RSC Sustainability

PAPER

Check for updates

Cite this: RSC Sustainability, 2025, 3, 836

Received 25th July 2024 Accepted 27th November 2024

DOI: 10.1039/d4su00416g

rsc.li/rscsus

Sustainability spotlight

To deal with the massive CO_2 emissions and nearly daily news on climate change, a scalable technology to repurpose this waste to the carbon backbone of fuels and chemicals could revolutionize society. Such routes have been plagued by technical and economic challenges. Our ongoing efforts focus on converting CO_2 to CO by chemical looping using a perovskite oxide with green hydrogen as a tandem reactant. The current research investigates the mechanism of efficient conversion, with which we can design better redox materials and processes to propel this technology forward. Our project goals align with the UN sustainable development goals of clean energy (SDG 7), industry innovation (SDG 9), responsible consumption (SDG 12), and SDG 13 (climate action).

1 Introduction

A potential approach to repurpose CO_2 emissions is the conversion of CO_2 , aided by intermittent renewable energy inputs, into valuable fuels and chemicals. An example of such an alternative pathway involves the conversion of CO_2 into CO, followed by the further conversion of CO into high-value chemical products, such as methane and other hydrocarbons with added value.¹⁻⁴ However, the pivotal step in this process, the conversion of CO_2 into CO, presents a significant challenge due to the exceptionally high energy requirement needed to

break a C=O bond. This is primarily attributed to the inherent stability of the symmetric CO₂ molecule ($\Delta G = -396$ kJ mol⁻¹). To address this energy barrier and reduce the associated energy costs, researchers have explored the utilization of catalytic and non-catalytic reaction processes. A promising process is the reverse water-gas shift chemical looping (RWGS-CL).

RWGS-CL operates through a two-step reaction process. In the first reduction step (eqn (1.1)) metal oxide materials (MO_x) are partially reduced by reacting with hydrogen to form the oxygen vacancy. In the second oxidation step (eqn (1.2)) the oxygen vacancy on reduced; $MO_{x-\delta}$ reacts with CO₂ to produce CO. Subsequently, the regenerated oxide returns to its original structure.

$$MO_x + \delta H_2 \rightarrow MO_{x-\delta} + \delta H_2O$$
 (1.1)

$$MO_{x-\delta} + \delta CO_2 \rightarrow MO_x + \delta CO$$
 (1.2)

Hanzhong Shi, Jiawei Guo, Prabhsimran Singh, Venkat R. Bhethanabotla * and John N. Kuhn *

Perovskite oxides, such as $La_{0.5}Ba_{0.5}FeO_3$ (LBF), facilitate CO₂ conversion by reverse water–gas shift chemical looping (RWSG-CL) at moderate conditions by employing an oxygen vacancy at the surface to aid CO₂ adsorption and then to scavenge an oxygen atom from it to fill the vacancy. The formation of composites with silica is also known to enhance the perovskite oxide's performance. To better clarify this, experimental and computational methods are now combined to probe CO₂ adsorption for both unsupported and silica-supported LBF. Chemisorption tests showed the CO₂ adsorption sites increased from 12.4 to 60.6 μ mol g_{LBF}⁻¹ after adding SiO₂ (75 wt%) to LBF (25 wt%). Spectroscopic studies (DRIFTS) indicated that the carbonate formation during CO₂ adsorption shifts from bidentate to monodentate because the surface morphology changes upon supporting on silica. Computational (DFT) results provide evidence for CO₂ adsorbed as a monodentate and a bidentate carbonate, respectively, on the (111) and (100) surfaces. Monodentate species required lower energy, as determined by DFT, to dissociate C–O bond than bidentate species. Since XRD results identified increases in the (111) relative to (100) planes upon supporting LBF on SiO₂, the combined DRIFTS and DFT approach revealed that the perovskite oxide restructures when in composite form, which explains the increased RWGS-CL process yield of CO.

ito this: DSC Sustain-

ROYAL SOCIETY OF CHEMISTRY

View Article Online

View Journal | View Issue

Department of Chemical, Biological, and Materials Engineering, University of South Florida, 4202 E. Fowler Ave., ENG030, Tampa, Florida 33620, USA. E-mail: bhethana@usf.edu; jnkuhn@usf.edu

[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d4su00416g

Paper

Compared to other CO₂ conversion reactions which often necessitate extremely high temperatures (generally above 1000 ° C) to break the C=O bond,¹⁻⁴ RWGS-CL offers an alternative energy requirement by utilizing perovskite oxides as agent materials. In our previous research, we proposed several perovskite oxide materials to operate at a moderate temperature (~550 °C) of RWGS-CL.⁵⁻⁷ Perovskite oxides (ABO₃) are composed of three essential components: an 'A' site element, a 'B' site element, and three oxygen atoms. Typically, 'A' site elements are chosen from Group II or the lanthanides, while 'B' site elements are primarily transition metals. The perovskite crystal structure exhibits stability across a wide range of compositions on each site, making it an ideal phase for tuning its properties to introduce oxygen vacancies and enhance its capacity for CO₂ adsorption.

In our previous research, La_{0.5}Ba_{0.5}FeO₃ (LBF) has emerged as a potential candidate for low-temperature RWGS-CL, as it is capable of converting CO₂ to CO at 480 °C with a CO yield of $0.2 \text{ mmol g}^{-1} \text{ LBF.}^7$ To facilitate the industrial-scale use of LBF material, LBF/SiO₂ composites were synthesized by adding SiO₂ (quartz, 75 wt%) to pure LBF (25 wt%) to enhance LBF dispersion.89 Subsequently, the LBF/SiO2 powder was formed into pellets using tableting or extrusion methods. The resulting LBF/ SiO₂ pellets converted CO₂ to CO at 550 °C with a CO yield of approximately 2.2 mmol g^{-1} LBF with long-term stability in 50cycle RWGS-CL reaction.¹⁰ Notably, the materials' CO yields were increased when compared to unsupported LBF. As summarized in Table 1, the addition of SiO_2 enriches the (111) plane of LBF crystals. This is noteworthy as previous studies have shown that the (111) surface of similarly related materials, such as MgO and NiO, exhibits higher adsorption strength compared to the (100) surface.^{11,12} Therefore, the enriched (111) plane in LBF/SiO₂ composites could potentially enhance the RWGS-CL reaction, providing a plausible explanation for the observed increase in CO yield after adding SiO₂ as a support material. However, a detailed explanation requires further investigation. In this study, we aim to elucidate the surface mechanism behind this phenomenon by combining experimental and computational methods.

The computational method employed in this study is density functional theory (DFT), which has gained popularity for its ability to compute molecular structures, vibrational frequencies, and energies of chemical reactions.^{13–16} DFT frequency calculations are known to be reliable and less computationally demanding.^{17,18} Therefore, we have used DFT to calculate the vibrational frequencies of the species adsorbed on LBF and LBF/SiO₂ during the RWGS-CL reaction. To validate the DFT frequency results, diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) was utilized to measure the vibrational frequencies of the same samples.^{19–21} These *in situ* DRIFTS experiments combined with DFT calculations were utilized to provide a comprehensive understanding of the vibrational frequencies and adsorption behavior of the species on LBF and LBF/SiO₂ materials. Mechanistic insights into factors contributing to the improved performance of the silica-supported LBF is gained from this study.

2 Materials and methods

2.1 Experimental method

2.1.1 Materials. Citric acid ($C_6H_8O_7$, CA; Aldrich, ACS reagent $\geq 99.5\%$), lanthanum(m) nitrate, (La(NO₃)₃, Aesar, 99.9%), barium carbonate (BaCO₃, Puratronic, 99.997%), iron(m) nitrate (Fe(NO₃)₃, Aldrich, ACS reagent $\geq 98\%$), ethylene glycol ($C_2H_6O_2$, EG; Millipore, $\geq 99.0\%$), silicon dioxide (SiO₂, Sigma-Aldrich quartz), and deionized water (H₂O > 18.0 M Ω cm⁻¹) were used as obtained.

2.1.2 Synthesis of LBF and SiO₂-supported LBF. LBF supported on SiO₂ was synthesized using a modified Pechini method.22 The aqueous solution was made with 12 g citric acid and deionized (DI) water. The metal sources, La(NO₃)₃, BaCO₃, and $Fe(NO_3)_3$ were dissolved in the prepared solution with 2 h stirring at 90 °C until the solution turned red. Following this, ethylene glycol was added to form the gel by mixing for 6 h at 90 °C. Then, 3.6 g of SiO₂ was added into the gel and mixed for an additional 1 h at 90 °C. The gel was transferred to an alumina crucible and heated at 450 °C for 2 h at a heating rate of 20 ° $C \min^{-1}$ in air. Then, after grinding the sample into fine powder it was calcined at 800 °C for 6 h in air with the same heating rate to form the LBF/SiO₂ sample. The precursor was measured to satisfy the element molar ratio of LBF (La : Ba : Fe : CA : EG =0.5:0.5:1:10:40), and SiO₂ was added at the mass ratio of LBF/SiO_2 25 wt% (LBF: SiO₂ = 1:3). The LBF sample was synthesized by the same method, but without adding SiO2 after forming the gel. Other experiments are described in ESI.†

2.1.3 DRIFTS-CO₂-adsorption experiments. For DRIFTS-CO₂-adsorption experiments, a mass of approximately 50 mg LBF or LBF/SiO₂ was placed into the reactor. The reactor was installed in the reaction chamber that was purged by dry air overnight before starting the experiments. A total gas flow rate of 50 sccm was used. The reactor was first heated to 500 °C for LBF or 550 °C for LBF/SiO₂ under Ar. Then the reduction process was conducted by flowing 10% H₂/Ar for 30 min. The reactor was cooled down to room temperature under Ar, and the background data were taken at 25 °C. The 10% CO₂/Ar was flowed for 1 h, then the reactor was purged by Ar for 30 min, the DRIFTS data were collected after gas CO₂ was removed.

Table 1	Materials conversion properties of LBF and LBF/SiO ₂ (ref. 7 and 10)					
	Reduction temperature (°C)	Oxidation temperature (°C)	$ m H_2O$ yield (mmol $ m g_{LBF}^{-1}$)	CO yield $(mmol \ g_{LBF}^{-1})$	Surface area $(m^2 g^{-1})$	
LBF LBF/SiO ₂	430 580	480 550	0.2 2.2	0.2 2.2	8.9 3.2	

Although both DFT and DRIFTS analyses in this study were conducted at temperatures lower than those used in the actual RWGS-CL reaction. The surface properties of the materials remain relatively stable as the temperature increases from room temperature to reaction temperature, given that both the perovskite and SiO₂ are stable materials which are synthesized at a temperature much higher than reaction temperature. Therefore, it is reasonable to assume that the nature of the surface sites and the form of CO₂ adsorption remain consistent across the different temperatures.^{23,24}

2.2 Computational methods

RSC Sustainability

The DFT calculations in this study were performed using Quantum ESPRESSO software (Version 6.2.1).^{25,26} The materials models used in the calculations are referenced from the materials project database, with corresponding index numbers.27 The plane wave basis set and gradient generalization approximation for electron densities were used.28,29 The projector augmented wave (PAW) potentials and Perdew-Burke-Ernzerhof (PBE) variant of exchange-correlation functional were employed.30,31 In this study, the energy convergence criterion was set to 10^{-6} eV per atom, and the force convergence criterion was set to 0.005 eV Å⁻¹. A constant cutoff of 40 Ry (544 eV) was selected. A k-point mesh of $(4 \times 4 \times 1)$ was used for the Brillouin zone sampling. The Hubbard U correction was not applied in this calculation, as the trend of energies did not show a significant shift with or without the application of the Hubbard U correction.^{32,33} The vibrational frequency was calculated from a 6 imes 6 Hessian matrix.³⁴ The details of DFT calculation and other experimental methods are described in ESI.†

3 Result and discussion

3.1 Materials characterizations

To better understand the reason for LBF/SiO₂ giving larger CO yield than LBF in the RWGS-CL process, XRF, XRD, and XPS experiments were performed on both materials. The metal elemental compositions of the samples were determined by XRF



Fig. 1 Metal elemental compositions of fresh samples in bulk calculated from XRF results, red solid bar is LBF result, and blue dotted bar is for LBF/SiO₂ 25 wt%.

(Fig. 1), showing that LBF has 26.1% La, 26.3% Ba, and 47.6% Fe, while LBF/SiO₂ (25 wt%) has 25.7% La, 25.4% Ba, and 48.9% Fe, which are close to the nominal compositions.

The XRD results (Fig. 2) indicate that LBF/SiO₂ structure is enriched with the (111) planes, as evidenced by the intensity ratio of diffraction lines ((111) to (110)) switching from 0.2 to 1, because the relative intensities generally correlate with a greater number of crystallites oriented along a specific plane.³⁵ This switch of crystallographic planes could be because SiO₂ influences the preferential growth or orientation of the active catalytic phase.³⁶ As demonstrated in our previous study, SiO₂ exhibits a wettability effect on the perovskite particles, potentially leading to the formation of a secondary plane associated with surface wetting.³⁷ Thus, the presence of SiO₂ and the synthesis method can stabilize (111) planes more than others (110), leading to an increased intensity in the XRD pattern.³⁸ LBF is used as the control sample for the (100) plane. The surface morphology difference is also observed from XPS results (Fig. 3). After adding SiO_2 to LBF, there is a noticeable shift in the difference of La3d_{5/2} binding energy from 4.1 to 4.0 eV, and a shift in the difference of La3d_{3/2} binding energy from 3.4 to 3.1 eV. The binding energy differences between two La3d peaks are close between LBF and LBF/SiO2. The binding energy of O1s exhibits a distinct difference due to the presence of oxygen bonds from SiO₂. Fe and Ba remain the same. Therefore, by combing the results from XRF, XRD, and XPS, the surface with enriched (111) is a potential reason that led to increase in the CO yield.

3.2 CO₂ chemisorption

To better understand the CO₂ adsorption on LBF and LBF/SiO₂, CO₂ chemisorption experiments were performed on these two reduced samples at room temperature (Fig. 4). Results indicate a similar volume of CO₂ adsorbed by LBF and LBF/SiO₂. However, a sample of SiO₂ support material was tested to find that SiO₂ is inert for CO₂ chemisorption, and the concentration of perovskite in LBF (100%) is 4 times than LBF/SiO₂ (25 wt%). Therefore, an increased CO₂ chemisorption upon supporting in SiO₂, increasing from 12.4 µmol g_{LBF}⁻¹ (LBF) to 60.6 µmol g_{LBF}⁻¹ (LBF/SiO₂) calculated by extrapolation method (eqn (S3)†).^{39,40} From these results, we conclude that supporting LBF on SiO₂ enhances of the number of CO₂ chemisorption sites, by *a* factor of 5 in this case. This enhancement can be considered as one reason the improved CO yield in the reduction process.

3.3 Temperature-programmed oxidation experiments

Oxygen vacancies are created during the reduction phase of the RWGS-CL process. The perovskite phase, $La_{0.5}Ba_{0.5}FeO_3$, is reduced to $La_{0.5}Ba_{0.5}FeO_{3-\delta}$, where δ presents oxygen vacancy amount that is 0.1 and 0.5 for LBF and LBF/SiO₂. respectively.^{7,10} The oxidation temperatures for samples of reduced LBF and reduced LBF/SiO₂ were examined under various oxidizers (O₂, H₂O, and CO₂) using TPO experiments. Comparing the results from TPO-O₂, TPO-H₂O, and TPO-CO₂ experiments (Fig. 5), it is evident that a lower oxidation temperature of 250 °C is required for O₂, which is the strongest oxidizer, compared to H₂O and



Fig. 2 XRD results of synthesized samples. (a) LBF/SiO₂, circular marker denotes the crystal planes from LBF, and rectangular marker denotes the crystal planes from SiO₂. (b) LBF. (c) Detailed results in the 2θ range of 30 to 40°.

CO₂, both of which require temperatures exceeding 400 °C. For the reduced LBF/SiO₂, a second oxidation peak appears at 530 ° C, indicating a shift in surface morphology due to the addition of SiO₂. Interestingly, the oxidation temperatures for H₂O and CO₂ are similar, deviating from the expectation that H₂O, typically a stronger oxidizer than CO₂, would oxidize the sample at a lower temperature. This anomaly is attributed to the oxygen vacancies on the perovskite oxide surface, which enhance CO₂ adsorption and lower the dissociation energy of CO₂ making CO₂ a strong oxidant in this reaction Consequently, this reduced energy requirement aligns with the lower temperature range (400 to 650 °C) for the CO₂ to CO reaction in the RWGS-CL process, compared to the higher temperatures (>1000 °C) required in conventional thermochemical processes.

3.4 DRIFTS-CO₂-adsorption

DRIFTS-CO2-adsorption experiments were conducted to reduce the LBF and LBF/SiO₂ (25 wt%) samples. The DRIFTS results (Fig. 6) show the vibrational frequencies of the adsorbed



Fig. 4 Isotherms for CO₂ adsorption at room temperature.

species. The experimental vibrational frequencies are $v_{as} =$ 1606 cm⁻¹ and $v_s = 1347$ cm⁻¹ for CO₂ adsorbed on reduced LBF, and $v_{as} = 1544 \text{ cm}^{-1}$ and $v_s = 1364 \text{ cm}^{-1}$ for CO₂ adsorbed on reduced LBF/SiO₂ (25 wt%). The difference between v_{as} and $v_{\rm s}$ decreases from 259 to 180 cm⁻¹. This is because CO₂ is



Fig. 3 XPS results for LBF (red, top) and LBF/SiO₂ (blue, bottom).

8



Fig. 5 TPO results for different oxidizers: (a) TPO- O_2 , (b) TPO- H_2O , and (c) TPO- CO_2 . The red lines (top) are the results for reduced LBF, and the blue lines (bottom) are the results for reduced LBF/SiO₂.



Fig. 6 DRIFT spectra of TPD-CO₂ experiments. Red solid line is the result for LBF, and blue dotted line is the result for LBF/SiO₂ 25 wt%.

adsorbed in the bidentate carbonate form on LBF whereas it is adsorbed in the monodentate carbonate form on LBF/SiO₂.⁴¹ This carbonate formation differences agree with the DFT calculations presented in the next section.

3.5 DFT calculations

3.5.1 Oxygen vacancy formation energy (E_{Ovac}). The surface constructions for LBF-111 and LBF-100, along with the possible sites of oxygen vacancy, are shown in Fig. S1.[†] The oxygen vacancy formation energy (Table 2) was calculated using the method described in ESI.[†] E_{Ovac} of LBF-100 with Fe surface (LBF-100(Fe)) was found to be slightly lower than that of La/Ba surface (LBF-100(La/Ba)), indicating that the Fe surface is more favorable for oxygen vacancy formation during the reduction process. Similarly, for the LBF-111 structure, Fe–O is the most likely site for reduction. Comparing the E_{Ovac} values of LBF-111 and LBF-100, it was found that E_{Ovac} of LBF-111 (1.62 and 1.80 eV) is significantly lower than E_{Ovac} of LBF-100 (3.01 and 3.24 eV), indicating that LBF-111 is the preferred surface for providing oxygen vacancies in the reduction step of RWGS-CL.

3.5.2 CO_2 adsorption structure. In the oxidation step of RWGS-CL, CO₂ molecules adsorb on the oxygen-vacated LBF surface. The physisorbed and chemisorbed CO₂ are identified based on a distance of 1.43 Å.42,43 Possible chemisorbed structures of CO₂ on LBF-111 are shown in Fig. 7(a)-(g), while one structure for LBF-100 (Fe-O) is shown in Fig. 7(i), based on the possible sites of oxygen vacancy as shown in Fig. S1.† The surface structure of (111) is named by surface atoms, for example, LBF-111(LaBa-La) means two La and one Ba atoms with one La atom on the center of the LBF (111) surface. The C=O bond distances between carbon and oxygen are presented in Table 3. C=O bond lengths falling within the range of 1.21 to 1.23 Å indicate the formation of a double bond between carbon and oxygen, while bond lengths of C-O_a and C-O_b (Table 3) in the range of 1.26 to 1.34 Å are considered as C-O single bonds.44 Therefore, the DFT calculations suggest that CO₂ adsorption occurs in the carbonate form. In (Fig. 7(g) & (h)), the bond lengths of 1.16 and 1.17 Å for LBF-111(BaLa-Ba) do not follow the same trend, which could be attributed to Ba not being a site for CO₂ adsorption.

3.5.3 Vibrational frequency of adsorbed CO₂. The CO₂ adsorption structure on the (111) plane was selected as LBF-111(BaLa–Fe) (Fig. 7(f)), while the adsorption structure on the (100) plane is shown in Fig. 7(i). The vibrational frequencies obtained from DFT calculations and DRIFTS results are shown in Table 4. The vibrational frequencies of CO₂ on LBF-100(Fe–O) are $v_{\rm as} = 1663$ cm⁻¹ and $v_{\rm s} = 1415$ cm⁻¹, while the vibrational frequencies of CO₂ on LBF-111(BaLa–Fe) are $v_{\rm as} = 1639$ cm⁻¹ and $v_{\rm s} = 1450$ cm⁻¹.

Table 2	Oxygen vacancy	formation energy results
---------	----------------	--------------------------

$E_{\rm Ovac}$ (eV)	Fe surface	La/Ba surface
LBF-111	1.62	1.80
LBF-100	3.01	3.24



Fig. 7 The CO₂ adsorption on each configuration. (a) & (b) are the adsorption results of LBF-111(LaBa-La). (c) & (d) are the adsorption results of LBF-111(LaBa-Fe), (e) & (f) are the adsorption results of LBF-111(BaLa-Fe). (g) & (h) are the adsorption results of LBF-111(BaLa-Ba), C-O_b in (g) is not marked because it is away to the ideal range. (i) Is the adsorption result of LBF-100(Fe-O).

Table 3 The C–O bond lengths of the adsorbed CO $_2$ on LBF-111 and LBF-100 surface

	C=O bond length (Å)	C–O _a bond length (Å)	C–O _b bond length (Å)
LBF-111(LaBa–La) ₁	1.23	1.48	1.28
LBF-111(LaBa-La) ₂	1.23	1.44	1.26
LBF-111(LaBa-Fe) ₁	1.21	1.43	1.32
LBF-111(LaBa-Fe) ₂	1.21	1.41	1.34
LBF-111(BaLa-Fe) ₁	1.22	1.42	1.32
LBF-111(BaLa-Fe) ₂	1.22	1.41	1.33
LBF-111(BaLa-Ba) ₁	1.16	1.16	_
LBF-111(BaLa-Ba) ₂	1.17	1.17	_
LBF-100(Fe–O)	1.22	1.40	1.37

Table 4 Vibrational frequency (cm $^{-1}$) results from DFT calculations and DRIFTS experiments

DFT	DRIFTS
1663	1606
1415	1347
248	259
1639	1544
1450	1364
189	180
	DFT 1663 1415 248 1639 1450 189

In comparison, the frequency differences (v_{δ}) between antisymmetric (v_{as}) and symmetric (v_s) vibrations from DFT calculation results are 248 cm⁻¹ and 189 cm⁻¹. Although there are slight differences between the calculated vibrational frequencies and the experimental data due to the limitations of DFT calculations,^{45,46} the v_{δ} values are in agreement. In this study, the calculated v_{δ} of CO₂ on reduced LBF is 189 cm⁻¹, which is close to the experimental value of 180 cm⁻¹, while the calculated v_{δ} of CO₂ on reduced LBF/SiO₂ (25 wt%) is 248 cm⁻¹, which is close to the experimental value of 259 cm⁻¹. Thus, it can be concluded that CO₂ is adsorbed in the form of carbonates during the oxidation step in RWGS-CL. The vibrational frequency difference between antisymmetric and symmetric vibrations decreases after adding SiO₂ as a support material, indicating that the adsorbed CO₂ on the (111) plane is monodentate carbonate different from bidentate carbonate on the (100) plane due to the difference of ν_{δ} in carbonate species.^{41,47,48}

3.5.4 Reaction intermediate energy and mechanism of oxidation step in RWGS-CL. The energy diagram in Fig. 8 illustrates the energy changes for each intermediate state during the RWGS-CL process by LBF-111 (BaLa–Fe) and LBF-100 (Fe–O). The initial state (IS, $E_{IS} = 0$) is the fresh LBF surface, which is also the final state (FS) after the reaction. The adsorption structures are shown in Fig. S2.† The total energy difference (E_{IMx}) between each intermediate state (IMx) and IS are as follows:

(1) IS – IM1: H₂ is physisorbed on the original surface, resulting in an energy change of $E_{IM1(100)} = -0.014$ eV for the (100) plane and $E_{IM1(111)} = -0.006$ eV for the (111) plane.

(2) IM1 – IM2: the physisorbed H₂ moves closer to the surface and forms a chemisorption state, resulting in an energy change



Fig. 8 Step reaction energy for RWGS-CL. Red solid line is the reaction on (100) plane, and blue dotted line is the reaction on (111) plane.

of $E_{IM2(100)} = 1.85$ eV for the (100) plane and $E_{IM2(111)} = 1.91$ eV for the (111) plane.

(3) IM2 – IM3: the samples are reduced with adsorbed H₂, resulting in the formation of oxygen vacancies on the surface and production of H₂O, with an energy change of $E_{IM3(100)} =$ 1.55 eV for the (100) plane and $E_{IM3(111)} =$ 1.62 eV for the (111) plane.

(4) IM3 – IM4: CO₂ is adsorbed on the reduced surface as carbonate form, resulting in an energy change of $E_{IM4(100)} =$ 1.63 eV for the (100) plane and $E_{IM4(111)} =$ 1.93 eV for the (111) plane.

(5) IM4 – IM5: adsorbed CO₂ is converted to CO by oxidizing the oxygen-vacated surface, and the produced CO is adsorbed on the surface, resulting in an energy change of $E_{IM5(100)} = 2.98$ eV for the (100) plane and $E_{IM5(111)} = 2.48$ eV for the (111) plane.

(6) IM5 – FS: CO is desorbed, and the surface regains its original structure.

The intermediate energy results indicate that the reaction on the (111) plane requires a lower maximum energy (2.48 eV) compared to the reactions on the (100) plane (2.98 eV), which suggests that the (111) plane has the potential to produce a higher amount of CO at the same temperature. The reduction steps occur from IS to IM3, while the oxidation steps are described in eqn (2.1) to (2.3).

$$\delta \text{CO}_2 + * \rightarrow \delta \text{CO}_2^* \tag{2.1}$$

$$\operatorname{CO}_{2}^{*} + \operatorname{ABO}_{3-\delta} \rightarrow \operatorname{ABO}_{3} + \operatorname{CO}^{*}$$
 (2.2)

$$\mathrm{CO}^* \to \mathrm{CO}^{+*}$$
 (2.3)

4 Conclusion

Previous studies reported the increases of CO yield upon supporting on SiO₂. To better describe the fundamental reason, this research focused particularly on the CO₂ adsorption on both LBF and LBF/SiO₂ surfaces during the oxidation step in RWGS-CL. X-ray diffraction (XRD) analysis revealed a morphology alteration upon SiO₂ addition, notably a relative increase in the exposed (111) planes. TPD-CO₂ combined with DRIFTS experiments further demonstrate CO₂ adsorption as carbonate on the reduced perovskite surfaces. CO₂ chemisorption shows CO₂ sites increased from 12.4 to 60.6 µmol g_{LBF}⁻¹ after adding SiO₂.

TPO experiments under various oxidizers highlight the role of oxygen vacancies on the reduced perovskite surface. DFT calculations provided insight into the adsorption energy and structure of carbonate on different planes, identifying monodentate carbonate on the (111) plane and bidentate carbonate on the (100) plane, distinguished by variations in the carbonate species' stretching modes.

In situ DRIFTS further confirmed these findings, correlating specific ν_{δ} to monodentate and bidentate carbonate structures on the (111) and (100) planes, respectively. DFT calculations of the energy profiles for reaction intermediates suggest that the

(111) surface offers a lower energy pathway for converting CO_2 to CO. Collectively, these results elucidate the role of SiO_2 in enhancing CO yield as the increased (111) planes facilitate monodentate carbonate formation, thereby reducing the energy requirement for converting CO_2 to CO.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

J. N. K. and V. R. B. declare patents and other financial interests in this technology.

Acknowledgements

This study was funded by NSF grant (IIP-1743623 and IIP-1913722), Florida High Tech Corridor, and a USF Fellowship (to HS).

References

- 1 Q. Jiang, J. Tong, G. Zhou, Z. Jiang, Z. Li and C. Li, *Sol. Energy*, 2014, **103**, 425–437.
- 2 A. Le Gal and S. Abanades, *J. Phys. Chem. C*, 2012, **116**, 13516–13523.
- 3 F. Fresno, R. Fernández-Saavedra, M. Belén Gómez-Mancebo, A. Vidal, M. Sánchez, M. Isabel Rucandio, A. J. Quejido and M. Romero, *Int. J. Hydrogen Energy*, 2009, 34, 2918–2924.
- 4 C. Agrafiotis, M. Roeb and C. Sattler, *Renewable Sustainable Energy Rev.*, 2015, **42**, 254–285.
- 5 Y. A. Daza, D. Maiti, B. J. Hare, V. R. Bhethanabotla and J. N. Kuhn, *Surf. Sci.*, 2016, **648**, 92–99.
- 6 A. E. Ramos, D. Maiti, Y. A. Daza, J. N. Kuhn and V. R. Bhethanabotla, *Catal. Today*, 2019, **338**, 52–59.
- 7 H. Shi, V. R. Bhethanabotla and J. N. Kuhn, *J. CO2 Util.*, 2021, **51**, 101638.
- 8 B. J. Hare, D. Maiti, Y. A. Daza, V. R. Bhethanabotla and J. N. Kuhn, *ACS Catal.*, 2018, **8**, 3021–3029.
- 9 J. C. Brower, B. J. Hare, V. R. Bhethanabotla and J. N. Kuhn, *ChemCatChem*, 2020, **12**, 6317–6328.
- 10 H. Shi, V. R. Bhethanabotla and J. N. Kuhn, *J. Ind. Eng. Chem.*, 2023, **118**, 44–52.
- 11 H. Onishi, C. Egawa, T. Aruga and Y. Iwasawa, *Surf. Sci.*, 1987, **191**, 479–491.
- 12 J. Hu, K. Zhu, L. Chen, C. Kübel and R. Richards, *J. Phys. Chem. C*, 2007, **111**, 12038–12044.
- 13 J. K. Labanowski and J. W. Andzelm, *Density Functional Methods in Chemistry*, Springer Science & Business Media, 2012.
- 14 P. Politzer and J. M. Seminario, *Modern Density Functional Theory: a Tool for Chemistry*, Elsevier, 1995.
- 15 R. G. Parr and Y. Weitao, *Density-Functional Theory of Atoms and Molecules*, Oxford University Press, 1995.

- 16 T. Ziegler, Chem. Rev., 1991, 91, 651-667.
- 17 M. W. Wong, Chem. Phys. Lett., 1996, 256, 391-399.
- 18 L. J. Bartolotti and K. Flurchick, *Rev. Comput. Chem.*, 1996, 7, 187–216.
- 19 N. H. Elsayed, D. Maiti, B. Joseph and J. N. Kuhn, *Catal. Lett.*, 2018, **148**, 1003–1013.
- 20 G. H. Smudde, T. L. Slager, C. G. Coe, J. E. MacDougall and S. J. Weigel, *Appl. Spectrosc.*, 1995, **49**, 1747–1755.
- 21 H. Gao, H. He, Y. Yu and Q. Feng, *J. Phys. Chem. B*, 2005, **109**, 13291–13295.
- 22 M. Popa and M. Kakihana, *Solid State Ionics*, 2002, **151**, 251–257.
- 23 M. Bersani, K. Gupta, A. K. Mishra, R. Lanza, S. F. R. Taylor, H.-U. Islam, N. Hollingsworth, C. Hardacre, N. H. de Leeuw and J. A. Darr, *ACS Catal.*, 2016, 6, 5823–5833.
- 24 L. F. Rasteiro, R. A. De Sousa, L. H. Vieira, V. K. Ocampo-Restrepo, L. G. Verga, J. M. Assaf, J. L. F. Da Silva and E. M. Assaf, *Appl. Catal.*, *B*, 2022, **302**, 120842.
- 25 P. Giannozzi, O. Andreussi, T. Brumme, O. Bunau, Buongiorno Nardelli, М М. Calandra, R. Car, C. Cavazzoni, D. Ceresoli, M. Cococcioni, N. Colonna, I. Carnimeo, A. Dal Corso, S. de Gironcoli, P. Delugas, R. A. DiStasio, A. Ferretti, A. Floris, G. Fratesi, G. Fugallo, R. Gebauer, U. Gerstmann, F. Giustino, T. Gorni, J. Jia, M. Kawamura, H. Y. Ko, A. Kokalj, E. Küçükbenli, M. Lazzeri, M. Marsili, N. Marzari, F. Mauri, N. L. Nguyen, H. V. Nguyen, A. Otero-de-la-Roza, L. Paulatto, S. Poncé, R. Sabatini, B. Santra, M. Schlipf, D. Rocca. A. P. Seitsonen, A. Smogunov, I. Timrov, T. Thonhauser, P. Umari, N. Vast, X. Wu and S. Baroni, J. Phys.: Condens. Matter, 2017, 29, 465901.
- 26 K. Burke, J. Chem. Phys., 2012, 136, 150901.
- 27 A. Jain, S. P. Ong, G. Hautier, W. Chen, W. D. Richards, S. Dacek, S. Cholia, D. Gunter, D. Skinner, G. Ceder and K. A. Persson, *APL Mater.*, 2013, 1, 011002.
- 28 P. Giannozzi, S. Baroni, N. Bonini, M. Calandra, R. Car, C. Cavazzoni, D. Ceresoli, G. L. Chiarotti, M. Cococcioni, I. Dabo, A. Dal Corso, S. de Gironcoli, S. Fabris, G. Fratesi, R. Gebauer, U. Gerstmann, C. Gougoussis, A. Kokalj, M. Lazzeri, L. Martin-Samos, N. Marzari, F. Mauri, R. Mazzarello, S. Paolini, A. Pasquarello, L. Paulatto, C. Sbraccia, S. Scandolo, G. Sclauzero, A. P. Seitsonen,

A. Smogunov, P. Umari and R. M. Wentzcovitch, *J. Phys.:* Condens. Matter, 2009, 21, 395502.

- 29 K. F. Garrity, J. W. Bennett, K. M. Rabe and D. Vanderbilt, *Comput. Mater. Sci.*, 2014, **81**, 446–452.
- 30 G. Kresse and D. Joubert, Phys. Rev. B, 1999, 59, 1758-1775.
- 31 J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1996, 77, 3865–3868.
- 32 D. Maiti, Y. A. Daza, M. M. Yung, J. N. Kuhn and V. R. Bhethanabotla, *J. Mater. Chem. A*, 2016, 4, 5137–5148.
- 33 I. W. Boateng, R. Tia, E. Adei, N. Y. Dzade, C. R. A. Catlow and N. H. de Leeuw, *PCCP Phys. Chem. Chem. Phys.*, 2017, 19, 7399–7409.
- 34 D. S. Sholl and J. A. Steckel, *Density Functional Theory: a Practical Introduction*, John Wiley & Sons, 2011.
- 35 L. Zhang, A. A. S. Gonçalves and M. Jaroniec, *RSC Adv.*, 2020, 10, 5585–5589.
- 36 R. A. McKee, F. Walker and M. Chisholm, *Phys. Rev. Lett.*, 1998, **81**, 3014.
- 37 D. M. Bryan, J. Hare, Y. A. Daza, V. R. Bhethanabotla and J. N. Kuhn, ACS Catal., 2018, 8, 3021–3029.
- 38 J. Arenas-Alatorre, M. Avalos-Borja and G. Díaz, *Appl. Surf. Sci.*, 2002, **189**, 7–17.
- 39 S. Brunauer, P. H. Emmett and E. Teller, J. Am. Chem. Soc., 1938, 60, 309–319.
- 40 C. Chen and W.-S. Ahn, Chem. Eng. J., 2011, 166, 646-651.
- 41 K. Coenen, F. Gallucci, B. Mezari, E. Hensen and M. van Sint Annaland, J. CO2 Util., 2018, 24, 228–239.
- 42 G. Pacchioni, Surf. Sci., 1993, 281, 207-219.
- 43 F. H. Allen and A. J. Kirby, J. Am. Chem. Soc., 1984, 106, 6197– 6200.
- 44 J. Demaison and A. G. Császár, *J. Mol. Struct.*, 2012, **1023**, 7–14.
- 45 A. A. El-Azhary and H. U. Suter, *J. Phys. Chem.*, 1996, **100**, 15056–15063.
- 46 M. L. Laury, M. J. Carlson and A. K. Wilson, *J. Comput. Chem.*, 2012, **33**, 2380–2387.
- 47 B. H. Solis, Y. Cui, X. Weng, J. Seifert, S. Schauermann, J. Sauer, S. Shaikhutdinov and H.-J. Freund, *Phys. Chem. Chem. Phys.*, 2017, **19**, 4231–4242.
- 48 J. Liu, Y. Li, H. Liu and D. He, *Biomass Bioenergy*, 2018, **118**, 74–83.