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Thermodynamic and First-principles Assessments of Materials for Solar-driven CO₂ Splitting Using Two-step Thermochemical Cycles

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Abstract: Carbon-neutral fuel production by solar-driven two-step thermochemical carbon dioxide splitting provides an alternative to fossil fuels as well as mitigates global warming. The success of this technology relies on the advancements of redox materials. Despite the recognition of the entropic effect, usually energy descriptors (enthalpy of formation or energy of oxygen-vacancy formation) were used for computational assessment of material candidates. Here, in the first step, the criteria was derived based on the combination of solid-state change of entropy and formation enthalpy, and was used to thermodynamically assess the viability of material candidates. In the thermodynamic map, a triangular region, featuring large positive solid-state changes of entropy and small enough solid-state changes of formation enthalpy, was found for qualified candidates. Next, a first-principles DFT+U method was presented to fast and reasonably predict the solid-state changes of entropy and formation enthalpy of candidate redox materials, exemplified for pure and Samaria-doped ceria, so that new redox materials can be added to the thermodynamic map. All above results highlight the entropic contributions from polaron-defect vibrational entropy as well as ionic (oxygen vacancies) and electronic (polarons) configurational entropy.

Key words: carbon dioxide splitting; two-step thermochemical cycle; first principles; entropy; solar fuel

Using solar energy to convert carbon dioxide (CO_2) into carbon-neutral fuels provides an alternative to fossil fuels and mitigates global warming. In analogy to conventional chemical plants, a solar-fuel plant is able to operate overnight addressing the intermittency of solar energy. Regarding this, concentrated solar thermochemical CO₂ splitting holds promise because cost-effective and high-density heat storage can be integrated into this technology^[1]. Direct CO₂ splitting by thermolysis, requires temperature as high as 3000 °C except for separation of oxygen from fuel products. Therefore, research converged to the two-step redox cycles based on partial reduction (or non-stoichiometric, oxygen-vacancy) and oxidation of non-volatile oxides (or redox materials), because this type of cycles features the combinations of practical operation temperature and high thermodynamic efficiency^[1-3].

$$CO_2 \rightarrow CO + \frac{1}{2}O_2$$
, at T_{th} (Reaction (1), CO_2 thermolysis)

 $\frac{1}{\delta} MO_x \rightarrow \frac{1}{\delta} MO_{x-\delta} + \frac{1}{2}O_2, \text{ at } T_H \text{ (Reaction (2), thermal reduction)}$ $\frac{1}{\delta} MO_{x-\delta} + CO_2 \rightarrow \frac{1}{\delta} MO_x + CO, \text{ at } T_L \text{ (Reaction (3), CO_2 splitting)}$

Although considerable materials^[4] have been examined including ceria^[5-8], ferrites^[9-11] and perovskites^[12-14], the state-of-the-art solar-to-chemical energy conversion efficiency was as low as ~5% for solar thermochemical CO₂ splitting^[5]. Therefore, computational assessments of materials are still of great importance^[15-17]. Intuitively, the thermodynamically suitable redox materials can be identified based on the fundamental concept that the change in the Gibbs free energy, ΔG , should be negative for the two cycle steps (Reactions (2) and (3)). Thereby, solid-state enthalpy and entropy of reduction excluding the contributions from gaseous species (whose thermodynamic property data is available for a wide range of conditions) can be used as the descriptors for thermodynamic assessments of redox materials. Particularly, Meredig^[18] and

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Shah, et al.^[19] raised the importance of entropic effects. Unfortunately, obtaining reliable thermodynamic data, especially the reduction entropy, of redox material candidates, either computationally or experimentally is often challenging^[11-12,20-24]. For example, Bork, et al.^[12] used CALPHAD models to access or extrapolate needed thermodynamic properties of perovskites, but models with this level of sophistication are not available for many redox material candidates^[2]. Michalsky, et al.^[22] found that the Gibbs free energy for formation of many bulk oxides (equivalent to solid-state enthalpy and entropy of reduction) scaled with their oxygen-vacancy formation energy at their most stable surfaces. However, this scaling feature was only demonstrated in the stoichiometric regime. Muhich, et al.^[11] developed an assessment method where only the enthalpy of reduction was used, but this method only worked for the assessments within the same class of materials (hercynite in their work) because the reduction entropy is approximately the same for the hercynite family.

Therefore, this study provided general descriptors and criteria for thermodynamic assessments of proposed or newly discovered redox materials, and to provide first-principles methods for fast predictions of these descriptors. Here, pure and Samaria-doped ceria were taken as benchmark materials because they own fast splitting kinetics, excellent high-temperature and cycling stability^[5-8], and also provide a platform for first-principles investigations of the effects of typical defects including oxygen vacancies, polarons and dopant ions^[25].

1 Thermodynamic derivation of descriptors and criteria

In this section, the descriptors was proposed, and then the criteria was derived for thermodynamic assessments of the viability of material candidates. In the derivation, complementary constraints were presented in consideration of non-stoichiometric reaction mechanism, theoretical efficiency and practical operating conditions of the solardriven two-step thermochemical process to extend the thermodynamic framework described by Meredig and Wolvertor^[18].

This study starts with the fundamental concept that the change in the Gibbs free energy, ΔG , should be negative for the two cycle steps (Reactions (2) and (3)) so that they are thermodynamically favorable. This gives Eq. (1–2).

$$\Delta G_2 = \frac{1}{\delta} (\Delta H_{\mathrm{f, MO}_{x-\delta}} - \Delta H_{\mathrm{f, MO}_x}) - T_{\mathrm{H}} \times \left[\frac{1}{\delta} (S_{\mathrm{MO}_{x-\delta}} - S_{\mathrm{MO}_x}) + \frac{1}{2} (S_{\mathrm{O}_2, T_{\mathrm{H}}} - R \ln P_{\mathrm{O}_2}) \right] \leq 0 \quad (1)$$

$$\Delta G_3 = \frac{1}{\delta} (\Delta H_{\mathrm{f, MO}_x} - \Delta H_{\mathrm{f, MO}_{x-\delta}}) + (\Delta H_{\mathrm{f, CO}, T_{\mathrm{L}}} - \Delta H_{\mathrm{f, CO}_2, T_{\mathrm{L}}}) - T_{\mathrm{L}} \times \left[\frac{1}{\delta} (S_{\mathrm{MO}_x} - S_{\mathrm{MO}_{x-\delta}}) + (S_{\mathrm{CO}, T_{\mathrm{L}}} - S_{\mathrm{CO}_2, T_{\mathrm{L}}}) \right] \leq 0 \quad (2)$$

where $\Delta H_{\rm f}$ represents the enthalpy of formation, *S* represents entropy, *P* represents pressure (kPa) and *R* is the ideal gas constant. Solid-state entropy and solid-state enthalpy of formation are assumed to be temperature-independent. Observed from Eq. (1) and (2), solid-state change of entropy, $\Delta S_{\rm solid}$, and change of enthalpy of formation, $\Delta H_{\rm solid}$

$$\begin{cases} \Delta S_{\text{solid}} = \frac{1}{\delta} (S_{\text{MO}_{x-\delta}} - S_{\text{MO}_x}) \\ \Delta H_{\text{solid}} = \frac{1}{\delta} (\Delta H_{\text{f, MO}_{x-\delta}} - \Delta H_{\text{f, MO}_x}) \end{cases}$$
(3)

depend on the redox material, while the rest quantities for gaseous species are independent and were well documented^[26]. Therefore, ΔS_{solid} with unit of J·(0.5 mol O₂)⁻¹·K⁻¹ and ΔH_{solid} with unit of kJ·(0.5 mol O₂)⁻¹, defined in Eq. (3), were used as descriptors for thermodynamic assessments in this work. By substituting Eq. (3) into Eq. (1) and (2), Eq. (4–5), two of the assessment criteria, are obtained.

$$\Delta H_{\text{solid}} \leq T_{\text{H}} \left(\Delta S_{\text{solid}} + \frac{1}{2} S_{\text{O}_2, T_{\text{H}}} - \frac{1}{2} R \ln P_{\text{O}_2} \right) \quad (4)$$

$$\Delta H_{\text{solid}} \geq \left(\Delta H_{\text{f, CO}, T_{\text{L}}} - \Delta H_{\text{f, CO}_2, T_{\text{L}}} \right) - T_{\text{L}} \left(S_{\text{CO}, T_{\text{L}}} - S_{\text{CO}_2, T_{\text{L}}} - \Delta S_{\text{solid}} \right) \quad (5)$$

In the next step, thermodynamic principles are applied to analyse the solar-to-chemical (STC) energy conversion efficiency of the two-step thermochemical process to provide complementary criteria. STC efficiency, $\eta_{\text{STC}} =$ $\eta_{\text{STT}} \times \eta_{\text{TTC}}$, where η_{STT} is solar-to-thermal efficiency, and $\eta_{\rm TTC}$ is thermal-to-chemical efficiency. Ideally, $\eta_{\rm TTC} = \Delta G_{1298K}$ ΔH_{solid} , where $\Delta G_{1298\text{K}}$ (~256 kJ·(0.5 mol O₂)⁻¹) is the change in the Gibbs free energy at 298 K for CO₂ thermolysis (Reaction 1). By assuming an achievable η_{STT} of 70%, a target η_{TTC} of 28.6% is required to achieve a practical η_{STC} of 20%, which is competitive to the technology that a solar photovoltaic device coupled with an electrolyser^[27]. Miller, et al.^[2] analysed that potential redox materials should be able to achieve a theoretical thermal-to-chemical efficiency which is twice of the target η_{TTC} (Eq. (6)).

$$\Delta H_{\text{solid}} \leq 450 \text{ kJ} \cdot (0.5 \text{ mol } \text{O}_2)^{-1} \tag{6}$$

Therefore, Eq. (4) to (6) were used to calculate the boundary values of (ΔS_{solid} , ΔH_{solid}). Eq. (4) and (5) tell that the (ΔS_{solid} , ΔH_{solid}) boundary varies with operating temperatures (T_{H} and T_{L}) and operating pressure (P_{O_2}). For the thermal reduction step of a practical solar thermochemical reactor, T_{H} =2000 K^[18] and P_{O_2} =0.101 kPa^[2]

were used as the upper limit for the reduction temperature and the lower limit for the pressure respectively. For the oxidation step, the exothermic reaction is thermodynamically favorable for low temperatures, but suffers from slow kinetics. Thus, $T_{\rm L}$ of 1000 K^[18] was used as the lower limit for the oxidation temperature. Now, the criteria is summarized as Eq. (7) with the descriptors, $\Delta S_{\rm solid}$ and $\Delta H_{\rm solid}$, defined in Eq. (3).

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$$\begin{vmatrix} \Delta H_{\text{solid}} \leqslant 450 \text{ kJ} \cdot (0.5 \text{ mol } \text{O}_2)^{-1} \\ \Delta H_{\text{solid}} \leqslant T_{\text{H}} \left(\Delta S_{\text{solid}} + \frac{1}{2} S_{\text{O}_2, T_{\text{H}}} - \frac{1}{2} R \ln P_{\text{O}_2} \right) \\ \text{at } T_{\text{H}} = 2000 \text{ K and } P_{\text{O}_2} = 0.101 \text{ kPa} \\ \Delta H_{\text{solid}} \geqslant (\Delta H_{\text{f, CO}, T_{\text{L}}} - \Delta H_{\text{f, CO}_2, T_{\text{L}}}) - \\ T_{\text{L}} (S_{\text{CO}, T_{\text{L}}} - S_{\text{CO}_2, T_{\text{L}}} - \Delta S_{\text{solid}}), \text{ at } T_{\text{L}} = 1000 \text{ K} \end{aligned}$$
(7)

As shown in Fig. 1, the $(\Delta S_{\text{solid}}, \Delta H_{\text{solid}})$ area below the purple curve features theoretical thermal-to-chemical efficiency which is twice higher than the target. $(\Delta S_{\text{solid}}, \Delta H_{\text{solid}})$ below the green curve and above the blue curve are thermodynamically favorable for thermal reduction step (Reaction (2)) and CO₂ splitting step (Reaction (3)) respectively. Therefore, the combination of ΔS_{solid} and ΔH_{solid} of qualified redox material should fall in the shaded triangular region.

These results highlight the importance of accurate predictions of ΔS_{solid} and ΔH_{solid} , which are investigated in Section 2. Fig. 1 also shows that a positive ΔS_{solid} opens the favorable region, so redox materials with large positive ΔS_{solid} benefits reaction thermodynamics. However, this is non-trivial for redox materials in the stoichiometric regime, because the reduced oxide has fewer atoms and thereby, fewer vibrational degrees of freedom.



Fig. 1 Thermodynamic assessment map

The combination of ΔS_{solid} and ΔH_{solid} of redox material candidates should fall in the shaded triangular region. The experimental data points (open scatters) of (ΔS_{solid} , ΔH_{solid}) were plotted for pure CeO₂/CeO_{2- δ} and 10% Samaria-doped Ce_{0.9}Sm_{0.1}O_{1.95}/Ce_{0.9}Sm_{0.1}O_{1.95- δ} redox pairs (δ =0.01–0.05)^[28-29]. The calculation results (solid scatters) in Section 2 were also plotted for these redox pairs (δ =0.03). Error bars represent standard deviations. Colourful figure is available on website

Fig. 1 plots the experimental data^[28-29] of $(\Delta S_{\text{solid}}, \Delta H_{\text{solid}})$ for pure CeO₂/CeO_{2- δ} and 10% Samaria-doped Ce_{0.9}Sm_{0.1}O_{1.95}/Ce_{0.9}Sm_{0.1}O_{1.95- δ} redox pairs (δ =0.01–0.05) as well. Results show that $\Delta S_{\text{solid}}, \Delta H_{\text{solid}}$ of both pure and doped ceria fall in the favorable region except for the pure ceria redox pair with δ =0.01. These results are reasonable because pure and doped ceria are usually used as benchmark redox materials in literature^[5-8].

The favorable region varies with operating temperature and pressure as shown in Fig. 2. Therefore, it is important to recognize that the developed thermodynamic framework can further be used for reactor testing to design optimal operating condition of a specific qualified redox pair. For example, results show that thermodynamically favorable thermal reduction step (Reaction (2)) can be performed at lower temperatures (Fig. 2(a)) and higher pressures (Fig. 2(b)) for pure and doped ceria redox pairs with larger ΔS_{solid} (smaller δ). These milder operating conditions can significantly reduce thermal radiation loss and reactor design complexity.



Fig. 2 Variations of the favorable regions with operating conditions of the thermal reduction step (Reaction (2)) (a) Temperature. $T_{\rm H}$ =2000 and 1773 K; (b) Pressure. $P_{\rm O2}$ =0.101, 1.01 and 101 kPa. Other conditions in (a, b) are the same with those in Fig. 1; Colourful figure is available on website

2 First-principles predictions of descriptors

In this section, first-principles calculations were applied to the predictions of the descriptors (ΔS_{solid} , ΔH_{solid}) so that new redox materials can be added to the thermodynamic assessment map of Fig. 1.

In the first step, fast and reasonable predictions of $(\Delta S_{\text{solid}}, \Delta H_{\text{solid}})$ were demonstrated. ΔS_{solid} in Eq. (8) includes both vibrational (ΔS_{vib}) and configurational (ΔS_{conf}) entropic contributions^[25].

$$\Delta S_{\text{solid}} = \Delta S_{\text{vib}} + \Delta S_{\text{conf}} \tag{8}$$

Vibrational entropic contributions originate from the formations of defects, including oxygen vacancies as well as polarons^[28,30], during the partial reduction step (Reaction (4)).

$$2Ce_{Ce}^{\times} + O_{O}^{\times} \rightarrow V_{O}^{\bullet} + 2Ce_{Ce}^{\prime} + \frac{1}{2}O_{2}$$
 (Reaction (4))

where Ce_{Ce}^{\times} and O_{O}^{\times} represent Ce and O ions on their respective sites, V_{O}^{\bullet} is a doubly ionized oxygen vacancy, and Ce_{Ce}^{\prime} is an electron localized on a Ce ion, *i.e.* polaron. Therefore, ΔS_{vib} is expressed in Eq. (9).

$$\Delta S_{\rm vib} = \Delta S_{\rm V_{\rm O}^{*}} + 2\Delta S_{\rm Ce'_{\rm Ce}} = (S_{\rm V_{\rm O}^{*}} - S_{\rm bulk}) + 2(S_{\rm Ce'_{\rm Ce}} - S_{\rm bulk}) (9)$$

Where S_{bulk} , $S_{V_0^-}$ and $S_{\text{Ce}'_{\text{Ce}}}$ are the entropies of a bulk CeO₂ supercell (Fig. 3(a)), a CeO₂ supercell with a single oxygen-vacancy defect (Fig. 3(b)) and a CeO₂ supercell with a single polaron defect (Fig. 3(c)), taking pure CeO₂ as an example. More details are available in Supporting Materials on the preparations of the supercells and on the calculations of S_{bulk} , $S_{V_0^-}$, $S_{\text{Ce}'_{\text{Ce}}}$ and corresponding ΔS_{vib} .

Configurational entropic contributions are from the random distributions of ionic (oxygen vacancies) and electronic (polarons) defects, so ΔS_{conf} was calculated using the ideal solution model (Methods in Supporting Materials).



Fig. 3 Supercells for DFT+U calculations

(a) Bulk CeO₂ supercell; (b) CeO₂ supercell with a single oxygenvacancy defect; (c) CeO₂ supercell with a single polaron defect(The charge density of the polaron is also shown); (d) Sm-doped CeO₂ supercell (Methods in Supporting Materials) ΔH_{solid} was approximated as the oxygen-vacancy formation energy^[11,21].

 ΔS_{vib} , ΔS_{conf} and ΔH_{solid} of the CeO₂/CeO_{2- δ} and Ce_{0.9}Sm_{0.1}O_{1.95}/ $Ce_{0.9}Sm_{0.1}O_{1.95-\delta}$ redox pairs (δ =0.03) were calculated before the comparison of these calculation results with experimentally measured ΔS_{solid} and $\Delta H_{\text{solid}}^{[28-29,31]}$. Table 1 shows that this applied theoretical method can predict ΔS_{solid} and ΔH_{solid} of pure and Sm-doped ceria pairs, and expectedly other redox materials with reasonable accuracy. Particularly for ΔS_{solid} of the CeO₂/CeO_{2- δ} redox pair (δ = (0.03), it was demonstrated that the relative differences between calculated and measured values are below 12%, although it is found in references[20, 23] that this method might over-estimate ΔS_{solid} . Regarding ΔH_{solid} of pure and Sm-doped ceria pairs (δ =0.03), the relative differences are also below 12%, demonstrating sufficient accuracy of this method which is expected to be more efficient enabled by the advancement and power of high-throughput computational tools^[15-16].

Table 1Comparison between calculated and measured
values of ΔS_{solid} and ΔH_{solid}

Redox pair	Ref.	$\Delta S_{\rm vib}{}^a$	$\Delta S_{\rm conf}^{a}$	$\Delta S_{ m solid}{}^a$	$\Delta H_{\rm solid}^{a}$
	Experimental ^[28]	-	-	~91.2 ^b	~429
$CeO_{2-\delta} (\delta=0.03)$	Experimental ^[31]	-	-	~105.9 ^b	~441
	This work	13.7	80.5	94.2	476.7
$\begin{array}{c} \text{Ce}_{0.9}\text{Sm}_{0.1}\text{O}_{1.95-\delta} \\ (\delta=0.03) \end{array}$	Experimental ^[29]	-	-	$\sim 67.7^{b}$	~400
	This work	9.1	70.3	79.4	376.5

^{*a*} Entropy and enthalpy units are $J \cdot (0.5 \text{ mol } O_2)^{-1} \cdot K^{-1}$ and $kJ \cdot (0.5 \text{ mol } O_2)^{-1}$; ^{*b*} Obtained from measured reduction entropy minus $0.5S_{O_2}$

Then physical insights were given into these calculation results to further demonstrate the reliability of the applied theoretical method. Fig. 4 shows the variations of $\Delta S_{V_{c}}$, $2\Delta S_{Ce'_{Ce}}$, ΔS_{vib} and ΔS_{solid} with temperatures. Since the results for pure and Sm-doped ceria pairs follow the same trend, the $CeO_2/CeO_{2-\delta}$ redox pair is taken as an example. It is observed that $\Delta S_{V_{u}}$ is negative, which can be intuitively understood from less atoms and resultantly lower vibrational degrees of freedom of the reduced oxides. This can also be quantitatively explained by a negative relaxation volume, ΔV_{rel} , due to the formation of an oxygen-vacancy defect, from Eq. (S2) in Supporting Materials. The surrounding O ions relax towards the vacancy for a distance longer than the Ce ions relax away, which renders a relaxation volume of -0.0170 nm³ (Table S1). On the contrary, $2\Delta S_{Ce'_{Ce}}$ is positive, which is resultant from a positive relaxation volume ($\Delta V_{rel}=0.0139 \text{ nm}^3$).

These results indicate that the change of volume, due to the formation of defects, dominates vibrational entropic contribution. The calculated ΔS_{vib} is 13.5 J·(0.5 mol O₂)⁻¹·K⁻¹



Fig. 4 Variations of $\Delta S_{V_0^{in}}$, $2\Delta S_{Ce'_{Ce}}$, ΔS_{vib} , and ΔS_{solid} with temperature Solid and dash curves are for CeO₂/CeO_{2-δ} and Ce_{0.9}Sm_{0.1}O_{1.95}/Ce_{0.9}Sm_{0.1}O_{1.95-δ} redox pairs (δ =0.03), respectively; Colourful figure is available on website

at 1000 K which agrees well with the experimental data^[28], and the calculated ΔS_{conf} is 80.5 J·(0.5 mol O₂)⁻¹·K⁻¹. For the Ce_{0.9}Sm_{0.1}O_{1.95}/Ce_{0.9}Sm_{0.1}O_{1.95- $\delta}$ redox pair, the fact that ΔS_{conf} (70.3 J·(0.5 mol O₂)⁻¹·K⁻¹) is smaller than that of pure ceria redox pair is reasonable because doping by an atom with a lower valence renders a reduction of possible microscopic states. ΔS_{solid} are 94.0 and 79.4 J·(0.5 mol O₂)⁻¹·K⁻¹ for the CeO₂/CeO_{2- δ} and the Ce_{0.9}Sm_{0.1}O_{1.95}/Ce_{0.9}Sm_{0.1}O_{1.95- δ} redox pairs (δ =0.03). These values are comparable to 0.5S_{O2}(121.8 J·(0.5 mol O₂)⁻¹·K⁻¹) at 1000 K. These high positive ΔS_{solid} , attributed to the configurational and polaron defective entropic contributions, explain the excellent thermochemical performance of pure and doped ceria reported in literature [5-8].}

3 Discussion

Based on the results of this work, a viable screening approach of materials for solar-driven CO_2 splitting using two-step thermochemical cycles was proposed as shown in Fig. 5.

The first step is to calculate the descriptors, the combination of ΔS_{solid} and ΔH_{solid} defined in Eq. (3) by the developed DFT+U based first-principles approach. As described in Introduction, despite the recognition of the entropic effect, energy descriptors (enthalpy of formation or energy of oxygen-vacancy formation) were usually used for the screening of material candidates in literature [11-12, 22]. Therefore, $(\Delta S_{\text{solid}}, \Delta H_{\text{solid}})$ represents a more general descriptor.

The second step is to screen material candidates by the derived criteria defined in Eq. (7) and find out qualified redox materials. As described in Section 1, these criteria were derived by comprehensively considering non-stoichiometric reaction mechanism, theoretical efficiency and practical operating conditions. It has to be pointed out that materials with $(\Delta S_{\text{solid}}, \Delta H_{\text{solid}})$ near the boundary of the favorable region (Fig. 1) should also be screened as qualified redox materials considering the accuracy of DFT calculations. In addition, Eq. (6) varies with target solar-to-chemical and thermal-to-chemical efficiencies as well as achievable solar-to-thermal efficiency. For example, $\Delta H_{\rm solid}$ is $\leq 600 \text{ kJ} \cdot (0.5 \text{ mol } O_2)^{-1}$ in Eq. (6) with the target solar-to-chemical efficiency reseted from 20% to 15%. Resultantly, the solid square scatter is inside the favorable region.

The third step is to perform reactor testing using qualified redox material. Thermodynamics of redox material candidates represents an initial screening criterion^[23,32], which was demonstrated in this work. Reaction kinetics of redox material candidates, which is characterized in reactor testing, is the important subsequent criterion^[33-34], although reaction kinetics is out of the scope of this work.

4 Conclusions

In summary, this work presented a rationale for thermodynamic and first-principles assessments of redox materials for solar-driven CO₂ splitting using two-step thermochemical cycles. The combination of solid-state change of entropy (ΔS_{solid}) and enthalpy of formation (ΔH_{solid}) was used as the descriptor. Comprehensive criteria based on it were derived to obtain the (ΔS_{solid} , ΔH_{solid}) map. It was found that a triangular region in this map identified qualified material candidates featuring large positive ΔS_{solid} and small enough ΔH_{solid} . Furthermore, a DFT+U based



Fig. 5 Illustration of a viable screening approach

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Supporting materials

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太阳能驱动的两步热化学循环二氧化碳裂解反应 活性材料的热力学与第一性原理评价

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摘 要:太阳能驱动两步热化学循环裂解二氧化碳可制备碳中性燃料,为替代化石燃料、缓解全球变暖提供了技术 途径。新型活性材料的开发对该技术非常重要。已有研究通常采用能量描述符(材料生成焓或氧空位生成能)评价候 选材料,忽略了材料熵的重要性。本研究采用活性材料的熵和生成焓的组合作为描述符,提出评价准则,开展材料 可行性的热力学分析。结果表明,活性材料应兼具较大的正的熵变与较小的生成焓变。在此基础上,本研究以氧化 铈和钐掺杂的氧化铈为例,发展了基于第一性原理的活性材料熵和生成焓的计算方法,为新型材料的筛选与开发 提供基础。计算结果揭示了极化子振动熵以及氧空位和极化子构型熵对活性材料熵变的贡献。

关键 词:二氧化碳裂解;两步热化学循环;第一性原理;熵;太阳能制燃料

中图分类号: TK51 文献标志码: A

Supporting materials:

Thermodynamic and First-principles Assessments of Materials for Solar-driven CO₂ Splitting Using Two-step Thermochemical Cycles

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Methods

General setups of first-principles calculations All first-principles density functional theory (DFT) calculations were performed using the Vienna Ab-initio Simulation Package (VASP)^[S1] where the Perdew-Burke Ernzerhof (PBE)^[S2] generalized gradient approximation (GGA) and the projector augmented wave (PAW)^[S3] method were used. The wave functions were expanded in plane waves with 500 eV energy cut-off. The 2×2×2 Monkhorst-Pack mesh was used for k-point sampling. Convergence parameters of 10^{-8} eV and 10^{-3} eV nm⁻¹ were used for the relaxation of electrons and ions respectively. The $5s^{2}5p^{6}6s^{2}5d^{1}4f^{1}$ electrons were treated as the valence electrons for Ce with 2s²2p⁴ for O and 5s²5p⁶6s²5d¹ for Sm. The +U approach (U=5 eV) was employed to describe the localized 4f-orbitals of Ce^[S4]. Sm has six f-electrons, so its localized f-electrons was treated with the standard model in GGA potential where five f-electrons were placed in the core (Sm adopts a valency of 3 in doped ceria)^[S5].

Preparations of bulk and defective supercells All $2 \times 2 \times 2$ fluorite supercells contain 96 lattice positions (Fig. 3). For defective supercells, the number of electrons was set corresponding to their charge states. The method reported by Zacherle, et al.[S6] was used to prepare the defective supercell with a single polaron (Fig. 3(c)). For the 10% Sm-doped ceria supercell (Fig. 3(d)), Sm atoms were placed symmetrically to reduce computational costs. The effect of random distribution was included in the calculations of ΔS_{conf} . For all supercells, imaginary frequencies were not observed in their phonon dispersion demonstrating the stability of these structures. Fig. S1 exhibits representative phonon dispersion results of a 10% Sm-doped ceria supercell comprising an oxygen vacancy. The calculated lattice constant of ceria is 0.5499 nm, which agrees well with the experimental value of 0. 5411 nm^[S7], and with the one (0.5494 nm) calculated using GGA+Umethod^[S6]. The slight difference from the experimental value is due to intrinsic overestimation of lattice constants by GGA+U method^[S6].

Phonon calculations The vibrational entropy of each supercell (Eq. (9)) was expressed as Eq. (S1) using the harmonic approximation.

$$S_{\text{bulk/V}_{0}^{\circ}/\text{Ce}_{\text{Ce}}^{\prime}} = Nk_{\text{B}} \int_{0}^{\infty} \left\{ \frac{\hbar\omega}{2k_{\text{B}}T} \coth\left(\frac{\hbar\omega}{2k_{\text{B}}T}\right) - \ln\left[2\sin h\left(\frac{\hbar\omega}{2k_{\text{B}}T}\right)\right] \right\} g(\omega) d\omega$$
(S1)

where, *N* is the number of degrees of freedom, $k_{\rm B}$ is the Boltzmann constant, \hbar is the Planck constant, *T* is the temperature. The normalized phonon density of states, $g(\omega)$, at each phonon frequency, ω , was calculated using the open source package of Phonopy^[S8]. Phonon calculations were performed for constant volume conditions $(V=V_0)$ where the stress is not relaxed due to defect formation, while entropies for constant pressure conditions $(p=p_0)$ in Eq. (9) were more concerned experimentally. They were obtained by Eq. (S2)^[S9].

$$\Delta S_{\text{V}_{0}^{\text{c}}/\text{Ce}_{\text{Ce}}^{\text{c}}}(p=p_{0}) = \Delta S_{\text{V}_{0}^{\text{c}}/\text{Ce}_{\text{Ce}}^{\text{c}}}(V=V_{0}) + \alpha_{\text{V}}K_{\text{T}}\Delta V_{\text{rel}} (\text{S2})$$

where α_V is the volumetric thermal expansion coefficient, K_T is the isothermal bulk modulus and ΔV_{rel} is the relaxation volume, *i.e.*, the difference between the volume of a defective supercell (oxygen-vacancy or polaron) and that of a bulk supercell. The free energy (*E*) of 11 unit cells with volumes (*V*) ranging from 0.03656 nm³ to 0.04656 nm³ was calculated for each temperature. These *E-V* curves for each temperature were fitted using the Birch-Murnaghan equation(Eq. (S3))^[S9] to obtain K_T and equilibrium volumes (*V*₀) for each temperature.

$$E(V) = E_0 + \frac{9V_0K_T}{16} \left\{ \left[\left(\frac{V_0}{V} \right)^{\frac{2}{3}} - 1 \right]^3 K'_T + \left[\left(\frac{V_0}{V} \right)^{\frac{2}{3}} - 1 \right]^2 \left[6 - 4 \left(\frac{V_0}{V} \right)^{\frac{2}{3}} \right] \right\}$$
(S3)

Then, α_V was derived from equilibrium volumes as Eq. (S4).

$$\alpha_{\rm V} = \frac{1}{V_0} \left(\frac{\partial V_0}{\partial T} \right)_{\rm p} \tag{S4}$$

Calculated $\alpha_{\rm V}$, $K_{\rm T}$ and $\Delta V_{\rm rel}$ are available in Fig. S2 and Table S1.

Calculations of configurational entropic contributions ΔS_{conf} was calculated using the ideal solution model. For a general $\text{Ce}_{1-y}\text{Sm}_y\text{O}_{2-0.5y}/\text{Ce}_{1-y}\text{Sm}_y\text{O}_{2-0.5y-\delta}$ redox pair, ΔS_{conf} was calculated as Eq. (S5)^[S10].

$$\Delta S_{\text{conf}} = -R \ln \left[\frac{0.5y + \delta}{2 - 0.5y - \delta} \left(\frac{2\delta}{1 - y - 2\delta} \right)^2 \right] \quad (S5)$$

Supporting figures and tables



Fig. S1 Phonon dispersion of a 10% samaria-doped ceria supercell comprising an oxygen vacancy



Fig. S2 Calculated variations of (a) α_V , (b) K_T and (c) $\alpha_V \times K_T$ with temperatures

Table S1 Relaxation volumes due to formations of an oxygen vacancy and a polaron for $CeO_{2-\delta}$ and $Ce_{0.9}Sm_{0.1}O_{1.95-\delta}$

Oxide	$\Delta V_{\rm rel, vacancy}/~{\rm nm}^3$	$\Delta V_{\rm rel,\ polaron}/{\rm nm}^3$		
${ m CeO}_{2-\delta}$	-0.0170	0.0139		
$Ce_{0.9}Sm_{0.1}O_{1.95-\delta}$	-0.0170	0.0144		

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