Direct Solid Oxide Electrolysis of Carbon Dioxide: Analysis of Performance and Processes

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Article

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Abstract: Chemical industries rely heavily on fossil resources for the production of carbon-based chemicals. A possible transformation towards sustainability is the usage of carbon dioxide as a source of carbon. Carbon dioxide is activated for follow-up reactions by its conversion to carbon monoxide. This can be accomplished by electrochemical reduction in solid oxide cells. In this work, we investigate the process performance of the direct high-temperature CO\textsubscript{2} electrolysis by current-voltage characteristics (iV) and Electrochemical Impedance Spectroscopy (EIS) experiments. Variations of the operation parameters temperature, load, fuel utilization, feed gas ratio and flow rate show the versatility of the procedure with maintaining high current densities of 0.75 up to 1.5 A cm\textsuperscript{-2}, therefore resulting in high conversion rates. The potential of the high-temperature carbon dioxide electrolysis as a suitable enabler for the activation of CO\textsubscript{2} as a chemical feedstock is therefore appointed and shown.

Keywords: carbon dioxide; solid oxide electrolysis; carbon dioxide utilization; CO\textsubscript{2}-electrolysis; high-temperature electrolysis; carbon dioxide reduction

1. Introduction

The world-wide chemical industry is currently depending on fossil resources for the production of carbon based chemicals. To reach full sustainability in the sector of chemical industry, the necessity of circular carbon economy is evident. This is not possible by exploiting fossil resources. An abundant compound containing carbon is carbon dioxide (CO\textsubscript{2}). Even in a scenario of 100% renewable energy generation, emissions of carbon dioxide by processes within the chemical sector is in the range of multiple hundred million metric megatons per year [1]. As CO\textsubscript{2} has a low energetic value and is therefore rather inert, it has to be activated to be used as a starting material in value-chains of the chemical industry. Carbon monoxide (CO) is already a bulk chemical in industrial chemistry. It is used as a building block in polymers like e.g., polyurethanes and polycarbonates, but can also be transformed to a vast range of products as the carbon containing part in syngas, a mixture of carbon monoxide and hydrogen [2–7]. Hence, a sustainable conversion of carbon dioxide to carbon monoxide is enabling the processes of carbon capture and utilization (CCU) and power-to-x [8].

Using renewable energy, the electrochemical reduction of CO\textsubscript{2} to CO is an interesting and seemingly simple process for the solution of the above mentioned challenges. This reaction has been examined both for low-temperature and high-temperature procedures with vast differences in performance, efficiencies and general operation parameters [9]. The high-temperature electrolysis of
CO₂ to CO on ceramic solid oxide cells seems superior in these regards with a faradaic efficiency of 100% and overall energetic efficiencies of over 75%. This conversion has been attempted mostly as a part of syngas production by using water and carbon dioxide simultaneously as starting feeds [10], which is not a direct electrolysis of carbon dioxide in particular. Concerning the electrolysis of pure carbon dioxide, a few aspects have been viewed at, however with a focus on materials [11–18]. We present the process investigation of the variation of different operation parameters temperature, load, fuel utilization, feed gas ratios and flow rates of the high-temperature solid oxide electrolysis of carbon dioxide to carbon monoxide.

2. Materials and Methods

The commercially available single button cells (Elcogen®, Tallinn, Estonia) used in the following experiments consist of a cathode-supported porous NiO-8YSZ-cermet fuel cathode, a 8%-Yttria-stabilized zirconia (8YSZ) electrolyte, a Gadolinium-doped ceria (GDC) and a porous La₀.₆Sr₀.₄CoO₃₋δ (LSC) oxygen anode. The cathode, electrolyte and GDC layer have a diameter of 20 mm and a totalled thickness of 300 ± 30 µm. The anode has a diameter of 10 mm and a thickness of 15 ± 5 µm, giving an active area of 0.79 cm².

The single-cells are installed in a two-electrode conductivity & impedance spectroscopy Probostat-setup (Norwegian Electro Cermaics® NORECS, Oslo, Norway). A gold ring is used as a sealing sample between the anodic side of the cell and the inner ceramic gas chamber to separate the air and fuel side. The current collector on the fuel side consists of a nickel and a platinum mesh. On the air side, only platinum is used for the current collection. The setup is enclosed and heated up with a furnace from the company Carbolute Gero® (Neuhausen, Germany). The heating rate is kept at 2 °C·min⁻¹ throughout all the measurements to protect the materials from thermomechanical stress. The reduction of the fuel electrode from NiO-YSZ to Ni-YSZ to activate the catalyst properties of Nickel is carried out at 900 °C with a stepwise increase of hydrogen concentration in the fuel feed gas starting at 0% up to 100% with nitrogen as balance gas.

In this work, we have chosen the temperatures 800 and 850 °C for the investigation of effects of varying operation parameters. This is a suitable compromise, by which cell performance and material stability can be guaranteed. Besides, a temperature series between 700–900 °C in equivalent T⁻¹/K⁻¹-steps with the volumetric fuel composition of 80:20 of CO₂:CO has been performed to apply the Arrhenius equation.

The supply of the gases H₂, N₂, CO₂, CO (fuel side) and air (air side) is controlled by mass flow controllers (MFC, Bronkhorst Nord®, Kamen, Germany). A constant volume flow of 6 L·h⁻¹ is applied to both gas chambers, when not specified differently. Current-voltage characteristics (IV) and Electrochemical Impedance Spectroscopy (EIS) experiments are managed via a Vertex 5A Potentiostat/Galvanostat (Ivium Technologies®, Eindhoven, Netherlands) using a two-electrode system. The cell performance is measured consecutively twice by applying a differential current with a scan rate of 10 mA·s⁻¹ from open cell voltage (OCV) up to 1.4 V. The area-specific resistance (ASR) is calculated using the numerical differential of the IV curves at OCV and 350 mA·cm⁻².

The process analysis of the cell is determined by AC characterization at OCV and 350 mA·cm⁻². The spectra are recorded in a frequency range from 110 kHz to 0.11 Hz with 73 measuring points and an applied amplitude of 20 mA.

3. Results

3.1. Performance

The performance of high-temperature CO₂-electrolysis was studied by dc current-voltage measurements. The resulting IV curves for various CO₂ partial pressures, temperatures and flow rates are shown in Figure 1a–c, respectively.
Analog to before, there is a factor of 1.6, only this time from 0.42 Ω·cm$^2$.

ASR also increased with each 10% step of CO$_2$ (ASR) and current densities at 1.4 V (i$_{1.4V}$) for variation of CO$_2$ partial pressure was balanced by CO. That means reducing CO$_2$ partial pressure was accompanied by increasing CO and vice versa. The obtained area specific resistances (ASR) and current densities at 1.4 V (i$_{1.4V}$) for variation of CO$_2$ partial pressure are presented in Figure 2a (values in Table A1 in Appendix A). For increasing the CO$_2$ partial pressure, the ASR at OCV increases with a factor of 1.6 from 0.34 Ω·cm$^2$ at 50% CO$_2$ to 0.56 Ω·cm$^2$ at 90% CO$_2$. The increase in ASR also increased with each 10% step of CO$_2$ partial pressure. From 50% to 60% it only increases by 0.02 Ω·cm$^2$, while from 80% to 90% the increase is 0.13 Ω·cm$^2$. At 0.35 A·cm$^{-2}$, however, the trend is opposite of the one observed at OCV. With increasing the CO$_2$ partial pressure, the ASR decreases. Analog to before, there is a factor of 1.6, only this time from 0.42 Ω·cm$^2$ at 90% CO$_2$ to 0.67 Ω·cm$^2$ at 50% CO$_2$. The current density at 1.4 V increases with increasing CO$_2$:CO ratio with a very similar increase of 0.08 or 0.09 A·cm$^{-2}$ for each 10 percentage point step of partial pressure. The experimentally determined OCV is comparable to the theoretical OCV, which is calculated by the Nernst equation. Deviations are up to 1%.

**Figure 1.** iV curves at various operating parameters: (a) variation of CO$_2$:CO ratio at 800 °C and 6 L·h$^{-1}$; (b) variation of temperature for a ratio of CO$_2$:CO of 80:20 at 6 L·h$^{-1}$; (c) variation of flow rate for a ratio of CO$_2$:CO of 80:20 at 800 °C.
Figure 2. (a) ASR, $|i|1.4$ V and OCV for various ratios of CO$_2$:CO at 800 °C. (b) ASR, $|i|1.4$ V and OCV for various temperatures for the ratio of CO$_2$:CO of 80:20. (c) ASR, $|i|1.4$ V and OCV for varied flow rate for a CO$_2$:CO ratio of 80:20 at 800 °C. Two values are given because of the hysteresis. The first denotes the forward scan and the second the backward scan of the $i$V curve.
The variation of temperature was performed at a gas composition of 80% CO\textsubscript{2} + 20% CO at temperatures in the range of 700–900 °C. The respective ASRs and \( i_{1.4\, \text{V}} \) are given in Figure 2b (values in Table A2) showing decreased ASR and increased \( i_{1.4\, \text{V}} \) with increased temperature. The decrease in ASR is steeper at OCV than at 0.35 A·cm\(^{-2}\). Moreover, the ASR increases at 700 °C going from OCV to 0.35 A·cm\(^{-2}\), while it decreases at 900 °C. The transition from this increase to decrease is at around 734 °C, where both values of ASR are the same. The OCV follows the trend given by the Nernst equation and decreases with increasing temperature.

For an investigation of the residence time, flow rates were varied in the range of 1–6 L·h\(^{-1}\). The obtained ASRs do not vary significantly, while the current density at 1.4 V increases by 0.14 A·cm\(^{-2}\) with increasing the flow rate from 1 L·h\(^{-1}\) to 6 L·h\(^{-1}\) (Figure 2c, Table A3). For the OCV it can be observed that for flow rates below 3 L·h\(^{-1}\) the initial OCV and the one measured after an \( iV \) cycle increasingly differ. At 3 L·h\(^{-1}\) the difference is only 10 mV. However, at 1 L·h\(^{-1}\) it is already a deviation of 30 mV.

3.2. Impedance Analysis

Electrochemical impedance spectra were taken for the same parameter variations as discussed for the performance analysis. Additionally, the comparison at variation of current is performed. The spectra at various current densities, CO\textsubscript{2}:CO ratios, temperatures and flow rates are shown in Figure 3a–d, respectively.

The variation of current density in Figure 3a shows increased total resistance at 700 °C resulting from a huge increase in high and middle frequency processes. At 900 °C the total resistance is first decreasing with current and starts increasing at about 0.35 A·cm\(^{-2}\). Compared to the spectra at 700 °C the increase in resistance is mainly caused by low frequency impedance contributions. In both cases the ohmic resistance is constant over the whole current density range. Especially the low frequency process is shifted to lower frequencies with increased current density causing the semicircle to be pushed out of the measurement range and only showing half of it in the spectra.

The CO\textsubscript{2}:CO ratio causes different trends at OCV and 0.35 A·cm\(^{-2}\) as can be seen in Figure 3b. At OCV the total resistance increases strongly with increasing the CO\textsubscript{2}:CO ratio. The major contribution is from the low frequency impedance parts. At 0.35 A·cm\(^{-2}\) the trend is opposite: an increased CO\textsubscript{2}:CO ratio causes the impedance to decrease. In this case, the ratio also influences the middle frequency region of the impedance. With increased CO\textsubscript{2}:CO ratio the low frequency process is also shifted towards higher frequencies and more into the measured frequency range.

The variation of temperature in Figure 3c is the first measurement to also influence the ohmic resistance. With increased temperature it is decreasing. Similarly, the polarization resistance is decreasing with increased temperature. The main contributions are from the high and middle frequency impedances. The low frequency process is shifted as well with variation of temperature. Higher temperatures shift it to higher frequencies and into the measurement range.

The last process parameter that was investigated by impedance spectroscopy is the flow rate. Here, the behavior is again different at OCV compared to 0.35 A·cm\(^{-2}\). At OCV the flow rate shows no significant influence on the spectra, while at 0.35 A·cm\(^{-2}\) a slight decrease of middle and low frequency impedance can be observed for increased flow rate.
As can be seen from the iV curves in Figure 1, the current density generally increases with increasing voltage. However, increasing the voltage above 1.4 V leads to severe and fast degradation according to literature [3] and our own previous studies. Compared to the experiments reaching up to 1.5 A·cm\(^{-2}\) at 1.4 V seen in Figure 1, experiments going up to 1.8 V (even for as short as 15 min), the correspondent current density at 1.4 V was permanently reduced to below 0.1 A·cm\(^{-2}\). As measure for the performance it is therefore the current density at 1.4 V that is compared.
However, due to the non-linearity at higher current density caused by transport limitations it might be more suitable to perform the electrolysis at lower voltages, as the benefit of higher currents do not justify the increased power input.

Another measure for the performance of a cell is the area specific resistance, which is a measure for the energy loss of the system. It includes the loss from multiple processes, which can be deconvoluted in impedance spectroscopy.

The analysis of the performance at different process parameters show the highest performance concerning the current density at 1.4 V for 90% CO$_2$ + 10% CO, 900 ºC and a flow rate of 6 L.h$^{-1}$. Reducing the amount of CO$_2$ results in less available reactant and this leads to earlier mass transport limitations. The limitations occur mostly at higher currents when not enough reactant can be transported to the cell for the conversion forced by the current to take place. The potential increases often abruptly at a limiting current density. Regarding the fuel utilization, however, it can be seen that it does not even reach 10% for 50% CO$_2$. Only by decreasing the flow rate the utilization can be increased up to 40% (Figure 4). It must be noted though that the two measurements were conducted on different cells and it seems they show different grades of degradation. The degradation may hinder the diffusion pathways through the cell, reducing the local concentration of reactant. This explains the limiting current already at such low overall utilizations.

![Figure 4](image-url)

**Figure 4.** Operating potential dependent on the fuel utilization, calculated according to Faraday, for various flow rates for a composition of 80% CO$_2$ + 20% CO at 800 ºC.

The comparison of ASR at OCV and 0.35 A-cm$^{-2}$ showed an increase from OCV to 0.35 A-cm$^{-2}$ for CO$_2$ concentrations of 80% and lower. For a concentration of 90% CO$_2$, however the ASR decreases. Observing a larger ASR at OCV than under load is typical for a pronounced activation overpotential, which is especially dominant at low currents. Another explanation for the behavior at low current density is the Nernst potential. As 90% CO$_2$ + 10% CO is in the boundary of the Nernst potential, a decrease in CO$_2$ and increase in CO, like it will happen during electrolysis, causes the Nernst potential to increase steep in the range of 90% CO$_2$ down to 80% CO$_2$. Below 80% CO$_2$, the increase is significantly lower. The small changes in CO$_2$:CO ratio during electrolysis will then cause the thermodynamic voltage to increase much more at 90% CO$_2$ in the feed than at 50% CO$_2$.

Varying the temperature shows a similar influence on the ASR comparing OCV to 0.35 A-cm$^{-2}$. Here the transition is already at around 734 ºC. The deviation observed at around 800 ºC in this experiment may be explained by the use of a different cell than for the composition measurement. It also indicates that probably both an activation overpotential and the Nernst potential are causing this decrease in ASR at low current densities.
The temperature dependence measurement is also performed to gain information on the activation energies of the processes. Figure 5 shows the Arrhenius representation of the ASR_{OCV}. It can be observed that for temperatures above 850 °C the ASR remains constant. Below 850 °C the logarithmic of the resistance shows a slightly curved trend with recursive temperature increase. The ASR includes all the underlying processes that may occur during electrolysis. Seeing a slightly curved Arrhenius behavior indicates the domination of different processes at lower temperatures than at higher temperatures. Additionally, it may indicate an increasing degradation with increased temperature. The series was conducted from low to high temperatures increasing the operation time of the cell with each temperature step. Especially the constant resistance above 850 °C is attributed to degradation as the LSC material of the anode is reported to be unstable above this temperature [19].

Figure 5. Arrhenius representation (\(\ln(R) \rightarrow T^{-1}\)) of the area specific resistance for a ratio of CO_2:CO of 80:20.

4.2. Process Analysis

The impedance spectra in Figure 3 were analyzed by an equivalent circuit model (ECM) containing an inductance, a serial resistance and four RQ elements (a resistor and constant-phase element in parallel). The obtained resistances are plotted against their respective varied parameter in Figures 6–9.

Figure 6. Arrhenius plot for the resistances for a ratio of CO_2:CO of 80:20 at 6 L·h^{-1} at (a) OCV and (b) 0.35 A·cm^{-2}.
The resistance $R_{Q4}$ is also strongly influenced by the current density $|i|$ and $\mathcal{R}$, while $R_{Q1}$ is not affected in the same way. The increase in performance comes from a decreased mass transport, which decreases the diffusion resistance and with it also the resistance $R_{Q4}$.

**Figure 7.** Resistances with respect to current density for a ratio of CO$_2$:CO of 80:20 at 6 L·h$^{-1}$ at (a) 700 °C and (b) 900 °C.

The flow rate is influencing the resistances $R_{Q1}$ and $R_{Q2}$ significantly. The highest reactant utilization can be reached for low flow rates. With suitable mass flow rates, the decrease in performance regarding the resistances can be explained if they are connected/coupled.

**Figure 8.** Resistances with respect to the flow rate for a ratio of CO$_2$:CO of 80:20 at 800 °C at (a) OCV and (b) 0.35 A·cm$^{-2}$.

The resistance $R_{Q4}$ increases with increasing the CO partial pressure at 800 °C than at 700 °C and therefore shows a similar trend as the ASR observation from dc measurements. The highly increased resistance at high current densities supports the assignment of the process connected/coupled. With increased CO partial pressure, the CO partial pressure is decreased causing the partial pressure increase of both resistances can be explained. The decreased mass transport, however, is more significant with increased current density. Even though the overall reactant utilization is not expected and therefore the increase of $R_{Q4}$–$R_{Q1}$ cannot occur.

**Figure 9.** Resistances with respect to the CO$_2$ partial pressure at 800 °C and (a) OCV and (b) 0.35 A·cm$^{-2}$.
The variation of temperature might reveal the most insightful analysis towards the assignment of the RQ element to physical processes. The resistances can be plotted in an Arrhenius diagram (ln(R) \( \rightarrow T^{-1} \)) and from the slope the activation energy can be obtained. The Arrhenius plots for a CO\(_2\)-CO ratio of 80:20 are shown in Figure 6 and the respective activation energies in Table 1. For the activation energies the resistances for temperatures above 850 °C were not considered due to the cell degradation falsifying the results.

| Process | \( E_{a, OCV} \) [kJ·mol\(^{-1}\)] | \( E_{a, 0.35 \text{ A cm}^{-2}} \) [kJ·mol\(^{-1}\)] |
|---------|-------------------------------|----------------------------------|
| \( R_{Q1} \) | 40.6 ± 1.5 | 38.1 ± 2.0 |
| \( R_{Q2} \) | 71.8 ± 5.4 | 18.5 ± 3.8 |
| \( R_{Q3} \) | 36.2 ± 8.0 | 80.2 ± 8.4 |
| \( R_{Q4} \) | 36.3 ± 7.3 | 71.8 ± 2.4 |
| \( R_{Q4} \) | 10.0 ± 2.1 | 19.1 ± 3.5 |

The obtained activation energies differ hugely comparing OCV and 0.35 A·cm\(^{-2}\). Only the activation of the ohmic resistance stays the same for increased current density. With ca. 40 kJ·mol\(^{-1}\) it is only half of the activation energy given in literature [20,21]. There it is compared to the activation energy of ionic conductivity in 8YSZ, given as 88 kJ·mol\(^{-1}\) [22]. In the cells used in this publication, however, the electrolyte material is significantly thinner than what is used in literature. Since not only the ionic conductivity of 8YSZ but also the ionic and electronic conductivities of the electrodes and wires as well as the conduction across material borders influence the ohmic resistance, a reduced electrolyte thickness increases the contribution of those other parts. This results in lowering the activation energy of the ohmic resistance presented here.

The activation energies for the resistances \( R_{Q1}, R_{Q2} \) and \( R_{Q3} \) seem to switch in magnitude from OCV to 0.35 A·cm\(^{-2}\). While the thermal activation of \( R_{Q1} \) is significantly reduced with current, the activation of \( R_{Q2} \) and \( R_{Q3} \) are increased. Values for activation energies of single processes in CO\(_2\) electrolysis were up to this time not reported in literature. Li et al. [23] investigated the influence of process parameters including temperature, however, did not show dependencies of single processes. Their obtained activation energy for the polarization resistance of around 60–70 kJ·mol\(^{-1}\) is larger than what we obtain (29 kJ·mol\(^{-1}\)). It has to be noted though that they measure at different temperatures (600–700 °C) and gas compositions employing different cell types. If only the temperature range of 700–750 °C is considered the activation energy for our polarization resistance increases to 46 kJ·mol\(^{-1}\), which is closer to what is reported by Li et al. showing a comparability.

Other literature studies of CO\(_2\) electrolysis don’t include Arrhenius analyses. Only some studies performed on fuel cell analysis or steam or co-electrolysis report activation energies for separate processes. Reported by modelling studies [17,24] an activation energy of 86.50 kJ·mol\(^{-1}\) and 86.50 kJ·mol\(^{-1}\) is given for the surface reaction of adsorbed CO\(_2\) to adsorbed CO and O. This would fit \( R_{Q1} \) at OCV or \( R_{Q2} \) or \( R_{Q3} \) at 0.35 A·cm\(^{-2}\). As also the current density has a huge impact on \( R_{Q2} \) (compare Figure 5) it is assumed that this process is connected to the triple-phase boundary reaction. In the fitting procedure it also turned out that \( R_{Q1} \) and \( R_{Q2} \) tend to exchange. Leonide/Sonn et al. [20,25] report for fuel cells that the two high-frequency processes are coupled processes including oxygen transport in YSZ of the fuel electrode, charge transfer at the triple phase boundary on the fuel electrode and gas diffusion inside the fuel electrode functional layer.

The process being expressed by \( R_{Q4} \) shows only a low thermal activation. In other experiments of co-electrolysis in our group the activation energy was even lower or showed negative values. Due to the low activation energy it has been assigned to gas diffusion as reported in literature by Primdahl [26], Ebbesen [27] and Leonide [20]. The results presented in this study also show that the activation energy of this process increases with increasing the operating current. This was also observed in other experiments for co-electrolysis in our group.
The resistance $R_{RQ4}$ is also strongly influenced by the current density as can be seen in Figure 7. At low current densities it slightly decreases and then increases almost exponentially with increase current. This behavior is more pronounced at 900 °C than at 700 °C and therefore shows a similar trend as the ASR observation from dc measurements. The highly increased resistance at high current densities supports the assignment to a diffusion process as transport limitations become more and more significant with increased current density. Even though the overall reactant utilization is not reaching high values, a local depletion of reactant is possible.

The flow rate is influencing the resistances $R_{RQ2}$ and $R_{RQ4}$ under load as can be seen in Figure 8. Below the flow rate of 3 L·h$^{-1}$ both resistances increase. $R_{RQ2}$ has been assigned to a process connected to the reaction on the fuel side, while $R_{RQ4}$ has been assigned to a diffusion process. At increased current more reactant is converted. This reactant amount needs to be transported to the triple phase boundary (TPB) for conversion. Decreasing the flow leading to a decreased mass transport is expected and therefore the increase of $R_{RQ4}$ can be explained. The decreased mass transport, however, also reduces the local reactant concentrations, which then also influences the reaction itself. Thus, the increase of both resistances can be explained if they are connected/coupled. The reaction cannot occur without diffusion preceding it and if diffusion is negatively affected by flow rate, so is the TPB reaction.

The partial pressure dependence reveals an assignment of $R_{RQ2}$, $R_{RQ3}$ and $R_{RQ4}$ to the fuel electrode (Figure 9). Especially the process RQ4 is influenced by the partial pressure. At OCV the resistance increases with increasing the CO$_2$ partial pressure, while it decreases at 0.35 A·cm$^{-2}$. This might be explained by the fact that at OCV the ac signal is in both the fuel cell and electrolysis direction. With increased CO$_2$ partial pressure, the CO partial pressure is decreased causing the increase in resistance at OCV. The assignment to diffusional processes also applies well, since the low concentrations of CO then cause local depletion for CO reduction. There also the trend of $R_{RQ4}$ fits in well, as it is similar to that of $R_{RQ3}$ only less pronounced. At 0.35 A·cm$^{-2}$ the trend for $R_{RQ4}$ that was expected is observed. Increasing the CO$_2$ partial pressure increases the reactant concentrations and decreasing the diffusion resistance and with it also the resistance $R_{RQ2}$, which is probably connected to the reaction on the fuel side.

5. Conclusions

Taking the increased degradation into account, the optimal operating temperature of a direct carbon dioxide electrolysis on solid oxide cells would be 800–850 °C. The topic of degradation will and has to be addressed in detail in future publications of our group. The highest performance can then be obtained for 80–90% CO$_2$ balanced with CO. At 1.4 V, also the current density is the highest. However, it is more beneficial to stay at around 1.1 V as the increased current density above that voltage does not follow a linear increase with voltage. Therefore, the increase in performance comes at a high cost in additionally needed power. The general trend of our outcomes will be observed in on-going and future long-term degradation experiments.

The variation of the feed gas composition CO$_2$:CO starting at a ratio of 90:10 with increments of 10% to 50:50 shows the simulated gradient of a gas flow through channels over a larger cell. With these results for operation at load (0.35 A·cm$^{-2}$), the lack of a critical drop in performance regarding ASR and polarization resistance indicates, that an operation with both a high starting concentration of CO$_2$ and a high fuel utilization above 40% is possible. The dominant part of the decrease in performance is identified as the process of diffusion. With a distinct design of gas flux within stacks and system, the possibility of high conversion rates is given even for elevated feeds of CO$_2$:CO ratios.

The highest reactant utilization can be reached for low flow rates. With suitable mass flow controllers, lower rates than 1 L·h$^{-1}$ are possible and probably increase the utilization even further. These parameters describe superior efficiencies for the electrochemical reaction of CO$_2$ to CO. Our results give an extensive outlook for scaling carbon dioxide electrolysis into stacks and systems. Especially regarding the moderate increase from 0.44 to 0.47 Ω·cm$^2$ of ASR at 0.35 A·cm$^{-2}$ with the decrease in flow rate, the fuel utilization is improved significantly resulting in elevated space-time velocities and
yields. With a suitable chemical engineering of the electrochemical system, desired high fuel utilization combined with low degrading operation conditions are achievable.

Overall, our study explicitly exhibits the electrochemical trends of the variation of all important parameters for the running process of pure CO\textsubscript{2}-electrolysis. Our work puts a stronger focus on CO\textsubscript{2} as a useable base chemical without having water involved in the process. Thereby, it allows the contemplation of electrolysis for the production of carbon monoxide by diversifying from co-electrolysis and pure water electrolysis coupled with water-gas shift reaction. Therefore, the potential of the conversion of carbon dioxide to carbon monoxide via solid oxide electrolysis to be a key enabling technology for the transformation of chemical industry from fossil resources to sustainable sources of carbon is disclosed.

**Author Contributions:** L.D., S.F. and T.D. conceived and designed the experiments; L.D. and T.D. performed the experiments; L.D., S.F. and T.D. analyzed the data; L.D., S.F. and T.D. wrote the paper; S.F., L.D., I.V., R.-A.E. and L.G.J.d.H. reviewed and edited the manuscript; S.F., I.V. and L.G.J.d.H. supervised the work; R.-A.E. and L.G.J.d.H. were responsible for funding acquisition. All authors have read and agreed to the published version of the manuscript.

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**Appendix A**

### Table A1. ASR, |\(i\)|1.4 V and OCV for various ratios of CO\textsubscript{2}:CO at 800 °C.

| CO\textsubscript{2}:CO | ASR\textsubscript{OCV} [\(\Omega\) cm\(^2\)] | ASR\textsubscript{0.35 A\textsubscript{cm}\(^{-2}\)} [\(\Omega\) cm\(^2\)] | |\(i\)|1.4 V [A cm\(^{-2}\)] | OCV\textsubscript{exp} [V] | OCV\textsubscript{theo} [V] |
|---|---|---|---|---|---|---|
| 50:50 | 0.34 | 0.67/1.05 | 0.58/0.50 | 0.95 | 0.94 |
| 60:40 | 0.36 | 0.54/0.91 | 0.66/0.56 | 0.93 | 0.93 |
| 70:30 | 0.39 | 0.51/0.82 | 0.74/0.66 | 0.91 | 0.91 |
| 80:20 | 0.43 | 0.46/0.70 | 0.83/0.75 | 0.89 | 0.88 |
| 90:10 | 0.56 | 0.42/0.61 | 0.92/0.83 | 0.85 | 0.84 |

1 Two values are given because of the hysteresis. The first denotes the forward scan and the second the backward scan of the iV curve.

### Table A2. ASR, |\(i\)|1.4 V and OCV for various temperatures for the ratio of CO\textsubscript{2}:CO of 80:20.

| T [°C] | ASR\textsubscript{OCV} [\(\Omega\) cm\(^2\)] | ASR\textsubscript{0.35 A\textsubscript{cm}\(^{-2}\)} [\(\Omega\) cm\(^2\)] | |\(i\)|1.4 V [A cm\(^{-2}\)] | OCV\textsubscript{exp} [V] | OCV\textsubscript{theo} [V] |
|---|---|---|---|---|---|---|
| 700 | 0.53 | 0.60 | 0.74 | 0.94 | 0.94 |
| 717 | 0.47 | 0.51 | 0.85 | 0.93 | 0.93 |
| 734 | 0.44 | 0.44 | 0.95 | 0.92 | 0.92 |
| 752 | 0.41 | 0.39 | 1.03 | 0.91 | 0.91 |
| 771 | 0.41 | 0.39 | 1.07/1.03 | 0.90 | 0.90 |
| 791 | 0.37 | 0.35 | 1.14/1.07 | 0.89 | 0.89 |
| 811 | 0.36 | 0.33 | 1.21/1.17 | 0.88 | 0.88 |
| 832 | 0.35 | 0.30 | 1.28/1.24 | 0.87 | 0.86 |
| 853 | 0.33 | 0.30 | 1.32 | 0.86 | 0.85 |
| 874 | 0.33 | 0.29 | 1.43 | 0.84 | 0.84 |
| 900 | 0.34 | 0.29 | 1.50 | 0.83 | 0.83 |

1 Two values are given because of the hysteresis. The first denotes the forward scan and the second the backward scan of the iV curve.
Table A3. ASR, $|i|1.4$ V and OCV for varied flow rate for a CO$_2$:CO ratio of 80:20 at 800 °C.

| Flow Rate [l h$^{-1}$] | ASR$_{OCV}$ [Ω cm$^2$] | ASR$_{0.35}$ [A cm$^{-2}$] | $|i|1.4$ V [A cm$^{-2}$] | OCV$_{exp}$ [V] | OCV$_{tho}$ [V] |
|------------------------|------------------------|------------------------|------------------------|----------------|----------------|
| 1.0                    | 0.44                   | 0.47                   | 1.11                   | 0.87/0.90      | 0.88           |
| 1.5                    | 0.44                   | 0.46                   | 1.12                   | 0.87/0.89      | 0.88           |
| 2.0                    | 0.43                   | 0.46                   | 1.14                   | 0.87/0.89      | 0.88           |
| 2.5                    | 0.42                   | 0.46                   | 1.17                   | 0.87/0.88      | 0.88           |
| 3.0                    | 0.43                   | 0.45                   | 1.19                   | 0.87/0.88      | 0.88           |
| 4.0                    | 0.42                   | 0.44                   | 1.22                   | 0.87/0.87      | 0.88           |
| 5.0                    | 0.42                   | 0.44                   | 1.23                   | 0.87/0.87      | 0.88           |
| 6.0                    | 0.42                   | 0.44                   | 1.25                   | 0.87/0.87      | 0.88           |

$^1$ Two values are given because of the hysteresis. The first denotes the forward scan and the second the backward scan of the IV curve.

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