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# The combined effect of temperature and pressure on the electrochemical CO<sub>2</sub> reduction on Cu

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## ABSTRACT

Research into the electrochemical CO<sub>2</sub> reduction (CO<sub>2</sub>RR) is often performed at ambient conditions. However, these are probably not the most industrial relevant conditions, eventually leading to insufficient insights. Using a high-pressure high-temperature electrochemical cell, we have investigated the combined effect of temperature and pressure on the CO<sub>2</sub>RR on Cu. We show that temperature has a larger effect on the selectivity than pressure and increasing temperature up to 75 °C can increase the C<sub>2</sub><sup>+</sup> formation. However, stability becomes a bigger issue at higher temperatures, while higher pressures increase the stability as evidenced by the time dependence of the ethylene selectivity. Moreover, elevated pressures can suppress the competing hydrogen evolution reaction and increase both activity and selectivity of CO<sub>2</sub>RR significantly, even at a few bars of pressure, as long as these pressures are CO<sub>2</sub> pressures and not reached by an inert gas.

## 1. Introduction

Electrochemical CO<sub>2</sub> reduction (CO<sub>2</sub>RR) provides a promising approach to convert CO<sub>2</sub> into valuable fuels and chemicals using renewable electricity [1–4]. Of all possible catalysts for this reaction, copper is the only one capable of forming C–C bonds with high Faradaic efficiencies [5–7]. Many factors can influence the CO<sub>2</sub>RR process on copper and for factors such as electrolyte composition [8–11] and electrode potential [7,10–13], their effects have been well explored. Recently, there has been an increasing focus on the impact of pressure [14–21] and temperature [21–29], which are key parameters that can enhance both the selectivity and activity of CO<sub>2</sub>RR [14–16,19–23,28,29], unlock new reaction pathways [17,30,31], and improve integration with other industrial processes [17,32–34]. Temperature and pressure are particularly significant because industrial electrolyzers typically operate at higher temperatures [35], to enhance kinetics but also to make use of thermal losses [36–39] or the use of hot feedstocks. However, much of the electrochemical research, including CO<sub>2</sub> reduction studies, is still conducted under conditions of ambient temperature and pressure.

Recently, we have shown the effect of temperature on the activity and selectivity of CO<sub>2</sub>RR on Cu at ambient pressure [23]. Initially, temperature has a positive effect on both the selectivity and activity towards C<sub>2</sub><sup>+</sup> products but above an optimum temperature at around 48

°C, the selectivity towards C<sub>2</sub><sup>+</sup> products and the CO<sub>2</sub>RR activity decreases with further increasing temperature. This optimum has been recently confirmed by Brandao et al. [29] Ahn et al. [25] observed an optimum in C<sub>2</sub><sup>+</sup> products as well, although at lower temperatures. Moreover, all abovementioned papers and the earlier work of Hori [26] observed that CH<sub>4</sub> selectivity decreases with increasing temperature, while HER selectivity increases.

Already in the nineties, Hara et al. have studied the effect of pressure on several metal electrocatalysts for CO<sub>2</sub>RR [14], including Cu [15]. At 1 bar, they observed only HER, but they were able to significantly suppress H<sub>2</sub> formation by increasing the pressure. Moreover, hydrocarbon selectivity was observed to have an optimum around 40 bar. Interestingly, the electrocatalysts studied hardly produced CO and also C<sub>2</sub><sup>+</sup> products were limited to maximum 7 % selectivity, whereas formic acid selectivity slowly increased with pressure. These older observations are in contrast to recent studies where formate formation dominates at elevated pressures, not only on Cu [16,17,40] but also on Au, Ag and Sn [16]. Interestingly, Girichandran et al. [17] observe 2-propanol as a product at elevated pressures. This product is not found under ambient conditions for CO<sub>2</sub>RR, as 1-propanol is the preferred product [41]. Proietto et al. [42] show that the FE towards CO<sub>2</sub>RR does not always rise monotonously but can also show an optimum with pressure, at least for Sn and Ag.

The C<sub>2</sub><sup>+</sup> formation on Cu at elevated pressures is not studied in

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much detail, although it seems that the dominant C2+ product can change from ethylene into ethanol [20] or acetate [19] at high pressures. Moreover, the C2+ selectivity has sometimes been observed to show an optimum with CO<sub>2</sub> pressure [17]. However, other studies have performed CO reduction at elevated pressures instead of CO<sub>2</sub>RR and observe a constant increase in C2+ selectivity and activity with increasing pressure [43,44]. This is attributed to an increase in CO coverage, which facilitates CO dimerization and thus the formation of C2+ products. Elevated temperatures have been shown to increase the CO coverage during CO<sub>2</sub>RR, at least at ambient pressures and temperatures up to 45 °C [23,29]. This illustrates that for CO<sub>2</sub> reduction at elevated pressures there might be a competition between C2+ products and HCOOH formation, which could be influenced with temperature.

Recently, we have developed a special electrochemical system in which both pressure and temperature can be regulated up to 200 °C and 140 bar [45]. On Au electrodes, we have shown that by increasing the pressure, selectivity for CO<sub>2</sub>RR was significantly improved, especially at elevated temperatures. Moreover, both temperature and pressure can increase the partial current density dramatically. Our system provides the opportunity to study the combined effect of pressure and temperature. In this study, we apply this opportunity to Cu electrodes and show that temperature can improve the selectivity of the traditional CO dimerization mechanism at temperatures up to 75 °C, and that pressure helps to keep the system stable at these temperatures. Elevated pressures can reduce HER activity and selectivity dramatically, even at pressures of only a few bar. However, it is not pressure itself, but specifically CO<sub>2</sub> pressure that enhances CO<sub>2</sub> reduction and C—C coupling.

## 2. Methods

### 2.1. Chemicals

The electrolyte was prepared from KHCO<sub>3</sub> (99.95 %, Sigma-Aldrich) with Milli-Q water ( $\geq 18.2$  MΩcm, TOC < 5 ppb) and stored with Chelex (100 sodium form, Sigma-Aldrich) to clean the electrolyte from any metal impurities [46]. H<sub>2</sub>SO<sub>4</sub> (95–98 %, Sigma-Aldrich), H<sub>2</sub>O<sub>2</sub> (35 %, Merck) and KMnO<sub>4</sub> (99 %, Sigma-Aldrich) were used to clean the cells. Ar (5.0 purity, Linde) and CO<sub>2</sub> (4.5 purity, Linde) were used for purging the electrolytes.

### 2.2. High-temperature high-pressure electrochemical cell

We used an adapted high-pressure, high-temperature cell from Parr instruments; this cell has been described in detail in a recent paper [45] and is shown in Figure S1. The cell was designed to work up to 140 bar and 200 °C. The cell was used in semi-continuous mode, where either CO<sub>2</sub> or a mixture of CO<sub>2</sub> and Ar were constantly purged through the cell. The outlet of the cell was coupled to a condenser and a cooler (Cool-Care, Van der Heijden Labortechnik GmbH) to keep all volatile products in the liquid phase in the setup. The outlet gas was analyzed every 6 min with a Micro-GC from Agilent with two TCD detectors. A combination of a MS6A and a CP-PORABOND Q column was used to detect H<sub>2</sub>, CO and CH<sub>4</sub>, and an Al<sub>2</sub>O<sub>3</sub> column was used to detect the C2 and C3 hydrocarbons. The pressure in the cell was maintained by controlling both the inlet and outlet flow with flow controllers (SLA5850, Brooks Instruments), while for the mixed gas experiments the flow was also checked with a flow meter (Defender 530+, Mesalabs). The temperature was controlled with an electric heating mantle. A homemade two-compartment PEEK cup was used inside the stainless steel chamber and all metal parts from the thermocouple and reference electrode were covered by Teflon tape. Moreover, the lid of the vessel was covered with Viton rubber to prevent any contamination from the stainless steel.

### 2.3. General electrochemistry

The homemade PEEK cup was cleaned prior to experiments by

storing in permanganate solution overnight (0.5 M H<sub>2</sub>SO<sub>4</sub>, 1 g/L KMnO<sub>4</sub>). Afterwards, the cell was rinsed, submerged in a diluted mixture of H<sub>2</sub>SO<sub>4</sub> and H<sub>2</sub>O<sub>2</sub> to remove any traces of MnO<sub>4</sub> and MnO<sub>2</sub>, rinsed again and boiled three times with Milli-Q water. A 1 mm thick Cu wire (99.99 %, Mateck) was used as working electrode. Before experiments, the working electrode was electropolished in 85 % H<sub>3</sub>PO<sub>4</sub> (Suprapure, Merck) by applying +3 V versus a graphite counter electrode for 20 s and subsequently rinsed with Milli-Q water. Due to the large difference in current at the various conditions used, 4 different sized wires were used to ensure the current stays between 10 mA (for product detection) and 200 mA (for stability of the system) during the experiment. The areas of the different working electrodes were normalized using their double layer capacitance, measured from −0.3 to 0.2 V vs SHE at scan rates from 200 to 1400 mV/s [47]. The areas of the electrodes can be found in Table S1.

During experiments, a DSA counter electrode (Magnet) was separated from the working and reference electrode by a PiperION membrane (A40, Versogen) [48,49]. The experiments were performed with a Gamry Interface 1010B potentiostat and current interrupt [50] was used to correct for the Ohmic drop. Experiments were performed for 60 or 90 min at constant potential. Gaseous products were determined every 6 min with GC as described above, while liquid products were determined afterwards by sampling the 65 ml catholyte and analyzing this using high performance liquid chromatography (HPLC, Shimadzu) with an Aminex HPX-87H column (Bio-rad). During experiments the pressure was maintained within an error of 0.1 bar, while temperature was maintained within an error of 2 °C. Before reaching the conditions used for the experiment (which typically takes up to an hour), the working electrode was always under potential control at a potential of −0.8 V vs Ag/AgCl.

### 2.4. Reference electrode corrections

In this study, all potentials are reported vs SHE(T). During experiments, an Ag/AgCl electrode (Ultradeg, 0.1 M KCl, Corr Instruments) was used as a reference electrode. However, with changing temperature, one should make several corrections to convert the potential from Ag/AgCl to SHE. The Nernst equation is used to determine the shift of the Ag/AgCl reference electrode against SHE:

$$E_{\text{Ag/AgCl}} = E_{\text{Ag/AgCl}}^0(T) - \frac{\ln(10) * RT}{nF} * \log(a_{\text{Cl}^-}(T)) \quad (7.1)$$

with  $E^0$  being the standard potential,  $R$  the gas constant,  $T$  the temperature,  $n$  the number of electrons,  $F$  Faradays constant and  $a_{\text{Cl}^-}$  the activity of the chloride anion. The standard potential of Ag/AgCl vs SHE is a function of temperature and can be calculated using the following formula[51]:

$$E_{\text{Ag/AgCl}}^0(T) = -0.111658 + 0.011134 * T - 0.001757268 * T * \ln(T) \quad (7.2)$$

The temperature dependence of the activity of the chloride ions can be determined from a formula from Bogaerts et al.[52]:

$$a_{\text{Cl}^-}(T) = \exp \left[ \log(a_{\text{Cl}^-}(25^\circ\text{C})) - \frac{\sqrt{I_{\text{Cl}^-}}}{1 + \sqrt{I_{\text{Cl}^-}}} * (A_T^T - A_T^{25^\circ\text{C}}) \right] \quad (7.3)$$

where  $a_{\text{Cl}^-}$  is the activity of the chloride anion at room temperature [53],  $I_{\text{Cl}^-}$  is the ionic strength of the chloride anion, and  $A_T$  is the Debye-Huckel parameter [52].

With these formulas, the shift between an Ag/AgCl reference and SHE could be determined at different temperatures as shown in Table S2. Even though the SHE potential is defined as zero at all temperatures, there is a difference in potential between SHE(T) and SHE (25 °C) [54]. One could correct for this difference too, but we decide to limit the corrections to the necessary corrections and therefore report on a SHE(T) scale. In previous studies we reported on the RHE(T) scale

[22–24,55], but in this study the potential could not be reported vs. RHE because the pH should be known to convert SHE to RHE. However, it was for us impossible to measure the exact pH of the electrolyte at elevated  $\text{CO}_2$  pressures and temperatures.

### 3. Results and Discussion

#### 3.1. Screening temperature, pressure and potential

As the research space increases tremendously when pressure and temperature are varied as parameters, we first screened the effect of these parameters at different potentials. We mainly focused on  $-1.4$  V vs SHE as shown in Figures 1 and S2. This potential corresponds to  $-1.0$  V vs RHE at ambient conditions in  $0.2$  M  $\text{KHCO}_3$ , a potential which is commonly used for electrochemical  $\text{CO}_2$  reduction on Cu. However, we also performed screening experiments at  $-1.3$  and  $-1.5$  V vs SHE (Figure S3 and S3), as the potential can change the effects of temperature and pressure [22,55]. Moreover, all experiments have been performed in  $0.2$  M  $\text{KHCO}_3$  because changing the electrolyte as well would increase the research space beyond reasonable measure.

Fig. 1 shows that temperature has a more significant effect on the selectivity (towards the major products hydrogen, carbon monoxide, formic acid, methane, ethylene, and ethanol) than pressure. With increasing temperature, more  $\text{H}_2$  is generated and the selectivity for all  $\text{CO}_2\text{RR}$  products decreases. With increasing pressure, this  $\text{H}_2$  enhancing effect of temperature can be suppressed and the  $\text{CO}_2\text{RR}$  stays active up to higher temperatures. At room temperature, there are only subtle effects on the selectivity, while the effect of pressure becomes more clear at  $50$  and  $75$  °C. These trends can also be observed at the other potentials. The exact values of the FE for the different products depends on the potential applied, but the overall trends for temperature and pressure seem independent of potential, at least in this potential window. Figure S2 shows the 2D maps for the minor products (methanol, acetate, ethylene glycol, acetaldehyde, propionaldehyde, propanol). In general, the trends are similar to the major products in Fig. 1, however the effect of pressure at room temperature is more clear, especially for methanol and acetate. Acetate and ethylene glycol seem to form even at very high

temperatures.

#### 3.2. Effect of temperature

The screening data shows that the most interesting parameters to study the  $\text{C}_2+$  formation in more detail are at  $-1.5$  V vs SHE at temperatures up to  $75$  °C. Therefore, these conditions have been studied in more detail with an improved setup in pure  $\text{CO}_2$ . Figures S5–7 show the effect of temperature on the activity and selectivity on Cu at different pressures. The trends at ambient pressure are comparable to the trends we reported before in a different setup [23], although not exactly the same. In general, HER selectivity increases with temperature, while the selectivity towards methane,  $\text{HCOOH}$  and  $\text{CO}$  decrease with increasing temperature. However, previously an optimum in  $\text{C}_2+$  formation was observed around  $50$  °C, which is now not the case at  $1$  bar but only at elevated pressures. In previous work we have discussed several contributions to this temperature optimum, of which the  $\text{CO}_2$  bulk concentration, which decreases with increasing temperature, is the main one affected by increasing the pressure.

To look into these differences in  $\text{C}_2+$  formation in more detail and illustrate more specifically the effect of temperature and time, Figs. 2 and S8 show the selectivity towards  $\text{C}_2\text{H}_4$  as a function of temperature and electrolysis time. Initially, at  $1$  bar, the effect of temperature is very similar to what we reported before [23]. There is an optimum in ethylene selectivity at ca.  $50$  °C. However, in the setup used in this work the Cu seems to be less stable than in the setup used previously, and with time the selectivity towards ethylene decreases, resulting in a negative trend with temperature over the entire experiment. The electrochemical cell used here is very different to the design used previously [23], for example the membrane, reference and electrolyte volume are different. Any of these aspects might influence the stability, but also the fact that the working electrode is now a wire instead of a disk. Additionally, the increased electrolyte concentration might also cause part of the differences in stability.

Pressure also influences the effect of temperature as at elevated pressures the FE towards ethylene decreases less with time compared to  $1$  bar. More interestingly, after  $20$  min, the selectivity towards ethylene

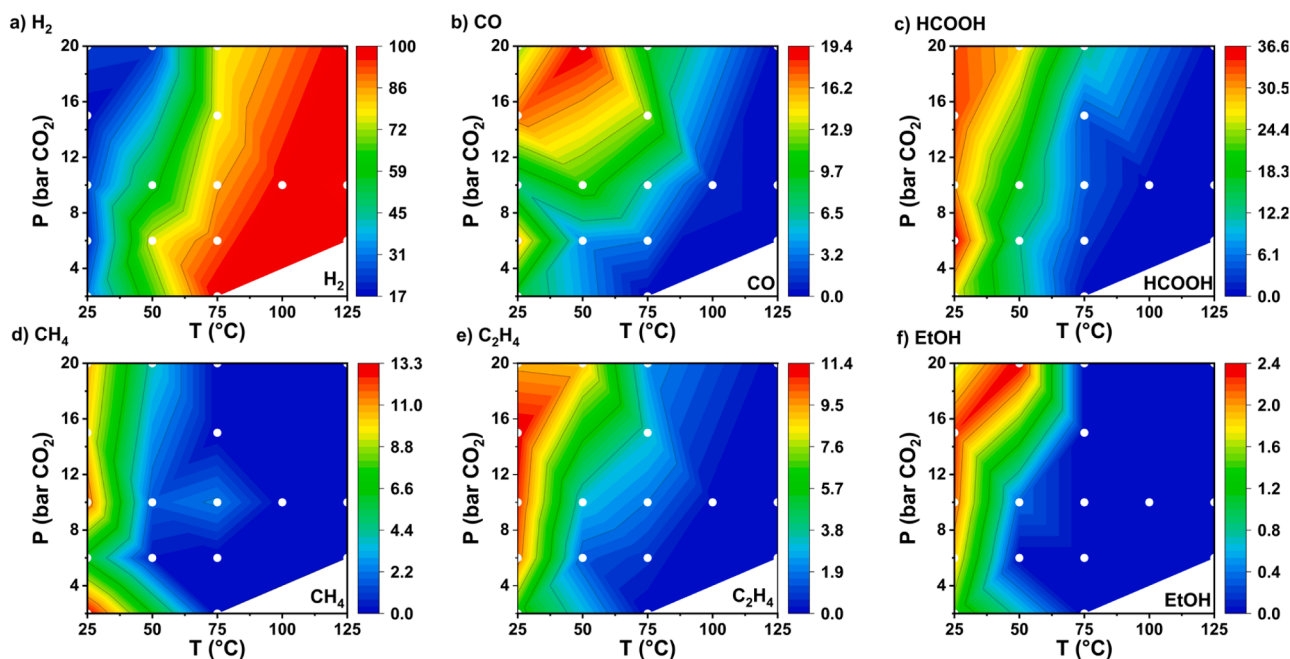


Fig. 1. Faradaic efficiency towards the main  $\text{CO}_2\text{RR}$  products as function of temperature and pressure at  $-1.4$  V vs SHE in  $0.2$  M  $\text{KHCO}_3$  with a 50/50 ratio of  $\text{CO}_2$  and Ar. The white dots give the measured data points and the maps are constructed via interpolation between these points. FE for a) hydrogen b) carbon monoxide c) formic acid d) methane e) ethylene and f) ethanol.

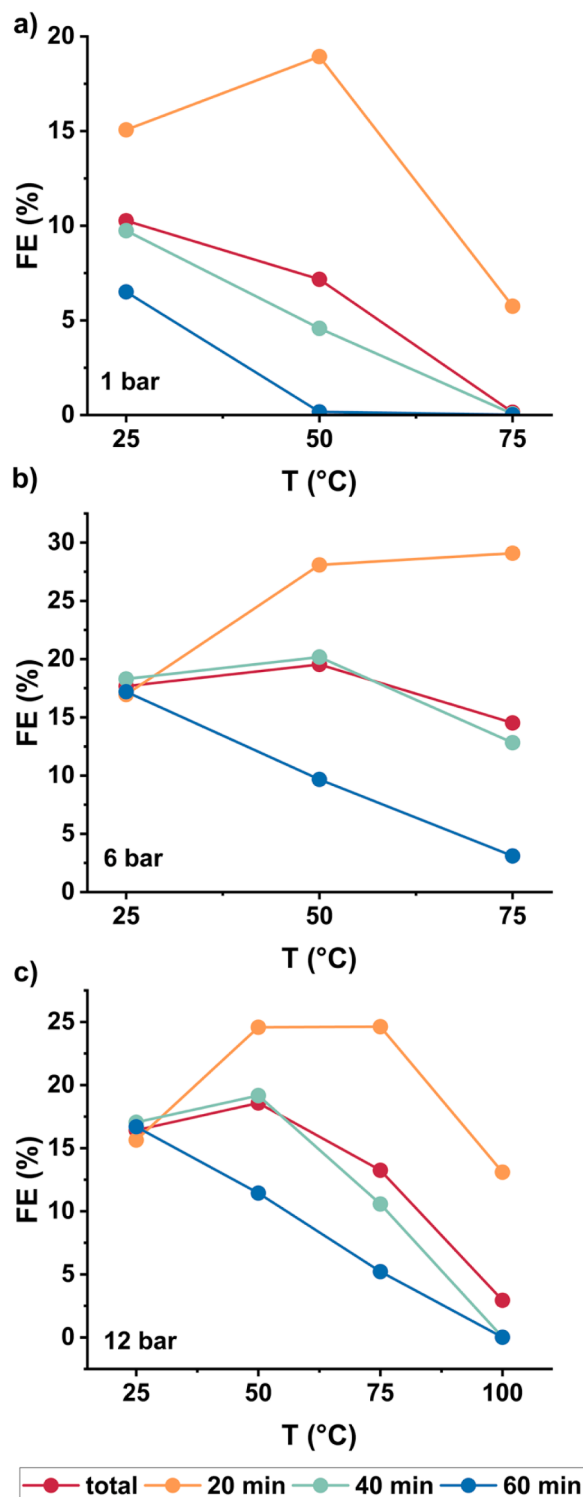


Fig. 2. Faradaic efficiency towards ethylene as function of temperature after different electrolysis times at  $-1.5$  V vs SHE in  $0.2$  M  $\text{KHCO}_3$  at a) 1 bar b) 6 bar and c) 12 bar.

remains high, even up to temperatures of  $75^\circ\text{C}$ . This shows that even a few bars of elevated pressure can enable the formation of  $\text{C}_2^+$  products at higher temperatures, most likely because the  $\text{CO}_2$  concentration in the electrolyte is maintained at a reasonable concentration (as shown by Figure S9), which could influence the CO coverage. However, a reasonable  $\text{CO}_2$  concentration in itself is not sufficient to maintain high FE towards ethylene. At  $100^\circ\text{C}$  and 12 bar, the  $\text{CO}_2$  bulk concentration is

higher than at  $75^\circ\text{C}$  and 6 bar, however the FE towards ethylene is lower.

We look in more detail at the effect of temperature and pressure on the stability of the Cu electrocatalyst in Fig. 3. This figure shows the selectivity towards ethylene at 40 and 60 min, plotted as a ratio compared to the initial FE at 20 min, as a descriptor for stability. This figure clearly shows that the ethylene selectivity on Cu becomes less stable with increasing temperature. We have shown that in the setup used in this study, no contamination such as Fe or Ni can be found that would explain the decrease in ethylene with time [31]. However, in previous study we have shown that the Cu surface deactivates at elevated temperatures, with the surface getting more smooth at elevated temperatures, which might cause this instability in ethylene formation [23]. Interestingly, increasing the pressure, at a given temperature, generally increases the stability of ethylene formation. To make  $\text{CO}_2\text{RR}$  on Cu feasible on an industrial scale, it is important to stabilize Cu at temperatures up to at least  $75^\circ\text{C}$ , which are reasonable industrial operation temperatures [56,57]. We hypothesize that the stabilization of Cu through pressure is due to the suppression of HER and an increase of the coverage of  $\text{CO}_2$  and/or CO. However, additional strategies are probably needed to obtain truly long term stability of the Cu catalyst at elevated temperatures. Some possible strategies may include pulsing of the potential or alloying the Cu electrode to make it more stable at high temperatures [55].

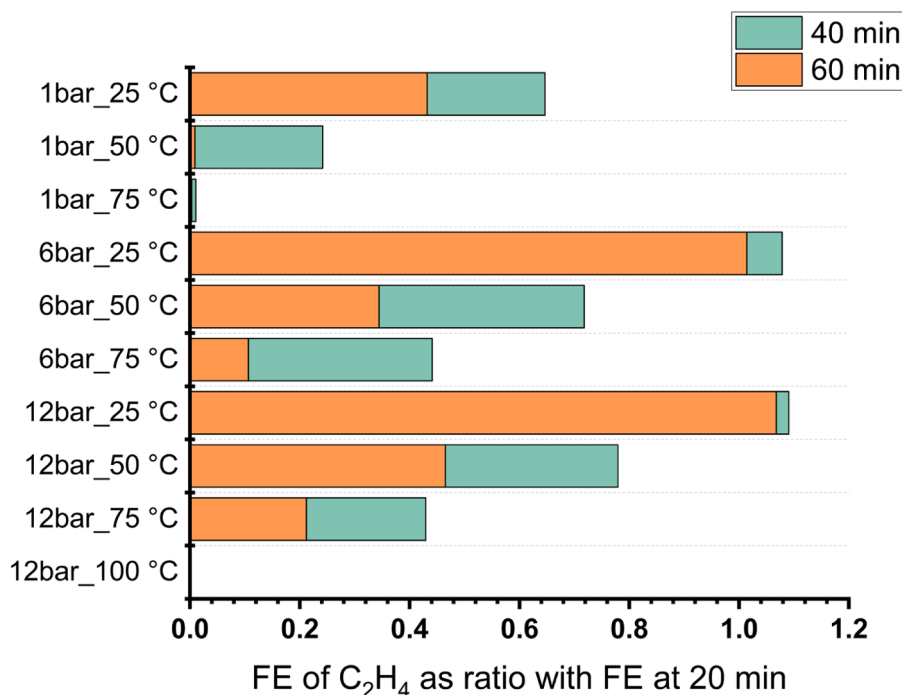
### 3.3. Effect of pressure

Pressure does not only influence the stability of the Cu catalyst or increase the operating temperature window, but can itself also affect the activity and selectivity of the  $\text{CO}_2\text{RR}$ . Fig. 1 showed that the effect of pressure on the selectivity is smaller than the effect of temperature, but it is not negligible. Fig. 4 shows the effect of pressure on the selectivity of the major products during  $\text{CO}_2\text{RR}$  at  $-1.5$  V vs SHE at room temperature, while the effect on the minor products is shown in Figure S23. The pressure has been increased both with  $\text{CO}_2$  and with Ar to be able to distinguish between the effect of pressure itself and of  $\text{CO}_2$  pressure specifically.

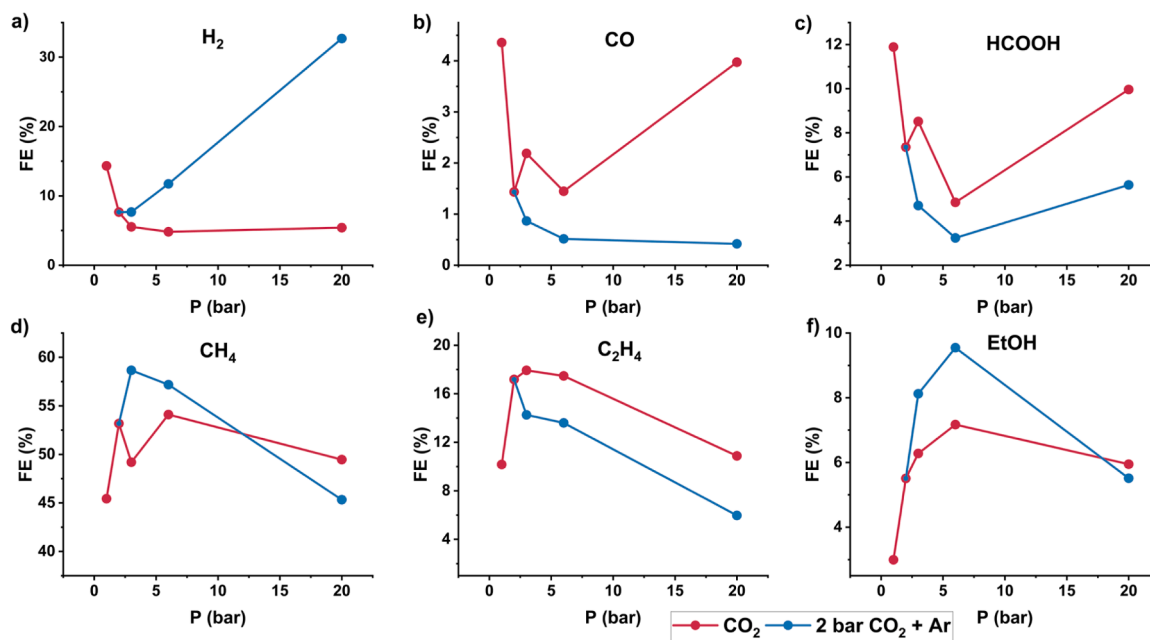
Figure 4 and S9 show that increasing  $\text{CO}_2$  pressure can decrease the selectivity towards  $\text{H}_2$  and that this effect already plateaus at a relatively low pressure of 3 and 6 bar at  $25^\circ\text{C}$  and  $75^\circ\text{C}$ , respectively. This same pressure is the optimum pressure for  $\text{C}_2\text{H}_4$  selectivity, and for  $\text{C}_2^+$  selectivity in general. At  $25^\circ\text{C}$ , the initial pressure increase from 1 to 3 bar leads to a rapid increase in  $\text{C}_2^+$  selectivity, but with further increasing pressure, the selectivity towards  $\text{C}_2^+$  slowly decreases in favor of the simple  $\text{C}_1$  products CO and HCOOH (although the selectivity of the minor  $\text{C}_2^+$  liquid products does keep increasing with increasing  $\text{CO}_2$  pressure). Even though the selectivity towards HCOOH increases slightly at high pressures, we never observe very selective  $\text{CO}_2\text{RR}$  towards HCOOH as described in some literature studies [16,17], although others did not observe this effect [42]. This difference could possibly be attributed to the differences in electrolyte pH [19], although Figure S24 indicates that simply a difference in electrolyte concentration between these studies does not explain the differences in HCOOH behavior with pressure. Possibly the observed differences could be ascribed to the electrode morphology as both references 16 and 17 use a copper foam working electrode, which could influence for example the local pH very differently than our Cu wire working electrode.

During CO reduction on Cu, a continuous increase in  $\text{C}_2^+$  selectivity with increasing pressure was observed [43,44], which was attributed to the increasing  $^*\text{CO}$  coverage. However, Fig. 4 shows this is not the case for  $\text{CO}_2$  reduction, especially at higher pressures of 20 bar. It would be interesting to compare  $\text{CO}_2$  and CO reduction at elevated pressures and temperatures but unfortunately, due to safety concerns, we are not able to perform CO reduction experiments at elevated pressures at this moment. Moreover, it would also be interesting to follow spectroscopically the  $^*\text{CO}$  coverage during  $\text{CO}_2\text{RR}$  at elevated pressures, which is





**Fig. 3.** Stability of ethylene formation as function of temperature and pressure. Stability of ethylene formation is here defined as the FE that remains after 40 and 60 min compared to the FE at 20 min at a given condition.



**Fig. 4.** Faradaic efficiency towards the main CO<sub>2</sub>RR products as function of CO<sub>2</sub> and total pressure at  $-1.5$  V vs SHE in  $0.2$  M KHCO<sub>3</sub> at  $25$  °C. Red lines are performed in pure CO<sub>2</sub> and blue lines are at  $2$  bar of CO<sub>2</sub> and pressure is increased with Ar. FE for a) hydrogen b) carbon monoxide c) methane d) formic acid e) ethylene and f) ethanol. Note that the (single) data point at  $3$  bar in b), c) and d) seems somewhat of a (random) outlier, but it does not interfere with the main trend.

unfortunately not possible with our current setup. However, from the product analysis, we hypothesize that the  $^*CO$  coverage does not continuously increase with increasing CO<sub>2</sub> pressure, which might be due to faster desorption of CO due to increased CO<sub>2</sub> adsorption [45,58]. This limit in  $^*CO$  coverage would explain the observed optimum in C<sub>2</sub>+ products at relatively low pressure and shift to simple C<sub>1</sub> products with higher pressure.

Fig. 4a demonstrates that the suppression of H<sub>2</sub> at elevated pressures is due to the increase in CO<sub>2</sub> pressure. When the pressure is increased

with Ar instead of CO<sub>2</sub>, the FE towards H<sub>2</sub> increases at  $25$  °C, although at elevated temperatures Ar pressure does suppress HER slightly (Figure S12a). This might be due to the very high initial HER selectivity of  $>90$  % or the improved stability, even under Ar pressure. Moreover, when Ar is used to increase the pressure, most CO<sub>2</sub>RR products show lower selectivities than in a pure CO<sub>2</sub> atmosphere, as shown in Figure 4 and S9. For ethanol and methane this is not the case at intermediate pressures of  $3$ – $6$  bar, but also for these products the activity is lower with Ar than in a pure CO<sub>2</sub> atmosphere, as shown in Figure S11 and S13.

Therefore the increase in selectivity for these two CO<sub>2</sub>RR products is due to a relatively smaller decrease in activity compared to for example ethylene. If the system is ideal and Ar would be completely innocent, it would be expected that increasing the Ar concentration while keeping the CO<sub>2</sub> partial pressure constant would have no impact at all as the adsorption and kinetics should not be affected. However, Figures 4 and S11 show that CO<sub>2</sub>RR performance decreases while HER is enhanced when the Ar pressure is increased. Therefore, the CO<sub>2</sub> partial pressure is not the only contributing factor, but there seems to be some dilution effect as well. However, it is impossible to completely decouple this dilution effect from the effect of the CO<sub>2</sub> partial pressure.

Figures 4 and S12 illustrate that the main reason for suppression of HER is not pressure itself, but CO<sub>2</sub> pressure, although Ar pressure might slightly help at elevated temperatures when HER is very active. We hypothesize that the HER suppression is due to competition of adsorption sites between CO<sub>2</sub>RR and HER intermediates, where elevated CO<sub>2</sub> pressures lead to a blockage of adsorption sites for HER [45].

#### 4. Conclusion

In this study we have investigated the combined effect of pressure and temperature on the electrochemical CO<sub>2</sub> reduction on Cu. We show that temperature has a larger effect on the selectivity than pressure, although the effect of the latter is not negligible. Increasing the temperature can increase the FE towards traditional C<sub>2</sub>+ products and increasing the pressure can broaden the temperature range in which Cu catalysts can operate, both by increasing the CO<sub>2</sub> available and the Cu stability. At ambient temperature, increased CO<sub>2</sub> pressure itself can significantly suppress the competing hydrogen evolution reaction, even at a few bar. It is important to make the distinction between pressure and CO<sub>2</sub> pressure, as these do not have the same effect on the CO<sub>2</sub>RR. For positive effects, it is in most cases crucial to increase the CO<sub>2</sub> (partial) pressure and not just the total pressure in the system.

This study has shown the importance of studying the effect of temperature and pressure. However, due to the large parameter space, more avenues should be explored, for example to the effect of other parameters such as electrolyte composition in combination with temperature and pressure. We also highlight the importance of studying the stability of the Cu electrocatalysts, especially at elevated temperatures, beyond simply the stability of ethylene formation, as was done here. Alloying Cu with more stable metals or pulsed experiments are examples of possible ways to increase both the stability of ethylene formation and the Cu catalyst itself. Furthermore, CO reduction instead of CO<sub>2</sub> reduction as well as experiments in different systems such as a flow cell and using a gas diffusion electrode at both elevated pressures and temperatures are interesting and important to gain more insight into the combined effects of pressure and temperature.

#### CRediT authorship contribution statement

**Rafaël E. Vos:** Conceptualization, Methodology, Validation, Formal analysis, Investigation, Writing – original draft, Visualization. **Daniel Schauermaun:** Validation, Investigation. **Marc T.M. Koper:** Conceptualization, Resources, Writing – review & editing, Supervision, Funding acquisition.

#### Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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#### Supplementary materials

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