Development of a Gas-Fed Plasma Source for Pulsed High-Density Plasma/Material Interaction Studies

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Abstract—A gas-fed capillary plasma source has been developed to study plasma–surface interactions under pulsed high pressure arc conditions, without the use of an exploding fuse wire or ablative liner. A nonintrusive preionization source has been developed to break down relatively large interelectrode gaps at low charge voltages of 2–6 kV. The preionization source comprises a nonequilibrium surface streamer discharge that forms a conducting channel through which the main thermal arc discharge is initiated. The arc electron temperature and number density are estimated to be $T_e \sim 1–2$ eV and $n_e \sim 10^{23}$ m$^{-3}$. Silicon and sapphire samples were exposed to the arc plasma and revealed deposition of electrode and wall materials. Substitution of Elkonite 50W3 for brass electrodes reduced plasma contamination to acceptable levels. The plasma–material interactions were examined and quantified using scanning electron microscopy and energy dispersive X-ray spectroscopy.

Index Terms—Arc discharge, atmospheric discharge, capillary discharge, plasma–material interaction, pulsed thermal plasma.

I. BACKGROUND

HIGH-POWER, high-pressure, capillary-plasma discharges have a variety of applications in pulsed space propulsion devices and as igniters for chemical propellant guns [1]–[5]. The plasma formed in these devices is sustained through the ablation of a sacrificial liner material (typically a polymer). Capillary plasmas are thermal arcs characterized by very high in-bore pressures ($\sim$1–10 MPa) with very high input power densities delivered to the plasma during a short pulse transient.

The high pressures, and correspondingly high collisionality, result in local thermodynamic equilibrium (LTE) conditions within the core of the arc. Typical LTE arc core temperatures are about 1–2 eV and charge particle densities of $\sim$10$^{23}$ – $\sim$10$^{28}$ m$^{-3}$ [6]. The arc column is typically spatially nonuniform and strong gradients in the arc properties exist. For example, the periphery of the arc column in the vicinity of the ablating surface has a much lower temperature of $\sim$0.1 eV [7].

Material surfaces exposed to thermal arc plasmas encounter extreme conditions. The charged particle flux to a surface under these conditions is estimated at $\sim$10$^{27}$ m$^{-2}$s$^{-1}$ within the core of the arc [6]. Surface heat flux for the same conditions is $\sim$1 GW/m$^2$ and can, therefore, cause substantial damage to plasma facing material components with repeated use [8].

An understanding of how thermal plasmas interact with material surfaces is key to the development of reliable, long-lifetime capillary arc based devices. However, plasma–material interaction studies in these discharges are extremely difficult due to the small capillary bore size (approximately few millimeters in diameter). Such small diameters preclude direct placement of a material sample within the chamber bore.

The easiest approach to studying how these plasmas interact with a material surface is by expanding the plasma from the bore into an external plasma jet, and allowing it to interact with a sample surface placed in its path. The expansion process, however, results in a drop in pressure and temperature, and the plasma composition can be vastly different from the conditions within the capillary bore. For example, expanding the plasma from a bore at a pressure of $\sim$10 MPa to ambient atmospheric conditions can result in a significant temperature drop from 1 to $\sim$0.1 eV [9], [10]. At this temperature, several chemical components of the plasma (originating from the ablating liner) condense out and are deposited on the target material surface. Furthermore, practical arc breakdown (ignition) schemes in capillary discharges typically involve a metal fuse wire connecting the electrodes that is exploded to produce an initial gaseous plasma conducting channel. The metal vapor contamination from the fuse wire is also a major problem for plasma–material interaction studies.


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The plasma chemical composition is controlled through use of high purity gas cylinders. The feed gases are regulated, metered, and mixed before flowing into the reaction chamber. Background feed gases, such as argon, are metered through a 0–50 SLPM digital flow meter (Omega FMA-1609A). Small amounts of additive gases, such as hydrogen, oxygen, or methane, are metered through a 0–1 SLPM meter (Omega FMA-1620A). Premixing of the background and additive gases occurs in a two meter long gas feed tube upstream of the chamber inlet. To ensure a uniform environment, the reaction chamber is purged for 5 s or more before initiating a discharge. Single component gases or gas mixtures are used to systematically investigate various plasma–surface reaction mechanisms. For example, a single species plasma composed of an inert noble gas is used to determine the importance of sputter mechanisms. Small amounts of oxidizing and reducing gases may be added to the inert background to investigate a chemically reactive plasma interacting with various materials. Materials exposed to the plasma include single crystal silicon, sapphire, and tungsten. Each of these materials is selected based upon their bond nature: covalent, ionic, or metallic, respectively.

The test samples are fixed to a substrate holder and inserted through the cathode into the discharge chamber. Samples are exposed to several pulsed, thermal plasma discharges, before surface analysis. The effects of various plasma environments on each benchmark material are experimentally investigated with various surface analysis tools, such as scanning electron microscopy (SEM) and energy dispersive X-ray spectroscopy (EDXS). The work presented in this paper focuses on the development and characterization of the plasma source. Details of the various experimental subsystems and diagnostics are presented in the remainder of this section.

B. Plasma Discharge Chamber and Feed Gas System

Fig. 2 shows a schematic of the plasma discharge chamber that consists of two concentric tubes, two annular end electrodes, and a helical copper wire used to initiate breakdown between the two electrodes.
substrate holder inserted through the cathode to which samples are affixed.

C. Power Supply

1) Pulsed Thermal Arc Discharge Supply: The pulsed power supply for the arc discharge is provided by 185 \( \mu \)F (nominal) electrolytic capacitors with a peak charge voltage of 3 kV (General Atomics part # 39504). A direct-current Spellman SL600 power supply is used to charge the capacitors. The charge circuit is isolated from the pulsed discharge circuit before ignition through high-voltage Ross relay switches. The capacitors are arranged in various series–parallel combinations for different operating voltages and energies. The system is operated between 2 and 5 kV, therefore two capacitors must be connected in series. Additional capacitors are added in parallel pairs to vary the stored energy. The system has been operated up to a maximum stored energy of 4.45 kJ with eight capacitors in series–parallel configuration at a charge voltage of 5 kV. To minimize circuit ringing, a low inductance strip line connects the capacitor bank to the plasma discharge chamber. With four capacitors in a series–parallel configuration, the inductance of the circuit is about 500 nH, resulting in a slightly underdamped pulse.

2) Nonthermal Preionization Trigger Supply: A high-voltage trigger approach was used to generate a streamer discharge that initiates breakdown of the arc plasma, as described in [11]. The high voltage pulse is generated using a solid-state capacitive discharge ignition (CDI) module (Mallory Hyfire 6A) that generates a 500 V pulse, which is then stepped up and inverted through a Mallory automotive transformer to the −50 kV pulse. We modified the CDI module to be triggered directly from a BNC model 555 pulse delay generator, which controls the sequencing of the diagnostics relative to the ignition of the plasma discharge. The output of the transformer secondary is connected to the helical trigger wire.

D. Instrumentation

Characteristics of the thermal arc discharge are determined from voltage and current traces, spectrum analysis, time resolved light intensity, and high-speed photography. Voltage and current traces, as well as high-speed photography are used to characterize the preionization source. The thermal discharge current and voltage are measured using passive Rogowski coils and a Tektronics P5210 differential high-voltage probe. Voltage and current of the high-voltage trigger wire are measured using a Tektronics P6015A high-voltage probe and a Pearson 2877 current monitor. The transient stagnation gauge pressure is measured using a piezoelectric sensor (PCB Piezotronics 113B21). The pressure gauge is epoxied into a 7-mm outer diameter, 8-cm long fused quartz tube to minimize charge accumulation and interference. The time-integrated emission spectrum of the discharge is recorded with an Ocean Optics HR2000+ spectrometer. Time resolved intensity (350–1100 nm) of the discharge is recorded by a ThorLabs PDA36A photodiode and a Dalsa charge-coupled device framing camera that provides a minimum exposure of 1 \( \mu \)s.
that leads to gas expansion and formation of an audible shock wave. The large current densities and high pressures result in a highly collisional plasma that is in thermodynamic equilibrium.

Fig. 4 shows an array of streamers inside the plasma chamber formed by the high-voltage trigger pulse. The streamers are highly branched, following a contorted path as they propagate in the discharge volume. The streamer array generates a volumetric source of seed electrons that provides sufficient ionization for thermal arc breakdown. The brightest streamer emission is observed at the left electrode, regardless of polarity. Enhanced light emission near the left electrode is due to asymmetry in the geometry of the electrode configuration. The left electrode protrudes 10 mm into the reaction chamber decreasing the breakdown threshold potential, whereas the right electrode is flush with the chamber exit.

The ignition threshold potential for a streamer that leads to the formation of a thermal arc depends on the applied potential across the main discharge electrodes. As the main interelectrode charge potential increases for an argon background, the trigger ignition threshold voltage for thermal arc formation decreases. Ignition threshold voltages may be defined as the voltage of the trigger pulse at which sufficient preionization has occurred to induce a thermal arc breakdown. The applied interelectrode potential reduces the required ignition voltage, and therefore, decreases the delay time for thermal arc formation for an argon background. Table I lists the ignition threshold voltages and transition to arc delay times as a function of charge voltage in an argon background. A charge voltage of 1.6 kV is the lowest potential for which breakdown was recorded. The dash indicates thermal breakdown is not reliable and the ignition threshold voltage occurred after the peak of trigger pulse.

**TABLE I**

IGNITION THRESHOLD AND ARC DELAY DEPENDENCE ON CHARGE VOLTAGE FOR ARGON BACKGROUND

<table>
<thead>
<tr>
<th>Charge Voltage</th>
<th>Ignition Threshold Voltage</th>
<th>Transition to Arc Delay</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.6 kV</td>
<td>-</td>
<td>45 µs</td>
</tr>
<tr>
<td>3.0 kV</td>
<td>40 kV</td>
<td>26 µs</td>
</tr>
<tr>
<td>5.0 kV</td>
<td>5 kV</td>
<td>7 µs</td>
</tr>
</tbody>
</table>

* Pressure exceeds limit of pressure transducer

**TABLE II**

RANGE OF OPERATING CONDITIONS WITH 99.5% PURE ARGON FEED GAS

<table>
<thead>
<tr>
<th>Operating conditions</th>
<th>Low</th>
<th>Typical</th>
<th>High</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of capacitors</td>
<td>2</td>
<td>4</td>
<td>8</td>
</tr>
<tr>
<td>Capacitance, µF</td>
<td>89</td>
<td>178</td>
<td>356</td>
</tr>
<tr>
<td>Charge voltage, kV</td>
<td>1.6</td>
<td>3.2</td>
<td>5.0</td>
</tr>
<tr>
<td>Peak current, kA</td>
<td>6</td>
<td>25</td>
<td>76</td>
</tr>
<tr>
<td>Plasma energy, J</td>
<td>85</td>
<td>670</td>
<td>3100</td>
</tr>
<tr>
<td>Peak power, MW</td>
<td>5</td>
<td>40</td>
<td>150</td>
</tr>
<tr>
<td>Pulse Duration, µs</td>
<td>40</td>
<td>70</td>
<td>140</td>
</tr>
<tr>
<td>Peak pressure, kPa</td>
<td>200±50</td>
<td>1500±500</td>
<td>&gt;4000</td>
</tr>
</tbody>
</table>

B. Thermal Plasma Arc Discharge

The pulsed thermal plasma discharge source operates over a range of conditions. The high pressure, and therefore the collisionality, regulates the plasma temperature to approximately 1–2 eV. The plasma temperature is relatively insensitive to the stored capacitive energy, therefore various capacitor configurations will alter the RC time constant and fluidic response even though plasma properties remain uniform.

The discharge has been operated with plasma durations of 40–140 µs, and with energies from 85 J to 3 kJ. Table II summarizes conditions for 99.5% purity argon plasma at the lowest and highest energies we have operated the device, with typical conditions in the middle column. The lower limit of operation is set by the minimum voltage at which breakdown is possible.

In high-power configurations, a shock wave is generated by rapid Joule heating. The sudden and large overpressure can fracture the quartz tube and destroy the test samples. The upper limit of operation of 150 MW is determined by mechanical failure of the chamber. A majority of the plasma/surface interaction studies were conducted at typical conditions. Typical conditions correspond to four capacitors arranged in a series–parallel configuration, charged to 3.2 ± 0.1 kV with argon flowing at 45 ± 2 SLP.

Fig. 5 shows typical normalized voltage, current, and light intensity waveforms for breakdown in 99.5% pure argon using four capacitors at 3.2 kV. Voltage and current traces are under-damped at typical operating conditions, and the light intensity
waveform lags the current. The light emission continues at a low level after the plasma current has reached zero.

C. Emission Spectra of the Thermal Discharge

Time integrated emission spectra of the discharge were recorded by the Ocean Optics spectrometer with a wavelength range from 200 to 1000 nm. The plasma was viewed with an optical fiber and lens that looked either across the diameter of the plasma source \((L \sim 32 \text{ mm})\) or along the length \((L \sim 100 \text{ mm})\) of the chamber. The light emission from the plasma source was dominated by lines of neutral \(\text{Ar I}\) and ionized \(\text{Ar II}\). In addition, emission lines from silicon and hydrogen \((\text{H} \alpha)\) were also observed. Table III summarizes the observed lines and associated energy levels for the argon arc at typical operating conditions.

Plasma temperatures were estimated between 1 and 2 eV assuming a Boltzmann equilibrium distribution for all excited species. The equilibrium Boltzmann assumption is valid for pressures of \(\sim 10^5 \text{ Pa}\) and ionization ratios of \(\sim 10^{-3}\) \[12\]. The spectrum of the discharge for typical operating conditions is shown in Fig. 6.

Some discharges showed line radiation from silicon neutral atoms in the spectral region \(\lambda \sim 250 \text{ nm}\). Silicon neutral lines in the spectrum indicate ablation of the quartz chamber walls. Surface analysis of the quartz chamber wall and test materials confirmed the presence of silicon from the plasma chamber. The ablation of silicon from the quartz wall is discussed in more detail in Section IV.

In all spectra, the radiated power was dominated by line radiation. Optical depth considerations were important when analyzing the plasma spectra because the device operates in a high-density and high-pressure regime. Stark broadening of argon ion lines was too small to be resolved with the Ocean Optics spectrometer. With the addition of a small amount of hydrogen, time-averaged measurements of electron number density from the Stark broadened hydrogen lines were estimated to be \(n_e \sim 10^{23} \text{ m}^{-3}\).

D. Time Evolution of the Thermal Plasma Discharge

High-speed imaging has provided additional insights into the plasma evolution. The most important findings are: 1) the plasma discharge is constricted and does not occupy the available chamber volume and 2) multiple arcs can form, with a high degree of randomness to their shape. Consequently, there is significant shot-to-shot variation in how the core of the plasma interacts with samples in the reaction chamber.

Fig. 7(a) and (b) shows high-speed image sequences for two separate shots of the arc discharge. Fig. 7(a) shows a time evolution of the plasma arc discharge. The arc initially exhibits multiple branched filaments from the anode (left) at 10 \(\mu\text{s}\). The arc filaments attach and track along the surface of the alumina sample-holder to the cathode. At 35 \(\mu\text{s}\) the arc filaments have coalesced into a single diffuse arc discharge. Fig. 7(b) shows the time evolution of another constricted arc filament bridging the interelectrode gap at 10 and 20 \(\mu\text{s}\). There is minimal interaction with the sample holder. Initially, the arc occupies a small localized volume. As the current and temperature rises, the arc volume grows while retaining the initial geometric configuration.

The images show the high degree of randomness in the initial formation and development of the arc filament. Once an arc is formed, the diameter increases until the arc occupies...
Fig. 7. High-speed photography of the thermal arc discharge in argon. (a) Time evolution of filamentary branching and surface tracking along the sample holder at 10 µs, and growth of the filamentary arc at 35 µs. (b) Time evolution of the constricted plasma arc filament that does not interact with the sample holder at 10 and 20 µs.

Fig. 8. 10× photomicrographs taken with a Nikon Optiphot 100 of (a) silicon and (b) sapphire showing deposits of copper from early tests using brass electrodes. Similar samples tested with Elkonite electrodes show no evidence of copper.

approximately a quarter of the chamber volume. Modifications to the design of the plasma chamber are likely required to create uniform and voluminous arcs that interact strongly with the samples. We are presently investigating smaller diameter reaction chambers where the quartz tube surface itself better defines and stabilizes the arc.

IV. INTERACTION OF HIGH-DENSITY PLASMA WITH SILICON AND SAPPHIRE SAMPLES

Presently, the device has been used to study the interaction of thermal plasmas with silicon, alumina, and sapphire test samples orientated orthogonal to the flow. Initial studies were conducted using a brass anode and a steel cathode. In those studies, the samples became coated with copper from the anode, as shown in Fig. 8(a) and (b). To decrease electrode material deposition on test samples, electrodes were fabricated from Elkonite 50W3, an infiltrated 10% copper and 90% tungsten contact material. The high-melting temperature of tungsten prevents evaporation of electrode material, when the arc is in contact with the surface. Copper possesses high thermal and electrical conductivity that allow large currents to flow and reduce local thermal loadings.

Before testing, the electrodes needed be conditioned. About 50 shots were required to evaporate the copper from the surface leaving only the tungsten. Fig. 9 compares the surface of the brass and Elkonite anodes. As expected, the brass shows evidence of gross melting from the arc, whereas the Elkonite shows only small spots that are presumably arc attachment spots on the Elkonite surface.

Evaporated silicon from the wall of the quartz chamber presented a second source of contamination. Silicon deposits were observed on the test samples. The quartz chamber and test samples were analyzed with a Hitachi S-4700 high-resolution SEM equipped with an Oxford Instruments EDXS microanalysis system. Fig. 10(a) shows an SEM photograph of a single-crystal sapphire sample after 15 exposures to the plasma. The sample was coated with a fine dusting of silicon particles with an average particle size of 30–40 nm. At the center of the photograph is an area that was gently touched with the edge of a set of fine tweezers, revealing the sapphire substrate that appears similar to the surface of unexposed sapphire. EDXS analysis of the sample, as shown in Fig. 10(b), confirms there was a significant amount of silicon on the surface of the sample as well as trace amounts of carbon contamination.
Fig. 11 shows a section of the quartz plasma chamber that was removed after approximately 300 exposures. The chamber was exposed to argon plasma at a charge voltage of 3.2 kV. The interior of the chamber closest to the gas inlet electrode (anode for this test sequence) was discolored and slightly opaque [left portion of quartz chamber Fig. 11(a)], whereas practically no damage was observed at the gas outlet [or cathode, right portion quartz chamber Fig. 11(a)]. SEM analysis of the sample from the damaged region showed the surface to contain a number of spheroidal pores on the order of 1 μm in diameter, as shown in Fig. 11(b). Analysis of a cleaved cross section, as shown in Fig. 11(c), showed three distinct zones: number 1 is the boundary between the unaffected quartz and the reacted zone above, number 2 indicates a vitrification zone, and number 3 indicates a highly porous zone.

The damaged zone extends approximately 3.2–3.4 μm into the inner surface of the quartz chamber. Cross-sectional analysis from samples on the cathode side of the chamber did not show any damage. EDXS analysis of the quartz chamber anode region showed a significant silicon deficiency with respect to the control. Carbon contamination was also documented with EDXS that corresponds to the discoloration observed in Fig. 11(a). To prevent silicon deposition on test materials due to the accumulation of damage, the plasma chamber must be replaced after 100 exposures.

Several factors have been proposed to explain the asymmetrical erosion of the plasma chamber. These include polarity of the electrodes, orientation of the trigger wire, and proximity of the gas inlet electrode to the chamber wall. Reversing polarity of the electrodes does not change the erosion pattern; erosion still occurs at the gas inlet side of the reaction chamber. Similarly, reversing the orientation of the trigger wire did not change the location of the damage. Damage of the quartz chamber is likely due to the protrusion of the gas inlet electrode and the associated enhancement of the electric fields on this electrode.

V. Conclusion

A gas-fed plasma source has been developed to study how pulsed, high-density plasmas interact with materials. Plasma parameters for the pulsed atmospheric thermal discharge were determined, including the estimated electron temperature using the Boltzmann plot method, $T_e \sim 1–2$ eV, and the estimated electron number density, $n_e \sim 10^{17}$ cm$^{-3}$. The electron temperature and density are within the expected regime for high-density LTE discharges.

The use of feed gases to form the plasma arc provides a higher degree of control over the plasma composition than can be achieved using consumable liners and exploding fuse wire systems. Although a high degree of control of the plasma chemistry and minimization of contaminant materials has been achieved in the gas-fed plasma chamber, electrode, and chamber wall deterioration was observed. Contamination from the electrodes was minimized using Elkonite 50W3 as the electrode material. Elimination of the quartz chamber wall erosion is difficult due to high-thermal loads and localized streamer surface tracking. High-speed images of streamers and arcs show that the damage to the walls correlates strongly with the location of streamers between the gas inlet electrode and trigger wire.

High-speed imaging has also shown a high degree of variability in the arc formation. Arcs typically evolve to occupy about a quarter of the reaction chamber volume. Consequently, there is a large shot-to-shot variation in how the plasma core interacts with samples. The random shape of the arc column is a consequence of the random nature of the streamers and the large chamber volume. Although the arc structure is random, the current, light intensity, pressure, and emission spectra are highly reproducible.

The gas-fed plasma source that has been developed provides a controllable environment for material interaction investigations and the ability to reliably breakdown at pressure–distance products exceeding the Paschen breakdown threshold. The novel triggering approach provides reliable ignition that does not require the use of ablative liners or exploding wire ignition techniques that would contaminate the system as well as reduce throughput. The plasma device is ideal for highly controllable, low contamination, and plasma–material interaction studies.

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Authors’ photographs and biographies not available at the time of publication.