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Numerical and experimental comparative study of microfluidic fuel cells with different flow configurations: Co-flow vs. Counter-flow cell

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Abstract

Microfluidic fuel cell (MFC) is a new type of fuel cell which utilizes two laminar flows as electrolyte for both reactant delivery and ionic conduction purposes. According to the flow configuration, two MFC designs, that is, the co-flow cell and the counter-flow cell, have been proposed in the literature, but the specific merits and demerits between them have not been well understood yet. Therefore, in this work both experimental and numerical comparative studies are conducted on this issue. It is found that the counter-flow MFC encounters an inherent drawback on its power output because the outer part of its electrodes contributes much less than the inner part due to the longer ionic transport path. In conclusion, short and wide electrodes are more appropriate for counter-flow MFCs rather than the conventional long and narrow ones. Nevertheless, the counter-flow MFC does exhibit great advantage on low flow rate tolerance, leading to much improved fuel utilization and energy density at the same time. Considering this huge superiority for real applications, structural optimization of the counter-flow MFC is further conducted by reducing the electrode distance and discarding the outer part of the electrodes. The
optimized counter-flow cell performance is successfully improved to the same level with its co-flow counterpart.

**Keywords**

Microfluidic fuel cell, co-flow, counter-flow, electrode configuration, fuel utilization

**Nomenclature**

**Symbol:**  
ρ: Density (kgm\(^{-3}\))  
u: Velocity (ms\(^{-1}\))  
P: Pressure (Pa)  
μ: Dynamic viscosity (Pa·s)  
ω: Mass fraction  
x: Molar fraction  
j: Diffusion flux (kgm\(^{-2}\)s\(^{-1}\))  
S: Production/consumption rate due to electrochemical reactions (kgm\(^{-3}\)s\(^{-1}\))  
M: Molar mass (kgmol\(^{-1}\))  
D: Diffusion coefficient (m\(^2\)s\(^{-1}\))  
c: Molar concentration (molm\(^{-3}\))  
M\(_{av}\): Average molar mass of the mixture (kgmol\(^{-1}\))  
σ: Conductivity (Sm\(^{-1}\))  
φ: Potential (V)  
i: Current density (Am\(^{-2}\))  
v: Stoichiometric coefficient  
n: Number of transfer electrons  
F: Faraday’s constant (96485 Cmol\(^{-1}\))  
i\(_0\): Exchange current density (Am\(^{-2}\))  
χ: Reaction order  
α: Charge transfer coefficient  
η: Activation overpotential (V)  
R: Gas constant (8.314 Jmol\(^{-1}\)K\(^{-1}\))
T: Temperature (K)
E_{eq}: Equilibrium potential (V)
Q: Charge source term (Am^{-3})

**Subscript:**
- a: Anode
- c: Cathode
- f: Fuel
- i, j, k: Species
- l: Electrolyte
- m: Main reaction
- o: Oxidant
- p: Parasitic reaction
- s: Electrode
- _0_: Standard, reference, or boundary value

**Superscript:**
x→1: Infinite solution

1. **Introduction**

Fuel cell is very promising as a future power source because of its distinct characteristics such as high energy efficiency, superior energy density, environmental friendliness, long-term energy storage and generation ability, and so on. Various applications of this technology are emerging in recent years, especially in the field of transportation and stationary power generation where high rated power is required. As for the market of small-power applications, the competitiveness of fuel cell is greatly impaired by its relatively high cost compared with other existing technologies such as batteries. To lower down the cost, a novel-type microfluidic fuel cell (MFC) has been proposed recently [1, 2], which utilizes the characteristics of laminar flow for fuel-oxidant separation purpose instead of the conventional solid membrane. As shown in Fig. 1(a), a typical MFC employs two laminar flow streams, one mixed with fuel (anolyte)
and the other mixed with oxidant (catholyte), to flow in parallel along the micro-channel. In this manner, the high-cost membrane in conventional fuel cells can be eliminated together with its dry-out and degradation issues [3, 4]. In addition, the reaction heat and water generated can be efficiently removed by the flowing electrolyte, leading to a much simplified balance of plant (BOP). As a consequence, the fabrication cost of MFC is much lowered, which makes it especially suitable as mini-watt power sources.

To date, there are plenty of MFC prototypes proposed in the literature, which have greatly contributed to the research and development of this technology. Most of these prototypes utilize the co-flow configuration as shown in Fig. 1(a), in which the diffusion direction of reactant crossover is orthogonal to the flow direction of electrolyte solution. This configuration has been the mainstream ever since the MFC technology was proposed, probably because of its simple structure and easy fabrication. During the cell operation, the convective electrolyte flow is much faster than the diffusive reactant crossover (that is, a high Peclet number), which can restrict the mixing zone of fuel and oxidant (as indicated by the white dash line in the figure) to a thin layer in the middle of the channel. In this manner, both the anode and cathode can work well so the co-flow MFC can generate very high power output with various types of fuels, some of which are even comparable to the PEMFC. Kjeang et al. [5] developed an all-vanadium co-flow MFC which can obtain a peak power density of 70 mWcm$^{-2}$ at room temperature. Lu et al. [6] employed the dual-electrolyte technique in their hydrogen-oxygen co-flow MFC with acid electrolyte for the cathode side and alkaline electrolyte for the anode side. A peak power density as high as 1.3 Wcm$^{-2}$ was achievable together with a high OCV of 1.89V. Hollinger et al. [7] proposed a direct methanol co-flow MFC which can generate a power output of 70 mWcm$^{-2}$ and a current output of about 650 mAcm$^{-2}$. As for the formic acid fuel, a peak power density of 55 mWcm$^{-2}$ was achieved by Jayashree et al. [8] using 1M formic acid and 50sccm oxygen. These high power outputs are mainly attributed to both the sufficient supply of reactant to the electrode surface and the effective suppression of reactant crossover, which generally requires a relatively high electrolyte flow rate. Nevertheless, the fuel utilization is consequently sacrificed due to the limited retention time of the fuel inside the channel. Moreover, a large amount of solution needs to be stored inside the co-flow MFC system, leading to a low energy density. These
shortcomings will greatly impair its application potential. In addition to the co-flow configuration, the counter-flow configuration as shown in Fig. 1(b) has also been proposed, in which the diffusion direction of fuel and oxidant crossover is opposite to the flow direction of catholyte and anolyte, respectively. This kind of MFC is less studied in the literature, although the fabrication process is as easy as its co-flow counterpart. When coupled with 3D flow-through electrodes, the counter-flow MFC can also achieve satisfactory power output. Ibrahim et al. [9] developed an all-vanadium counter-flow MFC and obtained a peak power density of 760 mWcm\(^{-2}\). Gurrola et al. [10] utilized 3M formic acid as fuel and the achieved peak power density was as high as 100 mWcm\(^{-2}\). As for the ethanol fuel, a peak power density of 99.4 mWcm\(^{-2}\) was generated by a dual-electrolyte counter-flow MFC developed by López-Rico et al. [11] Furthermore, the counter-flow MFC is considered to be highly tolerable to low electrolyte flow rates without the detrimental effect from aggravated reactant crossover. This is because that the convective electrolyte counter-flow can effectively prevent the diffusive reactant crossover from the electrode surface. As a consequence, high fuel utilization and low electrolyte consumption may be achievable by counter-flow MFCs. However, the cell power output may be sacrificed due to the insufficient reactant supply.

In addition to experimental approaches, numerical modeling is also an effective method for MFC study, which cannot only reveal the in-depth mechanisms behind the experimental phenomenon but also optimize the cell performance economically. A number of MFC modeling studies have already been published in the past decade, most of which focused on the co-flow configuration while only a few of them investigated the counter-flow configuration. For the co-flow MFC, great efforts have been made on the improvement of its fuel utilization by using various strategies, such as the increase of channel aspect ratio [12-14], the utilization of tapered electrodes [12] or very small electrodes [15], introduction of a third electrolyte [14, 16, 17], using multiple inlets [18], employing flow-through porous electrodes [19], using novel channel geometries [20], and so on. As for the counter-flow MFC, Xuan et al. [21] introduced micro-ridges into the microfluidic channel in order to generate chaotic flows, which can dramatically improve the cell current and power output. Xu et al. [22] investigated the possibility of counter-flow MFCs with low flow rate operation. With 1M formic acid as fuel, the highest fuel
utilization of 91.4% was achieved at 1 μlmin⁻¹. Other modeling studies focused on counter-flow MFCs with flow-through electrodes, which is especially suitable for the all-vanadium fuel cell [23-25].

Although there are already plenty of research works focused on the development of either co-flow or counter-flow MFCs, it is still unclear and questionable about which flow configuration is more suitable and promising for future MFC development. This issue cannot be addressed by the direct comparison between co-flow and counter-flow MFCs in the literature, since they were studied with different cell parameters and under different test conditions. Therefore, a comprehensive comparison of the merits and demerits between these two distinctive flow configurations under identical cell parameters and test conditions is very important for future MFC development, which has not been conducted yet to the best of our knowledge. In this study, we have conducted both experimental and numerical investigations on this issue. Two MFC prototypes, one with the co-flow and the other with the counter-flow configuration, have been designed and fabricated with similar cell parameters, and tested under identical conditions. The testing results obtained were further utilized to validate the corresponding numerical models. Based on the models, effects of electrode aspect ratio, channel thickness, and electrolyte flow rate on both cell performance and fuel utilization have been fully investigated. Meanwhile, the merits and demerits between the two different MFCs were also specifically compared with each other. Furthermore, several optimization strategies for the counter-flow MFC have been proposed and testified by modeling, which were found to successfully improve the cell performance of the original counter-flow prototype.
Fig. 1 Co-flow MFCs (left) and counter-flow MFCs (right): (a) (b) Schematic diagrams; (c) (d) Experimental prototypes; (e) (f) Computational domains.

2. Experimental

Two MFC prototypes, that is, the co-flow and the counter-flow cells, were designed and fabricated for the study. To achieve a cost-efficient fuel cell system, all cell components, except for the electrodes and current collectors, were made of plastic materials. As shown in Fig. 1(c), the co-flow MFC was mainly composed of five Poly(methyl methacrylate) (PMMA) layers, two GDEs, and two Cu foils. In its middle part, a 0.5mm-thick PMMA layer with half channel was sandwiched between two 1mm-thick PMMA layers with full channel (width: 10mm), which worked as a buffer layer to avoid vigorous mixing of the anolyte and catholyte when they first entered the cell. At their top, a 0.5mm-thick PMMA layer was used to seal the upper channel and to bear the anode; while at the bottom, another 0.5mm-thick PMMA layer was employed to seal the
bottom channel and to bear the cathode. Different methods were employed for fixing the electrodes. The anode (3mm x 10mm) was embedded into a groove (depth: 0.2mm) from the inside with its catalyst layer (CL) facing the channel, meanwhile a slit was cut out inside the groove to connect the anode with a Cu foil. As for the cathode (5mm x 12mm), it was simply pasted to the electrode window (3mm x 10mm) from outside the cell. This difference in fixing between anode and cathode was because of the air-breathing configuration of cathode, while the anode should be isolated from the ambient air. Fig. 1(d) illustrates the composition of the counter-flow MFC, which was mainly composed of three PMMA layers, two GDEs, and two copper foil current collectors. The micro-channel (width: 10mm) was cut out in the middle PMMA layer (thickness: 1mm). At the top, a 0.5mm-thick PMMA layer was employed to seal the channel, in which two inlets and two outlets were fabricated as electrolyte entrances and exits, respectively. At the bottom, another 0.5mm-thick PMMA layer was used to seal the channel as well as to bear the electrodes. The electrode fixing methods were exactly the same as those in the co-flow MFC as mentioned above. For both prototypes, double-layer adhesive tape was utilized to seal the adjacent PMMA layers, which was proved to be leakage-free and stable during the test, while plastic tubes were employed for electrolyte delivery.

The GDEs employed in this work were all commercial products. Pt–Ru Black (1:1 molar ratio, 10 mg cm\(^{-2}\), Johnson Matthey) and Pt/C (60wt.% Pt, 2 mg cm\(^{-2}\), Johnson Matthey) were chosen as anode and cathode catalysts, respectively, while carbon paper (HCP120, Hesen) with a microporous layer (MPL) served as the gas diffusion layer (GDL). The conductivity of the electrodes was reported to be 381.8 S m\(^{-1}\) according to the product specification. 0.5M sulfuric acid (H\(_2\)SO\(_4\), Sigma Aldrich) aqueous solution was utilized as electrolyte. For the catholyte, no oxidant additive was needed since air-breathing configuration was employed. As for the anolyte, ethanol (99.9% purity, Merck KGaA, Germany) fuel was added into the electrolyte with a concentration of 1M. All solutions were prepared by 18.2 M\(\Omega\) de-ionized water (Barnstead, NANOpure Diamond\textsuperscript{TM}). Table 1 summarizes the key cell parameters and test conditions, which were kept identical throughout the tests in order to make a fair comparison between the co-flow and the counter-flow MFCs.
A CHI660E electrochemical workstation (Shanghai Chenhua) was used to record the experimental results. When the OCV stabilized, cell polarization curve was obtained by measuring the potentiostatic currents at multiple potential steps from open-circuit to short-circuit. Sufficient time was given for current stabilization at each potential step. To obtain the single electrode potentials, a reference electrode (Ag/AgCl in saturated KCl, Shanghai Leici) was placed in the exit fluid and connected to the cathode through a multimeter (Fluke 116C).

Table 1 Comparison of the structural and operational parameters between the co-flow and the counter-flow MFCs

<table>
<thead>
<tr>
<th></th>
<th>Co-flow</th>
<th>Counter-flow</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrode area</td>
<td>3mm x 10mm</td>
<td>3mm x 10mm</td>
</tr>
<tr>
<td>Electrode distance</td>
<td>2.5mm</td>
<td>2.5mm</td>
</tr>
<tr>
<td>Anode catalyst</td>
<td>10 mgcm(^{-2}) Pt-Ru</td>
<td>10 mgcm(^{-2}) Pt-Ru</td>
</tr>
<tr>
<td>Cathode catalyst</td>
<td>2 mgcm(^{-2}) Pt/C</td>
<td>2 mgcm(^{-2}) Pt/C</td>
</tr>
<tr>
<td>Electrolyte</td>
<td>0.5M H(_2)SO(_4)</td>
<td>0.5M H(_2)SO(_4)</td>
</tr>
<tr>
<td>Fuel</td>
<td>1M ethanol</td>
<td>1M ethanol</td>
</tr>
<tr>
<td>Oxidant</td>
<td>Air-breathing</td>
<td>Air-breathing</td>
</tr>
<tr>
<td>Flow rate</td>
<td>300 μlmin(^{-1})</td>
<td>300 μlmin(^{-1})</td>
</tr>
</tbody>
</table>

3. Numerical model
3.1 Computational domain

The corresponding numerical models for both the co-flow and the counter-flow MFCs have also been developed in this work, in order to further study the differences between these two flow configurations and to optimize the cell performance, which adopted identical cell parameters and operational conditions as the previous experiment. As shown in Fig. 1(e) and (f), the computational domain in both models includes the electrolyte channel and two porous electrodes. In the co-flow model (Fig. 1(e)), the anolyte and the catholyte enter the cell from the same side but are initially separated by a buffer layer to avoid mixing. When they reach the electrodes, the buffer layer disappears and the mixing of fuel and oxidant begins, forming a mixing zone in the middle of the channel between the electrodes. Finally, the two streams flow out of the cell from a common outlet at the other end. As for the counter-flow MFC (Fig. 1(f)), the anolyte and the catholyte enter the cell from its two opposite sides and flow face-to-face passing the
corresponding electrodes, until they meet with each other in the middle of the channel. In addition, double outlets are employed as electrolyte exit. The electrode employed here includes both a gas diffusion layer (GDL) and a catalyst layer (CL), of which the latter is simplified to a 2-D plane next to the channel surface, considering the small thickness of CL (~20μm) compared with that of GDL (~200μm) in real cases.

During the previous experiment, no CO₂ bubbles were observed because of the relatively small current values. Therefore, gas bubble disturbance to the microfluidic flow is not considered in the present models. In addition, the gravity effect is also neglected considering the small size of the present models. According to Shaegh et al. [26], the oxygen concentration gradient through the thin GDL is negligible in MFCs. Therefore, the oxygen concentration at the cathode CL is assumed to be a constant, that is, the solubility of air dissolved in water at saturated level (0.25 molL⁻¹ [27, 28]).

3.2 Governing equations and boundary conditions

3.2.1 Hydrodynamics

The laminar flow in the micro-channel is governed by the continuity equation and Navier-Stokes equation:

\[ \nabla \cdot (\rho u) = 0 \quad (1) \]
\[ \rho(u \cdot \nabla u) = -\nabla p + \nabla \cdot (\mu(\nabla u)) \quad (2) \]

3.2.2 Mass transport

Mass transport inside the flowing electrolyte is governed by the conservation equation of species:

\[ \nabla \cdot (\rho \omega_i u) = -\nabla \cdot j_i + S_i \quad (3) \]

Maxwell–Stefan theory with a mixture-average approximation is utilized to describe the diffusion process of multiple components in water, including ethanol, oxygen and H₂SO₄.
\[ j_i = -(\rho D_i \nabla \omega_i + \rho \omega_i D_i \frac{\nu M_n}{M_n}) \quad (4) \]

\[ D_i = \frac{1 - \omega_i}{\sum_{k \neq i} \frac{x_k}{D_{ik}}} \quad (5) \]

\[ M_n = (\sum_i \frac{\omega_i}{M_i})^{-1} \quad (6) \]

Maxwell–Stefan diffusivities are calculated by the generalized Vignes equation [29]:

\[ D_{ij} = (D_{ij}^{x_i^{-1}})^x_i (D_{ij}^{x_j^{-1}})^x_j \prod_{k=1}^n x_{ik} (D_{ij}^{x_k^{-1}})^x_k \quad (7) \]

where “\( D_{ij}^{x_i^{-1}} \)” means the diffusivity of species “i” in species “j” at infinite dilution, which can be calculated by the Wilke-Chang Equation [30].

### 3.2.3 Electrochemical kinetics

Electrolyte potential is derived from the conservation equation of charge:

\[ -\sigma \nabla \phi = i \quad (8) \]

The species consumption or production rates due to electrochemical reaction are derived from:

\[ S_i = M_i \frac{\nu_{F_i} \nu_{P_i}}{n_i} \quad (9) \]

The current density is calculated by the Butler-Volmer equation. To consider the negative effect of species crossover, both the main reaction and the parasitic reaction are involved in the models [22, 31]. However, since the solubility of oxygen in water is rather low compared with the fuel concentration, it is reasonable to neglect the effect of oxygen crossover at the anode side [22]. Therefore, only the parasitic current of ethanol oxidation at the cathode side is considered here.
\[ i_a = i_{0,a} \left( \frac{c_f}{c_{0,f}} \right)^x \left( \exp \left( \frac{\alpha_a F \eta}{RT} \right) - \exp \left( -\frac{-\alpha_a F \eta}{RT} \right) \right) \] (10)

\[ i_{c,m} = i_{0,c} \left( \frac{c_a}{c_{0,c}} \right)^x \left( \exp \left( \frac{\alpha_a F \eta}{RT} \right) - \exp \left( -\frac{-\alpha_a F \eta}{RT} \right) \right) \] (11)

\[ i_{c,p} = i_{0,a} \left( \frac{c_f}{c_{0,f}} \right)^x \left( \exp \left( \frac{\alpha_a F \eta}{RT} \right) - \exp \left( -\frac{-\alpha_a F \eta}{RT} \right) \right) \] (12)

\[ i_c = i_{c,m} + i_{c,p} \] (13)

\[ \eta = \varphi_s - \varphi_l - E_{eq} \] (14)

At the electrode surfaces, current source Q is derived from:

\[ Q = \nabla \cdot i \] (15)

### 3.2.4 Boundary conditions

At the inlet and outlet of the micro-channel, velocity and pressure are set as follows, respectively:

\[ u_{inlet} = u_0 \] (16)

\[ P_{outlet} = 0 \] (17)

In addition, the non-slip condition is applied to the channel boundaries where the velocity is set to be zero. For the mass transport in the electrolyte, KOH and ethanol concentrations are set at the flow inlet and convective mass transport is set at the flow outlet. To simplify the cathode calculation, oxygen concentration at the cathode CL is set to be a constant as mentioned in Section 3.1. For the current distribution calculation, potential of the anode and cathode outer surfaces are set to be ground and cell voltage, respectively.

Key input parameters used in this study are summarized in Table 2. Considering the scarcity of the anode and cathode exchange current densities for the modeling of direct ethanol microfluidic fuel cells, these values are fitted from our own experimental results.

**Table 2** Key input parameters in the present MFC models
<table>
<thead>
<tr>
<th>Parameters</th>
<th>Values</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>$c_0$</td>
<td>0.25 molm$^{-3}$</td>
<td>[27, 28]</td>
</tr>
<tr>
<td>$c_{0,0}$</td>
<td>1.25 molm$^{-3}$</td>
<td>[27, 28]</td>
</tr>
<tr>
<td>$c_1$</td>
<td>1 molL$^{-1}$</td>
<td>-</td>
</tr>
<tr>
<td>$c_{0,f}$</td>
<td>0.1 molL$^{-1}$</td>
<td>-</td>
</tr>
<tr>
<td>$i_{0,a}$</td>
<td>0.002 Am$^{-2}$</td>
<td>Fitted from experiment</td>
</tr>
<tr>
<td>$i_{0,c}$</td>
<td>0.03 Am$^{-2}$</td>
<td>Fitted from experiment</td>
</tr>
<tr>
<td>$\alpha_a$</td>
<td>0.5</td>
<td>[32]</td>
</tr>
<tr>
<td>$\alpha_c$</td>
<td>0.5</td>
<td>[28, 33]</td>
</tr>
<tr>
<td>$\sigma_l$</td>
<td>20 Sm$^{-1}$</td>
<td>[26]</td>
</tr>
<tr>
<td>$\sigma_s$</td>
<td>381.8 Sm$^{-1}$</td>
<td>From the manufacturer</td>
</tr>
<tr>
<td>$E_{eq,a}$</td>
<td>0.084 V</td>
<td>[34]</td>
</tr>
<tr>
<td>$E_{eq,c}$</td>
<td>1.229 V</td>
<td>[34]</td>
</tr>
</tbody>
</table>

### 3.3 Numerical solution

A commercial FEM software, COMSOL$^\circledR$ 4.2, was employed in the present study. To deal with the above-mentioned processes (that is, hydrodynamics, mass transport and electrochemical kinetics), three physical interfaces were utilized correspondingly in the present model, including the Laminar Flow, the Transport of Concentrated Species, and the Secondary Current Distribution. To discretize the computational domain, structured grid with the number of cells ranging from 1420 to 101100 was used, with the corresponding computational time increased from 1min to 2.6h. Before commencing the study, grid independence check was conducted to ensure the accuracy of modeling results, and the variation of cell polarization curves were exhibited in Fig. 2. It was found that the relative difference of current densities under different cell voltages were all negligible when the grid number exceeded 8450. For instance, the relative difference of the short-circuit current density between the 8450 grid and the 101100 grid was less than 0.1%. Therefore, the grid with 8450 cells was selected for subsequent studies. To obtain the polarization curves, a parametric sweep solver based on cell voltage was utilized to calculate the current densities from 0.8V to 0V with an interval of 0.05V. The stationary segregated multifrontal massively parallel solver (MUMPS) was used for the calculation process.
4. Results and discussion

4.1 Experimental study and comparison

An effective control of the extent of fuel crossover is of paramount importance to the functionality of MFCs. Therefore, before the cell performance test, the mixing zone of the two prototypes were characterized first by using an alkaline and a phenolphthalein solution dissolved in ethanol, in order to determine a suitable electrolyte flow rate for the test. As shown in Fig. 3(a), the co-flow MFC encountered a disastrous mixing zone covering the whole area of its electrode surface at lower flow rates of 50-100 μlmin⁻¹, indicating a severe crossover contamination. This situation was gradually improved as the flow rate rose to 300-500 μlmin⁻¹, with the electrode surface safely away from the mixing zone. As for the counter-flow MFC, the mixing zone was well controlled in the middle of the channel and away from electrodes even at the lowest flow rate of 50 μlmin⁻¹. Consequently, 300 μlmin⁻¹ was chosen for the testing of both cells in order to make a fair comparison. Fig. 3(b) and (c) show the cell polarization curves and single electrode polarization curves of the co-flow and the counter-flow prototypes. In general, the co-flow MFC exhibited a much better cell performance than the counter-flow MFC, whose peak power density (that is, 2.9mWcm⁻²) was nearly two times as high as that of the latter (that is, 1.6mWcm⁻²). This huge difference is probably attributed to their different cell ohmic resistances as can be noticed from the different slope (absolute value) of the linear part of the polarization curves. Based on the slope, the ohmic resistances of the co-flow and counter-flow MFCs were found to be 9.7 and 18.2 Ωcm², respectively. Since all the
cell parameters and test conditions were kept identical, this difference in ohmic resistance should only be related to the different electrode configurations between them. The counter-flow MFC employs electrodes in the same plane which may lead to a higher ionic transport resistance especially for the outer part of the electrodes, while the co-flow MFC employs face-to-face electrodes which results in a uniform ionic resistance throughout the electrode surface. Nevertheless, in order to prove the above assumption, numerical study is requisite to further explore the differences and merit-demerit between the co-flow and the counter-flow MFCs.

![Fig. 3 Experimental study of the co-flow and the counter-flow prototypes: (a) Exhibition of the mixing zone; (b) Cell polarization curves; (c) Single electrode polarization curves.](image)

**4.2 Numerical investigation and comparison**

**4.2.1 Model validation and current density distribution**

Before any further computational studies, reliability of the present models was verified first by comparing the simulated polarization curves with the above experimental data. As shown in Fig. 4(a), the experimental and modeling results fit very well with each other for both the co-flow MFC and the counter-flow MFC, indicating a high reliability.
of the present models for simulating direct ethanol MFCs with acidic electrolyte. The modeling results also reveal the performance superiority of co-flow MFC compared with counter-flow MFC, while the reason behind is elucidated as follows. In addition, it is worth mentioning that since the main purpose of the present study is to compare the co-flow and the counter-flow MFC, their cell structures and operational conditions have not been optimized, leading to an inferior cell performance compared with other direct ethanol MFCs in the literature [11, 35, 36]. However, according to our previous experience this situation can be effectively improved by, say, using alkaline electrolyte with high-concentration, reducing the electrode distance, increasing the fuel concentration, and so on [37].

Both the above experimental and modeling results show that the co-flow MFC achieves better performance than the counter-flow MFC. To inspect the possible reason behind this difference, the distribution of current density at 0V on the electrode surface of the two cases were compared. As shown in Fig. 4(b), the distribution of current density of the co-flow case is quite uniform throughout its anode and cathode, which is within the range of 49.5-62.1 mAcm$^{-2}$ with an average of 53.5 mAcm$^{-2}$. Besides, since the current directions of the anode and the cathode are identical, both of them obtain positive current densities as shown in the figure. As for the counter-flow cell, since the current direction of anode and cathode are opposite to each other, only the color variation on anode is exhibited here by setting the color range from 0 to maximum, while the negative current densities on cathode is shown uniformly as dark blue in color (Fig. 4(c)). This action is taken to better visualize the current density distribution on counter-flow cell electrodes. From the figure, it is obvious that higher current densities are located at the inner part of the anode, while the outer part generates much lower current densities. The calculated range is 17.4 - 36.6 mAcm$^{-2}$ with an average of 22.5 mAcm$^{-2}$. As for its cathode, the corresponding range is (-17.4) - (-36.6) mAcm$^{-2}$ with an average of -22.5 mAcm$^{-2}$, which are exactly the opposite to the anode side.

Fig. 4(b) and (c) give an intuitional view of the difference between the co-flow and the counter-flow MFCs. To make a further quantitative comparison, the relative dispersion degree of current density is calculated via dividing the range by the average value, that is, 23.6% for the co-flow MFC and 85.3% for the counter-flow MFC. This huge difference
in current distribution should be the main reason behind their performance difference. In the co-flow MFC, since the electrodes are placed face-to-face, the electrode distance at every point is the same (2.5mm), leading to a uniform ionic resistance throughout the electrode surface. As for the counter-flow MFC, the electrodes are placed in the same plane, so the inner part of it has a shorter ionic diffusion path while the outer part has a much longer one. As a consequence, the outer part of electrode suffers from greater ionic resistance and poorer electricity generation ability. This drawback of counter-flow MFC is determined inherently by its special flow configuration, which can be improved by structural optimization as will be discussed in Section 4.3 below. To support the reliability of the optimization results, a more extended validation of the present counter-flow MFC model in terms of electrolyte flow rate has been further conducted as shown in Fig. 4(d1-d3). The simulation results can still fit the experimental data very well. In addition, it is worth mentioning that the difference in cell performance under the three different electrolyte flow rates is very limited. This phenomenon will be further investigated and elaborated in Section 4.2.4.
Fig. 4 Preliminary modelling results: (a) Model validation (1M ethanol as fuel, 0.5M \( \text{H}_2\text{SO}_4 \) as electrolyte, at the flow rate of 300\( \mu \text{l}\text{min}^{-1} \), under room temperature); (b) Current density distribution of the co-flow cell; (c) Current density distribution of the counter-flow cell; (d1-d3) Further validation of the counter-flow model under various electrolyte flow rates.

### 4.2.2 Effect of electrode aspect ratio

In this study electrode aspect ratio (AR) is defined as the ratio between the electrode length along the flow direction and the electrode width orthogonal to the flow direction. In the literature, long and narrow electrodes, that is, a high aspect ratio, were commonly adopted in both co-flow [5, 38, 39] and counter-flow MFCs [21, 22, 24], in order to obtain high fuel utilization. However, according to the previous section, such kind of electrodes may no longer be suitable for counter-flow MFCs as far as their cell performance is concern. To verify this conclusion, another two cases with a higher AR of
3.33 (10 mm length and 3 mm width) were studied and compared with the previous cases with a lower AR of 0.3 (3mm length and 10mm width). As shown in Fig. 5(a) and (b), the electrode AR has a negligible effect on the co-flow MFC, but can greatly impair the counter-flow MFC performance. The maximum current density drops dramatically from 22.5 to 8.0 mAcm$^{-2}$. Fig. 5(c) and (d) further exhibit the current density distribution in the two cases with a larger AR. Similarly, the distribution is still uniform for the co-flow MFC (25.7%) but is more polarized for the counter-flow MFC (352.1%). In conclusion, high electrode AR is suitable for the co-flow MFC but very disadvantageous for the counter-flow MFC, leaving the outer part of the electrodes greatly wasted. To solve this problem, either a low electrode AR or a partial modification of the electrodes [24] should be implemented for counter-flow MFCs. Nevertheless, it is also worth mentioning that if the electrode AR is too low, that is, the electrode width is extremely large, then the electrolyte flow may no longer be stable and uniform along the channel in practical operation. Therefore, a moderate value of 0.3 is still selected for the subsequent studies unless otherwise specified.

**Fig. 5** Effect of electrode aspect ratio on cell performance: (a) Polarization curve comparison of the co-flow MFCs; (b) Polarization curve comparison of the counter-flow
MFCs; (c) Current density distribution on the electrodes of the co-flow MFC; (d) Current density distribution on the electrodes of the counter-flow MFC.

4.2.3 Effect of channel thickness

A reduced channel thickness generally leads to a higher electrolyte flow speed under a fixed electrolyte volumetric flow rate, which is beneficial to both the crossover suppression in the channel middle and the reactant replenishment to the electrode surface. However, this may also bring different problems to different kinds of MFCs, which is studied in this section. As shown in Fig. 6(a), for the co-flow MFC the reduction of channel thickness from 2.5mm to 1mm leads to an improved maximum current density from 53.5 to 68.3 mA cm\(^{-2}\), which is mainly attributed to the shortened ionic transport path and, therefore, reduced ionic resistance. However, when the thickness further decreases to 0.5mm the initial part of the polarization curve drops dramatically. This is probably due to the intensified fuel crossover at the cathode side. As shown in Fig. 6(c), the mixing of anolyte and catholyte is more and more severe as the channel thickness decreases. Therefore, a channel thickness of 1mm is appropriate for the current co-flow MFC under 300 μl min\(^{-1}\). As for the counter-flow MFC, since the electrode distance does not vary with the channel thickness, a reduced channel thickness helps to better shrink the mixing zone and suppress fuel crossover as shown in Fig. 6(d). However, the cell performance is greatly impaired as shown in Fig. 6(b), with the maximum current density decreases from 22.5 to 5.2 mA cm\(^{-2}\). This is probably due to the elevated ionic resistance between the electrodes. The reduction of the channel cross sectional area may have impeded the proton transport from the anode to the cathode. Therefore, a channel thickness of 1mm is appropriate for the present counter-flow MFC. Moreover, when the channel thickness is further increased to 2mm, the maximum current density can be increased from 22.5 to 30.6 mA cm\(^{-2}\) (not shown in the figure). However, such a thick channel may bring several side effects during practical operation, such as flow instability and vigorous mixing especially at lower electrolyte flow rates. Therefore, 1mm thickness is still employed for the following studies unless otherwise specified.
Fig. 6 Different responses of co-flow and counter-flow MFCs to channel thickness: (a) Performance variation of co-flow MFC; (b) Performance variation of counter-flow MFC; (c) Mixing zone variation of co-flow MFC; (d) Mixing zone variation of counter-flow MFC.

4.2.4 Effect of electrolyte flow rate

Even though the counter-flow MFC achieves poorer cell performance than the co-flow MFC, it may be advantageous as far as fuel utilization is concern. Fuel utilization is closely related to the electrolyte flow rate in most MFCs, which favors lower rates in order to extend the fuel retention time inside the cell. However, a lower limit of the flow rate is requisite to prevent the fuel crossover problem. In the counter-flow MFC, since the direction of diffusive crossover is opposite to the direction of convective electrolyte flow, the lower limit of flow rate could be extremely small, leading to a much improved fuel utilization. Fig. 7(a) compares the crossovered fuel concentration at cathode CL under various flow rates between co-flow and counter-flow MFCs. The co-flow cell encounters a dramatically aggravated fuel crossover when the flow rate decreases to around 60
μlmin⁻¹, while the counter-flow MFC can tolerate a flow rate as low as 5 μlmin⁻¹. At an extremely low flow rate of 1 μlmin⁻¹, the crossovered fuel concentration at cathode CL is as high as 370 mM for the co-flow case, while the value can still be maintained less than 40 mM for the counter-flow case. Fig. 7(b) shows the fuel concentration distribution under four different flow rates. It is apparent that the mixing is more drastic in the co-flow channel than in the counter-flow one especially at lower flow rates. Furthermore, Fig. 7(c) and (d) exhibit the degradation of cell performance with the decrease of flow rate for the co-flow and the counter-flow cells, respectively. The former case encounters a much greater degradation in both its OCV and maximum current density due to the increasing severity of the fuel crossover, while the latter case achieves a fairly uniform cell performance from 300 to 5 μlmin⁻¹. Even at 1 μlmin⁻¹, the counter-flow cell can still function well which retains 90% of its maximum current density. The fuel utilization is calculated based on the following equation:

$$\xi_{fuel} = \frac{I_{max}}{nFv_c} \tag{18}$$

where $I_{max}$, n, F, v, c represent, respectively, the maximum current, the number of electron transferred in the reaction, the Faraday’s constant, the volumetric electrolyte flow rate, and the fuel concentration. If the crossovered fuel concentration at cathode CL is stipulated to be less than 1% of the fuel concentration (that is, 10 mM), then the minimum flow rate for the co-flow and the counter-flow cells are 30 and 2 μlmin⁻¹, and the corresponding fuel utilizations are calculated to be 2.7% and 16.6%, respectively. It is noteworthy that these moderate values can be further improved by using high-concentration alkaline electrolytes instead of the present low-concentration acid electrolyte. In conclusion, the counter-flow MFC can adapt a much lower electrolyte flow rate to maintain its normal operation, leading to a much improved fuel utilization and reduced parasitic energy loss. In addition, electrolyte storage in the MFC system is also greatly alleviated. All these merits make counter-flow MFCs more suitable for practical applications than its co-flow counterpart, which could be even more advantageous if their cell performance can be further optimized, as will be discussed in Section 4.3 below.
Fig. 7 Different sensitivity of co-flow and counter-flow MFCs to electrolyte flow rate: (a) Fuel concentration at cathode CL; (b) Extent of fuel crossover under four different flow rates; (c) Co-flow MFC performances under four different flow rates; (d) Counter-flow MFC performances under four different flow rates.

4.3 Optimization of the counter-flow MFC

4.3.1 Effect of electrode distance

Since the ionic resistance between the two electrodes is the most significant factor that restricts the counter-flow cell performance, it is straightforward to move its electrodes closer to each other in order to promote the ionic conduction. A series of different electrode distances from 2.5 to 0.5 mm have been studied under an electrolyte flow rate of 300 μl/min with their cell performances compared. As shown in Fig. 8(a), when the electrode distance decreases from 2.5mm to 1mm, the cell performance is gradually improved with the maximum current density increased from 22.5 to 28.9mA cm⁻². However, when the distance further decreases to only 0.5mm, even though there is a
great improvement to the maximum current density (33.3 mAcm⁻²), the initial performance at high cell voltages is slightly dragged down probably by the aggravated fuel crossover. To verify this, concentration of the crossovered fuel at the cathode CL was further inspected as shown in Fig. 8(b), which indicates that when the electrode distance is not less than 1mm the concentration of crossed over ethanol is close to zero. However, when the electrode distance is further reduced to 0.5mm, this value quickly rises up to 0.65mM. This may accounts for the initial drop of its polarization curve at high cell voltages. Therefore, an electrode distance of 1mm is selected for subsequent studies unless otherwise specified.

Fig. 8 Effect of electrode distance on counter-flow MFC performance (under 300μlmin⁻¹): (a) Polarization curves; (b) Concentration of crossovered ethanol at cathode CL.

4.3.2 Effect of electrode area

According to Section 4.2.1, the outer part of the electrode in a counter-flow MFC contributes much less to current generation than its inner part. Therefore, it can be beneficial to the current density by removing this “sluggish” part of electrode away from the cell. This section studies the polarization curve variation with different electrode areas from 0.3 to 0.1 cm² under both a high flow rate (300 μlmin⁻¹) and a low flow rate (1 μlmin⁻¹) condition. As shown in Fig. 9(a), the cell performance gradually increases with the decrease of electrode area at 300 μlmin⁻¹, which finally catches up with the co-flow cell. Maximum current density is improved from 28.9 to 54.0 mAcm⁻², which is almost
doubled. Fig. 9(b) shows that when the flow rate is decreased to 1 μlmin⁻¹, the optimized counter-flow MFC with an electrode area of 0.1 cm² achieves an even better performance than its co-flow counterpart, which is mainly attributed to its better suppression on fuel crossover as mentioned in Section 4.2.4. In conclusion, smaller electrodes located close to the flow interface are much favored in counter-flow MFCs, which can achieve comparable performance at higher electrolyte flow rates and even better performance at lower flow rates than its co-flow counterpart. Nevertheless, the absolute power output is inevitably reduced by the decreased electrode area, so the fuel utilization is greatly sacrificed, dropping from 37.2% to 21.3% when the electrode area decreases from 0.3 to 0.1 cm² under an electrolyte flow rate of 1 μlmin⁻¹. This fuel utilization loss leads to less superiority of the counter-flow cell compared with the co-flow cell. To tackle this dilemma, the 3D flow-through electrodes [40, 41] should be implemented instead of the present 2D flow-over electrodes, which can better utilize the “effective zone” (that is, the zone near the flow interface) in counter-flow MFCs. This will be studied in future to further improve both the cell performance and fuel utilization of counter-flow MFCs.

![Graphs showing cell voltage vs. current density for different electrode areas under different flow rates](image)

**Fig. 9** Effect of electrode area on counter-flow MFC performance: (a) Under 300 μlmin⁻¹; (b) Under 1 μlmin⁻¹.

### 4.3.3 Effect of outlet design

In conventional counter-flow MFCs, a “T” shape channel with two inlets and single outlet is commonly employed [22, 24]. As for the counter-flow prototype in the present
study, double outlets are utilized instead, which is proved to be better in crossover control especially under lower electrolyte flow rates. Fig. 10(a) shows the fuel concentration isoline in the middle of the channel at an electrolyte flow rate of 1 μlmin⁻¹, including both the single outlet case (left) and the double outlets case (right). The mixing zone in the double outlets case is a spindle shape, which is wider in the middle and narrower in the two ends near the outlets. As for the single outlet case, the mixing zone is convergent in one end near the outlet, and divergent in the other end away from the outlet. In general, the mixing zone of the double outlets case is more confined to the middle of the channel, leading to a better control of fuel crossover. Moreover, the crossovered fuel concentration at the cathode CL is calculated to be 182 mM for the double outlets case, which is lower than 245 mM for the single outlet case. Fig. 10(b) further compares the cell performance between a single outlet cell and a double outlets cell under an electrolyte flow rate of 1μlmin⁻¹, of which the latter achieves better cell performance than the former. Therefore, it is concluded that the double outlets configuration is more suitable for counter-flow MFCs especially operated under low electrolyte flow rate mode.

**Fig. 10** Effect of outlet number on counter-flow MFC performance: (a) Distribution of the fuel concentration (isoline under 1 μlmin⁻¹); (b) Polarization curves (under 1 μlmin⁻¹).

### 4.3.4 A counter-flow MFC prototype with optimized cell performance

The above-mentioned optimization strategies were further verified experimentally by implementing them into the original counter-flow prototype and comparing with its co-
flow counterpart, which were tested under the same conditions. As shown in Fig. 11, the optimized counter-flow MFC did achieve a much better cell performance than its original state and was comparable with the co-flow MFC, indicating the effectiveness of the optimization strategies and the reliability of the model. From the absolute slope of the polarization curve, the calculated ohmic resistance of the optimized counter-flow cell was 9.5 Ωcm², which was similar to the value of the co-flow case (9.7 Ωcm²). Nevertheless, the optimized counter-flow MFC provided a lower absolute power output than the co-flow MFC due to the reduction of electrode area. This issue will be addressed by utilizing the 3D flow-through electrode in the future, which can also benefit its fuel utilization.

Even though the power output as shown in Fig. 11 is relatively low, it is mainly due to the un-optimized cell structure and operation conditions, which is not the focus of the current study. Nevertheless, in the literature MFCs have already achieved a satisfactory power output ranging from several mWcm⁻² [42] to several hundred mWcm⁻² [43], some of which are even comparable to those of high-temperature PEMFCs [6]. Therefore, the application prospect of MFCs is very promising considering their cost advantage and easy fabrication. By conducting a comprehensive comparison, the present study has indicated that the counter-flow MFC may be a more promising choice for practical applications compared with the conventional co-flow type. By utilizing a counter-flow configuration, small electrolyte consumption rate and high fuel utilization are achievable at the same time, which greatly improve its energy density and system practicability. In addition, the inherent drawback on cell performance of the counter-flow MFC can be effectively addressed by the proposed optimization strategies in this study. In conclusion, a high-performance counter-flow MFC system with economic electrolyte supply and limited fuel wastage can be anticipated, which is very close to real applications such as powering portable electronic devices. Nevertheless, before the commercialization of this technology, several technical issues still need to be properly addressed, such as potential electrolyte leakage, efficient micro-pump, cell output stability in mobile working environment, etc.

Furthermore, it is noteworthy that although the above results and discussion are obtained from a single cell-level study, they are also applicable to MFC stacks. To realize a sufficient voltage and power output, multiple MFC single cells need to be unitized into
a MFC stack, which will consume much more electrolyte solution than the single cell. Therefore, it is believed that a MFC stack with counter-flow configuration will be more advantageous as a novel type of mini-watt power source than its co-flow counterpart. However, other related issues such as shunt current loss and uniform electrolyte distribution still need to be carefully tackled before the commercialization of this technology.

Fig. 11 Cell performance of the optimized counter-flow MFC (1M ethanol as fuel, 0.5M H₂SO₄ as electrolyte, at the flow rate of 300μlmin⁻¹, under room temperature)

5. Conclusion
In this study, both experimental and numerical investigations have been conducted to compare the co-flow MFC and the counter-flow MFC with identical cell parameters, in order to gain further understanding of this microfluidic power generation technology. Before developing the models, experimental study has been conducted first to obtain the cell performances as benchmarks for the model validation. The developed models successfully coupled the microfluidic hydrodynamics, species transport in the flowing electrolyte, and electrochemical reaction kinetics on the electrodes’ surface for predicting the MFC performance, which also considered the effect of fuel crossover under low flow rate operation mode. Primary conclusions from this study are summarized as follow:
• The counter-flow MFC is found to suffer an inherent disadvantage on its power output ability, whose peak power density is only half of that of the co-flow MFC. This is mainly attributed to the uneven ionic resistance across its electrode surface. Therefore, shorter and wider electrodes (that is, smaller aspect ratio) are more suitable for the counter-flow cell rather than longer and narrower electrodes (that is, higher aspect ratio).

• The counter-flow MFC achieves a much higher tolerance to extremely low flow rates than the co-flow MFC. When the concentration of crossovered fuel at cathode surface is stipulated to be less than 10 mM, the counter-flow cell can utilize a 15 times lower electrolyte flow rate and obtain a 6 times higher fuel utilization. This superiority in fuel utilization and energy density makes the counter-flow cell more competitive for practical applications, which is attributed to its special flow configuration in which the diffusive fuel crossover is impeded by the convective electrolyte flow.

• The inferior cell performance of the counter-flow cell can be optimized by employing narrower electrode distance, smaller electrode area close to the flow interface, and double outlets. With these strategies, the optimized counter-flow prototype achieves the same level power output with its co-flow counterpart. To further improve its power output, the 3D flow-through electrodes should be adopted to better utilize the effective zone inside the counter-flow MFC.

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References


