Hossain, Md Noor; Khakpour, Reza; Busch, Michael; Suominen, Milla; Laasonen, Kari; Kallio, Tanja

Temperature-Controlled Syngas Production via Electrochemical CO2 Reduction on a CoTPP/MWCNT Composite in a Flow Cell

Published in:
ACS Applied Energy Materials

DOI:
10.1021/acsaem.2c02873

Published: 09/01/2023

Document Version
Publisher's PDF, also known as Version of record

Published under the following license:
CC BY

Please cite the original version:
Temperature-Controlled Syngas Production via Electrochemical CO₂ Reduction on a CoTPP/MWCNT Composite in a Flow Cell

M. Noor Hossain, Reza Khakpour, Michael Busch, Milla Suominen, Kari Laasonen, and Tanja Kallio

1. INTRODUCTION

Anthropogenic activities generate more than 30 billion tons of CO₂ annually, and part of this gas accumulates in the atmosphere and contributes to the increase of atmospheric CO₂ concentration, which has reached a record high level of ca. 417 ppm recently. Among all the emission sources, chemical industries play a major role. For a sustainable future, actions are required to lower the atmospheric CO₂ concentration and cut the emission. One promising approach is to use industrial CO₂ waste streams as a feedstock for chemicals and fuel production as the conventional fossil fuel-based feedstocks need to be replaced with more environmentally friendly techniques. In this context, electrochemical reduction of CO₂ and H₂O to syngas is a promising and potentially viable approach to convert CO₂ waste streams into valuable commodity chemicals and fuels utilizing the Power-to-X (P-to-X) concept. Syngas is known as a building block for various high-value chemicals produced industrially, including alcohols, methane, dimethyl ether (DME), oxo-alcohols, ethylene, ethylene glycol, and many more. Currently, production of syngas utilizes environmentally detrimental techniques, for instance, coal gasification and steam reforming of fossil fuels or natural gas. These techniques are also among the major sources of industrial greenhouse gas emissions. Therefore, the eCO₂R to syngas approach can provide the society with multiple benefits, such as, cutting the CO₂ emissions by utilizing unavoidably emitted CO₂ waste gas streams, for example, from steel production plants and the cement industry. Controlling the purity and H₂/CO ratio of syngas is important for various downstream processes in the chemical industry, although the required ratios differ. For instance, H₂/CO ratio of 1.0–1.2 is required for hydroformylation, such as the reaction depicted in Scheme 1.

The H₂/CO ratio can be controlled by catalyst design. Numerous catalyst materials including metal alloys, metal nanoparticles, and molecular catalysts have been developed and investigated for the production of syngas via eCO₂R in aqueous media. To obtain a desirable ratio in syngas, a bifunctional electrocatalyst consisting of Au and Co has been investigated by Michael et al. In their study, a H₂/CO ratio of 0.6–2.1 was obtained in an aqueous environment. Varying amounts of Au nanoparticles loaded on a titane nanosheet (TiNS) substrate have been investigated by Filipe et al. to obtain the desired syngas ratio. They have found that with the decrease of the amount of Au nanoparticle loading on the TiNS substrate, the H₂/CO ratio increases. Along with the catalyst design, most of the approaches have also focused on...
investigating the applied potential for controlling the syngas H\textsubscript{2}/CO ratio.

Molecular catalysts supported on highly conductive carbon materials have been investigated mainly for CO production. For instance, Fe, Co, and Ni molecular catalysts supported on carbon show >90% selectivity for CO production in aqueous electrolyte media in a so-called H-cell.\textsuperscript{23\textendash}25 The potential-dependent eCO\textsubscript{2}R to CO conversion has been investigated in a flow-cell configuration on a ligand-modified cobalt phthalocyanine/carbon nanotube (CoPC/CNT) composite and on Fe-porphyrin and resulted in FE >94% for CO formation.\textsuperscript{26,27} Interestingly, syngas production on molecular catalysts has drawn less attention, though few works have been performed either in an H-cell\textsuperscript{28} or a three-electrode cell\textsuperscript{29} on Fe-porphyrin-based metal–organic framework (Fe-MOF) catalysts in aqueous media. Cobalt tetrphenylporphyrin (CoTPP) and its derivatives supported on carbon black also show >80% selectivity for CO production in the potential range from −0.5 \textit{V}_{\text{RHE}}\textsuperscript{26} to −0.65 \textit{V}_{\text{RHE}}\textsuperscript{30} in flow conditions. However, the current density shows a decreasing trend at −0.65 \textit{V}_{\text{RHE}} during constant potential electrolysis. The catalyst support is crucial in terms of, for example, electronic conductivity, chemical and mechanical stability, and surface area\textsuperscript{31} in this sense, temperature-controlled eCO\textsubscript{2}R to syngas production in a flow cell on CoTPP supported on carbon nanotubes (MWCNTs) has remained unexplored.

Indeed, temperature can affect the eCO\textsubscript{2}R by various ways such as increasing the pH and the conductivity of the electrolyte and decreasing the concentration of dissolved CO\textsubscript{2}, surface tension, viscosity of the electrolyte, and the adsorption strength of the intermediate species.\textsuperscript{32\textendash}35 Increasing temperature has been shown to switch the competition between the eCO\textsubscript{2}R and the hydrogen evolution reaction (HER). Product selectivity during the eCO\textsubscript{2}R on Cu has been shown to be largely affected by temperature, for example, CH\textsubscript{4} and C\textsubscript{2}H\textsubscript{4} production is favored at lower temperatures, while H\textsubscript{2} is dominant at higher temperatures in an H-cell.\textsuperscript{33} Also, CO formation selectivity has been observed to decrease, while that of H\textsubscript{2} increased with temperature increment in a flow-cell configuration on tin oxide and Ag catalysts, respectively.\textsuperscript{36,37} A highly selective eCO\textsubscript{2}R to the CO catalyst can strongly suppress the HER which inevitably takes place on the cathode according to reaction 1 when an aqueous electrolyte is used as the reaction medium. Applying temperature can reduce the degree of suppression of HER which would allow more H\textsubscript{2} production along with eCO\textsubscript{2}R to CO according to reaction 2.\textsuperscript{32} Hence, besides the catalyst design tactics, applying temperature could be a fundamentally interesting approach to produce syngas with controlled H\textsubscript{2}/CO ratios.

\textbf{Reaction 1:} \textit{H}_{2}\textit{O} + e\textsuperscript{−} \rightarrow \frac{1}{2}\textit{H}_{2} + \text{OH}\textsuperscript{−}

\textbf{Reaction 2:} \textit{CO}_{2} + 2e\textsuperscript{−} + \textit{H}_{2}\textit{O} \rightarrow \textit{CO} + 2 \text{OH}\textsuperscript{−}

Herein, we have investigated temperature-controlled syngas production by eCO\textsubscript{2}R in a flow cell on a composite comprising cobalt tetrphenylporphyrin supported on multiwalled carbon nanotubes (CoTPP/MWCNTs). To the best of our knowledge, systematic investigation of low-temperature-controlled syngas production has not been performed in a flow cell on CoTPP catalysts. Our results indicate that the composite is highly selective for CO at 20 °C. Interestingly, this selectivity is altered with increasing temperature in the range of 20\textendash}50 °C, resulting in an increase of the syngas H\textsubscript{2}/CO ratio. To gain insight on the mechanism, the origin of high selectivity of the composite toward CO formation is investigated through DFT modeling. These results underline that at −0.8 and −1.0 \textit{V}_{\text{RHE}} the eCO\textsubscript{2}R to CO becomes barrierless compared to its counterpart HER. In addition, the potential-induced oxidation state of the Co atom of the CoTPP molecule triggers the catalytic activities for CO formation. Moreover, the introduction of the \( pK_a \) value in addition to the activation barrier reveals the stability of the formed intermediate species such as Co−H for HER and Co−COOH for eCO\textsubscript{2}R. This approach combining experimental and theoretical studies is beneficial for
both investigating the production of syngas and for understanding the eCO$_2$R mechanism.

### 2. EXPERIMENTAL SECTION

#### 2.1. Electrode Preparation

A CoTPP-to-MWCNT ratio of 6:1 was chosen for these experiments based on earlier optimization.$^3$ Detailed description of the synthesis and structural characterizations of the CoTPP/MWCNT (6:1) composite can be found elsewhere.$^3$ Briefly, 50 mg of MWCNTs (9 nm diameter, >95% carbon, Nanocyl) was dispersed in 100 mL of 2 mM HCl (Merck) aqueous solution. 300 mg of CoTPP (dye content 85%, Sigma-Aldrich) was dispersed separately in 92 mL of 2 mM HCl (Merck) solution with the help of 8 mL of DMF (99.8%, Sigma-Aldrich). Both dispersions were sonicated for 1 h. After mixing, the MWCNT and CoTPP dispersion was sonicated again for 1 h followed by stirring for 48 h. Then, the supernatant solution was thrown away, and the precipitate was centrifuged for 1 h at 4000 rpm at 20 °C. This step was repeated multiple times. Finally, the precipitate was washed with Milli-Q water and freeze-dried for 24 h.

In all prepared electrodes for the flow-cell experiments, the CoTPP/MWCNT composite loading was kept at 1 ± 0.1 mg/cm$^2$. An appropriate amount of the composite was weighed and dispersed in a mixture with 300 μL of pure ethanol and 150 μL of Milli-Q H$_2$O. Subsequently, 5 wt % Nafion (Sigma-Aldrich) solution was added in the ink to obtain a 30 wt % Nafion ionomer content. The ink was sonicated for ca. 10–15 min and then stirred for 20–30 min and subsequently sprayed on 10.2 cm$^2$ microporous carbon paper (Sigracet 25 BC) gas diffusion electrode (GDE) by employing an air brush. Before deposition of the composite, the GDE was cleaned with ethanol and dried in a vacuum oven for 2 h at 60–70 °C and weighed. After spray painting, the electrode was dried again in a vacuum oven at 60–70 °C for 2–3 h to remove the solvents. The weight difference before and after spraying gives the amount of the loaded composite on the GDE.

#### 2.2. Chronoamperometry and Product Analyses

A custom-made electrochemical flow cell was employed to perform constant potential electrolysis. The schematic view of the flow cell consisting of a cathodic and anodic compartment separated with a pretreated Nafion 115 membrane is shown in Figure 1. The working electrode is mounted on a Ti current collector plate. An Ir mixed metal oxide frame (Ir MMO, Electrocell A/S) and a 1 mm leak-free Ag/AgCl reference electrode, respectively. Polytetrafluoroethylene (PTFE) was used as an electrolyte. A peristaltic pump was used to circulate the electrolyte at 15 mL/min in the front side of the gas diffusion electrode (GDE) by employing an air brush. Before deposition of the composite, the GDE was cleaned with ethanol and dried in a vacuum oven for 2 h at 60–70 °C and weighed. After spray painting, the electrode was dried again in a vacuum oven at 60–70 °C for 2–3 h to remove the solvents. The weight difference before and after spraying gives the amount of the loaded composite on the GDE.

The working electrode is a cathodic and anodic compartment separated with a pretreated Nafion 115 membrane is shown in Figure 1. The working electrode is maintained at the eCO$_2$R mechanism.

The working electrode is maintained at the eCO$_2$R mechanism. The applied potentials were converted to the reversible hydrogen electrode (RHE) using the following equation:

$$E_{\text{RHE}}(V) = E_{\text{Ag/AgCl}}(V) + \frac{2.303RT}{nF}\log([\text{H}^+]/\text{H}_2)$$

(1)

where $E_{\text{Ag/AgCl}}(V)$ is the applied potential, R is the universal gas constant (8.314 J K$^{-1}$ mol$^{-1}$), T is the applied temperature (in Kelvin), n is the number of electrons transferred in the reaction, and F is the Faraday constant (96485 C mol$^{-1}$).

Manual iR correction was performed. To compensate for the iR drop, electrochemical impedance spectroscopy (EIS) was employed to determine the ohmic resistance. The optimum frequency was chosen by applying EIS frequency ranging from 1 kHz to 1 Hz. The measured Ohmic resistance was then multiplied by the current obtained in cathodic cyclic voltammetry (CV) measurement, and the potential was corrected according to eq 2.

$$E_{\text{Ag/AgCl}}(V) = E_{\text{RHE}}(V) + 0.2(V) + \frac{2.303RT}{nF}\log([\text{H}^+]/\text{H}_2)$$

(1)

$$E_{\text{Ag/AgCl}}(V) = E_{\text{RHE}}(V) - \frac{2.303RT}{nF}\log([\text{H}^+]/\text{H}_2) + (+ iR)(V)$$

(2)

All potentials reported here are iR corrected. Before CA measurements, N$_2$ was purged for ca. 30 min to remove O$_2$ from the reaction environment, and then CV measurement was carried out. After that, CO$_2$ was purged for 30–40 min to saturate the electrolyte with CO$_2$, and CV measurement was repeated. CA measurements were performed at -0.6 V$_{\text{RHE}}$, -0.8 V$_{\text{RHE}}$ and -1.0 V$_{\text{RHE}}$ at selected temperatures (20, 30, 40, and 50 °C). The pH of the CO$_2$-saturated electrolyte was measured at each temperature equaling 6.76, 6.86, 7.17, and 7.23 for 20, 30, 40, and 50 °C, respectively. The temperature of the cell was controlled by placing catholyte and anolyte reservoirs in a temperature-controlled water bath. The cell temperature was determined from catholyte and anolyte outlet streams by a thermometer. During CA, CO$_2$ was continuously purged into the cathode through the GDE with a flow rate of 19 SCCM to keep the constant supply of the reactant gas in the electrode/electrolyte interface. The catholyte reservoir was connected to a micro gas chromatograph (mGC, Agilent 990). Between the catholyte reservoir and the mGC, an ice trap was used to lower the humidity of the outlet gas stream to protect the mGC columns. The effluent gases were passed through a flow meter before entering the mGC. The gaseous products were analyzed with the mGC equipped with CP-MOLESIEVE 5A and CP-PORAPLOT U 10 m long columns and micromachined thermal conductivity detectors (TCD). The permanent gases including CO, H$_2$, and N$_2$ were separated in the Mol sieve 5A column, while the hydrocarbons, for example, CH$_2$, C$_2$H$_4$, and CO$_2$ were separated in the PorapLOT U column. Liquid products were analyzed using a high-performance liquid chromatograph equipped with an Aminex 87H column and a refractive index (RI) detector.

The partial current density of the gaseous products was calculated by using eq 3.

$$i_{\text{gas}} = \frac{X_{\text{gas}}nF}{A_{\text{geo}}RT}$$

(3)

where $i_{\text{gas}}$ is the partial current density of gases, $X_{\text{gas}}$ is the mole fraction of the gases, $f_i$ is the flow rate of the effluent gas in L/s, n is the number of electrons, F is the Faraday constant, P is the atmospheric pressure, $A_{\text{geo}}$ is the geometric area surface of the electrode, R is the universal gas constant, and T is the temperature (in Kelvin). The mole fraction, $X_{\text{CO}}$, is calculated by using eq 4.

$$X_{\text{CO}} = \frac{\text{peak area of CO in mGC} \times \% \text{ of CO in standard gas mixture}}{\text{peak area of standard CO from the mGC} \times 100}$$

(4)

Faradaic efficiency (FE) was calculated by using eq 5.

$$\text{FE}_{\text{CO}} = \frac{I_{\text{FE}}}{I_{\text{th}}} \times 100\%$$

(5)

where $I_{\text{CO}}$ is the FE of CO and $I_{\text{th}}$ is the total current density. The FE of the liquid products was calculated by using eq 6.

$$\text{FE} = \frac{n_iFV}{I_{\text{th}}} \times 100\%$$

(6)

where $n_i$ is the number of electrons required to produce the $i_{th}$ product, $c_i$ is the concentration of the product obtained from HPLC analysis, F is the Faraday constant, V is the volume of the catholyte, and $I$ is the average current during CA measurements.

The turn-over frequency (TOF) calculation was performed by using the following formula:

$$\text{TOF} = \frac{Q \times FE}{nF \times m_x \times w_x \times M_x \times t}$$

(7)
where $Q$ is the total charge during chronocoulometry, $FE$ is the Faradaic efficiency, $F$ is the Faraday constant, $n$ is the number of electrons exchanged for the product formation, $m_\text{c}$ is the mass of the composite loaded on the GDE, $w_\text{c}$ is the weight % of the catalyst (e.g., Co for CoTPP) from XPS, $M_\text{c}$ is the molecular weight of the catalyst, and $t$ is the time used to express the TOF unit.

The rate of electrochemical CO and H$_2$ production is calculated by following the Arrhenius equation

$$\ln(j_i) = \ln K - \left(\frac{E_a}{R}\right)\frac{1}{T}$$

where $j_i$ is the partial current density of the reaction products, $K$ is a constant, $E_a$ is the activation energy, $R$ is the universal gas constant, and $T$ is the temperature.

### 2.3. Computational Details

A triple-$\zeta$Def2-TZVP$^{38}$ functional as implemented into Gaussian 16 Rev C.01 [4] in combination with an “ultrafine” grid size were used to optimize the structures of all compounds. This functional has been reported to perform the best for transition-metal-doped porphyrins considered in this study. $^{40}$ Solvation effects were included through the implicit SMD solvation model. $^{41}$ Redox potentials of proton-coupled electron-transfer (PCET) $^{42}$ steps were computed using the computational normal hydrogen electrode which uses H$_2$ in the gas phase as discussed in ref 43. Electron-transfer potentials were calculated using the effective absolute potential method which relies on the absolute potential of the SHE computed for the used computational setup. $^{44}$ $pK_a$ values were obtained using the iso descriemic method in combination with forminic acid as a reference compound and an appropriate scaling relationship to correct for the shortcomings of the implicit solvation model. Following an earlier work, eq 8 was used as correction. $^{45}$

$$pK_a(\text{exp}) = 0.49pK_a(\text{DFT}) + 3.2$$

Calculations of transition-state (TS) structures were performed using the quadratic synchronous transit (QST3) method. Owing to significant instabilities in the forces, this method failed for several structures. In these cases, manual potential energy surface (PES) scans along the relevant reaction coordinates were performed. Also, barrierless transitions were confirmed by PES scans. All reported TS structures were tested for the presence of exactly one imaginary frequency along the reaction coordinate. Ground states were assumed to be converged if no imaginary frequencies were present.

The CoTPP/MWCNT composite was estimated by a Co-porphyrin (CoTPP) molecular model. The most favorable spin states were determined through explicitly testing of all reasonable spin states. This small model system was chosen as it allows for the use of high-level DFT methods while retaining the central structural features of the active site of the catalyst. Moreover, using a molecular system we can easily model different oxidation states by charging the system. If we would use periodic boundary conditions, the charges could interact if the system is too small, resulting in wrong total energies. Potentials have been calculated based on an RHE reference at pH = 7. (For more information regarding computations, please refer to the Supporting Information.)

### 3. RESULTS AND DISCUSSION

#### 3.1. Electrocatalytic Activities of the Composite

To investigate product distribution on the CoTPP/MWCNT composite, electrolysis was performed in a flow cell at a constant potential, while CO$_2$ was continuously supplied in the backside of the GDE (Figure 1). The gas flow generates a convective force, helping CO$_2$ molecules to reach the electrode/electrolyte interface at the three-phase boundary where the electroreduction reaction occurs. In the investigated single-atom type electrocatalyst, Co$^{2+}$ atoms in the CoTPP molecule are isolated by tetraphenylporphine (TTP) ligands and are expected to function as the active sites. At sufficiently negative potentials, CO$_2$ molecules should adsorb on the Co centers and convert to CO, which is then released from the surface before further reduction, as will be shown later with DFT modeling. Catalytic activities of the composite have been evaluated by CA measurement for 2 h, and the results are presented in Figure 2.

Figure 2. Chronoamperometry curves for the CoTPP/MWCNT composite at four studied temperatures of (a) $-0.6$ V$_\text{RHE}$, (b) $-0.8$ V$_\text{RHE}$, and (c) $-1.0$ V$_\text{RHE}$.

The curves show two distinguishable features of the total current density ($j_T$), the increment over the time and the oscillation. Figure 2a shows that during the 2 h long measurements at $-0.6$ V$_\text{RHE}$, the activity of the composite remains rather steady except at 50 °C where the activity slightly increases, indicated by the increase in $j_T$ over time.

Applying more negative potentials enhances the overall catalytic activities of the composite as indicated by increasing $j_T$ for all studied temperatures presented in Figure 2b,c. At $-0.8$ V$_\text{RHE}$ (Figure 2b) and at $-1.0$ V$_\text{RHE}$ (Figure 2c), the $j_T$ increased over time in all four temperatures studied. This may occur either from demetallation of the molecular catalysts or increasing wettability of the electrode. Although it is hard to completely exclude demetallation of the catalysts as a possible reason for current density increment over time, there is an alternative explanation based on the scanning electron microscopy (SEM) images, X-ray diffraction (XRD) pattern, CV measurement, time vs mol % of product formation analysis, 1 h CA measurement on pristine MWCNTs at $-0.8$ V$_\text{RHE}$ in a CO$_2$ environment, and earlier studies on catalyst demetallation.

https://doi.org/10.1021/acsaem.2c02873
ACS Appl. Energy Mater. 2023, 6, 267–277
t. The SEM images of fresh and used electrodes (see Figure S2) show similar morphologies. The XRD analysis performed for a fresh and a used composite electrode, pristine CoTPP, and MWCNTs (Figure S3) can reveal structural changes in the used electrodes as shifting of the major 2θ peaks or change in the peak intensity ratios. In the XRD pattern of the composite, the 2θ peaks at 20 and 26.5° correspond to the main peak of CoTPP and MWCNT, respectively. The intensity ratio of these two peaks is 3.9 for both fresh and used electrodes. In addition to this, other 2θ values of the fresh and used composite electrodes show similar characteristics indicating that major changes do not take place on the catalyst under the selected reaction conditions. In case of catalyst demetallation, the most pronounced indication would be the decrease of current density over time due to increase of resistance. To investigate this, CV measurements were performed before and after the 2 h CA measurement at −0.8 V RHE (see Figure S4). In accordance with the CA measurements, the CV measurements show increasing current density after 2 h of CA. Demetallation may also cause a decrease in the formation of one of the products. As discussed below in more detail, only CO and H2 are detected as the major products, and their time versus mol % curves at −0.8 V RHE and −1.0 V RHE (see Figures S5 and S6) show that the formation of both products increase over time.

To investigate the contribution of MWCNTs on the overall electrode activity, 1 h CA measurement on pristine MWCNTs was carried out in the flow cell at −0.8 V RHE at the four studied temperatures (Figure 3). These measurements show a similar trend of current density increment over time. Earlier studies have proved that in the case of demetallation, catalysts promote the HER rather than eCO2R. For example, Hailiang et al. showed that due to demetallation of CoPC/MWCNT hybrid catalysts, the selectivity of CH3OH formation decreases from the FE of 44 to 0.6%, while the FE of H2 increases to 80%. In this case, current density also shows a decreasing trend. Thus, all experimental evidence and earlier findings support the above discussion and suggest that catalyst demetallation is unlikely.

The other possible reason for the current density increment is the increased wettability of the electrode resulting from gradual reduction of hydrophobicity of the electrode. The reduced hydrophobicity intrinsically causes perspiration of the electrolyte, which may contribute to increase the rate of hydrogen formation. Notably, electrode wettability strongly depends on the applied potential. This is also evident from the CA measurement at −1.0 V RHE where current density increment at −1.0 V RHE is higher than at −0.8 V RHE. Likewise, a higher temperature also contributes to an increase in the wettability of the electrode during the operation which is evident from the CA measurement at 50 °C in Figure 2 and also from previous studies. Hence, during the CA measurements, the deactivation of the composite is unlikely which underlines the stability of the composite under reducing conditions. The oscillation of jC is ascribed to the so-called bubble nucleation, growth, and detachment effect on the electrode surface. Whenever the produced gas bubbles occupy a fraction of the electrode surface, the bubbles block the electrode surface and induce a decrease of the current density. When the bubbles detach from the electrode surface, the current density starts to increase again.

The FE of a reaction product is the input and output charge balance during the constant potential electrolysis, while partial current density (jP) represents the amount of products after a certain duration. The FE and partial current densities of CO and H2 under different potentials and temperatures are presented in Figure 4. Figure 4a shows the FE of CO and H2 at −0.6 V RHE at the studied temperatures. The highest FE of CO (ca. 78%) is obtained at 20 °C, while the FE of H2 is ca. 17%. At 30 °C, the FE of CO slightly decreases to 63%, while the FE of H2 increases to ca. 35%. However, at 40 and 50 °C, the FE of H2 reaches 52 and 61%, respectively, while the FE of CO decreases to ca. 43 and 36%, respectively. These

![Figure 3. CA measurement at −0.8 V RHE on pristine MWCNTs in a CO2 environment.](https://doi.org/10.1021/acsaem.2c02873)

![Figure 4. FE and partial current densities for CO and H2 production on the CoTPP/MWCNT composite at four studied temperatures of (a) −0.6 V RHE, (b) −0.8 V RHE, and (c) −1.0 V RHE.](https://doi.org/10.1021/acsaem.2c02873)
changes in the product distribution clearly show that $\text{H}_2$ production is favored at the elevated temperatures. The influence of temperature is also observed in the partial current density for CO and $\text{H}_2$. At 20 °C, the CO partial current density ($j_{\text{CO}}$) is ca. $-0.18 \text{ mA/cm}^2$ and it increases to $-0.32 \text{ mA/cm}^2$ at 50 °C. Similarly, for $\text{H}_2$ formation, the partial current density ($j_{\text{H}_2}$) increases to $-0.58 \text{ mA/cm}^2$ at the highest studied temperature, showing 10 times higher value compared to its current density at the lowest studied temperature. This indicates that temperature plays an important role during the eCO$_2$R by also enhancing the reaction kinetics. It is noteworthy that neither other gaseous nor liquid products are detected at this potential.

The FE and $j_p$ of the reaction products at $-0.8 \text{ V}_{\text{RHE}}$ are presented in Figure 4b. At this potential, the FE of CO reaches 96% at 20 °C, indicating that the composite is very selective toward CO$_2$ to CO conversion under these conditions. However, the FE of CO decreases with increasing temperature also at this potential. Conversely, the FE of $\text{H}_2$ is ca. 7% at 20 °C but 26% at 50 °C. Only HCOO$^-$ is identified as the liquid product in low amounts as its FE is ca. 1%. Interestingly, $j_p$ values of both CO and $\text{H}_2$ increase with temperature. The $j_{\text{CO}}$ at 20 °C is ca. $-2 \text{ mA/cm}^2$, while at 50 °C, it increases to ca. $-11 \text{ mA/cm}^2$. The $j_{\text{H}_2}$ at 20 °C is ca. $-0.2 \text{ mA/cm}^2$, whereas at 50 °C, it is $-4.5 \text{ mA/cm}^2$. Thus, it is clear that the rate of the reaction for both products at this potential increase with increasing temperature even more than at $-0.6 \text{ V}_{\text{RHE}}$. Applying a more negative potential of $-1.0 \text{ V}_{\text{RHE}}$ also results in a higher FE and $j_p$. The FE and $j_p$ of the reaction products at $-1.0 \text{ V}_{\text{RHE}}$ are presented in Figure 4c. The FE of CO reaches 98% at 20 °C, confirming that the composite is highly selective for CO$_2$ to CO conversion also under these reaction conditions. Likewise, for the measurements at $-0.8 \text{ V}_{\text{RHE}}$, the selectivity toward CO formation decreases when the temperature increases from 20 °C to 50 °C. At the latter temperature, the FE of CO is ca. 72%. The opposite trend is observed for $\text{H}_2$ formation as the FE is ca. 6% at 20 °C and ca. 30% at 50 °C. For the only minor product, HCOO$^-$, the FE is still less than 1%. Furthermore, the $j_{\text{CO}}$ at the lowest studied temperature 20 °C is ca. $-9 \text{ mA/cm}^2$, while it increases to $-12 \text{ mA/cm}^2$ at the highest studied temperature. The $j_{\text{H}_2}$ at 20 °C is ca. $-0.6 \text{ mA/cm}^2$, while at 50 °C, it is $-5.2 \text{ mA/cm}^2$. The $j_{\text{H}_2}$ at $-1.0 \text{ V}_{\text{RHE}}$ is higher than the values at $-0.6 \text{ V}_{\text{RHE}}$ and $-0.8 \text{ V}_{\text{RHE}}$, indicating significant $j_{\text{H}_2}$ increases with increasing applied potential along with the temperature.

Interestingly, for all studied potentials, at 20 °C the FE of $\text{H}_2$ and $j_{\text{H}_2}$ are quite low, but with increasing temperature both are increasing. This trend of $\text{H}_2$ production suggests that at 20 °C, the HER presumably takes place on the MWNT site which is supported by the 1 h CA measurement on the MWNT electrode (see Figure 3) in a CO$_2$ environment, producing $\text{H}_2$ as only reaction product with a FE of ca. 100% at the four studied temperatures at $-0.8 \text{ V}_{\text{RHE}}$. This is consistent with the earlier studies where H$_2$O may function as an $\text{H}^+$ source. 23,35,36 DFT analysis also shows that HER is unlikely to proceed on the CO active sites due to a high activation barrier (for details, see Section 3.4). However, at elevated temperature a fraction of CO active sites may participate in $\text{H}_2$ production due to a decrease of the activation barrier. The opposite trend is observed for the CO generation which also suggests the CO participation for the HER at elevated temperatures.

In summary, the lower temperatures are favorable for eCO$_2$R, while higher temperatures promote the $\text{H}_2$ evolution reaction more. At more negative potentials, for example, $-0.8 \text{ V}_{\text{RHE}}$ and $-1.0 \text{ V}_{\text{RHE}}$, the composite is more selective for CO production at all studied temperatures, while at $-0.6 \text{ V}_{\text{RHE}}$ the selectivity for CO is quite low. The origin of these selectivity differences has been analyzed by DFT and discussed below in Section 3.4. Importantly, the temperature contributes to switching the selectivity between CO and $\text{H}_2$ formation which can be used to produce varying ratios of $\text{H}_2$/CO and is further discussed in the following section. The continuous increase of $j_{\text{CO}}$ from lower to higher temperatures suggests that direct supply of CO$_2$ behind the working electrode in these reaction conditions is sufficient. This is consistent with other works performed in flow-cell systems. 27,28,77

3.2. Electrochemical Cell-Dependent Product Distribution. The product distribution of eCO$_2$R strongly depends on the reaction conditions, which is seen by comparing our previous 37 and current studies. In the previous study, an H-cell was used for constant potential electrolysis with the same electrolyte and composite loading on the electrode and the same temperature and potential range, that is, under similar conditions to the current study. In the H-cell, CH$_3$OH and CO production is observed at all studied potentials and temperatures, whereas formate production is observed at $-0.8 \text{ V}_{\text{RHE}}$ and $-1.0 \text{ V}_{\text{RHE}}$ at all temperatures. Lower temperatures favor CH$_3$OH formation in all studied potentials, while conversely higher temperatures favored $\text{H}_2$ formation. CO formation is favorable at $-0.6 \text{ V}_{\text{RHE}}$ and $-0.8 \text{ V}_{\text{RHE}}$ in the temperature range 20–40 °C, while at $-1.0 \text{ V}_{\text{RHE}}$, it is only favorable at 20 °C. CH$_4$ formation is observed at $-1.0 \text{ V}_{\text{RHE}}$ at 40 °C and 50 °C temperatures only.

As described above, the product distribution clearly shows potential and temperature dependency but is also affected by the measurement arrangements. In the current study, the only liquid product is formate which is detected at $-0.8 \text{ V}_{\text{RHE}}$ and $-1.0 \text{ V}_{\text{RHE}}$ with a low FE<1%. CO and $\text{H}_2$ are detected at all temperatures and potentials as the major products, while CO is the main product except at the highest measured temperatures of 40 and 50 °C at $-0.6 \text{ V}_{\text{RHE}}$. The difference in the product distribution in comparison to the H-cell is suggested to rise due to the following reasons: (1) Mass transport. In the H-cell, a stagnant electrolyte layer is formed on the electrode surface because of absorption between the electrode and solvent molecules. Sparely dissolved CO$_2$ diffuses to the electrode surface, and CO$_2$ together with the generated intermediate products can stay a longer time in the vicinity of the electrode because of the stagnant layer. Hence, the intermediates can react further on the electrode. In contrast, in the flow cell, the continuously forced CO$_2$ gas flow removes the formed CO, regenerating the electrode surface, thus preventing potential further reactions. (2) In the H-cell electrode, the ionomer loading is 10 wt %, while in the flow cell, it is 30 wt %, and this hydrophobicity difference may cause the difference in CO$_2$ adsorption and coverage on the catalyst surface. 58 (3) The properties of the GDE, that is, porosity, surface area, PTFE loading, resistance, and so forth, are different in the H-cell and flow-cell electrodes prepared by using carbon cloth (GDL-CT, FuelCellsEtc) and carbon paper (Sigracet 25 BC, FuelCellEtc), respectively. The GDE properties can combinedly affect the mass transport, conductivity, and active surface area of the composite materials.

3.3. Evaluation of Temperature Influence on Syngas and Intrinsic Activity of the Composite. In case of syngas, the ratio between CO and $\text{H}_2$ is clearly dependent on the
studied temperature as shown in Figure 5a. Generally, syngas is the mixture of CO and H\textsubscript{2} which is the building block for many industrially important chemicals. In the case of the Fischer-Tropsch synthesis of chemicals and fuels, the desired syngas H\textsubscript{2}/CO ratio is 0.3-4.\textsuperscript{59} Figure 5a shows that at −0.6 V\textsubscript{RHE}, the H\textsubscript{2}/CO ratio is 0.3 at 20 °C but it increases to ca. 1.2 at 40 °C. The 1.2 H\textsubscript{2}/CO ratio is important for the synthesis of oxo-alcohols and aldehydes, for example, propionaldehyde, pentanols, 1-octanols, and many more.\textsuperscript{11,60} At 50 °C, it reaches ca. 1.70 which is close to the ratio 2 required for methanol synthesis from syngas. Hence, the increase of the ratio with temperature is a very important indication for achieving industrially important chemicals via eCO\textsubscript{2}R. The reaction temperature is undoubtedly controlling the formation of syngas also at the more negative potentials, although the syngas H\textsubscript{2}/CO ratio is much lower at these potentials than at −0.6 V\textsubscript{RHE}. At −0.8 V\textsubscript{RHE} the syngas ratio increases from ca. 0.1 at 20 °C and reaches a maximum of 0.4 at 50 °C. At −1.0 V\textsubscript{RHE} it increases from ca. 0.06 at 20 °C and reaches a maximum of 0.4 at 50 °C. Hence, for both −0.8 V\textsubscript{RHE} and −1.0 V\textsubscript{RHE} potentials, the H\textsubscript{2}/CO ratio at 50 °C could be utilized for the Fischer-Tropsch process.\textsuperscript{18}

For the industrial application of syngas, the mixture needs to be completely free from CO\textsubscript{2} gas. Such mixture could be achieved by cell engineering but is beyond the scope of the current study. Moreover, there is ongoing research on CO\textsubscript{2} separation from syngas, for example, absorption and desorption by ionic liquids.\textsuperscript{61}

The intrinsic catalytic activity of the composite is revealed by the TOF of CO, which is presented in Figure 5b for all applied potentials and temperatures. Importantly, it shows that the TOF of CO formation increases from 20 to 50 °C at −0.6 V\textsubscript{RHE} and −0.8 V\textsubscript{RHE}. However, at −1.0 V\textsubscript{RHE} it reaches the maximum at 40 °C and then drops at 50 °C. Thus, increasing TOF with increasing temperature indicates that the rate of the CO\textsubscript{2}-to-CO conversion is additionally influenced by the temperature. Also, the TOF of H\textsubscript{2} (see Figure S7) increases with temperature for all applied potentials, suggesting that the composite is more active for H\textsubscript{2} formation at elevated temperatures.

The production rates of CO and H\textsubscript{2} are presented in Figure 5c,d. It shows that with increasing temperature, production rates of both gases increase at −0.6 V\textsubscript{RHE} and −0.8 V\textsubscript{RHE}. However, at −1.0 V\textsubscript{RHE} the rate of CO formation increases from 20 to 40 °C, while that of H\textsubscript{2} increases from the lowest to the highest temperature. This similar trend is observed also for the TOF, FE, and \(j_p\) of CO and H\textsubscript{2} formation. The production rates of CO at −1.0 V\textsubscript{RHE} at 20 and 40 °C are comparable with the most active nanoporous Au catalyst reported (see Table S1).

Figure 5. Temperature-controlled electrocatalytic activities of the composite. (a) Ratio of H\textsubscript{2} and CO in the produced syngas, (b) TOF of CO, (c) CO production rate (\(\mu\)mol/cm\textsuperscript{2} h\textsuperscript{−1}\)), and (d) H\textsubscript{2} production rate (\(\mu\)mol/cm\textsuperscript{2} h\textsuperscript{−1}\)) for all studied potentials and temperatures.

Figure 6. Gibbs free-energy profiles of (a) HER and (b) eCO\textsubscript{2}R on Co(0)TPP (starting point) at pH = 7; potentials are versus RHE.

https://doi.org/10.1021/acsaem.2c02873
ACS Appl. Energy Mater. 2023, 6, 267−277
The influence of temperature on the electrochemical reaction rates of CO and H\textsubscript{2} formation is evaluated by Arrhenius activation energy (E\textsubscript{a}) calculation. In relation to this, potential vs apparent E\textsubscript{a} (Figure S9) was calculated to understand the influence of potential on the energy barrier for the rate-determining step of CO and H\textsubscript{2}. When comparing the CO (Figure S8a) and H\textsubscript{2} formation (Figure S8b) rates, temperature appears to influence less in the former. The E\textsubscript{a} for H\textsubscript{2} formation is 70.2 kJ/mol at −0.6 V\textsubscript{RHE}, 88.4 kJ/mol at −0.8 V\textsubscript{RHE}, and 58.0 kJ/mol at −1.0 V\textsubscript{RHE}. A high E\textsubscript{a} for H\textsubscript{2} formation at all studied potentials (see Figure S9) is expected in a CO\textsubscript{2} environment as it depends on the magnitude of the energy barrier for the rate-limiting step and H\textsubscript{2}O molecule reorganization.

For CO formation, E\textsubscript{a} is 14.2 kJ/mol at −0.6 V\textsubscript{RHE}, 43.0 kJ/mol at −0.8 V\textsubscript{RHE}, and 10.8 kJ/mol at −1.0 V\textsubscript{RHE}. The highest E\textsubscript{a} of CO ca 43 kJ mol\textsuperscript{−1} is observed for −0.8 V\textsubscript{RHE} and considering this, the rate-limiting step is postulated to involve a chemical step.\textsuperscript{64} The lowest E\textsubscript{a} at −1.0 V\textsubscript{RHE} is understandable from the DFT results (see Figure 6b), indicating that the reaction path is barrierless at this potential. However, the explanation for E\textsubscript{a} at −0.6 V\textsubscript{RHE} remains currently unclear.

### 3.4. Understanding CO Formation Selectivity Trends by DFT

The experimental results show that the CoTPP/MWCNT composite is highly selective for CO production at more negative potentials. Furthermore, the H\textsubscript{2}/CO ratio of syngas is increased when the temperature increases. To understand the origin of these trends, DFT modeling is employed. In general, the eCO\textsubscript{2}R starts with the direct adsorption of CO\textsubscript{2} at the catalyst surface from where it is transformed to either CO or more reduced products through stepwise protonation.\textsuperscript{65} An alternative to this direct reduction path is the reaction of CO\textsubscript{2} with a cobalt hydride (Co−H) intermediate to form formic acid.\textsuperscript{66}

The cobalt hydride is typically formed through the discharge of a proton or H\textsubscript{2}O to form a Co−H hydride (Volmer reaction). Naturally, this intermediate can also react further to produce hydrogen through the reaction with a second proton or water molecule (Heyrovsky reaction). Due to the very low concentration of H\textsubscript{2}O\textsuperscript{+} of only 10\textsuperscript{−7} mol/L under the reaction conditions (pH = 7), H\textsubscript{2}O is the most likely proton source under experimental conditions.\textsuperscript{57}

Depending on the potential, the reactions will proceed either over Co(II), Co(I), or Co(0). Co(I) is only relevant at the potential of −0.6 V\textsubscript{RHE}. With decreasing the potential to −0.8 V\textsubscript{RHE} and −1.0 V\textsubscript{RHE}, Co(I) can be further reduced to Co(0). Interestingly, Co(II)−H is not stable under the reaction conditions. This can be realized by the comparison of the HER and the CO\textsubscript{2}RR on Co(II). The Volmer reaction to form a Co(II)−H species through a PCET requires a redox potential of −0.84 V\textsubscript{RHE}. Thus, it cannot occur in the potential range where Co(II) would be stable. Furthermore, the formed hydride is a very strong acid with a pKa of −14 and will therefore readily dissociate into the bare CoTPP complex and a proton. The presence of a strongly acidic hydride might appear surprising at first glance. However, it is well established in organometallic chemistry that pKa of a hydride can be comparable to that of strong acids in the presence of strongly electron-withdrawing ligands or metal centers in higher oxidation states.\textsuperscript{68} Accordingly, any follow up reactions to form H\textsubscript{2} or formic acid are blocked. Similarly, CO\textsubscript{2} is also unstable on Co(II)TPP and readily desorbs since it cannot provide the necessary electrons to convert linear CO\textsubscript{2} to a bent Co−COO carbonylate intermediate. Hence, both the eCO\textsubscript{2}R and the HER are blocked on Co(II)TPP.

On Co(I)TPP, the redox potential for forming the Co(I)−H hydride is −0.61 V\textsubscript{RHE} and thus, in principle, feasible at all considered potentials. However, this reaction is, identical to Co(II), blocked by the rather high acidity of the “hydride” proton, which with a pKa of 4 will readily dissociate under reaction conditions, for example, a pH of approximately 7. Note that proton-transfer reactions are subject to only a very minor activation barrier of the order of 0.1 eV\textsuperscript{69} and thus can be expected to outcompete any follow-up reaction. Furthermore, the reaction is subject to a very high activation barrier of 1.7 eV for forming Co(I)−H from water (Figure S1), thus effectively blocking the HER and the eCO\textsubscript{2}R to formic acid through the hydride route.

Co(0) finally forms with a pKa of 10, a stable hydride under the reaction conditions. Forming this hydride requires a potential of −0.36 V\textsubscript{RHE} (Figure 6a) and can therefore be easily facilitated at the potentials of −0.8 V\textsubscript{RHE} and −1.0 V\textsubscript{RHE} where at higher temperatures significant amounts of H\textsubscript{2} are observed. However, this reaction is again blocked by a very high activation barrier of 1.4 eV for water as a proton source (Figure 6a). It must be cautioned that this activation can be subject to significant errors of ca. ±500 meV, owing to neglecting of explicit water molecules in the model. Furthermore, H\textsubscript{2}O\textsuperscript{+} is a more reactive proton source\textsuperscript{67} and it is available in very small concentrations. Thus, the formation of Co(0)−H may become feasible at rather low reaction rates. This in turn would, in line with experimental evidence (see Figure 4b,c), enable the formation of minor amounts of hydrogen. Naturally, also increasing the temperature will increase the rate of hydrogen formation further. Additionally, the Co(0)−H path for the eCO\textsubscript{2}R to form formic acid is, owing to the consistently low activation barrier,\textsuperscript{70} in principle, also possible but is unlikely, owing to the very low concentrations of CO\textsubscript{2} in the electrolyte.

The unfavorable formation of hydrides leaves the direct reaction of CO\textsubscript{2} with cobalt as the only viable reaction route. This reaction is initialized by the direct adsorption of CO\textsubscript{2} at the active site to form a Co−COO carboxylate (Figure 6b). As a result of the significant loss of entropy when binding a gas phase CO\textsubscript{2} to the catalysts, the direct CO\textsubscript{2} adsorption on Co(0)TPP and Co(1)TPP are found to form Co−COO bonds which are slightly endergonic by +0.20 and +0.45 eV, respectively (Figure 6b for Co(0) and Figure S1 for Co(1)). Hence, increasing the temperature will increase the entropy costs for this reaction even further and thus render this reaction step less likely. Thus, the selectivity will, in line with experimental evidence, shift toward increased hydrogen in the gas mixture (see Figure 4a,b,c). The reactions are subject to rather high activation barriers of +0.77 eV for Co(0)TPP and +0.67 eV for Co(1)TPP (TS2 in Figures 6b and TS S1). This high barrier is most likely due to the energy cost associated with bending of the linear CO\textsubscript{2} molecule in a carboxylate-like geometry with an O−C−O bond angle of roughly 130°. Carboxylate formation is followed by the protonation to Co−COOH. Again, water serves as a proton source for this reaction. The redox potential of the reaction is −0.60 V\textsubscript{RHE} on Co(1)TPP. However, the obtained acid has a pKa of only 4 and thus readily dissociates in the pH-neutral electrolyte. Moreover, the PCET reduction of Co(1)TPP to form Co(0)−COOH requires a comparable potential of −0.57 V\textsubscript{RHE}. This Co(0)−COOH species has a high pKa value of 10 which renders it stable toward the deprotonation under reaction.
conditions. Thus, the only route forward for Co(I)–COO is the reduction through the addition of an electron, leading to the Co(0)–COO route which requires a potential of \(-0.71 \ V_{\text{RHE}}\). It is important to note that Co–COOH is a crucial intermediate in the pathway since two different products can be obtained from it. A second proton can electrochemically attack the OH group in Co(0)-C-COOH and eliminate the water molecule, leading to the formation of Co(0)–CO. The reaction requires a potential of only \(-0.22 \ V\) and is surprisingly also barrierless after activation through protonation as evident from the PES scan (Figure 6b). Considering a moderately strong binding energy, CO can be released as the final product, for example, the final release is endergonic by 0.46 eV. Again, the second possible pathway is the attack of the proton on the carbon atom of Co–COOH in an electrochemical reaction on Co(0)TPP. This would result in the breaking of the Co(0)-C bond and the release of HCOOH. Energetically, this reaction is highly favorable and requires a redox potential of +0.27 eV. However, since the carbon atom in the Co(0)-COOH intermediate is electron poor and additionally a Co(0)-C bond also needs to be broken, the proton attack will result in a significant energy barrier. This in turn renders the formation of formic acid, in line with the experimental evidence, unfavorable.

Overall, the high selectivity toward CO₂ reduction to form CO is the result of the very high activation barriers associated with the formation of the Co(0)–H hydride intermediates which could facilitate hydrogen evolution, while CO₂ adsorption, which is the rate-determining step toward CO formation, requires a significantly lower barrier of the order of 0.7 eV. In line with experimental evidence, our computations suggest an onset potential of \(-0.6 \ V_{\text{RHE}}\). Thus, CO₂ reduction would in principle be possible at all experimentally considered potentials. However, in reality, it is blocked by the almost complete absence of Co(I) and Co(0) at the lowest overpotential of \(-0.6 \ V_{\text{RHE}}\) where the experiment also shows negligible partial current densities (Figure 4a). For \(-0.8 \ V_{\text{RHE}}\) and \(-1.0 \ V_{\text{RHE}}\) the active Co(0)TPP complex is finally available, resulting in the experimentally observed formation of significant amounts of CO (Figure 4b,c).

4. CONCLUSIONS

In this work, we have investigated the temperature influence on eCO₂R on a CoTPP/MWNT composite in a flow cell and discovered that temperature can be a parameter which controls the ratio of H₂ and CO in the formed syngas. This finding is beneficial for industrial applications where different H₂/CO ratios are required to produce various important chemicals and fuels. Importantly, when the temperature is increased, the selectivity for CO formation decreased accompanied with increased selectivity for H₂ at all studied potentials. Although the selectivity for CO production decreases with increasing temperature, the partial current density increases, suggesting that higher temperatures accelerate the CO production by increasing the reaction rate. In addition to the experimental work, DFT modeling has been used to find the origin of high selectivity of the composite for CO formation in these experimental conditions and identifying the key oxidation states of the active site, for example, at \(-0.6 \ V_{\text{RHE}}\) the mostly dominating active site is Co²⁺ but at \(-0.8\) and \(-1.0 \ V_{\text{RHE}}\) it is Co⁰. We believe that this investigation of the temperature-controlled syngas production would contribute an alternate way for the production of syngas.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsaem.2c02873.

Computation details, electrode preparation, SEM analysis, XRD analysis, CV analysis, product distribution with time, turnover frequency for H₂, CO production-rate comparison data, ln(j/p) versus time graph, and potential versus activation energy graph (PDF)

AUTHOR INFORMATION

Corresponding Author

Tanja Kallio — Department of Chemistry and Materials Science, Aalto University School of Chemical Engineering, Espoo FI-00076 AALTO, Finland; orcid.org/0000-0001-6671-8582; Email: tanja.kallio@aalto.fi

Authors

M. Noor Hossain — Department of Chemistry and Materials Science, Aalto University School of Chemical Engineering, Espoo FI-00076 AALTO, Finland
Reza Khakpour — Department of Chemistry and Materials Science, Aalto University School of Chemical Engineering, Espoo FI-00076 AALTO, Finland
Michael Busch — Department of Chemistry and Materials Science, Aalto University School of Chemical Engineering, Espoo FI-00076 AALTO, Finland; Present Address: Institute of Theoretical Chemistry, Ulm University, Albert-Einstein-Allee 11, D-89081 Ulm, Germany
Milla Suominen — Department of Chemistry and Materials Science, Aalto University School of Chemical Engineering, Espoo FI-00076 AALTO, Finland
Kari Laasonen — Department of Chemistry and Materials Science, Aalto University School of Chemical Engineering, Espoo FI-00076 AALTO, Finland; orcid.org/0000-0002-4419-7824

Complete contact information is available at: https://pubs.acs.org/10.1021/acsaem.2c02873

Author Contributions

M.N.H.: Conceptualization, methodology, investigation, formal analysis, and writing—original draft, review, and editing; R.K.: Methodology, investigation, formal analysis, and writing—review and editing; M.B.: Supervision and writing; A.K.: Methodology, supervision, funding acquisition, and writing—review and editing; M.S.: Writing—review and editing; K.L.: Supervision and writing—review and editing; T.K.: Methodology, supervision, funding acquisition, and writing—review and editing.

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

We are thankful for the financial support from European Union under the project ELCOREL ITN Horizon 2020 (721624-ELCOREL), Academy of Finland (Profi-5), and Jane and Aatos Erkko Foundation (USVA), Finland. We also acknowledge the generous computer resources provided by the Finnish National Supercomputer Centre CSC and Aalto University Raw Materials Infrastructure (RAMI) and OtaNano Nanoscopy Center.
REFERENCES


