to a combustible mixture by following the example of the spherical mixing element solutions in combustible mixtures provided by Topham et al. (1986). This model predicted kernel behaviour remarkably well, the most striking result being the existence of two distinct classes of model solutions. In one class, the "extinction branch", are solutions that resemble the non-combusting solutions, where the density approaches the ambient level. In the second class, the solutions attain a low density, corresponding to a high bulk temperature which is maintained at a constant level. This self-sustaining, high-temperature state is referred to as the "combusting branch."

Section 6.1 below, we extend the toroidal mixing model to combustible mixtures by including a simple Arrhenius type of heat addition term to simulate combustion in an element propagating into a combustible mixture.

6.1 Toroidal Mixing Model with Heat Addition

In the non-combusting case, mixing caused the element's density to rise to the ambient level. In the combusting case, the cold, ambient mixture is introduced into the interior of the element by entrainment, and this
acts to control the rate of heat release. To examine the behaviour of the combusting case, we modify the conservation equations of Section 5.4 to include heat input in the energy equation, and add a reactant concentration rate equation. The equations are:

Mass:
\[
\frac{d}{dt}(2\pi^2 r^2 b^2 \rho) = \alpha 4\pi^2 \rho_o r b \frac{dr}{dt} \quad (6.1)
\]

Momentum:
\[
2\pi^2 r b^2 (\rho + k \rho_0) \frac{dr}{dt} = I_0 \quad (6.2)
\]

Energy:
\[
\frac{d}{dt}(rb^2(\rho(\rho - h_0))) = rb^2 qk c^2 \quad (6.3)
\]

Concentration:
\[
\frac{d}{dt}(rb^2 c) = -rb^2 k c^m + 2\alpha r b^2 c_0 \frac{dr}{dt} \quad (6.4)
\]

where the quantities are as defined in Section 5.4, with the addition that \( c_0 \) is the ambient reactant concentration, and \( m \) is the order of the reaction, \( m=1,2,\ldots \).

Taking a simple Arrhenius reaction rate of the form:
\[
k = A\exp\left(-\frac{E}{RT}\right) \quad (6.5)
\]

and defining some new parameters as:
normalized specific exothermicity: 
\[ q^* = \frac{q c_0}{\rho_a h_0} \]

characteristic chemical reaction time: 
\[ t_{ch} = \frac{1}{A c_0^{n-1}} \exp\left( \frac{E}{RT_0} \right) \]

reduced ambient temperature: 
\[ \epsilon = \frac{RT_0}{k} \]

the conservation equations (6.1) to (6.4) can be put into normalized form, with \( c^* = c/c_0 \), as:

\[
\frac{d}{dt} \left( r^* b^* \beta^* \right) = 2a r^* b^* \left( \frac{dr^*}{dt} \right) \quad (6.6)
\]

\[
r^* b^* c^* \frac{dr^*}{dt} (\beta + k) = 1 \quad (6.7)
\]

\[
\frac{d}{dt} (r^* b^* (1 - \beta)) = q^* \left( \frac{l_m}{t_{ch}} \right) r^* b^* c^* m \exp\left( \frac{1 - \Lambda \beta}{\epsilon} \right) \quad (6.8)
\]

\[
\frac{d}{dt} (r^* b^* c^*) = -\left( \frac{l_m}{l_{ch}} \right) r^* b^* c^* m \exp\left( \frac{1 - \Lambda \beta}{\epsilon} \right) + 2a r^* b^* \left( \frac{dr^*}{dt} \right) \quad (6.9)
\]

The solution of (6.6) to (6.9) requires careful attention to initial conditions as well as to the real temperature profile of the element, assumed uniform in the model, due to the exponential function in the Arrhenius term. These details are treated in Topham et al. (1986).
Instead of a numerical exploration of the solutions to (6.6) to (6.9), we anticipate, in analogy with the spherical element case, that the combustion solution will produce a self-sustaining element which maintains a near constant density, corresponding to a high-temperature state. Setting \( \beta \) to \( \beta_c \) in (6.1) and (6.2), we obtain the constant density solutions (see Appendix III for details):

\[
b'^* = \Psi p'^* + (1 - \Psi_c)r'^* \frac{1}{2} \tag{6.10}
\]

\[
\frac{1}{4} \Psi_c^2 (r'^* - 1) + \frac{4}{5} \Psi_c (1 - \Psi_c) \left( \frac{1}{r'^*} - 1 \right) + (1 - \Psi_c^2) (r'^* - 1) = \frac{1}{\beta_c + k'} \tag{6.11}
\]

where:

\[
\Psi_c = \frac{1}{\beta_c} \Psi = \frac{1}{\beta_c} \left( \frac{2}{3} \alpha b_0 \right)
\]

As anticipated, the fully combusting solutions mirror the spherical element case because the effect of heat addition leads to an increase in the entrainment rate by the factor \( 1/\beta_c \).

### 6.2 Ignition Criteria

The appearance of two distinct classes of solutions in the spherical element combustion model of Topham et al. (1986) is an important result. Numerical exploration of the equations indicated that the parameter:
was critical in determining the kernel behaviour. High values lead to combustion, low values lead to extinction.

For the toroidal element, the same sensitivity to the above parameter can be seen in equation (6.9). If the fully combusting state is to be achieved, then the density ratio attains a constant level, \( \beta_c \), and the concentration, \( c^* \), must go from an initial value of zero to some maximum, \( c_{\text{max}}^* \), and then quickly to zero as reactant is consumed within the element. Thus, the time derivative of the concentration must go through a zero, and using this we can obtain from equation (6.9) the condition:

\[
\frac{1}{\alpha} \frac{t_e}{t_s} \left[ c_{\text{max}}^* \exp \left( \frac{1 - \Delta \beta}{\varepsilon} \right) \right] \simeq O(1)
\]

Using values of \( c_{\text{max}}^* \) in the range of 0.05 - 0.10 and \( \beta_c \) in the range of 0.2 - 0.4, which are taken from the numerical solutions of Topham et al. (1986), estimates for the combustion parameter (6.12) can be obtained from (6.13) to determine whether a self-sustaining or extinction solution will occur. Thus, (6.13) can be viewed as an ignition criterion for the very early phase of spark ignition. The further development of the kernel from the early phase (where mixing controls the dynamics) to a fully
developed flame front propagating into the remaining mixture (where thermal diffusion controls the dynamics) has not been addressed here. This would require perhaps using the mixing model developed here to provide initial conditions for the more recent thermal models of Ko et al. (1991). This is left as future work.

Measurements of real ignition kernels are lacking, so comparison of the toroidal combustion model is not possible yet. However, the comparison of the spherical element model with plasma jet data in combustible mixtures is sufficiently encouraging that we can draw several conclusions from the toroidal model with some degree of confidence.

First, the value of the combustion parameter (6.12) is critical in determining the evolution of the kernel. Unlike strictly thermal models of ignition, this parameter combines the fluid mechanical properties of the kernel, with the properties of the chemical reaction.

Second, for a fixed energy input, a device which imparts high momentum has a short fluid time scale, $t_m$, which inclines the element towards extinction. On the other hand, low momentum is favourable for combustion. Plasma jet devices, for example, can vary the impulse delivered by changing the dimensions of the cavity. Axial discharges can reduce the delivered impulse by increasing the discharge time. In the case of the fast discharge kernels measured by Maly and Vogel (1978) and used in Section 5.5, we speculate that, as the central temperature rapidly
felled, the kernels had too high an impulse to achieve self-sustaining conditions. Hence, the non-combusting model was more appropriate.

Third, given that high momentum devices are not apparently favourable for ignition, why do plasma jets and fast discharge igniters demonstrate a clear ability to ignite lean mixtures? For a given mixture ratio, say a lean mixture, the critical combustion parameter indicates that the mixing time $t_m$ would have to be sufficiently long to achieve the self-sustaining state. Apart from numerical constants, the mixing time $t_m$ is essentially the same for both the spherical and toroidal cases and, more importantly, we can write:

$$t_m \propto \frac{H^3}{I}$$

Thus, the energy imparted to the kernel is still very important to the establishment of ignition. Even though plasma jets and fast discharge devices impart a relatively high momentum to the kernel, they also deliver high energy, compared to a conventional axial discharge or common spark plug. In the case of the plasma jet, the discharge is bounded, except at the orifice, and measurements (Smy et al. 1983) indicate that about 10% of the 1-2 J of the total discharge circuit energy is imparted to the kernel. This is true when the discharge time is on the order of twice the time it takes an acoustic wave to travel the length of the cavity and the radial expansion of the arc has just contacted the cavity walls.
This latter condition arises from the bounded nature of the discharge. The hot gases are expelled from the cavity on the order of the acoustic travel time and, if the arc contacts the cavity walls, high energy losses occur. Thus, discharges lasting longer than the time it takes to expel the cavity, or long enough to contact the cavity walls, waste the discharge energy.

In the case of fast discharge devices, where the discharge is open or unbounded, it is the low inductance of the discharge circuit that promotes the coupling of the available energy, typically 10 - 30 mJ, to the kernel. Circuits that discharge their energy in 100 - 500 ns are delivering it to a circuit element, the spark gap, whose impedance more nearly matches the rest of the circuit. This impedance match only lasts a short time, as discussed in Chapter 2. Thus, plasma jet and fast discharge devices couple much higher energies to their spark kernels than do conventional igniters which typically deliver 1 - 2 mJ to their ignition kernels. This makes the former more effective igniters in lean mixtures.

6.3 Ignition in Turbulent Mixtures

Ignition in most practical systems takes place against a background of ambient turbulence. Given that ignition kernels are now to be viewed as regions of concentrated vorticity, their interaction with a general background of turbulence would be valuable knowledge. The following
discussion is treated in more detail in the paper by Pitt et al. 1991, which appears in Appendix I.

Our observations of spark kernels from axial discharges indicate that the kernels start out as coherent toroidal structures and then an instability occurs which causes a transition to a more rapid, turbulent mixing state. This can be seen in the Schlieren photographs of Section 4.4. We propose that the intrinsic mixing properties of the element itself dominate its evolution for a significant period and that the role of ambient turbulence is to control the transition of the element to the more turbulent state. For given discharge conditions, the more intense the ambient turbulence, the earlier transition occurs and the greater the proportion of the total momentum incorporated directly into the now turbulent kernel. As the mixing rate of the kernel is proportional to the inverse of the momentum, higher ambient turbulence leads to more rapid evolution of the kernel. A comparison between the mixing model and published results on the spreading kernels in turbulent mixtures lends some support to this view (Pitt et al. 1991).

The spark kernel model presented in the previous chapter, and further developed in this chapter to include the effects of combustion, represents a primitive state of knowledge. The reason is the lack of suitable data which accurately measure the energy and momentum of kernels or the temporal progress of the major and minor radii. Such measurements are experimentally challenging. The model more closely resembles the observed kernel structures and, for the first time, links the types of
structures produced by common electrical discharge igniters to the same underlying physical process. The traditional view of an ignition kernel as a point heat source whose evolution is governed by diffusive processes overlooks the essential mechanism of fluid dynamics which dominate immediately following the initial shock wave and persist until a freely propagating flame is established. Consequently, the traditional view fails to predict the observed kernel structures and the critical processes involved.
7 Summary

This thesis presented a new view of spark ignition that links all electrical discharge ignition devices through a common physical mechanism—hydrodynamic mixing. Our viewpoint manifested itself in the following way.

We began by establishing the motivation for lean-burn or alternative fueled IC engines. Reduced emissions and fuel efficiency are the dual goals behind this move. Ignition difficulties arise, however, when the operating point of an engine is moved to leaner conditions or an alternative fuel such as natural gas is substituted for gasoline. We demonstrated, in Chapter 2, some practical means by which the ignition delay, an annoying problem of lean mixtures, can be reduced. Results from quiescent combustion bomb experiments and IC engines led us to conclude that, aside from demonstrating the reduction of ignition delay, ignition devices affect only the early phase of ignition in IC engines. The bulk turbulence in an engine combustion chamber dominates the overall combustion event. Effects seen in quiescent combustion vessels disappear in real engines, except for the 1-2 ms beginning phase of the event.

With the early phase of spark ignition established as the critical period at which ignition devices can influence IC engines, the focus of the thesis shifted to a discussion of the conventional thermal theory of ignition—a theoretical framework that has shaped most of the
theoretical, experimental, and practical thinking on ignition for nearly 50 years. The thermal approach to spark ignition views the spark discharge as a point or line heat source that raises a critical volume of mixture to the flame temperature. Heat loss, by conduction, to the surrounding mixture is the critical mechanism that determines the minimum spark energy for ignition. Furthermore, the thermal approach predicts that the ignition kernel would develop as a spherically symmetric region possessing an exponentially decaying radial temperature profile. This prediction was challenged by experimental observations, some new and others from existing literature, which clearly showed that spark kernels from axial discharges initially develop as sharply bounded, toroidal regions of hot gas.

We discussed the importance of the temporal characteristics of the electrical discharge. Energy delivered to the spark gap in the first $10^{-6}$ s was seen to drive the spark kernel's behaviour. The energy delivered at later times is less effective in feeding the spark kernel, as more of the discharge energy is going into supporting the arc phase of the discharge, which suffers heavy energy losses, mostly to the electrodes. Thus, conventional automotive engineering wisdom of long, typically 1-10 ms discharges is not well founded.

In Chapter 5, we stood back and looked at the evidence of Chapter 4 and argued that the toroidal structures produced by axial discharges are a result of underlying hydrodynamic circulation patterns which
are strikingly similar to flows in plasma jet spark kernels. Armed with this insight, we proposed a classification scheme that links the practical ignition sources of Chapter 2 to a fundamental process: hydrodynamic mixing. This classification scheme sees plasma jet, puff jet, and surface discharge ignition devices producing spark kernels with spherically symmetric circulation patterns and structures. Axial discharges produce toroidally symmetric circulation patterns and structures.

A mixing model, developed by Topham et al. (1984) for interpreting plasma jet spark kernel behaviour, was modified to the toroidal geometry of axial discharge spark kernels. The mixing model required knowledge of the momentum imparted to the developing spark kernel. Such measurements are rare since the dominant theoretical framework ignored the kernel's impulse and subsequent flows. However, we were able to make an initial comparison of some limited data with the mixing model. The results are encouraging, but must be interpreted cautiously because of the limited data set available. Obviously, further measurements are required before more definite claims can be made.

In Chapter 6, our mixing model was extended to the realm of combustible mixtures following the experience of the spherical element solutions developed for plasma jet kernels (Topham et al. 1986) A distinct combusting solution was sought which featured a constant, below-ambient level element density, reflecting a high-
temperature, self-sustaining state. An ignition criterion was discussed which identified a common combustion parameter:

\[ \frac{1}{\alpha} \frac{t_m}{t_{ch}} \]

This parameter combines the energy, momentum, and resulting vorticity distribution of the kernel, together with the exothermic properties of the mixture. The combustion parameter is the essence of the mixing models discussed, because it identifies the balance between heat release and hydrodynamic mixing as the controlling mechanism in the early phase of ignition by electrical discharges. The ignition system designer is then encouraged to take into account the role of the momentum imparted to the spark kernel, which manifests itself through the mixing time \( t_m \), since:

\[ t_m \propto H^\frac{4}{3} \]

For a given energy, low momentum devices produce kernels that are more effective for ignition in very lean mixtures.

Chapter 6 concludes with a reinterpretation of measurements of spark kernels in turbulent combustible mixtures, using the mixing model as a guide. It is proposed that the observed transition of toroidal elements from coherent to fully turbulent structures is affected by ambient turbulence. The greater the turbulent intensity, the earlier
the transition occurs, the greater the proportion of total momentum incorporated directly into the now more turbulent element, and the faster the mixing time. A comparison between the model and published data on the spreading rate of kernels in turbulent mixtures lends some support to this view.

**Serendipity**

We cannot end without mentioning the striking similarity between the photographs of toroidal spark kernels and the photographs of several planetary nebulae. For example, photographs of the Ring Nebulae, M 57, the Helix Nebulae, NGC 2244, and many other astronomical objects show similar, well-defined toroidal structure, but on vastly larger scale than the spark kernels discussed in Chapter 4. We wonder, if we modify the mixing model...

_Everything of importance has been said before by somebody who did not discover it._—A. N. Whitehead
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Appendix I Papers published by the author

The University Librarian has recommended that published papers which I have co-authored not be reproduced in this appendix out of respect for copyright protection laws. I therefore present references to specific papers so that an interested reader can pursue them.

Work on plasma jet ignition in natural gas fueled engines can be found in:


Work on the development of the puff-jet ignition concept can be found in:


Work on the toroidal mixing model of spark ignition can be found in:

Appendix II  Non-combusting model solutions

The scaled conservation equations developed in Chapter 5 are repeated here:

\[ \frac{d}{dt} (\beta r^* b^{**}) = 2\alpha \frac{r_a}{b_0} r^* b^* \frac{dr^*}{dt} \]  \hspace{1cm} (A1)

\[ r^* b^{**}(\beta + k) \frac{dr^*}{dt} = 1 \]  \hspace{1cm} (A2)

\[ r^* b^*(1 - \beta) = 1 \]  \hspace{1cm} (A3)

To obtain an expression for \( b^* \) in terms of \( r^* \), the calculation proceeds as follows. Equating (A2) and (A3), we obtain

\[ \frac{dr^*}{dt} = \frac{1 - \beta}{\beta + k} \]  \hspace{1cm} (A4)

Rewriting (A3) as

\[ r^* b^{**} = \frac{1}{1 - \beta} \]  \hspace{1cm} (A5)

and using this, the LHS of A1 becomes

\[ \frac{d}{dt} (\beta r^* b^{**}) = \frac{d}{dt} (\frac{\beta}{1 - \beta}) = \frac{d}{dt} \chi \]

where
\[ x = \frac{\beta}{1 - \beta} \]

The RHS of (A1) is, using (A4) and (A5),

\[ 2\alpha \frac{r_0}{b_0} b^* \frac{dr^*}{dt} = 2\alpha \frac{r_0}{b_0} \frac{1}{b^* \beta + k} \]

Thus (A1) can be put into the following form:

\[ \frac{dx}{dt} = 2\alpha \frac{r_0}{b_0} \frac{1}{b^* \beta + k} \quad (A6) \]

Again, using (A3), we can express \( b^* \) as

\[ b^* = \left( \frac{1}{r^*} \frac{1}{1 - \beta} \right)^{\frac{1}{2}} \]

Using this result in (A6), we obtain

\[ \frac{dx}{dt} = 2\alpha \frac{r_0}{b_0} r^* \frac{1}{2} \left( \frac{1 - \beta}{\beta + k} \right)^{\frac{1}{2}} = 2\alpha \frac{r_0}{b_0} \frac{1}{2} \left( \frac{1 + \chi}{1 + k} \right)^{\frac{1}{2}} \quad (A7) \]

Equation (A2) can be re-written as

\[ \frac{dr^*}{dt} = \frac{1}{(1 + k)^{\chi + k}} \quad (A8) \]
Using (A6) and (A8), an expression for \( \frac{dr^*}{d\chi} \) can be calculated:

\[
\frac{dr^*}{d\chi} = \frac{dr^*}{dt^*} \cdot \frac{1}{2\alpha \frac{r_0}{b_0} r^{*\frac{1}{2}} (1 + \chi)^{\frac{1}{2}}} 
\]

(A9)

Equation (A9) can be integrated, using the initial conditions listed in Table 5.3:

\[
2\alpha \frac{r_0}{b_0} \int r^{\frac{1}{2}} dr = \int_{\chi}^{\chi'} \frac{dx'}{(1 + x')}^{\frac{1}{2}}
\]

Using \( 1 + \chi = \frac{1}{1 - \beta} \), the results of the integration can be rearranged to yield

\[
b^* = \frac{2}{3} \alpha \frac{r_0}{b_0} r^* + \left( 1 - \frac{2}{3} \alpha \frac{r_0}{b_0} \right) r^{*\frac{1}{2}}
\]

(A10)

Setting \( \Psi = \frac{2}{3} \alpha \frac{r_0}{b_0} \), then (A10) can be put into the form used in Chapter 5 as equation (5.14):

\[
b^* = \Psi r^* + (1 - \Psi) r^{*\frac{1}{2}}
\]

(5.14)

To calculate \( r^*(t^*) \), we integrate (A8) after computing \( \chi(r^*) \). Using (A4) and (A5), we obtain
\[
\frac{dr^*}{dt} = \frac{1}{r^*} \frac{1}{b^*} \frac{1}{\beta + k} 
\]
(A11)

From (A6) we have

\[
\frac{1}{\beta + k} = \frac{b^*}{2a} \frac{d\chi}{dt} 
\]

(A12)

Substituting (A12) into (A11) yields

\[
\frac{d\chi}{dr^*} = 2a \frac{r_0}{b_0} b^* 
\]

which can be integrated, using (5.3), resulting in

\[
\chi = \Psi^2(r^* - 1) + 2\Psi(1 - \Psi)(r^* - 1) \quad (5.15)
\]

Substituting (5.15) into (A8) produces an equation which can be integrated directly to yield equation (5.16) of Chapter 5

\[
t^* = \left( \frac{1+k}{4} \right) \Psi^2(r^{*4} - 1) + \frac{4}{5}(1+k)\Psi(1 - \Psi)(r^{*2} - 1) + \{k - (1 + k)[2\Psi - \Psi^2]\}(r^* - 1)
\]

which completes the solution of the non-combusting model.
Appendix III  Constant Density Solutions

For these solutions, we set \( \beta \rightarrow \beta_c \), where

\[ \beta_c = \text{constant density ratio}. \]

In (A1) and (A2) of Appendix II. The constant density model is thus described by

\[
\beta_c \frac{d}{dt} (r^* b^*) = 2 \alpha \frac{r_0}{b_0} r^* b^* \frac{dr^*}{dt} \tag{B1}
\]

\[
r^* b^* (\beta_c + k) \frac{dr^*}{dt} = 1 \tag{B2}
\]

Expanding the LHS of (B1) and collecting terms yields

\[
\frac{db^*}{dr^*} = \frac{\alpha}{\beta_c b_0} \frac{r_0}{2r} \tag{B3}
\]

If we assume that a solution for (B3) has a functional form similar to the non-combusting case, then

\[
b^* = a_r r^* + a_r r^{*\frac{1}{2}} \tag{B4}
\]

where \( a_1 \) and \( a_2 \) are undetermined coefficients. Substituting (B4) into (B1) and comparing terms yields
From the initial conditions \( b^* = 1 \) when \( r^* = 1 \), equation (B4) gives

\[
a_1 = \frac{2}{3} \frac{\alpha}{\beta_c} \frac{r_0}{b_0}
\]

Therefore

\[
l = a_1 + a_2
\]

and (B4) becomes

\[
\begin{align*}
b^* &= \frac{2}{3} \frac{\alpha}{\beta_c} \frac{r_0}{b_0} r^* + \left(1 - \frac{2}{3} \frac{\alpha}{\beta_c} \frac{r_0}{b_0} \right) r^{\frac{1}{2}} \\
&= \psi_c r^* + (1 - \psi_c) r^{\frac{1}{2}}
\end{align*}
\]

Equation (B5) resembles the non-combusting solution with \( \alpha \) replaced by \( \alpha/\beta_c \). To complete the solution, we solve (B2), using (B5). Thus (B2) becomes
\[
\frac{dr^*}{dt^*} = \frac{1}{\beta_c + k} \frac{1}{\psi_c^2 r'^{2} + 2 \psi_c (1 - \psi_c) r'^{2} + (1 - \psi_c)^2}
\]

which can be integrated directly to give

\[
\frac{1}{4} \psi_c^2 (r'^2 - 1) + \frac{4}{5} \psi_c (1 - \psi_c) \left( r'^2 - 1 \right) + (1 - \psi_c^2) (r'^2 - 1) = \frac{1}{\beta_c + k} t^* \quad (B6)
\]