Microfabricated electrospray emitter arrays with integrated extractor and accelerator electrodes for the propulsion of small spacecraft

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Abstract
Microfabricated electrospray thrusters could revolutionize the spacecraft industry by providing efficient propulsion capabilities to micro and nano satellites (1–100 kg). We present the modeling, design, fabrication and characterization of a new generation of devices, for the first time integrating in the fabrication process individual accelerator electrodes capable of focusing and accelerating the emitted sprays. Integrating these electrodes is a key milestone in the development of this technology; in addition to increasing the critical performance metrics of thrust, specific impulse and propulsive efficiency, the accelerators enable a number of new system features such as power tuning and thrust vectoring and balancing. Through microfabrication, we produced high density arrays (213 emitters cm⁻²) of capillary emitters, assembling them at wafer-level with an extractor/accelerator electrode pair separated by micro-sandblasted glass. Through IV measurements, we could confirm that acceleration could be decoupled from the extraction of the spray—an important element towards the flexibility of this technology. We present the largest reported internally fed microfabricated arrays operation, with 127 emitters spraying in parallel, for a total beam of 10–30 µA composed by 95% of ions. Effective beam focusing was also demonstrated, with plume half-angles being reduced from approximately 30° to 15° with 2000 V acceleration. Based on these results, we predict, with 3000 V acceleration, thrust per emitter of 38.4 nN, specific impulse of 1103 s and a propulsive efficiency of 22% with <1 mW/emitter power consumption.

Keywords: electrospray, spacecraft propulsion, wafer-level integration, small satellites

(Some figures may appear in colour only in the online journal)
potential \( V \) the spray. An accelerator electrode can provide an additional extraction electrode aligned with the capillary emitter to generate negative, is applied between the conductive liquid propellant and an extraction and acceleration stage. A potential (1) and (2), where \( g_0 \) is the gravity constant, \( I_T \) is the emitted beam current, \( V_b \) the beam potential, \( \frac{q}{m} \) is the charge over mass ratio of the emitted particles and \( m \) is the propellant flow rate.

\[
T = I_T \sqrt{\frac{2 V_b m}{q}} \quad (1)
\]

\[
I_{q} = \frac{T}{mg_0} = \frac{1}{g_0} \sqrt{\frac{2 V_b q}{m}} \quad (2)
\]

An attractive configuration is to operate in PIR with arrays of emitters (thousands), effectively multiplying the thrust while retaining the high \( I_{q} \).

Externally fed [10] and porous [11] emitters have been shown to operate principally in PIR using a variety of propellants. Internally fed emitters generate a plume often formed by a mix of both types of species, with the PIR much more difficult to reach. Romero-Sanz et al [12] achieved this in 2003 with the ionic liquid EMI–BF\(_3\) and 20–40 \( \mu \)m inner diameter needles. By actively controlling the pressure drop across the capillary emitter, they showed that PIR could be reached by enforcing very low flow rates, in the order of \( 20 \times 10^{-12} \) Kg s\(^{-1}\) for their configuration. An alternative means to reduce the propellant flow rate \( \dot{m} \) respectively) miniature satellites driven by commercial off-the-shelf smartphones. Yet such missions remain limited by the current lack of a critical technology: a flexible and efficient miniaturized propulsion system capable of carrying the spacecraft beyond their initial orbit.

Electrospray thrusters are acknowledged as one of the most promising technologies to fill this gap, as demonstrated by their inclusion in NASA’s Space Technology Roadmap for In-Space Propulsion Systems [1]. The electrostatically generated spray of high velocity particles creates thrust by direct momentum transfer with few sources of efficiency loss—a fundamental advantage of this technology. Several electrospray thruster systems are in or approaching commercialization [2, 3], yet none are microfabricated.

Microfabrication provides a natural evolution of the technology, building on its strengths by allowing the creation of highly integrated, compact systems ideal for miniature spacecraft. The small feature sizes are key to achieve the high fluidic impedance and thereby low flow rates required for high propellant usage efficiency (with the metric of specific impulse \( \text{ISP} \)), an advantage over conventional systems. Such small emitting features typically generate low thrust (T) levels (tens of mN). However, microfabrication permits large arrays of high specific impulse emitters on a single die; critical to achieving useful (hundreds of \( \mu \)N) thrusts at moderate voltages and within typical small satellite power budgets. The MicroThrust project was an European initiative aimed at developing a thruster module based on these microfabricated electrospray thrusters.

In electrosprays, a conductive liquid propellant (often an ionic liquid) is transported to an extraction site where a large electrical field is applied by precisely aligned electrodes. Increasing the electrical pressure, the liquid’ surface tension can be overcome, deforming the liquid meniscus into a conic shape (Taylor cone [4]), which to a first order stabilizes with a half-angle of 49.3°. The sharpened liquid tip generates a large electric field concentration, in the order of 1 GV m\(^{-1}\), sufficient to atomize the liquid and generate a high velocity particle spray [5]. The extraction field is provided by a single electrode stage, and later stages can provide acceleration and focusing of the spray. We label the potentials supplied by each electrode as \( V_{\text{ext}} \) and \( V_{\text{acc}} \), as shown in figure 1.

Typically, electrospray emitters can be externally [6], porously [7] or internally [8, 9] fed (figure 2). We use the internal approach with passive feeding, relying on capillarity to deliver the liquid. Passive feeding implies that no active pressure control is enforced and that the internal liquid pressure will be driven by the specific configuration of the emitters/reservoir system. It is a system where the liquid is initially static and where flow is entirely induced and controlled by the applied electric field. In a complete system, we envision grooved reservoir walls and porous filling to deliver the liquid to the emitter. As long as the capillary pressure of the small emitters is sufficiently larger than that of these guiding features, the emitters should remain the driving factor, pulling propellant from the reservoir as they get depleted. In this way, we remove the need for pressurized tanks and lines—a great advantage from a system perspective. The present work does not include such a reservoir and we employ gravity to bring the propellant in contact with the emitters, delivering small amounts of liquid on the backside of the thruster. It is important to note that this pressure, due to gravity, is not sufficient to overcome the capillary holding pressure once filled. Hence a flow is only induced, and controlled, by the emission processes after applying a voltage.

Depending on the operating conditions and emitter design, the nature of the emitted species can be modified, from single ions to nanodroplets. In the purely ionic regime (PIR), the thruster boasts optimal specific impulse, though at the cost of low thrust per emitter. This tradeoff is highlighted in equations (1) and (2), where \( g_0 \) is the gravity constant, \( I_T \) is the emitted beam current, \( V_b \) the beam potential, \( \frac{q}{m} \) is the charge over mass ratio of the emitted particles and \( m \) is the propellant flow rate.

![Figure 1. Basic operation of an electrospray thruster with an extraction and acceleration stage. A potential \( V_{\text{ext}} \), positive or negative, is applied between the conductive liquid propellant and an extraction electrode aligned with the capillary emitter to generate the spray. An accelerator electrode can provide an additional potential \( V_{\text{acc}} \) to increase the thruster performance.](image-url)
is to increase the fluidic impedance of the emitters \( (m = \Delta P / Z) \), where \( Z \) is the fluidic impedance and \( \Delta P \) is the pressure drop), which can be achieved through microfabrication. For microfabricated emitters, PIR could be reached by our group [13] and De La Mora’s [9], in both cases using very high impedance, bead-filled micro-capillaries yet still with active pressure control.

PIR has never been achieved using passively fed, internally wetted emitters, as targeted by our recent work. An unpressurized and passive feeding system has the advantage of being more compact, since no high pressure reservoir or valve is needed. More importantly, it removes the risk of flooding due to over-pressurization. Reaching the necessary high fluidic impedance without requiring the bead filling step is also an advantage, as it improves the uniformity of the fluidic impedance. The devices presented in this paper could nearly be operated in PIR, with typical plumes composed of 95% ions (monomers and dimers) using the ionic liquid EMI–BF₄ as propellant.

Previous thruster prototypes produced by our group were comprised of microfabricated arrays of 19 bead-filled capillary emitters [13, 14] with a single extraction stage. Our latest generation uses a simpler, improved emitter fabrication process and wafer-level bonding of the electrodes, which we had proposed earlier [15]. The new process permits the large scale fabrication of high aspect ratio capillaries with the smallest <10 μm inner diameters to date. For the first time, we report on the wafer-level integration of individual acceleration electrodes—a critical step in the development of this technology. The addition of accelerators increases simultaneously the specific impulse, thrust and system efficiency, and introduces new system features, such as power tuning and thrust vectoring and focusing. Integrating these electrodes at wafer-level, we can provide individual accelerators for each extractor/emitter pair with high density and excellent alignment. Individual accelerators are advantageous over the one accelerator/multiple emitter configuration, since each spray is accelerated uniformly and there are no edge effects. Uniformity and alignment of the acceleration field is important: errors may lead to off-axis spraying which elevates the risk of spraying onto the electrodes, reducing the propulsive efficiency and thruster lifetime. Furthermore, off-axis spray might also lead to errors in thrust vectoring, since any asymmetry in spray produces spurious lateral torque.

In sections 2 and 3 of this paper, we discuss the effect of these electrodes on the thruster performance through modeling and simulation. The overall design of the thrusters is described in section 4 and the process allowing the fabrication of large (127 emitters) and uniform arrays, is presented in section 5. In section 6, we study the performance of the accelerator electrodes through I–V curves, time-of-flight measurements and beam shape characterization, discussing the results in section 7.

2. Impact of acceleration electrode

The propulsive efficiency \( \eta_T \) is defined as the ratio between the propulsive beam power \( P_e \) and the input electrical power \( P_e \) required to generate it ((3), where \( m \) is the mass flow rate, \( I_{emi} \) is the source current and \( V_{app} \) is the applied potential, equal to \( V_{ext} + V_{acc} \) which are shown in figure 1).

\[
\eta_T = \frac{P_e}{I_{emi}V_{app}} = \eta_p\eta_I\eta_\theta.
\]

(3)

The polydisperse efficiency \( \eta_p \), the energy efficiency \( \eta_I \) and the angular efficiency \( \eta_\theta \) are, assuming sufficient physical clearance of the electrodes and low propellant evaporation, the principal sources of propulsive efficiency loss [16]. Equations (4) and (5) highlight how thrust and specific impulse can be increased by supplying additional electrical power to the thruster:

\[
T = \sqrt{2\eta_T mP_e}
\]

(4)

\[
I_{sp} = \frac{1}{g_0} \sqrt{\frac{2\eta_T P_e}{m}}
\]

(5)

Yet for a given thruster geometry, stable spray only occurs in a relatively small extraction voltage range, limiting the ability to boost performance by a direct increase of \( V_{ext} \). The introduction of the acceleration electrode enables this feature, since the applied potential \( V_{app} \) can now be increased while keeping \( V_{ext} \) at an optimal level. Of course, the square root relationship between the performance increase and electrical input power indicates that the advantages of acceleration will diminish with increasing voltage, limiting the scale-up potential. Yet at the modest voltages of a few kV, which can well be supplied by miniaturized power sources [17], the increase in thrust and \( I_{sp} \) can have a major impact on the mission design and feasibility—entirely justifying the additional power requirement.
points, corresponding to droplet size. Larger droplets lead to a sharper decline in efficiency.\(\frac{b}{1+\kappa}\) that will be reached by an emitted particle\(V\) the energy and the angular efficiencies. Increasing the propulsive efficiency, in particular in terms of chemical degradation of the ionic liquid propellant \([18]\). Balancing schemes necessary to suppress reported electro-current control \((emi)\), this arrangement will ensure that variance\(I\) going through an electric field provided by electrodes at\(V\) loss is not related to the post-acceleration and is constant throughout the thruster lifetime. Through decoupling thrust and current control \((I_{\text{em}})\), this arrangement will ensure that variable thrust can be achieved without compromising charge balancing schemes necessary to suppress reported electro-chemical degradation of the ionic liquid propellant \([18]\).

Acceleration provides the additional advantage of increasing the propulsive efficiency, in particular in terms of the energy and the angular efficiencies.

The energy efficiency, given by \((6)\) \([19]\), relates the equivalent potential \(V \text{app}\) that will be reached by an emitted particle going through an electric field provided by electrodes at potential \(V \text{app}\).

\[
\eta_p = \left[ \frac{V_p}{V_{\text{app}}} \right] = 1 - \frac{V_{\text{loss}} + |RI|}{V_{\text{app}}} \tag{6}\]

In \((6)\), \(V_{\text{loss}}\) is related to the electrical energy required to form the cone and extract particles from the liquid while \(RI\) is the potential drop due to ohmic losses in the liquid.

For a highly conductive ionic liquid, we approximate \(R\) to 0. \(V_{\text{loss}}\) is not related to the post-acceleration and is constant regardless of the acceleration potential applied, such that the efficiency will approach unity as the acceleration voltage is increased. The value of \(V_{\text{loss}}\) is difficult to predict and can be most easily measured through the use of retarding potential analysis \((\text{RPA})\). For this model, we use the value of 100 V, which is generally large \(\eta_p\), the droplet mode inevitably leads to low specific impulse \((\text{equation } (2))\). To increase the \(I_{\text{sp}}\), the droplet size must be reduced, which often leads to a gradual increase in ion content and lower efficiency. Operation in the PIR, with little to no droplets \((\text{well below } 1\%)\) present, offers a solution to this tradeoff. In this case, both high efficiency and specific impulse are achieved.

Combining the effect of increased beam potential and propulsive efficiency, and assuming that acceleration does not affect the propellant flow rate and emitted current, we can define an acceleration performance factor, \(\kappa_a\), as the ratio between the thrusts of an accelerated and unaccelerated beam:

\[
\kappa_a = \frac{T_{\text{acc}}}{T_0} = \frac{I_{\text{sp,acc}}}{I_{\text{sp,0}}} = \left\lfloor \frac{V_{\text{app,accelerated}}}{V_{\text{app,0}}} \right\rfloor \frac{\eta_{\text{accelerated}}}{\eta_{\text{nonaccelerated}}} \tag{9}\]

Table 1 lists computed values of \(\kappa_a\) and its constituents with applied acceleration ranging from 0 to 3000 V. We assumed an extraction voltage \(V_{\text{ext}} = 1000 \text{ V}\) and a linear decrease in the plume half angle so that the angle is decreased 30° to 20° between 0 V and 3000 V acceleration.

With the above assumptions, a 2.16 increase in thrust and \(I_{\text{sp}}\) can be obtained with 3000 V acceleration, independently

\[
\eta_p = \cos^4 \left( \frac{\theta}{2} \right) \tag{7}\]
of beam composition and emitted current. This gain, however, comes at the cost of a 4x increase in electrical power.

To compute the efficiencies in table 1, we made two major assumptions. The first is that the acceleration did not have an affect on the nature or amount of charge being emitted (mass flow rate). Second is the beam half angle reduction from 30° to 20°. In the next section, we will review these assumptions through simulations.

3. Simulations

3.1. Extraction field and electrospray onset

Modeling of the electrical field in the emission region was done using COMSOL, extending the model of Krpoun [26] to study the effect of the acceleration electrode on onset. The iterative procedure takes a 49.3° truncated cone topped by a spherical tip. The cone is modeled as a solid, perfectly conductive surface which is gradually sharpened, keeping equilibrium between the liquid surface tension and electrical pressure. We compared the electrical field induced at the tip with and without accelerators, using a fixed potential of 1000 V between the emitter and extractor. For the accelerators, we varied two geometric parameters: the accelerator inner diameter, ranging from 150 µm to 1000 µm and the extractor/accelerator separation, from 100 µm to 1500 µm (see figure 4).

In both cases, and with up to 3000 V acceleration applied, we obtained less than 5% variation in electrical field, negligible compared to the effects of the extractor design. This result indicates that the accelerator has little impact on the electrical field in the extraction region, supporting the argument that acceleration can be treated independently from emission.

3.2. Ion-beam optics

SIMION® was used to simulate the focusing effect of the accelerator. Conceptually, we assume that the particles are distributed uniformly and with equal energy in a cone of half-angle BHA. We generate a particle at the exit point of the extractor electrode. This particle has an energy of 1000 eV, representing the acceleration of a single charge through the 1000 V extraction field, and an initial angle BHA_initial ≈ 28.7°. This angle was chosen so that the particle would not physically impact the acceleration electrode in any simulated configurations, allowing us to compare the effects of accelerator inner diameter and extractor-accelerator separation. Note that the nature of the species (ion or droplet) is irrelevant in this case, as only the particle energy (q · Vbeam) affects the trajectory. The extractor electrode, 50 µm thick and spaced 50 µm from the capillary emitter has an inner diameter of 168 µm. We probe the particle direction at the exit of the accelerator (BHA_final) and compare it with the angle at the exit of the extractor to get the relative focusing. Again, the accelerator inner diameter and the extractor/accelerator spacing were varied, respectively from 370 µm to 1000 µm and from 100 µm to 400 µm.

For these two design parameters, figure 5 shows how the beam half angle is reduced when introducing acceleration. Typically, a 30–35% reduction seems achievable with 3000 V acceleration voltage (BHA_final≈19°). The accelerator inner diameter has an inverse relationship with the final beam half angle, showing that smaller accelerators are more effective at focusing the beam. The accelerator-extractor spacing also has an effect on the focusing, with larger spacing being beneficial. In both cases, though, the dimensions have, at least in the range studied, almost negligible effects. Indeed, the extreme cases in all simulations, for the best and worst focusing, would lead to less than 1% variation in κq.

4. Electrospray thruster design

The thruster design is a progression over previously published devices [14], which included only the extraction stage. We now introduce an acceleration electrode, completely integrated in the fabrication process. From the above simulations, we find that the extractor and accelerator design can be de-coupled and optimized individually, with the criteria of performance, reliability and manufacturability. The end device results from the wafer-level integration of an emitter and electrode wafer using a laminated thin film. This enables the optimisation of the emitters and electrodes separately, providing valuable flexibility in the design. The emitter

http://simion.com
wafer contains the capillary emitters and feedthrough reservoirs while the electrode wafer includes the extractor and accelerator electrodes. Figure 6 is a cross-section diagram of the thruster stack, with dimensions listed in table 2. With limited information on the real focusing effects of the accelerator electrode, we fabricated chips of three accelerator diameters, which allow increasing beam clearances. The baseline design allows approximately 40° half-angle clearance, while the daring and safe variants allow 30° and 50° respectively. A smaller clearance, or accelerator diameter, is not only beneficial for its improved focusing effect, but most importantly to allow tighter array packing, which ultimately results in increased thrust density.

4.1. Capillary emitter design

A key dimension for the capillary emitter design is its inner diameter, $D$, which in turn determines its fluidic impedance. A high impedance will contribute to a low propellant flow rate which is critical to reaching the high $I_{sp}$ PIR. Prior to this work, our devices relied on the filling of wider capillaries (24 $\mu$m inner diameter) with silica microbeads to tailor the hydraulic impedance and operation mode of the device [13], as also done by Lenguito et al [27]. However as our work increasingly focused on the fabrication of large arrays, this method became impractical due to the random arrangement of the beads which could lead to non-uniformities in operation. Instead, our group has focused on improving the etching step of the emitter interiors to allow narrower emitters with a more uniform impedance.

We aimed to achieve similar impedance to the bead-filled capillaries [13], which could be operated in PIR. A significant difference remains that these earlier emitters were operated with active pressure control and so it is difficult to guarantee that the new emitters, even with equal impedance would be equally ionic. Yet one can expect that a passively fed capillary operated in the same conditions and having no backpressure would necessarily have a flow rate lower or equal to a pressurized one, which would be consistent with PIR.
To compute the impedance of the bead-filled emitters, we follow a model adapted from Ergun [28], as suggested by Lenguito et al [9]. Using (10), we compute the effective inner diameter \( D_{\text{eff}} \) of the emitter, which can be inserted in the Hagen-Poiseuille equation (11).

\[
D_{\text{eff}} = \sqrt[3]{\frac{128 \cdot \mu \cdot L \cdot D^3 \cdot \epsilon^4}{600 \cdot (1-\epsilon)^2}} \tag{10}
\]

\[
R_{\text{fl}} = \frac{128 \mu L}{\pi D_{\text{eff}}^4} \tag{11}
\]

where \( \mu \) is the dynamic viscosity of the fluid (0.038 Pa·s for EMI–BF4 [29]), \( L \) is the length of the capillary, \( D_i \) is the diameter of the microspheres and \( D \) is the inner diameter of the capillary. The void fraction \( \epsilon \), which ranges from 0 to 1 describes the packing of the beads inside the cylinder. For macroscopic cylindrical beds, \( \epsilon \) stabilizes to 0.4 starting 5 \( D_i \) from the wall (both through model and experiment [30–32]). Closer to the wall, the packing varies considerably, with \( \epsilon \) oscillating between approximately 0.25 and 0.65. In our earlier bead filled devices, the cylinders themselves measured 24 \( \mu \)m and the beads 5 \( \mu \)m, so that the ‘stabilized’ value of 0.4 cannot be directly used. Instead, we assume a range of \( D_{\text{eff}} \) resulting from void fractions \( \epsilon \) ranging from 0.25 (\( D_{\text{eff}} = 3.0 \) \( \mu \)m) to 0.65 (\( D_{\text{eff}} = 9.1 \) \( \mu \)m).

Figure 7 shows how the bead filled emitters compare with the emitters presented in this work. We aimed for emitter inner diameters of 5.0 \( \pm /-0.5 \) \( \mu \)m, but lateral attack in the etch process led to slightly wider fabricated devices (7.9 \( \pm /-0.5 \) \( \mu \)m), as will be discussed later. Considered in the coordinates of figure 11, our targeted impedance was therefore comparable to the low range predicted using bead filling. For our fabricated devices we estimate an order of magnitude lower impedance than our target and the high range of the bead-filled emitters.

The emitter tip is also sharpened using an isotropic etch which attacks the perimeter of the cylindrical emitter. This etch effectively reduces the width of the annular plateau at the tip of the capillary. This sharpening reduces the operation voltage and is favorable for liquid containment at the tip. We aim for 15 \( \mu \)m lateral etch, leading to a 5 \( \mu \)m plateau. This value is the maximum that can be safely achieved without risk of breaching the capillary tube considering the fabrication tolerances in terms of emitter inner and outer diameter as well as alignment.

### 4.2. Electrode design

A driving requirement is to minimize the extraction voltage. While a higher \( V_{\text{ext}} \) may be useful to increase the beam energy and thruster performance, we find that a lower voltage at this level leads to a more reliable chip and simpler system design. Post-acceleration remains the preferred choice to increase the performance. From earlier simulations [26], we see that the key drivers to reduce \( V_{\text{ext}} \) are the extractor electrode diameter and the extractor-emitter gap. The former is designed at 168 \( \mu \)m, enough to allow 40° beam half-angle clearance while the latter is kept at 50 \( \mu \)m. Earlier devices used a 15 \( \mu \)m film to separate the emitter and extractor, but this provided insufficient electrical insulation. Although the bulk film’s breakdown strength was measured at 2.3 kV, leakage current problems forced the use of a thicker, 50 \( \mu \)m film. Experiments showed that the leakage occurred at the edge of the die where conductive silicon dust generated at dicing contaminated the film. Insulation between the extractor and acceleration electrode levels is provided by 250 \( \mu \)m of glass. This dimension should be as small as possible as it drives, given the non-zero beam half-angle, the packing density of the emitters. From the ion beam simulations of section 3, we also see that this thickness, which forms the gap between extractor and accelerator, has minimal impact on beam focusing.

Thinner glass insulation may have been sufficient but led to low fabrication yield due to the difficult handling of thin glass wafers. The current thickness, expected to insulate upwards of 15 kV, allows significant flexibility in the system, effectively permitting any acceleration voltage to be applied within the limitations of typical small spacecraft. The opening of the glass on the top side, follows the allowed beam half-angle clearance and has diameter of 370 \( \mu \)m, 540 \( \mu \)m or 765 \( \mu \)m. The final baseline thruster includes 127 emitters packed with a density of 213 emitters \( \text{cm}^{-2} \). Variants have 91 (125 emitters \( \text{cm}^{-2} \)) and 167 emitters (257 emitters \( \text{cm}^{-2} \)).

### Table 2. Design parameters of MicroThrust electrospray thrusters (All dimensions in micrometers.)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Identification</th>
<th>Nominal dimension (( \mu )m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emitter inner diameter</td>
<td>Cap ID</td>
<td>5</td>
</tr>
<tr>
<td>Emitter wall thickness</td>
<td>Cap Wall</td>
<td>20</td>
</tr>
<tr>
<td>Emitter pitch</td>
<td>Cap Pitch</td>
<td>670, 737, 963</td>
</tr>
<tr>
<td>Emitter isotropic etch radius</td>
<td>Cap isoEtch</td>
<td>15</td>
</tr>
<tr>
<td>Extractor electrode diameter</td>
<td>Ext ID</td>
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<tr>
<td>Extractor electrode thickness</td>
<td>Ext dev H</td>
<td>50</td>
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<tr>
<td>Extractor-emitter gap</td>
<td>Ext gap H</td>
<td>50</td>
</tr>
<tr>
<td>Glass thickness</td>
<td>Ext handle H</td>
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</tr>
<tr>
<td>Accelerator electrode diameter</td>
<td>Acc ID</td>
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</tr>
<tr>
<td>Accelerator to accelerator spacing</td>
<td>Acc gap</td>
<td>150</td>
</tr>
</tbody>
</table>

![Fluidic impedance vs capillary inner diameter](image)

*Figure 7. Estimated fluidic impedance of capillary emitters.*
5. Thruster fabrication

Although the end structure is complex, microfabrication and wafer-level integration techniques allowed decoupling of many structural complexities. In this particular case, the fabricated capillaries and integrated extractor/accelerator electrodes are fabricated on separate substrates, respectively the emitter wafer and electrode wafers.

5.1. Capillary emitter wafer

Principal fabrication steps of the emitter wafer are shown in figure 8. A 2.2 μm thermal oxide layer is grown on the starting SOI wafer (100/2/350 μm) (a). The oxide layer is patterned using a 4.8 μm thick 10:1 AZ9260:PGMEA mixture photoresist spun at 3500 rpm. The emitter interior is then patterned by DRIE (b). Control of this etch is critical as it will drive the fluidic impedance of the emitter. Fabricated emitters have an inner diameter measured at 7.9 ± 0.5 μm, a result of the slight widening typically seen in high aspect ratio DRIE. Next, the liquid reservoir is patterned by DRIE on the backside of the wafer, stopping on the buried oxide (c). Third, the exterior of the emitters is patterned using the same resist mixture described above (c). The resist exhibits sufficient viscosity to cover the emitter orifices cleanly and provide a uniform surface. In some cases, resist residues remained inside the emitters, even after several attempts at plasma O₂ cleaning, indicating that further optimization is required at this stage. The resist and oxide hard mask are then patterned before a final, two-step dry etch is performed. The first step, isotropic, sharpens the emitter tip, reducing the onset voltage and restricting the ionic liquid from overflow. The second step is anisotropic and provides clearance from the emitter tip to the silicon bulk ((d)–(e)). A final vapor HF etch is done to remove the BOX, connecting the backside reservoir to the capillary emitter (f). SEM images of completed single emitters (top side) are seen in figure 9.

5.2. Extraction and acceleration electrode wafer

The novel electrode process provides each emitter with an individual extractor and, for the first time, accelerator electrode.
The process begins with the anodic bonding of a 300 µm thick silicon wafer to a 250 µm thick Borofloat 33 wafer patterned by micro sandblasting (figure 10(a)). With its low thermal expansion coefficient (3.25 · 10^{-6} K^{-1}), Borofloat 33 is a viable alternative to Pyrex for anodic bonding.

The silicon wafer is patterned through a single DRIE, while the glass is patterned by a novel micro-sandblasting method [33] offered by Icoflex. Unlike with silicon, deep high-aspect ratio patterning of glass is challenging. While laser machining allows precise structuring, it is time consuming and expensive. Photo-patternable glasses (PPGs) such as APEX 10 and Foturan 11 offer good alternatives but are relatively difficult to process and even more difficult to be used in anodic bonding, due to their high thermal mismatch with silicon. HF based wet etches [34] are fast and well developed, but their isotropic nature limits the packing density that can be achieved. Micro sandblasting, on the other hand, is a fast and inexpensive wafer-level technique. Although limited aspect ratios can be achieved with this technique (of the order of 1:1) and despite being less precise than PPGs, it offers reasonable side-wall angles (15–20°) and is well suited for this application (See also Lenard et al [35]).

After bonding, the silicon is ground and polished to a thickness of 50 µm to reveal the extractor pattern and create through-wafer openings. 2 µm silicon dioxide is then sputtered on the bottom of the wafer to provide electrical insulation between the extractor and emitter levels, and metal is evaporated on the front side through a shadow mask to create the accelerator electrodes (c). Due to the insular nature of the patterns to be masked, two depositions (200 nm thickness) are required to generate the adequate electrode pattern, each rotated by 180°. An SEM image of a cleaved electrode, before metal deposition, is shown in figure 11.

5.3. Wafer-level integration

In a final microfabrication step, the emitter and electrodes are wafer-level bonded using a dry photo-patternable negative resist (DuPont MX5050). The resist is laminated on the bottom of the electrode wafer and patterned by standard lithography and development. Alignment of the two wafers is done in a Suss Microtech MA6/BA6 in bond alignment configuration and the stack bonded by thermo-compression at 150°C and 0.3 MPa in a Suss Substrate Bonder SB6.

Figure 10. Fabrication process of the electrode wafer. (a) The patterned glass, 250 µm thick, is patterned by micro sandblasting before anodic bonding to a patterned silicon wafer. The wafer is ground and polished (b) before oxide and metal are deposited on the bottom and top, respectively (c).

Final alignment is typically better than 10 µm. Figure 12 shows SEM images of the assembled thruster stack while figure 13 is an optical image of a finished die.

6. Test and characterization

The test setup (figure 14) consists of two chambers: a main chamber for spraying and a second for the storage of the ionic liquid. The liquid is itself transferred using a glass capillary by applying a pressure differential and dropped on the backside of the assembly, where capillarity transports it to the emitter tip. The spray direction is vertically downwards.

Electrical connections to the extractor and accelerator are made from the front side while connection to the emitter is through the ionic liquid, via a central metal section within the PEEK holder (not shown). Generally, the accelerator is grounded, while both extractor and emitter potentials are raised above ground to generate extraction and/or acceleration. The ionic liquid 1-ethyl-3-methylimidazolium tetrafluoroborate (EMI−BF₄) is used as the propellant. This ionic liquid is commonly used to demonstrate thrust and has been known to operate an a variety of regimes: purely ionic [11], mixed droplet/ion [3] or droplet.

The emitted spray is collected either by a large (220 mm) plate situated 500 mm downstream or by a small Kimball Physics Faraday cup mounted on a 280 mm translation stage at the same distance. The stage can move laterally on one axis,
mapping the plume which is assumed to be axisymmetric. All the emitters of the array are contained within a 6 mm radius of the center, so that the effect of their finite position on the measured beam profile is negligible. The first configuration is used for time-of-flight (ToF) and IV measurements, while the second allows characterization of the plume shape. For the collection of ToF data, a DEI PVX-4150 high speed switch is used to rapidly (20 ns) turn off spraying by connecting the extractor to the emitter potential. Monitoring the flight time of the particles, it is possible to determine their nature and consequently the generated thrust, specific impulse and propulsive efficiency [36].

In all cases, the current is measured on the emitter, extractor and accelerator by the measurement of the voltage drop across a resistor. More details on the test setup, in addition to extended characterisation of the thruster mounted in its breadboard assembly, can be found in [36].

Figure 16 plots the current measured at the emitter using the potential divider system against the applied voltage difference between the emitters and the extractor ($V_{ea}$). The emitted current is proportional to the plume current. Data

![Figure 12](image12.png)

**Figure 12.** SEM images of thruster chips showing arrays of emitters aligned to individual extractor and accelerator electrodes.

![Figure 13](image13.png)

**Figure 13.** Optical image of diced thruster chip. The top right and bottom left metal pads allow contact with the accelerator layer, with the top left and bottom right contact the extractor level.

![Figure 14](image14.png)

**Figure 14.** The test setup. For beam shape measurements, the large (220 mm) target plate is replaced by a small plate mounted on a small translation stage.

![Figure 15](image15.png)

**Figure 15.** Emitter array installed in test assembly. Contact to the extractor and accelerator electrodes can be verified in situ by measuring the resistance between two opposite pins.
is plotted for various acceleration voltages \(V_{\text{acc}}\) defined as the voltage applied to the extractor to raise it above ground. The actual accelerator on the THC is at ground throughout. Data is shown for the baseline design, i.e. with 540 \(\mu\)m accelerator diameter and 127 emitters. There is a strong linear trend of emitter current, independent of the acceleration voltage. This independence demonstrates that the electric field within the emitter-extractor region is not affected by field conditions downstream, as predicted by the onset voltage simulations. The gradient of the IV relationship in figure 16 is \(\sim0.4\ \mu\text{A per emitter-extractor volt}\). This strong relationship is in agreement with previous studies using large colloid thruster arrays [7].

Figure 17 is an example of the plume profile from a single emitter as the applied acceleration voltage is increased from 0 to 2000 V. A clear focusing effect is seen, with the plume profile reduced from \(\sim30^\circ\) to \(\sim15^\circ\), corresponding to an angular efficiency increase from 87% to 97%. Figure 18 plots the measured BHA against acceleration voltage.

We use time of flight measurements to determine the nature of the emitted species and calculate the specific impulse and thrust. A sample ToF spectra is presented in figure 19. The plot displays a large initial drop corresponding to the arrival of ions followed by a quickly decreasing droplet tail. Typical results of 127 emitter arrays show that the beam is composed at 95% of ions (monomers and dimers) with the remaining droplets being clusters of \(\approx230\) ions. Table 3 summarizes these results and lists, for 850 V extraction, the resulting calculated flow rates, efficiency, specific impulse, thrust and power consumption. All values have been been divided by the array size and are consequently average values per emitter.
Characterization of the emitter arrays has shown very promising results in terms of the usability of acceleration electrodes. It was successfully shown that the acceleration and extraction stages could be decoupled, an important element to simplify the system and mission design. Beam shape focusing proved better than expected from simulations, with measured beam half-angles being focused from approximately 30° to 15° with 2000 V acceleration.

The composition of the beam, with 95% ions, is approaching the desired PIR, although the small amount of droplets drastically reduced the achieved specific impulse and efficiency. With the droplet size seen here, the minimum theoretical polydisperse efficiency occurs at 94%, close to the present situation. Yet it is encouraging that the curve minimum (figure 3) was passed and that any further increase in the ion content will rapidly lead to better performance. We expect that this can be achieved by further reducing the inner diameters of the emitters, thereby increasing their hydraulic impedance and reducing the flow rate.

We can compare the performance of acceleration for the beam composition seen here with positive and negative ionic regimes. Table 4 lists the calculated thrust, specific impulse, total efficiency and power for unaccelerated and accelerated beams. Cases 1 and 2 describe, for comparison purposes, calculated performance parameters based on the current levels and focusing observed. Case 1 refers to a 50%/50% mix of positive monomers and dimers while case two is the equivalent mix of negative monomers and dimers. Case 3 corresponds to the composition of a typically observed beam (95% ions, with $M_{\text{droplet}} = 230M_{\text{ion}}$). In all cases, we use $V_{\text{ext}} = 850$ V, $I_{\text{em}} = 250$ nA and assume 30° to 15° focusing.

To illustrate the potential benefits of acceleration, we take as a reference a miniature HV power supply [17] developed for this application and capable of delivering 3800 V in the positive and negative polarities. With ≥800 V required for emission, we can use up to 3000 V acceleration, which increases all metrics by a factor of 2.32x over the unaccelerated case, at the cost of a 4.5x increase in power. If 100 nN total thrust was necessary for a given propulsion configuration, arrays of 2600 emitters would be required. The total power consumption, increased to 2.5 W, is well within the range that can be provided by even a single unit CubeSat. With the currently achieved packing density (213 emitter cm⁻²), this array would require approximately 12 cm² of surface, still just 12% of a CubeSat face.

### Table 3. Measured beam composition using time of flight analysis.

<table>
<thead>
<tr>
<th>(I_{\text{em}}) (nA)</th>
<th>(\langle \mu \rangle_1) (C kg⁻¹)</th>
<th>(f_1)</th>
<th>(\langle \mu \rangle_2) (C kg⁻¹)</th>
<th>(f_2)</th>
<th>(\epsilon)</th>
<th>(\dot{m}) (kg s⁻¹)</th>
<th>(\eta_T)</th>
<th>(I_{\text{op}}) (s)</th>
<th>(T) (nN)</th>
<th>(P) (µW)</th>
</tr>
</thead>
<tbody>
<tr>
<td>250</td>
<td>8.68e5</td>
<td>0.95</td>
<td>3.81e3</td>
<td>0.05</td>
<td>4.4e⁻³</td>
<td>3.55e⁻¹²</td>
<td>0.18</td>
<td>474</td>
<td>16.5</td>
<td>212.5</td>
</tr>
</tbody>
</table>

### Table 4. Expected thruster performance improvement with acceleration. All values are given per emitter.

<table>
<thead>
<tr>
<th>Spray composition</th>
<th>Unaccelerated</th>
<th>3000 V Acceleration</th>
</tr>
</thead>
<tbody>
<tr>
<td>(T) (nN)</td>
<td>(I_{\text{sp}}) (s)</td>
<td>(\eta_T)</td>
</tr>
<tr>
<td>Case 1&lt;sup&gt;a&lt;/sup&gt;</td>
<td>12.9</td>
<td>2421</td>
</tr>
<tr>
<td>Case 2&lt;sup&gt;b&lt;/sup&gt;</td>
<td>12.0</td>
<td>2550</td>
</tr>
<tr>
<td>Case 3 (Observed)</td>
<td>16.5</td>
<td>474</td>
</tr>
</tbody>
</table>

<sup>a</sup> Positive beam composed of 50% [EMI]<sup>+</sup> and 50% [EMI–BF<sub>4</sub>]<sup>+</sup>

<sup>b</sup> Negative beam composed of 50% [BF<sub>4</sub>]<sup>-</sup> and 50% [EMI–BF<sub>4</sub>]<sup>+</sup>

<sup>c</sup> Typical mix found when operating in ‘mixed mode’ with present emitters

### 7. Discussion

Due to their high achievable specific impulse, efficiency and miniaturizability, microfabricated electrospray thrusters are one of the most promising technologies for the propulsion of small spacecraft. The inclusion of integrated acceleration electrodes is a key step in their development, allowing not only significant performance increases but also a number of new device features. Acceleration improves the thrust and specific impulse of the device not only by raising the energy of the emitted particles, but also by increasing the thruster efficiency. Models of the acceleration electrode indicate that, within the design space of interest, the effect of these electrodes on the extraction of the spray is negligible, an element confirmed by experiments. This decoupling is critical as it permits tuning of the acceleration potential while keeping an optimal extraction voltage.

We developed a process with allows the wafer-level fabrication of large arrays of emitters, each with integrated extractor and accelerator electrodes. The emitter’s critical dimensions are highly repeatable, with carefully isotropically etched tips and high aspect-ratio interiors (7.9 ± 0.5 µm inner diameter, 100 µm deep). The electrodes, insulated by 250 µm of glass patterned by a novel micro-sandblasting process, are wafer-level bonded to the emitters to improve device reliability and minimize the footprint.

Arrays of 127 capillary emitters were successfully tested using the ionic liquid EMI–BF<sub>4</sub> and presented beams composed of 95% ions, with current levels in the range of 0.25 µA per emitter. From plume shape measurements, we confirmed that the individual accelerator electrodes could effectively focus the plume, bringing the beam half angle from 30° to 15°. Next steps will include a further reduction of the emitter inner diameters, with the aim of lowering flow rates and increasing ion transport efficiency.
content. In this way, we hope to achieve purely ionic emission, maximizing the specific impulse and efficiency of the thruster.

Acknowledgments

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