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## 1 Introduction

The hydrogenation of  $\text{CO}_2$  to  $\text{CH}_3\text{OH}$  is of great significance, given potential applications in sustainable chemical processes and its important role in mitigating  $\text{CO}_2$  emissions.<sup>1,2</sup> Cu-based catalysts have shown outstanding performance and are widely used in industrial applications.<sup>3–5</sup> To optimize the catalytic efficiency, extensive research has been conducted to identify the active sites of copper species and understand the thermodynamic factors influencing the reaction kinetics, such as temperature, pressure and feedstock composition.<sup>6–8</sup> First principles calculations have played a pivotal role in this fundamental research.<sup>9,10</sup> The computational approach is highly effective in analyzing the interactions of various chemical species with the catalyst surface, locating active sites, exploring reaction pathways, and elucidating the underlying electronic structure and reaction mechanisms.<sup>11–13</sup>

The hydrogenation of  $\text{CO}_2$  on metallic Cu has been extensively studied using electronic density functional theory (DFT).<sup>14–17</sup> The theoretical work has helped establish a widely accepted reaction pathway for methanol synthesis, involving a series of intermediates such as  $\text{HCOO}$ ,  $\text{HCOOH}$ ,  $\text{H}_2\text{COOH}$ ,

$\text{H}_2\text{CO}$ ,  $\text{H}_3\text{CO}$ , and  $\text{H}_3\text{COH}$ .<sup>13,18</sup> In addition to catalyst development, numerous studies have focused on investigating thermodynamic effects on  $\text{CO}_2$  hydrogenation, including the influence of temperature, pressure, and feed composition on the reaction rate and conversion efficiency.<sup>19–22</sup> In general, the methanol yield can be maximized at low temperature, high pressure, high  $\text{H}_2:\text{CO}_x$  and  $\text{CO}:\text{CO}_2$  ratios.<sup>23</sup> However, different kinetic behaviors have been reported for the same catalyst under varying reaction conditions, and the underlying mechanisms remain not fully understood.<sup>20,24,25</sup>

The environmental effects on reaction rates can be investigated using Kinetic Monte Carlo (KMC) simulation, typically with the DFT results as input. To understand the kinetic behavior of  $\text{CO}_2$  hydrogenation to methanol under different conditions, Liu *et al.*<sup>26</sup> employed DFT and KMC simulations to investigate the turnover frequencies (TOF) of methanol synthesis at various  $\text{CO}/\text{CO}_2/\text{H}_2$  ratios on Cu (111) at 553 K and 80 bar. Their findings revealed that the  $\text{CH}_3\text{OH}$  TOF initially increases with the  $\text{CO}_2$  to  $(\text{CO}_2 + \text{CO})$  ratio and then decreases beyond an optimal composition. Sauer and coworkers<sup>25</sup> developed a three-site mean-field extended microkinetic model based on DFT calculations from the literature. They also demonstrated that methanol concentration initially increases and then decreases with the  $\text{CO}_2$  to  $(\text{CO}_2 + \text{CO})$  ratio on Cu (211) surfaces at 41 bar and temperatures ranging from 483 to 553 K. While these simulations accurately captured the reaction kinetics under certain conditions, discrepancies were noted

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when compared to experimental results. As reported by Sauer *et al.*,<sup>25</sup> the experimental data indicated that methanol concentration only increases with the  $\text{CO}_2$  to  $(\text{CO}_2 + \text{CO})$  ratio when the temperature is reduced below 493 K. The optimal  $\text{CO}_2$  to  $(\text{CO}_2 + \text{CO})$  ratio predicted by the micro kinetic model was not observed in experiments. This inconsistency likely stems from the interactions of reaction intermediates with gas molecules from the bulk, which are not adequately accounted for by conventional DFT calculations.

While first principles methods have achieved remarkable success in catalysis research, their applications for studying environmental factors, such as temperature, pressure, and gas-phase compositions, remain limited. Conventional DFT calculations typically assume a vacuum environment at 0 K. Although free-energy corrections are often applied to account for the bond vibrational entropy of the adsorbate, it ignores temperature effects on the thermodynamic properties of the entire system. Besides, conventional DFT calculations do not consider pressure and composition effects that influence the interaction of the adsorbate with gas molecules in the surroundings. Such interactions become increasingly significant at high pressure, leading to a “pressure gap” between theoretical predictions and experimental measurements.<sup>27</sup> Because of the limitations of conventional DFT methods, many discrepancies have been reported in the hydrogenation activity of  $\text{HCOO}^*$  on different Cu surfaces.<sup>11,12</sup> The pressure gap is responsible for differences between DFT-calculated energy barriers and experimental apparent activation energies,<sup>18</sup> the local reactant compositions, and uncertainties in reaction pathway selection.<sup>17,28</sup>

To bridge the pressure gap, we need to consider the interaction of catalyst and chemically adsorbed intermediates with gas molecules in an open environment. Toward that end, *ab initio* molecular dynamics (AIMD) simulations have been widely used to predict reaction processes under specific thermodynamic conditions.<sup>29–31</sup> AIMD allows for the inclusion of environmental molecules, enabling a precise analysis of their interactions with the catalyst surface. However, the applicability of AIMD to thermocatalytic reactions is limited, not only by the computational intensity of first principles calculations but also by the relatively low number density of gas molecules in the bulk phase. For instance, at 503 K and 50 bar, the bulk density of  $\text{CO}_2$  is only about 1 molecule per  $\text{nm}^3$ . Such a low density makes it challenging to simulate the gas composition near the catalyst surface accurately.

It should be noted that the impact of solvent effects in quantum chemical calculations is well established and is typically accounted for using either explicit or implicit solvent approaches. Implicit solvent models commonly employ the Poisson–Boltzmann equation to simulate the microenvironment around catalysts. Recently, Song *et al.* introduced a hybrid AIMD/cDFT model, demonstrating that classical DFT (cDFT) simulations can offer a more accurate description of solvent effects, thereby highlighting the potential of cDFT.<sup>32</sup> At high pressure, similar effects are possible in gas-phase reactions; however, these effects remain largely underexplored in current computational studies.

In this work, we propose a grand potential theory that combines DFT for electronic structure calculations with cDFT to describe the thermodynamic properties of the entire reaction system. By embedding the catalytic sites in an open thermodynamic environment, the hybrid DFT allows for an explicit description of the inhomogeneous distributions of gas molecules as well as the grand potential of the heterogeneous system at different stages of chemical reactions. Equipped with this new theoretical capability, we have explored diverse thermodynamic factors influencing  $\text{CO}_2$  hydrogenation to methanol, including the effects of temperature, pressure,  $\text{H}_2 : \text{CO}_2$  and  $\text{CO} : \text{CO}_2$  ratios, and the impact of CO and  $\text{H}_2\text{O}$  on  $\text{HCOO}^*$  hydrogenation. Through the grand potential simulations, we uncovered the molecular mechanisms for the requirement of high  $\text{H}_2$  pressure and elucidated the intercorrelated roles of  $\text{CO}_2$ , CO and  $\text{H}_2\text{O}$  in the hydrogenation process. The theoretical framework can be readily generalized to account for environmental effects in other thermocatalytic systems.

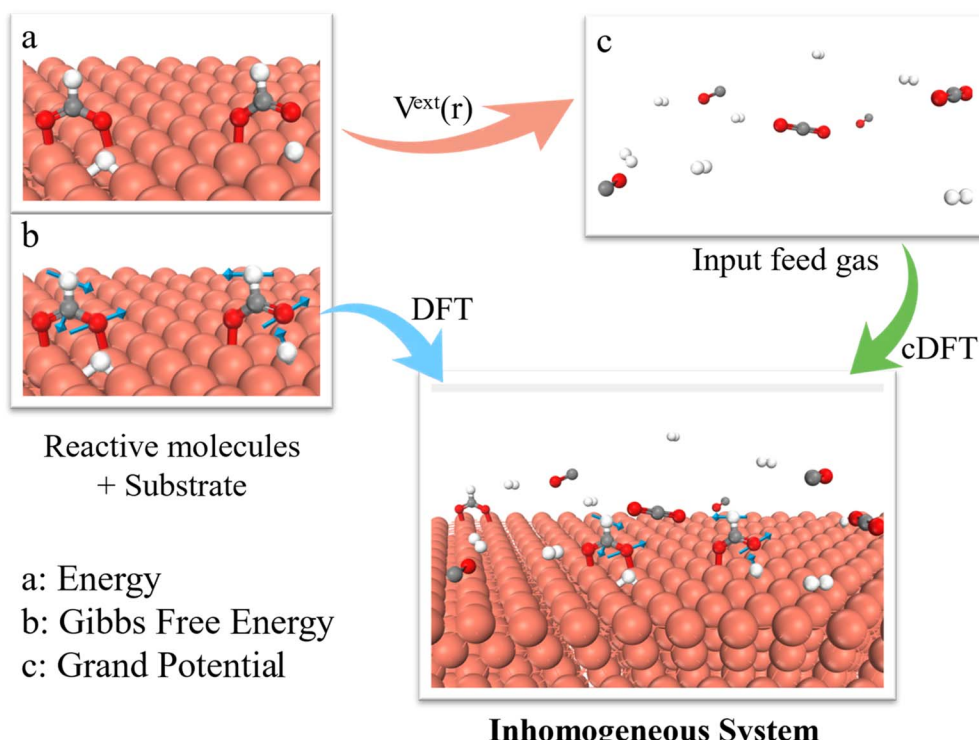
## 2 Results and discussion

The theoretical details for our grand potential theory are described in ESI.† Briefly, Fig. 1 illustrates the hybrid DFT approach to simulate  $\text{CO}_2$  hydrogenation to  $\text{CH}_3\text{OH}$  on Cu (111) and Cu (211) surfaces. The thermocatalytic process is governed by the grand potential of the entire system at different stages of the reaction, rather than the energy of individual intermediates (or the single molecular free energy) as used in conventional DFT calculations. To calculate the grand potential, we first identify the configuration of intermediate species at the catalyst surface through DFT calculations. The thermodynamic properties are then described using cDFT with a coarse-grained model for the surface species and environmental gas molecules. Notably, the grand potential simulations were conducted under a grand canonical ( $\mu\text{VT}$ ) ensemble, allowing for the variation of gas molecules in alignment with experimental conditions. While conventional KS-DFT methods are used to evaluate the electronic energies of reaction intermediates, cDFT accounts for physical interactions among gas molecules and their adsorption on the catalyst surface under reaction conditions. The grand potential calculations thus consider both chemical bonding and the equilibrium between the catalyst surface (including chemically adsorbed species) and the gas phase, incorporating all intermolecular interactions including molecules in the near-surface region and bulk phase.

### 2.1 Grand potential landscape

The grand-potential approach provides insight into the reaction mechanism from two distinct perspectives: the grand potential of the reactants and the catalyst as an open system and the local concentrations of molecular species near the catalyst surface. Specifically, Fig. 2 presents the grand-potential profile for  $\text{CO}_2$  hydrogenation to  $\text{CH}_3\text{OH}$  as well as CO hydrogenation to CHO on Cu (111) and Cu (211) surfaces. For comparison, Fig. 2 also shows the corresponding energy ( $E$ ) and free energy ( $G$ ) landscapes calculated from conventional methods. It should be





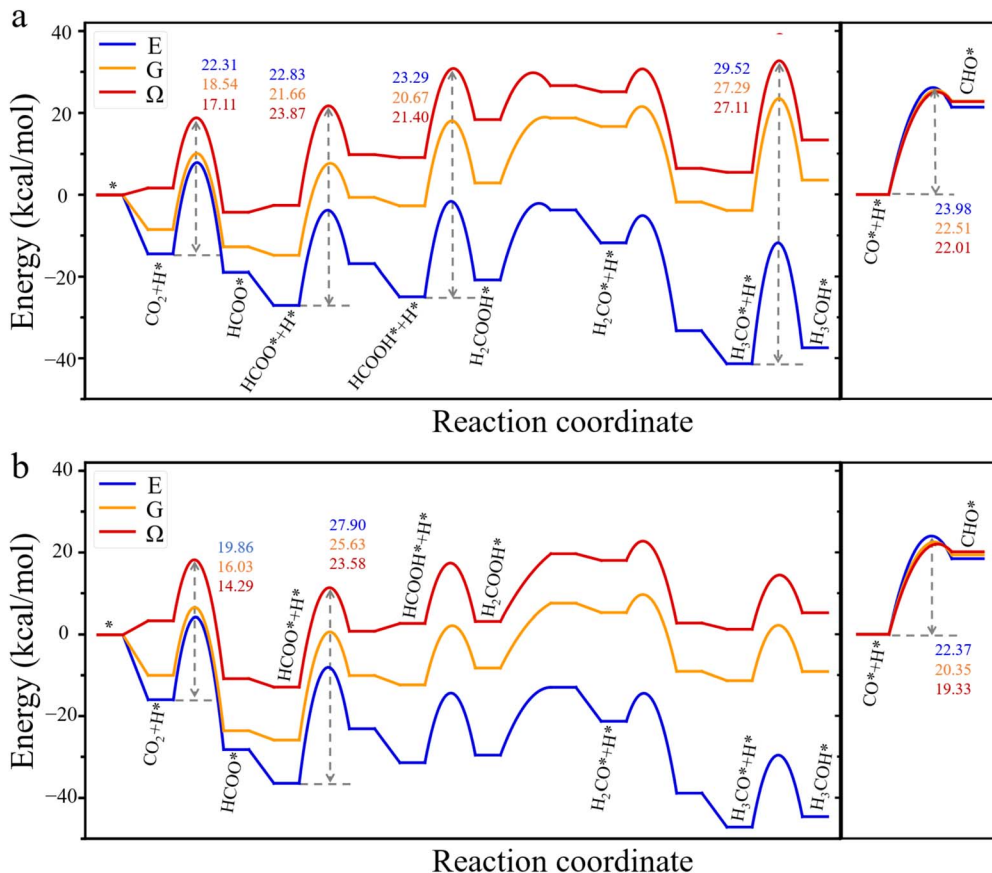
**Fig. 1** Scheme of the grand potential approach to predict environmental effects on heterogeneous catalysis. (a) Calculation of substrate energy ( $E$ ) with electronic DFT. (b) Free energy correction to account for the bond vibrational entropy, which is referred to as the Gibbs free energy in this work. (c) Calculation of the grand potential ( $\Omega$ ) of the entire inhomogeneous system with cDFT.

noted that in conventional DFT calculations, the energy  $E$  is defined as the difference between the surface with an adsorbate and the bare surface, along with a gas molecule in an isolated state (see eqn (S4)<sup>†</sup>). The free energy  $G$  corrects only the bond vibration entropy of individual intermediates. By contrast, the grand potential calculation naturally incorporates gas molecules in the environment at 503 K and 60 bar, with a feed gas composition of  $H_2:CO_2:CO = 6:1:1$ . This condition is commonly used in industrial applications of  $CO_2$  hydrogenation. Because the grand-potential corrected energy is defined relative to that of a bare surface in contact with the gas mixture under the reaction condition, it has a positive value for each intermediate. The positive energy does not imply exothermic adsorption of the intermediates, it only indicates that the adsorbates reduce the surface attraction for gas molecules in the bulk. In other words, the positively shifted energy profile originates from a negative energy in the reference state due to the strong attraction between Cu atoms and gas molecules.<sup>2,18,33</sup>

As the electronic energy is significantly larger than the corrections for bond vibrations and intermolecular interactions, the conventional and hybrid DFT methods predict similar energy diagrams. Nevertheless, the thermodynamic effects are significant, particularly for predicting the intermediate steps of  $CO_2$  hydrogenation. Fig. 2 shows that grand potential corrections are as important as free energy corrections related to bond vibrations, with both being of the same order of magnitude. On the Cu (111) surface (Fig. 2a), both conventional DFT methods and the grand potential theory predict four high-energy barriers

in the reaction pathway from  $CO_2$  to  $CH_3OH$ . These include  $CO_2$  hydrogenation to  $HCOO^*$ ,  $HCOO^*$  to  $HCOOH^*$ ,  $HCOOH^*$  to  $H_2COOH^*$ , and  $H_3CO^*$  to  $H_3COH^*$ . Notably, KS-DFT predicts that the highest energy barrier,  $29.52\text{ kcal mol}^{-1}$ , occurs in the elementary step from  $H_3CO^*$  to  $H_3COH^*$ . This is the rate-determining step, mostly responsible for the low activity of  $CO_2$  hydrogenation at the Cu (111) surface. After applying the free-energy corrections of bond vibrations, the energy barriers decline for all four elementary steps. In particular, the energy barrier for  $CO_2$  hydrogenation falls to  $18.54\text{ kcal mol}^{-1}$ , implying that the adsorbed  $CO_2$  becomes more easily hydrogenated to form  $HCOO^*$  at high temperature. These theoretical predictions are consistent with experimental results,<sup>34</sup> showing  $HCOO^*$  as the most prevalent species on Cu surfaces. With the grand-potential corrections, the reaction barriers for  $CO_2$  and  $H_3CO^*$  hydrogenation further decrease, while those for  $HCOO^*$  and  $HCOOH^*$  hydrogenation increase, narrowing the energy gap between  $HCOO^*$  and  $H_3CO^*$  hydrogenation. Given the high concentration of  $HCOO^*$  intermediate species,  $HCOO^*$  plays a key role in the conversion of  $CO_2$  to  $CH_3OH$  on the Cu (111) surface.

On the Cu (211) surface (Fig. 2b), all theoretical methods predict that  $HCOO^*$  hydrogenation to  $HCOOH^*$  is the most important rate-determining step. The energy barrier predicted by the KS-DFT decreases from  $27.90\text{ kcal mol}^{-1}$  to  $25.63\text{ kcal mol}^{-1}$  after the free-energy correction for bond vibrations. Nevertheless, this energy barrier is still much higher than  $21.66\text{ kcal mol}^{-1}$ , the energy barrier for  $HCOO^*$



**Fig. 2** Energy profiles for the hydrogenation of CO<sub>2</sub> and CO predicted by hybrid DFT (grand potential  $\Omega$ ) and conventional DFT (electronic energy  $E$  and free energy  $G$ ). (a) CO<sub>2</sub> hydrogenation to CH<sub>3</sub>OH and CO hydrogenation to CHO on Cu (111) surface. (b) Hydrogenation on Cu (211) surface. The simulations were conducted at 503 K and 60 bar, with a feed gas ratio of H<sub>2</sub> : CO<sub>2</sub> : CO = 6 : 1 : 1. Different from conventional DFT calculations for  $E$  and  $G$ , the grand potential is defined relative to that of the catalyst surface in contact with the gas phase as the reference system. The numbers represent energy barriers corresponding to the transition between various intermediate states predicted from different theoretical models.

hydrogenation to HCOOH\* on the Cu (111) surface. However, after applying grand-potential corrections, the energy barrier for HCOO\* hydrogenation increases on the Cu (111) surface but decreases on the Cu (211) surface. Consequently, the grand-potential barrier on Cu (211) is slightly lower (23.58 kcal mol<sup>-1</sup>) than that on Cu (111) (23.87 kcal mol<sup>-1</sup>), indicating that the HCOO\* hydrogenation on Cu (211) is comparable to that on Cu (111). While this slight difference is insufficient to explain why Cu (211) exhibits higher activity than Cu (111) for methanol production observed in experiment, the opposite trends in variation of the energy barriers on Cu (111) and Cu (211) surfaces underscore the importance of grand-potential corrections in understanding the reaction kinetics. As discussed later, the grand potential theory also provides information on the local concentrations of reactants, which have significant impacts on the reaction rate.<sup>12</sup> For CO hydrogenation, the conventional and hybrid DFT methods predict a similar energy profile on both Cu (111) and Cu (211) surfaces, indicating that the reaction energy is relatively insensitive to vibrational entropy and grand-potential corrections. However, environmental effects can still play a significant role in the kinetics of CO hydrogenation because the reaction rate depends

not only the energy barrier but also temperature and the local concentrations of reactants. In fact, temperature and gas composition are known to have strong influence on the water-gas shift (WGS) reaction.<sup>35</sup>

Through cDFT simulations, we have the density profiles of the feed gases near the catalyst surface. Fig. 3a and b show the gas density profiles on HCOO\*-adsorbed Cu (111) and Cu (211) surfaces, while the profiles for bare Cu (111) and Cu (211) surfaces are presented in Fig. S2.† These figures reveal a significant disparity between the surface densities of the input gases and their bulk values. Notably, the surface density of CO<sub>2</sub> is two orders of magnitude higher than its bulk density. Meanwhile, the surface densities of H<sub>2</sub> and CO oscillate near the catalyst surface and show significant deviations from their respective bulk values. The local composition of gas molecules near the catalyst surface is markedly different from the input composition, with CO<sub>2</sub> being much more prevalent than the other two gases. The H<sub>2</sub> : CO<sub>2</sub> ratio on the catalyst surface is much lower than the input ratio, implying that a higher H<sub>2</sub> : CO<sub>2</sub> ratio in the feed gas is necessary to increase surface H<sub>2</sub> density. Besides, CO<sub>2</sub> adsorption is more favorable than CO, leading to CO<sub>2</sub> the dominant carbon source in alignment with experimental observations.<sup>18</sup>

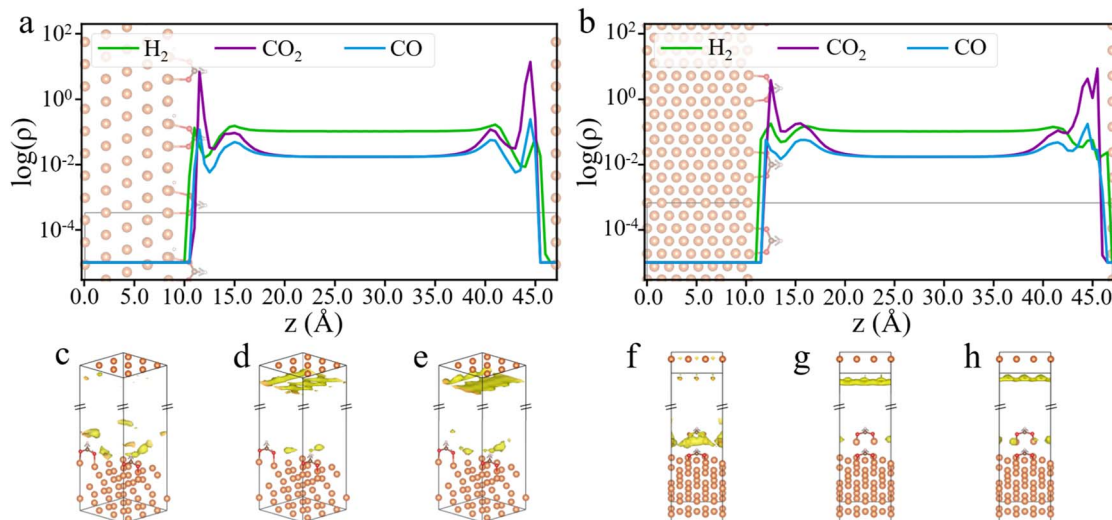


Fig. 3 Density profiles and 3D density maps of gas molecules on Cu surfaces. Average gas densities along the direction (z-axis) perpendicular to HCOO\* adsorbed Cu (111) (a) and Cu (211) (b) surfaces. 3D density maps of H<sub>2</sub> (c and f), CO<sub>2</sub> (d and g), and CO (e and h) on HCOO\* adsorbed Cu (111) and Cu (211) surfaces, respectively. Different isosurface values were used for better visualization of each gas molecule. The simulations were conducted at 503 K and 60 bar, with a feed gas ratio of H<sub>2</sub> : CO<sub>2</sub> : CO = 6 : 1 : 1.

As will be discussed later, using surface reactant densities rather than their bulk densities provides a more accurate description of the hydrogenation kinetics. Fig. 3 shows that the first peak of the feed gas density occurs within about 5 Å of the surface, prompting us to define the gas concentration at the surface within this range. Additionally, by contrasting the differences between the HCOO\*-adsorbed surface and the pristine surface, we observe that the presence of HCOO\* reduces the surface density of CO<sub>2</sub> and CO, while increasing the surface density of H<sub>2</sub> (the bottom panel of Fig. S2b†). These differences are likely attributed to larger excluded volume effects due to the interactions of HCOO\* with CO<sub>2</sub> and CO molecules, whereas H<sub>2</sub>, with its smaller molecular size, is relatively less affected by the surface occupancy of the HCOO\* species. Furthermore, from the 3D density maps of the feed gases on Cu (111) and Cu (211) surfaces (Fig. 3c–h), it is evident that H<sub>2</sub> tends to accumulate around the HCOO\* species rather than on the bare Cu surface. In contrast, CO<sub>2</sub> and CO show greater adsorption on the bare Cu surfaces.

## 2.2 Effects of temperature and pressure

To investigate the environmental effects on CO<sub>2</sub> hydrogenation to CH<sub>3</sub>OH, we calculated the grand potential barriers and surface gas densities over a wide range of temperatures and pressures. Fig. 4 presents the changes in the grand potential barrier ( $\Delta\Omega$ ) for HCOO\* and H<sub>3</sub>CO\* hydrogenation on Cu (111), as well as HCOO\* hydrogenation on Cu (211), while Fig. S3† shows the simulation data for HCOOH\* hydrogenation on Cu (111). For HCOO\* hydrogenation on the Cu (111) surface, the variation in the grand potential barrier is around 2.5 kcal mol<sup>-1</sup>, with the minimum appeared at high temperature and low pressure. In the case of HCOOH\* hydrogenation, the grand potential barrier varies within the range of approximately 0.8 kcal mol<sup>-1</sup> under most conditions. In contrast, for H<sub>3</sub>CO\*,

the grand potential correction is highly sensitive to both temperature and pressure, generally increasing with temperature and decreasing with pressure, with the changes ranging from -1.6 to 0.4 kcal mol<sup>-1</sup>. On the Cu (211) surface, HCOO\* hydrogenation shows a similar trend, with the grand potential increasing with temperature but decreasing with pressure. At low temperature and high pressure, the grand potential correction can reduce the energy barrier by up to 3 kcal mol<sup>-1</sup>.

The impact of temperature and pressure on the grand-potential corrected energy barrier is primarily driven by their influence on the surface-phase composition. Fig. 4d–i presents the surface densities of the feed gas at various temperatures and pressures. A higher gas-phase density results in stronger interactions between the gas-phase molecules and surface intermediates. While it is somewhat expected that the surface density of each gas increases with pressure and decreases with temperature, several noteworthy points emerge from the grand-potential simulation: first, CO<sub>2</sub> adsorption becomes easily saturated due to its high concentration in the gas mixture. Second, the surface density of H<sub>2</sub> initially decreases and then increases with temperature. At high pressure, the surface density of H<sub>2</sub> on Cu (111) is greater at 703 K than at 303 K. This seemingly counterintuitive result can be attributed to CO<sub>2</sub> saturation at low temperature, which inhibits hydrogen adsorption. Third, the surface densities of CO<sub>2</sub> and CO on Cu (111) and Cu (211) surfaces are similar, while the H<sub>2</sub> surface density on Cu (211) is approximately twice as high as that on Cu (111). The stepped Cu (211) surface is thus more favorable for H<sub>2</sub> adsorption than the Cu (111) surface, further supporting that Cu (211) has a higher reactivity.

Based on the grand potential landscape and surface densities, we predicted the reaction rate of HCOO\* hydrogenation using the transition state theory (TST). As shown in Fig. S4a and b,† the reaction rate increases monotonically with temperature, and it initially rises with pressure before reaching a plateau.



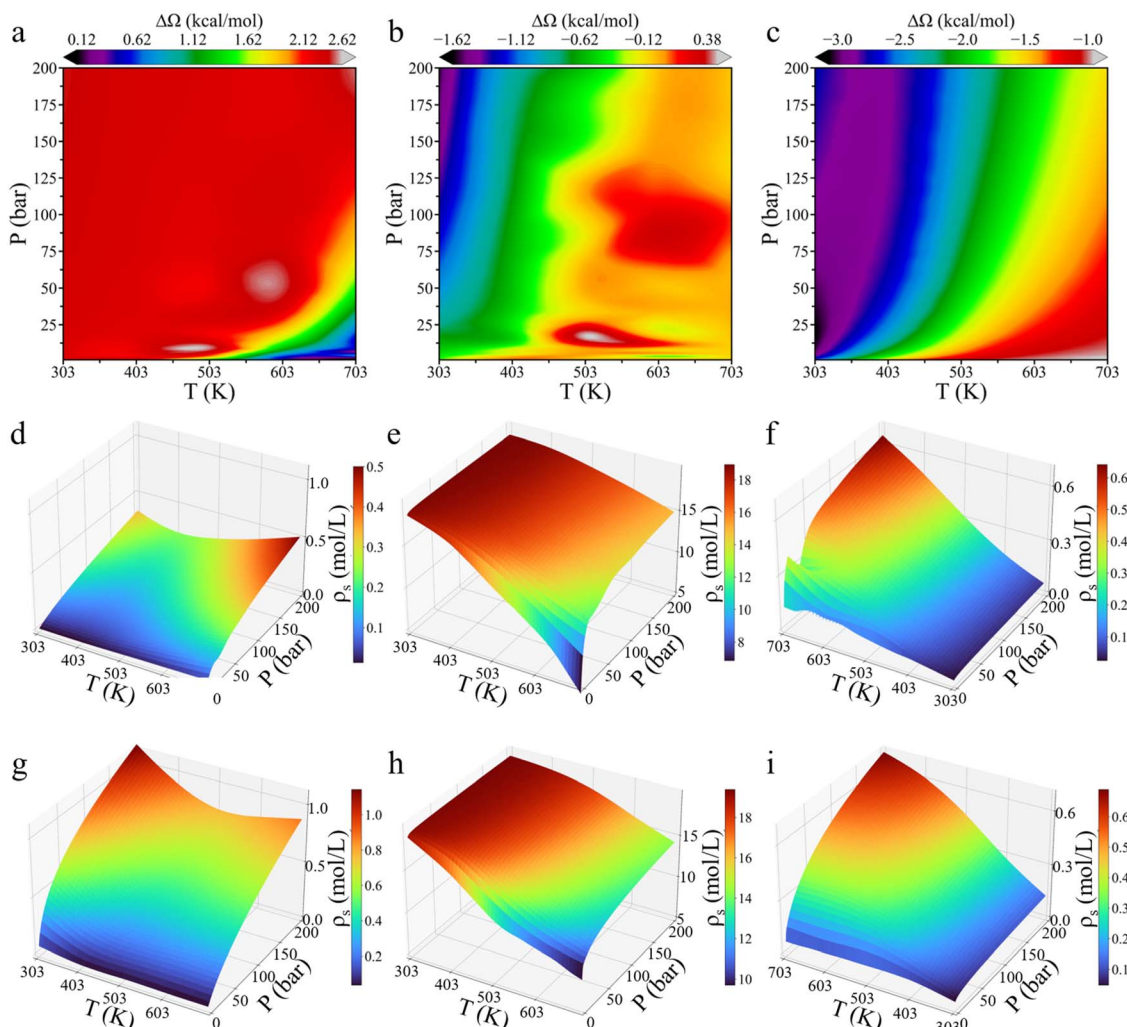


Fig. 4 The influences of temperature and pressure on the grand potential barrier and surface densities of various gas molecules. Contour plots of the change in the grand potential barrier ( $\Delta\Omega$ ) for  $\text{HCOO}^*$  (a) and  $\text{CH}_3\text{O}^*$  (b) hydrogenation on Cu (111), and for  $\text{HCOO}^*$  (c) hydrogenation on Cu (211). The surface density ( $\rho_s$ ) versus temperature and pressure for  $\text{H}_2$  (d and g),  $\text{CO}_2$  (e and h), and  $\text{CO}$  (f and i) on Cu (111) and Cu (211), respectively. The coordinate orientation has been adjusted for better visualization. The feed gas ratio was set to  $\text{H}_2 : \text{CO}_2 : \text{CO} = 6 : 1 : 1$ .

Fig. S4c and d† present the ratio of reaction rates calculated *via* the grand potential theory to those calculated using the conventional DFT method (*viz.*, through the free-energy barrier and bulk densities). After accounting for corrections of the surface densities and the grand potential, the reaction rate decreases by 1–2 orders of magnitude on the Cu (111) surface at most conditions. However, at 703 K and low pressure, the thermodynamic correction raises the reaction rate by almost 3 orders of magnitude. The drastic increase in the reaction rate can be attributed to the enhanced  $\text{HCOO}^*$  coverage, which can be estimated from the quasi-equilibrium condition (see ESI†). As shown in Fig. S5,† the conventional approach predicts a small  $\text{HCOO}^*$  coverage at high temperature and low pressure. The grand potential correction reduces the reaction energy for  $\text{HCOO}^*$  formation. Besides, it predicts that the surface density of  $\text{CO}_2$  on Cu (111) significantly exceeds its bulk density. As a result, the grand-potential simulations yield an extremely high  $\text{HCOO}^*$  coverage, close to 100%, explaining the drastic increase

of the reaction rate. On the Cu (211) surface at low temperature and pressure, the grand potential theory predicts a reaction rate approximately 1–2 orders of magnitude higher than that calculated using the conventional method. At high temperature and pressure, the grand potential theory still predicts a higher reaction rate, but the relative difference between the two methods decreases. Because of the strong adsorption of  $\text{HCOO}^*$  on Cu (211), the  $\text{HCOO}^*$  coverage approaches near 100% according to both grand potential and free energy simulations. The high  $\text{HCOO}^*$  coverage implies that the hydrogenation rate of  $\text{HCOO}^*$  is independent of the surface concentration of  $\text{CO}_2$ , challenging conventional understandings based on the  $\text{CO}_2$  pressure.

### 2.3 Effects of feed gas composition

**2.3.1.  $\text{H}_2 : \text{CO}_2$  ratio.** In this section, we explore the influence of the  $\text{H}_2 : \text{CO}_2$  ratio on the hydrogenation kinetics. Previous research suggests that a high  $\text{H}_2 : \text{CO}_2$  ratio favors the



conversion of  $\text{CO}_2$  to  $\text{CH}_3\text{OH}$ . For example, Lin and Bhan<sup>20</sup> showed that, at a constant  $\text{CO}_2$  pressure, increasing  $\text{H}_2$  pressure linearly increases the methanol formation rate. Conversely, at 523 K and constant  $\text{H}_2$  pressure, the reaction exhibits Langmuir-type saturation kinetics with respect to  $\text{CO}_2$  pressure on Copper-Zinc-Alumina (CZA) catalysts. Nielsen *et al.*<sup>24</sup> reported a similar Langmuir-type behavior concerning the effect of  $\text{CO}_2$  pressure on  $\text{CH}_3\text{OH}$  formation rates on  $\text{Cu}/\text{ZnO}$  catalysts. These findings suggest that maintaining a high  $\text{H}_2 : \text{CO}_2$  ratio is crucial for optimizing the kinetics of methanol synthesis. This is often attributed to the dearth of surface  $\text{H}^*$  and the saturation of  $\text{HCOO}^*$  species on the catalyst surface. However, the upper limit of  $\text{H}_2 : \text{CO}_2$  ratio remains uncertain. Lin and Bhan<sup>20</sup> demonstrated that methanol production continues to increase linearly with  $\text{H}_2$  pressure, even when the  $\text{H}_2 : \text{CO}_2$  ratio exceeds 10. This raises the question: why is such a high  $\text{H}_2$  pressure necessary? Moreover, even after the inflection point of the Langmuir-type curve, a further increase in the  $\text{CO}_2$  partial pressure can still raise the methanol production rate. This brings up another question: what role does  $\text{CO}_2$  partial pressure play once  $\text{HCOO}^*$  saturation is reached?

The answers to these questions are related to the low  $\text{H}_2 : \text{CO}_2$  ratio on the surface and the interactions between  $\text{CO}_2$  molecules and the reaction intermediates. As shown in Fig. 5, the grand potential barrier for  $\text{HCOO}^*$  hydrogenation on the  $\text{Cu} (111)$  surface increases with rising  $\text{CO}_2$  pressure, while it

remains nearly unchanged with varying  $\text{H}_2$  pressure. On the  $\text{Cu} (211)$  surface, however, the grand potential barrier falls with increasing  $\text{CO}_2$  pressure but increases with  $\text{H}_2$  pressure at higher  $\text{CO}_2$  pressure. Additionally, the surface density of  $\text{H}_2$  increases linearly as  $\text{H}_2$  pressure rises, with the increment being more pronounced at lower  $\text{CO}_2$  pressure. Notably, the surface density of  $\text{H}_2$  on  $\text{Cu} (211)$  is more than twice that on  $\text{Cu} (111)$ . For  $\text{CO}_2$ , the surface density rises sharply at low  $\text{CO}_2$  pressure before saturation. The inflection point occurs around 10 bar  $\text{CO}_2$  pressure, indicating that a further increase in  $\text{CO}_2$  pressure offers little additional benefit. As shown in Fig. S6,<sup>†</sup> at low  $\text{CO}_2$  pressure, the surface density of  $\text{CO}_2$  is two orders of magnitude higher than its bulk density, but the ratio of its surface density to the bulk density decreases sharply as the  $\text{CO}_2$  pressure increases to several bars. However, experimental results by Nielsen *et al.*<sup>24</sup> indicate that the inflection point for the  $\text{CO}_2$  partial pressure in relation to  $\text{CH}_3\text{OH}$  formation rate occurs at only 1–2 bar. This discrepancy can be attributed to the limited availability of active sites. As illustrated in Fig. S7,<sup>†</sup>  $\text{HCOO}^*$  coverage becomes saturated even at low  $\text{H}_2$  and  $\text{CO}_2$  pressures, resulting in the  $\text{HCOO}^*$  coverage reaching a plateau before the  $\text{CO}_2$  surface concentration becomes saturated.

Fig. 6 shows the reaction rate for  $\text{HCOO}^*$  hydrogenation predicted by the transition state theory. On the  $\text{Cu} (111)$  surface, the hydrogenation rate increases with  $\text{H}_2$  pressure while decreasing with  $\text{CO}_2$  pressure. The  $\text{HCOO}^*$  hydrogenation rate

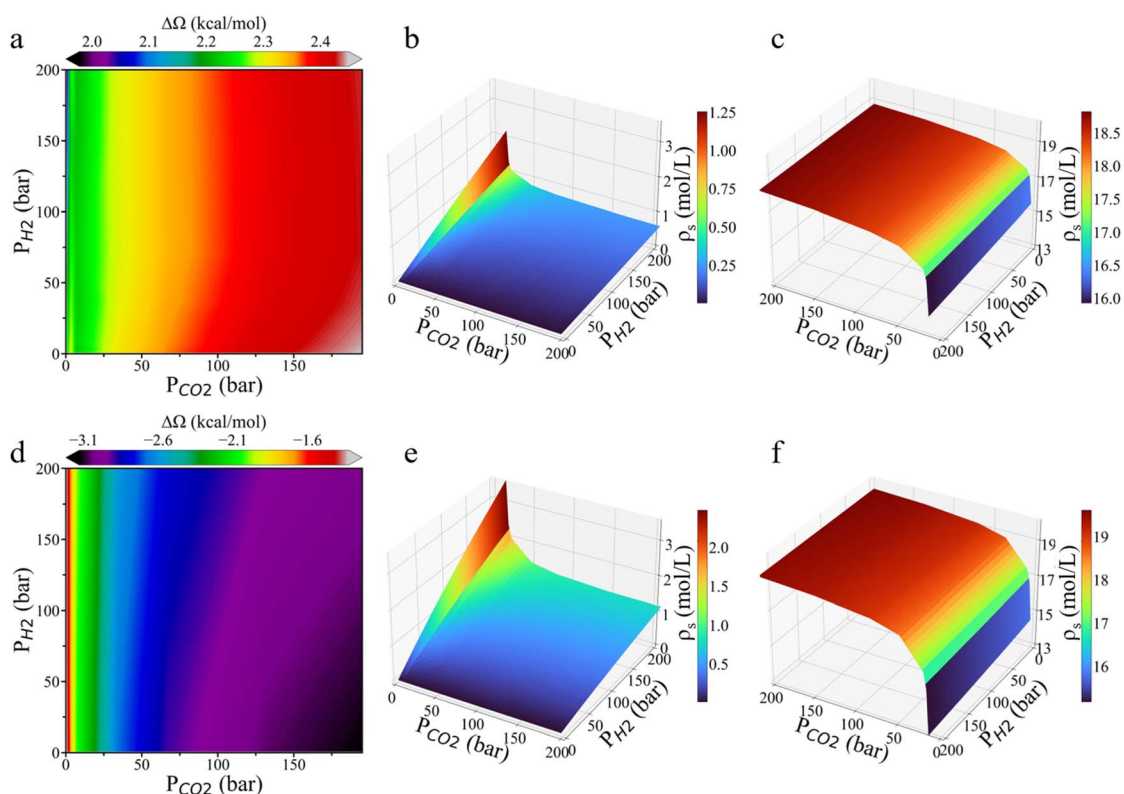


Fig. 5 Variations of the grand potential barrier and surface gas densities with respect to the partial pressures of  $\text{H}_2$  and  $\text{CO}_2$  in the feed gas. Contour plots of the changes in the grand potential barrier for  $^*\text{HCOO}$  hydrogenation on  $\text{Cu} (111)$  (a) and  $\text{Cu} (211)$  (d) surfaces. Average surface densities ( $\rho_s$ ) of  $\text{H}_2$  (b and e) and  $\text{CO}_2$  (c and f) on  $\text{Cu} (111)$  and  $\text{Cu} (211)$ , respectively. The grand potential simulation was performed at 503 K. The coordinate orientation has been adjusted for better visualization.



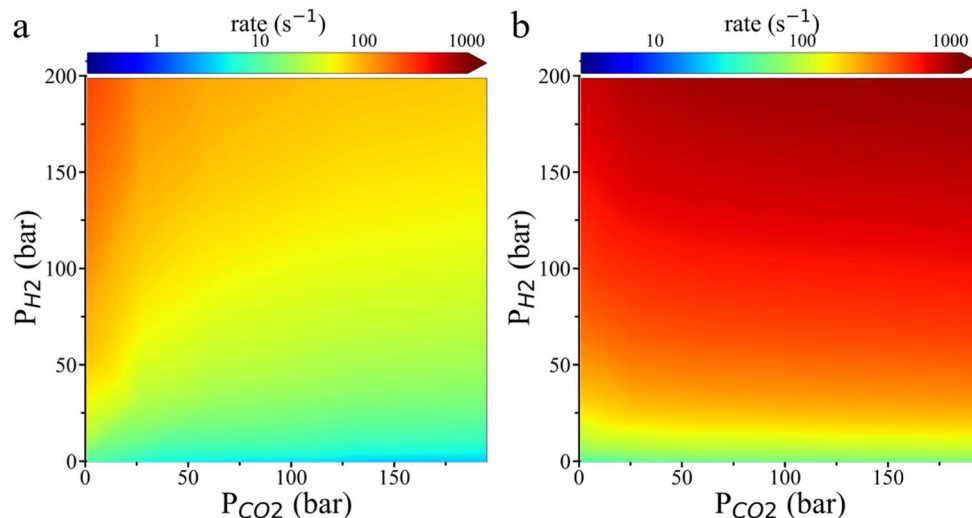


Fig. 6 Contour plot of the  $\text{HCOO}^*$  hydrogenation rate as a function of  $\text{CO}_2$  and  $\text{H}_2$  partial pressures on Cu (111) (a) and Cu (211) (b) surfaces. The reaction rate was predicted by the transition state theory with grand potential simulation at 503 K.

is faster on the Cu (211) surface than on the Cu (111) surface. The theoretical result aligns well with the experimental observation that increasing  $\text{H}_2$  pressure significantly enhances the reaction rate, but increasing  $\text{CO}_2$  pressure has a minimal effect only when  $\text{H}_2$  pressure is high while  $\text{CO}_2$  pressure is low. Besides, as shown in Fig. S8,<sup>†</sup> the grand potential simulation indicates that the environmental effects become more pronounced on the rate of  $\text{HCOO}^*$  hydrogenation on the Cu (211) surface at low  $\text{H}_2$  pressure and high  $\text{CO}_2$  pressure. The rate increase predicted by the grand potential theory agrees with the experimental finding that the  $\text{CH}_3\text{OH}$  formation rate increases slightly with rising  $\text{CO}_2$  pressure, even when the surface coverage of  $\text{HCOO}^*$  is saturated.

Based on the grand potential simulation, we found that the surface density of  $\text{H}_2$  is two orders of magnitude lower than the bulk density (Fig. 5b and e). This explains why the surface density of  $\text{H}_2$  increases linearly with bulk density without reaching saturation. In contrast,  $\text{CO}_2$  does not influence the  $\text{HCOO}^*$  hydrogenation rate through surface concentration because the  $\text{HCOO}^*$  coverage is already saturated. Instead, the  $\text{HCOO}^*$  hydrogenation rate is influenced by both the energy barrier and the surface  $\text{H}_2$  concentration, these opposing effects cancel each other, leading to a stable  $\text{HCOO}^*$  reaction rate with varying  $\text{CO}_2$  pressure. From this, we deduce that, at low temperature, the effect due to the reduction in energy barrier is more dominant, whereas at high  $\text{H}_2$  pressure, the reduction in surface  $\text{H}_2$  density has a greater impact. Therefore, the optimal  $\text{H}_2 : \text{CO}_2$  ratio should decrease with temperature while increase with rising  $\text{H}_2$  pressure.

Additionally, our grand-potential simulations may help resolve the ongoing debate regarding the formate pathway *versus* the carboxy pathway in  $\text{CO}_2$  hydrogenation.<sup>3,11,17,35</sup> The reaction mechanism has been extensively debated over the past decade, with the formate pathway now widely accepted. However, certain observations continue to support the possibility of a carboxy pathway. For example, Yang *et al.*<sup>28</sup> reported

that surface reaction involving formate-containing adlayers on Cu (both supported on  $\text{SiO}_2$  and unsupported) failed to produce methanol in hydrogen atmospheres. Furthermore, some researchers questioned that surface  $\text{HCOO}^*$  crowding at high coverages could enhance the reaction rate through the formate pathway.<sup>36</sup> Based on the grand potential simulations, we observed that  $\text{CO}_2$  in the gas phase reduces the  $\text{HCOO}^*$  hydrogenation energy barrier (Fig. 5d), indicating that  $\text{HCOO}^*$  can hydrogenate more readily under a high-pressure atmosphere of  $\text{CO}_2$  gas. In contrast, without  $\text{CO}_2$  in the bulk gas, the grand potential theory predicts that  $\text{HCOO}^*$  cannot undergo hydrogenation due to the high energy barrier (Fig. 5d), confirming that the  $\text{HCOO}^*$  pathway is the dominant route for  $\text{CO}_2$  hydrogenation.

**2.3.2. Effect of CO.** The influence of CO to  $(\text{CO}_2 + \text{CO})$  ratio on the kinetics of hydrogenation to methanol has been a controversial issue. It is generally accepted that CO can reduce the surface binding of oxygen on  $\text{Cu}^+$ , inhibit the reverse water-gas shift (RWGS) reaction, and remove  $\text{H}_2\text{O}$  from the catalyst surface, thereby making a high  $\text{CO} : \text{CO}_2$  ratio favorable for methanol production.<sup>2,21,36,37</sup> However, Kunkes *et al.*<sup>38</sup> observed a continuous rise in methanol production rate with increasing the  $\text{CO}_2$  to  $(\text{CO}_2 + \text{CO})$  ratio on CZA at 413 K and 30 bar. Similar trends were reported by Liu *et al.*<sup>39</sup> on CZA at 523 K and 50 bar. Meanwhile, Studt *et al.*<sup>33</sup> noted that, while the intrinsic rate of methanol formation increases with the  $\text{CO}_2$  to  $(\text{CO}_2 + \text{CO})$  ratio on CZA, it decreases on  $\text{Cu}/\text{MgO}$  at 503 K and 30 bar. To minimize the effect of byproduct  $\text{H}_2\text{O}$ , Nielsen *et al.*<sup>24</sup> carried out experiment under low conversion conditions (<0.3 mol% methanol). They found that CO is purely inhibitory on RANEY® Cu and a variety of supported Cu-catalysts due to its competitive adsorption to the catalyst surface. However, as previously mentioned, when  $\text{CO}_2$  pressure exceeds 2 bar, the surface species  $\text{HCOO}^*$  becomes saturated, suggesting that an additional increase of the  $\text{CO}_2$  pressure may not enhance methanol production significantly. This raises the question: why does CO



still inhibit methanol formation even at high  $\text{CO}_2$  pressure? Moreover, recent experimental studies<sup>3,12,18</sup> indicate that  $\text{CO}_2$  remains the dominant carbon source, even at low  $\text{CO}_2$  to  $(\text{CO}_2 + \text{CO})$  ratios, because CO tends to desorb more easily rather than undergo hydrogenation. This allows  $\text{HCOO}^*$  derived from  $\text{CO}_2$  to reoccupy the active sites, as evidenced by the high coverage of  $\text{HCOO}^*$ . This suggests that CO adsorption does not significantly poison the active sites or reduce TOF of methanol formation. Another question then arises: what is the precise role of CO in this process?

To address the above questions, we carried out grand-potential simulations at different ratios of CO to  $(\text{CO}_2 + \text{CO})$  while keeping the  $\text{H}_2$  pressure constant. As shown in Fig. 7, the grand-potential correction of the energy barrier falls with increasing CO ratio for  $\text{HCOO}^*$  hydrogenation on the Cu (111) surface, whereas the opposite trend is observed on the Cu (211) surface. This difference arises because the interaction between surface intermediates and CO is weaker than that with  $\text{CO}_2$ . As the CO ratio increases, the absolute value of the grand-potential correction diminishes for both surfaces. Notably, the grand-potential curve at 8 bar  $\text{H}_2$  pressure exhibits a distinct inflection point, likely due to a shift in the interaction of  $\text{HCOO}^*$  at the Cu (111) surface with the input gas molecules—from  $\text{H}_2$  sub-dominance to CO sub-dominance—which causes an unexpected increase in the grand-potential barrier. From Fig. 7c and

d and S9,<sup>†</sup> we observe that the surface density of  $\text{H}_2$  increases slightly with the rising  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratio, leading to the increase of the  $\text{HCOO}^*$  hydrogenation rate on the Cu (111) surface. On the Cu (211) surface, the rate of  $\text{HCOO}^*$  hydrogenation is nearly unchanged at 8 bar  $\text{H}_2$  partial pressure, until the  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratio reaches above 0.7, at which point the rate increases (Fig. 7f). At 42 bar  $\text{H}_2$  pressure, however, the reaction rate first decreases and then increases with the  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratio. The nonmonotonic trend corroborates the experimental observation that CO does not directly compete with  $\text{CO}_2$  for adsorption on the active sites.<sup>24</sup> With the increasing  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratio, the accumulation of CO on the Cu surface increases the energy barrier of  $\text{HCOO}^*$  hydrogenation as well as  $\text{H}_2$  density on Cu (211) surface. These two opposing effects together control the  $\text{HCOO}^*$  hydrogenation rate. At low  $\text{H}_2$  pressure, the increase in  $\text{H}_2$  surface density is the dominant factor, while at high  $\text{H}_2$  partial pressure, the effect on the energy barrier becomes more influential. Therefore, at low  $\text{H}_2$  pressure, the  $\text{HCOO}^*$  hydrogenation rate increases with the CO ratio, but at high  $\text{H}_2$  pressure, the rate decreases with increasing CO ratio. This conclusion aligns well with the experimental findings that  $\text{CH}_3\text{OH}$  formation rate remains unchanged with varying CO pressure at  $P_{\text{CO}_2} = 4.7$  bar and  $P_{\text{H}_2} = 14$  bar, but decreases with CO pressure at  $P_{\text{CO}_2} = 4.7$  bar and  $P_{\text{H}_2} = 23$  bar.<sup>20</sup>

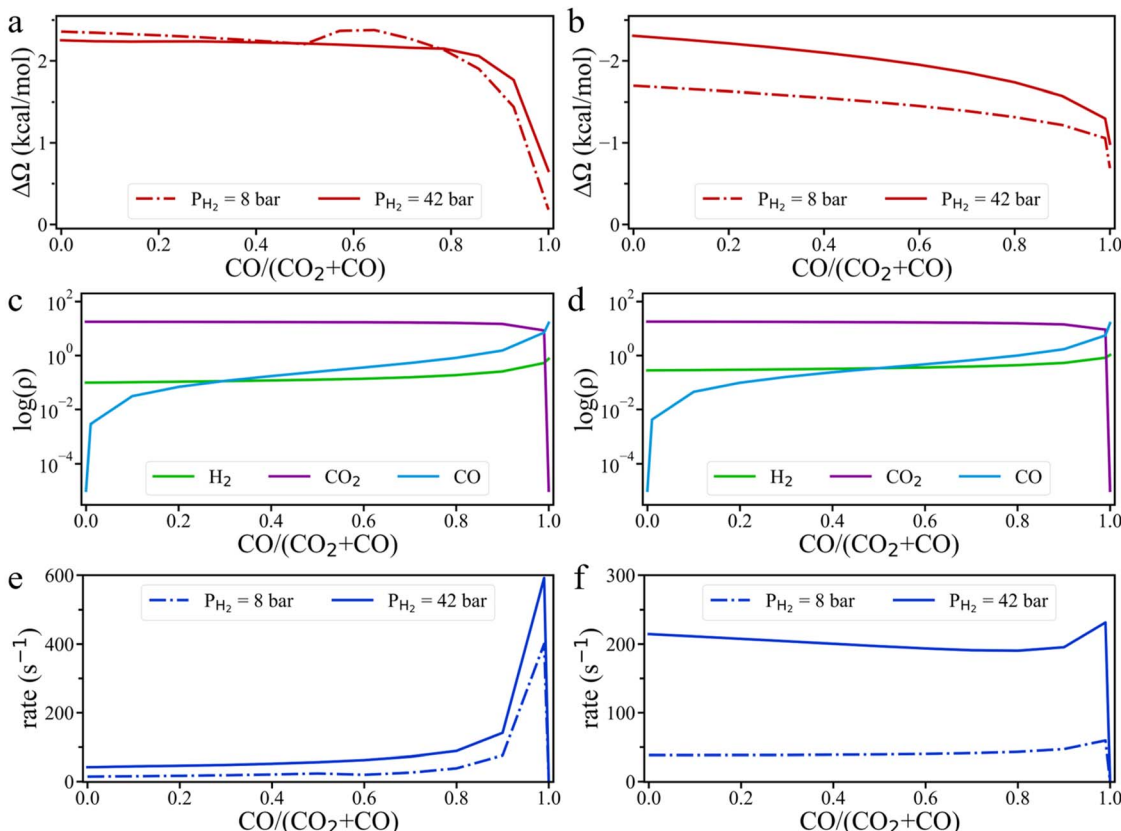


Fig. 7 The influence of the  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratio on the kinetics of hydrogenation. Variation of the grand potential barrier ( $\Delta\Omega$ ) for  $\text{HCOO}^*$  hydrogenation on Cu (111) (a) and Cu (211) (b) surfaces. The surface densities of the feed gas on Cu (111) (c) and Cu (211) (d) under 42 bar  $\text{H}_2$  pressure. The rate of  $\text{HCOO}^*$  hydrogenation on Cu (111) (e) and Cu (211) (f). All simulations were conducted at 503 K, with a fixed feed gas ratio of  $\text{H}_2 : (\text{CO}_2 + \text{CO}) = 3 : 1$ .



Fig. 7f also shows that the  $\text{HCOO}^*$  hydrogenation rate unexpectedly increases when the  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratio reaches 0.99. In this case, the surface density of  $\text{CO}_2$  remains sufficiently high to form a substantial amount of  $\text{HCOO}^*$  at the catalyst surface, while the surface density of  $\text{H}_2$  also increases significantly. The hydrogenation rate grows with the increasing  $\text{H}_2$  surface density because it has a favorable effect on  $\text{HCOO}^*$  hydrogenation outweighing the negative effect due to the increased energy barrier. Experimental results from 40 years ago indicated that methanol formation rate in  $\text{H}_2/\text{CO}$  mixtures is optimized when a small amount of  $\text{CO}_2$  is included in the feed, which supports our conclusion.<sup>40,41</sup> However, the subsequent interpretation suggesting that 'the RWGS reaction is likely autocatalyzed by water or water-derived species' was incorrect and has largely been overlooked. Based on our proposed mechanism, CO undergoes both the water-gas shift (WGS) reaction and  $\text{CO}_2$  hydrogenation pathways to produce  $\text{CH}_3\text{OH}$ . The fact that  $\text{HCOO}^*$  is not readily detected can be explained by the prevalence of the Cu (111) surface, which is the most common facet for the Cu catalyst. When the  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratio is greater than 0.9, the  $\text{HCOO}^*$  hydrogenation rate on Cu (111) is hundreds of times faster than that at lower  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratios. Therefore,  $\text{HCOO}^*$  detected in experiments is likely presented on the Cu (111) surface, where it is rapidly converted to other intermediates at high  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratios.

This mechanism also explains why CO hydrogenation can be accelerated by the addition of  $\text{H}_2\text{O}$ ,<sup>28</sup> and why CO hydrogenation on Cu/MgO oxide supports is significantly faster than  $\text{CO}_2$  hydrogenation.<sup>34</sup>

**2.3.3. Effect of  $\text{H}_2\text{O}$ .**  $\text{H}_2\text{O}$  is an inevitable byproduct during  $\text{CO}_2$  hydrogenation to  $\text{CH}_3\text{OH}$  and has been shown to significantly impact the kinetics of methanol synthesis. However, the influence of  $\text{H}_2\text{O}$  partial pressure on  $\text{CO}_2$  hydrogenation remains controversial. Numerous studies have reported that a small amount of  $\text{H}_2\text{O}$  can drastically reduce methanol formation rate. For instance, Thrane *et al.*<sup>42</sup> found that adding 1500 ppmv  $\text{H}_2\text{O}$  to the syngas feed led to a 60–70% reduction in the methanol formation rate over CZA at 493–523 K and 41 bar. Similarly, Sahibzada *et al.*<sup>41</sup> observed a 90% loss in the catalyst activity for methanol synthesis at 523 K and 50 bar when 2 vol%  $\text{H}_2\text{O}$  was added. In contrast, Yan *et al.*<sup>43</sup> reported that  $\text{H}_2\text{O}$  had no significant effect on  $\text{CO}_2$  conversion within the temperature range of 453–573 K for Cu/ZnO, Cu/Al<sub>2</sub>O<sub>3</sub>, and Cu/SiO<sub>2</sub> catalysts, while Zhao *et al.*<sup>3</sup> found that a small amount of  $\text{H}_2\text{O}$  could even have a positive effect under certain conditions. Additionally, DFT and KMC simulations performed by Liu *et al.*<sup>26</sup> suggested that the effect of  $\text{H}_2\text{O}$  on TOF is minimal. These conflicting results raise the question: why does a small amount of  $\text{H}_2\text{O}$  drastically reduce the methanol formation rate in some cases,

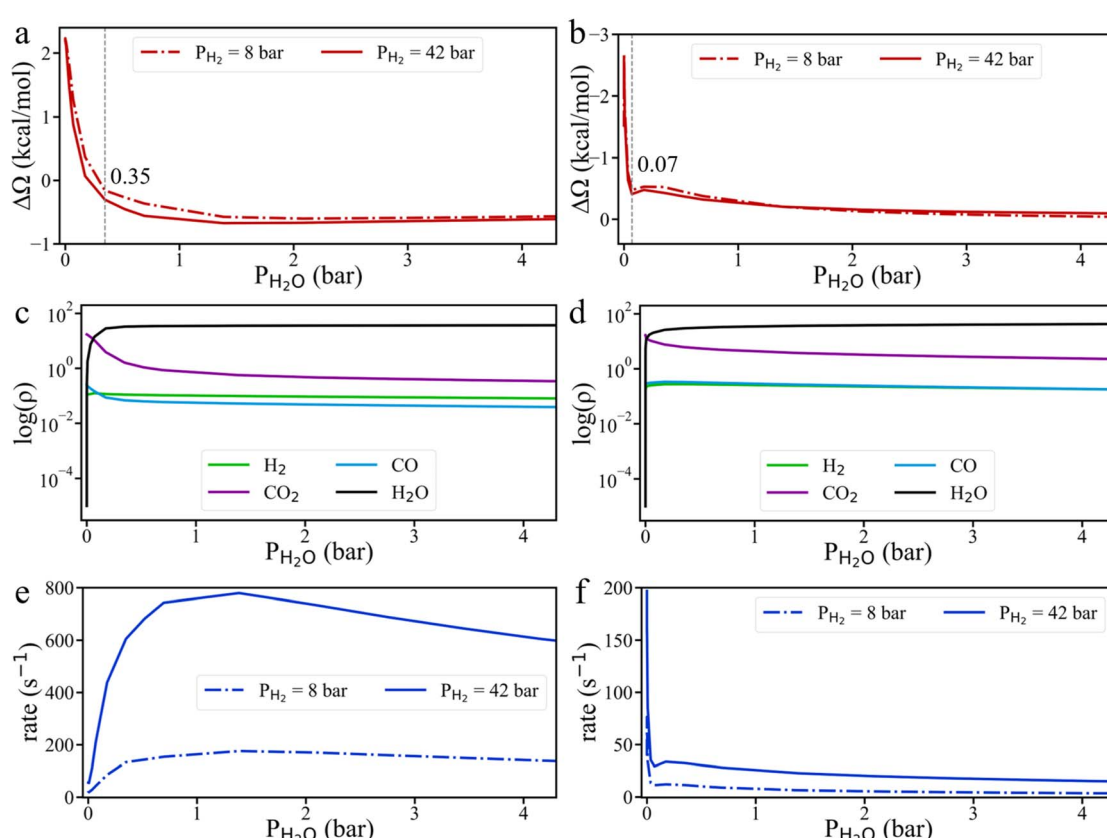


Fig. 8 The influence of  $\text{H}_2\text{O}$  partial pressure on the kinetics of  $\text{CO}_2$  hydrogenation. Variation of the grand potential barrier for  $\text{HCOO}^*$  hydrogenation on Cu (111) (a) and Cu (211) (b) surfaces. Surface gas densities on Cu (111) (c) and Cu (211) (d) under 42 bar  $\text{H}_2$  pressure.  $\text{HCOO}^*$  hydrogenation rates on Cu (111) (e) and Cu (211) (f) surfaces. All simulations were conducted at 503 K, with a fixed gas ratio of  $\text{H}_2 : \text{CO}_2 : \text{CO} = 6 : 1 : 1$ .



while in others, it has no significant effect or even exerts a positive influence?

To resolve these conflicting observations, we conducted grand potential simulations to explore the effect of  $\text{H}_2\text{O}$  pressure on  $\text{HCOO}^*$  hydrogenation. As shown in Fig. 8 and S10,<sup>†</sup>  $\text{H}_2\text{O}$  tends to accumulate more readily on the Cu surface than  $\text{CO}_2$ . Even a small amount of  $\text{H}_2\text{O}$  can significantly impact the grand potential barrier and the surface densities of gas compounds on both Cu (111) and Cu (211) surfaces. On the Cu (111) surface, the  $\text{HCOO}^*$  hydrogenation rate initially increases before declining, while on the Cu (211) surface, the  $\text{HCOO}^*$  hydrogenation rate sharply decreases with the addition of  $\text{H}_2\text{O}$ , consistent with current understanding. However, due to the strong adsorption of  $\text{H}_2\text{O}$ , even a small  $\text{H}_2\text{O}$  pressure (0.07 bar) leads to surface saturation. Beyond this point, the  $\text{HCOO}^*$  hydrogenation rate remains nearly unchanged, regardless of additional  $\text{H}_2\text{O}$ . Therefore, when the methanol yield is high and CO is lacking, the byproduct  $\text{H}_2\text{O}$  can reach its saturation point, meaning that further addition of  $\text{H}_2\text{O}$  does not change the reaction rate. In fact, the strong interaction of  $\text{H}_2\text{O}$  molecules with intermediate species might even positively influence the reaction under certain conditions. As a result, when the  $\text{H}_2\text{O}$  pressure in the system is near zero, adding CO and  $\text{H}_2\text{O}$  may reduce the  $\text{CH}_3\text{OH}$  production rate. However, at high  $\text{H}_2\text{O}$  partial pressure, the addition of CO and  $\text{H}_2\text{O}$  could potentially increase the  $\text{CH}_3\text{OH}$  production rate. Additionally, it should be noted that water can directly participate in the reaction. This effect should be explicitly captured through KS-DFT calculations involving  $\text{H}_2\text{O}$  as a reactant in the reaction network.<sup>44,45</sup>

### 3 Conclusions

In this study, we combined electronic DFT and cDFT calculations to investigate the effects of various reaction conditions on  $\text{CO}_2$  hydrogenation to  $\text{CH}_3\text{OH}$  over Cu (111) and Cu (211) surfaces. The grand potential approach enables to bridge the “pressure gap” between first principles calculations and experimental observations at conditions relevant to industrial practice. From the perspectives of grand potential landscape and surface concentrations, our findings provide several key insights into the kinetics of  $\text{CO}_2$  hydrogenation under the influence of temperature, pressure, and gas composition.

First, the hybrid DFT calculations account for the thermodynamic effects on the grand potential barriers and local gas concentrations. The grand potential corrections have a significant impact on the formation of different intermediates, particularly at low temperature and high pressure, with the absolute values comparable to conventional corrections for the bond vibrational entropy. We demonstrated that the grand potential profiles align better with experimental data in terms of both surface composition and reaction kinetics. It is the surface density, rather than bulk concentration, determines the hydrogenation rates.

Our grand-potential simulations revealed that the surface density of  $\text{H}_2$  is lower than its bulk density, increasing linearly without reaching saturation, whereas the surface density of  $\text{CO}_2$  is two orders of magnitude higher than its bulk density and easily

becomes saturated.  $\text{CO}_2$  does not influence the  $\text{HCOO}^*$  hydrogenation rate through its surface concentration, as the  $\text{HCOO}^*$  coverage is already saturated. Instead, two opposing effects—the reduction in the  $\text{HCOO}^*$  hydrogenation energy barrier and the decrease in the surface  $\text{H}_2$  density—collectively influence the hydrogenation rate. This also explains why  $\text{HCOO}^*$  could not continue to hydrogenate in a  $\text{CO}_2$ -free gas atmosphere. Consequently, we conclude that the ideal  $\text{H}_2 : \text{CO}_2$  ratio decreasing with temperature and increasing with  $\text{H}_2$  pressure.

We clarified that CO does not directly compete with  $\text{CO}_2$  for active sites but instead competes for physical adsorption on the Cu surface. On the Cu (211) surface, increasing the  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratio leads to both an increased energy barrier for  $\text{HCOO}^*$  hydrogenation and a higher  $\text{H}_2$  surface density. At low  $\text{H}_2$  pressure, the  $\text{HCOO}^*$  hydrogenation rate is unaffected by the  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratio. However, at high  $\text{H}_2$  pressure, the reaction rate falls as the  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratio increases. Notably, the  $\text{HCOO}^*$  hydrogenation rate rises significantly when the  $\text{CO}/(\text{CO}_2 + \text{CO})$  ratio reaches 0.99. From this, we infer that CO participates in both the water–gas shift (WGS) reaction and  $\text{CO}_2$  hydrogenation pathways to produce  $\text{CH}_3\text{OH}$ .

While  $\text{H}_2\text{O}$  is an inevitable byproduct of  $\text{CO}_2$  hydrogenation, it may exhibit either positive or negative effects on the hydrogenation rate. Although a small amount of  $\text{H}_2\text{O}$  significantly reduces the reaction rate,  $\text{H}_2\text{O}$  adsorption becomes saturated at very low partial pressures, after which the  $\text{HCOO}^*$  hydrogenation rate remains nearly unchanged. Therefore, when the product yield is high and CO is scarce, the byproduct  $\text{H}_2\text{O}$  can reach its saturation point, meaning that further addition of  $\text{H}_2\text{O}$  does not reduce the reaction rate.

In summary, the grand-potential theory provides a comprehensive understanding of  $\text{CO}_2$  hydrogenation to  $\text{CH}_3\text{OH}$  under industrially relevant conditions. By accounting for the influence of gas-phase interactions, surface adsorption effects, and variations in temperature and pressure, our study addresses inconsistencies between theoretical predictions with conventional DFT and experimental observations, offering new insights for optimizing catalyst performance in methanol synthesis. As the thermodynamic properties of inhomogeneous fluids can be systematically described using cDFT, the grand-potential framework can be readily extended to other gas-phase reactions involving heterogeneous catalysis, such as highly active  $\text{CO}_2$  hydrogenation catalysts like CuO and PdZn alloys.

### Data availability

Additional data can be obtained from the authors upon request.

### Author contributions

J. S. and J. W. designed the research, discussed the results, and wrote the paper. J. S. performed the calculations.

### Conflicts of interest

There are no conflicts to declare.

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