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Introduction

thermochemical splitting cycles.

Solar thermochemical splitting cycles have been gaining scientific and technological relevance as an environmentally sustainable route to produce synthetic fuels from H₂O and CO₂. Actually, they enable straight solar energy harvesting and conversion into synthesis gas (syngas), a CO/H₂ mixture, that can be further processed to gaseous or liquid fuels.1-5

Among available technologies, two-step thermochemical redox cycles based on non-stoichiometric metal oxides (MO_r) hold huge promise, exhibiting the lowest complexity as well as the highest theoretical solar-to-fuel energy conversion efficiency $(\eta_{\text{solar-to-fuel}})$ as a result of the use of the whole solar spectrum.

In the first step of a typical red-ox cycle, MO_x undergoes high temperature self-reduction to $MO_{(x-\delta)}$, according to the following reaction:

$$MO_x \rightarrow MO_{(x-\delta)} + \delta/2O_2$$

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Redox behavior of potassium doped and transition metal co-doped Ce_{0.75}Zr_{0.25}O₂ for thermochemical H₂O/CO₂ splitting[†]

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CeO₂ slow redox kinetics as well as low oxygen exchange ability limit its application as a catalyst in solar thermochemical two-step cycles. In this study, Ce0.75Zr0.25O2 catalysts doped with potassium or transition metals (Cu, Mn, Fe), as well as co-doped materials were synthesized. Samples were investigated by X-ray diffraction (XRD), N_2 sorption (BET), as well as by electron paramagnetic resonance (EPR) and X-ray photoelectron spectroscopy (XPS) to gain insight into surface and bulk features, which were connected to redox properties assessed both in a thermogravimetric (TG) balance and in a fixed bed reactor. Obtained results revealed that doping as well as co-doping with non-reducible K cations promoted the increase of both surface and bulk oxygen vacancies. Accordingly, K-doped and Fe-K codoped materials show the best redox performances evidencing the highest reduction degree, the largest H₂ amounts and the fastest kinetics, thus emerging as very interesting materials for solar

> Then, in the second step, $MO_{(x-\delta)}$ is oxidized by CO_2 and/or H₂O to form CO and/or H₂ as described by the following equations:

> > $MO_{(x-\delta)} + \delta CO_2 \rightarrow MO_x + \delta CO$ $MO_{(x-\delta)} + \delta H_2O \rightarrow MO_x + \delta H_2$

Ceria (CeO_2) is considered the most attractive material for these processes, combining good thermal stability, high oxygen storage capacity (OSC) without any structural changes as well as faster reduction and splitting kinetics than other oxides.^{1,6} Notably, CeO₂ based materials, particularly doped samples, often can outperform perovskites, affording higher solar-to-fuel efficiency above all in the CO₂ splitting reaction.⁷

However, several major drawbacks still prevent their technological application. These include the limited amount of oxygen vacancies as well as high reduction temperature, which involves poor cyclability due to structural modifications.

Partial substitution of Ce(IV) species and doping with metal cations in the fluorite structure of ceria is a successful strategy to address these issues, enhancing the redox activity of the material.8-19

Density-functional-theory (DFT) calculations¹³ support experimental results²⁰ indicating that Ce(IV) substitution by Zr(w) ions could improve ceria splitting properties and

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particularly its self-reduction behavior. Among the investigated compositions, Ce_{0.75}Zr_{0.25}O₂ showed the best catalytic cyclability.^{21–25} Moreover, bi- or trivalent cations doping, including Cu(π), Mn(π), Fe(π), can improve both self-reducibility and splitting activity of bare CeO₂.^{16–18} Notably, Terribile *et al.* (1999) analyzed the reduction and catalytic behavior of Mn- and Cudoped ceria–zirconia solid solutions. They found that manganese and copper substitution created ionic defects in ceria fluorite structure, by a charge compensation mechanism, which promoted low-temperature redox activity, involving reduction of Cu²⁺ to Cu⁰ and of Mn^{3+/4+} to Mn^{2+,26}

Recently, Takalkar *et al.* (2020) showed that doping with alkali (Li), alkaline earth (Mg, Ca, Sr, Ba), and post-transition (Sn) metal improved both O_2 release and CO production during CO_2 red-ox splitting cycles, due to the formation of mixed phases neglected by the DFT calculations.²⁷ Moreover, Ruan *et al.* (2017) reported that the formation of Ce₂Sn₂O₇ pyrochlore phase in CeO₂–SnO solid solution, boosted both self-reduction and splitting activity.²⁸ Similarly, according to Pappacena *et al.* (2016 and 2017), the formation of nanostructured Zr₂ON₂-like phase in CeO_{1.5}Zr_{0.85}O₂ promoted splitting.^{29,30}

An intimate mixing at the nanoscale of two different active sites involved in oxygen evolution and splitting reactions has been emerging as successful strategy for the design of high-performance catalysts.^{31,32}

In order to improve ceria thermochemical red-ox performance, Charvin *et al.* (2009) proposed a three-step cycle.³³ After the usual solar-driven self-reduction step, the process involved reaction between reduced oxide and an alkali hydroxide (NaOH or KOH), producing hydrogen and a mixed oxide. Its subsequent hydrolysis regenerated the starting oxide and the alkali hydroxide. Furthermore, potassium cation doping has been reported to improve perovskite-based catalysts performance towards water gas shift (WGS) reaction.³⁴ Moreover, for the same process, Zugic *et al.* (2014) suggested that the addition of the electropositive alkali metal cations promoted the formation of active –OH species, thus increasing adsorption active sites number with high thermal stability.³⁵

Inspired by literature data, in this study, we investigated the effect of K(1) doping on Ce_{0.75}Zr_{0.25}O₂ (CeZr) red-ox activity towards both CO2 and H2O splitting. K was also investigated as co-dopant with transition metal cations (M = Cu, Mn, Fe)which were found to improve CeZr thermochemical performance and were also used as reference to compare the performances of K-doped samples. Prepared materials were submitted to in-depth physical-chemical characterization using different techniques. Furthermore, thermogravimetric analysis (TGA) was carried out to assess redox activity towards CO₂ splitting during repeated cycles, whereas flow reactor tests were performed to investigate catalytic activity towards H₂O splitting. Particularly, the outcomes of catalytic experiments were combined with XPS and EPR spectroscopy results, providing key information on the molecular features underlying redox behaviour, thus drawing strategic guidelines for the design of high-performance catalysts for thermochemical splitting cycles.

Experimental section

Materials

The following reagents were employed for the catalyst synthesis: cerium(III) nitrate hexahydrate, Ce(NO₃)₃·6H₂O; zirconium(IV) oxynitrate hydrate, ZrO(NO₃)₂·xH₂O; iron(III) nitrate nonahydrate, Fe(NO₃)₃·9H₂O; potassium nitrate, KNO₃; copper(II) nitrate tetrahydrate, Cu(NO₃)₂·4H₂O; manganese(II) nitrate tetrahydrate, Mn(NO₃)₂·4H₂O. All products were purchased from Sigma-Aldrich and used as received.

Catalyst synthesis

To achieve high reduction behaviour, Ce/Zr molar ratio was kept constant at 3 ($Ce_{0.75}Zr_{0.25}O_2$ as general formula), since it ensured high reducible behaviour.³⁶

Bare ceria–zirconia, M-doped (M = Cu, Fe, Mn) and K-doped ceria–zirconia materials, as well as co-doped samples were prepared according to the co-precipitation method described elsewhere.³²

Briefly, stoichiometric amounts of cerium and zirconium precursors were dissolved in 75 ml bi-distilled water and then stoichiometric quantity of the doping metal was added and stirred for 3 h. Afterwards, solutions were heated in a microwave oven (CEM SAM-155) and the resulting gel was calcined in air at 1100 °C for 4 h. Table 1 reports obtained material with their nominal and ICP measured compositions as well as the label used thereof, whereas Fig. S1† shows the co-precipitation synthesis steps of the doped ceria materials.

Materials characterization

ICP-MS analysis (Agilent 7500CE instrument) was used to assess the actual composition of the samples. The measured compositions differed from the nominal ones within the experimental error ($\pm 5\%$).

X-ray diffraction (XRD) measurements were performed with a PANalytical X'Pert Pro XRD diffractometer (step size: 0.02° ; counting time: 80 s per step). Scherrer equation was employed to calculate the average crystal size (τ):

$$\tau = \frac{K\lambda}{\beta\cos\theta} \tag{1}$$

where τ is the mean size of the crystallite domains, *K* is a dimensionless shape factor, λ is the X-ray wavelength, β is the broadening at half the maximum intensity (FWHM), θ is the Bragg angle (in radiant). Bragg's law was used to determine interplanar spacing (*d*):

$$n\lambda = 2d\sin(\theta) \tag{2}$$

Interplanar spacing values were employed to evaluate the cell parameter (a), using the equation:

$$d = \frac{a}{\sqrt{h^2 + k^2 + l^2}}\tag{3}$$

 N_2 adsorption at 77 K with a Quantachrome Autosorb-1C instrument was employed to measure BET specific surface

Table 1Materials tested for the thermochemical splitting process, their nominal and ICP measured compositions (molar ratios), generalformulas, and the highest oxygen amount (mmol g^{-1}) theoretically produced in the reduction step

	Theoretical		ICP				n _{O2} ,max
Sample	M/(Ce + Zr)	K/(Ce + Zr)	M/(Ce + Zr)	K/(Ce + Zr)	Ce/Zr	General formula	
CeZr	_	_			3.02	Ce _{0.75} Zr _{0.25} O ₂	1.17
Fe-CeZr	0.05	—	0.0523		3.14	Fe _{0.05} Ce _{0.71} Zr _{0.24} O _{1.98}	1.22
Mn-CeZr	0.05	_	0.0479		2.97	Mn _{0.05} Ce _{0.71} Zr _{0.24} O _{1.98}	1.29
Cu-CeZr	0.05	—	0.0521		2.89	Cu _{0.05} Ce _{0.71} Zr _{0.24} O _{1.95}	1.3
K-CeZr	—	0.05		0.0518	3.11	K _{0.05} Ce _{0.71} Zr _{0.24} O _{1.93}	1.16
K-Fe-CeZr	0.05	0.05	0.0485	0.0522	3.05	$K_{0.045}Fe_{0.045}Ce_{0.68}Zr_{0.23}O_{1.91}$	1.2
K-Cu-CeZr	0.05	0.05	0.0509	0.0488	2.94	K _{0.045} Cu _{0.045} Ce _{0.68} Zr _{0.23} O _{1.88}	1.28

areas (SSA); samples were degassed at 150 $^\circ \mathrm{C}$ for 1.5 h before each measurement.

X-ray photoelectron spectroscopy (XPS; XPS PHI 5000 Versa probe instrument) analyses were performed on both pristine and used samples. The band-pass energy, the take-off angle, and the X-ray spot size for survey spectra were set at 187.85 eV, 45° , and 100.0 μ m diameter respectively. High-resolution XPS spectra were recorded using the following experimental conditions: pass energy 20 eV, resolution 1.1 eV and step 0.2 eV. The C 1s line with a binding energy (BE) value of 284.8 eV was used as internal reference in order to compensate sample charging effects. XP-spectra were deconvoluted by means of a commercial software (CasaXPS, version 2.3.17), using mixed Gaussian–Lorentzian (70–30%) profiles.

Electron paramagnetic resonance (EPR; X-band (9 GHz) Bruker Elexys E-500 spectrometer (Bruker, Rheinstetten, Germany), equipped with a super-high sensitivity probe head) measurements were carried out on selected as-prepared and used samples. Solid samples were placed into flame-sealed glass capillaries which, in turn, were coaxially put into a standard 3 mm quartz sample tube. EPR spectra were recorded with an attenuation of 15 dB and by accumulating 128 scans to improve the signal-to-noise ratio. Details of the measurements settings are reported elsewhere.³⁷ A quantitative analysis was performed by determining the *g*-factor and ΔB values, as previously described.³⁷

Thermochemical CO₂ splitting tests

Thermochemical CO₂ splitting cycles were carried out in a TGA/ DSC apparatus (TA Q600SDT), according to the procedure reported elsewhere³² and briefly described hereafter. Fine powdered samples (about 30 mg) were put into an alumina crucible, first pre-treated in air at 1200 °C (heating rate: 20 °C min⁻¹) to remove any chemisorbed CO₂, cooled down to room temperature and purged under a nitrogen flow for 1 h. Then, the samples were submitted to five consecutive reduction (1350 °C) and CO₂ splitting steps (1000 °C). During each selfreduction, the sample was heated up to 1350 °C (heating rate: 20 °C min⁻¹), and then kept at this temperature for 20 min under a nitrogen flow (0.1 l (STP) min⁻¹). Afterwards, the sample was cooled down to 1000 °C (cooling rate: 40 °C min⁻¹), the gas flow was fast changed to pure CO_2 and kept for 60 min.

The following equations were used to evaluate the amount of gaseous products (mol g^{-1}) released during the self-reduction and oxidation step respectively:

$$n_{\rm O_2} = \frac{\Delta m_{\rm loss}}{m_{\rm cat} \times M_{\rm O_2}} = \frac{m_{\rm loss(\%)}}{100 \times M_{\rm O_2}} \tag{4}$$

$$n_{\rm CO} = \frac{m_{\rm gain(\%)}}{100 \times M_{\rm O}} \tag{5}$$

where Δm_{loss} is the mass loss (g), $m_{\text{loss}(\%)}$ is the % mass change evaluated by TGA starting from 200 °C to exclude any weight loss due the moisture release, M_{O_2} indicates O_2 molecular weight (g mol⁻¹), m_{cat} is the catalyst mass (g) used during each experiment, $m_{\text{gain}(\%)}$ is the mass gain percentage during the splitting step, and M_O is oxygen atomic weight (g mol⁻¹).

The performance of the samples was assessed by evaluating the reduction degree ($x_{red,i}$) after each reduction or oxidation step (*i*), the oxidation yield (α_i) and the reduction yield (β_i) as follows:

$$x_{\text{red},i} = x_{\text{red},i-1} + \frac{n_{\text{O}_2,i}}{n_{\text{O}_2,\text{max}}} \quad (\text{reduction step}) \tag{6a}$$

$$x_{\text{red},i} = x_{\text{red},i-1} - \frac{n_{\text{CO},i}}{2 \times n_{\text{O}_{2,\text{max}}}}$$
 (oxidation step) (6b)

$$\alpha_i = \frac{n_{\text{CO},i}}{2 \times n_{\text{O}_2,i-1}} \tag{7}$$

$$\beta_i = \frac{2 \times n_{\mathrm{O}_2,i}}{n_{\mathrm{CO},i-1}} \tag{8}$$

where $n_{O_2,max}$ is the maximum O_2 amount (mol g⁻¹) that could be released if reducible cations were completely reduced to their lower oxidation state (Ce⁴⁺ to Ce³⁺, Fe³⁺ to Fe²⁺, Cu²⁺ to Cu⁰, Mn⁴⁺ to Mn²⁺; amounts reported in Table 1) and $x_{red,i-1}$ is the reduction degree of the previous (oxidation or reduction) step. The oxidation (α_i) and reduction (β_i) yield are calculated through the mass change during the current step (re-oxidation or reduction) and the amount of evolved gaseous product (O₂ or CO) during the previous step (reduction or re-oxidation). The schematic of TGA apparatus is reported in Fig. S2.[†]

Thermochemical H₂O splitting tests

 $\rm H_2O$ splitting activity of prepared samples was assessed following an experimental already employed in previous papers.^{32,37,38} Briefly, powdered samples (500 mg; 170–300 μ m) were submitted to consecutive temperature programmed reduction (TPR) and oxidation (TPO) runs in a tubular quartz reactor externally heated by an electric tubular furnace (Lenton). Temperature was measured by a K-type thermocouple hosted inside a co-axial tube.

Evolved gases were continuously analysed by a Fisher-Rosemount NGA2000 analyser which analyses CO and CO_2 by infrared detectors, O_2 by a paramagnetic detector and hydrogen by a thermal conductivity detector. The analyser is provided with a cross-sensitivity system.

Samples were heated up to about 1000 °C at 10 °C min⁻¹ in H₂ mixture flow (TPR; 10 l (STP) h⁻¹; 5 vol% H₂/N₂)^{37,38} and then cooled down to 60 °C under N₂ flow. TPO was then performed switching the atmosphere to 3 vol% H₂O/N₂ mixture (10 l (STP) h⁻¹), heating up to 1000 °C (heating rate: 10 °C min⁻¹) and keeping the sample at this temperature for 20 min. The use of diluted water mixture was previously reported³⁹ to enable reaction rate control, particularly in the case of H₂-reduced samples. Reduction and oxidation profiles obtained with the gas analyser were deconvoluted using OriginPro 8.5 software.

Oxygen and hydrogen amounts (mol g^{-1}), n_{O_2} and n_{H_2} , respectively, were evaluated as follows:

$$n_{\text{O}_{2,i}} = \frac{2 \times n_{\text{H}_2_\text{cons},i}}{m_{\text{cat}}} = \frac{2 \times \frac{\text{area}_{\text{H}_2_\text{cons},i}}{100} \times F_{\text{red}} \times \frac{P}{R \times T}}{m_{\text{cat}}} \quad (9)$$

$$n_{\mathrm{H}_{2,i}} = \frac{n_{\mathrm{H}_{2}\mathrm{-prod},i}}{m_{\mathrm{cat}}} = \frac{\frac{\operatorname{area}_{\mathrm{H}_{2}\mathrm{-prod},i}}{100} \times F_{\mathrm{ox}} \times \frac{P}{R \times T}}{m_{\mathrm{cat}}}$$
(10)

where m_{cat} (g) is the sample mass, P (atm) and T (K) indicate standard pressure and temperature, respectively, R (atm l mol⁻¹ K⁻¹) is the ideal gas constant. F_{red} and F_{ox} (l (STP) s⁻¹) express the gases flow rates during H₂ reduction and splitting step, respectively, $n_{\text{H}_2-\text{cons},i}$ and $n_{\text{H}_2-\text{prod},i}$ are the amount of hydrogen consumed and produced during reduction and splitting step, respectively, and area_{H_2-}cons,*i* and area_{H_2-}prod,*i* (% s) are the areas of reduction and oxidation H₂ profiles, respectively. During these experiments, n_{O_2} indicate the oxygen amount removed from the materials during TPR rather than a real molecular oxygen evolution. Fig. S3† reports the schematic of the quartz reactor setup is reported.

Results and discussion

Materials characterization

Prepared samples were submitted to an in-depth physicochemical characterization to elucidate the microscopical features accounting for their red-ox behaviour.

XRD analysis was carried out on investigated samples both before (fresh samples, treated up to 1100 $^{\circ}$ C) and after a redox cycle in the TG apparatus (used samples, treated up to 1350 $^{\circ}$ C). Fig. 1 shows the XRD patterns of the used samples, while lattice

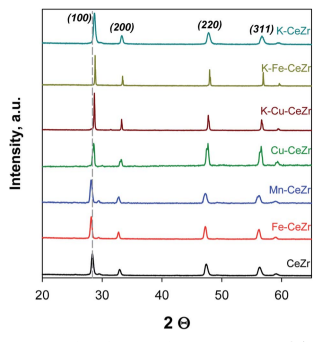


Fig. 1 XRD profiles of samples after a redox cycle in TG (used samples). Miller indices relative to fluorite CeO_2 from Terribile *et al.*²⁶

parameters are reported in Table S1.† All samples show a fluorite structure typical of CeO₂ lattice, no additional phases due to doping species being detectable. Moreover, the absence of any other significant diffraction peak in the XRD profiles confirms the formation of a solid solution, through the incorporation of Zr⁴⁺ into CeO₂ lattice. In Fe-CeZr and Mn-CeZr materials the reflection of the (111) plane at $2\theta = 28.4^{\circ}$ shifts to lower angles than bare CeZr; this could indicate the formation of solid solutions. However, it is known that lower valence ions such as Fe³⁺ and Mn³⁺ are extremely difficult to dissolve into the ceriazirconia lattice, especially when treating at high temperature.40 Thus, some iron and manganese ions could also be encapsulated in ceria-zirconia lattice as free mixed (Fe,Mn)₂O₃ or (Fe,Mn)O nanostructured oxides, not detectable by XRD analysis.⁴¹ Due to their smaller size than Ce^{4+} (ionic radius = 0.97 Å), Fe^{3^+} (ionic radius = 0.55 Å) and Mn^{2^+} (ionic radius = 0.67 Å) species are expected to be incorporated as interstitial defects, which should produce an expanded lattice with a consequent shift of XRD peaks to lower angles, as appreciated in XRD patterns of doped samples. Conversely, potassium doping induces a decrease in cell parameter, as confirmed by the shift of (111) reflection at higher angles. This could be due to the formation of K₂O species segregated into CeO₂-ZrO₂ lattice.⁴²

Furthermore, a small peak at $2\theta = 29.4^{\circ}$ can be appreciated in CeZr, Fe-CeZr and Mn-CeZr XRD patterns probably due to the presence of some segregate tetragonal ZrO₂ phase. XRD spectra of fresh samples are quite similar to those of used materials. However, some of them, particularly fresh K-Fe-CeZr, show wider and less intense peaks (Fig. S4†), indicating that the reduction step at 1350 °C causes an increase in the average crystal size. BET surface areas (Table S1[†]) are very low, as expected for samples treated at high temperature.

XPS and EPR spectroscopies were carried out on prepared samples in order to assess surface and bulk oxygen vacancies, respectively, which determine oxygen exchange ability, governing CeO₂ catalytic performance.⁴³ Notably, surface properties of fresh and used samples were investigated by means of XPS to assess any changes in surface composition after splitting tests. Table 2 reports Ce/Zr ratio, Ce oxidation states as evaluated from high-resolution XPS spectra and the curve-fitting of the Ce 3d region, respectively. The relative amount of surface labile oxygen (O_{α}) and bulk oxygen (O_{β}) moles) are reported in Table S2.[†]

The nominal bulk Ce/Zr atomic ratio is equal to 3. In general, all fresh samples show a decrease in this ratio, indicating a zirconium surface enrichment. This segregation is more evident for the CeZr sample and gradually decreases in the doped samples, with the following order: CeZr > Mn-CeZr > Fe-CeZr > K-Cu-CeZr > Cu-CeZr > K-CeZr > K-Fe-CeZr. This can be attributed to the surface dispersion of the metal cation species which partly replace the Zr atoms.44 The highest depletion in Zr content is disclosed by K-Fe-CeZr sample, where the Ce/Zr ratio is almost 4, thus suggesting a Zr depletion of the surface. After the splitting tests, the Ce/Zr ratio increases for CeZr sample, in agreement with our previous work,37 whilst it is considerably smaller than the fresh catalyst for the doped samples, in accordance with Trovarelli et al.30 From the analysis of XPS data it can be inferred a zirconium enrichment, which grows in the following order: K-CeZr < Fe-CeZr < Mn-CeZr < K-Cu-CeZr < Cu-CeZr < K-Fe-CeZr. In particular, the K-Fe-CeZr sample shows the biggest decrease in surface Ce content, since it passes from an initial superficial Zr depletion to a final remarkable enrichment. As reported in the literature, the formation of Zr-rich phases enhances the splitting activity.30 Eventually, this could explain better red-ox performance of doped samples than bare CeZr that will be investigated in the following sections.

The high-resolution XPS spectra of the Ce 3d region and the relative curve-fitting are illustrated in Fig. S5.[†] According to the literature, 10 components were selected, "u" and "v", corresponding to Ce $3d_{3/2}$ and Ce $3d_{5/2}$ spin-orbits, respectively.⁴⁵ The u", u, v", v, v peaks were associated to the Ce⁴⁺ state while the u', u₀, v' and v₀ peaks were attributed to the Ce³⁺ state.⁴⁶⁻⁴⁸ As can be appreciated in Table 2, the fresh doped samples show

a significant increase in superficial Ce³⁺ with respect to the CeZr sample. This suggests that the atoms of the transition metals can enter into the CeZr lattice and then Ce4+ can be partially converted to Ce³⁺.⁴⁴ As reported in previous studies, Ce³⁺ ions concentration is associated to the formation of oxygen vacancies,49 which can promote oxygen migration during selfreduction as well as the dissociation of the O-H bonds in the H₂O splitting reaction.⁴³ Ce³⁺/Ce⁴⁺ atomic ratio after the splitting tests follows different trends: it keeps constant in CeZr, K-CeZr and K-Fe-CeZr, and significantly decreases in Cu-CeZr and Mn-CeZr samples. On the other hand, Fe-CeZr and K-Cu-CeZr show a marked increase of this ratio, evidencing the ability to keep a high concentration of surface Ce³⁺ ions that can be available for next cycles. This behavior is envisaged to promote CO/H₂ production. Indeed, surface enrichment in Zr species (decrease in the Ce/Zr ratio) often determines an increase of the ${\rm Ce}^{3+}\!/{\rm Ce}^{4+}$ atomic ratio since smaller ${\rm Zr}^{4+}$ ions could itself promote the conversion of Ce⁴⁺ ions into larger Ce³⁺ species in order to compensate for lattice distortion.

Since splitting reaction often involves bulk sites, bulk properties were investigated through electron paramagnetic resonance (EPR) analysis. EPR spectra of CeZr samples doped with different metals were recorded at room temperature and are shown in Fig. 2. Different behaviors were observed, also due to

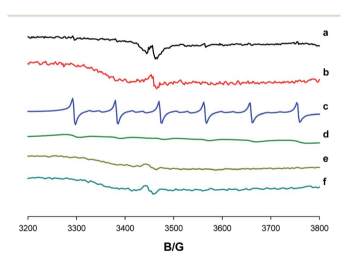


Fig. 2 EPR spectra of K-Cu-CeZr, Mn-CeZr and Fe-CeZr samples, fresh (a, c and e) and treated at 1350 $^\circ\text{C}$ (b, d and f).

Sample	Fresh				Used			
	Ce/Zr	M/(Ce + Zr)	K/(Ce + Zr)	Ce ³⁺ /Ce ⁴⁺	Ce/Zr	M/(Ce + Zr)	K/(Ce + Zr)	Ce ³⁺ /Ce ⁴⁺
CeZr	1.84			0.24	2.80			0.30
Fe-CeZr	2.46	0.31		0.31	1.93	0.28		0.73
Mn-CeZr	2.29	0.32		0.59	1.36	0.26		0.44
Cu-CeZr	2.73	0.60		0.70	1.41	0.47		0.39
K-CeZr	2.86		0.15	0.36	2.70		0.06	0.41
K-Fe-CeZr	3.92	0.30	0.06	0.55	2.17	0.40	0.15	0.58
K-Cu-CeZr	2.48	0.08	0.14	0.53	1.43	0.37	0.35	1.01

 Table 2
 Surface atomic composition of fresh and used samples by XPS analysis

the presence of various paramagnetic species. For the fresh K-Cu-CeZr, a single asymmetric broad peak with a g-factor value of \sim 1.968 \pm 0.003 is observed (Fig. 2a) and presents a line-shape amplitude $\Delta B = 8.4 \pm 0.2$ G. This peak is commonly assigned to surface Ce³⁺ ions that contain unpaired electrons.⁵⁰⁻⁵² After the treatment at 1350 °C, a slight increase in the peak intensity is observed confirming increase of Ce³⁺ species (Fig. 2b), in accordance with XPS results. For the fresh K-CeZr sample no signal is appreciated (spectra not shown), suggesting that no paramagnetic species can be clearly revealed. On the other hand, a weak signal at g-factor value of \sim 1.971 \pm 0.004, ascribable to Ce³⁺ ions,⁵⁰ is observed for fresh Fe-CeZr (Fig. 2e), which gets slightly more intense after treatment (Fig. 2f), confirming an increase in the paramagnetic species as also indicated by XPS results. For the Mn-CeZr sample, the EPR spectrum shows the characteristic six-line hyperfine splitting pattern of Mn^{2+} at g-factor value of 2.004 \pm 0.003 (Fig. 2c), due to the interaction of electron spin (S = 5/2) of Mn²⁺ ions with nuclear spin I = 5/2.⁵³ Being the EPR signal of Mn²⁺ characterized by a high intensity, it is not possible to discriminate the presence of any Ce3+ signal. However, after treatment at 1350 °C, a strong decrease in Mn²⁺ peaks intensity is detected (Fig. 2d), suggesting a change in its oxidation state.

Interestingly, even K^+ species which cannot undergo any reduction, are able to influence the Ce^{3+}/Ce^{4+} ratio, once employed as dopants. This behavior has been more extensively investigated in a previous study.⁵⁴

Thermochemical CO₂ splitting

Redox activity towards CO_2 -splitting was investigated by repeated thermochemical cycles performed in TG apparatus.^{37,38} The reduction of samples was appreciated through a mass loss in TG curve due to evolved O_2 (Fig. S6†), and the corresponding onset temperatures are displayed in Table 3.

Bare CeZr and Fe-CeZr samples show an onset temperature of about 1150 °C, whereas the reduction process starts at lower temperatures for all the other investigated materials. Notably, Cu-doped samples are reduced in two steps the former of which occurs at about 750 °C, producing 156 μ mol g_{cat}^{-1} of oxygen. TG curves of all doped samples display higher slope change than bare CeZr in the reduction step suggesting faster self-reduction kinetics (DTG curves, Fig. S7†). Overall, Cu containing materials featured higher weight losses than bare CeZr indicating both larger evolved oxygen amount and higher reduction degree (Table 3).

After reduction, the samples underwent a consecutive oxidation step in TG apparatus, that was carried out at 1000 °C under a CO_2 flow (Fig. S8†). The amount of produced CO was driven from the recorded mass gain in TG curve, due to material oxidation through eqn (5) (Table 3).

Cu-CeZr showed poor activity towards CO₂ splitting and overall oxidation yield comparable to bare CeZr in spite of its high self-reduction extent, the highest among transition metal doped samples. Conversely, the other doped materials have better overall redox performance than bare CeZr, resulting into higher oxidation yield (Table 3). Actually, K-containing samples (K-CeZr, K-Fe-CeZr, K-Cu-CeZr) showed the best activity in the redox cycle, featuring both the lowest reduction onset temperature and the highest oxidation yield (Table 3). K and Cu appear to have a synergistic effect, providing an outstanding reduction degree, although it results partially irreversible giving an oxidation yield of about 40%.

Red-ox activity of the most promising catalysts was also evaluated in five consecutive cycles in the TG apparatus, to assess their cyclability. Fig. 3 reports obtained TG curves for K-Fe-CeZr, K-Cu-CeZr and K-CeZr samples. The amounts of oxygen and carbon oxide production were evaluated by mass changes in TG curves and are reported in Table 3. Since bare CeZr sample showed a marked decrease of redox activity after two redox cycles (Fig. S6†), it was not investigated over 5 repeated cycles. K-Fe-CeZr, and K-CeZr samples show very good and stable activity during five consecutive cycles (Fig. 3a), displaying high oxidation yield, about 60%, as well as complete reversibility, confirmed by a reduction yield of about 100%, whereas K-Cu-CeZr sample evidences a poor stability due to a heavy decrease of mass changes in TG apparatus over repeated

Table 3	Reduction onset temperature (T_{OR} , °C), released O ₂ (n_{O2} , μ mol g ⁻¹) and produced CO (n_{CO} , μ mol g ⁻¹) during thermochemical cycles in
TG^a	

		CeZr	Fe-CeZr	Mn-CeZr	Cu-CeZr	K-CeZr	K-Fe-CeZr	K-Cu-CeZr
	$T_{\rm OR}$	1150	1150	1100	750, 1100	870	900	725, 880
I cycle	n_{O_2}	300	313	297	406	307	279	553
5	n _{co}	83	219	237	125	332	294	331
II cycle	n_{O_2}	63	_	_	_	72	46	161
	n _{co}	25	_	_	_	160	122	59
III cycle	n_{O_2}	_	_	_	_	73	54	109
	$n_{\rm CO}$	_	_	_	_	134	99	19
IV cycle	n_{O_2}	_	_	_	_	71	51	65
	n _{co}	_	_	_	_	110	97	0
V cycle	n_{O_2}	_	_	_	_	70	56	81
-	$n_{\rm CO}$	_	_	_	_	107	90	12

^{*a*} Expected error (\leq 10%). In the case of K-CeZr and K-Fe-CeZr during the II cycle, the CO is more than twice the released oxygen, probably due to a reorganization of the catalytic active site during the first cycle. In the case of K-Cu-CeZr during the III–V cycles, the β value is abnormal due to the baseline drift. In these conditions, the evaluation of this parameters is quite useless.

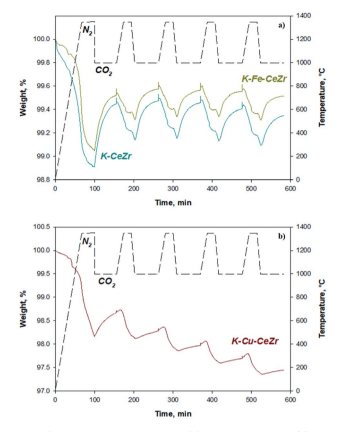


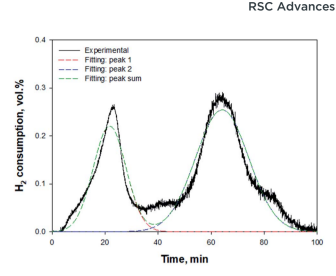
Fig. 3 TG profiles of five consecutive CO_2 splitting cycles on (a) K-Fe-CeZr and K-CeZr samples and (b) K-Cu-CeZr sample.

redox cycles (Fig. 3b). Reduction degree after each step (x_{red} , %), oxidation yield (α , %), and reduction yield (β , %) are reported in ESI (Table S3†).

H₂O splitting tests

In order to further assess the splitting properties of the investigated materials, these were tested towards H_2O splitting following a procedure, which submitted the samples to alternated flows of H_2 and H_2O mixtures.^{29,30} Actually, H_2 driven reduction is expected to lead to higher reduction degrees than self-reduction. However, recent studies proved that the qualitative trends of red-ox performance were closely related to doping and not influenced by reduction pathway, thus allowing for effective comparison between H_2O splitting (lab-reactor) tests and CO_2 oxidation measurements in TG apparatus.³² Fig. 4 reports a typical TPR profile together with its deconvolution by Gaussian functions obtained for K-Fe-CeZr sample, which can be considered as representative of TPR curves of the other samples (Fig. S9†).

Overall amounts of released oxygen and produced hydrogen were assessed by integrating the obtained peaks in TPR (Fig. S9†) and TPO profiles (Fig. S10†). Fig. 5 reports evolved oxygen and hydrogen amounts at each main peak *vs.* peak temperature in two consecutive reduction–oxidation cycles. Overall oxygen and hydrogen amounts produced in each cycle are reported in Table 4.



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Fig. 4 TPR profile obtained on the K-Fe-CeZr during the first cycle and related curve-fitting.

Reduction degree after each step (x_{red} , %), oxidation yield (α , %), and reduction yield (β , %) are reported in ESI (Table S4†).

During the first reduction cycle the TPR profile of CeZr sample shows three peaks at about 530, 660, and 870 $^{\circ}$ C, while only two peaks, at about 670 and 835 $^{\circ}$ C can be detected in the

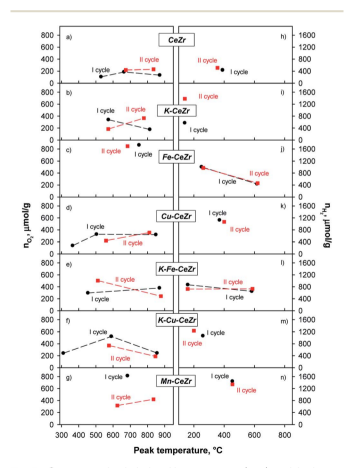


Fig. 5 Oxygen evolved during H_2 treatments (a–g) and hydrogen produced during H_2O splitting tests (h–n) in two reduction–oxidation cycles for investigated materials.

Table 4 Released O₂ (n_{O_2} , μ mol g⁻¹) and produced H₂ (n_{H_2} , μ mol g⁻¹) during TPR and TPO cycling, reduction degree after each step (x_{red} , %), oxidation yield (α , %), and reduction yield (β , %)

	Sample	CeZr	Fe-CeZr	Mn-CeZr	Cu-CeZr	K-CeZr	K-Fe-CeZr	K-Cu-CeZr
I cycle	n_{O_2}	428	866	814	785	516	676	1008
	n_{H_2}	436	1440	1445	1127	573	1517	1064
II cycle	n_{O_2}	450	847	738	576	549	747	559
	n_{H_2}	509	1437	1344	1070	1380	1458	1237

second reduction run (Fig. 5a and S9†). For CeZr low temperature reduction peaks in the range 300–600 °C are usually ascribed to surface sites, whereas those occurring at high temperature, $T \ge 800$ °C, are expected to involve bulk sites.⁵⁵ The decrease in the number of reduction peaks after the first run could be due to sintering phenomenon, which might likely convert surface sites into bulk ones.^{41,55}

K, Fe, Cu and Mn doping significantly modifies the reduction profiles, which disclose at least two superimposed (Fe-CeZr sample) or well-resolved peaks involving surface and bulk sites reduction, respectively. Notably, the changes of both peak temperature and peak shape recorded in the second run for most samples suggest that they undergo a restructuring during thermal treatments (Fig. 5b–g).

TPR profile of Cu-CeZr shows a peak at low temperature (about 365 °C) that is no longer evident in the second cycle (Fig. 5d). This could be due to reduction of some copper (or copper-ceria) species, which are not re-oxidized by H₂O during the splitting step, in accordance with the red-ox behavior assessed through TG analysis. In fact, it has been reported that different copper species in CuO/CeO2-based catalysts can be reduced at low temperature (100-250 °C).55,56 K-Cu-CeZr shows a similar behavior in which the reduction peak is further shifted to lower temperatures. Actually, the low temperature peak (about 310 °C) exhibits very similar shape and area in Cu-doped and K-Cu co-doped samples, which share the same Cu content, confirming its attribution to reduction of copper (or coppercontaining) species. In general, all doped samples show better reduction performance than bare CeZr, as already evidenced by TG measurements. Indeed, peak edge in TPR profiles correspond to the highest reaction rate. Thus, since in copper and iron doped samples, these maxima occur at lower temperature than any other investigated material (Fig. S9[†]), it can be driven that copper and iron doping results in faster reduction kinetics (especially for Fe starting from the second cycle), in agreement with previous studies. On the contrary, Mn-CeZr does not show any improvement in reduction kinetics after its re-organization during the first redox cycle, as driven from the shift of the larger reduction effect towards higher temperatures in the second cycle (Fig. S9[†] and 5g).

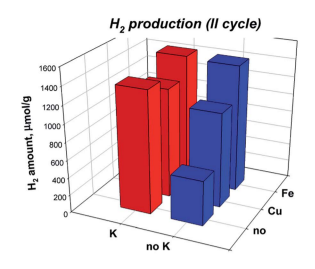
Moreover, K-containing samples (K-CeZr, K-Fe-CeZr) show stable TPR profiles during two consecutive red-ox cycle (Fig. 5b), evidencing comparable peak temperature (Fig. 5) as well as evolved O_2 amount (Table 4), differences lying within expected range of uncertainty ($\leq 10\%$). These features suggest that Kdoping improves thermal stability of CeZr samples preventing the sintering phenomenon, in accordance with TG results. Re-oxidation occurs through one or two phenomena which can involve surface and sub-surface sites or bulk sites. TPO profiles of bare CeZr (Fig. 5h and S10[†]), Cu-CeZr (Fig. 5k and S10[†]) and Mn-CeZr (Fig. 5n and S10[†]) samples evidence a broad oxidation effect, at 350 °C, 400 °C and 450 °C, respectively. This suggests that Mn doping gets splitting kinetics decreased (Fig. 5n). On the contrary, two effects characterize the oxidation profiles of Fe-CeZr in both cycles (Fig. 5j). The first oxidation phenomenon occurs in a wide temperature range, with a main peak at about 100 and 300 °C. K-CeZr, K-Fe-CeZr. Moreover K-Cu-CeZr samples show faster oxidation kinetics than the corresponding K-free materials. In particular, H₂ production occurs at lower temperature; only K-Fe-CeZr shows a second oxidation phenomenon at intermediate temperature (at about 590 °C, Fig. 5l).

Doping by transition metals significantly improves the reducibility of the ceria–zirconia, as well as hydrogen production and oxidation yield (α). Interestingly, both reduction (β) and oxidation (α) yields of the second cycle are very high and close to 100% (Table 4).

Notably, K-doping promotes red-ox performance, since it enables reduction at lower temperature than bare CeZr and improves water splitting activity, as evidenced by the marked increase of evolved H_2 , starting from the second red-ox cycle (Fig. 3b and i).

K addition to the lattice does not significantly affect the quantitative redox performance of the Fe and Cu doped materials (Fig. 5 and Table 4). Interestingly, K-Fe-CeZr shows negative reduction degrees after TPO; this means that the atomic oxygen captured by the sample during the first TPO is larger than the oxygen evolved during first TPR. This behavior can be addressed to the presence of oxygen vacancies after the high temperature calcination that can be fully oxidized by water during TPO.

The effect of doping and co-doping on hydrogen production and oxidation temperature is better illustrated in Fig. 6, where average temperatures are reported for the best performing samples, *i.e.* those containing either Fe(m) or Cu(n) and/or K(n) species. Fig. 6 clearly evidences that hydrogen production is markedly influenced by doping either with transition metals (Fe(m), Cu(n)) or potassium ions. Moreover, K-addition to the lattice improves the oxidation kinetics, as evidenced by lower reaction temperature (Fig. 6b). The positive influence of Kdoping on oxidation kinetics is confirmed by results of a mathematical model which recently formulated and evidenced that K species can increase the number of active sites, promoting redox performances.⁵⁴



Average oxidation temperature (II cycle)

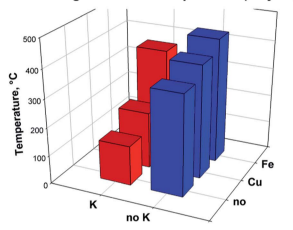


Fig. 6 H_2 production (top) and average oxidation temperature (bottom) during the oxidation steps of the second cycle for transition metals (Cu and Fe) co-doped and K-doped materials.

K-doped and co-doped materials showed the best redox performances in terms of (i) higher reduction degrees, reduction and oxidation yields (*i.e.* larger H_2 amounts), (ii) lower reduction and oxidation temperatures. These results are in accordance with red-ox performance towards CO₂ splitting evaluated through TG analysis. Obtained materials were further cycled showing stable performance, especially the co-doped samples; Fig. S11[†] shows the H_2 production measured during five consecutive cycles. It can be noticed that from the second cycle H_2 production keeps the same value within the 6% error range.

Combining results of thermochemical splitting tests towards CO_2 and H_2O with physico-chemical features driven from EPR and XPS analysis, it can be inferred that red-ox activity is positively affected by surface segregation of Zr species, as well as by a high fraction of Ce^{3+} both on the surface and in the bulk, which should account for a high concentration of oxygen vacancies. Notably, Ce^{3+}/Ce^{4+} surface amount and related oxygen vacancies exert a heavy influence on overall redox behavior; actually, this parameter either increases after use or

keeps constant and higher than bare CeZr in all the best performing compositions: K-CeZr and K-Fe-CeZr, enabling relevant and stable activity towards both CO_2 and H_2O splitting.

Conclusions

In this study, $Ce_{0.75}Zr_{0.25}O_2$ mixed oxides were doped with both transition metal (Cu, Mn, Fe) and potassium cations and tested in CO₂ and H₂O thermochemical splitting cycles. Furthermore, on the basis of XPS and EPR spectroscopic evidence bulk and surface properties accounting for red-ox behavior were enlightened. K doping enhances splitting properties towards both CO₂ and H₂O improving evolved oxygen amounts and lowering activation temperature (*i.e.* faster kinetics) during reduction step. Even Fe and Cu doping and Fe/K or Cu/K co-doping improves redox performance, reducing reduction/ oxidation temperatures.

Physico-chemical characterization evidences a higher fraction of both surface and bulk oxygen vacancies (OV) in K-doped and co-doped samples. Both govern the catalytic activity, promoting oxygen diffusion through the sample during selfreduction as well as the dissociation of the O–H bonds in H_2O splitting. The improved OV migration in the bulk phase and between the bulk and the surface, account for enhanced thermo-catalytic properties.

Oxygen vacancies can be tuned by modifying the catalyst with reducible as well as non-reducible cations. In particular, the use of alkali metals appears a promising route to enhance the redox properties. Moreover, co-doping can produce a synergistic effect as shown by K-Fe-CeZr and K-Cu-CeZr materials.

Overall this study provides strategic guidelines to tailor catalytic performance in thermochemical splitting cycles.

Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 G. Takalkar, R. R. Bhosale and F. AlMomani, *Fuel*, 2019, **256**, 115834.
- 2 F. A. Costa Oliveira, M. A. Barreiros, S. Abanades, A. P. F. Caetano, R. M. Novais and R. C. Pullar, *J. CO2 Util.*, 2018, **26**, 552–563.

- 3 R. R. Bhosale, G. Takalkar, P. Sutar, A. Kumar, F. AlMomani and M. Khraisheh, *Int. J. Hydrogen Energy*, 2019, 44, 34–60.
- 4 Y. Mao, Y. Gao, W. Dong, H. Wu, Z. Song, X. Zhao, J. Sun and W. Wang, *Appl. Energy*, 2020, **267**, 114860.
- 5 S. Chuayboon and S. Abanades, *Int. J. Hydrogen Energy*, 2020, **45**(48), 25783–25810.
- 6 Y. Chen, X. Zhu, K. Li, Y. Wei, Y. Zheng and H. Wang, ACS Sustainable Chem. Eng., 2019, 7, 15452–15462.
- 7 C. L. Muhich, S. Blaser, M. C. Hoes and A. Steinfeld, *Int. J. Hydrogen Energy*, 2018, **43**, 18814–18831.
- 8 B. Zhao, C. Huang, R. Ran, X. Wu and D. Weng, *J. Mater. Sci.*, 2016, **51**, 2299–2306.
- 9 R. R. Bhosale, A. Kumar, F. Almomani, U. Ghosh, S. Al-Muhtaseb, R. Gupta and I. Alxneit, *Ceram. Int.*, 2016, 42, 9354–9362.
- 10 G. D. Takalkar, R. R. Bhosale, A. Kumar, F. AlMomani, M. Khraisheh, R. A. Shakoor and R. B. Gupta, *Sol. Energy*, 2018, **172**, 204–211.
- 11 R. R. Bhosale and G. D. Takalkar, *Ceram. Int.*, 2018, 44, 16688–16697.
- 12 T. Cooper, J. R. Scheffe, M. E. Galvez, R. Jacot, G. Patzke and A. Steinfeld, *Energy Technol.*, 2015, **3**, 1130–1142.
- 13 C. Muhich and A. Steinfeld, J. Mater. Chem. A, 2017, 5, 15578–15590.
- 14 S. Mostrou, R. Büchel, S. E. Pratsinis and J. A. van Bokhoven, *Appl. Catal.*, A, 2017, 537, 40–49.
- 15 M. Takacs, J. R. Scheffe and A. Steinfeld, *Phys. Chem. Chem. Phys.*, 2015, **17**, 7813–7822.
- 16 N. Gokon, T. Suda and T. Kodama, *Thermochim. Acta*, 2015, 617, 179–190.
- 17 N. Gokon, T. Suda and T. Kodama, *Energy*, 2015, **90**, 1280–1289.
- 18 F. Lin, V. A. Samson, A. O. Wismer, D. Grolimund, I. Alxneit and A. Wokaun, *CrystEngComm*, 2016, 18, 2559–2569.
- M. Kang, J. Zhang, C. Wang, F. Wang, N. Zhao, F. Xiao, W. Wei and Y. Sun, *RSC Adv.*, 2013, 3, 18878–18885.
- 20 R. Jacot, R. Moré, R. Michalsky, A. Steinfeld and G. R. Patzke, J. Mater. Chem. A, 2017, 5, 19901–19913.
- 21 A. Le Gal, S. Abanades, N. Bion, T. Le Mercier and V. Harlé, *Energy Fuel.*, 2013, **27**, 6068–6078.
- 22 S. Abanades, A. Legal, A. Cordier, G. Peraudeau, G. Flamant and A. Julbe, *J. Mater. Sci.*, 2010, **45**, 4163–4173.
- 23 A. Le Gal and S. Abanades, *Int. J. Hydrogen Energy*, 2011, **36**, 4739–4748.
- 24 S. Abanades and A. Le Gal, Fuel, 2012, 102, 180-186.
- 25 D. Arifin, A. Ambrosini, S. A. Wilson, B. Mandal, C. L. Muhich and A. W. Weimer, *Int. J. Hydrogen Energy*, 2020, 45, 160–174.
- 26 D. Terribile, A. Trovarelli, C. De Leitenburg, A. Primavera and G. Dolcetti, *Catal. Today*, 1999, **47**, 133–140.
- 27 G. Takalkar, R. R. Bhosale, S. Rashid, F. AlMomani, R. A. Shakoor and A. Al Ashraf, *J. Mater. Sci.*, 2020, 55, 11797–11807.
- 28 C. Ruan, Y. Tan, L. Li, J. Wang, X. Liu and X. Wang, *AIChE J.*, 2017, **63**, 3450–3462.

- 29 A. Pappacena, M. Boaro, L. Armelao, J. Llorca and A. Trovarelli, *Catal. Sci. Technol.*, 2016, **6**, 1–9.
- 30 A. Pappacena, M. Rancan, L. Armelao, J. Llorca, W. Ge, B. Ye,
 A. Lucotti, A. Trovarelli and M. Boaro, *J. Phys. Chem. C*, 2017,
 121, 17746–17755.
- 31 C.-K. Yang, Y. Yamazaki, A. Aydin and S. M. Haile, *J. Mater. Chem. A*, 2014, **2**, 13612–13623.
- 32 G. Luciani, G. Landi, A. Aronne and A. Di Benedetto, Sol. Energy, 2018, 171, 1–7.
- 33 P. Charvin, S. Abanades, E. Beche, F. Lemont and G. Flamant, *Solid State Ionics*, 2009, **180**, 1003–1010.
- 34 T. Maneerung, K. Hidajat and S. Kawi, *Int. J. Hydrogen Energy*, 2017, **42**, 9840–9857.
- 35 B. Zugic, D. C. Bell and M. Flytzani-stephanopoulos, *Appl. Catal.*, B, 2014, 144, 243–251.
- 36 A. Le Gal and S. Abanades, J. Phys. Chem. C, 2012, 116, 13516–13523.
- 37 G. Luciani, G. Landi, C. Imparato, G. Vitiello, F. A. Deorsola,
 A. Di Benedetto and A. Aronne, *Int. J. Hydrogen Energy*, 2019, 44(33), 17565–17577.
- 38 M. Portarapillo, A. Aronne, A. Di Benedetto, C. Imparato, G. Landi and G. Luciani, *Chem. Eng. Trans.*, 2019, 74, 43–48.
- 39 N. D. Petkovich, S. G. Rudisill, L. J. Venstrom, D. B. Boman, J. H. Davidson and A. Stein, *J. Phys. Chem. C*, 2011, **115**, 21022–21033.
- 40 E. Aneggi, C. de Leitenburg, G. Dolcetti and A. Trovarelli, *Catal. Today*, 2006, **114**, 40–47.
- 41 P. S. Barbato, S. Colussi, A. Di Benedetto, G. Landi, L. Lisi, J. Llorca and A. Trovarelli, *Appl. Catal.*, A, 2015, **506**, 268–277.
- 42 K. Asano, C. Ohnishi, S. Iwamoto, Y. Shioya and M. Inoue, *Appl. Catal., B*, 2008, **78**, 242–249.
- 43 Z. Su, W. Yang, C. Wang, S. Xiong, X. Cao, Y. Peng, W. Si, Y. Weng, M. Xue and J. Li, *Environ. Sci. Technol.*, 2020, 54, 12684–12692.
- 44 G. Li, Q. Wang, B. Zhao and R. Zhou, Fuel, 2012, 92, 360-368.
- 45 D. R. Mullins, Surf. Sci. Rep., 2015, 70, 42-85.
- 46 E. Bêche, P. Charvin, D. Perarnau, S. Abanades and G. Flamant, *Surf. Interface Anal.*, 2008, 40, 264–267.
- 47 Y. Polyak and Z. Bastl, Surf. Interface Anal., 2015, 47, 663–671.
- 48 F. Larachi, J. Pierre, A. Adnot and A. Bernis, *Appl. Surf. Sci.*, 2002, **195**, 236–250.
- 49 L. Zhou, X. Li, Z. Yao, Z. Chen, M. Hong, R. Zhu, Y. Liang and J. Zhao, *Sci. Rep.*, 2016, 6, 23900.
- 50 M. Zhao, M. Shen and J. Wang, J. Catal., 2007, 248, 258-267.
- 51 J. Wang, J. Wen and M. Shen, *J. Phys. Chem. C*, 2008, **112**, 5113–5122.
- 52 J. Xu, J. Harmer, G. Li, T. Chapman, P. Collier, S. Longworth and S. C. Tsang, *Chem. Commun.*, 2010, **46**, 1887–1889.
- 53 P. I. Archer, S. A. Santangelo and D. R. Gamelin, *Nano Lett.*, 2007, 7, 1037–1043.
- 54 M. Portarapillo, D. Russo, G. Landi, G. Luciani and A. Di Benedetto, *RSC Adv.*, 2021, **11**, 39420–39427.
- 55 A. Di Benedetto, G. Landi and L. Lisi, Catalysts, 2018, 8, 209.
- 56 T. Caputo, L. Lisi, R. Pirone and G. Russo, *Appl. Catal., A*, 2008, **348**, 42–53.