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Bridging the Catalyst Reactivity Gap Between Au and Cu for the Reverse Water Gas Shift Reaction

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Abstract

The reverse water gas shift reaction (rWGSR) is highly relevant for CO₂ utilization in sustainable fuel and chemical production. Both Au and Cu are interesting for rWGSR catalysis, but it turns out that the reactivities of Au and Cu are very different. In this study, we consider alloys made from Au, Ag, Cu, Pt, and Pd to identify surfaces with reactivities for CO₂ dissociation in between Cu(111) and Au(111). Additionally, interesting alloy surfaces should have activation energies for CO₂ dissociation that are only a little higher than the endothermic reaction energy. We find that certain Cu based alloys with Ag and Au meet these criteria, whereas alloys containing Pt or Pd do not. The low additional cost in activation energy occurs when the transition state and final state configurations are made to look very similar due to the placement of the different metal elements in the surface. Finally, we construct a kinetic model that compares the rate of the rWGSR to the estimated rate of unwanted side reactions (i.e. methane formation or coking) on Ag-Cu alloy surfaces with varying composition and random placement of the Ag and Cu atoms. The thermodynamics favor methane formation over rWGSR, but the model suggests that Ag-Cu alloy surfaces are highly selective for the rWGSR.

TOC Graphic



1. Introduction

CO₂ hydrogenation to CO via the reverse water gas shift reaction (rWGSR, eq 1) could be important for making the chemical industry more sustainable. The rWGSR is both a way to convert CO₂ into CO, which can be used directly as feedstock in syngas chemistry,¹ and a way to utilize hydrogen, which is the high energy chemical most straightforwardly produced by renewable powered electrochemistry.^{2,3}

$$CO_2(g) + H_2(g) \rightarrow CO(g) + H_2O(g)$$
⁽¹⁾

CO₂ hydrogenation can also form CH₃OH and CH₄ instead of CO.^{4,5} Figure 1 shows reaction free energies to form the three products at the standard state pressure (i.e. 0.1 MPa pressure for all species) and with 2 MPa H₂ pressure. The thermodynamic data in Figure 1 is taken from ref [⁶⁻⁸]. Formation of the gas-phase CO is allowed at high temperatures, with Δ_r G for rWGSR being downhill above 1097 K at standard state pressure and above 629 K at 2 MPa H₂ pressure. On the other hand, CH₃OH is only stable (Δ_r G < 0) at lower temperatures, i.e. below 267 K at standard state pressure and below 452 K with 2 MPa H₂ pressure. The formation of CO rather than CH₃OH can therefore largely be achieved by controlling the reaction conditions. More importantly, the equilibrium strongly favors CH₄ formation rather than CO formation below ~900 K or at increased H₂ pressure.^{9,10} It is therefore important to have a catalyst that is selective towards CO instead of CH₄ when the goal is the rWGSR.



Figure 1: Reaction free energies^{6–8} for CO₂ hydrogenation to CO (rWGRS), CH₃OH, and CH₄ at the standard state pressure (solid lines) and with 2 MPa H₂ pressure (dashed lines). The dashed lines are shifted from the solid lines by $\Delta n_{\text{H2}} \cdot k_{\text{B}} \text{T} \cdot \ln(20)$, where Δn_{H2} is the change in H₂(g) molecules in the reactions.

The main reaction step of interest is the initial CO_2 dissociation on the catalyst surface (reaction eq 2). $CO_2(g) + 2^* \rightarrow *CO + O^*$ (2) In addition, two reaction steps concerning the fate of *CO are worth considering, namely the possible dissociation of *CO to *C and O* (reaction eq 3) and desorption of *CO to the gas phase (reaction eq 4) and desorption (reaction eq 4

4).

$$*CO + * \rightarrow C* + O* \tag{3}$$

$$*CO \rightarrow CO(g) + * \tag{4}$$

Under normal rWGSR conditions the rate of CO_2 conversion to CO on transition metal catalysts is mainly controlled by the ability of the metal surface to catalyze reaction eq 2.^{11–14} In Figure 2a, we have calculated reaction energy and activation energy for reaction eq 2 on pure transition metal (111) surfaces and Cu(100). The reactive (strong binding) Pt, Pd, and Cu have low reaction energy cost (ΔE) and low activation energy (E_{TS}) compared to the inert (weak binding) Au and Ag metals. However, the selectivity towards the different CO₂ hydrogenation products is more strongly determined by the interaction between the metal surface and CO. Strong interaction could lead to C-O bond dissociation (reaction eq 3), which favors coking and CH₄ formation, whereas weak interaction favors CO desorption (reaction eq 4).¹ Metals are separable based on the energy cost for *CO desorption on one axis and the energy cost for *CO dissociation to C* and O* on the other, which we have calculated and plotted in Figure 2b. The inert Au(111), Ag(111), and to some extent Cu(111) favor *CO desorption and are expected to be more selective towards CO formation than the reactive Pd(111), Ni(111), Pt(111), Rh(111), and Ru(111) metals where C-O bond dissociation is not exceedingly costly. Indeed ref ¹⁵ found that Ag has lower rWGSR activity than Pd, Ni and Cu, but that Ag and Cu have higher selectivity towards CO than Pd and Ni. It was also recently shown that the Ni selectivity for the rWGSR can be increased by incorporating Zn. The presence of Zn weakens the CO adsorption energy on the Ni atoms, allowing for faster CO desorption.¹⁶

Experiments¹⁷ have found that Cu(100) is better at facilitating CO₂ dissociation than Cu(111), which is also what we would expect from Figure 2a. However, Figure 2b suggests that Cu(100) also has a higher tendency to dissociate CO, which is unwanted, since the CO dissociation energy is much less costly on Cu(100) than on Cu(111).

With the thermodynamic and catalytic aspects in mind, experiments have obtained 50% or more conversion of CO₂ and >95% selectivity towards CO on both Au based (TiO₂-supported Au nanoparticles)¹⁸ and Cu based (nanoceria-supported Cu nanoparticles)¹⁹ catalysts at 673 K.¹ SiO₂-supported Pd-Fe nanoparticles are almost as good with 45% CO₂ conversion and 97% CO selectivity at 723 K,²⁰ but most other catalysts^{1,21} either lack CO₂ conversion, CO selectivity, or utilize high temperatures where the equilibrium is shifted more towards CO formation.



Figure 2: (a) Reaction energies (ΔE) and activation energies (E_{TS}) for CO₂(g) dissociation to *CO + O* on pure metal (111) surfaces and Cu(100). The ΔE and E_{TS} are explained by the insert. (b) Reaction energies for *CO + * \rightarrow C* + O* versus reaction energies for *CO desorption on pure metal (111) surfaces and Cu(100). The dashed vertical lines show the large reactivity gap between Cu(111) and Au(111).

Cu^{12–15,19} and Au^{18,22–24} are both interesting for the rWGSR catalysis, so it is curious that there is such a large reactivity gap between these two metals. This is especially the case for reaction eq 2 on (111) surfaces (Figure 2a), where the reaction energy differs by 1.82 eV. Candidate catalysts with reactivities in this reactivity gap would likely be interesting for the rWGSR. According to our calculations, Ag(111) is slightly more reactive than Au(111), but there is no other pure metal in between Cu(111) and Au(111). One possibility is to use Au nanoparticles, which are expected to have increased reactivity compared to larger Au particles.²⁵ Actually, Au seems to be so inert that it works best as nanoparticles supported on oxide co-catalysts, where the oxide helps facilitate the CO₂ dissociation.^{26,27} Unfortunately, nanoparticles often suffer more than larger particles from metal

oxidation, particle sintering and restructuring at high temperatures, which negatively affect their catalytic performance.^{28–31}

Another possibility is to alloy two or more metals. Alloys are especially interesting because it might be possible to modify their activity and selectivity for rWGSR by changing their precise metal composition. Many bimetallic nanoparticles have recently been studied for the rWGSR, for instance, unsupported FeCo,³² unsupported CuNi,³³ CuNi on Al₂O₃,³⁴ NiAu on SiO₂,³⁵ PtCo on TiO₂,³⁶ and CuIn on ZrO₂.³⁷

In this study, we consider CO₂ dissociation (reaction eq 2) on (111) surfaces of alloys with up to four elements picked from Au, Ag, Cu, Pt, and Pd. We find multiple alloys with reactivities in between those of Au and Cu, indicating that the discrete reactivities of pure metals become a continuous reactivity distribution in the broad alloy composition space. This is not too surprising as it has already been shown that the adsorption energies of *CO and O* have broad distributions on multimetallic alloys.^{38,39} However, we also find that the activation energy for CO₂ dissociation can be lowered in a very intuitive way by using an alloy surface that makes the transition state and final state look more similar. Alloys made from Ag, Cu, and Au are found to be most interesting for the rWGSR, as they span the reactivity gap between Cu and Au, have activation energies only slightly higher than the energy cost for CO₂ dissociation, and have low tendency to facilitate C-O bond dissociation. The low rate to facilitate C-O bond dissociation compared to the CO₂ to CO conversion rate on Ag-Cu alloy surfaces is confirmed by a kinetic model that considers both processes.

2. Methods

2.1 Computational details

We have performed density functional theory (DFT) calculations at the generalized gradient approximation (GGA) level with the Grid-based Projected Augmented Wave (GPAW) code.⁴⁰ The

RPBE functional⁴¹ is used to describe exchange and correlation effects. The wave functions are expanded in plane waves with a 440 eV energy cutoff and the Brillouin zone is sampled with a Monkhorst-Pack grid of $2 \times 2 \times 1$ k-points for surface slabs and Γ -point for gas-phase molecules.

We consider the reactivity of four types of FCC alloy slabs with the compositions A_3B , AB, A_2BC and ABCD, in addition to the pure metal slabs. The slabs have (4×4) surface cells exposing the (111) facet (Figure 3) and four atomic layer thicknesses. The A_3B , AB, A_2BC and ABCD bulk unit cells that the slabs are constructed from are shown in Supporting Information section S6. We use the Atomic Simulation Environment (ASE) package⁴² to construct structural motifs such as cutting the alloy slabs from the bulk structures and adding adsorbates to the surfaces. We note that the B atoms in A_3B are sitting in a different surface pattern than the B/C atoms in A_2BC and the A/B/C/D atoms in ABCD (Figure 3). A consequence of this is that CO_2 can dissociate at four adjacent A atoms in A_3B , which is not possible on the AB, A_2BC , and ABCD surfaces.

We add 16 Å of vacuum in the z direction between the periodic images of the slabs. The atoms in the two bottom layers of the slabs are fixed in the bulk FCC positions, while the top two atomic layers are allowed to relax during geometry optimization. All the surfaces are optimized until the maximum force is lower than 0.03 eV/Å, while gas-phase molecules are relaxed to a maximum force of 0.01 eV/Å. The alloy lattice parameters are approximated as the composition weighted average of the DFT calculated lattice parameters for the pure metals (see Supporting Information section S6 for details).⁴³⁻⁴⁵

The computational details are similar to those used in as in ref [⁴⁶], and the DFT energies can be directly compared. All structures and scripts for plots presented in this work are stored in a database, which can be found online at <u>https://nano.ku.dk/english/research/theoretical-electrocatalysis/katladb/</u> <u>bridging-the-catalyst-reactivity-gap</u>.



Figure 3: Illustration of (4×4) surface cells exposing the (111) facet for A₃B, AB, A₂BC and ABCD.

Figure 4 shows the DFT energy landscape for the rWGSR on Cu(111), Au(111) and AuCu. With our DFT parameters, the rWGSR has a DFT energy cost of 0.73 eV (independent of the surface), which is somewhat overestimated given the experimentally derived 0 K reaction enthalpy of $\Delta H^{0}_{0K} = 0.42 \text{ eV}.^{6}$ The DFT error in the rWGSR reaction energy is well-known and present in all the commonly used GGA functionals.⁴⁷

We mainly focus on the $CO_2(g) \rightarrow *CO + O^*$ part of rWGSR, for which, the reaction energy (ΔE) and activation energy (E_{TS}) are important. On Cu(111), we calculate ΔE and E_{TS} to be 1.30 eV and 1.87 eV (Figure 4a). With the PBE exchange-correlation functional instead of RPBE, the same values are 1.27 eV and 1.60 eV.⁴⁸ Consequently, we will predict slower CO₂ dissociation kinetics compared to studies using PBE, because we have a larger E_{TS} value. On Au(111), we calculate ΔE and E_{TS} to be 3.12 eV and 3.41 eV (Figure 4b). Again, our values are higher than the values calculated with PBE (2.86 eV and 3.16 eV).⁴⁸ Finally, we show the DFT energy landscape for rWGSR on AuCu, as an example of an alloy surface with ΔE between Cu(111) and Au(111) (Figure 4c). The ΔE and E_{TS} values are 2.13 eV and 2.30 eV on AuCu. We note that since CO₂ dissociation is uphill for the catalysts of interest, we have that $E_{TS} > \Delta E > 0$. A catalyst with a desirable ΔE , therefore always has E_{TS} at least as big as ΔE no matter how optimized the catalyst is.



Figure 4: DFT energy diagram for rWGSR on (a) Cu(111), (b) Au(111) and (c) AuCu surfaces. E_{TS} is the energy of the transition state compared to the energy of the reactants (CO₂(g) and H₂(g)). ΔE is the reaction energy of the CO₂(g) \rightarrow *CO + O* reaction step. Dashed lines indicate reaction barriers that we have not investigated in this study.

2.2 Computational methodology

Our first goal is to find alloy surfaces with ΔE in the reactivity gap between Au(111) and Cu(111). We therefore utilize a simple screening model developed for the water gas shift reaction, which uses

adsorption energies of O* and *CO on pure metal surfaces to estimate ΔE and E_{TS} for the possible alloy surfaces.⁴⁶ The alloys that are estimated to have ΔE in the reactivity gap and small $E_{TS} - \Delta E$ are further analyzed (see Supporting Information section S1 for details).

For each interesting alloy surface, we look for the most stable dissociated CO₂ configuration with the *CO and O* species at adjacent surface sites (*CO at an ontop site and O* at a hollow site). As an example, 24 configurations with co-adsorbed *CO and O* have been calculated on the AuCu₂Pd(111) surface to find the most stable configuration (Figure S18 in the Supporting Information). The path from the initial state (IS) with CO₂ in the gas phase to the most stable dissociated final state (FS) will according to the Brønsted-Evans-Polanyi (BEP) relation,^{49–52} often have the most stable transition state (TS) and therefore the lowest activation energy. The TS between molecular CO₂ and dissociated *CO and O* is obtained with the Climbing Image Nudged Elastic Band (CI-NEB) method.⁵³ In addition to the pathway to the most stable FS, we investigate pathways to other FS that are within ~0.02 eV of the most stable FS, if we suspect that their TS are more stable. We only present the pathway with the most stable TS.

We also use the insight presented in this study to obtain two pathways that have significantly less stable FS, but (in the case of Ag₃Cu) a more stable TS or (in the case AgCu₃) a TS within 0.05 eV of the most stable TS. These are denoted with the superscript "StableTS".

Finally, we use the Ag_3Cu , AgCu, and $AgCu_3$ alloy surfaces to calculate E_{TS} for a range of possible Ag-Cu containing active sites and use these values in a kinetic model for Ag-Cu alloy surfaces with varying composition and random placement of the Ag and Cu atoms.

3. Results and discussion

3.1 Bridging the reactivity gap between Au and Cu.

Even though we have limited our study to five different elements and four types of (111) alloy surfaces, we find multiple surfaces with reactivities in between that of Au(111) and Cu(111). In Figure 5 we

mark the considered alloy surfaces with circles based on their ΔE and $E_{TS} - \Delta E$ values. We chose to plot $E_{TS} - \Delta E$ instead of E_{TS} , because the optimal value of $E_{TS} - \Delta E$ is 0, whereas the optimal value of E_{TS} is ΔE . The largest gap in ΔE between alloy (or pure) surfaces is reduced to 0.22 eV (between Ag₃Cu and AgAuCu₂), and interestingly, it is possible to change ΔE significantly without affecting $E_{TS} - \Delta E$. This is especially true for alloys composed of Cu, Ag, and Au. For instance, AgCu, AgAuCu₂, AuCu, Ag₂AuCu, AgAu₂Cu, and Ag₃Au span 1.16 eV in ΔE but all have $E_{TS} - \Delta E$ of less than 0.2 eV. This is a substantial break from the linear scaling relation between ΔE and E_{TS} found for the CO₂ dissociation on pure metal surfaces.⁵⁴⁻⁵⁶ Alloys containing Pt and Pd generally seem to be less optimized for the CO₂ dissociation and have $E_{TS} - \Delta E$ of at least 0.46 eV.



Figure 5: DFT reaction energies (ΔE) versus ($E_{TS} - \Delta E$) DFT energies calculated for the CO₂(g) \rightarrow *CO + O* reaction step on all the considered alloy surfaces. Blue circles show ΔE and ($E_{TS} - \Delta E$) for the pathway to the stable FS on each alloy surface. Orange circles show ΔE and ($E_{TS} - \Delta E$) for pathways where the TS is the most stable (in the case of Ag₃Cu^{StableTS}) or within 0.05 eV of the most stable TS (in the case of AgCu₃^{StableTS}), but where the FS is not the most stable.

We now take a closer look at the role of the four adjacent metal atoms involved in the CO₂ dissociation step to understand why many Cu, Ag, and Au containing alloy surfaces have small $E_{TS} - \Delta E$ over a large range of ΔE values. The CO₂ molecule dissociates through a TS where the O atom interacts with two metal atoms (designated B and C) and the CO interacts with one metal atom (designated D).^{48,54,57} When CO₂ is fully dissociated the *CO is adsorbed at atom D, while the O* is adsorbed at atoms B, C and a third metal atom (designated A). The following discussion outlines how the metal atoms can be picked in a way to make TS and FS look more similar and thereby make $E_{TS} - \Delta E$ small.

We start with all four adjacent metal atoms being Ag, e.g. as found on the pure Ag(111) surface (Figure 6a). The Ag atoms are very inert and bind both *CO (on top) and O* (hollow) weakly. The energy to dissociate CO₂ is large (E_{TS} = 2.89 eV), but since there is barely any stabilization by binding the fragments to the Ag(111) surface, ΔE is almost as large as E_{TS} , and $E_{TS} - \Delta E$ becomes small (0.11 eV).

Next, we use the Ag₃Cu alloy surface to consider CO₂ dissociation at a site with three Ag atoms and one Cu atom (Figure 6b). When the Cu atom substitutes an Ag atom at position B or C, ΔE is lowered from 2.78 eV to 2.25 eV, while $E_{TS} - \Delta E$ remains low (0.12 eV). Cu binds O* significantly stronger than Ag, so at the TS where O* is sitting in a bridge configuration at positions B and C, the interaction between O* and Cu is optimized. The O* binds to an additional Ag atom at the FS, but that does not provide strong stabilization. We note that the CO₂ dissociation site with three Ag atoms and one Cu atom has the most stable TS but not the most stable FS on the Ag₃Cu surface, i.e., it does not follow the BEP relation. We discuss the CO₂ dissociation pathway to the stable FS site on Ag₃Cu later.

We continue substituting Ag atoms with Cu atoms and get two Cu at positions B and C (Figure 6c). This site is the most stable *CO + O* adsorption site on the AgCu alloy surface. The second Cu atom lowers ΔE from 2.25 eV to 1.77 eV, but since the extra Cu atom stabilizes both ΔE and E_{TS} , the difference remains small ($E_{TS} - \Delta E = 0.11$ eV).

We substitute again to get three Cu atoms and one Ag atom (Figure 6d). Such a site is found on the AgCu₃ alloy surface. Having two Cu atoms at positions B and C and the third Cu atom at the *CO adsorption site (position D) lowers ΔE a little from 1.77 eV to 1.65 eV. However, not all of this stabilization is not present in E_{TS} giving a larger $E_{TS} - \Delta E$ (0.19 eV). The AgCu₃ alloy surface contains a CO₂ dissociation site with four Cu atoms and a more stable FS than the site with three Cu atoms and one Ag atom, however, the E_{TS} for the three Cu atoms and one Ag atom are within 0.05 eV of the most stable TS.

Substituting the last Ag for Cu to obtain a site with four Cu atoms (as on the Cu(111) surface) changes ΔE from 1.65 eV to 1.30 eV (Figure 6e). However, it barely changes the E_{TS} (from 1.84 eV to 1.87 eV) since the fourth Cu atom interacts very limited with the O* atom in the TS. The $E_{TS} - \Delta E$, therefore, increases to 0.57 eV. We generally find that the element at position A, which mostly binds the O* atom in the FS, influences the TS energy very little. Consequently, (111) alloy surfaces can be optimized for catalyzing the normal water gas-shift reaction by having reactive atoms at position B and C, and more inert atoms at position A.^{46,57}



Figure 6: Illustration of TS and FS for CO₂ dissociation on (a) Ag, (b) Ag₃Cu^{StableTS}, (c) AgCu, (d) AgCu₃^{StableTS}, and (e) Cu. The FS shown for Ag, AgCu, and Cu are the stable FS on these alloy surfaces. The FS for Ag₃Cu^{StableTS} and AgCu₃^{StableTS} are not the most stable, since these surfaces have sites with one extra Cu atom, however the TS for Ag₃Cu^{StableTS} is the most stable and the TS for AgCu₃^{StableTS} is within 0.05 eV of the most stable TS found.

As mentioned earlier, the CO₂ dissociation site with three Ag atoms and one Cu atom (Figure 6b) does not have the most stable FS on the Ag₃Cu surface. Due to the makeup of the A₃B type surface cell (Figure 3), the Ag₃Cu surface also contains a CO₂ dissociation site, where the four adjacent metal atoms are two Cu and two Ag atoms, with the two Ag at positions B and C binding to O* in the TS state (Figure 7a). This adsorption site has a more stable FS than the CO₂ dissociation site with one Cu and three Ag atoms, but a less stable TS, since it is the Ag atoms and not the Cu atoms that binds to O* at the TS. This is an additional example that the scaling relation between ΔE and E_{TS} is subject to the precise makeup of the CO₂ dissociation site, but in this case, it is broken in to give a higher E_{TS} .

Figure 7b and Figure 7c illustrate what happens when Ag is exchanged for Au. In Figure 7b the *CO binding D position is changed from Ag to Au compared to Figure 6b. ΔE increases slightly from

2.25 eV to 2.41 eV, but the destabilization from Au is present both in the FS and the TS, so the ($E_{TS} - \Delta E$) energy only changes from 0.12 eV to 0.09 eV. The situation is similar when a second Ag at position B is changed to Au (Figure 7c). ΔE increases from 2.41 eV to 2.58 eV, but $E_{TS} - \Delta E$ is unchanged.

Figure 7d and Figure 7e illustrate what happens when Ag is exchanged for Pt or Pd. Both Pt and Pd bind the CO molecule very strongly, so the most stable FS is obtained by having Pt or Pd at position D. The substitution of the Ag for Pt (or Pd) lowers ΔE from 2.25 eV to 1.07 eV (or 1.38 eV), however not all the stabilization is present at the TS so $E_{TS} - \Delta E$ increases from 0.12 eV to 0.51 eV.



Figure 7: Illustration of TS and FS for CO₂ dissociation on (a) Ag₃Cu, (b) Ag₂AuCu, (c) AgAu₂Cu, (d) Ag₂CuPt, and (e) Ag₂CuPd. All the shown FS are the stable FS on these alloy surfaces.

It has been found that the reaction and activation energy for O₂ dissociation on multimetallic surfaces with random placement of the elements show scaling between the two sets of values, but with points being scattered around the optimum scaling line.³⁹ Our results for CO₂ dissociation indicate that the breaking of scaling relations can be understood by analyzing the makeup of the catalytic surface

sites. Consequently, the scaling relation is likely only a property that arises as the average behavior of all the possible surface site combinations.

3.2 Kinetic model for the rWGSR on Ag-Cu alloy surfaces.

The Ag-Cu alloy surfaces seem interesting for the rWGSR given that they can dissociate CO₂ with a range of reaction energies, have activation energies only slightly higher than the energy cost for CO₂ dissociation, and are made of elements with low tendencies to facilitate C-O bond dissociation. To validate these statements, we construct a kinetic model to obtain both the CO₂ dissociation rate and the CO dissociation rate on Ag-Cu alloy surfaces, hereby estimating both how active and selective the surfaces are for CO formation rather than methane formation or coking. The model includes CO₂ adsorption / desorption (eq 5), CO_2 dissociation / recombination (eq 6), CO desorption / adsorption (eq 7), and O* hydrogenation / dehydrogenation (eq 8). We follow the example for CO oxidation in [⁵⁸] and assume that reactions eq 5, 7, and 8 are in quasi-equilibrium and that eq 6 is the rate determining step for the rWGSR. We ignore H* species and collect O* hydrogenation in (eq 8). It should be acceptable to neglect H* species, because the coverage of H* is low on Cu(111) and Ag-Cu alloys, and because the presence of H* doesn't seem to facilitate C-O bond breaking in CO₂ with lower barrier than the direct CO₂ dissociation to *CO and O* (see Supporting Information section S5 for details).⁵⁹ Finally, the kinetic model also includes *CO dissociation (eq 9), which we assume is irreversible and results in unwanted coking or methane formation.

$$CO_2(g) + * \rightleftharpoons *CO_2$$
 (5)

$$*CO_2 + * \rightleftharpoons *CO + O*$$
 (rate determining step) (6)

$$*CO \rightleftharpoons CO(g) + * \tag{7}$$

$$O^* + H_2(g) \rightleftharpoons H_2O(g) + *$$
(8)

*CO + * \rightarrow C* + O* (irreversible) (9)

Most experiments are conducted at atmospheric pressure with high concentration of H₂ rather than high pressure of H₂.¹ We therefore assume that the rWGSR is carried out at 700 K and a total pressure of 0.1MPa, that the inlet gas mixture is 9:1 H₂ to CO₂, but that 40% of the CO₂ has been converted to CO and H₂O giving partial pressures of $p_{H2} = 0.86$, $p_{CO2} = 0.06$, $p_{CO} = p_{H2O} = 0.04$ (the equilibrium is $p_{H2} = 0.84$, $p_{CO2} = 0.04$, and $p_{CO} = p_{H2O} = 0.06$, as derived in the Supporting Information section S4).

We also need to decide the surface makeup of the Ag-Cu alloys. We note that the Ag-Cu phase diagram shows that at 700 K (and below) only bulk Ag with a small concentration of Cu and bulk Cu with a small concentration of Ag is thermodynamically stable.⁶⁰ However, several experimental groups have managed to synthesize Ag-Cu alloy nanoparticles with varied compositions and used these for catalysis.^{61,62} An interesting limiting case is a solid solution, where the Ag and Cu atoms are randomly distributed in the surface with x_{Cu} and x_{Ag} molar fractions (where $x_{Ag} = 1 - x_{Cu}$). The CO₂ dissociation rate is mainly determined by the four metal atoms at the dissociation site, so to describe an Ag-Cu solid solution we need to obtain ΔE and E_{TS} for all 16 combinations of Ag or Cu at the A, B, C, D sites. We have done this by conducting CI-NEB calculations for CO₂ dissociation at different positions on the Ag(111), Ag₃Cu, AgCu, AgCu₃, and Cu(111) surfaces and collected the results in Table 1. We ignore changes in entropy and zero-point energies (ZPE) between the IS and TS, but account for this in the backwards reaction by using ΔG instead of ΔE in the calculation of the backwards activation energy. Furthermore, we note that the experimental enthalpy for the full rWGSR is poorly reproduced with DFT and we estimate a DFT reaction energy error (ΔE_{error}) of 0.45 eV at 700 K (derived in the Supporting Information section S4). We also correct for this error in the backwards activation energy such that the activation energy for CO₂ recombination becomes E_{TS} - (ΔG - ΔE_{error}). This ensures that we get the correct overall thermodynamic for the rWGSR. We can now estimate the forward (k_6^+) and backward (k_6) rate constants for the reaction eq 6 at each A, B, C, D site using the formula from transition state theory (eq 10).⁶³

$$k_6^{+} = \frac{k_B T}{h} \cdot \exp\left(\frac{-E_{TS}}{k_B T}\right) , \qquad k_6^{-} = \frac{k_B T}{h} \cdot \exp\left(\frac{-(E_{TS} - (\Delta G - \Delta E_{error}))}{k_B T}\right)$$
(10)

The reaction rate for eq 6 (r_6 , eq 11) further depends on the coverages of CO₂ (θ_{CO2}), *CO (θ_{CO}), and O* (θ_0), which due to the quasi-equilibrium also can be expressed by the equilibrium constants for reaction eq 5, 7, and 8 (K_5 , K_7 , and K_8 , respectively), and the p_{H2} , p_{CO2} , p_{CO} , p_{H2O} partial pressures.

$$r_6 = k_6^+ \theta_{\rm CO2} \theta_* - k_6^- \theta_{\rm CO} \theta_{\rm O} = k_6^+ K_5 p_{\rm CO2} - k_6^- K_7^{-1} p_{\rm CO} K_8^{-1} p_{\rm H2O} p_{\rm H2}^{-1}$$
(11)

We assume that $*CO_2$ has 0 eV DFT adsorption energy on all surfaces in the calculation of K_5 , and use the adsorption energies for *CO and O* adsorbed at the 16 different A, B, C, D sites to calculate K_7 and K_8 . The *CO is adsorbed at position D, the O* is adsorbed at positions A, B, C and we ignore coadsorption interaction between the two species. Entropy, heating, and ZPE corrections are obtained from [^{6,64,65}] and applied to all DFT reaction energies (see Supporting Information section S4 for details). The kinetic values for CO_2 dissociation at the 16 different A, B, C, D sites are collected in Table 1.

Table 1: CO₂ dissociation (eq 6) reaction rates (r_6) and backward and forward activation energies (E_{TS} , E_{TS} - (ΔG - ΔE_{error})), rate constants (k_6^+ , k_6^-), and *CO (θ_{CO}) and O* (θ_0) coverages for every combination of the A, B, C, D sites. The properties are calculated on the specified alloy surfaces.

Alloy surface	A, B, C, D site E	E _{TS} (eV)	k_6^+ (s ⁻¹)	E_{TS} - (ΔG -	k_6^- (s ⁻¹)	$ heta_{ m CO}$	$ heta_{ m O}$	$r_{6}(s^{-1})$
				ΔE_{error}) (eV)	I			
Ag(111)	Ag, Ag, Ag, Ag	2.89	2.27E-08	0.21	4.49E+11	1.08E-10	1.55E-17	4.68E-15
Ag ₃ Cu	Ag, Ag, Ag, Cu	2.67	8.72E-07	0.35	4.41E+10	3.64E-07	4.83E-18	1.31E-13
Ag ₃ Cu	Ag, Ag, Cu, Ag	2.37	1.26E-04	0.22	3.80E+11	1.08E-10	7.24E-14	2.71E-11
AgCu	Ag, Ag, Cu, Cu	2.09	1.31E-02	0.27	1.66E+11	1.35E-07	7.24E-14	1.50E-09
Ag ₃ Cu	Ag, Cu, Ag, Ag	2.37	1.26E-04	0.22	3.80E+11	1.08E-10	7.24E-14	2.71E-11
AgCu	Ag, Cu, Ag, Cu	2.09	1.31E-02	0.27	1.66E+11	1.35E-07	7.24E-14	1.50E-09
AgCu	Ag, Cu, Cu, Ag	1.88	4.25E-01	0.21	4.49E+11	4.08E-10	1.26E-10	7.85E-08
AgCu ₃	Ag, Cu, Cu, Cu	1.84	8.25E-01	0.29	1.19E+11	3.10E-07	4.56E-12	2.86E-08

Ag ₃ Cu	Cu, Ag, Ag, Ag	2.8	1.01E-07	0.6	6.98E+08	1.08E-10	7.24E-14	1.87E-14
Ag ₃ Cu	Cu, Ag, Ag, Cu	2.58	3.88E-06	0.81	2.15E+07	3.64E-07	7.24E-14	3.60E-13
AgCu	Cu, Ag, Cu, Ag	2.2	2.11E-03	0.52	2.63E+09	4.08E-10	1.26E-10	3.69E-10
AgCu ₃	Cu, Ag, Cu, Cu	2.29	4.75E-04	0.85	1.11E+07	3.10E-07	4.56E-12	9.78E-11
AgCu	Cu, Cu, Ag, Ag	2.2	2.11E-03	0.52	2.63E+09	4.08E-10	1.26E-10	3.69E-10
AgCu ₃	Cu, Cu, Ag, Cu	2.29	4.75E-04	0.85	1.11E+07	3.10E-07	4.56E-12	9.78E-11
AgCu ₃	Cu, Cu, Cu, Ag	2.12	7.95E-03	0.6	6.98E+08	1.51E-10	4.08E-09	1.47E-09
Cu(111)	Cu, Cu, Cu, Cu	1.87	5.02E-01	0.67	2.19E+08	3.10E-07	6.61E-10	7.50E-08

In the solid solution where the Cu and Ag atoms are randomly placed, each A, B, C, D site has a probability to occur given by the product of the molar fractions ($P_{A,B,C,D} = x_A x_B x_C x_D$). As an example, the A, B, C, D = Cu, Cu, Cu, Cu site has a probability of 1 on a pure Cu surface ($x_{Cu} = 1$) and a probability of 0.0625 on a surface with $x_{Cu} = 0.5$. The CO₂ dissociation rates on the Ag-Cu solid solution surfaces are then given by the rates of the different A, B, C, D sites weighted by their probability (eq 12).

$$r_{6,tot} = \sum \mathbf{x}_{A} \mathbf{x}_{B} \mathbf{x}_{C} \mathbf{x}_{D} \cdot r_{2} (\mathbf{x}_{A} \mathbf{x}_{B} \mathbf{x}_{C} \mathbf{x}_{D})$$
(12)

In Figure 8 we plot the CO₂ dissociation rate as a function of the Cu molar fraction in the surface. The plot shows a strong improvement in activity by adding a small amount of Cu to pure Ag, followed by a more gradual additional improvement when going towards pure Cu. We also plot the smallest and highest activity of (A, B, C, D) sites with 3:1 Ag:Cu, 2:2 Ag:Cu and 1:3 Ag:Cu. At $x_{Cu} = 0.50$ the solid solution is a little less active than the (Ag, Cu, Cu, Ag) site and at $x_{Cu} = 0.75$ it is a little more active than the (Ag, Cu, Cu, Site. However, interestingly, at $x_{Cu} = 0.25$ the solid solution is significantly more active than the (Ag, Ag, Cu, Ag) site. This highlights that in a solid solution with a small amount of Cu, there is still a small occurrence of sites with more Cu and higher activity (such as (Ag, Cu, Cu, Ag)) that due to the exponential nature of activity strongly improves the activity of the solid solution.

Overall, the CO₂ dissociation rates predicted by our kinetic model are very slow even for the pure Cu(111) surface. Some of this may be due to the high E_{TS} predicted by RPBE compared to PBE. However, it may also be that more reactive Cu facets are required for faster rates, for instance the Cu(100) surface¹⁷ or even the steps⁶⁶ on Cu(100) surfaces. Although these facets are also likely to dissociate CO faster.

We can estimate the tendency of the Ag-Cu solid solution to dissociate *CO (eq 9) in much the same way as for *CO₂ dissociation, however, we must add two extra approximations. The first is that *CO dissociation is irreversible. This approximation is needed because we do not consider reactions with *C to give methane or coke and therefore do not know how fast the *C species are depleted. The second is to use the reaction energies ($\Delta E_{(*CO \rightarrow C^*+O^*)}$) for *CO dissociation instead of the activation energies. We do this because we have not performed CI-NEB calculations for *CO dissociation. Together these two approximations will give an upper limit for the *CO dissociation rate.

Both C* and O* adsorb in hollow sites on the surfaces, so six metal atoms (denoted A, B, C, D, E, F) are involved in the *CO dissociation reaction. However, we choose to divide the six metal atoms into two subgroups of three metal atoms by assuming that *CO adsorb at position D, that O* adsorb at positions A, B, C, that C* adsorb at positions D, E, F, and that O* and C* do not interact. With this separation $\Delta E_{(*CO_{-}C^{*+O^{*}})}$ is then given by eq 13.

$$\Delta E_{(*CO \to C^{*}+O^{*})}(A, B, C, D, E, F) = \Delta E_{O^{*}}(A, B, C) + \Delta E_{C^{*}}(D, E, F) - \Delta E_{*CO}(D)$$
(13)

There are 64 possible (A, B, C, D, E, F) combinations, but $\Delta E_{(*CO \rightarrow C^{*+O^{*}})}$ is obtained from just four $\Delta E_{0^{*}}(A, B, C)$ values, four $\Delta E_{C^{*}}(D, E, F)$ values, and two $\Delta E_{*CO}(D)$ values. There is some ambiguity in which alloy surfaces we use to represent the different A, B, C, and D, E, F sites, but we choose to use the least stable adsorption energies for *CO and the most stable adsorption energies for O* and C* to again get an upper limit for the CO dissociation rates (see Supporting Information section S4 for details).

The CO dissociation rate on the Ag-Cu solid solution surfaces is also given by the rates of the different (A, B, C, D, E, F) sites weighted by their probability. In Figure 8 we plot the CO dissociation rate as a function of the Cu molar fraction in the surface. The CO dissociation rate is more than 5 orders of magnitude slower than the CO_2 dissociation rate, which strongly indicates that the Ag-Cu solid solution surfaces will be highly selective towards the rWGSR rather than methane formation or coking. However, the two rates have the same overall trends, highlighting that it is difficult to increase the rate for CO_2 dissociation without also getting more CO dissociation.



Figure 8: CO₂ and CO dissociation rates on Ag-Cu solid solution surfaces with varying composition and random placement of the Ag and Cu atoms. The plot also shows the activity of pure Ag(111) (labeled (Ag, Ag, Ag, Ag)) and pure Cu(111) (labeled (Cu, Cu, Cu, Cu)) and the smallest and highest activity from specific A, B, C, D sites with 3:1 Ag:Cu, 2:2 Ag:Cu, and 1:3 Ag:Cu composition. The reaction conditions modeled are 700 K and p_{H2} = 0.86, p_{CO2} = 0.06, p_{CO} = p_{H2O} = 0.04 partial pressures.

4. Conclusion

The Au(111) and Cu(111) reactivity for CO₂ dissociation differ by almost 2 eV even though both Au and Cu are used for rWGSR catalysis. We find that multimetallic (111) alloy surfaces made from Au,

Ag, Cu, Pt and Pd can largely fill the reactivity gap. Alloys consisting of Cu, Ag, and Au are most interesting because they both span more than 1 eV in reaction energies for CO₂ dissociation and have activation energies less than 0.2 eV larger than the reaction energies. This is a substantial break from the linear scaling relation between the two quantities found for CO₂ dissociation on pure metal surfaces. The similar activation and reaction energies can be understood at the atomic level because the catalytic sites with this property have metal atoms that stabilize both the transition state and final state. The Ag-Cu (111) alloy surfaces are further analyzed with a kinetic model, which shows that the surfaces are likely highly selective towards the rWGSR rather than methane formation or coking.

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Supporting Information Available

Description of the simple screening model used to identify interesting alloys, identification of the most stable *CO + O* final state on each alloy surface, CI-NEB calculated potential energy surfaces for the CO₂ dissociation pathways, description of the kinetic model for reverse water gas shift and CO dissociation, discussion on why it's acceptable to neglect hydrogen adsorption, details regarding bulk alloy structures and lattice constants.

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