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Broader context

Despite the greater role of methane in the global energy blueprint, many methane sources, both natural and artificial, have been untapped due to their low methane concentrations (typically <20%) and correspondingly low heating values. Generally known as low calorific value gases (LCVGs), they are not useful unless the CH₄ concentration can be increased, which requires a significant cost even with state-of-the-art technologies. Therefore, efficient utilization of LCVGs constitute significant economic and environmental challenges. In this study, we demonstrated that a ceramic fuel cell with commercially available materials could yield promising performances (>2 W cm⁻²) even with 10–20% methane in the fuel. The new 'micro-monolithic' conceptual design has displayed not only the advantages in minimizing concentration polarization, but also a unique mechanics in suppressing coking with >500 h stability test conducted. Therefore, with the depletion of existing fossil fuels and the urgency to mitigate climate change, this work demonstrates a promising technology for both efficient and environmentally benign use of LCVGs.

Introduction

Ceramic fuel cells offer the potential of environmentally sustainable and high-efficiency (typically >60%) power generation

High-performance fuel cell designed for cokingresistance and efficient conversion of waste methane to electrical energy[†]

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Utilization or emission of low calorific value gases (LCVGs) containing <20% CH₄ constitute economic and environmental challenges. Ceramic fuel cells offer a possible solution, but their performance is hindered by carbon formation ('coking'). Herein, we report a novel fuel cell designed to mitigate coking, yielding superior performances but using conventional commercially-available materials. The new micro-monolithic design has an extraordinary geometrical asymmetry that separates the mechanical support and anode current collector from the electrochemically active region and results in significantly facilitated mass transport, yielding power densities of 1.77–2.22 W cm⁻² using LCVGs. In addition, the effluent containing only H₂, CO and CO₂ is of great industrial interest for methanol synthesis, if their ratios are adjusted appropriately. The new fuel cell developed was almost free from coke deactivation and was stable for over 500 h, indicating great promise for both efficient and environmentally benign use of LCVGs.

compared to natural gas combustion-based electrical power plants, as they are not subject to constraints of the Carnot cycle. Moreover, they provide the critical advantage of fuel versatility over other types of fuel cells and enable fuelling by hydrocarbons *via* internal reforming, due to their high operating temperatures (700–1000 °C). Among the various types of hydrocarbons, natural gas, consisting primarily of methane, presently meets *ca.* 27% of global energy demand,^{1,2} and is preferred to coal due to its significantly lower carbon footprint.

Despite the great importance of methane in the global energy blueprint, many methane sources have been untapped due to their comparatively low methane concentrations, and correspondingly low heating values, so are known as low calorific value gases (LCVGs). There is no rigid LCVG definition as it was associated initially with the products of coal



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gasification, with heating values below 7 MJ m^{-3} .^{3,4} There are other sources of LCVGs, e.g. a significant number of natural gas reservoirs with CO₂ contents up to 87% across the world that remain unexplored, despite trillions of cubic metres of recoverable reserves, due to economic concerns.⁵⁻⁸ Such gas is not useful unless the CH₄ concentration can be increased, which typically requires a significant cost, even with state-of-the-art technologies. Another important source of LCVGs is from the upgrading procedures necessary to obtain marketable biomethane from biogas, produced from domestic and agriculture wastes. Though an important source of renewable energy, they could cause tremendous amount of off-gas emissions of 10–15% methane, the balance being CO₂.^{9,10} Methane in these LCVGs is uneconomic to use, and would contribute to climate change, since methane is one of the main greenhouse gases with a global warming potential (GWP) 25-34 times higher than that of CO₂.^{11,12} Presently, these LCVGs commonly are treated by flaring and in some cases, it is necessary to have further addition of fuel to sustain the combustion, which has economic and environmental implications, *i.e.* greater CO₂ emissions. Therefore, with the depletion of existing fossil fuels and the urgency to mitigate climate change, a technology that can achieve clean and efficient use of these low utility LCVGs is in great need.

As examples of a well-studied technology, ceramic fuel cells are promising candidates to utilize LCVGs *via* internal dry reforming of methane (DRM) reaction (1),¹²⁻¹⁴ as the high operating temperatures favour high equilibrium conversion (endothermic reaction, $\Delta H_{298K} = 247$ kJ mol⁻¹). DRM reaction converts the two greenhouse gases into a mixture of H₂ and CO (syngas), which is the fundamental feedstock for valuable chemicals *via* processes such as the Fischer–Tropsch synthesis,¹⁵⁻¹⁸ making the whole concept promising for CO₂ sequestration to mitigate global warming.

$$CH_{4(g)} + CO_{2(g)} \rightarrow 2CO_{(g)} + 2H_{2(g)}$$
(1)

One major challenge for hydrocarbon-fuelled ceramic fuel cell is carbon formation or coking, the mechanism of which depends on both thermodynamic and kinetic considerations. The carbon formed mainly *via* reactions (2)–(4), not only result in the de-activation of the anode by occupying active sites, but also lead to physical blockage of percolated pore structures, impeding gas transport:

Gas phase pyrolysis: $CH_{4(g)} + C_{(s)} \rightarrow 2H_{2(g)}$ (2)

Boudouard reaction: $2CO_{(g)} \rightarrow + CO_{2(g)} + C_{(s)}$ (3)

Carbon gasification: $CO_{(g)} + H_{2(g)} \rightarrow H_2O_{(g)} + C_{(s)}$ (4)

Despite the system complexity due to the multiple reactions involved, the thermochemistry of the gas phase equilibria, depicted by a C-H-O ternary diagram, is well established and accepted as a useful indication of the thermodynamic driving force for coking.^{12,19} As for coking mitigation, one major strategy is to replace conventional Ni-cermet with alternative materials that do not catalyze carbon formation, *e.g.* Cu-based electrode and some perovskites.²⁰ The other main approach involves operating Ni-based electrode under well-controlled conditions by addition of reforming agents or even oxygen into hydrocarbon fuels to adjust the C:H:O ratio and so suppress coking.²¹ Despite much research focused on methane conversion using solid oxide fuel cells,¹² state-of-the-art ceramic fuel cells cannot be used for efficient LCVG recovery, because of their cell designs, operating costs, lack of coking resistance and electrochemical performance.

Hence, a new strategy was developed to achieve high utilization efficiency of LCVG methane, with improved coking resistance and long-term stability. Fabricated from commercially available materials, the ceramic fuel cell had an evolved 'micro-monolithic' geometry with droplet-shaped channels. These exhibited a new gas transport configuration than conventional designs by offering a unique feature of geometrical 'asymmetry', *i.e.* the electrochemically active region (EAR) was deliberately separated spatially from the electronically conducting mechanical support, dramatically shortening the diffusional length for gaseous fuels to reach the EAR and enabling enhanced control over location where carbon deposition is favourable. This design helps to deliver high electrochemical performance operating with methane concentrations as low as 10%, by minimizing concentration polarization and creating an area-specific atmosphere that suppressed coking in the EAR.

X-ray computed tomography (X-ray CT) was used to reconstruct the three-dimensional structure, the results of which were used for CFD modelling to confirm the atmosphere variation across the cell. It was established that coking occurs only on the inert current collector/mechanical support at the fuel inlet (upstream) region, which had negligible influence on the structural integrity and electrochemical performances. Even with a $1:1 \text{ CH}_4$ -CO₂ feed ratio, predicted thermodynamically to favour coking at <800 °C, the fuel cell operated with negligible performance degradation for >500 h.

Results and discussion

Morphology

Fig. 1a shows the cross-section of the micro-monolithic anode support incorporating 6 sub-channels with droplet shapes. As described in the 'Method' section, the fabrication was based on the phase inversion process, which involves the precipitation of a polymer binder phase as a result of solvent/coagulant exchange. By controlling the rate of precipitation and utilizing the hydraulic pressure of the internal coagulant, the shape of the sub-channel was tailored, and converted from the circular shape of conventional designs, to a droplet shape predicted to achieve increased electrochemical performance. The micromonolithic anode had an outer diameter of 2.2 \pm 0.1 mm, so can be considered as a micro-tube. Such successful miniaturization is essential for high surface areas per volume and consequently improved volumetric power densities. As illustrated in Fig. 1b, the thickness of the exterior electrochemical active region (EAR) in the anode was decreased to $<50 \mu m$, which is



Fig. 1 SEM photomicrographs of 6-channel micro-monolithic anode supports, after sintering at 1400 °C. (a) Central-axis-view; cross-sectional images of (b) exterior electrochemical active region (ERA) between the shell and two channels in the 6-channel anode, (c) porous region between the two channels and outer surface, (d) central porous region (shaded area denotes mechanical support/current collector); (e) schematic diagram of a complete single cell and compositions for each component.

close to the well-accepted thickness of EARs of 20-40 µm.^{22,23} Moreover, the design of the droplet shape ensured a similar degree of curvature in between the interior and exterior surfaces of the region, as highlighted in Fig. 1b. Therefore, compared to micro-monolithic designs reported elsewhere, the electrochemically active region of the cell in Fig. 1 had a higher ratio to the overall circumference, so achieving more efficient utilization of the geometric surface area. As for the mechanical support, it was deliberately distributed in between neighbouring EARs (Fig. 1c), as well as the hexagonal region in the middle (Fig. 1d). In most conventional ceramic fuel cell designs, the electrochemically active region in the anode typically was sandwiched between the support and the electrolyte. Hence, gaseous fuels had to be transported through the support prior to participating in the electrochemical reactions occurring in the active zone, introducing additional mass transport resistance, whereas such additional resistance was eliminated in this new micromonolithic anode design. In addition, a plurality of microchannels was introduced during the fabrication by the phase inversion process and well-preserved after sintering, as depicted in Fig. S1 (ESI[†]). Fig. 2 illustrates the macroscopic morphology of the micro-monolithic anode imaged and reconstructed using micro-CT. The micro-channels were segmented using grayscale-based



Fig. 2 (a) 3D reconstruction of micro-monolithic anode by X-ray computed tomography (X-CT); (b) 3D visualization of the distribution of microchannels. The micro-channels are highlighted in red and the solid layer is displayed in light blue (semi-transparent).

thresholding method and are visualised in Fig. 2b (red phase), with the anode body rendered semi-transparent. The presence of such micro-channels further facilitates gas transport, which has been proved previously to lead to significantly decreased concentration polarization.^{24,25}

After the preparation of the anode substrate, both yttriastabilized zirconia (YSZ) electrolyte and gadolinium-doped ceria (GDC) were deposited by dip-coating (Fig. 1e), each having a thickness of 4.5 (\pm 0.5) µm; the YSZ electrolyte was highly dense after the co-sintering at 1400 °C (Fig. S3, ESI†). The porous GDC-LSCF cathode was dip-coated to complete a single cell and had a thickness of 25–30 µm after sintering at 1000 °C for 2 hours. Both electrodes were wrapped with silver wires (0.2 mm diameter) as current collector.

Electrochemical performance with hydrogen

Firstly, the micro-monolithic ceramic fuel cell was characterized using hydrogen fuel as a base case, with 50 cm³ min⁻¹ of dry hydrogen fed to the anode, while atmospheric oxygen was reduced at the cathode. The cell's active area of 0.69 cm² was calculated based on the outer diameter. Fig. 3a shows the cell potential difference (*U*)-current density (*j*)-power density data for operating temperatures from 550 °C to 700 °C, which is the generally accepted intermediate-temperature range. At those temperatures, the measured open-circuit potential difference (OCPD) in pure hydrogen was 1.13–1.15 V, comparing well with values calculated from Nernst's equation, so indicating the YSZ-GDC dual-layer electrolyte was gas-tight after sintering.

Maximum power densities of 0.93, 1.75 and 2.27 W cm⁻² were measured at 600, 650 and 700 °C, respectively. To the best of our knowledge, 2.27 W cm⁻² at 700 °C using hydrogen fuel is the highest power density reported in the literature for any SOFC design, as shown in Table 1. In addition, the anode's geometrical asymmetry and deliberate separation of mechanical support from the EAR demonstrated effectiveness in facilitating gas transport, as illustrated by electrochemical impedance spectroscopy (EIS) in Fig. 3b. The Nyquist curves were fitted using a classical equivalent circuit $LR_s(R_1||CPE_1)(R_2||CPE_2)$, where *L* stands for an inductor, *R* for a resistor and CPE for constant phase element. The fitted impedance spectra are



Fig. 3 (a) Effect of current density (*j*) on cell potential difference (*U*) and power densities of micro-monolithic SOFC at 550–700 °C, with 50 cm³ min⁻¹ H₂ as the fuel and ambient air as oxidant. (b) Effect of temperature on impedance spectra (10^{-1} to 10^{5} Hz) at open circuit potential difference. The inset illustrates equivalent circuit diagram for EIS analysis.

 Table 1
 Comparison of electrochemical performance for ceramic fuel

 cells operating at intermediate and high-temperature range

Type of fuel	Temperature/°C	Max. power density/W cm^{-2}	Ref.
Hydrogen	700	2.27	This work
	850	1.70	29
	800	0.98	25
	900	1.32	30
	600	0.93	This work
	600	0.62	31
	600	0.39	32
	600	0.431	33
	600	0.65	21
	600	0.8	34
LCVG	750	2.2	This work
LCVG	750	0.14	28
Biogas	800	0.19	35
Biogas	875	0.07	36
Biogas	800	0.2	37
Propane	850	1.3	29

shown in Fig. S4 (ESI \dagger) and the fitting results are summarized in Table S2 (ESI \dagger). The low-frequency (1–100 Hz) arc of the impedance spectrum, implied an area-specific diffusion resistance of approximately 0.16 Ω cm², which is *ca.* 1/3 the values for other micro-monolithic counterparts reported previously,²⁶ suggesting the diffusional resistance inside the porous anode had been decreased significantly by this design's tailored morphology.

Due to the extraordinary geometrical asymmetry involved in the complex micro-monolithic electrode design, integrated computed fluid dynamics (I-CFD) modelling was undertaken on the cell's real 3D structure to further illustrate the influence of the design on facilitating gas transport. As shown in Fig. 2, the 3D reconstruction of the electrode obtained from a customdeveloped multi-length scale X-ray CT technique, which includes the micro-CT for the extraction of macro-structural features, as well as the nano-CT for determining micro-structural parameters.^{25,27} Integrated computed fluid dynamics (I-CFD), which was developed based on such multi-length scale X-CT technique, has been demonstrated to be capable of quantifying effective mass transport parameters on the hierarchically-structured porous anode, with the pore size spanning over two orders of magnitude. This is often challenging due to the balance between imaging resolution and field of view.^{25,27} This is the first time that such an I-CFD technique was employed to investigate the new micro-monolithic electrode, providing a more vivid and straightforward insight into gas transport in such a complex, hierarchical micro-structure. As shown in Fig. S5a-c (ESI⁺), when the gas feed was pure hydrogen, the hydrogen molar fraction in the EAR (Fig. 1b) was very close to the bulk concentration under typical operating current densities of 1 and 2 A cm⁻². Even at 4 A cm⁻², the hydrogen molar fraction decreased from 100% to only ca. 80%, due to hydrogen depletion by its electrochemical oxidation producing steam, suggesting well-optimized fuel transport. Such observations agree well with the U-j curves shown in Fig. 3a, in which the concentration polarization started to dominate at current densities up to 5.5–6 A cm $^{-2}$.

Electrochemical performance of cells fuelled by simulated LCVGs

Fig. 4 shows the electrochemical performances of the cells fuelled by simulated LCVGs with methane concentrations of 10%, 15% and 20%. Despite the high CO₂ content as an inert diluent in the feed, maximum power densities of 1.77, 2.03 and 2.22 W cm⁻² still were achieved at 750 °C. These values compare quite well with those reported in the literature for pure hydrogen and demonstrated direct and highly efficient conversion from 'valueless' LCVGs to electrical power over the intermediate temperature range. The power density of 2.22 W cm⁻² obtained with 20% CH₄ in CO₂ is several times higher than values reported previously with similar materials,²⁸ as shown in Table 1.

The superior power densities can be attributed to the welldesigned cell structure that optimizes fuel transport inside the EAR, eliminated the additional transport resistance from the support. It is understood that generally fuel conversion/ utilization for typical SOFCs should be limited to 70% to minimize concentration polarization losses and avoid Ni



Fig. 4 (a) Schematic diagram of a 6-channel micro-monolithic SOFC utilising LCVGs with reactions at triple-phase boundaries; (b) effect of current density (j) on cell potential difference (U) and power density at 750 °C, with various LCVGs fed as fuels.

oxidation at low H_2 : H_2O ratios, though such conversions may vary depending on specific geometries and operating conditions.

For simulated LCVGs with low fuel contents, concentration polarization becomes a dominant factor limiting cell performance. Therefore, reducing the diffusive path length from several hundred micrometres to $<50 \ \mu$ m, together with the micro-structure tailoring during fabrication, proved essential to achieving efficient conversion of simulated LCVGs. Such explanations could be confirmed further by the I-CFD modelling, results for which are shown in Fig. S5d and e (ESI[†]). When the inlet gas contains high percentages of inert diluent (set at 80%), the driving force for fuel diffusion decreased accordingly. It is noteworthy that although CO is also a gaseous fuel, H₂ had a slightly lower equilibrium potential difference (0.9839 V) than that of CO (0.9907 V) relative to oxygen reduction at 1027 K,38 and a much greater self-diffusion coefficient $(11.3 \text{ vs. } 1.7 \text{ cm}^2 \text{ s}^{-1} \text{ at } 1027 \text{ K})$.³⁹ Under the operating conditions used, hydrogen was expected to be oxidized preferentially, so CO was assumed to be 'sluggish or inert' during CFD modelling. Hydrogen concentrations decreased radially from 20% to approximately 15% and 8% under operating current densities of 1 and 2 A $\rm cm^{-2}$, respectively; fuel starvation was predicted at 4 A cm⁻². The permeation flux density (*J*, mol m⁻² s⁻¹) driven by a concentration gradient, can be derived by a modified Fick's first law shown in eqn (S1) (ESI[†]). As the hydrogen concentration in the initial feed was decreased, micro-structure tailoring, as discussed in the Morphology section, becomes critical to facilitate gas transport, such as by decreasing electrode thickness, increasing the porosity or decreasing the tortuosity. As we reported previously, introducing micro-channels could decrease the tortuosity effectively by several times.25 The structural tailoring, together with the thickness control in the EAR, significantly decreased concentration polarization and contributed to achieving increased electrochemical performances when simulated LCVGs were used as fuel.

As discussed in the Introduction, an important benefit of utilizing LCVGs as fuels in SOFCs is their relatively high operating temperatures (>600 °C) that favour the DRM reaction, according to thermodynamic predictions (Fig. S6 ESI,† endothermic reaction, $\Delta H_{298K} = 247 \text{ kJ mol}^{-1}$). Based on results of our calculations of thermodynamic equilibria, shown in Fig. S8 (ESI^{\dagger}), complete conversion (conversion rate > 99.9%) of methane could be achieved theoretically at temperatures of 660, 720 and 950 °C with 10%, 20% and 50% initial methane contents, respectively. However, due to the miniaturization of the dimensions of the single cell, the residence time for the DRM reaction was constrained to approximately 0.1 s. Thus, the actual methane conversion can be guite far from thermodynamic equilibrium conditions if the cell performs solely as a catalytic reactor without any current loads, as shown in Fig. S9 (ESI[†]). On the other hand, once the DRM reaction was coupled with electrochemical reactions, during which both hydrogen and carbon monoxide were formed by reaction (1) then consumed continuously by electrochemical oxidation, the methane conversion could be increased dramatically.

As mentioned previously, gas flaring, which is the conventional route to dispose of natural gas via controlled combustion, is the most common approach for LCVG abatement in order to prevent direct emission of methane as a major greenhouse gas. However, presently it is also one of the most detrimental for energy wastage and climate change. For LCVG flaring, the high CO₂ concentration frequently necessitates addition of extra fuel to reach the ignition limit and maintain the flame stability.^{3,4} Therefore, the disposal of LCVGs via flaring is not only a waste of primary energy, but also a significant contributor to global climate change, as shown in Fig. S10 (ESI⁺). By comparison, operating ceramic fuel cells with LCVGs could result in highly efficient recovery of their calorific values by conversion to electrical energy, with greatly decreased environmental consequences. As predicted by the techno-economic analysis (Fig. S10, ESI[†]), a minimum electricity price of 6.5 ¢ kW⁻¹ h⁻¹



Fig. 5 Concentrations of different compositions in the effluents of LCVG mixtures (10–20% CH₄) at 650 °C under open-circuit condition (residual CH₄ mole fractions not shown as values were < 0.1%).

could generate revenue to off-set capital expenditure, maintenance and operation cost by selling generated electricity. As provide by U.S. Energy Information Administration (EIA), the latest average national electricity price is approximately 13 ¢ kW⁻¹ h⁻¹.⁴⁰ Therefore, the utilization of LCVGs using ceramic fuel cells could well be profitable.

The effluent gas mixtures were characterized using gas chromatography (GC) to quantify mole fractions of the components, as exemplified in Fig. 5 for 650 °C under open-circuit conditions. Due to the relatively low methane content, the conversion of methane at 650-750 °C for all three LCVG mixtures was close to their thermodynamic equilibrium composition (Fig. S7, ESI[†]), confirmed by the negligible methane content measured downstream of the reactor. The effluents contained approximately 45 to 61% syngas (CO + H₂), the balance being CO_2 . The H₂/CO ratio was slightly less than 1 (stoichiometric ratio in DRM reaction (1)) due to the reverse water-gas shift reaction. With applied current, the effluent compositions changed according to the additional oxygen content. In addition to efficient electricity generation, the effluents of LCVG utilization are still of considerable industrial interest, e.g. for methanol production via catalytic CO₂ hydrogenation.^{41,42} As demonstrated in several industrial plants, methanol manufactured from these H2-CO-CO2 mixtures potentially can be used as a fuel or adopted as feedstock to produce further value-added chemicals. The highly efficient energy conversion resulting from the well-designed cell structure, together with the significant industrial importance of the downstream effluents, provides a new route to utilize LCVGs and to decrease their carbon footprints effectively.

Longer-term coking resistance test using LCVGs

Coking is a critical concern for operating ceramic fuel cells with hydrocarbon fuels, particularly at intermediate temperatures (500–800 °C). According to the well-established ternary diagram of thermodynamic equilibrium of C–H–O system,¹² in which the boundary contours of the coking region have been defined as a function of temperature, it has been reported that the coking region expands as the temperature decreases from 800



Fig. 6 Effect of time on current densities of a single cell at a constant cell potential difference of 0.7 V fuelled by CH_4-CO_2 mixture (1:1 ratio) and ambient air as the oxidant.

to 600 °C; all three feed compositions used were at the risk of coking if the cells were operated in the intermediate temperature range. Fig. S6 (ESI[†]) shows the Gibbs energy changes and equilibrium constants of the three side reactions (2)–(4) that lead to coking. As shown in the figure, both the Boudouard reaction and methane cracking have equilibrium constants (K_T) larger than 1 within the intermediate temperature range used (650–750 °C); the formation of carbon is more likely when operating at even lower temperatures. To investigate the longer-term coking resistance of the cell, a CH₄:CO₂ ratio of 1:1 was selected as the feed. As evident in the C–H–O ternary diagram, the mixture containing 50:50 of CH₄–CO₂ is predicted thermodynamically to favour coking at all temperatures used in this study, as has been confirmed experimentally.²⁸

Fig. 6 shows the time dependence of current densities of the novel micro-monolithic ceramic fuel cell operating at 0.7 V and 700 °C with 50 cm³ min⁻¹ of CH₄-CO₂ mixture as the fuel (1:1 ratio), exhibiting a degradation rate over 500 hours of <2% (0.02 A cm⁻² kh⁻¹). Fig. 7 shows SEM images around the inner contour after operation. As shown in Fig. 3a, the bulk composition of gaseous fuel varies axially as a result of H₂/CO oxidative depletion, with a corresponding increase in steam concentrations. Therefore, the surfaces of the support region (Fig. 7c, region A1) and active region (Fig. 7c, region A2) at both upstream and downstream of the cell were examined. The EAR (A2) along the whole axial direction was free from coking, due to the relatively high steam concentration generated from hydrogen oxidation (Fig. 4a) that maintained the local C:H:O ratio below the coking threshold (Fig. 7a). In contrast, the steam content at fuel inlet was almost negligible and the feed composition $(CH_4:CO_2 = 1:1)$ was well within the coking region. Hence, carbon filaments were observed on the support (A1) at the upstream region (Fig. 7e and f), in agreement with previous reports,¹⁷ whereas the downstream side was free of coking. However, the influences of such coking on current densities at constant potential difference and on structural integrity were almost negligible. Thus, it can be concluded that even with a conventional Ni-based electrode, which is well known to be prone to coking, relatively stable operation could



Fig. 7 Post-mortem SEM images of SOFC after 500 hours operation at 700 °C fuelled by 50 cm³ min⁻¹ CH₄–CO₂ mixture (1:1): (a) surface of exterior electrochemical active region (EAR); (b) concentration profile of steam in EAR from CFD modelling; (c and d) schematic diagram of cell structure on *X*-axis. (e and f) Surfaces with coking on the central support region.

be achieved as result of controlling area-specific C:H:O ratios *via* morphology design and optimization. Potentially, this provides a new and more cost-effective approach to suppress the long-standing coking problem and enable efficient utilization of LCVGs as an overlooked energy source.

Conclusions

A new strategy was proposed to realize cost-effective and efficient recovery of LCVGs as a long-overlooked energy source, using a novel high-performance, coking-resistant ceramic fuel cell. Various parameters of the fabrication process were controlled and manipulated to optimize the micro-monolithic design in the anode substrate. The unique geometrical asymmetry, as well as the micro-structure induced from the phase inversion process, dramatically increased mass transport rates, confirmed by CFD modelling assisted by X-ray CT characterization, ensuring adequate fuel supply despite the high concentration of inert diluent in the LCVG feed. The electrochemical characterizations were conducted with both pure hydrogen and simulated LCVGs. A maximum power density of 2.27 W cm⁻² was obtained at 700 °C, which is the highest ever reported for any fuel cell geometry. Impedance spectroscopy analysis confirmed the minimization in concentration polarization, the value of which is *ca.* 1/3 of sub-optimal counterparts. Superior performances were also been obtained when simulated LCVGs were adopted, showing maximum power densities of 1.77, 2.03

Paper

and 2.22 W cm⁻² with 10, 15, and 20% methane contents, respectively. These performances were also several times higher than reported reference values. The geometrical asymmetry brings additional benefits of enabling control over area-specific gas atmospheres, which helps to suppress coking. A longer-term stability test was conducted with a CH₄–CO₂ feed ratio of 1:1, predicted thermodynamically to favour coking. The cell was operated for more than 500 hours with negligible degradation in current output. Post-mortem examination confirmed the entire EAR was free from coking, and the gradually increased steam concentration in the axial direction protected most of the cell support from carbon formation. This novel fuel cell design represents a new strategy to tackle the long-existing coking issue in a much cheaper and effective way.

Author contributions

The new coking-resistant design was conceived by TL and KL. TL and BW performed the electrode fabrication and morphological analysis. MFR prepared the cells for testing and performed the electrochemical tests and long-term dwelling test with TL, NF and GK. OM and NB performed impedance analysis and equivalent circuit fitting. XL, PS and DB conceived X-CT characterization and CFD modelling. TL, GK and KL prepared the manuscript with feedbacks provided from other authors. KL directed all aspects of the project contributions at Imperial, while PS and DB at UCL. They provided research oversight and funding to support the work.

Conflicts of interest

There are no conflicts to declare.

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